



ACTIVITY REPORT 2010

Edited by U. Johansson, K. Lilja, A. Nyberg, R. Nyholm

Preface

It has been a very eventful year for MAX-lab. There have been many great research achievements and the development of the laboratory continues. 2010 also represents the start of the build-up of MAX IV. The number of annual users is now well over 800. Due to the increasing number of users, the expanding range of research activities and in particular due to the start-up of the MAX IV project there is presently a rapid increase of the personnel at the laboratory.

A new organization, the MAX IV laboratory, has been established which will operate the present facilities as well as build up the new ones. During the year the funding for the MAX IV facility has been released and several major parts of the accelerator systems have been ordered. A proposal to fund a first set of beamlines was submitted to the Knut and Alice Wallenberg Foundation during 2010.

The 22nd of November there was a groundbreaking ceremony at the MAX IV site with among others the Swedish Deputy Prime Minister and Minister of Education Jan Björklund. Several important steps for the MAX IV building project have been taken during the year. One critical issue has been to ensure that the site is sufficiently good in terms of stability towards vibrations. This is important due to the nearby road and the fact that there will be other building activities in the vicinity. There have been careful measurements at the site, a mockup has been built to test the foundation and theoretical modeling has been carried through. The result of this careful work has been very positive. There has also been a parallel commission to create the exterior of the building and the landscaping, and the result was presented in the autumn. The outcome of this process was that the architect companies Fojab and Snøhetta were asked to work together to produce the final design concepts.

There is intense and fruitful collaboration with several other synchrotron radiation laboratories in connection with the MAX IV project. The Jagiellonian University has during the year got the decision to build the Polish synchrotron radiation source in Krakow. The Polish source will be a copy of the MAX IV 1.5 GeV storage ring. This is an excellent example of European collaboration and it is the first case when two identical machines are built. Several persons from the Jagiellonian University are involved in the common project, some of which are stationed at MAX-lab at present.

Finally I want to acknowledge the support by VR, the Knut and Alice Wallenberg foundation (KAW), VINNOVA, the European Commission, Lund University, Region Skåne, the Swedish Foundation for International Cooperation in Research and Higher Education (STINT) as well as other funding organizations. I would also like to take this opportunity to thank all MAX-lab staff and all the users for their excellent work during all the years I have been the Director and for the good atmosphere that they have created at the laboratory. I would also like to thank all others who have contributed to the development of the laboratory as members of the Board, members of the PAC, SAC, and MAC expert committees, etc.

Lund 13 June, 2011

Nils Mårtensson

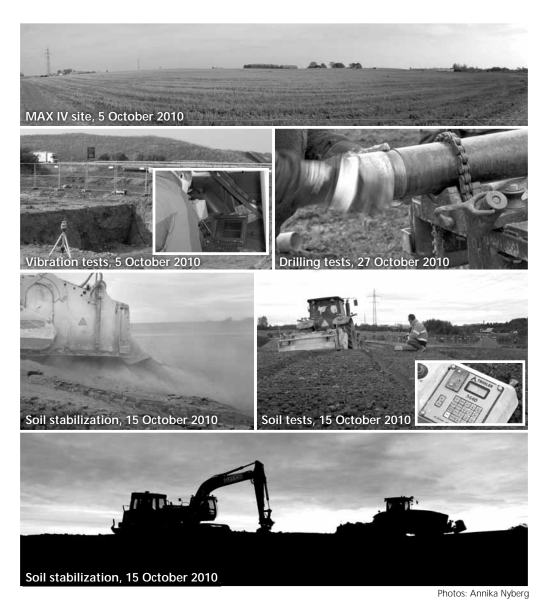
Director MAX-lab, 1997 - 2010

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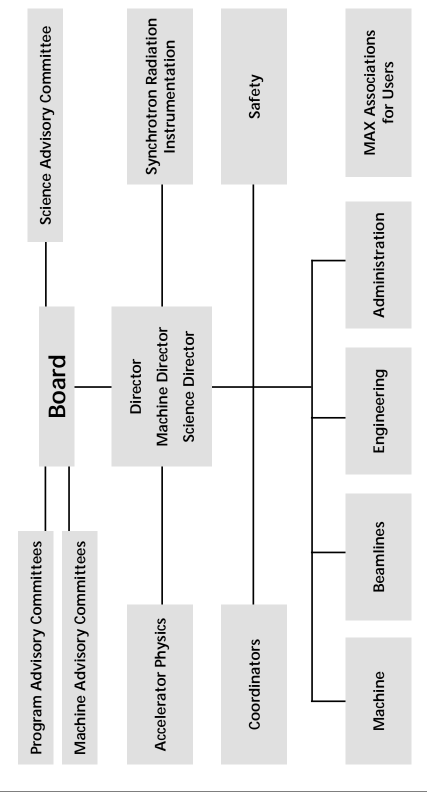


The Machine Advisory Committee (MAC); Peter Kuske, Massimo Cornacchia, Klaus Balewski, Søren Pape Møller, and Lenny Rivkin, at the meeting that took place 1 September 2010. Photo: Annika Nyberg



MAX-lab Activity Report 2010

Organisation December 2010 - an overview



Please note:

This is the organisation chart for 2010. During 2011, a reorganisation of the MAX IV Laboratory will be made to meet the new demands imposed by the MAX IV project. An up-to-date organisation chart will be made available at www.maxlab.lu.se.

MAX-lab Personnel From January 2010 – December 2010

Department	of Acce	elerator	Physics
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Position Name Area of responsibilities

Professor Mikael Eriksson Head of the accelerator physics department Professor Sverker Werin Research and development in accelerator physics Research and development in accelerator physics Lecturer Lars-Johan Lindgren (Universitetslektor) and coordinator for accelerator physics research,

until 31 July 2010

Research and development in accelerator physics

Erik Wallén

Researcher (Forskare)

Ph.D. student Nino Čutić

(Doktorand)

Ph.D. student Anders Hansson

(Doktorand)

Department of Synchrotron Radiation Instrumentation

Position Name Area of responsibilities Professor Ralf Nyholm Head of the synchrotron radiation instrumentation department Rami Sankari Research and development of beamlines Lecturer (Universitetslektor) and instrumentation Research Associate Sophie Canton Research and development of time-resolved (Forskarassistent) X-ray methods Ph.D. student Jörg Schwenke

Operating Staff

Doomlings

(Doktorand)

Beamlines		
Position	Name	Area of responsibilities
Professor	Jesper Andersen	Science director
Professor	Ralf Nyholm	Manager and coordinator for user operations
Research engineer (Forskningsingenjör)	Ulf Johansson	Deputy manager
Research engineer (Forskningsingenjör)	Kurt Hansen	Head of the user support group
Researcher (Forskare)	Johan Adell	Maintenance of beamlines and experimental stations, especially beamline I4
Research engineer (Forskningsingenjör)	Thiagaraian Balasubramanian	Coordinator for low energy beamlines, (73, I3, I4)
Researcher (Forskare)	Matts Björck	Design, installation and maintenance of beamlines, especially beamline I1011
Researcher (Forskare)	Jason Brudvik	Tagging facility MAX I
Researcher (Forskare)	Stefan Carlsson	Design, installation and maintenance of beamlines, especially beamline I811

Beamlines, cont.		
Position	Name	Area of responsibilities
Project coordinator (Projektsamordnare)	Yngve Cerenius	Coordinator for beamline projects, MAX IV
Researcher (Forskare)	Jeppe Christensen	Design, installation and maintenance of beamlines, especially beamline I711
Research engineer (Forskningsingenjör)	Anders Engdahl	Maintenance of beamlines and experimental stations, especially beamline 73
Researcher (Forskare)	Henrik Enquist	Design of the short-pulse facility for MAX IV
Researcher (Forskare)	Johan Forsberg	Design of the new beamline I511
Researcher (Forskare)	Yury Gaponov	Software development for hard X-ray beamlines until 31 December 2010
Researcher (Forskare)	Carsten Gundlach	Design, installation and maintenance of beamlines, especially I711, I811, and I911 Postdoctor until January 2011
Research engineer (Forskningsingenjör)	Dörthe Haase	Maintenance of hard X-ray beamlines and experimental stations
Researcher (Forskare)	Franz Hennies	Design, installation and maintenance of beamlines, especially beamline I511
Researcher (Forskare)	Lennart Isaksson	Coordinator for Nuclear physics research and radiation safety
Research engineer (Forskningsingenjör)	Mikael Johansson	Electronics for the experiments
Research engineer (Forskningsingenjör)	Mats Leandersson	Maintenance of beamlines and experimental stations, especially on beamline I3
Research engineer (Forskningsingenjör)	Filip Lindau	FEL test experiments and laser safety
Research engineer (Forskningsingenjör)	Magnus Lundin	Computers for the experiments and radiation safety
Research engineer (Forskningsingenjör)	Anders Månsson	Maintenance of vacuum and cryo systems
Research engineer (Forskningsingenjör)	Björn Nilsson	Nuclear physics
Researcher (Forskare)	Katarina Norén	Coordinator for Hard X-ray (I711, I811, and I911) and chemical safety
Engineer (Ingenjör)	Anders Olsson	Beamline engineer at I311 and I511
Researcher (Forskare)	Annette Pietzsch	Design, installation and maintenance of beamlines, especially I511. Postdoctor until August 2010
Researcher (Forskare)	Tomás Plivelic	Design, installation and maintenance of beamlines, especially 1911-4. Postdoctor until May 2010
Researcher (Forskare)	Alexei Preobrajenski	Maintenance of beamlines and experimental stations, especially beamline D1011
Researcher (Forskare)	Janusz Sadowski	MBE system since 1 June 2010
Research engineer (Forskningsingenjör)	Peter Sondhauss	Design and simulation of X-ray optics

Beamlines, cont.		
Position	Name	Area of responsibilities
Engineer (Ingenjör)	Bertil Svensson	Maintenance of hard X-ray beamlines and experimental stations
Project manager (Projektledare)	Christer Svensson	Software development for hard X-ray beamlines
Researcher (Forskare)	Maxim Tchaplyguine	Design, installation and maintenance of beamlines, especially beamline I411
Researcher (Forskare)	Karina Thånell Schulte	Coordinator for soft X-ray beamlines (I311, I411, I511, D1011, and I1011) Postdoctor until March 2010
Research engineer (Forskningsingenjör)	Johan Unge	Design installation and maintenance of beamlines, especially 1911. Postdoctor until May 2010
Research engineer (Forskningsingenjör)	Thomas Ursby	Design, installation and maintenance of beamlines, especially beamline I911
Research engineer (Forskningsingenjör)	Stefan Wiklund	Maintenance of beamlines, experimental stations; safety manager
Researcher (Forskare)	Alex Zakharov	Maintenance of beamlines and experimental stations, especially the SPELEEM at I311
Researcher (Forskare)	Gunnar Öhrwall	Design, installation and maintenance of beamlines, especially beamline I1011

Machine

Position	Name	Area of responsibilities
Professor	Mikael Eriksson	Machine director
Lecturer (Universitetslektor)	Lars-Johan Lindgren	Deputy manager and coordinator for accelerator physics research until 31 July 2010
Researcher (Forskare)	Åke Andersson	Deputy machine director
Project manager (Projektledare)	Pedro Fernandes Tavares	s Project manager for storage rings at MAX IV
Research engineer (Forskningsingenjör)	Lars-Gösta Johansson	High-tension current systems for the accelerators and coordinator for electronics and electrical installations
Project engineer (Projektingenjör)	Martin Johansson	Design and construction of magnets MAX IV
Research engineer (Forskningsingenjör)	Dionis Kumbaro	Maintenance and operation of accelerators
Researcher (Forskare)	Simon Leemann	Beam dynamics
Research engineer (Forskningsingenjör)	Per Lilja	Maintenance and operation of accelerators
Research engineer (Forskningsingenjör)	Lars Malmgren	Micro-wave electronics for the accelerator system
Project coordinator (Projektsamordnare)	Jonas Modéer	MAX IV accelerator coordinator
Research engineer (Forskningsingenjör)	Robert Nilsson	RF and diagnostics

Machine, cont. Position	Name	Area of responsibilities
Research engineer (Forskningsingenjör)	Jerry Schmidt	Development and maintenance of the accelerator system, especially insertion devices
Researcher (Forskare)	Magnus Sjöström	Beam dynamics
Researcher	Sara Thorin	Project manager for the LINAC at MAX IV

Engineering
Docition

(Forskare)

	rea of responsibilities
Berglund He	ead of Engineering group
lbäck Va	acuum system design, MAX IV
ndersson W	/orkshop, mechanics
jermo De	esign and construction
stiansson W	/orkshop, mechanics
El Afifi De	esign and construction
ırsztka Al	lignment and installations
ägneryd Ele	ectrical installations
	ability, tolerances and vibrations; ability group manager
nngren Re	esponsible for electrical installations
dkvist Au	utomation/PLC
ilsson W	orkshop, mechanics
on W	/orkshop, mechanics, retired 31 October 2010
mmarin Al	lignment and installations
rensson De	esign and construction
ånell Au	utomation/PLC since 28 January 2010
k Wingren M	laintenance, service
	ilbäck Vandersson Walbäck Vandersson Walbäck Vandersson Walbarsztka Alägneryd El Walbarsztka Alägneryd El Walbarsztka Alägneryd El Walbarsztka Alägneryd Walbarsztka Alägneryd Walbarsztka Alägneryd Walbarsztka Alässon Walbarszt

Computing Services and Control Systems

Position	Name	Area of responsibilities
Research engineer (Forskningsingenjör)	Krister Larsson	Head of computing services and control systems

Computing Services and Control Systems, cont.		
Position	Name	Area of responsibilities
Ph.D. student (Doktorand)	Thilo Friedrich	MAX-lab and KTH
Software developer (Programutvecklare)	Mirjam Lindberg	Software development
Web developer (Webbutvecklare)	Tobias Lundquist	Web development
Research engineer (Forskningsingenjör)	Mats Nilsson	Computers for the accelerator system
Software developer (Programutvecklare)	Andreas Persson	Software development
Project manager (Projektledare)	Darren Spruce	Software development coordinator
Research engineer (Forskningsingenjör)	Jüri Tagger	Accelerator control system developer
Computer engineer (IT-tekniker)	András Váncsa	Computer support

Administration

Administration		
Position	Name	Area of responsibilities
Procurement advisor (Upphandlingssamordnare)	Göran T Andersson	Procurements
Administrative officer (Ekonomiadministratör)	Ann Barthel	Office work, invoicing
Administrative officer (Enhetsadministratör)	Elisabeth Dahlström	Office work
Administrative officer (Enhetsadministratör)	Carolina Ingvander	Office work, reception
Project assistant (Projektassistent)	Markus Johannesson	IRUVX-PP project
Communication manager (Kommunikationsansvarig)	Karin Lilja	Communication
Custodian (Lokalvårdare)	Catarina Nilsson	Cleaning
Information officer (Informatör)	Annika Nyberg	Communication
Custodian (Lokalvårdare)	Lisbeth Olsson	Cleaning
HR officer (Personalsamordnare)	Therese Oppliger	Human resources
Janitor (Vaktmästare)	Johan Rosdahl	Janitor's office
Financial manager (Ekonomichef)	Mats Roxendal	Finances
Administrative coordinator (Administrativ koordinator)	Helena Ullman	Executive support, meetings coordinator

Personnel with Administrative Assignments at MAX-lab

Position	Name	Area of responsibilities
Professor	Jesper Andersen	Science director since 1 October 2010
Professor	Mikael Eriksson	Machine director
Guest Professor	Åke Kvick	Senior advisor
Professor	Ingolf Lindau	Senior advisor
Project coordinator	Caj Lundquist	MAX IV building project
Professor	Nils Mårtensson	Director until February 2011
Professor	Bent Schröder	Senior advisor
Professor	Svante Svensson	Deputy director until 31 December 2010
Expert	Sten Wennerström	Administrative director

Beamline

Affiliation

Postdoctors and Visiting Scientists		
Position	Name	
Researcher (Forskare)	Marcus Agåker	

		204	
Researcher (Forskare)	Marcus Agåker	I511	Uppsala University, 1 July-31 December 2010
Ph.D. student (Doktorand)	Tomas Andersson	I411	Uppsala University
Researcher (Forskare)	Olle Björneholm	1411	Uppsala University
Researcher (Forskare)	Kevin Fissum	Nuclear physics	Lund University, Faculty of Technology
Visiting scientist (Gästforskare)	Piotr Goryl	Accelerators	Jagiellonian University
Postdoctor	Maher Harb	D611	Lund University, Faculty of Technology
Research engineer (Forskningsingenjör)	Maria Håkansson	Protein crystallization facility	SARomics Biostructures AB
Ph.D. student (Doktorand)	Andrius Jurgilaitis	D611	Lund University, Faculty of Technology
Postdoctor	Brian Kennedy	I511	MAX-lab
Postdoctor	lwona Kowalik	I1011	MAX-lab until 14 January 2010
Postdoctor	Rajesh Kushwaha	Rebecca Project	From November 2010
Researcher (Forskare)	Ana Labrador	1911-4	MAX-lab since 1 November 2010
Researcher (Forskare)	Jörgen Larsson	D611	Lund University, Faculty of Technology
Researcher (Forskare)	Andreas Lindblad	IRUVX-PP	Uppsala University
Researcher (Forskare)	Derek Logan	1911	Lund University and MAX-lab
Researcher (Forskare)	Bengt Nelander	73	MAX-lab
Ph.D. student (Doktorand)	May Ling Ng	D1011	Uppsala University, Thesis defence 5 November 2010
Ph.D. student (Doktorand)	Ralf Nüske	D611	Lund University, Faculty of Technology

Postdoctors and Visiting Scientists, cont.

Position	Name	Beamline	Affiliation
Ph.D. student (Doktorand)	Jesper Nygaard	1711 and 1911	University of Copenhagen and MAX-lab
Postdoctor	Prabir Pal	13	Uppsala University
Researcher (Forskare)	Pål Palmgren	13	October-December 2010
Project manager (Projektledare)	Christian Riekel	Nanofocus beamline	Until 31 July 2010
Research Engineer (Forskningsingenjör)	Conny Såthe	I511	Uppsala University
Research engineer (Forskningsingenjör)	Keld Theodor	I911	University of Copenhagen and MAX-lab
Researcher	Marjolein Thunnissen	1911	Lund University and MAX-lab
Postdoctor	Wimal Ubhayasekera	1911	University of Copenhagen
Ph.D. student (Doktorand)	Intikhab Ulfat	MBE at 41 and I3	Chalmers University of Technolog
Postdoctor	Samuli Urpelainen	13 and 1411	Oulu University and MAX-lab
Researcher (Forskare)	Per Uvdal	73	Lund University and MAX-lab
Visiting scientist (Gästforskare)	Adriana Wawrzyniak	Accelerators	Jagiellonian University
Ph.D. student (Doktorand)	Nikolay Vinogradov	D1011	Uppsala University
Ph.D. student (Doktorand)	Chaofan Zhang	I411	Uppsala University

General Information

Introduction

MAX IV Laboratory is a Swedish National Laboratory for research using synchrotron radiation and high-energy electrons and for research in accelerator physics. The organisation is comprised of the present MAX-lab and the MAX IV project. MAX-lab is based upon a linac injector, a storage/pulse-stretcher ring – MAX I – and two third generation storage rings: MAX II and MAX III.

During 2010 MAX-lab has accommodated about 820 scientists representing scientific groups from 130 industry, academic and government laboratories from 35 countries.

Synchrotron radiation based research at MAX I, MAX II and MAX III is done in a large variety of disciplines including surface science, semiconductor physics, materials science, atomic and molecular physics, chemistry, biology, medicine and environmental science.

At the 550 MeV MAX I storage ring there are four beamlines for synchrotron radiation research in the infra-red through VUV photon energy range using various spectroscopic techniques. At the 1.5 GeV MAX II storage ring thirteen beamlines are now in operation. Three planar undulators and one EPU for the VUV and soft X-ray regions are used for various spectroscopic techniques. Three multipole wigglers, two of which have superconducting magnets, are used for absorption and diffraction experiments in the X-ray region. In addition to these insertion device beamlines also two bending magnet ports are utilized. At the MAX III ring one planar undulator and one EPU are used for spectroscopy in the UV and VUV regions.

In the pulse-stretcher mode the MAX I ring is used for experiments in nuclear and hadron physics. The electron energy range available is at present 145 to 200 MeV, resulting in monochromatic photons from 15 to 185 MeV. The program used 18 weeks of beam time in 2010.

Contact Persons MAX IV Laboratory

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Direc	tor	

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 MAX IV Laboratory

Lund University P.O. Box 118 SE-221 00 Lund SWEDEN

Visiting address: Ole Römers väg 1, Lund, SWEDEN

Web site: www.maxlab.lu.se

Organisation

MAX IV Laboratory is operated by Lund University under a contractual agreement with the Swedish Research Council (Vetenskapsrådet). The Research Council is responsible for the main operating budget, the scientific programs and periodic reviews of the activities. As host university, Lund University is responsible for employee relations and the conventional facilities.

From 1 March 2004 until 28 February 2009, MAX-lab was a part of the Integrated Infrastructure Initiative (I3) "Integrating Activity on Synchrotron and Free Electron Laser Science" (IA-SFS). Since 1 March 2009, this program has been followed by the I3 program "European Light Sources Activity" (ELISA). This means that a number of European projects within synchrotron radiation research are sponsored, including support for travel and subsidiary costs for users from EU and other associated countries.

An outline of the internal organisation of MAX IV Laboratory is found on page 1 in this report.

MAX IV Laboratory is governed by a Board, consisting of eight members and a chairperson since October 2010. The Board is appointed by Lund University in consultation with the Research Council and Vinnova. The Board appoints the Director and a Managerial group of MAX IV Laboratory.

The members of the Board during 2010 are listed in table I a-c.

The Board of MAX IV Laboratory is charged with the responsibility of working towards national and international use of the laboratory, to prepare and monitor the operating budget, to prioritize the research programs and facility developments, to appoint personnel, and to promote the overall growth and development of the laboratory. The MAX IV Laboratory Board also decides how to divide the operating time between research with synchrotron radiation, energetic electrons, accelerator physics research and system maintenance and development. To facilitate these decisions the Board has appointed program advisory committees (PACs), a machine advisory committee (MAC), and a scientific advisory committee (SAC) to guide it on matters connected with the scientific programs and facility developments at the laboratory. The committees consist of national and international scientists whose backgrounds encompass the various areas of research which are possible at MAX-lab.

The members of the program advisory committees are listed in table IIa and b, the members of the scientific advisory committee in table III, and the members of the machine advisory committee in table IV.

Table la

Board of MAX-lab			
July 2007 – Ju	July 2007 – June 2010		
Member Representing			
I. Skogö	Chairperson		
A. Borg	Research Council		
L. Kloo	Research Council		
M. Fahlman	Users		
S. Svensson	Users		
B. Söderström	Lund University		
S. Sörensen	Lund University		

Table Ib

Interim Boar July - Septe	rd of MAX IV Laboratory mber 2010
Member	Representing
L. Börjesson	Chalmers University of Technology, Chairperson
P. Eriksson	Lund University

Table Ic

October 201	0 –
Member	Representing
L. Börjesson	Chalmers University of Technology, Chairperson
H. Dosch	DESY, Hamburg, Germany
K. Edström	Uppsala University, Sweden
L. Hultman	Linköping University, Sweden
S. Larsen	University of Copenhagen, Denmark (until Feb 2011)
A. I'Huiller	Lund University, Sweden
I. Reineck	Sandvik Tooling AB, Sweden
G. Schneider	Karolinska Institutet, Sweden
S. Sörensen	Lund University, Sweden

Table IIa

MAX IV Laboratory Programme Advisory Committees Synchrotron Radiation (in 2010)

B. Johansson Uppsala, Sweden, Chairperson

H. AkselaM.C. AsensioD. ChandesrisOulu, FinlandParis, FranceParis, France

R. Feidenhans'l Copenhagen, Denmark

R. Fourme Paris, France

M. Gajhede Copenhagen, Denmark

Å. KvickLund, SwedenC. MironParis, FranceL.H. TjengKöln, GermanyK. WilsonYork, UKD.P. WoodruffWarwick, UK

Energetic Electrons (in 2010)

B. Höistad Uppsala, Sweden, Chairman

J. Ahrens Mainz, Germany
D. Phillips Athens, Ohio, USA
D. Watts Edinburg, UK

Table IIb

MAX IV Laboratory Programme Advisory Committees Synchrotron Radiation (from 2011)

L. Johansson Karlstad, Sweden, Chairperson A. Borg Trondheim, Norway

N. Brookes Grenoble, France
J. Daillant Paris, France
A.N. Fitch Grenoble, France
M. Gajhede Copenhagen, Denmark
C. Hirschmugl Milwaukee, USA
K. Horn Berlin, Germany
B. Hunter Dundee, Scotland UK

M. Kiskinova Trieste, Italy
E. Kukk Turku, Finland

Å. Kvick Lund, Sweden (spring meeting 2011)

R. Mathiesen Trondheim, Norway
C. Miron Paris, France
L. Patthey Villigen, Switzerland
H. Friis Poulsen Roskilde, Denmark
W. Wurth Hamburg, Germany

Table IIIa

MAX IV Laboratory Scientific Advisory Committee until end of 2010

S. Lidin Stockholm, Sweden, Chairperson

L. Braicovich Milano, Italy
C. Nave Daresbury, UK
B. Schoch Bonn, Germany
A. Wrulich Villigen, Switzerland

Table IIIb

MAX IV Laboratory Scientific Advisory Committee from end of 2010

B. Johansson Uppsala, Sweden, Chairperson

H. Aksela Oulu, Finland M.A. Carrondo Lisbon, Portugal W. Eberhardt Berlin, Germany Lausanne, Switzerland G. Margaritondo L. McCusker Zürich, Switzerland A Molenbroek Lyngby, Denmark H. Reichert Grenoble, France I Robinson London UK P. Woodruff Warwick, UK

Table IV

MAX IV Laboratory Machine Advisory Committee

L. Rivkin Villigen, Switzerland, Chairperson

K. Balewski Hamburg, Germany

M. Cornacchia Palo Alto, California, USA (during 2010)

P. Kuske Berlin, Germany
S. Pape Møller Aarhus, Denmark
R. Walker Didcot, UK (from 2011)

Research

MAX IV Laboratory supports three distinct areas of research: Accelerator physics, research based on the use of synchrotron radiation, and nuclear physics research using energetic electrons. The use of the facility is shared by the groups working in these three research areas as is the responsibility for different aspects of the facility.

The laboratory is operated primarily in a user oriented fashion. Those interested in making experiments using synchrotron radiation or the electron-beam facility should obtain more information from the coordinators (see table V) about requirements for experiments and available equipment. Technical information on beamlines and experimental stations can also be found on the MAX IV Laboratory web site, www.maxlab.lu.se.

Table V

MAX-lab Research Coordinators

Synchrotron Radiation

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Prof. Ralf Nyholm

Energetic Electrons
Lennart Isaksson

Phone: +46-(0)46-222 77 15 lennart.lsaksson@maxlab.lu.se

Accelerator Physics
Prof. Mikael Eriksson

Phone: +46-(0)46-222 76 96 mikael.eriksson@maxlab.lu.se

Postal address: MAX-lab, Lund University, P.O. Box 118, SE-221 00 Lund, SWEDEN

Visiting address: Ole Römers väg 1, Lund, SWEDEN

Those who wish to make experiments using synchrotron radiation or energetic electrons should apply in the form of a research proposal to the appropriate coordinator. Proposals are periodically reviewed by the MAX IV Laboratory Program Advisory Committees (see Table II), and scheduling of time on beamlines is made by the research coordinators based upon user requests.

Most of the beamlines are equipped with experimental chambers, analysers and detectors available to all users. In addition some user groups provide their own experimental set-ups. More information about the storage ring parameters and the beamlines may be found in the introductions to the sections on accelerator physics research, synchrotron radiation research and research with energetic electrons in this report.

The machines are usually operated 24 hours per day, six days per week. Each week one day is devoted to machine maintenance and development of storage ring instrumentation.

The users should take full responsibility for carrying out their research projects and thus should become familiar with the experimental equipment and data-taking systems which are available at the laboratory before they begin the experiments in order to optimize the use of beam time. User equipment which is used in the ultra-high vacuum environment required for connection to the storage ring and monochromators must be checked and approved by the laboratory.

Safety guidelines pertaining to samples and experimental procedures must also be followed by users. Prior to experiments all participating researchers must register their beamtime through our web-server. Also a declaration of substances and hazardous equipment has to be submitted to the MAX IV Laboratory for approval.

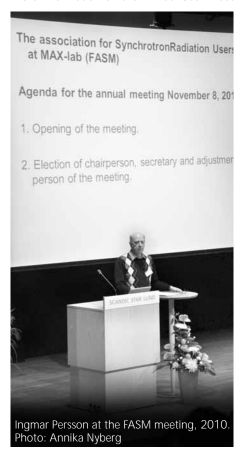
Much of the design of monochromators and experimental equipment is the result of collaborations with research groups outside MAX IV Laboratory. If a user group is seeking funding for equipment which is intended for use the laboratory, the proposal should be discussed with the MAX IV Laboratory so that planning can be made together with the laboratory personnel. Funding organizations in Sweden may appeal to the laboratory staff for information on the feasibility of projects connected to the laboratory. This ensures that new equipment will be compatible with the standards of the laboratory and with the existing beamline structure.

User Associations

MAX Association for Synchrotron Radiation Users at MAX-lab - FASM

The 23rd annual meeting of "Föreningen för Användare av Synkrotronljuset vid MAX-laboratoriet" (The association for Synchrotron Radiation Users at MAX-lab), FASM, took place on 8 November, 2010, at hotel Scandic Star in Lund, in connection with the Annual User Meeting. A new FASM board for the up-coming three years was elected, see below.

More information on the Annual User Meeting is presented on page 17.



All users of the MAX-lab facility are by definition members of the FASM organization, which aims to have a broad participation involving all parts of the user community and efficient communication with the MAX-lab board and management. It is thereby possible to communicate current and future needs within the user community and the MAX-lab Board. At the same time the association disseminates information to the users about MAX-lab plans and prospects. FASM's mission is increasingly important as the user community is growing rapidly and more scientific disciplines are actively using the MAX-lab facility. In view of this, FASM will introduce a number of proposals aimed to strengthen the activities at MAX-lab and be of further aid to new users.

MAX-lab users who want to bring up ideas to improve MAX-lab as scientific tool and organization are welcome to contact:

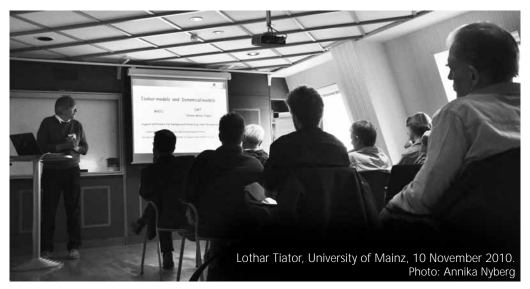
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Table VI

Board of Association for Synchrotron Radiation Users (FASM)

I. Persson Uppsala, Sweden, Chairperson

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R. Neutze
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B.-M. Steenari
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MAX Association for Nuclear Physics Users

The Association of Nuclear Physics Users at MAX-lab met on 8-9 November at a combined PAC and User meeting in conjunction with the MAX-lab Annual User Meeting at Hotel Scandic Star in Lund. Four new proposals and letters of intent were presented, as well as status reports from ongoing experiments, facility-related reports and talks from invited speakers. The detailed program of the meeting is found on http://www.maxlab.lu.se/usermeeting/2010/sessions/nuclear_pac_meeting.html Election of the board of the Association resulted in the following composition:

Table VII

Board of MAX Association for Nuclear Physics Users

J.R.M. Annand Glasgow, UK, Chairperson W. Briscoe Washington DC, USA

P. Grabmayr Tübingen, Germany, Election coordinator

L. Isaksson Lund, Sweden

Commercial Utilisation of the MAX IV Laboratory

The MAX IV Laboratory is selling synchrotron radiation beamtime as well as consulting services in connection to this. Several companies are regular customers at the laboratory. The Laboratory can furthermore provide consulting services in accelerator technology and synchrotron radiation instrumentation. Further information can be obtained from our director Sine Larsen.

A macromolecular crystallization facility is available at tha lab. The facility is operated under a contractual agreement with SARomics Biostructures AB and is at present used approximately equally for commercial and academic projects.

Workshops and Schools

Beamlines at MAX IV, Workshop, Lund, 22-23 February.

This workshop gathered over 170 users of Synchrotron Radiation from Sweden, the Nordic/Baltic countries, Poland and Europe. The program of the workshop can be found at www.maxlab.lu.se.

Prof. Ulf Karlsson held an introduction, where he explained the application process of the MAX IV beamlines from the Swedish Research Council's point of view. After presentations of the MAX IV project from Prof. Nils Mårtensson and Prof. Mikael Eriksson two talks were given on subjects of great importance for the future MAX IV facility:

- Dr. Christian Riekel, MAX-lab, gave a presentation on nanobeams, where the unique properties of the ultra-brilliant radiation from MAX IV was discussed in context with international research.
- Dr. Anders Madsen, ESRF, gave a presentation of X-ray coherence where the ultra-low emittance of the MAX IV facility will open unique possibilities.



Ulf Karlsson, 22 February 2010. Photo: Annika Nyberg

After this the meeting was divided into three parallel sessions:

- 1. VUV, IR and Soft X-rays
- 2. Hard X-rays
- 3. Life Sciences

In all these fields there has been a strong activity from the User Community and a large number of beamline proposals were put forward.

The discussions were very fruitful and reflected the strong user community around the MAX laboratory. Several cases of synergy were found and it was possible to find opportunities where the different user groups can make a consorted effort to make use of the most ultra-brilliant facility in the world.

Second Workshop on High Harmonic Seeding for Present and Future Short Wavelength Free-Electron Lasers (FELs), Lund, 5-7 May.

Hosted by MAX-lab, the Lund Laser Centre, Sincrotrone Trieste, ENEA, INFN-LNF and University of Rome, this second workshop was held in Lund, Sweden, 5-7 May 2010. The first workshop, held in December 2008 in Frascati, Rome, brought together the accelerator and high power laser communities. This second workshop permitted both continued interactions of participants as well as an expanded participation roster to focus upon the need in the free-electron laser community for coherent, short wavelength seed pulses.

During the workshop results from High Harmonic Generation (HHG) laser seeding at SCSS (Japan) and SPARC (Rome) were reported. Also the status of the seeding projects at sFLASH (DESY) and Fermi@Elettra were reported. The main results of the workshop lay in the area of stating the requirements and the need for development of the HHG source. Issues on intensity, pulse length, polarization, position and pointing, timing and stability were addressed. Another urgent field was how the APT (Attosecond Pulse Train) will influence the seeding efficiency and also the tuneability of the HHG source.

The workshop attracted over 50 participants from all the European FEL laboratories involved in seeding of FELs.



Participants at the workshop, 14-16 June 2010.
Photo: Annika Nyberg

Lärardagar på MAX-lab, Lund, 14-16 June.

Encouraged by the feedback from the study day arranged for local high-school teachers in August 2009, a three-day workshop was organized 14-16 June 2010 in collaboration with MAX-lab and National Centre for Education in Physics.

A total of 30 physics teachers (29 from Sweden and 1 from Finland) came to MAX-lab in order to learn how synchrotron radiation is produced and what kind of studies can be done with it. The teachers also got an opportunity to perform experiments on beamlines with MAX-lab's staff, and they presented their results the last day.

Debra Belsey – from Lloydminster Comprehensive High School in Lloydminster, Canada – was also invited to tell the participants about her experiences from working with high-school students at Canadian Light Source, and this the participants considered particularly inspiring. The feedback collected afterwards from participants was outstanding and detailed plans to continue outreach activities aimed for high-school teachers and students are underway.

FEL 2010, Malmö, 23-27 August.

The 32nd International Free Electron Laser conference was organized by MAX-lab in Malmö 23-27 August 2010. Over 300 registered participants gathered for five days at Hilton Malmö City to hear and follow the latest development on Free Electron Lasers. 63 presentations were held, half of them by invited speakers and also including four tutorials on focus area in the field. Highlights included reports of the first lasing of FELs around the globe during the past year, reports from the key projects such as LCLS, FLASH, SPring8, SwissFEL and Fermi including their upgrade programs. Other areas that were covered included synchronization issues, diagnostics, low charge operation and much more. The previous conference was held in Liverpool, UK, and after Malmö the event will move to first Shanghai in 2011 and Kyoto 2012.

Proceedings from the conference are available at: http://fel2010.maxlab.lu.se/FEL2010_proceedings/index.htm

The 23rd MAX-lab Annual user meeting, Lund, 8-10 November.

The 23rd Annual User Meeting for MAX-lab users was held at Scandic Star Hotel in Lund on 8-10 November 2010. The meeting was as previous years jointly organized by MAX-lab and the MAX-lab Association for Synchrotron Radiation Users (FASM). This year there was a new record of 342 registered participants and 26 commercial exhibitors.



The first day of the user meeting was primarily dedicated to the present activities at the lab, and included status reports, user highlights and two plenary lectures. Prof. Peter Schurtenberger from Lund University gave a talk on Responsive Colloids – From Model Atoms to Novel Hybrid Nanomaterial, while the Director of SOLEIL, Prof. Michel van der Rest, talked about Current Status and Future plans for SOLEIL. In the evening a poster session was held at MAX-lab. At the latter, a Best Student Poster Award sponsored by SPECS GmbH, was rewarded to Mr. Mo Segad at the Department of Theoretical Chemistry, Lund University. The title of his poster was: Structure and swelling properties: experimental studies of bentonite (MX-80)-Water Systems.

The next two days were very much focused on MAX IV. The current status of the project was presented in several talks. In addition there were two more plenary lectures; Dr. Rolf Follath from Helmholtz-Zentrum Berlin talked about Beamlines for Low Emittance Rings, a subject of greatest relevance for MAX IV. Prof. Marek Stankiewicz, at the Jagiellonian University in Kraków talked about the new Synchrotron Radiation Facility in Poland. This storage ring will have the same design as the 1.5 GeV storage ring of the MAX IV facility.

After the common session the meeting was divided into several parallel workshops related to issues of importance for MAX IV. A majority of these workshops were focused on some of the different suggested first phase beamlines on MAX IV. In addition two more general workshops were organised. A workshop titled Pixel Detectors for MAX IV covered different aspects of position sensitive detectors for synchrotron radiation applications. A second session was the IDMAX2010 workshop, with the heading Insertion Devices for Rings and Linacs.

Further information: www.maxlab.lu.se, search for User Meeting

IDMAX2010, Insertion Devices for Rings and Linacs, Lund, 9-10 November.

A workshop called IDMAX2010 about insertion devices for synchrotron radiation production was organised 9-10 November 2010 in connection with the MAX-lab Annual User Meeting.



The workshop was a success and the list of speakers included in total 23 experts from world leading laboratories in Asia, Europe and the USA. There were also presentations from 5 different companies manufacturing insertion devices during the workshop.

Among the topics discussed during the workshop were existing insertion devices, plans for future installations, research and development activities on novel insertion devices, fast polarization switching, chicanes and canted undulators, operational aspects, and manufacturing aspects of insertion devices.

The presentations given at the workshop can be found on http://www.maxlab.lu.se/usermeeting/2010/sessions/IDmax2010/index.html







Mikko-Heikki Mikkelä, University of Oulu, and Samuli Urpelainen, fine tuning the evaporation oven of the exchange metal cluster source at beamline I411, 11 October 2010. Photo: Annika Nyberg

List of Users at MAX-lab January 2010 – December 2010

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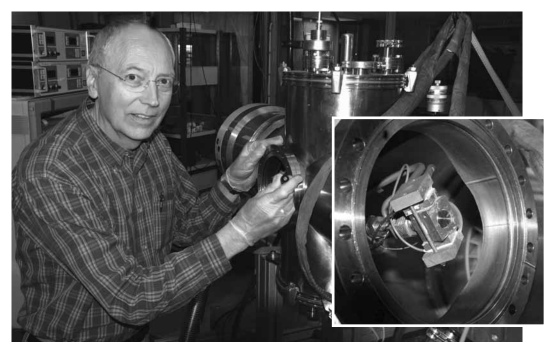
Electronics

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Leif J. Sætre, from the University of Bergen, setting up a cluster experiment at beamline I411, 11 October 2010.

Photo: Annika Nyberg



Knut Børve, Leif Sætre's colleague from University of Bergen, is switching on the pumps on the cluster source, with which they produce and study hydrogen-bonded clusters of simple inorganic and organic molecules.

Photo: Annika Nyberg

Accelerator Physics and Machine Development



Caj Lundquist (MAX-lab), Göran Hellqvist (Fojab), Janis Kursis (Fojab), and Sara Thorin (MAX-lab) discussing final adjustments on the Linac building at a project group meeting at ML4, 6 December 2010. Photo: Annika Nyberg

Machine and Accelerator Physics

There are three storage rings at MAX-lab; MAX I, MAX II and MAX III. All three rings are used for synchrotron radiation production and the MAX I ring is also used for experiments in nuclear physics, working as a pulse-stretcher. All three rings are fed by the 400 MeV MAX injector.

A test-FEL is placed inside the MAX II storage ring which also uses the MAX injector as the electron source.

The construction of **the MAX IV facility** has started in the fall of 2010. A necessary condition for this was the finalization of the first version of the Detailed Design Report (http://www.maxlab.lu.se/maxlab/max4/DDR_public/index.html) which defines the accelerator complex. The Machine group and the department of Accelerator Physics have been heavily involved in the construction work of the MAX IV facility.

The characterization of the electron gun in the FERMI (Trieste) project has been finalized.

The MAX injector is of recirculated linac type and is primarily used for injection into the three storage rings and for free electron laser (FEL) experiments.

The MAX I ring is injected at 190 MeV and the MAX II and MAX III rings at 380 MeV.

This injector consists of an RF gun and two S-band linac sections 5.2 m long equipped with SLED cavities. A recirculation magnet system is used to double the electron energy. These linacs are now conditioned to a little more than 100 MeV energy gain each and the maximum energy gain for both linacs is 210 MeV. By recirculating the electron beam once through the linacs, a maximum electron energy of 420 MeV can be reached.

The RF electron gun used for injection is equipped with a thermal cathode. This gun is quite reliable, but the beam quality is rather poor, due to space-charge effects during the early acceleration in the gun. The performance of this thermionic gun is however quite sufficient for injection into the rings.

For the FEL runs, the thermionic BaO cathode is used as a laser-driven photo-cathode.

The MAX injector has been quite reliable and negligible time is lost due to malfunctioning of this accelerator.

The MAX I ring is aging and suffers from reduced beam life-time in the storage mode due to vacuum problems. Since most of the SR experiments now have been transferred to the MAX III ring and also due to man-power limitations, a vacuum upgrade is of lower priority.

In the pulse-stretching mode for nuclear physics, the deterioration of the vacuum system is of no importance and the properties of the electron beam in this application are now rather close to the design values and both the ring and injector are working under stable conditions.

The MAX II ring was working as well as a low-energy injected storage ring can do until November 2010 with a down-time around 2%. In mid November, we had a break-down of the transformer in the dipole power supply, which stopped operation for five weeks until a new transformer could be delivered and installed.

The electron beam stability is in the micrometer range, 300 mA can be injected twice a day and with a beam life-time of 5-6 Ah, some 160 mA remains at the next injection. The mean current is now typically 180-200 mA. 8% of the total time is spent on injection and ramping of the lattice magnets, undulators and the superconducting wigglers.

The MAX III ring is now operating with two beamlines. Typically over the year the availability was around 95%, where 3% of the total time were taken away by injections and 2% by down-time. However, during two weeks in October the availability was down to 50% due to a tricky vacuum leak. The beam life-time has improved to some 1-1,5 Ah, due to an active blow-up of the vertical emittance. Work is in progress increasing the longitudinal emittance as well.

The test Free Electron Laser at MAX-lab has produced first light. The system is built around the 400 MeV linac injector utilizing two undulators placed inside of the MAX II storage ring. The system is seeded by a 263 nm Ti:Sapphire laser and can produce fully coherent radiation in harmonics of the seed. During 2010 coherent harmonics up to 6th order (42 nm) have been produced in linear mode and up to 4th order (65 nm) in circular mode. Photon pulse lengths are in the order of 500 fs. The peak intensities in the harmonics exceed the spontaneous emission by more than an order of magnitude and a significant line width narrowing is observed.

References:

First Results of Coherent Harmonic Generation at the MAX-lab Test FEL, S. Werin et al, Proceedings FEL2010, Malmö, Sweden (2010).

On-Line Arrival Time and Jitter Measurements Using Electro-Optical Spectral Decoding, N. Čutić et al, Proceedings FEL2010, Malmö, Sweden (2010).

Machine Parameters

Inject	tor	linac
HHE.	l CH	LIIIAI.

Max. energy420 MeVPulse current50 mAPulse length50 nsEnergy spreadNot verifiedEmittanceNot verified

MAX I RING Storage mode

Max. energy 550 MeV
Max. circ. current 300 mA
Hor. emittance 40 nm rad
RF 500 MHz
Bunch length (FWHM) 80 ps

Beam lifetime 4 h (now decreased to some 2 h)

Pulse-stretcher mode

Electron energy 144, 188 MeV

Duty factor 75 % Stretched pulse current 20 nA

MAX II RING

Max. energy1.5 GeVMax. circ. current290 mAHor. emittance8.8 nm radRF100 MHz

Beam lifetime 5-6 Ah (25-30 h at 200 mA)

MAX III RING

Max. energy 700 MeV
Max. circ. current 300 mA
Hor. emittance 14 nm rad
RF 100 MHz
Beam lifetime 1-1.5 Ah

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Coherent Harmonic Generation at the MAX-lab test FEL

N. Čutić, F. Lindau, S. Thorin, S. Werin, Max-lab, Lund University, Sweden, C. Erny, A. L'Huillier, E. Mansten, Dep. of Atomic Physics, Lund University, Sweden, J. Bahrdt, K. Holldack, Helmholtz Zentrum Berlin, Germany

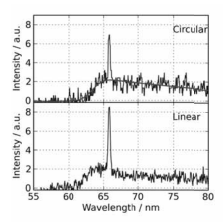


Figure 1. Coherent 4th harmonic signal in both linear and circular mode with fitted spontaneous undulator radiation in circular mode (red solid line).

INTRODUCTION

The test FEL is a collaboration project between MAX-lab and BESSY (now the Helmholtz Zentrum Berlin) to study, develop and build knowledge on seeding as a technique for free electron lasers. The existing injector at MAX-lab is used as the electron source, with the gun operated in RF photocathode mode. Pulses containing 25-40 pC and compressed to a duration shorter than 1 ps are normally used.

Two undulators, provided by BESSY, and a magnetic chicane form an optical Klystron. In the first undulator, an energy modulation is seeded on the accelerated electrons with a 0.5 ps long laser pulse with a wavelength of 263 nm. The second undulator - the radiator - is of APPLE II type, and can emit coherent radiation at harmonics of the seed wavelength with variable polarization. A horizontally focusing collection mirror and a spectrometer (2400 l/mm grating) with a liquid nitrogen cooled CCD array are used to record the CHG signal.

See [1] for details on the setup.

RESULTS

Operation

Coherent Harmonic Generation (CHG) was set up in a multi-step process. The spontaneous radiation from the modulator undulator and radiator undulator was recorded to assure the

proper gap settings calibrated to the seed laser wavelength. This was necessary as a precise electron beam energy calibration was not at hand. The electron beam orbit was also optimised for best spectrometer signal. This was followed by transverse overlap where the seed laser was placed on top of the electron beam on two YAG screens placed before and after the modulator undulator. Temporal overlap was adjusted via the Electro Optical system (EO). After this the CHG signal was sought with small adjustments to the temporal overlap by use of the seed laser delay line. This was followed by an optimisation of the CHG signal by adjusting the transverse overlap. Unfortunately, the system has a tendency to drift out of overlap over a 10 minute period. A temporal feed-back system based on the EO signal partly cures the problem [2]. Relative phase drifts between the RF gun and the linac remain though, which makes prolonged operation troublesome without tuning.

Coherent Harmonic Generation

CHG has been recorded from the fundamental (1st harmonic at 263 nm) to the 6th harmonic (45nm) in linear mode (see figures 3 & 4). The 2nd and 4th harmonics (133 nm & 66 nm) have also been recorded in circular polarization mode

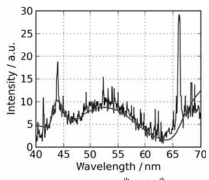


Figure 2: Linear coherent 4th and 6th harmonic signal with fitted spontaneous undulator radiation. (Radiator set to 131 nm and the harmonics are thus 2nd and 3rd of the radiator wavelength).

(figure 1). The spectra show the spontaneous radiation background and the CHG peak placed at exactly the harmonic of the seed laser. In figure 2 the radiator undulator was put to a fundamental wavelength of 133 nm where the 2nd and 3rd harmonic of the undulator equalled the 4th and 6th harmonic of the seed.

The spontaneous undulator radiation was calculated using SPECTRA [3] where the gap and

opening angles where fitted to achieve the recorded spontaneous signal. Thus the CHG flux could be calculated. The results show that in the 4th harmonic (66 nm) we have 390.000 photons per pulse (at 25 pC in circular mode) which corresponds to 1.1 pJ/pulse.

A modulator undulator gap scan was made (figure 3) and the resulting CHG signal at the 2nd harmonic (133 nm) was recorded. The CHG signal is present within a larger range than the full undulator linewidth which can be explained that the modulation is efficient even if it only works on a shorter part of the undulator, thus an increased linewidth.

Laser energy and modulation

The laser energy and the chicane magnet system strength were scanned. This influences the microbunching in the CHG process. The seed laser energy can be adjusted by a rotatable polariser. It is favourable to induce a large energy spread by the seed laser compared to the natural energy spread in the electron beam. With a larger induced energy spread the microbunching in the chicane will be quicker and a lower strength is needed. Also over bunching will occur quickly. Theoretically the chicane setting should be a sensitive knob to optimise the CHG output. However instabilities and drift in the system makes these kind of optimisations difficult.

In figure 4 the qualitative results of this kind of scans are shown. A stronger chicane (3 A) quickly reaches higher CHG signal when the laser energy is increased. The signal then drops when the laser is further increased, which is a sign of overbunching.

A lower chicane setting (1.3 A) requires more laser energy to microbunch the electrons, but the laser was not intense enough to over bunch the system.

The intermediate chicane settings give

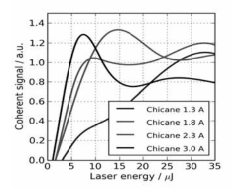


Figure 4: Scan of the laser energy (in the undulator) for different excitations of the microbunching chicane.

intermediate results, but only qualitatively.

There is no definite sign of a higher CHG signal using more laser energy. This is an indication that the natural energy spread is not a limiting factor at the 2nd harmonic (133 nm).

SUMMARY

The collaboration between MAX-lab and HZB/Bessy has successfully built and commissioned the test FEL at MAX-lab for Coherent Harmonic Generation. An existing linac system has been upgraded and CHG in linear mode in the 1-6th harmonic (263-42 nm) and (2nd and 4th harmonic (133 and 66 nm) in circular mode has been achieved. We believe that circular polarised coherent radiation at 66 nm has never been observed from an accelerator source before.

The main goals of the system have been achieved and some features even beyond the goals. One of the main purposes of the facility was to gain experience of operation, diagnostics and FEL technology to build a base for a future FEL facility within the MAX IV project.

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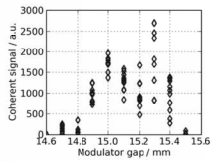
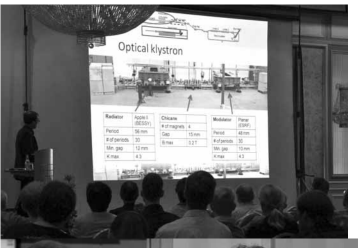


Figure 3. Coherent 2nd harmonic signal as a function of modulator gap.

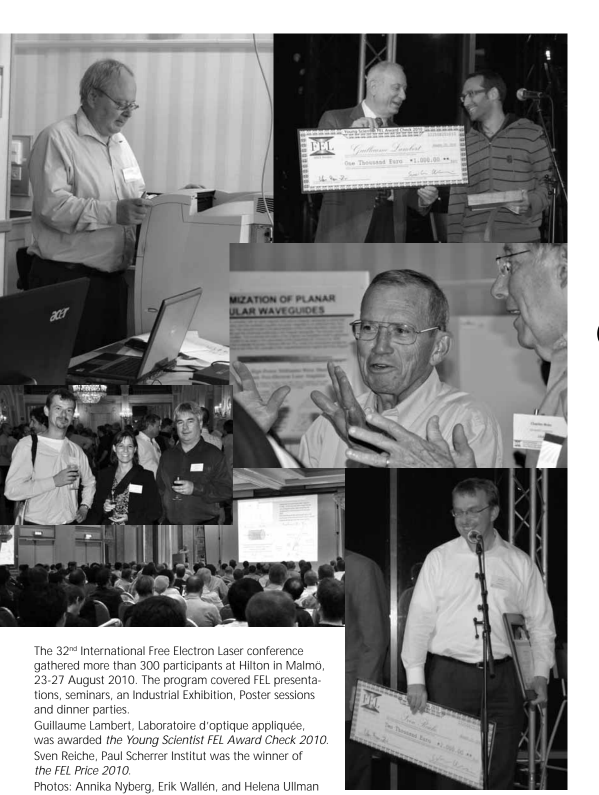




- MMX Sweden -

http://fel2010.maxlab.lu.se





ON-LINE ARRIVAL TIME AND JITTER MEASUREMENTS USING ELECTRO-OPTICAL SPECTRAL DECODING

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INTRODUCTION

A crucial component of the test-FEL is a device based on electro-optical spectral decoding (EOSD) which allows on-line monitoring of the bunch compression and the electron bunch arrival time relative to the seed laser pulse. Since the electron bunch and the seed laser pulse are of subpicosecond duration a technique that allows timing measurements with better precision than what can be achieved with photodiodes (200 ps) is needed. Electro-optical schemes using the interaction of a terahertz field created by the traveling electron bunch and a laser pulse passing through a crystal have been developed (first in laser based sources of THz radiation) and later modified and applied for measurements of bunch arrival time and duration at accelerators [1, 2]. These technique have shown to be robust enough and comparatively cheap since a required laser pulse for seeded facilities is readily available without major extra costs.

EXPERIMENTAL SETUP

The EOSD system consists of 3 major parts and operates using a small sample ($< 1 \mu J$) of the amplified seed laser pulse (before it is tripled to UV). The first part of the system is in the seed laser hutch and it consists of a separate stretcher built for IR pulses, focusing optics (telescope focusing onto the crystal inside the EOSD chamber, 6.5 m focus), delay stages (a main delay stage controlling the delay of both pulses relative to the electron bunch, and a UV delay stage shifting the UV pulse relative to the IR pulse) and a UV-IR overlap monitor based on difference frequency generation (DFG). The second part is the EOSD chamber which is the only part of the EOSD system placed in vacuum. The chamber holds the crystal(1 mm thick ZnTe cut in the (100) plane), mirror and photodiode (for rough timing) on a translation stage (used to position the crystal transversally arbitrarily close to the passing electron bunch). The third part is the EOSD detector consisting of polarization optics and a spectrometer which is placed outside of the vacuum system right next to the chamber. Figure 1 shows what is described as the EOSD chamber and EOSD detector.

To stabilize the drifts a feedback routine is built that controls the position of the peak of the signal so that it is always on desired position (the laser pulse is at desired timing relative to the electrons). This is done with a PI controller moving the main delay stage (Thorlabs 150 mm).

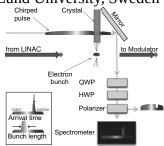


Figure 1: The EOSD chamber and detector. The IR chirped pulse (3.3 ps FWHM, 5 nm FWHM bandwidth, 790 nm central wavelength) passes through the crystal and is reflected out of the chamber to the detector. The detector consists of polarization optics (Quarter-wave plate, half-wave plate and Glan-laser polarizer) which filters the polarization of interest and the spectrometer. Based on the position of the peak in the signal on spectrometer it is possible to determine the arrival time of the electron bunches relative to the IR pulse and estimate the length of the electron bunch from the width.

To reduce jitter caused by the 50 Hz grid frequency, the triggering of devices should be synchronized to this frequency. The synchronization is done by a microcontroller sending a trigger pulse every 5 grid cycles to obtain the desired 10 Hz pulse rate for the laser.

MEASUREMENT AND RESULTS

Jitter measurements and feedback Figure 2 on left shows three images each showing a series of spectra through time. The frequencies on the right span to 1 Hz because the repetition (sampling) frequency is 2 Hz. The top two images belong to measurement done without any feedback. The middle two are with the 50 Hz lock is active. And the lower two are with 50 Hz lock active and a PI controller controlling the position of the peak at certain point. Noticeable improvement in stability is visible. Turning on the 50 Hz lock reduces the RMS jitter from 915 fs to 425 fs. It also removes the 0.45 Hz peak which was an influence of the power grid to the system probably on higher frequencies that was undersampled. This still

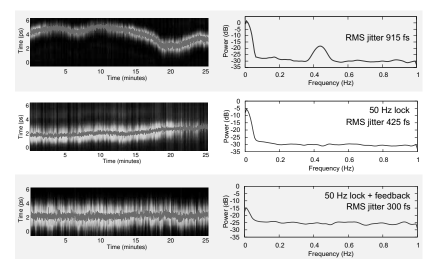


Figure 2: Measurements of the long term drifts and jitter (left) and the corresponding power spectra of the arrival time (right). On the left side each spectrum captured (about 3300 of them, corresponding to 25 minutes) is converted to time and shown vertically using the false coloring of the signal's intensity. The central position is calculated and shown in red overlaid curve. The right side shows power spectra of these red curves. Top - without feedback; middle - with 50 Hz lock; bottom - with EOSD feedback and 50 Hz lock.

leaves majority of long term drifts below $0.05~{\rm Hz}$ which are lowered $10~{\rm dB}$ more by the delay stage feedback giving final RMS jitter of $300~{\rm fs}$.

Drifts and bunch length Measurements of the bunch length are done simply by determining the width of the EOSD signal. Their main limitation is the large thickness of the ZnTe crystal. Larger thickness of the crystal influences the effective cutoff frequencies of the system (makes the EOSD "slower" due to mismatch between the propagation of the THz and optical pulses through the crystal) but improves the signal strength (larger thickness means more polarization rotation).

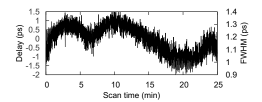


Figure 3: Correlation between the arrival time of electron bunches and the width of the EOSD signal (approximately bunch length, see text). This correlation points to the accelerator as the main cause of drifts.

Figure 3 shows the correlation between the measured pulse width and arrival time for the case without any feedbacks (overlaid red curve from top left in Fig. 2). Assuming the Gaussian shape of the bunch, simulations show that our setup overestimates the real bunch length by 6% for shorter measurements (1 ps) and 3% for longer (implying the bunch width to be ranging from 0.95 ps to 1.26 ps FWHM). Obvious correlation between the arrival time of electrons and their compression is a sign that the drifts are coming from the linac and not the laser system. If the laser system was the main cause of drifts the compression would not correlate so good with the drifts.

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Upgrade of the magnetic structure of undulator I311

Erik Wallén

INTRODUCTION

In the summer of 2010 the original hybrid type magnet arrays of the I311 undulators were replaced by pure permanent magnet arrays. Figure 1 shows the upgraded I311 undulator. The minimum gap has been reduced from 22 mm to 16 mm and the period length reduced from 66 mm to 54.4 mm.



Figure 1: The I311 undulator with the upgraded magnetic structure in July 2010.

THE ORIGINAL MAGNETIC STRUCTURE OF 1311

The I311 undulator [1] at the the 1.5 GeV MAX II storage ring [2] was installed in 1995-1996. The original I311 undulator was of hybrid type with a period length of 66 mm and total total length of 2.6 m. The magnetic blocks have shown signs of cracking and some steel shims have either fallen off or moved during the last few years.

The I311 undulator was removed from the MAX II ring tunnel on July 5, 2010 and set up at the magnet measuring benches to be measured before the old magnet assemblies are removed to document the status of the undulator after 15 years of operation. Measurements were done at 23, 30, 85 and 66 mm gap as reported in [3]. Figure 2-6 contain the measured phase errors and multipoles of the original magnetic structure. Figure 2-6 also include the case when the

first and second field integrals are removed by correction coils in front of and behind the undulator in order to better represent the way the undulator was used in the MAX II ring.

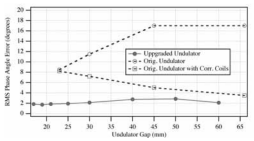


Figure 2: The phase errors for the original and upgraded 1311 undulator.

THE UPGRADED MAGNETIC STRUCTURE OF 1311

The upgraded I311 undulator is of pure permanent magnet type to avoid the non-linearities of the iron poles in a hybride type undulator. The size of the magnetic blocks have been adjusted to get the same forces at minimum gap in the new and original configuration [4]. The minimum gap is 16 mm and the period length is 54.4 mm with the upgraded magnetic structure. The main parts of the support structure has not been changed. The total number of poles is 98 and the fundamental harmonic of the emitted synchrotron radiation covers the energy range 40-350 eV.

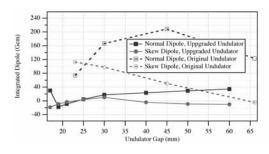


Figure 3: The dipole field integrals for the original and upgraded I311 undulator.

The shimming process was carried out in July 2010. Af-

ter the shimming the undulator properties at 17, 19, 21, 25, 30, 40, 50 and 60 mm gap were measured as reported in [5]. Figure 2-6 contain the measured phase errors and multipoles of the upgraded magnetic structure. The phase angle error is below 3° at all gaps. The fundamental photon energy at 17 mm gap is 42.68 eV calculated with a filament electron beam and with the gap lowered to 16 mm the fundamental photon energy will be below 40 eV.

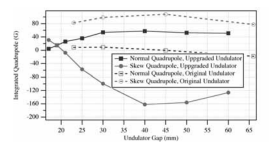


Figure 4: The quadrupole strength for the original and upgraded I311 undulator.

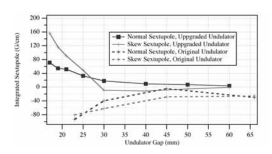


Figure 5: The sextupole strength for the original and upgraded I311 undulator.

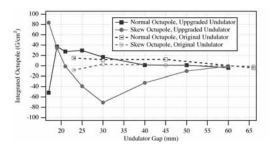


Figure 6: The octupole strength for the original and upgraded I311 undulator.

PHOTON FLUX AT THE EXPERIMENTAL STATION AT BEAMLINE 311

The photon flux at the sample at the experimental station at beamline 311 with the original and upgraded magnetic structure is shown in Figure 7 [6]. As can be seen in Figure 7, the flux at the sample has increased an order of magnitude over a wide range of the spectrum with the upgraded compared to the original magnetic structure.

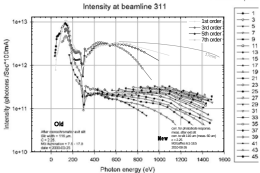


Figure 7: Flux at the sample at the experimental station at beamline 311[6].

ACKNOWLEDGEMENTS

I would like to thank Ingvar Blomqvist for the extensive and highly appreciated help and advice during the work with the magnetic and mechanical design, shimming, and software development during the work at the magnetic measurement system.

I would like to thank Håkan Svensson for advice on the mechanical design, help with drawings, contact with companies, and taking care of orders, deliveries, and transports during the project.

I also would like to thank Bengt Sommarin for mounting 400 magnet blocks in magnet holders and mounting the holders on the girders with high precision.

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The I311 undulator got new magnets in July 2010. The shimming, or magnetic field strength tuning, was carried out at the magnetic measurement system in the MAX II hall. Erik Wallén next to the upgraded undulator.

Photo: Bengt Sommarin

Synchrotron Radiation

Synchrotron Radiation Research

General

During 2010 the MAX I, MAX II and MAX III storage rings have been operated for synchrotron radiation research for 22, 40 and 40 weeks, respectively. About 760 scientists have been performing experiments during this time period. Of these about 60 % came from abroad. The scientific projects include experiments in atomic and molecular physics, solid state physics, surface physics, material science, chemistry, life science and environmental science. Reports from the users on recent experimental results are given on the following pages.

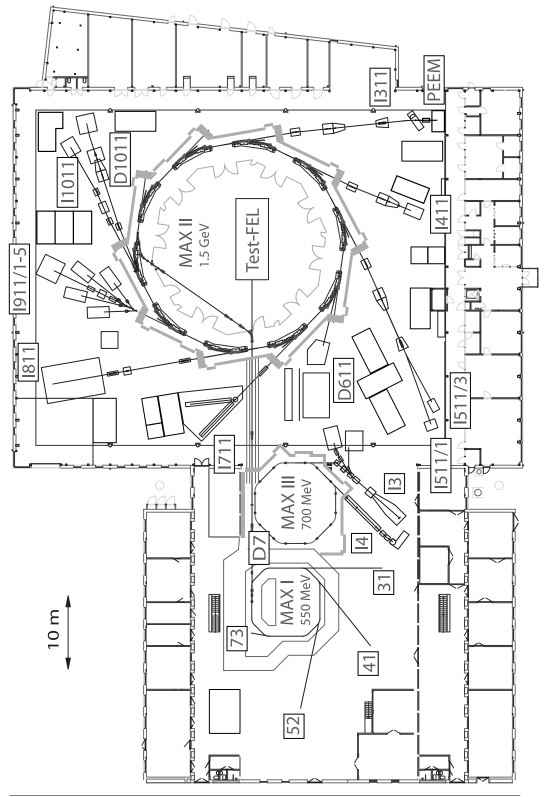
Most beamlines on MAX I utilize radiation from bending magnets which provide radiation with a critical energy of 300 eV. In addition a hybrid undulator (period length 75 mm) covering the photon energy range 15 to 150 eV is used for a spectromicroscopy beamline. In total there are four beamlines in operation. The characteristics of the beamlines are summarized in the table on page 59 and a short description of each beamline is given below.

The MAX II storage ring is equipped with three planar undulators and one EPU for the VUV and soft X-ray regions and three multi-pole wigglers for the X-ray region. One conventional multi-pole wiggler beamline is used for small molecule crystallography and powder diffraction. Of the two superconducting multi-pole wigglers, one is used for a materials science beamline for absorption and diffraction experiments and one is used for a system of five independent beamlines mainly for protein crystallography and small angle X-ray scattering. The undulators serve beamlines with a variety of spectroscopic techniques such as X-ray absorption (including circular dichroism), X-ray emission, X-ray photoelectron spectroscopy and photoemission electron microscopy in the VUV and soft X-ray regions. A fourth spectroscopy beamline is installed on a bending magnet port where circularly polarized radiation can be used. A bending magnet beamline is also used for time resolved X-ray diffraction.

Two undulator beamlines are operational at the 700 MeV MAX III storage ring; one EPU covers the low energy region 5 – 50 eV and planar undulator covers the energy range 13 – 200 eV. The undulators serve beamlines for angular and spin resolved photoelectron spectroscopy. A third beamline for infra-red microspectroscopy, utilizing bending magnet radiation, is under construction.

More detailed descriptions of these beamlines are found at the end of this chapter and in the tables on pages 60 - 62.

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MAX I Beamlines

(see www.maxlab.lu.se for more details).

Beamline 31 utilizes undulator radiation in the energy range 15 to 150 eV for scanning photoelectron microscopy with a lateral resolution in the micrometer range.

The experimental system consists of separate analyzer and preparation chambers accessible via a transfer arm. The analyzer chamber, which also holds an ellipsoidal focusing mirror, is equipped with a hemispherical sector electron energy analyzer (VG CLAM2) and a piezo-driven sample scanning stage. The preparation chamber is equipped with LEED, ion sputtering gun, sample storage magazine and a number of optional ports for user owned sample preparation accessories.

Beamline 41 is used for angle resolved photoelectron spectroscopy on solids in the photon energy range from 15 to 200 eV using a toroidal grating monochromator (TGM). The beamline is suited for measurements of both valence bands and shallow core levels.

The experimental station consists of an analyzer chamber with a goniometer mounted electron energy analyzer (VSW HA50), a sample storage chamber and a sample introduction chamber. The analyzer chamber is equipped with LEED, ion sputtering gun, gas-inlet system and a number of optional ports for user owned sample preparation accessories.

In addition to this basic set-up a molecular-beam epitaxy (MBE) system with six Knudsen cells and a RHEED optics is available for the growth and in situ studies of III-V compound semiconductors.

Beamline 52 consists of a normal incidence monochromator working in the photon energy range 5 to 30 eV. The beamline is equipped with a differential pumping stage that makes it useful for measurements on gases as well as on solids. During 2010 no user projects have been carried out on this beamline, they have migrated to the new I3 beamline on MAX III. Instead beamline 52 has been used for outreach activities for high-school teachers and students.

Beamline 73 is used for spectroscopy and microscopy in the far, mid and near infrared region (10-12 000 cm⁻¹). It is equipped with two different Fourier transform spectrometers.

A Bruker HR 120 is used for high resolution, 0.001 cm⁻¹, spectroscopy. The set up has several options for introducing gases, liquids and solid samples. A gas cell, usable between 90 and 300 K, with a variable optical path length (maximum 120 m) is connected to the spectrometer.

A Bruker 66v/S spectrometer along with a Hyperion 3000 microscope is used for chemical imaging and mapping with a spatial resolution down to the diffraction limit. A 128x128 element focal plane array detector is available. The microscope can operate both in transmission and reflection mode. Additional objectives for gracing angle and ATR measurements are available as well.

MAX II Beamlines

(see http://www.maxlab.lu.se for more details).

Beamline I311 is an undulator based VUV, soft X-ray beamline for high resolution X-ray Photoemission Spectroscopy (XPS) and X-ray Absorption Spectroscopy (XAS) and photoemission electron microscopy. The monochromator is a modified SX-700 type PGM with spherical optics and a movable exit slit, which allows a high flexibility concerning the interplay between photon flux, resolving power and higher order suppression. In summer 2010 the twelve year old magnetic structures in the undulator were replaced by new ones with 45 periods and a period length of 54.4 mm. The undulator now covers the energy range from 43 eV up to about 1500 eV.

The main experimental station consists of separate analyzer and preparation chambers accessible via a long-travel manipulator. The preparation chamber includes the usual equipment for preparation and characterization of surfaces (ion sputtering gun, LEED optics etc.). A hemispherical electron energy analyzer (SCIENTA SES200) is used for photoelectron spectroscopy and XAS in Auger yield mode.

A SPELEEM instrument for photoemission electron microscopy is installed downstream from the main experimental station. This microscope has a spatial resolution better than 10 nm in the LEEM mode and 30 nm in the PEEM mode. It can also perform energy filtered XPEEM with a bandwidth of 300 meV in imaging mode, routinely achieving a lateral resolution of 30 nm.

Beamline 1411 is based on an SX-700 type PGM and an undulator source (43-period, 59 mm period length) that covers the photon energy range 40 eV to about 1500 eV. The end-station has the unique versatility of being able to handle solid, liquid and gaseous samples. Thus the beamline is well suited for high-resolution electron spectroscopy on free atoms and molecules as well as for studies of liquids and non-UHV compatible solids.

The experimental system consists of separate analyzer and preparation chambers accessible via a long-travel manipulator. The analyzer chamber is equipped with a hemispherical electron energy analyzer (SCIENTA R4000) which can be rotated around the incoming beam for polarization dependent measurements. In front of the experimental station a one-meter long section of the beamline can host other types of equipment for atomic and molecular spectroscopy, e.g. ion-electron coincidence detectors.

A laser system for two-colour experiments is also available.

Beamlines I511/1 and I511/3 are used for XAS, XPS, and X-ray Emission Spectroscopy (XES) in the VUV and soft X-ray range. The two beamlines utilize a common undulator and monochromator with a flip-mirror placed immediately after the exit slit to direct the radiation alternately into two experimental stations. The undulator has 49 periods and a 52 mm period length originally giving a photon energy range of 50 to about 1500 eV. The monochromator is the same type of modified SX-700 monochromator as used on beamline I311.

During 2010 beamline I511/1 has been used for surface studies under UHV using an end station equipped with a hemispherical electron energy analyzer (SCIENTA R4000) for XPS and an electron yield detector for XAS measurements. In December 2010 this end station was replaced by a new station aimed at surface studies under near-ambient pressure conditions. The new station is equipped with a SPECS Phoibos 150 NAP analyzer for XPS, and XAS in Auger yield mode. Experiments at pressures up to some mbar are carried out using a dedicated high pressure cell, which can be moved in and out of the analysis UHV chamber.

Beamline I511/3 is equipped with a grazing incidence grating spectrometer for XES and can handle non-UHV compatible solids. The analysis chamber can be rotated around the incoming beam which makes it possible to utilize the linear polarization of the radiation.

Beamline D611 is a bending magnet beamline dedicated to time-resolved studies. We are exploiting the fact that MAX II is a pulsed source operating at 100 MHz. The duration of the pulses has been measured to be approximately 350 ps using a streak camera. The beamline has a toroidal focusing mirror and a double crystal mononchromator. A laser providing pulses with durations of 20-30 fs has

been synchronised to the ring, and a streakcamera yielding a temporal resolution of about 500 fs is available. The temporal resolution in the experiments does not depend on the relative jitter between the laser and the synchrotron (10 ps) but mainly on the jitter between the streak camera and the laser. The laser operates at a maximum repetition rate of 10 kHz which sets the data accumulation rate. Experiments can be performed in air or in vacuum (10-6 mbar). More information about these activities can be found at: http://www.atomic.physics.lu.se/research/ultrafast_x_ray_science/

Beamline I711 has been used for powder diffraction and small angle X-ray scattering (SAXS) during 2010. It utilizes a 13-period, 1.8 T, multipole-wiggler and is designed to operate in the 0.8 Å (15.5 keV) to 1.6 Å (7.8 keV) region. The beamline has a vertical focusing mirror and a focusing single crystal monochromator working in the horizontal plane. This design sacrifices easy tunability and high energy resolution for high flux at the sample.

In autumn 2010 a new experimental station was installed. This se-tup includes a 4-circle diffractometer with kappa geometry, capable of doing both single crystal and powder diffraction using a large area CCD detector. The SAXS experiments have been moved to a new set-up on beamline 1911-4 with user operation starting in spring 2011.

Beamline I811 is intended for materials science research using X-ray absorption spectroscopy (XAS) and X-ray diffraction (XRD). It is based on a super-conducting multi-pole wiggler insertion device that produces high-flux photons in the energy range 2.4 - 20 keV (0.6 - 5 Å). The design is based on adaptive optics where the beam is collimated and focused vertically by cylindrical bendable first and second mirrors, respectively. Horizontal focusing is obtained by sagittal bending of the second monochromator crystal. The typical flux in a 1 x 1 mm² beam spot on the sample is 5 x 1011 photons/ sec. One experimental station is used for XAS research with detectors for transmission and fluorescence yield techniques. A second station is equipped with a diffractometer for surface, interface and thin-film crystallography.

Beamlines 1911/1-5 are used mainly for macromolecular crystallography and small-angle X-ray scattering (SAXS) but also some other diffraction experiments and for courses. The beamlines use a superconducting multi-pole wiggler. The central beamline 1911-3 is tunable in the range 0.75 – 2.0 Å and is optimized for MAD experiments. The beam is vertically collimated by a Rh-coated mirror, monochromatized by a Si(111) double-crystal monochromator and focused by a Rh-coated toroidal mirror. The optics for the four side stations consists of horizontally focusing monochromator crystals and vertically focusing curved multilayer mirrors providing fixed wavelength beams. 1911-3 has a new experiment set-up since 2010 with a microdiffractometer, a mini-kappa, a large-capacity automatic sample changer and a 225 mm CCD detector. Two of the side stations, 1911-2 (wavelength 1.04 Å) and 1911-5 (wavelength 0.97 Å) are equipped with single axis diffractometers and 165 mm CCD detectors. The new SAXS station, 1911-4, has a fixed wavelength beam (0.91 Å) and is equipped with a mar165 detector.

Beamline D1011 is a bending magnet beamline covering the energy range 30 to 1500 eV. An adjustable local bump of the electron beam provides out of plane radiation. This makes magnetic circular dichroism (MCD) possible in addition to photoemission and photoabsorption using linearly polarized light.

The monochromator is a modified SX-700 PGM of the same design as the monochromator on beamline I411. The experimental system consists of separate analyzer and preparation chambers accessible via a long-travel manipulator. The analyzer chamber is equipped with a SCIENTA SES200 electron energy analyzer (with a lens of SES2002 type) and an MCP detector for electron yield measurements. The preparation chamber is equipped with LEED, ion sputtering guns, gas-inlet system and a number of optional ports for user owned sample preparation accessories.

A second experimental station receives radiation that is let through the first station and re-focused by a KB mirror system. This station is specifically designed for NEXAFS, XMCD and soft X-ray reflectivity experiments. Measurements can be performed under static magnetic fields of up to 500 G. One of the unique features of the station is to offer element specific reflection-based hysteresis measurements.

Beamline I1011 is used for studies of magnetic materials using magnetic circular dichroism and related techniques. An elliptically polarizing undulator (EPU), with variable polarization (linear and circular), in the energy range 200 to 2000 eV is the source of the soft X-rays.

The undulator radiation is monochromatized by an SX-700-type of PGM with vertical collimation and focused into an experimental chamber for magnetic circular dichroism measurements.

There are currently two different chambers available to users. One being a chamber equipped with an octupole magnet allowing for work under applied magnetic fields in an arbitrary direction in space with, presently, a field of 0.5 T. The chamber is also designed to conduct soft x-ray magnetic reflectivity measurements. The second experimental system consists of separate preparation and analysis chambers. The analysis chamber is equipped with an UHV electromagnet providing a peak field of 0.1 T in pulsed mode and 35 mT with a continuous field. The preparation chamber is equipped with a LEED and an ion sputter gun as well as a number of extra ports for user supplied auxiliary equipment

MAX III Beamlines

(see http://www.maxlab.lu.se for more details).

Beamline 13 is an undulator beamline for the low energy region (5 - 50 eV). It is equipped with a normal incidence monochromator with a very high energy resolution (resolving power greater than 105). The undulator is of the "apple-type" providing variable polarization. There are two branchlines, one with a fixed end-station for high resolution angle- and spin-resolved photoemission on solids, equipped with a rotatable Scienta R4000 analyzer and a Scienta 2D spin detector. Angle resolved photoemission can be made in ± 15 , ± 7 and ± 3.5 degree mode. An on-line MBE system provides the possibility for studying in situ grown samples. The second branch-line, with a differential pumping stage, is designed for an easy exchange of end-stations for atomic and molecular spectroscopy and luminescence measurements.

Beamline 14 is an undulator beamline used for angle resolved photoemission. It is equipped with a spherical grating monochromator covering the energy range 13 to 200 eV.

The end station analyzer chamber hosts two electron energy analyzers: An in vacuum rotatable VG ARUPS 10 analyzer and a fixed mounted PHOIBOS 100 mm CCD analyzer from SPECS.

The SPECS analyzer has an ultimate resolution of less than 3 meV. There are three angular dispersion modes namely MAD (medium angular dispersion), LAD (low angular dispersion) and WAM (wide angular dispersion). The MAD mode has angular acceptance of \pm 3 deg with angular resolution of less than 0.1 deg. The LAD mode has angular acceptance of \pm 6 deg with angular resolution of about 0.15 deg. The WAM mode has angular acceptance of \pm 10.5 deg with angular resolution of about 0.4 deg.

Beamline D7 is an infrared microspectroscopy beamline presently under construction. It will receive radiation from a bending magnet port. During the design and construction work for the beamline the microscope is used at the existing infrared beamline, 73, at MAX I.

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Beam port	Beam port Source type	Beamline/Monochromator	Energy or wavelength range	Experimental techniques
31	Undulator	PGM with Kirkpatrick-Baez objective and ellipsoidal focusing mirror.	15 – 150 eV	Scanning photoelectron microscopy.
41	Bending magnet 18 mrad	4.7m-TGM, 162°	15 – 200 eV	Angular resolved photoemission. On-line MBE system.
52	Bending magnet 20 mrad	1m NIM	5 – 30 eV	Presently used for education/ test station.
73	Bending magnet 60 * 100 mrad²	Two ellipsoidal mirrors, 1:1	12000 – 10 cm ⁻¹	Infrared microspectroscopy. Infrared spectroscopy using a high
N M	Tagged photon beam	Tagged photon beam Tagging spectrometer	15 – 185 MeV	Nuclear physics.

	Experimental techniques	High resolution XPS. X-ray absorption spectroscopy. Photoemission microscopy, PEEM.	High resolution XPS. X-ray absorption spectroscopy. Coincidence spectroscopy.	High resolution XPS. X-ray absorption spectroscopy. New end station aimed at studies under near-ambient	Pressure Conditions X-ray absorption spectroscopy. X-ray emission spectroscopy. Non-UHV compatible.	Time resolved X-ray diffraction.	Small molecule crystallography. Powder diffraction.	EXAFS, XANES. Surface, interface and thin film crystallography.
	Energy or wavelength range	43 – ~1500 eV	40 - ~1500 eV	50 - ~1 500 eV	50 - ~1500 eV	2 – 10 keV	0.8 – 1.6 Å	2.4 – 20 keV
	Beamline/Monochromator	PGM (modified SX-700, with spherical focusing mirror).	PGM (modified SX-700 with plane-elliptical focusing mirror).	PGM (modified SX-700, with spherical focusing mirror).	PGM (modified SX-700, with spherical focusing mirror).	Be-windows, Double-crystal monochromator.	Be-window, Bent Si(111) crystal.	Double-crystal monochromator. Exchangable Si(111) & Si(311) crystals.
	Operational status	Operative	Operative	Operative New end station for high pressure XPS under commissioning.	Operative	Operative	Operative	Operative
It MAX II	Beam port Source type	Undulator	Undulator	Undulator	Undulator	Bending magnet	Multi-pole wiggler	Superconducting multi-pole wiggler
Beamlines at MAX II	Beam port	1311	1111	1511/1	1511/3	D611	1171	1811

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Experimental techniques	Presently used for education/ test station.	Protein crystallography.	Protein crystallography. MAD technique.	Small Angle X-ray Scattering.	Protein crystallography and other scattering experiments.	MCD and related techniques for studies of magnetic materials.	High resolution XPS. X-ray absorption spectroscopy. Circular dichroism.
Energy or wavelength range	Quasi-fixed wavelength, 1.2 Å	Fixed wavelength, 1.04 Å	0.75 – 2.0 Å	Fixed wavelength, 0.91 Å	Fixed wavelength, 0.97 Å	200 – 2 000 eV	40 – ~1 500 eV
Beamline/Monochromator	Diamond crystal, multi-layer mirror.	Bent SI crystal, multi-layer mirror.	Collimating mirror Double-crystal monochromator. Focusing toroidal mirror.	Bent SI crystal, multi-layer mirror.	Diamond crystal, multi-layer mirror. Fixed wavelength, 0.97 Å	Collimated PGM.	PGM (modified SX-700 with plane-elliptical focusing mirror). Off-plane radiation for circular polarized radiation.
Operational status	Presently test station	Operative	Operative	SAXS station under commissioning 2010. In user operation April 2011	Operative	Operative	Operative
Source type	Superconducting multi-pole wiggler	Superconducting multi-pole wiggler	Superconducting multi-pole wiggler	Superconducting multi-pole wiggler	Superconducting mulfi-pole wiggler	Undulator with variable polarization	Bending magnet
Beam port	1911/1	1911/2	1911/3	1911/4	1911/5	11011	D1011

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Beam port	Beam port Source type	Operational status	Beamline/Monochromator	Energy or wavelength range Experimental techniques	Experimental techniques
<u> </u>	Undulator with variable polarization.	Operative	6.65 m Off-axis Eagle Type NIM.	5 – 50 eV	High resolution (meV) angleand spin-resolved photoemission on solids, atomic and molecular spectroscopy, and luminescence. On-line MBE system.
4	Undulator	Operative	5.5 – 5.8m-SGM, 162°	13 – 200 eV	Angle-resolved photoemission.
D7	Bending magnet	Under construction	Transfer optics, FTIR spectrometer and IR microscope.	12 000 – 10 cm ⁻¹	Infrared microspectroscopy.



Yiqiang Zhan, Fudan University, and Mats Fahlman, Linköping University, preparing their experiment on hybrid organic spintronic interfaces at the beamline I1011, 25 October 2010. Photo: Annika Nyberg

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Mats Leandersson is discussing ideas with Jan-Olof Forsell at the Poster Session during the MAX-lab Annual User Meeting at MAX-lab, 8 November 2010. Photo: Annika Nyberg

Electronic structure of Ge_{1-x-v}Mn_xEu_vTe epilayers

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GeTe-based diluted magnetic semiconductors attract considerable interest due to recently discovered features advantageous in view of spintronic applications. The highest Curie temperature of Ge_{1-x}Mn_xTe reported in the literature strongly depends on Mn contents or

carrier concentration and it can be as high as 190^o K.¹ This inspired extensive investigations of this system and related materials. Among others, Ge_{1-x-v}Mn_xEu_vTe was considered as a system with ferromagnetism promoted by introduction of Eu, presumably reducing number of antiferromagneticaly coupled pairs of Mn atoms.² The band structure of Ge_{1-x-v}Mn_xEu_vTe was studied by the angleintegrated photoemission only $Ge_{1-x-v}Mn_xEu_vTe$ polycrystals and the Ge_{1-x-v}Mn_xEu_vTe surface alloy (prepared by deposition of Mn on the surface of a Ge₁₋ _xEu_xTe layer and annealing of the system).³ For the polycrystals grown by a Bridgman method, some features corresponding to precipitates of other phases were detected. The formation of a Ge_{1-x-v}Mn_xEu_vTe surface alloy on Ge_{1-x}Eu_xTe was followed during the Mn deposition and spectral features corresponding to Mn and Eu were observed at subsequent stages of the system preparation. However, corresponding data from a crystal with manganese and europium ions built during the growth into the proper sites in the crystal lattice were still missing.

The reported experiments consisted in photoemission measurements of epitaxial layers of Ge_{0.88}Mn_{0.1}Eu_{0.02}Te, doped with both manganese and europium during the growth

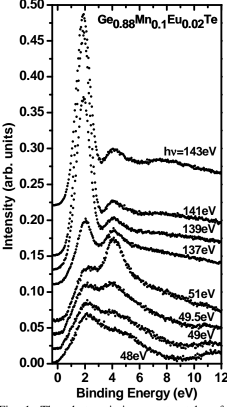


Fig. 1. The photoemission spectra taken for $Ge_{0.88}Mn_{0.1}Eu_{0.02}Te$ at the photon energies close to Mn 3p-3d (48-51 eV) and Eu 4d-4f (137-141 eV) resonances.

process. The samples were grown by an MBE technique on BaF_2 substrates. The crystalline quality was assessed by XRD measurements. The clean surface, suitable for the photoemission experiments was prepared by repeated cycles of Ar^+ ion sputtering and annealing at $250^{\circ}C$.

The photoemission experiments were carried out for two photon-energy ranges corresponding to the Fano-type⁴ resonances of two open-shell constituents of the system – Mn and Eu. The range of 31-55 eV covered the Mn 3p-3d resonance, the range of 131-150 eV –

Eu 4d-4f. The selected curves, taken for the photon energies close to the resonances are shown in Fig. 1. Since the Fano resonance leads to strong increase of the emission from the open shell (Mn 3d or Eu 4f) electronic states, the comparison of the spectrum taken at the resonance with that measured for the antiresonance allows us to reveal the spectral features related to these states. The difference curves for both resonances are shown in Fig. 2. They represent the Mn 3d and Eu 4f contributions to the photoemission from the valence band of Ge_{0.88}Mn_{0.1}Eu_{0.02}Te. They show that Eu is built into the system as Eu²⁺ ions, as expected for a IV-VI compound⁵. The weak feature of Eu³⁺ can be ascribed to Eu-related defects, in particular to those created on the surface during the cleaning process.

The Mn 3d-related feature (Fig. 2) corresponds well to that found for the $Ge_{1-x}Mn_xTe$ epilayers grown by the same technique as the investigated $Ge_{0.88}Mn_{0.1}Eu_{0.02}Te$ sample.⁶ The shape of this feature enables us to interpret it as the emission from the manganese ion bound to the ligands with the octahedral coordination. These observations indicate, within the sensitivity limits of the method, that

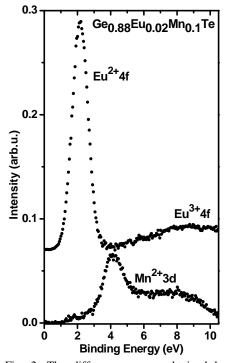


Fig. 2. The difference curves obtained by subtraction of the anti-resonance spectrum from the resonance one for Mn 3p-3d and Eu 4d-4f.

the presence of Eu ions in the layer does not change markedly the charge state of the manganese ions and their interaction with the surrounding crystal as compared with those in $Ge_{1-x}Mn_xTe$.

The research leading to these results has received funding from the European Community's Seventh Framework Programme (FP7/2007-2013) under grant agreement no 226716.

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Tin-stabilized (1×2) and (1×4) reconstructions on GaAs(100) and InAs(100) studied by scanning tunneling microscopy, photoelectron spectroscopy, and *ab initio* calculations

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Previous experiments have demonstrated beneficial effects of tin (Sn) stabilized (1×2)-reconstructed III-V(100) surfaces on the epitaxial growth of device materials. For example, deposition of the Sn prelayer in the epitaxial growth of GaAs eliminates effectively interfacial defects. In order to understand the reasons for the beneficial effects of the Sn-terminated III-V -surfaces and to controllably utilize these reconstructions, it

is essential to know the atomic structures of these III-V(100)(1×2)-Sn surfaces. To our best knowledge, no combined experimental and theoretical core-level photoemission and scanning tunneling microscopy (STM)

study, which would allow the identification of detailed surface structure, has been performed for the III- $V(100)(1\times2)$ -Sn surfaces previously.

We have addressed this issue by combining corelevel photoelectron spectroscopy (BL scanning tunneling microscopy, and ab initio calculations of the GaAs(100)-InAs(100)(1×2)-Sn surfaces.4 The Vienna ab initio simulation package (VASP) calculations demonstrate the stability of two different structures on GaAs(100)-Sn surface: Sn-As and Sn-Ga dimer models, where the corresponding atoms occupy the top surface sites. For InAs(100), only the Sn-In model is energetically favourable at 0 K. Furthermore, we have found a new (1×4) structure (Fig. 1) which, in turn, seems to violate the common supposition or guiding rule according to which the occupied dangling bonds tend to avoid each other. The (1×4) structure is formed rotating every second dimer row by 180° and it was found that the (1×4) becomes gradually more stable as the size of the group IV atom increases. The comparison of simulated and measured STM images also supports the presence

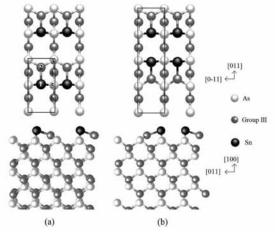


Figure 1. (a) Atomic model for the IIIAs(100)(1 \times 2)-Sn surface. (b) Atomic model for the IIIAs(100)(1 \times 4)-Sn

of these structures but the similarities in calculated images of energetically favourable models hinders the final identification of the Sn-induced surface structures by the means of STM measurements.

The Sn4d, Ga3d and As3d core-level spectra of the GaAs $(100)(1\times2)/(1\times4)$ -Sn reconstruction are shown in Fig. 2. The fitting of Sn4d emission shows only one spin-orbit split component which is an indication of one bonding environment, within the energy resolution limits, for Sn atoms at the considered surface. Some

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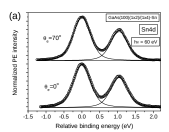
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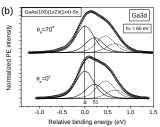
broadening of the Sn peak arises from the co-existence of the (1×2) and (1×4) -areas (Table1) Similar Sn4d spectra (not shown) were obtained for the InAs $(100)(1\times2)/(1\times4)$ -Sn surface. The both Ga3d and As3d include at least one shifted component separated by +0.22 eV and +0.18 eV, respectively, from the bulk emission. The comparison of measured and calculated core-level shifts, calculated for energetically most favourable atomic models from the first principles, is shown in Table 1. Calculated core-level shifts were obtained within the initial state model by calculating the average electrostatic potential at core and averaging from different group V and III atomic layers of the slab to obtain the bulk reference values. From Table 1 it can be seen that the calculated shifts for Ga-As (In-As) and Sn-As dimer models are too large compared to the experimental shifts. This is especially true for Sn-As model. However the existence of Sn-Ga (Sn-In) dimer model is nicely supported by the comparison of experimental and theoretical shifts.

Calculated (1×2)SnGa	+0.11, +0.06	+0.03, -0.08, +0.07	0
Calculated (1×2)GaAs	+0.41, -0.30	+0.12, +0.13	
Calculated (1×2)SnAs	-0.24, +0.10	+0.24, +0.55, -0.19	0.07
Calculated (1×4)SnGa	+0.11, -0.09, +0.20	-0.10, +0.14	-0,02
Calculated (1×4)GaAs	+0.43, -0.30, +0.32	-0.04, +0.17, +0.25	-0.03
Calculated (1×4)SnAs	-0.24, +0.20, -0.05	+0.27, +0.60, -0.22, +0.17	-0.02
Measured	+0.22	+0.18	0
Calculated (1×2)SnIn	+0.12, +0.07	0.08	0
Calculated (1×2)InAs	+0.38, -0.20	+0.16, +0.14	-0.05
Calculated (1×2)SnAs	-0.23, +0.11	+0.19, +0.40	0.1
Calculated (1×4)SnIn	+0.13. +0.18	0.14	-0.02
Calculated (1×4)InAs	+0.40, -0.21, +0.31	+0.13, +0.18, +0.26 -0	
Calculated (1×4)SnAs	-0.16, +0.14, +0.22	+0.54, +0.13, +0.19, +0.10 -0.2	
Measured	+0.17	+0.17	0

have studied the GaAs(100)-Ιn summary, we $InAs(100)(1\times2)/(1\times4)-Sn$ surfaces by LEED, photoelectron spectroscopy, STM/S, and ab initio calculations. Calculations demonstrate the stability of the Sn-Ga and Sn-As dimer models on the GaAs-Sn. For the InAs-Sn, only the Sn-In model is stable at 0 K. A new (1×4) reconstruction was found in which the filled dangling bonds are closer to each other than in the (1×2) . The comparison of measured and calculated STM images supports the presence of energetically favoured Sn-III models on the considered surfaces. Furthermore, the measured and calculated surface core-level shifts support the Sn-III model and allow the exclusion of III-As and Sn-As models.

We would like to thank the MAX-lab staff for all support.





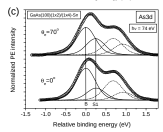


Figure 2. Fitted (a) Sn4d, (b) Ga3d and (c) As3d core-level spectra of the GaAs(100)(1×2)/(1×4)-Sn surface, measured with different emission angles (0° and 70°). Photon energy was 60 eV for Sn4d and Ga3d and 74 eV for As3d. All the spectra are normalized to their maxima.

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Photoelectric Properties of ion Implantation Induced Ge-Mn Layer Regrown at 400 and 600 °C on Ge(100)

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Introduction: Our work is the continuation of a previous one presented in this book series [1]. We use Mn as transition metal (TM) impurity in Ge(100) in order to obtain diluted magnetic semiconductor (DMS). In Ge the solubility is much better, than in III-V semiconductors. For DMS applications the relatively high concentrations (1-10 at%) of Mn are promising, substituting some Ge atoms by Mn. For the DMS fabrication we use Mn implantation into Ge. In this case the surface regrowth from the implantation induced radiation damage is the critical point, similarly to the doped semiconductor devices. On the other hand, the regrowing with the required high doping concentration must be careful, as we have to avoid intermetallic formation. The elevated temperature during the regrowth process may result in more ordered, or even single crystalline and reconstructed surface.

Experimental: The undoped Ge(100) wafers were implanted with Mn at 100 keV, 8x10¹⁶ ion/cm² at room temperature (at Institute of Ion Beam Physics and Material Research, Dresden-Rossendorf, Germany). The regrowth process at the surface was in situ investigated in UHV by photoemission and LEED techniques at BL41 in MAX-lab (Lund University, Sweden), with special attention to the damaged-undamaged interface. This case the temperature of the heat treatment (HT) was higher, namely 400 °C and 600 °C. After the regrowthing, the crystalline structure was investigated by Transmission Electron Microscopy-Electron Diffraction, while the magnetic properties were measured by SOUID.

Photoemission Results: In the spectrum of the as received samples the oxidized state of Ge 3d was seen together with some oxygen and carbon 2s contamination. After a short Ar⁺ bombardment (15 min, 0.5 keV), the Ge 3d and the valance band levels were clean, and strong reduction of the oxidized Ge 3d has appeared. After a longer sputtering (60 min, 0.5 keV) the Ge 3d emission was not changed further.

It is well known, that there is a resonance photoemission at 51 eV photon energy of the Mn 3d valence state. Now, similarly, after the HT at lower temperature (400 °C) the photoemission spectra show the resonance photoemission of Mn 3 d states [Fig. 1], but surprisingly, after a higher temperature (600 °C) annealing the Mn 3d states could not be identified [Fig. 2].

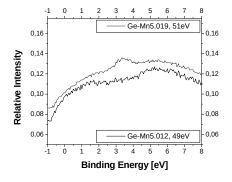


Figure 1. The valence band of Ge-Mn having polycrystalline structure. The resonance photoemission excitation of Mn 3d states were measured at 51 eV and 49 eV excitations.

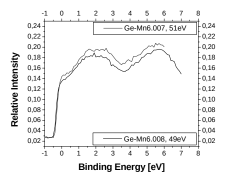


Figure 2. The valence band of Ge-Mn having single crystal structure. The resonance photoemission excitation of Mn 3d states were measured at 51 eV and 49 eV excitations.

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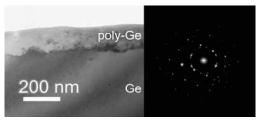


Figure 3. Poly-crystalline Ge sample, heat treated at 400 °C temperature.

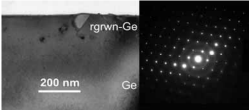


Figure 4. Fully crystallized Ge sample regrown at 600 °C as single crystal to the substrate.

According to the TEM crystal patterning, after the lower temperature heat treatment (at 400 °C) a polycrystalline film was formed (Fig. 3), with rather large polycrystallites. The effect of the interface regrowth could be neglected, which is rather unusual in the regrowth of ion implantation damaged surface layers. Moreover, there is some nucleation in the damaged layer. The relatively high defect concentration may be responsible for the blocked atomic regrowth and relaxation/reconstruction. At higher temperature (at 600 °C) the regrowth was found (Fig. 4.), possibly regulated by the damaged/undamaged interface. The growth mechanism of this layer can be similar to the one generally accepted for the epitaxial regrowth of the implantation damaged surface layers. The solid phase epitaxy resulted dominantly single crystal, though, the quality of this single crystal is far from being high.

The magnetic properties of the Mn implanted and regrown Ge are interesting, because these materials have a rather new atomic structure, therefore it can suggest new magnetic metal systems.

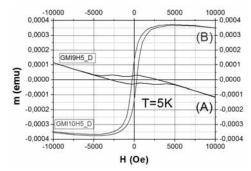


Figure 5.

- (A): Polycrystalline sample with small ferromagnetic clusters and with a blocking temperature of about 270 K
- (B): Single crystalline sample with ferromagnetic phase with a Curie point of about 160 K

(The negative magnetization at higher fields is the consequence of the diamagnetic contribution of Ge.)

According to TEM measurements, the structure of the Mn implanted Ge was amorphous after the implantation and the surface cleaning. The annealing of this structure produced some grains in the amorphous background. The magnetic properties could be characterized by super-paramagnetic clusters in the as-implanted case. After the annealing at 400 °C some ferromagnetic component and crystalline clusters were observed. The LEED pattern was not apparent, nor the ARUPS spectrum. Probably the annealing temperature of the thermal budget was too low, and the crystallization has only started. The pre-crystallization process was not localized to the damaged-undamaged interface, instead the whole damaged layer was involved. This indicates an unusual regrowing mechanism.

Acknowledgements: We would like to acknowledge the help from MAX-lab staff and wish to thank the financial support from ARI.

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Katarina Norén, Adriana Wawrzyniak, and Janusz Sadowski at the December dinner party at Grand Hotel in Lund, 1 December 2010.
Photo: Bengt-Erik Wingren



Mikael Eriksson is congratulating Bo Persson to his retirement after impressively 26 years at MAX-lab, 16 December 2010. Photo: Bengt-Erik Wingren

Studies of Carbon Microparticles by Raman and Infrared Microscopy in an Electrodynamic Balance

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A feasibility study concerning the use of Raman and Fourier Transform Infrared (FTIR) spectroscopy in studies of activated carbon particles levitated in an Electrodynamic Balance (EDB) is presented. Activated carbon particles treated with different substances $(O_3, HNO_3, NO_x, H_2SO_4)$ were studied in bulk samples, as single particles on a glass plate, and levitated using the EDB. FTIR of motionless particles on a glass plate showed a presence of different functional groups on the surface of the particles. Spectra taken on levitated particles were disturbed by oscillating motion of the particle. The obtained results and experimental challenges are discussed. The use of Raman technique did not reveal any difference between the samples, and it is therefore discarded as a possible technique for the studied particles.

The growing demand for energy production is a strong motivation behind research concerning more efficient and less emissive ways of fuel combustion. One strategy to achieve this goal is oxy-fuel combustion, a field of research that have seen intense activity during the last few years. In this type of combustion the oxidizing atmosphere consists of O_2 preliminary separated from air and mixed with recycled flue CO_2 . This means that at the end of the coal combustion process CO_2 will be the only exhaust gas. It enables efficient post-combustion capture and storage of CO_2 and also increases the combustion efficiency in comparison to combustion in air. Moreover, coal fuels, which are mostly planned to use in this type of processes, could be replaced by for example biochar particles. Neither gas phase nor surface chemistry occurring during combustion of coal and biomass are well understood, especially for the case of O_2+CO_2 atmosphere. Thus there is a particular interest in studies aimed to advance the knowledge of single coal or biochar particle combustion under oxy-fuel conditions. Observing the properties of solid phase particles under laboratory conditions presents two major challenges: particles need to be isolated from the influence of any supporting surfaces and the measurement techniques should not disturb the processes of interest. When considering single particle studies the former might be provided by using a levitation method while the latter can be solved by implementing light scattering or absorbing techniques.

In the present research the Electrodynamic Balance (EDB) method was used to suspend a carbon particle in air. In an EDB the levitation force originates in the electrical field, and this requires the particle to be charged. But it does not require for example non-absorbance or transparency like in optical levitators. This makes the EDB suitable for the study of carbon particles. Two non-intrusive spectroscopic methods which may provide extensive information about the presence of different functional groups in a studied sample are Raman and Fourier Transform Infrared (FTIR) microspectroscopy. In case of combustion studies they could be used to determine chemical composition of the particle surface before and after oxidation; this will result in a better understanding of the process in terms of its kinetics.

Experimental

The particles used were Norit CA3 activated carbon particles. This carbon was prepared by phosphoric acid activation and has a size distribution in the range of $7-100 \, \mu m$ with a median in 35 μm .

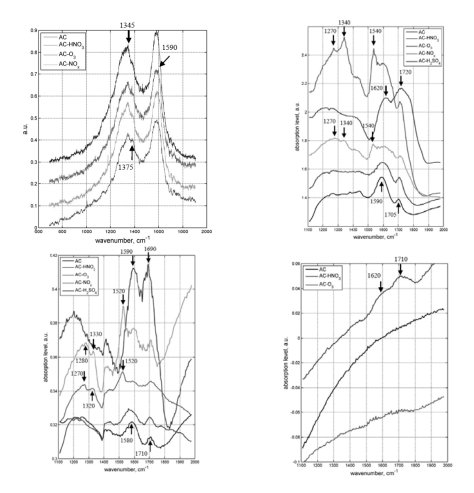
Prior to measurements 5 glass flasks with samples were prepared. First pair was exposed to a flow of ozone and nitrogen oxides respectively for several minutes and stored in flasks in air. Second pair was dissolved in concentrated nitric and sulphuric acids and treated for 24 hours. The samples exposed to liquids were filtered and washed with methanol, then dried overnight and stored in air. One sample was left untreated and used as reference. The untreated sample is hereafter denoted as AC, and treated samples as AC-O₃, AC-NO₅, AC-HNO₃ and AC-H₂SO₄ respectively.

FTIR measurements were performed using a Bruker IFS66V/S spectrometer coupled to a Bruker Hyperion 3000 microscope with 15x objective at Maxlab, Lund, Sweden. The microscope works either in optical or infrared regime. Raman spectra were collected at the University of Copenhagen, Denmark, by illuminating the particle by a 633 nm laser beam focused with a 100x Olympus microscope objective and collecting backscattered light.

For trapping the particle an EDB setup developed at the University of Copenhagen was used. The EDB chamber has classical configuration with three hyperbolic-shaped electrodes – two cap electrodes to which DC voltage is applied and one ring electrode which generates AC field. The two main features of the present setup are: (a) it has eight windows in horizontal plane and two in vertical that allows for optical access in all directions, and (b) the whole chamber can be isolated from ambient air, which makes it possible to change the oxidizing atmosphere to for example O_2+CO_2 .

In the present experiments the main source of periodic oscillations of the particle was the AC field in the chamber. The field needed to be strong enough in order to prevent any other motions of the particle due to laser irradiation,

displacement from the null-point etc. It is believed that increase of the particle charge-to-mass ratio could provide better stability which should result in better spectral information from its surface.



Upper left: Normalized Raman spectra of a variety of samples.

Upper right: FTIR spectra of bulk samples.

Lower left: FTIR microscopy spectra of single particles on a window.

Lower right: FTIR microscopy of single particles suspended in the electrodynamic balance.

Conclusions

The main conclusion of this feasibility study is that synchrotron FTIR microscopy is a promising technique in identifying different functional groups on treated activated carbon samples. Nitrogen or oxygen containing groups are distinguishable in the spectra of the treated particles. The particles can easily be injected to the EDB and once a particle is trapped it can be kept levitated for as long time as needed. Combining the electrodynamic levitation and FTIR microscope techniques seems to be a promising approach for studies of single particles. The main challenge ahead is to increase the particle stability in the electrical field. When particles can be kept levitated with small or no oscillation, it is expected that FTIR spectra with a quality similar to that for single particle on a glass plate will be obtained. Using the Raman technique the difference between samples could not be confirmed.

Studies of Small Water Complexes in Inert Matrices at Low Temperatures ≥ 2.8 K, by Infrared Vibrational Spectroscopy

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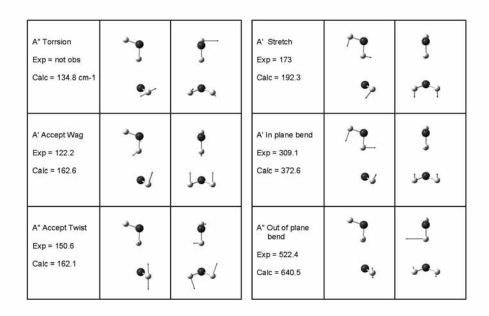
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Water dimers have been studied at low temperatures in different inert matrixes. Both noble gas matrices, Ne, Ar and Kr and different hydrogen matrices, e.g. parahydrogen and orthodeuterium have been utilized. The experiments are interpreted with the aid of density functional theory calculations, using the Gaussian package at a Linux cluster provided by Lunarc, the center for scientific and technical computing for research at Lund University. In particular the calculated harmonic isotopic shifts were used. A large set of different water isotopes have been studied to allow for identifications of different complexes as well as assignments different vibrational modes. All six intermolecular vibrational modes of the water dimer and the fully deuterated water dimer was assigned based the isotopic shifts induced. 31 of a total of 42 intermolecular fundamental modes of the seven different H, D, and ¹⁸O containing water dimers was experimentally observed and assigned accordingly. The over all agreement between the calculations and the experiments of all isotopologues allows for the first time for a complete and consistent description of these modes. Furthermore, a fine structure, which is more or less matrix independent and very similar for different intramolecular fundamentals of the same isotopologic dimer, is present on the high energy side of the main component. The fine structure and temperature dependency is interpreted as evidence for acceptor switching and rotation of the water dimer around its O-O axis in the matrices studied here. We also observe that for a given monomer concentration, the dimer concentration is significantly higher in solid Ne (or Ar) than in solid p-H₂. In p-H₂ the dimer concentration is only slightly higher than expected for a random distribution of water in the matrix. The dimer concentration in o-D₂ matrices is intermediate between p-H₂ and noble gas matrices. This strongly suggests that most

90

dimers form on the surface of the growing matrix, and not as the result of diffusion in the bulk of the matrix.



Figure, The six intermolecular vibrational modes in the water dimer, side and top views.

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FTIR spectra of propanol and isopropanol in gas phase

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The clustering phenomena and structural peculiarities of partially ordered liquids are of great interest in the scientific community. This interest is even growing in context of recent trends and developments in studies on modern multifunctional materials, heterogeneous systems and nanotechnologies. Monohydric alcohols, that usually build broad variety of H-bond aggregates, are quite simple and convenient models to investigate properties of molecular systems sized over the mezoscopic scale ($\sim 1-100$ nm). The importance of the problems connected with the alcohol clustering, structure and, in particular, with the mechanisms of the diffuse absorption band formation reflects in the great number of experimental [1, 2], theoretical [3 - 5] and combined works [6, 7] published in the recent years.

The most part of the investigations is naturally devoted to the simplest monohydric alcohol – methanol. In the present contribution we report the infrared absorption spectra of two isomers of propyl alcohol – n-propanol or propan-2-ol CH₃CH₂CH₂OH and isopropanol or propan-2-ol (CH₃)₂CHOH in gas phase. The spectra were recorded with the Bruker 120 FTIR HR spectrometer and the infrared synchrotron radiation component offered by the MAX-I storage ring. Spectra were registerd in the spectral region from 500 to 400 cm⁻¹ with resolution 1 cm⁻¹. In Fig. 1 the recorded spectrum of propanol is presented.

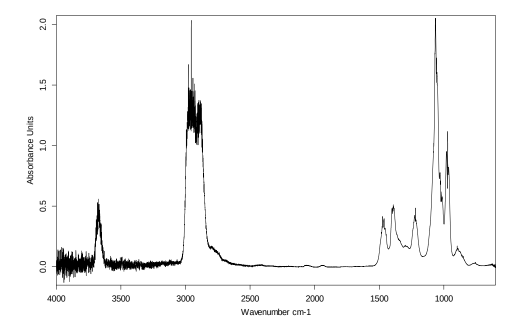
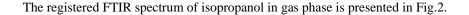


Fig. 1 The registered infrared absorption spectrum of propanol in gas phase



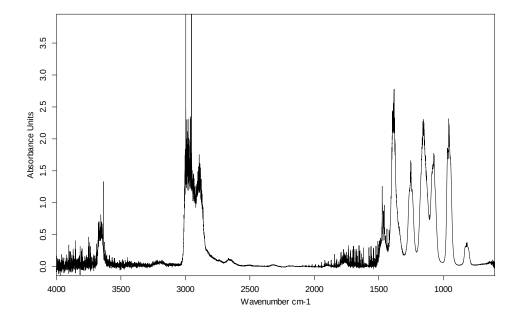


Fig. 2 The registered infrared absorption spectrum of isopropanol in gas phase

As it is seen from Figs.1 and 2, in the both isomers there is rather intense absorbtion at about 3660 cm⁻¹ corresponding to the vibrations of the free hydroxyl group. In the spectrum of isopropanol there is a broad band near 3200 cm⁻¹, which in absent in the spectrum of n-propanol. This band usually indicates the existence of the intermolecular hydrogen bonding. It means that in the case of isopropanol the individual molecules begin to aggregate into the larger association – so called clusters. This fact may be considered as the first step towards the formation of the cluster structure which is typical for the monohydric alcohols in the condenced state.

Thus, the direction of the farther investigations is obvious – the increasing of the pressure should increase the probability of the cluster formation in the studied objects and the process of the aggregation of the individual alcohol molecules into the hydrogen-bonded clusters during the phase transition from the gas phase to the liquid one can be traced.

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Continuous-wave laser annealing of Si-rich oxide: A microscopic picture of macroscopic Si--SiO₂ phase separation.

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We report on the first observation of the macroscopic (long-range) Si--SiO₂ phase separation in Si-rich oxide SiO_r (x<2)obtained by continuous-wave laser annealing of free-standing SiO_x films. The effect is analyzed by a unique combination of microscopic methods Raman, transmission, photoluminescence, and infrared spectroscopy, transmission electron microscopy, electron energy loss spectroscopy, and x-ray photoelectron spectroscopy. Three regions can be distinguished on a SiOx free-standing film after 488 nm laser annealing at intensities above ~104 W cm⁻²: central spot, ring around the central spot, and pristine film outside the irradiated area. In the pristine SiO_x material, small Si nanocrystals (Si-nc) (diameters of a few nanometer) are surrounded by SiO₂ with an addition of residual suboxides, the Si-nc being produced by annealing at 1100 °C in a furnace. The central spot of the laser-annealed area (up to ~30 µm wide in these experiments) is practically free of Si excess and mainly consists of amorphous SiO2. The ring around the central spot contains large spherical Si-nc (diameters up to ~100 nm) embedded in amorphous SiO₂ without the presence of suboxides. Laserinduced temperatures in the structurally modified regions presumably exceed the Si melting temperature. The macroscopic Si--SiO₂ phase separation is connected with extensive diffusion in temperature gradient leading to the Si concentration gradient. The present work demonstrates the advantages of high spatial resolution for analysis in materials research.

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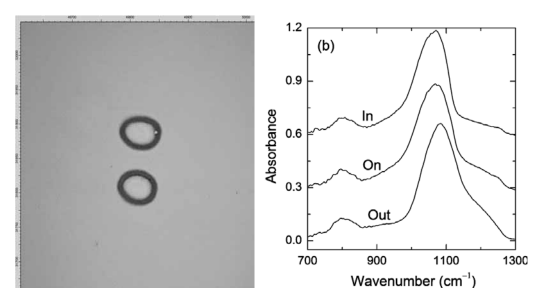
For the FTIR measurements, a free-standing sample was placed on a 2-mm-thick KBr window in the focal point of a Bruker Hyperion 3000 microscope with a condenser and objective both with 15x magnification giving a total visible magnification of 215x. In the microscope, a fixed square 5 x 5 μm aperture was used for the measurements. The microscope was coupled to a Bruker IFS 66 v/s FTIR instrument which used synchrotron light from the MAX-I ring , beamline 73, at MAX-lab in Lund. 2048 scans were coadded with a resolution of 4 cm $^{-1}$. These studies were performed with a 0.5 μm SiO $_x$ free-standing film exposed to $\sim\!110$ mW of power through a lens with a 150 mm focal length. The investigated laser-annealed areas are about 40 μm wide so that the spatial resolution of 5 μm is sufficient to study selectively absorption at different key positions.

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The optical transmission picture of two laser spots and spectra measured at different positions. Measured in an Hyperion 3000 microscope with an aperture of $5x5~\mu m$ and a spectral resolution of $4~cm^{-1}$ at beamline 73 MAX-lab by Anders Engdahl.

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Microspectroscopic analysis of potential bone proteins from the Cretaceous of Angola

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During the summer of 2009 limb elements and vertebrae from mosasaurs¹ and plesiosaurs² were collected for molecular and biochemical analyses at Bentiaba³ (Cretaceous: Maastrichtian), south-western Angola, using a protocol⁴ to optimize the chances of recovery of potential primary biomolecules while minimizing the risk of introducing contaminants. Under sterile conditions small bone tissue samples were decalcified using ethylenediaminetetraacetic acid (EDTA). In some cases the acid-resistant residues contained fragmented, vessel-like structures associated with a fibrous substance that, in modern bone, would represent the organic phase of the extracellular matrix (the osteoid). The fibrous organization of the partially organic matter was demonstrated by optical and scanning electron microscopy. Additionally, under transmission electron microscopy regions of cross-striated, fibril-like structures were observed.

To test the possibility of endogenous biomolecular preservation, isolated fibre bundles recovered from a plesiosaur propodial were examined using infrared microspectroscopy at beamline 73, MAX-lab, Lund University. The majority of the absorbance spectra obtained showed peaks corresponding to carboxylic acid and carboxylate groups (Fig. 1A), to suggest that the bulk of the organic matter was oxidized. However, a few fibre bundles also exhibited peaks indicating the possible presence of amino acid containing molecules (Fig. 1B). Putative amide bands were found at the frequencies of 1651–1647 (Amide I), 1558 (Amide II), and 1255 (Amide III) cm⁻¹, respectively. In modern proteins these bands are associated with stretching and bending vibrations of the peptide (CO–NH) bonds, and their positions and heights are sensitive to structural changes of the three-dimensional conformation of the molecule. Vibrational bands corresponding to inorganic materials were also indentified, and ascribed to, among other things, A-B carbonate of hydroxyapatite. These results are somewhat ambiguous and additional analyses are required in order to establish whether or not any endogenously derived organics remain in the fossil bones from the Cretaceous of Bentiaba.

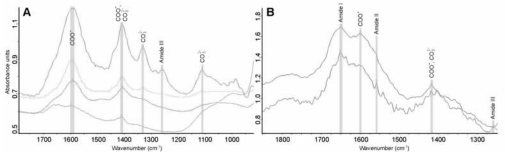


Figure. 1. A, Infrared spectra of isolated fibrous tissues from a plesiosaur propodial. Note presence of bands attributed to COO and carbonate groups. B, Infrared spectra of isolated fibrous tissues from a plesiosaur propodial. Note presumptive amide bands.

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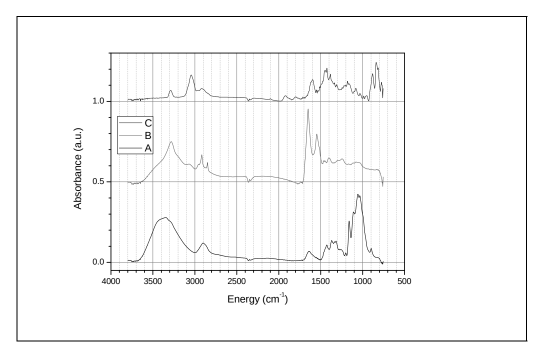
Johan Lindgren, department of Earth and Ecosystem Sciences, Lund University, is holding a piece of fossilized bone from a Mosasaur.
Photo: Annika Nyberg

Investigation of the spectroscopic properties of combustion generated particles in the mid infrared region using synchrotron radiation at MAX-lab, beamline 73.

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The project is a study of carbon-based particles formed under different conditions of combustion and their interaction with the environment, radiation and biological systems. Structural and chemical information is obtained from the identification and composition of spectral signatures, which are related to different types of bonds, e.g., C-C, C-H, and C-O. In combustion it is mostly accepted that soot precursors are dominated by PAH, but more recent results also suggest that particle formation may be induced by more polymeric heterogeneous structures including elements of aliphatic and aromatic units and experiments have shown interesting results [1,3]. Thus the C/H ratio can be a significant indicator of the history of the soot particles. The oxidation of soot particles in the atmosphere is also important to understand. The level and the types of bonds oxygen makes to the carbon can tell us about the aging history of the soot particles. The first implications on the level of toxicity can also be obtained from a chemical analysis based on the IR-spectral properties.



Experiments have been performed on diffusion-like propane Bunsen and ethylene flames. The species produced were metal plates inserted at different heights of the flames. The plates were exposed to the flame gases for different periods of time ranging from 30 seconds to 2 minutes. Samples were then extracted and cooled before being inserted into the Bruker IFS 66 spectrometer and then analyzed. Data from these measurements is currently being analyzed and will be presented during 2011. Examples of spectra obtained for different soot-related particles are shown in the figure below. One can observe the different aromatic/aliphatic content, polarity and types of oxygen bonds present. A variety of spectra are thus obtained that reflects the specific chemical and physical path that the group of molecules and particles has been subjected to in the flame. We have much more data to evaluate. More targeted experiments directed towards the understanding of the correlation between local combustion conditions and spectral properties will be performed during 2011.

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Collagen contents changes in vascular endothelial cells deprived of copper as revealed by SR-FTIR imaging.

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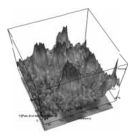
FTIR Beamline n°73 – 3 weeks of measurements using synchrotron radiation source

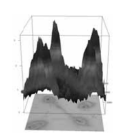
Summary: This study has been conducted to determine the effect of copper chelation on vascular endothelial cells (HCMEC) undergoing tubular formation as found during tumor angiogenesis. This study was based on FTIR imaging and chemometrics to analyze collagens contents in the extracellular matrix (ECM) produced by HCMEC cells in culture on Si₃N₄ substrate, which is compatible with X-Ray fluorescence microscopy (XRF) for elemental analysis (notably Cu⁺⁺). Cells were cryofixed prior to all measurements for ensuring comparison of results obtained by both FTIR imaging and X-Ray fluorescence microscopy (experiments to come in 2011). Three cell culture conditions were used for determining the effect of Cu⁺⁺ deprivation on ECM collagens at cell-cell junctions: cells were cultured in copper free culture medium (Cu0) or 50 and 100 µM CuCl₂ culture medium (Cu50 and Cu100, respectively). FTIR images of cells were used for spectral data treatments, notably for determining changes in ECM contents at cell-cell junctions. Although the IR signal on so weak organic material appeared well below the linearity scale of the detector, qualitative analysis of proteins secondary structure parameters could be performed. The main results obtained in this study are presented in the table below:

Conditions	Parameters	Cu0	Cu50	Cu100
24 hours	Junctions (n/mm ²)	12 ± 4	15 ± 3 (+ 25%)*	16 ± 5 (+33%)*
	1637/1656 cm ⁻¹	0.3 ± 0.1	$0.8 \pm 0.2 \ (+266\%)^*$	$0.8 \pm 0.3 \ (+266\%)^*$
36 hours	Junctions (n/mm ²)	19 ± 7	23 ± 5 (+21%)*‡	25 ± 3 (+32%)*;
	1637/1656 cm ⁻¹	0.5 ± 0.2 ‡	$1.1 \pm 0.4 (+220\%)$ *;	$1.3 \pm 0.4 \ (+260\%)$ *†‡
48 hours	Junctions (n/mm²)	39 ± 13‡#	45 ± 10 (+15%)*; [#]	46 ± 8 (+18%)*; [#]
	1637/1656 cm ⁻¹	0.5 ± 0.2 ‡	1.1 ± 0.5 (+220%)*;	1.5 ± 0.6 (+300%)*;†*

Table: Number of cell-cell junctions and collagen-to-proteins ratio for HCMEC cultured with gradual concentration of $CuCl_2$. All data are average of 24 different cell cultures (1 cell culture = 1 Si_3N_4 window with a 3*3 mm frame for cell counting and spectral image acquisitions; n = 3 FTIR images per window). Collagen-to-proteins ratio was calculated by integration of the 1637 (triple helix absorption from collagens) and the 1656 (α-helix absorption of proteins) cm⁻¹ bands after spectral curve fitting of the 1750-1600 cm⁻¹ spectral interval. * = significantly different from Cu0 condition; † = significantly difference between Cu50 and Cu100 conditions; ‡ significantly different from 24h-culture condition; # = significant difference between 36h- and 48h-culture conditions.

The first result obtained is that the number of cell-cell junctions were found reduced by 15 to 30% only at equivalent cell culture durations (at 24, 36, and 48 hours – corresponding to early, mid, and advanced phases of cell development on support) when using copper-free culture medium. There was no major difference between the Cu50 and Cu100 conditions. Therefore, even if copper appears as a significant factor for allowing HCMEC tubular formation, the absence of this element is not aborting the process. On the other hand, it appeared that ECM collagen contents in HCMEC deprivated from Cu⁺⁺ was diminished by 50 to 75 % on average for all culture durations, with a concentration effect observed between the Cu50 and Cu100 conditions. Therefore, our results show clearly that there is a direct correlation between intracellular copper concentration and collagen production in ECM of HCMEC undergoing tubular formation.





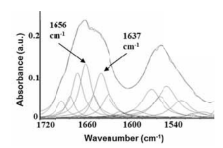
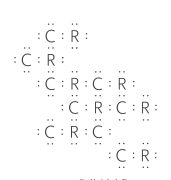


Figure: FTIR imaging of human cerebral microvascular endothelial cells. HCMEC images using a FPA detector truncated at 32*32 pixels, a 15X magnification (3*3 µm resolution), 256 scans at 8 cm⁻¹ spectral resolution. Right, spectral curve fitting of spectrum corresponding to the cell-cell junction location as found in the image in the center.

These results raise fundamental questions about initial steps of angiogenesis, i.e., the tubular formation of migrating vascular endothelial cells under angiogenic stress. Copper deprivation is not sufficient to abort completely the cell-cell junction and the production of ECM, but just delay the process. The correlation found between intracellular copper concentration and extracellular matrix production is thus likely to re-define the anti-angiogenic strategy for brain tumors since copper chelating agents cannot abort alone the process of cell-cell junction and anti-MMP agents alone cannot abort the process of ECM production.

These results will be presented for publication in the journal Analytical Chemistry once X-Ray fluorescence microscopy measurements will have confirm the absence of Cu⁺⁺ at cell-cell junctions in copper-free cultures.

User report 2011-02-22



Colloidal Resource

Colloidal Resource User report at MAX lab 2010

Colloidal resource AB is a consultant company in the field of chemistry, especially physical chemistry and surface chemistry. In our work we assist industry and start up companies with their chemistry related questions and problems. We use the facilities at Lund University including MAX lab. During 2010 we used the IR facility together with Anders Engdahl (experimental station nr 73) and the X-ray facility (I911-2) with Dörthe Haase.

IR is used both in transmission mode and with the ATR crystal. IR is used in projects to determine absorption on filters and the permeability of membranes. During 2010 medically associated products were studied.

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High Resolution Far-Infrared Absorption Spectroscopy of Halogenated Hydrocarbons: Rovibrational Analysis for the 1/4 Band of CH⁷⁹BrF₂

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Halogenated hydrocarbons play important roles in stratospheric ozone depletion and global warming. Since rotational and rovibrational spectroscopy are among the most widely applicable and accurate methods for remote sensing, highly accurate spectroscopic data of halocarbons in the atmospheric windows are crucial for atmospheric modeling studies. Reliable spectroscopic data for many halocarbons are still missing in the literature in particular for those involving bromine. One important species is CHBrF₂ (halon 1201) which has been proposed as an interim replacement of fully halogenated halons due to its short atmospheric lifetime and good fire-suppression properties.

The infrared spectrum of CHBrF₂ exhibits an extremely dense rotational structure due to both the very small rotational constants and absorption originating from cold as well as hot bands of the two abundant bromine isotopes in the natural sample (50.7/49.3% of $^{79/81}$ Br). The present contribution reports the first infrared absorption spectrum of an isotopically enriched CH₂D⁷⁹Br sample (purity >95%). This spectrum has been obtained with a spectral resolution of 0.0025 cm⁻¹ employing the Bruker IFS 120 HR Fourier transform infrared spectrometer with the highly brilliant MAX-I storage ring as the external radiation source. The upper far-infrared spectral region contains the fundamental band ν_4 at 718.77 cm⁻¹ corresponding to the HCBr deformation mode.

The observed spectrum reveals that the ν_4 band is composed mainly by a-type transitions. The spectrum shows a scarcely structured Q-branch degrading towards lower wavenumbers with a noticeable line density in the P- and R-branches. In these dense P- and R-branches the spectrum shows distinct band heads separated by about 2B approximately equal to 0.20 cm^{-1} . These absorptions consist of series of lines differing by one unit in J and each involving the almost degenerate levels with K_a^+ and $(K_a + 1)^-$, where the + and - signs correspond to even $(K_a + K_c = J)$ and odd $(K_a + K_c = J + 1)$ components, respectively. A typical rovibrational structure is depicted in Fig. 1, where a portion of the P-branch near 705.5 cm⁻¹ shows the resolved lines in the qP_K (J = 74 - 76) band heads.

The majority of assigned transitions is a-type and the maximum J and K_a values are 98 and 22, respectively. The assigned rovibrational transitions have been fitted to the A-reduced Watson-type Hamiltonian in the I' representation, keeping the ground-state constants fixed at the values determined from the complementary microwave study of the present work [1]. The obtained rotational and centrifugal distortion constants, along with those derived for the ground state, are listed in Table 1.

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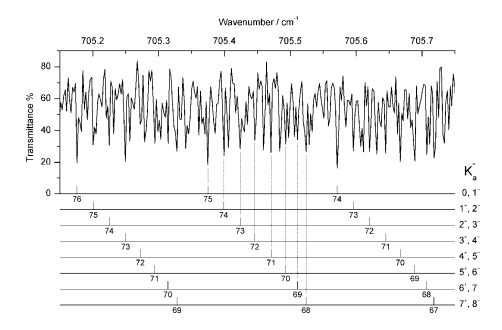


Fig. 1: Rotational *P*-branch assignment for the v_4 band of CH⁷⁹BrF₂ near 705.5 cm⁻¹ showing the resolved transitions in the ${}^{q}P_{K}$ (J=74-76) band heads. The lines from degenerate levels with different values of K_a^+ and $(K_a+1)^-$ are labelled; the + and – signs of K_a refer to $(K_a+K_c=J)$ and $(K_a+K_c=J+1)$, respectively.

Table 1: Spectroscopic constants (cm⁻¹) of the ground and $v_4 = 1$ states of CH¹⁹BrF₂^a.

			Ground state	$v_4 = 1$				
ν_0				718.771191(120)				
A			0.34022500986(117)	0.34012333(173)				
В			0.09684740828(103)	0.096430882(163)				
C			0.07872615218(100)	0.078451420(92)				
Δ_{J}	×	10^{7}	0.2225513(43)	0.220306(160)				
Δ_{JK}	×	10^{7}	0.964875(29)	0.9928(31)				
Δ_K	×	10^{6}	0.2455442(40)	0.2403(49)				
δ_I	×	10^{8}	0.445420(20)	0.43698(98)				
δ_{K}	×	10^{6}	0.1044172(87)	0.10483(37)				
Φ_{I}	×	10^{14}	0.4400(60)	0.4400 ^b				
Φ_{IK}	×	10^{12}	0.1538(33)	0.1538 b				
Φ_{KI}	×	10^{12}	0.1017(120)	0.1017 ^b				
Φ_{K}	×	10^{12}	0.1938(93)	0.1938 ^b				
ϕ_J	×	10^{14}	0.2091(30)	0.2091 ^b				
ϕ_{JK}	×	10^{13}	0.8446(160)	0.8446 ^b				
ϕ_K	×	10^{11}	0.2879(67)	0.2879 ^b				
No.	of	Data	658	4225				
σ	×	10^{3}	0.00045	0.749				

^aThe quoted errors in parenthesis are one standard deviation in units of the last significant digit. ^bConstrained to ground-state values.

Infrared absorption spectra of conformational mixture of 1-butene SOZ in gas and in non-equilibrated nitrogen matrices

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Secondary ozonides (*SOZ*) are intermediate products of ozonization reaction of alkenes, which constantly occurs in the troposphere. The final fate of the reaction depends on several factors: conditions of the reaction and particularly on the alkene structure. The ozonization reaction proceeds in a few steps and various unstable intermediate products of the reaction is formed. Some of the products are not characterized yet. This is particularly true for the primary and secondary ozonides, because of the short life times of these molecular species [1-2]. The intermediate products are chemically highly reactive and actively participate in photochemical smog formation.

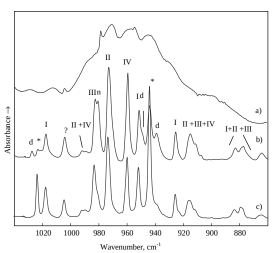


Fig. 1. FTIR absorption spectra of 1-butene secondary ozonide in gaseous state (a) and in N_2 solid matrix prepared using hot nozzle technique: (b) – 800 K and (c) – room temperature 300 K. Notation of the bands belonging to the different conformations is the same as in table 1. * - denotes ethene secondary ozonide spectral bands, d - denotes the bands of the decomposition products. The matrix spectra is corrected for the matrix effects by red shifting of the spectra by 18 cm $^{-1}$.

The intermediate products can be stabilized carrying out ozonization reaction in laboratory conditions at low temperature (T = 60 -75 K). From the previous works at Max-lab, we have made a conclusion that gaseous 1-butene ozonide sample consists from 4 conformers, which slightly differ in geometry. Present experiments are carried out in order to find some additional arguments for assigning the spectral vibrational bands to the different conformers of 1-butene SOZ. For this purpose hot nozzle technique was used. This technique allows to

form matrices with conformational equilibrium corresponding different temperatures in 300-800 K range. The matrix spectra were combined with corresponding spectra of gaseous 1-butene SOZ.

In 1040 – 860 cm⁻¹ spectral region of gaseous 1-butene SOZ spectral bands at 986, 977 and 965 cm⁻¹ are clearly visible and some spectral bands are partly hidden in the slopes of these three bands (see fig. 1). Comparison of the matrix isolation and gaseous spectra allows to assign the

bands to four different conformers. The assignment is in a good agreement with the results of the theoretical calculations (see Table 1).

Table 1. DFT calculated B3LYP 6-311++G(3df, 3pd) positions (scaling factor – 0.994) and intensities of the strongest infrared spectral bands in the spectral region of the five membered ring vibrations (860 – 1040 cm⁻¹) for four most stable staggered structures of 1-butene secondary ozonide.

•	equatorial gauche (≈ -60°) I			equatorial anti II		equatorial gauche (≈ 60°) III		axial IV	Appr. vibra-
v _s cr	n ⁻¹	I, m ³ *mol ⁻¹	$ \nu_{\rm sc}^{}, $ $ {\rm cm}^{-1} $	I, m ³ *mol ⁻¹	$ \nu_{\rm sc}^*, \\ {\rm cm}^{-1} $	I, m ³ *mol ⁻¹	$ \nu_{\rm sc}^*, $ cm ⁻¹	I, m ³ *mol ⁻¹	tional mode
9:	31	13	918	5	915	16	917	11	νCO asym- metric
9:	58	24	980	49	948	28	997	55	νCO _p sym- metric
10)22	24	997	12	990	59	998	7	ν CC _t

The calculated structures of the four conformers are presented in fig. 2. From the theoretical calculations and from matrix experiments it can be concluded that 965 cm⁻¹ spectral band consists of two conformers spectral bands – the most stable conformer *equatorial II* and *axial IV*. Spectral band 977 cm⁻¹ belongs to second stable conformer – *equatorial III* and spectral band 986 cm⁻¹ – to third conformer *equatorial III*.

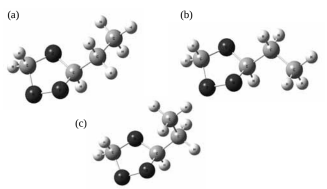


Fig. 2. 1-butene secondary ozonide structures corresponding to three most stable staggered conformers found by DFT calculations performed at B3LYP 6-311++G(3df, 3pd) theory level. (a) equatorial I, τ_{OCCC} = -66.1°; (b) equatorial II, τ_{OCCC} = -177.5°; (c) equatorial III, τ_{OCCC} = 56.2°.

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Diffusion in mechanically stressed bone using infrared microspectroscopy

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The interaction between mechanical load and bone growth in mammals is investigated. It is well known repeated mechanical load increases the mass and changes the morphology of the bone. Various experiments show that mechanical load directly or indirectly promote bone growth. Detailed clinical studies have been made by, e.g., Lanyon and Rubin¹, and Isaksson $et\ al.^2$. Models of the phenomena based on an assumed direct influence of load sometimes lead to confusing results and unexpected load rate dependences. The conclusion has been that the interplay between mechanical load and transport of nutrients and signal substances play an important role which is not covered in the existing models. In the proposed project an indirect mechanical influence is suggested (cf. Banks-Sills $et\ al.^3$). The model is based on the hypothesis that the primary condition leading to bone growth is a change of the chemical environment caused by stress driven diffusion.

The most important substances that are believed to be transported through stress driven diffusion are the prostaglandins, e.g. prostaglandin E2 (PGE2), which according to results from in vivo studies stimulate osteoblast activity. An alternative could be nitric oxide (NOS) is a strong inhibitor of bone resorption through processes which decrease the recruitment of osteoclasts.

The role of stress gradients as driving forces of diffusion or other mechanisms of mass transport is studied in the present project. The long term goal is to improve medical treatment of skeletal bone fractures and disorders like e.g. osteoporosis. Nutrients and signal substances are known to move through the bone, via a system of canals, *i.e.*, via Haversian canals and canaliculi to segments of compact bone. The motion is supposed to be a combined convective flow and diffusion, where both are believed to be affected by mechanical straining. Thus, experimentally determined diffusivity and its stress sensitivity is required. In a previous study unaffected diffusion was studied using infrared microspectroscopy operating in ATR mode. As then, heavy water was selected for its visibility in the infrared spectrum. The samples were around 2.5 to 3 mm thick cross-sections of the ulna long-bone of a sheep. Three different experiments were performed. A typical series of spectra is shown in Fig. 1. First, the evaporation rates from the sample surfaces were studied for water and heavy water. The water and heavy water content at the surface was followed for fully saturated samples, that were placed

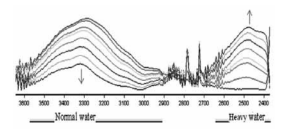


Figure 1: In a typical spectral profile is observed from measuring at some distance from where heavy water is injected into the bone. The concentration of normal water is observed as the concentration of heavy water is observed to increase with time.

in the observation chamber. Three different points This monitored. background to the measurements following thereafter. Second. heavy water injected along a part of the bone surface and the diffusive transport of heavy water in the wet bone was monitored. Third, mechanical load was applied using a micrometer. The result was compared with a numerical result for diffusion in bone computed using a finite volume

method (see Fig. 2). The process is assumed to be plane. The governing equation for stress enhanced diffusion is as follows:

$$\frac{\partial C}{\partial t} + \nabla (D\nabla C) - \nabla (BC\nabla S) = 0,$$

where C is the concentration of heavy water, D the diffusivity in wet bone, B the stress sensitivity and S the hydrostatic stress. proximate solution for a more realistic geometry computed using a finite element method.

A fair resemblance was obtained between experimental and computed results was obtained. The influence of stress could not be compiled because of unsatisfactory measurements. Additional experiments are needed. However, interesting enough, total suppression of the flux was observed in at least one case which seems promising.

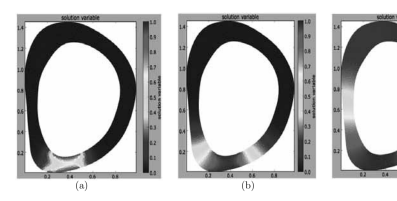


Figure 2 a-c: Computed distribution of heavy water during almost 3 hours. The heavy water is injected at the lower edge as indicated in fig (a). The heavy water is observed to spread symmetrically along the sides of the cross section. In (c) one may note that significant amounts of heavy water is spread through the entire cross section while most of the heavy water still remains in the lower half of the cross section.

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High quality graphene layers on commercial available cubic-SiC(001)/Si wafers: a perspective for mass production of graphene-based electronic

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Astonishing electronic properties of graphene make it the most promising candidate for replacing silicon in future electronic devices. A best method of graphene preparation so far, i.e. annealing of alpha-SiC bulk substrate in an inert atmosphere of argon at 900 mbar, has a dramatic improving influence on domain size and film homogeneity [1, 2]. However such graphene production on the surface of bulk alpha-SiC, although resulting in high quality graphene layers, does not meet the requirements of industrial mass-production because of the limited size and the costly nature of alpha-SiC wafers sliced from the single crystal ingots. Indeed, so far there are commercially available only alpha-SiC wafers with small size (about 2 inches in diameter), which are very expensive (about 2000 US dollars per wafer).

If instead of using a bulk SiC crystal one can fabricate a graphene film onto a thin SiC film grown on a largediameter Si wafer (SiC virtual substrate), its industrial impact would be enormous. SiC has more than 200 polytypes. However, the only polytype that grows on Si wafer is the cubic-SiC. It is their strong belief that one must utilize a proper orientation of the cubic-SiC surface in order to accommodate with the 6-fold symmetry of the graphene crystal. In this context, the 3C-SiC(111) surface is definitely the best choice, however there are no commercial available 3C-SiC(111) films on Si substrate so far.

We have demonstrated already the feasibility of graphene synthesis on the surface of cubic-SiC(001) thin film (about 1 µm) deposited on standard Si wafer in UHV conditions [3]. The later serves so far as the basis (substrates) for integrated circuits in microelectronics. The cubic-SiC(001) thin films (about 1 µm) deposited on standard Si wafer are commercially available up to 300 mm in diameter, not expensive, are wide gap (2.3 eV) single crystal semiconductor. Therefore, such cubic-SiC(001)/Si wafer is the perfect substrate for graphene layers and graphene/cubic-SiC(001)/Si wafer could be easily adapted for graphene-based electronic technologies and thus could be directly patterned by standard Si-electronic lithographic processes. However it is well known that decomposition of SiC during annealing in UHV conditions yields graphene layers with small grains, typically 30-100 nm in diameter [1, 2]. Using LEEM we have observed similar result for graphene grown in vacuum on SiC(001) substrate. The small domain size and inhomogeneous distribution of film thickness self-evidently result from the fact that the annealing process in UHV conditions is taking place far from equilibrium. A few recent publications shows that the flak size of graphene grown in UHV on a SiC surface could not be bigger than one-two hundred nanometers [4] because the strong sublimation rate of Si in vacuum conditions prevents the carbon atoms from proper rearrangement, leading to the heterogeneity and formation of small graphene domains. Taking into account these conclusions, we have covered the outer surface by a thin film, which strongly prevent the Si sublimation in UHV annealing and have managed to substantially improve the graphene layers grown in vacuum at high temperature. The high quality of the graphene was proved by number of techniques like LEEM, PEEM, AR-PES, NEXAFS and Raman spectroscopy. Some of the results are presented in Figs 1-4.

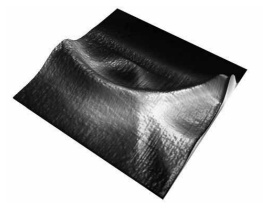


Fig. 1. Electronic states in graphene on cubic-SiC(001) surface: overview ARPES intensity map taken along the ΓK direction.

Fig. 2. Electronic states in graphene on cubic - SiC(001) surface: ARPES intensity map taken at the K point along the black dotted line.

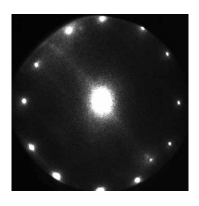


Fig. 3. μ -LEED pattern from graphene on cubic-SiC(001)surface; sampling area is 1 $\mu m,$ electron energy is 54eV.

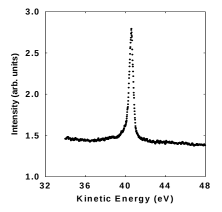


Fig. 4. C1s photoelectron spectrum (photon energy is $h\nu=330eV$) collected from a 10 μm graphene area on cubic-SiC(001) surface.

Finally we believe that our results represent a realistic way of bridging the gap between the outstanding graphene properties and their technological applications.

Acknowledgements: This work was supported by the DFG under grants KN393/9 and KN393/14, by the Finally we believe that our results represent a realistic way of bridging the gap between the outstanding graphene properties and their technological applications.

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Unconventional zwitterionic state of cysteine

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Introduction: Amino acids form a particularly interesting class of adsorbates from a surface science point of view. Typically, they are sufficiently small to be sublimated by standard thermal methods in ultrahigh vacuum (UHV), but at the same time they offer a large chemical variability. In addition to the amine and carboxylic acid groups other chemically relevant sidegroups (OH, SH, aromatic rings) subtly influence the properties of the molecules. This variability leads to a large number of possible bonding geometries. And indeed, in structural biochemistry the 20 universal amino acids form the building blocks of an endless variety of proteins, as their funtional groups allow them to link to one another aided by the easy formation of hydrogen bonds. The different moieties can act either as acceptors or donors, depending on the chemical environment.

L-cysteine (HS-CH₂-CH(NH₂)-COOH, Fig. 1, top) offers a high degree of chemical complexity due to the presence of three different functional groups. Further complexity is added by the fact that L-cysteine may assume both the neutral and zwitterionic forms. In the gas phase the non-ionic form is the most stable form of L-cysteine^{1,2}, while both in aqueous solution and in the solid state it is found in the zwitterionic state with the carboxylic group deprotonated^{3,4} (Fig. 1, left). Two polymorphs have been found for the crystal at adiabatic pressure⁴, and phase transitions at 70K (orthorombic)⁵ and 150K (monoclinic)⁶ have been observed. transitions are subtle and smeared out over a larger temperature range. The use of different experimental methods has shown that a series of small changes in dihedral angles involving, amongst others, the -CH₂SH sidechain can affect the hydrogen bonding network and hence the crystal structure^{6,7}. Similarly, they report a dependence on the thermal history of the crystals and on the heating/cooling velocity during experiments.

Recently, Tian and coworkers presented hydrogendeuterium exchange and gas phase acidity measurements alongside calculations which indicate that the thiol group should actually be more acidic than the carboxylic group by 3.1 kcal/mol. Deprotonation of cysteine would therefore most likely lead to a thiolate ion, even though they predicted that a zwitter ion involving the thiol group (see Fig. 1, right) should be more unfavourable by 10.1 kcal/mol⁸. Multilayers grown at RT and around 200 K show only the normal zwitter ion, and a ratio of zwitter ion/neutral molecule that increases with thickness, being close to 9:1 for the thickest multilayers⁹.

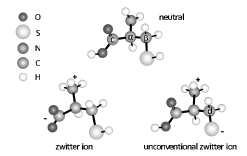


FIG. 1: Neutral and zwitterionic forms of cysteine.

Experimental: L-cysteine molecules (powder, 99.5% pure, Sigma-Aldrich) were sublimated at $\sim 120^{\circ}\mathrm{C}$ for 2 h, after degassing up to $120^{\circ}\mathrm{C}$ for 12 h. During deposition the pressure in the preparation chamber increased to around 5×10^{-9} mbar (from low tens). Deposition onto a cleaned TiO₂ substrate held at 100 K was employed to ensure a thick layer growth. Afterwards, no signal from the substrate could be observed, and the layer was determined to be at least 40 Å thick. All spectra were calibrated with respect to the position of the carboxylate O 1s peak at 531.9 eV obtained for thicker films deposited at 200 K⁹. During measurements the sample was kept moving at a speed, such that no difference between succesive scans (of around 30 sec each) was observed due to x-ray irradiation damage 12 .

Results: Figure 2 displays the O 1s, C 1s, N1s, and S 2p x-ray photoelectron spectra for a multilayer of L-cysteine deposited onto the $\text{TiO}_2(110)$ surface at 100 K. The best fit for the C 1s spectrum is obtained by using four components, the carboxylic carbon contribution denoted by the label c, that of the amino carbon by α . The two remaining peaks imply the presence of both carbon atoms related to hydrogen dissociated thiol groups (\mathbf{C}_d) , as well as those bound to intact thiol groups $(\mathbf{C}_{\theta})^9$. In the S 2p region two doublets are observed. The high and low binding energy components are attributed to thiol and deprotonated thiol groups, respectively. Both carbon and sulphur spectra indicate that 45% of the thiol groups are deprotonated.

The O 1s spectrum was curve fitted with three components, which, in order from low to higher binding

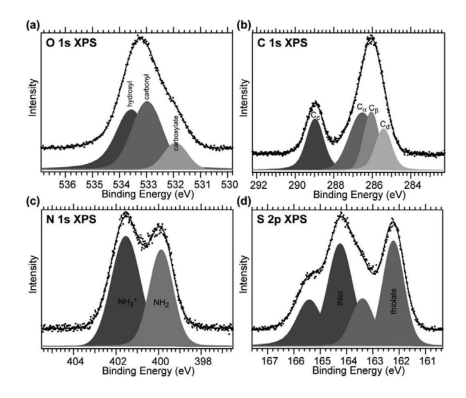


FIG. 2: a) Oxygen, b) carbon, c) nitrogen and d) sulphur x-ray photoelectron spectra with component fits as indicated. For carbon labels, see Fig. 1.

energies, are attributed to deprotonated carboxylic, carbonyl, and hydroxyl oxygen atoms, respectively. From the intensity ratio we deduce that only 13% of all carboxylic groups were deprotonated. We find a binding energy (BE) difference of 0.6 eV between the carbonyl and hydroxyl oxygen peaks. In literature somewhat larger values are reported: from 1.8 eV (amino acid molecules, gas phase 10) to 0.8 eV for glycine on Si(111) 11. Hydrogen bonding, where the hydroxyl and carbonyl groups act as donors and acceptors, respectively, could contribute to a lowering of the BE difference. We did consider the inclusion of a significant amount of water molecules in the grown layer as an explanation, but the intensity ratio of the oxygen peaks to that of nitrogen, measured at the same photon energy and corrected for

flux and cross-sections, is entirely conform stoichiometry.

The low and high binding energy peaks observed in the N Is spectrum can be attributed to photoemission from the amino and protonated amino groups, respectively. The intensity ratio shows that the amino groups of 57% of the molecules were protonated. When we now look at all the relevant percentages deduced from these measurements (45% S $^-$, 13% COO $^-$ and 57% NH $^+_3$) we find that a much larger fraction of the thiol groups than of the carboxylic groups is deprotonated. The thiol groups must therefore act as the primary donors of protons to the amino groups.

This means that, surprisingly, the unconventional zwitter ion is the favoured state in this case!

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XPS study of Rh bulkoxide on Rh(111) and 21 nm Rh nanoparticles

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Single crystal metal surfaces have been intensively studied by electron based UHV techniques for many years and XPS has been a major contributor to our present understanding of gas-surface interactions. Spectra obtained from single crystal surfaces are well known and the chemical shifts induced by gas molecules adsorbing on the surface can be determined with high precision and can be related to the exact adsorption structure on the surface. However, such measurements are performed ex situ and at low pressures while the latest development has utilized in situ High Pressure XPS studies [1] on more complex systems and at high pressures. These studies are relevant for applied surface science, such as investigations of model catalysts where the surfaces of nanoparticles are active during reactions under high pressure reaction conditions. One of the main problems when transferring from UHV to high pressure XPS is a dramatically decreased signal yield. A supported nanoparticle sample shows a high degree of disorder as well as a low coverage of the active material. The decrease of the signal, the low coverage and the disordered particles has a profound effect on the resolution of the spectra and may therefore inhibit the interpretation and decomposition of the spectra.

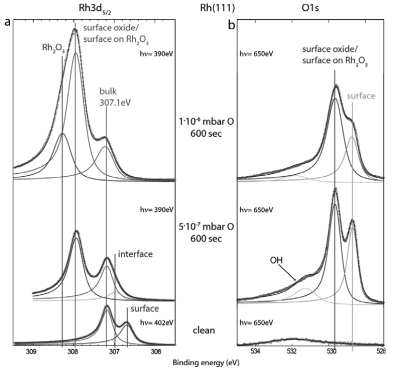


Figure 1. UHV XPS spectra of the oxidation of Rh(111) using atomic oxygen and 500°C. **a)** Rh 3d and **b)** O 1s.

In this report we compare results from UHV XPS from a Rh(111) single crystal and HPXPS from 21nm Rh nanoparticles [2]. We show that a consistent interpretation of the spectra from nanoparticles exposed to high pressure can be obtained using input from UHV based measurements and a well ordered single crystal surface.

The ex situ UHV experiments have been performed on beamline I311 at Maxlab using atomic oxygen produced by a gas cracker. Fig. 1 shows UHV XPS results from the oxidation of Rh(111). When the crystal is exposed to $5\cdot10^{-7}$ mbar O at 500° C, the LEED pattern shows the characteristic moiré pattern, and the spectra reveal the formation of the well-known trilayer surface oxide [3]. At higher oxygen coverage the moiré pattern disappears in LEED and the Rh $3d_{5/2}$ spectrum develops a clear shoulder

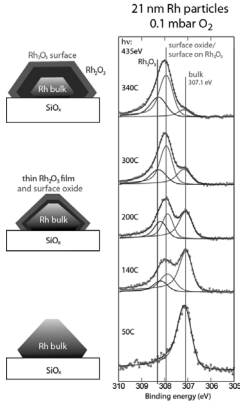


Figure 2. The Rh $3d_{5/2}$ spectra measured in-situ in 0.1 mbar O_2 as the temperature is increasing.

towards higher binding energy indicating the formation of a Rh bulk oxide. The surface core level shifts for the surface oxide is 0.75 eV and for the bulkoxide 1.12 eV with respect to the Rh bulk. Concurrently, the overall appearance of the O 1s spectrum is broadened. The high resolution of the UHV spectra makes it possible to distinguish between these two stages of the oxidation.

The above information has been of significant importance for the interpretation of the 21 nm Rh nanoparticels spectra measured *in situ* at 0.1 mbar O₂, see figure 2. The High Pressure XPS experiments were performed at beamline 9.3.2 at the ALS. By increasing the temperature stepwise, the oxidation of the particles can be followed. The appearance of two components in the Rh 3d_{5/2} spectrum shifted by 0.75 eV and 1.12 eV from the bulk Rh 3d_{5/2} component shows that the particles start to form a thicker oxide already at 140°C at 0.1mbar O₂. Further, a well-ordered surface oxide is never observed for the particles. At a temperature of 340°C, the metal bulk component has almost disappeared and the oxide is at least a couple of layers thick.

To conclude, the information gained by the UHV high resolution XPS has been used for the decomposition of spectra obtained at higher pressures and for more complex model systems. We show that

the oxidation of the 21nm Rh particles is similar to the Rh(111) single crystal, although the bulk oxidation appears to happen at lower temperatures for the nanoparticles.

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Production and characterization of PdAg aerosol nanoparticles on SiO_x

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Due to the huge economic and environmental rewards, one major goal in catalysis related research is to create cheaper catalysts. As catalysis happens on the surface of the catalyst, one possible way to realize this would be to dilute the more expensive active material with a less costly one, providing that the active material stays at the surface. This could be achieved by using a material which is less prone to interact with the reactant gases, such as a noble metal. In most catalysts, the active material is dispersed in a high area complex oxide support as nanoparticles. In order to maintain the high activity, it would be necessary to ensure that the active material is at the surface of the nanoparticle.

We have initiated a project to investigate whether this approach towards cheaper catalysts is viable. In the present contribution we report on our initial findings from attempts to produce PdAg alloy particles using an aerosol deposition technique [1, 2]. The use of PdAg is motivated by the fact that Pd segregates to the surface in the presence of a reactive gas while Ag segregates in UHV and that Ag is considerably less expensive than Pd.

In Fig. 1(a) we show a Scanning Electron Microscope (SEM) image from aerosol produced PdAg particles with a diameter of 10 nm deposited on an etched SiO_x substrate. To produce the PdAg particles, one Pd rod and one Ag rod were used as electrodes for the discharge sublimation of both materials with the aim to produce fully alloyed nanoparticles. It can be seen from the SEM image that the particles all have the same size, however obviously no information on the composition can be obtained. In Fig. 1(b) we show a Transmission Electron Microscope (TEM) image from a PdAg particle with a diameter of 15 nm. It can be seen that the particle has an approximately hexagonal shape as indicated, however it can also be seen that part of the particle is less well ordered marked in the upper right corner of Fig.1(b). The distance between the rows correspond to the distances between (111) planes. Unfortunately these distances are very similar for Pd and Ag (2.24 vs 2.35Å), making it difficult to distinguish wether the particle is Pd, Ag or a PdAg alloy which would have a value in between the Ag and Pd distances. The X-ray Energy Dispersive Spectroscopy (XEDS) analysis indicates that approximately 66-75% of each particle consists of Pd. However XEDS do not reveal whether the particle is an alloy or a mixed metal particle with separate Pd and Ag phases.

In order to gain further information we turn to High Resolution Core Level Spectroscopy performed at I311 at Max-lab and the HPXPS beamline 9.3.2 at ALS, Berkeley. In Fig. 1(c) we show the aerosol PdAg nanoparticles (top) after oxidation using 1 mbar of O_2 and reduction by CO following the cleaning procedure in Ref. [3]. For comparison is also the Ag $3d_{5/2}$ spectra from a clean Ag(110) single crystal surface (bottom), the clean $Pd_{75}Ag_{25}(100)$ surface (middle) shown. The Ag $3d_{5/2}$ bulk binding energy displays a shift of almost 0.5 eV between the pure metal and the $Pd_{75}Ag_{25}$ alloy. Density Functional Theory (DFT) calculations confirm the large alloy induced Ag $3d_{5/2}$ shift. Comparing the Ag $3d_{5/2}$ bulk binding energy in $Pd_{75}Ag_{25}(100)$ with the bulk Ag $3d_{5/2}$ binding energy from the PdAg particles, it is immediately clear that they are almost identical. From this observation, we therefore conclude that the aerosol PdAg alloy particles consist of a proper alloy and not Ag and Pd in separate

phases. Although the resolution is lower, the Ag $3d_{5/2}$ spectra from the particles also indicate that no significant amount of Ag is present at the surface of the PdAg particle in the presence of CO. Turning to Fig. 1(d) we show the Pd $3d_{5/2}$ core level from the Pd(100) (bottom), the Pd₇₅Ag₂₅(100) (middle) and from the PdAg nanoparticles (top). Here the shift between the bulk binding energy of the pure metal and the alloy is less prominent than in the Ag $3d_{5/2}$ level, however the surface core level shift of the alloy is opposite to that of the pure metal, an observation confirmed by theory. Again, the bulk binding energy of the Pd $3d_{5/2}$ from the PdAg particles coincides with that from the $Pd_{75}Ag_{25}(100)$ and deviates from the pure Pd metal. Further, most of the intensity from the spectra originates from a surface related signal, indicating that Pd segregates to the surface in the presence of CO, consistent with observations from the $Pd_{75}Ag_{25}(100)$ single crystal surface.

To conclude we have shown that it is possible to produce size selected PdAg nanoparticles with a high degree of alloying by an aerosol discharge deposition technique. In particular a comparison of the Ag $3d_{5/2}$ bulk binding energy from the particles and a single crystal $Pd_{75}Ag_{25}(100)$ allows for significant confidence in the characterization. Further, our results indicate that indeed the active Pd is present at the surface of the PdAg particles in the presence of reactant gases.

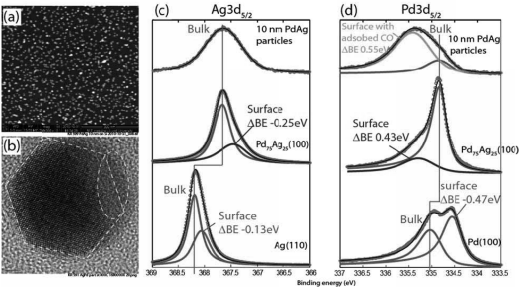


Figure 1 (a) SEM image from 10 nm large aerosol deposited PdAg particles on SiOx. (b) TEM image of a 15 nm large PdAg particle on a TEM grid. The hexagonal structure is indicated as well as a disordered region. (c) Ag $3d_{5/2}$ core level spectra from Ag(110), $Pd_{75}Ag_{25}(100)$ and the aerosol PdAg particles. Note the large shift between the bulk binding energy of the pure metal and the alloy. (d) Pd $3d_{5/2}$ core level spectra from Pd(100), $Pd_{75}Ag_{25}(100)$ and the aerosol PdAg particles.

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Large area quasi-free standing monolayer graphene on 3C-SiC(111)

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With its unique and appealing two-dimensional electron gas properties, graphene holds great potential for future electronics. Several different techniques are presently used to obtain graphene. Among those, by offering the possibility of growing large area graphene on semi-insulating substrates, thermal decomposition of silicon carbide (SiC) presents the most promising route towards a future of carbon-based electronics. Because they provide an ideal template for graphene growth, hexagonal SiC crystals, namely 4H and 6H-SiC, have evolved as the substrates of choice in the past years. In contrast, limited attention has been given to the possibility of growing graphene on the cubic SiC polytype (3C-SiC), although the [111] orientation of this crystal would also naturally accommodate the six-fold symmetry of graphene. Because of its extreme robustness and proven biocompatibility [1] cubic SiC is an extremely appealing platform for the growth of graphene that could lead to a new generation of microelectromechanical systems (MEMs) and advanced biomedical devices. Moreover, cubic SiC can be epitaxially grown on Si crystals and – obviously – this could significantly reduce the production costs of graphene. In the present work we report significant advances in the state-of-the-art production and characterization of graphene on 3C-SiC(111). We demonstrate that on cubic substrates it is indeed possible to obtain homogenous quasi-free standing monolayer graphene with domains extending over areas of hundreds of micrometers. We characterize the morphological, structural and electronic properties of these layers via atomic force microscopy (AFM), low energy electron microscopy (LEEM) and angle resolved photoemission spectroscopy (ARPES).

The 3C-SiC(111) samples used in this work were grown by hetero-epitaxy on a 4H-SiC(0001) substrate with the continuous feed-physical vapour transport (CF-PVT) method [2]. The 4H-SiC substrate was subsequently polished away. The high quality of the samples obtained was demonstrated by the almost complete absence of double positioning boundaries (DPBs) in optical microscopy. A graphene buffer layer was grown on these substrates by adopting the atmospheric pressure graphitization method presented in [3]. AFM analysis of the buffer layer's surface revealed a morphology similar to that reported in [3] with step heights of about 8 nm and terrace widths ranging typically between 2 and 7 µm (see Figure 1(a)). The sample was subsequently hydrogen intercalated as explained in [4]. The success of the intercalation process was demonstrated by sharp monolayer bands measured via ARPES in the home-lab in Stuttgart. However, different to quasi-free standing graphene on hexagonal SiC crystal [4], the π -band dispersion of the quasi-free standing monolayer graphene on 3C-SiC(111) indicated a slight ntype doping (see panel (c)). The origin for this electron doping is currently under investigation. LEEM inspection with a field of view (FOV) of roughly 57 µm was performed on different areas of the sample and consistently revealed an extremely homogeneous graphene thickness (indicated by the lack of contrast in the grayscale) all over the surface. Panel (b) displays a characteristic LEEM micrograph recorded from this surface. A darker contrast, symptom of a different graphene thickness, is only observed at the step edges and in restricted domains. The homogeneity of our graphene layers is highly remarkable considering that, up to date, the best LEEM data reported for graphene grown on SiC(111) presented domains with lateral dimensions of less than 1 µm [5]. The electron reflectivity spectra extracted from LEEM image series for regions of different contrast as indicated in panel (d) confirm that the sample is mostly covered by monolayer graphene (representative regions labeled A and B) with bilayer graphene being only

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present at the step edges (region C). Notably, in the present case of graphene grown on 3C-SiC(111), the minimum indicative of quasi-free standing monolayer graphene is positioned at an energy that is roughly 2 eV lower than that characteristic of graphene grown on 4H- or 6H-SiC(0001). Also an additional peak is visible at around 6 eV, possibly imputable to the different substrate layer stacking. The areas of lighter gray contrast that are running along certain crystal orientations (region B) are clearly identified as monolayer graphene also by the LEEM intensities. Hence, the contrast difference could be caused by defects in the substrate, which might be mediated by strain.

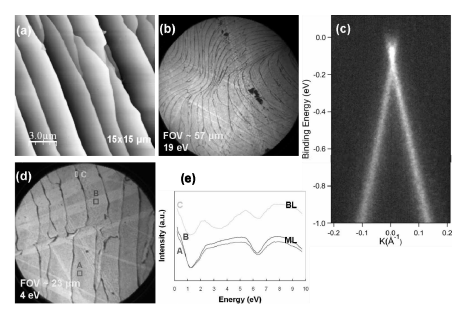


Figure 1. (a) AFM micrograph with lateral dimension of 15 μ m showing the morphology of a buffer layer graphene grown on SiC(111). (b) LEEM micrograph with FOV of ca. 57 μ m recorded with an electron energy of 19 eV. (c) Dispersion of the π -bands measured with ARPES perpendicular to the Γ K direction of the graphene Brillouin zone for quasi-free standing monolayer graphene. (d) LEEM micrograph with FOV of ca. 23 μ m recorded with an electron energy of 4 eV and labeled representative region A, B and C. (e) Electron reflectivity spectra measured for A, B and C (note that the curves are shifted on the y-axis for better display).

In conclusion, we demonstrate that large-area quasi-free standing monolayer graphene can be produced on 3C-SiC(111) substrates and investigate its morphological, structural and electronic properties. The high quality of the graphene obtained suggests that 3C-SiC(111) might be an appealing and cost effective platform for the future development of graphene technology.

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Large area quasi-free standing trilayer graphene on SiC(0001)

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Graphene epitaxially grown on SiC substrates is an appealing candidate for a wide variety of electronic applications, provided that large area, charge neutral layers can be produced. We have recently reported that hydrogen intercalation can be used to obtain technologically promising undoped quasi-free standing monolayer and bilayer graphene [1]. The growth of large area homogenous domains of trilayer graphene on SiC(0001) is indeed challenging as evident from the scarce amount of experimental data that can be found literature. At present steps required aim towards the production of high quality trilayer graphene so that its electronic properties can be experimentally investigated. In this work we demonstrate that by intercalating hydrogen we can obtain large-area undoped quasi-free standing trilayer graphene on SiC(0001). The structural and electronic properties of these layers are studied via low energy electron microscopy (LEEM) and angle resolved photoemission spectroscopy (ARPES).

Bilayer graphene was initially grown on semi-insulating hexagonal SiC(0001) substrates using the process described in [2]. The morphology of the obtained graphene was monitored via atomic force microscopy: well-ordered atomically flat surfaces not affected by stepbunching and presenting terraces as large as a few hundreds of nanometers could be observed. Subsequently, H-intercalation was performed at the home-lab in Stuttgart as described in [1]. The structural properties of the intercalated sample were studied via LEEM using the ELMITEC3 instrument at the end-station of beamline I311 at MAX-LAB. Figure 1(a) shows a characteristic LEEM micrograph with a field of view (FOV) of 15 µm for a hydrogen intercalated bilayer. At the reported energy of 1.2 eV, regions of different graphene thickness can be distinguished by the differences in reflected intensity. Even if surface domains with three different grayscale contrasts can be identified, the sample is highly homogeneous with light-gray and dark-gray domains occupying only a small amount of the overall area. The number of dips in the electron reflectivity spectra in panel (d) identifies the dark-gray and the light-gray regions as bilayer and tetralayer graphene, respectively [3]. The majority of the area is indeed covered by quasi-free standing trilayer graphene. Subsequently the sample was annealed in a stepwise fashion to gradually remove the hydrogen and hence gain insight on the desorption dynamics. After annealing above 900 °C the hydrogen starts to desorb at the edges of the atomic steps of the surface. In panel (b) the terrace structure of the surface is drastically emphasized by the additional dark-grey contrast present at the step edges which indicates how these regions have already transformed into as-grown bilayer graphene. By annealing at temperatures higher than 1100 °C the hydrogen completely desorbs as suggested by the reversed contrast in the LEEM micrograph in panel (c). The electron reflectivity spectra reported in panel (e) confirm that the sample is now for a large percentage covered by bilaver graphene.

The same sample was again hydrogen intercalated and measured via ARPES at the end-station of the SIS beamline at the Swiss Light Source synchrotron facility. The ARPES spectra reported in Figure 2 were measured perpendicular to the Γ K direction of the graphene Brillouin zone with a photon energy of 90 eV. An extremely sharp electronic band structure could be measured for the quasi-free standing trilayer graphene after outgassing between 400 °C and 600 °C (Figure 2(a)). Similar to what observed for hydrogen intercalated monoand bilayer graphene, also quasi-free standing trilayer graphene appears to be initially slightly p-type doped. The high quality of the measured band structure allows for a precise identification of the trilayer stacking sequence. By comparing the experimental bands with those theoretically calculated for graphene trilayer [4] it can be concluded that in this sample

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both domains with Bernal (ABA) and rhombohedral (ABC) stacking are present. As reported in [1], also in this case subsequent annealing at higher temperatures yields charge neutrality within a few meV (see panel (b)). Complete desorption of hydrogen reveals sharp bilayer bands (panel (c)) confirming that the small percentage of domains of different thicknesses observed via LEEM does not cause significant contributions to the measured band structure.

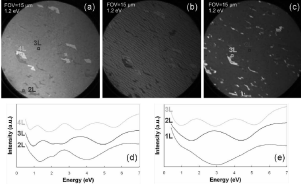


Figure 1. LEEM micrographs with FOV of 15 μ m recorded with an electron energy of 1.2 eV for the same area of (a) quasi-free standing trilayer graphene annealed at 400 °C, (b) annealed at above 900 °C and (c) at above 1100 °C. The electron reflectivity spectra measured for the regions of different contrast labeled in panel (a) and (c) are reported in panels (d) and (e), respectively.

In conclusion, we report on the structural and electronic properties of quasi-free standing trilayer graphene. LEEM analysis shows homogenous trilayer domains on areas of tens of squared micrometers and yields to the estimate that at least 90% of the sample is covered by three layers of graphene. High resolution ARPES provides evidence that quasi-free standing trilayer graphene can reach, upon UHV annealing, charge neutrality within a few meV. Moreover, thanks to the high homogeneity of the sample, the electronic structure of Bernal and rhombohedral stacking could be experimentally measured with high resolution.

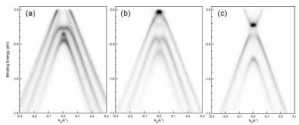


Figure 2. Dispersion of the π -bands measured with ARPES perpendicular to the ΓK direction of the graphene Brillouin zone for (a) quasi-free standing trilayer graphene after annealing at 600 °C, (b) at 700 °C and (c) after complete hydrogen desorption.

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Adsorption and decomposition of Alanine on Ni{111}

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The ability to selectively produce just one enantiomer of product is very desirable industrially, especially in the area of pharmaceuticals where two enantiomers can have vastly different therapeutic properties. Enantioselective catalysis allows the multiplication of chirality; large quantities of a chiral product can be synthesised using a catalytic amount of a chiral source. Stereochemical control can be induced on a non-chiral metal surface by the adsorption of a chiral auxiliary. For example the hydrogenation of β -ketoesters over nickel catalysts can be stereochemically controlled by modifying the catalyst with alanine [1]. These reactions have been optimised to give enantiomeric excess values near to 100% [2]. Several mechanisms have been suggested to account for the observed stereoselectivity but none have been proven.

In this experiment L-alanine was evaporated in UHV onto a clean Ni{111} single crystal. By performing this experiment in UHV the adsorption behaviour of alanine on this surface can be closely monitored unlike more complicated solution based studies. The Ni{111} surface was chosen because this is the most abundant surface termination found on supported Ni nanoparticles used in catalysis.

This experiment was performed on beamline I311 on MAX II. XPS (X-ray photoemission spectroscopy) was used to determine the coverage and to investigate the adsorption. The resolution of the spectra was calculated to be 0.6 eV. TP-XPS (temperature programmed XPS) was used to monitor changes in the structure and bonding and ultimately the desorption/decomposition routes of alanine. NEXAFS (near edge X-ray absorption fine structure) at the oxygen and nitrogen K-edges was recorded at three angles of incidence to determine the geometry with respect to the surface.

Alanine was found to adsorb onto Ni{111} (sample held at 250 K) as an intact molecule, as indicated by the XPS spectra in figure 1. In the O1s region, figure 1(b), only one peak is present indicating that both oxygen atoms are in the same chemical environment and hence that the carboxylate group is deprotonated. Both oxygen atoms make bonds to the Ni surface. The larger peak at 285.8 eV in the C1s XPS spectrum, figure 1(c), is due to the two alkyl carbon atoms and the smaller peak at higher binding energy is due to the carboxylate carbon. At this temperature, two peaks are observed in the N1s region with approximately equal intensity (1.00:1.05 on the lower binding energy side). For lower coverages, two peaks are also observed but the peak at lower binding energy is more intense. This suggests that there are two alanine species on the surface that differ only by the state of the N atom. The signal at 399.3 eV is due to a neutral NH₂ group whereas the signal at 401.5 eV is due to the zwitterionic form of the molecule, which involves an NH₃⁺ group. Upon heating this layer, figure 1(a), there is some conversion from the zwitterionic form to the neutral NH₂ form. After 345 K, there is no zwitterionic form left on the surface and a new peak appears at a binding energy of 397.5 eV. This peak is due to a decomposition product of alanine, most likely a CNH_x species. This signal disappears after annealing to 700 K at which point the O1s and C1s signals have also disappeared. On

the $Cu\{110\}$ and $\{531\}$ surfaces, alanine was only found to exist in a deprotonated state with a neutral NH₂ group [3,4].

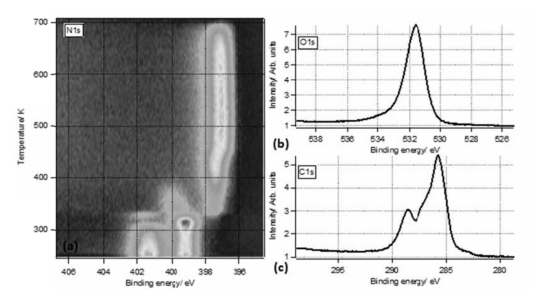


Figure 1(a) TP-XPS in the N1s region (b) XPS spectrum in the O1s region (c) C1s region of a saturated alanine layer on Ni{111} at a sample temperature of 250 K. All recorded at normal emission.

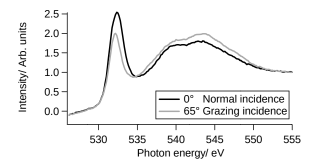


Figure 2 NEXAFS spectra at the O K-edge at two angles of incidence after a low dose of L-alanine on Ni {111}

Oxygen NEXAFS allows us to determine the orientation of the carboxylate O-C-O triangle [3-6]. A sharp π^* -resonance can be seen in figure 2 at 532 eV, and a broad peak due to σ -resonances starting at 535 eV. By measuring the angular dependence of the π^* -resonance intensity the angle of the O-C-O triangle with respect to the surface was calculated to be approximately 55°.

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Microscopic study of quasi-free standing epitaxial graphene and graphene *p-n* junctions formed on SiC by atomic intercalation of germanium

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Epitaxial graphene (EG) grown on SiC single crystal wafers appears to be a suitable candidate for graphene based electronics [1-3]. For further implementations it is, however, required to be able to control the electronic and structural properties of EG which are predominantly governed by the heterointerface with the substrate. Generally, solid heterointerfaces are quite difficult to influence since they remain inaccessible after growth. The heterointerface graphene-SiC(0001) is unique as its electronic properties can be manipulated on an atomic scale. For example, hydrogen can be used to passivate interface states [4]. Alternatively, the interface can be modified by atomic intercalation. We prepared quasi-free standing epitaxial graphene on SiC(0001) by intercalating atomically thin germanium film underneath the $(6\sqrt{3}\times6\sqrt{3})R30^{\circ}$ -reconstructed interface layer. The latter consists of a carbon layer with graphene-like topology and bond lengths but with strong covalent bonds to the SiC substrate [5]. These bonds are broken upon in-diffusion of Ge atoms to the interface so that the C-layer is structurally decoupled from the SiC surface. This leads to a complete recovery of the electronic structure of graphene (compare Fig. 1(a) and (b)). Depending on the annealing temperature graphene exhibits moderate p- or n-doping (Fig. 1(b-d)). Interestingly, an intermediate stage shown in Fig 1(c) is characterized by the coexistence of the p- and n-doped graphene regions, i.e. at this stage the surface splits into lateral graphene p-n junctions. In the present work the formation of graphene p-n junctions was studied in situ by low energy electron microscopy (LEEM) and photoelectron microscopy (PEEM) at the beamline I311.

In Fig. 2(a) we show the LEEM micrograph of the initial quasi-free standing graphene layer (p-phase, see Fig. 1(b)) obtained after intercalation of germanium. The graphene layer is quite homogeneous and covers the entire surface. Graphene domains are determined by the size of the SiC substrate terraces, that is, of the order of $3-5~\mu m$ in width.

Initial and later stages of the n-doped graphene phase formation are presented in Fig. 2(b) and in Fig. 3, respectively. The corresponding PEEM image in Fig. 3(b) reveals a significant contrast in intensity of the Ge 3d core level signal for the two graphene phases. Contrast in PEEM is determined by the concentration of Ge atoms located beneath the epitaxial graphene layer and coincides with the contrast obtained in LEEM. Apparently, the n-phase is induced upon a partial desorption of germanium from under the surface, i.e. partial deintercalation. Surprisingly, the process is not initiated at the step edges but rather on the terraces (see Fig. 2(b)). This leads to the formation of n-doped induced graphene islands with the size as small as 100 nm embedded into the p-doped graphene sheet. With further annealing the islands grow in size and coalesce forming extended n-doped graphene areas. We would like to comment that Ge atoms are mobile at quite low temperatures and the eventual pattern of p-n areas might depend on the cooling speed.

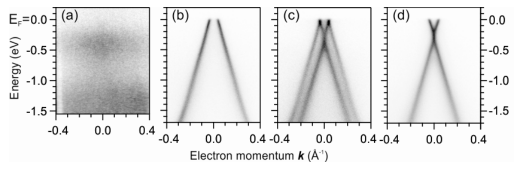


Fig. 1. Photoemission valence band maps (energy vs electron momentum) in the vicinity of the K-point (k=0) of the graphene Brillouin zone taken from: (a) the ($6\sqrt{3}\times6\sqrt{3}$)R30° surface and after deposition of 5 ML of Ge followed by vacuum annealing at (b) T=720°C, (c) 820°C, and (d) 920°C. The photon energy was 90 eV.

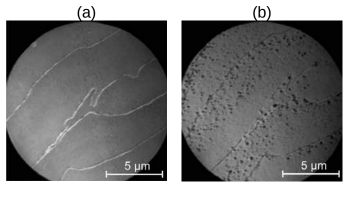


Fig. 2. LEEM micrographs of the quasi-free standing graphene layer obtained by intercalation of Ge atoms at the interface with SiC(0001) surface: (a) homogeneous p-doped graphene phase, and (b) initial stage of the graphene p-n junction formation. Dark inclusions in (b) correspond to the n-doped graphene islands embedded into the p-doped graphene.

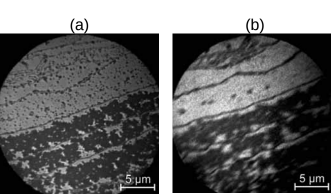


Fig. 3. (a) - LEEM micrograph and (b) - PEEM Ge3d intensity map of the graphene *p-n* coexistence stage (see also Fig. 1(c)). Bright (*dark*) regions in (a) correspond to p-doped (*n-doped*) quasi-free standing graphene regions.

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Oxidation of metal nanoparticle arrays

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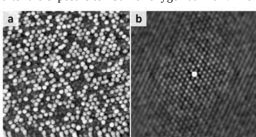
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The emergent phenomena in ordered arrays of surface supported oxide quantum dots represent a new challenge in nanoscience. An equally challenging endeavour is the fabrication of such nanodot model systems. Here we report the successful creation of a system of ordered oxide nanodots by using the directed assembly of metallic nanodots on a nanostructured surface followed by a morphology conserving oxidation process. The system of study has been Ni-oxide nanodots on an alumina thin film substrate.

The ultra-thin alumina substrate has been prepared by controlled high-temperature oxidation of a $Ni_3Al(111)$ single crystal surface, following the recipe detailed in the literature [1-3]. The complex AlO_x structure [1] provides an ordered lattice of hole sites, which act as anchoring sites for the Ni nanodots condensing them into a regular superlattice with 4.2 nm dimensions. The oxide nanoparticle superlattices have been prepared according to a two step process: (i) self-assembly of metal nanoparticles on the $AlO_x/Ni_3Al(111)$ substrate; (ii) oxidation of the metal nanoparticles under preservation of their nanoscale ordering. Nanodot arrays of Ni clusters have been formed by PVD at 473K on to the alumina template, using a "seeding" procedure [1, 3], in which Pd seeds promote metal cluster nucleation. The final and crucial step in the preparation is the oxidation of the metal nanoparticles under preservation of their ordered nanoscale morphology.

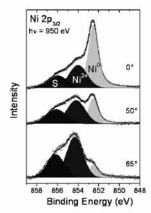
Using Scanning Tunneling Microscopy, we have demonstrated in our laboratories that a morphology conserving oxidation procedure is possible and that the resulting NiO_x nanodots form a highly regular superlattice. This is illustrated in Fig. 1. In the panel (a) a STM image of the Ni clusters on $\text{AlO}_x/\text{Ni}_3\text{Al}(111)$, after exposure to $1\text{x}10^{-6}$ mbar O_2 partial pressure at 470K for 15 min, is presented. The regular periodicity of the clusters is highlighted by the auto-correlation image depicted in the panel (b). The nanodots form a hexagonal superlattice with a periodicity of 4.2 nm.

The chemical identity of the oxidation products in terms of oxidation state has been investigated by high-resolution X-ray photoemission spectroscopy (HR-XPS) with the use of synchrotron radiation at MAX-Lab, beamline I311. Fig. 2 shows the core level spectra of Ni $2p_{3/2}$ from 0.1nm Ni clusters (nominal deposition amount) on the alumina film on Ni₃Al(111) after the exposure to 180L of oxygen at 470K. The top spectrum represents the measurement



<u>Fig.1</u>: (a) STM image $(100 \times 100 \text{ nm}^2)$ of Ni clusters on the alumina film on Ni₃Al(111) after exposure to 1×10^{-6} mbar O₂ partial pressure at 470K for 15 min; (b) auto-correlation plot of the image (a) to highlight the order of the clusters, with a periodicity of 4.2 nm.

at normal emission. Besides the metallic Ni⁰ peak, an oxide Ni²⁺ component is present with its satellite S, indicating a NiO stoichiometry [4]. In order to explore whether the metal nanoparticles can be converted completely into oxide particles or whether a metallic core remains, we have performed measurements as a function of the emission angle with respect to the normal direction. Increasing the angle, the contribution of the surface with respect to the bulk rises. In the present case, the oxide component significantly increases with



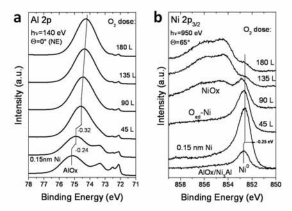
the angle whereas the metallic component is progressively reduced, as apparent from Fig. 2. This suggests that the Ni clusters on the surface are fully oxidized to NiO; note that there is a metallic component also from the Ni₃Al substrate, which is still detectable at 65°.

We have also found important changes in the AlOx substrate as a consequence of the Ni deposition and of the subsequent exposure to the oxygen. The evolution of Al 2p core level spectra at normal emission during the NiO nanodot fabrication process is depicted in Fig. 3, panel (a). Two shifts of 0.24eV and 0.32eV occur on the AlO_x peak, respectively, after the deposition of 0.15nm Ni and after the exposure to 45L of oxygen. A similar shift of 0.25eV is also present in the core level spectra of Ni 2p_{3/2} at grazing emission, as shown in the panel (b). This effect indicates a charge transfer from Ni₃Al to the Ni overlayer, due to the difference of work function between the alloy and the metal separated by the alumina, which plays the role of a barrier [5]. The second shift in the AlO_x peak is due to the initial oxygen chemisorption on the Ni clusters, which induces a further change in the overlayer work

function [5]. Moreover, the oxygen exposure increases the intensity of the Al^{3+} component, indicating that the alumina film becomes thicker. This is probably due to the catalytic role played by the Ni clusters, which dissociate the O_2 molecules [6]. The atomic oxygen penetrates then into the substrate and reacts with the Al atoms of the alloy at the interface, consequently increasing the alumina thickness.

conclusion. we have demonstrated that it is possible to fabricate ordered arrays of Ni oxide nanodots by oxidation of a superlattice of Ni nanoclusters. The clusters are fully oxidized and the stoichiometry is close to NiO. The Ni clusters on top of the alumina and the subsequent exposure to oxygen produce a charge transfer between the alloy and the Ni nanodots and influence the alumina layer by increasing its thickness.

Financial support was provided by the ERC Advanced Grant SEPON. Support during experiments by MAX-Lab is gratefully acknowledged.



<u>Fig. 3</u>: Evolution of (a) Al 2p normal emission (NE) and (b) Ni $2p_{3/2}$ grazing emission core level spectra during the nanodot fabrication process: pristine alumina, deposition of 0.15nm Ni, exposure to 45L, 90L, 135L, 180L O_2 at 470K.

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Gas adsorption on Pt clusters supported by graphene

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Single crystal surface studies do not fully capture the complexity of supported nanoparticles found in real catalysts. For example the adsorption properties can depend on both the size and shape of the active nanoparticle. To bridge this material gap more complex model systems such as well-characterized in-situ deposited nanoparticles can be used, but with these systems new problems arise. Often, the particles have a broad distribution of size and shape, making it difficult to correlate the adsorption properties unambiguously to the atomic structure of the nanoparticles with averaging techniques. Metal nanoparticles grown on a graphene/Ir(111) moiré film, however, show exceptionally well ordered arrays with a narrow size distribution [1, 2], thus making them an ideal model system for adsorption studies.

Here we present the first results of our studies of the interaction between Pt-cluster and the supporting graphene layer, and the effects of CO adsorption using photoemission X-ray spectroscopy (XPS) and density functional theory (DFT) calculations [3].

Figure 1 shows the C 1s spectra for the pristine graphene (black), after deposition of 0.18 ML Pt (dark grey), and after adsorption of 10 L CO (light grey). For the pristine graphene film a single peak at 284.15 eV is observed. After Pt-deposition we observe a broad shoulder shifted 0.3-1.1 eV with respect to the C 1s peak from pristine graphene. DFT calculations reveal that this shoulder is due to carbon atoms displaced towards the Ir(111) surface and positioned below and in the vicinity of the clusters. A schematic drawing of the graphene film before (a) and after Pt-deposition (b) is shown in figure 2 with the atoms assigned to the shoulder component highlighted.

Upon CO adsorption the Pt-induced shoulder diminishes in the C 1s spectrum (light grey curve in figure 1). We interpret this as

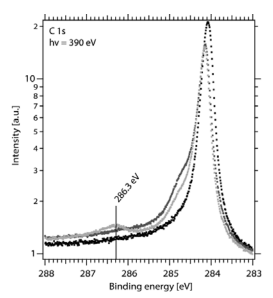


Figure 1: The C 1s photoemission spectra for the pristine graphene film (black), after deposition of 0.18 ML of Pt (gray), and after adsorption of 10 L CO (light grey). Notice the log-scale!

unpinning of the graphene film when CO adsorbs on the clusters. We also observe a new feature at 286.3 eV, coinciding with the peak position reported for CO adsorbed atop of steps on Pt(322) [4].

Consistent with atop CO adsorption the O 1s spectrum after CO exposure, shown in fig. 3, has its main peak positioned at 532.4 eV, agreeing well with the position reported for a top adsorption of CO on Pt(111) [5]. A schematic drawing of the unpinning and CO adsorption is shown in figure 2 c.

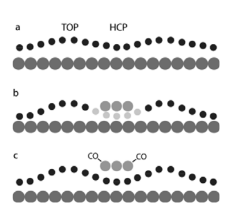


Figure 2: Schematic drawings of a) pristine graphene on Ir(111), b) Pt-cluster on graphene, and c) CO adsorption on the Pt-clusters.

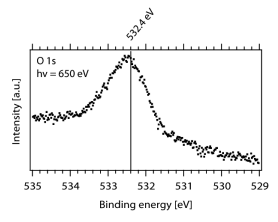


Figure 3: The O 1s spectrum after deposition of 0.2 ML Pt and subsequent adsorption of 10 L CO.

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The effect of Au particles on the morphology and Ga droplet dynamics on the GaP(111)B surface

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We have found that Au nano particles (50-100 nm diam., 0.1- $1/\mu m^2$) affects the surface morphology of GaP(111)B. This changes the dynamics of the self-propelled Ga droplets that form on the surface when annealing above the maximum temperature for congruent evaporation.

After deoxidation at around 600°C and cooling down of GaP(111)B with Au particles, patches that appear bright for certain electron beam energies in low energy electron microscopy (LEEM) are observed (fig. 1A). Low energy electron diffraction patterns from micrometer sized areas (μ -LEED) within and outside the patches were recorded. The μ -LEED pattern from the dark area outside the patches indicates a high density of pyramid shaped crystallites (fig 1B) which we have seen also on the clean GaP(111)B surface after oxide desorption [1]. The μ -LEED pattern from the bright patches is that of the so called Hattori reconstruction which indicates a low density of crystallites (fig 1C). In each bright patch there is at least one Au particle which appears as bright spots in UV-PEEM (inset of fig 1A). We therefore believe that the Au particles enhance the evaporation of pyramid shaped crystallites in their close vicinity.

Further annealing above the temperature for congruent evaporation leads to the formation of Ga droplets. On the clean surface these self-propelled droplets move all in the same direction up the steps introduced by the wafer miscut [2]. It is energetically favourable for the droplets to cover as many step sites as possible and the droplets will become pinned to the steps. The steps move faster under the droplet as the rate of evaporation is enhanced there and the droplet will follow as the steps move. When the steps are arranged along a particular direction across the sample the droplets will all move in this direction. The pyramid shaped crystallites also evaporate faster under the droplets and a trail of smooth well ordered surface is found where the droplets have passed.

On the surface with Au particles where the density of pyramid shaped crystallites is not homogenous as on the clean surface it is more energetically favourable for the droplets to sit on an area with many pyramids. If a droplet is sitting on the boundary between an area with few and an area with many pyramids it will feel a force towards the area with more pyramids. The balance between the force from the moving steps and the force towards areas with a

higher pyramid density will lead to an irregular movement of the droplets but with an average direction up the surface steps. On the sample shown in the LEEM image of fig 1D Au particles has been deposited only on the left half. On the right side with no Au particles the droplet trails are straight whereas on the left side with Au particles the trails are more irregular.

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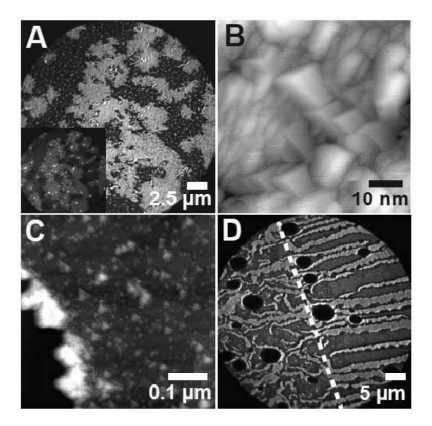


Figure 1

- **A)** LEEM image of the GaP(111)B surface with Au nano particles after oxide desorption and cooling down. The bright areas have a lower density of pyramid shaped crystallites. There is an Au particle in each bright area as can be seen in the UV-PEEM image (inset).
- **B)** STM image of an area with a high density of pyramid shaped crystallites.
- **C**) STM image of an area with a low density of pyramid shaped crystallites.
- **D**) After the formation of Ga droplets. The droplet trails are more irregular on the left side of the sample that has Au particles.

nin-InP nanowires studied by x-ray photoelectron emission microscopy

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III-V semiconductor nanowires (NWs) have long been proposed as future key components within a wide range of areas, e.g. sensing, (opto)electronics and photovoltaics. Because the nanowires allow integration of III-V materials directly with the existing Si platform used in industry, they enable high performance at low cost. Much of the research performed so far has been focused on growing perfect single crystal nanowires, but a prerequisite for most devices is the ability to in a controlled manner, both spatially and magnitude wise, dope the structures.

We have performed x-ray photoelectron emission microscopy (XPEEM) on doped InP nanowires at beamline I311 at MAX-lab. The nanowires were grown using Metal Organic Vapor Phase Epitaxy (MOVPE) and an Au-seed particle was used to define the size of the wires. The precursors for the growth were trimethylindium (TMI) and phosphine (PH $_3$) while H $_2$ S was used as precursor for n-type dopants. During growth the supply of dopants was switched rendering a segment stacking of n-type – intrinsic – n-type (nin) as can be seen in the schematic in figure 1 a. After growth the nanowires were transferred to a HF-etched Si-wafer and put into vacuum. The nanowires were cleaned in an atomic hydrogen atmosphere at 400°C for 8 min, removing most of the native oxide.

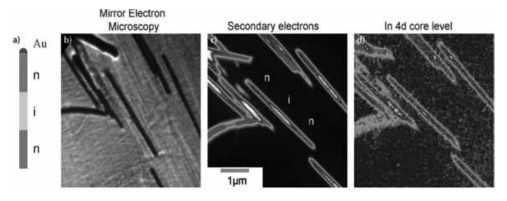
In fig 1 b. a mirror electron micrograph of the nin-NWs can be seen. In mirror mode, the sample is illuminated by low energy electrons that are reflected before they reach the surface. This mode is mainly used in order to get a (fast) overview of the NWs and to inspect the surface for possible contaminants. In fig 1 c. x-ray photons are incident on the sample and it is immediately observed that dopant induced contrast can be observed along the NW when secondary electrons are collected. It was found that the intrinsic segment did emit more secondary electrons than the n-parts at all photoelectron energies tested (0-3 eV) and at both photon energies tuned in (70 and 130 eV). The contrast can be explained using a model first proposed by Sealy et al. assuming a locally varying vacuum level potential just outside the sample surface [1]. If the ionization potential is equal in both the n and i-parts $(E_n=E_n)$, which would be fair to assume due to the low amount of dopants incorporated in the lattice, then the vacuum potential as it appears just outside the sample has to vary, reaching a uniform level (E_{detector}) first at a macroscopic distance. In order for an electron from the n-type side to not only leave the sample surface but also to reach the detector it requires an energy equal to $E_n + \Delta E_n$, that is ΔE_n more energy than a corresponding electron from the intrinsic side would need. This causes a higher secondary electron flux from the intrinsic part since those electrons generally need less energy to surmount the barrier and reach the detector.

With the SPELEEM one can also tune in and make spatial maps of different elements in the sample. In fig 1 d. electrons from the In4d core level has been tuned in and shows were In can be found in the sample.

[1] C.P. Sealy et al., J. Electron Microsc. 49 (2000) 311

Figure 1

- a) Schematic showing the grown nin-structure
- b) Mirror electron micrograph of the NW shown in (a). Start voltage 0.39 eV.
- c) XPEEM image showing the NW in (a). The image is acquired using secondary electrons and the n and intrinsic segments are indicated by n and i respectively. Start voltage0.6 eV, photon energy 70eV.
- d) XPEEM image showing the NW in (a). The image is acquired using In4d core electrons and thus showing a spatial map of In in the sample. Start voltage 47.5 eV, photon energy 70 eV.



Formation of Carbonyl Complexes by CO Adsorption on Monolayers of Iron Phthalocyanine on Au(111)

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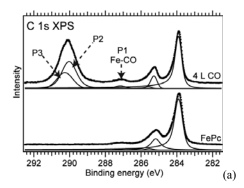
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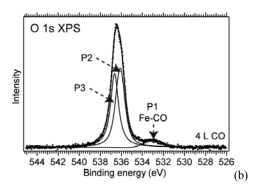
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The interaction between monolayers of iron phthalocyanine on a Au(111) support and carbon monoxide (CO) molecules is studied by photoelectron spectroscopy. The CO molecule is a very common ligand in coordination chemistry and the Metal-CO complexes are most often of extreme biological importance. Since the formation of such biologically active complexes involves the interaction between the CO ligand molecules and the metal centre of the metalloprotein cofactor, it is important to develop a fundamental understanding of such interactions.

The adsorption results in the formation of several carbon monoxide adsorbate species, as can be seen from the Figure 1 (a) and (b) showing the C 1s and O 1s spectra at saturation. Among these species of central importance for the present study is the formation of the FePc-carbonyl complex (peak labeled P1, showing up at ~287 eV in the C 1s spectrum and at ~533 eV in the O 1s spectrum). It has to be mentioned that the saturation represents the gas dose at which all the iron centers are coordinated to CO molecules (this we are able to show by comparing the Fe 2p spectra with increasing the amount of gas; from saturation onwards there is no change in the shape of the Fe 2p line shape). The formation of FePc-carbonyl complexes is particularly important, because it leads to a redistribution of the electrons in the iron 3d levels resulting in a change of spin state, as can be seen from a comparison of the Fe 2p photoelectron spectra before and after adsorbing CO at saturation (Figure 1 (c)). It is well known that the broad structure in the Fe 2p spectrum is caused by the open shell structure of the iron ion.¹⁻⁴ The decrease in the width of the Fe 2p line upon ligand adsorption is a consequence of reduction of the valence spin. The increase in the binding energy of the low binding energy Fe 2p component also shows that the formation of FePc-carbonyl bonds leads to a decrease of electron density on the iron ion. Since the metal-CO bond formation is known to involve both direct electron donation from the ligand to the metal and back donation from

the metal orbitals of appropriate symmetry to empty ligand orbitals, this implies that in the case of CO adsorption the back-donation is stronger than the direct donation.





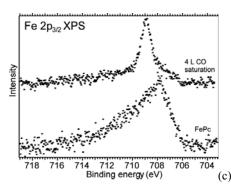


Figure 1. (a) C 1s photoelectron spectra before and after CO adsorption on a monolayer of FePc on Au(111); (b) O 1s photoelectron spectra after adsorption of CO at saturation (4 Langmuirs); Fe $2p_{3/2}$ photoelectron spectra before and after CO adsorption at saturation on a monolayer of FePc.

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Spin Change and Adsorbate-Substrate Decoupling as a Consequence of NO Adsorption on Iron Phthalocyanine Monolayers

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The NO molecule is a very reactive, free radical species. It is a signalling molecule in many physiological processes in the human body (neurotransmitter, cytoprotective molecule, regulation of cardiovascular function), being one of the smallest biologically active molecules. The adsorption of nitric oxide (NO) molecules on iron phthalocyanine (FePc) monolayer networks on Au(111) and their influence on both the iron's valence spin and the FePc-Au(111) coupling was investigated by means of photoelectron spectroscopy.

The NO molecules were adsorbed on the FePc monolayer in increasing amounts. The Fe $2p_{3/2}$ spectra before and after adsorption of increasing amounts of NO are shown in Figure 1. Two major changes can be observed in the evolution of the spectra: first, the lines become significantly narrower, and second they shift in binding energy. More quantitatively, the broad, multiplet structure the Fe 2p spectrum of the monolayer with a FWHM of 3.7 eV narrows down 1.3 eV for NO at "iron saturation dose". This dose corresponds to the point where any increase in the amount of adsorbed gas does not lead to any further changes in the shape of the Fe 2p spectrum. At the iron saturation dose all the iron centres are bound to NO ligand molecules and there are no more binding sites available on the iron. Figure 2 shows the results of least square fitting for the Fe $2p_{3/2}$ photoemission spectrum of the clean FePc monolayer, as well as the corresponding spectrum at saturation (4 Langmuirs), after NO adsorption. The monolayer spectrum was fitted by two main multiplet lines (P1 and P2) and a satellite (P3), while the spectrum after NO adsorption was fitted by one main line only (P2) and a satellite (P3). We assign the low binding energy line (P1) of the FePc/Au(111) to a component arising from charge transfer from the Au(111) substrate to the Fe atom in either the initial or final state. The fact that the low binding energy P1 peak is vanished after NO adsorption is a clear indication for the NO induced FePc-Au(111) decoupling. The middle component P2 shows up at 708.28 eV for FePc/Au(111) and it has a FWHM of 2.39 eV,

while after NO adsorption P2 is shifted to 708.81 eV and narrowed down to 1.25 eV. Concerning the decrease in width, it is known from the literature that the broad structure in the Fe 2*p* spectrum is caused by the open shell structure of the iron ion.²⁻⁵ We attribute the decrease in width of P2 to a decrease in the spin state on the iron atom, while the increase in P2 binding energy basically indicates that the electronic density on the iron atom is decreased by the adsorption of NO ligands.

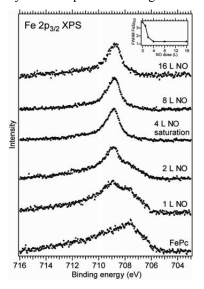
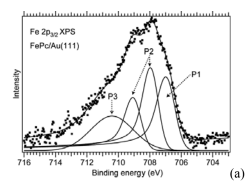


Figure 1. Fe $2p_{3/2}$ photoemission spectra with increasing amounts of NO. The spectra become narrower with increasing the amounts of NO gas. The NO amount of 4 Langmuirs represents the saturation dose.



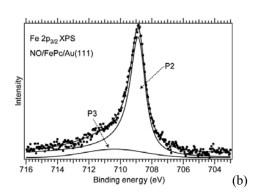


Figure 2. Fe $2p_{3/2}$ photoemission lines (experimental data and fit) for (a) monolayer FePc/Au(111), (b) NO adsorption on monolayer FePc/Au(111) at saturation.

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Evolution of surface compounds during the initial stages of high temperature oxidation of FeCr alloys

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FeCr-based alloys with low Ni content (ferritic stainless steels) have gained recent interest due to their excellent thermal properties, mechanical strength, and corrosion resistance, which make them an attractive choice for many energy applications ranging from exhaust systems of combustion engines to solar thermal collectors and fuel cells. The enhanced properties of advanced ferritic grades are often achieved through careful optimization of the alloy composition, even at parts-permillion level for certain minor elements. The influence of minor alloying elements is particularly crucial at the surface of the material, where they may become enriched through segregation and oxidation phenomena under corrosive or high temperature conditions. Better understanding of these phenomena is essential for developing novel surface treatment methods for stainless steels and improving their performance under demanding application environments.

In the present work, we studied segregation phenomena and formation of surface compounds on Fe-17Cr alloy at temperatures up to $800\,^{\circ}\text{C}$ upon annealing in UHV and during kinetically controlled oxidation by O_2 and H_2O . We performed high-resolution PES and XAS measurements, which provided information on the chemical states of the segregating elements and their behavior in high temperature surface phenomena in much greater detail than any preceding work based on XPS and AES measurements. In particular, we obtained previously unavailable spectroscopic information on the evolution of the CrN surface compound and its electronic properties.

The Fe-17Cr surface was initially cleaned *in situ* by cycles of 2.0 keV Ar^+ ion sputtering and subsequent annealing at 800 °C. Surface segregation was induced by annealing the sputter-cleaned sample in UHV at selected temperatures up to 800 °C for 5 min, and then allowing it to cool freely (within 1 h) to 50 °C. Isochronal oxidation experiments of 10 - 100 Langmuirs were performed on the pre-annealed surfaces by backfilling the UHV chamber with $5.3 \times 10^{-8} - 5.3 \times 10^{-7}$ mbar of either O_2 or H_2O while maintaining the sample at the oxidation temperature for 250 s. Chemical composition of the surface was subsequently analyzed by PES employing photon energies of 494 eV and 700 eV for the main part. Relative atomic concentrations and chemical states of elements were determined from the spectra by least-squares fitting of asymmetric Gaussian-Lorentzian lineshapes to the photoelectron peaks after subtracting a Shirley background.

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Figure 1 shows the complex evolution of surface compounds on the Fe-17Cr alloy resulting from annealing under UHV conditions. At 527 °C and above, the outermost surface consists mainly of CrN compounds, although minor alloy constituents such as Sn also produce strong signals due to their relatively high photoelectric cross-sections. The high-resolution PES data allows for the differentiation of surface and subsurface N species, thus facilitating the study of chemical bonding and compound formation on Fe-17Cr.

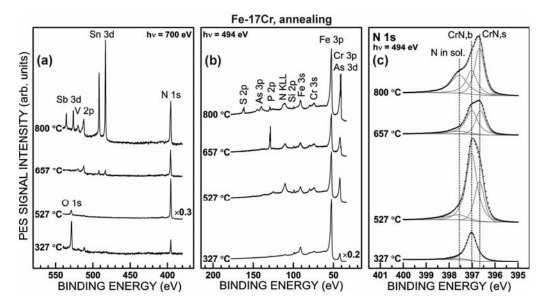


Figure 1. Evolution of surface compounds on the free Fe-17Cr alloy surface resulting from thermally induced segregation at 327 °C - 800 °C. The PES measurements were carried out by utilizing photon energies of (a) 700 eV and (b) 494 eV, which reveals many features that could not be detected by conventional XPS employing Al Kα radiation. In particular, the high-resolution spectra of N 1s (c) can be utilized for identification of different N species: two-dimensional CrN surface compound (CrN,s), CrN precipitates (CrN,b), and interstitial N in solid solution (N in sol.).

Subsequent oxidation experiments (not shown) indicate that upon O_2 exposure at 327 °C, an Fe-rich mixed oxide layer develops on the surface, whereas at 800 °C, the oxide consists of protective Cr_2O_3 - and SiO_2 -type oxides instead. In addition, the formation of protective oxide species is initially enhanced by the cosegregation of Cr and N, and by adsorption of H_2O . N remains at the oxide/metal interface as the oxidation proceeds, which calls for further investigation due to possible effects on the growth kinetics of oxides and their adhesion to the alloy surface. These phenomena are of great importance in processing and application environments involving low pO_2 conditions, such as surface finishing of FeCr alloys under reducing atmospheres (bright annealing).

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Carbonate formation on p(4x4)-O/Ag(111)

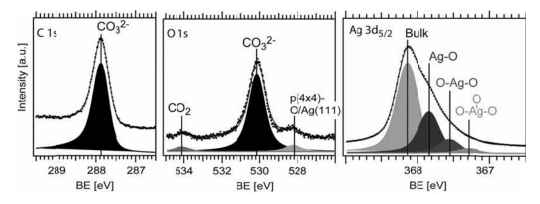
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One way to reduce fuel consumption and CO_2 emissions is to operate engines at oxidizing conditions with a high air to fuel ratio. Under such oxidizing conditions the traditional Pt/Pd based three Way Catalysts are unable to efficiently reduce nitrogen oxides (NO_x) to nitrogen (N_2) and other catalysts have to be used instead. Ag/Al₂O₃ is one promising catalyst for NO_x removal in an oxidizing environment, since it is able to selectively reduce NO_x using unburned hydrocarbons (HC) from the exhaust gas as reducing agent. The oxidation state of Ag in the working catalyst is not known, but it is likely that silver is present in an oxidized state due to the oxidizing environment. As a first step towards an atomic scale understanding of the catalytic activity of the Ag/Al₂O₃ system we initiated a study of adsorption of CO and NO on oxidized silver surfaces representing extremely simplified model systems of the real catalysts.

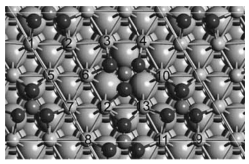
In this report we focus on CO adsorption on the p(4x4)-O/Ag(111) surface [1, 2], which is known to form on the Ag(111) surface under oxidizing conditions. Using the interplay of high resolution photoemission spectroscopy (HR-XPS) and density functional theory (DFT) we will show that carbonates are formed on the p(4x4)-O/Ag(111) surface at low temperatures, resembling the high pressures in the real catalysts.



Figur 1: C 1s, O1s, and Ag 3d5/2 after the p(4x4)-O/Ag(111) surface oxide has been saturated with CO at 100 K

Figure 1 shows the XP-spectra C 1s, O 1s and Ag $3d_{5/2}$ recorded after the p(4x4)-O/Ag(111) has been saturated with CO at 100 K. Both the C 1s peak at 287.9 eV and the dominant O 1s peak at 530.2 eV are characteristic for carbonates [3]. The O 1s peak at 528.2 eV assigned to oxygen in the p(4x4)-O/Ag(111) structure almost disappears upon CO dosing

indicating that all O-atoms on the surface are bonded in carbonate molecules. Further, our experiments suggest that each CO molecule



Figur 2: Proposed structural model of the carbonate structure with 1 monolayer of ${\rm CO_3}^2$ adsorbed. Colorcode: C gray, O black, Ag light gray.

react with two oxygen atoms on the surface, since the total O 1s intensity after CO dosing corresponds to 0.6 ML, which should be compared with 0.4 ML for the p(4x4)-O/Ag(111) structure.

Using this information as input for extensive DFT calculation we calculated the total energy of several model structures of the carbonate structure. Figure 2 shows the structural model with the lowest total energy in the set of tested structures. Further validation of our structural model comes from the calculated Ag $3d_{5/2}$ core level shifts (CLS), which show that surface Ag atoms bound to one (1-9 in fig. 2), two (10-12 in fig. 2), and three oxygen atoms (13 in fig. 2) are shifted -0.41 eV, -0.68 eV, and -0.95 eV, respectively, using the value for Ag bulk atoms as reference. These values are in good agreement with the measured Ag $3d_{5/2}$ spectrum shown in fig. 1, which can be deconvoluted into 3 surface components having an intensity ratio of 9:3:1 and shifted -0.30 eV, -0.58 eV, and -0.85 eV, respectively.

In contrast to the carbonate formation observed upon CO exposure at 100 K, room temperature exposure leads to titration of surface oxygen in p(4x4)-O/Ag(111) structure due to formation of CO_2 , which directly desorbs to the gas phase. Low temperature and low pressure close to UHV is thermodynamically equivalent to working at high temperatures near atmospheric pressure. The proposed atomic scale structure of carbonate formed on the p(4x4) structure is therefore not at all unrealistic for oxidized silver surfaces exposed to CO near atmospheric pressure. In this respect, it is interesting to note that we observed carbonate formation already at 100 K, clearly indicating that the carbonate reaction barrier is low compared to temperatures in the Ag/Al_2O_3 catalyst connected to the exhaust gas.

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Surface versus interface composition by element specific microscopy: Co doped ZnO

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ZnO doped with transition metal ions is the most studied oxide dedicated to spintronics applications. Zinc oxide doped with Co was the first diluted magnetic semiconductor (DMS) for which room temperature (RT) ferromagnetism (FM) was reported [1]. However, the reports from different research groups, contradict each other and despite the intensive studies of this material room temperature ferromagnetism in (Zn,Co)O is far from being fully understood. Despite the controversy on the magnetic response of (Zn,Co)O, a direct element specific mapping of this materials composition is lacking in the literature. We therefore, present here some X-ray Photo Electron Emission Microscopy (X-PEEM) and spectromicroscopy results with the associated Low-energy Electron Emission Microscopy (LEEM) micrographs. This experiment is performed at the I311 beam line of MAX-lab.

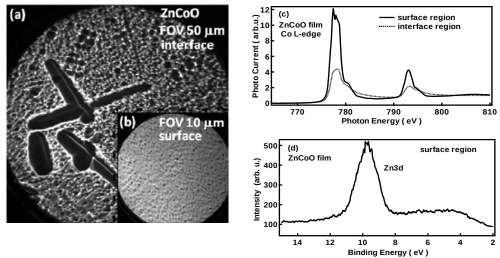


Figure 1. (a) LEEM micrograph taken in the mirror mode on ZnCoO film. The Field of View (FOV) is indicated. In the interface region needles of about 2000 nm long and 200 nm wide are observed. (b) LEEM micrograph of the surface region. The sample appears homogeneous in the 10 micron range. (c) The Co $L_{3,2}$ -edge measured with XAS (BL I1011) varies between surface and interface region. At the surface the shape of Co edge is typical for DMS ZnCoO films. At the interface we observe metallic Co, which exhibits also dichroic magnetic contrast. (d) The Zn3d states are homogeneously distributed in the surface region probed.

The samples presented in this report were grown on silicon substrates by means of the Atomic Layer Deposition (ALD) technique at the IF PAN (Warsaw, Poland) [2]. Prior to the X-PEEM work, the samples have been characterized by means of SQUID magnetometry and electron microscopy (SEM). The samples were chosen at various ALD external growth control parameters, and are exhibiting different magnetic properties. In order to best exploit

the surface sensitivity of X-PEEM, but also to study the inner of the material, some (Zn,Co)O films were Ar ion etched at the PAN under well defined conditions. The etching process allows to create samples with a variable and well defined (Zn,Co)O thickness profile, ideal for soft x-ray X-PEEM investigations. First results on the etched samples have been recently presented [3]. Prior to the work at I311, X-ray Absorption Spectroscopy data were taken at the I1011 beam line of MAX-lab. Based on several laboratory based techniques at IFPAN, such as a combination of SEM, SIMS, cathodo-luminescence, SQUID and XPS a model for the element specific profile as a function of film thickness for the ALD grown samples was proposed, according to which, Co rich regions are forming at the "inner" (Zn,Co)O/Si interface of the films [3]. X-PEEM experiments in combination with spectro-microscopy and XAS can probe directly the element specific profile of the etched films.

We studied both, as grown, as well as etched samples. For the etched samples a variable thickness profile is obtained across the sample surface, allowing to investigate the (Zn,Co)O films as a function of the layer thickness. The XAS results at I1011 are in agreement with this model. Some XAS spectra are shown in Figure 1(c). The spectra taken at the "outer" surface region exhibit much stronger white lines at the Co L-edges. The various ALD growth conditions do not appear to influence the elemental composition of the near surface region as probed by XAS using the sample photo-current. At the "inner" interface, the much lower white line intensity and the higher inter peak region are indicative of Co in close to a pure metallic state. The topography of the samples is characterized in the LEEM and X-PEEM modes. At the surface, for several samples probed no clear topological fine structure is identified in this mode, within 50 nm. For a few samples, in mirror mode (MEM) some features are visible in the 100 nm range, in agreement with the SEM results. For the nonetched samples all elements, Zn Co and O were identified in the micro-XPS mode. An example is shown for the Zn 3d states in Figure 1 (d). The samples appear homogeneous in the 50 microns range. Finally, for the etched samples the elemental composition is found to vary close to the inner interface region. Quasi circular islands of up to 4.5 µm in diameter, as well as needles up to 12 μm long and 4 μm wide, are clearly imaged. A micrograph is shown in Figure 1 (a). Depending on the precise etching process, some of the islands identified in our study, appear to contain Zn in two different chemical states. Some other nano crystals appear to be Co rich in agreement with the XAS results of Figure 1(c).

In conclusion, we provide by means of LEEM and X-PEEM direct evidence of topological micro-structure for the "inner" interface of ALD grown (Zn,Co)O films on Si. The reported measurements provide microscopic data which allow to explain some conflicting results in the literature, and support a novel interpretation of the magnetic properties for this material [3]. For some of the samples quasi metallic Co islands appear to form in the "inner" interface in agreement with XAS measuremets at I1011 on the same samples.

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High resolution core level spectroscopy study of the ultrathin aluminum oxide film on NiAl(110)

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Well-ordered oxide films grown on single crystal substrates have attracted considerable attention in the past 20 years mainly because of various applications in microelectronics, catalysis, anticorrosion, nanotemplates, coatings and sensors [1-7]. One of the most common model substrate for model catalysts is a thin aluminum oxide film grown on NiAl(110). It has previously been shown that well-ordered ultrathin oxide layers of about 0.5 nm thickness may be grown by oxidation of NiAl(110) [8-14]. The thin aluminum oxide film grown on NiAl(110) has been extensively studied in the literature but its structure has only recently been solved by Kresse et al.[1], and so far no independent experimental study in the literature supports the model by Kresse et al. [1] the ultrathin aluminum oxide film grown on NiAl(110) consists of a double oxide layer with NiAl-Al_i-O_i-Al_s-O_s stacking.

We have studied the ultrathin aluminum oxide film on NiAl(110) by a combination of high resolution core level spectroscopy (HRCLS) and density functional theory calculations (DFT).

In Fig. 1a we show the Al 2p and O 1s core levels recorded at normal emission with photon energy of 130 eV and 900 eV, respectively, from the ultrathin aluminum oxide on NiAl(110). It is immediately clear that the broad appearance of the core level spectra related to the oxidized Al atoms is due to a large number of shifted components originating from Al atoms in the oxide film with slightly different local environment. The Al 2p recorded with a photon energy of 130 eV is presented in Fig 1a right. The spectrum can be decomposed into 4 components as shown in Fig. 1a, each of them splitted by 0.4 eV and with a ratio of 1/2. The component at the lowest binding energy (72.4 eV, grey) is the so-called interface component, electron emission from the Al atoms in the NiAl(110) surface directly below the ultrathin Al oxide. The two Al 2p components shifted 1.00 eV and 2.27 eV are assigned to photoemission from Al atoms in the Al₁₀O₁₃ film. To distinguish the photoemission from Al_i and Al_s atoms we used energy dependent photoemission measurements, as shown in Fig. 1b and d. It can be seen that at higher energies, the 2.27 eV shifted component is decreasing more rapidly in intensity as compared to 1.00 eV shifted component which is also decreasing but not so fast. The measurements reveal that the 1.00 eV shifted component should be assigned to Al atoms from the interface of the oxide layer Ali and the 2.27 eV shifted component to the Al atoms at the surface of the oxide Als.

The O 1s spectrum recorded with photon energy of 900 eV is presented in Fig. 1a left. It can be seen that the emission is broad and asymmetric and exhibits a shoulder on the high binding energy side. The spectrum can be decomposed into two components. The energy dependent measurements shown in Fig. 1 c and e reveal strong diffraction effects at lower kinetic energies, however at higher energies, the smaller component decreases more rapidly in intensity as compared to the larger component suggesting that the small component is from the atoms in the O_s layer. However, it is clear that not all of the oxygen atoms in the O_s layer can contribute to the intensity, since a significant higher intensity would in that case be

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expected. By comparing our experimental findings to the calculated binding energy shifts the small component could be assigned to surface oxygen atoms which do have Ali atoms underneath and do not reside very close to another oxygen atom corresponding to only 30% of the surface oxygen atoms. The present HRCLS study supports the model by Kresse et al. and allows for a correct assignment of the different HRCL components to specific atomic sites.

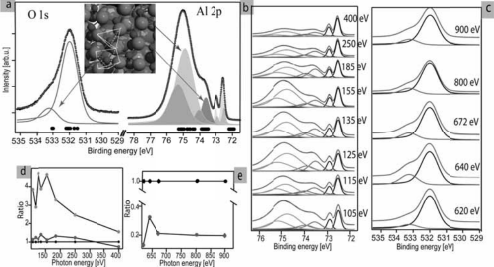


Figure 1: High resolution core level spectra from the ultra thin aluminum oxide on NiAl(110). a) Al 2p spectrum (right) recorded at 130 eV. Four components are clearly visible, each with a 0.4 eV spin-orbit split with the ratio of 1/2. The components can be identified as Al in the NiAl bulk (blue), Al at the interface between the Al oxide film and the NiAl(110) substrate (grey), Al in the Al_i layer (violet) and Al in the Al_s layer (orange). O 1s spectrum (left) recorded at 900 eV. The more intense component originates from the O atoms in the O_i layer and the O atoms in the surface layer which reside very close to another Os atom. The weaker component is due to O atoms in the O_s layer which have the Al_i atoms underneath and do not reside very close to another oxygen surface atom (see inset and text). The calculated core level binding energies are indicated by filled spheres. Energy dependent measurements from b) the Al 2p level and c) the O 1s level. The integrated area from d) the Al 2p components and e) the O 1s components.

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Lowering the Barrier for Graphene Formation by Alloying Transition Metal Carbides

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Graphene growth on SiC has been shown to result in wafer sized single layers, but very high temperatures (>1500 K) and minute process control are required^{i,ii}. On similar systems, such as e.g. single crystal TiC(111), has graphene been shown to grow epitaxially by chemical decomposition of hydrocarbons such as ethyne and ethene at high temperatures of about 1500 K^{iii} , hence they represent potential candidates for the production of high quality graphene. However, the thermodynamic barrier for out-diffusion of carbon to the surface can be lowered if weakly carbide forming elements such as e.g. Ni are alloyed in the TiC^{iv}.

In this project we investigate graphene formation on thin films of ternary alloys of (Ti,Ni)C grown on alumina. Thin films of (Ti,Ni)C carbides were synthesized through non-reactive unbalanced magnetron sputtering from elemental targets of Ti and C and a segmented Ni/C target. The material was grown on single crystal Al_2O_3 (0001) substrates. A homogeneous coating with the composition ($Ti_{0.86}Ni_{0.14}$)C_{0.80} was achieved and no oxygen could be observed within the films except for a thin oxidized surface layer. Coating thickness was controlled by deposition time, chosen so to produce thickest possible coating with retained crystalline quality and found to be 14 nm thick determined by X-ray reflectivity measurements. X-ray diffraction measurements reveal that the sample is a solid solution of Ni in the TiC-lattice, where it substitutes the larger Ti atoms, the lattice parameter is 4.33 Å with a very slight tetragonal distortion and that the growth is epitaxial, the relationship to the substrate is Al_2O_3 (0001)[110] // TiC(111) [10 $\overline{1}$]. The structure of the surfaces and the out-diffusion of carbon from the bulk were investigated at the PEEM/LEEM endstation at Max-lab.

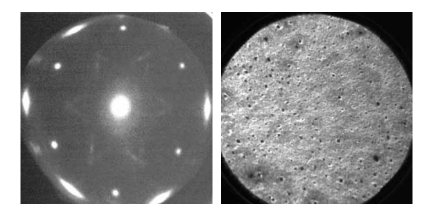


Figure 1: a) LEED pattern (E_p =50eV) and b) LEEM micrograph (25 μ m) from the TiNiC alloy taken after thermal treatment at 1250K.

Upon annealing the sample, profound changes occur in the sample in that the carbon and Ni are diffusing.

The LEED pattern presented in Figure 1a, shows a 1x1 pattern from bulk TiC (inner hexagon) and a star-like pattern which is believed to originate in a surface reconstruction of TiC. Also present is a streaky pattern with a 30° rotational relationship to the substrate crystallographic directions. This we attribute to a combination of carbon on Ni which has segregated from the alloy. The lattice parameter is virtually the same for C(0001) and Ni(111) so they cannot be distinguished. This pattern emerges already at annealing temperatures as low as 975 K. Depending on heating method, there is a risk of depleting the sample of Ni and if that happens, the surface carbon layer will not form. However, Ni is a vital component in order to achieve segregation of carbon to the surface at lower temperatures. Reference samples of TiC_{0.75} made in a similar fashion with the same magnetron sputtering system showed a $\sqrt{3}\times\sqrt{3}R30^{\circ}$ reconstruction upon annealing to 1075 K which was replaced by a disordered 1×1 phase at 1360 K but formation of a surface carbon layer was not observed up to 1400 K, the maximum temperature used. This indicates that the cause for driving carbon to the surface is the Ni incorporation. Figure 2 is a photoelectron spectrum of the sample annealed to 1250K showing that there is indeed a carbon layer at the surface. This corresponds to the signal at 40.5 eV while the shoulder structure at 41 eV is related to amorphous or disordered carbon species at the surface. The spectral feature at 43.5 eV is attributed to carbon atoms in TiC.

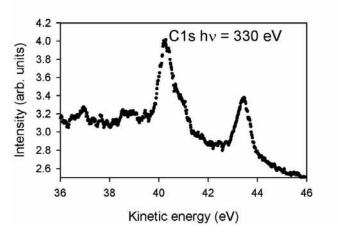


Figure 2:C1s photoelectron spectrum of TiNiC annealed to 1250K.

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Spectroscopic photoemission and low-energy electron microscopy study of nitridation of aluminum nanopatterns

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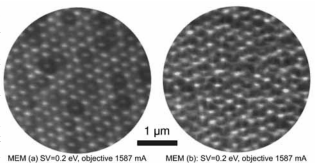
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In recent years, the interest in growth of III-nitrides quantum structure using AlN anostructures as assembling templates by selective area molecular-beam epitaxy technique has been increasing $^{1\text{-}3}$. In this contribution, the spectroscopic photoemission and low-energy electron microscope (SPELEEM) instrument at beamline I311 at MAX II was applied to study the spatial resolved effects of nitridation on the chemical, topographic and crystalline structural property of the aluminum nanopatterns (~100 x100 nm²) fabricated by e-beam lithography. The nitridation was carried out in the preparation chamber with NH $_3$ gas around 5.0 x10 4 Pa through three consequential stages: first at 680 °C for about 1.5 hour, then 730 °C for about 2 hours and finally around 800 °C for 45 minutes. All the SPELEEM measurements were carried out in the analysis chamber which holds the ultra high vacuum conditions.

Fig.1 shows the mirror-mode images (MEM) of the nanopatterns at a field of view (FOV) of $10~\mu m$ (a) before and (b) after nitridation. The contrast in mirror-mode was the contributions of sample surface work function difference and topography. The bright dots arranged in a hexagonal pattern in Fig.1a represent the individual Al nanopatterns, with the gray background representing the native SiO₂ layer on the Si substrate.



inclin (a) 5v-0.2 ev, objective 1557 file

Fig.1 MEM images of the nanopatterns (a) before and (b) after nitridation.

In addition, some dark ring perpetuated nano-dots indicate there exist either some topographic peculiarity⁴ or a sharp work function difference between the nanopatterns and their immediate surroundings. After nitridation, as shown in Fig.1b, the nanopatterned surface appeared more corrugated, with the individual nanopetterns becoming sharpened and previous black rings transformed into black holes or simply disappeared. The results in Fig.1 as a whole indicate that both the work function difference between the nanopatterns and their immediate surroundings was reduced and some perpetual topographic peculiarities of the nanopatterns were diminished by nitridation.

Fig.2 shows the x-ray photo-emission electron microscopy (xPEEM) images of the nanopatterns at a FOV of 20 µm and illumination photon energy of 130 eV (a) before and (b) after nitridation. In xPEEM. both topographic chemical/element contrast be resolved. As shown in Fig.2a, nanopatterns appeared separated, bright parallel lines (owing to an off-normal illumination of the sample surface by x-ray) over the background of native SiO2 layer,

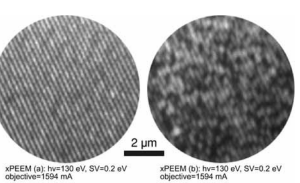


Fig.2 xPEEM images of the nanopatterns (a) before and (b) after nitridation.

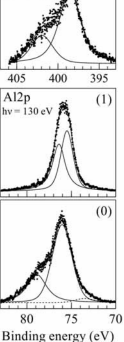
N1s

indicating the Al nanopatterns were geometrically well confined by the naopatterned SiO₂ mask layer. After nitridation, the line patterns were broken into elongated nano-bars or desecrated nano-dots. In consistent with the MEM image in Fig.1b, the surface as a whole appeared more corrugated, the

contrast between the bright nano-bars and the surrounding background was of less regularity⁵, with some black holes/structure vacancies frequently seen among the bright nano-features and gray background. This indicates that the nanopatterns became less regularity after nitridation.

Fig.3 shows the core level Al2p and N1s photo-emission spectroscopy (PES) of the nanopatterns before (0) and after (1) nitridation. The results clearly show that two major Al compounds presented in the nanopatterns before nitridation, with Al2p peak at binding energy (BE) of 75.9 eV representing AlO compound and the Al2p peak at BE of 78.8 eV representing AlF compound. After nitridation, the AlF compound disappeared, the AlO compound was converted to a mixture of AlN (Al2p BE 75.4 eV) and AlON (Al2p BE 76.3 eV) compound.

Fig. 4 shows the low-energy electron microscope (LEEM) image of the nitridated nanopatterns at FOV of 20 um and electron energy of 1.9 eV. In LEEM mode the incoming electrons experience diffraction (if the surface is ordered) and based on the intensity distribution of (0,0) diffracted spot the LEEM image is conformed. In LEEM differently ordered areas have different I (V) curves and by tunning SV, the optimized contrast between the differently ordered structures can be obtained. The bright nano-dots in LEEM image in Fig.4a indicates that surfaces of the majority nitridated nanopatterns were ordered. The μ-LEED pattern in Fig.4b recorded from the probing area of 0.4 µm (approximately 1-2 nanopatterns) at electron energy of 7.9 eV suggests that the surface of nitridated nanopatterns was structurally ordered, but consisting of very small (less then 100 nm) grains Fig.3 Core level Al2p and N1s differently oriented.



(1)

PES spectra of nanopattems

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Investigation of TiO_x phases grown on Au(111) by chemical vapor deposition

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Titania on gold is a widely studied system for its interesting properties, e.g. as an inverse catalyst¹. We are currently investigating titania loading of nanoporous gold to enhance and modify its catalytic properties². For this purpose chemical vapor deposition (CVD) is a promising technique because the gaseous precursor can efficiently penetrate the pores. In parallel, we are conducting studies of titania deposition on Au(111) by way of CVD to gain a deeper understanding of the properties of that system. As it turns out, titania CVD on Au(111) is an interesting system in its own right.

We have begun to investigate CVD of titania on Au(111). The experiments have been carried out at I311. In the first attempt, we discovered that it is possible to form an atomically ordered structure with (2x2)periodicity vs. the substrate as witnessed by LEED. Furthermore, Xray photoemission spectroscopy (XPS) showed a component at a binding energy (BE) of 459.0 eV indicative of fully oxidized Ti (Ti⁴⁺) along with (at least) two additional components at lower binding energies, assigned to reduced forms of Ti. The results thus suggest an inhomogeneous situation.

The studies were continued in our home laboratories in Uppsala, using Scanning Tunneling Microscopy (STM), LEED and XPS. It soon stood clear that also other ordered structures were possible to form. Our aim was to characterize the phases in more detail

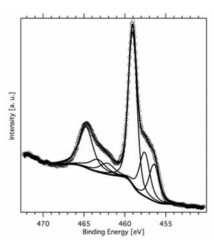


Figure 1. Ti 2p spectrum for TiOx on Au(111).

at I311 in November last year. However, photoemission was not possible as MAX was "down" during this period. Still, we could use Low-Energy Electron Microscopy (LEEM) and μ -LEED, which proved to be extremely informative and helpful to understand the nature of the TiO_x phases.

At present we have been able to pinpoint and characterize at least 3 phases:

- 2x2: a monolayer thick phase with a honeycomb structure. Its lattice parameter is twice the one of Au(111);
- wagon-wheel: the overlayer has a coincidence relationship with the substrate. This gives rise to a

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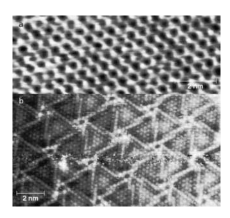


Figure 2. High resolution STM pictures of (a) the 2x2 phase and (b) the wagonwheel phase.

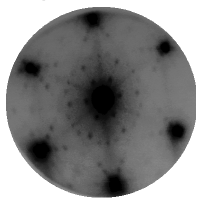


Figure 3. μ -LEED pattern of the "star" phase.

- Moiré pattern that resembles a wagon wheel;
- "star": most likely made of three-dimensional islands with a rectangular unit cell. Its LEED pattern looks like a David's star, therefore the name. Its precise nature is not fully understood yet.

Ongoing and future work aims at a detailed description on how to prepare the various phases. Of importance with respect to studies at I311 is to optimize the growth parameters in order to obtain larger islands (at least 100 nm x 100 nm) to be able to use LEEM to investigate the growth of the different phases.

Once the nature of these ultrathin phases will be clarified we will study their interaction with small "probe" molecules (e.g. CO, CH₃OH) and the CVD growth process.

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Cleaning and oxidation of nanoporous gold. A detailed photoemission study.

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Nanoporous gold (np-Au) is a fascinating class of bulknanostructured material. It is made up of a threedimensional mesh of ligaments which can be between some tens of a nanometer up to several microns depending on the preparation conditions chosen¹. This material is coined by its extended metallic surface and its high surface to volume ratio. Many of the unique properties of this material are a direct consequence of its sponge like morphology making its entire surface accessible to fluids and gases.

In recent years several applications of this material in optics, sensors, and heterogeneous catalysis have been deployed and only a few studies are cited in the following. The key to many of these applications is the extended metallic surface being reversibly oxidized by molecular oxygen or ozone. As for example the oxidation of the surface by ozone was shown to alter the macroscopic dimensions of np-Au². Recent studies covering the thermal stability of np-Au showed that surface oxygen plays a crucial role in stabilizing its structure and prevent coarsening³. Besides its impact on the mechanical properties of np-Au surface bonded oxygen can also be used as a reactant in heterogeneous catalysis⁴⁻⁵. As for example np-Au was shown to be an excellent catalyst for low temperature oxidation of CO and highly selective oxidation of alcohols such as methanol4,6-7

Though all these applications are strongly linked to the reversible oxidation of the np-Au surface the picture derived so far remains elusive. Silver which is still contained in the material of preparation seems to play an important role especially when using molecular oxygen as a source for surface oxygen. 8-9 Further on, different oxide species present at the beginning of oxide formation as for higher coverages have been revealed in surface science studies of gold single crystals. However, a detailed chemical characterization of possible oxide species present at the np-Au surface is missing.

The aim of the studies conducted at beamlines I411 and I311 was to investigate in detail a typical np-Au sample after dealloying by free corrosion in nitric acid. Additionally another sample was subjected to heat treatment and a cleaning procedure employing an O_2/O_3 gas stream admitted through a gas doser [~7-10 vol% O_3 , $2x10^{-7}$ mbar background pressure].

For the sample measured at I411 two weeks after dealloying in nitric acid (pristine sample) the existence of a "native" oxidic gold species could be proven for the first time (fig. 1a). Moreover, a detailed spectrum of the

carbon 1s region (not shown) revealed the presence of different oxidized (hydro-) carbon species, containing hydroxyls and carbonates amongst others. Heating such a sample to 140 °C for 19h decomposed the oxidic gold species and lead to desorption of the oxidized carbon species. A broadening of the Au 4f signal to lower BE values indicates a slightly higher amount of free lowcoordinated Au atoms (fig. 1b). However, a significant amount of carbon was still detected. In an attempt to clean the surface ozone treatment was applied. Indeed the carbon impurities at the surface could be removed. Additionally the gold surface was oxidized, as is evidenced in fig 1c). Interestingly the nature of the oxide generated by ozone treatment seems to be different from the initially observed oxidic gold since the BE of the oxide signal in the Au 4f spectrum appears to be 0.41 eV lower. A further broadening towards lower BE values is a result of further cleaning and exposition of undercoordinated Au atoms from terraces and steps (kinks). A new signal appears at even lower BE values in the Au4f spectrum. Since the low BE regionhas not been investigated with high resolution so far no report of that signal is found in literature.

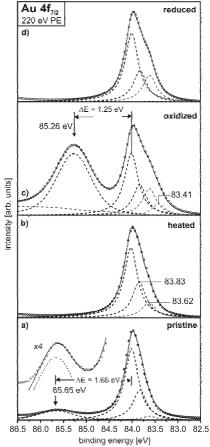


Fig. 2: Au4f_{7/2} core level spectra a) pristine sample b) after heating c) after oxidation by ozone d) after three days in UHV

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It is likely that the signal is caused by Au atoms at the interface to a 2D gold surface oxide probably in similarity to Rh(111) substrates¹⁰. More detailed considerations are under way to substantiate this conclusion.

The observed oxide is not stable in UHV for a longer period of time as was revealed after measuring the Au4f region again after storing the sample three days in UHV (fig. 1d). The oxide signals has disappeared as has the low BE signal, providing further evidence that those two signals are related. Additional insight into the nature of the gold oxide is provided by oxygen 1s core level spectra. At first, a spectrum from the pristine sample was recorded as shown in fig. 2a. At least two contributions can be discerned, which most likely are caused by oxidized carbon and hydrocarbon species (around 531.75 eV) and hydroxylated components. However, no significant intensity of a signal corresponding to the "native" gold oxide observed in the Au 4f spectrum is present. Such a signal would be expected around 530 eV BE and below.

The oxygen spectrum of the heated sample (2b) also consists of two contributions but a shift to lower BE (530.69 and 532.97 eV) occurred compared to the spectrum of the pristine sample (2a). If now the ozone cleaning procedure described above is applied to the heated sample spectrum 2c) results. The signal at highest BE has disappeared and only a small contribution at 530.6 eV is left. The more striking change is the appearance of two new and sharp components at 529.08 eV and 529.94 eV, respectively. In literature three different oxygen species on single crystal gold existing after different oxidation treatments have been suggested: a mobile chemisorbed oxygen species without long range order, a surface oxide forming ordered two-dimensional structures and a bulk oxide forming crystallites with three-dimensional networks¹¹. In our case we can identify the low BE signal as caused from chemisorbed oxygen and the signal at 529.94 eV as caused by oxygen atoms bond in a surface oxide structure. By heating annealing experiments (not shown) we got strong indication that there is an equilibrium between chemisorbed oxygen and the surface oxide, and that the rate limiting step for oxygen desorption is the transformation from surface oxide to chemisorbed species which then desorb. This behaviour has been suggested for single crystal surfaces earlier.

Finally, as mentioned in the introduction the np-Au structures contain roughly 1 atm% of residual silver. At the surface, this content is higher, up to ~10%, as has been revealed by XPS in the past⁶ Upon the ozone cleaning procedure the silver is oxidized as well and enriched at the surface while the silver concentration decreases again upon reduction (heat + CO treatment and UHV storage) as shown in fig. 3. This can have implication on catalytic reactions during which atomic oxygen can be present since residual Ag is supposed to have important influence on the overall catalytic behaviour of np-Au.

Further combined studies on the pre-cleaning and modification of np-Au in UHV and ex-vacuo are in progress.

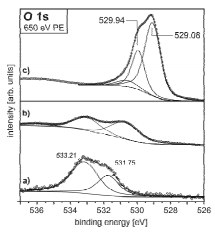


Fig. 2: O Is core level spectra a) pristine sample b) after heating c) after oxidation by ozone

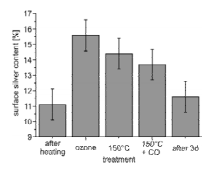


Fig. 3: Silver content at the np-Au surface as obtained from curve fitting of the Ag3dAu4d region

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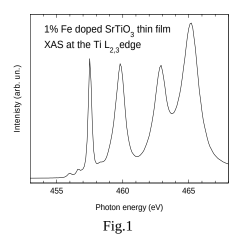
Resonant photoemission study of functional materials for terabit resistive switching memories

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A large class of binary and ternary oxides shows the resistive switching effect. By applying an appropriate threshold voltage one can reversibly switch between low and high resistance states. Doped ABO₃-perovskites are suitable functional materials for storage devices with a density in the terabit range since extended defects such as dislocations or defect clusters with nanoscale dimensions are considered to be the single resistive switching units [1,2]. One of the main goals of the project was to recognize the nature of conductance within the filaments and processes accompanying electroforming and switching for the samples of thin films of Fe doped SrTiO3 deposited on the Nb doped SrTiO3 single crystalline substrate. The resonant photoemission studies performed at the I311 beamline were aimed at clarifying the nature and origin of the states in the electronic gap which can be related with the well conducting filaments. Three samples were studied - epitaxial thin films of SrTiO3 doped with 1 % Fe, 2 % Fe and the undoped SrTiO3 film. The samples were annealed in UHV prior to studies. XAS spectra were obtained for all samples at the Ti, O and Fe low energy thresholds.



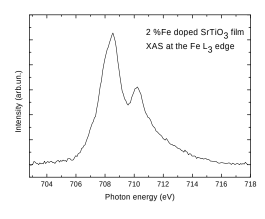


Fig. 2

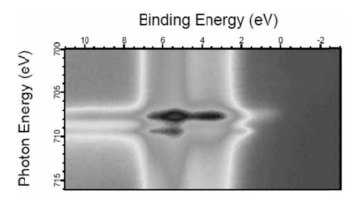


Fig. 3 Photoelectron intensity map of the valence band for the 2 % Fe doped SrTiO₃ thin film in the photon energy range of the Fe 2p-3d resonance.

Fig. 1 shows the spectrum obtained for the 1% Fe doped film. The shape of the spectrum is the same as reported for the bulk SrTiO₃. The XAS spectrum at the Fe L3-edge (Fig. 2) differs from the known Fe containing perovskites. The binding energy versus photon energy photoemission intensity maps were obtained for the interesting resonance regions. Fig. 3 shows the valence band region for the 2% Fe film in the photon energy range of the Fe 2p-3d resonance. A clear enhancement of photoemission is visible in the main part of the valence band – at energies of about 4 eV and 6 eV. Some enhancement of the intensity in the energy gap was also detected.

We found clear contributions from Ti, Fe and O states to the various regions of the valence band. However, the effect is different to that described recently for Nb doped SrTiO₃ [4]. The results will be confronted with the calculations (partial densities of states) of the electronic structure. The knowledge about the Fe contribution to the valence band should enable us to elucidate the role of that dopant in resisitive switching behaviour.

For the 2 % Fe sample we performed electroformation in the preparation chamber with the use of a voltage applied to a retractable Al foil electrode. We could observe increasing current through the sample indicating the proper formation process. The selected spectra were obtained from the electroformed sample. They exhibited very slight but clear variation of the valence states caused by the electroformation process.

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Interface composition of atomic layer deposited HfO₂ and Al₂O₃ thin films on InAs studied by XPS

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Thin high- κ oxide films on InAs, formed by atomic layer deposition (ALD), are the key to achieve high-speed metal-oxide-semiconductor devices [1]. Thereby the quality of the interface between the semiconductor and the high- κ material is crucial for the device performance for bulk InAs substrates [2] as well as for InAs nanowires [3]. X-ray photoemission spectroscopy (XPS) is a well-suited tool to investigate the chemical composition of the interface between III-V semiconductors and high- κ thin films, as has been shown in many cases for GaAs and InGaAs [4]. However, although InAs has an even higher electron mobility and is an important binary reference material, only a few XPS studies on high- κ thin films on InAs(100) substrates have been reported yet [5].

We have performed a detailed XPS investigation on the interface between InAs and 2-nm-thick ${\rm Al_2O_3}$ or ${\rm HfO_2}$ layers, deposited by ALD at different temperatures, for InAs substrates with different surface orientations as well as for InAs nanowires with different crystal structure. From XPS core-level spectra we can reveal the composition of the native oxide and obtain how the high- κ layer deposition reduces the different oxide components.

Planar InAs(100) substrates have been prepared by a heat treatment in an As-rich environment to remove the epi-ready layer, followed by 30 s dip in HF and rinse in isopropanol. Al_2O_3 and HfO_2 layers were deposited in a Cambridge NanoTech Savannah 100 ALD reactor, except for reference samples containing only a native oxide layer. XPS was performed at beamline I311, taking In 3d, In 4d, As 3d, O 1s, Hf 4f, and Al 2p core-level spectra at photon energies between 70 and 1330 eV. Spectra from several samples and various photon energies were fitted in a self-consistent way with the FITXPS package assuming a Doniach-Sunjic line form, with obtained peak parameters agreeing nicely with literature data [5].

Figures 1 and 2 show As 3d and In 3d spectra of an InAs reference sample compared with InAs/Al₂O₃ and InAs/HfO₂ samples, demonstrating the strong oxide reduction. Two arsenic oxidation states (As+3 and As+5) can be distinguished, and their peaks in both high- κ samples have typically less than 10% the size of the corresponding peak in the reference sample. The In-oxide peak is only reduced to about half of its reference size in InAs/HfO₂ samples of various depositions temperatures [Fig. 2(b)]. The

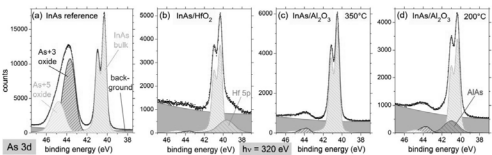


Figure 1: Fitted As 3d spectra of (a) an InAs reference, (b) an InAs/HfO₂, and (c, d) two InAs/Al₂O₃ samples with different deposition temperatures. In (b), a constant part of the actually larger background is subtracted for clarity.

same is valid for $InAs/Al_2O_3$ samples with an ALD temperature of 350°C [Fig. 2(c)], while for temperatures of 250°C or lower an In-oxide reduction down to 10% of the reference peak size is achieved [Fig. 2(d)]. In this case, however, the corresponding As 3d spectrum shows evidence of a thin AlAs film at the interface [Fig. 1(d)]. By simulating the relative oxide

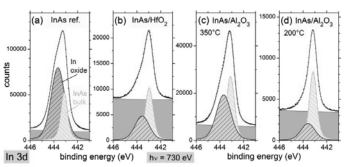


Figure 2: Fitted In 3d spectra of (a) an InAs reference, (b) an InAs/ HfO₂, and (c, d) two InAs/Al₂O₃ samples with different deposition temperatures.

and bulk peak sizes for different samples and photon energies, the thickness of the native oxide film on the reference samples is obtained, which can then be compared with the reduced thickness of the oxide interface, as it is shown in Fig. 3. It should be noted that the native oxide film is not completely homogeneous, but that some sub-Oxide components seem to be located closer to the InAs material and others closer to the surface, as is discussed in more detail in reference [6].

The effectiveness of the oxide reduction upon high-κ deposition depends on the substrate surface orientation: We have compared InAs(100), InAs(111)A, and InAs(111)B samples with and without Al₂O₃ or HfO₂ layers. InAs reference samples have the largest oxide peak for the (100) surface and smaller peaks for (111) surfaces, with more In-oxide at the In-rich (111)A surface and more As-oxide

at (111)B. The strongest oxide reduction upon deposition of HfO_2 or Al_2O_3 is obtained for (100) samples, while both In- and As-oxides are less effectively reduced on (111)A or (111)B surfaces.

Beside planar InAs samples, even the interfaces of HfO_2 and Al_2O_3 layers on InAs nanowires were studied. Typically, nanowire samples exhibit larger oxide peaks than the corresponding bulk samples, which can only partly be explained by the different geometry of planar surfaces and nanowires in the XPS experiment. Importantly, also for the InAs nanowires, a significant reduction of the native oxide upon deposition of HfO_2 and especially Al_2O_3 is obtained.

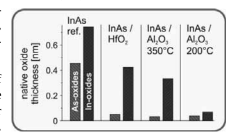


Figure 3: Average thickness of the surface or interface oxide layer, respectively, of InAs(100) reference samples, InAs/HfO₂ samples, and InAs/Al₂O₃ samples with different deposition temperatures.

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Charge transfer dynamics of several Ru(II) complexes which could form the charge transfer centre of a multi-centre water splitting dye complex

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a) O
$$\stackrel{\text{OH}}{\longrightarrow}$$
 N $\stackrel{\text{C}}{\longrightarrow}$ N

Dye sensitized solar cells (DSCs) are a cost effective method of generating renewable energy, using DSC architecture it may be possible to use energy from sunlight to perform chemical reactions. One of these reactions is the water splitting reaction which produces hydrogen and oxygen molecules from water molecules. By using a DSC to split water molecules we can produce hydrogen fuel using low cost materials and a free source of energy. Past studies have shown that a multi-centre water splitting dye complex is more effective than a single-centre water splitting dye complex.[1] The dye complexes studied in this investigation are non-volatile and cannot withstand thermal evaporation, so in situ UHV electrospray deposition was used in order to study the molecules with high resolution synchrotron-based spectroscopy. The deposition solution is passed through a high voltage capillary, the electric field causes the formation of charged droplets. These droplets undergo a succession of Coulomb fission events and solvent evaporation to leave a beam of desolvated molecular ions which are transported through 4 stages of differential pumping before arriving at the sample surface at pressures in the 10⁻⁸ mbar range [2-4].

In this experiment we successfully deposited the Ru 535 (N3), Ru 455 and Ru 470 dye complexes shown in figure 1 onto atomically clean $TiO_2(110)$ surfaces under UHV conditions at beamline I311. These dye complexes were chosen due to their potential suitability as the basis for a charge transfer centre of a multi-centre water splitting dye complex. The adsorption geometry and charge transfer timescales of the molecules were investigated using core-level photoemission and the core-hole clock implementation of resonant photoemission spectroscopy. The molecules were deposited using a commercial UHV-compatible electrospray deposition source (Molecularspray Ltd, UK) [5] from a 3:1 methanol:water solution at an emitter voltage of +2kV and a flow rate of 500 nlmin⁻¹. The pressure during the deposition was $\sim 5 \times 10^{-7}$ torr.

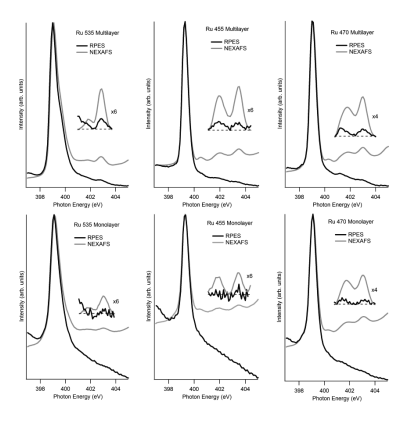


Figure 2. N *1s* RPES and N *1s* NEXAFS spectra of multilayer (top) and monolayer (bottom) coverages of Ru 535 (left), Ru 455 (middle) and Ru 470 (right).

Figure 2 shows the N *Is* NEXAFS and N *Is* RPES spectra of each dye complex at monolayer and multilayer coverages. Energy alignments revealed that the LUMO of each molecule is incapable of charge transfer, whilst other unoccupied orbitals are capable. There is a reduction in intensity of the monolayer RPES peaks corresponding to orbitals which are capable of charge transfer, this effect can be attributed to charge transfer from the adsorbed molecules to the substrate.[6] The NEXAFS peaks are used for normalizing the intensity of the RPES peaks. Charge transfer timescales were computed for the LUMO+2 and LUMO+3 of each dye complex where possible, which reveal that each complex has an orbital capable of charge transfer on the 12-21fs timescale. The Ru 470 complex appears to show the slowest charge transfer timescale, this information gives insight into the charge transfer process.

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Towards graphene on Fe(110)

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One of the most serious technological challenges on the way towards graphene-based electronics is the process of inexpensive fabricating the large scale areas of high-quality graphene. Both monolayer and multilayer graphene can be grown by thermal decomposing hydrocarbons on hot d-metal surfaces like Ni(111) or Co(0001), and even on polycrystalline foils. In the context of high technological relevance of inexpensive routes towards large-scale graphene it is highly attractive to try graphene synthesis on iron. It has been shown recently that hexagonal BN can grow as a single monolayer on Fe(110) [1]. A close structural similarity between graphene and h-BN allows one to expect a similar (graphitic) growth mode for carbon at certain conditions, although a very rich C/Fe phase diagram is a challenge here, as it implies an easy formation of different carbides. Here we present first results showing the crystalline structure of carbon films formed on Fe(110), as obtained by micro-LEED and LEEM.

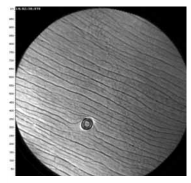


Figure 1. LEEM image of Fe/W(110) sample demonstrating high quality of the epitaxial iron film. The field of view is 15 μ m.



Figure 2. A LEED pattern from the C/Fe(110) sample. E_p =102 eV.

All measurements were performed *in situ* at the beamline I311, MAX-Lab, on the electron microscopy end station. The Fe(110) substrate was prepared by epitaxial growth of iron films (thickness ca. 20 nm) on the W(110) surface. The W single crystal was cleaned by several cycles of Ar^+ -sputtering, annealing in oxygen atmosphere and subsequent flashing in UHV. The cleanliness of the substrate was checked by LEED. After iron deposition the sample was gently annealed to improve the quality of the resultant Fe(110) surface, which was controlled by LEED [2,3] and LEEM. Carbon films were grown by thermal cracking of vaporized propene (C_3H_6) in contact with the hot metal surface at the temperature of $^{\sim}630^{\circ}\text{C}$ in the propene partial pressure of 5×10^{-8} mbar, the quality of the sample was verified by the homogeneity of the LEED pattern over the sample surface.

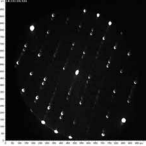
Figure 1 shows a typical LEEM image of the Fe/W(110) surface ensuring the high structural quality of the iron film. This can be deduced from the visibility of atomic steps of iron film imaged as horizontal stripes, as well as from a low amount of randomly scattered electrons which typically create a strong background.

Figure 2 shows the LEED pattern of the C/Fe(110) surface. Analogically to the hBN/Fe(110) interface [1], a number of spots attributed to the similar moirè surface structure is observed. However, the LEED pattern of C/Fe(110) is much more complex than that of hBN/Fe(110). We attribute it to the poor quality of the sample probably due to both too high temperature of graphene growth which led to formation of multiple domains of carbidic C/Fe phase with random orientations instead of graphene and to

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the stress in the iron film due to the noticeable mismatch between Fe(110) and W(110) lattices. Nevertheless primary orientations of carbidic structures are still defined by the crystallographic directions of lattice match between graphene and Fe(110) as in the case of hBN/Fe(110).



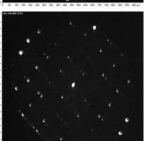
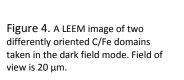
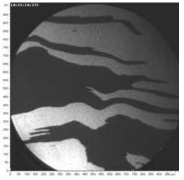


Figure 3. µLEED patterns from two different domain orientations of the C/Fe(110) sample.

Figure 3 shows $\mu LEED$ patterns from different domain areas of the surface. Since the illumination aperture size is 10 μm and no spots of differently oriented domains could be detected on the successive $\mu LEED$ patterns, the size of a single domain can be estimated as at least 10 μm . As can be seen from the LEEM image in the mirror mode (not shown) C/Fe domains with single orientation tend to occupy one or more terraces of Fe(110) surface, however boundaries of these domains perfectly match to the step edges.

Figure 4 shows a typical dark field mode image of domains with two different orientations. These domains tend to spread along the atomic steps on the iron surface. The LEEM image shows a rather high crystallographic quality of the C/Fe phase; however LEED and μ LEED patterns suggest that the sample is rather iron carbide than graphene on iron. More studies are planned for the determination of exact structure of carbon adsorbed on Fe(110), as well as for revealing the prospects for the growth of real *graphene* on Fe.





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Constant energies imaging of π -cones from a buffer layer and after intercalation

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An X-ray Photoelectron emission microscope (XPEEM) equipped with a hemispherical energy analyzer is capable of fast acquisition of momentum resolved photoelectron angular distribution patterns in a complete cone. We applied this technique to observe the 3D (E,k_x,k_y) electronic band structure from 0 and 1 ML graphene samples on SiC(0001) before and after Li intercalation [1]. For 0 ML samples no π -cones (bands) were detected close to the Fermi level, see Figs. 1 (a)-(b), while after Li deposition and heating clearly resolved π -cones appear at the K-points in the BZ, see Figs. 1 (c)-(d). The 0 ML graphene sample less wrinkles/cracks were observed to form on the surface after Li deposition and heating compared to on 1 ML samples. The observed deterioration in quality of the graphene upon Li intercalation may be connected to an explanation why the efficiency in fuel cell reduces with usage.

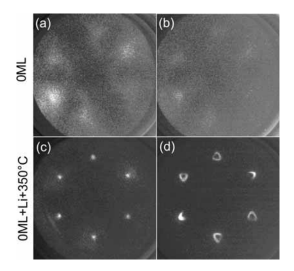


Fig. 1. Photoelectron angular distribution patterns (E,k_x,k_y) collected from 0 ML graphene at an initial state energy of (a) 1.2 eV and (b) 0.2 eV below the Fermi level, using a photon energy of 35 eV. (c)-(d) Patterns collected from the 0 ML graphene sample after Li deposition and heating at 350°C at the same initial state energies as in (a)-(b), respectively.

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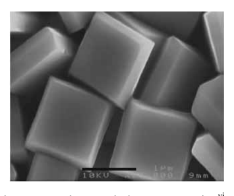
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Micro LEED on Single Micro Crystal of Anatase TiO₂

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Shape-controlled nanostructures are of great interest due to their often new physicochemical properties i,ii,iii. Among metal oxides, anatase TiO₂ is of special importance in the field of energy iv and catalysis due to their outstanding photoelectrochemical properties. Theoretical calculations predict that the (001) surface should be the most active one because of the low atomic coordination numbers and the wide Ti-O-Ti bond angle. However, it is thermodynamically less stable than (101) which naturally dominates over (001). Recently Lu's group has successfully developed a method to synthesize nanoplates with up to 64% (001) plane, using fluorine to stabilize this surface vi,vii,viii.

The fluorine-based synthesis utilizes the fact that F effectively lowers the surface energy making (001) and (101) growth rates comparable. In addition, on the (001) surface a (1x4) reconstruction lowers the surface compared to pristine (1x1) surfaces 1x. clear However, it is not reconstruction is created at high F concentrations. Theoretically it is predicted that formation of O-F and Ti-F bond may



pin the atomic movement on the surface and consequently quench the reconstruction vi . The completely removal of F can be realized by thermal treatment at 600 o C for 90 min. Whether desorption of F facilitates the (1x4) reconstruction or not will be a significant evidence to justify the theoretical interpretation.

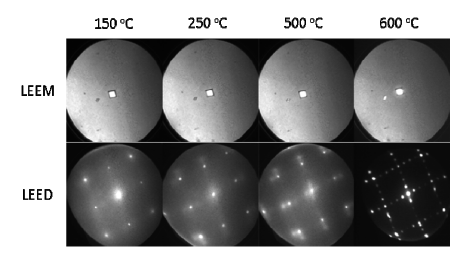
The TiO_2 microcrystals were suspended in an aqueous solution only by deionized water to avoid further organic contamination, dipped by droplets on natural silica covered Si(100) substrates and dried in air. After being transferred into the UHV system, the crystal was first outgassed at 150 °C for around 15 min. Micro LEED pattern was taken on the same crystal at round 60 °C after each 5 min annealing. For 150 °C, 250 °C and 500 °C, the start voltage is around 48 V; for 600 °C, the start voltage is around 28 V.

After outgassing, LEED showed a clear 1×1 pattern, which is bulk related. By increasing the annealing temperature, surface related spots become clearer. However,

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until 500 °C, no well resolved surface pattern can be resolved. A greatly improved LEED pattern was observed after 600 °C annealing for 5 min when a sharp two domain (1×4/4×1) pattern was collected. However, even though the LEED pattern evolution matches the theoretical prediction and other experimental evidence that fluorine desorption might induce a 1×4 recontruction, other aspects should also be taken into account. For example, surface contaminations due to the ex-situ preparation method. In that case, the contamination covered surface will either modify the surface structure or simply block the signal from the surface, leaving only the bulk contribution. This type of influence should be minimized in the future by improving sample preparation condition.



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Nano-scale 3D(E,kx,ky) band structure imaging by XPEEM

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Mapping of the electron band structure or the Fermi surface of a solid using angle resolved photoelectron spectroscopy is commonly quite tedious and time consuming. The technique requires the mechanical scanning of the sample at least in one direction and many photoelectron spectra have typically to be collected in order to fully determine the energy and momentum of the electron states in the Brillouin Zone. Here we show that an energy filtered X-ray Photoelectron emission microscope (XPEEM) is capable of fast acquisition of momentum resolved photoelectron angular distribution patterns. By imaging the diffraction plane of the objective lens¹ an energy slice of the photoelectron angular distribution pattern from a very small (<1µm) area on the surface can be collected in a complete cone in one shot and projected on the screen. To restrict the aberrations we use an extremely small selected area aperture and probe a sub-micrometer (800nm) area of the sample. We apply this technique to observe the 3D $(E_x k_x, k_y)$ electronic band structure of 0, 1 and 2 monolayer graphene grown ex-situ on 6H-SiC(0001) substrates where a carbon buffer layer (0ML) forms underneath the graphene layer(s). Fig.1a shows a typical LEEM image from such a surface. Diffraction patterns (micro-LEED) originating from 1 and 2ML areas (Fig.1b,d, correspondingly) show distinct decrease in spot intensities around the main graphene spots for a 2ML island. Area selected azimuthal distributions of the photoelectron intensity can be recorded by placing the selected area aperture (SAA) either on the 1ML surface or a 2ML island. In Fig.1c,e two-dimensional momentum resolved photoelectron angular distribution patterns shown at a 2.5eV binding energy from a small (800nm) area for 1ML (Fig.1c) and 2ML(Fig.1e) island. For 2ML island the second π -band in the electronic band structure becomes clearly observable (Fig.1e). In Fig.2 two dimensional (k_x,k_y) photoelectron angular distributions are shown as a function of electron energy down to 4eV below the Fermi level. The shape of the well known graphene band structure with the Dirac points approximately 0.4eV below the Fermi level is clearly reproduced. The results presented demonstrate the uniqueness and the versatility of high energy and momentum resolved XPEEM and broadens the applicability and impact of this technique within surface science and nanotechnology.

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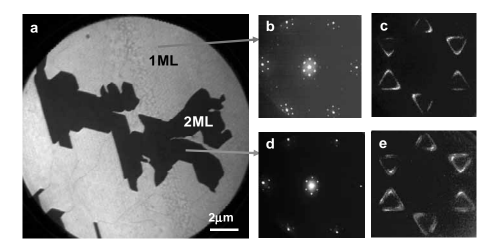


Fig.1. a): LEEM image from an ex-situ grown sample of 1ML graphene with 2ML islands; electron energy is 4.6eV. b) and d) Electron diffraction: μ -LEED images (400nm sampling area) acquired at 60eV electron energy from 1ML (b) and 2ML (d) areas. c) and e) Photoelectron diffraction: high energy resolution 2D (k_x , k_y) maps for 1ML (c) and 2ML (e) graphene taken at an electron binding energy of 2.5eV below the Fermi level. Two π bands are clearly seen from the 2ML island. Photon energy is 35eV.

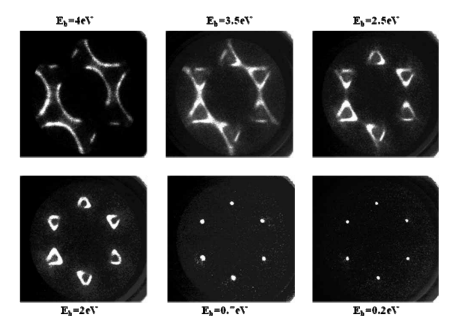


Fig.2. Energy resolved 2D (k_x,k_y) maps for 1ML graphene given by the angular distribution of photoelectrons. The binding energy is indicated at each image. The images are collected from an area of 800nm diameter on the surface.

Photoelectron spectroscopy studies of the 5s and 4d bands in free silver clusters

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We have probed the 5s and 4d valence bands of free Ag clusters using photoelectron spectroscopy. The clusters were produced in two different ways, 1) with a gas aggregation source[1] where silver was first vaporized using magnetron-based sputtering[2], and the vapour atoms were then condensed to clusters in a liquid nitrogen-cooled cryostat, and 2) with a pick-up source[3] (a.k.a. EXMEC source) where argon clusters were sent through silver vapour created in an oven with induction heating. Photoelectron spectra recorded with 40 and 60.5 eV photon energies are presented in figure 1 (gas aggregation source) and 2 (pick-up source) below.

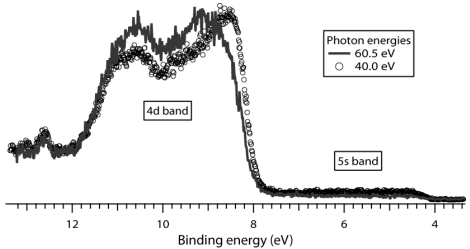


Figure 1. Photoelectron spectra of the 5s and 4d valence bands in free silver clusters, produced by a gas aggregation source, recorded with 40 and 60.5 eV photon energies. Absolute calibration was done using the water peak at 12.62 eV binding energy.

Apart from the higher intensity of the 5s band for the 40 eV spectrum, one can also see differences in the 4d feature between the two spectra. The relative intensity between the two maxima is clearly different, and also the lower-energy maximum is shifted towards lower binding energies in the 40 eV spectrum by more than half an eV. With the gas aggregation source, we typically produce clusters with a number of constituent atoms at least in the $\sim 10^3$ range. As the cluster on-set of the 5s band is indistinguishably close to the solid state on-set, the clusters are probably quite large (possibly in the $\sim 10^5$ range).

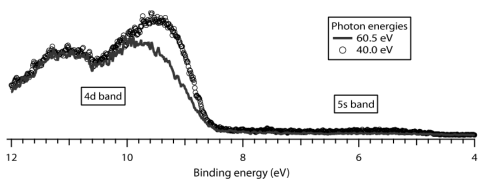


Figure 2. Photoelectron spectra of the 5s and 4d valence bands in free silver clusters, produced by a pick-up source, recorded with 40 and 60.5 eV photon energies. Absolute calibration was done using the argon 3p peaks.

Also in this case the 5s signal is stronger for the 40 eV photon energy. The difference in the 4d feature is also similar to the case of larger clusters, the relative intensity between the two maxima is clearly different, and also the lower-energy maximum is shifted towards lower binding energies in the 40 eV spectrum, but this time by less than half an eV. From the difference of the 5s band on-sets for the clusters and solids, we can estimate the clusters to have a mean radius of ~1 nm and to contain a few hundred atoms.

A noticeable dependence of the spectral shape on the photon energy has been observed also for free copper clusters containing $\sim 10^3$ - 10^4 atoms [4], where the lower binding energy part of the valence density of states was gaining in relative intensity with the photon energy increase from 60 eV up to 130 eV. In ref. 4 this was, at least partially, assigned to the photon energy dependence of the electron escape depth, facilitating the response of the bulk density of states at certain photon energies.

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Auger parameter in free nanoscale metal clusters: Does the old approach hold?

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In early days of electron spectroscopy C.D.Wagner proposed a method to characterize the changes in the chemical environment of a certain element in different compounds using the so-called Auger parameter α [$^{1.2}$]. This magnitude is calculated as a sum of the core-electron binding energy E_b and of the subsequently emitted Auger-electron kinetic energy E_k : $\alpha = E_b + E_k$. The advantage of such a characteristic is that the Auger-parameter value is free of the energy-reference problem - in contrast to either core-ionization energy or Auger energy treated separately. In numerous studies it has been shown that for the same core levels in different compounds the largest α has been the case for the free-electron substances - metals, and the smallest α -in the compounds where the screening of the core-vacancy should be the weakest. Within reasonable assumptions based on electrostatics the difference $\Delta \alpha$ between the Auger parameters for two chemical states, a free atom and a solid compound, gives the extra-atomic (ea) relaxation/screening energy[1] in the corresponding solid: $\Delta \alpha = 2R^{ea}$. In the present work the authors have attempted to consider the Auger parameter method for free metallic nanoparticels/clusters, at the dimension scale when the number of constituent atoms (and thus of free electrons) is finite and relatively small.

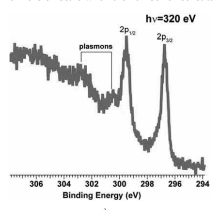


Fig.1 K 2p cluster (<N> \sim 10³) spectrum recorded at 320 eV photon energy. K 2 $p_{1/2}$ "line" overlaps with a plasmon-loss feature due to the kinetic energy loss of the 2 $p_{3/2}$ photoelectrons

For such particles with a countable number of free electrons the task has been to determine the Auger parameter, and to analyze the consistency of the experimental cluster Auger spectrum and the expected extra-atomic relaxation energy extracted from the so-called XPS-shift for free clusters and from the Auger parameter difference $\Delta \alpha_{cl-at}$. Experimentally the task has been realized in a study on a beam of unsupported potassium clusters containing several thousand atoms, by measuring binding energies for two core levels -2p and 3p - and recording the corresponding Auger spectrum due to the transition between them. Using a beam of free nanoparticles allows avoiding uncertainties introduced by the substrate, the interaction with which changes nanoparticle geometry and energy-levels.

The so-called gas-aggregation cluster source [3] creating a cluster beam propagating in vacuum, has been attached to the permanent experimental station of the I411 beamline. The x-ray photoelectron spectra and Auger spectra have been recorded with an electrostatic energy analyzer Scienta R4000 placed at 90 degrees to the horizontal polarization plane of the beamline radiation. Photon energies have been chosen to combine the

highest possible ionization cross-sections of the 2p and 3p potassium core-levels under investigation and the more intense regions of the beamline radiation: 320-330 eV for the K 2p and 100 eV for the K 3p. In the clusters created in the present work the K 2p response with two main "lines" centered at \approx 296.6 eV and 299.3 eV (Fig.1) has practically reproduced that of solid potassium [4], being just 0.2 eV above the latter. Also a similar layout of the solid/cluster spectrum and the atomic one[5]-with two main lines due to the $2p_{3/2}$ and $2p_{1/2}$ states- justifies the estimation of the atom-solid energy shift - in the core-ionization transition- by just subtracting the corresponding energies from each other. This gives $\Delta E_{a-c} = E_{ar} E_{cl} \approx 4$ eV. Since both cluster and atom energies are measured relative to the same, vacuum level this difference is then the change in the binding energy mainly due to the corehole extra-atomic screening [6], and should be the same as R^{ea} . (For the 3p cluster states with the cluster feature at \approx 21 eV [3] one gets the same \approx 4 eV separation as for the 2p level between the atom and the cluster responses in the spectra.

The cluster Auger spectrum recorded in the present work and shown in Fig.2 has a continuous shape, with at least three distinct maxima - at around 245 eV, at \approx 247, and \approx 249 eV. The cluster spectrum practically reproduces the shape of the solid-case spectrum from C.D.Wagner's Handbook of X-ray Photoelectron Spectroscopy [7]. The solid K spectrum has its maximum at 247.7 eV kinetic energy with the Fermi edge as a zero

point in the original spectrum [7], and thus \approx 245.4 eV relative to vacuum. The Auger parameter α_{sol} calculated for the $2p_{3/2}$ state with the values of Wagner, occurs to be equal to \approx 542 eV.

In order to interpret the cluster and solid Auger spectra it is helpful to look at the results for the $K2p^{-1}$ level decay in crystalline potassium halides [8]. Due to the 4s-electron absence in K ions in alkali-halide crystals the K2p

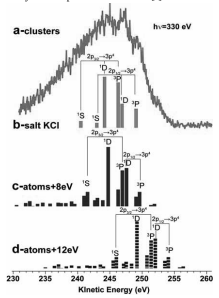


Fig. 2 a- Cluster Auger spectrum due to $2p^{-1}3p^{6}4s \rightarrow 2p^{6}3p^{-2}4s$ transition, four maxima are seen- at \approx 245, 247, 249, 252 eV; b- K 2p Auger spectrum for solid KCl; e-free-atom spectrum due to the same transition- shifted up in energy by 8 eV;

d - free-atom Auger spectrum shifted up in energy by 12 eV

Auger decay occurs to be analogous to the Ar-atom 2p corevacancy decay [8]: the same as in Ar final doubly ionized states ${}^{1}S$, ${}^{1}D$, and ${}^{3}P$ of the K^{3+} $3p^{4}$ configuration are populated. The experimental Auger kinetic energy spectrum of solid KCl [8], has practically the same spectral shape and even the same absolute energy position as the K metal cluster spectrum. Also in free K atoms the Auger spectrum layout is very similar to that of K salts and metal/cluster - probably because of the weak coupling of the 4s electron to the ionic core. It is tempting then to relate the most intense atomic Auger line at 236.7 eV -due to the $2p_{3/2}$ $\rightarrow 3p^4(^1D)4s^2D_{5/2,3/2}$ transition $[^9]$ - and the cluster feature maximum at ≈245 eV. There are three additional arguments for that: 1) \approx 245 eV is also the solid Auger energy [7]; 2)K $3p^4$ state with the same ${}^{1}D$ ionic-core term in the solid KCl is at \approx 245 eV; 3) Since the cluster and solid 2p binding energies are also very similar, the cluster Auger parameter would be very close to that reported for the solid [2], as it should be. The correspondence assumed for the atomic and cluster Auger spectra would mean the shift for the atomic Auger lines for ≈8eV up in kinetic energy (Fig.2). The other atomic lines when shifted up for 8 eV would also match well the overall cluster profile.

For the most intense atomic Auger $L_{3,2}$ - $3p^4$ (1D) $4s^2D_{5/2,3/2}$ lines one calculates: $\alpha_{at}(2p_{3/2})$ =300.6 eV+236.7 eV=537.3 eV, $\alpha_{at}(2p_{1/2})$ =303.4 eV+239.5 eV=542.9 eV. Within the 8 eV-shift assumption for the cluster transitions corresponding to the above used atomic ones, the Auger parameter for two- $2p_{3/2}$ and $2p_{3/2}$ -initial states are obtained as: $\alpha_{ct}(2p_{3/2})$ =296.5 eV + $E^{Aug}(2p_{1/2})$ =541.2 eV, and $\alpha_{ct}(2p_{1/2})$ = 299.2 eV + $E^{Aug}(2p_{1/2})$ =546.6eV. The difference between the atom and cluster parameter $\Delta\alpha_{ct-at}$ would

be 3.9 and 3.7 eV, respectively. As mentioned above, the cluster $\alpha_{cl}(2p_{3/2})$ value occurs to be very close to Wagner's 542 eV for solid K, what gives $\Delta \alpha_{sol-al}(2p_{3/2}) \approx 4.7$ eV.

However, $\Delta \alpha_{cl-at}$ and $\Delta \alpha_{col-at}$ estimated within the assumption of ≈ 8 eV Auger shift are too small to be twice the screening energy R^{ea} , which is shown above to be about 4 eV. Indeed, the derivation of $\Delta \alpha_{sol-at}$ assumes the two times larger charge of the final-state ion giving a 4 times larger energy of interaction with the surrounding electron density: $R^{ea}(3p^{-2})=4R^{ea}(2p^{-1})$. Then the difference between $R^{ea}(3p^{-2})$ and $R^{ea}(2p^{-1})$ is $3R^{ea}(2p^{-1})$. This relation would mean a ≈ 12 eV atom-solid Auger shift and correspondingly different cluster Auger spectrum interpretation. The most intense *atomic* Auger line at 236.7 eV due to $2p_{3/2} \rightarrow 3p^4(^1D)4s^2D_{5/2,3/2}$ transition would correspond to the cluster maximum at 249 eV, the latter being then due to the $3p^4(^1D)$ final state. I assignment oft is this assignment of the cluster Auger spectral features which would be consistent with the electrostatic considerations. A lot of intensity below 245 eV one can try explaining as due to the Auger electrons experiencing inelastic losses –in plasmon excitations and valence –electron excitations.

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Investigating the solid electrolyte interphase with functional electrolytes

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Electrolyte additives are one of the cheapest and most effective ways to improve safety and performance of the Li-ion battery. A small amount of an additive is added to the electrolyte to improve its characteristics. Here, film forming additives are investigated: vinylene carbonate (VC), lithium bis(oxalato)borate (LiBOB) and propargyl methane-sulfonate (PMS) (Fig. 1).

Fig. 1 Electrolyte additives used in our study.

When the battery operates, a surface layer is formed on the electrodes: it is called the Solid Electrolyte Interphase (SEI). This layer causes some capacity lost due to irreversible electrolyte reduction. However, the formation of a thin and stable SEI is needed to maintain passivation of graphite and protect it from, e.g., solvent co-intercalation. Electrolyte additives are being reduced prior to other electrolyte components; they change the composition of the SEI and the battery performance (Fig. 2).

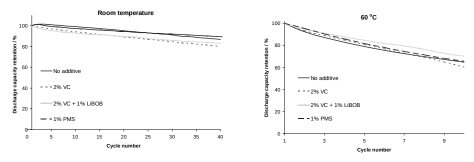


Fig. 2 Capacity retention in a LiFePO₄/graphite cell studied without additives and with VC, LiBOB and PMS.

The changes at electrode-electrolyte interfaces are studied with photoelectron spectroscopy (PES). Ion etching is commonly used to obtain depth profiling, however it can lead to decomposition of unstable surface products [1]. Here, synchrotron radiation at Max IV in Lund and Bessy in Berlin is used to obtain a unique non-destructive PES depth profiling. Four different excitation energies, in the 260-6900 eV range, are used. The excitation energy for

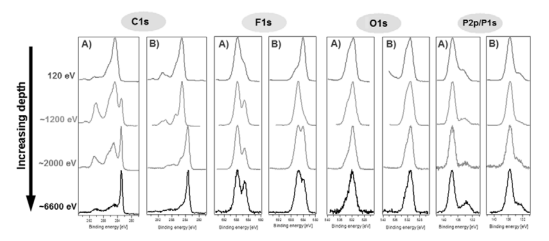


Fig. 3 Depth profiles of delithiated graphite measured without additives (A) and with PMS (B) at 4 different kinetic energies.

different core levels is varied for most surface sensitive measurements in order to obtain similar mean free path.

In figure 3 depth profiles of delithiated graphite cycled without additives and with PMS are shown. The bulk graphite peak is observed at 1200 eV for the non-additive sample and at 2000 eV for the PMS sample, which indicates that the battery cycled with the additive has thinner SEI. It has also more LiF, which in unwanted element of SEI due to low lithium permeability.

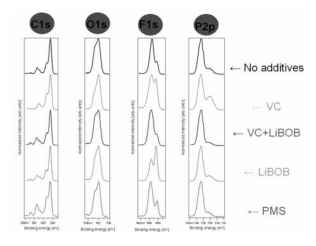


Fig. 4 Comparison of lithiated graphite spectra measured at 120 eV.

In figure 4 spectra measured for lithiated graphite at 120 eV (the most surface sensitive measurement) are compared. VC and LiBOB additives lead to formation of LiF in the surface film, however when they are added together to one electrolyte the formation of LiF decreases significantly.

Surface film formed on an electrode cycled with PMS shows more ether and carbonate containing products.

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Effects of solvation in the fragmentation of ammonia molecules

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The last decades Van der Waals clusters have been subject of growing interest to investigate matter in soft X-ray regime (e.g [1-2]). Ammonia (and water) molecular clusters are prototypes for studying the transition from isolated ions to solvation in the liquid phase [3]. The inter atomic/molecular forces encountered in the bonding of these clusters such as, the dispersion interaction, the dipole-dipole interaction and hydrogen interaction are expected to play a major role in redistributing internal energies.

"Coulomb explosion" of multiply charged clusters enables examining the various channels of energy dissipation in matter. Our results on atomic clusters demonstrate that nuclear motion takes actively part in thermo-cooling [4]. Such a mechanism in molecular clusters would activate vibrational modes resulting in new fragmentation pathways. To our knowledge experimental and theoretical studies on multiply charged ammonia molecular clusters are scarce. Only recently, delayed Coulomb fission of doubly charged stable ammonia cluster has been studied [5], and the fission mechanism is found to be well described within the framework of a dielectric liquid drop model.

Control of the fragmentation dynamics requires knowing the initial conditions that lead mother clusters to undergo fragmentation. This can be partially achieved by photo-ionization in the vicinity of the N-1s edge.

Experiment part

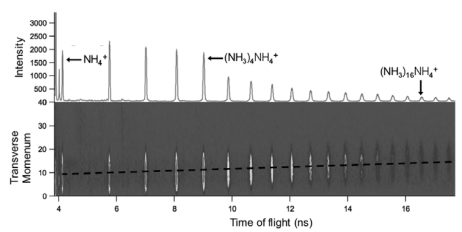


Figure 1: Fragmentation spectra at 420 eV. (Upper Panel) Ammonia clusters fragments time-of-flight. (Lower panel) Transverse momentum of the fragment. The dash line indicates the momentum drift due to the initial velocity in the cluster beam.

The experiment was carried out in the 1-meter section of the beam line I411 with a 3D imaging momentum spectrometer capable of high mass resolving power (m/ δ m~1500). We access the complete kinetics of ionic fragments by measuring individual ions in coincidence using a Position Sensitive Detector (PSD) to derive their momenta from the time of flight and positions. The clusters were produced with an adiabatic expansion source that allows changing cluster size from a few molecules to more than fifty molecules. With this

method of production, neutral clusters have an initial velocity in the order of 900 m/s that leads to an increasing drift of the momentum for larger fragments as can be seen in figure 1.

Results

In contrast to free molecules, the fragmentation of ammonia clusters does not produce small fragments such as H^+ , N^+ , NH^+ . All modes of fragmentation in the molecule are inhibited due to solvation. At 390 and 420 eV photon energies, singly and multiply charged clusters undergo fragmentation and mass spectra exhibit a simple pattern with only protonated $(NH_3)_nNH_4^+$ species in comparison to homogeneous species $(NH_3)_n^+$ observed at 10~eV photon energy [6]. These protonated species are attributed to a fast intra-cluster proton transfer with subsequent neutral evaporation.

There is a relatively high abundance of small fragments up to $(NH_3)_4NH_4^+$, where the first closed shell system with a charged ammonium molecule is in the center the cluster fragment. The four ammonia molecules are relatively tightly bound due to the hydrogen bonding interaction [7]. The intensity of fragments continuously decreases for larger fragments, without noticeable change when the second shell, $(NH_3)_{16}NH_4^+$, is closed. This suggests that interaction energy between the first-second shells and second-third shells, due to dispersion interaction and dipole-dipole interactions, are comparable with the internal energy left in nuclear motion.

Figure 2 shows an average angular correlation diagram between protonated fragments measured in coincidence. This diagram shows a preferential back-to-back emission, i.e. ions repel each other during the fragmentation. It suggests that the dissociation is fast and Coulomb repulsion is a primary driving force.

This behavior is quite general and does not depend on the photon energy and the fragment sizes. A very efficient charge

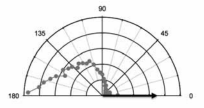


Figure 2: Angular correlation diagram between two protonated fragments. The arrow indicates the emission of the first fragments.

migration mechanism must occur. This (ultra) fast migration/delocalization of the charges before break up is further supported by the decrease of the kinetic energy released when the fragment size increases. Indeed this decrease means that charges are getting further away when the fragment size increases.

However, the angular correlation diagram is comparable to that of neon van der Waals clusters where the high mobility of ionic atoms is assumed to be the reason for charge migration [2]. Therefore a comparison with molecular clusters having heavier atoms is under progress to examine the interplay between these two competing factors, namely the charge delocalization and mobility of the ions.

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Soft X-ray photofragmentation of free protonated leucine enkephalin

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The response of peptides and its constituent aminoacids upon VUV and soft X-ray photoabsorption is of great interest in the context of astrobiology [1] and radiobiology [2]. Key questions concern the possibility of transport of intact gas phase peptides from space to earth and the molecular mechanisms underlying biological radiation damage. It is a well known fact that in most molecules the photodissociation efficiency, i.e. the ratio between the number of broken bonds and the total number of bonds, increases upon crossing of the K-edges. This is not only due to removal of the inner shell electron itself, but also due to the subsequent Auger decay of the core hole leading to the emission of a second electron [3]. Note that in a biological environment, this Auger electron will in turn lead to a whole cascade of secondary ionization events in the close vicinity of the initial photoionization site.



FIG. 1. a) Sketch of the Setup. b) Structure of leucine enkephalin

Here we report on a systematic study for backbone cleavage and subsequent fragmentation of the free protonated peptide leucine enkephalin (leu-enk) as function of photon energy $E_{photon}=40$ - 430 eV. The amino acid sequence of leu-enk is YGGFL as can be seen in fig. 1 b). For the case of this peptide, a comparison of C and N 1s-ionization is of particular interest since N is found in the peptide bond only whereas C is also present in the sidechains. The N 1s ionization site is thus neccessarily localized on the backbone of the peptide.

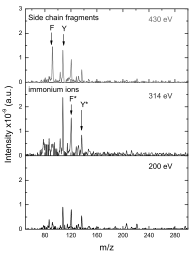


FIG. 2. Fig 1. Soft X-ray mass spectra of protonated leu-enk at several photon energies of the 411i MaxLab beamline.

In this report, we only give a brief sketch of the experiment (the details can be found in [1]). In fig. 1 a sketch of the setup is shown. A solution of methanol with $30\mu\mathrm{M}$ of leu-enk was introduced into the electrospray ion source (ESI) to generate protonated leu-enk (m= 556). The ions were transported and mass filtered by a linear and a mass filter quadrupole, respectively.

 $(\text{Leu-enk} + \text{H})^+$ ions were accumulated and cooled in a RF-ion trap to reach sufficient target density. The trap content was then exposed to XUV/soft X-rays photons from the MaxLab II I411 beamline. Intact molecules and fragments were extracted into a time-of-flight system equipped with a MCP detector.

Normalized photofragmentation mass spectra for (leu-enk + H)⁺ after synchrotron irradiation at 3 energies are given in fig. 2. Note the intensity increase when crossing the C K-edge by increasing photon energy from 200 to 314 eV.

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For a quantitative comparison of the spectra at different photon energies from 40 eV to 430 eV, in fig. 3 the yields of the most important photofragments are plotted versus the photon energy. Over the whole range of photon energies under investigation, the tyrosine (Y) side chain (m/z=107) and the immonium ions from leucine L (m/z=86), phenylalanine F (m/z=120) and tyrosine Y (m/z=136) were the most abundant photofragments. In our recent photofragmentation study in the VUV range of 8 to 40 eV the very same fragments dominated the spectra [1]. Up to photon energies of 200 eV, similar to the VUV studies larger fragments due to single backbone scission where observed, e.g. GF (m/z=205), b₂ (m/z=221), b₃/y₂ (m/z=278/279), and c₃ (m/z=295) fragments. Fragments with masses exceeding m/z=300 that are common in CID experiments [4] were very weak. Note, that due to a low m/z cutoff of the RF trap, the glycine (G) immonium ion (m/z=30) and other light fragments cannot be detected.

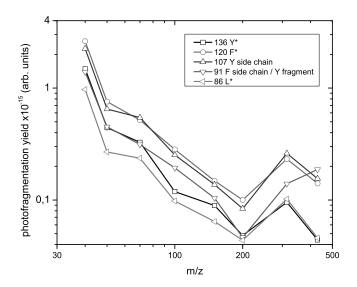


FIG. 3. Photofragmentation yield for leu-enk. The symbol * denotes immonium ions.

From fig. 3 it is obvious that all fragment yields decrease continuously between 40 eV and 200 eV. This is in line with the expectations since the cross sections for photoionization of atomic as well as molecular valence electrons are known to decrease strongly with photon energy in this energy range. All yields show a clear increase at 314 eV, due to the fact that the C K-edge at about 288 eV is passed. Interestingly, only moderate changes in the fragmentation pattern are observed. Peak ratios are only weakly affected by the dramatic change in ionization mechanism from valence shell photoionization to core-level ionization followed by Auger decay. At 430 eV the N K-edge is passed. Overall, here the fragment yields are smaller than for C 1s ionization which reflects the fact that leu-enk contains 28 C atoms, but only 5 N atoms. Assuming additive photoionization cross sections, K-shell photofragmentation should be 5-6 times weaker for N than for C. An obvious exception is the m/z = 91 fragment (C₇H₇) that can be due to a part of the phenylalanine (F) or of the tyrosine (Y) side chain. Due to N 1s photoionization m/z = 91 becomes the dominating peak in the spectrum which is remarkable in view of the few N atoms in the peptide. Formation of this side chain fragment is thus necessarily due to a localized photoionization that triggers a fast scission process. For the F case, Auger decay of the K-shell vacancy of the adjacent N atom could for instance involve one or two electrons from C_{α} - C_{β} bond, leading to a fast scission of this bond and release of the respective side chain.

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Soft X-ray photofragmentation of free protonated oligonucleotides

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Most biological effects of ionizing radiation are triggered by direct and indirect DNA damage. The latter is mainly due to DNA interaction with free radicals from radiolysis of nearby water molecules. Direct damage to DNA on the other hand can either be due to the primary quanta of radiation themselves or due to secondary particles such as electrons and ions. The interaction of soft X-rays with DNA in the condensed phase has been studied recently by Ptasinska et al. [1] who applied X-ray photoelectron spectroscopy to investigated chemical DNA modifications such as dehydrogenation and oxygen loss. However, the assignment of such reaction pathways to a particular damage mechanism remains ambiguous because of the interplay of direct and indirect damage.

In order to study the direct effect, only, over the last decade, numerous studies have been devoted to the investigation of ionization and fragmentation dynamics of isolated gas phase DNA building blocks upon interaction with energetic photons [2, 3], low energy electrons [4, 5] and keV ions [6, 7]. The gas phase studies have led to a much better understanding of the molecular mechanims underlying biological radiation damage. However, a major drawback of the gas phase studies was the fact that in order to reach sufficient density, effusive targets produced by evaporation were employed.

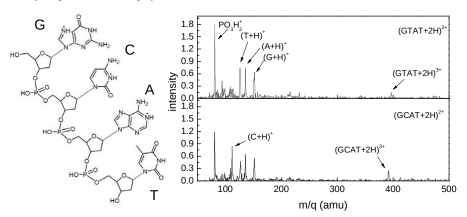


FIG. 1. a) Structure of GCAT. b) Photofragment mass spectra of $(GCAT+2H)^{2+}$ and $(GTAT+2H)^{2+}$ at 70 eV photon energy.

More complex biomolecules such as proteins and oligonucleotides already decompose thermally upon very moderate heating. To overcome this limitation we have recently commissioned an apparatus based on an inhouse built electrospray ionization (ESI) source, used to produce a beam of protonated biomolecules. After passing a radiofrequency (RF) ion funnel and a collisionally focussing quadrupole RF ion guide, the doubly protonated tetranucleotides GCAT and GTAT were mass selected by an RF quadrupole mass filter. Eventually a target of the selected biomolecular ions was accumulated in a 3D RF ion trap. A pulse of He buffer gas was injected to collisionally cool the trapped ions. The trap was interfaced with the MAXLab II I411 beamline. The trap content was exposed to soft X-ray photons for several 100 ms, before the trap content was extracted into a time-of-flight (TOF) mass spectrometer. The ions were detected by a silhouette-type micro-channel-plate detector (El-Mul) operated in analog mode and interfaced with a 1 GHz digitizer (Z-Tec). To remove contributions of buffer gas and neutral molecules from the ESI source to the mass spectra a cycling technique was used. Mass spectra obtained for ESI source off and ion beam off were subtracted from the inclusive spectra sequentially.

A typical photofragmentation spectrum with sufficient statistics required acquisition of about 1000 acquisition cycles. A typical mass spectrum obtained after 70 eV photoionization of the doubly protonated oligonucleotides GCAT and GTAT is shown in figure 2. In both cases, the strongest peak is due to $PO_3H_2^+$ from the sugarphosphate backbone. In cellular DNA the loss of such a phosphate group will induce the formation of a

single strand break. Pan et al [8] observed desorption of OH⁻ from solid NaH₂PO₄ upon low energy electron irradiation, which are expected to arise from POH bond cleavage by dissociative electron attachment, giving rise to the same fragment. Note, that while the RF-trap has a cutoff for m/z < 70, we do not observe (more or less) intact sugar fragments from the backbone, viz fragments in the 99 to 115 amu range, which most likely disintegrate and contribute to the unobservable fragment signal. This supports the notion that in DNA the sugar moiety is the most fragile, as noted elsewhere [9, 10]. At higher masses, strong peaks due to protonated nucleobases are found. For GCAT, the intensities follow the sequence C, A, G, T - for GTAT the cytosine moiety is obviously absent. We note, however that while for GTAT the intensities scale as T = A > G, for GCAT A > T. A more salient aspect of the present measurements is that while the molecular mass of the bases in the oligo is 1 amu less than the mass of the canonical free base (due to the glycosidic bond), here we always observe a base+1 amu cation. This means that base release following photoinization involves not just mere proton pickup from within the parent oligo (e.g. the original proton from electrospray ionization that often localizes on the bases), which would lead to the observation of a molecular mass equal to a base cation. Instead it is found that it involves both concerted hydrogen abstraction and proton transfer from within the oligo during dissociation, namely the observation of a base+1 amu cation. (GCAT+2H)³⁺ and (GTAT+2H)³⁺ due to non-dissociative photoionization can be clearly identified. In general, we must note that for all photon energies no larger fragments such as intact nucleosides, nucleotides, or half oligos, etc., are observed, which suggests that the protonated oligos are very sensitive to fragmentation.

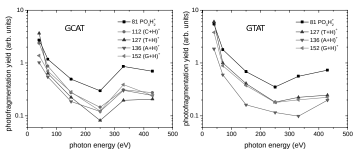


FIG. 2. Photofragment yields for the dominant dissociation products from GCAT (left) and GTAT (right) as a function of photon energy

In figure 2, the yields of $PO_3H_2^+$ and protonated nucleobases as a function of photon energy are plotted for both GCAT and GTAT. With increasing energy, the yields first decrease strongly, reflecting the decreasing photoionization cross sections for electrons from the valence band. The decrease continues up to photon energies of 250 eV, crossing P 2s and 2p absorption edges at 142 eV and 199 eV. At higher photon energies, the K-edges of $C \approx 288 \text{ eV}$) and $N \approx 410 \text{ eV}$) are crossed. It is obvious that photoionization of P does not contribute strongly to the yields whereas for K-shell photoinization of C and N, a clear increase in yield is observed for all fragment ions under study.

For all fragments under study, the photon energy dependence is similar. This finding implies that the location of the ionization site (P is only found in the backbone, N is only found in the nucleobases) is not affecting fragmentation. Such an independence could be due to relatively long fragmentation time scales. If all excitation energy deposited in the molecule by the photoionization process would distribute over the entire molecule by internal vibrational redistribution, the fragmentation would be site-independent. However, another explanation for our observation could be the fact that protonated oligonucleotides are relatively unstable, as obvious from the small fragments observed. Possible site effect would then be more pronounced in de-protonated oligonucleotides.

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Atom-solid binding energy shifts for K 2p and Rb 3d sublevels

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Binding energy shifts between free and solid state atoms for K 2p and Rb 3d photolines have been determined by measuring the vapor and solid state spectra simultaneously in similar experimental conditions applying synchrotron radiation excited photoelectron spectroscopy. This method has the important benefit that the work function is not needed to correct for different reference energy levels, therefore much more accurate values for binding energy shifts are obtained.

The measurements were carried out at the I411 beamline of the MAX II synchrotron radiation storage ring. A resistively heated oven was used to evaporate the solid samples. The 2p photoelectron spectra of K were measured with the photon energy of 365 eV and the 3d photoelectron spectra of Rb with the photon energy of 200 eV. In these measurements a needle like surface was located into the vapor beam and the spectra from the vapor around the tip and from the condensed solid layer were simultaneously observed. The vapor atoms in the vicinity of the solid tip are very accurately in the potential of the solid surface which form the joint reference potential. The needle was made from a copper wire which was covered with the tantalum wire wrapped around the aluminium oxide tube. Using this experimental setup both vapor and solid state spectra could be observed simultaneously in identical experimental conditions. No work function correction is needed because the spectra have the same reference energy level (work function potential).

K 2p and Rb 3d photoelectron spectra from simultaneous measurements are shown in figures 1 and 2. In the 2p photoelectron spectrum of K also bulk plasmons are seen, the plasmon satellites are broad features centered at 300.0 eV and 302.8 eV. The obtained energy values for the plasmons energies are 3.72 eV. The binding energy shifts between atom and solid for K 2p photolines is 4.2 eV and for Rb 3d photolines 3.7 eV. Photolines of solid K and Rb are thus several eV in lower binding energies relative to photolines of vapor. This is mainly done by the extra atomic relaxation (screening) of the ionized final state, which lowers the total energy of the

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emitting system and thus increases the kinetic energy of the emitted electron (decreaces the binding energy). The second contribution to the observed shifts arises from the initial state chemical shift of atoms in solids. This is however much smaller in elemental solids. The solid photolines are separated into surface and bulk components because the atoms are in two different chemical environments. Bulk sites experiences a stronger screening than surface sites and therefore binding energy of a core electron is smaller in bulk region than in surface region.

Accurate binding energy shifts between atoms and solids for 2p photolines of K and 3d photolines of Rb are reported. Vapor and solid measurements were carried out simultaneously in the same potential and determination of work function is not needed.

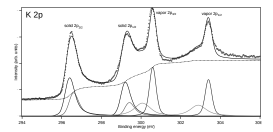


Fig. 1. K 2p PES taken simultaneously from vapor and solid sample. Blue lines corresponds to $2p_{1/2}$ and $2p_{3/2}$ vapor lines. Red line for solid is the surface and black line the bulk peak. Green lines correspond the bulk plasmons. Red circles are measured data points, solid black line is the total fitted spectrum and dashed black line is the Shirley background.

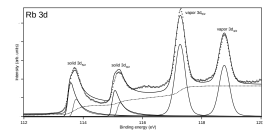


Fig. 2. Rb 3d PEs taken simultaneously from vapor and solid sample. Blue lines corresponds to $3d_{3/2}$ and $3d_{5/2}$ vapor lines. Red line for solid is the surface and black line the bulk peak. Red circles are measured data points, solid black line is the total fitted spectrum and dashed black line is the Shirley background.

Molecular fragmentation of pyrimidine derivatives following site-selective carbon coreionization

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Site-dependent fragmentation of three cyclic biomolecules - uracil, 5-bromouracil and thymine - into pairs of cations has been studied using electron-energy-resolved photoelectron-photoion-photoion coincidence (PEPIPICO) spectroscopy [1-3]. Previous studies concerning ionization site dependent fragmentation have mainly been carried out with linear molecules. We found out that all studied molecules have dicationic fragmentation channels whose intensity depends on the initial core-ionization site, although these channels cover only a relatively small fraction of the total fragment yield. The present study shows on one hand, that it is often the surrounding bond(s) of the ionized atom that will break following the initial core ionization, and on the other hand, that some specific fragmentation channels can display strong site-dependency where there is no direct correlation between the ionization site and the bond breakage locations.

The apparatus for the present PEPIPICO measurements consists of a modified Scienta SES-100 electron energy analyzer and a homemade Wiley-Mclaren type ion time-of-flight detector. More detailed description about the PEPIPICO method and the measurement equipment has described in detail in Ref. [4]. The C 1s photoelectron spectra measured coincidence with all photoions are presented in Fig 1. The peak positions are good agreement with earlier measurements thymine [5] and uracil [6], and their assignment is according to these works. As seen, different carbon sites are well resolved in all cases except for the overlapping C5 and C9 peaks in thymine.

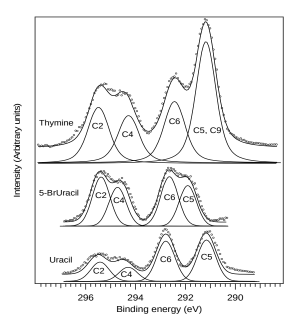


Figure 1. C 1s photoelectron spectrum (background subtracted) of thymine and 5-bromouracil measured with 330eV and of uracil measured with 315eV photon energy. Solid lines represent a least-squares curve-fitting decomposition of the spectrum. Structures of the studied molecules are shown next to the spectra.

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As the data analysis is explained and results can be found in more detail in Ref. [7], only a brief description is given here. In order to extract the site selective effects, so called PEPIPICO maps [8], where the ion pairs appear as tilted patterns, are used. For each photoline, also a PEPIPICO map is made, which allows one to see possible site selective effects as intensity variations between the same patterns in different maps. In Figure 2 there is an example of a significant and non-existent site selectivity.

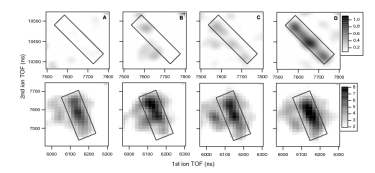


Figure PEPIPICO 2. patterns of the fragmentation Thymine. Top row: fragment pair (HNCOH+, $C_4H_4ON^+$), bottom fragment pair (HNCH⁺. HNCO+). Columns A-D correspond to the coreionization of carbons C2, C4, C6 and C5 C9, respectively.

Our study shows the suitability of electron energy resolved PEPIPICO for investigating the initial ionization site-dependent fragmentation in cases where individual atoms are separable by sufficient chemical shifts in the XPS spectra. All three molecules exhibit site-dependent fragmentation effects, which are, however, restricted to the weaker fragmentation channels. This is quite understandable, as the dominant pathways are likely to involve a multitude of electronic states thus being not sensitive to their exact population distribution by Auger transitions, whereas for the minor pathways the role of the ionization site and the population of specific dicationic states can be much more important.

Some of the channels where site-dependency was observed involved light fragments such as C⁺ and/or CO⁺. In these cases the results indicate that the bonds adjacent to the core-ionized atom tend to weaken, enhancing the production of the corresponding fragments. The most interesting pathway was observed in thymine and uracil, producing large fragments in two-body fragmentation ((HNCOH⁺, C₄H₄ON⁺) for thymine and (HNCOH⁺, C₃H₃ON⁺) for uracil). It is a clear example that even relatively complex cyclic molecules can exhibit strong site-dependent behavior in specific cases. According to our interpretation, in this particular case the site-dependent enhancement appears clearly, since (a) the Auger decay populates selectively a particular dicationic state only at certain core-hole sites and (b) due to low excess energy in that state, the variety of available fragmentation pathways is much restricted.

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Valence Photoionization of Neutral Metal Clusters

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Clusters are nanometer scale objects that contain few to few thousand atoms or molecules. These objects pose peculiar properties from gaseous and solid states of matter that vary as a function of cluster size. Therefore, for some time clusters have been of keen interest in studies of basic as well as applied physics. Since the 80's numerous studies of ionization of vapor phase clusters using laser sources have been published, but only recently cluster sources providing target densities that are sufficient for photoionization studies using synchrotron radiation have been developed. The use of synchrotron radiation allows one to probe the energy regions that are free from various valence excitations that hinder the data analysis. VUV and X-ray energy region photons may also ionize clusters from core levels, thus opening Auger decay channels.

In a recent study [1], our lately constructed exchange metal cluster source (EXMEC) [2] was used to study valence ionization of initially neutral K and Rb alkali-metal clusters. The experiment was carried out at MAX-lab at beamline I411. The ionizing photon energy used was 40 eV and the size range of the clusters varied from 30 to 650 atoms. This allowed us to study for the first time valence ionization of medium sized initially neutral metal clusters in the photon energy region free from any disturbing resonances.

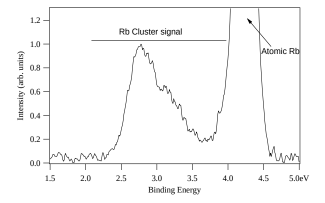


Figure 1: Valence photoelectron spectrum of medium sized (~ 500) Rb clusters. The spectrum shows also signal from 5s valence ionization of atomic Rb.

As an example, figure 1 shows an experimental valence photoelectron spectrum of initially neutral Rb clusters measured at MAX-lab. The average size of the clusters in this case is approximately 500 atoms. The photon energy was used 40 eV. The cluster signal is seen between the atomic valence ionization energy of about 4.18 eV and the bulk work function of 2.16 eV. The position of the most intensive point of the cluster signal was seen to move towards the atomic line as a function of decreasing cluster size. In addition, in the case small clusters (size < 100 atoms) the photoelectron spectrum start to show discrete structures due to less dense array of occupied valence energy levels. On the other hand, the photoelectron spectra obtained from large clusters was seen to approach the shape expected for solid Rb [1].

Theoretical *ab initio* description of photoionization of clusters is a very demanding problem. The size of the calculation increases exponentially as a function of cluster size, and reliable description

of photoelectron wavefunctions even for the smallest clusters is far from a trivial task. Therefore one need to rely on approximative descriptions of different levels. In our recent study we applied so-called jellium model to describe the experimental results [1,3]. In this simple model the cluster's valence electrons are assumed to move independently in a spherically symmetric potential created by the other electrons and nuclei. Figure 2 shows the result from an example calculation for Rb₅₈ cluster together with two continuum wavefunctions.

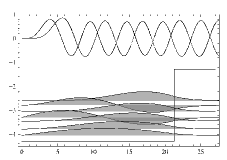


Figure 2: Calculated Rb₅₈ cluster's valence jellium orbitals in a spherical potential well and two continuum partial wavefunctions.

The model was used to simulate our experimental findings. As a result we found that the valence ionization of molten alkali-metal clusters can be qualitatively described surprisingly well using such a simple model. Our approach allowed the calculation of continuum wavefunctions and the ionization dipole matrix elements. From these calculations we found that the use of continuum wavefunctions in the description of valence ionization in clusters is crucial. The result thus demonstrated that the commonly used density of states approximation for relative ionization probabilities may yield misleading results due to lack of proper description of matrix elements. Figure 3 depicts example calculations for photoelectron spectra of three different sizes of Rb clusters. Comparing the largest size (Rb $_{558}$) to the experiment shown in figure 1 a very nice agreement can be seen.

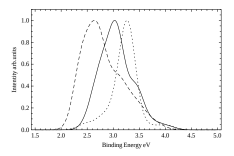


Figure 3: Simulated valence photoelectron spectrum of Rb_{40} (dotted), Rb_{58} (solid) and Rb_{558} (dashed) clusters.

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An XPS study of dolomite

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Dolomite (CaMg(CO3)2) is a carbonate mineral that consists of alternating layers of calcium (Ca^{2+}), magnesium (Mg^{2+}) and carbonate (CO_3^{2-}) ions, which are perpendicular to the crystallographic c-axis [1 – 3]. Usually dolomites are formed from already existing calcites ($CaCO_3$) in a process called dolomitization. In the process calcites are infiltrated by magnesium rich solutions, which can be formed in variable environments. The dolomitization process needs efficient hydrological circulation, and usually not all of the calcite is converted to dolomite. Dolomites can also precipitate straight from the solution into the pore spaces of the sediments [4]. Dolomites are studied widely perhaps due to the "dolomite problem" [5,6]. It is generally known that precipitation of dolomite occurred in a large scale in the past, but for some reason it is very rare at present. An environmental aspect of dolomite and other carbonates lies in their role in the storage of carbon dioxide. For this reason, the dissolution and precipitation of dolomite has been studied widely [7 – 11]. In this study, we have measured the Mg 2p and Ca 3s spectra of a metamorphosed dolomite from Reetinniemi, Finland.

Prior to experiments, the dolomite sample was mechanically sawed to fit the sample holder and cleaved in UHV. The experiments were carried out at the I411 beamline of MAX II storage ring in Lund, Sweden. The beamline is equipped with a modified SX-700 monochromator and a hemispherical Scienta R4000 electron energy analyzer. The pressure was typically around 10⁻⁸ mbar. A pass energy of 100 eV was used for both measurements shown in Figures 1 and 2. The photon energies can be seen in the Figures. Dolomite is an insulator, which makes it difficult to study with photoelectron spectroscopic method. Its indirect electronic band gap has been reported to be ~5.0 eV [12]. We therefore used a special home-made sample holder with integrated dipole type electron gun. The filament and the extraction aperture were from a PHI 04-085 specimen neutralizer. The flood gun was about 10 mm from the sample surface. The kinetic energy of the electrons was set to 4.2 eV. The voltage over the gun filament was typically between 2 and 3 V, while the current was kept at ~ 3 A. All photoelectron spectra measured at MAX II were analysed with a Unifit program [13]. The Shirley background correction method and a Gaussian-Lorentzian peak shape were used.

Figures 1 and 2 show the Mg 2p and Ca 3s spectra of the dolomite with the excitation energies of 780 and 400 eV, respectively. The Mg 2p doublet is fitted with spin-orbit splitting $\Delta_{s\circ o}$ = 0.28 eV [14]. These results suggest that the cleavage process might have left magnesium on the surface, because the area under the Mg 2p peak is significantly greater with the excitation energy of 400 eV than with the excitation energy of 780 eV when compared to the area of the Ca 3s photoelectron line. Our comparison measurement with conventional XPS gave a result of 36 % for the area under the Ca 3s peak, which is consistent with the synchrotron experiments. The Mg enrichment on the surface cleaved in vacuum may be different from a cleaved dolomite exposed to air, which, according to Hu et al. [8], shows close to stoichiometric surface. Surface sensitivity was enhanced in their measurements by observation of electron emission from a 60-degree angle from surface normal. In addition, Hu et al. discovered that exposure to aqueous solution caused surface enrichment of Mg or Ca depending on the level of saturation.

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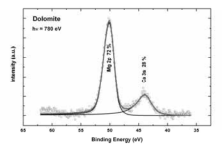


Figure 1. Mg 2p and Ca 3s spectra of the dolomite from Reetinniemi, Finland.

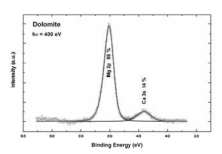


Figure 2. Mg 2p and Ca 3s spectra of the dolomite from Reetinniemi, Finland. The portion of magnesium has increased with surface sensitivity.

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Electron-ion coincidence study of photofragmentation of the CdCl₂ molecule

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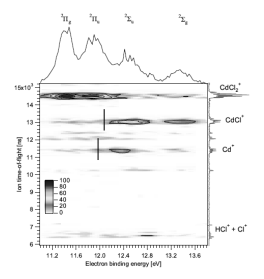
Electron-ion coincidence (PEPICO) methodology is nowadays often employed for studies of various molecular systems [1-2], as it provides means to separate electronic processes via their link to specific ions. The combined measurement of electron spectra and ion production can be used to obtain high resolution information about molecular processes. The ion data may be used to correlate fragments and, depending on the detection method, gain additional information about the dissociative processes. These experimental studies are also applicable to many theoretical studies such as the verification of molecular modeling calculations, and relevant applications exist among different fields of research such as atmospheric sciences, where breakdown of molecules can have large scale effects. The electron spectroscopy group of University of Oulu and collaborators, especially at University of Turku, have performed several PEPICO studies [3-5] with synchrotron radiation provided by MAX-lab, in UV and soft X-ray regimes.

Cadmium is a group 12 metal, sharing chemical properties with other metals of this group, zinc and mercury. Cadmium has previously been heavily used in applications such as batteries. Nowadays cadmium's use is discouraged due to its toxicity to human health and the environment [6] – for example the European Union has negotiated a directive in 2006 banning the sale of batteries, which contain more than trace amounts of mercury and cadmium. Cadmium, alike to mercury, is typically found in +2 oxidation state so it forms, among others, the linear cadmium dichloride (CdCl₂) molecule.

In this work, we studied the UV-induced photodissociation of $CdCl_2$ with UV lamp (He I α , hv = 21.218 eV) and synchrotron excitation at beamline I411 [7] in MAX-lab (hv = 40 eV), for molecular valence and 4d, respectively. The electron spectra for valence and Cd4d are well-known from previous studies. [8,9] The photoionized molecule, its photoelectrons and photofragments were observed, and the connection between the ionized states and produced fragments were made. The energetics and possible fragmentation pathways were examined with ab initio Multi-configurational self-consistent field (MCSCF) and Coupled Cluster (CC) calculations. The calculated results are in qualitative agreement with the observed data.

Figs. 1 and 2 depict the recorded data in an overview style. These maps are extracted from the coincidence data – the horizontal axis represents the observed electrons, and the vertical axis represents the flight time of the cations. The total electron and TOF spectra are displayed on top and right sides, respectively, of both figures. The actual extraction of parameters such as appearance energies was done by inspecting the coincident ion yields (CIYs), which contain the ion-specific yields as a function electron binding energy, linking the internal energy changes by photoemission to fragment yields [10]. The vertical black bars of Fig. 1 represent the fragment energies from CC calculations.

The inspection of the appearance energies of the fragments reveals which photoionized states are energetic enough to allow for different fragments to be produced. The results are in agreement with the traditional, tentative view of the linear molecules and their σ and π bonding – it can be seen that the π bonds, due to their non-axis orientated charge distribution, can be essentially seen as non-bonding as a contrast to σ bonds. The σ^{-1} ionized states were found to be competing between production of CdCl $^+$ and Cd $^+$. The 4 σ^{-1} photofragmentation was found to be dominated by production of Cl $^+$, which suggests that some charge-transfer process takes place. The observed appearance energies are in agreement with the calculated fragment energies.



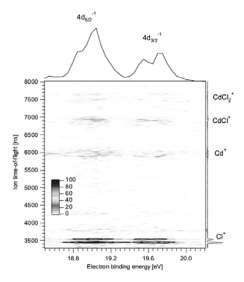


Fig 1. PEPICO map for CdCl₂ valence photoionization. [11]

Fig 2. PEPICO map of photoionized *Cd4d* ¹ states for CdCl₂. [11]

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Dissociation of C₆₀ fullerenes studied with a linear momentum coincidence spectrometer

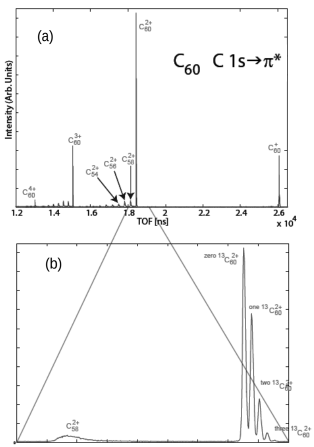
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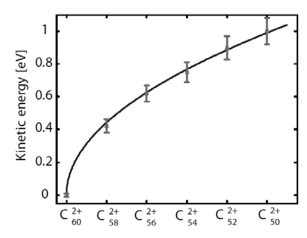
Since it was discovered in 1985 by Kroto et al. [1] the C60 molecule has attracted much attention and been subjected to the study of its electronic structure as well as dissociation dynamics [2-4]. Photo-fragmentation of C_{60} molecules has been investigated after electron impact by Foltin et al [5]. They showed that after ionization with 200 eV electrons, production of C_{56}^{2+} and C_{56}^{3+} preferentially occurs by stepwise ejection of neutral carbon-dimers rather than direct ejection of C4 even though the latter is energetically favorable. Synchrotron induced resonant core excitation and ionization of fullerenes studied by TOF mass-spectroscopy by Aksela et al. showed a significantly stronger tendency for the multiply charged fullerene to dissociate at the C $1s-\pi^*$ resonance compared to C $1s-\sigma^*$ resonance or far above resonance, which they attributed to the difference in valence hole state populations [6].

The objective of this study is to probe the dissociation sequence after Auger decay of the C_{60} molecule which we aim to achieve with careful analysis of KER distributions. We have carried out momentum imaging spectroscopy on cationic products formed by resonant coreexcitation. Our setup is a Wiley McLaren type TOF spectrometer equipped with a position sensitive delay line detector.



Figur 1. (a) C 1s- π^* state at 286.2 eV. Up to tetracationic ions are detected of both complete and fragmented fullerenes. (b) For undissociated fullerenes, we can even resolve the isotopeshift.

Access to complete kinematic information combined with a time to digital converter with a very high sampling rate (25 ps) allows us to extract kinetic and anisotropic distributions more reliable than previous studies. Fig. 1 (a) shows a TOF mass spectrum after core-excitation to the C 1s- π * state at 286.2 eV. Energy calibration was made by comparing total ion yield with that in Ref. [6]. Up to tetracationic fullerenes are detected. To the left of each undissociated C₆₀ peak a progression of even-atomic fragments are visible. This feature is a well-known characteristic since geometry considerations impose a constraint for only even-atomic fullerenes to be stable. Fig. 1 (b) gives an expanded view of the dication and its



Figur 2. Kinetic energy of fragmented fullerens have a square root dependence which indicates a sequential dissociation process.

primary dissociation channel, C582+. The peak splitting of the dication is due to the isotope distribution. The main difficulty TOF spectroscopy on large molecules is that resolution decreases with mass. However, our design that utilizes a long drift tube as a dispersive element and the high sampling rate of the TDC clearly allows us to even resolve the isotopic shift for the undissociated For fragmented dication. species however, this fails due to the KER during breakup. As a means to retrieve the KER distribution despite the isotopic overlap we use the following approach: The TOF peak for C_{60}^{2+} can be seen as a convolution of a line-broadening

mechanism, F_{exp} due to thermal energy

and experimental limitations and a discrete binomial function, Bin(60; 0.011) where 60 is the number of atoms and 0.011 is the natural abundance of 13C

$$TOF(C_{60}^{2+}) = Bin(60; 0.011) \oplus F_{exp}$$

The TOF peak for C_{60-2n}^{2+} however also has a kinetic energy contribution, F_{KER}

$$TOF(C_{60-2n}^{2+}) = Bin(60 - 2n; 0.011) \oplus F_{exp} \oplus F_{KER}$$

Multiplying each isotope peak in the undissociated fullerene with the corresponding factor $\frac{Bin(60-2n)}{Bin(60)}$, we retrieve the TOF for C_{60-2n}^{2+} for zero kinetic energy released. Deconvoluting this from the actual TOF distribution, results in the kinetic energy contribution

$$F_{KER} = TOF \ C_{60-2n}^{2+} \ \ominus \frac{Bin \ 60-2n}{Bin \ 60} \cdot TOF \ C_{60}^{2+} \ .$$

By assuming an isotropic distribution of ejected fragments, we estimated the mean kinetic energies of the fragmented fullerenes. At 100 eV photon-energy for the C_{58}^{2+} dication we found the peak at 0.4 ± 0.1 eV, in agreement with previous estimates [2]. We investigated the KER of C_{60-2n}^{2+} fragments for n={1-5} at the C $1s-\pi^*$ resonance (see Fig. 2) and found a square root dependence of KER on fragment size. Assuming the internal energy before breakup is the same irrespective of resulting fragments, this pattern can be explained with a five step sequential dissociation process of carbon dimers where the energy released in each step is convoluted together which results in the behavior of Fig. 2.

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Depth Profiling the Li-ion Battery Solid Electrolyte Interphases

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The Solid Electrolyte Interphase (SEI) [1] is a passivating layer of electrolyte reaction products deposited on lithium-ion battery electrodes. It is believed to play an important part in the ageing of lithium-ion batteries which is one among many reasons to aim for a more complete understanding of the composition, morphology and thickness of the SEI. The focus of this work has been to develop the method of non-destructive PES depth profiling for the study of the lithium-ion battery SEI.

LiFePO₄/graphite batteries with 1 M LiPF₆ EC:DEC 2:1 electrolyte cycled for 3 or 3.5 cycles and left in discharged or charged state, respectively, were examined at the I411 beamline at the MAXLAB synchrotron, Lund, the KMC1 beamline at the BESSY synchrotron, Berlin, as well as at an in house PHI 5500 XPS system equipped with an AlK α 1486.6 eV x-ray source. At the MAXLAB synchrotron, the excitation energies were varied so that the kinetic energy of emitted electrons was approximately the same for all probed core levels, 135 eV \pm 15 eV. At the BESSY synchrotron, fixed excitation energies of 2300 eV and 6900 eV were used. The batteries were dismantled in an argon filled glovebox and transferred to the XPS system vacuum without exposure to air using specially built transfer equipment. The resulting depth profiles of lithiated anodes and cathodes are shown in figure 1.

Figure 1 shows that the intensity of the bulk electrode peaks increase with probing depth. This is clearly seen in the anode C1s and cathode P2p/P1s spectra by that as the probing depth increases, so does the intensity of the lowest binding energy core level peaks interpreted as lithium intercalated graphite and phosphate, respectively. The depth profiles were used to calculate the SEI thicknesses which were found to be 10 nm on a delithated and 30 nm on a lithiated anode. The cathode SEI thickness was found to be a few nm, only. The charged anode SEI was found to be richer in ether bonds and phosphorous-fluorine bonds than the discharged anode SEI.

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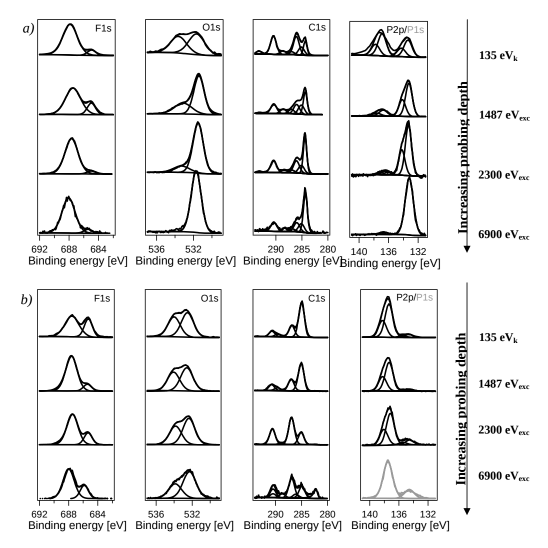


Figure 1. Deconvoluted PES depth profiles of a) a lithiated LiFePO₄ cathode and b) a lithiated graphite anode. The most surface sensitive measurements were made with approximately the same kinetic energy, E_k of electrons emitted from all probed core levels. The more depth sensitive measurements were performed with fixed excitation energies, E_{exc} . The phosphorous depth profile shows both P2p (black) and P1s (grey) spectra, though only the P2p binding energy scale is shown in the picture.

C1s photoelectron spectroscopy and modeling of ethane clusters

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Carbon 1s photoelectron spectroscopy may serve as a valuable tool in estimating the mean cluster size present in a cluster beam. In this study, we present modeling and experimental results for free, neutral clusters of ethane. Theoretical spectra of ethane clusters of different sizes were constructed and compared to an experimental carbon 1s (C1s) photoelectron spectrum. Based on the shift in ionization energy between the cluster and the monomer peak and the detailed shape of the carbon 1s (C1s) photoelectron spectrum, we estimate the mean cluster size produced at a stagnation pressure of 1740 mbar at room temperature. In addition, the mean free path of the escaping photoelectron in the cluster is approximated.

Molecular dynamics (MD) simulations were performed on clusters containing 13, 57, 147, 311, 469 and 803 molecules. The simulations started out from structures prepared as spherical cuts from crystalline ethane at 90K and 60K. The simulations employed a polarizable force field, amoeba, included in TINKER. Both for the neutral and the C1s ionized state, electrostatic multipole moments were obtained by means of distributed multipole analysis (GDMA) of electron densities. A model photoelectron spectrum can be constructed by the procedure described in Ref 3.

The lineshape of the photoelectron spectrum is influenced by the mean free path of the escaping photoelectron, λ . Figure 1 shows the effect of increasing λ from 5 to 1000 Å. The pronounced shoulder at the low-energy side of the spectrum grows with increasing λ , and by fitting the model spectrum to the experimental spectrum one may obtain a good estimate of λ in the experiment.

From the simulations, one may calculate a theoretical shift in C1s ionization energies, ΔIE , between the cluster and monomer peak. Figure 2 shows theoretical predictions of shifts in ionization energies for different cluster sizes, for two values of λ and for two simulation temperatures.

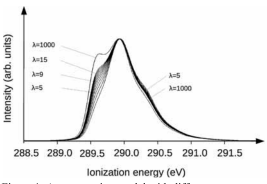


Figure 1. An attenuation model with different attenuation lengths, $\lambda/\text{Å} \in \{5, 15, 100, 1000\}$, is applied to a theoretical C1s photoelectron spectrum of ethane.

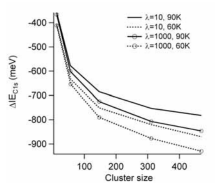


Figure 2. Theoretical predictions of shifts in ionization energy, Δ IE (meV), for different cluster sizes. Here shown for λ /Å=10 and 1000 and T= 60K and 90K.

Free, neutral clusters of ethane were produced by adiabatic expansion from a sample at high stagnation pressure through a narrow nozzle and into a low-pressure area. The expansion leads to supersaturation followed by condensation and cluster growth. Several combinations of stagnation pressure and temperatures at the nozzle were employed to affect the distribution of cluster sizes. Moreover, to aid the clustering process, a 20/80 % mix of ethane and helium gas was used. The clusters were characterized by C1s photoelectron spectroscopy at beamline I411 at MAX-lab, using a photon energy of 330 eV.

Here, we present results for the spectrum acquired at a stagnation pressure of 1740 mbar and room temperature, shown in Figure 3. The spectrum is fitted with theoretical lineshapes for the 311-membered cluster at 90K with $\lambda=9$ Å. By mapping the variation in the goodness-of-fit parameter with λ , we estimate λ to be 9±2 Å for ethane. This agrees well with inelastic mean free paths for a selection of organic compounds and with 50 eV kinetic energy of the outgoing photoelectron.⁴

From the fit, we obtain a shift in ionization energy, Δ IE, between the monomer and the cluster contributions of 0.76 eV. This agrees well with the shift calculated from the MD simulations⁵ for the 311-cluster and suggests that the mean size of the clusters thus produced, at the given experimental conditions, is approximately 300.

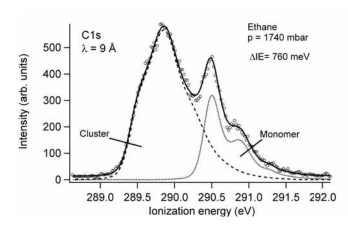


Figure 3. Experimental C1s photoelectron spectrum of ethane clusters and free ethane monomers (at RT). Experimental data points are shown as circles and the solid black line represents a least-squares fit to the data. The contributions from monomers and clusters are shown as dotted and dashed lines, respectively.

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Evaluating the binding energy shifts on iron group atoms

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Tunable synchrotron radiation makes possible selectively to measure binding energies of the electrons in atoms and solids by monitoring photoelectrons and measuring their respective kinetic energies with known photon energy. The experiment carried out at Max-lab, we measured binding energy shifts of transition metals which are interesting in point of process metallurgy, where the high power electrical arcs are commonly used for melting process in which the metals are excited and ionized in bulk and vapor form.

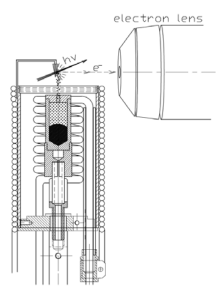


Figure 1. Experimental setup.

Measurements were carried out at soft X-ray beamline I411 at MAX-II storage ring [1]. The measurement setup (figure 1) consists of an inductively heated oven [2] and a needle located directly above the oven. This allows a vapor stream from the oven to condense on the surface of the needle, continuously keeping it clean and the same elemental composition as the vapor. The setup is adjusted so that the synchrotron beam is slightly absorbed by the needle and simultaneously by the vapor around the surface. Following photoelectrons are measured and resulting spectra are combinations of bulk and atomic spectra.

The vapor near the surface of the needle experiences the same potential as the surface thus both are referenced to the same level. Simultaneous measurement of the spectra allows direct comparison of the bulk and atom spectra. The photoelectrons from the both phases of matter are also affected equally by the different inaccuracies of the measurement setup, which increases accuracy of binding energy comparison. Binding energy shift between the phases can be determined directly in these measurements unlike in the

separate measurements, where usually not so accurately know work function of the solid phase must be taken into account, due to separate measurements referenced differently in bulk and vacuum [3].

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Photoelectron spectroscopy of solvated salts in water clusters

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In this work, the large water clusters doped with alkali halides were studied by the photoelectron spectroscopy technique. Clusters were produced by recently developed Exchange Metal Cluster Source (EXMEC) which was preliminary designed for electron-spectroscopic studies of neutral metal clusters¹. Previous studies with the EXMEC set-up show its capability for cluster production in the size regime from few tens to few hundred of atoms per cluster¹. This study shows that the set-up is able to produce also water clusters which are doped by salt molecules vaporized in the oven system. A schematic view of the cluster source is shown in Fig. 1. Liquid water is evaporated inside the cluster source in the first oven. High pressure He gas is let through the water oven to take water vapor out via a conical nozzle and to facilitate water cluster formation. In the nozzle the water clusters are formed via an adiabatic expansion process. The water cluster beam is skimmed downstream by a copper skimmer. Further downstream the water clusters are doped by alkali halide monomers which are created inside the second resistively heated oven with entrance and exit orifices on the axis of the water cluster beam. Leaving the second oven the clusters are ionized by synchrotron radiation and the photoelectrons emitted are recorded by a hemispherical electron analyzer, mounted in 90° angle with respect to the polarization direction of the radiation.

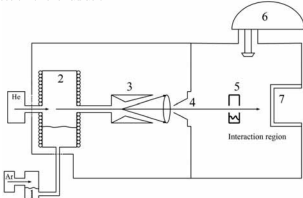


Figure 1: Schematic view of experiment:

- (1) water storage,
- (2) water oven,
- (3) nozzle, (4) skimmer,
- (5) oven and interaction region, (6) Scienta R4000 hemispherical electron analyzer and (7) liquid nitrogen cooled cold trap.

When alkali halides are dissolved in water, the molecule dissociates as anions and cations. There are both experimental and theoretical evidence that large polarizable anions are more attracted by water liquid/vapor interface whereas the cations are enhanced just below the surface in bulk water (see e.g. ² and references therein). Thus, the surface-sensitive method like low-kinetic energy photoelectron spectroscopy, used in this work, suites particularly well in studies of anions and cations in water. Alkali halides studied here are NaBr, NaCl, NaF, KBr, KCl, and KI. The valence region photoelectron spectrum in the binding energy region of 6–45 eV was measured for salt-water clusters using a photon energy of 79.6 eV. Additionally, the Br 3d region was measured with 156.7 eV, Cl 2p region with 227.7 eV and 234.0 eV (for NaCl and KCl, respectively), and I 4d region with 79.4 eV photon energies. The photoelectron spectral features of salt-water clusters were separated from those originated from pure alkali halides, water molecules, and water clusters without dissolved salt by measuring also the corresponding photoelectron spectra separately. The Na 2p region photoelectron spectra of NaCl and NaBr are shown in Fig. 2. The spectra show the Na⁺ 2p photoline of salt-water clusters, the Na 2p

Table 1: Binding energies of anions and cations in water clusters E_{cl} and in aqueous ions E_{aq} . The binding energies of different spin-orbit states are separated by slash.

ion		E_{cl}	E_{aq}^a
K^{+}	3p	22.3 ± 0.08	22.2 ± 0.06
Na^{+}	2p	35.5 ± 0.06	35.4 ± 0.04
Cl^-	2p	$203.02/204.66 \pm 0.05$	$202.1/203.6^b$
${ m Br}^-$	3d	$73.3/74.4 \pm 0.05$	$74.3 \pm 0.09 / 73.2 \pm 0.07$
I-	4d	$54.3/56.0 \pm 0.05$	53.8/55.8

^aFrom ref. 3, ^bFrom ref. 5

photolines from a pure alcali halide molecule and the He 1s photoline. A broad structure between the He and cluster lines is from water molecule and water clusters.

In table 1 the measured binding energies of core electrons for alkali and halide parts of salts in water clusters are given. In addition the corresponding binding energies of the ions in aqueous solutions are given. A remarkable result is that the obtained binding energies of salt-water clusters are very close to the binding energies of aqueous solutions reported earlier by Winter et al. 3 , 4 . Some discrepancies may arise from the different binding energy calibration. Helium 1s photoline serves as a reliable calibration peak in our spectra of cations, but the binding energy calibration for the higher binding energy spectra of anions may contain some uncertainties due to the lack of reported photoelectron spectra for pure molecular salt vapors. The binding energy data reveals that when a salt molecule is picked up by a water cluster, the salt molecule dissolves in water, and it is dissociated. The ions are then solvated and dispersed like in liquid water. The same E_{cl} and E_{aq} binding energies mean that the water surrounding of anions and cations is very similar both in liquids and clusters.

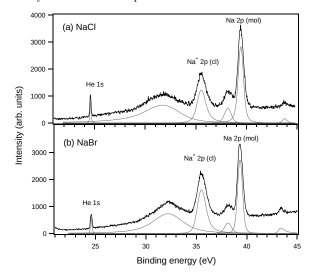


Figure 2: The photoelectron spectra of NaCl and NaBr measured at Na 2p region.

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4d¹ photoelectron and subsequent N_{4,5}OO Auger electron spectra of Sb

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Synchrotron radiation excited $4\sigma^1$ photoelectron and subsequent $N_{4,5}OO$ Auger electron spectrum of Sb have been measured [1] at beamline I411. An inductively heated oven was used to evaporate the solid-phase Sb, which evaporates predominantly as Sb₄ molecules, which can be further fragmented to Sb₂ and Sb by strongly increasing the temperature of the upper part of the specially designed oven system [2]. The electron configuration of Sb is $[Kr]4\sigma^95s^25p^3$ and complex features in the spectra created by an open shell electronic structure have been interpreted using multiconfigurational Dirac-Fock calculations utilizing GRASP92 and GRASP2K packages [3]. The photoelectron spectrum (PES) (in Fig. 1) was interpreted satisfactorily by a single non-relativistic configuration and 26 lines was identified. For the subsequent Auger electron spectrum (AES) (in Fig. 2) both single and multiconfigurational Dirac-Fock calculations were carried out. Previously published optical data of the Auger final states was also used for interpreting the structure of the Auger spectrum. The $N_{4,5}O_1O_{2,3}$ transitions lie in low kinetic energy region, which caused some challenges to the data handling, mainly because of the exponentially increasing background towards zero kinetic energy and non-linearly behaving transmission. It was found that

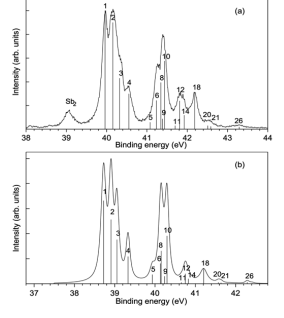


Figure 1. (a) Experimental (hv=90eV) and (b) calculated 4d photoelectron spectrum of Sb.

despite of complex 4d⁻¹ PES displaying a lot of transitions, the most intense structures in the N_{4.5}OO AES are created by transitions from the initial states ⁵D₄, ⁵D₃. ⁵D₂. and ³D₁ (configuration $[Kr]4d^95s^25p^3$) to final states ${}^4P_{1/2}$, ${}^4P_{3/2}$, and ${}^{4}P_{5/2}$ (configuration [Kr]4d ${}^{10}5s^{1}5p^{2}$). It noticed that also was configurations having 5d and 6s orbitals occupied do not mix with the 4P final states.

The comparison to the previously published corresponding Sb₄ study by Urpelainen et al. [4] was made, and it was noticed that the final states of the N_{4,5}OO Auger transitions are very differently populated in Sb₄ compared to the atomic Sb. In the case of Sb₄, the transitions to the final states, having one hole in the molecular orbital with significant contribution of the 5s atomic orbital and one hole in the MO with significant contribution of the 5p AO, were as

important as the transitions to the final states, having two holes in the MOs with significant contribution of 5p AO. In atomic Sb, the final states having $5s5p^2$ configuration (region B in Fig. 2) were much more populated than the states with $5s^25p$ configuration (region A in Fig. 2). A tentative explanation to this was offered by counting the number of possible single configuration final states:

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in the atomic case there are 8 and 2 states originating from the $5s5p^2$ and $5s^25p$ configurations respectively. The calculated intensity ratio between $N_{4,5}O_1O_{2,3}$ and $N_{4,5}O_{2,3}O_{2,3}$ Auger groups is 9:1. In the case of Sb_4 , the number of the final states of the allowed Auger transitions was estimated to be the configurations that have at least one of the holes in the orbital with the same symmetry as the orbital having the hole in the initial state. The Sb_4 molecule has T_d symmetry, and in this symmetry group the d-orbitals in full rotational symmetry group transform as irreps a_1 , a_2 , and a_3 . The a_4 -orbital transforms as irreps a_4 and a_4 -orbitals as irreps a_4 , a_4 -orbitals and a_4 -orbitals as irreps a_4 , a_4 -orbitals as irreps a_4 , a_4 -orbitals and a_4 -orbitals as irreps a_4 , a_4 -orbitals and a_4 -orbitals as irreps a_4 , a_4 -orbitals and a_4 -orbitals and a_4 -orbitals as irreps a_4 , a_4 -orbitals as irreps a_4 , a_4 -orbitals and a_4 -orb

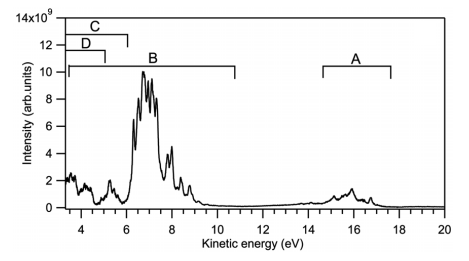


Figure 2. Background subtracted and transmission corrected AES. Labels A–D show the range of the transitions to the final states A = $5s^25p^7$, B = $5s^35p^2$, C = $5s^26s^3$, and D= $5s^25d^3$.

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XPS study of the electrode/electrolyte interface on silicon electrodes for Li-ion batteries and formation of the Li-Si alloy

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Metals and semimetals that can electrochemically form alloys with lithium are interesting alternative materials to replace the carbonaceous materials currently used as the negative electrode in lithium-ion batteries. New emerging applications require higher performance in terms of rechargeable capacity, energy density, power and safety. Silicon appears to be an ideal candidate for the next generation of negative electrodes. At room temperature it can alloy with 3.75 Li atoms per Si atom, resulting in a maximum capacity of 3579 mAh.g⁻¹ (compared to 372 mAh.g⁻¹ for graphite)¹. Silicon is also inexpensive and environmentally friendly.

During the 1st cycle, the surface of the negative electrode is passivated with a layer which is composed of the decomposition products of liquid electrolyte. The stability of this layer named SEI (Solid Electrolyte Interphase) is crucial for good battery performance. Electrode/electrolyte interface reactivity has been widely studied mainly for carbon electrodes, but less attention has been directed to the SEI formation in silicon electrodes, and to the understanding of the limitations due to the interface phenomena.

Non-destructive depth-resolved analysis by varying the energy of the X-ray beam instead of using destructive argon ion etching was used here to allow us to investigate the Li-Si alloying process and the passivation layer formation occurring upon lithiation/delithiation.

A step by step study of the 1st cycle of Si/Li electrochemical cells was performed at the I-411 beamline at the MAXIV laboratory in Lund. Soft X-ray was used to analyze the outermost surface of our system by selecting two fixed kinetics energies (variation of the excitation energies for each analyzed core peak); 130 and 610 eV. The kinetic energy is directly linked to the mean free path of the electrons which gives the depth of the analysis (Figure 2, a-b).

A similar study has been done with an in-lab XPS, Kratos Axis Ultra spectrometer using a focused monochromatized Al K α radiation ($h\nu$ = 1486.6 eV) and to complete the results obtained with the synchrotron radiation (MAX-lab) and AlK α radiation (in-lab XPS), experiments were carried out with hard X-ray ($h\nu$ = 2300 and 6900eV) at the KMC-1 beamline of the Helmoltz-Zentrum Berlin, Bessy II allowing a deeper analysis than could be obtained at the other used facilities. Figure 2, c shows the evolution of the Si2p peak of our starting electrode for the 5 photon energies. The evolution of the ratio [SiO₂(surface)/Si-Si(bulk)] is linked to the depth of the analysis.

This study allowed us to follow the formation of the Li-Si alloying process, step by step, upon the first lithiation thanks to the high energy photons. The variation of the analysis depth is probing the nature and composition of the SEI as a function of its thickness. We have determined that the main species of the SEI was formed during the 1st part of the 1st discharge and a stable composition and thickness seems to be reached after the 1st charge/discharge cycle.

Acknowledgements

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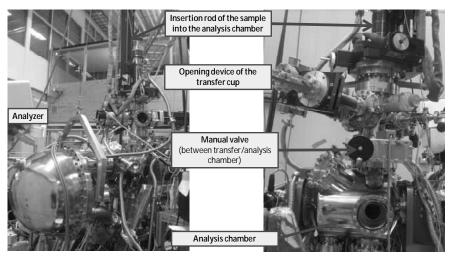


Figure 1: Experimental facilities used in MAXlab.

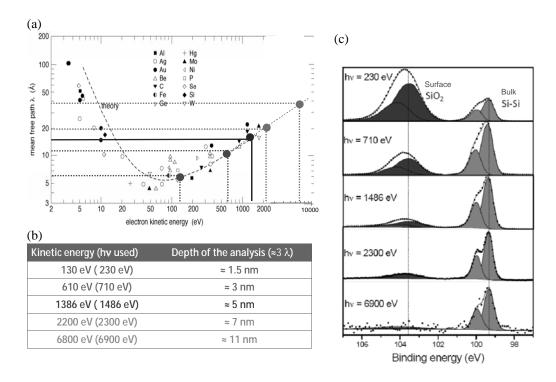


Figure 2: a, Mean free path of the electrons as a function of the kinetic energy.
b, Depth of the analysis for the Si2p peak for 5 photon energies
c, Evolution of the Si2p peak for this 5 photon energies (starting electrode). Surfacic SiO₂ and Si-Si from the bulk

Final electronic state dependence of fragmentation pathways in water molecules during the O $Is \rightarrow 4a_I$ resonant Auger decay

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Properties of water have been extensively studied for years using several experimental and theoretical methods since it is vital for all known forms of life. Being abundant and widely available, water has been considered as the most potential source for future energy systems based on H_2 combustion. This has aroused interest in understanding dissociation mechanisms of water molecule (see e.g. [1] and references therein).

The lowest unoccupied molecular orbital (LUMO) in water is an anti-bonding state, $4a_1$. A core-excitation of O $1s \rightarrow 4a_1$ is known to cause an ultra-fast dissociation with a neutral H and an excited O*H fragments before the molecule has time to undergo a resonant Auger decay [1,2]. We were interested in final state-dependent tendencies of fragmentation after core-excitation to the $4a_1$ state and wanted to study this with the help of electron-energy resolved photoelectron-photoion coincidence (PEPICO) measurements.

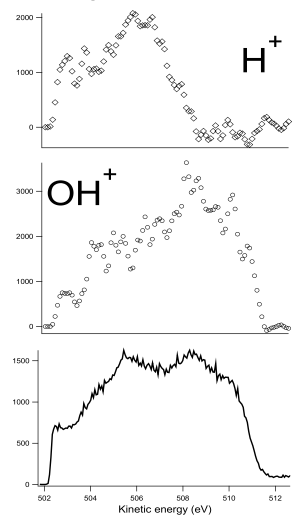
Experimental setup

Fragmentation details were investigated at the gas-phase beamline I411 on one-meter section using a photoelectron-photoion coincidence apparatus built in collaboration by Electron Spectroscopy group in University of Oulu and Material Sciences group in University of Turku. The kinetic energies of photoelectrons are analyzed using a modified Scienta SES-100 hemispherical electron analyzer with a resistive anode position-sensitive detector. The home-made Wiley-McLaren -type ion spectrometer is operated in the pulsed extraction field mode and the ion extraction pulse is triggered by either the arrival of an electron at the electron analyzer or a random signal, which is used in data analysis to subtract the random coincidences from the true ones. [3]

The sample was prepared by several freezing, pumping, and melting cycles. Pressure in the experimental chamber was approximately $2 \cdot 10^{-6}$ mbar. Photon energy of 534.0 eV corresponding to core-excitation O $1s \rightarrow 4a_1$ was used to define the intermediate electronic state in our experiment. Pass-energy of 100 eV was chosen in order to monitor a broad range of the resonant Auger spectrum, which is shown in the bottom panel of Fig. 1. The experimental resolution was not optimal as we tried to maximize the transmission of electrons instead.

Results and discussion

In the coincidence ion spectrum, we could hardly see any parent ions H₂O[>] in coincidence with the measured resonant Auger electrons. Earlier, these electronic states have been identified to correspond spectator decays from fragments [2]. Instead of H₂O[>] cations, the prominent fragment in the PEPICO spectrum are OH[>] and H[>] ions, whose electron-energy dependence (coincident ion



yield, CIY) is shown in Fig. 1 on top o the resonant Auger electron spectrum in the middle and upper panel, respectively We could also see traces from O[>] ions in the PEPICO map but the statistics fo these coincidences was not good enough to draw conclusions about their fina electronic state dependence.

According to the earlier study by Hjelta et al. [2], part of the resonant Auge electrons around 507 eV would complete from molecule and these should be observed in coincidence with paren ions. This we could not confirm in ou experiment. We could only see at enhanced production of H[>] cations a well as a weak hint of O[>] ions around this kinetic energy region.

On the other hand, our experiment agree well with their interpretation and calculations of resonant Auger spectrun originating from O⁵H species after

HO^5HBH+O^5-

pre-Auger dissociation in the kinetic energy region close to 510 eV.

Figure 1. Coincident ion yields for H⁵ OH⁵, and O⁵ fragments as a function o kinetic energy of resonant Auge electrons. The resonant Auger electror spectrum is shown in the bottom Intensities are in arbitrary units.

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Understanding the role of water at the ZnO/N719 interface

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Dye-sensitized solar cells are widely studied as an alternative to conventional solar cells. The most common semiconductor material is TiO_2 but others are also of interest, where ZnO is one of the best candidates. Earlier studies using Ru-dyes and ZnO show a decrease in solar cell performance with increasing immersing time of the ZnO electrodes in the dye bath for ruthenium dyes. ii,iii According to these studies, the dye sensitization process on ZnO electrodes entails a small amount of Zn^{2+} ions dissolving into the dye bath, with a subsequent aggregation of the dye and Zn^{2+} ions occurring. In new studies, the efficiency of solar cells based on the ZnO/N719 system has shown a dramatic increase when adding water in the sensitisation process. A detailed photoelectron spectroscopy study was therefore performed on these dye-sensitized ZnO electrodes in order to further understand the dependence of water for the dye aggregation on ZnO electrodes. The ruthenium dye N719 (cis-diisothiocyanato-bis(2,2'-bipyridyl-4,4'-dicarboxylato)-ruthenium(II)bis(tetrabutylammonium)) was studied. The dye was dissolved in ethanol and ethanol/water (1:1) mixture, and the sensitization were performed by dipping the oxide films into the dye solutions.

Table 1 shows the ratio between the Ru $3d_{5/2}$ and Zn 3p intensities after 15 h and 10 min of sensitization with N719, with and without including water to the dye solution. This ratio can be used to compare the amount of dye on the ZnO surfaces. As can be seen in the table, the ratio is much larger for the longer sensitization time if only ethanol is used as solvent. Thus, adding water to the dye solution, this surface induced aggregation mechanism is prevented. The coverage is similar to those obtained on TiO₂.

Detailed analysis of the nitrogen N Is and sulphur S 2p signals was made for the samples sensitized for 10 min. The nitrogen N Is spectra for N719 shows mainly three peaks corresponding to the nitrogen atoms in the thiocyanate ligands (N_{NCS}) at a binding energy of 397-398 eV, the bipyridyl ligand (N_{bpy}) at a binding energy of 400 eV, and the TBA⁺ counter ions (N_{TBA}) at a binding energy of 402.5 eV iv , see figure 1. Comparing the N Is spectra with and without water, some differences are observed. For example, the N_{TBA} decreases when water is included in the solvent. This indicates that less TBA⁺ ions are present. The same effect has previously been observed for similar measurements on TiO₂. v

The N_{NCS} structure is broad and shows clear signs of multiple contributions, a lower binding energy peak and a higher binding energy peak. This result is different from that obtained when adsorbing the molecule at a TiO_2 surface were one main peak is observed. Moreover, when measuring a multilayer of N719, there is only one contribution from N_{NCS} and the position of this peak, relative the bipyridyl peak is indicated with a line in figure 1 at about 398 eV. The multilayer N_{NCS} peak coincides well with the higher binding energy peak seen in this study, while the lower binding energy peak seen here, must arise as a result from the adsorption at the ZnO surface. This lower binding energy peak cannot be seen when N719 is adsorbed onto TiO_2 . Moreover, this spectroscopic result show that the molecular arrangement is different at ZnO compared to TiO_2 . Moreover, this difference is due to a strong interaction with the S atom as observed by a large shift (1 eV) in the N_{NCS} signal.

Sulphur S 2p spectra are shown in figure 2. The sulphur S 2p level has a spin orbit split of 1.18 eV with intensity ratio 1:2 between the S $2p_{1/2}$ and S $2p_{3/2}$ peak areas. Using this as fitting parameters, two chemically inequivalent sulphur features can be seen for both solvents. Since a single dye molecule contains only one chemical state of the sulphur atoms, the two states observed here indicate some specific interaction at the surface. The relative intensity of the peaks is different for samples the two samples. Interesting however, also for the samples sensitized in the presence of water, still the lower binding energy peak has almost 50% of the total intensity.

Thus together the results reported here show that water assist to form close to monolayer coverage, but that the surface organisation of these layers is different from those obtained at TiO_2 surfaces. At ZnO a large fraction of the molecules interact not only via the carboxylic groups but also include some specific interaction via the sulphur atoms. Still, however, also these molecules will efficiently convert visible light into current as observed from light to current conversion measurements having a maximum of about 40% (not shown).

Table 1. Ratio between Ru $3d_{5/2}$ and Zn 3p intensities (Ru/Zn) measured with a photon energy of 758 eV.

Sample	15 h	10 min
N719 H ₂ O	0.19	0.10
N719 no H ₂ O	1.72	0.13

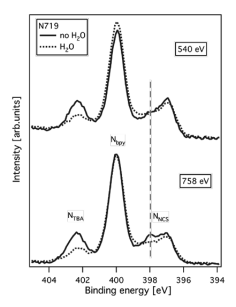


Figure 1. N *Is* spectra of N719 on ZnO measured with photon energies 540 and 758 eV respectively. The dashed line indicating the position of the N_{NCS} peak in a multilayer of the dye.

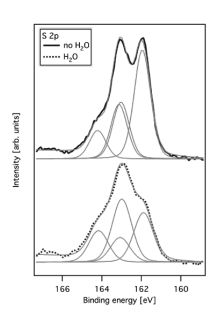


Figure 2. S 2p spectra measured using photon energy 454 eV.

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Insulator to metal transition in Sb clusters studied using core-level and valence photoelectron spectroscopy

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The transition from large band gap insulators to metallic properties as a function of size has been studied for clusters of Group VA element antimony (Sb) using core-level and valence photoelectron spectroscopy. The experiments were performed at the undulator beam line I411 with two different cluster sources: the exchange metal cluster source (EXMEC) built in collaboration between the universities of Oulu and Uppsala [1] and a gas aggregation source (GAS) built in Turku. For the experiments with the EXMEC source the permanent Scienta R4000 electron analyzer was used and for the experiments with the GAS a modified Scienta SES-100 electron analyzer [2,3] was employed. In both the EXMEC and GAS experiments the core-level and valence photoelectron spectra were recorded at identical conditions of the respective cluster source. In the EXMEC experiments the size of the clusters was controlled by varying the temperature of the resistive heated evaporation oven. The 4d core-level photoelectron spectra with the parent Sb₄ clusters are presented in Fig. 1.

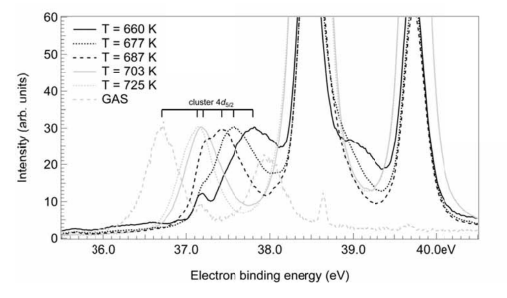
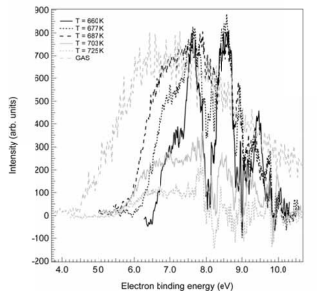


Figure 1. The 4d core-level photoelectron spectra of various sized Sb clusters. The large lines reaching over the intensity scale are from the parent Sb_4 clusters of which the antimony vapor consists.

The valence photoelectron spectra, with the well known contribution subtracted away, thus consisting only of the signal from the larger clusters, are presented in Fig. 2. The size of the clusters was estimated using the 4d photoelectron spectra using the Born-Haber cycle approach. The average size range using this method was determined to be from < N > = 17 to < N > = 102 for the EXMEC source, with the lowest temperature corresponding to the smallest size, and <N>=850 for the GAS. The smallest sizes show a clear discrete line structure in the valence photoelectron spectrum. which gradually disappears as the size of the clusters becomes larger. The apparent discrete structure in the



izes with

two largest sizes in the EXMEC experiments is most likely due to the errors in the subtraction process of the Sb_4 components. The ionization potentials that are obtained for the smallest sizes can be compared with the work of Rayane et al. [4] and show indeed a large deviation (up to 2 eV) from the ionization potential values obtained from the metallic sphere approximation. As the size of the cluster grows, the ionization potential approaches the metallic sphere case indicating that the transition from an insulator to metal takes place in clusters consisting of approximately $100 \ Sb$ atoms.

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Is there a Simple Relationship between C1s Ionization Energies and Hybridization?

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Introduction

When assigning spectra of adsorbed states, it is frequently assumed that carbon 1s (C1s) ionization energies group according to the hybridization state of the carbon atom and, moreover, that saturated (sp³ hybridized) carbons have lower ionization energies than do unsaturated (sp², sp hybridized) carbons.¹ We argue that this is an unjustified assumption. Here, we present the analysis of the gas-phase photoelectron spectrum of cyclopentene as an example. The results are backed by a large number of accurately measured C1s ionization energies.

Experimental and Results

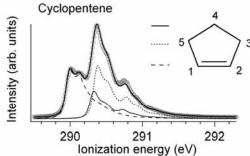


Figure 1. C1s photoelectron spectrum (circles) of cyclopentene. Fit of the spectrum is prepared as a sum (solid line) of three atom-specific theoretical lineshapes, one for each inequivalent carbon atom (broken lines).

The C1s photoelectron spectrum of gaseous cyclopentene was measured at beamline I411 at an instrumental resolution of 72 meV and a photon energy of 332 eV, c.f. Fig 1. The spectrum was fitted using atom-specific theoretical lineshapes. Since the two sp² carbons (C1 and C2) are equivalent and also C3 and C5, cyclopentene has three inequivalent carbon atoms. The agreement between the fit and the experimental spectrum leaves no doubt about the assignment.

The measured adiabatic ionization energies are 289.988 eV (C1,C2), 290.372 eV (C3,C5) and 290.318 eV (C4). ²

Our interpretation of the spectrum is in strong contradiction to how several authors interpret the photoelectron spectrum of cyclopentene physisorbed on various semiconducting surfaces 1,3,4 . As stated above, it seems common to assume that C1s ionization energies group according to hybridization, and moreover, that unsaturated (sp², sp) carbons have higher ionization energies than do saturated (sp³). This assumption seems to have been motivated by gas-phase measurements of the three C_2 hydrocarbons, namely ethane (H_3C-CH_3), ethene ($H_2C=CH_2$) and ethyne ($HC=CH_3$).

Since physisorption leads only to a small perturbation of the molecule, there is no reason to believe that this would radically change the internal positions of the ionization energies measured in the gas phase. Hence, it is relevant to look to high-resolution gas phase ionization energies when assigning physisorption spectra.

Firstly, albeit the ionization energies of the two sp³-carbons of cyclopentene have similar ionization energies, this is not a rule applicable for hydrocarbons in general. E. g. in propene ⁵ (H₂C=CH-CH₃), the central sp² carbon and the sp³ carbon have very similar ionization energies, while the terminal sp² carbon has about 0.5 eV lower energy. Secondly, in both cyclopentene and propene (and also many other unsaturated hydrocarbons), the lowest ionization energy within the molecule is found to be that of an unsaturated carbon atom.

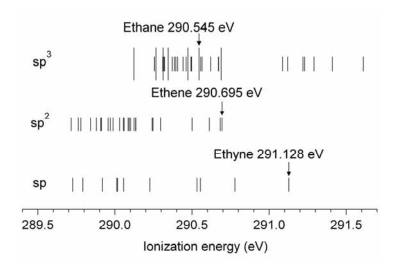


Figure 2. C1s ionization energies for 25 hydrocarbons. The ionization energies are grouped according to the hybridization of the ionized carbon. Long bars represent saturated compounds, i.e. compounds having only sp³ carbons.

Fig. 2 shows how C1s ionization energies of 25 hydrocarbons group according to the hybridization of the ionized carbon atom. The compounds included are methane, ethane, ethene, ethyne, propane, propene, propyne, butane, 1,3-butadiene, 1-butyne, 2-butyne, cyclopentene, 1-pentyne, 2-pentyne, 1,3-pentadiene, trans-3-hexene, 3-hexyne, cyclohexane, cyclohexene, 1,3-cyclohexadiene, 1,4-cyclohexadiene, benzene, toluene, 1,3-dimethylbenzene (m-xylene), and 1,4-dimethylbenzene (p-xylene). For each hybridization state, we see that the ionization energies are distributed over a fairly wide energy range (1-1.5 eV) which largely is overlapping with the energy regions for other hybridization states. Evidently, the choice of ethane, ethene and ethyne as model compounds is not justified since their ionization energies are not typical for their hybridization states. In fact, both ethene and ethyne have the highest ionization energy within their respective groups. A more thorough discussion of this topic can be found in Ref. 2.

Conclusions

Photoelectron spectra of hydrocarbons physisorbed to semiconducting surfaces cannot be assigned using simple rules based on the hybridization of the ionized carbon. However, since physisorption only to a small extent will influence relative shifts, one possible solution is to use ionization energies obtained in the gas phase for the very same compound as a tool in the assignment.

Acknowledgments

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Size dependent properties and interactions in free nanoscale potassium halide clusters as seen by XPS

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Geometric structure and chemical properties of alkali halide clusters have been extensively investigated by various methods, also because these clusters are good models for studying ionic bonding in compounds. Mass spectroscopic results suggest that this kind of clusters is formed with a similar crystalline structure as their solid counterpart and shaped as cubes or cuboids. Photoelectron spectroscopy is an established tool to directly investigate the evolution of

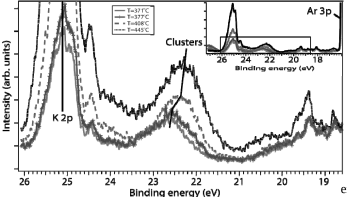


Figure 1. K 2p core level photoelectron spectra of varied size KCl clusters recorded at 70eV photon energy.

formation of clusters before their fragmentation. Core level photoelectron spectroscopy of clusters provides insight into the interactions inside clusters revealing the information on the size dependent bonding mechanism.

The EXMEC cluster source in a junction with an adiabatic-expansion argon-cluster source has succeeded earlier in producing the metal clusters,

and in the present work it was

employed to produce the ionic-bonding clusters out of alkali-halide salts. A detailed description of the EXMEC-source is given in Ref [1]. All the experimental apparatuses were mounted at the permanent end-station of the I411 soft x-ray undulator beamline at Max-lab. The core level photoelectron spectra were recorded using Scienta R4000 hemispherical analyzer fixed at 90 degrees to the polarization plane of the beamline radiation. By changing the temperature of the crucible with the solid salt inside, different sizes of salt clusters were produced, and K 2p spectra were recorded at 70eV photon energy for each cluster-size produced (Fig.1). Figure 1 shows the K 2p monomer signal at around 25eV binding energy with two spin-orbit components overlapped. At four different clustering conditions, as the crucible was heated up from ≈370°C to

445°C, unambiguous cluster peaks were obtained at 2.5 to 2.8 eV lower binding energy relative to the monomer.

The K 2p spectra of KI clusters were also recorded using the same setup but in a different series of experiments. The difference in binding energies of K 2p between KCl and KI monomer is only 0.2eV, which matches very well the experimental results obtained by A. W. Potts in 1970s [2]. Inspection of figure 2 discloses the difference in the monomer-cluster shift between KI and KCl of around 0.8eV.

In our former studies on NaCl clusters [3], an ionic model was employed for interpreting the big binding energy shift (from monomers to clusters) both at the metal and halogen sites. The calculated values have satisfactorily matched the experimental results. It has been found that the Coulomb interaction and polarization interaction are vital contributions to the shift. When the clusters became bigger, the steps of shifts became smaller. This phenomenon has also been found for the other kinds of clusters like rare gas clusters and metal clusters. From the experimental and model studies on the electronic structures of Na and K halide clusters, it becomes clear that the interatomic distance and polarizability of halogen ions do affect substantially the energy structure of the clusters.

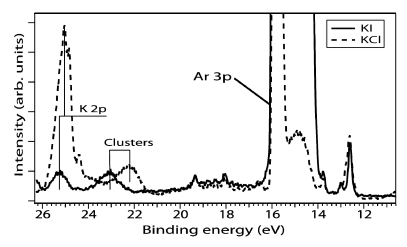


Figure 2. K 2p core level photoelectron spectra of KCl and KI clusters recorded at 70eV and 40eV respectively.

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Solvent-mediated inter-molecular Coster-Kronig decays in aqueous ions

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The degree of association between a cation and solvent molecules in a solution can be probed in various ways. One characteristic that conceivably could be affected by chemical interaction between the ion and the solvent, and provide an experimental observable, is the lifetime of a vacancy in the outermost core orbital, which decays primarily via radiationless Auger decay, in which the core hole is filled by one outer electron while another outer electron is emitted. The influence of the surrounding on the valence electrons can affect the rate of core-valence-valence Auger decays, which indeed has been observed in free small molecules [1]. An extreme situation occurs when all the valence electrons of a solvated cation has been donated to the counterion, which in principle would prevent an Auger decay from the outermost core level. Auger-like decays involving orbitals from the surrounding solvent molecules will still be possible, however. An example of such a decay has been found in aqueous solutions of KCl, where Auger signal from final states involving orbitals on water molecules surrounding the \hat{K}^+ ion has been observed after ionization of the $L_{2,3}$ edges [2]. In the case of the aqueous ions Na⁺, Mg²⁺, and Al³⁺, the 3s and 3p orbitals are usually considered empty, and the 2s and 2p levels are the outermost ones. In the corresponding metals, which can be described as consisting of such ions surrounded by delocalized valence electrons in s- and p-bands, a 2p vacancy will predominantly relax by Auger decay. A 2s hole predominantly relaxes by Coster-Kronig decay, which is much faster than normal Auger decay. For the corresponding free ions, these decays are forbidden, due to the lack of available valence electrons to fill the 2p hole, and for energy reasons for the 2s vacancy. One can compare to the situation in solid sodium halides, where it is known that interatomic Auger and Coster-Kronig processes occur after Na 2p and 2s ionization, respectively, and the electrons that take part in the relaxation originate from the neighboring halide ions [3]. This could also be expected for crystalline magnesium and aluminum salts, but in a solution, where the interaction between the cation and the solvent molecules is weaker than in an ionic compound, the situation is by no means self evident. We have therefore made an investigation of the widths of the cation 2s and 2p photoelectron lines from solutions of NaCl, MgCl₂, and AlCl₃, using a liquid micro-jet setup, to see whether a difference in lifetime related to the different ion-solvent interaction of the three differently charged ions can be observed [4].

The measurements were performed at the undulator beamline I411. Aqueous solutions of sodium chloride, magnesium chloride, and aluminum trichloride were prepared by mixing commercially obtained salts with de-ionized water. The concentrations were 3.0, 3.0, and 0.5 m (mol / kg solvent) for NaCl, MgCl₂, and AlCl₃, respectively. The experimental setup used has been described in detail in Ref. [5]. Briefly, the sample was introduced into vacuum as a jet formed by a nozzle with a diameter of $15~\mu m$, backed by a HPLC pump. The jet was injected into a differentially pumped compartment, with a 1 mm diameter hole to allow the synchrotron radiation into the ionization region, and with a skimmer with a 0.5 mm diameter opening towards the electron spectrometer. After passing the soft X-ray radiation, the liquid was collected by a liquid nitrogen-cooled trap. The measurements were performed at a distance of 2-3 mm downstream from the nozzle, at a point before the jet spontaneously breaks into a train of droplets. The jet propagation axis was perpendicular to the exciting radiation and to the detection axis of the spectrometer. The spectrometer was mounted with the lens axis at an angle of 54.7° relative to the plane of polarization from the undulator radiation, to minimize angular distribution effects.

In Fig. 1, example photoelectron spectra of the Mg 2p and 2s levels from an aqueous solution of MgCl₂ are shown A direct comparison shows that the 2s line is significantly broader than the 2p feature, even though the latter is split by the spin-orbit interaction in the final state. This was also the case for Na and Al from aqueous NaCl and AlCl₃ [4]. We have attempted to determine the lifetime of the 2p and 2s vacancies from the spectral widths using curve fitting of symmetric Voigt profiles. In our fittings, we have fixed the spin-orbit splitting to values for metallic sodium, magnesium and aluminium: 0.169, 0.28 and 0.40 eV, respectively [6, 7]. We have assumed a zero contribution to the Lorentzian width for the 2p lines, and under this assumption, the Gaussian widths in the fitting of the 2p spectra will be maximized. Using the same Gaussian widths when fitting the 2s spectra will then give a minimum for the lifetime width of the 2s vacancies. From such fits, we obtain Lorentzian widths Γ_{2s} of 0.21, 0.45, and 0.67 eV for the 2s lines of aqueous sodium, magnesium and aluminum ions, respectively. These lifetime widths show the same trend and are actually quite similar to those found in the metallic cases, 0.28, 0.46, and 0.78 eV, respectively [6, 7].

The reason for the larger lifetime width of the 2s compared to the 2p level in the metals is the fast Coster-Kronig decay of the 2s hole, involving a 2p electron filling the 2s vacancy, and a 3s or 3p electron being ejected, on a time scale in the femtosecond range. In free ions, this decay would be impossible, as there are no 3s or 3p electrons to carry away the excess energy. A super-Coster-Kronig decay, involving two 2p electrons, one filling the 2s core hole and one being ejected, is impossible for energetic reasons in the metals as well as for the free or solvated ions. The

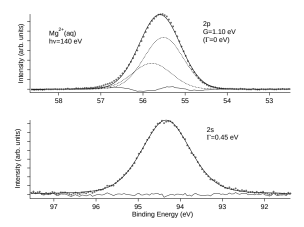


FIG. 1: Photoelectron spectra of the 2p (upper) and 2s (lower) levels from an aqueous solution of MgCl₂. The solid lines (black) correspond to the total intensities of the fits, and the dot-dashed lines (blue) represent the residues. The data points are represented by filled circles (red). A linear background has been subtracted in the spectra. In the 2p spectrum, the dashed lines (black) represent the Voigt profiles of the 2p spin-orbit components. Γ and G refer to the Lorentzian lifetime width of the lines and the corresponding Gaussian width obtained from curve fitting. The scale on the vertical axis differs between the spectra, but the scale of the energy axis is the same.

most obvious explanation of the Lorentzian width of the 2s lines for the solvated ions is that an intermolecular Coster-Kronig process takes place, where instead of the 3s or 3p electron, an electron from the surrounding water molecules or an associated anion is emitted after the 2s vacancy has been filled by an electron from the 2p shell. This process is similar to ICD, in which a vacancy in an inner valence shell of an atom or molecule in a weakly bound complex is filled by an outer valence electron, while a valence electron from a neighboring site in the complex is emitted into the continuum [8].

The observed widths correspond to lifetimes of the $(2s)^{-1}$ inner valence state of 3.1, 1.5 and 0.98 fs for Na⁺, Mg²⁺, and Al³⁺ respectively. The decrease of the lifetime from Na⁺ to Al³⁺ correlates well with the increased ion-water interaction. This interaction is dominated by the first solvation shell, which contains approximately the same amount of water molecules (5.4 for Na⁺, 6 for Mg²⁺ and Al³⁺ [9], but the ion-water distance decreases with increasing ionic charge (2.4 Å for Na⁺, 2.1 Å for Mg²⁺, and 1.9 Å for Al³⁺ [9]). First of all, a smaller ion-water distance means a larger overlap between the ion and water orbitals, which would facilitate this ICD-like Coster-Kronig process. Secondly, the solvent polarization induced by the positive ions, where the increased charge in the core-ionized intermediate state should be considered, increases the electron density in the vicinity of the ion with the core hole. Since the degree of water polarization in the first solvation shell is very much dependent on the charge of the ion the discussed ICD-like decay processes should be more efficient for the more highly charged ions, just as observed. These polarization effects, and, particularly for the case of Al³⁺, the possibility of hybridization between the outer water valence orbitals and those of the ions, are thus certainly factors that determine the decay transition rates. To make quantitative estimates of the transition rates to gain insight into these phenomena would most likely require a very complex and most demanding theoretical treatment, but would be an exciting issue to investigate further.

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Charge transfer and chemical interaction of ZnPc and FePc

on Au(111) and Au(111)-I

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The dye-sensitized solar cell, also known as the Grätzel cell contains many components whose mutual interaction determines transfer of photo-excited electrons, current and voltage from the cell, and thus efficiency of the cell. The detailed interaction and energy level alignment at each interface will affect the output from the cell.

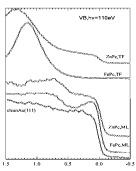
Phthalocyanines have been widely used in photo-electronic devices and are potential candidates for solar cell applications. In this work we have studied two different phthalocyanines; FePc and ZnPc adsorbed on Au(111) and iodated Au(111).

Iodine modified Au is used to represent the cathodic part of the solar cell [1]. Two different surface structures occur on Au(111)-I, a ($\sqrt{3}x\sqrt{3}$) reconstruction at 1/3 monolayer coverage and a ($\sqrt{7}x\sqrt{7}$)R19° reconstruction just below ½ monolayer coverage.

The charge transfer across the weakly interacting gold-organic interface (both magnitude and direction) was shown to depend on the d-filling of the metallic center (Co, Ni, Cu) in the molecule [2]. We have extended their study to include ZnPc and FePc monolayers on Au(111). The presence of iodine at the interface modifies the bonding and charge transfer situation. This depends both on the metal center atom and the iodine reconstruction.

At monolayer coverage there is a substrate mediated interaction between phthalocyanine molecules and Au substrate which is mixture of molecular and metal states.

In figure 1, spectra from the "near Fermi level" region are shown. MPc induced structures are clearly seen whose energy distribution depends on the central metal atom.



BindingEnergy(eV)

Figure 1. Photoemission spectra (hv=110eV)for clean
Au(111),FePc/ZnPc on Au(111), monolayer and thick film

The structures at about 0.8eV for monolayer (ML) spectra are due to the interface HOMO which possesses the highest intensity at 1 ML. At higher coverage these interface structures decrease in intensity while the bulk like molecular HOMO appears at higher binding.

In monolayer of FePc we also observe a state close to Fermi level at about 0.15 eV. This state decreases at higher coverage after ML and disappears in thick molecular film. It probably is related to central metal atom states interacting with Au. Being very close to the Fermi level, this state is said to be due to charge transfer from metal states to formerly LUMO.

This state is absent in the ML film of ZnPc which has filled 3d metal orbitals in the centre. This is in agreement with literature where this state exists for FePc and CoPc but disappears by increasing the number of electrons in 3d-metal atom [3].

Figure 2a and b present valence band spectra of one monolayer of FePc and ZnPc on Au(111), Au-I $(\sqrt{7}x\sqrt{7})R19^{\circ}$ and Au-I $(\sqrt{3}x\sqrt{3})$. Iodine in the $(\sqrt{7}x\sqrt{7})R19^{\circ}$ structure reacts with Zn in ZnPc and oxidize the molecule while this reaction is not visible on the $(\sqrt{3}x\sqrt{3})$ structure. The chemical interaction between iodine and FePc is weaker. The different interactions affect the energy level alignment, the HOMO position with respect to the Fermi level are different.

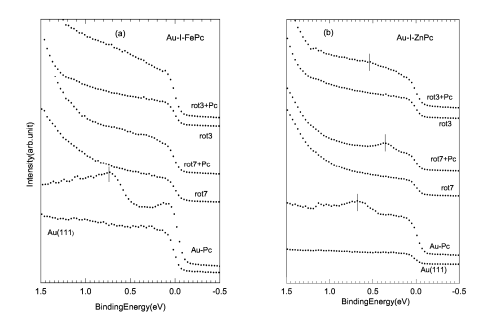


Figure 2. Photoemission spectra (hv=110eV) for monolayer of a)FePc and b) ZnPc on clean Au(111), Au-I ($\sqrt{7}x\sqrt{7}$)R19 and Au-I ($\sqrt{8}x\sqrt{3}$)

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Electronic Properties of Lutetium Phthalocyanine Thin Films

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In a preoccupation with a reduction of the production costs as well as to decrease the environmental impact during manufacture; many hybrid devices containing active macromolecules, such as phthalocyanine, made their appearance since the year 2000. These hybrid devices have real potentialities of development in industrial sectors such as environmental protection ^{1,2}. Lutetium phthalocyanine (LuPc₂) molecules is of interest for gas sensors based on field effect transistor system and related devices, where these molecules are used as an active medium sensitive to polutant exposures (NO₂, NH₃). However the processes that govern the

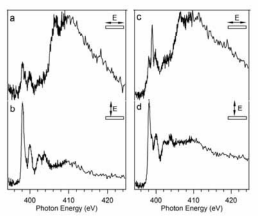


Figure 1 Polarization dependent N1s NEXAFS on thin layer of LuPc2 adsorbed on vicinal Si(001)-(2x1) surfaces. Spectra on Si-H (left, a,b) and clean Si (right, c,d) at two different sample positions with respect to the polarization of the incoming light which are shown by arrows.

reactivity are not well understood yet, motivating the interest to investigate these processes.

In this experiment we examined the organization of LuPc2 on one of the technologically most largly used electronic's base materials - namelly Si(100) reconstructed 2x1 substrate. We compared the organization of LuPc2 on clean silicon (clean Si) substrate and on a passivated Si (Si-H) substrate by means of fully polarized Near Egde Absorption Fine Structure (NEXFAS) carried out at the nitrogen K edge. The Si(100) reconstructed 2x1 surface was performed by removing the native oxyde layer by flashing the substrate to 1050°C for 15s. The passivation of Si substrate was performed by exposing the reconstructed 2x1 silicon surface to atomic hydrogen while keeping the sample at 300°C. During that process the atomic hydrogen

bonds to Si dangling bonds making the substrate surface less reactive and therefore allowing the self-organization of the adsorbed molecules. The evaporation of LuPc₂ was performed *in situ* under ultra high vacuum conditions and varying the thickness of the evaporated layer from few layers to multi-layer sample.

We performed Near Edge X-ray Absorption Fine structure (NEXAFS) measurements in two geometries with respect to the polarization of the incoming light – paralel to the substrate and perpendicular to the substrate; allowing us to get information on organization of the molecules on

the surfaces.

The graphs shown in figure1 represent NEXAFS spectra for thin layer of LuPc₂ evaporated on clean Si (c,d) and on Si-H(a,b) for the two geometries. All the spectra show sharp peaks at lower energy which correspond to pi transitions perpendicular to the plane whereas at high energy broad features represent sigma transitions happening in the plane of the molecule. On both substrates an intense sigma transition is observed in geometry when the light polarization is in the

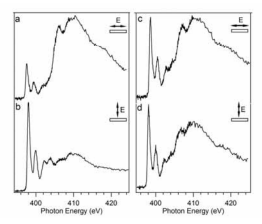


Figure 2 Polarization dependent N1s NEXAFS on thick layer of LuPc2 adsorbed on vicinal Si(001)-(2x1) surfaces. Spectra on Si-H (left, a,b) and clean Si (right, c,d) at two different sample positions with respect to the polarization of the incoming light which are shown by arrows.

surface plane, together with weak signal from pi transitions; opposite behavior is seen for out of plane polarization indicating a rather flat orientation of the molecules on both surfaces One can notice an additional feature at 399eV for graph c on clean Si which is not present in the same thickness sample on Si-H. The same feature is slightly observed also when the light polarization is perpendicular to sample surface. This clearly indicates the presence of a Si-N bond when LuPc2 is deposited on the clean surface whereas to bound are present on the hydrogenated surface. Additionally in figure2 where we

compare the two substrates at large thicknesses one can observe that the additional peak at 399 eV is not present any more, corroborating that this contribution comes from the interactions between the first layer of molecules and the substrate.

Now if one compares the spectra from samples with thick layer of $LuPc_2$ (figure2) one can observe that spectra for both orientations on clean Si (figure2, c,d) have practically the same structure which gives us an indications that the molecules are oriented in an angle close to 45° with respect to substrate. However the case for Si-H (figure2, a,b) is slightly different where they seem to be more lying down with two deckers parallel to substrate. However for more precise understanding of orientation shift with increasing thickness additional experiments are needed.

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The grafting of styrene on water-saturated Si(001)-2×1: Evidences for a radical chain reaction induced by isolated silicon dangling bond

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Recent experiments revealed that the Si(001)–2×1 surface modified by exposure to water vapor can be used as a reactive surface for the attachment of organic molecules such as carboxylic acidsⁱ and ethoxysilanes.ⁱⁱ In a preceding work combining x-ray photoelectron spectroscopy (XPS) and scanning tunneling microscopy (STM), we reported on the nature of the silicon dangling bonds, left after saturation of the surface by water.ⁱⁱⁱ STM shows that these defects, whose density is in the range

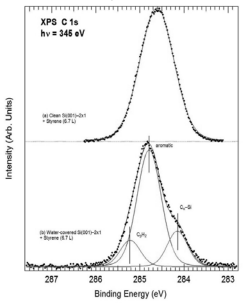


Figure 1: XPS C 1s spectra measured at hv = 345 eV. The takeoff angle of the electrons, with respect to the surface normal, is 45°. The experimental curves (dots) are fitted with one or more Voigt components (solid lines). (a) Clean Si(001)–2×1 exposed to 6.7 L of styrene at room temperature. (b) Water-terminated Si(001)–2×1 surface exposed to 6.7 L of styrene at room temperature.

 1.2×10^{-2} - 1.7×10^{-2} defects/Si atom are isolated (hence the term isolated dangling bond, IDB). Are those IDB reactive? Does an alkene react with this surface? To clarify this effect, we chose to study the adsorption of styrene with the silicon water covered surface.

The high resolution C 1s spectrum, performed on I511-1 (surface) (hv = 345eV) of the water-covered exposed 6.7 L styrene $(H_2C_\alpha=C_\beta H-C_6H_5)$ is reported in figure 1 (b). The spectrum is fitted with three Voigt components, positioned at 284.22, 284.79 and 285.16 eV, respectively (18, 69 and 13% of the spectral weight, respectively). There is evidence in the experimental literature for a chemical shift difference of 0.22 eV in polystyrene between aromatic carbons (binding energy of 284.77 eV) and aliphatic carbons (284.99 eV). iv This is supported by a recent DFT Δ SCF calculation of the ionization energies of ethylbenzene v. Therefore we can attribute the main peak at 284.79 eV, and its satellite at 285.22 eV, to the aromatic carbons and the β

carbon ($C_\beta H_2$ unit), respectively. On the other hand, we attribute the lowest binding energy component at 284.15 eV to the α carbon, bonded to silicon. In fact, the binding energy of Si–CH₃ units resulting from the dissociation of methyliodide on Si(001)–2×1 is found at 284.10 eV. So the C 1s spectrum is consistently interpreted assuming that the β carbon abstracts one H to give a $C_\beta H_2$ unit. We can also definitely eliminate the possibility of an OH abstraction by the β carbon and the formation of a monohydroxyl carbon, that should be strongly shifted to higher binding energy than the "phenyl" main peak (284.79 eV) and its $C_\beta H_2$ satellite (285.22 eV). The addition of the molecule to a silanol group, leading to a Si–O–C linkage, is also excluded as we do not see any C 1s component at ~286.5 eV ii. The Si 2p spectra of the water-covered surface (measured in surface sensitive conditions at hv = 150 eV, on I511-1), before and after exposure to 6.7 L of styrene, are given in figure 2 (a) and (b), respectively. The Si 2p spectrum of the water-covered surface exposed to 6.7 L of styrene, shown in figure 2 (b), is practically identical to figure 2 (a), apart from a small rigid shift of 0.03 eV towards

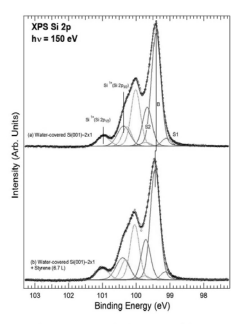


Figure 2: XPS Si 2p spectra measured at hv = 150 eV of the water-covered doped Si(001)−2×1 surface (1 L of water) before (a) and after (b) exposure to 6.7 L of styrene. The takeoff angle of the electrons, with respect to the surface normal, is 45°. The experimental curves (dots) are fitted with sums of Voigt components. The Si 2p_{3/2} (Si 2p_{1/2}) fitting components are the solid (dotted) lines

higher binding energy. We do not see the appearance of a distinct line related to the formation of a Si-C bond. This is not surprising as the Si-C component has a SCLS of +0.26 eV. The Si¹⁺ (SiOH) peak weight does not vary appreciably (it remains at ~10% of the spectral weight), which is consistent with the C 1s spectrum indicating that surfaces hydroxyls are not abstracted. The main information provided by the Si 2p spectra concerns the surface band bending, directly related to the areal density of surface acceptor states in the case of the water-covered n^+ -doped $Si(001)-2\times1$ surface. iii The bands are bent upward by 0.45 eV in regards of the clean surface. The above-mentioned rigid shift in binding energy indicates that the band bending slightly diminishes (by 0.03 eV) after styrene absorption. This means that the overall negative surface charge, viii and hence the areal density of negatively charged IDB, is only slightly decreased (by ~5%) after exposure to 6.7 L styrene.

Stimulated by recent observations (Ref. iii) that isolated dangling bonds have an analogous structure on H-terminated

Si(001)–2×1 and water-covered Si(001)–2×1 surfaces, we have examined if chain reaction schemes may also induce the grafting of alkenes on the latter surface. Choosing styrene as a test molecule, and core-level photoemission as a spectroscopic tool, we have observed that the molecule does react with the water-covered n^+ -doped Si(001)–2×1 surface at room temperature. The C 1s spectrum of the resulting product is significantly different from that of the adduct di- σ bonded to a silicon dimer of the clean surface. In particular, it exhibits a clear, low binding energy component, characteristic of the Si–C bond. The C 1s spectral shape also excludes the formation of a C–OH bond (OH abstraction by the radical adduct) or of a Si–O–C bond (OH addition to a surface silanol). Therefore we have gathered evidences that styrene makes a *single* σ bond with a silicon surface atom via the α -carbon, and that the β -carbon captures a nearby H atom, as it is the case for defective H–terminated surfaces. Due to the intensity of the Si–C component in the C 1s spectrum, the oligomerization of the molecule via a chain reaction is also excluded.

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Bi-isonicotinic acid on rutile TiO₂ (110): resonant inelastic x-ray spectroscopy

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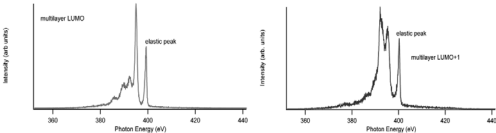


Figure 1. X-ray emission spectra for incident energy of 399.18eV (N 1s to LUMO transition) on the left and for 400.27eV (N 1s to LUMO+1 transition) on the right for a multilayer of bi-isonicotinic acid on TiO₂(110).

Bi-isonicotinic acid is an important molecule as it forms ligand groups through which many important dye molecules, such as N3, anchor to surfaces found in dye-sensitized solar cells (DSSC). One type of DSSC involves the dye molecule N3 adsorbed onto a TiO_2 surface [1]. In this type of DSSC, incoming photons promote the electrons from the highest occupied molecular orbital (HOMO) to the lowest unoccupied orbital (LUMO) of the dye and the excited electrons tunnel into the TiO_2 [1]. The electrons are resupplied to the HOMO via an electrolyte solution. The charge transfer between potential ligand groups, such as bi-isonicotinic acid, and the TiO_2 is very important to determine the theoretical efficiency of such devices.

The experiment was carried out on the bulk station of beamline I511-3 to which we simply added a simple argon gas-line for sputtering the sample clean between surface preparations. The 10mm square TiO₂ (110) sample was held onto a circular sample holder by tantalum wire at the corners. After cleaning the sample, bi-isonicotinic acid was deposited onto the sample using a Knudsen cell type evaporator at a temperature of 230°C. For multilayer coverages the sample was held at room temperature during the deposition, while for a monolayer coverage the sample was heated to 200°C to prevent multilayer growth [2].

Previous investigations of the charge transfer interaction between adsorbed biisonicotinic acid and a rutile $TiO_2(110)$ surface employed the core-hole clock implementation of resonant photoemission (RPES) combined with x-ray adsorption spectroscopy (XAS) to clock the average timescale for injection from the molecule into the conduction band of the substrate at less than 3 femtoseconds (fs) [3]. In this study we set out to see if we could use an analogous approach with resonant inelastic x-ray scattering (RIXS) to probe this ultrafast charge injection from different unoccupied molecular orbitals of the molecule. The technique relies on the fact that x-rays will be emitted from the molecule due to the transition of electrons into the core holes left by

electrons excited to the unoccupied molecular orbitals. Previous studies have firmly established that in the core-excited state the LUMO of the molecule here lies below the conduction band and therefore cannot take part in the tunnelling process to the substrate as it lies energetically within the band gap. The LUMO+1 however, overlaps with the conduction band and is the level from which we are expecting ultrafast charge transfer to occur. We used x-ray absorption spectroscopy to determine the position of the LUMO and LUMO+1 of the molecule and measured the RIXS at these two energies for both the multiayer and monolayer of bi-iso on the surface. The spectra were calibrated using the positions of several elastic peaks.

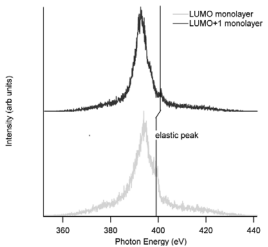


Figure 2. X-ray emission spectra for N 1s to LUMO transition (top) and N 1s to LUMO+1 transition (bottom) photon energies for the monolayer of bi-isonicotinic acid on $TiO_2(110)$. The solid line shows the expected position of the elastic peaks.

Figure 1 shows the RIXS for the multilayer showing clearly the elastic peak for both the photon energies of the LUMO and LUMO+1 transitions from the N 1s core level. The elastic peak represents electrons that have been promoted to the LUMO (or LUMO+1 depending on the incident photon energy) and consequently fell straight back into the core hole that they created and so losing no energy in the process. Unlike the multilayer case, the monolayer spectra in figure 2 showed a much larger elastic peak for the LUMO transition than for the LUMO+1. The most likely explanation for this large reduction is that the electron which has been promoted to the LUMO+1 has consequently tunnelled into the TiO₂(110) substrate from the LUMO+1 and so can not drop back into the core state. We are currently investigating these effects on larger dye molecules such as organometallic dye complexes which have the bi-isonicotinic acid binding ligand in order to develop of photons-in-photons-out implementation of the core-hole clock applicable to the study of dye-sensitised solar cells.

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The valence band electronic structure of the distorted rutile-like WO_2 – a study of chemical bonding via resonant soft x-ray emission at the O K edge.

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Rutile and rutile-like transition metal dioxide MO_2 compounds are interesting from both fundamental and technological viewpoints, encompassing TiO_2 and SnO_2 (wide bandgap semiconductor oxides suitable for transition metal doping to obtain ferromagnetic semiconductors); as well as the ferromagnetic half-metallic CrO_2 . WO_2 is an example of a rutile-"like" monoclinic metal oxide. It differs from the rutile shape along its would-be rutile c-axis (monoclinic a-axis), here the metal-metal bonds dimerise with alternating bond lengths in a similar fashion to the well known VO_2 at room temperature. It is these metal-metal bonds that give rise to the dimerisation, specifically where the W ion has a $5d^2$ occupation. The inherent structural repeat units of edge-sharing WO_6 octahedra and W_3O planes, a feature of all rutile systems, persists in the monoclinic structure with octahedral edge sharing along the monoclinic a-axis. Of importance for these spectroscopic measurements are the W_3O planes or trigonal planar coordination of the oxygen atom; this results in oxygen sp^2 hybridisation. All these planes have central oxygen atoms that form the corners of every WO_6 octahedra and thus the W_3O planes common to the apical oxygens are at 90° to the W_3O planes arising from any and all of the oxygen atoms around the waist of the octahedron. Nevertheless each W_3O plane shares the monoclinic a-axis within their planes.

Previously, WO₂ has been studied by UPS [1, 2], core level photoemission, EELS [3] as well as W $L_{2,3}$ XAS and O K-edge non-resonant x-ray emission [4] but no study has yet exploited the linear dichroism at the O K edge which is expected due to the arrangement of W₃O planes. The electronic structure of WO₂ and of similar distorted rutile-like oxides such as MoO₂, NbO₂ and VO₂, are of interest due to the variety of the metal-metal interactions and metal-insulator transitions that these materials exhibit. To understand the electronic structure and chemical bonding within this system we have applied a combination of x-ray absorption spectroscopy (XAS) and resonant x-ray emission spectroscopy (RXES) at the O K edge to a single oriented bulk crystal of WO₂. These measurements occurred at the I511 soft x-ray undulator beamline, specifically the I511-3 bulk branch at MAX-lab.

All measurements were carried out on a single crystal of WO₂. The crystal was grown by the vapour transport method [5] from commercially available WO₂ powder and this crystal had previously been characterised by ultraviolet photoemission spectroscopy, LEED and STM [1, 2]. The crystal itself measures approximately 4 x 5 x 2.5 mm and has a single flat smooth growth face which experimental Laue x-ray back reflection supported by LEED and STM has identified as the (012) plane [2]. All the measurements shown here are recorded in an "out-of-plane" scattering geometry with the XES spectrometer rotated so that the plane of scattering is at 90° to the plane of polarisation of the incident excitation.

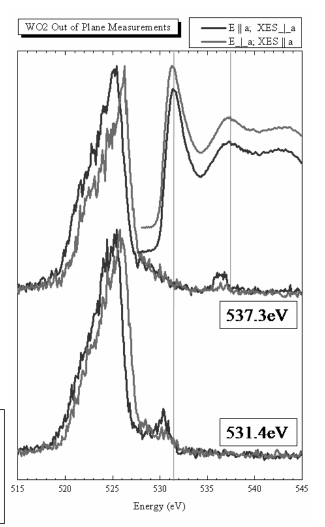
Like rutile compounds, the local planar coordination of oxygen atoms in the W_3O planes result in planes containing a common axis, in this case the monoclinic a-axis [6]. In general the sp^2 hybridised oxygen 2p states are separated into π - and σ -like molecular orbitals involving the metal cation, perpendicular to the W_3O plane or within the plane respectively, giving rise to a natural linear dichroism. In the case of WO_2 , due to the monoclinic distortion and W-W dimerisation along the monoclinic a-axis due to the $5d^2$ configuration, there now arises two inequivalent oxygen sites O1 and O2, where O2 is centred in a W_3O triangular plane with the longer W-W distance along one-edge and where the average W-O bondlength is increased by about 3.3%. Thus the O 2p π -like orbital on O2 is essentially non-bonding, whereas the equivalent O 2p orbital (\bot local W_3O plane) is much more effectively hybridised with W 5d orbitals. When the monoclinic a-axes of WO_2 is oriented parallel to the linearly polarised incident synchrotron radiation, it then excites into primarily σ^* states, when oriented perpendicular to the a-axis then a combination of σ^* and σ^* are selected thus giving a symmetry selected oxygen 2p conduction band partial density of states (PDOS) although averaging over the O1 and O2 sites. Exploiting polarisation dependent resonant soft x-ray emission spectroscopy (RXES) at the oxygen K-edge can then probe the occupied oxygen 2p π and σ states to measure the symmetry and site selected oxygen 2p valence band PDOS.

The figure shows the oxygen K-edge x-ray absorption spectrum of WO₂ where the electric field vector is parallel to the a-axis (blue lines) and parallel to the a-axis (red lines). Two sets of oxygen K-edge RXES spectra are similarly compared at these two orientations of, where the excitation energies are respectively 531.4 and 537.3 eV as indicated by the orange lines. The difference in the RXES propagating

along the monoclinic *a*-axis in each instance can be attributed directly to the dichroism present or anisotropy in bonding around the oxygen sites.

As observed, the leading edge of the dominant O 2p portion of the valence band emission in the E || a spectrum trails significantly behind that of E_|_a spectrum; this occurs both at the threshold excitation energy of 531.4eV and at the higher excitation energy of 537.3 eV but to a lower degree. A difference spectrum at the threshold energy essentially reveals the O2 2p π like PDOS as more holes are created on the O2 site at these excitation energies while at the higher energy this difference spectrum has transformed to a more equal linear combination of the O1 and O2 2p π -like PDOS. This is evident in the context of density functional theory calculations which are not shown here for brevity. In addition, a dispersion of the features in this difference spectrum can be observed as the excitation energy increases. These RXES tools allow for further insight into the W-O bonding as well as into the W 5d-O2p hybridisation and the W-W bonding along the monoclinic a-axis.

Figure 1: Out of plane O *K*-edge XAS and RXES spectra of metallic WO₂. Two orthogonal crystal orientations are compared for a near threshold excitation and one higher excitation energy. The excitation energies for each set of spectra are indicated by the orange lines.



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Electronic and Surface Structure of Light-harvesting Quantum Dots

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There is an urgent requirement both to improve the efficiency of solar cells and to produce a step change in the cost of solar technology. Promising candidates for 'next-generation' cell technology include colloidal semiconductor quantum dots (QDs) that harvest the incident light, creating an electron-hole pair (exciton), and n-type and p-type conductors that separate the charge. These QDs have the advantages of a cheap, wet chemical synthesis¹, and a band gap that may be tuned to match the solar spectrum. Of particular interest are so-called 'Type

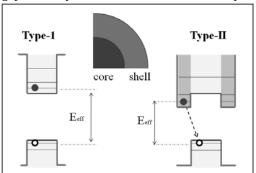


Figure 1: Schematic diagram of Type I and Type II quantum dots. In Type II, the hole is localised in the core, while the electron is trapped in the shell.

II' systems (Figure 1), that have 'core-shell' structure with a staggered band gap that allows the electron and hole to be localized in different regions. The result is a greater degree of control over the excitonic properties, such as the recombination lifetime. Crucial to the exploitation of these QDs is control over the surface chemistry of the QD, as carriers may be trapped there. It is also essential to establish that the required 'core-shell' structure has been synthesized – and there are few techniques capable of doing this. Using beamline I511/1 at MAX II, we have used the tunability of the SR

radiation to carry out depth-profiling XPS of core-shell Type II ZnTe/ZnSe/ZnS and ZnTe/ZnSe QDs, aimed at investigating the internal structure of the QDs.

The synthesis of the ZnTe QD cores (of around 2.2 nm diameter) follows a modified hot injection method using zinc oleate and tellurium trioctylphosphine (TOP) precursors². The ZnSe shell structure (in this case, of 4 monolayers' thickness) is added by the SILAR procedure (successive ionic layer adsorption and reaction)³ giving an overall diameter of 4.5 nm. A further ZnS capping layer can be added to reduce oxygen sensitivity, and in this case the resulting ZnTe/ZnSe/ZnS dots had a diameter of 5.0 nm. For photoemission measurements, the initial oleic acid/TOP ligands were exchanged with the shorter butylamine to prevent sample charging. The butylamine-capped QD samples were deposited from hexane solution onto conducting glass slides.

Figure 2 shows the Se 3d and Te 4d core level spectra obtained from a ZnTe/ZnSe/ZnS sample, as a function of photon energy sampling electron kinetic energies of approximately 200 eV - 700 eV, and thus sampling depths of around 2.2 - 5.0 nm (assuming a sampling

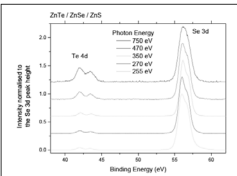


Figure 2: Se 3d and Te 4d core level peaks for a ZnTe/ZnSe/ZnS Type II QD as a function of photon energy. For convenience, the spectra are normalised to a constant height at Se 3d.

thickness. This gives a value of 1.3 nm for the ZnSe shell thickness, in excellent agreement with the shell thickness estimated from TEM, and provides strong evidence for production of

a 'core-shell' structure with little alloying of the phases within the dot. We also observe the presence of a second Se component at the surface, to higher binding energy than the ZnSe signal, that may be due to oxidation⁴, and some surface-localised P, due to incompletely-exchanged TOP ligands. For the 'core-shell-shell' ZnTe/ZnSe/ZnS QDs, we find that S is, as expected, localized at the surface of the QDs, and a second Zn species is present, that may again be associated with surface oxidation (to ZnSO₄). information about the surface and core composition of the QDs is important in device optimization, and is not easy to extract by

depth of 3λ , where λ is the inelastic mean free pathlength.) We thus expect the spectrum taken at 255 eV photon energy to probe mainly the ZnSe shell (of estimated thickness 1.1 nm), as confirmed by the low intensity of the Te 3d

On increasing the sampling depth to around 5 nm (similar to the dot diameter), the Te:Se elemental ratio increases markedly from 2.5 to 4.7, providing unambiguous confirmation that the core of the dot is markedly more Te-rich than the shell (Figure 3). As the flux of photoelectrons is attenuated according to the Beer-Lambert law as it emerges from the sample, it is possible to use a simple 'twolayer' model to obtain an effective shell

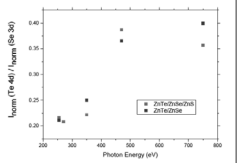


Figure 3: Te 3d/Se 3d core level intensity ratios (normalised for photoelectron cross-section) for ZnTe/ZnSe/ZnS and ZnTe/ZnSe Type II QDs as a function of photon energy.

other means. This work is in preparation for publication.

During this beamtime allocation, we were also able to obtain similar data from aged InP/ZnS QDs, previously studied in a pristine state during experiment 1511/1-129; this allows us to complete a paper on surface oxidation of these QDs. We also attempted studies if polymershelled QDs with catalyst grafts in connection with the use of these dots in artificial photosynthesis, but we were unable to solve sample charging problems during the beamtime.

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Energy level alignment at organic semiconductor interfaces by intramolecular degrees of freedom: transition metal phthalocyanines

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Engineering of the energy level alignment at organic semiconductor interfaces is of tremendous importance for the performance of the corresponding devices. Means to control and adjust the energy levels at particular interfaces have been put forward, among them the introduction of additional interfacial layers, appropriate pretreatments of the metal contact surfaces, and doping of the organic semiconductor. These procedures in common increase the complexity of the device fabrication process which, among other aspects, is unfavorable in view of the anticipated low-cost of organic electronic devices. In this study, we demonstrate using photoemission spectroscopy (PES) studies that an intramolecular degree of freedom, the type of transition metal (TM: Mn, Fe, Co, Ni, Cu, Zn) in TM-phthalocyanines (TMPc's), can be successfully used to adjust the energy level alignment at interfaces to Au without having to increase the number of fabrication steps of the contacts.

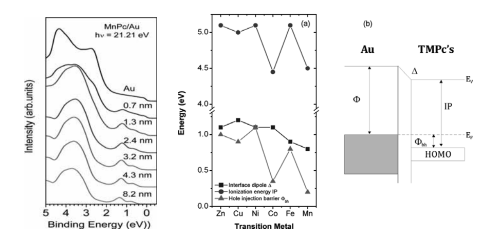


Fig. 1. Valence-band photoemission data of MnPc deposited onto polycrystalline gold as a function of the MnPc film thickness. The data were taken with a photon energy of 21.2 eV.

Fig. 2. Evolution of the interface dipole Δ , the ionization energy IP, and the hole injection barrier Φ_{bh} at TM-Pc/Au interfaces.

In our study we take advantage of the fact that the electronic properties of the TM-Pc's near the energy gap is increasingly influenced by the metal 3d states of the central TM. While for the late TM-Pc's (ZnPc, CuPc, NiPc) important properties such as the ionization energy or the character of the highest occupied molecular orbital (HOMO), which forms the valence band in an organic film or crystal, are exclusively determined by the ligand π states, the metal 3d states come very close to the energy of the HOMO and the

lowest unoccupied molecular orbital (LUMO) when going to CoPc, FePc, and MnPc, and there is also an increasing hybridization of the ligand and TM wave functions. A direct consequence of this variation is a decrease of the oxidation potential (ionization energy) as probed via electrochemistry, and the first oxidation states of MnPc and FePc for instance have been assigned to metal oxidation. One now can expect that such changes of the ionization energy are also reflected in the interfacial electronic properties of the materials when in contact to metals because of the equivalency of the TM-Pc's in many other respects.

In Fig. 1, we show the evolution of the valence band spectra of MnPc deposited on polycrystalline Au as a function of film thickness. Equivalent data have been measured for all TM-Pc's discussed here. From these data one can derive important interface parameters shown in Fig. 2. The ionization energies and in particular the hole injection barriers as determined using PES vary significantly. While for ZnPc, CuPc, NiPc, and FePc this hole injection barrier is rather large and quite similar (0.8-1.1 eV), it is substantially reduced in the case of CoPc and MnPc (0.2-0.35 eV), i.e. via the choice of the transition metal, the energy level alignment can be adjusted. Our results suggest that it is much easier to inject holes into MnPc and CoPc as compared to the other TM-Pc's.

In the following, we demonstrate that the first ionization state of CoPc is indeed of almost pure Co 3d character and thus strongly localized in the center of the CoPc molecule, while the situation is different with respect to the photoemission for MnPc. Fig. 3 shows, that using potential energy surfaces (PESs) it is possible to discriminate between different orbital contributions to a molecular level due to the photoemission cross section variation of the orbital contributions as a function of the applied photon energy

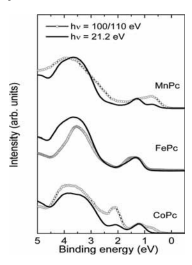


Fig. 3. Comparison of the valence band photoemission spectra of MnPc, FePc, and CoPc taken with photon energies of 21.2 and 110 eV (MnPc, CoPc) or 100 eV (FePc). The thickness of the TM-Pc films was about 70 $\rm \mathring{A}$.

Fig. 3 clearly shows the difference for the first ionization states of the three phthalocyanines. For CoPc the photoemission intensity from this state is negligible at a photon energy of 21.21 eV but reaches about 2/3 of that of the a_{1u} ligand orbital at hv = 110 eV. The situation is very different for MnPc, where the lowest ionization state is clearly visible for both photon energies, but grows with increasing photon energy. For FePc however, intensity changes are observed at the high binding energy side of the first emission feature in direct agreement to what has been published previously. Consequently, while for FePc the electronic states at lowest binding energy can be assigned to ligand a1u states, the situation is quite different for MnPc and CoPc.

Finally, our results demonstrate that the transition metal center has a strong influence on the electronic properties of the phthalocyanine films as well as their interfaces with gold. In particular, the energy level alignment, which has direct consequences for the contact resistance of these interfaces, can be chosen in a large energy range by the choice of the appropriate TM-Pc, and the interfaces CoPc/Au and MnPc/Au are characterized by rather small hole injection barriers. These are directly related to the presence of metal 3d states closest to the chemical potential; a fact that is also reflected in smaller ionization potentials for MnPc and CoPc as compared to other phthalocyanines. Further, we have discussed the nature of the molecular orbitals (metal 3d or ligandlike), that form the states closest to the chemical potential, and it results that they differ between MnPc and CoPc.

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XAS and RIXS measurements of aqueous micelle solutions of Dodecyl trimethyl ammonium chloride

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Dodecyl trimethyl ammonium chloride/water mixtures can be seen as simple models of surfactant/lipid self-assemblies (micelles) [1-4] or proteins [5]. Identifying the factors controlling the formation and dissolution of such complexes and describing water confined in these systems are of fundamental importance for understanding biological processes such as DNA condensation in the chromosome and protein secretion. The state-of-the-art theory is that, in the absence of specific interactions, the complexes are stabilized by attractive electrostatic ion-ion correlation or polyion bridging forces [6]. However, there is very little experimental data showing how the degree of dissociation of polyacids and how polyions and small ions are distributed in the aqueous domains between the particles.

Our experiments were carried out at Beamline 511-3 with a flow cell described in [7]. XAS and RIXS spectra at the O K-edge were measured on dodecyltrimethyl ammonium chloride/water mixtures at rather high water content (70% water), where the system is forming micelle solution (Fig 1). These measurements were compared to the results of an earlier XAS and RIXS study [8] of water-poor dodecyltrimethyl ammonium chloride/water mixture (25% and 10% water respectively) where the system is forming liquid crystalline gels and where the water is more confined in the liquid crystalline lattice. In figure 1 spectra measured on the aqueous liquid crystalline system, micelle solution and pure water are shown. A distinct peak is seen around 532 eV was earlier noted on the liquid crystalline system and is still remaining in the micelle solution system. This reflects the interaction between the water molecules and the ions of the system. Further comparing the RIXS spectra excited at 537 eV, the emission peak at 526 eV is more or less identical to the pure water spectra while it in the liquid crystalline system is narrower and has by quantum chemical calculation been attributed to surrounding trimethyl ammonium groups and chlorine ions. Comparing these three systems the different surrounding of water molecules are reflected in the local electronic structure.

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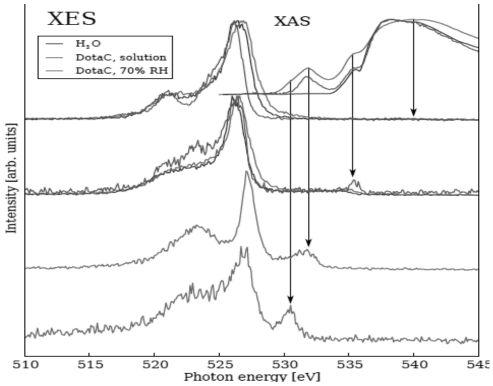


Figure 3: Oxygen K edge spectra of $C_{12}TAC$ liquid crystalline gel (green) and solution (red). The FY spectra are shown in the upper right, and the excitation energies used for the SXE spectra are indicated. The spectra of bulk liquid water (blue) are shown for comparison.

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Polarization dependent RXES at the O K-edge of anatase-TiO₂

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Anatase TiO_2 possesses a M_3O strutural unit, where each oxygen is trigonally coordinated by three metal cations, similar to the rutile structure, but with different bond lengths and angles compared to the rutile TiO_2 M_3O unit. The differences in the O 2p partial density of states in TiO_2 , between anatase and rutile TiO_2 related to these structural changes has been studied using O K-edge XAS and RXES at MAXIab beamline I511-3. Figure 1 includes a diagram of the anatase structure that highlights the M_3O plane for both the anatase and rutile structure, indicating the increased metal-metal distances in the anatase M_3O unit. The spatial anisotropy of the charge density in rutile-type transition metal dioxides gives rise to a natural linear dichroism in the absorption of radiation, and combining this dichroism with the geometric selectivity of the dipole emission, essentially by switching the scattering plane upon rotation of the spectrometer axis with respect to the incident x-ray polarization, allows the polarization dependence of the O K-edge RXES to be described for the rutile oxides for the first time. The approach to the bonding in rutile oxides that we adopt in the present study is based on the description by Sorantin and Schwarz [1].

Compared to rutile, anatase TiO_2 has surfaces that are more photocatalytically active, with a higher surface energy than the rutile TiO_2 surfaces. In its pure and doped form it is the focus of research towards this end: both photocatalytic decomposition of molecules and photolytic production of hydrogen gas. For further discussion of the properties of anatase TiO_2 see Chen et al. [2]

In previous experiment at ALS beamline 7.0.1, which formed part of the current study, Ti L-edge and O K-edge XAS and RIXS was measured from samples of nanostructured anatase TiO₂, with a predominant surface area for the (001) surface, which has been found to have the highest surface energy. Details of the nanostructured anatase TiO₂ samples have published by our collaborators who produced them [3].

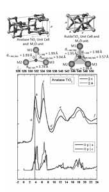


Fig. 1: O K-edge XAS from an atase ${\rm TiO_2}$ and the calculated O 2p crystalline axes-projected PDOS in the conduction band showing the excitation energies for the experimental and simulated RXES.

We acquired a bulk single crystal of an atase ${\rm TiO_2}$ from SurfaceNet GmbH, the main surface of which was also (001) and performed a polarization dependent O K-edge XAS and RIXS study on it. The out-of-plane measurements were obtained at MAXlab beamline I511-3 in March and April 2010.

The RXES spectra were compared to simulated spectra based on the results of electronic structure calculations for anatase ${\rm TiO_2}$ carried out by Dr. Cormac McGuinness using the Wien2k formulation of density functional theory. In general there was excellent agreement between the experimental and simulated spectra with particular respect to the development of the k-selective features in the RXES spectra.

The octahedral crystal field experienced by the Ti ion at the centre of the TiO_6 octahedron lifts the degeneracy of its 3d orbitals with the e_g levels higher in energy than the t_{2g} levels by an amount referred to at the crystal field splitting.

The further distortion of the ${\rm TiO_6}$ octahedron present in anatase, where the oxygen anions are displaced such that they lie, consecutively, both above and below the equatorial plane in turn lifts the degeneracy

of the Ti $3d t_{2g}$ orbitals. This distortion is visible at the centre of the unit cell depicted in the inset to figure 1. As mentioned previously, the experiments at MAXlab I511-3 were performed in the out-of-plane orientation

of the analysis chamber with respect to the polarization of the synchrotron beam. The anatase TiO_2 crystal was mounted such that for a grazing angle of incidence in the out of plane geometry, the polarization of the exciting photons were parallel to the one of the crystalline a axes. The axis of the emission spectrometer was parallel to the crystalline c axis, and sampled the emission from the O 2p PDOS projected in the plane defined by the two crystalline a axes. In this scattering geometry the states formed by non-bonding or metal-oxygen (M-O) π -bonding O $2p_y$ orbitals oriented perpendicular to the M₃O planes are populated upon absorption, as well as the M-O σ -bonding states due to the O $2p_x$ and $2p_z$ orbitals lying within the M₃O planes.

The O K-edge XAS was carried out with an experimental monochromator resolution of 0.1 eV, while the resolution of the monochromator was set to a calculated value of $0.4 \, \mathrm{eV}$ for the RXES spectra.

Figure 1 shows the results of the O K-edge XAS for E||a and E||c excitation geometries, alongside the projection of the calculated O 2p conduction band PDOS along the same crystalline axes. The features in the conduction band associated with the selected RXES excitation energies labeled a-f are also shown in figure 1.

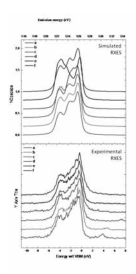


Fig. 2: Experimental and simulated O K-edge RXES from a natase ${\rm TiO_2}$ for the excitation energies indicated in figure 1.

Figure 2 presents the results of the both the in-plane and out-of-plane RXES, for both the experimental and simulated spectra.

The comparison of the experimental and simulated spectra (with the aid of the calculated band structure character plots which are not shown here) allow the k-dependence of the RXES spectra to be observed quite clearly, at least for the resonant energies a-d inclusive.

The simulated spectra for energies higher above resonance do not reproduce the trends in the experimental spectra as agreeably as those with excitation energies closer to resonance. This may indicate that the calculated electronic structure preserves the k-dependence too far above the absorption threshold, compared to the k-states that are selected by the combination of experimental geometry and excitation bandwidth which the simulated spectra approximate. A more accurate treatment of the valence hole broadening applied to the simulated spectra, which is modeled in the present spectra as a linear dependence with valence hole binding energy, may also produce a better agreement between the experimental and simulated spectra.

One set of RXES spectra remain to be measured from an atase TiO_2 in the out-of-plane geometry at MAXlab I511-3. These are the spectra with an $\mathrm{E}||c$ excitation geometry, and XES along a spectrometeral lignment. At this point a detailed comparison between the rutile and an atase TiO_2 polarization-dependent O

K-edge XAS and RXES will be complete, and the polarization-dependent study of the XAS and RXES at the Ti L-edge will also be attempted. This, in turn, forms part of a systematic polarization-dependent soft x-ray spectroscopic study of the electronic structure of rutile transition metal oxides and fluorides by the present authors, including SnO_2 , RuO_2 , WO_2 , IrO_2 , MnF_2 and MgF_2 .

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RIXS in Li₆YB₃O₉ and Li₆GdB₃O₉

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We report resonant X-ray inelastic scattering (RIXS) spectra of Li₆YB₃O₉ (LYBO) and Li₆GdB₃O₉ (LGBO). Borate crystals are widely used for non-linear optical, isolation, luminescence and phosphorescence purposes. High efficiency of borate crystals as scintillation materials as well as in neutron detection has been demonstrated [1, 2].

Recently special attention has been given to complex alkali rare-earth borates, which have anion groups consisting of boron and oxygen. LYBO doped with rare earth ions is considered to be a promising laser medium [3]. LYBO crystals have a band gap of 7.04eV and are transparent in 176-900 nm region [4].

Experimentally the electronic structure of borates has been studied by X-ray photoelectron spectroscopy. However the standard methods based on electron emission (resonant and nonresonant photoemission and Auger spectroscopies) suffer due to the strong charging effects in the wide band gap crystals. In case of the insulating materials it is timely to apply a charge neutral spectroscopic probe. Resonant inelastic x-ray scattering is a photon-only technique performed at soft x-ray absorption resonances, making it atomic and orbital symmetry specific, bulk sensitive and thus complementary in many respects to electron spectroscopies.

The experiments were performed at the "bulk" branch of the beamline I511 (I511-3) of MAX-lab's synchrotron MAX II, Sweden. The resolution of the beamline monochromator was set to better than 25 meV at the B 1s threshold absorption measurements. The X-ray spectrometer had 0.2 eV resolution at the B 1s edge. The absorption spectra were measured in TFY (total fluorescence yield) mode using an MCP detector with a repulsive field for electrons. The samples were monocrystals.

The absorption spectra of the crystals (top panels of Fig. 1.-2.) display strong sharp peaks at 193.55 and 193.7 eV. These peaks can be attributed to boron core exciton states.

The RIXS spectra excited in the vicinity of the B 1s core resonance (bottom panels of Fig. 1.-2.) show two principal features. The first feature is the emission in the energy region 174-187 eV and it is quite wide. The second feature - the elastic peak with the energy loss shoulder to its low energy side - occurs at about 193 eV.

There are two main radiative decay channels in the core exciton state of boron: the "participator" channel, where the electron that was excited by absorbing the X-ray photon decays back to the core level and the "spectator" emission channel where a valence electron fills the empty core orbital and the previous core electron remains in an excited state.

195 205 210 193.55 193.75 Li₆YB₃O₆ 194.0 193.2 201.5 194.2 203.5 194.4 193.0 192.8 190 Excitation energy (eV)

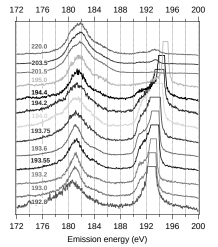


Fig. 1. XAS spectrum of Li₆YB₃O₉ measured in TFY mode (top). RIXS spectra of $Li_6YB_3O_9$ (bottom). The RIXS spectra have been normalized to equal spectator emission maxima and the top of the elastic emission peak has been cut to prevent the overlapping of the spectra.

The first feature (approx. 174-187 eV) arises from the scattering on a valence excitation, which at higher excitation energies verges into the characteristic $K\alpha$ emission. Until the excitation energy is in resonance with core exciton level, this emission band is the spectator emission.

234 MAX-lab ACTIVITY REPORT 2010 The emission around 193 eV arises from elastic scattering and also from inelastic scattering on the exciton state. The proposed origin of the energy loss shoulder to the elastic peak is lattice relaxation in the absorption

site, similar to BeO [5]. This process was first observed by Ma et al. [6] in the emission spectra of graphite and diamond, while in the absorption spectrum only a relatively sharp core exciton peak was observed. The authors suggested that strong vibronic coupling leads to a local lattice distortion and to the corresponding shifts of the potential energy in the excited electron state.

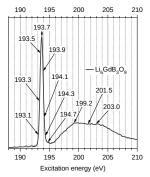
The participator emission process is only possible when the core excited electron remains bound to the absorbing atom — in the case of the crystals considered here the participator process is possible when a core exciton is created. At higher excitation energies, when the excitation is no longer in resonance with the core exciton level, the energy loss shoulder disappears and only a elastic scattering peak is observed.

The participator emission band reappears when the excitation energy is increased further (hv > 203 eV) although its intensity then is much weaker. This is a second threshold phenomena, where a core exciton together with a valence band hole are created by an X-ray photon directly or indirectly.

The absorption spectra of LYBO and LGBO are quite similar. There is a difference at which energy the excitation of the core electron to the conduction band has the highest intensity – LYBO has a maximum at 203 eV, while LGBO has nearly constant intensity at 199-204 eV.

The RIXS spectra of both crystals are also similar. The low energy cut-off of the low energy tail to the elastic peak is at nearly constant photon energy - 190.5 eV in both crystals. In both cases the energy loss tail extends up to the elastic peak.

The similarity of the spectra can be explained because of the fact that the structures of the crystals and the local symmetries of boron sites are similar in both crystals [7].



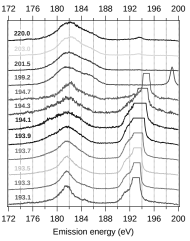


Fig. 2. XAS spectrum of $Li_0GdB_3O_9$ measured in TFY mode (top). RIXS spectra of $Li_0GdB_3O_9$ (bottom). The RIXS spectra have been normalized to equal spectator emission maxima and the top of the elastic emission peak has been cut to prevent the overlapping of the spectra.

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Passivation and stability of infrared photodetectors based on type-II GaAs quantum dots in InAs matrix

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Infrared (IR) photodetectors and focal plane arrays with operating temperatures above 77 K are highly requested since it allows simpler, more reliable and lower-cost cryogenic cooling systems. The longer-term goal is thermo-electric cooling or even operation at room-temperature which will enable very cost-effective and ultra-compact IR systems. IR detector systems presently under intense development are type-I InAs/GaAs quantum dot photodetectors (QDIP) and type II In(Ga)As/GaSb strain layer superlattices (SLS) photodetectors. They have both been demonstrated to be promising for operation at moderately high temperatures. ¹⁻⁴ However, there are still challenges in their realization that have hampered their development. For instance, it is difficult to obtain high response from QDIPs due to the limited quantum dot (QD) density and numbers of QD layers, while the antimony-based SLS photodetectors are very challenging from an epitaxial growth standpoint and also suffer from high surface leakage current in mesa-etched devices.

In this work, different passivation techniques of photodetectors based on GaAs quantum dots embedded in an InAs matrix have been investigated with the target on long wavelength IR (8-12 µm; LWIR) imaging.⁵ In such structure, a type-II band alignment is created between the highest valence band energy level in the GaAs QD and the conduction band edge of the surrounding InAs material. The operating wavelength can then be tuned by adding Al or Sb incorporation into the GaAs QDs.⁵ As compared to the extensively investigated type-I QDIPs, there has only been a few reports on type-II QDs for IR detection, for example in the InSb/InAsSb, InSb/InGaAs and Ge/Si systems.⁶⁻⁸ A key advantage of this device is that it enables a longer carrier lifetime due to the spatial separation of the different types of carriers, and possibly higher operating temperatures.^{5,6}

However, the matrix material InAs is a narrow band gap semiconductor where the surface Fermi level is pinned in the conduction band due to large density of surface states, resulting in a high conductive n-type surface layer. ⁹⁻¹¹ In the present study, the dark current (that could be directly related to the surface leakage current) of the detectors treated by various surface passivations was evaluated using current-voltage (I-V) measurements at 77 K and room-temperature (RT), respectively. In particular, device stability was observed during eight months after the devices were passivated. X-ray photoelectron spectroscopy (XPS) was used to study the surface chemical composition before and after the passivation step.

The device structure consisted of five stacked undoped GaAs QD layers embedded in the InAs matrix grown on undoped (100) InAs substrates by metal-organic vapor phase epitaxy. The QD layers were separated by 50 nm InAs spacer. A 0.5 μ m InAs p-contact layer (2.5×10¹⁸ cm³) was first grown on the substrate, followed by the growth of a 200 nm undoped InAs buffer layer. The nominal growth rate for GaAs QDs was 0.03 nm/s with V/III ratio of 10, while the InAs layers were grown at a growth rate of 0.37 nm/s under the V/III ratio of 150. The QD layers were finally terminated by a 300 nm n-doped (5×10¹⁷ cm³) InAs contact layer.

The devices were fabricated using standard photolithography, inductively coupled plasma (ICP) etching, metal deposition, and lift-off procedures. The mesas were etched using ICP with Cl₂/H₂/CH₄ plasma at powers of ICP=1 kW, RF=80 W followed by a citric acid-based wet etch to remove etching damage and residues.

As fabricated devices showed extremely high dark current, and rather behaved as typical conductors. Their I-V characteristics at 77 K show a typical Ohmic behavior. Furthermore, the I-V characteristics of the unpassivated devices were nearly size and temperature insensitive. It indicates that the surface leakage (shunt current) is dominating the high dark current, otherwise the dark current should be lower in smaller size devices.

In order to resolve the surface-leakage problem, different surface passivation treatments were carried out and investigated. Although the dark current of the devices improved slightly after these treatments, the expected pin diode characteristics were still not obtained. However, it was discovered that a simple annealing step at a temperature higher than 180°C for longer than 5 minutes in vacuum can reduce the dark current dramatically. To further optimize the thermal passivation approach, a series of heat treatments were explored as varying temperature from 180°C to 400°C. The results showed that the dark current decreases with about two orders of magnitudes when the annealing temperature increases from 180°C to 330°C.

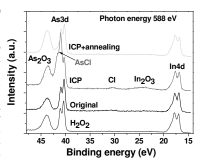
To investigate the stability of the passivated devices, the I-V characteristic of several selected devices was measured at both RT and 77 K every second weeks for eight months. During the whole period, the devices were kept in atmospheric ambient conditions. Encouragingly, the I-V characteristic remained almost identical during the entire investigation period, indicating an excellent stability. The good device stability in atmospheric ambient condition is a somewhat surprising finding considering what is known about performance degradation induced by oxides for InAs and

other narrow band semiconductor devices. When the devices were exposed to air, AsO and InO related oxides are formed on the InAs surface.

To understand mechanism of the annealing process leading to efficient passivation and good stability of the devices, the surface chemical compositions after the different treatments were analyzed by XPS at beamline I511 at the national Swedish synchrotron radiation Laboratory MAX-lab. Four samples were thereby prepared. One is as-grown sample (original), the second one was dry etched by ICP, the third one was also etched by ICP but then annealed at 255°C for 5 min (ICP+annealing), whereas the last sample was treated in H₂O₂ for 6 min.

Overview XPS spectra of these samples were recorded at 730 eV photon energy, and dominating peaks were observed corresponding to O1s at 530 eV, In3d at 450 eV, C1s at 285 eV, In4p at 110 eV, As3d at 45 eV and In4d at 18 eV. The results revealed contaminations of oxygen, carbon and chlorine on the surface for all samples.

As 3d and In 4d spectra were also recorded at photon energy of 588, 250 and 110 eV, respectively. One of them measured at 588 eV is shown in the figure. The peak at 44.5 eV is attributed to the oxidized arsenic species, and the intensity of the ICP+annealed sample is the highest. There are two peaks at about 41 eV corresponding to the well-resolved As3d spin–orbit splitting, except for the ICP etched sample. These spin–orbit split peaks in the ICP etched sample were disturbed by an additional chemically shifted peak from arsenic chloride (AsCl). Apparently these AsCl related surface impurities might be electrically active on the surface, and played a dominant role for the observed shunt current. $\ln_2 O_3$ could be another factor for surface leakage, but its intensity is low for all measured samples. It can be explained by the As-rich surface determined by V/III ratio of 150 as growing the top InAs layer epitaxially. Also, In4d related peaks are quite similar for all samples.



After annealing, the As3d peak of the ICP+annealing sample were almost the same as the original sample, but its As_2O_3 concentration was enhanced. It can be explained by the chloride desorbed when the temperature was higher than a threshold (180 °C), which apparently favored formation of As_2O_3 . This result provides an understanding why annealing can eliminate the surface states, thereby reducing surface leakage. In addition, the annealing process would also realize As dimers owing to surface reconstruction by the thermal energy.¹³ It is worth to point out that the As_2O_3 intensity is high for both annealed and H_2O_2 treated samples, however their dark current level was still low. It seemed that although the As_2O_3 covered and finally saturated on the device surface, they won't influence the device electrical properties. This finding is useful to interpret the good device stability as they were stored in ambient condition.

In summary, we have presented a simple annealing approach to passivate type-II GaAs/InAs QD photodetectors owing to efficiently elimination of AsCl related surface states induced by ICP dry etching. The good stability and device reliability during long term operation demonstrated a promising potential for commercialization of such devices

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Hybrid systems comprising Au nanoparticles in organic matrix

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There are enormous worldwide efforts to develop new memory devices, which would bring together the best characteristics of today's principal technologies: dynamic random access memory, flash memory and hard-disk drives and would get over the disadvantages of those. Such a "storage class memory" would revolutionize the information technology industry as outlined in [1, 2]. A number of physical effects have been proposed to serve as a basis for new memory elements of future technologies. New devices based on switchable resistive materials are commonly ranked as resistive random-access memory (RRAM). One type of RRAM are hybrid organic—inorganic systems, mainly consisting of nanoparticles (NP) blended into an organic matrix.

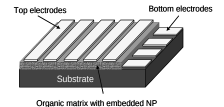
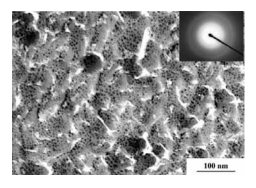


Fig. 1. Crosspoint memory architecture.

Fig. 1 shows possible memory architecture is obtained if the organic thin film containing metal nanoparticls is sandwiched between a cross-point array of electrodes – that consist of narrow metal stripes, running in perpendicular directions above and below the film. The resistivity at "crosspoints" can be switched either a high- or low-conductivity state by applying a voltage, suitable to *write* or *erase*. The resistivity in the high- and low conductivity states can differ by 6-8 orders of

magnitude. The corresponding state can then be probed by measuring the current across the crosspoints at some lower voltages. New devices based on switchable resistive materials are commonly ranked as RRAM. One type of RRAM are hybrid organic–inorganic systems, mainly consisting of inorganic nanoparticles (NP) blended into an organic matrix.



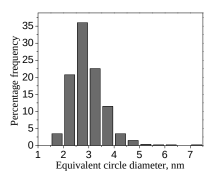


Fig. 2. Electron micrograph of nano-composite thin-film composed of gold particles embedded in molecular large gap crystal CuPc for nominal Au deposition of 4 Å (left panel). Electron diffraction patterns for corresponding depositions are presented as insets. Histogram showing size distribution of gold nanoparticles from electron micrograph (right panel).

The embedded nanoparticles were self-assembled from gold atoms deposited on the outer surface of the organic film due to surface and bulk diffusion, resulting in a three-dimensional nanoparticle distribution in the bulk of the organic semiconductor. The evolution of microstructure, i.e. the size, concentration, bulk- and size-distribution of the resulting nanoparticles as a function of nominal metal content was studied using transmission electron microscopy (TEM) with a JEOL JEM 100 CX operated at 100 kV, while the evolution of electronic structure of the hybrid organic-inorganic system was studied by surface- and bulk sensitive spectroscopic methods.

It was found that at a nominal coverage of 4 Å gold the averaged diameter of the particles amounts to <d> = 3 nm with a standard deviation of 0.7 nm (see Fig. 2). This nanoparticle size distribution (psd) corresponds to an average cluster nuclearity of about 950. It is amazing that for nominal Au deposition of 4 Å the diameter distribution of the gold NP's in the CuPc film is very narrow. For nominal coverage of 16 Å gold the averaged diameter of the particles amounts to <d> = 6.3 nm and for 32 Å the averaged diameter amounts to <d> = 9.8 nm. We also provide evidence that the Au particles do not agglomerate near the surface but their diffusion through the entire CuPc film occurs.

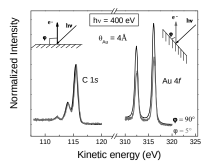


Fig. 3. C 1s and Au 4f core-level spectra, measured for two different emission angles from CuPc film after nominal gold coverage of about 4 Å. All spectra were normalized

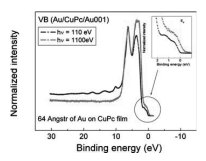


Fig. 4. The VB spectra acquired using soft x-ray hv=110eV (black line) and hard x-ray hv=1100 eV (grey line) photon energies on the Au/CuPc system for nominal gold coverage of about 64 Å.

The results presented in Fig. 3 mean, that the Au concentration in the bulk is higher than in the subsurface region as a result of strong diffusion processes of individual Au atoms or whole clusters within the organic matrix.

In order to verify that the gold atoms/clusters diffuse deep inside the organic matrix we also compared valence-band spectra taken at different photon energies (Fig.4). The more bulk sensitive VB spectrum (hv=1100 eV) is practically characteristic of a Au bulk sample, which means that in this wider thickness region of our hybrid organic- inorganic system, the gold is assembled in large nanoparticles, which show metallic properties, including the formation of a distinct Fermi-level step and probably demonstrate some percolation.

Finally, we should emphasize that there is no formation of a continuous metallic film on top of organics up to large nominal coverage of about 130 Å considered in the present study. The gold is assembled in well defined NP's with metallic properties.

Acknowledgements: This work was supported by the DFG under grant KN393/14, and by the RFBR under grant 10-02-00269.

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Investigations of valence band electronic structure of Pr_{0.5}Sr_{0.5}MnO₃

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The charge ordering (CO) phenomena in colossal magneto resistance (CMR) materials have been attracting a great deal of research interest owing to the intriguing physics involved in the dynamics of conduction electrons. This phenomenon occurs in those perovskite manganese oxides in which the on-site coulomb interaction is stronger than the kinetic energy of the charge carriers [1]. Pr $_{0.5}$ Sr $_{0.5}$ MnO $_3$ is a good candidate, which exhibits CO state across its ferromagnetic metallic (FMM) to antiferromagnetic insulating (AFMI) transition. This CO state is intimately related to the CMR properties.

Pr $_{0.5}$ Sr $_{0.5}$ MnO $_3$ also exhibits multiple magnetic phase transitions upon cooling. From a paramagnetic metallic (PMM) state at room temperature, this composition turns to a FMM state below T_C (~270 K) and finally to a CO - AFMI state at T_N (~150K). The PMM - FMM phase transition is isostructural (tetragonal with c/a > 1) while the FMM - AFMI transition leads to monoclinic change in lattice symmetry [2]. These magnetic phase transitions are expected to modify the near Fermi level (E_F) electronic structure of these materials, especially when they are accompanied by a change in the lattice symmetry. It is, thus, important to study the surface electronic structure of the energy scales involved in the CO phenomena and the associated changes in the near E_F electronic structure.

The single-crystal samples of Pr $_{0.5}$ Sr $_{0.5}$ MnO $_3$ were prepared by the floating zone method in a mirror furnace. The compositional homogeneity and the tetragonal structure of the crystal were confirmed using energy-dispersive spectroscopy analysis and room temperature powder X-ray diffraction respectively. To validate the multiple transitions temperature dependence of magnetization and resistivity were measured.

The high-resolution photoemission (PES) measurements were performed at BL I-511 at MAX-lab using a Scienta R4000 analyzer. The photon energy 50 eV was used for these measurements. To calibrate the binding energy Fermi edge of the gold reference sample was used. The energy resolution was set at 20 meV for 50eV photons.

Fig. 1(a) shows the valence band photoemission spectra of the sample across its PMM-FMM-AFMI transitions showing four distinct features marked A (1.1 eV), B (2.1 eV), C (3.1 eV) and D (5.4 eV). The spectra from the PMM and FMM phases

show a clear Fermi edge. It is also clear from the figure that the width of the feature A (which corresponds to the e_g states) decrease as we go down in temperature across the FMM-AFMI transition. Further, the intensity of the near E_F was found to decrease as the material goes through this transition [Fig.1 (b)] attributed to the localization of the e_g electrons in the CO-AFMI phase. Further, a small but finite intensity at E_F was observed in the insulating state of the sample. Considering the difference in the electron mean free paths between the bulk and surface sensitive photoemission spectra, our results indicate that the surface Mn 3d states are somewhat different from that of the bulk Mn 3d states in $Pr_{0.5}Sr_{0.5}MnO_3$. Thus, the CO state in the surface is suppressed from that of the bulk $Pr_{0.5}Sr_{0.5}MnO_3$. The finite intensity at the Fermi level insinuates the rotation and distortion of the MnO_6 octahedra in the insulating phase resulting in changes in the crystal structure and reflected strongly on its surface.

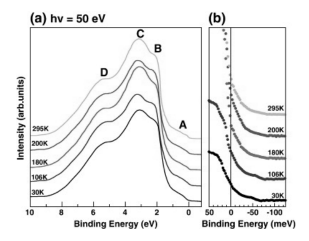


Fig 1: (a) Valence band photoemission spectra of $Pr_{0.5}Sr_{0.5}MnO_3$ obtained using 50 eV photon energy across the PMM-FMM-AFMI phase transition (b) Expanded spectra showing the spectral changes in the vicinity of E_F

We have further analyzed the near EF spectral behavior across the PMM - FMM - AFMI transition to show the formation of a pseudo gap in the PMM and FMM phases [3] and finite photoemission intensity at EF in the CO-AFMI state. In Ref 3, the formation of pseudogaps has been further discussed with reference to strong electron-phonon interaction, consequent charge localizations and possible formation of JT polarons.

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Investigating the local partial density of states in swelling clay in dry and humid environments

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Clay is an interesting material in many fields such as geosciences, pharmacy, and biology. Swelling clay consists of sandwiched SiO2/Al2O3 sheets carrying a net negative layer charge which in turn are electrostatically bound by cations, see Fig. 1. When the clay is soaked in water, the water molecules enter between these sheets, the layers separate and the material swells. Due to this, large amounts of water can be stored in the clay.

The valency and the hydration properties of the cations between the sheets control both swelling and colloidal behavior of the clay [1]. Interaction between the water and the interlayer is affected by changes in the ion concentration in the water and by changes in the mineral structure of the clay. Thus, the behavior of the clay material in water will also yield information about the interaction of water with metal oxide interfaces.

Near edge x-ray absorption fine structure (NEXAFS) measurements in photon-in/photon-out mode and Resonant inelastic soft x-ray scattering (RIXS) have unique features which make them an ideal tool for the investigation of complex samples as they yield information about the local electronic structure at specific atomic sites and thus reflect the local chemical environment of the probed atom. Additionally, they can be applied for the investigation of matter in humid environments since the large penetration depth of soft x-ray photons of this photon-in/photon-out technique allows insitu investigation of the properties of gases, liquids and suspensions enclosed behind ultrathin windows.

The experiments were performed at the beamline I511-3 at MAX-lab in Lund, Sweden using a button cell with a 100 nm thin Si_3N_4 membrane window. Both the incidence and emission angle were 45 deg.

NEXAFS and RIXS spectra of natural swelling clay are shown in Fig. 2 together with a reference spectrum of clean de-ionized water. The clay has been measured as dry sample as well as soaked in de-ionized water. In the NEXAFS spectrum, the low energy shoulder

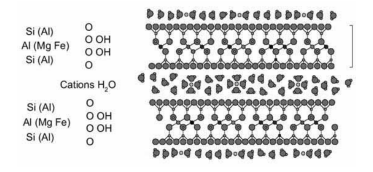


Figure 1: Schematic structure of swelling clay [2]

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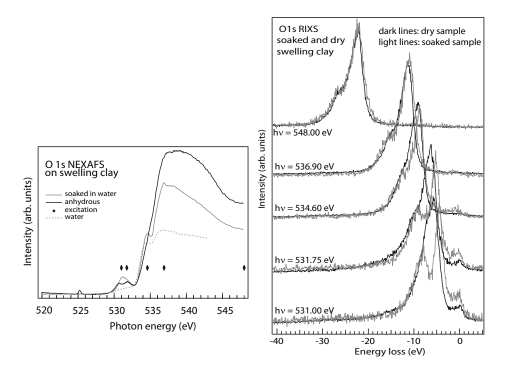


Figure 2: NEXAFS (left) and RIXS (right) measurements on dry and humid natural swelling clay show clear differences originating from changes in the electronic structure of the oxygen atoms due to gelation of the clay.

around 531 eV is characteristic for clay materials and is associated with quasi-free O_2 molecules in amorphous alumina and silicate [3].

In the dry sample, the oxygen signal originates from OH inside the layers. The pre-peak at 525.25 eV is assigned to an amorphous defect which is temporally healed by interaction with water. In the soaked sample, an additional signal from water stored in the clay is observed at 534.6 eV which corresponds with the OH bond signal from the water reference spectrum.

We acknowledge support from the MAX-lab staff.

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Photoemission studies of a single-centre water splitting complex and chlorine containing analog adsorbed on rutile TiO₂(110)

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Figure 1. Chemical structures of the dye complexes studied in this investigation, (a) is the single centre water splitting complex (WSC) and (b) is its chlorine containing analog (CISC).

Natural reserves of fossil fuels will eventually become exhausted, hydrogen fuel has the potential to replace fossil fuels for many of their applications including transportation. To make hydrogen fuel a cheap method of production must be found, ideally one that uses renewable energy. A reaction that produces hydrogen molecules is the water splitting reaction in photosynthesis which uses the energy from absorbed sunlight. A cost-effective method of absorbing sunlight is the dye-sensitized solar cell (DSC), by adapting the structure of the dye molecules we could cost-effectively produce hydrogen fuel. Past studies have shown that a single-centre dye complex is less effective than a multi-centre dye complex at the water splitting reaction.[1] In situ electrospray deposition (ESD) was used to deposit the dye complexes, as the molecules would not survive the thermal evaporation process, this allows us to study the dye complexes using UHV synchrotron techniques. The deposition solution passes through a high voltage capillary where the electric field causes the formation of charged droplets. The droplets pass through 4 differential pumping stages and lose solvent molecules through evaporation, the desolvated molecular ions arrive at the sample surface at pressures in the 10^{-7} mbar range [2-4].

In this experiment we successfully deposited the water splitting complex (WSC) and its chlorine containing analog (CISC), with the structures shown in figure 1 onto atomically clean $TiO_2(110)$ surfaces under UHV conditions at the surface end-station of beamline I511. The WSC is not only a single-centre water splitting complex but also a model of the reaction centre of a multi-centre water splitting dye complex. The adsorption geometry of the molecules on the rutile $TiO_2(110)$ was investigated using photoemission spectroscopy and energy level alignments were obtained by comparing valence photoemission and NEXAFS spectra. The molecules were deposited using a commercial UHV-compatible electrospray deposition source (Molecularspray Ltd, UK) [5] from a 1:1 methanol:water solution for the WSC and a pure methanol solution for the CISC, the deposition was conducted with an emitter voltage of +2kV and a flow rate of 500 nlmin⁻¹. The pressure during the deposition was $\sim 5x10^{-7}$ torr.

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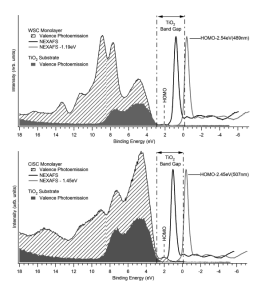


Figure 2. Energy level alignments of monolayers of the WSC (top) and CISC (bottom). The diagrams show valence photoemission spectra of the clean ${\rm TiO_2}$ and with a monolayer of the dye complex adsorbed, also shown are the experimental N *Is* NEXAFS spectra and a shifted NEXAFS to remove the influence of the core-hole.

Figure 2 shows the energy level alignments of the dye complexes adsorbed on the rutile $TiO_2(110)$ substrate. Valence photoemission spectra give information on the occupied states of the molecule, whilst the NEXAFS spectra provide information on the unoccupied orbitals in the system. The NEXAFS spectra are taken under the influence of a core-hole which causes a shift in binding energy of the orbitals compared to normal excitation with visible light. This effect can be removed by comparing the HOMO-LUMO gap from the experiment with optical data.[6] The alignment reveals that the LUMO of both complexes is incapable of charge transfer in the core-hole excited case but charge transfer is possible from the LUMO using optical excitation. This information means that the core-hole clock implementation of resonant photoemission can be used to examine charge transfer from the adsorbed molecules to the substrate. Also this information shows that a rutile $TiO_2(110)$ surface is a viable substrate for use in a water splitting DSC.

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Interaction of Zinc Phthalocyanine thin films with TiO₂(110)

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The development of novel phthalocyanine (Pc) based organic devices demands a better understanding on the interaction between molecules and other components, like electrode and buffer layer, etc, in order to improve the performance. Previously, tremendous work has been devoted into the characterization of Pc/metal interface. However, as the functionality of oxide becomes more and more intriguing, the study on Pc/oxide system is of great necessity.

In this work, we investigate the adsorption of zinc phthalocyanine (ZnPc) on top of rutile $TiO_2(110)$. The molecule usually behaves like non-transition metal-phthalocyanine but with a

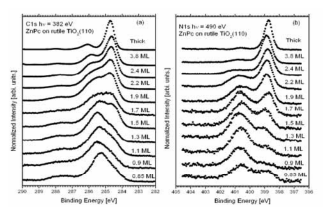


Figure 1 C1s (left) and N1s (right) of ZnPc on TiO₂(110)

filled d-orbital center atom. TiO2 is one of the most well characterized materials oxide with wide applications in catalysis, solar cell and electronic devices. In figure 1 both C1s and N1s spectra are present as the coverage of ZnPc increase on top of TiO₂. The thick film spectra display the typical Pc line profile: C1s consists of three which are peaks assigned Benzene carbon (B, 284.4 eV), pyrrole carbon (P) with B related shake-up structure (286.0 eV) and P

related shake-up structure (287.8 eV); N1s shows a single dominant peak at 389.8 eV and shake-up around 400.6 eV. However, in the monolayer region, the spectra display a completely different structure. At low coverage, the dominant peak of C1s is located around 285.3 eV with a shoulder on the lower binding energy side at 284.5 eV and another broad feature around 287.5 eV. As the coverage increases, the shoulder eventually develops into a distinct peak which is assigned to the thick film related feature, while the dominant feature becomes smeared, which should come from the interface layer. For N1s, double peaks are recorded at low coverage, a stronger peak at 400.7 eV and a weak one at 398.8 eV. The coverage dependent development is similar as C1s: the weak feature gradually evolves into the thick film peak, while the peak at 400.7 eV vanishes. The different peak shifts of core level spectra, 0.8 eV for C1s and 1.9 eV for N1s, on one hand indicates that probable different the electronic densities are distributed around benzene carbons and nitrogen atoms (mainly the pyrrole part) after adsorption. On the other hands, it might also indicate a possible

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molecular plane bending with the carbon towards the surface while nitrogen part further out. Subsequently, the nitrogen atom will have less efficient screening from the substrate than the carbon counterparts.

The observation of core level change can be well supported by the counterparts of highest occupied molecular orbital (HOMO) in Figure 2. The HOMO structure cannot be clearly observed until the coverage reaches 1.9 ML. The shoulder at the higher binding side of the defects states of TiO₂ gradually develops into the HOMO peak. This fact suggests that the HOMO electron is depleted from the interfacial ZnPc into the substrate. Similar discovery has been found on FePc/TiO₂ system¹. This also explains that the big shift of core level is most likely due to the molecular oxidation.

NEXAFS spectra at nitrogen K-edge are shown in Figure 3. Two types of polarizations have been used to probe the empty orbital of molecule and help to determine the molecular configuration. Clearly, it requires more photon energy to excite the

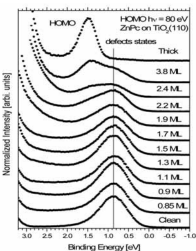


Figure 2 HOMO structure of ZnPc on TiO₂

NEXAFS N k-edge 3.8 MI 2.4 M units.] [arb] 401 402 403 404 399 400 401

Figure 3 NEXAFS at N K-edge of ZnPc on TiO2 under p polarization (left) and s polarization (right)

N1s electron of ZnPc in the interfacial into the lowest unoccupied molecular orbital (LUMO) since N1s orbital has been pulled to higher binding energy due to oxidation. Meanwhile, the transition from N1s to LUMO+1 is barely seen at low coverage, which might indicate that the molecular symmetry has been broken in this case. As the coverage increases, p polarization can resolve more contributions from the ZnPc adlayer, where the N1s-LUMO transition needs 0.7 eV less photon energy. However, almost no such resonance can be observed at s polarization be 2 ML,

indicating that the second layer adopts a very flat geometry.

To summarize, we have characterized the initial steps of the adsorption of ZnPc on TiO₂. A strong charge transfer was found from the first layer molecule to the substrate. The changes in NEXAFS also support this observation and point to a possible molecule symmetry change.

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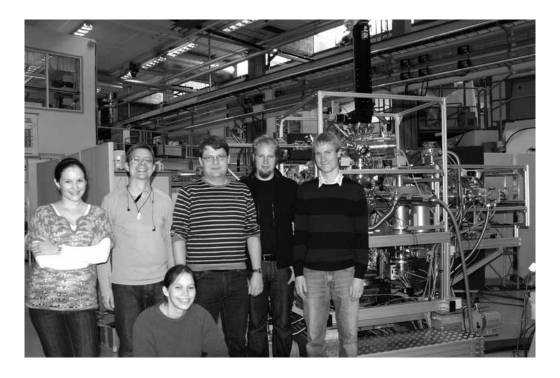
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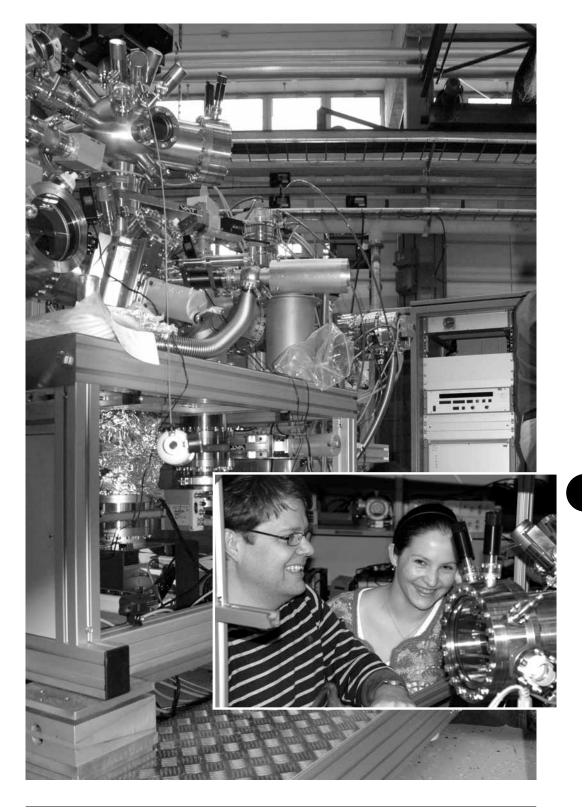
New Endstation at Beamline I511

Olesia Snezhkova, Franz Hennies, Joachim Schnadt, Niclas Johansson, Jan Knudsen, and Annette Pietzsch are all part of the team behind the new HP-XPS Endstation at the beamline I511, 5 October 2010.

Photos: Annika Nyberg



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Time resolved X-ray diffraction from semiconductor nanostructures

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We have investigated propagation of laser induced strain and heat in InSb nanowires grown on an InAs substrate. The strain was measured with time-resolved X-ray diffraction. The scattering from the excited sample was measured using a streak camera. The observed oscillatory reflectivity pattern was attributed to coherent acoustic phonons in the InSb nanowires.

The experiment was carried out at beam line D611. It has a double-crystal InSb monochromator with a bandwidth of $\Delta E/E = 4 \cdot 10^{-4}$, which operates in the spectral range between 1.8 and 8 keV. A 400

 \times 200 μ m² X-ray focal spot size can obtained with $7 \times 0.7 \text{ mrad}^2$ divergence (horizontal × vertical). The Bragg angle for the experiment was chosen to be 59 deg ((111) crystal planes for InSb nanowires, were parallel to the surface of the substrate), while the laser excitation The geometry was normal illustrated in Fig.1. At this geometry the X-ray reflectivity from sample was optimized for the

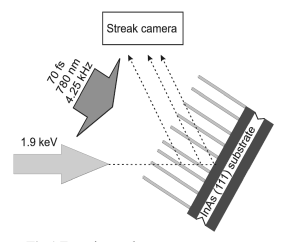


Fig.1 Experimental geometry.

experiment. The whole volume of the nanowires is probed. Short laser pulses were generated by a passively mode-locked, titanium-doped sapphire oscillator followed by a cryogenically cooled Ti:Al₂O₃ multipass laser amplifier. The amplifier was operated at 4.25 kHz and an average power of 2.5 W. The wavelength was 800 nm. The laser system was synchronized to a single electron bunch in the MAX II storage ring with a

jitter of less than 50ps. The diffracted signal from the sample was sent to a streak camera, equipped with a CsI photocathode and triggered by the laser via a GaAs photoconductive switch. The investigated InSb nanowires had a length of 1.7 μ m and a width of 80 nm. These nanowires were grown by MOVPE using trimethylindium, trimethylantimony and arsine precursors on an InAs (111) substrate prepared with Au nanoparticles. The density of the grown nanowires on the surface was 5 nanowires per μ m².

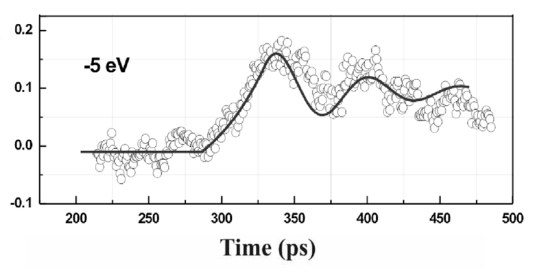


Fig.2 Coherent acoustic phonons in InSb nanowires. The solid line is a guide to the eye.

The frequency of the coherent acoustic phonons as function of detuning from the ideal Bragg condition can produce part of the dispersion curve for the acoustic branch and enable determination of the sound speed of materials [1,2]. Mingo et. al [3] calculated lattice thermal conductivity in various materials and found a reduction in speed of sound, that is following the diameter of the wire. The observed frequency of 17 GHz for 5 eV detuning supports the fact that the velocity of the longitudinal acoustic mode is different from that of the bulk material.

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Formation of nanoscale diamond by femtosecond laser-driven shock

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Formation of cubic diamond from graphite following irradiation by a single, intense, ultra-short laser pulse has been studied. Highly oriented pyrolithic graphite (HOPG) samples were irradiated by 100 fs pulses at a wavelength of 800 nm. Following laser exposure the highly oriented pyrolythic graphite samples have been studied using Raman spectroscopy of the sample surface. In the laser-irradiated areas nanoscale cubic diamond crystals have been formed. We have studied the exposed areas using grazing incidence powder diffraction at beamline D611. We observed a restacking of planes from hexagonal graphite to rhombohedral graphite. The intensity of the rhombohedral graphite powder peaks get more pronounced as the fluence is increased due to restacking at larger depths.

An HOPG sample (12× mm * 12× mm * 2 mm, ZYA grade with mosaic spread of 0.4° was used in this study. It was irradiated by a single laser pulse. The laser was a titanium-sapphire laser with wavelength centered around 800 nm. The duration of the laser pulses were 100 fs. The laser radiation was focused to a peak intensity of up to 2.1×*10¹⁵ W/cm². The fluence range 80-210 J/cm² was investigated. To increase the effective spot size for the x-ray measurements, an array of non-overlapping spots were irradiated by a single pulse each. After laser irradiation, the crystal structure was analyzed using grazing incidence x-ray powder diffraction performed at beamline D611 at the MAX-laboratory synchrotron radiation facility in Lund. The X-ray spot size on the sample was 0.2*2.5 mm² at the incidence angle of 0.8° used in the experiment. This footprint is comparable to the spatial extent of the array of single laser pulse irradiated spots which has a size of 0.15*3.0 mm². Each individual laser spot size was 0.15*0.15 mm². The separation between the laser spots was 0.25 mm. The laser spot-size was measured by a knife-edge scan whereas the X-ray spot was measured using a phosphor and a CCD camera. The photon energy of the X-rays was 9 keV and the bandwidth ΔΕ/Ε=0.1%. Scattered X-rays incident on a phosphor screen were imaged onto a CCD camera. The images were recorded and analyzed in order to produce the powder patterns. The set-up was calibrated with silicon powder. The set-up allowed for coverage of a q-range from (2.9-4.4 Å⁻¹) with a q-resolution of (0.02 Å⁻¹). The result can be seen in figure 1 where we show powder diffraction data from laser-exposed areas as well as from areas with pristine HOPG. As can be seen, all peaks can be identified as reflections from either hexagonal or rhombohedral graphite. The rhombohedral modification of graphite can be found only after the laser irradiation. The structure of rhombohedral graphite (ABCABC) can be understood as an extended stacking fault of the hexagonal configuration (ABAB), and is known to be generated when hexagonal graphite is deformed

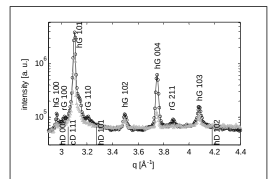


Figure 1: grazing incidence powder XRD pattern of single laser pulse irradiated HOPG. The laser peak intensity: 2.1×10^{15} W/cm², x-ray energy 9 keV, x-ray grazing angle: 0.2° . Δ , \circ : before and after laser exposure; hG: hexagonal graphite, rG: rhombohedral graphite, hD: hexagonal diamond and cubic diamond cD.

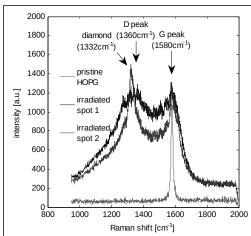


Figure 2: Raman spectra of HOPG after femtosecon laser irradiation and pristine HOPG in comparison. Peak positions for graphite G- and D-peaks, and the diamond peak are marked as reference.

mechanically. What is clear is that a single laser pulse can restack the graphite and the higher the fluence the larger volume fraction of the restacked material is obtained.

Raman spectroscopy was used to investigate the sample surface. This method provides high spatial resolution and surface sensitivity. The Raman spectra from both pristine and laser-irradiated HOPG are shown in figure 2. Within the laser-exposed area, the measured Raman spectra vary locally. This is shown in figure 2 with two typical spectra from different spots within the area exposed to a single laser pulse with an incident fluence of 90 mJ/cm². The most striking difference is the peak at 1332 cm⁻¹, which is the characteristic Raman peak for the T_{2g} phonon mode in cubic diamond. This clearly shows formation of nanodiamonds in the laser-exposed area. The feature at 1580 cm⁻¹ can be identified as the E_{2g} mode of graphite and is present in all spectra. In laser irradiated areas we find a broad feature at 1360 cm⁻¹, which originates from the graphite A_{1g} mode. This mode becomes Raman active due to laser-induced disorder in the graphite structure.

Nanostructural features of lipid-based liquid crystal drug delivery systems

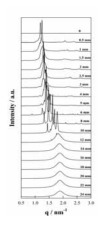
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Depending on various molecular and ambient conditions lipids can form a variety of different liquid crystalline (LC) phases, where the lamellar phase is simplest and the most common in lipid systems. In contrast, nanostructural architecture of reversed non-lamellar phases (like hexagonal and cubic) is much more complex possessing fascinating two- and three-dimensional periodicity. Due to their unique key property to exist in excess water, non-lamellar reversed LC have recently gained an increasing interest in medical uses, e.g. in the delivery of active substances. In this respect, nanoporous lipid LCs have very large interfacial inner area and serve as a matrix for the incorporation (loading), protection and controlled release of drug molecules.

The nanostructural changes as a function of lipid composition of two novel drug delivery systems, SPC/GDO and SPC/VitE in both water and saline solutions have been investigated. The results have shown that a common feature of both lipid systems is that they form non-lamellar liquid crystals over a broad compositional range which exist in excess water solutions. Both systems form reversed hexagonal and reversed micellar cubic LCs at high and low SPC content range, respectively. NaCl and temperature on the phase behavior in excess water conditions have been investigated. From the obtained results it may be concluded that 0.9% of NaCl strongly affects the phase transition boundary between hexagonal and cubic phases. Thus, with NaCl the phase transition boundary for both lipid systems is shifted towards lower SPC content by almost 10 wt% (in respect to SPC/oily lipid ratio). Temperature effects on the phase behavior and the LCs nanostructure have been studied in the physiologically relevant interval (25 – 42 °C). It may be concluded that increased temperature have practically no effect on the reversed hexagonal LCs. Contrary, in both lipid systems reversed micellar cubic LCs are rather unstable and transforms into reversed micellar solution at 37 °C. Non-equilibrium phase behavior of SPC/GDO/saline mixtures has been also investigated in order to get more information about swelling kinetics of different SPC/GDO mixtures. The aim was to determine the swelling profiles of various SPC/GDO depot pre-formulations as a function of lipid composition and time. To this end, mixtures of different SPC/GDO composition were prepared by filling liquid lipid pre-formulation into approx. 1 mm (i.d.) and about 6 cm in length glass capillaries and slowly adding saline solution on top of the air/lipid interface. The capillaries were immediately flame sealed and left to equilibrate for various periods of time at room temperature before examination by SAXS. The swelling profile of the mixtures was determined by measuring X-ray diffraction patterns as a function of beam position along the swelling direction relative to the lipid/saline interface. Figure 1 shows typical results of the obtained swelling profiles as a function of time. Altogether, the data are being complemented with the in vitro release studies of the small molecular weight compounds from both lipid LC systems.

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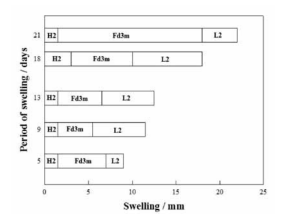


Figure 1. An example of the swelling profiles and the phases identified of SPC/GDO pre-formulation of 50/50 lipid composition as a function of swelling (left) distance and (right) time studied by SAXS at I711.

For the further utilization of novel lipid-based LC delivery systems it is of utmost importance to optimize and control the LC nanostructure for better performance as drug delivery system. We have started to investigate new synthetic lipids suitable to serve as drug delivery matrices. At this stage of the project we have synthesized 4 novel oleyl- and linoleyl-amide derivatives: derivatives (N-(2,3-propanediol)-oleylamide (Compound 1), N-(1,3-propanediol)-oleylamide (Compound 2), N-(2,3-propanediol)-linoleylamide (Compound 3), and N-(1,3-propanediol)-linoleylamide and investigated their temperature-composition phase behavior. Although Compounds 1-3 did not form non-lamellar liquid crystals in excess water conditions, the phase transitions between L_{β} and L_{α} appears to be rather complicated with the formation of intermediate lipid aggregates (Figure 2). Investigations in this direction are still very much in progress; new lipid derivatives are being synthesized.

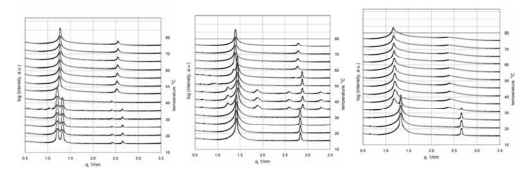


Figure 2. Temperature induced phase transitions of aqueous dispersions of Compounds 1-3 studied by SAXS at I711.

Investigation of the dehydrogenation of $Ca(BH_4)_2$ as potential hydrogen storage material

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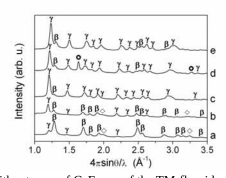
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Light metal borohydrides are considered as promising materials for solid state hydrogen storage. Among them, Ca(BH₄)₂ represents a potential candidate, due to its high gravimetric (11.5 wt. %) and volumetric (~130 kg m⁻³) hydrogen content. Furthermore, the dehydrogenation enthalpy is calculated to be 32 kJ mol⁻¹H₂, if CaH₂ and CaB₆ are the decomposition products, which is within the optimal range for mobile applications. ^{2,3}

The formation of $Ca(BH_4)_2$ was shown to be partially reversible by using suitable additives. *In-situ* SR-PXD was employed to obtain a better understanding of the reactions involved in the samples with NbF₅ and TiF₄. The experiments were carried out at the Synchrotron MAX II, Lund, Sweden, at the beamline I711 with a MAR165 CCD detector. For comparison, all XRD data are reported referring to the scattering vector $4\pi \sin\theta/\lambda$. The results are reported in Figure 2 and Figure 3 respectively.

X-ray diffraction patterns for all the milled materials are presented in Figure 1. All the samples indicate presence of γ -Ca(BH₄)₂ and β -Ca(BH₄)₂ in different abundance. The spectra (a) and (b), related to the samples milled with TiF₄ and NbF₅, besides the two polymorphs (γ and β), show reflections of CaF₂ (PDF # 35-819).



neither traces of CaF₂ nor of the TM-fluorides.

Fig. 1: XRD of the samples after ball milling: $Ca(BH_4)_2 + TiF_4$ (a), $Ca(BH_4)_2 + NbF_5$ (b), $Ca(BH_4)_2 + VF_3$ (c), $Ca(BH_4)_2 + TiF_3$ (d), $Ca(BH_4)_2 + VF_4$ (e). γ - $Ca(BH_4)_2$ (γ); β - $Ca(BH_4)_2$ (β); $CaF_2(\diamondsuit)$; $TiF_3(\blacksquare)$.

Its presence hints to a reaction between the TM-fluoride and the borohydride already during milling. In contrast, in pattern (d) we can still find peaks of TiF_3 (ICSD # 28783), indicating that no reaction has taken place in the vial. Patterns (c) and (e), with VF_3 and VF_4 respectively, present

Desorption reactions were studied under static vacuum by heating from room temperature up to 450 °C. Rietveld refinement of the pattern collected at 30 °C indicates that the initial powder is composed of 72 wt. % low temperature polymorph γ -Ca(BH₄)₂, 24 wt. % high temperature β -Ca(BH₄)₂ and 4 wt. % CaF₂ (PDF # 35-819). The latter one must have been formed by a reaction between Ca(BH₄)₂ and NbF₅. This might partly explain why the total amount of hydrogen desorbed from the samples with additives is lower than in case of the

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pure calcium borohydride. The pattern measured at 282 °C shows that there are no traces of γphase anymore, since it is already transformed into β-phase. At this temperature another calcium borohydride polymorph appears, called δ -phase in a previous work. In that study, it was reported that hydrogen desorption from β-Ca(BH₄)₂ occurs in the range of 330 °C to 380 °C, whereas δ -Ca(BH₄)₂ releases hydrogen at higher temperature, in the range of 380-500 °C. When TM-fluorides are added to calcium borohydride, its kinetics of desorption is modified and the thermal events are shifted. As can be seen in Figure 2, the first hydrogen desorption step takes place between 282 °C and 361 °C and involves only the β-phase. In this range, most of the hydrogen is desorbed with some residual being present inside the δ -phase. This phase fraction grows continuously up to 361 °C, as can be inferred from the increase of the intensity of its diffraction peaks. At this temperature, CaF2 is still present and coexists with the δ -Ca(BH₄)₂ phase. The residual hydrogen is released in the 361-450 °C temperature range. CaH₂ should be formed after the first desorption event, however, our measurements do not show its formation up to 361 °C. At 450 °C CaO and an unidentified phase form. The latter decomposes after a short time. Later on, the last recorded XRD pattern at 450 °C shows an increase of the peak intensity at the scattering vector value of 2.23. This peak corresponds to the (200) peak of the CaF_{2-x}H_x phase⁸ which is formed by a reaction of CaH₂ and CaF₂ and can explain the absence of any CaH₂ reflections. The ICSD database matches those peaks with an exact stoichiometry CaF_{0.76}H_{1.24}, however, the current experimental results are not conclusive, since not all of the peaks are present.

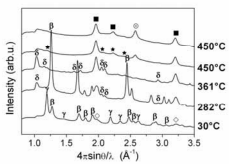


Fig. 2 SR-PXD patterns of Ca(BH₄)₂ milled with NbF₅. The experiment was carried out by heating in vacuum from RT up to 450 °C with 5 K/min as heating rate and 10 minutes isotherm (at 450 °C). γ -Ca(BH₄)₂ (γ); β -Ca(BH₄)₂ (δ); CaF₂ (δ); δ -Ca(BH₄)₂ (δ); unidentified phase (\bigstar); CaO (\otimes); CaF_{2-x}H_x (\blacksquare).

This study showed that when TM-fluorides were added to Ca(BH₄)₂, CaF₂ was formed as side product contributing to reduce the hydrogen content of the mixtures.

Formation of $CaF_{2-x}H_x$ was detected in the decomposition products due to the reaction between CaH_2 and CaF_2 . Furthermore, *in-situ* XRD represented a key investigation tool for the understanding of the multistep decomposition reaction of the calcium borohydride/TM-fluoride system.

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Peptide Self-Assembly

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The self-assembly and aggregation of peptides have been widely studied in recent years. $^{1-3}$ Much of the interest can be traced to the fact that many neurodeseases are consequences of protein or peptide aggregation. As the bottom-up approach through e.g. self-assembly appears to be most reasonable route for the fabrication of nanostructures, peptides self-assembly is also considered in the design of nanomaterials of desired functionality. 4,5

In this study, we demonstrate our results for the study of self-assembly structures of short synthetic oligopeptides A_8K and $A_{10}K$ in water over a wide range of concentrations. In our previous work⁶, we studied the self-assembly structures, including the formation of nematic ordering. SAXS data show that the thickness of the nanotube wall, δ , is less than 1 nm indicating that the peptides form a monolayer in the nanotube wall. The nanotubes have a very large aspect ratio and form an ordered phase nematic or hexagonal phase. Following that work over high concentrations⁷, it was shown that because of the low δR ratio, the nanotube volume fraction grows very rapidly with increasing peptide concentration, ϕ , and reaches close packing already at ϕ =0.15. When increasing the concentration further, there is a phase transition to a novel lamellar phase where the peptide molecules form bilayers consisting of two, presumably oppositely oriented, monolayers.

In this part of the work, the studies were extended to peptides that are slightly more hydrophobic. Two peptides, A_8K and $A_{10}K$ are explored over a range of concentrations and the self-assembly structures were investigated using cryo-transmission electron microscopy (Cryo-TEM) and small angle X-ray scattering (SAXS). Small-angle X-ray scattering experiments were conducted on beamline I711 at the MAX II storage ring in Lund, Sweden, using a sample-to-detector distance of 1.5 m and a wavelength, λ =1.025 Å

As expected from hydrophobicity, the apparent critical aggregation concentrations for these peptides are also much lower than that of A_6K . SAXS patterns of A_8K obtained at peptide concentrations from $\phi = 0.1$ to 0.25 are measured.

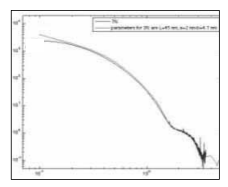


Figure 1. Theoretical Model showing the scattering pattern of a system with elliptic cylinders (green) and the the actual data of 7% (blue).

The observed pattern can be modeled with an elliptic cylinder form factor where an elliptic cross-section with dimensions of 2.0 nm and about 4.5 nm is suitable for concentrations up to 10% as presented in Figure 2. It is noteworthy that within this range, the size of the cross-section remains constant.

The length of the tubes are not distinguishable in this q-range which suggest a length longer than about 40 nm, which gives a high aspect ratio for the cylinders. These results are in good agreement with cryo-TEM for concentrations that can be imaged.

SAXS patterns of $A_{10}K$ obtained at peptide concentrations from $\phi = 0.1$ to 0.15, are also measured. This data can also be modeled using an elliptic cylinder form factor where an elliptic cross-section with dimensions of 2.0 nm and about 4.5 nm is suitable for concentrations up to 10%.

This data, coupled with cryo-TEM, reveal that both A_8K and $A_{10}K$ form elliptic cylinders with a finite size, yet high aspect ratio; exhibiting behavior very different from the previously investigated peptide, A_6K . Although different only with a slight increase in the hydrophobicity, the self assembly structures assumed by these peptides are totally different from the infinitely long hallow nanotubes observed previously by A_6K or any other structure peptides assume such as twisted ribbons or fibrils that are commonly seen in the literature.

In summary, we have investigated self assembly of oligopeptides with varying hydrophobicities and have shown that this plays an important role in the type of self assembly structure that forms. This simple system lends itself for systematic investigations of peptide self-assembly including e.g. investigation of slightly different peptides such as A_6R (R: Argenine; cationic) and A_6D (D:Aspartate; anionic) where the end charge is varied.

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Towards Understanding Peptide Self-Assembly

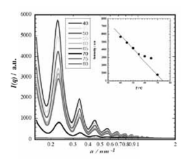
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Peptides can self-assemble into various structures due to hydrophobic interactions. There is a big interest in the subject, not only because these structures have a wide range of applications¹⁻⁶, but also peptide self-assembly is a naturally occurring phenomenon in the body. In some cases, these aggregated structures accumulate in a particular part of the body, causing irregularities, which lower the quality of life significantly, and even causing death sometimes. For example, recent studies showed that amyloid, which is a fibrous protein aggregate, is linked to Alzheimer's Disease^{7,8}, an incurable neurodegenerative disease. Therefore, there is a growing interest in understanding the mechanisms behind peptide self-assembly.

Our study is an ongoing study of investigation of a model peptide system. We investigate the self-assembly behavior of the peptide A_6K . Our main tool is Small Angle X-ray Scattering (SAXS), supported by Cryogenic Transmission Electron Microcopy (Cryo-TEM). We performed our SAXS experiments in MaxLab, in I711 beamline station at the MAX II storage ring in Lund, Sweden, using a sample-to-detector distance of 1.2 m and a wavelength, λ =1.1 Å.

 A_6K peptide self-assembles into very long, hollow nanotubes of radius 26 nm when the concentration is above a certain aggregation concentration, $\phi_{cac} = 0.10$. We studied a series of different concentrations ($\phi = 0.07$ –0.53) to see how the scattering changes as concentration changes in the previous years and constructed a phase diagram. Apart from concentration studies, we also studied the temperature effect on the system. The dissolution of nanotubes was observed upon increasing the temperature. For a sample of $\phi = 0.12$, we recorded the SAXS patterns at different elevated temperatures, from 40 to 80 °C in 5°C steps. The results are presented in Figure 1. From 40 to 70 °C we see the characteristic Bessel function scattering pattern, showing that the nanotube structure is retained as the temperature is increased. However, the scattered intensity decreases with increasing temperature. This means that the nanotube concentration is decreasing, which in turn is a consequence of an increase of ϕ_{cac} with increasing temperature. In the insert of Figure 1, the scattered intensity is plotted at q=0.23 nm⁻¹, corresponding to the second side maximum of the nanotube form factor.



As can be seen, the scattered intensity decreases approximately linearly with increasing temperature and vanishes at circa T = 74 °C. At 75 °C and 80 °C we only observe the weak scattering from monomers and hence our concentration is below ϕ_{cac} at these temperatures. From this data we can thus conclude that ϕ increases with temperature approximately linearly.

Another interesting result was obtained when we attempted to align our nanotubes and do WAXS on a sample of ϕ = 0.13. We worked at the same station; I711 with the same wavelength of 1.1 Å but the sample to detector distance was 25.5 cm. By this approach, we aimed to get Bragg peaks, which would provide information on the peptide orientation within the nanotube. We already had results with X-ray Diffraction (XRD) method⁹, which was performed in the University of Reading in the UK. We simply centrifuged a capillary, filled with our sample, for 2 minutes in a centrifuge machine at 4000 rpm. The results we obtained were very similar to XRD results. As a result, we were able to show that we could perform diffraction experiments at I711 station in MaxLab with appropriate experimental set-up.

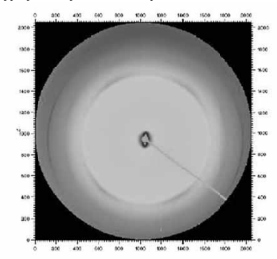


Figure 2. 2D image of WAXS result from the centrifuged sample of ϕ = 0.13. By aligning the nanotubes with centrifuging, we were able to get information about the local orientation of the peptides within the nanotube wall.

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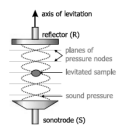
Silk assembly in acoustically levitated droplets

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At the heart of protein's aggregation are specific molecular events; and how they are modulated by changes in solvent conditions is key to our understanding and control of fibrillogenesis. A certain class of structural fibrous proteins may provide the answer. Indeed, spider silk and silkworm silk proteins have evolved to readily form 'insoluble' ordered structures. But, to date little is known about the series of association steps involved in the nucleation and growth of silk protofilaments. To characterise this process we must clearly establish silk proteins inter- and intra-molecular behaviour in solution. One major hurdle to achieve this goal is the handling difficulty inherent to silk. We therefore, turn our attention to acoustically levitated droplets. Suspending a low concentration silk droplet and leaving it to gently dry provides the perfect setup to monitor silk association and assembly in solution. In the present contribution we demonstrate that in-situ SAXS on levitated silk proteins is possible and that silk proteins as a function of concentration can assemble into well ordered structures.

Acoustic levitation is a proven technology and has found numerous areas of application in recent years. In figure 1 we show the working principle of an acoustic levitator, at the pressure nodes droplets can be suspended and maintained stable for at least 20min in air. To standardize the SAXS-Levitator we used the protein lysozyme as a model. Figure 2 shows the scattering intensity of a drying drop of lysozyme. The intensity was comparable to the known and reported lysozyme scattering intensities. Note in figure 2, the appearance and shift of the structure peak in the middle q range.



Α

frequency range of 15 to

wave

standing

with

Figure

acoustic

100kHz.

Figure 2. Standard protein, lysozyme, used to test the levitation setup. SAXS intensities of the drying drop show the emergence of a structure peak in the middle q range as the concentration of

protein increases (bottom to top).

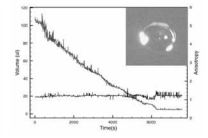
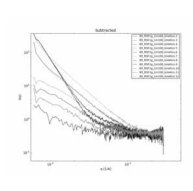


Figure 3. Typical drying and concentration plot for lysozyme. A concentrating factor of 25 was calculated by estimating the drop volume during the experiment (decaying curve). Additionally the drop anisotropy was monitored (flat curve)

We further monitored the levitated drop morphology by filming it during the experiment. Figure 3 shows a composite plot of the estimated drop volume (red curve) and its anisotropy (black curve) as a function of time. In the inset, an image of the starting drop is shown. We found that the drop volume decreased by a factor of 20-30 fold whilst retaining its shape. This meant a progressive increased to final concentration 20-30 times higher than the initial one. Having determined the working conditions for the levitator on the I711 beamline, we investigated the behaviour of silk protein solutions. For this experiment we used regenerated silk proteins from silkworm cocoons prepared as in reference ². Sample proteins were extensively dialysed to remove ionic species.

In figure 4 we show the radial averaging of a drying droplet of silk proteins. Under normal circumstances such a plot would suffice to examine the association and assembly. We found, however, that as the concentration increased silk proteins adopted an anisotropic alignment. Figure 5 summarises the drying kinetics, this time seen from the 2D scattering images.



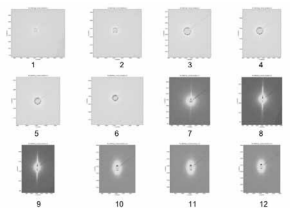


Figure 4. Radial averaging plot of the SAXS intensities of a silk droplet drying over a 20min period.

Figure 5. 2D SAXS images of the same silk droplet drying over 20min as in figure 4. Frame 7 to 9 show a likely edge effect from the drop relatively to the beam size. Overall, it is clear that upon increasing concentration silk molecules go from an isotropic distribution to an anisotropic ordered assembly at high concentration.

A closer look at the drop shape (anisotropy) showed a slight flattening in the horizontal plane. This change in shape can be correlated to an anisotropic alignment in the vertical plane of the 2D images (figure 5, frame 10 to 12). Interestingly, this alignment of silk proteins suggest an inbuilt concentration driven mechanism that allows silk proteins to organise themselves prior to spinning.

The results presented will shed some light on two central questions of in silk research: (i) what are silk storage structures and shapes; and (ii) what is the mechanism of silk fibre formation. It is interesting to note that until now it was believed that to process silk one needed 'colloidal aggregate'. ^{3, 4} The results we present suggest a liquid crystalline phase to promote fibre formation.

Ultimately, the mechanism of silk fibrillogenesis is not clear.⁵ The two current views, one involving liquid crystallinity ⁶ and the second involving micellar assembly ³ rather than being conflicting can be reconciled by the present results. It is indeed conceivable that oriented domains exist within larger micellar assemblies.

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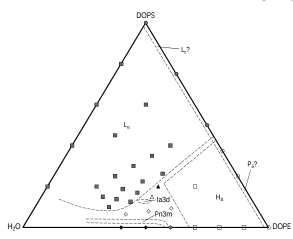
CUBIC PHASES IN THE DOPS/DOPE/WATER SYSTEM

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The project aims at mapping the lyotropic and thermotropic phase behavior of the DOPS/DOPE/water system, with particular focus on finding and characterizing any potential bicontinuous mesophases.

Dioleyl phosphatidyl ethanolamine (DOPE) is known to form mostly an inverted hexagonal structure (H_{II}) in aqueous systems (lamellar liquid crystalline (L_{α}) at low temperatures and medium hydration)¹⁻³, but reports also exist of cubic phases at low temperatures.²⁻⁴ No cubic phases at room temperature have been reported so far. Dioleyl phosphatidyl serine (DOPS) is less studied, but is known to form L_{α} structures upon hydration.



Synchrotron SAXS light and microscopy with crossed polarizers have been used to construct a tentative phase diagram, see Figure 1. Two bicontinuous cubic phases, Pn3m and Ia3d, have been identified, the position tentatively which have calculated from a swelling analysis of the cubic parts of the 2-phase samples. The Pn3m resides close to the DOPE side and at medium hydration, while the Ia3d is closer to DOPS/DOPE 35/65, also at medium hydration. Unfortunately, no single phase cubic samples have been identified so far.

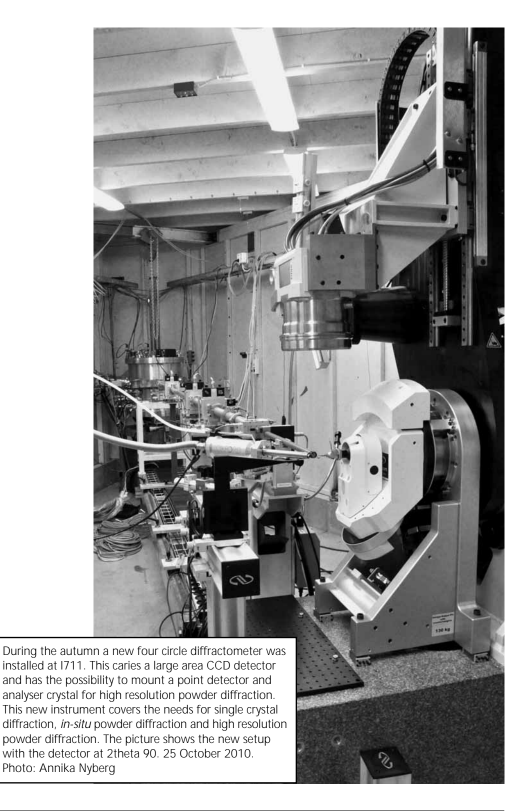
For DOPS/water a L_{α} phase with a maximum swelling above 80% water is found, and for DOPE/water a H_{II} phase with a maximum swelling at 32% water.

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Structural dynamics of the LiFePO₄ lithium-ion battery cathode material

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The properties of the cathode material LiFePO₄ used in lithium-ion batteries have been well studied ever since the first experiments showing the lithium intercalation capabilities of this material [1]. However, the intercalation mechanic is still not well understood. To be able to study this, an experimental technique capable to study this phase transition as it happens in an electrochemical cell is of great importance.

One possible way is to use *in-situ* X-ray diffraction [2]. We have used a synchrotron light source in transmission mode, λ =0.9084Å, MarCCD at the I911-5 beamline at MAXlab, Lund, Sweden and beamline I711, λ =1.06Å with an Oxford Diffraction Titan CCD.

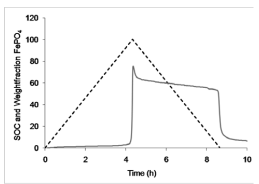


Figure 1: Comparison between the state of charge dashed black line (SOC, calculated from the amount of charge delivered to the battery), and the phase amount of FePO₄ grey line, calculated from diffraction data.

Batteries were assembled as coffee-

bag cells (75% PhosThech LiFePO₄ 10% carbon black 15% PVdF, Solupore separator, 1M LiPF₆ EC:DEC 2:1 electrolyte, lithium foil counter electrode) which were then pre cycled with C/10 for three cycles between the cut off voltages 4.2V and 2.7V (measured capacity about 150mAh/g). The cells were then examined by *in-situ* X-ray diffraction.

Slow discharge and charge

Cells were cycled galvanostatically (C/4.3 or C/4.5) while monitoring the diffraction pattern with a frequency of about one diffraction pattern per minute. Rietveld refinement (Fullprof software [3]) of the diffraction patterns was used to determine the fraction of each crystalline phase (calculated from scale-factors) in the battery as a function of time. The results for experiments with the shorter wavelength can be seen in figure 1. There is a great difference between the state of charge (SOC), calculated from the amount of charge delivered to the battery (dashed line in figure 1), and the amount of the phase FePO₄, determined from the diffraction pattern. Previously Chang *et al.* have suggested the interface between the phases to be metastable[4]. However, when the in –situ experiments are repeated with at longer wavelength this rapid change in phase composition is not detectable. At longer wave-lengths the fraction of FePO₄, when inspecting the diffraction data, follow the state of charge (as could be expected as LiFePO₄ loses its lithium and converts to FePO₄ continuously during charge) as previously seen by Andersson *et al.* [5].

Fast discharge and charge

Experiments were also done by applying constant potential to the cells monitored the current response diffraction on the I911-5 beamline. Fully charged cells were discharged under a 2.7V potential and fully discharged cells were charged by applying 4.2V). To diffraction patterns were recorded with a frequency of about 4 each minute. Almost all of the material change phase during the first few minutes of the potential step. A selection of the diffraction patterns from this time period are seen in figure 2. The difference between the peak width changes of both phases can be seen as an indication the difference between the lithiation and delithiation processes. The broad peaks of the growing LiFePO₄ phase during discharge can be explained by larger strains in the materials than during the charge or by multiple nucleation of the new phase and thus the creation of many small LiFePO₄ domains instead of one big in each crystallite.

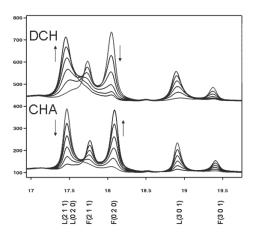


Figure 2. Diffraction pattern for potential step to 2.7V, discharging the battery (upper) and potential step to 4.2V, charging (lower) for a LiFePO $_4$ electrode. The axis in the figure $are2\theta$ with peak indices marked (L=LiFePO $_4$, F=FePO $_4$) and intensity. The difference in peak width during the charge and discharge is clearly visible and also that the peak width for the two phases are different. Most notable are the broad peaks of the growing LiFePO $_4$ during discharge, as compared to the growing FePO $_4$ during charge.

Conclusions

There is an interesting difference between

the measurements done at different wavelengths. This is previously not reported in the literature and might together with computational chemistry give better understanding of the properties of the LiFePO₄ cathode material. We are also currently investigating how rapid charge and discharge of the batteries can best be examined by *in-situ* X-ray diffraction and how to fully utilize the new Oxford Diffraction Titan CCD on the I711, as it can easily circumvent the limitations of the MAR system when used on highly textured and high background samples as fully functionally batteries.

Acknowledgements

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Clay delamination in saline water during heating

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Clay mineral particles in the form of layered, plate-shaped smectites receive attention in areas ranging from geology and petrology to material science and basic physics. A central property of smectites is their capacity for uptake of water and other species into the nanopores that are formed when several unit clay layers stack to form lamellar particles. Because smectite clays are hydrated at ambient temperature, pressure and humidity conditions – and also often in geological environments – the interaction of smectites with water is crucial. The present report concerns x-ray scattering studies conducted at I711 on delamination and swelling of the synthetic smectite Na-fluorohectorite suspended in saline water as a function of temperature.

When subjected to an atmosphere of sufficient water vapor content, or to direct contact with liquid water, layered smectite particles intercalate water molecules, causing the particles to swell in the stacking direction. This water uptake is associated with a stepwise increase in the spacing of layers, giving rise to a nomenclature where hydration states are referred to as intercalating zero, one, two or three water-layers [1]. However, when clay is dispersed in water, a point may eventually be reached beyond which no well-defined hydration states are stable. At this point the system transitions into the regime of osmotic swelling, as stacked particles delaminate and their unit layers become separate colloids surrounded by double layers of counter-ions.

In the current project, changes to hydration state and structure in 10 mM NaCl aqueous dispersions of Na-fluorohectorite were investigated in situ by transmission mode diffraction as a function of temperature. Figure 1 shows the changes in scattered intensity obtained upon heating from 25 to 78 $^{\circ}$ C.

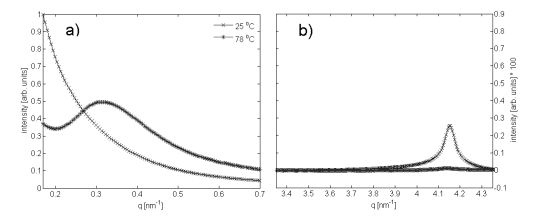


Figure 1: Azimuthally integrated scattering recorded at 25 and 78 $^{\circ}$ C: a) the effects of heating at small scattering angles, and b) the changes in the two water-layer 001 Bragg reflection that originates from the structural periodicity along the stacking direction of layered particles.

From Figure 1b), it is evident that the overall 001 Bragg diffraction intensity has decreased as the sample was heated. The position of the 001 peak, at $q=4.152~\mathrm{nm}^{-1}$ at 25 °C, corresponds to the two water-layer hydration state [2, 3]. The disappearance of this peak with temperature could either result from a change in the hydration state of the clay particles or from a degradation of particle crystallinity.

An approximate structure factor can be obtained by dividing the overall intensity by a fitted Aq^{-x} form factor contribution found for $q \gtrsim 0.4 \, \mathrm{nm^{-1}}$. Preliminary analysis indicate a structure factor characteristic of a 'dense' system with repulsive interactions [4], especially at the highest temperatures. We interpret the loss of two water-layer 001 peak intensity along with the relaxation of the scattering curves for $q \gtrsim 0.4 \, \mathrm{nm^{-1}}$ towards a q^{-2} behaviour, as evidence of particle delamination at elevated temperatures. The significant structural changes in the dispersions accompanying delamination shows characteristics associated with a repulsive Wigner glass. We are currently continuing the study of this temperature induced transition with scattering at smaller angles and rheological studies.

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Alizarin red S modifies apatite nanocrystal size and formation kinetics

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The hard parts of the human body, bone and teeth, contain an inorganic biominerals related to the geological apatite with the idealized formula $Ca_{10}(PO_4)_6(OH)_2$. In bone, the mineral is present in the form of nanocrystalline plates of roughly $45\times20\times3$ nm 1 . The extremely thin dimension of only ~3 nm is parallel to the hexagonal a-axis of the bioapatite. There is a large interest in synthesis of synthetic apatite nanoparticles for use in biomaterials. Therefore we have initiated a research program where additives are used to control the crystal formation and crystallite size. Herein we report on how the dye molecule alizarin red S (ARS) modifies apatite formation 2 . ARS, see Figure 1, is used to dye bone and/or calcifications *in vitro* and *in vivo* in experimental animals.

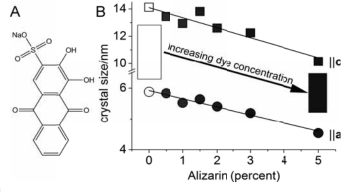
The nanoparticles were synthesized by mixing aqueous solutions of $CaCl_2$ and $Na_3PO_4/NaOH$ according to 3

$$10 \ CaCl_{2}(aq) + 6 \ Na_{3}PO_{4}(aq) + 2 \ NaOH(aq) \rightarrow Ca_{10}(PO_{4})_{6}(OH)_{2} \ (s) + 20 \ NaCl \ (aq)$$

giving a final $CaCl_2$ concentration of 0.2 M. ARS was added to the phosphate solution prior to mixing. The final concentration of ARS will be reported as a percentage of the calcium concentration. The resulting powders were characterized by in house powder X-ray diffraction (PXRD), TGA and

FTIR while the reaction kinetics were probed by *in situ* pH measurements and *in situ* synchrotron PXRD. Full details can be found in reference 2.

During the synthesis, an amorphous gel is formed first, which then crystallizes leading to a suspension of nanocrystal aggregates. *Ex situ* PXRD showed that ARS reduced the size of the nanocrystals. These are rod-shaped and the size, found by full-pattern Rietveld refinement, is reduced essentially linearly with ARS concentration as shown in Figure 1B. Above 5% ARS led to



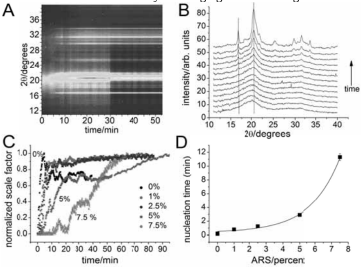
amorphous powder after 24 h reaction time at room temperature but prolonged reaction time showed that the samples eventually became nanocrystalline. This suggested that ARS predominantly functions by inhibiting crystal nucleation rather than hindering growth. Therefore we undertook *in situ* time resolved synchrotron PXRD measurements.

The *in situ* PXRD measurements were performed at ID7-11 at MAXLAB using a 0.7 mm diameter single crystal sapphire capillary as reactor. The solutions were pumped by a peristaltic

pump and mixed in a linear mixer just prior to entry into the capillary. The measurements were performed at 70±2 °C to speed up the reactions thus allowing several measurements to be made with a reasonable amount of measurement time. The diffracted X-rays were detected with a MARCCD detector using 4 or 15 s exposure times resulting in a best time resolution of 6.5 s when accounting for detector readout. Typically the time resolution was a little worse than this due to data transfer times over the computer network; all measurements were assigned a measurement time based on the actual time stamp of the data file on the computer. The wavelength was 0.9983 Å.

Figure 2 shows a summary of the *in situ* data. Data were collected for ARS concentrations of 0, 1, 2.5, 5 and 7.5%. The measurements are very challenging due to the large diffraction

background of water that A dwarfs the diffraction signal from the minute nanocrystal sample. In spite of these challenges, quality data were obtained as illustrated in Figures 2A and 2B that display background subtracted data. Initially, the diffraction signal displays broad peaks around 20 and 30° 2θ as seen in the first 6 graphs displayed in Figure 2B. Then broad diffraction peaks consistent with the apatite lattice appear and grow. The data were fitted Rietveld refinement. The scale factors, which provide a measure of the amount of crystalline material, are shown in Figure 2C. For 0% ARS, there is a very short nucleation time before the scale factor be-



gins to rapidly rise. This time gets longer for higher ARS concentrations. After nucleation, the scale factor curves present many bumps. This is not due to poor data quality but rather due to the fact that the initially formed amorphous gel breaks down into clumps of high calcium phosphate concentration in a depleted solvent. These nanocrystal containing high concentration clumps can slowly diffuse in and out of the beam leading to the observed jumps in apparent scale factor. Due to this physical phenomenon, we do not analyze the crystal growth kinetics in detail here. Instead we plot the nucleation time in Figure 2D. It is observed to increase exponentially with added ARS content confirming that the primary effect of ARS is to hinder nucleation. These studies combined with others currently underway will be used to map the roles of additives on apatite formation and will hopefully shed light on how the crystals form in bone.

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Soluble Complex Salts of Surfactant Ions and Polymeric Counterions: Composite Macromolecular Self-Assembly

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The aim of this project is to understand the physical chemistry of soluble complex salts in aqueous solution and to gain information about how the balance of the intermolecular interactions will affect the structures, the packing and the water miscibility of the resulting mixed surfactant aggregates. A complex salt is defined as the neutral salt of surfactant ions (aggregated into highly charged micelles) that interact with a polyelectrolyte chain, which in turn acts as a large counterion (a polyion). The complex salts studied in this work consist of polyacrylate (PA_v) and cationic C₁₆TA⁺ surfactant ions aggregated into micelles [1]. They are denoted C₁₆TAPA_v, where y is the degree of polymerization (y=25 or 6000). Both are insoluble in water. To make these complex salts soluble, poly(ethylene oxide) (PEO)containing nonionic surfactants of the type C_iE_i (E stands for PEO) are added and C₁₆TA⁺/C_iE_i mixed micelles surrounded by PA⁻_v chains are formed. Phase studies on systems containing C₁₆TAPA_v complex salts in water mixed with either C₁₂E₅ or C₁₂E₈ have been carried out by visual inspection of the samples and small-angle Xray scattering (SAXS) at the beam line I711 at MAX II been employed to characterize the different structures found in the ternary phase diagrams [2]. We used a wavelength of 1.1 Å and a 2D CCD camera. At I711 we had several different measurement sessions and the sample to detector distance varied somewhat, 1403 mm and 1245 mm respectively. All measurements were performed at 25 °C and under vacuum. Typical measurement times were 300-600 seconds. The data obtained analyzed using the software FIT2D [3].

The results revealed that decreasing polyion length and increasing the PEO-chain length of the nonionic surfactant were important factors for increasing the solubility of the complex salt [2]. We also found that the curvature effects are quite small at low water content when gradually exchanging $C_{12}E_8$ with either one of the complex salts while there is a gradual change in curvature for the systems containing $C_{12}E_5$. Another interesting observation was the possibility for relatively large amounts of complex salt to be incorporated into a V_1 (la3d, bicontinuous) phase in the $C_{12}E_8$ -containing systems. In the dilute regime several different liquid crystalline phases were found in equilibrium with a dilute liquid phase containing the nonionic surfactant. The $C_{16}TAPA_{25}/C_{12}E_8/H_2O$ ternary phase diagram is given in Figure 1. Figure 2 presents a SAXS curve showing the hexagonal structure in a two-phase sample of the same system.

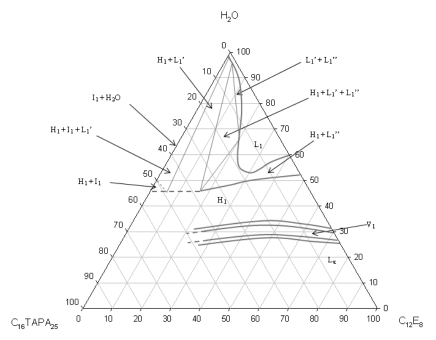


Figure 1. Ternary phase diagram of $C_{16}TAPA_{25}/C_{12}E_8/H_2O$. One-phase areas (bold lines), boundaries to three-phase areas (full lines), boundaries not exactly determined (dashed lines). Compositions in wt %. Notations: liquid phases (L_1 and L_1), liquid micellar phase (L_1 "), hexagonal phase (H_1), bicontinuous) phase (V_1), micellar cubic phase (I_1) and lamellar phase (I_2).

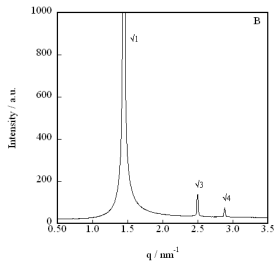


Figure 2. SAXS data demonstrating the hexagonal structure from the $H_1 + L_1$ ' two-phase area in the $C_{16}TAPA_{25}/C_{12}E_8/H_2O$ system. Compositions in wt %: (20.1 : 10.0 : 69.9).

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Anisotropic crystal growth kinetics of anatase TiO₂ nanoparticles synthesized in a non-aqueous medium

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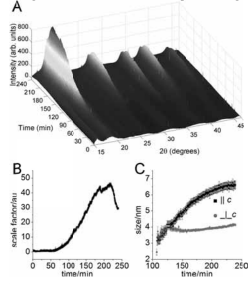
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Titania nanoparticles are the subject of intense investigation due to their use in catalysts. There is a wealth of synthetic methods yielding titania nanoparticles. One convenient method employs benzyl alcohol as the solvent^{1,2}. To improve our understanding of the mechanisms of crystallization in general and of non-aqueous metal oxide syntheses in particular, we have undertaken studies of the crystallization and aggregation kinetics of anatase titania nanoparticles synthesized in benzyl alcohol using *in situ* synchrotron powder diffraction, *in situ* laboratory small angle X-ray scattering and turbidity measurements³. Herein we will discuss the powder diffraction (PXRD) data only and refer to reference 3 for full details.

The synthesis was performed by pre-hydrolyzing TiCl₄ in anhydrous ethanol prior to addition to benzyl alcohol and then formation of the tetragonal anatase form of TiO₂ upon heating.

This was originally done to reduce the violence of the hydrolysis reaction⁴. The presence of ethanol leads to the formation of anisotropic crystals as evidenced by both TEM4 and *ex situ* PXRD3. Thus, the present study allows insights into the factors governing anisotropy in crystal growth. The *ex situ* diffraction data clearly showed that the crystals were anisotropic in shape with the size along the *c*-axis being larger than that in the *a*,*b*-plane.

We performed in situ PXRD at beamline I7-11 at MAXLAB. The liquid sample was placed in a 1 mm borosilicate capillary and heated to 85 °C using a home-built furnace employing a hot air stream. An 1.1072 Å X-ray beam was employed in combination with a MAR165 CCD detector. With an exposure time of 30 s, the time resolution was 34.3 s per data frame when accounting for detector read out. The diffraction patterns could be refined by Rietveld refinement with the exception of the first few frames that essentially did not contain any diffraction information therefore only allowing scale factor refinement. Figure 1A displays background substracted raw data. After an initial induction time, crystals nucleate and grow.



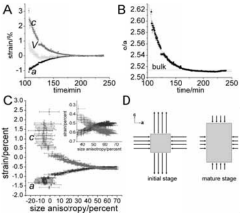
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Rietveld refinement allowed for extraction of very detailed information on the growth behavior. The scale factor, which reflects the amount of material in the crystalline state, is shown in Figure 1B. After the nucleation time it increases steeply and reaches a plateau. Thereafter it falls slightly due to sedimentation of particles in the capillary caused by the onset of aggregation of crystallites. More interestingly the Rietveld refinement also gave detailed information on the crystallite size and shape as shown in Figure 1C. The crystallites are initially spherical (note that Figure 1C only displays data after the time point where crystallite size information could be refined) but after \sim 125 min whereafter the size in the a,b-plane essentially stops increasing and the crystals become increasingly needle shaped.

The observation of a transition from spherical to needle crystal shape makes one wonder what driving forces are responsible for this behavior. Figure 2A shows the time evolution of the lattice constants plotted as relative deformation (with respect to long times). Initially the *a*-axis displays a compressive strain as expected for nanoparticles. Interestingly, the *c*-axis displays a positive and large strain that relaxes exponentially towards a stable value for the mature crystals. The unit cell anisotropy ratio *c/a*, shown in Figure 2B, starts being ~4% larger than in bulk anatase and relaxes exponentially towards the bulk value as the crystals grow.

The surprisingly large positive strain along the *c*-axis suggests that there is a penalty to pay for the initial essentially spherical crystal shape and that relaxation of this positive strain is related to the onset of anisotropic crystal shape. This is confirmed by considering the strain along the two axes as a function of the size anisotropy, which is shown in Figure 2C (plotted with respect to literature reference lattice constant values). During the initial stages of growth, the particles



are spherical with size anisotropy close to zero and large strains. The onset of positive size anisotropy is very close to the cross-over from positive to negative strain along the c-axis. The situation is recapitulated in Figure 2C that schematically illustrates the large positive strain in the initial isotropic crystals and the large strain reduction seen when the crystals mature. Thus the driving force for crystal shape anisotropy is found to be related to lattice strain. The lattice strain is likely to be governed by the presence of two solvents ethanol and benzyl alcohol that bind with different preference to the (001) and $\{100\}$ planes.

The insights obtained through these studies highlight the role of solvents in wet chemical syntheses of nanocrystalline materials and beautifully illustrates how mechanistic insights can be obtained by time resolved in situ synchrotron powder X-ray diffraction.

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High pressure X-ray scattering experiments of polyfluorenes at I711

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We carried out on an X-ray scattering experiment of bulk poly[9,9-bis(2ethylhexyl)fluorene] under quasi-hydrostatic pressure from 1 to 11 GPa at room temperature¹. We found that the scattering pattern of high molecular weight (HMW) polyfluorene (>10 kg/mol) undergoes significant changes between 2 and 4 GPa in bulk phase (Fig. 1). The 110 reflection of the hexagonal unit cell disappears, indicating a change in equatorial intermolecular order. The intensity of the 00 21 reflection drops, with a sudden move towards higher scattering angles. Beyond these pressures, the diminished 00 21 reflection tends to return towards lower angles. These changes may be interpreted as a transition from crystalline hexagonal to glassy nematic phase (perceiving order only in one direction) and rationalized by density arguments and underlying theory of phase behavior of hairy-rod polyfluorene. Also the possible alteration of the 21-helical main chain towards more planar main chain conformation can be discussed. We find, moreover that the scattering of low molecular weight polyfluorene (<10 kg/mol), which is glassy nematic in ambient pressure, is reminiscent with that of HMW polymer above 2-4 GPa. The transition is concomitant with the previously reported transition seen by Raman scattering².

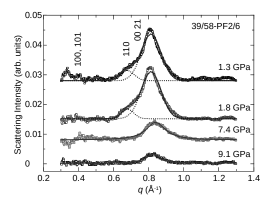


Figure. 1. Scattering patterns of PF2/6 for selected pressures measured on 711 beamline.

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Robert Corkery, YKI, Institute for Surface Chemistry, Stockholm, is preparing his samples to be measured at beamline I711, 5 May 2010.

Photos: Annika Nyberg

Structural Analysis of Wheat Gluten Biomaterials containing Urea and Nanoclays using SAXS

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Wheat gluten (WG), a biobased polymer, in blend with glycerol and other additives, i.e. clays, showed a number of advantages to be used in biomaterials^{1,2}, which are good alternative to synthetic polymers in *i.e.* packaging applications.

Additives, sodium hydroxide³ and ammonium hydroxide⁴, as well as temperature and mechanical energy input⁵ have been shown to affect molecular interactions and polymerisation of the WG proteins, and end use quality of biomaterials. Also, natural montmorillonite (MMT) clays (Cloisite® Na⁺) slightly increased stiffness and strength of the materials², while urea (protein denaturant) is known to be an effective pH regulator for preparation of protein films. The aim of our study was to study protein polymerization and structure in WG films containing urea, nanoclays and urea-nanoclays.

In this report we show the structural characterization of WG materials produced in various temperatures and containing the following additives: 1) urea (10%, 15% and 20%) produced by extrusion (Fig.1a); 2) natural and organically modified MMT clays, Na⁺ and Cloisite 15A, respectively, (5%) wt (Fig.1b) and 3) 10% urea+5% of each natural and organically modified MMT clays (Fig. 1c) produced by injection moulding. The small angle X-ray scattering (SAXS) was performed at the beamline I711 at MAX-lab Synchrotron. The wavelength was 1.1 Å and the sample to detector distance was 1403 mm.

All 2D SAXS images for the WG-urea samples, extruded at 130° C, were found being isotropic, while one dimensional intensity plots were influenced by the urea concentration in the sample: higher urea concentration correlated with greater inter-domain distance a, 10% urea with 68Å, while 20% urea with 72Å, respectively (Fig. 1a). We have found the specific WG protein assemblies in urea containing films which seemed to present a well defined supramolecular structure. The film morphologies could be well fitted with a hexagonal close packed (HCP) structure (Fig. 1a, three reflections indicated by arrows).

For the WG-clay injection moulded composites (Fig. 1b, black line) with natural MMT clay, anisotropic SAXS pattern was observed indicating the preferential orientation of the silicate layers due to the processing conditions (i.e. injection moulding). However, no temperature effect on the composite structure was observed. Similar SAXS patterns were also obtained for the WG-modified clay samples (Fig. 1b, red line).

Clay containing samples (Na+ and Cloisite 15A series) present characteristic peaks, which can easily be associated with an intercalated gluten/clay stacking morphology (red and black arrows in the Iq^2 vs q plot for the first order reflection). The d-spacing, a distance between silicate layers, was estimated 17 and 48 Å for the WG/Na⁺ and WG/Cloisite 15A samples respectively. SAXS pattern of Cloisite 15A presented a more regular and spaced structure

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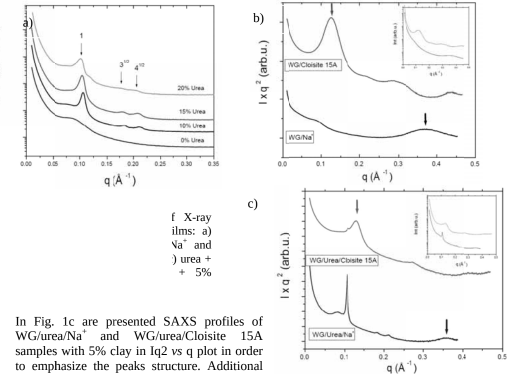
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morphology for the intercalation of gluten, and this indicated a greater order of structures and larger intercalated distances comparing with a natural unmodified clay (Na⁺). This structure complexity seems to be related with good mechanical properties of the WG containing modified MMT clay composites comparing with WG samples².



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intensity curves are plotted in inset. The arrows (blue - the WG/Na⁺ and purple - WG/Cloisite 15A) indicate the first reflection of an intercalated gluten/clay structure. Since, urea induces a hexagonal structure, peaks for this structure are observed together with the peaks of the intercalated structure of gluten/clays. The SAXS profile intensity of both structures seems to be highly dependent on the sample composition.

With this study we conclude that the observed a partially organized WG protein arrangement in urea containing samples is due to urea and its concentration, as well as processing conditions temperature effect. The WG/urea/Cloisite 15A composite structure morphology seems to be greatly influenced by the nature of Cloisite 15A, and urea effect on WG proteins. The final structure morphology seems to be better dispersed and more homogenous in the WG/urea/Cloisite 15A composite comparing to the WG/urea/Na⁺. Further studies are in progress in order to understand WG protein properties in urea/clay systems and tailor biomaterials for specific end uses.

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SAXS studies on insulin self-association

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Insulin is a small protein and a hormone that is part of the body's blood sugar regulation. Diabetes Mellitus is a metabolic disease characterised by inability to regulate the body's blood sugar level properly most often due to either insufficient insulin production or that the insulin receptors have become insulin resistant which decreases the effect of the insulin present in the blood. Worldwide more than 220 million people are suffering from diabetes and in 2004 3.4 million people were estimated to have died from high blood sugar. The number of deaths related to diabetes is expected to double between 2005 and 2030 [1]. A way to improve the treatment of the diabetic patient is to use slow acting insulin to mimic the flat basal level of insulin present in the blood between meals and fast acting insulin to mimic the insulin level just after food intake [2]. The flat basal level of insulin can be mimicked by injecting a type of insulin that has self-associated to larger structures in under the skin. The idea is that the slow release of this insulin to the blood will give a constant insulin background in the blood.

With the idea of slow acting insulin in mind per fluorinated insulin is investigated. It is known that the addition of perfluorinated chains to organic molecules changes their physical and chemical properties [3]. Here we try to direct intermolecular fluorous self-assembly by placing perfluoroalkyl chains of different length to desB30 human insulin by acylation of the ε-mino position of the sidechain of LysB29. Six different perflouroalkyl-insulin analogues were synthesized (Figure 1).

Figure 1: Top Human insulin (DesB30) with a fluoroalkyl chain associted to it (-R). Bottom: Perfluoro alkyl chains of varying lengths.

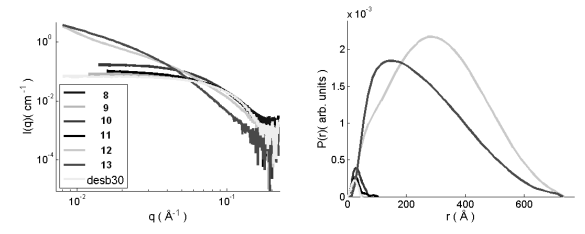


Figure 2: Left: 1-D background subtracted scattering profiles of the 6 types of perfluoroalkyl-insulin compared with the scattering from DesB30 (human insulin) hexamers. Right: the p(r) function, of same measurements.

Our investigation of the perfluoroalkyl-insulin analogues is performed by use of SAXS to estimate their degree of self-assembly and analysed the way they self-associate.

As can be seen from figure 2(left) the scattering from samples 8, 9, 11 (see figure 1 for nomenclature) and to some extend sample 10 corresponds well with the scattering from the hexamer human insulin (DesB30). Thus the structure of these four insulin analogues is made up of perfluoroalkyl-insulin hexamers that self-associates to form larger structures. From the pair distribution function, p(r), in figure 2(right) it can be seen that samples 8 and 9 have a D_{max} of 54 Å.

The scattering from samples 12 and 13 does not resemble the scattering from DesB30. It is therefore likely that samples 12 and 13 do not form hexamers but rather (in the case of sample 12) self-associates as monomers. The p(r) function for both samples 12 and 13 show large degree of self-assembly. For both samples 12 and 13 the D_{max} is larger than what is measurable at I711.

A full analysis of the data sets is in progress [4] and will be described in the scientific literature shortly.

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Improved Hydrogen Storage Kinetics of Nanoconfined NaAlH₄ Catalyzed with TiCl₃ Nanoparticles

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Increasing energy demands, the depletion of fossil fuels in the 21st century and increasing levels of atmospheric carbon dioxide, which may induce global climate changes, call for a transition towards utilization of renewable and sustainable energy sources.¹ One of the major problems in a society based on sustainable energy sources, such as solar, wind and water, where the temporal energy production varies significantly, is to develop a robust, efficient, inexpensive energy storage scheme. Hydrogen has indeed been suggested as a carrier of renewable energy in a future carbon free energy system, but a requirement is that hydrogen can be stored with high gravimetric and volumetric density to reach sufficiently high energy capacities, *e.g.* for mobile applications.² To date, no material simultaneously fulfills all of the above mentioned requirements.³

Nanoconfinement is a new concept where nanoparticles of hydrides can be synthesized or melt-infiltrated in a nanoporous inert scaffold material where the metal hydride particle size is limited by the average pore size of the scaffold allowing for the direct synthesis of small nanoparticles. The scheme of nanoconfinement has several advantages: (i) increased surface area of the reactants, (ii) nanoscale diffusion distances, and (iii) increased number of grain boundaries, which all facilitate the release and uptake of hydrogen and enhance the reaction kinetics.⁴ Furthermore, particle growth and agglomeration of the nanoparticles may be hindered by the compartmentalization within the scaffold template. The design of new nanomaterials is expected to have a major impact on the development of novel sustainable energy and especially efficient energy storage technologies.⁵⁻⁷

In this work, Nanoparticles of NaAlH₄ have been embedded in nanoporous carbon aerogel with TiCl₃ nanoparticles (sample Na-Ti-X) in order to explore possible favorable synergetic effects between nanoconfinement and a functionalized catalytic scaffold. Resorcinol-formaldehyde carbon aerogels with an average pore size of 17 nm and total pore volume of 1.26 g/mL were infiltrated with TiCl₃ dissolved in acetone to obtain an aerogel doped with 2.5 wt% TiCl₃ nanoparticles. NaAlH4 was melt-infiltrated into the functionalized carbon aerogel at 189 °C and $p(H_2) \sim 186-199$ bar. Energy-dispersive spectrometry (EDS) combined with focused ion beam (FIB) techniques has revealed the presence of Na, Al, Ti and Cl inside the

aerogel scaffold material. The infiltrated NaAlH₄ was X-ray amorphous, see Figure 1A, whereas ²⁷Al magicangle spinning (MAS) NMR spectroscopy confirmed the presence of nanoconfined NaAlH₄. Temperature programmed desorption mass spectroscopy (TPD-MS) and Sieverts' measurements demonstrated significantly improved hydrogen desorption kinetics for this new nanoconfined NaAlH₄-TiCl₃ materials as compared to nanoconfined NaAlH₄ without the catalysts TiCl3 (sample Na-X) and to bulk ball-milled samples of NaAlH₄-TiCl₃ (sample Na-Ti-BM), see Figure 1B. We find that the onset temperature for hydrogen release was close to room temperature (Tonset = 33 °C) and the hydrogen release rate reached a maximum value at 125 °C, which demonstrates favorable synergetic effects between nanoconfinement and catalyst addition.

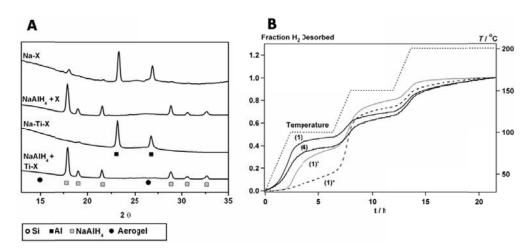


Figure 1. (a) Synchrotron radiation powder X-ray diffraction diagrams of samples Na-X and Na-Ti-X after melt infiltration of NaAlH₄ at $T \sim 190\text{-}200$ and $p(\text{H}_2) \sim 150\text{-}160$ bar, $\lambda = 0.93813$ Å. Maxlab beamline I711. (b) Normalized Sieverts' hydrogen desorption profiles. The hydrogen release kinetics for samples Na-Ti-X (solid lines) cycle (1) and (4), Na-X (dashed line), and Na-Ti-BM (dotted line) are compared.

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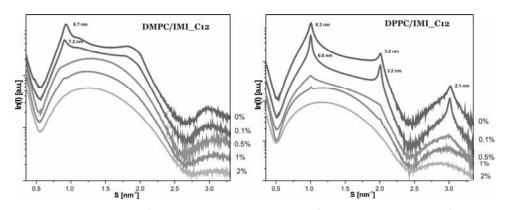
SAXS Studies of DMPC and DPPC-Dimeric Surfactant Lipoplexes.

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Dimeric surfactants are new class of surfactants. They show significantly improved properties in comparison to conventional (monomeric) surfactants. The dimeric surfactants are characterised by critical micellization concentration (CMC) much lower than that of classic surfactants with equivalent hydrophobic chain length. Dimeric surfactants are also good surface tension reducers, much more efficient than the corresponding monomeric surfactants ¹⁻³.

The aim of this study was to analyse the effect of different concentrations of dimeric surfactant on the disturbance or stabilisation of particular phases of phospholipids in its mixtures. The study was performed on 1,2-dimyristoyl-sn-glycero-3-phosphocholine (DMPC) and 1,2-dipalmitoyl-sn-glycero-3-phospho-choline (DPPC) and Gemini type surfactant IMI_C12 (with dodecyloxy- hydrophobic chain). Series of the SAXS data sets were collected in MaxLab, at the Beam Line I711 at the MAXII storage ring. The data were collected at temperatures from 279 to 303 K for DMPC and from 283 to 318 K for DPPC using the synchrotron radiation (λ =0.107 nm) and the Mar 165 CCD detector. The scattering vector range was $0.05 < s < 3.42 \text{ nm}^{-1}$.

In the Figure 1 are presented the scattering data collected for DMPC/IMI_C12 (left) and DPPC/IMI_C12 (right) mixtures. As expected, in reference samples (pure DMPC or DPPC) the observed curves are typical for lamellar structures. For pure DPPC suspension, the diffraction pattern shows three strong peaks characterized by d_{001} =6.3 nm, d_{002} =3.2 nm and d_{003} =2.1 nm. For low concentration of the surfactant (0.1%, 0.5%) in the system, the corresponding lamellar lattice constant increased up to 6.7 nm.



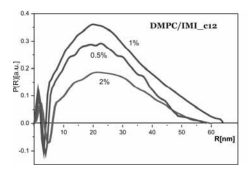
 $Figure~1.~SAXS~results~for~DMPC/IMI_C12~in~291~K~(right)~and~for~DPPC/IMI_C12~in~293~K~(left).$

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With the increased surfactant concentration in the systems (0.5% and 1% IMI_C12) the gradual disappearance of the lamellar structure can be observed. For DMPC/2% IMI_C12 system the SAXS data shows the characteristic broad maximum, which is typical for single bilayers in the absence of stacking⁴. Similar very broad peak is observed also in scattering patterns of DMPC with 2% concentration of IMI_C12.

The SAXS results implied a gradual disappearance of the lamellar phase typical for DMPC and DPPC and probable formation of the mixed liposomes. In case of DMPC we can notice that the phase change occurs in lower surfactant concentration, so we can that their phase behaviour is more dependent on presence of surfactant IMI_C12.



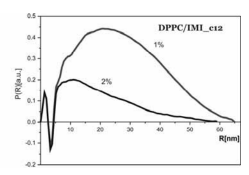


Figure 2. The pair distribution function calculated for DMPC/IMI_C12 (right) and for DPPC/IMI_C12 (left).

Presented in the Figure 2 pair distribution functions, calculated for DMPC and DPPC/IMI_C12 systems, also reveal significant differences in size of formulated nanostructures. For example for DPPC/1% IMI_C12 the maximal size (D_{max}) was 65 nm and for DPPC/2% IMI_C12 D_{max} =58 nm .

Also the temperature range of the main phase transition of DMPC and DPPC was shifted towards lower temperatures. The observed scattering patterns of both phospholipid/surfactant systems are very similar to the SAXS patterns reported for the bicelar DMPC/DHPC systems⁵. The surfactants studied induce changes of the DPPC and DMPC phase transition temperatures and promote a gradual disappearance of the lamellar phase of DMPC and a probable formation of the bicellar phase

This research was supported in part by research grant (No N N202 248935) from the Ministry of Science and Higher Education (Poland). The data collection (SAXS studies at MaxLab Lund, Sweden) leading to these results has also received funding from the European Community's Seventh Framework Programme (FP7/2007-2013) under grant agreement no 226716.

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SAXS Studies of Zwitterionic Surfactants Interactions with DMPC

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Sulfobetaines (also known as sultaines) are sulphur analogs of betaines. Sulfobetaines with long hydrophobic chains adsorb at charges surfaces in the wide range of pH without forming hydrophobic films. They are also good lime soap dispersants and are widely used in cosmetic formulations. It should be noted, however, that a little skin irritation attributed to sulfobetaines was also reported¹.

In this work we studied the effect of two novel zwitterionic surfactants (Figure 1), prepared in our laboratory, with the sulfobetaine structure (4-(N-hexylmorpholine)-1-butansulfonate - HMBS and 4-(N-dodecylmorpholine)-1-butansulfonate - DDMBS), on the stability of the model system of biological membrane based on DMPC.

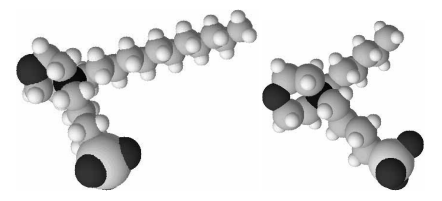


Figure 1 Chemical structures of 4-(N-dodecylmorpholine)-1-butansulfonate (left) and 4-(N-hexylmorpholine)-1-butansulfonate (right)

The small angle X-ray scattering data were collected on the beam line I711^{2,3} at the MAXII storage ring using the synchrotron radiation (λ =0.107 nm). The phospholipid-surfactant mixtures (with molar ratio 100:1, 50:1, 10:1, 5:1 and 3:1) were placed in thermostated flow-cell with quartz capillary. The sample-to-detector distance was 1.76 m, which leads to the scattering vector range 0.1 < s < 3.42 nm⁻¹ (where s=4 π sinθ/ λ , 20 was the scattering angle). The transmission SAXS measurements were performed at temperatures ranging from 279 – 303 K. The background scattering was collected before and after data collection for all samples.

The SAXS data sets were normalized to the incident beam intensity, corrected for detector response and integrated using SAXS data collection and processing software Bli711³. The final scattering curve was obtained after subtraction of the buffer scattering using the program package PRIMUS⁴.

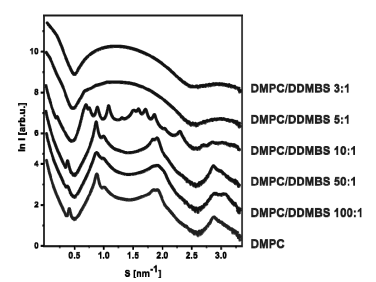


Figure 2. SAXS data recorded for DMPC/DDMBS systems at 287 K

The zwitterionic sulfobeitaine derivatives interacting with DMPC disrupted the formation of stable lamellar phases. 4-(N-hexylmorpholine)-1-butansulfonate (HMBS) with shorter hydrophobic group only induced small changes in d-spacing characterising the lamellar phases. The second surfactant - 4-(N-dodecylmorpholine)-1-butansulfonate (DDMBS) promote a formation of the stable cubic Pn3m phase at temperatures below main phase transition of DMPC(in DMPC:DDMBS molar ratio 10:1). For higher concentrations of DDMBS (5:1 and 3:1) probably the bicellar phase is formed. The exemplary scattering curves recorded for DMPC/DDMBS systems are presented in Figure 2.

This research was supported in part by research grant (No N N202 248935) from the Ministry of Science and Higher Education (Poland). The data collection (SAXS studies at MaxLab Lund, Sweden) leading to these results has also received funding from the European Community's Seventh Framework Programme (FP7/2007-2013) under grant agreement no 226716.

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Effect of NaH/MgB $_2$ ratio on the absorption kinetics of the system NaH-MgB $_2$

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Recently, based on the inspected kinetic effect of the MgB₂, Barkhordarian et al. [1] and Vajo et al. [2] reported on the possibility of reversibly store hydrogen in tetrahydroborates when mixed with MgH₂. As confirmed by the numerous works published in the last years, this discovery ignited new interest in the tetrahydroborates as potential material for hydrogen storage. In particular, the systems LiBH₄-MgH₂, Ca(BH₄)₂-MgH₂ and NaBH₄-MgH₂ have been subject to intensive investigations. Among these mixed hydride systems NaBH₄-MgH₂ in the molar ratio of 2NaBH₄/MgH₂ is considered as a model system for the study of the hydrogen sorption properties. This system is expected to reversibly exchange an amount of hydrogen equal to 7.8 wt.% according to the following reaction: $2NaBH_4+MgH_2 \leftrightarrow 2NaH+MgB_2+4H_2$. The overall reaction enthalpy for this reaction is calculated to be 62 kJmol⁻¹H₂. [3] This enthalpy value entails for the system 2NaBH₄+MgH₂ an equilibrium hydrogen pressure of 1 bar at 350°C. Lately, the reaction mechanism of the hydrogen absorption for the composition 2NaBH₄/MgH₂ has been investigated in detail. [3, 4] It was reported that the reaction paths of the hydrogen absorption take place in several single steps. Hereunder the hydrogenation reaction for the composition 0.5NaBH₄/MgH₂ is described. The measurement was carried out at 50 bar of hydrogen pressure, in scanning temperature from RT to 350 °C and than keeping the system under isothermal conditions at 350 °C for several hours (Figure 1). During heating at roughly 230 °C the appearance of an unknown crystalline phase is observed. Later on at 270 °C the appearance of this phase is followed by the formation of NaMgH₃. With the appearance of the NaMgH₃ phase the NaH phase disappears completely. At 325 °C the formation of NaBH₄ and little later of MgH₂ is observed together with the simultaneous disappearance of both the NaMgH₃ and the unknown crystalline phase. In contrast to the system 2NaH/MgB₂ [3] the system 0.5NaH/MgB₂ (figure 1) does not show the formation of an amorphous background. In addition, for the composition 0.5NaH/MgB₂ the formation of MgH₂ was achieved.

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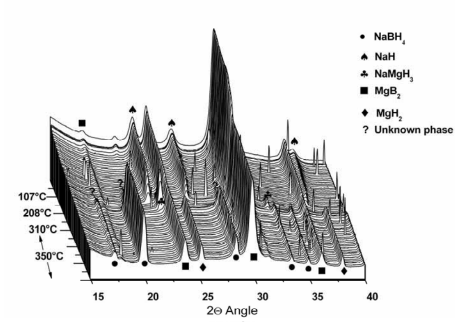


Figure 1: Series of SR-PXD patterns of the $2NaH+MgB_2$ system heated at 50 bar hydrogen pressure from RT to 350 °C and kept under isothermal condition (5 °C/min, wavelength = 1.097 Å).

This work shows that *in situ* SR-PXD at variable temperatures and gas pressures is a powerful tool for studying gas/solid reactions. These investigations prove the dependency of the hydrogen absorption reaction in the NaBH₄-MgH₂ on the NaBH₄/MgH₂ ratio and help to reveal the reaction mechanisms in these new and rather complex class of hydrogen storage materials.

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Novel metal borohydrides: Studies of synthesis, crystal chemistry and thermal decomposition

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The world today is facing increasing energy demands and a simultaneous demand for cleaner and more environmentally friendly energy technologies. Hydrogen is recognized as a possible renewable energy carrier, however its large-scale utilization is mainly hampered by insufficient hydrogen storage capabilities. Complex metal hydrides, such as metal borohydrides, is an interesting class of materials for hydrogen storage, since they exhibit high gravimetric hydrogen densities and a variety of decomposition temperatures. Unfortunately, no single material has yet been found, which fulfil all the criteria for mobile hydrogen storage. Therefore, there is an urgent need for discovery and studies of novel classes of materials, such as transition metal-based or mixed metal borohydrides as potential future hydrogen storage materials.

The past few years have seen a significant increase in the number of new borohydrides and of experimental and theoretical studies on their structural and solid-state chemistry. 3,4 In our lab we have synthesized and characterized a wide range of novel borohydride materials based on alkali metals and alkali earth or transition metals e.g. $MZn_2(BH_4)_5$ (M=Li or Na), $NaZn(BH_4)_3$, $KZn(BH_4)Cl_2$, $Y(BH_4)_3$, $NaY(BH_4)_2Cl_2$ and $MSc(BH_4)_4$ (M=Na or K). Here, we will show selected results from *in situ* SR-PXD studies on $KSc(BH_4)_4$.

Samples were prepared from KBH₄ and ScCl₃ mixed in molar ratios of 2:1, 3:1 and 4:1 and ball milled for 120 min under argon atmosphere. In-situ time-resolved SR-PXD was measured at the MAX II Synchrotron in Lund, Sweden, at Beamline I711 using a MAR165 CCD detector system. 11 Samples were mounted in sapphire single-crystal tubes (1.09 mm o.d., 0.79 mm i.d., Al₂O₃), which were attached on a specially designed sample cell for studies of gas/solid reactions at elevated temperature and pressure. The temperature was controlled thermocouple placed in the sapphire tube next to the sample. 12 All sample handling was perform in an argon-filled glove box $(p(O_2, H_2O) < 0.3 \text{ ppm})$. The selected wavelength was 1.097 Å and the X-ray exposure time was 30 s per powder diffraction pattern. The FIT2D program was used to transform raw data to powder patterns.

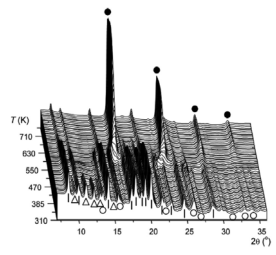


Fig. 1 In situ synchrotron radiation powder X-ray diffraction measured for KBH₄-ScCl₃ (2:1) heated from RT to 770 K with a heating rate of 7 K/min and p(H₂) ~ 2.5 bar (λ = 1.097 Å). Only peaks which are at least partly resolved are marked by the symbols: Δ KSc(BH₄)₄, | K₃ScCl₆, O KBH₄, and ● KCl.

After synthesis the KBH₄-ScCl₃ samples all contain two sets of unidentified Bragg peaks. One of these is assigned to a new metal borohydride, KSc(BH₄)₄, and the other to a new ternary potassium

scandium chloride K₃ScCl₆. This suggests that the reaction during ball milling can be described according to reaction scheme eq. 1 and that the optimal starting ratio between KBH₄ and ScCl₃ is 2:1.

$$4KBH_4 + 2ScCl_3 \rightarrow KSc(BH_4)_4 + K_3ScCl_6 \tag{1}$$

In situ SR-PXD as a function of temperature (Fig. 1) proves very important in cases like this, where the synthesis product contains two structurally unknown compounds. The novel borohydride, KSc(BH₄)₄ is observed to melt at ~405 K, whereas the diffraction from K₃ScCl₆ is unchanged from *RT* up to ~500 K. Hence, only *T*-ramping allowed the assignment of observed peaks to individual compounds, and led to a successful indexing of powder patterns and to solving both crystal structures.

The study of the thermal decompostion shows that a series of simultanous and coupled reactions takes place (Fig. 1). As $KSc(BH_4)_4$ decompose from the melt at ~460 K, an increase in the amount of KBH_4 is observed, suggesting that this is formed as a decomposition product along side hydrogen and possibly amorphous scandium borides. As the temperature reaches 500 K, K_3ScCl_6 reacts with KBH_4 to form KCl, release more hydrogen and possibly form amourphous acandium borides. In the same temperature interval the formed KCl dissolves in the remaining KBH_4 and forms a solid solution of composition $K(BH_4)_{1-x}Cl_x$ according to reaction scheme eq. 2, similar to what have previously been observed for $LiBH_4$ -LiCl.¹³

$$xKCl + (1 - x)KBH_4 \rightarrow K(BH_4)_{1-x}Cl_x$$
 (2)

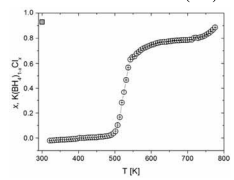


Fig. 2 Substitution degree, *x* for K(BH₄)_{1-x}Cl_x obtained from Rietveld refinement of *in situ* SR-PXD data. Symbols: O values during heating, □ values after heating/cooling.

This reaction was investigated in further detail by Rietveld refinement of the *in situ* SR-PXD data. A structural model was developed where Cl substitute for BH₄, i.e. Cl was constrained to the same set of x,y,z coordinates as B, and the sum of their occupancies was constrained to 1. Fig. 2 shows the change in the substitution degree, x in K(BH₄)_{1-x}Cl_x as a function of temperature. From RT to ~500 K, no substitution in KBH₄ is observed. At ~500 K x increases at a high rate and at ~610 K a plateau is reached where x ~ 0.78. Upon further heating at ~730 K, the substitution continues likely due to slow decomposition of the KBH₄ contained in the K(BH₄)_{1-x}Cl_x solid solution.

Acknowledgement

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Tailoring the properties of lithium tetrahydridoboranate by anion substitution.

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Borohydrides are considered interesting materials for hydrogen storage in mobile applications due to their theoretical high hydrogen content, 1,2 *e.g.* 18.5 weight percent (wt.%) for lithium tetrahydridoboranate, LiBH₄. Unfortunately, the borohydrides are either too stable or unstable for practical hydrogen storage systems. LiBH₄ release hydrogen at a temperature of 410 °C ($p(H_2) = 1$ bar) but the system can be modified by adding suitable compounds, ³ e.g. SiO₂, TiF₃ or TiCl₃, or by substitution of ions aiming to destabilize the structure. ⁴

The aim for this investigation is to alter the properties of LiBH₄ by substitution of the anion BH₄⁻ with Br⁻ in LiBH₄ which is feasible due to the similar size in radii.⁵
The substitution is explored by *in situ* synchrotron radiation powder X-ray diffraction (SR-PXD) and by differential scanning calorimetry (DSC) on a sample of LiBH₄ + LiBr (1:1) synthesized by a mechano-chemical method, e.g. ballmilling for 2 hours effectively.

In situ time-resolved SR-PXD data was measured at the MAXII synchrotron beamline I711 at MAX-lab, Lund, Sweeden ($\lambda = 1.07200$ Å) with a MAR165 CCD detector system. The samples were mounted in sapphire single crystal tubes (1.09 mm o.d., 0.79 mm i.d., Al₂O₃) in an argon filled glovebox ($p(O_2, H_2O) < 0.5$ ppm). The sample holder was constructed so that hydrogen pressures from vacuum up to 200 bar and temperatures from RT to 800 °C can be used during the experiment. The system was flushed with argon and evacuated three times before the valve to the sample was opened prior to the X-ray experiment. The X-ray exposure time was 30-60 s per powder diffraction pattern. This *in situ* SR-PXD experiment was conducted with a constant hydrogen pressure of $p(H_2) = 1$ bar and with a steadily increasing temperature (rate of 5 °C/min) from RT-427 °C.

Differential scanning calorimetry (DSC) and thermogravimetric analysis (TGA) was measured simultaneously with a Netzsch STA449C Jupiter instrument at heating rate of $\nu = 2.0$ °C/min from RT to 420 °C in a flow of He (50 mL/min). The samples were mounted in Al₂O₃ crucibles.

The *in situ* SR-PXD-data of a sample of LiBH₄ + LiBr ballmilled in the ratio 1: 1 and preheated to 245 °C for 3 days is shown in Figure 1 in the temperature range RT – 420 °C. There is no indication of substitution at RT. At 300 °C the substitution of LiBr initializes and the intensity of LiBr decreases. At 360 °C a reaction of LiBr with the h-LiBH₄ phase is observed with the formation of a new hexagonal structure, denoted Li(BH₄)_{1-x}Br_x. The new phase must consist of an approx. equal amount of Br⁻ and BH₄⁻ anions since the LiBr diffraction disappears with the formation of the Li(BH₄)_{1-x}Br_x phase.

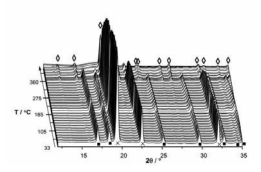


Figure 1 *In situ* SR-PXD on LiBH₄ + LiBr (1:1) preheated to 245 °C for 3 days, sample **S4**. The sample was heated from *RT* to 427 °C with a heating rate of 5 °C/min (λ = 1.07200 Å). Symbols: the crosses (X) represents the LiBr; the filled squares (■) represents *h*-LiBH₄; the empty diamonds (◊) represents the new phase, Li(BH₄)₁. $_x$ Br_x. At T ≈ 185 °C the beam was cut for a minute. At T ≈ 420 °C no crystalline materials was left in the sample due to melting.

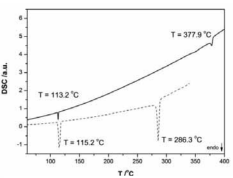


Figure 2 Differential scanning calorimetry shown in the temperature range from 60 to 400 °C. The reference sample of LiBH₄ is represented with a broken line and the sample of LiBH₄ + LiBr (1:1), **S3**, is represented by the solid line.

Further experiments with TGA and DSC reveals a stabilization of the hexagonal phase of LiBH₄ by approximately 90 °C altering the melting point of LiBH₄ at T = 286 °C (pure LiBH₄) to T = 378 °C (Li(BH₄)_{1-x}Br_x), see Figure 2. However, the stabilization of the hexagonal phase has no apparent impact on the hydrogen release temperature determined by sieverts measurements to be unchanged (not shown here).

The next step will be the investigation of the reversibility of the material upon hydrogen release and uptake.

Aknowledgements

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Hydrogen absorption and desorption properties of ScNiAl, a novel material for hydrogen storage

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The potential use of hydrogen as energy carrier in carbon-free energy systems have been extensively studied in the past 20 years. Due to promising properties, such as high capacities and good reversibility, metal hydrides are claimed to be among the most suitable hydrogen storage materials and considerable research efforts have been devoted to these compounds[1].

Scandium is interesting from a fundamental point of view by combining the rare-earthmetal properties with a low weight. Many scandium-based alloys, such as Mg_{0.65}Sc_{0.35} [2-3] and ScAl_{1-x}Mg_x [4], have shown promising hydrogen storage properties. ScNiAl crystallises in the hexagonal MgZn₂-type structure, *P*6₃/*mmc* [5], and here we present this compound as a hydrogen storage material.

In situ synchrotron radiation powder X-ray diffraction (SR-XRD) measurements were performed at the MAX-II synchrotron in Lund, Sweden, using the instrument at beamline I711 [6]. The wavelength was determined to be $\lambda = 0.94608$ Å. A Mar165 CCD detector was used for intensity measurement of the diffracted beams. The X-ray exposure was 30 s/scan. The sample holder consisted of a single crystal sapphire tube connected to a vacuum pump and pressurized hydrogen gas source, due to X-ray absorption problems the sample was placed in a thinner capillary inside the sapphire tube. The sample was heated by resistive heating (tungsten wire) and the temperature was measured with a thermocouple placed inside the sapphire tube.

ScNiAl was hydrogenated *in situ* at 5 MPa and a linear temperature increase of 5 °C/min up to 180 °C, the temperature was then kept constant at 180 °C for the rest of the experiment. *In situ* SR-XRD showed that ScNiAl absorbs hydrogen at temperatures below 180 °C, shown in Figure 1. Hydrogen absorption started at ~140 °C. After hydrogenation the pressure was released to vacuum whereupon the material instantly desorbed virtually all the hydrogen. A second hydrogenation cycle was performed by resetting the hydrogen pressure to 5 MPa followed by another release to vacuum. No significant changes of the hydrogen absorption behaviour could be seen during cycling.

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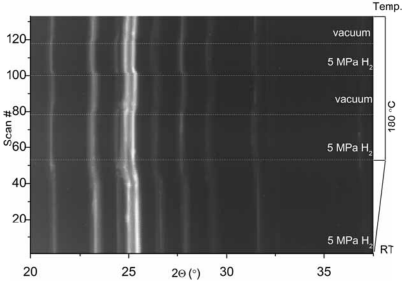


Figure 1. Hydrogen absorption and desorption investigated with in situ SR-XRD (λ = 0.94608 Å, heating rate 5 °C/min).

The absorbed amount of hydrogen was estimated by studying the unit cell volume expansion of samples hydrogenated ex situ at similar conditions as for the in situ measurements. It was shown that the compound absorbs ~0.5 H/f.u., leading to a hydrogen storage capacity of 0.40 wt.%. These estimations are based on Peisl's finding that the solid solution of hydrogen in a variety of metals and alloys causes an volume expansion, ΔV , averaging 2.9 Å³ per hydrogen atom [7].

In summary, we have shown that ScNiAl can absorb and desorb hydrogen interstitially up to ~0.5 H/f.u. Also, ScNiAl is easy to synthesize and safe to handle in air which is of great importance for possible practical applications of the material.

Acknowledgements

Financial support from The Swedish Research Council and The Nordic Energy Research Programme is gratefully acknowledged.

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When do Water-Insoluble Polyion-Surfactant Ion Complex Salts "Redissolve" by Added Excess Surfactant?

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Oppositely charged polymer-surfactant systems are used as rheology modifiers, gelling agents, colloidal stabilizers and surface deposition agents in a wide range of applied contexts, including pharmaceutics, foods, personal care products and waste water treatment [1-7]. The fundamental feature of the system that is utilized in all these contexts is the strong attraction between the charged polyions and the oppositely charged surfactant ion aggregates, both containing multiple charges, which leads to extensive association in the system. Obviously, an essential property to control is the phase behaviour of the mixture. Typically, a concentrated polyion-surfactant ion phase separates out under mixing conditions when a stoichiometric, neutral polyion-surfactant ion complex is formed. However, the situation for off-stoichiometric mixtures may vary widely, depending on the system, as will be amply exemplified below. Certain offstoichiometric systems produce single-phase solutions, whereas other systems remain phase separated at practically all mixing ratios. These differences are widely utilized. In some applications off-stoichiometric single-phase solutions are desired, for instance in the form of highly viscous physical networks. Under other circumstances one may want a complex that remains insoluble, for instance as a surface coating, for wide variations of the system composition. Indeed, important applications, such as hair care products or laundry detergents, use formulations that effectively switch from one-phase mixtures to phase-separated mixtures by a simple operation such as a dilution of the system [6]. Typically, the initial one-phase formulation in the latter applications contains a large excess of surfactant.

The above background serves to justify why the present study focuses on the phenomenon that we will refer to as "surfactant redissolution", that is, when the addition an excess of surfactant to an associatively phase-separated stoichiometric polyion-surfactant ion complex leads to its transformation into a single-phase solution. As we have indicated, the capability for the surfactant to redissolve the complex varies greatly from system to system, but the underlying physics is poorly understood. The ability to understand and control this variation is of considerable interest for applications, such as those mentioned above. The phase behaviour of polymer-surfactant mixtures is also a problem of fundamental importance in polymer and colloid science.

The purpose of the work presented here was to gain further insight in the phenomenology and the mechanism of surfactant redissolution. We have in this study determined additional phase diagrams for mixtures of complex salt containing linear polyions based on polyacrylate that are either very long or very short homopolyions, or long copolyions (PA¯-co-NIPAM or PA¯-co-DAM, containing *N*-isopropylacrylamide or *N*,*N*-dimethylacrylamide comonomers, respectively), in mixtures with C_nTAX

surfactants of different alkyl chain lengths and with either bromide or acetate counterions. We have also compared the results obtained experimentally with the outcome of calculations using a simple Flory-Huggins model.

The present phase studies confirmed that the outcome of an experiment, where excess surfactant was added to a phase separated mixture of a complex salt in water, may vary widely: from a wide miscibility gap, covering practically all mixing ratios of complex salt and surfactant, to an efficient "surfactant redissolution", where a limited amount of excess surfactant is capable of closing the miscibility gap into a single-phase micellar solution. Compared to previously available complex salt/surfactant/water phase studies, the present study has succeeded to isolate various factors responsible for this transition, and to show that a range of behaviours previously encountered for chemically different systems can be covered using minimal chemical variations in a single class of experimental systems.

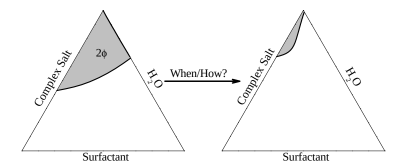


Figure 1 – Representation of the efficient redissolution of polyion-surfactant ion complexes.

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Phase diagrams come alive: understanding how to create, destroy or change ordered surfactant structures by polymerization reactions

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There is a strong and long-standing interest in concentrated associating polymer-surfactant mixtures. The associative phase separation that gives rise to concentrated mixtures in aqueous polymer-surfactant systems of opposite charge is particularly well documented¹⁻⁸. Owing to the self-association of the surfactant component, the concentrated phases that separate out from such mixtures often feature ordered liquid crystalline structures^{5,6,9-14}. Important applications that stimulate the research on concentrated associating polymer-surfactant mixtures include templating for preparation of mesoporous ordered materials, surface deposition of concentrated polymer-surfactant layers, and the creation of soft structured particles¹⁵⁻¹⁸.

Materials based on associating polymer-surfactant pairs can differ in structure, water uptake and mechanical properties depending on factors such as the lengths of the polymer and surfactant chains and the overall compositions. The materials can be produced using different routes, e.g. chemical synthesis, sequential addition of components, dilution and simple mixing. An important question is to what extent different routes of preparation give the same results, indicating that the systems have reached equilibrium. Of particular concern here are systems produced by polymerization reactions. Most approaches to polymerization reactions in surfactant systems so far seems to have been based on trial and error, and the resulting structures have often been found to be different from the original surfactant template 19,20. That is, polymerization changes the equilibrium structure of system. One reason for such changes is the development of a repulsive depletion interaction between the formed polymers and the surfactant aggregates¹⁹. In fact, not much attention has been given to the fact that, as it happens for silica and its templates in hard mesoporous materials, the interaction between the different components in the system should be attractive in order to create long-ranged ordered structures containing polymer and surfactant²¹. The situation is further complicated since many systems, and certainly inorganic mesoporous materials, become kinetically arrested at some point during the polymerization.

Specifically, equilibrium phase diagrams have been established for associating polymer-surfactant systems involving aqueous cationic alkyltrimethylammonium surfactant ions with two different counterions ($C_{16}TAAc$, $C_{12}TAAc$ and $C_{16}TABr$) (Ac=acetate) mixed with the respective complex salts $C_{16}TAPA_{30}$, $C_{16}TAPA_{6000}$ and $C_{12}TAPA_{30}$, $C_{12}TAPA_{6000}$ (PA=polyacrylate)^{22,23,24}. In the complex salts, the counterions to the surfactant ions were polyacrylate polyions of either very high or very low degrees of polymerization (25, 30 or 6000). Interestingly, the obtained ternary phase diagrams illustrate what happens, at equilibrium, when monomeric counterions to the surfactant are gradually replaced by polymeric counterions - a process closely analogous to a polymerization reaction.

In the present work, we built on and extended our previous studies by investigating

polymerization reactions in systems of similar minimum complexity, by the use of alkyltrimethylammonium surfactants with polymerizable acrylate counterions. Two aspects made our approach to polymerization in surfactant media unique: i) it is based on our *a priori* knowledge of the phase behavior of ionic surfactants with monomeric versus polymeric counterions and ii) the synthesis was done under conditions where the surfactant aggregates are the sole counterions to the growing polyion chains. The latter is a strong constraint for the structure resulting from the polymerization. The present investigation indeed showed that the results of the polymerization reactions were predicted by the relevant equilibrium phase diagrams. The results also indicate general principles relevant for more complex systems, such as mesoporous silicates.

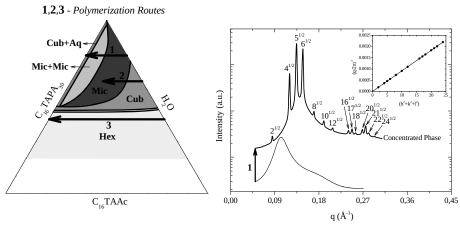


Figure 1 – *Left*: equilibrium phase diagram of C₁₆TAPA₃₀/C₁₆TAAc/H₂O; *Right*: SAXS profiles of a micellar phase (bottom profile - before reaction) and a cubic phase (top profile - after reaction) of a two phase sample, ascribed as pathway number 1 in the left figure.

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Investigation of the hydrogen reaction mechanism in CaF2+3CaH2+4 MgB2 composites

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The reactive hydride composite (RHC) CaF₂+3CaH₂+4MgB₂ was selected to perform synchrotron radiation powder X-ray diffraction (SR-PXD, beamline I711) experiments during the first cycle of hydrogenation/ dehydrogenation. These experiments are suitable to investigate the reaction pathways and structural features of RHC for hydrogen storage. In the hydrogenation setup, the sample was heated at 5 K min⁻¹ from room temperature to 350°C and kept isothermal for 4 hours under 130bar of hydrogen pressure. For dehydrogenation reaction, a previously hydrogenated sample was heated at 5 K min⁻¹ from room temperature to 350°C and kept isothermal for 1 hour under dynamic vacuum.

Figure 1 presents the characterization of the first hydrogenation of $CaF_2+3CaH_2+4MgB_2$ RHC. The peaks observed at room temperature correspond to the raw materials CaF_2 , CaH_2 and MgB_2 in addition to an unidentified peak at 3.09 Å ⁻¹. The first change observed with heating is the increase of peak intensity due to recrystallization or coarsening of the micro/nano-particles. It is particularly clear for CaF_2 . The evolution of two unidentified peaks at 1.56 Å ⁻¹ and 2.55 Å ⁻¹ was observed at 280°C. As the temperature increased, these peaks shifted to higher q [Å ⁻¹] values and two changes in intensity were observed in those peaks. The first of them was an increase in peak intensity as the temperature reaches 350°C and then a decrease in peak intensity until a complete depletion after some time under isothermal (350°C) conditions. Those peaks suggest the formation of reaction intermediaries, however none of the proposed intermediates or phase changes for the hydrogenation/ dehydrogenation reactions in the related $Ca(BH4)_2$ [1, 2, 3, 4, 5, 6] or $Ca(BH4)_2 + MgB_2$ [2] composites fit them. The formation of β- $Ca(BH_4)_2$ phase started at 300°C as indicated by the most clear peak at 1.27 Å ⁻¹. As indicated by SR-PXD patterns the hydrogenation products show a complex mixture of phases. In addition to the identified products $Ca(BH_4)_2$ and β- $Ca(BH_4)_2$, there was the formation of unidentified phase(s) with peak positions at 1.68, 1.90, 2.28, 2.53, 2.83, 3.36 and 3.52 Å ⁻¹; most of them disappeared after cooling.

The subsequent dehydrogenation reaction is presented in Figure 2. At room temperature it can be observed the predominance of α - Ca(BH₄)₂, MgH₂, unreacted CaF₂ and unidentified peak at 1.60 Å ⁻¹. At about 140°C the α to β Ca(BH₄)₂ phase transition is observed. At about 275°C, subtle changes in the peak intensity of Ca(BH₄)₂ phases occur indicating the beginning of the dehydrogenation reaction. The Ca(BH₄)₂ peaks vanish before reaching 350°C while the unidentified peak at 1.60 Å ⁻¹ disappears under isothermal conditions at 350°C. Finally, peaks corresponding to CaH₂, MgB₂, CaF₂, and Mg are observed indicating the reversibility of the system.

Conclusions: The hydrogenation and dehydrogenation reaction of the $CaF_2+3CaH_2+4MgB_2$ composite was demonstrated by means of in-situ SR-PXD. The hydrogenation reaction proceeds by means of a still unknown intermediate. For the dehydrogenation reaction there is no clear evidence of an intermediate, a direct reaction pathway is proposed.

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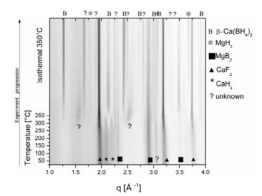


Fig. 1. In situ SR-PXD patterns of as milled CaF₂+3CaH₂+4MgB₂ composite, hydrogenation at 130 bar and heating from room temperature to 350°C (5 K min⁻¹, λ=0.979Å, Max Lab Synchrotron).

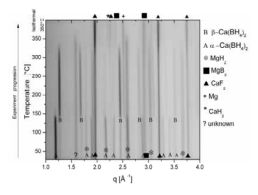


Fig. 2. In situ SR-PXD patterns of first dehydrogenation of CaF₂+3CaH₂+4MgB₂ (prehydrogenated) composite, dehydrogenation at dynamic vacuum and heating from room temperature to 350°C (5 K min⁻¹, λ=0.979Å, Max Lab Synchrotron).

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FREEZING OF MONTMORILLONITES IN BRINE STUDIED BY SYNCHROTRON X-RAY POWDER DIFFRACTION

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Introduction

The Swedish repository for high-level radioactive waste will be located in Forsmark. The current KBS-3 approach [1] includes several barriers: (i) the ceramic fuel itself, (ii) the copper canister, (iii) compressed rings of smectite-rich, swelling clay, and (iv) a suitable rock. The clay barrier functions as mechanical support for the copper canister and protects it from movements in the rock. It also functions as a diffusion barrier for ions and water. It is of fundamental importance that the clay retains its properties over a period of time as long as 100 000 years. A deep repository is not expected to freeze during glacial periods, while levels closer to the surface can be affected by freeze-thaw cycles [1]. This will be of importance for back-filled shafts and sealed bore holes, when swelling clay is used as a sealing material. The aim of our project is to study fundamental processes of importance in this context [2,3], for example variations in the salt content of the ground water is one factor that is expected to have great influence [1].

Experimental and results

The freezing and thawing of smectite in different aqueous salt solutions was further studied after our initial study with pure water [3]. Na-montmorillonite was frozen in aqueous solutions of NaCl of varying concentration (0-3 M) and Ca-montmorillonite was frozen in CaCl₂ solutions (0-3 M). Samples containing 70 wt% salt solution and 30 wt% montmorillonite powder were put into glass capillaries. Before mixing with the aqueous solution, the montmorillonite was present as a mono-layer hydrate (Na) and a two-layer hydrate (Ca). The stationary capillaries were cooled from + 20 $^{\circ}$ C to -50 $^{\circ}$ C using a stream of flowing nitrogen with adjustable temperature (Cryojet, Oxford Instruments). X-ray powder diffraction data was collected on the beam-line I711 or I911-5 at MAX-lab using a Mar system with a flat CCD detector (Mar 165, 2048 x 2048 pixels). Each frame was monitored for 20 seconds and the integrated intensity, scattering angle and width of the recorded diffraction rings were evaluated using the software Fit2d (A. P. Hammersley, ESRF).

The presence of a strong 001 basal reflection at around $2\theta = 3^{\circ}$ in the diffraction patterns in Fig. 1, showed that crystalline hydrates were present in all cases except for the Namontmorillonite at the lowest salt concentrations (0 and 0.25 M). For the latter, osmotic swelling occurred and the basal spacing was too large to be observed in the present experiments (though it was measured in a separate SAXS experiment). At -50 °C, the presence of ice was indicated by the two intense peaks at $2\theta = 13$ -14.5°. In the Na-montmorillonite-case, the basal basing decreased both when the salt concentration increased and the temperature decreased. As expected, lower hydrates formed when the water vapour pressure of the system was lowered. In the Ca-case, the basal spacing at room temperature decreased in a similar way with increased salt

concentration, while it unexpectedly increased at -50 °C. In a frozen system, the water vapour pressure will be that of pure ice at the specific temperature and this indicated that there was a change in the properties of the Ca-mineral taking place with increasing concentration of salt. An interesting possibility was that the phenomenon was related to the occlusion of neutral CaCl₂ into the montmorillonite according to the Gibbs-Donnan effect [4].

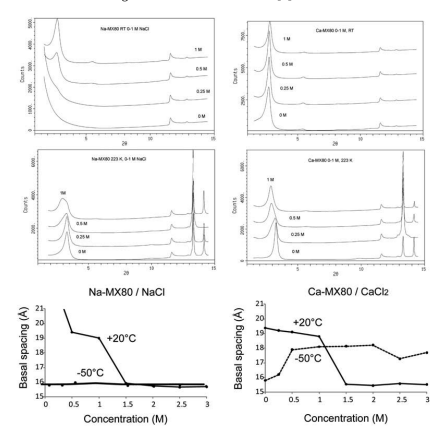


Fig.1. Diffraction data for Na-montmorillonite (left column) and Ca-montmorillonite (right column). X-ray powder diffraction patterns recorded at $+20\,^{\circ}\text{C}$ (top row) and at $-50\,^{\circ}\text{C}$ (central row). Basal spacing 001 at different temperatures and concentrations of salt (bottom row).

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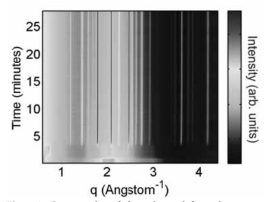
In situ synchrotron radiation studies of inorganic nanomaterials syntheses

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In Situ Study of the Defect Formation During Hydrothermal Synthesis of LiFe_{1-x}Mn_xPO₄

Materials for Li-ion battery cathodes and anodes have attracted much attention during the last decade.1 One compound which has received enormous interest is LiFePO42 as it possesses promising properties in terms of capacity, cycle life and safety, as well as being both cheap and nontoxic.³ Li-ion batteries with LiFePO₄ cathodes are already on the market, but there is still a need to develop new production methods in order to reduce synthesis costs and improve product control⁴. Hydrothermal synthesis, in batch continuous mode, constitutes an interesting option, since it is a cheap, environmentally benign and easily scalable method. Yang et al. (2001)were the first to show nanostructured LiFePO4 could be synthesized this way. However, the compound contained a high concentration of Li-Fe anti-site defects which have a large impact of the capacity of the final battery due to a blocking of the 1D Li-diffusion channel.⁵ By means of in situ PXRD methods we have studied the formation of the compound "live". Rietveld refinements of the data clearly show that the primary nanoparticles contain a large concentration of defects. Ordering of the anti-site defect is brought about by increasing the residence time and/or the temperature. However, even at supercritical synthesis conditions, the ordering of the compound takes several minutes. This is an important piece of insight when designing new synthesis methods for the compound.



In Situ Synchrotron Radiation Study of Formation and Growth of Crystalline $Ce_xZr_{1-x}O_2$ Nanoparticles Synthesized in Supercritical Water

The solid solution of CeO₂ and ZrO₂ exhibits materials properties applicable for many areas, ceramics and catalysis among others. The supercritical synthesis of crystalline Ce_xZr_{1-x}O₂ nanoparticles facilitates fast and easy production of the material and obviates the normal procedure of high-temperature solid state synthesis.

In situ studies where performed to elucidate the effect of the doping-level x of the solid solution on the formation reactions and final product. The *in situ* studies showed that crystalline size-stable particles less than 10 nm in size formed within 5 min at the supercritical conditions of water (P=250bar, T=375°C).

The doping-level had a marked effect not only on the phase of the crystal structure but_7 also on the morphology of the particles. The growth of the nanoparticles could be modelled and explained as being limited by either diffusion in the solvent or surface reactions on the particles. It was shown that nanoparticles with a high Ce $(x_> 0.5)$ content followed a different reaction mechanisms than nanoparticles with a high Zr content $(x_< 0.5)$.

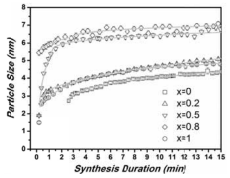


Figure 2: Growth curves of $Ce_xZr_{1-x}O_2$ particles formed under supercritical conditions

Formation and Growth of Bi_2Te_3 in Biomolecule-Assisted Near-Critical Water: In Situ Synchrotron Radiation Study

Alloys based on Bi₂Te₃ are state-of-the-art materials for thermoelectric applications. The common chemical routes for the preparation of nanostructured Bi₂Te₃-based compounds rely on toxic reductants such as hydrazine, NaBH₄, or DMF.10 In this study, a chemical "green" approach was used, utilizing the biomolecule alginic acid as a reductant. The in situ measurements gave direct evidence elemental Te (and not Bi), being an intermediate in the synthesis of Bi₂Te₃ from BiCl₃ and Te(OH)₆, which dismisses earlier theories on the reaction mechanisms.¹¹ The kinetics of the particle growth, while being very dependent on temperature, did not affect the final size of the particles. The alkaline conditions, on the other hand, had a profound influence on the stable particle sizes but had a negligible effect on the kinetics.

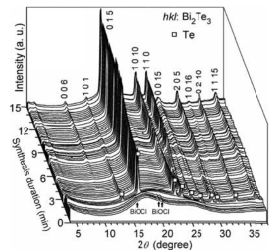


Figure 3: 3D plot of X-ray scattering data obtained from the formation of Bi₂Te₃ under near-critical conditions

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Effect of Hydration on Structural and Thermodynamic Properties of Mucin

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The mucus barrier and its transport properties are essential for proper functioning of the digestive, respiratory and reproductive systems of vertebrates, including humans. The principal components of mucus are the glycoprotein mucin¹ and water. Mucin forms the macromolecular matrix of mucus and dominates its rheological properties. From an engineering point of view, mucin is an outstanding water-based lubricant.

Mucin molecules obtained from different sources have different structures that can be described by models ranging from bottle brush to dumbbell type. It is known from literature that mucin can form liquid crystalline phases and that the phase behavior of mucin is dependent on the hydration level. However, the exact phase behavior of mucin at different temperatures and hydration levels has yet to be determined.

We studied hydration of mucin using several experimental techniques. Samples were prepared from porcine gastric mucin (PGM), from bovine submaxillary gland mucin (BSM) (both from Sigma) and from a recently slaughtered pig. To characterize the molecular structure of PGM, we used atomic force microscopy (AFM) operated in the tapping mode. AFM indicates the presence of a dumbbell structure as well as a fiberlike structure at higher concentrations. Synchrotron SAXS measurements indicate the presence of lamellar structure at concentrations of mucin higher than 15 wt% and show that repeat distance is dependent on mucin concentration. A phase transition was observed at temperatures above 60°C. A slight birefringence was observed in most samples, but polarized light microscopy discovered that birefringence arises mostly from insoluble particles of PGM. Strongly increased birefringence was observed close to the fluid – air interface which indicates that the presence of interfaces induces structuring of mucin. To observe the phase behavior of the mucin - water system differential scanning calorimetry (DSC) and isothermal sorption calorimetry were used. To characterize the glass transition of mucin at isothermal conditions, the water content and the change of the partial molar enthalpy of mixing of water at the transition point were measured. Sorption calorimetric results suggest that at low water contents mucin is in a glassy state and at higher water contents mucin is in an elastic state. Using DSC data we calculated the amount of non-freezing water as a function of concentration. Combining results obtained with different techniques we constructed a phase diagram of mucin covering a wide range of mucin concentrations and temperatures.

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Redesign of the hutches around and at the beamline I711 itself. The present experimental hutch was raised 30cm and an additional office area was built on top of the redesigned beamline. Katarina Norén and Anders Månsson outside the hutch, 7 July 2010. Photos: Jeppe Christensen

Probing the Local Structure of ZrO₂:Yb³⁺,Er³⁺ Up-Conversion Luminescence Materials by X-Ray Absorption Spectroscopy

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One of the most sophisticated and complex applications of the lanthanide luminescence is in medical diagnostics. There are, however, major problems in the use of photoluminescence based on the UV excitation in immunoassays [1], since human blood absorbs strongly the UV radiation and the emission of the phosphor in the visible [2]. A promising manner to overcome these problems is to use red up-conversion luminescence [3] resulting from the absorption of two (or more) low energy photons followed by the emission of a higher energy photon [4].

ZrO₂:Yb³⁺,Er³⁺ is one of the best up-converting materials providing red emission. Moreover, it shows considerable red afterglow, *i.e.* persistent up-conversion luminescence [5], due to the trapping of the excitation energy to lattice defects and its subsequent delayed release [6]. This suggests great potential for *in vivo* imaging without the need for any damaging excitation.

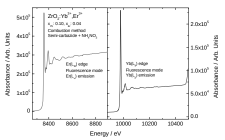
In this work, the valences and surroundings of Yb³⁺ and Er³⁺ in ZrO₂:Yb³⁺,Er³⁺ upconversion luminescence materials prepared by combustion synthesis were studied with room temperature XANES and EXAFS measurements using the beamline I811 at MAX-lab (Lund, Sweden). The measurements were carried out in the fluorescence mode using an energy dispersive solid state detector (Vortex, Si Nano Technology USA). The extraction of interatomic distances from the EXAFS data was carried out with the EXAFSPAK program package [7].

The XANES results show only the trivalent form for both the Yb and Er dopants (Fig.). This indicates that charge compensation must take place as the dopants enter a tetravalent site. Most probably, oxygen vacancies are created for this purpose according to the following scheme (using the Kröger-Vink notation): $2 Zr_{Zr}^{\times} \rightarrow 2 R_{Zr}' + V_{0}^{\bullet \bullet}$.

The distance distributions calculated from EXAFS correspond well (Fig.) to those calculated from the structural data [8] for the cubic rare earth stabilized ZrO_2 . The similarity of the Zr-Zr, Yb-Zr/Er/Yb and Er-Zr/Yb/Er distances (3.63 Å for each) confirms that Er^{3+} and Yb³⁺ occupy the Zr^{IV} sites in the structure. The R-O and R-M distributions are rather broad partly due to the high measuring temperature and partly to the multisite nature of the R positions, *i.e.* the slightly different spatial positions of the oxygen vacancies created by the charge compensation induce several slightly different R'_{Zr} sites [9].

The observed Yb-O (2.30) and Er-O (2.26 Å) distances are too short when compared with the calculated Zr-O distance (2.23 Å) when taking into account the ionic radii (Zr: 0.84, Yb: 0.985, and Er: 1.004; CN: 8 [9]). This indicates the rigidity of the structure but also the presence of oxygen vacancies around Er^{3+} and Yb^{3+} . The oxygen vacancies ($V_0^{\bullet\bullet}$) resulting from the charge

compensation are expected to agglomerate with R'_{Zr} due to electrostatic reasons. This effectively reduces the coordination number 8 around Zr to 7 (or even 6) around R^{3+} . In terms of the R-O distances, this decrease in the coordination number by one unit, corresponds to ca. 0.06 Å [9]. The R-O distances calculated from the ionic radii should be 2.375 and 2.394 Å for Yb³⁺ and Er³⁺, respectively. Thus, the average coordination number of the R^{3+} ions in ZrO₂ is not lower than 7. It may be even higher if the rigidity of the lattice is taken into account.



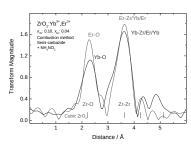


Figure. XANES spectra of the $Er(L_{III})$ and $Yb(L_{II})$ edges (left) and distance distributions calculated from EXAFS data (right), in ZrO_2 : Yb^{3+} , Er^{3+} .

Finally, according to the ionic radii, the Yb-O distance should be shorter than Er-O, but the opposite is observed. This is serendipitous, since the higher charge density of Yb^{3+} should attract more the oxide ions. Thus, the reason could be that there are more oxygen vacancies around Er^{3+} than Yb^{3+} . On the other hand, taking into account the persistent luminescence exhibited by these materials this may be explained as follows: the persistent luminescence seems to require the (at least virtual) change of the oxidation state of one of the species involved. In the Yb^{3+} , Er^{3+} system, the only change possible at ambient conditions, is the formation of the divalent Yb^{2+} (or Yb^{3+} -e $^-$) species which both demand more space than the simple Yb^{3+} ion.

Unfortunately, during the latest (May and October 2010) beamtimes at the I811 no measurements could be carried out due to the malfunction of the beamline monochromator. The present data is thus from an earlier beamtime.

Acknowledgments: The help of Dr. Katarina Norén and Dr. Stefan Carlson (MAX-lab) during the measurements is gratefully acknowledged. This work was supported by the European Community Research Infrastructure Action under the FP6 "Structuring the European Research Area" Programme (through the Integrated Infrastructure Initiative "Integrating Activity on Synchrotron and Free Electron Laser Science") as well as The Academy of Finland and the Conselho Nacional de Desenvolvimento Científico e Tecnológico (CNPq), Brazil.

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XANES characterization of $Fe_{(3-x)(1-z/3)}V_{x(1-z/3)}\Box_zO_4$ catalysts for methanol oxidation

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In the metal oxide catalyzed process for industrial production of formaldehyde from methanol the catalyst is a mixture of MoO_3 and $Fe_2(MoO_4)_3$. The catalyst is superb in many aspects, showing very high selectivity to formaldehyde (>93%) at high methanol conversion. However, in operation the molybdenum is volatile limiting the lifetime of the catalyst [1]. Consequently, alternative more stable catalysts are of potential interest provided they show comparable selectivity to formaldehyde. It is known that also vanadium based catalysts are selective for this reaction [2], but since vanadia is toxic, low volatility is required. In previous studies we have shown that the presence of iron causes stabilization of vanadia towards volatilization [3] and, moreover, $FeVO_4$ transforms into a $Fe_{1.5}V_{1.5}O_4$ spinel-type phase when being used in methanol oxidation [2]. Considering our previous results, we prepared a series of cation vacant $Fe_{(3-x)(1-z/3)}V_{x(1-z/3)}\Box_zO_4$ spinel-type catalysts with different Fe:V ratios, which are denoted $Fe_{3-x}V_x$ for short.

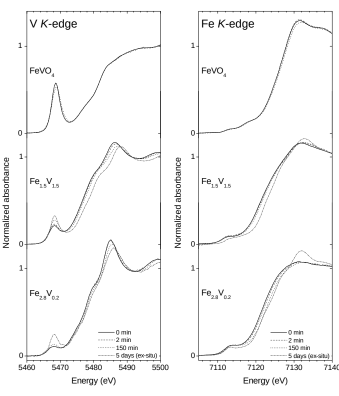


Fig. 1. a) V K-edge and b) Fe K-edge XANES spectra for FeVO₄, Fe_{1.5}V_{1.5} and Fe_{2.8}V_{0.2} as measured in situ after 0, 2 and 150 minutes and ex situ after five days on stream in methanol oxidation at 300°C (feed: 10% O₂ and 10% CH₃OH in N₂).

XANES spectra were recorded both before and after use of the samples in methanol oxidation, as well as in situ during methanol oxidation using a home-made reaction cell. Figure 1 displays V K-and Fe K-edge spectra recorded in situ immediately before (0 min) and after (2 min) the catalytic

reaction started as well as after 150 min on stream. In addition, to give information about the changes occurring over a longer period of time, ex situ spectra recorded after five days in methanol oxidation are shown for $Fe_{2.8}V_{0.2}$ and $Fe_{1.5}V_{1.5}$. At the start of the in situ experiment (0 min), the V K-edge positions ($E_{1/2}$) for $Fe_{2.8}V_{0.2}$, $Fe_{1.5}V_{1.5}$ and $FeVO_4$ are 5477.1, 5476.6 and 5480.8 eV, respectively and the Fe K-edge ($E_{1/2}$) positions are 7121.4, 7121.7 and 7124.3 eV, respectively, showing that both V and Fe in the spinel phase catalysts are less oxidized than in $FeVO_4$. During methanol oxidation, both $Fe_{2.8}V_{0.2}$ and $Fe_{1.5}V_{1.5}$ show a shift of the V and Fe main edge positions towards higher energies, revealing that both catalysts are oxidized during the reaction. As shown by the ex situ spectra, after five days on stream both V and Fe in the spinels have become even more oxidized. However, for $FeVO_4$ only negligible adjustments of the edge positions are observed after 150 min, indicating that the initial reduction of the catalyst is slower than the oxidation of the more reduced spinel phase catalysts.

Extended V K pre-edge analysis was performed following the procedure described in the literature [4,5]. In Fig. 2 are plotted the centroid position and the pre-edge intensity as obtained from the normalized and background subtracted V K pre-edge spectra together with data for several reference compounds with known coordination and valence. Since the spinel-type phase has four and six coordinated cation positions only, five coordinated positions are not likely to be found. Hence, the results suggest that trivalent vanadium in octahedral positions $^{[VI]}V^{3+}$ is oxidized to predominantly tetravalent vanadium $^{[VI]}V^{4+}$ during the reaction. The lower initial value of both the centroid energy (5468.9 eV) and the intensity (0.10) for $Fe_{2.8}V_{0.2}$ compared to the values for $Fe_{1.5}V_{1.5}$ (5469.4 eV and 0.19, respectively) reveals that V is less oxidized in the former catalyst. However, after five days in methanol oxidation vanadium in $Fe_{2.8}V_{0.2}$ has been oxidized to a slightly larger extent (+0.5 eV) than is the case in $Fe_{1.5}V_{1.5}$ (+0.3 eV), and the oxidation states have become more similar.

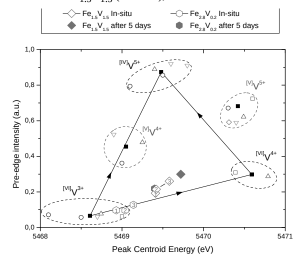


Fig. 2. Correlation between the intensity and the peak centroid energy of the background subtracted V K pre-edge spectra for Fe_{2.8}V_{0.2} and Fe_{1.5}V_{1.5}. The data encircled by dashed lines are for reference compounds with V in different oxidation states and coordinations (see ref. [6] for references). The notations 1, 2 and 3 represent the centroid position and the intensity obtained in situ after 0, 2 and 150 minutes in methanol oxidation, respectively.

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Characterization of metals (Cu and As) in mine tailings

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During the processing of the ores the mining industry produces large quantities of mine waste/tailings that can contain relatively high concentrations of metals (Cu, As, Zn etc.). The mobility of metals in these systems is dependent on the chemical speciation, which in turn is greatly influenced by the mineralogical composition and the physical and geochemical conditions prevailing in the mine waste. However, the knowledge about the specific forms of metals that occurs in mine tailings of different mineralogical composition and age is limited. The aim of this project is to use X-ray Absorption Fine Structure (XAFS) Spectroscopy to investigate the local structure and oxidation state of metal ions (Cu, As and Zn) associated with mine tailings, and secondary iron minerals, of different age collected at different mining sites.

Materials and methods

Cores of tailings were collected at three different mines in northern Sweden (Kristineberg, Aitik and Zinkgruvan) with varying mineralogical composition. At Kristineberg secondary iron mineral samples were also collected. In addition fresh tailings were sampled in the concentrator in Boliden. The tailings deposited in Kristineberg contain high concentrations of sulphide minerals in contrast to the sulphide poor tailings at Aitik. The tailings deposited at Zinkgruvan are characterized by substantial concentrations of carbonate minerals. Unmanipulated tailing and secondary iron mineral samples and manipulated tailing samples were analyzed. The manipulated samples were prepared by adding metals (Cu and As) to the tailings, to reach concentrations of ca. $15-80~\mu mol$ Me g⁻¹ on a dry mass basis. Solid phase reference compounds for Cu and As (e.g. CuS, As₂O₃) were diluted with an appropriate amount of powdered boron nitride before analysis.

As and Cu K-edge XAFS data were collected at the superconducting beam line i811 at MAX-lab. For the mine tailings data were collected on moist samples in fluorescence mode (I_f/I_0 against energy), using a Lytle detector, in the energy range from 150 eV before the edge to ca. 800 eV after the edge. However, due to problems with the beam line we only got usable data for the XANES region, which mainly gives information about the oxidation state of the metals and some information/indications about the speciation.

Results and discussion

Our Cu XANES data show that the unmanipulated sample from Boliden is dominated by CuS (Figure 1. A). In the other unmanipulated samples, both the tailing samples and the secondary iron minerals, the concentrations of both Cu and As were below the detection limit. The tailing samples with added Cu (15–80 μ mol Cu g⁻¹; pH 5.5–8.9) probably contain a mixture of CuS and Cu(OH)₂ (Figure 1 A). For arsenic the XANES data indicates that the unmanipulated sample from Boliden is dominated by arsenopyrite, FeAsS (As⁰), or arsenic sulfide, As₂S₃ (As[III]) [1, 2]. In the manipulated samples there is either a mixture between FeAsS (or As₂S₃) and AsO₄³⁻ (As[V]) adsorbed to goethite/gibbsite or as a precipitate e.g. scorodite (FeAsO₄ × 2H₂O) as in the sample from Boliden (50 μ mol As g⁻¹) or only As(V) (adsorbed or as a precipitate) as in the sample from Kristineberg (Figure 1 B). Thus, for the mine tailing sample from Boliden our XANES results clearly shows that the added metal

(Cu and As) occurs in a different form than the metal originally present in the tailing. Furthermore, our results indicate that all the tailing samples contain CuS to some degree. The speciation for arsenic in the unmanipulated tailings seems to differ more. The sample from Boliden, a fresh tailing directly from the concentrator, is dominated by FeAsS while the aged tailing from Kristineberg probably is dominated by As(V) (adsorbed or as a precipitate).

The XAFS data we have collected show that it should be possible to collect usable EXAFS data for these samples with a functioning beam line. Good quality EXAFS data is needed in order to determine the solid phase speciation of Cu and As in these mine tailing samples, e.g. it is not possible to distinguish between scorodite and As(V) adsorbed to goethite or gibbsite based on data from the XANES region alone [1, 2]. Determining the speciation of Cu and As in mine waste is of great importance because the mobility and toxicity is different for different species of the metals. For example, As bound in precipitates are considered to be less bioavailable than As adsorbed to mineral surfaces [1].

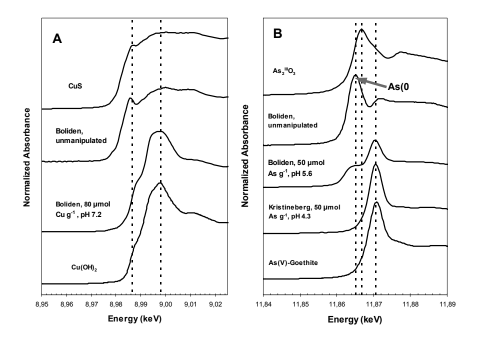


Figure 1. X-ray absorption near edge structure (XANES) spectra for ($\bf A$) copper and ($\bf B$) arsenic in reference compounds and in mine tailings from Boliden and Kristineberg. Spectra for unmanipulated mine tailings (Boliden) and for mine tailing samples where Cu and As (80 and 50 μ mol) has been added are shown.

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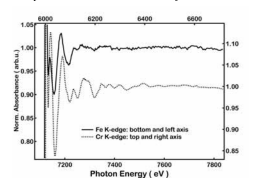
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Extended X-ray Absorption Fine Structure of Prussian blue analogues

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Molecular magnetism is at present an active area of research, given the large number of potential applications. Molecular magnet samples can be seen as an assembly of identical nanometric objects, produced at low cost and high efficiency by chemical means. They are attractive candidates for qubits, the quanta of information employed in quantum computing.^{1,2} Another interesting characteristic of molecular magnets lies in the fact that the quantum coupling between pairs of molecules can be controlled through the intermolecular interactions, by adequately tuning the spatial arrangement of the molecules on a solid substrate. Among the different classes of molecular magnets being investigated nowadays, Prussian blue and its derivatives constitute model compound suitable for basic research studies. Pure Prussian blue, contains iron and cyanide ions, and exhibits long-range ferromagnetic ordering below 5.6 K.4 However, Curie temperatures well above 300 K have been reported for several analogues containing also Cr ions. 5-7 The magnetic behaviour of these compounds can be explained using a localized orbital approach.⁸ They also show a high flexibility for tailoring their magnetic properties, since metal substitutions induce only small lattice constant changes. From the electronic point of view, Prussian blue analogues are characterized by electronic de-localization between the metal sites across the bridging cyanide anion. The combination of charge transfer effects with the strong magnetic interactions mediated by the cyanide anion gives rise to interesting properties. For instance, reversible magnetization switching mediated by external agents such as irradiation with light, 9,10 temperature 11 or pressure, 12,13 among others, has been observed; electron transfer plays a key role in all these cases. In the case of light irradiation, for Fe and Co Prussian blue analogues, photo-induced magnetism is linked with photo-induced electron transfer and a Co-N bond length variation.¹⁴ This strong correlation between electronic and structural effects motivates Extended X-ray Absorption Fine Structure (EXAFS) measurements on Prussian blue analogues. These hard x-ray data is to be correlated to work using soft x-rays on the same samples where the photo-magnetic effect is probed.

Here we present first results of EXAFS measurements on a $Fe_{II}Cr_{III}(Cr_{III}(CN)_6)$ Prussian blue analogue, at the Fe and Cr K-edges at 300K at BL I811. The samples measured were thin films grown by electrodeposition onto conducting substrates (Au-covered mylar). Previous SQUID measurements on this type of samples have revealed a Curie temperature of 200 K for the non-irradiated sample, which rises up to 225 K under laser irradiation. Here we show first structural results without laser irradiation. The measurements were performed in the fluorescence yield mode. For the Cr K-edge both the photodiode detector (Passivated



Implanted Planar Silicon from CANBERRA) and the seven-element Gresham Si(Li) detector could be used. The photodiode detector was used at the Cr K-edge in combination with a V filter for the secondary fluorescent radiation. For the Fe K-edge, a tighter filtering of the secondary radiation was required and it was necessary to use the seven-element detector.

Figure 1. The normalized x-ray absorption coefficient for a Fe, Cr Prussian blue analogue is shown versus photon energy. Data are shown at both the Fe (bottom and left axis) and Cr (top and right axis) K-edges.

The measurements were done in a He atmosphere. No time dependent effects due to beam damage were observed. The samples did not exhibit any signs of radiation damage after the data collection was completed.

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The normalized EXAFS oscillations after spline subtraction are shown in Figure 1 for both the Fe and Cr K-edges. The quality of the data allows resolving features much smaller than in the per cent range of the absorption coefficient. The next step of the EXAFS analysis is shown in Figure 2, where the normalized EXAFS oscillations are shown versus photoelectron wave vector (or k-vector). Interestingly, the amplitude of the EXAFS at the Fe K-edge is smaller versus the one at the Cr K-edge. This amplitude reduction appears to be in particularly pronounced at high k values, not allowing to have a clear EXAFS signal beyond k=10 Å^{-1} . In contrast for the Cr K-edge one can obtain data up to k=13 Å^{-1} . A possible explanation for the reduction of the EXAFS at the Fe K-edge is an increased Debye Waller factor in this case.

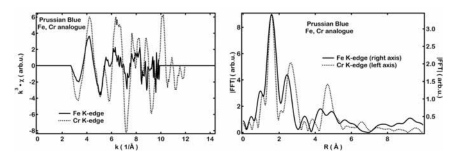


Figure 2. Left: k^3 weighted EXAFS oscillations versus photoelectron wave vector after normalization and spline subtraction at both the Fe and Cr K-edge. Right: Fast Fourier Transform modulus for the data shown in the left panel in real space.

The Fourier transform of the EXAFS data is also shown in Figure 2. Three main peaks can be identified in real space, in agreement to earlier EXAFS work¹³. The peaks around 1.8-2 Å are due to N or O. The peak around 2.5 Å is due to the C atoms. Finally the peak around 4.5 Å is due to the Cr or Fe neighbour atoms. There is good agreement in the position of the nearest neighbour atom peaks for the Fe and Cr atoms but not for the more distant neighbour atom shells. Despite the fact that the noise level is higher for the Fe Kedge data, the present data set indicates that the local environment of the Fe and Cr atoms present measurable differences. Further analysis and simulations will allow confirming these preliminary observations.

In conclusion, the present set of data indicates that it is possible to obtain high quality EXAFS on molecular magnet films. This is the first step in making further investigations in view of correlating structural effects with the electronic structure and the magnetic properties of these materials.

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Speciation of sulfur compounds in natural matrices using sulfur K edge XANES

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Sulfur XANES as an analytical tool

Sulfur K edge XANES (X-ray Absorption Near Edge Structure) spectroscopy is an excellent tool to quantitatively determine the speciation of sulfur compounds in complex matrices using internally calibrated reference spectra of model compounds [Almkvist et al., J. Synchrontron Rad. 17, 683-688, 2010]. Due to significant self-absorption of the formed fluorescence radiation in the sample itself the fluorescence signal displays a non-linear correlation with the sulfur content in a wide concentration range. Self-absorption is a problem at low total absorption of the sample when the sulfur compounds are present as particles. Particles of elemental sulfur need to be smaller than 2 µm to be give a signal linear to the concentration [Morgan et al, J. Phys.: Conf. Ser. 190, 012144, 2009]. This method developed by measurements at MAX-lab, beam-line I811, is published, and ready to be used. As beam-line I811 has not been available for users during 2010, the planned experiments in the projects "Sulfur speciation in preserved water-logged archaeological wood from the Vasa" and "Sulfur mineralization in arable soils" has been postponed.

Structural characterization metal complexes in applied and fundamental systems

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Structure of N,N-dimethylthioformamide solvated gallium(III) and indium (III) complexes

The structure of the *N*,*N*-dimethylthioformamide (DMTF) solvated gallium(III) ion has been determined in solution by means of extended X-ray absorption fine structure (EXAFS) spectroscopy. The gallium-(III) ion is four-coordinate in tetrahedral fashion with a mean Ga–S bond distance of 2.233(2) Å in DMTF solution. At the dissolution of indium(III) perchlorate or trifluoromethanesulfonate in DMTF coordinated solvent molecules are partly reduced to sulfide ions, and a tetrameric complex with the composition [In₄S₄(SHN(CH₃)₂)₁₂]⁴⁺ is formed. The structure of the solid tetrameric complex in the perchlorate salt was solved with single crystal X-ray diffraction. Four indium(III) ions and four sulfide ions form a highly symmetric heterocubane structure where each indium binds three bridging sulfide ions and each sulfide ion binds three indium(III) ions with a mean In–S bond distance of 2.584(1) Å, and S–In–S angles of 90.3(1)°. Each indium(III) additionally binds three DMTF molecules at significantly longer mean In–S bond distance, 2.703(1) Å; the S–In–S angles are in the range 80.3–90.4°. Large angle X-ray scattering data on a DMTF solution of indium(III) trifluoromethanesulfonate show that the same tetrameric species characterized in the solid state is also present in solution, whereas the EXAFS measurements only give information about the In–S bond distances due to the short core hole lifetime.

Publication

Ö. Topel, I. Persson, D. Lundberg and A.-S. Ullström, Reactions and Structures in the Gallium(III)/Indium(III)-*N*,*N*-dimethylthioformamide Systems. Inorg. Chim. Acta 365, 220-224, 2011.

XANES INVESTIGATION OF IRON REDOX CHEMISTRY IN THE ABM AND TBT EXPERIMENTS AT THE ÄSPÖ HARD ROCK LABORATORY

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Introduction

Clay barriers, rich in swelling smectite minerals, will be utilized in repositories for high level radioactive waste. The properties of the smectite are affected by the oxidation state of iron in the mineral [1]. In the field experiment Alternative Buffer Material (ABM) [2], started at Äspö Hard Rock Laboratory during November 2006, eleven clays in the form of compacted rings with an outer diameter of 0.3 m were placed in contact with metallic iron at a temperature of up to 130 °C at the -450 m level. In some cases iron cages with pellets were used and the height of the stacked rings was 3 m. The first package, of totally three, was excavated in May 2009. The field experiment Temperature Buffer Test (TBT) was started at Äspö Hard Rock Laboratory in March 2003. Compacted blocks (outer diameter 1.8 m) of MX80 clay were placed in contact with metallic iron at a temperature of up to 150 °C at the -420 m level. The hole was of full size KBS-3 type and 8 m deep. The experiment was excavated in 2010.

Experimental and results

During the whole chain consisting of excavation, transport, storage, specimen preparation and measurement, effort was put into avoiding the exposure of the clay to air. The clay was sampled radially from the iron heater towards the rock and studied using FeK XANES at beamline I811, in order to obtain spatially resolved information about the Fe redox chemistry in the deposition holes. Some results obtained on the MX80 clay in the ABM test have been summarized in Fig. 1 and in the TBT test in Fig. 2.

The position of the FeK edges in Fig. 1(a) indicated that the clay samples collected close to the heater had been reduced in the ABM test, while the samples further out were similar to the clay before deposition. The Fe pre-edge peak, which can yield useful information concerning both oxidation state and coordination, was not well defined for the excavated samples, see Fig. 1(c), and the pre-edge peak was not evaluated in the present study. Instead, the position of the edges were used to determine the relative amount of divalent iron in the samples, see Fe(II)/Fetot in Fig. 1(b). The degree of water saturation was determined by thermogravimetry and the total amount of iron by ICP analysis. The results indicated that the samples were fully water saturated and that anaerobic corrosion of the iron heater had caused an increase in the amount of Fe(II) about a centimeter into the clay close to the heater in the ABM test. Further out, the oxidation state of the clay was unaffected.

The situation was different in the TBT test, where no highly reduced samples were observed, cf. Figs 2(a) and 2(b). Instead, the samples close to the heater contained no Fe(II) at all, while the original clay (reference) contained at least a small amount of Fe(II). The clay close to the heating element was black, and magnetic particles could be collected using a permanent

magnet. The X-ray absorption spectrum of the separated particles in Fig. 2(c), indicated a pure Fe(III) compound, possibly maghemite (γ -Fe₂O₃) considering the magnetic properties. This indicated aerobic oxidation of the iron heater in the TBT test, which is compatible with the much larger scale of the clay blocks containing pores filled with air in their dry state.

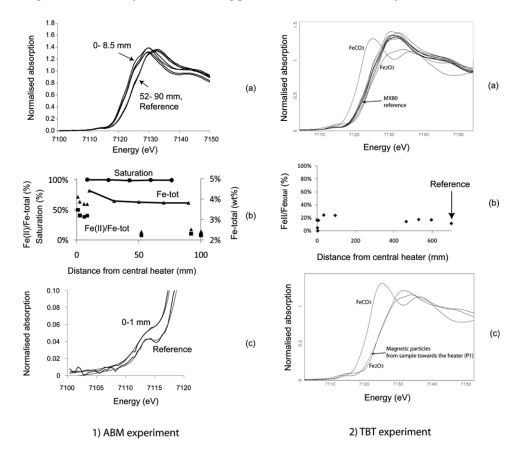


Fig. 1. The ABM test [left]. (a) FeK edges as a function of distance from the heater with the original clay as the reference. (b) Radial characteristics of the clay ring. (c) Pre-edge features of the FeK edge.

Fig. 2. The TBT test [right]. (a) FeK edges of clay samples and standard compounds. (b) Radial characteristics of the clay ring with the original clay as the reference. (c) FeK edges of magnetic particles collected from clay close to the heater and standard compounds.

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Student visits are very common at MAX-lab. Katarina Norén is giving a guided tour explaining the beamline I811 for a group of students from Polhemskolan, Lund, 18 January 2010. Photo: Annika Nyberg

Structural Basis for Receptor Recognition of Vitamin-B₁₂-Intrinsic Factor Complexes.

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Cobalamin (Cbl, vitamin B_{12}) is an essential coenzyme in mammals taken up from the diet by combined action of the gastric intrinsic factor (IF) and the ileal endocytic *cubam* receptor formed by the 460 kDa protein *cubilin* and the 45 kDa transmembrane protein *amnionless*. We have determined the crystal structure of the complex between IF-Cbl and the cubilin IF-Cbl-binding region (CUB₅₋₈) at 3.3 Å resolution. The structure provides the first insight into how multiple CUB domains collectively function as modular ligand-binding regions and how two distant CUB domains in a Ca^{2+} dependent fashion bind the two IF domains embracing the vitamin B_{12} molecule. This dual-point model provides a likely explanation of how vitamin B_{12} indirectly induces ligand-receptor coupling. Anomalous data collected at MAX-lab 911-3 was essential for confirming the presence of calcium ions in each of the four CUB domains of the receptor fragment used in this study. The comparison of Ca^{2+} -binding CUB domains and the LDL receptor-type A modules suggests that the electrostatic pairing of a basic ligand arginine/lysine residue with Ca^{2+} coordinating acidic aspartates/glutamates is a common theme of Ca^{2+} dependent ligand-receptor interactions.

Andersen, C.B.F, Madsen, M., Storm, T., Moestrup, S.K., and Andersen, G.R. (2010) *Nature*, 464, 445-8. Structural Basis for Receptor Recognition of Vitamin- B_{12} -Intrinsic Factor Complexes.

Structural studies of an Hfq orthologue from the cyanobacterium, Synechocystis sp.

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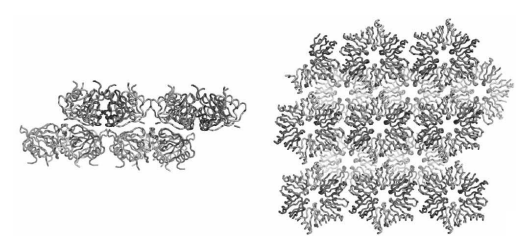
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Hfq proteins are common in many species of enterobacteria where they participate in RNA folding and translational regulation through pairing of small RNAs and messenger RNAs. The proteins share the distinctive Sm-fold and form ring-shaped structures similar to the Sm/Lsm proteins regulating mRNA turnover in eukaryotes. Bacterial Hfq proteins are usually homohexameric while eukaryotic Sm/Lsm proteins are heteroheptameric. Recently, an Hfq orthologue was identified in the freshwater cyanobacterium *Synechocystis* sp. PCC 6803, here termed Syn-Hfq¹. We have determined the crystal structure of Syn-Hfq at 1.3 Å resolution, and analysed RNA binding using synchrotron radiation at MAX-lab².



Syn-Hfq appears as stable hexamer in solution as determined by gel filtration during purification of the protein. The crystals obtained belong to the facecentred space group F222 and diffract beyond 1.3 Å. The structure was determined by molecular replacement using poly-alanine models of the known bacterial Hfq structures and refined iteratively with manual rebuilding resulting in R/R_{free} = 15.2%/20.1%. The high resolution electron density map for Syn-Hfq is exceptionally clear, except in the loop region 51-53, which appears to have alternate, un-modelled

conformation and therefore contributes to the R factors being slightly on the high side for a 1.3 Å structure. The final model has three monomers of the hexamer in the A.S.U. and covers residues 5 to 69 in chain A and residues 5 to 70 in chains B and C. In the crystals, Syn-Hfq packs in double layers of two hexamers, with the proximal sides against each other. Between the layers, the hexamers are shifted and interact with their distal sides. Hfq proteins bind RNA and are implicated in RNA remodelling and pairing of mRNAs with anti-sense regulating RNAs. We are currently studying the RNA binding of Syn-Hfq by soaking specific RNA oligos into the crystals and collecting synchrotron diffraction data.



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In-situ XRD studies on Ceramic Bone Substitutes

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In-situ X-ray scattering experiments on CSH/HA bone cement were carried out at beam line 911-5 at the MAX-laboratory in Lund. The aim was to expand earlier work and study the setting of the cement with and without vancomycin. The underlying assumption was that studying the phase transitions over time would reveal what step of the reaction vancomycin affects.

The results of the studies are presented and discussed in the Master of Science Thesis with the title "Addition of Vancomycin to a Ceramic Bone Substitute – the Impact on Gypsym Setting" by Oskar Börjesson.

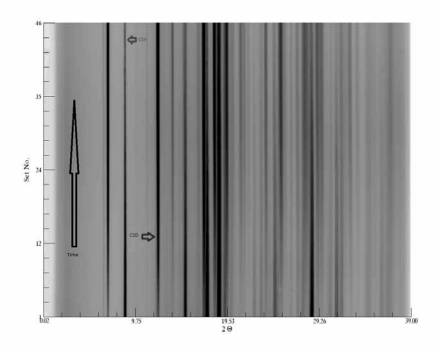


Figure. 1. 2d-plot of one series of measured samples, representing 46 single scans.

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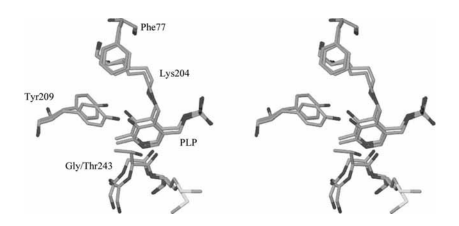
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Structural analysis of mycobacterial branched-chain aminotransferase: implications for inhibitor design

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The structure studies were based on data collected at MAX-lab beam lines 911. Data from ESRF were aslo used. The structures, apo and holo forms, were refined to 1.9 and 2.2 Å resolution. An intermediate binding site for an inhibitor was indicated. Mean $I/\sigma(I)$ were 5.3 and 4.9. The structures were refined with Rwork/Rfree values of 0.19/0.24 and 0.20/0.23.



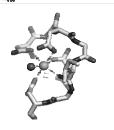
Stereoview of the structural differences between the active sites of Ms-BCAT (gold) and the mitochondrial Hs-BCAT (green; PDB code 1ekf). At the position of Gly243 in the M. smegmatis enzyme the human BCATenzymes contain a threonine residue. This difference in turn has an effect on the orientation of the side chain of Tyr209, which binds to the PLP pyridine plane on the Si face in Ms-BCATand on the Re face in Hs-BCATs. Gly243 is one of the substrate pocket-lining residues.

The structure of a GH13_31 α-glucosidase from the probiotic bacterium Lactobacillus acidophilus NCFM

Folmer Fredslund¹, Marie Sofie Møller¹, Maher Abou Hachem¹, Birte Svensson¹ and Leila Lo Leggio²

Probiotics are live microorganisms that administered in adequate amounts will confer a beneficial health effect on the host. Many beneficial effects of the probiotic *Lactobacillus acidophilus* NCFM (*La*NCFM), commercially available since the 1970's, have been reported¹. The annotated genome of *La*NCFM revealed the presence of 37 glycoside hydrolases (GHs) assigned into 13 GH families² (http://www.cazy.org), thus reflecting the significant carbohydrate catabolism potential of this bacterium. Nine genes classifiable under GH13 were identified in the genome of *La*NCFM, among which two genes encode putative intracellular enzymes, tentatively identified

Statistics for L	aGH13_31A
Data Collection	
Beamline	1911-2, MAXIab
Wavelength (Å)	1.038
High resolution	2.05
Space group	P 2 ₁ 2 ₁ 2
Unit cell parameters (Å)	a=55.83 b=107.25 c=103.61
Unique reflections	39496
Resolution (Å)	30-2.05 (2.10-2.05)
Completeness (%)	97. 2 (91.2)
Redundancy	5.3 (4.7)
Mean I/o(I)	25.55 (9.71)
R _{vym}	5.4 (20.8)
R _{rim}	5.9 (23.1)
Refinement	
Reflections work set	37 499
Reflections test set	1997
Protein atoms	4430
Hetero atoms	49
Waters	517



0.196

0.151

as oligo- α -1,6-glucosidase (O16G) and glucan- α -1,6-glucosidase (G16G), which are believed to be part of the isomaltooligosaccharide (IMO) catabolism.

The O16G from LaNCFM (LaGH13_31A) has been crystallized by optimizing initial screening conditions from a JCSG+ screen (20% glycerol, 16% PEG8000, 0.1 M MES pH 6.5 as precipitant). The structure was determined by Molecular Replacement (MR) using data collected at beamline I911-2, with statistics (from XDS³) as shown in the Table below. The model used for MR was the G16G from S. mutans⁴. The final refinement statistics (from PHENIX⁵) are shown in the Table. One of the structural features is a calcium binding site stabilizing a loop the N-terminal part of the structure (residues 20-28) and shown in the figure.

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Supported by a grant from the Danish Strategic Research Council, Committee of Health and Nutrition to the project "Gene discovery and molecular interactions in prebiotics/probiotics systems. Focus on carbohydrate prebiotics" and the Danish Research Council for Natural Science.

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Crystal structure of the copper pump

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Paradoxically some metals such as copper are essential for our bodies while being poisonous at elevated concentrations. As a consequence, heavy metal homeostasis and detoxification is crucial for cell viability. Class IB P-type ATPases play an essential role in these processes through the active extru-

	Native P1	Native P2 ₁ 2 ₁ 2 ₁	Derivative 1	Derivative 3	Derivative 4
			SeMet	Ta ₆ Br ₁₂	Na ₃ IrCl ₃
Data collection	SLS, PXI	SLS, PXI	BESSY	SLS, PXIII	MAXLAB, 911-5
Space group ^a	P1	P2 ₁ 2 ₁ 2 ₁	P2 ₁ 2 ₁ 2 ₁	P2 ₁ 2 ₁ 2 ₁	P2 ₁ 2 ₁ 2 ₁
Cell dimensions:					
a, b, c (Å)	44.2, 73.0 , 330.0	44.1, 72.9, 329.6	43.7, 72.3, 334.0	43.6, 72.5, 339.3	43.8, 72.3, 327.8
α,β,γ(°)	89.97, 90.04, 90.22	90, 90, 90	90, 90, 90	90, 90, 90	90, 90, 90
Resolution (Å) ^b	80-3.2 (3.3-3.2)	80-3.1 (3.2-3.1)	40-5.5 (6.0-5.5)	70-6.8 (7.0-6.8)	40-4.0 (4.1-4.0)
R _{sym} (%)	15.3 (104.2)	10.6 (88.4)	17.9 (77.0)	12.3 (74.8)	12.8 (53.1)
I/ơI	12.2 (1.5)	19.8 (2.0)	9.7 (2.0)	14.5 (2.1)	13.6 (3.0)
Completeness (%)	96.0 (93.0)	100.0 (99.8)	98.6 (95.8)	99.6 (96.9)	99.7 (99.9)
Redundancy	3.5 (2.7)	12.6 (9.1)	4.3 (4.3)	8.6 (8.3)	6.1 (5.5)
Data statistics for a sub-set of the data sets collected to solve the reported structure.					

sion of heavy metal ions from cells. Humans have two such pumps, ATP7A and ATP7B that both two transport Cu(I) ions across our cellular membranes and when defect these give rise to the rare

but severe Menkes and Wilson diseases, respectively.



Architecture of the copper pump. The transmembrane spanning part, with eight helices, is displayed under the characteristic P-type ATPase head piece composed of the soluble domains. Electron density maps shown at σ=2 (black), σ=1 (blue) and σ=0.5 (purple) are derived from experimental phases. Orange electron density peaks demonstrate anomalous SeMet positions at σ=4.

Using X-ray crystallography, we have determined the first high-resolution (3.2 Å) structure of the entire protein family: a Cu(I) Class IB P-type ATPase from the bacterium *Legionella pneumophila* caugh in a copper-free conformation (Figure) (Gourdon et al., Nature, *accepted for publication*). MAXLAB allowed for most of the screening of (optimised) crystals as well as for collection of some of the data sets used for solving the structure (Table).

The structure displays the three cytoplasmic domains that are common to all P-type ATPases associated the transmembrane domain. The latter contains only eight transmembrane helices of which the two most N-terminal are unique to the class IB P-type ATPAses. In addition, electron density derived from experimental phases indicate the approximate position of the so-called heavy-metal-binding-domain; a characteristic feature of the protein class. In addition, the structure suggests a putative copper transport pathway that allows the ions to be transported through the membrane. Ions are initially captured by so-called protein-chaperones that bind almost all free copper in the cell. We propose that the copper then may be transported through the protein in three stages with 1) an ion entry site (for ions from the proteinchaperones) including a conserved Asp-Met pair on the inside of the cell, 2) ion binding sites in the membrane spanning part of the protein and 3) a subsequent extracellular exit site. In addition to the mechanistic insights that our structure has yielded, the structure provides a unique framework for understanding the molecular mechanisms behind new as well as previously described missense mutations in the human ATP7A and ATP7B.

Optimization of SR beamline alignment

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Based on Synchrotron Radiation (SR) beamline optical element-by-element alignment with analysis of the alignment results an optimized beamline alignment algorithm has been designed and developed.

The alignment procedures have been designed and developed for the MAX-lab I911-4 fixed energy beamline. It has been shown that the intermediate information received during the monochromator alignment stage can be used for the correction of both monochromator and mirror without the next stages of alignment of mirror, slits, sample holder, etc. Such an optimization of the beamline alignment procedures decreases the time necessary for the alignment and becomes useful and helpful in the case of any instability of the beamline optical elements, storage ring electron orbit or the wiggler insertion device, which could result in the instability of angular and positional parameters of the SR beam.

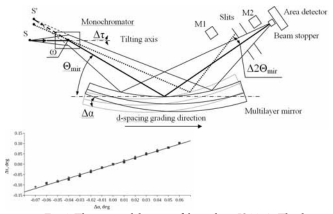


Fig.1 The general layout of beamline I911-4. The layout, drawn by dotted lines, shows the mirror when it is not aligned with the slit system. $\Delta2\Theta_{mirr}$ is the angle between the aligned and misaligned beams. S' is the SR source image at aligned position. The plot represents the functional dependency between the changes of the monochromator tilting $\Delta\tau$ and the mirror Bragg angle $\Delta\alpha$.

The I911-4 fixed energy beamline (Fig.1) is a SAXS side station beamline with a horizontally focusing monochromator (in a tilted vertical plane) and a vertically focusing curved multilayer mirror with longitudinally graded d-spacing and with several beam shaping and antiscatter slits [1]. At the first beamline alignment all slits should be opened to let the beam pass through the complete beamline unaffected. The first approximations of the parameters for the monochromator and the mirror are calculated from the beamline geometry. The monochromator theta is used to adjust the beam in the horizontal plane. The monochromator tilting together with the mirror angle and z-position are used to adjust the beam in the vertical plane. During the monochromator tilting scan the beam cone (cone-shaped diverging SR beam with the source image S') is turned around the ω -axis (which is perpendicular to tilting axis and which lies in the horizontal plane). The angles of beam cones with the fitted source images might be different, which means that only common part of beam cones can pass through the slit system. The monochromator is bent so that the focus of the SR beam is at the area detector plane. Afterwards all slits, sample holder and beam stopper are adjusted to fit the center of the beam. After finishing

the first beamline alignment all corrected optical element parameters are stored for the selected beamline setup.

Due to different reasons (unstable temperature or high heat load of optical element, unstable storage ring orbit, not stable mechanical properties of optical element) it is necessary to realign the beamline time-by-time. At least three types of realignment procedure can be identified: a) a-type - complete beamline element-by-element alignment with adjustment of all optical elements; b) b-type - adjustment of the SR beam to the slit system (sample-holder and beam stopper); c) c-type - adjustment of the slit system (sample-holder and beam stopper) to the SR beam.

The monochromator and the mirror at I911-4 cannot be considered as independent optical elements. The angle of the mirror is defined by the incident beam, which depends on the tilting angle of the monochromator, which ultimately depends on the incident beam angle in the vertical plane at the front-end. Within the rocking curve of the mirror the reflected beam passes through the aligned slit system and can be monitored. During the monochromator tilting scan monitor M2 (see Fig.1) monitors the intensity after the slit system. The functional dependence between the monochromator tilting and the mirror Bragg angle is approximately linear at small values of monochromator tilting angle. So, for deviations one can write: $\Delta\tau\sim\Delta2\Theta_{mir}\approx k\cdot\Delta\alpha$ or $\Delta\alpha\approx\Delta\tau/k$. For I911-4 the linear coefficient k is about $k\approx1.64$ (Fig.1). After completing the monochromator tilting scan the mirror correction value $\Delta\alpha$ is calculated using the measured $\Delta\tau$ and used to move the mirror to the calculated Bragg angle position. In such a way the beamline can be realigned without the realignment of all optical elements downstream the mirror (optimized b-type realignment).

A general purpose software package for manual, semi-automatic and automatic SR beamline alignment has been designed and developed using the developed algorithm. Multi-processing with Unix Interprocess Communications and RunEngines were used to make software safe to access in parallel to the beamline equipment and being protected from human mistakes and equipment failures [2]. The TANGO control system [3] is used as the middle-ware between the standalone beamline control applications BLTools (Fig.2), BPMonitor and the beamline equipment.

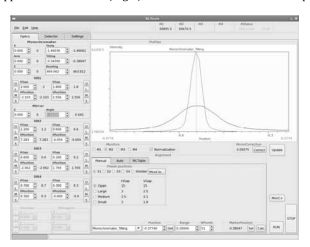


Fig.2 The view of the BLTools application during the optimized b-type realignment procedure.

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Two Novel Layered Germanates with Extra-Large Rings

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Porous oxide materials with uniform pores and channels have been of continuous interest for applications regarding size and shape selective adsorption, ion exchange, separation, and catalysis. From the assembly of large clusters such as Ge_7X_{19} (Ge_7), Ge_9X_{26-m} (Ge_9) and $Ge_{10}X_{28}$ (Ge_{10}) (X=0). OH, F, m = 0-1) a growing number of germanium oxide structures have been discovered with extra-large rings (>18-membered rings). Here we report the synthesis and crystal structures of two layered germanates, entitled SU-63 and SU-64, prepared under related conditions as ASU-21. Varying the water and/or HF content of the starting mixture resulted in the formation of the three phases. The phases each formed under well defined phase boundaries, and each phase consists of various germanate clusters.

SU-63, SU-64, and ASU-21 were prepared by hydrothermal synthesis from a gel of germanium dioxide, water, pyridine, 1,6-diaminohexane (DAH), and HF (40 wt%) with a molar ratio 1:x:39.1:20.0:y. The range of water/GeO₂ and HF/GeO₂ ratios for each phase are indicated on the phase diagram in figure 1. The mixtures were stirred for two hours, placed in 22mL Teflon-lined autoclaves, and then heated at 160°C for seven days. All product crystals were filtered, washed with distilled water, and dried at room temperature overnight.

Single crystal X-ray diffraction data of SU-63 were collected on a MarCCD at 100(2) K using a synchrotron radiation (λ=0.91Å) at the Beam line 1911:5, Max Lab, Lund University, Sweden. Data reduction and numerical absorption correction were applied by the TwinSolve software.4 Single crystal X-ray diffraction data of SU-64 were collected on a Saturn 724+ detector at 150(2) K using a synchrotron radiation $(\lambda=0.69\text{Å})$ at the Beam line I19 (EH1), Diamond Light Source, UK. Data reduction and numerical absorption correction were applied with d*TREK within CrystalClear.⁵ The structures were solved and refined by SHELX.

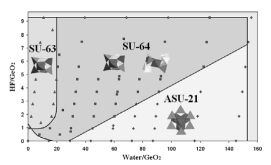


Figure 1 The phase diagram of SU-63, SU-64, and ASU-21 with water content and HF as variables. Polyhedral representations of the clusters that compose the various phases are illustrated with octahedral GeO_6 in red, trigonal bipyramidal GeO_5 in yellow, and tetrahedral GeO_4 in green. Gray points indicate absence of a porous crystalline product.

Crystal data for SU-63 (|1.5H₂DAH|[Ge₇O₁₄(OH)₃]·2H₂O): P6₃cm, a = 28.794(2) Å, c = 20.603(4) Å, V = 14793(3) Å³. 150 × 80 × 20 µm³. A total of 51182 reflections, of which 8692 are unique. $R_{\text{int}} = 0.0954$, R = 0.0851 and wR2 = 0.2163 for reflections with I > 2 σ (I). Crystal data for SU-64 (|11H₂DAH|[Ge₉O₁₈OH₄][Ge₇O₁₄(OH)₃]₆·16H₂O): P-I, a = 12.101(4) Å, b = 18.113(6) Å, c = 22.444(8) Å, a = 87.86(1) °, $\beta = 89.50(2)$ °, $\gamma = 83.17(1)$ °, V = 4881(3) Å³. 30 × 8 × 2 µm³. A total of 42228 reflections, of which 15210 are unique. $R_{\text{int}} = 0.1435$, R = 0.1296 and wR2 = 0.3363 for reflections with I > 2 σ (I).

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SU-63 is built of stacked layers of Ge₇ clusters decorating a 3.6.3.6 Kagomé lattice. The clusters form 6 and 18-membered rings in the *ab*-plane. 18-rings of neighboring layers stack resulting in one-dimensional channels. DAH was disordered between the layers with their presence confirmed by CHN elemental analysis and thermogravimetric analysis. SU-64 is a slab structure with strips of Ge₇ clusters joined by Ge₉ clusters. Intersecting 10 and 18-ring channels run through the lengths of the slab. DAH was located within the 18-ring channels and between slabs. In situ XRPD revealed changes in the unit cell angle from 88.21 to 79.57° at 30°C and 200°C respectively suggestive of a shift in the stacking of layers.

As indicated in the phase diagram, within this system the rough concentration of HF affected the type of cluster building unit used to fabricate the resulting crystalline phase. Low concentrations of HF lead to ASU-21, a 3D Ge_{10} framework built from hexagonal packing of rigid cylinders. High concentrations lead to SU-63 with Ge_7 clusters, while relatively intermediate HF concentrations lead to SU-64 combining Ge_7 and Ge_9 clusters.

We have presented two novel germanate structures that have been prepared by modifying the synthesis conditions of ASU-21. Linking synthesis conditions with the final crystalline product may lead to better control over the formation of germanate clusters and consequently lead to the discovery of other novel structures.

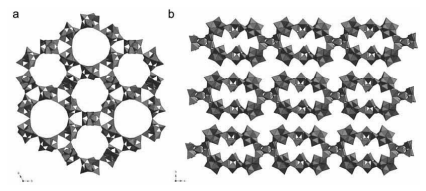


Fig. 2 Polyhedral representations of (a) SU-63 and (b) SU-64.

Acknowledgement

We thank M. Peskov for his initial topology analysis of SU-63 and SU-64. C. Bonneau and J.-L. Sun were supported by post doctoral grants from the Wenner-Gren and Carl-Trygger Foundations, respectively. This project is supported by VR and VINNOVA through the Berzelii Center EXSELENT and Göran Gustafsson Foundation.

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Structural studies on the human mitochondrial

acetoacetyl-CoA thiolase (T2)

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Thiolases have many important roles in different biochemical pathways. Depending on subcellular locations they participate either in the degradative or biosynthetic pathways. Currently, in humans, six different thiolases have been identified. Three of those are mitochondrial enzymes, two can be found within peroxisomes, and one is a cytosolic thiolase. The peroxisomal thiolases are homodimeric enzymes; the rest are homotetrameric enzymes, except the mitochondrial protein complex which is composed of four α and four β subunits; the β subunit is thiolase. Two of the characterised thiolases, mitochondrial acetoacetyl-coenzyme A (CoA) thiolase (T2) and cytosolic acetoacetyl-CoA thiolase (CT), are short-chain specific acetoacetyl-CoA thiolases and are classified as biosynthetic thiolases (EC 2.3.1.9). Enzymatically T2 can be distinguished from the other thiolases by its requirement for potassium ions for full activity². The kinetic measurements showed that in the presence of K⁺ the catalytic efficiency of T2 increased approximately six fold³. Furthermore, T2 degrades 2-methyl-acetoacetyl-CoA exclusively.

The biosynthetic thiolases catalyse the nondecarboxylating Claisen condensation of two acetyl-CoAs to form acetoacetyl-CoA⁴; this involves the catalytic triad of Cys-Cys-His. In the first half of the reaction, one of the active site cysteine becomes acetylated (Figure 1). In the second half of the reaction another acetyl-CoA molecule enters the active site and the above mentioned Claisen condensation reaction, the formation of C-C bond, takes place (Figure 1). Finally, acetoacetyl-CoA is released.⁵

$$C_{0A-S}$$
 + H_{S} $C_{ys_{126}}$ C_{0A-SH} + H_{S} $C_{ys_{126}}$ C_{0A-S} + H_{S} $C_{ys_{126}}$

Figure 1: The Claisen condensation reaction, as catalysed by T2. In the first part of the reaction cycle (the top panel) the catalytic cysteine, Cys_{126} , becomes acetylated. During the second part of the cycle another acetyl-CoA enters the active site and acetoacetyl-CoA is released (the bottom panel).

Even though the reaction mechanism of bacterial biosynthetic thiolase (bCT) has been studied extensively, the reaction mechanism is not yet fully understood^{5,6}. Our ultimate aim is to understand the fine details of these reaction mechanisms which will give us more information on how these enzymes work. Especially, we are interested in reaction intermediates and transition state intermediates of the reactions catalysed by thiolases. This would then allow us to design new inhibitors of these enzymes, for instance.

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The macromolecular crystallisation facility at MAX-lab

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An efficient facility for low-volume screening of crystallisation conditions has been in operation at MAX-lab for five years. The facility is open for all academic users and is run by SARomics Biostructures AB on behalf of MAX-lab. The equipment consists of a TECAN Freedom EVO 150 robot for general liquid handling, a Mosquito nanolitre pipettor from TTP Labtech that is used to set up the crystallisation drops, and a CrystalPro imaging system from Tritek Corporation for photographic documentation and database storage of the experiments. We also have a Stratagene Mx3500p qPCR machine for differential scanning fluorimetry. The Stratagene equipment is used to find stabilising buffer conditions and additives.

During 2010 the facility was used by 23 scientists from 14 research groups: 11 at Lund University (including the Departments of Biochemistry and Structural Biology, Pure and Applied Biochemistry, Biophysical Chemistry, Organic Chemistry, Cell and Organism Biology, Glycobiology and Biotechnology), one from Copenhagen University, one from the Carlsberg Laboratory in Copenhagen and one from Uppsala University. In addition SARomics Biostructures has carried out 6 academic projects (in collaboration with the Departments of Immunotechnology, Biotechnology, BMC, Clinical Chemistry and Pure and Applied Biochemistry at Lund University, as well as researchers from the University of Hannover in Germany), resulting in many new crystallisation conditions and five new crystal structures.

Results from the macromolecular facility have been presented in four publications (Diehl et al. 2010, Helgstrand et al. 2010, Johansson et al. 2010 and Leinartaité et al. 2010) and two dissertations (G. Svensson, 2011 and R. Yengo, 2011). The generally long lag time between crystallisation and publication of crystal structures means that several more publications are in the pipeline.

In late 2010 the crystallisation facility received a grant of 3.4 MSEK from the Swedish Research Council in order to undertake a major upgrade of the equipment. In the first hand the grant will be used to improve the experimental imaging system to include a "plate hotel" that will improve throughput. We also aim to extend the range of biophysical characterisation techniques that we offer to include high-throughput dynamic light scattering. With these improvements the facility will take important steps to becoming a national laboratory integrated with the structural biology beamlines at MAX IV and with a pipeline for protein production within Lund University.

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Our annual Christmas card



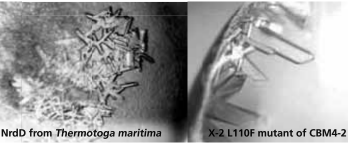
Some highlights from the Macromolecular Crystallization Facility 2010





Human galectin-3C in complex with inhibitor





E. coli aldehyde dehydrogenase in the apo form and in complex with NAD^{\dagger}

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Aldehyde dehydrogenase is an enzyme which catalyses the conversion of aldehydes to acids. In the *E. coli* variant of aldehyde dehydrogenase (ALDH) the co-factor NAD(P) participates in the oxidation process. The mechanism involves a nucleophilic attack by a cysteine followed by hydrogen transfer and release of the carboxylic acid, see figure 1.

Figure 1: Reaction mechanism of aldehyde dehydrogenase

This ALDH was reported to be active with 3-hydroxy propionladehyde (3-HPA), which is of industrial importance in producing biopolymers after oxidation to 3-hydroxy propionic acid. Propionaldehyde was used as a substrate for obtaining the kinetic data. The ALDH was shown to be active using NAD+ / NADP+ with higher specific activity with propionaldehyde in presence of NAD+ as cofactor (Jo et al., 2008). The kinetic data with ALDH wih propionaldehyde in presence of NAD+ showed in Table 1.

Table 1. Kinetic data of ALDH with propional dehyde as a substrate (manuscript in preparation)

V_{max}	11,72 U/mg
k _m	0,61 mM
K _{cat}	11,57 s ⁻¹
K _{cat} /k _m	19,09 x10 ³ M ⁻¹ s ⁻¹

In this project one of the aims is to get help from the structure to design a "better" functioning enzyme with respect to K_m and K_{cat} by mutagenesis. Aldehyde dehydrogenase from E. coli (ALDH) was overexpressed in E. coli with a N-terminal His-tag followed by a 50 amino acid expression vector tag and then the 495 amino acid ALDH construct. His-trap columns were used to purify the protein and crystallization trials were set up at the MAX-lab macromolecular crystallisation facility. Data on apo ALDH were collected at 1911-2 to 1.9 Å resolution. The structure was determined using molecular replacement and the PHASER program (McCoy et al., 2007) using the 42 % sequence identical structure of bovine aldehyde dehydrogenase (PDB code 1AG8) as a template. Two copies of the enzyme were found in the asymmetric unit. The biological assembly of the protein is a tetramer but in this case the tetramer is formed by crystallographic symmetry, all four molecules connected by two β sheets including β strands from the four monomers. One calcium site per monomer was found in the apo ADH structure, however only partially occupied.

The NAD+ form of ALDH was formed by soaking crystals over night in stabilising solution containing 2 mM NAD+ and 2 mM CaCl₂. In this way full occupancy of Ca²⁺ sites and NAD+ sites was achieved. Data on the NAD+ form were collected at I911-2 to 1.75 Å resolution. The reactive cysteine, Cys302, changes conformation upon NAD+ binding since its apo conformation collides with the nicotinamide ring of NAD+. With NAD+ bound the side chain of Cys302 has the appropriate conformation to readily form the thioacyl intermediate (step 2 in figure 1) since it is in close proximity to NAD+ and Asn168, which probably stabilises the intermediate, as has been shown for a homologous enzyme (D'Ambrosio et al., 2006).

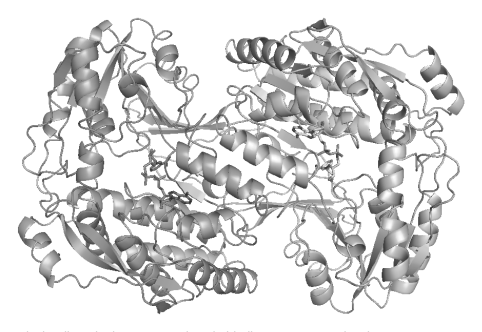


Figure 2) The dimer in the asymmetric unit, binding one NAD molecule per monomer

Current status: manuscript on structure is in preparation

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Non-specific CBM4-2 variant X-2 L110F - apo, xylopentaose, cellopentaose structures

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Carbohydrate binding modules can be used for developement of specific binders for use e.g. in bioanalysis. A key aspect of the use of carbohydrate-specific probes in a variety of applications is their ability to differentiate between the many different carbohydrates that exist in biological samples. Specific antibodies are often used as probes to detect biomolecules in complex samples but they are often not that specific for individual carbohydrates (Pattathil et al., 2010). The carbohydrate-binding module CBM4-2 derived from Rhodothermus marinus, has also been explored to develop carbohydrate-specific probes (Cicortas Gunnarsson et al., 2004) for use in a variety of applications (Filonova et al., 2007; von Schantz et al., 2009; Sandquist et al., 2010). In the work presented here we have elucidated at the molecular level features of carbohydrate recognition of highly similar specific and cross-reacting probes. A single amino acid mutation L110F turns the specific xylose binding CBM4-2 variant X-2 (Cicortas Gunnarsson et al., 2007) into a more non-specific binder of not just xylose oligomers but also linear glucose oligomers as well as branched carbohydrates, such as xyloglucan (von Schantz et al., in preparation). The structures of X-2 alone and in complex with xylose oligomers have been determined previously (Håkansson et al., 2009; von Schantz et al., in preparation) within this project. The structures of the apo variant, the xylopentaose and cellopentaose co-crystal structures of X-2 L110F were now determined to 1.1, 1.4 and 1.3 Å resolution respectively. Crystals were made in MRC plates or NEXTAL plates at the macromolecular crystallization facility at MAX-lab. Diffraction data were collected at beamline 1911-2. The structures were determined using molecular replacement using the previously determined structure of the X-2 variant of CBM4-2 (von Schantz et al., in preparation).

The overall structure of X-2 L110F and X-2 is the same (r.m.s. deviation of $C\alpha$ = 0.17 Å for 152 positions compared). The mutation L110F that confers cross-reactive binding properties to the module is situated in the central cavity of the carbohydrate binding cleft. The larger hydrophobic surface of the phenylalanine (compared to a leucine) in combination with plasticity of side chains in the carbohydrate binding cleft allows it to accommodate sugar oligomers based on monomers other than xylose, see figure 1. However only two or three out of five sugar rings are visible in the cellopentaose-containing structure probably due to absence of surface complementarity and clashing side chains.

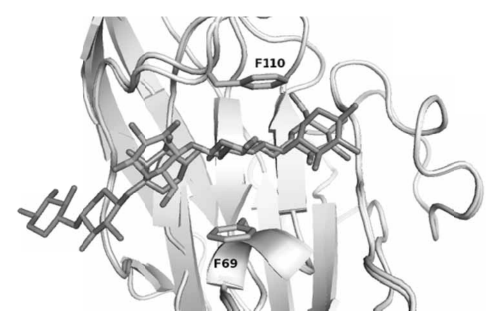


Figure 1) Xylopentaose bound to X-2 L110F aligned with cellopentaose bound to X-2 L110F. The three sugar rings (out of five) possible to model in the electron density maps are sandwiched between F110 and F69. The xylose makes more interactions to X-2 L110F and is therefor not so dependent on the sandwich interaction between the phenylalanines (all five xylose rings visible).

Current status: manuscript on structures is in preparation

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Crystallography and microsecond multiple molecular dynamics simulation in concert: Structure of an intrinsically disordered nerve agent antidote in complex with sarin-AChE

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Organophosphorous (OP) nerve agents and insecticides are some of the most poisonous chemicals known to mankind. They inhibit the essential enzyme acetylcholinesterase (AChE) by a rapid phosphylation of the catalytic serine residue. Compounds such as fenamiphos and methamidophos are widely used in modern agriculture, and the nerve agents continue to be a threat in the world due to their possible use in terrorist acts. The OP insecticides is estimated to cause between 2 to 5 million intoxications per year, of which around 300 000 are fatal.

Design of effective nerve agent antidotes to various OP compunds requires information on how the reactivators interact with the phosphonylated AChEs. However, antidotes (*i.e.* reactivators), are preferably are flexible or mobile to be able to convert from the ground state to the transition state for reactivation. This mobility makes reactivators intrinsically disordered in crystal structures. Moreover, complexes of nerve agents and antidotes are chemically unstable, further complicating crystallographic studies.

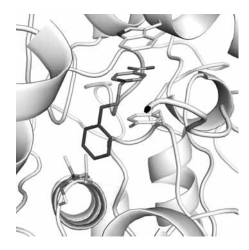


Figure 1. The 2.2 Å crystal structure of sarin-AChE in complex with HI-6. Sarin is shown in yellow while the ordered and disordered portion of HI-6 is shown in orange and magenta, respectively. The reactive part of HI-6 could not be built in the electron density map, but was analysed using MMDS.

A strategy to address the challenges associated with structural studies of reactivators is exemplified in our work with a ternary complex of acetylcholinesterase (AChE), inhibited by the nerve-agent sarin in complex with the reactivator HI-6¹. In this study, the ordered portion of HI-6 was determined by X-ray crystallography (Figure 1). The crystal structure was used as a starting structure in a Multiple Molecular Dynamics Simulation (MMDS) where the reactive (disordered) portion of HI6 was investigated. In addition, enzyme kinetics was employed to cross-validate the structural data. Based on these studies, we suggest that the nucleophilic part of the reactivator (*i.e.* the oxime moiety) enters the active site gorge of AChE in its neutral (protonated) form and that a hydrogen bonding network among a tetrad (Glu334-His447-wat-oxime) is necessary to deprotonate the oxime prior to reactivation. The importance of a hydrogen bonding network between Glu334, His447 and the oxime is supported by the complex between AChE, the insecticide fenamiphos and the reactivator ortho-7². The study of fenamiphos also implicates that entrapment of the reactivator at the ground state may contribute to a low reaction rate by preventing the oxime to reach the transition-state for reactivation. In our opinion, these studies show that mobility or flexibility is as important as affinity for reactivator design.

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Flexibility and communication within the structure of the

Mycobacterium smegmatis methionyl-tRNA synthetase (metRS)

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The structure studies were based on diffraction data collected at the 911 beam lines at MAX-lab in Lund. Two complexes of the enzyme were included in the study; metRS in complex with the substrates methionine (alone) and the methionine and adenosine. Data were collected with a beam wavelength of 1.038 Å, and the diffraction pattern are obtained to 2.3 and 2.8 Å resolution. Mean $I/\sigma(I)$ were 11.8 and 17.8 for the two data sets and $R_{p.i.m}$ 5.1 and 3.3. The structures were solved with molecular replacement and could be refined to the limits of the data collections. The structures demonstrate the flexibility of the multidomain enzyme. A new conformation of the structure has been identified in which the connective peptide domain binds closer to the catalytic domain than described earlier. The potential use of the adenosine binding site for inhibitor binding has been evaluated and a potential binding site for a specific allosteric inhibitor has been identified.



Overall structure of metRS from *Mycobacterium smegmatis*



Ana Labrador presenting the new multipurpose SAXS station for Marc Obiols-Rabasa, Lund University, at the poster session during the Annual User Meeting, 8 November 2010. Photo: Annika Nyberg

MAX-lab ACTIVITY REPORT 2010 343

Zebra fish interferons

Ole Jensen Hamming and Rune Hartmann

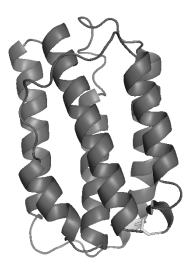
Centre for Structural Biology, Department of Molecular Biology, Aarhus University, Denmark

Cells respond to viral infections by producing hormone-like compounds called interferon's (IFNs), which are danger signals that warn surrounding cells of the approaching danger (the virus). We are investigating how these cytokines interact with their cognate receptor and induce signalling across the cell membrane.

We are currently engaged in an effort to determine the evolutionary relationship between the interferons of mammals and zebra fish. The zebra fish has homologs all the internal signalling and transcriptions factors involved in interferon signalling in mammals. It is however not known if the interferons of zebra fish corresponds to mammalian type I or III interferons. Determining this will help to establish to what extend the zebra fish can be used as a model organism of interferon action. During 2009 we collected data on zebra fish IFN-1, (IFN Φ 1). We managed to get a dataset with a resolution of 2.1. For further details see table 1. The structure was solved (Figure 1) using molecular replacement with a model of zebra fish interferon 2 that was solved using data collected at maxlab in 2008. Both these structures form part of my PhD thesis (1). A paper of the structures of the Zebra fish interferons is currently being written.

Table 1: Details of data collected on Zebra fish interferon crystal.

Data collection		
Beam line	Lund I-911	
Soak	-	
Anomalous signal (f´´)	-	
Wavelength (Å)	0.87420	
Space group	P2 ₁	
Cell dimensions a/b/c (Å)	48.834/62.247/51.306	
Cell angels α/β/γ (°)	90.000/91.424/90.000	
Resolution (A°)	2.10	
Reflections	60560	
Completeness (outer shell) (%)	96.3 (78.9)	
Data redundancy (outer shell)	3.5 (2.4)	
Mean I/σI (outer shell)	2.1 (2.7)	
Rsym (outer shell) (%)	9.1 (32.4)	
Resolution range used (A°)	48.818-2.086	
R (%)	23.72	
Rfree (%)	18.13	
Number of atoms/solvent	2776/179/2	
molecules/ions		



Figur 1: Structure of zebra fish interferon 1.

The structure of Zebra fish interferon 1 solved to 2.1 Å.

The structure shows the classical type II cytokine fold with 5 long and 1 short helix forming an elongated "American football" type structure.

Based on the structure of the Zebra fish interferons we determined that the zebrafish interferon system is closly related to the mammalian type I interferons. This means that the zebrafish is a convinient modelorganism for the mammalian type I interferon system.

1: Hamming, O.J PhD thesis:"The Interferon System of Humans and Fish – A Structural Point of View".

High-resolution crystal structures of the flavoprotein Nrdl in oxidised and reduced states: an unusual flavodoxin

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The small flavoprotein NrdI is an essential component of the class Ib ribonucleotide reductase system in many bacteria¹. NrdI interacts with the class Ib radical generating protein NrdF. It is suggested to be involved in rescue of inactivated diferric centres or generation of active dimanganese centres in NrdF². Though NrdI bears a superficial resemblance to flavodoxin, its redox properties have been demonstrated to be strikingly different³. In particular NrdI is capable of two-electron reduction, while flavodoxins are exclusively one-electron reductants. Biochemical studies

	oxidised	reduced
Unit cell dimensions (Å)	a = 42.8	a = 42.83
	b = 45.6 c = 56.3	b = 45.26 c = 55.66
Data collection		
X-ray wavelength (Å)	0.7300	0.9077
Resolution range (Å)	23.9-0.96 (0.98-0.96)	23.7-1.4 (1.44-1.4)
Completeness (%)	97.3 (77.8)	99.3 (98.7)
R _{merge} (%)	3.9 (60.1)	9.3 (74.4)
<i>/<σ(I)></i>	17.1 (2.0)	9.1 (1.5)
No. of observations	283726	102202
No. of unique reflections	65490	21820
Wilson B-factor (Å ²)	11.1	21.5
Refinement		
Resolution range (Å)	25-0.96 (1.0-0.96)	26.7-1.4 (1.46–1.40)
R _{model} (%)	12.5 (25.9)	14.7 (23.8)
R _{free} (%)	15.2 (-)	19.2 (32.5)
Mean isotropic B-factor (\mathring{A}^2)	13.4 (protein) 7.3 (FMN) 34.2 (water)	14.3 (protein), 10.5 (FMN) 39.2 (water)
Rms deviations from ideal geo	ometry	
Bond length (Å)	0.016	0.024
Bond angles	$0.035~\textrm{\AA}^{1}$	1.99°

on NrdI from *Escherichia coli* led to the conclusion that these unusual redox properties depended on a lesser destabilisation of the negatively charged hydroquinone state of the FMN cofactor than what is common in flavodoxins, as NrdI was predicted to be much less negatively charged than flavodoxin³. We have determined the crystal structures of NrdI from *Bacillus anthracis*, the causative agent of anthrax, in the oxidised and semiquinone forms, at resolutions of 0.96 Å and 1.4 Å respectively, using data from stations I911-3 and I911-5 at MAX-lab respectively⁴ (Table 1).

The structure of NrdI revealed an unusually compact flavodoxin fold, defining a new structural subfamily within the flavodoxins (Fig. 1). The conformational behaviour of NrdI in response to FMN reduction is very similar to that of flavodoxins, involving a peptide flip in a loop near the N5 atom of the flavin ring. NrdI from B. anthracis is indeed much less negatively charged than flavodoxins, which can be expected to affect its redox properties significantly. However analysis of 199 NrdI sequences demonstrated a remarkable spread in the predicted isoelectric points (pI) of NrdIs, from approximately pH 4-10, whereas the pI values of classical flavodoxins almost all lie between pH 4.0-4.5. The reason for this is not

known, but the results imply that the redox chemistry of NrdI is not as simple as one might expect from analysis of the *E. coli* protein alone.

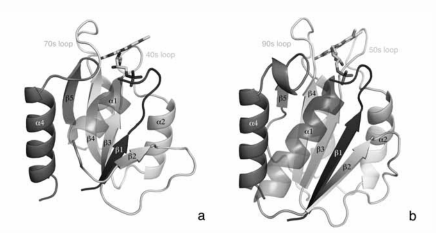


Fig. 1: Comparison of the overall structures of a) NrdI and b) the most structurally homologous standard flavodoxin, the short chain protein from *D. desulfuricans* (PDB ID 3F6R). Secondary structure elements are labelled.

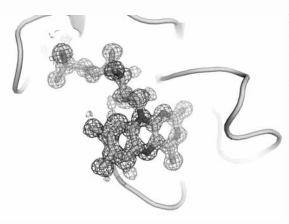


Fig. 2: Electron density for the FMN cofactor in NrdI at 0.96 Å resolution. Difference electron density (lighter shade) can clearly be seen for several non-exchangeable protons on the cofactor.

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Structural and mechanistic basis for new modes of inhibition of glycosyltransferases

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Glycosyltransferases (GTs) are a large class of enzyme, encoded by 1-2% of the genes in all genomes sequenced to date (www.cazy.org). They are responsible for the biosynthesis of all glycosidic linkages in Nature- in polysaccharides, oligosaccharides and glycoconjugates which are carbohydrates linked to non-carbohydrate molecules. Complex glycoconjugate biosynthesis requires the concerted action of numerous of glycosyltransferase enzymes that catalyze the transfer of a mono- or oligosaccharide from a glycosyl donor (most frequently a sugar-nucleotide) to an acceptor. Acceptors are carbohydrates, lipids, proteins, DNA, natural products (antibiotics, flavonols, sterols etc) and unnnatural compounds such as drugs and xenobiotics ¹⁻⁴. GTs produce the three most abundant biopolymers, cellulose, chitin and starch. GTs play a key role in many fundamental biological processes underpinning human health and disease, such as cell signaling, cellular adhesion, carcinogenesis and cell wall biosynthesis in human pathogens⁵⁻⁸. The development of small-molecule GT inhibitors is therefore of considerable scientific interest in chemical glycobiology and drug discovery^{9,10}.

We have developed several new, base-modified UDP-Gal derivatives with an aromatic or heteroaromatic substituent in position 5 of the uracil base as chemical tools for the investigation of glycosyltransferases and other UDP-Gal dependent glycoprocessing enzymes (Figure 1)¹¹. The most potent of these new derivatives with a formylthienyl substitutent, acts on five different GTs, as an inhibitor of glycosyl transfer, with K_i values in the low micromolar to nanomolar range. Other substituents (phenyl, furanyl, 4-MeO-phenyl) are not as potent¹¹.

Fig. 1. UDP-Gal derivative where different substituents are placed at position 5 of the uracil base.

To understand the molecular basis for the enzymological behavior of these inhibitors we chose a mutant of the ABO(H) blood group A and B glycosyltransferases which catalyze the final step in the synthesis of the A and B antigens. This mutant is a cis-AB mutant capable of transferring both Gal and GalNAc to the H-antigen with equal efficiency. We have solved high-resolution crystal structures of several of the for inhibitors bound to the cis-AB mutant, which crystallizes particularly well, and for direct comparison, structures of the enzyme in its unliganded apo form and in complex with UDP. Several of the data sets for these structures were collected at beamline I911 at MAX II of which many

of the crystals diffracted to near-atomic resolution (1.2–1.4 Å resolution).

Surprisingly the inhibitors block the closure of a flexible loop in the active site of a human blood group GT by preventing the stacking of two amino acid residues where one is placed in the flexible loop and one in the C-terminus. This is a new mode of inhibition for GTs that, given the strong mechanistic similarities between many GTs^{1-4,12}, and should be applicable to other GTs. The differential potency of inhibitors with various substituents is attributed to the extent to which loop closure is blocked or by abortive interactions with the loop to give non-productive complexes.

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Crystal Structure of the DEAD-Domain of Human RNA Helicase DDX10/Dbp4

Karlberg, T., Schütz, P., Berglund, H., Collins, R., Flodin, S., Flores, A., Gräslund, S., Hammarström, M., Johansson, I., Kotenyova, T., Moche, M., Nielsen, T.K., Nordlund, P., Nyman, T., Persson, C., Siponen, M.I., Svensson, L., Thorsell, A.G., Tresaugues, L., Welin, M., Weigelt, J. and Schüler, H.

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Introduction

RNA helicases of the DExD/H-box superfamily are critically involved in all RNA related processes. DDX10 is critically involved in ribosome assembly, likely by assisting the folding of RNA constituents of the ribosome. DDX10 gene fusion with the nucleoporin NUP98 causes leukemia by an unknown mechanism. Many human DEAD-box helicases are considered drug targets for cancer treatment, since their inhibition affects general cell growth. Some DEAD-box proteins are involved in the processing of viral and other pathogen RNA, and these are considered drug targets for infectious disease. As a consequence, understanding the structures and functions of human DEAD-box helicases is of explicit medical importance, as it will guide the development of selective drug therapies.

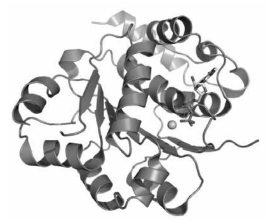


Figure 1. Human DDX10 in complex with magnesium and ADP.

Results

We solved a crystal structure of the conserved domain-1 of the human RNA helicase DDX10. It consists of a RecA-like core, and contains the DEAD-box motif with the nucleotide binding site occupied by magnesium and ADP. The crystal structure shows how a divergent tyrosine in the active site contributes to nucleotide binding.

Together with crystal structure of 10 other DEAD-box domains, the structure of DDX10 enabled us to carry out a structural analysis of the events associated with ATP hydrolysis and RNA remodeling (Ref. 1). We propose that consensus α -helix 8 in the DEAD-domain partially blocks the RNA binding site, and that ATP binding releases this blockage by pulling consensus α -helix 8 out of the RNA binding site.

Materials and Methods

Protein Expression and Purification

The cDNA encoding full length human DDX10 was retrieved from the Mammalian Gene Collection (BC093656). *E. coli* strain BL21(DE3) R3 pRARE cells transfected with expression vector encoding human DDX10⁴⁷⁻²⁸⁰ including a TEV protease cleavable N-terminal hexahistidine tag were grown in TB medium supplemented with 8 g/l glycerol and 50 μg/ml kanamycin. After induction with 0.5 mM IPTG, culture were grown at 18°C for 20 h. Cell pellets were resuspended in 50 mM HEPES pH 7.8, 500 mM NaCl, 10 mM imidazole, 10 % glycerol, 0.5 mM TCEP, and Complete EDTA-free Protease Inhibitor (Roche Biosciences). Protein was purified using IMAC (HisTrap HP, GE Healthcare), gel filtration (HiLoad Superdex-200, GE Healthcare) and the hexahistidine tag was removed by TEV cleavage. Protein was concentrated using Vivaspin (Sartorius) cartridges and stored as aliquots at -80°C.

Crystallography

Crystals of DDX10⁴⁷⁻²⁸⁰ were obtained by vapor diffusion in sitting drops at 4°C (for details, see **Ref. 1**). Diffraction data were collected at MAXIab II beamline I911-2. Data were indexed and integrated in space group P6122. The structure was solved by molecular replacement with MOLREP, using PDB entry 2gxs as search model. Refinement was done using Refmac5, and Coot was used for model building.

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Human poly (ADP-ribose) polymerase family, member 14 - macro domain 2

Karlberg, T., Schütz, P., Berglund, H., Collins, R., Flodin, S., Flores, A., Gräslund, S., Hammarström, M., Johansson, I., Kotenyova, T., Moche, M., Nielsen, T.K., Nordlund, P., Nyman, T., Persson, C., Siponen, M.I., Svensson, L., Thorsell, A.G., Tresaugues, L., Welin, M., Weigelt, J. and Schüler, H.

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PDB Code: 3Q71

ADP-ribose transferases use NAD⁺ as a substrate to add ADP-ribose to substrate proteins. Some of the ADP-ribose transferases add long chains of ADP-ribose (so called poly(ADP-ribose) polymerases- PARPs) whilst other transfer only one molecule. The human family consists of 17 proteins that all contain a catalytic domain in the context of different domains conferring interactions with other proteins and subcellular localization (1 - 2). PARPs are attractive drug targets and there are already drug candidates in advanced stages of clinical trials (3).

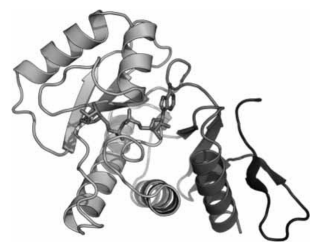


Figure 1. PARP14 macrodomain 2 in complex with its target ADP-ribose

PARP-14 belongs to the sub-class of macro-PARPs, including also PARP-9 and PARP-15. These proteins have not been extensively studied but are thought to function as transcription cofactors (4). PARP14 was recently identified as a co-factor for Stat6 (Signal Transducer and Activator of Transcription 6), as a result PARP14 has the ability to alter interleukin 4 (IL-4) and Stat6 dependent transcription (5 - 6). IL-4 activates Stat6 via Janus kinase phosphorylation and is an important factor in T helper cell immunity and B cell responses.

In addition to a catalytic PARP domain these proteins have up to three macro-domains in sequence at their N-terminal. Macro-domains are small globular domains found in more than 200 bacterial and viral proteins as well as in histone macro H2A-protein in mammals (7 - 8). Macro-domains bind ADP-ribose with high affinity and selectivity (9) and are recruited to sites of PARP1 activation following DNA damage (10).

Here, the crystal structure of the second macrodomain of human PARP-14 is presented in complex with ADP-Ribose at 2.2Å resolution (Figure 1). The structure consists of a seven-stranded mostly parallel β -sheet, except for β 1 and β 7 that are anti-parallel (strand-order β 1- β 2- β 7- β 6- β 3- β 5- β 4), and a total of six α -helices, three on each side of the β -sheet. The overall structure is globular and measures approximately 35Å × 35Å × 50Å. ADP-ribose was clearly seen in the electron density located in a binding cleft mainly made up from two seemingly flexible loops, β 3-loop- α 2 and β 6-loop- α 5.

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STRUCTURAL STUDIES OF HUMAN PROLYL 4-HYDROXYLASES (P4Hs)

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Prolyl 4-hydroxylases (P4Hs) play central roles in two important processes in biology and medicine, the synthesis of collagens and the response of cells to hypoxia [1]. They are 2-oxoglutarate dioxygenases and require Fe^{2+} , 2-oxoglutarate, O_2 and ascorbate. The 2-oxoglutarate is decarboxylated during hydroxylation, with one atom of the O_2 being incorporated into 2-oxoglutarate (producing CO_2 + succinate) and the other into the hydroxy group of the peptidyl 4-hydroxyproline [1].

Collagen prolyl 4-hydroxylases (C-P4Hs), enzymes residing within the lumen of the ER, catalyze the formation of 4-hydroxyproline in collagens and more than 20 other proteins with collagen-like sequences by the hydroxylation of -X-Pro-Gly- triplets. The 4-hydroxyproline residues formed are essential for the formation of collagen molecules with stable triple helices. All known vertebrate C-P4Hs are $\alpha_2\beta_2$ tetramers. The α subunit has the catalytic activity but is highly insoluble. Therefore it needs for solubility the β subunit, which is identical to the enzyme and chaperone protein disulphide isomerase [1].

Detailed knowledge of the properties and structures of the P4Hs will be valuable for the development of inhibitors that may have a major impact on the treatment of fibrotic and ischemic diseases, respectively. Because the X-ray data obtained from large protein complexes can be initially at relatively low-resolution, the high-resolution crystal structures of the individual subunits or domains of such enzymes become crucial during the structure determination. Therefore, while working with the full-length C-P4H, we have also expressed and purified some of its α -subunit domains / domain homologues for structural studies (Figure 1). We have solved the high resolution crystal structures of a peptide-substrate-binding (P-S-B) domain [2], and a homolog to the catalytic domain, namely a P4H from green alga *Chlamydomonas reinhardtii* (referred to as Cr-P4H) [3,4].

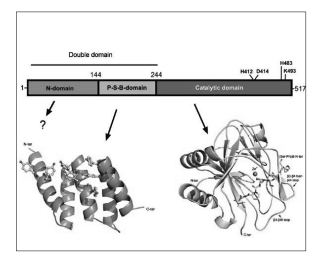


Figure 1.

The α-subunit of human C-P4H with its domain assignments. The active site residues of the α -chain important for the prolyl hydroxylation reaction are also indicated. The crystal structures of the peptide-substrate-binding (P-S-B) domain (PDB code 1TJC) showing the 5 α -helices, which form the tetratricopeptide repeat fold, and the Cr-P4H (PDB code 3GZE), showing the mode of binding the proline-rich (Ser-Pro), peptide to the jelly-roll core structure, are also shown.

The function and the structure of the N-domain region of the α -subunit is still unknown, but we have managed to crystallize a truncated construct consisting the N-domain and the P-S-B domain of the C-P4H α -subunit (referred to as double domain). These crystals have an additionally value, because they have been obtained in the presence of peptide-substrate (Pro-Pro-Gly)₃. Crystals were

tested using the in-house diffractometer at Oulu, and they revealed to belong to the space group $P2_12_12$ with most probable two molecules per asymmetric unit. More data were collected using synchrotron radiation at beamline I911-3 MAX-lab, Lund. We managed to collect a high resolution data set to 1.98 Å, as well as a highly redundant (=24) data set to 3 Å (using 1.5 Å wavelength). By using these two data sets we have initiated a structure solution determination process of the double domain. Our aim is solve the phases using the molecular replacement technique using crystal structure of the P-S-B domain as a model, and/or a single anomalous dispersion method using the weak anomalous signal from the natural sulphur atoms.

Meanwhile, while doing the X-ray crystallographic analysis, we have recently also collected synchrotron-SAXS data from the different C-P4H constructs and complexes. Most of the SAXS data have been collected at I711 [5], MAX-lab, Lund. Soluble double domain concentrated to 8 mg/ml has been one of our samples, but we have also managed to collect SAXS data from the full-length C-P4H-1. The SAXS data has revealed, for example, that the double domain is an elongated molecule with longest dimension of 18 nm (Figure 2). The volume of the map corresponds to a dimeric protein with a molecular weight estimation of 60 kDa.

Using all the structural information obtained using different techniques, we can, for the first time, to construct a complete model for the domain organization of the $\alpha_2\beta_2$ C-P4H complex. The preliminary model of C-P4H, mainly based on the SAXS maps and the high-resolution crystal structures of its domains/subunits, will be very useful for the subsequent crystal determination complete of the assembly. Furthermore, SAXS experiments have indicated that the complete C-P4H tetramer is a rigid and stable molecule. This is the ultimate requirement for the crystallization of the full C-P4H-1, which we hope to happen in near future.

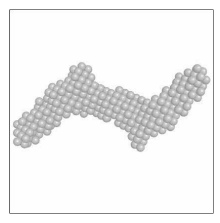


Figure 2. The experimental *Ab inito* model (made by DAMMIN program [6]) of the double domain construct of C-P4H α-subunit.

References

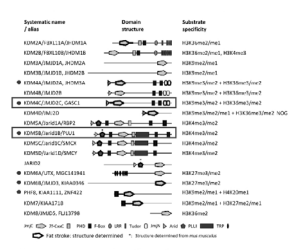
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Histone demethylases as targets for cancer therapy

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An important mechanism in the dynamic regulation of all gene expression is the reversible posttranslational modification of the N-terminal tails of histones (Mosammaparast & Shi, 2010). The modifications include acetylations, phosphorylation, methylations, hydroxylations, ubiquitination, SUMOylation, and biotinylation. There are many possible modifications on each of the four H3 and H4 tails of the approximately 20 million nucleosomes in human chromatin. Considering only methylation of lysines there are five major modification sites: H3K4, H3K9, H3K27, H3K36 and H4K20. These can be



JmjC proteins in the BR-pipeline

unmodified, mono-, di- and trimethylated, giving a total of 1024
different labels. Combination with the
other modifications makes unique
labels for each nucleosome possible.
The total set of such labels of a
genome is often referred to as the
epigenetic code. A given set of
modifications then determine whether
a gene is silenced or actively
transcribed. The state of chromatin is
then maintained by enzymes that can
write and erase the modifications. In

addition binding domains that can recognize (read) modifications are known. They often guide the writers and erasers to the correct genetic loci.

Histone demethylase (HDM) enzymes consequently appear to play a central role in the regulation of known oncogenes (Chi et al., 2010). These enzymes are strong candidate anti-cancer drug targets (Lohse et al., 2011). One family of histone demethylases (HDM) is characterized by containing a catalytic Jumonji C (JmjC) domain with specificity for methylated lysines on histone 3. The first member was discovered in 2006 (Tsukada, 2006) and today the 28 members shown in figure 1 are known.

The structure of the JmjC domain of KDM4A has been determined previously in apo form (Chen et al, 2006) and in complex with various substrates (Ng, et al., 2007). During the last year we have established a pipeline for expression, purification and crystallization of the human JmjC enzymes, and we have established methods to optimize co-crystallization. In summer 2010, we collected the first data set to 2.7 Å resolution at MAX-lab (I911-2) of a complex of KDM4A with an peptide substrate with previously uninvestigated covalent modifications. Although the peptide could be recognized in the electron density we are currently optimizing co-crystallization conditions to achieve more information about the binding mode.

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Structure of the dimeric form of CTP synthase from Sulfolobus solfataricus

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CTP synthase catalyzes the last committed step in *de novo* pyrimidine nucleotide biosynthesis. Active CTP synthase is a tetrameric enzyme composed of a dimer of dimers. In the presence of the substrate nucleotides, ATP and UTP, the tetramer is favoured. The crystal structure of a dimeric form of the CTP synthase from *Sulfolobus solfataricus* enzyme has been determined to 2.5 Å resolution [1]. All previous crystal structures of CTP synthase are fairly similar and showed a homotetramer, independent of whether there were nucleotides bound. The tetramer interacts through the N-terminal synthase domain with the C-terminal amidotransferase domains located far from the tetramer interfaces. The ATP binding site and CTP binding site in the synthase domain are located at the tetramer interface. A comparison of the dimeric interface of *S. s.* CTP synthase with the intermolecular interfaces in the tetrameric structures of *Thermus thermophilus* CTP synthase [2] and *Escherichia coli* CTP synthase [3,4] shows that the dimeric interfaces are almost identical in the three systems and that the residues forming the tetramer interfaces are moving substantially.

Figure 1

a b

The overall fold of the *S.s.* protein is similar to the fold found in the CTP synthases from *E. coli* (pdbentries: 1S1M, 2AD5 and *T. Thermophilus* (pdb-entries 1VCN, 1VCM and 1VCO). The subunit of *E. coli* apo CTP synthase is shown in Fig. 1a and the *S. solfataricus* structure is shown in Fig. 1b. The most

pronounced difference is the change in the loop going from residues 11-16, the loop+helix from residues 180-200 and the helix formed by residues 215-230.

We have superposed the tetramer formed by *E. coli* CTP synthase (pdb-entry 1S1M) with two copies of the *S. solfataricus* CTP synthase model to form a pseudo-tetramer of the *S. solfataricus* structure. The pseudo-tetramer is shown in Fig. 2.

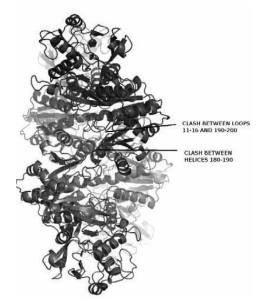
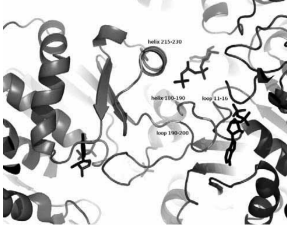


Figure 2

In the *S. solfataricus* pseudo-tetramer the helix from 190-200 clashes with the same helix from the other dimer; the loop from 180-190 also clashes with the loop from 11-16 from the other dimer. It is therefore clear that these regions have to move substantially to facilitate tetramerization.

In Fig. 3 we have superposed the *E. coli* 2AD5 structure with the *S. solfataricus* pseudo-tetramer and zoomed in on the ADP binding site. Only the peptide chain from *S. solfataricus* is shown in the figure. The loop from residues 11-16 would indeed clash with ADP leading to a reposition of this loop to displace the loop and helix that runs from residues 180-200, which in turn would reposition the neighbouring helix spanning residues 215-230.

Figure 3



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Temperature-Responsive Inclusion Complex of Cationic PNIPAAM Diblock Copolymer and γ-Cyclodextrin

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Pseudopolyrotaxanes are supramolecular assemblies formed by a linear molecule (e.g. a polymer chain) and ring shaped molecules such as cyclodextrin (CD) [1]. Pseudopolyrotaxanes are widely studied because the structure of the assembly can readily be tuned by using external stimuli as well as the molecular architecture. A precise control of such supramolecular structures is a challenge for designing smart nanomaterials and for development of applications based on molecular recognition, e.g. drug delivery [2]. CDs are cyclic oligosaccharides formed by glucopyranose units. They have a truncated cone shape with a hollow cavity, which may incorporate more or less hydrophobic solutes such as surfactants or polymer chains to form an inclusion complex. Several CDs rings can thread a polymer chain assuming either a compact or loose structure depending on the nature of the CD.

In this project, we investigate the inclusion complex formation between γ-CD and polymers containing poly(N-isopropylacrylamide) (PNIPAAM) chains in solid state and in aqueous solution. PNIPAAM belongs to the class of temperature-responsive polymers. The characteristic feature of such polymers is that they undergo a physical change when exposed to external temperature-stimuli. PNIPAAM possesses an inverse solubility in water upon heating with a lower critical solution temperature (LCST) between 32-38 °C. Both a cationic diblock copolymer, PNIPAAM-b-poly((3acrylamidopropyl)trimethyl ammonium chloride (PNIPAAM24-b-PAMPTAM(+)18) and a PNIPAAM homopolymer (PNIPAAM₄₇) have been investigated using various experimental methods, such as ¹H-NMR, Fourier transform infrared spectroscopy, synchrotron radiation powder X-ray diffraction (SR-PXD), steady-state fluorescence spectroscopy, differential scanning calorimetry, dynamic and static light scattering and cryo-transmission electron microscopy [3]. The NMR measurements performed on γ-CD-polymer mixtures in D₂O demonstrated that the threaded γ-CD molecules form a close-packed structure over the PNIPAAM chains and that the bulky cationic block of the copolymer is not incorporated into the γ-CD cavity. The fluorescence experiments using pyrene as a fluorescent probe showed that increasing the temperature to 40 °C (above the LCST of PNIPAAM) causes the CD molecules to dethread.

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Solid γ -CD-polymer inclusion complexes form at ambient temperature in fairly concentrated CD solutions. The formation of a precipitate can be considered as an evidence of the formation of crystalline pseudopolyrotaxanes. The SR-PXD spectra of the solid inclusion complexes (Figure 1) indicate a compact columnar structure of CD molecules threaded onto the PNIPAAM chains, which corroborate the NMR results.

The SR-PXD data were collected at beamline l911-5 at MAX II. The X-ray exposure time was 10 s and the wavelength was 0.9077 Å. All raw images were analyzed using the FIT2D program [4]. Calibration measurements were carried out using a LaB₆ sample. Sample-to-detector distances of 50, 100, 200, 300, 380 mm were used to cover a wide range of the scattering vector $q = 4\pi\lambda^{-1}\sin(\theta/2)$, where λ denotes the wavelength and θ the scattering angle.

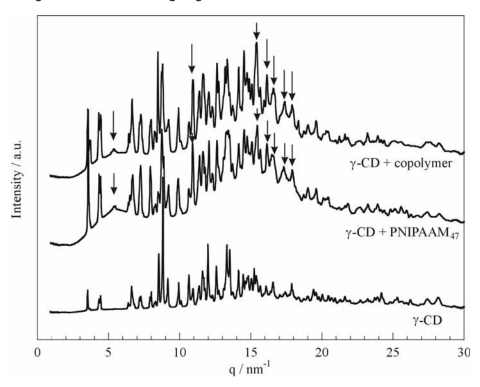


Figure 1. Synchrotron radiation X-ray diffraction spectra of γ -CD and pseudopolyrotaxanes in the solid state. The arrows indicate the peaks that appear in the presence of the polymer and which are related to a columnar structure of γ -CD.

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In situ XRD characterization of carbon-coated Ni_{0.5}TiOPO₄ electrode material

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Extensive research is devoted to understanding and improving existing anode materials for today's lithium ion (Li-ion) batteries. For commercial Li-ion batteries, the most commonly used negative electrode material is graphite. There are, however, many incentives to find new negative electrode materials that can store more lithium or function more safely than graphite. One example is the spinel $\text{Li}_4\text{Ti}_5\text{O}_{12}$ which has become an attractive alternative due to the insertion of lithium above the potential of lithiated graphite, thereby improving safety. The experimental discharge capacity of this material is ~160mAh/g. For the positive electrode, lithium transition metal phosphates are mainly used. The framework of $M_n(\text{PO}_4)_y$ provides an excellent stability and long term cycling compared to lithium transition metal oxides. Indeed, the oxygen—phosphorous bond is more covalent in nature than polar oxygen—metal bonds. Thus, no loss of oxygen occurs from the framework and the reactivity with the electrolyte is low. Also for the negative electrode an enhanced safety could be achieved by using the $M_n(\text{PO}_4)_y$ framework tuned for low potentials vs. Li/Li^+ .

In this study the structure and the lithiation process of negative electrode material $Ni_{0.5}TiOPO_4$ has been studied *in situ* at MAX IV-laboratory beam-line 711. The compound crystallizes in the monoclinic system with the space group $P2_1/c$ as shown in Fig. 1. A carbon-coated sample and a non-carbon coated sample were synthesized using a co-precipitation method.

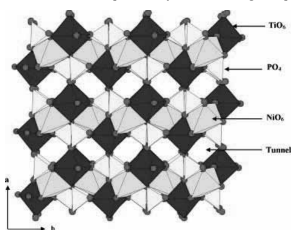
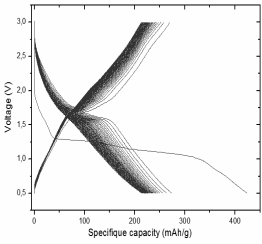


Fig. 1. The crystal structure of Ni_{0.5}TiOPO₄.

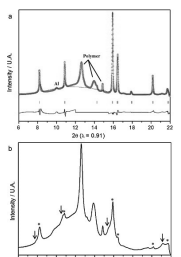
The electrochemical reactions for a carbon-coated sample showed better battery cycling stability than a non-coated sample. Cyclic voltammetry showed that during the lithiation process lithium is expected to first insert at a potential 1.2 V vs. Li/Li⁺ and a second phase at 1.07 V vs. Li/Li⁺. The cycling curve for the carbon-coated sample is shown in Fig. 2.



Ni_{0.5}TiOPO₄:.(bottom)

The cycling stability for the carbon coated material is superior to that of the non-coated. One explanation can be the different morphology obtained with smaller particles and a more homogeneous particle size distribution than for the non-coated.

The *in situ* X-ray diffraction patterns are complex and shown in Fig. 3. The data was collected on beamline I911-5 using a Mar 165 detector. The monochromator $(\lambda = 0.91 \text{ Å})$ was equipped with a bent Si $(2\ 2\ 0)$ crystal.



The diffraction pattern for a pristine cell is shown in Fig. 3 (top) and for a fully discharged cell (bottom). The starting material is highly crystalline. During lithiation a complex pattern involving both solid solution and first order phase changes is observed. However, during the lithiation process the crystallinity is decreasing. A more detailed study of the *in situ* structural response during discharge-charge of this and similar samples are now underway.

Acknowledgements

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Fig 3. In situ XRD of a pristine carbon-coated Ni_{0.5}TiOPO₄ sample (top) and a fully lithiated sample (bottom)

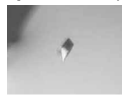
Characterization of a new crystal form of the *B. subtilis* YkuD protein.

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The protein YkuD has been suggested to be involved in redesign of cell walls based on the presence of an N-terminal LysM domain and an L,D-transpeptidase catalytic domain in the C-terminal. The structure has previously been determined to high resolution as part of a structural genomics project. In order to investigate possible substrate binding we have screened for crystallization conditions both in the absence of ligands and presence of mono-, tetra-, and penta-N-acetyl-glucosamine. Crystallization conditions similar to the previously reported conditions for the apoform yielded plate like crystals whereas three dimensional crystals were obtained in the presence of carbohydrate.

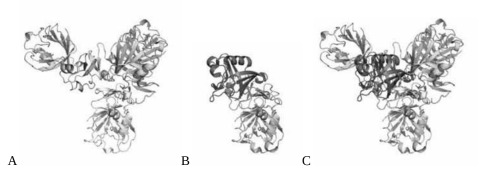




Left: Cluster of thin plate crystals of YkuD formed in absence of carbohydrate. Right: Bipyramidal crystal of YkuD formed in presence of carbohydrate.

The very thin crystals obtained in the absence of carbohydrate were soaked in excess N-acetyl-glucosamine and flash frozen prior to data collection at I911-3. A nearly complete dataset was collected to 3.8 Å resolution. The space group of the crystals is P2₁, a = 58.1 Å, b = 82.7 Å c = 74.3Å, $\gamma = 99.1^{\circ}$ and 4 molecules per asymmetric unit in contrast to the previously reported crystals of YkuD, which belong to space group P2₁2₁2₁, a = 56.3 Å, b = 63.9 Å, c = 93.7 Å with two molecules per asymmetric unit (Bielnicki et al., *Proteins*, **62**, p144-51, 2006).

Phasing of the data was done by Molecular Replacement and revealed a different packing of YkuD molecules. Even when symmetry related molecules were examined the interface observed in the orthorhombic crystal form could not be found in the monoclinic form. Unfortunately no electron density was observed, which could be interpreted as the presence of carbohydrate.



A): the four molecules of YkuD in the P2₁ asymmetric unit. B): The two molecules of YkuD in the P2₁2₁2₁ asymmetric unit (PDB:1Y7M). C): Overlay of molecules in A and B.

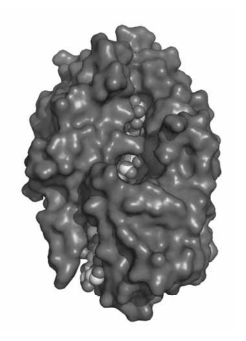
Structure determination of cellulase Cel7A - The major secreted protein of the tree-killing fungus *Heterobasidion annosum* (Rotticka)

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Fungal cellobiohydrolases of glycoside hydrolase family GH7 are key players in cellulose degradation, in nature as well as in industrial applications. Structure-function studies form the basis for protein engineering and discovery of new enzymes with enhanced performance.

In connection with the recent sequencing of the genome of the Root-rot fungus, *Heterobasidion annosum*, the most serious pathogen in our forests in economic terms [1], we have identified major proteins in cultures on spruce wood powder. The most abundant protein was unambiguously identified as a product of the only gene in the genome that codes for a GH7 enzyme. The protein, named Ha_Cel7A, was purified from culture broth and initial crystals were obtained with the JSCG+ Suite sparse matrix screen (Qiagen).



The structure was solved by molecular replacement using *Phanerochaete chrysosporium* Cel7D (1GPI; [2]; 69% sequence identity) as search model. Using X-ray diffraction data from MAX-lab beamlines I911-2 and I911-5, the structure of the apo enzyme (space group $P2_1$; unit cell 60.5, 84.1, 75.9 Å) and that of a ligand complex after soaking with β -thio-xylopentaose (space group C2; unit cell 134.8, 49.5, 73.5 Å; β =118.8°) have been refined at 1.8 and 1.6 Å resolution, respectively (manuscript in preparation).

Ha_Cel7A is clearly a cellobiohydrolase rather than an endoglucanase, and has a cellulose-binding tunnel that is more closed than in *P. chrysosporium* Cel7D and more open than that of *Hypcorea jecorina* Cel7A (eg. 6CEL; [3]) suggesting intermediate properties between these extremes; something that we are now investigating further. The figure shows the surface of Ha_Cel7A, superimposed with a cellulose chain modelled in *H. jecorina* Cel7A (8CEL; [3]).

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Enzymatic Starch Hydrolysis

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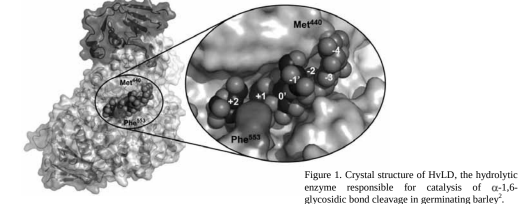
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Starch is by far the most common storage carbohydrate in plants. Globally, starch crops are more important than all other industrial and food crops, and starch crops cover fifty percent of the world's average daily calorie intake. Starch is used by the plants themselves, but also by micro- and higher organisms, which need to hydrolyze the starch to simple sugar molecules by means of enzymatic catalysis to make use of the carbon. Chemically, the major components of starch are amylose to amylopectin. Both molecules are α -1,4-linked glucose polymers, but amylopectin is unique by having α -1,6-linked glucose branches decorating the α -1,4-linked glucose backbone. Different plant species have different ratios of amylose to amylopectin content; most commonly, 20-25% amylose and 75-80% amylopectin. The amylose and amylopectin are packed in starch granules, which vary markedly in size and morphology between plant species.

Of the two starch components, amylopectin presents the largest challenge to hydrolytic enzyme systems due to the α -1,6-glycosidic branch points that are found on 4-6% of the glucose units. Most hydrolytic enzymes are specific for α -1,4-glucosidic links, but to get a complete conversion of amylopectin to glucose also the α -1,6-glucosidic bonds must be cleaved.

Barley limit dextrinase (HvLD) catalyses hydrolysis of the α -1,6-glycosidic linkages of amylopectin derived limit dextrins in germinating barley but the enzyme is also involved in starch biosynthesis. Absence of limit dextrinase or debranching activity can give rise to changed starch grain morphology and even abolished starch grain synthesis. HvLD catalyses hydrolysis via a general acid catalysis. This mechanism requires a catalytic nucleophile and a catalytic acid/base proton donor to interact with the substrate. Hydrolysis results in the retention of the anomeric conformation through a double displacement mechanism characteristic of the glycoside hydrolase 13 family, to which HvLD belong. We previously determined the crystal structure of HvLD (Figure 1)².



Besides the role in germination and starch synthesis, HvLD is critical for the malting and mashing steps in beer production. Here, limit dextrins are hydrolysed to fermentable sugars for ethanol production. In malt, LD is present in a free, active form and in an inactive form, bound to its endogenous proteinaceous inhibitor, HvLDI.

To understand the molecular basis for the substrate specificity of HvLD and its activation, HvLD has been recombinantly expressed in *Pichia pastoris*³ for crystallisation. Crystals of LD in complex with carbohydrate ligands and inhibitory ligands have been obtained and diffraction data have been collected to 2.0 Å and 4 Å, respectively at the I911-2 beam line at MAX-lab II.

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Structural Studies of the Surface Adhesin SpaP from Streptococcus mutans

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Streptococcus mutans is the major causative agent of caries and its surface adhesin SpaP is important for its incorporation into the oral biofilm. SpaP is a member of the Antigen I/II (AgI/II) family of proteins that are expressed on the surfaces of oral streptococci. The AgI/II proteins are key molecules for networking with other microorganisms and host cells in the oral biofilm. AgI/II proteins are approximately 1500 residues and consist of several domains. The structure of AgI/II is unique in the sense that a proline-rich region and an alanine-rich region wrap around each other forming an elongated fibrillar structure, presenting a putative adhesin at the tip (1). The structure further consists of a C-terminal domain which contains an LPXTG motif that anchors the protein to the cell-wall. We have previously solved the crystal structure of the major part of the SspB (AgI/II) C-terminal domain from the commensal bacteria, Streptococcus gordonii (2). It has been shown that the SspB C-terminal domain, but not S. mutans SpaP, functions as an adherence site for the oral pathogen Porphyromonas gingivalis, associated with adult periodontitis. The region that functions as an attachment site for P. gingivalis constitutes a helix that protrudes like a handle. This region is also where the sequence differences comparing SspB and SpaP is the largest. In order to compare the S. mutans SpaP to the previously known structure of SspB from S. gordonii, we determined the crystal structure of the SpaP C-terminal domain (residues 1136-1489). The structure was solved and refined to 2.2 Å resolution with six molecules in the asymmetric unit. Similar to the related AgI/II structure, SpaP is stabilized by isopeptide bonds between lysine and asparagine side chains (3). The structure is overall very similar, as is the secondary structure of the helix corresponding to the P. gingivalis recognition handle. However, the amino acid composition in the helix results in a very different recognition surface. The SspB helix represents a very promising template for the design of substances to exclude *P. gingivalis* from dental plaque.

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Structural Studies of a Fimbrial Sortase from Actinomyces oris.

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The oral bacteria Actinomyces oris, is one of the first colonizers of the oral cavity. As such, it is important in the formation of the oral biofilm. The bacteria is Gram-positive and can express two forms of fimbriae, type-1 and type-2 (1). Type-1 mediates adhesion of the bacteria to salivary proteins that coat the surface of the tooth enamel (2). Type-2 fimbriae contribute to biofilm formation by binding to carbohydrate structures on oral streptococci and host cells (3). The two fimbriae are encoded by two separate operons, both encoding an adhesin, a fimbrial shaft protein, followed by a fimbriae-specific sortase (4, 5). The sortase is responsible for the covalent polymerization of the shaft protein. Sortases are cysteine transpeptidases and can be divided into two classes; housekeeping (SrtA) and pilus sortases (SrtC), both of which catalyze related but distinct reactions. Housekeeping sortases are membrane-associated transpeptidases that recognize and cleave proteins between the threonine and glycine of their peptidoglycan anchor motif LPXTG (6). During the reaction the threonine of the surface protein becomes covalently bound to the nucleophilic cysteine of the sortase after which the protein is transferred to the peptidoglycan of the cell-wall. The polymerization of fimbriae requires specific sortases that catalyze a similar reaction by cleaving the fimbrial shaft protein at its LPXTG motif and linking the threonine to a conserved lysine of another subunit. This results in a stepwise polymerization of the fimbriae. We have solved the structure of the type-1 associated sortase, AcSrtC-1, to 2.4 Å resolution (7). The structure comprise an eight stranded β -barrel with three helices on one side. The active site, containing a His-Cys-Arg catalytic triad, is protected by a flexible lid held in a closed position by the presence of a calcium ion.

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Solving the structure of the A1 protein from bacteriophage QB

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Bacteriophage $Q\beta$ is a small RNA virus that infects *E.coli* bacteria. The virion consists of a single RNA molecule about 4200 nucleotides long and an icosahedral protein shell of T=3 symmetry. In addition to the coat protein, the virion contains a single copy of the maturation (A2) protein and a few copies of the minor coat protein A1, which is a C-terminally prolonged version of the coat protein and is formed when ribosomes occasionally read-through the leaky stop codon of the coat protein¹. Both A1 and A2 proteins are required for infectivity, although their exact functions are not known².

The read-through domain of the A1 protein was crystallized and the crystals diffracted to 1.8 Å resolution. The amino acid sequence of the A1 extension is not similar to any other protein, except some closely related phages, and the three-dimensional structure of none of them was known at the time. Therefore, to solve the structure, it was necessary to collect additional diffraction data from heavy atom derivatives of the crystals. To employ our method of choice, MIRAS (multiple isomorphous replacement with anomalous scattering), crystals were soaked in different iodine, mercury, platinum and gold-containing compounds and their diffraction data at different wavelengths were collected at MAX-Lab beamline 1911-3. The mercury derivatives turned out to be too non-isomorphous to be useful, in platinum and gold-containing soaks there were no heavy atoms bound, and only the iodine derivative was of value for phasing and subsequent solving of the structure.

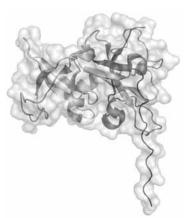


Figure 1. The three-dimensional structure of the A1 domain

The solved structure revealed a rather heavily deformed five-stranded β -barrel at one end of the protein and two anti-parallel sheets at the other (Fig.1). The N-terminal part of the protein contains a prominent polyproline type II helix. Several short α -helices and well ordered loops are also present throughout the protein. The overall fold of the domain is not similar to any published structure in the Protein Data Bank. Although the structure of the A1 domain does not provide immediate answers about its function, our work provides a good starting point for further studies that could eventually lead to the understanding of the molecular mechanism by which the RNA of the phage enters the bacterial cell.

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The new automatic sample changer at 1911-3. The central part is an industrial robot that can move with 10 m/s and 30 μ m precision and automatically mount anyone out of 90 prepared samples. The robot will help the users finding the best crystal, which can be the decisive point in elucidating unknown biological functions by determining molecular structures, 20 October 2010.

Photo: Annika Nyberg

The binding of 1-(β-D-glucopyranosyl) pyrimidine derivatives to Glycogen phosphorylase b

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Type 2 diabetes is characterised by insulin resistance and/or abnormal insulin secretion. Intensive control of blood glucose levels is critical in minimizing the debilitating effects of diabetes and the increasing prevalence of type 2 diabetes has been the driving force for the development of new therapeutic agents. Glycogen phosphorylase (GP) catalyses the degradative phosphorolysis of glycogen to glucose-1-phosphate (Glc-1-P) and offers a potential target for such compounds because of its essential roles in glycogen metabolism and control of liver glucose output^{1, 2}. The catalytic site of the enzyme is a deep cavity 15 Å from the protein surface accessible via a channel blocked by the inhibitor site and the 280s loop (residues 282-287). From T state to R state transition (activation of the enzyme), the 280s loop becomes disordered and displaced, opening this channel allowing the entrance of the substrate and inducing the conformational change of a crucial residue, Arg569, to enter the catalytic site in place of Asp283 in order to create the phosphate recognition site¹. The binding of several compounds to GPb active site has been reported and it is detailed in a number of comprehensive reviews¹⁻³.

Within a collaborative program devoted to the design, synthesis, and evaluation by biochemical and crystallography methods of glucose-based molecules designed as glycogen phosphorylase inhibitors (GPis) we have recently demonstrated that 1-(3-deoxy-3-fluoro- β -D-glucopyranosyl) pyrimidine derivatives are promising GPis⁴. The inhibitory efficiency of seven new 5-substituted pyrimidine glucopyranonucleosides for glycogen phosphorylase b (GPb) was tested in kinetic experiments. All seven compounds were found to be competitive inhibitors of GPb with respect to Glc-1-P with IC₅₀ values ranging from 15 μ M to 6.1 mM. The most potent compound is the one, with an ethinyl group at the 5' position of the pyrimidine ring.

To elucidate the mechanism of inhibition for these compounds, the crystal structures of GPb in complex with each ligand were determined and refined to high resolution. The structures demonstrated that the inhibitors bind preferentially at the catalytic site by anchoring their β -D-glucopyranose moiety at the α -D-glucose binding site. They all promote the less active T-state conformation of the enzyme by making several favourable contacts with residues of the 280s loop. The pyrimidine group is located between residues 284-286 of the 280s loop, Ala383 of the 380s loop, and His341 of the β -pocket. These interactions appear important in stabilizing the inactive quaternary T-state of the enzyme.

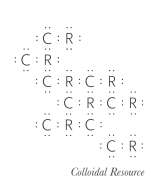
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User report 2011-02-22



Colloidal Resource User report at MAX lab 2010

Colloidal resource AB is a consultant company in the field of chemistry, especially physical chemistry and surface chemistry. In our work we assist industry and start up companies with their chemistry related questions and problems. We use the facilities at Lund University including MAX lab. During 2010 we used the IR facility together with Anders Engdahl (experimental station nr 73) and the X-ray facility (I911-2) with Dörthe Haase.

IR is used both in transmission mode and with the ATR crystal. IR is used in projects to determine absorption on filters and the permeability of membranes. During 2010 medically associated products were studied.

X-ray was used only once during 2010 and this was to study an iron complex for a medical purpose.

Contact person at Colloidal Resource is Dr. Anna Stenstam +46 (0)702 599 755 anna@colloidalresource.se www.colloidalresource.se



Janina Sprenger from department of biochemistry and structural biology, Lund University, and her colleague, are discussing their project during beamtime at the beamline 1911-2, 20 October 2010. Photo: Annika Nyberg

Structural studies of glutathione transferases A2-2 and A3-3

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Glutathione transferases (GSTs) are abundant enzymes, which catalyze conjugation of various hydrophobic, toxic substrates to glutathione (GSH) 1 . The resulting conjugates are generally more soluble and less toxic than substrate. Human genome encodes 17 different GST, divided in 7 classes 2 . In addition to their detoxification function, several alpha class GSTs perform isomerization reaction during steroid biosynthesis 3,4 . Namely, human GST A3-3 catalyzes conversion of Δ^5 -androstene-3,17-dione into Δ^4 -androstene-3,17-dione (AD) (Figure 1).

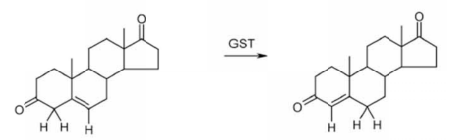


Figure 1. Steroid isomerization reaction catalyzed by human GST A3-3

Human GST A2-2 is 80% identical to A3-3, however, it has neglible steroid isomerase activity⁵, although it binds Δ^5 -androstene-3,17-dione. In order to explain the lack of GST A2-2 activity, we have solved the three-dimensional structures of both GST A2-2 and A3-3 enzymes in complex with AD. In both cases we observed good electron density for AD. In case of A3-3, AD was in a position, suitable for GSH-catalyzed divalent bond rearrangement. In case of A2-2, however, AD was bound in a completely different orientation, explaining the lack of catalytic activity (Figure 2).

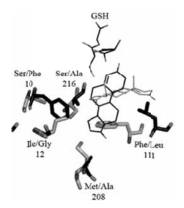


Figure 2. Superimposed active sites of GST A2-2 (green carbon atoms) and GST A3-3 (black carbon atoms) showing five key residues crucial to steroid isomerase activity, as demonstrated by mutagenesis studies6. AD and GSH molecules are rendered as thin lines.

Diffraction data from above structures were collected both at ESRF and MAX-lab synchrotrons. Results are published in Journal of Molecular Biology.

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Acetylcholine binding protein as a model system for nicotinic acetylcholine receptors

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Nicotinic acetylcholine receptors (nAChRs) form ligand-gated ion channels and are directly linked to an ion channel. nAChRs are activated by binding of the neurotransmitter acetylcholine. nAChRs are present in many tissues and are some of the best studied ionotropic receptors. The neuronal receptors are found in the central nervous system and the peripheral nervous system, whereas the neuromuscular receptors are found in the neuromuscular junctions of somatic muscles. The nAChR family comprises 17 different subunits in humans and the assembly of combinations of subunits results in a large number of different receptors.

Therapeutic intervention in receptor signaling has proven beneficial in several neurological and psychiatric disorders. In addition to being targeted by marketed smoking

Figure 1. Structure of a complex of LS-AChBP with an agonist from NeuroSearch A/S

cessation aids, the acetylcholine receptors constitute promising targets for the treatment of a wide range of neurodegenerative and psychiatric disorders, such as Alzheimer's disease, Parkinson's disease, schizophrenia and depression. Therefore, much effort has been and needs to be directed toward understanding the structure, function and basis for subtype selective drug targeting of these receptors.

3D-structural information are available from a low resolution cryoelectron microscopy image of a

torpedo electric ray nAChR (1), a high resolution X-ray structure of the mouse alpha1 nAChR monomer (2), two different species of distantly related bacterial ion channels (3,4) as well as from the acetylcholine binding proteins from the water snails *Lymnaea stagnalis* (5), *Bulinus truncatus* (6) and *Aplysia californica* (7). Although not ion

channels, the snail proteins have the dimension and shape of the *N*-terminal domain of the nAChRs and furthermore, they have been shown to reproduce the relative binding affinities of a number of nAChR ligands. Despite more than 25 AChBP-ligand co-crystal structures deposited in the Protein Data Bank, there is a need for structures addressing *e.g.* the differences between a good binder and a good agonist, subtype selectivity and binding sites for allosteric modulators.

We have established a pipeline for expression, purification and crystallization of the AChBPs from *Lymnaea stagnalis* (LS) and *Applysia californica*, as well as screening assays to guide the selection of ligands for co-crystallization. In 2010, we have collected full data sets of LS-AChBP with four compounds synthesized at NeuroSearch A/S and two in-house synthesized compounds. Of these six data sets, four have been collected at MAX-lab (beamlines I911-2 and I911-3). The resolution of the data sets ranges from 2.3-2.7 Å. All structures have been solved by molecular replacement and contain 2 or 4 pentamers (ca. 2000 or 4000 amino acid residues) within the asymmetric unit of the crystal (figure 1). The structures unravel important details of ligand binding.

We consider LS-AChBP to be the best surrogate of nAChRs for studying binding of agonists and partial agonists, due to a high sequence identity and good correlation between binding affinities when compared to nAChRs.

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Crystal structure of odorant binding protein 4 from *Anopheles gambiae* complexed with *N* -Phenyl-1-naphthylamine

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Malaria is the most common parasitic tropical disease, caused by the parasite *Plasmodium falciparum* which is transmitted to humans by female mosquitoes, particularly *Anopheles gambiae*, during blood feeding. Hydrophobic odor molecules that enter in the sensillum lymph cavity from the outside air space are captured by the odorant-binding proteins (OBPs), present in the olfactory sensilla of mosquito antenna, which facilitate to the odor transportation to the odorant receptors. OBPs may serve as molecular targets for the rational development of better, environmentally-friendly repellents thus preventing mosquitoes from identifying their human hosts and transmitting the malaria parasite in the process.

Only 2 of the 60 odorant binding proteins of *A. gambiae* mosquito [1], AgamOBP1 [2] and AgamOBP4 [Tsitsanou *et al.*, *in preparation*] (Figure 1) have been crystallised until today.

The structure of AgamOBP4 in complex with the fluorescence probe N-Phenyl-1-naphthylamine (1-NPN), determined at 1.6 Å resolution and refined to an R-factor of 0.191 ($R_{free} = 0.229$). Single crystal diffraction data were collected on the MaxLab-ID911-2 station. Data collection statistics are summarized in Table 1.

Table 1. Data collection and refinement statistics

Experiment	AgamOBP4-1-NPN
Space group	P2 ₁ 2 ₁ 2 ₁
Unit cell dimensions (Å, °)	59.95, 60.13, 67.26, 90.0, 90.0, 90.0
Resolution range (Å)	20.24-1.60 (1.69-1.60)
No. of observations	129,721
No. of unique reflections	31,053
<i o(i)=""></i>	18.9 (8.8)
Completeness (%)	95.5 (98.9)
R _{merge}	0.058 (0.14)
Redundancy	4.2 (4.0)
Molecules/asymmetric unit	2





Figure 1. AgamOBP4 crystals

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AgamOBP4 binding site is a hydrophobic cavity located in the center of the molecule and is accessible to the solvent through a narrow opening between a4 and a5 helices. A PEG molecule, present in the crystallisation reagent, occupies the remaining area of the cavity and interacts with 1-NPN. AgamOBP4–1-NPN complex structure analysis revealed the location of the binding site and provided insight into the role of the amino acids that participate in ligand recognition.

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1911-3 Upgrade: A new State-of-the-art Experimental Set-up

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I911-3 beamline¹ is one of five experimental stations at the I911 (or Cassiopeia) beamline. In 2010 the experiment set-up of the I911-3 beamline was completely rebuilt. The beamline was closed during March-May and opened again for users in June 2010 though not all features of the upgrade were yet available for the users. The new set-up together with ongoing software developments includes standards that can be expected of a state-of-the-art MX beamline even in the perspective of MAX IV. At MAX IV the beam qualities will be much higher allowing e.g. to work with micro-crystals but already at present we have taken important steps towards handling smaller crystals and allowing screening of large number of crystals to the benefit of our users but also preparing and learning for the MAX IV beamlines. This is in particular true for the software environment we are developing and implementing, which should be transferable to MAX IV.

The new set-up includes a EMBL/ESRF MD2 micro-diffractometer that reduces the x-ray background significantly and therefore delivers a much better signal to noise ratio for weak diffraction. In addition, the MD2 is equipped with a triple aperture allowing the user to select a 50 um, 100 um or a 200 um beam defining aperture. For small crystals (50 um or below) this combination dramatically improves the signal to noise ratio. The on-line crystal and beam viewing is an additional important tool for working with small crystals.

The MD2 is equipped with a mini-kappa allowing to orient the sample specifically which can be important for e.g. optimizing the



Figure 1: MD2 with single-axis head (left), fluorescence detector (top), cryo (top right) and beam stop (centre).



Figure 2: Sample holder on the mini-kappa (lower left) with sample changer tool (upper right), cryo nozzle (top), on-line viewing and sample illumination (upper left) with the triple aperture in the centre of the lens.

anomalous signal and for optimally completing the data from radiation sensitive crystals.

The new sample changer CATS² is based on an industrial robot. It can select and mount one of 90 pre-frozen samples from EMBL/ESRF standard sample containers stored in a liquid nitrogen dewar. The sample changer can also handle crystallization plates, which allows screening of crystals before any sample manipulation helping in the optimisation of the

Figure 3: The new 1911-3 experiment set-up with the robot of the sample changer in the centre. The area detector can be seen to the left on its granite support.

crystallization conditions and the sample handling.

The HC1³ is a device that can control the humidity around crystal samples. It has been developed at EMBL-Grenoble is now further developed in collaboration with EMBL-Grenoble, Diamond Light Source and MAX-lab⁴. HC1 will be added as a user option at the I911-3 beamline. The HC1 can in some cases improve the diffraction quality of crystals but it is also important in allowing room-temperature tests without difficult sample handling and is in this way complementing the crystallization plate handling of the sample changer robot.

The new experiment set-up also includes additional slits, new intensity detectors, a new fast shutter, a BPM positioned close to the sample and a new Leica microscope. The fluorescence detector has been upgraded while the area detector is the same marmosaic225 as before the upgrade.

Unfortunately, the water-cooled crystal of the double crystal monochromator cracked during the summer shutdown 2010 causing the beamline to be closed during the autumn. Now in Februrary 2011 the beamline is again open for users. We expect the full implementation of the upgrade to be in place later during the spring, including making the sample changer available for users, while the work on improving the beamline, especially concerning software, will continue in the autumn and beyond.

The beamline has obtained funding from the Knut and Alice Wallenberg foundation, the Danish Biotechnology Instrument Center DABIC, the Swedish Research Council VR, the Swedish Foundation for Strategic Research, Danscatt, Bioxhit, IA-SFS and from AstraZeneca and Novo Nordisk A/S. We would like to acknowledge the generous support by the MAX-lab personnel. For further information and updates please see the beamline homepage⁵.

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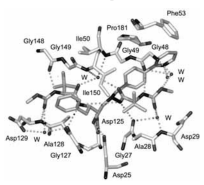
⁵ http://www.maxlab.lu.se/beamlines/bli911

X-ray Structure Studies of potent inhibitors agains the drug resistance due to the mutations V82T and I84V of HIV-1 Protease

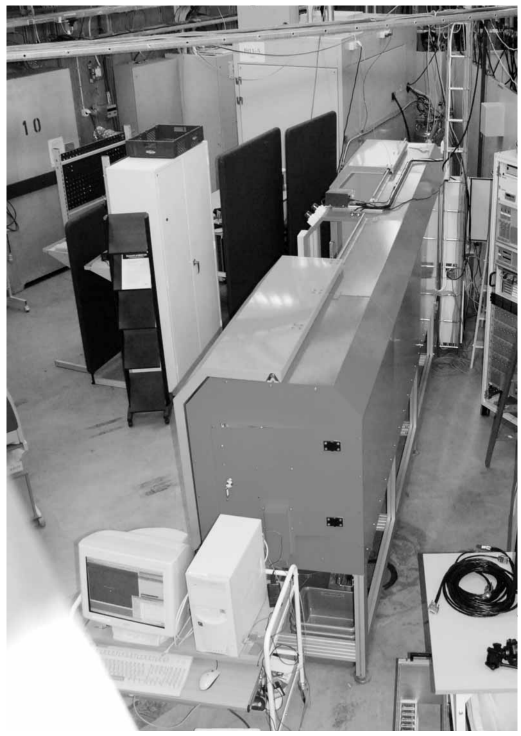
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A new generation of HIV-1 protease inhibitors with potency against the drug resitance mutations V82T and I84V has been developed. This study includes inihibitors base on new synthetic methods and also the effects of different lengths of the central core of the inhibitors. X-ray data were collected at the beam lines 911.1-3. Crystal of HIV-1 protease were produced in complex with a number of inhibitor compounds under optimization development. In this work 3 data sets were collected of the complexes with three different inhibitors. Data were collected at highest resolutions of 1.5 to 1.7 Å resolution. The crystal space group was P21212, which contains the entire protease dimer in the asymmetric unit.



Binding mode of one the inhibitors in the active site of HIV-1 protease. The catalytic residues are Asp25/Asp125.



During the summer months the new hutch of I911-4 SAXS beamline was put into place. The commissioning activities were under development in 2010. 24 June 2010. Photo: Annika Nyberg

Structural fundament for structure-based design of ligands for ionotropic glutamate receptors

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Ionotropic glutamate receptors (iGluRs) are responsible for the major part of the fast excitatory synaptic transmission in the mammalian brain. The iGluRs are divided into 2-amino-3-(3-hydroxy-5-methyl-4-isoxazolyl)propionic acid (AMPA), kainic acid and *N*-methyl-D-aspartic acid (NMDA) receptors based on selective agonist-binding properties and sequence similarity of the receptor subunits. The three classes of iGluRs each consist of a number of subunits: GluA1-4 for AMPA receptors, GluK1-5 for kainate receptors and GluN1, GluN2A-D and GluN3A-B for NMDA receptors. The iGluRs share a similar tetrameric structure where the subunits assemble as a set of two dimers. Each subunit contains a bi-lobed ligand-binding domain attached to the transmembrane regions

forming the ion channel pore.

Since the first crystal structure of the genetically engineered form of the GluA2 ligandbinding domain (LBD) was reported (1), several structures of this protein and other subunits have been determined. The majority of the structures are co-crystals of the ligandbinding domains with agonists, antagonists and positive allosteric modulators. Even though a representative structure has been reported for all classes of iGluRs, structures of several iGluR subunits are still lacking. The crystal structure of the rat full-length homotetrameric GluA2 receptor was recently determined to 3.6 Å resolution in complex with the competitive antagonist ZK 200775 (2). A comparison of the full-length GluA2 structure with the GluA2-LBD structure

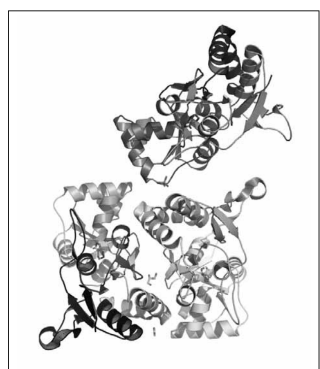


Fig.1 Structure of the ligand binding domain of the AMPA receptor GluA2 in complex with the agonist 7-HPCA (pdb code 3PD8). Data were collected at I911.

in complex with the antagonist UBP282 shows that the soluble LBD is a good model system of the full length-receptor for studying binding of agonists and antagonists binding at the glutamate binding site as well as allosteric modulators binding at the LBD dimer interface as the dimeric unit is very similar.

In 2010, we have successfully worked on this project at MAX-Lab (I911). The studies have focused on the LBDs of GluA2, GluA4, GluK1 and GluK3. We have succeeded to determine four new structures. Structures of the GluA2 LBD have been determined in complex with two allosteric modulators from NeuroSearch A/S and the GluK1 LBD in complex with an in-house antagonist. Also, in 2010 we have established a method for expression and purification of the GluK3 LBD for which no structures have been reported. Data were successfully collected at I911 to 2.3 Å resolution which allowed structure determination. Data have recently been optimized to 1.7 Å resolution elsewhere. Furthermore, the structures of the in-house synthesized agonists 5-HPCA and 7-HPCA (fig. 1) in complex with GluA2 LBD have been published (3).

The iGluR project work is heavily dependent on the use of synchrotron radiation.

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Structural Studies of the G3BP1 and Rin NTF2-like domains.

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Being over-expressed in breast cancer tumours (French *et al.*, 2002), Ras-GTPase activating Protein SH3 domain Binding Protein (G3BP) is considered a significant cancer marker protein. It was identified in a screen for proteins binding to the SH3 domain of Ras-GTPase activating protein (RasGAP) (Parker *et al.*, 1996). Initially, multiple PxxP motifs found in a proline rich region of the protein was thought to be responsible for binding to the RasGAP SH3 domain (Parker *et al.*, 1996), but later the binding site has been located to the N-terminal Nuclear Transport Factor 2 (NTF2)-like domain of G3BP (Kennedy *et al.*, 2002). Besides binding to RasGAP, G3BP is managing multiple tasks: it takes part in c-myc mRNA turnover (Gallouzi *et al.*, 1998), interacts with ubiquitin specific proteases (Soncini *et al.*, 2001), and is involved in NF-κB regulation (Prigent *et al.*, 2000) and stress granule assembly (Tourrier *et al.*, 2003).

The NTF2-like domain of human G3BP1 was sub-cloned, overexpressed in *Escherichia coli*, purified and crystallized. Diffraction data was collected to 3.6 Å resolution using synchrotron radiation (Vognsen *et al.*, 2010). The crystals belonged to the hexagonal space group P6₃22 with unit cell dimensions a=b= 89.84 Å, and c=70.02 Å. Initial phases were obtained by molecular replacement (MR) and by experimental heavy-atom substitution (Figure 1). Preliminary data, to a resolution of 2.8 Å, were also collected from a weakly diffracting crystal of the orthologue Drosophila Rin NTF2-like domain. The structure was solved by MR in space group C2 and with unit cell parameters of a=38.08, b=71.33, c=79.75 Å and β =90.27°.

Recently, a molecular model of the NTF2-like domain of G3BP in complex with the RasGAP SH3 domain has been published (Cui *et al.* 2010). Cui and co-workers based their model on a combination of homology modelling, protein-protein docking, and molecular dynamics simulations. However, obtaining an experimental structure of the NTF2-like domain is important in order to increase the reliability of molecular modelling attempts and to investigate the potential of G3BP as a drug target.

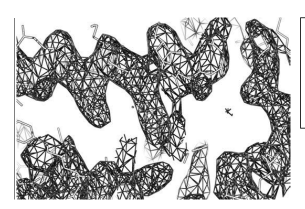


Figure 1. Experimental electron density map for the G3BP1 NTF2-like domain partly derived from phases obtained using data collected at the MaxLab beamline I911-2. Model building and refinement is in progress.

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Synthesis and NEXAFS characterization of Gd₂O₃ nanoparticles

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Approximately 50 % of the Magnetic Resonance Imaging (MRI) exams are today performed with different kinds of contrast agents to enhance the resolution and increase the ability to discriminate between diseased and healthy tissues in the body. Among the most frequently used contrast agents are Gd complexes, in which the Gd ion is bound to a chelating agent, for example DOTA or DTPA. To further increase the sensitivity, contrast agents based on for example nanoparticles are studied. The main advantage of using a nanoparticle based system instead of ion based chelates is that the former is expected to yield a considerably higher relaxivity due to a higher density of magnetic ions^{1,2}.

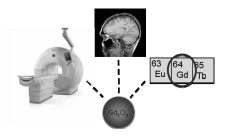


Figure 1. A schematic figure showing the general purpose of the project. The image of the MRI scanner is used with permission from Philips.

The aim of this project is to design new nanoparticle based contrast agents with for MRI signal enhancement The material presented in this work is synthesized at room temperature and the end product after centrifuge washing is highly crystalline and shows an extraordinary solubility and stability in water solution. Characterization of the nanoparticles

were done at beamline D1011, MaxLab, Lund using X-ray Photoelectron Spectroscopy and Near - Edge X-ray Absorption Fine Structure (NEXAFS) spectroscopy.

The C K-edge spectra of Gd acetate and Gd_2O_3 nanoparticles are shown in Figure 2 A and B, respectively. The spectrum for the Gd_2O_3 nanoparticles shows two major peaks originating from acetate- and carbonate groups and several minor features. The very sharp peak at 288.3 eV is assigned to C1s \rightarrow π^* C=0 transitions in the acetate unit in consistence with the main peak in Gd acetate spectrum.

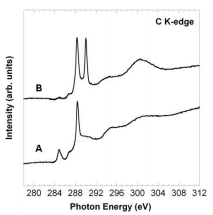


Figure 2 NEXAFS C K-edge spectra of Gd acetate (A) and Gd₂O₃ nanoparticles (B).

Complex binding of the acetate unit to metal ions have earlier shown to alter the NEXAFS spectra of the acetate group^{3,4}. Complexation is thought to induce an intensity decrease in the main peak at 288.3 and at the same time a shoulder appears at slightly lower energy. Our results show a small feature observed as a shoulder

approximately 1 eV below the π^* C=O acetate peak in the C K-edge acetate spectrum Such holder is also observed in the C K-edge spectrum for Gd₂O₃ nanoparticles . In both cases the attributed to acetate shoulder is complex binding of the Gd ion. Gd₂O₃ nanoparticles readily react atmospheric CO₂ to form carbonates on the surface⁵ and the sharp peak at around 290 eV is assigned to C1s → π^* C=O originating from carbonate species⁶⁻⁸. At higher energy states broad structures originating from σ* resonances are observed. associated with C-C bonds ($\sigma *_{C-C}$) are found at 296 eV and states associated with $\sigma *_{\text{C-O}}$ in acetate are found and at $300~\text{eV}^9.$

Carbonate formation is further confirmed in the O K-edge spectrum of the Gd₂O₃ nanoparticles see Figure 3B. The Gd₂O₃ nanoparticle O K-edge spectrum involves 5 peaks, Two of them; peak 1 and 2, are overlapping.

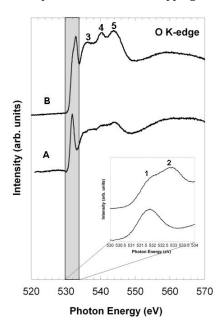


Figure 3 NEXAFS O K-edge spectra of Gd acetate (A) and Gd₂O₃ nanoparticles (B).

The spectrum of Gd-acetate reveals that peak 1 corresponds to the oxygen in the acetate unit. Peak broadening in this area has earlier been observed by Koprinarov et al when studying the stepwise degradation of carbonates to carbonyls⁶. The carbonate π^* _{C=O} transition (peak 2) partly coincides with the acetate π^* C=O as can be observed in the magnified inset in Figure 3. The broad peaks numbered 3, 4 and 5 are assigned to σ * resonances. Splitting of the σ^* system, caused by interaction of two adjacent C=O bonds, together with transitions due to presence of acetate as well as carbonate groups are most likely part of peak 4 and 5 in the spectrum¹⁰.

This project is supported by grants from the Swedish Research Council (VR), The Swedish Governmental Agency for Innovation Systems (VINNOVA) and Carl Tryggers Foundation. The authors are also grateful for all help from the Max-lab staff during the measurements.

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Probing vertical phase separation in the surface region in spincoated films of polymer: PCBM blends – a NEXAFS study

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Polymer photovoltaics is one of the most promising technological solutions to the challenge of low-cost mobile power supply. Solution-processing and bulk-heterojunction (BHJ) architectures turn low-cost, mass-produced solar cells into a not too distant reality. The conversion efficiency of polymer-based photovoltaic devices has now reached 8.3%.

The active layer of such a solar cell is typically prepared from a blend solution of a conjugated polymer and a fullerene derivate, [6,6]-phenyl-C61-butyric acid methyl ester (PCBM). One of the main challenges is to control the morphology in this active layer, which is a result of solvent extraction during film formation. Characterizing the composition in the film is a very important factor to improve the understanding of the device performance. Spontaneous vertical phase separation has been demonstrated for several polymer blend systems through a variety of characterization techniques. The observed self-stratification, with polymer-enrichment of the free surface and PCBM-enrichment of the interface region, can be explained by surface-directed spinodal decomposition. Near-edge x-ray absorption fine structure (NEXAFS) spectroscopy at the C K-edge makes it possible, due to its high chemical specificity, to determine the polymer/fullerene composition ratio at the surface.

Spincoated thin films of (1:4) blends of PCBM with low-bandgap polymers, such as poly[(9,9-dioctylfluorenyl-2,7-diyl)-co-5,5-(4',7'-di-2-thienyl-2',1',3'-benzothiadiazole] (APFO3) and a set of new polyfluorene copolymers (APFO-Green11, APFO-Green12 and APFO-Green13) were studied by near-edge x-ray absorption fine structure (NEXAFS) spectroscopy at beamline D1011 at MAX-lab. APFO-Green11, APFO-Green12 and APFO-Green13 were designed to tune the miscibility with PCBM, and include 0%, 5% and 10% of segments with modified side chains that give the polymer an enhanced ability to form H-bonding with PCBM. Spectra for the pure components were also measured and differences in the π^* - and σ^* - resonance peaks in those spectra served as labels for the analysis of the composition of the blend films. Spectra obtained in partial (PEY) and in total (TEY) electron yield modes, which probe the surface and sub-surface regions of the films respectively, were compared.

The results for APFO3:PCBM show the existence of compositional gradients in the vertical direction in the 1:4 w/w blend. The PEY spectra of the 1:4 blend was fitted with a linear combination of the spectra of the pure components, taking into account the densities of the components. The fitting provides evidence that the surface composition of the 1:4 blend is close to 1:1. The sub-surface composition contains a higher PCBM fraction than the surface. These results are in good agreement with results from our dSIMS studies on this system. Both PEY and TEY spectra for the films of APFO-Green11, APFO-Green12 and APFO-Green13 blended with PCBM (1:4) w/w are shown in Figure 1. The composition ratios at the surface, obtained from the PEY spectra of these blends, are 1:1.9 (34% polymer), 1:1.6 (38% polymer) and 1:1.4 (42 % polymer), respectively. For all cases, this proves a strong polymerenrichment of the surface compared to the 1:4 bulk ratio. Besides, the results show that increased polar character of the polymer, and hence increased miscibility with PCBM, leads to more extended vertical phase separation.

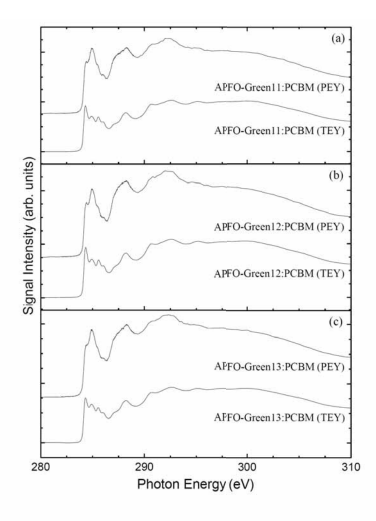


Figure 1. NEXAFS spectra of thin films of APFO-Green: PCBM blends in 1:4 w/w ratios collected in PEY and TEY modes: (a) for APFO-Green11: PCBM (b) for APFO-Green12: PCBM (c) for APFO-Green13: PCBM.

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Materials and interfaces in organic electronics and spintronics

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In the fields of organic electronics and spintronics, there is a need for a multiscale theoretical approach to model organic-organic and hybrid-organic interfaces, ranging from the atomistic to mesoscopic scale (using a combination of quantum-chemical and force-field calculations, solid-state physics and coarse-grained approaches, and classical microelectrostatic models). The EU FP7 project MINOTOR is focused on this issue and we are pursuing experimental characterization of a number of model interfaces to assist in the model development.

In one set of experiments we first prepared clean rutile TiO₂ following the procedure described in literature [1]. We then deposited *in situ* a monolayer of 2,3,5,6-Tetrafluoro-7,7,8,8-tetracyanoquinodimethane, F4-TCNQ, a strong acceptor molecule. Strong bonding interaction with the cyano nitrogens and the TiO₂ surface was observed with resulting charge transfer from the TiO₂ to the F4-TCNQ monolayer causing the formation of an interface dipole. Finally, angle-dependent NEXAFS measurements were performed to determine the molecular orientation at the interface.

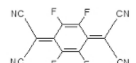


Fig.1 Geometrical structure of F4-TCNQ.

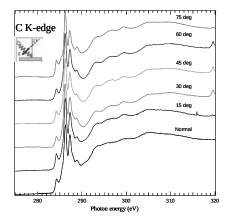


Fig.2 C K-edge NEXAFS of F4-TCNQ/TiO₂

In a separate project, carried out in a collaboration led by M.P. de Jong (University of Twente), we assisted in a study of the electronic and magnetic properties of the interface between C_{60} molecules and a Fe(001) surface. X-ray absorption spectroscopy and x-ray magnetic circular dichroism of C_{60} monolayers on Fe(001) surfaces showed that hybridization between the frontier orbitals of C_{60} and continuum states of Fe leads to a significant magnetic polarization of C_{60} π^* -derived orbitals.

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Nikolay Vinogradov mounting the current feedthrough on the UHV manipulator at beamline D1011, 18 October 2010. Photo: Annika Nyberg

The evolution of the electronic properties for FeHal@SWNT (Hal=I, Br, Cl) during filler decomposition studied by core-level spectroscopies.

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Encapsulated single-walled carbon nanotubes (SWNTs) with inner channels filled by different compounds present the new class of composite materials. Such CNTs are of a big interest because of an opportunity to form 1D nanocrystals as well as quantum nanowires with new physical and chemical properties inside the tubes. Electronic properties of modified CNTs determine substantially the field of their application. Therefore, the main goal of the present project was to perform studies of electronic structure for new materials formed by encapsulation of iron halides FeHal (FeCl, FeBr, FeI) into SWNTs, using x-ray absorption and photoelectron

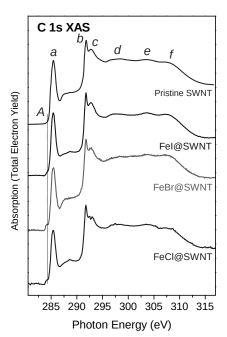


Fig. 1. C 1s absorption spectra of FeHal@SWNT and pristine SWNTs.

spectroscopies, and of possibilities for formation of Fe nanowires during filler decomposition.

All measurements have been performed at beamline D-1011 at MAX II. The FeHal@SWNT nanocomposites were produced by the filling of metallic single-walled carbon nanotubes with inner diameter of 1.1-1.4 nm by FeHal nanocrystals using capillary technique [1]. Approximate loading value of FeHal equals ~70 wt. %.

The C 1s absorption spectra of FeHal₂@ SWNT composites (Fig. 1) demonstrate the existence ofthe interaction between FeHal and nanotube This is evidenced appearance of the low-energy absorption band A (284.1 eV) in the spectra of FeHal@SWNT. The band A is caused by the appearance of a new vacant state, which is probably a result of a substantial Fermi level downshift caused by the doping [2]. Relative intensity of A

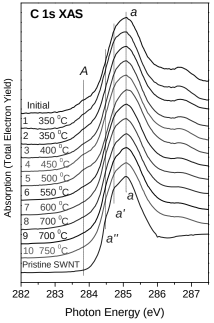


Fig. 2. C 1s absorption spectra near π -resonance for FeBr₂@SWNT during annealing.

formation of metallic Fe. Indeed after the final annealing the C 1s spectrum is practically identical to that of the pristine SWNTs and the Fe 2p spectrum coincides with that of metallic iron. C 1s core-level PE spectra during annealing of FeBr₂@SWNT were measured also. Evidently these findings are caused by the changes of the nanocomposite during the annealing.

The research leading to these results has received funding from the European Community's Seventh Framework Program (FP7/2007-2013) under grant agreement n° 226716 and Russian Foundation for Basic Research (project no. 09-02-01278).

We would like to thank A. Preobrajenski and N. Vinogradov (MAX-lab) for help during experiments.

increases in the row $FeI_2@SWNTs \rightarrow FeBr_2@SWNTs \rightarrow FeCl_2@SWNTs$. This means the increasing of charge transfer upon switching from $FeI_2@SWNT$ to $FeCl_2@SWNT$.

In the course of the annealing of the FeBr₂@SWNT composite the band A in C 1s spectra decreases initially in intensity and then disappears totally. Such a behavior of the band A is indicative of the gradual weakening of the chemical bonding between FeBr₂ and SWNT in the nanocomposite and the separation of the latter into individual phases of FeBr₂ and SWNTs. The practically complete disappearance of the band A occurs after annealing at T ~ 600 °C.

Further vacuum annealing leads to decomposition of the FeBr₂ with the

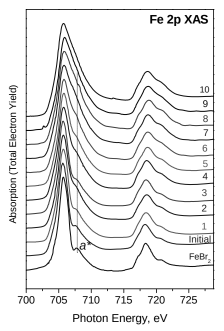


Fig. 3. Fe 2*p* absorption spectra for FeBr₂@SWNT during annealing.

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Adsorption of water and ammonia on graphene/Ni(111)

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Graphene is a single layer of carbon atoms arranged in a honeycomb lattice with two crystallographically equivalent atoms (C1 and C2) in its primitive unit cell [1]. The π and π^* bands of graphene touch in a single point at the Fermi energy (E_F) at the corner of the hexagonal graphene's Brillouin zone, and close to this so-called Dirac point the bands display a linear dispersion and form perfect Dirac cones. Thus, undoped graphene is a semimetal ("zero-gap semiconductor"). The linear dispersion of the bands results in quasiparticles with zero mass, so-called Dirac fermions. The unique "zero-gap" electronic structure of graphene, however, leads to some limitations for application of this material in real electronic devices. In order, for example, to prepare a practical transistor, one has to have a graphene layer where energy band gap is induced via application of electric field or via modification of its electronic structure by means of functionalization. There are several ways of the modification of the electronic structure of graphene with the aim of gap formation. Here we present an attempt to modify the electronic structure of graphene via contact of this material with metal (ferromagnetic Ni substrate) and via adsorption of polar molecules (H2O, NH3) on top of the graphene/metal system. These studies of water and ammonia adsorption on graphene/Ni(111) were

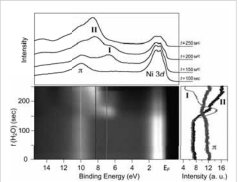


Fig. 1. (Central panel) Photoemission intensity map shows the modification of the valence band of the graphene/Ni(111) system at the Γ point upon adsorption of water molecules (partial water-pressure $p=5\times10^{-8}$ mbar; t is the deposition time). (Upper panel) Photoemission intensity profiles are shown for time-points demonstrating the photoemission features. (Right panel) Photoemission intensity profiles as a function of water deposition time (t) taken at particular binding energies: red solid line, blue solid circles, and green open squares show intensity profiles at 7 eV, 8.3 eV, and 10 eV of the binding energy.

performed via combination of experimental [angle-resolved photoelectron spectroscopy (ARPES), x-ray absorption spectroscopy (XAS)] and theoretical methods [density-functional theory (DFT) calculations]. These studies were performed at the BESSY UE56/2-PGM-1 and UE56/2-PGM-2 beam-lines and MAX-lab D1011 beam-line. Details of experiments and calculations can be found elsewhere [2,3]. In order to study the growth modes of water or ammonia, the time sequences of the photoemission maps around the Γ point of the Brillouin zone (sampling angle of $\pm 10^{\circ}$ with respect to the normal emission) were recorded. The extracted photoemission intensity map showing the modification of the valence band at the Γ point of the graphene/Ni(111) system upon adsorption of water molecules (t is the deposition time) is shown in Fig. 1 (central panel). Photoemission intensity profiles for several time-points demonstrating the main photoemission features of spectra [Ni 3d states, graphene π states, and water induced states (I and II)] as well as intensity profiles as a function of water deposition time (t) taken at particular binding energies (red solid line, blue solid circles, and green open squares show intensity profiles at 7 eV, 8.3 eV, and 10 eV of the binding energy, correspondingly) are shown in the upper and right panels, respectively. The behavior of the water-related photoemission features, I and II, allows to conclude that islands-type growth of water on graphene/Ni(111) takes the place: (i) these features start to grow simultaneously at t = 130 sec, but slopes of the intensities growth are different; (ii) after t = 170

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sec the intensity of feature I decreases via the exponential law and there is a small plateau for the feature II (1st ML is complete); (iii) at t = 230 sec, when the thickness of deposited water is more than 2ML,

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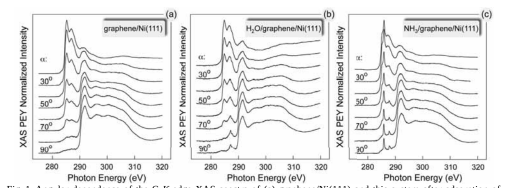


Fig. 1. Angular dependence of the C K-edge XAS spectra of (a) graphene/Ni(111) and this system after adsorption of one-half of the molecular layer of (b) H_2O and (c) NH_3 , respectively. (α =90° corresponds to the normal incidence of the linearly polarized light).

probably, the structural phase transition takes the place – formation of ice. Due to the fact that ice is an insulator, the rapid decrease of the photoemission intensities of the Ni-related features and the shift of some states to higher binding energies can be explained by the formation of an insulating thin film of ice on top of the graphene/Ni(111) system. The general trend in the observation of the ammonia-related photoemission features in the similar experiments is the same. In further XAS and ARPES experiments we chose the thickness of water and ammonia layers to be 1/3-1/2 ML (see discussion about the structure earlier in the text).

The effect of the possible orbital mixing of the valence band states of the graphene layer on Ni(111) and orbitals of water and ammonia molecules was studied by XAS (Fig. 2). This figure shows the angular dependence of the C K-edge XAS spectra of (a) graphene/Ni(111) and this system after adsorption of 1/2 of the molecular layer of (b) H₂O and (c) NH₃, respectively (α =90° corresponds to the normal incidence of the linearly polarized light). These results demonstrate the controllable way of the graphene functionalization by water and ammonia and the corresponding adsorbate-induced states in the region of the unoccupied valence band states were detected (Fig. 2: the photon energies region of 280-290 eV corresponds to the C $1s \rightarrow \sigma^*$ transitions; the photon energies region of 290-320 eV corresponds to the C $1s \rightarrow \sigma^*$ transitions). Here we would like to emphasize that the presented x-ray absorption spectra were recorded at the C K absorption edge and reflect (to some extent) the partial density of states of the carbon atoms in the system and they are clearly demonstrate the appearance of the orbital hybridization of the graphene- and water- and ammonia- related states, respectively. The absence of the strong angular variation of the water- and ammonia-induced XAS signal might be explained by a statistically uniform distribution of the orientations of H₂O and NH₃ molecules on graphene/Ni(111).

The interpretation of the XAS spectra measured after water or ammonia adsorption can be performed on the basis of the peak-assignment, which is discussed on Ref. [2]. For water adsorbate the new structure in the XAS spectra appears at the photon energy range corresponding to the hybrid state in the electronic structure of graphene/Ni(111) involving both carbon atoms in the unit cell of graphene and interface Ni atom. This leads to the assumption that water molecules are adsorbed either in the *center* or in the *on-bond* position on graphene/Ni(111). Ammonia-induced spectral features in the C K XAS spectra are observed in the photon energy range corresponding to the hybrid state which is a result of hybridization of the p_z orbital of the C-top atom and the $3d_{z2}$ state of the Ni interface atom. On the basis of this analysis one can conclude that ammonia molecules are placed in the *on-top* position on graphene/Ni(111) with the lone-pair toward carbon atoms and N-H bonds along C-C bond of the graphene layer. Our DFT analysis of these systems supports these experimental observations.

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Monomolecular films of perfluoroterphenyl-substituted alkanethiols: molecular conformation and odd-even effects

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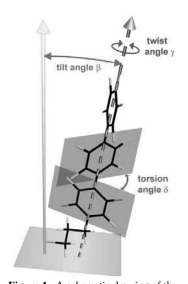


Figure 1. A schematic drawing of the FTP3 molecule. The aromatic backbone has a helical conformation. The orientation of the backbone is given by the tilt angle β , twist angle γ , and torsion angle δ . The torsion results in different values of γ for the inner ring and outer rings.

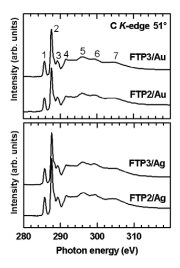
The integrity, chemical identity, packing density, and molecular structure of self-assembled monolayers (SAMs) formed by perfluoroterphenyl-substituted alkanethiols (FTPn) with variable length of the aliphatic linker (n = 2 or 3 methylene units) on Au(111) and Ag(111) were studied by a combination of several complementary spectroscopic and microscopic techniques. The FTPn molecules were found to form well-defined and highly ordered SAMs on these substrates. A characteristic feature of these SAMs is the helical conformation of the FTP moieties (see Figure 1) - in contrast to the planar and close-to-planar conformations of oligophenyl moieties in the analogous non-fluorinated systems. In spite of the helical conformation and larger atomic volume of fluorine as compared to hydrogen, FTPn SAMs exhibit very high packing densities, which are $\approx 3.9 \times 10^{14}$ cm⁻² in FTP3/Au and FTP2/Ag, in particular. Such densities are close to the values for the analogous nonfluorinated systems ($\approx 4.6 \times 10^{14}$ cm⁻²) and can only be possible at a correlated twist of the FTP helices of the neighboring molecules.

The FTPn SAMs exhibit pronounced odd-even effects, i.e. dependence of the molecular orientation and packing density on the length of the aliphatic linker in

the target molecules, with the parity of n being the decisive parameter (as e.g. follows from the NEXAFS spectra in Figures 2 and 3 and from the STM-derived molecular structures in Figures 4 and 5). The relation between the properties of the FTPn films and the parity of n is on Au opposite to that on Ag. FTP3 SAMs on Au and FTP2 SAMs on Ag exhibit a smaller molecular inclination and a higher packing density than those in FTP2/Au and FTP3/Ag - by $\approx 12\%$ if one compares the ordered domains only and by ≈ 15 -30% if one compares overall packing densities. Note that FTP2/Au is characterized by a small portion of the ordered domains due to competitive balance of structure-building interactions in this system.

The odd-even behavior of the FTPn SAMs correlates precisely with the analogous performance of non-fluorinated oligophenylalkanethiolates and -selenolates and, similar to

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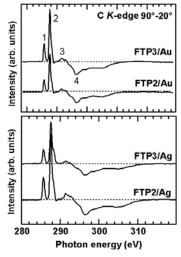


Figure 2. Carbon K-edge NEXAFS spectra of FTPn/Au and FTPn/Ag acquired at an X-ray incident angle of 51°. The most prominent absorption resonances are marked. The most prominent iresonances are marked by numbers.

Figure 3. Difference between the carbon K-edge NEXAFS spectra of FTPn/Au and FTPn/Ag acquired at X-ray incident angles of 90° and 20°. The most prominent difference peaks are marked by numbers. Dashed lines correspond to zero.

the latter systems, can be explained by the presence of a strong substrate—headgroup—C bending potential, which is different on Au and Ag. This quite "rigid" bending potential favors a certain substrate—headgroup—C bonding angle, which is transferred to the substituent moiety by the sufficiently rigid aliphatic linker. A difference of the FTPn system as compared to the non-fluorinated case is the helical conformation of the FTP moiety, characteristic of the molecular state and persistent in the dense phase, what excludes a variation of the intramolecular torsion and molecular twist as the mechanism behind the odd-even effects.

The high quality of the FTPn SAMs makes them promising candidates for use as semiconductor SAMs or for controlling energy barriers between organic semiconductors and metal electrodes. In addition, aromatic SAMs, such as FTPn films, generally should provide an even more efficient charge transport through the molecular backbone than the aliphatic films, which are frequently used for this purpose. This work has been supported by DFG (ZH 63/10-1).

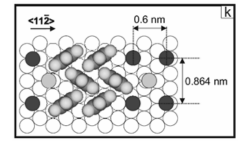


Figure 4. A schematic illustration of the adsorption structure of FTP3 on Au(111) derived on the basis of the STM data. The respective area per molecule is about 26 Å^2 , which corresponds to a packing density of $3.9 \times 10^{14} \text{ cm}^{-2}$.

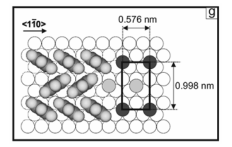


Figure 5. A schematic illustration of the adsorption structure of FTP2 on Au(111) derived on the basis of the STM data. The respective area per molecule of 28.7 Å², which corresponds to a packing density of 3.44×10^{14} cm⁻².

Preliminary XPS results on alkylated silicon nanocrystals produced at two different etching current densities

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Over the past twenty years room temperature optical emission from the nanostructures contained within porous silicon (PSi) layers has attracted considerable interest owing to its potential for use in applications within silicon-based optoelectronic devices [1, 2]. However the source of PSi luminescence remains under debate. Extensive studies upon the silicon nanocrystals (SiNCs) obtained from porous silicon have been geared towards greater understanding of the nature of the efficient photoluminescence (PL) [3, 4] where it is believed that the physical and electronic structure of the PSi surface plays a strong role in determining the photoluminescence wavelength range. PL from porous silicon is observable over wavelengths across the optical spectrum from blue to red [5].

Although it is generally accepted that the visible luminescence originates from the defects in silicon oxide compounds on PSi surface, the band gap enlargement due to quantum confinement effect, or surface defects states, the wavelength range of PL from porous silicon had been shown to be closely determined by the initial etching parameters [6]. In this study, we have studied the effect of etching current density on the photoluminescence of alkylated SiNCs.

Alkylated SiNCs were formed by refluxing porous silicon chips in solutions of undecene at 230 °C for five hours according to the procedure described in [7]. We prepared SiNCs from porous silicon chips with current densities of 400 mA/cm⁻² and 10 mA/cm⁻² with diameters of approximately 2.2 nm. SiNCs from PSi etched at the higher current density display luminescence at \approx 600 nm ('orange') and those at the lower density show luminescence at \approx 380 nm ('blue'). The SiNCs were dispersed in dichloromethane and drop cast in separate films onto the experimental chamber sample holder. Photoemission spectra were collected over the silicon 2p, carbon 1s and oxygen 1s edges at beamline D1011 (MAX II) with an incident photon energy of 150 eV and a spectrometer pass energy of 50 eV.

While the results are still undergoing analysis, preliminary results from the Si2p core level region are shown in Figure 1. Although the samples were in good electrical contact with the sample holder, the thickness of the films meant that the samples suffered from the effects of progressive charge trapping over time during X-ray irradiation. For thick films of SiNCs as in this case, it becomes difficult for electrons to leave the sample and as a result, the spectrum is shifted towards higher binding energies. Thus the data presented are shifted to a lower binding energy by 10 eV in order to bring spectral features to into energy ranges expected for these samples.

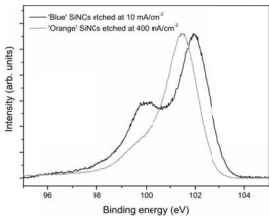


Fig. 1 Photoelectron spectra on 'blue' and 'orange' silicon nanocrystals at 150 eV

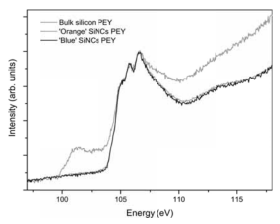


Fig. 2 Near edge absorption spectra (partial electron yield) for 'blue' and 'orange' silicon nanocrystals. Shown with bulk silicon for comparison.

The photoemission spectrum from SiNCs obtained from PSi wafers etched at 400 mA/cm 2 shows a peak at \approx 101.46 eV with evidence of a small shoulder on the leading edge at \approx 99.5 eV. In contrast, the main peak in the spectrum from the 'blue' nanocrystals is at \approx 101.95 eV with a clear second peak at \approx 99.5 eV. This additional peak at the lower binding energy may be attributed to sample oxidation brought about through exposure to the X-ray source.

Near edge adsorption (NEXAFS) profiles in partial electron yield (PEY) mode with a retarding bias of -20 V over the silicon L-edge are shown in Figure 2 for the 'blue' and 'orange' SiNCs alongside the equivalent for bulk silicon for comparison. The absorption features for both samples of SiNCs appear identical to each other with peaks at 105.0 eV, 105.7 eV and 106.5 eV. The latter two features are typical signatures for an oxide-related component. Similar doublet features have been observed close to 105 eV and identified as such within porous silicon surfaces [8].

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In-situ metalation of free-base porphyrins by substrate atoms

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Metalloporphyrins are common in biological systems, with chlorophyll, haem and vitamin B12 as typical examples. These molecules provide essential cofactors in a vast number of proteins and enzymes, while their interesting structural, optical, and electrical properties have led to important applications in fields ranging from biology to sensors to materials chemistry [1]. Since 1960 many papers dealing with metalloporphyrin formation have appeared and several reviews are available [2, 3]. An important step in the formation of these molecules is the insertion of the metal ion into the metal-free porphyrin core. This step has been studied extensively in solution, in biological systems, and more recently on metal surfaces where it has been demonstrated that the metalloporphyrins can be synthesized directly on a surface by metalation of an adsorbed metal free porphyrin with vapour-deposited metal atoms, for example Fe, Co, and Zn [3]. For some metals the metalation reaction proceeds at room temperature (Fe, Co) while others require elevated temperatures (Zn) [3].

Here we report the *in situ* metalation of a free-base porphyrin by metal atoms extracted from the supporting metal surface instead of a vapour deposited metal. We also provide direct evidence for the existence of an intermediate complex formed between surface atoms and the molecule. We apply X-ray absorption (XA) and X-ray photoemission spectroscopy (XPS) to study the mechanism of *in situ* metalation of the free-base 5,10,15,20-tetra(4-bromophenyl)porphyrin (H₂TBrPP) on the Cu(111) surface.

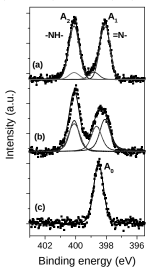


Figure 1. N 1s XPS of (a) 5 ML, (b) 2 ML of H_2TBrPP on the Cu(111) after deposition at room temperature and (c) after annealing at 420 K.

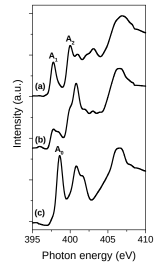


Figure 2. N 1s XA spectra of (a) 5 ML, (b) 2 ML of $\rm H_2TBrPP$ on the Cu(111) after deposition at room temperature and (c) after annealing at 420 K.

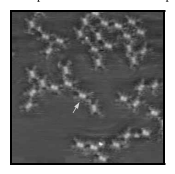
XA and XPS measurements were performed at the beamline at the MAX II storage ring in Lund, Sweden. The XPS spectra were measured with a Scienta SES-200 electron energy analyzer. The kinetic energy resolution was set to 150 meV for the N 1s spectra. The photon energy resolution was set to 100 meV at the N K-edge (~400 eV). The XA spectra were recorded in the partial electron yield mode (U = -100 V) by a multichannel plate detector and normalized to the background curves recorded from the clean substrate. The H₂TBrPP was evaporated onto the clean Cu(111) surface at a rate of about 0.1 monolayer (ML) per minute.

The H₂TBrPP molecule

contains two chemically different types of nitrogen atom, two pyrrolic (-NH-) and two iminic (=N-) nitrogens. Fig. 1a shows the N 1s photoemission spectra from \sim 5 ML of H₂TBrPP on the Cu(111) surface recorded at room temperature. Fitting of the spectrum was carried out using a deconvolution with the minimum number of peaks. The respective signals appear at 400.0 eV

(-NH-) and 397.9 eV (=N-) and the ratio of the peak intensities is 1:1 in agreement with the stoichiometry of the molecule. Fig. 1b shows the N 1s XPS from ~2 ML of H₂TBrPP deposited on clean Cu(111). The iminic peak now appears broader and deconvolution confirms the presence of a third peak at an intermediate binding energy of 398.7 eV. We attribute this third peak to the formation of an intermediate complex in which surface Cu atoms are coordinated with the iminic nitrogen atoms, while the pyrrolic nitrogen atoms are not involved in the bonding with the surface. The fact that not all iminic nitrogen atoms are coordinated to Cu atoms in the surface, as evidenced by the presence of two iminic peaks in the spectrum, is attributed to the existence of second layer molecules which do not interact with the Cu(111) surface. In this intermediate state the planar configuration of the porphyrin is distorted into a saddle conformation with the two iminic nitrogen atoms displaced downwards towards the surface while the two pyrrolic nitrogen atoms are bent away from the surface. This distorted geometry represents an intermediate state of the metalation reaction which cannot be completed at room temperature due to unfavourable energetics. Such intermediate sitting atop (SAT) complexes have been observed during porphyrin metalation in solution [4]. Full copper metalation of the porphyrin is achieved by heating the 2 ML system to 420 K, which also results in the desorption of the second layer. After full metalation the porphyrin has four chemically identical nitrogen atoms which produce a single N 1s peak at 398.5 eV (Fig. 1c).

Evidence for the formation of the intermediate complex is also provided by X-ray absorption. In the N 1s XA spectrum of ~5ML of H₂TBrPP, shown in Fig. 2a, there are two



strong low-energy absorption structures A_1 and A_2 that are related to the π molecular orbitals and arise due to the two different N sites in the centre of the free-base porphyrin macrocycle. The structure A_1 observed at a photon energy of 397.8 eV is produced by the iminic (=N-) nitrogens, while the structure A_2 at 400.0 eV results from the pyrrolic (-NH-) nitrogens [5]. The corresponding XA spectrum from 2 ML of H_2 TBrPP (Fig. 2b) shows a reduction in the iminic nitrogen peak (A_1) relative to the pyrrolic peak (A_2) and the appearance of an additional absorption feature at 398.5 eV. The XA spectrum recorded after heating to 420 K shows an absorption structure (A_0) at 398.6 eV (Fig. 2c), and loss of the features associated with the iminic and pyrrolic nitrogen atoms. The intensities and positions of the observed structures correlate well with the N 1s XPS data presented above.

To extend and strengthen these findings scanning tunnelling microscopy (STM) measurements were performed. Fig. 3 shows an STM image of a sub-monolayer coverage of H_2TBrPP molecules deposited on the Cu(111) surface and reveal that the molecules are stable on the Cu(111) surface. The STM image shows that the molecules are distorted having a rectangular rather than the expected square shape. The distortion of the molecules arises from the formation of Cu-Br bonds between the porphyrin and the surface. This bonding also induces a stress in the Cu(111) resulting in a local (2×2) reconstruction of the surface which produces the protrusions observed on opposite sides of each porphyrin molecule, as indicated by the arrow in Fig. 3. The Cu-Br bonding has the combined effect of distorting both the molecule and the surface resulting in a configuration that favours the formation of the intermediate complex.

This work was supported by Science Foundation Ireland, the Irish Research Council for Science, Engineering and Technology and by the Swedish Research Council.

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Fabrication of a mixed aminopropyltriethoxy and mercaptopropyltrimethoxy silane layer with biofunctional properties on hydroxylated stainless steel

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Biofunctionality of surfaces and surface coatings plays a critical role in biomedical applications involving biomedical implants, biosensors and aerosol based drug delivery techniques. Stainless steel is a low-cost and durable material that is to an extent inherently biocompatible. In this investigation, we fabricated a nanoscale, covalently bound mixed silane layer on stainless steel and bind fluorine labeled maleimide (MAL-F) to it. We employed (3-mercaptopropyl)trimethoxysilane (APS) and (3-aminopropyl)trimethoxy-silane (MPS). MAL-F binds preferentially to the thiol groups. Eventually maleimide terminated biotin molecules will be used to bind avidin proteins to the surface. The activity of the surface can be controlled by varying the concentration of MPS. The majority of the surface is covered by APS which will act as a spacer between MPS sites, thus facilitating unhindered adsorption of large proteins.

In our previous investigations we have found that the deposition of high quality silane films on stainless steel requires a high pre-coverage of hydroxyl species. In this study, we have employed a chemically and mechanically cleaned ASTM 201 (Fe-18Cr-6Mn-4Ni) sample that was subsequently passivated electrochemically as described in Ref. 1. APS and MPS were deposited *ex situ* from EtOH:H₂O solution after 60 s hydrolysis. Several samples were prepared by varying the relative concentration of APS and MPS in the solution. After the deposition the surface was dried of excess solution with dry N₂ and annealed at 373 K for 10 min (Figure 1).

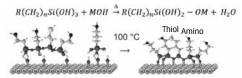


Figure 1: A schematic overview of the formation of the mixed silane layer from hydrolysed solution. Hydroxyl groups covalently bind the silane (gray/light blue) compounds to the surface leaving the amino (blue) and thiol (yellow) termini free. Annealing the surface results in a uniform, high quality silane layer.

For the adsorption of biotin, and subsequently avidin protein, it is essential that the layer is uniform and that the amino and, in particular, the thiol termini of the silane layer are free and perpendicular to the surface plane. It is also important to confirm that the concentration of MPS on the surface correlates with MPS concentration in the solution.

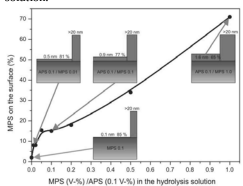


Figure 2: Morphology of the APS/MPS layer after annealing as a function of MPS concentration in the solution.

Figure 2 shows the morphology of selected samples. The morphology was determined from the inelastic electron background by employing the theory developed by Tougaard

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¹ P. Jussila, H. Ali-Löytty, M. Hirsimäki, and M. Valden, Surf. Interf. Anal. 42 (2010) 157-164.

et. al.² At all MPS concentrations, approximately 80% of the surface area is covered by a thin, uniform APS/MPS layer.

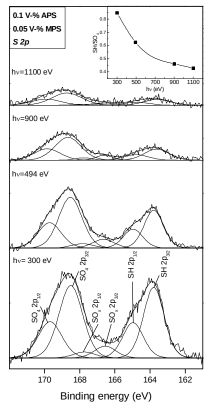


Figure 3: S 2p signal measured after silane deposition from a 0.1 V-% APS / 0.05 V-% MPS solution as a function of photon energy. The inset shows the ratio of SH and SO₄ peaks.

The preferential orientation of the thiol groups (SH) was deducted from the dependence of S 2p photoelectron spectra on the photon energy. Figure 3 shows high resolution photoelectron spectra of S 2p transition measured at photon energies of 300 eV, 494 eV, 900 eV and 1100 eV. As can be seen, the signal originating from the thiol groups is significantly enhanced in comparison to the sulfate signal as the photon energy is decreased and the surface sensitivity

is increased. This indicates that the thiol groups are, indeed, pointing away from the surface.

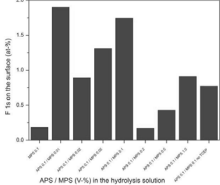


Figure 4: MAL-F uptake as a function of APS and MPS concentrations in the solution. TCEP treatment significantly enhanced the MAL-F uptake.

The uptake of MAL-F as a function of the APS/MPS concentration ratio in the solution is shown in Figure 4. Surprisingly, the MAL-F uptake was initially relatively low. One of our major findings was that in a mixed APS/MPS layer, adjacent sulphur tends to form disulphide bonds that deactivate the thiol groups. Treating the surface with 0.05 M Tris(2-carboxyethyl)-phosphinehydrochoride (TCEP) solution breaks the disulphide bonds, leading to significantly enhanced MAL-F uptake. The results indicate that the optimal silane layer is obtained with a solution of 0.1 vol-% APS and 0.01-0.1 vol-% MPS.

We have shown that a mixed silane layer can be covalently bound to stainless steel in a way that results in a relatively uniform layer. Furthermore, we find that the silane layer can be functionalized with maleimide. In the future investigations, MAL-F will be replaced by maleimide terminated biotin to facilitate experiments on avidin protein adsorption kinetics.

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Thin ferromagnetic Mn-germanide film on Ge(111)

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New materials that possess both magnetic and semiconducting properties are of great interest today. Magnetic semiconductors have a great potential to provide a new type of conduction control. A magnetic semiconductor would not only control the charge carriers as in traditional electronics but also the spin state of the electron. The possibility to use the spin degree of freedom in the standard charge based electronics could lead to a tremendous improvement in memory handling and data storage [1].

Magnetic materials based on Mn and Ge have received a lot of attention lately. There *are* two important facts that make MnGe alloy interesting. First, thick Mn germanide films show magnetic properties at a temperature close to 300 K. Secondly, germanium based alloys should be easy to integrate into the modern semiconductor technology.

In this work a thin film of manganese germanide with a $\sqrt{3}\times\sqrt{3}$ surface reconstruction on

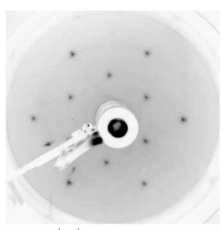


Figure 1. $\sqrt{3} \times \sqrt{3}$ LEED diffraction spots from the thin Mn₅Ge₃ film.

Ge(111)c2x8 has been investigated by means of x-ray magnetic circular dichroism (XMCD). The Ge(111) sample was outgassed in situ by direct current heating at 500 °C for several hours. This was followed by repeated sputtering (Ar+, 1kV) and annealing cycles, 5 min at 600 °C. This resulted procedure in well ordered Ge(111)c2x8 surface as seen by LEED. Sixteen monolayers (ML) of manganese were evaporated from an electron beam evaporator (Omicron/Focus). The annealing of the sample at 200 °C for 15 minutes resulted in intense $\sqrt{3} \times \sqrt{3}$ LEED diffraction spots as seen in figure 1.

The in-plane XMCD spectra were obtained by recording two XAS (x-ray absorption

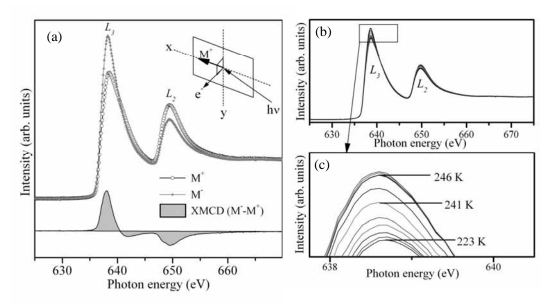


Figure 1. (a) XMCD spectrum measured at the L_2 and L_3 absorption edges of the MnGe(111) film at 100 K. (b) XAS spectra recorded at different sample temperature. (c) Close up image of the L_3 absorption edge.

spectroscopy) spectra at liquid nitrogen temperature. First the sample was magnetized with an magnetic field pulse of 450 Gauss along the horizontal plane. The L-edge XAS spectrum was measuring with 70% circularly polarized light at an incident angle of 75° in the positive x-direction (M+) (see inset in figure 2(a)). Then an magnetic field pulse was applied in the opposite direction (M-) to the sample and the absorption spectrum was measured with the same geometry. The intensity of the XMCD signal of the L_2 and L_3 absorption edges is shown as the difference between the XAS spectra recorded with opposite magnetization (M+ and M-), as seen in figure 2(a). The Curie temperatur (T_C) was found by following the L_3 absorption edge in the XAS spectra during sample warm up, figure 2(b) and 2(c). T_C was estimated to 246 K which is 50 K lower than that found in thicker Mn germanide films. To conclude this part, the Mn/Ge(111) film shows a clear ferromagnetic property in contrast to the case of Mn/Si(111).

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Femtosecond dynamics of the charge transfer in alkanethiolate self-assembled monolayers via resonant Auger electron spectroscopy

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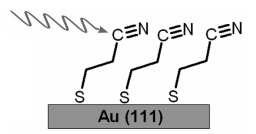


Figure 1. Resonant excitation of the terminal nitrile group by X-ray. The CT pathway is unambiguously defined by the molecular architecture.

We have applied resonant Auger electron spectroscopy (RAES) within the framework of the core hole clock (CHC) method to study dynamics of charge transfer (CT) processes in alkanethiolate (AT) self-assembled monolayers (SAMs) on gold. For this purpose, we prepared a series of high-quality SAMs of nitrile-substituted ATs with variable length of the alkyl chain, NC-(CH₂)_n-SH (CnCN). Due to such a molecular architecture, the CT pathway was unambiguously defined – from the nitrile

tailgroup, which could be resonantly excited by narrow bandwidth synchrotron radiation at the N K-edge, to the substrate over the alkyl chain and the thiolate-gold anchor (see Figure 1).

After the primary resonant excitation of the nitrile group, which was the $[N1s]\pi^*$ in our case, different scenarios for the excited state decay are possible (see Figure 2), including the CT to the substrate within the lifetime of the excited state. These scenarios result in the different final states and can be associated with different spectral features. Significantly, the final state associated with the CT to the substrate (CT) is identical to the final state of a non-resonant Auger process (A). The overall decay spectra can then be decomposed into the contributions related to the P/SP and CT scenarios and the CT time (τ_{CT}) be obtained from the relation $\tau_{CT} = \tau_{core} (I-P_{CT})/P_{CT}$, where τ_{core} is the known lifetime of inner shell vacancy and P_{CT} is the integrated post-CT portion of the decay spectrum.

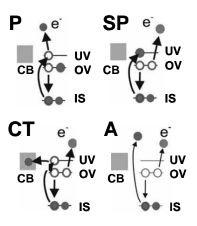


Figure 2. A scheme of core excitation / de-excitation routes in a functional group weakly coupled to a continuum (substrate conduction band - CB), with IS, OV and UV denoting inner shell, occupied and unoccupied valence levels, respectively. Following the excitation of a core electron (curved arrow) into a bound state, either the excited or another, OV electron can be emitted, described as participator (P) and spectator (SP) decay, respectively. Both these processes result in characteristic final states with one hole in OV. Alternatively, charge transfer (CT) of the excited electron to CB can occur. This route leads to the same final state with two holes in OV as non-resonant Auger process (A). The letter process involves the excitation of a core electron into a continuum state (curved arrow) followed by the emission of UV electron. The hole in IS is in every case filled by the electron transition from

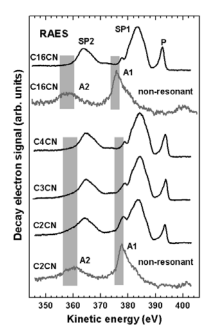
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Within this framework, we measured the RAES spectra of the target SAMs (see Figure 3) and decomposed them into the contribution related to the P/SP and CT scenarios. On the basis of the relative weights of theses two processes, we determined the characteristic CT time to the substrate (see Figure 4). It was found that, similar to the static conductance, the dependence of the CT time on the number of methylene units in the alkyl chain can be coarsely described by exponential function with an attenuation factor of 0.93 per the unit, which corresponds to 0.72 Å⁻¹ with respect to the length. As a result, the CT time is quite long even for a relatively short alkyl chain; it particular, it is ca. 100 fs for the chain consisting of only four CH₂ units. In contrast, the CT time associated with the thiolate headgroup anchor was found to be quite short, viz. 2.3 fs, which suggests that thiolate-gold anchor provides an efficient interfacial electronic coupling between the aliphatic backbone and the substrate.

The advantages of CHC method in its specific application to SAMs are, except for the direct access to the dynamic characteristics of CT, a clear knowledge of the number of molecules participating in CT and the lack of contact problem. Nitrile, as used here, is seen to be a good candidate for CHC studies because of a suitable core hole lifetime and the large contrast between resonant and non-resonant core decay spectra, in particular the large spectator shift.

This work has been supported by DFG (ZH 63/9-3 and ZH 63/14-1).



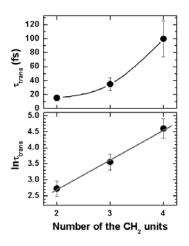


Figure 3. [N1s]π* RAES spectra of C2CN/Au, C3CN/Au, C4CN/Au, and C16CN/Au (black curves), along with nonresonant AES spectra of C2CN/Au (representative of C3CN/Au and C4CN/Au as well) and C16CN/Au (gray curves). P, SP1, and SP2 denote the participator and two spectator features, respectively. A1 and A2 denote two characteristic features in the non-resonant spectra. The kinetic energy ranges where a contribution from these features (CT trace) in the RAES spectra can be expected are highlighted by light gray.

Figure 4. CT time (top panel) from the excited nitrile group to the substrate and its natural logarithm (bottom panel) versus the number (n) of the CH_2 units in the aliphatic chain of CnCN/Au (black filled circles). The gray solid line in the bottom panel represents the best fit. The slope and intercept points corresponding to this fit are 0.93 (0.72 Å⁻¹; a tunneling along the chain was assumed) and 0.83 respectively.

Pulsed laser deposition of gold nitride: Photoemission spectroscopic study of its electronic structure

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Gold nitride is of great interest as a material with properties which potentially offer practical applications. Incorporation of gold nitride into gold films makes them 50% harder without losing it's metallicity [1,4]. Gold nitride containing films promised to be a better material for electrical contacts in electronic circuits and for gold coatings. The first clear observation of the formation of solid gold nitride was made with nitride formation detected upon nitrogen ion implantation of the Au (11 0) surface using ions with an incident ion energy of 500 eV [2, 3]. More recently, techniques based on nitrogen reactive ion sputtering (RIS) and low energy plasma processing have been developed [5]. Both these methods are based on the interaction of low energy nitrogen ions with gold. The RIS technique allows the production gold films containing gold nitride in the bulk. Despite these simplicities there is a remarkable lack of information on the electronic properties of these materials, which are significantly dependant on the preparation conditions of these gold nitride films.

Thin films of gold and gold nitride were produced by pulsed laser deposition (PLD) in an ultra high vacuum chamber. Films were deposited on Silicon, ITO and Sapphire substrates maintained at 450 °C during deposition in a nitrogen atmosphere with a varying pressure between 10⁻⁷ Torr to 10⁻¹ Torr. A Nd:YAG laser 266nm, 10 Hz repetition rate, and fluence of 1.8 Jcm-2 was used to ablate the metal target. All the films are approximately 120 nm thick. In seeking to verify the chemical bond formation and electronic structure of this material in detail, we used soft x-ray absorption and high resolution photoemission spectroscopy to understand the element specific conduction band density of states and to study the valence and core level photoemission. We report here the first measurements of this group on the gold nitride preformed at D1011 beamline. Max-lab.

Shown in Figure 1 is the comparison of gold and gold nitride thin films grown on silicon substrates. The Au 4f spectrum shows a marked difference in structure. The solid line represents the gold thin film of Au 4f_{7/2} and Au 4f _{5/2} spin orbit splitting represented as (A) and (C) respectively. In the case of gold nitride films there is a new feature (B) around 85.8 and 88.7 eV have been observed previously but not resolved [4] started to appear. This new feature is the signature of chemically bonded nitrogen on gold. However, this feature

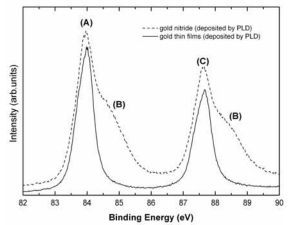


Figure 1 comparison of Au4f photoemission of gold (solid line) and gold nitride (dotted lines) thin films deposited on silicon substrates

B, varies in intensity with varying growth

conditions (not presented here)

In Figure 2 N1s spectra obtained from gold nitride thin films taken at 500 eV excitation energy. The intense peak at 398.5 ± 0.1 eV corresponds to the nitrogen chemically bonded to gold. This is in agreement with our previous results [2,3]. The lineshape shows inhomogeneous broadening and there may be a peak at 397.5 eV.

The intensity of N1s lines varies with varying preparatory conditions. Hence the formation of gold nitride is very sensitive to the growth conditions

In Figure 3 we report the changes in the valence band features of the gold nitride thin films with respect to gold thin films. It can be clearly seen that there is a tail appearing between 0.1 to 2.8 eV and considerable broadening features of d bands. This could be attributed to the incorporation of defects or vacancies induced as a function of nitrogen incorporation in gold. More studies have to be done in order to understand the nature of defects.

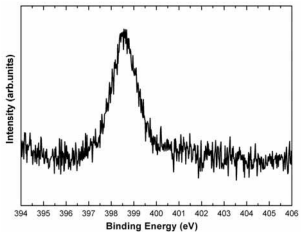


Figure 2 show the N1s photoemission from the gold nitride thin film

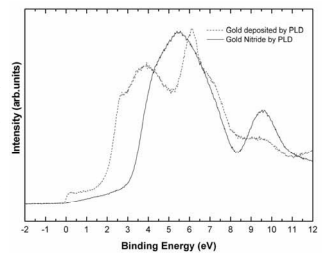


Figure 3 show the N1s photoemission comparison of gold and the gold nitride thin films on silicon substrates

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The structure and magnetic properties of Fe nanoclusters grown on the Ge(001) surface

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Semiconductor surfaces such as the Ge(001)- (2×1) reconstructed surface are suitable templates for the growth of well-ordered, uniformly-sized metal nanoclusters [1]. The size, shape, and spacing between clusters are dictated by the substrate. Such metal-nanocluster-semiconductor systems are considered exposed dilute magnetic semiconductors (DMS) [2]. DMS are promising materials for use in many technological applications and their study is important for future developments in nanotechnology, as well as from the fundamental point of view. Such systems have attracted much attention recently since they can be utilized as essential building blocks in the field of spin-dependent electronic (spintronic) devices, providing a link between magnetism and semiconductor technologies [3, 4].

Here we apply X-ray magnetic circular dichroism (XMCD) and scanning tunnelling microscopy (STM) to study the nucleation, structure and magnetic properties of Fe nanoclusters on the Ge(001) surface.

XMCD measurements were performed at the D1011 beamline at MAX II storage ring in Lund, Sweden. Measurements were conducted at room temperature as well as at 150 K. Fe 2p X-ray absorption (XA) spectra were recorded using sample drain current and normalised to the background curves measured from a clean substrate. The photon energy resolution was set to 200 meV at the Fe L_3 -edge (\sim 710 eV). For XMCD measurements a switchable magnetic field of 0.05 T was applied. Fe was evaporated onto the clean Ge(001) surface at a rate of about 0.1 monolayer (ML) per minute.

When deposited onto the Ge(001) surface at room temperature, the Fe atoms self-assemble into two-dimensional nanoclusters of uniform size. A typical low temperature STM image of 0.2 ML of Fe on the Ge(001) surface is presented in Fig. 1a, where bright protrusions correspond to Fe nanoclusters. All nanoclusters have a square shape and a

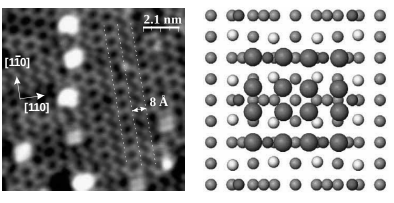


Fig. 1. Low-temperature STM image of 0.2 ML of Fe on the Ge(001) surface: $V_b = -1.5 \ V$, $I_t = 0.50 \ nA$, size 11.0 nm \times 11.0 nm, 78 K (a). The calculated model for the Fe nanocluster, which consists of 16 Fe atoms, on the Ge(001) surface (b). The Fe atoms are denoted by large spheres, the Ge atoms forming the surface layer by red, orange and yellow spheres, while the bulk Ge atoms by green spheres.

uniform size of $9.0 \text{ Å} \pm 0.5 \text{ Å},$ corresponding to sixteen Fe atoms per cluster. These nanoclusters are single monolayer height, having an apparent corrugation height of 1.1 Å \pm 0.1 Å with respect to the underlying substrate, which does not depend

on the bias voltage. These two-dimensional nanoclusters follow the [1–10] direction of the substrate dimer rows, although the separation between the clusters varies throughout the image.

In order to confirm the number of Fe atoms in the nanoclusters formed on the Ge(001) surface, DFT calculations were performed using the Vienna *Ab-initio* Simulation Package. Fe clusters with different numbers of atoms were simulated. The cluster with sixteen atoms was found to have the lowest energy and a size of $9.0~\text{Å} \times 9.0~\text{Å}$, which is in excellent agreement with the experimental data. The resulting model of the Fe nanocluster on the Ge(001) surface is shown in Fig. 1b.

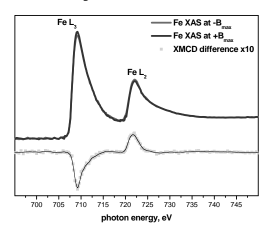


Fig. 2. Fe 2p XA spectra measured from 0.5 ML of Fe on the Ge(001) surface at 150 K with the magnetic field of 0.05 T applied in opposite directions ($-B_{max}$ and $+B_{max}$).

To study the magnetic properties of Fe nanoclusters, **XMCD** the measurements were carried out. The Xray absorption (XA) spectra, measured at 150 K from the 0.5 ML of Fe on the Ge(001) surface with the magnetic field of 0.05 T applied in two opposite directions are shown in Fig. 2. The XA spectra have a metallic shape, indicating that no significant intermixing occurs at the Fe/Ge(001) interface, which is in agreement with previous studies [1, 5]. The XMCD spectrum shown in Fig. 2 is a result of subtraction of the XA spectrum taken at the maximum magnetic field (+B_{max}) applied in one direction (blue curve) from the other spectrum taken at the maximum magnetic field (-B_{max}) applied in the opposite direction (red curve). The XMCD spectrum taken at

150 K shows a prominent difference between the two opposite magnetic directions. In turn, XMCD measurements taken from the same Fe/Ge(001) system at room temperature show almost no difference between the spectra obtained for a magnetic field applied in opposite directions. This indicates a superparamagnetic behaviour of the Fe nanoclusters (consisting of sixteen Fe atoms) grown on the Ge(001) surface. Free Fe clusters of similar size were also shown to exhibit superparamagnetism [6].

Thus, by applying XMCD and STM to Fe nanoclusters grown on the Ge(001) surface, it was found that the nanoclusters consist of sixteen Fe atoms and show superparamagnetic behaviour.

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Adsorbed graphene as a non-volatile platform for chemical reactions

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A non-volatile thin film platform is the fundamental requirement for chemical sensing. Conversely, the system must be chemically, physically and thermally stable. In addition, chemical reactions taking place on the sensors must result in detectable property changes and these reactions have to be reversible. Calculations [1] have shown that the band structure (gap size) of carbon nanotubes (an allotrope of graphene) changes uniquely/specifically upon interaction with chemical functional groups (-CH, -C-O, -C-OH and -C=O). Experimental studies [2] have proven that there is a profound conductance change when graphene oxide is reduced.

Previously, we have demonstrated that monolayer graphite (MG) adsorbed on transition metal (TM) such as Ir and Pt is more susceptible for chemical reaction with atomic H and the adsorption process is reversible upon soft annealing, i.e. the perfect MG surface can be recovered. [3] However, not all chemical processes on such systems are perfectly reversible by thermal treatment due to the strong chemical bonds, e.g. -C-O-C- that can rip off and cause defects in the MG layer. [2,4] Here, our preliminary x-ray spectroscopic studies indicate that MG adsorbed on Ir can be reduced and oxidized reversibly and repeatedly by atomic H and O respectively at room temperature (RT).

All samples were prepared and characterized *in-situ* at Beamline D1011. The MG/Ir(111) sample was prepared and hydrogenated to saturation according to the description in Ref. [3]. This system was oxygenated stepwise with atomic oxygen and then with atomic hydrogen, whereby the exposure time is as described in Fig. 1.

Fig. 1 shows the C 1s and O 1s PE spectra recorded from the clean and various treated MG/Ir. The evolution of chemical states of the overlayer due to the stepwise chemical treatments occurs in the following sequence.

- A single component C 1s represents the pristine C-C bond in graphene (Fig. 1[left,a]).
- Triple components C 1s (Fig. 1[left,b]) where two components are due to the H adsorption on pores and wires of the slightly corrugated MG/Ir surface while the third is the chemically intact graphene. These results are published in Ref. [3].
- After 1 min (Fig. 1[left,c]) of exposure to atomic O, O atoms adsorb on the H adsorption sites while H atoms desorb from the surface. This plausible inteterpretation is based on the increasing intensity of the epoxy-related components and the decreasing of the H adsorption related components in relation to the O exposure (Fig. 1[left,d]). The epoxy (-C-O-C-) related features are also observed in the O core level (Fig. 1[right,a-b]).
- After the second hydrogenation treatment, the hydrogen atoms first form hydroxyl groups (-OH) with adsorbed O, as observed in Fig. 1[left,e] and subsequently form water molecules and desorb away when more H atoms arrive on the surface (Fig. 1[left,f]). The system gradually recovers back to the state of the hydrogenated one (Fig. 1[left,g]. These statements are in agreement with the O 1s spectra (Fig. 1[right,c-e]), where the hydroxyl component appears and the intensity of the epoxy and

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carbonyl components falls upon hydrogenation. This observation is also confirmed by the results in Ref. [5,6].

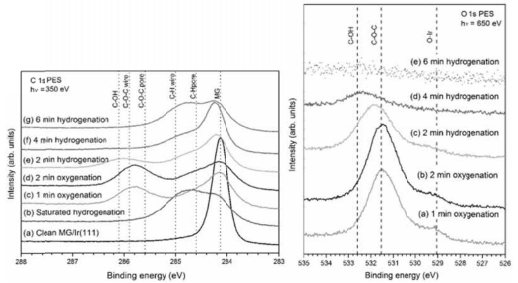


Figure 1: C 1s [left] and O 1s [right] PE spectra of the stepwise hydrogenation and oxygenation treatments of MG/Ir(111).

It is plausible to state that the graphene layer is stable throughout all chemical treatments because the total intensity of C 1s relative to Ir 4d remains almost the same after each treatment (spectra not shown here).

It can be concluded that adsorbed graphene is indeed a non-volatile platform for chemical reactions at RT. The system also exhibits signature properties (core level chemical shifts) that are unique to the adsorption of each functional group. By adsorbing O on hydrogenated graphene, the O adsorption process can be reversed by simply exposing the system to H at RT.

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Electronic Structure of Graphene Grown on Copper

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The interaction between graphene and a transition metal substrate varies across the periodic system in that the a strong bond is evident for early transition metals (e.g. Ru and Ni) while the late transition metals (e.g. Pt and Ir) display a weaker interaction between the graphene and the substrate. A system that is weakly interacting ought to result in a graphene layer that retains its physical properties to a larger degree and also that removing it from the substrate for further processing is possible. Graphene can be grown on Cu with Chemical Vapor Deposition (CVD) at rather extreme conditions: very high temperature close to the melting point of the metal is necessary and also the presence of a stabilizing hydrogen atmosphere is seemingly warranted to achieve high quality films [1]. It has been shown that the graphene on Cu forms domains that display a Moiré pattern, i.e., there is a small angle between the crystallographic directions of the substrate and that of the graphene overlayer [2]. Formation of domains is generally a sign of a weak interaction.

In the present study, a graphene layer is formed by deposition of a 6 Å thick carbon layer onto epitaxially grown Cu(111) which is in turn grown on Al₂O₃(0001). All steps in the growth process is done in a magnetron sputtering chamber, the 1400 Å thick Cu film is deposited at 900K, annealed at the same temperature to achieve a high degree of crystalline order and finally carbon is sputtered onto the surface while maintaining the same temperature. The resulting system is then analyzed with Low Energy Electron Diffraction (LEED), X-ray Photoelectron Spectroscopy (XPS) and Near Edge X-ray Absorption Fine Structure (NEXAFS) at D1011.

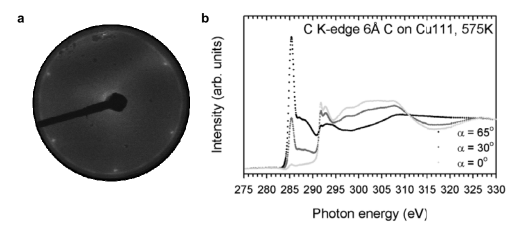


Figure 1: a) LEED pattern showing a 1x1 hexagonal pattern from Cu(111) and a superimposed ring from the graphene. b) NEXAFS spectrum from the graphene layer on Cu, measured in partial yield with a retarding potential of -120V.

We find that domains are formed on the Cu substrate with the ring shaped diffraction pattern from the graphene, see Figure 1a. The inner hexagon is the diffraction pattern from Cu(111), which has a lattice constant of 2.55Å in the (111) plane, very close to the 2.46Å lattice constant found for graphene. Figure 1b display the NEXAFS spectrum over the Carbon K-edge and a clear difference between the π (at 285 eV) and σ (at 292 eV) peaks for the different angles is evident. This means that the carbon layer on the surface is largely bonded in a sp² hybridized

C1s $h_V = 350 \text{ eV}$ 30° take-off 880K ntensity (arb. units) 780K 575K as is 290 284 282 292 288 286 Binding energy (eV)

Figure 2: XP spectra taken at different annealing temperatures and at a take-off angle of 30 degrees.

fashion and that it is flat lying on the Cu surface.

The spectral evolution with temperature is shown in Figure 2, which is a series of C1s XP spectra measured at a take-off angle of 30° to achieve the highest surface sensitivity. The as is sample is just introduced in the vacuum chamber and there is a clear state at 288.5 eV which is associated with carboxylic groups on the surface. The main line is broadened by contributions from carbon atoms in a non-sp² type environment or bonded to hydrogen. Upon thermal treatment to 575K, much lower than the temperature for formation of the film, the C1s line profile is much narrower indicative of a well ordered film. There is also a shift to lower binding energy compared to the as is situation, this is attributed to a change in screening that the graphene experience due to an oxidation of the Cu surface. The O1s signal strength does not change between the two preparations but there is a chemical shift of about 2 eV (not shown here) which hints of a modification of the Cu surface beneath the graphene. At 780 K only a small fraction of the original O1s intensity remains and the shift is reversed because of the similar screening that was present from the beginning. With increasing temperature, the integral intensity of the C1s peak is steadily decreasing, which we interpret as a loss of carbonaceous species on the surface.

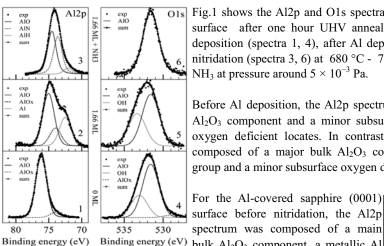
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Room temperature deposition of self-assembled Al nanoclusters on stepped sapphire (0001) surface and subsequent nitridation

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A great amount of interest has shown on the sapphire (0001) surface as a prototype oxide surface to study the interaction between an oxide and metals. In this work, we focused on using the atomic terraces of the stepped sapphire (0001) surface as the initial template for room temperature selfassembling growth of Al nanoclusters, which were then transformed to AlN nanostructures as an advanced template for further growth of other III-nitrides nanostructures. In this contribution, room temperature deposition assembled Al nanoclusters on stepped sapphire (0001) surface and subsequent nitridation were assessed by a high resolution x-ray photoemission spectroscopy (XPS) and a lowenergy electron diffraction (LEED) at beam line D1011 at MAX II.

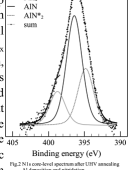


Ols Fig.1 shows the Al2p and Ols spectra of clean sapphire (0001) surface after one hour UHV annealing at 700 °C before Al deposition (spectra 1, 4), after Al deposition (spectra 2, 5) and nitridation (spectra 3, 6) at 680 °C - 780 °C for about 3 hours in NH₃ at pressure around 5×10^{-3} Pa.

Before Al deposition, the Al2p spectrum shows a typical bulk Al₂O₃ component and a minor subsurface AlOx compound in oxygen deficient locates. In contrast, the O1s spectrum was composed of a major bulk Al₂O₃ component, a surface OH group and a minor subsurface oxygen defects.

surface before nitridation, the Al2p spectrum was composed of a main bulk Al₂O₃ component, a metallic Al Fig.1 A12p and O1s core-level spectra after UHV-annealing before (1, 4), after A1 deposition (2, 5) and nitridation (3, 6). component and an interfacial AlO_x component. As for the O1s spectrum,

besides a major bulk Al₂O₃ component, the persistent OH group was unambiguously resolved. This suggests that at least the OH groups observed here were not originated from the weakly bound surface OH absorbents, but strongly bound structural OH groups of the surface owing to surface structural defects. There have been strong evidence in the literature and were 405 confirmed by scanning force microscopy (SFM) experiments that atomic structure disorder and defects were the important centers of hydroxylation



N1s

and for hydroxyl groups be condensed into clusters 1-3. Especially the Al-terminated surface was

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pointed out to be highly reactive with water⁴.

Obvious changes were detected in both Al2p and O1s spectra, accompanied by emerging of nitrogen components in N1s spectrum (shown in Fig.2) after nitridation. The metallic Al component disappeared, while the intermediate component grew to an intensity almost comparable to the main component peak at 74.6 eV, but slightly shifted to a lower BE side at 73.7 eV. This intermediate peak could be assigned either to the AlN component as the result of nitridation of surface metallic Al or to the persistence of surface and or interfacial Al-OH compounds. However, taking the O1s spectrum into consideration, it contained only one narrow symmetric peak, undoubtedly corresponding to O in bulk Al_2O_3 , with no sign of OH groups after nitridation. Therefore the intermediate Al2p peak should be assigned to AlN compound. Interestingly a minor peak with BE at 72.1 eV, even lower than the BE of the earlier metallic Al2p peak was still resolved. We ascribed it to a small fracture of AlH component integrated within AlN phase owing to penetration of H radicals through the surface to react with the Al metals of the nanoclusters during nitridation 5 . The N1s spectrum was fairly wide and was resolved into

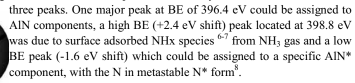


Fig. 3 shows the LEED pattern of the sapphire (0001) surface after UHV annealing before (1), after (2) Al deposition and nitridation. Before Al deposition, the LEED pattern demonstrated an incomplete (1×1) hexagonal structure, suggesting some extent of oxygen-deficiency and the evidence of Al-rich surface.

After Al deposition and nitridation, some hints of extra sub-spots appeared besides the main hexagonal pattern after Al deposition and nitridation. This indicates that the nitridated nanoclusters possessed a certain degree of single crystallinity. The bright sharp LEED spots were possibly from the underlying sapphire (0001) surface because the Al coverage was nominally less than 2 ML. However, because of a small number of extra spots and bright background probably due to surface charging, the LEED diagram was not definitive enough to infer the crystallographic relationship between the AlN nanoclusters and the underlying sapphire (0001) surface.

The results as a whole indicate that the Al nanoclusters were uniformly nitridated from surface downwards through 3D structures of the nanoclusters. The LEED pattern indicated a certain degree of crystallinity on the nitridated surface with a nominal Al coverage less than 2 ML.

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Electronic structure of Zn doped copper chloride probed by soft x-ray spectroscopies

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CuCl is a wide, direct band gap (3.39 eV), zincblende compound semiconductor with great potential in linear and non-linear optical applications due to its large free excitonic binding energy (190 meV) [1]. In order to make use of the potentially strong luminescence properties of CuCl in photonic device applications, there should be a reliable technique for doping of this material system. Since, Zn acts as an n-type dopant for CuCl, the electronic and optical properties of CuCl varies significantly as a function of percentage of Zn doping in the copper lattice.

So far, there are several reports on the electronic band structure of CuCl [2, 3]. Kono $et\ al.$ investigated the valence band spectra of CuCl using x-ray photoelectron spectroscopy and reported that it consists of two branches [4]. Furthermore, the s, p, and d valence states of CuCl have been reported by Goldmann $et\ al.$ [5]. The electronic structure and energy gap have been studied by comparing the valence band of CuCl with band structure calculations [3]. It was reported by Novakov that satellites are formed when the emission of a 2p photoelectron takes place in parallel with a valence band-conduction band excitation in CuCl [6]. The K β emission and K absorption spectra of Cl in CuCl have already been discussed in terms of the energy band structure [7].

However, there is no report on the changes in the electronic structure under the influence of Zn doping in CuCl. In order to verify the chemical bond formation and electronic structure of this material in detail, we used soft x-ray absorption and high resolution photoemission spectroscopy to understand the element specific conduction band density of states and to study the valence and core level photoemission of various Zn doped γ -CuCl samples.

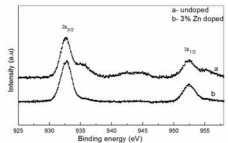


Figure 1: Cu 2p core level spectra of the (a) undoped and (b) 3 wt % Zn doped CuCl films

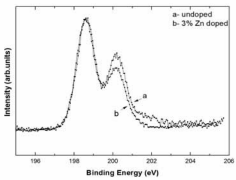
Thin films of CuCl:Zn were deposited onto Si substrates by pulsed dc magnetron sputtering of CuCl/Zn target, with the weight percentage of Zn varying from 0-5%. The doped films exhibit maximum n-type carrier concentration of the order of $\sim 10^{18}$ cm⁻³ (3 wt % Zn doped). We report here the first measurements of Zn doped γ -CuCl performed at the D1011 beamline, Max-lab, Sweden. Figure 1 shows the comparison of Cu $2p_{1/2}$ and $2p_{3/2}$ core level spectra of undoped CuCl and 3% Zn doped CuCl. In figure 1(a) there is evidence of satellite peaks in the undoped CuCl, which has been reported as being due to the presence of cupric species on the surface of the sample as a result of exposure to oxygen or moisture [8]. Moreover, shoulder peaks are also observed on the higher binding energy side of the 2p peaks. These can also be considered as supporting information for the existence of cupric species in the pristine sample. In Figure 1(b) the 2p line shape varies significantly under the influence of Zn doping, the intensity of satellite peaks and shoulder peaks decreases, although

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there is no change in the copper peak position. This could be attributed to the disappearance of cupric species under the influence of Zn doping.

Fig. 2 shows the effect of Zn doping on the Cl 2p level of CuCl. The increased intensity of the higher binding energy Cl 2p peak is most likely due to the presence of the cupric species in the pristine sample. Similar observation has been reported by Kishi $et\ al$. for Au and Pd chlorides [9]. There is an increase in the intensity of Zn 2p core levels as function of Zn doping (not shown) and the oxidation state of Zn is $2p^{2+}$, this being the species that replaces Cu in the lattice.

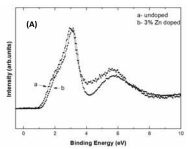
The valence band spectra of the undoped and the 3 wt % Zn doped CuCl samples are illustrated in figure 3(A). The influence of Zn doping can be observed as a shift of ~0.18 eV to the higher binding energy side for the valence band edge of the 3 wt % Zn doped sample with respect to the undoped material. This is assumed to be an indication of filling of the conduction band in the doped samples and perhaps a number of defects are produced by the doping. In figure 3(B)



NEXAFS comparison of the Cu L edge of both undoped and doped are shown. Significant

Figure 2: Cl 2p core level spectra of the (a) undoped and (b) 3 wt % Zn doped CuCl films

differences in the white line intensity and fine structures were observed and are attributed to the multiple valency, copper vacancies and the influence of chlorine species.



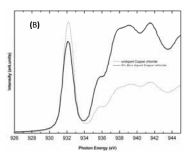


Figure 3(A) Valence band spectra, (B) unoccupied states of Cu 2p of (a) undoped and (b) 3 wt % Zn doped CuCl film.

These are the preliminary observations of the effect of Zn doping on the core, and valence band of CuCl. Further studies are in progress.

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Single molecule magnets on a Au(111) surface: *in-situ* electrospray deposition and a comparison of different ligand shells

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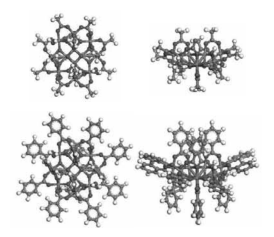


Figure 1. (top) Chemical structure of $Mn_{12}(acetate)_{16}$ viewed from above and from the side, (bottom) Structure of $Mn_{12}(benzoate)_{16}$ as viewed from above and the side.

Single molecule magnets (SMMs) represent an example of a self-contained molecular building block, with magnetic properties, which has the potential to be incorporated as an active component within nano-scale devices. The intrinsic magnetic properties of these fascinating and complex molecules arise from the specific oxidation states of the metal atoms at their core. SMMs are so-called as they exhibit a range of magnetic properties which originate from a purely molecular basis [1] and it is their potential use as components within quantum computation devices which has caused great interest. Some of the more widely studied molecular architecture, and the focus of this experiment, are those based on the mixed valence dodecamanganese (III, IV) cluster which contains four Mn⁴⁺ ions within the core and eight Mn³⁺ ions around the periphery. The magnetic core of such SMMs is protected by an organic shell, with one of the simplest examples using acetate ligands (Mn₁₂(acetate)₁₆) molecule. An obstacle to the formation of well defined monolayers or network-isolated clusters however is that SMMs tend to be non-volatile and therefore cannot be sublimed onto a surface under UHV conditions. In order to overcome this hurdle and to facilitate the study of non-volatile molecules with high resolution synchrotron-based spectroscopy, we developed, a few years ago, a novel in-situ UHV deposition system, based on electrospray ionisation. Essentially, the system forms nanoscale droplets of a liquid solution at the tip of a high voltage capillary. These droplets undergo a succession of Coulomb fission events and solvent evaporation to leave a beam of desolvated molecular ions which are transported through 4 stages of differential pumping before arriving at the sample surface at pressures in the 10^{-8} mbar range [2-4].

In this experiment we successfully deposited the $Mn_{12}(acetate)_{16}$ and $Mn_{12}(benzoate)_{16}$ single molecule magnet molecules shown in figure 1 *in-situ* onto atomically clean Au(111) surfaces under UHV conditions at beamline D1011. The purpose of this experiment was to find out if a

larger ligand shell would be more effective at protecting the magnetic core of the molecules from the reduction that has previously been attributed to charge transfer from the surface or distortion in the adsorbed molecules. The SMMs were deposited using a commercial UHV-compatible electrospray deposition source (Molecularspray Ltd, UK) [5] from a methanol solution at an emitter voltage of +2kV and a flow rate of 500 nlmin⁻¹. The pressure during the deposition was $\sim 4x10^{-8}$ torr.

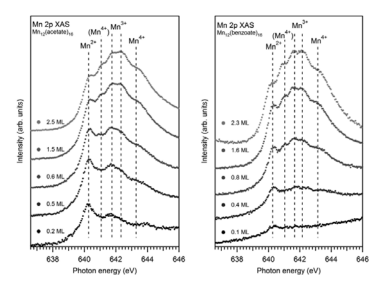


Figure 2. NEXAFS spectra measured at the Mn 2p adsorption edge (Mn L-edge) for a range of SMM coverages (0.2 ML - 2.8 ML)

Figure 2 shows the Mn 2*p* (L-edge) NEXAFS for a range of coverages of both SMMs between 0.2 ML and 2.5 ML from which the oxidation state of the manganese atoms in the core of the SMM may be identified. For multi-layer coverages both sets of spectra are consistent with a mixture of Mn³⁺ and Mn⁴⁺ oxidation states similar to that previously observed [6] for the undamaged Mn₁₂(acetate)₁₆ molecule (4 Mn⁴⁺ and 8 Mn³⁺ atoms). However for sub-monolayer coverages (0.1 ML to 0.8 ML) the spectra observed are consistent with that of a sample consisting predominantly of the Mn²⁺ oxidation state. Thus for sub-monolayer coverages of both magnets the majority oxidation state for the Mn atoms is 2+. This suggests that the larger ligands of the benzoate SMM are not sufficient to protect the core from the surface-induced reduction. We are currently investigating SMMs with even bulkier ligands and the adsorption of a range of single molecule magnets on oxide surfaces.

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Oxidation of h-BN monolayer on Ir(111) studied by core-level spectroscopies

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Recently the possibility of patterned oxygen adsorption on monolayer graphene (MG) grown on 5d metal substrates has been demonstrated [1]. In this connection a characterization of the oxidative process of h-BN monolayer (an isostructural and isoelectronic analog of MG) on transition metal substrates is of considerable interest. In this work the interaction of atomic oxygen with a h-BN monolayer grown on Ir(111) was studied by means of near-edge x-ray-absorption fine structure spectroscopy (NEXAFS) corelevel photoelectron spectroscopy (CLPES) and low-energy electron diffraction (LEED).

All the spectroscopic measurements were carried out *in situ* at the beamline D1011. The surface of Ir(111) crystal was cleaned by several cycles of Ar^+ sputtering, heating in oxygen atmosphere and subsequent annealing in UHV. The cleanliness of the substrate was verified spectroscopically by CLPES

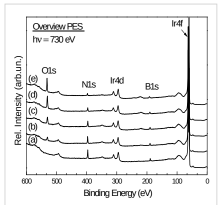


Fig.1. The overview PE spectra taken for: (a) – clean h-BN/Ir(111); (b) – 3 min. O; (c) – 10 min. O; (d) – 40 min. O; (e) – after further annealing at 600° C

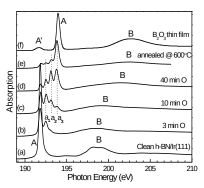


Fig. 2. Evolution of the B K-edge NEXAFS spectra of h-BN on Ir (111) upon oxidation and further annealing (a-e). (f) - B K-edge NEXAFS spectrum of B₂O₃ thin film [3] with photons incident on the surface at an angle of 50° relative to the substrate.

and LEED. Boron nitride films were grown by thermal cracking of vaporized borazine in contact with the hot metal surface at the temperature of around 1050 °C at exposures of 90L (the borazine partial pressure was of 1×10^{-7} mbar). The samples were treated at RT by atomic oxygen produced in a commercial oxygen cracker (from MBE-Komponenten GmbH). The source was operated at the temperature of 1650 °C and oxygen partial pressure of 3×10⁻⁸ mbar. All samples were characterized by PES, NEXAFS and LEED. The B K and N K NEXAFS spectra were measured in the partial electron yield mode (retarding potential U = -100 V and U = -200 Vrespectively) in order to increase the signal-tobackground ratio. The photoelectron spectra were measured relative to the Fermi level in the normal emission geometry.

The weakness of the h-BN/Ir(111) interfacial bonding along with the lattice mismatch between Ir(111) and h-BN result in the formation of weakly corrugated h-BN overlayer on Ir(111) [2]. The perfect quality of the resulting h-BN monolayer was confirmed by sharp LEED patterns showing a characteristic moiré structure. Subsequent oxidation produced only a gradual blurring of the patterns with increasing O coverage, implying that the oxygen adsorption occurs randomly. The overview PE spectra (Fig. 1) corresponding to the pristine sample (a), every stage of its oxidation (b-d) and the sample after annealing (e) show that the intensity of the N 1s core level peak is decreasing upon oxidation while the intensity of the B 1s peak remains mainly unchanged. Clearly, the oxidation process is accompanied by removing N atoms from the h-BN lattice. The B K-edge NEXAFS spectra (Fig. 2) show that already at the first step of oxidation (b) the main sharp peak A corresponding to the B 1s $\rightarrow \pi^*$ transition is decreasing in intensity significantly and new high-energy structures a1, a2 and a3 start to develop. Moreover, band B (representing transitions to the σ^* states) is smeared out. At the

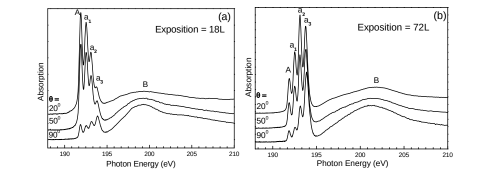


Fig. 3. Angle-dependent B 1s NEXAFS spectra of oxidized BN overlayer on Ir(111) where θ is an angle between the light polarization vector and the substrate surface normal. (a) exposure of 18L, (b) exposure of 72 L (oxygen saturated) sample.

following oxidation steps (c-d) a further decay of the peak A intensity occurs, while structures a_1 , a_2 and a_3 are enhanced non-equivalently. The B K-edge spectrum of the saturated O/h-BN/Ir(111) sample is characterized by the dominant a_3 and a_2 structures. An annealing at 600°C (e) results in the suppression of peaks A, a_1 , a_2 and the domination of peak a_3 . The spectrum of an annealed sample (e) is very similar

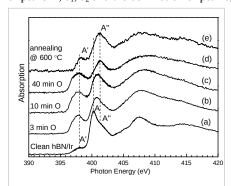


Fig. 4. Evolution of the N K-edge NEXAFS spectra of h-BN on Ir (111) upon oxidation and further annealing (a-e) with photons incident on the surface at an angle of 50° relative to the substrate.

to that of B_2O_3 (f) [3]. The B $1s^4\pi^*$ core excitations in B_2O_3 are of a quasi-molecular nature, being determined by the BO₃ planar units of the D_{3h} symmetry [3]. It is plausible to suggest that peak a_3 corresponds to the B atoms for which all of the neighboring nitrogens are replaced by oxygen.

The rapid growth of peak a₃ points at an active process of embedding O atoms into the h-BN lattice by replacing N atoms in agreement with PES data. We believe that this process has to proceed through the intermediate stages corresponding to a replacement of either one (a₁) or two (a₂) N atoms in the BN₃ unit of h-BN. That is why at initial steps of oxidation the appearance of peaks a₁ and then a₂ precedes the development of a₃. An analysis of the B 1s NEXAFS spectra of the h-BN/Ir(111) (not shown) upon the Ar⁺ion treatment reveals only a smearing of band A, thus excluding defects as an origin of peaks a₁, a₂ and a₃. From the angle-dependent B 1s spectra (Fig. 3) it is

apparent that the states corresponding to peaks a_1 , a_2 and a_3 are of the π^* symmetry, being caused by the B $2p_z$ π - O $2p_z$ π covalent bonding. An incomplete suppression of peaks a_1 , a_2 and particularly a_3 in going from the grazing to the normal incidence (for both oxidation steps) can be due to strong atomic buckling at the corresponding $BN_{3-x}O_x$ (x=1,2,3) sites. This fact is easy to understand taking into account a difference between the B-N and B-O bond lengths. The main trends in the evolution of N 1s spectra with oxidation (Fig.4) are their decrease in intensity and a considerable suppression of band A associated with the π^* states of pristine h-BN overlayer. This suppression is a direct consequence of destroying the B-N bands due to the replacement of N atoms by oxygen.

In conclusion, the adsorption of atomic oxygen on h-BN/Ir(111) occurs mainly through the embedding of O atoms into the h-BN lattice. This is different from the case of MG/Ir(111), for which oxidation occurs predominantly through the formation of epoxy groups [1]. On the other hand, the simultaneous formation of epoxy-like B-O-N chains at the initial oxidation stages (as proposed in [4]) cannot be excluded.

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Growth of the first water layer on rutile TiO₂(110)

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In this work we address a most central and straightforward question in surface science: How does the very first layer of water form on a defect free metal oxide surface? With respect to fundamental studies, water adsorption on single crystalline rutile $TiO_2(110)$ can be considered as the prototype system, and the $H_2O/TiO_2(110)$ system has indeed received more attention than any other comparable system [1-3].

So far, experimental studies addressing the formation of a water monolayer on the defect free TiO₂(110) surface have been lacking, while addressed frequently in the literature by theory. Early studies predicted dissociative adsorption at all coverages while more recent studies suggest a delicate balance between dissociated, partial dissociated and molecular configurations, a balance that most probably depends on the coverage, e.g. [4,5]. Recently we presented the first experimental work to give a conclusive answer to the long-standing question whether water dissociates or not on a surface free from oxygen vacancies [6]. Using the set-up at beamline D1011, we found evidence for mixed molecular and dissociative water adsorption at monolayer coverage on the stoichiometric rutile TiO₂(110) surface. At monolayer coverage the OH:H₂O ratio is close to 0.5 and reducing the coverage by heating was found to result in an increased OH:H₂O ratio. The OH species of the monolayer recombines and leaves the surface at much lower temperatures than OH formed by water dissociation on oxygen vacancies.

In this contribution we present new data on the $H_2O/TiO_2(110)$ system, also collected at D1011. The formation of the first layer of water on the stoichiometric rutile (110) surface is here unraveled in real-time measurements of O 1s photoelectron spectra, giving a detailed picture of how the OH: H_2O ratio depend on the coverage. The O 1s spectra were measured during water uptake at 210 K. Figure 1 (left) shows a 2D plot of the O 1s spectral intensity vs. elapsed time (=exposure) attained at a pressure of $1.5 \cdot 10^{-10}$ mbar. Each one of the spectra was delineated into components from TiO₂, OH and H_2O . Figure 1 (right) gives an example of a fitted spectrum.

Figure 2 shows H_2O and OH coverages as a function of time for growth at (a) $3\cdot10^{-10}$ mbar, (b) $1.5\cdot10^{-10}$ mbar and (c) $5\cdot10^{-11}$ mbar (background pressure), respectively. The coverage is obtained by first adding the O 1s intensities as $0.5\cdot[O\ 1s(OH)]+[O\ 1s(H_2O)]$ and then normalize these values to the monolayer intensity, defined as that attained by heating a water multilayer to $210\ K\ [6]$. For the two higher pressures, saturation was observed. The growth at background pressure had to be terminated before saturation due to time limitations.

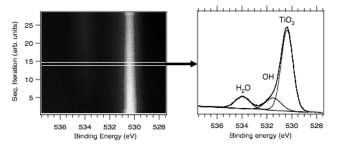


Fig. 1. Left: Real-time O1s spectra for water adsorption on TiO₂(110). Right: decomposition of one of the spectra.

At 3.10^{-10} mbar we reach saturation coverages of 0.82 ± 0.05 ML H_2O and 0.35 ± 0.05 ML OH and at 1.5·10⁻¹⁰ mbar the saturation 0.80 ± 0.05 ML coverages are H₂O 0.40±0.05 ML OH. It is also important to note that the composition is very similar to that reached by either heating of a multilayer to 210 K or by dosing 5 L at a pressure of $5.5 \cdot 10^{-8}$ mbar at 210 K. The composition found for these two preparations is 0.79±0.05 ML H₂O and 0.42±0.05 ML OH. This gives strong support to the absence of radiation induced effects since the same composition is observed under very different preparation conditions and measuring also demonstrates This composition of the complete first layer is not significantly affected by kinetic effects.

The three growth series show that both H₂O and OH are formed from a very early stage. Particularly intriguing is that Figs. 2 (b) and (c), i.e. the two lowest pressures investigated, show very similar growth rates of the H₂O and OH coverages from 0.04 ML up to about 0.40 ML total coverage. A constant H₂O:OH ratio during growth is a strong indication of cluster formation with a specific composition. In all three series it is observed that the OH coverage levels out well before the H₂O coverage

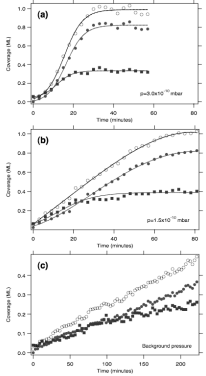


Fig. 2. Coverage of H_2O (filled circles), OH (filled squares) and total H_2O uptake (open circles) as function of esposure time.

does. The point at which this happens is rather similar for all three situations, suggesting a critical total coverage of 0.40±0.05 ML. That is, this coverage corresponds to the maximum coverage of the characteristic mixed water-hydroxyl clusters. Above this coverage, water only adsorbs intact. Ongoing ab initio calculations on the formation of H₂O-OH complexes at a series of coverages will help to understand the growth mechanism up to the critical point of 0.4 ML.

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Mixed dissociative and molecular water adsorption on anatase TiO₂(101)

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The adsorption of water on a metal oxide surface defines a scientific problem of high general importance [1]. The archetypical system in this respect is water on the (110) surface of rutile titanium dioxide (TiO₂) [2]. The adsorption state of water (molecular vs. dissociated) on a rutile (110) surface free from defects has been highly controversial over the years [3-5]. Very recently, we demonstrated for the first time that the first water layer on rutile (110) contains a significant fraction of dissociated species, even in a total absence of surface oxygen vacancies [6].

In this contribution, we present another study, also conducted at beamline D1011, that addresses water adsorption on a related system, the (101) surface of anatase TiO₂. Again a set of O 1s photoemission spectra optimized for maximum surface sensitivity is used to unravel the temperature dependent adsorption state of water on the stoichiometric surface.

Figure 1 shows O 1s spectra obtained after adsorption and subsequent annealing of a multilayer of water. The spectra have been delineated into individual contributions from the TiO₂ substrate, OH and H₂O. After adsorption of water at 120 K followed by heating to 160 K in order to desorb multilayer water two O 1s features are observed apart from the TiO₂ substrate peak at 530.5 eV: The state at 531.5 eV, assigned to OH, and the state at 534 eV, assigned to H₂O [6,7]. Thus, the first layer of water consists of both molecular and dissociated species. The amount of OH is far too high to be attributed to oxygen vacancies. OH recombination and desorption of the dissociated species on the stoichiometric surface is furthermore found to occur at much lower temperature than expected for water dissociated on oxygen vacancies [8]. This observation suggests the formation of a pseudo-dissociated form of water [9].

The water and hydroxyl coverages are shown in Fig. 2a. For comparison the corresponding results for rutile (110) are

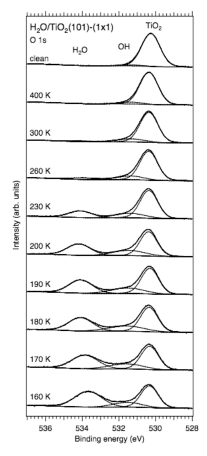


Figure 1. O 1s spectra for water adsorption on the anatase $TiO_2(101)$ -(1x1) surface. The spectra show the results after progressive heating of a multilayer of water.

shown in Fig. 2b. For both anatase (101) and rutile (110), one monolayer (1 ML) corresponds to a density of 5.2·10¹⁴ cm⁻², that is, the density of 5-fold-coordinated Ti [Ti(5)] ions on the surface.

The temperature dependences of the H_2O and OH coverages are qualitatively very

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similar for the two surfaces with a decrease in the H₂O coverage that is more rapid than the decrease in the OH coverage.

By comparing to the O 1s intensity at the monolayer point for rutile (110), we interpolate that a coverage of 1 ML on anatase (101) is attained at about 220 K. This is close to the monolayer temperature of 210 K found for rutile. The composition of the water monolayer on anatase (101) is estimated to be 0.77 ± 0.05 ML H_2O and 0.47 ± 0.05 ML OH. This composition is within the error bars identical to that found on rutile (110) [6].

In conclusion, we find clear evidence for a water monolayer that comprises comparable amounts molecular of dissociated species. The behavior is essentially the same as that found on rutile (110), with a similar H₂O/OH ratio of the monolayer and a H₂O/OH ratio that decreases with decreasing coverage. The finding of a mixed hydroxylmolecular water layer is in stark contrast to previous work, e.g. [10]. The previous studies advocate molecular adsorption only, a view that is based on STM in conjunction with DFT calculations and standard XPS. Our new results demonstrate convincingly that highquality spectroscopic input is necessary in order to obtain a correct picture.

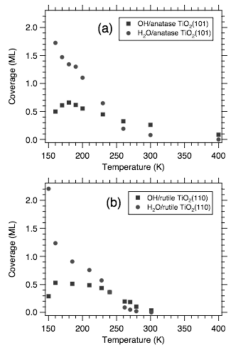


Figure 2. Estimates of the OH and H_2O coverages in monolayers on the (a) anatase $TiO_2(101)$ -(1x1) and (b) rutile $TiO_2(110)$ surface as a function of temperature.

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X-ray magnetic circular dichroism studies of the graphene/Fe/Ni(111) intercalation-like system

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Recent demonstration of spin injection in graphene [1] opens a new road in application of this 2D material in spintronics questioning electronic, magnetic, and interfacial properties of the graphene/ferromagnet layered system, which have to be studied in details. In our latest works we have demonstrated by means of x-ray magnetic circular (XMCD) and dichroism spin-resolved photoelectron spectroscopy that the net magnetic moment of about 0.05-0.1µ_B per carbon atom is induced in the graphene layer via its contact with ferromagnetic Ni(111) substrate [2,3]. magnetic properties of the graphene layer (net magnetic moment as well as exchange interaction) in this system are expected to be improved via intercalation of thin Fe layers underneath graphene on Ni(111) due to the larger magnetic moment of an Fe atom.

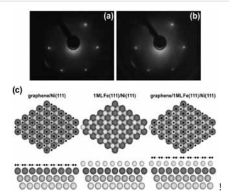


Fig. 1. LEED images of the (a) graphene/Ni(111) and the (b) graphene/Fe(111)/Ni(111) systems. (c) Top- (upper panel) and side-views (lower panel) of graphene/Ni(111), Fe(111)/Ni(111), and graphene/ Fe(111)/ Ni(111).

The present experiments on the graphene/Fe/Ni(111) intercalation-like system were performed at the D1011 beamline of the MAX-lab. The procedure of the sample preparation and experimental conditions are identical to the one described in Refs. [2-4].

Fig. 1 shows the LEED images of (a) graphene/Ni(111) and (b) graphene/1 ML Fe(111)/Ni(111) with the corresponding crystallographic structures obtained from our DFT calculations: the *top-fcc* configuration for graphene/Ni(111) and the *top-hcp* configuration for the graphene/Fe/Ni(111) system (these notations are always considered with respect to the underlying metal layers, Ni or Fe).

The magnetic properties of the graphene layer on the ferromagnetic Ni(111) surface were studied in details in Refs. [2,3] and it was shown that strong hybridization of the graphene π and Ni 3d states leads to the appearance of the induced magnetic moment of carbon atoms with a value of 0.05-0.1 μ B per carbon atom. The respective reduction of the magnetic moment of the Ni interface atoms, compared to the bulk value, was predicted and observed in the former experiments.

The results of the investigation of the magnetic properties of the graphene/1 ML Fe(111)/Ni(111) system are summarized in Fig. 2. The XMCD spectrum collected at the Ni $L_{2,3}$ edge in the TEY mode is in perfect agreement with the previously published data [2,3]. The bulk values of the spin- and orbital-magnetic moments μ_S =0.69 μ_B and μ_L =0.07 μ_B of Ni calculated from the spectra on the basis of sum-rules are in very good agreement with previously published experimental values. The Fe $L_{2,3}$ XMCD spectrum is in good agreement with previously presented data for the fcc Fe [5]. The spin- and orbital-magnetic moments can be estimated from these data assuming the number of holes in the iron layer in the graphene/Fe(111)/Ni(111) system to be n_h =3.7 (for bulk Fe n_h =3.4; transfer of 0.15e from Fe to Ni and the same value from Fe to graphene layer; the present theoretical calculations give value of n_h =3.691 for Fe 3d states). This number leads to μ_S =2.56±0.1 μ_B and μ_L =0.31±0.05 μ_B for the spin- and orbital-magnetic moments of Fe atoms in the intercalated layer which are in very good agreement with value of spin-magnetic moment 2.469 μ_B for Fe obtained in calculations for graphene/1 ML Fe(111)/Ni(111). The relatively large uncertainty in the value of magnetic moment extracted from experiment arises mainly from

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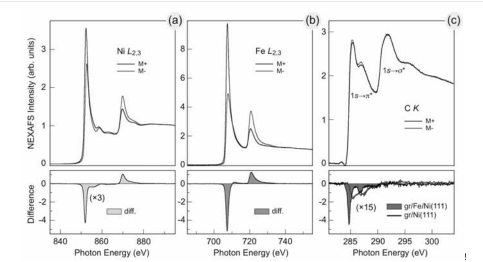


Fig. 2. XMCD spectra of the graphene/Fe(111)/Ni(111) system measured at the (a) Ni $L_{2,3}$, (b) Fe $L_{2,3}$, and (c) C K absorption edges. The upper panels show the absorption spectra measured with the circularly polarized light for two opposite magnetization directions and the lower panels present the corresponding. The XMCD difference obtained for the graphene/Ni(111) system (multiplied by factor 15) is reproduced from Ref. [3] in the lower panel of (c).

the estimation of the number of Fe 3d holes and from the error for the degree of circular polarization of light.

The C K-edge XMCD spectrum of graphene/Fe(111)/Ni(111) is strongly modified compared to the one measured for the graphene/Ni(111) system [2,3]. The most important observation is the increasing of the magnetic contrast at the C K edge by factor of 2.7 which correlates with the theoretical predictions about spin-magnetic moment of carbon atoms in the graphene layer on the Ni(111) and Fe(111)/Ni(111) substrates. As in the previous case [2] the relatively strong XMCD contrast is detected for C $1s \rightarrow \pi^*$ transitions whereas there is almost no variation of the absorption signal upon magnetization reversal is visible for the C $1s \rightarrow \sigma^*$ transitions. These observations are explained well by the theoretically predicted strong hybridization of the out-of-plane graphene π and Fe 3d states and the existence of the relatively weak hybridization between in-plane graphene σ and Fe 3d states, similar to the graphene on Ni(111) [2,3].

The C K-edge XAS spectrum of the graphene/Fe(111)/Ni(111) system in the energy range corresponding to the $1s \rightarrow \pi^*$ transition consists of two peaks which can be assigned, similar to Refs. [2,3], to transition of 1s electron on the interface states which are result of the hybridization of C p_z orbitals of graphene layer and Fe 3d orbitals. The modification and the increasing of the XMCD contrast going from graphene/Ni(111) to graphene/Fe(111)/Ni(111) can be explained by the larger energy splitting between spin-up and spin-down C-projected density of states. XMCD spectra measured at the C K-edge can only provide information on the orbital moment. From the negative sign of the XMCD signal one can conclude that the averaged orbital moment of carbon atoms of the graphene layer is aligned parallel to both, the spin and orbital moments of the substrate layer. It is noteworthy that the orientation of individual spin and orbital moments of both Fe and C at different sites cannot be determined from the experimental XMCD data. On the basis of comparison of the dichroic signals measured at the C K-edge for the graphene/Ni(111) and graphene/Fe(111)/Ni(111) systems one can estimate the increasing of the spin-magnetic moment of carbon atoms up to $0.2\mu_B$ which is consistent with theoretical results [2,6].

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The impact of atomic oxygen on graphene weakly bound to metal substrates

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The oxygen adsorption on the monolayer graphite (MG or graphene) grown on the metal substrates has been investigated by X-ray core level photoelectron spectroscopy (CLPES or CLS), near-edge x-ray adsorption spectroscopy (NEXAFS) and scanning tunneling microscopy (STM)[1]. The goal of the study was to provide deeper insight into the mechanism of graphene's interaction with active adsorbates such as atomic oxygen and give information about a possibility of fine tuning of graphene's electronic structure by patterned adsorption of oxygen, which was recently reported for hydrogen [2,3]. The Ir(111) and Pt(111) single crystals were used as substrates for graphene growth since both of them provide periodically corrugated graphene due to the lattice mismatch between the latter and the substrate, and shows a well-known moirè superstructure [4,5]. The highly oriented pyrolitic graphite (HOPG) sample was used as a reference to model freestanding graphene with no corrugation.

All the spectroscopic measurements were carried out *in situ* at the beamline D1011. The surface of Ir(111) and Pt(111) crystals was cleaned by several cycles of Ar^+ -sputtering, annealing in oxygen atmosphere and subsequent flashing in UHV. The cleanliness of the substrate was checked by CLS and low energy electron diffraction (LEED). Graphene was grown by thermal cracking of propylene (C_3H_6) in contact with the hot metal surface at the temperature of ~1000°C in the propylene partial

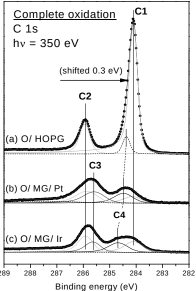


Figure 1. Impact of atomic oxygen on C 1s PE spectra from O-HOPG (a) and and O-MG on Pt(111) (b) and Ir(111) (c). All spectra correspond to saturated oxidation. The spectrum from HOPG is shifted by 0.3 eV to lower BE to facilitate the comparison.

pressure of 1×10^{-7} mbar, the quality of the sample was verified by CLS and LEED. Cleaning of the HOPG sample was performed by cleaving it in situ with subsequent annealing in UHV to 600°C. The samples were treated at RT by atomic oxygen produced in a commercial oxygen cracker (from MBE-Komponenten GmbH). The cracker was operated at the temperature of 1650°C and oxygen partial pressure of 1×10⁻⁷ mbar. All samples were characterized by CLS, NEXAFS and LEED. The C K-edge NEXAFS spectra were measured in the partial electron yield mode (retarding potential U = - 100 V). The photon energy resolution of the C 1s NEXAFS spectra was set to 75 meV, the resolution of the C 1s PE spectra was 125 meV, at hv =350 eV. The STM measurements were performed at RT on the samples prepared in a separate chamber equipped with a commercial UHV STM (STM1 from Omicron GmbH), in the Division of Synchrotron Radiation, Institute of Physics, Lund University.

Figure 1 shows the C 1s photoelectron spectra of saturated oxidation of samples, i.e. no changes in the spectra were detected upon further oxidation. The spectrum from pristine HOPG can be fitted with just one peak (C1) at the binding energy (BE) of 284.42 eV, while at low and intermediate O coverage an additional peak (C2) develops at the BE of 286.22 eV (relative shift +1.8 eV). The detail C2 is assigned to the formation of epoxy groups on

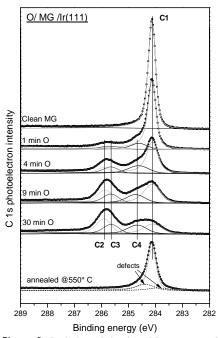


Figure 2. Evolution of the C 1s PE spectrum of graphene on Ir(111) as a function of increasing oxidation time up to the saturation level, and further annealing at 550° C.

the basal plane of graphite, while the broadening of peak C1 (which was handled by an additional component at higher BEs) is associated with the slight buckling of those graphite atoms in the uppermost layer, which are not directly bound to oxygen. Formation of other functional groups than epoxy can be excluded on the basis of O 1s PE spectra analysis. The peak assigned to epoxy groups on O-HOPG (C2) is considerably broadened in the samples of oxidized graphene on Pt(111) and Ir(111), Fig. 2 (b), (c) respectively. This peak can be described in terms of two contributions: one, analogous to oxidized graphite, at 285.9 eV (C2) and a new peak at 285.6 eV (C3). Peak C4 associated to the buckling of graphene close to sites of oxygen adsorption is also found on the metal substrates. Components C3 and C2 are attributed to the epoxy groups formed in the inequivalent sites of the patterned geometrical corrugation of graphene, namely bonding (closer to metal surface, "pores") and less bonding (further from metal surface, "wires"). The components C2 and C3 then correspond to epoxy groups formed in these higher and lower sites respectively. This assignment is based on the observation that C2 is separated from the original C1 component by 1.8 eV, just like in the case of epoxy groups on HOPG.

The transformation of the C 1s photoelectron spectrum in MG/Ir(111) upon successive oxidation steps is presented in Figure 2 together with a spectrum obtained after annealing of a fully oxygenated sample. Upon oxygen adsorption features C2, C3 and C4 become more pronounced, at the expense of the signal from the pristine sp^2 -bonded graphene (C1), indicating an increase in the number of carbon atoms involved into the oxidation process. The rate of growth/decline of these peaks is not the same for all four peaks, which indicates a patterned adsorption of oxygen onto the sample surface. Upon annealing the system tends to release oxygen and after annealing at 500°C no oxygen signal could be detected, however the resultant carbon peak is broadened and reduced in area to 50-60% compared to the C 1s obtained on pristine graphene on Ir(111). This means that part of the graphene layer has been etched away upon heating, most likely desorbing in the form of CO/CO2, and thus the remaining layer contains defects that broaden the C 1s line. The CLS on the 4f core level of Pt(111) and Ir(111) substrates has revealed a gradual pinning of graphene to the metal substrates upon oxidation. Our STM studies support these spectroscopic findings, emphasizing the role of the patterned adsorption – upon annealing "wires" are more sensitive to oxygen etching, resulting in the formation of punched holes in the otherwise reasonably restored graphene lattice [5]. In conclusion, the adsorption of atomic oxygen on graphene was investigated and was found to occur mainly via formation of epoxy groups. For the plane samples with (HOPG) it occurs randomly and does not introduce significant geometrical distortion to the sample. For periodically corrugated samples (MG/Ir(111), MG/Pt(111)) patterned oxygen adsorption takes place which causes substrate-mediated buckling of the graphene adlayer and upon annealing produces punched holes in less-bonding sites of the moirè corrugation pattern.

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The hBN nanowaves on the Fe(110) surface

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A detailed experimental study of growth and structure of the single layer hexagonal boron nitride (*hBN*) grown on the Fe(110) surface was performed by means of core level photoelectron spectroscopy (CLPES or CLS), near-edge x-ray adsorption fine structure (NEXAFS), low energy electron diffraction (LEED), low energy electron microscopy (LEEM) and scanning tunneling microscopy (STM) [1]. It was found that a monolayer of hBN forms on the Fe(110) surface a ribbed superstructure of strips lifted above the surface, "waves" and regions lying close to the surface, "grooves". This structure arises from a strong interfacial interaction between iron substrate and hBN adlayer and the lattice mismatch between those in the certain direction, similarly to the "nanomesh" hBN/Rh(111) [2].

All spectroscopic measurements were carried out *in situ* at the beamline D1011. The surface of Fe(110) was prepared by epitaxial growth of iron films (thickness ca. 20 nm) on the surface of W(110) single crystal. The W(110) single crystal was cleaned by several cycles of Ar⁺-sputtering, annealing in oxygen atmosphere and subsequent flashing in the UHV. The cleanliness of the substrate was checked by

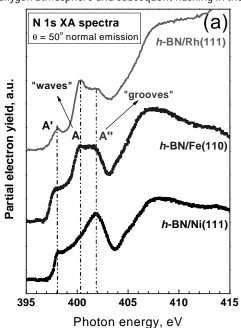


Figure 1. NEXAFS N K-edge (a) and PE N 1s (b) spectra of hBN/Fe(110). The same spectra of hBN/Rh(111) and hBN/Ni(111), the top and bottom curves on both figures, correspondingly are shown for comparison.

CLS and low energy electron diffraction (LEED). After deposition of iron the sample was gently annealed to improve the quality of the resultant Fe(110) surface, which was controlled by the LEED pattern [3,4]. hBN adlayer was grown by thermal cracking of vaporized borazine (B₃N₃H₆) in contact with the hot metal surface at the temperature of ~750°C in the borazine partial pressure of 1×10^{-7} mbar, the quality of the sample was verified by CLS and LEED. All samples were characterized by CLS, NEXAFS and LEED. The N and B K-edge NEXAFS spectra were measured in the partial electron yield mode (retarding potential U = -100 V). The photon energy resolution of the N 1s NEXAFS spectra was set to 125 meV, the resolution of the N 1s PE spectra was 150 meV, at hv =470 eV. The LEEM measurements were performed in situ at the beamline I311. The STM measurements were performed at RT on the samples prepared in a separate chamber equipped with a commercial UHV STM (STM1 from Omicron GmbH), in the Department of Synchrotron Radiation, Institute of Physics, Lund University. Base pressure in the STM preparation chamber was better than 5×10^{-10} mbar and in the analytical chamber about 1×10^{-10} mbar.

The Figure 1 shows NEXAFS (a) and PE (b) spectra of nitrogen in hBN/Fe(110), compared to those of hBN/Rh(111) and hBN/Ni(111). The NEXAFS spectrum of N K-edge in the hBN/Fe(110) shows the two

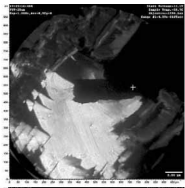




Figure 2. An island of the *hBN* on the Fe(110) surface imaged with the dark field LEEM imaging mode, which allows discrete imaging of structures with different spatial orientations or different symmetry. Domains with two primary spatial orientation are shown. The field of view is 20 μm , the scale bar is 2 μm .

between Fe and h-BN and a lattice matching in only one direction of the surface unit cell, h-BN monolayer grows in the form of strongly corrugated one-dimensional waves in two domain orientations. The corrugation pattern of hBN/Fe(110) can make it a good template for the nanowire growth or self-assembling of linear structures.

bands, A and A" in the region of π^* -resonance. The A band is characteristic for the regions of an hBN adlayer lifted above the surface as seen in hBN/Rh(111) [5], and the A" band is attributed to the strong interfacial coupling of an hBN adlayer and the substrate similar to the case of hBN/Ni(111) [6]. Thus it is natural to suggest strong corrugation of a hBN monolayer on the iron surface i.e. co-existence of both lifted and down regions, i.e. "waves" and "grooves". The LEEM studies in the dark field mode shown in the Figure 2, have revealed two major orientations of hBN domains, corresponding to the directions of lattice match between the hBN and Fe(110) lattices, namely [I \(\bar{I} \) I] and [\(\bar{I} \) I] directions. However due to the considerable misfit between surface W(110) and Fe(110) lattices the surface of epitaxial iron film might be stressed, which could lead to the formation of hBN domains slightly misoriented respectively to the major growth directions, or stretched/shrunk with respect to the lattice constant of the relaxed hBN film.

The STM studies confirm the suggested model of a single hBN layer corrugated in a wavy pattern. Figure 3 represents an STM image of hBN/Fe(110) sample which shows a wavy corrugation pattern. The mean corrugation of the adlayer i.e. the mean difference between highest and lowest point of the

relief is ca. 1 Å, the period of structure is 2.6

In conclusion
we report on a
growth of
hexagonal boron
nitride on the
Fe(110) surface.
Due to the strong
interaction

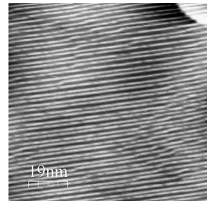


Figure 3. A typical STM image of hBN/Fe(110). The image size is 100x100nm, the scale bar is 19nm.

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Electronic Structure of PTCDA on Pb/Si(111) $\sqrt{3} \times \sqrt{3}$

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Molecular systems that are interesting for molecular electronics applications have attracted an attention in the fields of the surface and solid-state physics. Recently, we have studied the electronic structure of PTCDA (3,4,9,10-perylene tetracarboxylic dianhydride) on the Ag/Si(111) $\sqrt{3}\times\sqrt{3}$, Sn/Si(111) $\sqrt{3}\times\sqrt{3}$, and Pb/Si(111) $\sqrt{3}\times\sqrt{3}$ surfaces by near edge X-ray absorption fine structure (NEXAFS) and core-level photoelectron spectroscopy. Contrary to Ag/Si(111) $\sqrt{3}\times\sqrt{3}$ [1], we found that PTCDA strongly interacts with the Sn/Si(111) $\sqrt{3}\times\sqrt{3}$ surface. Because of the large interaction strength, the molecules no longer lie parallel to the substrate to form a layer-by-layer structure, as shown in the Ag/Si(111) $\sqrt{3}\times\sqrt{3}$ case. Instead the PTCDA molecules grow in a random fashion and form low-ordered phases on the Sn/Si(111) $\sqrt{3}\times\sqrt{3}$ substrate. However, on the other hand, we found that sub-monolayers (ML) of PTCDA weakly interact with the Pb/Si(111) $\sqrt{3}\times\sqrt{3}$ surface and gradually cover the whole substrate with increasing PTCDA coverages. The molecules lie parallel to the substrate and form a self-assembled thin film.

A deposition of 1.5 ML Pb on the Si(111)7×7, following by 340 °C annealing, resulted in a nice Pb/Si(111) $\sqrt{3} \times \sqrt{3}$ surface, evidenced by a sharp $\sqrt{3} \times \sqrt{3}$ LEED (low-energy electron diffraction) pattern. A series of coverages (0.3-10) of PTCDA were evaporated on the Pb/Si(111) $\sqrt{3} \times \sqrt{3}$ surface at room temperature. As shown in Fig. 1, there is an interesting change in the LEED pattern just after 0.3 ML deposition of PTCDA. The LEED pattern shows a multiple ring-structure at a low electron energy. The ring-structure is quite common in the cases of PTCDA on Ag(111), Au (111), and Ag/Si(111) $\sqrt{3} \times \sqrt{3}$. It originates from the herringbone phase of PTCDA molecules. Thus LEED patterns clearly points to a highly-ordered growth model. Only the ring-structure could be observed in LEED after 1 ML evaporation of PTCDA.

Figure 2 shows high-resolution C K-edge NEXAFS spectra recorded from the 2 ML PTCDA on the Pb/Si(111) $\sqrt{3}\times\sqrt{3}$ surface with three incidence angles. The Pb/Si(111) $\sqrt{3}\times\sqrt{3}$ surface has a high coverage of Pb (1 ML), resulting in an incommensurate phase. This surface is very much different from the 1/3 ML $\sqrt{3}\times\sqrt{3}$ phase with dangling bonds. The high-coverage Pb/Si(111) $\sqrt{3}\times\sqrt{3}$ surface is quite inert. Under such interaction strength, one could expect that PTCDA molecules might grow into a layer-by-layer structure. Indeed, NEXAFS measurements from the normal, 40° and 70° incidence angles show clear spectral differences in the π - and σ -resonance region. The π -resonance peaks located between 283-290 eV are very strong at 70° incidence angle; while they decrease in intensity very much at 40°, and become very weak at the normal incidence angle. On the other hand, the σ -resonance peaks located between 290-300 eV remain almost the same at three different geometries. These findings clearly indicate that the π -orbitals of PTCDA molecules are perpendicular to the surface. Thus PTCDA molecules are flat-lying on the surface, and form a thin film with layer-by-layer structure.

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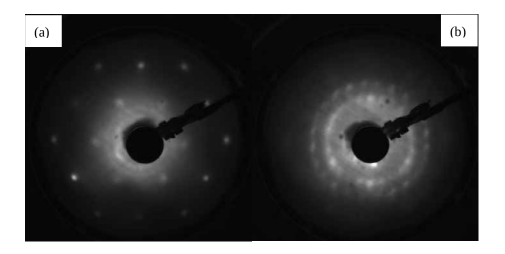


Fig. 1 LEED patterns of 0.3 ML PTCDA on the Pb/Si(111) $\sqrt{3}\times\sqrt{3}$ surface at room temperature. (a) 35.7 eV; (b) 11.7 eV.

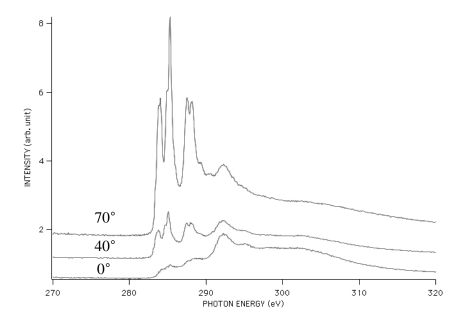


Fig. 2 High-resolution C K-edge NEXAFS spectra recorded from the 2 ML PTCDA on the Pb/Si(111) $\sqrt{3} \times \sqrt{3}$ surface with three incidence angles: 0° , 40° , and 70° .

Soft x-ray spectroscopy on (Ga,Mn)N/GaN digital ferromagnetic heterostructures: first results at MAX-lab

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Recently, extensive efforts have been undertaken to increase the Curie temperature (T_c) of diluted magnetic semiconductor (DMS) materials due to their potential applications for many promising spintronic devices operating at high temperatures. As alternative DMS materials with high T_c , (Ga,Mn)N materials are of current interest because their T_c values can be as high as room temperature, based on theoretical calculations [1]. Theory predicts that the ferromagnetic properties observed in a (Ga,Mn)N material system depend on the occupancy of the Mn energy band in (Ga,Mn)N and the position of the Fermi level relative to this band. Carriers (holes) in the Mn energy band are needed to mediate the ferromagnetic interaction: the depletion and enhancement of the carrier concentration in the band will change the ferromagnetic properties of (Ga,Mn)N. GaNbased DMSs have attracted particular interest because the GaN host has been considered as a promising host material for DMS having high T_c based on theoretical predictions [2]. Although (Ga,Mn)N random alloy thin films were grown by molecular-beam epitaxy (MBE), a high concentration of Mn atoms cannot be incorporated into the GaN host because nitrogen defects and/or ferromagnetic precipitates are inevitably created in the crystals. Therefore, doping high concentration Mn into the GaN host without degrading its quality has been desired to realize Mndoped GaN having a high T_c .

The samples with various composition of Mn to be used in this study are a multiple heterostructure of GaN/(Ga,Mn)N of multi periods grown on the sapphire substrate with a GaN buffer layer by the rf-plasma assisted MBE [3]. The thickness of each (Ga,Mn)N layer and GaN layer is 5 and 15 nm, respectively. On top of it, a thin GaN capping layer (1 nm) is deposited to prevent oxidations of the top (Ga,Mn)N layer. An interesting and promising route to realize high T_c is to synthesize DMS embedded in two-dimensional hetero-structures by digital doping such as digital ferromagnetic hetero-structures (DFH) which consist of nonmagnetic semiconductor layers and highly doped thin magnetic semiconductor layers. This allows the doping of high concentration transition-metal ions into spatially localized regions in the semiconductor host without degrading its quality. We adopted the DFH technique to the growth of (Ga,Mn)N to enhance the local Mn concentration and reported that a considerable enhancement of magnetization and T_c was achieved compared with those of the (Ga,Mn)N random alloy.

The X-ray Absorption Spectroscopy (XAS) measurements were carried out in the electron yield mode measuring the photocurrent of the sample at BL I1011 of MAX-lab. For these first

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experiments we focus on linearly polarized x-rays and attempt a characterization of sample stoichiometry and of the symmetry of electronic states probed by XAS. The samples exhibit as inserted in the Ultra High Vacuum chamber traces of surface contamination, which are removed by Ar ion sputtering. The typical spectra obtained at the N K-edge and the Mn L-edges are shown in Figure 1 (left). We observe the typical features of GaN in the N K-edge XAS spectra. Furthermore, the strong angle dependence obtained as the angle of polar incidence is varied testifies of a high degree of crystallinity for these samples. A strong XAS absorption signal is also obtained at the Mn L-edges. The observation of a strong Mn XAS signal testifies of the successful Mn incorporation within the GaN lattice. For the Mn XAS, in Figure 1 (right), we observe both typical multiplet features on the Mn L-edge white lines, as well as a high inter-peak intensity. Multiplet features are characteristic of the localized part for the final 3d states in XAS, a high inter-peak continuum also testifies of the existence of strong delocalized character as well. Theoretical calculations will allow us to interpret the XAS spectra and eventually assign these spectral features to specific locations of the Mn atoms with the DFH.

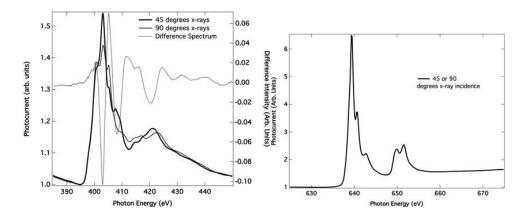


Figure 1. X-ray absorption spectroscopy spectra for (Ga,Mn)N/GaN digital ferromagnetic heterostructures. *Left:* N K-edge data with linear x-rays. A strong angle dependent absorption intensity is found, the difference spectrum between the two angles is also shown. *Right:* Mn L-edge spectra. Interestingly, here no strong angle dependence is found.

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Investigation of magnetism in a solid solution of two antiferromagnets: $\alpha\text{-Fe}_2O_3$ and $\text{FeTi}O_3$

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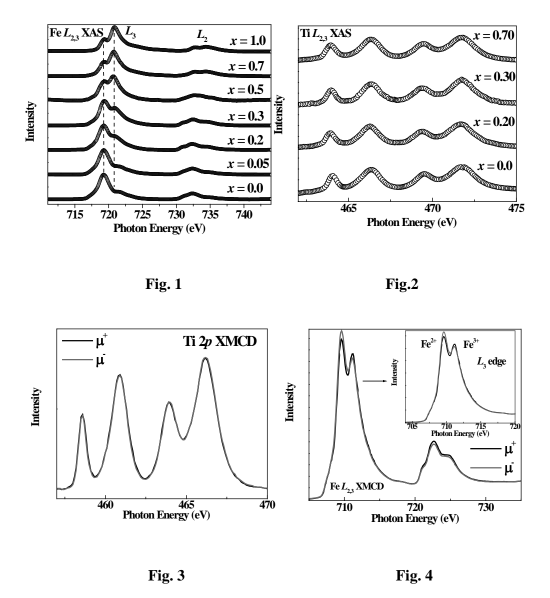
There is a huge scientific and industrial interest in magnetic semiconductors with magnetic transition temperatures above room temperature. One way to achieve this is through doping of transition metal ions into traditional semiconductors, such as Mn doped in GaAs, however, this route has not met with success so far. One unconventional way of achieving this was realized in the solid solution of two antiferromagnets, α -Fe₂O₃ and FeTiO₃ [1], which were found to be strongly magnetic with high transition temperatures.

 α -Fe₂O₃ has a space group of R-3c and is an antiferromagnet with a high Neel temperature of 950 K, and FeTiO₃ has a space group of R-3 and is an antiferromagnet with a Neel temperature of 55 K. The origin of magnetism is intriguing in this solid solution series. Magnetization of this solid solution series is supposedly dictated by the Fe ions, with the relative ratio of Fe²⁺ and Fe³⁺ expected to play a deciding role, however, there has been no systematic study to determine these relative ratios. Further, the role of Ti in the magnetization of this solid solution series cannot be ruled out, especially with the recent observation of Ti XMCD in FeTiO₃, arising from the presence of Ti³⁺ in FeTiO₃ [2].

We have performed detailed x-ray absorption spectroscopic (XAS) studies and x-ray magnetic circular dichroism (XMCD) studies under an applied magnetic field of 0.3 Tesla on carefully prepared single phase samples of solid solution members of (x) Fe₂O₃ – (1-x) FeTiO₃ series with x = 0, 0.05, 0.2, 0.3, 0,5, 0.7, 1 at the I1011 beamline in MAXLAB, Sweden. The Fe $L_{2,3}$ XAS line-shapes are very characteristic of the valency of Fe ions, with Fe L_3 exhibiting a shoulder to the left for Fe³⁺ ions and having a shoulder to the right for Fe²⁺ ions, and thus can be used to characterize the Fe valencies very effectively. We find a gradual evolution of Fe ion valencies from (2+) to (3+) as a function of x across the solid solution series, as shown in Fig. 1. We also find Ti ions to remain as Ti⁴⁺ throughout the solid solution series, as shown in Fig. 2. One of the characteristic XMCD spectra at the Ti $L_{2,3}$ edge is shown in Fig. 3 for two circularly polarized photon beams μ^+ and μ^- and clearly shows the absence of any Ti related XMCD,

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thus clearly establishing that magnetism in this solid solution series is only mediated by Fe ions. Further, we observe a clear XMCD at the Fe $L_{2,3}$ XAS edge, shown in Fig. 4 for x=0.30, thus providing a direct evidence of the role of Fe in magnetization in this solid solution series [3].



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Probing the proposed half-metallic antiferromagnetism in LaSrVMoO₆

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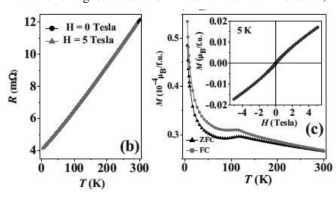
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Half-metallic antiferromagnets (HMAFM) proposed by van Leuken and de Groot, have attracted enormous attention in recent times as they offer an unique possibility of realizing an exotic state that has large degree of spin polarization of conduction electrons but vanishing macroscopic magnetic moment. Unlike a conventional antiferromagnet, in a HMAFM the nullification of magnetic moment occurs between the antiparallel spins of same magnitude, residing on different ions of distinct symmetry, and as a result a perceptible exchange splitting between the up- and downspin channels is observed. Due to this unusual situation, HMAFM materials are also sometime termed as "compensated half-metals."

However, existence of a true HMAFM appeared elusive until Uehara $et\ al.^1$ proposed the curious case of LaSrVMoO6 (LSVMO). LSVMO has been suggested to be a HMAFM system by considering antiparallel alignment of alternately occupied $V^{3+}(3d^2)$ and Mo^{4+} ($4d^2$) spins at the B site of the perovskite structure. Although ab-initio calculations always predicted LSVMO to be a half-metallic ferrimagnet, based on the magnetic and transport measurements, it was claimed that the system can indeed become HMAFM for a perfectly B-site ordered state. Our samples showed identical magnetic and electrical transport behaviours, as can be seen in Figure 1.

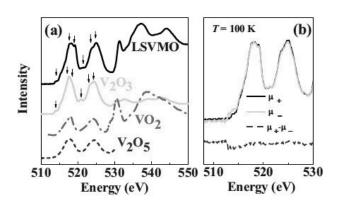
Following this we have attempted to check the proposed half-metallic antiferromagnetic behaviour of the sample using neutron powder diffraction (NPD), x-ray absorption (XAS), and x-ray magnetic circular dichroism (XMCD) measurements. As is seen from the magnetization measurements, there is a weak peak at around 125 K, which was identified as a clear

antiferromagnetic transition in earlier works. 1,2 However, in our 300 K and 2 K NPD



measurements, no indication of any antiferromagnetic ordering obtained. Further, in order to understand 'half-metallicity' of the compound better, XAS and **XMCD** measurements have been

performed on LSVMO. XAS of V L and O K-edges collected at room temperature with finer structures, clearly establishes that vanadium is in 3+ oxidation state. The Mo M-edge XAS spectrum is featureless and does not provide useful information



regarding the Mo charge state. Now, for a true HMAFM state in LSVMO. compensation of spin moments could only he realized by considering AFM coupling between ferromagnetic V³⁺ and ferromagnetic Mo⁴⁺ sublattices.

and therefore, clear XMCD signal from the polarized V *L*-edge spectra can be expected. But surprisingly, no XMCD signal at vanadium L edge with reverse polarizations at 100 K with a field of 0.3 Tesla was observed (see Fig. 2). This observation is in accord with the disordered structure and clearly refutes the HMAFM model proposed for a perfectly ordered structure.

In order to achieve insight about the local chemical structure, which might be critical, XAFS measurements were performed on LSVMO. However, the XAFS results revealed an unusual segregation of every Mo (V) ion toward another Mo (V) ion as the nearest neighbor and toward Sr (La) ion as its next-nearest neighbor. Thus, a picture of an intrinsic chemical fluctuation within very small spatial domains, giving rise to very local LaVO₃ and SrMoO3-like regions could be evoked in this case, which along with *ab initio* calculations based on this chemically inhomogeneous structure succeeded to explain all the unusual electronic and magnetic properties of this compound.

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Structural and magnetic characterization of Co₂MnGe/Rh₂CuSn Heusler multilayers

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It is well known that the performance of electronic components has increased rapidly during the last four decades. The exponential increase in computational power is described by Moore's law. The basic scheme is to decrease the size of electronic components in order the enhance the performance. While this has been very successful in the past, we are now approaching the atomic scale in device engineering, which could severely limit the possibility for further increase in performance. Within magnetic read heads technology several changes has occurred during the last 15 years, the first anisotropic magnetoresistance (AMR) read heads were surpassed by giant magnetoresistance (GMR) in 1996 after which tunneling magnetoresistance (TMR) took over the after 2005. The TMR structure used today has an MgO tunnel barrier which is about 1 nm thick and further substantial decrease is not realistic. It has been suggested that currentperpendicular-to-plane (CPP) GMR structures, as opposed to current-in-plane (CIP) used in the first generation GMR, are candidates to become the next type of magnetoresistance (MR) sensors.

Half metallic ferromagnets1 which have a 100% spin polarized conduction band have been suggested for realizing high performance CPP-GMR where Heusler compounds have shown to be strong candidates.^{2,3} The Co based compounds are of especial interest due to their high Curie temperatures. Full-Heuslers are well ordered ternary compounds with the composition X₂YZ where, in the case of an half metallic compound, X and Y are two different transition metal atoms. Spin valve structures composed of Co₂MnGe (CMG) and various non-magnetic layer has been reported where MR values of 3.6% at room temperature have been achieved4. The half metallic character has been theoretically predicted for perfect crystals while for realistic compounds a certain amount of disorder is expected which are detrimental for the high spin polarization⁵. Ambrose and Mryasov^{6,7} proposed a combination of ferromagnetic (FM) and non-magnetic (NM) Heusler alloys to maximize the spin asymmetry at the FM/NM interface, giving a high spin polarization even if a perfect half metallic character is not obtained. Nikolaev et al.8 constructed an all Heusler CPP-GMR structure according to this scheme using Rh₂CuSn (RCS) as the NM layer and CMG as the FM layer. An MR of 6.7% at RT was achieved which is higher than what has been

TABLE I: Samples annotation according to the ratio between thickness of CMG and RCS layers.

Annotation	CMG (Å)	RCS (Å)	x (repetitions)
S0.66	12	18	16
S1	18	18	13
S1.33	24	18	11
S1.5	18	12	16

reported for transition metal CPP-GMR but lower than for TMR structures which might be an effect of intermixing at the NM/FM interface, a few percent of Cu, Rh or Sn can severely lower the spin polarization in CMG.⁵

We have studied the Heusler compound Co₂MnGe (CMG) as the magnetic layer between non-magnetic layers of the full Heusler Rh2CuSn (RCS) in a multilayer structure. The samples were grown using a commercial magnetron sputtering system (Canon Anelva C7100). The aim of this study is to understand the effects of annealing and its dependence on layer thicknesses. The magnetic properties and in particular the quality of the interfaces are studied using x-ray magnetic circular dichroism (XMCD). We have also conducted experiments using x-ray resonant magnetic scattering (XRMS) and ferromagnetic resonance (FMR), these results will be presented elsewhere. The XMCD experiments were performed at beamline I1011 using total electron yield with 90% circularly polarized light, incident on the sample at an angle of 45°. The samples were magnetized in-plane with a magnetic field of 630 Oe, which is far above the coercive field of these samples, after which they were measured in remanence at room temperature. The dichroic difference was obtained after measuring the Co and Mn L2.3 edges with both left and right circularly polarized light. The samples are presented in Table I and are denoted according to the ratio between their CMG and RCS thicknesses. The number of layer repeats (x) is varied so that the total thickness is approximately equal. The samples were heated up between room temperature (RT) and 400° C, after which they were cooled down to RT before the measurements were performed.

In Fig. 1 we show the dichroic difference obtained at Co L_3 for sample S1.5 which has the highest ratio of

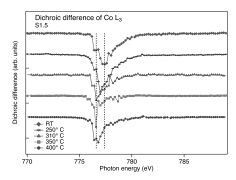


FIG. 1: XMCD difference spectra of Co L₃ obtained after annealing at various temperatures. The vertical lines corresponds to the energies plotted in Fig. 2.

CMG. A shift in the peak position is observed where two peak energies appears to have the strongest intensities, shown by dashed vertical lines. A similar shift is not found for the Mn L3 edge. The two samples with the smallest ratio of CMG, S0.66 and S1, have no remanence at RT after annealing to 310°C as evident from Fig. 2. In Fig. 2, we have plotted the magnitude of the dichroic difference at two energies (marked with vertical dashed lines in Fig. 1) at the Co L₃ edge and also difference at the Mn L_3 . The Co intensities (left axis) are shifted by 0.3 between samples for clarity and the Mn (right axis) is shifted by 0.45. It is readily seen that the Mn intensity follows the Co 777.4 eV intensity which would suggest that this peak corresponds to the bulk of the CMG layers. The intensity at this energy for sample S1.33, which has the thickest CMG layer, is 57% larger than for sample S0.66, which has the thinnest CMG layer. Sample S1 is intermediate between S1.33 and S0.66. Note that all three samples have 18 Å RCS. This discrepancy can be explained by a 6 - 8 Å magnetically dead layer in the CMG. Sample S1.5 which has a thinner RCS (12 Å) appears to have 1 nm thicker magnetically dead layer. This is well in accordance with XPS data recorded at several keV kinetic energy (not shown) which show a clear increase of intermixing of the RCS layer as it becomes thinner. Also, a sample with CMG(6 Å)/RCS(18 Å) showed no

dichroism by XMCD. In the overview spectra of all samples (not shown) it is observed that the Co and Mn intensities increase compared to the Cu lines for annealing temperatures of 310° C and higher, which indicates a migration of Co and Mn to the surface. This effect occurs at a surprisingly low temperature and it is confirmed by XRMS where the multilayer structure is practically destroyed after annealing to 400°C. This makes it difficult to analyze the Co component at 776.6 eV after annealing at high temperatures. After annealing to 250° C this component increases for the samples which become non-magnetic while decreases for

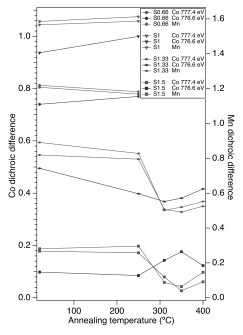


FIG. 2: Dichroic difference of both Co (left) and Mn (right). The intensities are obtained at the energies indicated by the vertical dashed lines in Fig. 1.

the other two. This could be due to a partially intermixed region near the interfaces where the Co is still ferromagnetically ordered.

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X-ray Absorption and Magnetic Circular Dichroism on Prussian blue analogues

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Molecular magnets has become a very active and increasingly exciting area of research lately. They constitute truly nanometric objects and massive numbers of totally identical entities can be easily produced by chemical synthesis techniques. These objects typically show magnetic hysteresis at low temperatures and magnetization reversal by quantum tunneling. They are attractive candidates for qubits, the quanta of information employed in quantum computing. ^{1,2} Another interesting characteristic of molecular magnets lies in the fact that the quantum coupling between pairs of molecules (needed to obtain the superposition of pure quantum states) can be controlled through the intermolecular interactions by adequately tuning the spatial arrangement of the molecules on a solid substrate.

Figure 1: Schematic representation of Prussian Blue molecules.³

Among the different classes of molecular magnets being investigated nowadays, Prussian blue and its derivatives deserve special attention. The pure compound, containing iron and cyanide ions (see Fig. 1), shows long-range ferromagnetic ordering at 5.6 K.⁴ However, Curie temperatures well above 300 K have been reported for several analogues containing also Cr ions.⁵⁻⁷ The magnetic behavior of these compounds can be explained using a localized orbital approach.⁸ They also show a high flexibility for tailoring their magnetic properties, since metal substitutions induce only small

lattice constant changes. From the electronic point of view, Prussian blue analogues are characterized by electronic delocalization between the metal sites across the bridging cyanide anion. The combination of charge transfer effects with the strong magnetic interactions mediated by the cyanide anion gives rise to interesting properties. For instance, reversible magnetization switching mediated by external agents such as irradiation with light, 9,10 temperature 11 or pressure, 12,13 among others, has been observed; electron transfer plays a key role in all these cases.

X-ray magnetic circular dichroism (XMCD) measurements at MAXlab are being used to study the electronic processes that determine the magnetic characteristics of these systems. The data depicted in Figure 2 were obtained at the octupole end of beamline I1011 station $K_{0.4}Fe_4[Cr(CN)_6]_{2.8}\cdot 16H_2O$ Prussian blue analogue containing Fe and Cr. The dichroic pair of spectra shown in this figure were measured at the Fe L-edge for normal x-ray incidence at room temperature. The Cr edge data (not shown here) exhibit more complex multiplet features, in agreement to earlier work.13 The samples measured were thin films grown by electrodeposition onto conducting substrates (Au-covered mylar). Their Curie temperatures lie around 200 K, as determined by Magneto-optic Kerr effect (MOKE) and SQUID measurements. For

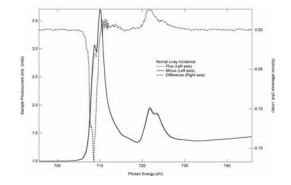


Figure 2: X-ray absorption spectra measured in total electron yield mode on a Prussian blue analogue containing Fe and Cr. Left: X-ray absorbed intensities at 300 K under an applied magnetic field of 0.28 T along the x-ray propagation direction. Right: Dichroic difference between the two absorption spectra. The degree of circular polarization is close to 0.8.

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this particular sample, MOKE ex situ measurements indicated the occurence of ferromagnetic order with a Curie temperature of 220 K. Still, a clear XMCD response is observed around 300 K under an applied magnetic field at the Fe L-edges. The Fe L-edge spectra shown in Figure 2 indicate a mixture of at least the "unisomerized" and the "metastable" state containing isomerized Fe^{II}-CN fragments. Our measurements clearly show the two components previously reported for the Fe L₃ white line, one around 707 eV and another one around 710 eV; nevertheless, this latter component seems to be more intense in our data, in contrast to the previous results. The XMCD difference (dashed line, right scale) which is obtained from the dichroic pair by reverting the magnetic field, presents similarities with both the "unisomerized" and the "metastable" states. Here at the L₃ white line the main contribution to the XMCD dichroic intensity appears to come from the first peak around 707 eV but being asymmetric with a broad cointribution towards the low energy side.

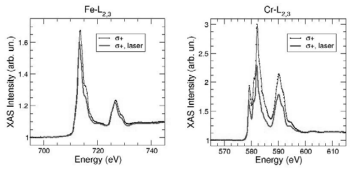


Figure 3: XMCD measurements on a $Fe_{II}Cr_{II}(Cr_{III}(CN)_6)$ Prussian blue analogue thin film showing the reduction in electron emission at the Fe and Cr L edges upon laser irradiation.

In a second set of experiments performed at BL I1011 the photomagnetic characteristics of these compounds have been explored. XMCD measurements at the Fe and Cr L adsorption edges were carried out on a thin film of an Fe_{II}Cr_{III}(Cr_{III}(CN)₆) Prussian blue analogue, with and without laser beam irradiation. Previous SQUID measurements on this type of samples have revealed a Curie temperature of 200 K for the non-irradiated sample, that rises up to 225 K under laser irradiation. A preliminary analysis of the XMCD measurements (presented in Figure 3) reveals a substantial reduction in the electron emission at both the Fe and Cr L edges when the sample is irradiated with a 5mW, 635 nm red laser. This is an effect that had not been reported before on these compounds; a detailed investigation of this behavior might thus provide some valuable understanding of the electronic mechanisms responsible for the photomagnetic response of these highly interesting materials.

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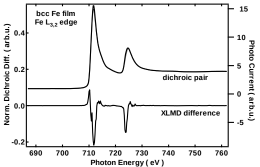
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Linear x-ray dichroism at BL I1011: first results on bcc Fe

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An element specific characterization of the magnetic properties is often based on Xray Magnetic Circular Dichroism (XMCD) spectroscopy in the x-ray absorption mode. Using XMCD one obtains an element specific measure of the spin and orbital moment on a per atom basis. At MAX-lab routine XMCD work has been possible since the late 1990's at the BL D1011 of MAX II using out of plane bending magnet radiation delivering sufficient circular x-ray flux in the soft x-ray range, in combination with a plane grating monochromator based beamline. Recently at the new BL I1011 it has been possible to perform XMCD experiments on a variety of systems both concentrated as well as dilute. This beam line uses a different concept, the x-ray source is an Elliptically Polarizing Undulator (EPU) which produces bright circular x-rays of variable helicity. [1] Here still a plane grating monochromator is used. [1] One advantage of using an EPU instead of out of plane bending magnet radiation is a better control of the x-ray polarization state. The EPU produces both $\sigma(+)$ and $\sigma(-)$ x-rays of a high degree of circular polarization. In the soft x-ray range of interest here, between 0.5 and 1 keV to cover the L-edges of the 3d transition metals, given the energy of the stored electron beam at the MAX II storage ring, the third harmonic is used, reaching a degree of circular polarization of order 80-90%. [1]



1.5 bcc Fe film Fe L_{3,2} edge

3 Photo Current

1.5 dichroic pair

1.6 dichroic pair

1.7 dichroic pair

1.8 dichroic pair

1.9 dichroic pair

1.9 dichroic pair

1.0 dichroic pair

Figure 1. Dichroic pair obtained for Fe L_{3,2} edge for bcc Fe film (right axis). Here linear light is used, the sample photocurrent is measured. The sample is at normal x-ray incidence, the electric field is rotated to be parallel and perpendicular to the magnetization. The XMLD difference (left axis) is normalized to the edge jump ratio for a comparison with the literature.

Figure 2. Dichroic pair obtained for Fe $L_{3,2}$ edge for bcc Fe film (right axis). Here circular light is used, with an angle of x-ray incidence of 45 degrees. The sample photocurrent is measured. The XMCD difference is plotted versus the left axis and the original spectra are plotted versus the right axis.

Furthermore, using the EPU as an x-ray source allows for still a different type of experiment, for which we present here first results. The EPU allows the user to produce linearly polarized x-rays with a variable polarization plane. Here we use linearly polarized x-rays in the plane of the storage ring and perpendicular to it. By comparing the x-ray absorption upon rotation of the electric field plane, by keeping the magnetic geometry of the sample fixed one can perform X-ray Linear Magnetic Dichroism (XLMD) experiments.

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XLMD effects in the x-ray absorption mode are much smaller which makes them much harder to measure. Earlier studies of this small effect in x-ray absorption often settled for the geometry at normal incidence for the sample and rotated the sample magnetization by keeping the electric field fixed. [2] Using this geometry for bulk Fe the normalized XLMD effect was found to be of order 8% for bcc Fe as compared to the XMCD effect which was found of order of 150%. [2] Despite its small intensity, the XLMD effect is worth measuring, as it is non-zero for anti-ferromagnets, scaling with the square of the magnetization. In comparison, XMCD scales linearly versus the sample magnetization and vanishes for an anti-ferromagnet. For ferromagnetic systems, it has furthermore been suggested that the XLMD signal also allows one to measure the orbital moment anisotropy. [3,4]

Here we focus on data taken on a bcc Fe film, sputtered in situ to remove surface contamination. The data of Figure 1, following earlier work in the literature, show that also at BL I1011, it is possible to obtain a non zero XLMD difference for a bcc Fe standard. To make sure that the small XLMD signal is not an artifact we fixed, as suggested earlier, the sample geometry at normal incidence and rotated both the sample magnetization as well as the plane of linear polarization using the EPU. One obtains in such a manner four spectra to compare in pairs and determine eventual experimental artifacts. Such a combination of experimental parameters is not easily obtainable using bending magnet radiation. In Figure 2 we show on the same sample an XMCD pair and the associated difference. Also here, four pairs of spectra at least were obtained, reverting both the magnetization and the light helicity by means of the EPU x-ray source. From the XMCD difference, in Figure 2, using the spin sum rule, a degree of circular polarization of 0.90(5) is obtained.

To allow for a comparison with the available literature data on Fe, we have normalized the XLMD difference by means of the high energy signal to background (the spectrum edge jump). We observe that the maximum XLMD difference is about 15 times less than the maximum XMCD difference once normalized to the spectrum edge jump. In the literature for Fe the correspondent figure is about 18 as given earlier [2]. Given the fact that this estimate is qualitative and does not take into account the differences in degrees of helicity or linear polarization between the present setup and literature values the agreement can be judged as satisfactory. The spectral shape of the XLMD difference obtained is not exactly the same as the one published earlier [2], however the literature data refer to Fe in multilayer structures which does not have the same local atom environment as the bcc Fe sample investigated here, so that differences in the spectral shape can be expected, as the local distortions will affect the XLMD signal [3,4].

In conclusion, we demonstrate here the possibility to measure the XLMD of bcc Fe. The small XLMD signal exploits fully the possibilities offered by the EPU at I1011 and allows for investigations also in anti ferromagnets.

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Interfacial effects in the magnetism of metastable fcc FeCu alloy thin films

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Fe is a very suggestive element with a rich magnetic phase diagram^{1,2} which underlines the tight relationship existing between magnetism and crystal structure, and thus allows us to manipulate its electronic and magnetic properties by appropriately tuning its chemical and structural environment.

This project was prompted by previous results such as the one illustrated by the XMCD-PEEM image presented in Figure 1. There, a 15 ML thick film of a Fe_{0.50}Cu_{0.50} alloy has been grown by UHV co-deposition³

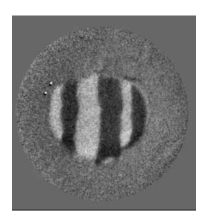


Figure 1: XMCD-PEEM image taken at 300K with circularly polarized light at the Fe L_3 edge on a 15 ML thick Fe_{0.50}Cu_{0.50} alloy film grown on Cu(111). The structure showing dichroic contrast at the image center is a Pb island. FOV: 4 um.

onto a Cu(111) single-crystal surface; a surfactant monolayer of Pb is employed to reduce the surface diffusivity of the deposited species and prevent their segregation.⁴ The structure visible at the image center is a Pb island, several nanometers thick. The magnetic contrast in this image was achieved using circularly polarized light tuned at the Fe L_3 absorption edge at the Nanospectroscopy beamline of synchrotron Elettra in Trieste (Italy); a clear pattern of in-plane magnetized domains can be observed at room temperature in the FeCu alloy film deposited on top of the bulk-like Pb island, whereas no dichroic signal is found in the area surrounding the island, where the same alloy film lies directly on top of the Cu(111) substrate.

The crystal structure of the pure Fe and FeCu alloy thin films has been characterized previously by means of low-energy electron diffraction. LEED I-V measurements demonstrate that the alloy films remain fcc for several nanometers in thickness, depending on their precise composition. The in-plane and out-of-plane interatomic distances in the alloy are the same, irrespective of whether the film is grown on the Cu(111) substrate or on the Pb island. These observations thus suggest that the presence of ferromagnetic order in the alloy film, which occurs only when it is grown on the Pb but not on the Cu substrate, must be of electronic origin. The purpose of the experiments carried out at

beamline I1011 of MAX-lab, was to use x-ray magnetic circular dichroism to measure the magnetic moments and anisotropy at the Pb/alloy interface, to determine the influence of the Pb substrate on the magnetic properties of the Fe grown directly on it.

The experiments were carried out at the UHV end station of beamline I1011, using a Pb(111) single-crystal as substrate. The FeCu alloy films were grown in situ by simultaneous co-deposition from previously calibrated electron-bombardment evaporators; the deposited thicknesses were cross-checked by measuring the jump ratio J_R in the XAS spectra at both the Fe and Cu $L_{2,3}$ absorption edges.

Rather strikingly, XMCD measurements with circularly polarized x-rays revealed that the Fe_{0.50}Cu_{0.50} alloy films already display dichroic contrast at submonolayer Fe coverages at a temperature of 100 K. Figure 2 shows a representative spectrum measured at 45° incidence at the Fe L_{2.3} edges; the alloy film had a total thickness of

0.8 ML (0.4 ML equivalent of Fe) and was grown at room temperature on the clean Pb(111) surface; after that, the sample was cooled down to 100 K for the measurements and magnetized by applying a magnetic pulse of ~650 Oe. A clear in-plane dichroic asymmetry (shown in the lower panel of the image) can be observed. Similar measurements were performed for different alloy film thicknesses, both at in-plane and out-of-plane geometries in order to probe the magnetic anisotropy at the interface between the alloy and the Pb substrate.

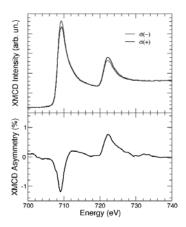


Figure 2: XMCD spectrum (above) and dichroic asymmetry (below) measured at 100 K at the Fe $L_{2,3}$ edges on a 0.8 ML thick $Fe_{0.50}Cu_{0.50}$ alloy film on Pb(111).

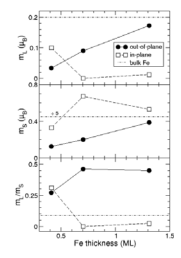


Figure 3: Orbital and spin moments and their ratios, in-plane and out-of-plane, as derived from XMCD measurements on $Fe_{0.50}Cu_{0.50}$ alloy films of different thicknesses at 100 K.

From this series of spectra the orbital and spin components of the magnetic moment have been determined by application of the XMCD sum rules.⁶ The resulting values are summarized in Figure 3. It is interesting to note that for

are summarized in Figure 3. It is interesting to note that for 0.8 ML of alloy the magnetic anisotropy seems to favor in-plane magnetization, turning out-of-plane for film thicknesses above 1 ML. Besides the XMCD measurements, some EXAFS scans were measured at the Cu L₂₃ edge to check for possible structural changes localized at the Pb/alloy interface. These measurements are currently under study.

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Time resolved XMCD combined with FMR study on a Fe₂₀Ni₈₀ single film

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INTRODUCTION

Spin dynamics in magnetic materials plays an increasingly important role in the study of spintronic devices [1]. Ferromagnetic resonance (FMR) has been employed for several decades to investigate important fundamental phenomena in magnetism such as gyromagnetic ratios, damping, relaxation, and anisotropy energies. In modern magnetic materials, e.g. magnetic thin films and multilayers, this method has been developed to study collective spin reorientation transitions and interlayer coupling. FMR provides information on the dynamical properties of a material. Complimentary information can be obtained by X-ray magnetic circular dichroism (XMCD), a core-level spectroscopic technique that provides information on the chemical constituents. Stroboscopic probing of the magnetization using XMCD gives access to the time domain and by combining this with FMR a versatile technique has been obtained that allows time resolved measurements on the forced magnetization precession in a Permalloy thin film with element specificity.

METHODS

A 20 nm film of polycrystalline Permalloy was deposited onto 100 nm thick SiN membrane supported by a 0.5 mm thick Si frame. The frame is adhered to a coplanar waveguide (CPW). The CPW is made of Au-plated laminate with a 0.500 mm wide conductor for the RF-signal and a clearance hole of 0.250 mm in the center conductor. The CPW is designed with a shorted end, allowing for measurements of the reflected RF-signals.

This study was carried out at MAX-II, MAX-Lab, at the soft x-ray beam line I1011. The beam line is equipped with an octupole magnet end-station designed to generate fields up to approximately 1 T in arbitrary directions.

The time resolved measurements rely on harmonic generation of microwaves that are phase locked to the electron bunches of the synchrotron ring. The timing is controlled by passing signals from the bunch clock through a delay generator. A comb generator generates a harmonic spectrum of the 100 MHz input signal and a sequentially connected tunable band pass filter (BPF) is used to select a suitable harmonic. This output signal is amplified and passed through a directional coupler, which directs the signal to co-planar waveguide onto which the sample is mounted. The reflected signal is passed to an RF diode for conventional FMR measurements. The harmonic microwaves are used to excite the precession of the magnetization in the sample. X-rays are transmitted through the sample as well as the tailored membrane and detected by photodiodes. The octupole vector electromagnets produce a bias field \mathbf{H}_{A} , oriented orthogonal to both the driving field \mathbf{H}_{driv} and the incident light.

RESULTS

A phase locked RF signal with frequency of 1.1 GHz was used to excite magnetic precession in the Permalloy sample. Figure 1 (top) shows in-situ FMR. A magnetic bias field was scanned between 0 and 1 A (horizontal axis) and the reflected differential response from the sample was measured (vertical axis). A clear resonance is indicated by the zero crossing about 0.35 A. This zero crossing increases linearly with the square of the frequency. The bottom part shows a single time-resolved measurement at the resonance point, using x-rays at the Fe L3 edge. Oscillations are observed, with a period of 910 ps. Despite the fact that the beam line was not fully operational due to misalignment

during our first dynamic measurement, we were still able to use the bunch clock of the synchrotron ring to generate a phase locked RF-signal to drive precession of the spins in a Permalloy film.

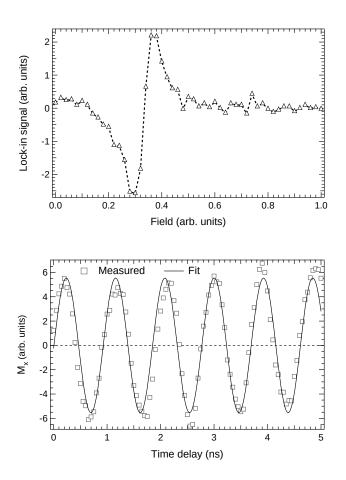


FIGURE 1: (Top) Conventional FMR measurement, performed in-situ at the i1011 beamline, MAX-Lab on a Permalloy thin film, 20 nm thick deposited on a SiN membrane. The excitation frequency is 1.1 GHz and phase locked to the synchrotron bunch clock. The graph shows the differential reflectivity response of the RF-signal (vertical axis) as a function of applied current to the electromagnet (horizontal axis). The resonance can be found at the zero-crossing, occurring at about 0.35 A. (Bottom) A time resolved scan showing the intensity of the transmitted photons at the Fe L3 edge (vertical axis) versus the time delay.

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Investigation of Magnetic Properties of the Multiferroic Layer System Cobalt on Bariumtitanate

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Systems with multiferroic properties are of interest for research since the 1960s. ^[1] In the beginning single phase multiferroics were in the focus of research, whereas nowadays systems with a ferroelectric and a ferromagnetic component are investigated. ^[2] The work reported here covers the layer system Cobalt on Bariumtitanate (001) within SFB 762. In particular, XAS measurements of 3, 6, 9, 12, and 21 monolayer thick Co films prepared by e-beam evaporation were taken into account. The films show in-plane magnetization starting from 6 ML.

At the Co L-edge X-ray magnetic circular dichroism (XMCD) experiments had been performed at beamline I1011 at MAX II to determine the magnetic properties.

In Fig. 1 we show a raw sum spectrum of 6 monolayers Cobalt on Bariumtitanate. It becomes obvious that the calculation of the magnetic properties applying the sum

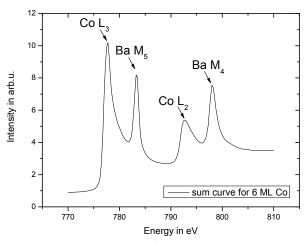


Figure 1: Raw XAS at the Co $L_{2,3}$ -edges and the Ba $M_{4,5}$ -edges for 6 ML Co on $BaTiO_3(001)$.

rules cannot be perfored easily by using the integral of the sum curve since there is a great influence of the Barium. In Fig. 2 the difference curve for the same system resulting from measurements with opposite helicity is shown. In signal there influence of the Barium peak leaving the opportunity integrate it directly for the calculation of the magnetic moments.

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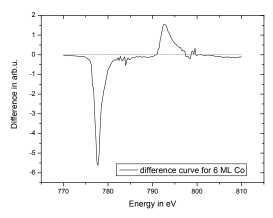


Figure 2: Difference curve for 6 ML Co on BaTiO₃(001).

In order to resolve the problem with the sum spectrum the thickest film of Cobalt was used as starting point. There is no Barium signal anymore so that this spectrum can be used in a scaling procedure. The sum curve of the 21 monolayers is normalized to the intensity of the sum curve at the certain film thicknesses and integrated after subtracting a step background. In Fig. 3 the quality of this scaling procedure is shown.

After the described steps it was possible to calculate the magnetic moments. These calculations gave total magnetic moments in the range of known values for Co (hcp) bulk reported in literature, which are around 1,63 μ_B to 1,78 μ_B . [3], [4]

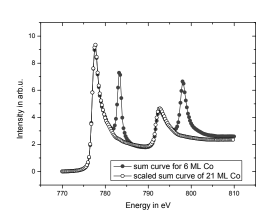


Figure 3: Sum curve for 6 ML Co and scaled sum curve for 21 ML Co on BaTiO₃(001) to demonstrate the vaildity of the scaling procedure.

However, there are increased angular magnetic moments for all observed film thicknesses. In literature, there are several cases – even for other elements – where this increase is observed in thin films. [5], [6]

Changes to the Bariumtitanate substrate by heating in vacuum up to 600 °C prior to Co deposition, resulting in O vacancies and increased conductivity, do not yield significant differences for the magnetic moments.

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XMCD studies of the MnSb inclusions formed in GaSb matrix

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In the pursuit of novel ferromagnetic materials, which can be used in spintronic devices, heterostructures created by introducing ferromagnetic inclusions in a semiconductor matrix became promising. In this kind of material, small ferromagnetic nanoparticles are immersed in the semiconductor host lattice providing a built-in local magnetic field. Main advantage of MnSb as a material to form such precipitations is its high Curie temperature (T_C). For the $Mn_{1-x}Sb_x$ layers grown on GaAs substrate a T_C can reach 620 K [1]. What's more, studies on the MBE grown thin layers showed that already in pure material MnSb does not form smooth surface but tends to form big islands or columns [2]. This leads to the other advantage which is possibility to form the hexagonal MnSb precipitates directly during the MBE process without subsequent annealing.

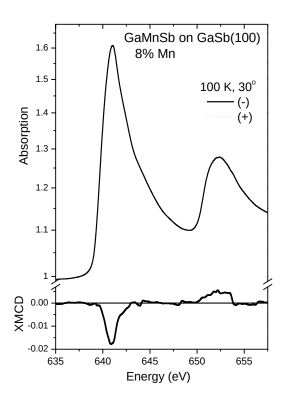


Fig. 1) The XMCD and signal XANES spectra of Mn $L_{3,2}$ edges measured at the LN_2 temperature for opposite polarities of magnetic field with X-ray beam at 30°.

The MnSb hexagonal inclusions were formed directly the epitaxial procedure without the post growth annealing. The GaSb(100) and GaAs(111)A substrates were used for the MBE growth of the GaSb:MnSb layers. In order to check how the type of substrate influences the dots' formation, the samples on both substrates were grown in the same process. First, the GaSb buffer of 40-45 nm was grown and then the GaMnSb layer. The samples were grown at 450°C with different nominal Mn content.

The X-ray Magnetic Circular Dichroism (XMCD) experiment was performed at the beamline I1011. X-ray absorption spectra were collected in the Total Electron Yield (TEY) mode by recording the sample photocurrent. Since the TEY detection mode is sensitive to the near surface region, the samples were Ar⁺ sputtered in situ in order to remove oxides and other impurities from the surface. The measurements were carried out on the remanently magnetized

samples. The magnetic field used to magnetize samples was around 0.035 T and it was oriented along the X-ray path. The samples were measured at room temperature (RT) and at 100K (LN₂). At the RT the magnetic field was too weak to order the domains. The dichroic signal was registered at LN₂ only.

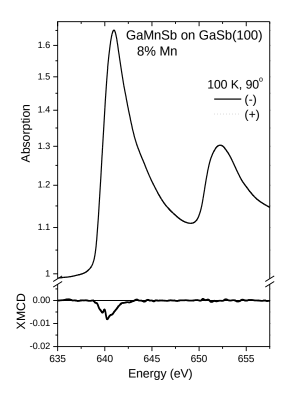


Fig. 2) The XMCD and signal XANES spectra of Mn L_{3,2} edges measured at the LN₂ temperature for opposite polarities of magnetic field with X-ray beam at 90°.

In case of the samples grown on the GaAs(111)A substrate, the obtained XMCD signal was very weak, probably due to the large blocking energy and weak magnetic field available at the station which was not sufficient to magnetically order the sample.

For the samples grown on the GaSb(100) substrate the obtained **XMCD** signal was more pronounced. The results for the GaSb:MnSb sample with nominal Mn concentration equal to 8% are presented in Figs. 1 and 2. The signal shown in Fig. 1 was measured in the grazing incidence, with the X-ray beam at 30° to the surface. sample The spectra gathered from the same part of the sample but measured in the normal incidence, with the X-ray beam perpendicular to the sample surface, are shown in Fig. 2. In both cases the scale was left the same in order to facilitate the comparison. It can be noticed that the XMCD signal shows the angular dependence. This observation agrees with the measurements carried out with a superconducting

interference device magnetometer which revealed that for this kind of samples the easy magnetization axis is in the sample surface plane. The results of performed studies are summarized in the paper [3].

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Creation of long-lived Frenkel defects via hot electron-hole recombination in pure NaCl crystals at 12 K

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High radiation resistance of many wide-gap materials (WGMs) on the basis of binary and complex metal oxides can be explained by the fact that the adiabatic formation energy ($E_{\rm FD}$) of a pair of Frenkel defects (FD) significantly exceeds the energy gap, $E_{\rm FD} > E_{\rm g}$. In such WGMs, the energy released at the recombination of holes and cold (relaxed to the bottom of a conduction band) electrons is insufficient for FD creation. However, the recombination of hot (non-relaxed) conduction electrons with holes can tentatively cause the creation of FD even in WGMs with $E_{\rm FD} > E_{\rm g}$ [1]. This hypothesis is experimentally confirmed for highly pure NaCl single crystals. The creation spectrum of long-lived F-H pairs has been measured at 12 K using synchrotron radiation of hv = 7-40 eV at the undulator beamline I3 at MAX-III.

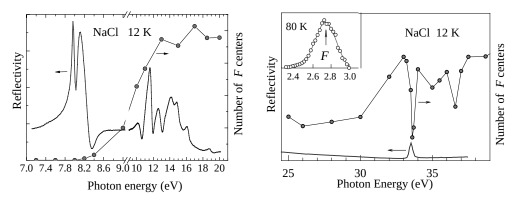
The sample was irradiated perpendicular to the (100) plane by a prescribed number of photons at each of several energies. The reference signal for normalization to equal quantum irradiation dose was recorded from a sodium salicylate coated mesh. The irradiated crystal was then stimulated through a double prism monochromator by 2.7-eV photons along the [100] direction in the maximum of the absorption band of radiation-induced F centres (an electron in a field of an anion vacancy). At 12 K, the 2.7-eV photons cause the excitation (not ionization!) of F centres. In thirty seconds after the end of NaCl irradiation, the time dependence of photostimulated luminescence (PSL) selected through a combination of optical filters (3-5 eV) was recorded. The light sum of this rapidly damping PSL was taken as a measure of F-H pairs formed by SR. This low-temperature luminescent method was elaborated at the Institute of Physics (Tartu) [2] and later used for the measuring of the creation spectra of F and α centres (an anion vacancy) in KBr and KCl crystals at beamline BL52, MAX-I [3]. In highly pure KBr and KCl crystals with $E_{\rm FD} < E_{\rm g}$, FDs are efficiently created at both room temperature and at 4-10 K due to the decay of anion excitons or the recombination of relaxed conduction electrons with self-trapped holes. The first creation spectrum of F centres by VUV-radiation in pure NaCl at 300 K was measured already in 1964 [4]. Selective photo-formation of either anion excitons or separated electrons and holes via interband transitions causes the creation of F centres. The annealing of these F centres occurs in the same temperature region as that of F and V₂ centres induced in similar samples by Xrays or fast fission neutrons [4]. However, the creation efficiency of FDs decreases by dozens of times at 10 K, thus impeding the detection of long-lived F-H pairs in highly pure NaCl crystals at low temperatures.

According to our experimental results (see figures 1 and 2), the exciting photons in the region of exciton absorption (7.8–8.4 eV) do not cause the creation of F centres. A very weak PSL (probably impurity-related) is detected in the spectral region slightly above the energy gap ($E_g = 8.9 \text{ eV}$), when s-conduction electrons are formed. However, a sharp rise of the PSL light sum takes place when the exciting photons of hv = 12-16 eV form already hot d-conduction electrons, but do not yet cause the formation of secondary anion excitons via the

multiplication process [5,6]). Therefore, the energy released at the recombination of *d*-conduction electrons and *p*-valence holes is sufficient for the creation of stable F-H pairs with a relatively large interdefect separation. The PSL arises via the following reaction

$$F...H + h v_{\text{stim}} \rightarrow F^*...H \rightarrow \alpha...I + h v_{\text{emis}}$$

while an α -I pair, formed during this recharging process, undergoes non-radiative recombination with the reconstruction of a perfect lattice. The number of F-H pairs (i.e. PSL light sum) changes only slightly with the rise of the exciting photon energy from 12.5 to 27 eV (see figs. 1 and 2), while the creation efficiency of FDs doubles at 28-33 eV. According to our previous data on the multiplication of electronic excitations in a number of alkali halides, such photons can form up to three electron-hole pairs in NaCl [5]. However, only two hot e-h pairs contribute to the creation of stable F-H pairs.



Figs. 1 and 2. The reflection spectra and creation spectra of F-H pairs measured at 12 K for a highly pure NaCl single crystal. Inset shows the stimulation spectrum of PSL at 80 K.

Of particular interest is the creation of F centres by 33-34 eV photons that form cation excitons. It was shown earlier [5] that a cation exciton (\sim 33.4 eV) decays into an anion exciton and several e-h pairs. The energy excess of about 24 eV remained after the formation of an anion exciton is enough to form two e-h pairs, but only one of them will be sufficiently hot for the creation of a pair of FDs. A deep dip of the creation efficiency of F-H pairs is observed in the region of cation exciton absorption. This dip remains even if the reflection losses (narrow reflection peak at 33.4 eV – see figure 2) are taken into account. So, only a hot e-h pair, formed together with an anion exciton and a cold e-h pair at the decay of a cation exciton, is able to create an F-H pair at 12 K. Concluding, the recombination of a hot conduction electron with a hole causes the creation of a pair of anion FDs in a NaCl crystal with $E_{\rm FD} > E_{\rm g}$ (as it was already suggested for WGMs in [1]).

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Absorption cross section measurements of CO₂ isotopologues ¹³CO₂ and C¹⁸O₂ in the wavelength range 150-187 nm

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Introduction

VUV photolysis of CO_2 is important for planetary atmospheres. Preferential photolysis of one isotopic species over another leads to isotopic enrichment. We conducted measurements of the absorption cross-sections of isotopically enriched samples, $^{13}CO_2$ and $C^{18}O_2$, in the wavelength range 150–187nm.

It has been suggested that (nuclear) spin (electronic) orbit coupling may give rise to preferential photo-dissociation of isotopic species with non-zero nuclear spin, such as 17 O (5 2) or 13 C ($^{-1}$ 2). Photo-dissociation experiments, CO₂ + $hv \rightarrow$ CO + O, conducted with a mercury lamp (185nm) led to products with 17 O enrichment of more than 100% [1]. A similar effect could, however, arise from a weak vibronic structure in the absorption cross-section. We aim to investigate the mechanism responsible for the anomalous enrichment.

Experimental Set-Up

Measurements were performed on the gas phase branch of the I3 beamline. The beam was passed through a pressure-monitored sample cell and collected at a PMT detector. Background measurements were collected on the evacuated cell. Commercially produced isotopically enriched samples, ¹³CO₂ produced by Fluka (99% atom ¹³C) and C¹⁸O₂ from ICON (85% atom ¹⁸O), were used at pressures between 2–150 mbar. As background and sample could not be measured simultaneously, measurements were conducted in 10nm blocks alternating between background and sample.

Preliminary Results

Recorded sample and background intensities, I and I_0 respectively, were corrected for decaying beam current. The absorption cross-section, σ , was calculated following,

$$\sigma = \frac{\log_{10}(I_0/I)}{n \cdot I} \tag{1}$$

where *n* represents the gas concentration [molecules·cm⁻³] and *l* the length of the sample cell [cm]. Where multiple measurements were available, the average intensities were calculated

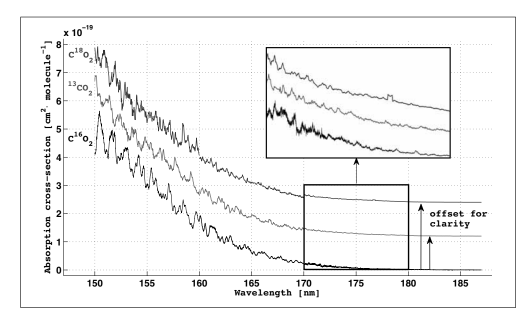


Figure 1: **Absorption cross-sections.** Reference CO_2 (bottom) [2, 3]; measured $^{13}CO_2$ (middle) and $C^{18}O_2$ (top). The upper spectra have been offset from CO_2 (bottom) for clarity.

for each isotopologue. Drift between measurement regions was corrected for each measurement interval: calculated cross-sections were multiplied by a factor to give the same slope as a linear fit to CO₂ reference data [2, 3], and then shifted to give the same mean. Weak vibronic structure is seen that shifts in a regular way with isotopic substitution, Fig. 1.

Acknowledgements

We thank the MAX-lab staff for their support and assistance.

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Investigation of VUV transitions of iron group elements for astrophysical applications using I3 at MAX III

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Abstract We are developing an experimental station at the gas-phase branch line at the 6.65 m normal incidence beam line at I3, MAX III. It is used to investigate free metal atoms and ions in plasma phase. The sample ions are produced in a hollow cathode discharge lamp, which is illuminated by the synchrotron radiation producing absorption when in resonance with the atomic lines. The synchrotron light is scanned in energy building up an absorption spectrum. Contributions from intrinsic light of the discharge are minimized through geometrical, spectral and temporal filters.

The project's major goal is to measure transition rates for specific lines in the spectrum of ionized iron, Fe II lines, at vacuum UV wavelengths, 100-130 nm. These transitions are important for many astrophysical applications, e.g. in objects where line fluorescence is prominent, as well as stars observed in the ultraviolet *FUSE* and *HST/STIS* region. In this wavelength region many other iron-group element ions have important transitions, which will be measured in future projects.

Iron lines are of great importance in analyses of stars and nebulae, not only for abundance studies, but also for diagnostics of the plasma conditions and derivation of physical properties such as temperature, electron density and radiation field.

We describe the experimental setup at the I3 beam line at MAX III, and the early results.

Introduction In spectroscopic studies of astrophysical objects iron is an important tracer, and its relatively high cosmic abundance and complex atomic structure result in many spectral lines. Iron is often used as a probe of metal abundance in distant objects, such as quasars and Active Galactic Nuclei (AGNs). Many stars and nebulae have representative temperatures around 10 000 K, and for this temperature the favored ionization stage is singly-ionized iron, Fe⁺, whose spectrum is denoted FeII. The lines of FeII are thus of great importance in analyses of stars and nebulae.

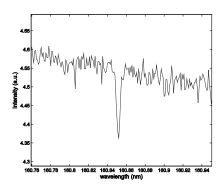
Fluorescence lines can be used to probe active plasma regions. Since these lines are optically thin, they offer more possibilities for plasma diagnostics than most other lines, which in general are low-excitation lines and therefore optically thick. To model the radiative pumping process, a number of atomic parameters need to be known, among them the transition rates (*A*-values) or oscillator strengths (*f*-values).

The present project aims at measuring the intrinsic line strengths for these transitions by letting the synchrotron light pass through a plasma hosting the ions. When proved successful this technique will enable us to measure a wealth of lines important for several astrophysical applications.

Experimental setup Synchrotron light from MAX-III is used as a background source to study the absorption from gas-phase iron atoms. The plasma responsible for the absorption is produced in a hollow-cathode (HC) discharge, which is run with neon as carrier gas. The electronic transitions in iron produce narrow absorption lines in the continuous light from the synchrotron. Comparisons of different absorption lines from a common lower level give the relative internal strengths for the lines,

and the values are put on an absolute scale by using known reference lines. The 6.65m high spectral resolution (R>100,000) normal incidence monochromator (NIM) on the I3 beamline is used to produce monochromatic light, and with its performance in terms of resolution and photon intensity ideal for the purpose. The energy of the monochromator is scanned over the transition to obtain the absorption line. Since the intrinsic width of the absorption line, determined by the thermal motion of the atoms in the plasma, is of the same order magnitude we need the full resolution to obtain reliable and accurate results.

The HC producing the plasma is a source of strong line emission in the same spectral region, which is orders of magnitudes stronger than the synchrotron light studied. This emission light is removed by modulating the synchrotron light with a frequency of up to 1000 Hz. The signal from the photomultiplier tube (PMT) and the chopping frequency are fed into a Lock-In amplifier to extract the absorption signal in the synchrotron light. In addition, geometrical and low-resolution spectral filters are used to suppress the light from the discharge.



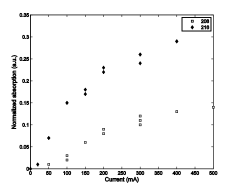


Figure 1: *left*: Absorption line from ionized iron, Fe II, at 160.85 nm. *right*: Dependence of the absorption as a function of current through the discharge for two Fe I lines from the same lower level but with different intrinsic line strengths (*f*-values). The behavior is clearly non-linear, and the ratio of the two lines depends on the current.

Early results. We are pushing the measurements into the VUV region. Absorption originating from the ground level of Fe II at 160 nm (Fig1a) has been measured at different currents. Lines from the same lower level a ${}^6D_{9/2}$ at shorter wavelengths (120 nm) are however not observed.

Measurements under changing experimental conditions are important for the investigation of saturation effects, and the correction for the same in the derivation of the true intrinsic line strengths. Figure 1b shows the dependence of the absorption as a function of current through the discharge, for two transitions, from the same lower level but with different strength. The saturation is clearly observed as a non-linear dependence. The saturation is dependent of the strength of the line, and varies between the lines.

Acknowledgment: We are grateful to the MAX-lab personnel, in particular Mats Leandersson and Balu Thiagarajan, for providing superior experiment environment and excellent user support. We acknowledge the support from the Swedish Science Council through grant 621-2006-3085, and the financial support from Royal Physiographical Society (Fysiografen) in Lund.

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Top of the valence band of the GaAs(100)-c(4x4) surface

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Numerous surface reconstructions can be prepared on GaAs(100) surface by molecular beam epitaxy (MBE). Among others, the c(4x4) reconstruction represents a stable, technologically important atomic arrangement and its surface properties has been investigated for many years [1-4]. In spite of it, there are still unclear points between theoretical and experimental results. One of them is the existence of a surface state around 0 eV of binding energy, i.e. in proximity to the valence band top. To clear it, we have studied the GaAs(100)-c(4x4) β surface state near the top of the valence band using enhanced both energy and angular resolution at BL I3. In the same time, this measurement could provide a necessary testing on known sample surface.

To this aim, the GaAs(100)-c(4x4) β reconstruction was prepared in SVT MBE growth chamber attached to the BL I3. After UHV sample transport and a LEED check, the photoelectron spectra have been measured by the VG Scienta R4000 electron analyzer. Experimental conditions for recording the photoelectron spectra were the following: energy of

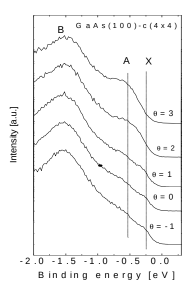


Fig.1 Photoemission spectra measured in direction [1-10] round about surface normal. Energy of the synchrotron radiation was 21.2 eV.

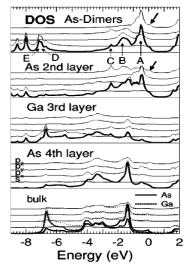


Fig.2 Density of electronic states at Γ point for GaAs(100)-c(4x4) β with resolved orbital contributions (thin lines) from four topmost atomic layers. Thick lines are for total DOS [5].

synchrotron radiation 21.2 eV and angle of incidence of the synchrotron light 73° with respect to the surface normal. Photoelectrons were collected from the acceptance angle +/- 15° . The photoelectron spectra were recorded with a step of 1° in the [1-10] direction.

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Fig. 1 presents the photoemission spectra near the top of the valence band around normal direction, showing two shoulders (A) and (X). The shoulder at around 0.6 eV corresponds to the position of surface state, detected in previous experiments [1,3], and is also found theoretically [5]. The other shoulder at ~0.2 eV below the top of the valence band has not

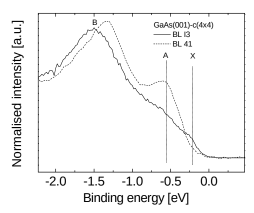


Fig. 3 Comparison of the normal photoemission spectra of the GaAs(100)-c(4x4) surface, measured at BL I3, and BL41 [1]. Excitation energy 21.2 eV.

been experimentally observed before and can well be assigned to a position in DOS spectrum in Fig. 2, marked by oblique arrows. According to this calculation the signal may be composed of contributions from p_z orbitals on As atoms in surface (As dimers) and on As atoms in subsurface layer.

However, the intensity of the measured surface state around 0.6 eV is not so clear as compared with earlier experiment [1] (Fig. 3). In this connection it is to mention that the studied GaAs(100)-c(4x4) sample was the very first growth performed in

the new MBE apparatus and the quality of the prepared surface might be influenced by this fact, in spite of perfect RHEED and LEED patterns taken during and after the growth. To verify the existence of the surface state near 0 eV of binding energy, supplementary experiments have to be done at BL I3.

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UV-induced photofragmentation of 1,1,1,2-tetrafluoroethane (CF₃-CH₂F)

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Electron-ion coincidence (PEPICO) methodology is nowadays often employed for studies of various molecular systems [1-2], as it provides means to separate electronic processes via their link to specific ions. The combined measurement of electron spectra and ion production can be used to obtain high resolution information about molecular processes. The ion data may be used to correlate fragments and, depending on the detection method, gain additional information about the dissociative processes. These experimental studies are also applicable to many theoretical studies such as the verification of molecular modeling calculations, and relevant applications exist among different fields of research such as atmospheric sciences, where breakdown of molecules can have large scale effects. The electron spectroscopy group of University of Oulu and collaborators, especially at University of Turku, have performed several PEPICO studies [3-5] with synchrotron radiation provided by MAX-lab, in UV and soft X-ray regimes.

Fluorinated ethanes such as hydrofluorocarbons (HFCs) were found as an alternative to chlorofluorocarbons (CFCs) [6] due to the greater ozone-depletion potential of chlorine compounds with respect to fluorinated compounds, which lead to CFCs being phased as a result of the Montreal Protocol on Substances That Deplete the Ozone Layer. CFC and HFC substances have been and are used as, for example, refrigerants, propellants, blowing agents and solvents. In addition, CFCs and HFCs have been subject to research for their Global Warming Potential (GWP), which in turn is likely to affect their future usage even more. In this work, we have examined the photofragmentation of the 1.1.1.2-tetrafluoroethane molecule in its gaseous form. We consider this to be important, especially in the UV range, which may provide useful information of real-life chemical reactions that these substances undergo in the atmosphere through UV ionization by sunlight. Previous work exists for tetrafluoroethane, where the fragment production has been inspected by following the cationic yields as a function of photon energy with TPEPICO. [7]

The experiment was conducted at the gas-phase branch – FINEST [8] – of the I3 beamline UV at MAX-lab. The gas-phase sample was radiated with photons of 30 eV for the PEPICO data. In addition, more detailed time-of-flight spectra were obtained as a function of photon energy, and the inner and outer valence photoelectron spectra were recorded. The data

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Fig. 1. Photoelectron spectra of 1,1,1,2-tetrafluoroethane measured with three different photon energies. [9]

processing is in progress, and we are approaching the study with the aid of molecular *ab initio* calculations. Fig. 1 displays the recorded photoelectron spectra, and Figs. 2 and 3 depict the coincidence data. As compared to previous studies [7], we have identified smaller fragments, and thus we are able to form a more complete picture

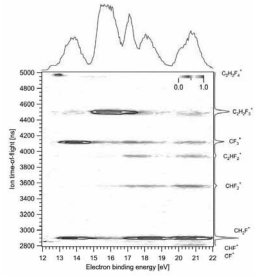


Fig 2. A PEPICO map of 1,1,1,2-tetrafluoroethane's valence region. [9]

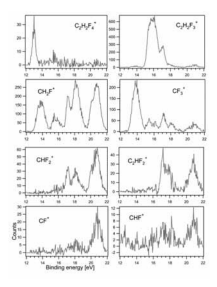


Fig 3. Coincident ion yields (CIYs) for the fragment cations observed with valence photoionization ($h\nu = 30 \text{ eV}$). [9]

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Size selective study of the Se microclusters induced by direct vacuum evaporation

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The chemistry of the chalcogene clusters like Se is an area of growing relevance because of their ability to form polyatomic complex chains and rings of a variety of sizes and shapes. These structural preferences influence the physical properties of their microclusters[1, 2]. Here is presented the results of experimental study of the ionization and fragmentation of Se_n clusters (n<9) after Se valence photoionization, using the Partial Ion Yield (PIY) and PhotoElectron-PhotoIon COincidence (PEPICO) techniques.

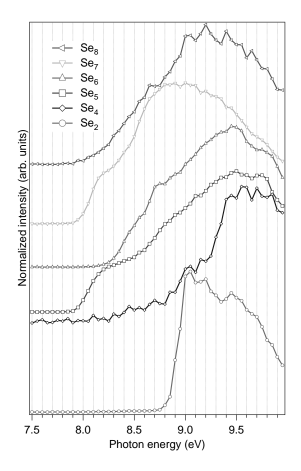


Figure 1. The PIY spectra of Se_n clusters measured at the valence region threshold .

The measurements were performed at FinEst branchline of the beamline I3 at MAX-III. In this work have been used the experimental PEPICO setup which consists of a modified Scienta SES-100 electron analyser and a homemade Wiley-McLaren type ion TOF detector [3]. The Se sample was evaporated into the interaction area of the spectrometers from the boron nitride crucible (T \approx 235 °C) of effusion cell . The base pressure of the system was below 6 \times 10⁻⁷ mbar and the measurements were performed at room temperature.

The normalised partial ion yields of Se_n clusters measured in the photon energy range from 7.5 eV to 10 are shown in the Figure 1. In the photon energy range up to 10 eV have been detected the Se clusters Se_2 , Se_4 , Se_5 , Se_6 , Se_7 and Se_8 . However, the PIYs of Se_4 and Se_8 clusters have very weak signal on absolute scale. The PIYs show strong dependence on the size of cluster.

In Figure 2 are represented the PEPICO maps of Se_n clusters in coincidence with Se valence photoelectrons. The detected ionised clusters are marked on the maps and the patterns have been recorded in two adjacent energy windows ($E_{center} = 9$ and 12 eV) of the electron spectrometer. The ionizing photon energy was chosen equal to 22 eV. The appearance of coincidence patterns of Se_1 , Se_2 , Se_3 and Se_4 cations in higher binding energy range refers to the fragmentation of larger clusters.

Further data analysis is in progress and a systematic discussion of electronic structure and fragmentation of Se microclusters will be published as a regular article.

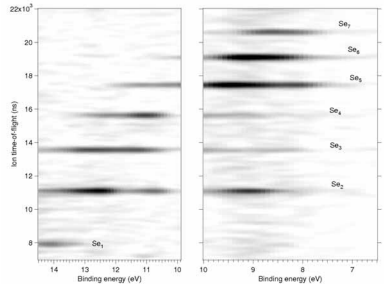


Figure 2. PEPICO maps of selenium clusters recorded in coincidence with Se valence photoionization.

This work was supported by Nordforsk Infrastructure Network "Advanced spectroscopy using MAX-laboratory in Lund" and the EU "Transnational Access to Research Infrastructures" programme. Financial support from the Academy of Finland is also acknowledged. The staff of MAX-lab is acknowledged for their help during the experiments. We thank the Electron Spectroscopy Group of Oulu University for the opportunity to share the experimental equipment.

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Angle-resolved electron momentum imaging in the vicinity of the surface Plasmon resonance of C_{60}

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In the last decades collective electronic properties of C_{60} have attracted continuing interest. The cloud of the outer-most valence electrons of carbon atoms distributed "uniformly" on the surface of a hollow sphere exhibits surprising behavior upon photoemission. Historically a giant surface plasmon resonance (SPR) was discovered in 1992 above the ionization threshold, near 20 eV, through measurements of the photoionization cross-section [1]. This feature was interpreted as a collective oscillation of delocalized electrons relative to the ionic carbon cage. Recently, a second collective resonance, associated with a dipole-excited volume plasmon, has been observed experimentally near 40 eV [2]. The study of collective excitation of electrons, or plasmons, in such model system enables to understand electronic correlation and coupling effects in systems with a large number of interacting fermions, especially in C_{60} , where the collective excitation can relax through various channels, like ionization and/or fragmentation.

In a recent publication [3], we demonstrate in the framework of the TDLDA (Time Dependent Local density Approximation) as well as with a semiclassical model that the nature of the SPR in C_{60} can be studied using angle-resolved momentum electron imaging. We show that while the imaginary part of the complex polarizability, $\text{Im}[\alpha]$, is related to the photoabsorption cross-section, the real part, $\text{Re}[\alpha]$, is associated with the energy-dependent angular distribution. In particular, we expect a drastic variation of the photoelectron angular distribution (characterized by its asymmetry parameter β) in the vicinity of the SPR. This variation is related to the correlated electron motion. Therefore, we are interested in the investigation of this resonance using the velocity map imaging technique [4].

Experimental setup

The experiment was carried out at the gas phase branch of the beamline I3 at MAXIII. The undulator delivers intense photon beam of 10^{12} ph/s with vertical polarization, with a wavelength tuned around the SPR. An effusive beam of C_{60} was produced with a ceramic oven, developed at LASIM, and resistively heated to a temperature of 550°C. The beam was collimated to get a well-defined spatial overlap with the monochromatic photon beam (few mm).

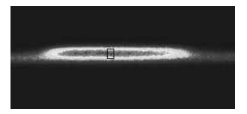


Figure 1a): Ion spatial imaging mode used to monitor the spatial overlap between the molecular beam of C_{60} and the light beam.

The velocity map imaging apparatus from the Atomic-Physics Division of Lund University was installed for the first time at the MaxLab beamline. This system allows reconstructing the 3D electron(or ion) velocity distribution from its measured 2D projection, meaning that both the energy and angular distribution of the emitted electrons are accessible with a typical energy resolution of a few percent.

The spectrometer can work in two distinct modes:

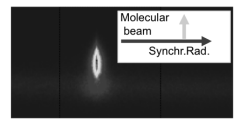


Figure 1b) Ion velocity map imaging mode used to characterize the molecular velocity of 170 m/s, $\Delta v=50$ ms/s.

- i) spatial imaging mode that can used to optimize the spatial overlap (Figure 1a),
- ii) velocity map imaging mode to characterize the beam dynamics (figure 1b) or for electron/ion spectroscopy.

Photoemission measurements were performed scanning the photon energy from 15 to 30 eV.

Results

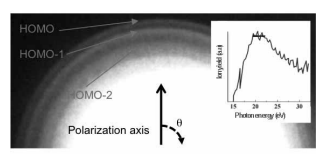


Figure 2) Electron image on the detector of the spectrometer obtained at 20 eV exhibiting the C_{60} shell structure of the valence orbitals (HOMO, HOMO-1...). In the inset is presented the total ion yield in the vicinity of the SPR as a function of the photon energy.

In Figure 2 we present a typical electron image obtained at 20 eV around the maximum of the SPR. The rings map out the distinct lines corresponding to the different molecular orbitals. The radius scales with the velocity of the photoelectrons. The first ring corresponds to the ionization of a single orbital $(6h_u)$, the second ring is related to the ionization of two orbitals $(10h_g)$ and $(6g_g)$.

The polarization axis is indicated as a guideline to inspect the angular distribution as function of θ . A clear difference between the HOMO and HOMO-1 is observed at $|\theta|$ =90°, where the signal almost vanishes for the outer ring. We are currently analyzing the variation of the angular distribution of these various orbitals both from the experimental data and theoretical point of view. This will lead to a reconstruction of the energy dependent real part of dynamical polarizability of C_{60} .

In addition, we have also studied anthracene molecules for which similar measurements have been performed around the the σ -> σ^* resonance. These results pave the way towards experiments where we can also access the temporal evolution of plasmonic processes.

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Angle- and spin-resolved photoemission on ferromagnetic Mn₅Ge₃(0001) thin films grown on Ge(111)

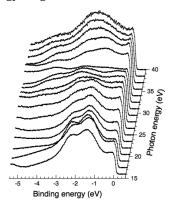
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Mn-doped germanium has recently attracted a lot of attention due to its capability to form either diluted magnetic semiconductors with a high Curie temperature ($T_{\rm C}$) or to develop ferromagnetic (FM) compounds in the Ge semiconducting matrix, which are opportunities to achieve spin injection for the emergent spintronics. Recently, the structure and the magnetic/electronic properties of FM Mn₅Ge₃(0001) films epitaxially grown on Ge(111) have been investigated (see, e.g. Ref. 3 and the references cited therein). A degree of spin polarization (DSP) of 42% has been reported from point-contact Andreev reflection measurements. Recent spin-resolved photoemission in angle-integrated mode using He I radiation show a DSP of 15% at the Fermi energy ($E_{\rm F}$). This result is not consistent with the value of -41% predicted by band structure calculations, although theoretical predictions of the DSP are extremely sensitive to the details of the crystal structure in materials having a complicated Fermi surface. Thus further experimental measurements of the valence band electronic structure and of the DSP of FM Mn₅Ge₃ are highly needed.

We have performed angle- and spin-resolved photoemission measurements on beamline I3 (MAX III) on *in situ* prepared $Mn_5Ge_3(0001)$ Ge(111) films held at 80 K, i.e. well below the T_C of bulk Mn_5Ge_3 (≈ 300 K). In Figure 1 (left) we show a waterfall picture of the energy distribution curves in normal emission (Γ -A direction) in the 13-40 eV photon energy range.



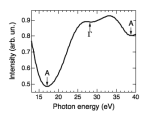


Figure 1: Left: valence band photoemission spectra of Mn_5Ge_3 recorded at normal emission and photon energies in the 15-40 eV energy range; right: intensity variation at E_F as a function of the photon energy.

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Each spectrum is normalized to its integral. By following the photon energy dependence of the spectral weight at $E_{\rm F}$ [see Figure 1 (right)] and comparing to the calculations⁶ we were able to determine the Γ and A points in the Brillouin zone. We then performed angle-resolved measurements (not shown here) along the A–L direction which shows clear band dispersion close to $E_{\rm F}$.

Preliminary spin-resolved photoemission measurements have also been performed. The spin-up and spin-down spectra recorded in normal emission for a photon energy of 15 eV, i.e. close to the A point of the Brillouin zone, as well as the corresponding DSP are shown in Figure 2. The DSP exhibits significant binding energy dependence and reaches a value of \approx 20% at $E_{\rm F}$ (it has been checked that the DSP related to the out-of-plane magnetization is zero). Note that the change we observe in the DSP in the 0-1 eV binding energy range has not been detected in previous angle-integrated measurements. This highlights the interest of the present angle- and spin-resolved study in scrutinizing the spin-polarized band structure of Mn₅Ge₃.

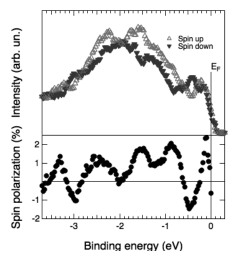


Figure 2: Valence band spin-resolved photoemission spectra of Mn_5Ge_3 (normal emission; photon energy of 15 eV) and the corresponding degree of spin polarization.

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In situ diagnostics of the thermal decomposition of partially hydrated hafnium tetrachloride by time-of-light mass spectrometry.

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HfCl₄ is the most common precursor for atomic layer deposition (ALD) of hafnium dioxide, used as high-k dielectrics in manufacture of modern metal-oxide-semiconductor based nanoelectronic devices. *In-situ* vapor analysis of the hydrated hafnium tetrachloride in the process chamber was performed by using time-of-light (TOF) mass spectroscopy and monochromatized synchrotron radiation in order to understand the chemistry and reaction channels of ALD. The measurements were carried out on the FINEST branchline of beamline I3 at the MAX-III storage ring, utilizing the photon energy range 7-40 eV¹. Using the Wiley-McLaren type ion TOF spectrometer² the investigated mass range was from 10 to 1000 amu. For the vapor production of the HfCl₄ (98%; Sigma-Aldrich) a resistively heated oven was used. The temperature in the evaporation zone and a temperature range at which the thermolysis was investigated were 35 – 120 °C. The pressure of the residuals was in the range of 10⁻⁶ mbars during the measurements.

Hafnium tetrachloride is very hygroscopic due to the strong affinity of the Hf^{4+} cation for O_2 and reacts spontaneously with H_2O to produce oxyhalide $HfOCl_2$ evolving hydrogen chloride. Anhydrous hygroscopic oxochloride (and hydrated oxochloride $HfOCl_2 \cdot 8H_2O$) decompose above 65 °C, yielding the dioxides³: $2HfOCl_2 \rightarrow HfO_2 + HfCl_4$.

Hafnium dioxide, HfO_2 , is very stable oxide of hafnium (melting point 2900 $^{\circ}C^3$), in contrast to the $HfCl_4$ which sublimes at 317 $^{\circ}C$ (pressure 1 atm) or at 212 $^{\circ}C$ (0.01 atm)⁴. The $HfCl_4$ compound studied in the present work was partially hydrated. Thermogravimetric analysis⁵ has shown that the partially hydrated $HfCl_4$ powders consist in particles with $HfCl_4$ core and a hydrated outer layer of $HfOCl_2 \cdot nH_2O$ with n in the range of 0–8. The low-temperature (<200 $^{\circ}C$) weight-loss was attributed to decomposition of $HfOCl_2 \cdot nH_2O$, for which n > 4.

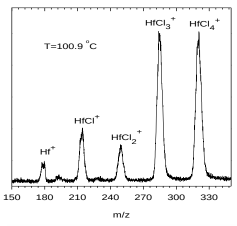


Figure 1. Mass intensity signals of Hf species for the thermal decomposition of hafnium tetrachloride, at 100.9 °C.

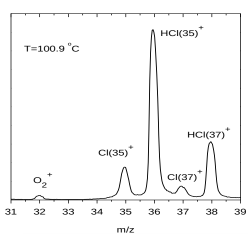


Figure 2. TOF mass spectra in the region of hydrogen chloride, at 100.9 °C.

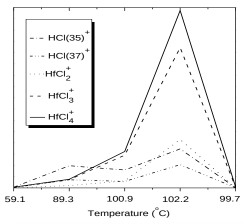


Figure 3. Temperature dependence of the main gaseous products of thermal decomposition at the photon energy of 31 eV.

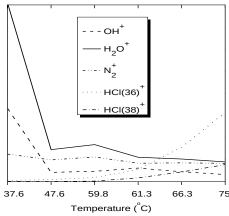


Figure 4. Temperature dependence of TOF signals of the residuals at low-temperature heating.

Figs. 1 and 2 demonstrate typical TOF mass spectra in the region of hafnium and hydrogen chloride at the photon energy of 31 eV. The mass spectra exhibit intense spectral features assigned chlorides of hafnium and hydrogen chloride. The fact that we observe these features at relatively low temperatures (about $100~^{\circ}\text{C}$) is clear evidence the complexity of the HfOCl₂·8H₂O dehydration processes. Note, that mass spectra features of Hf species in Fig. 1 are significantly broadened due to the natural isotopic distributions.

In Figs. 3 and 4 we present temperature dependences of the products of thermal decomposition. It is noteworthy that the HCl^+ intensity change nonmonotonically on heating. So at the low temperature region (<90 °C) the intensity of Hcl^+ grows faster than $HfCl_4^+$ signal. It is also worthwhile to note that intensity of H_2O^+ as well as OH^+ decreased remarkably with increasing temperature (Fig. 4). One can assume that the water and the hydroxyl radicals (OH) originate from the outward gaseous diffusion of these molecules from the porous, hydrated outer layer of $HfCl_4$ particles. It is natural to suppose that drastic decrease of TOF signals, which occurs at temperature of about 40-50 °C, indicate the temperature when H_2O and OH become highly active for the dehydration reactions. The total dehydration reaction of the hydrated $HfCl_4$ powders at higher temperatures can be considered, according to the global reaction: $HfCl_4 + 2H_2O \rightarrow HfO_2 + 4HCl$.

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Dissociation of Sb₄ clusters following 4d core excitation and the subsequent resonant Auger decay

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The fragmentation patterns of Sb₄ clusters following the 4d core excitation and the subsequrent resonant Auger decay process were studied on the FINEST branch of the the undulator beamline I3 of the MAX-III storage ring. The fragmentation of the Sb₄ cluster was studied using electronion coincidence (PEPICO) and partial ion yield (PIY) techniques. The PEPICO events were recorded using a PEPICO apparatus consisting of the SES-100 and a Wiley-McLaren-type ion time-of-flight (TOF) spectrometer built in Turku. The same ion TOF spectrometer was used also for recording the PIY spectrum. The Sb₄ vapor was generated using a resistively heated oven. The recorded PIY spectrum is presented in Fig. 1. The PIY spectrum shows the highly dissociative nature of the final states of the resonant Auger decay following the 4d excitation.

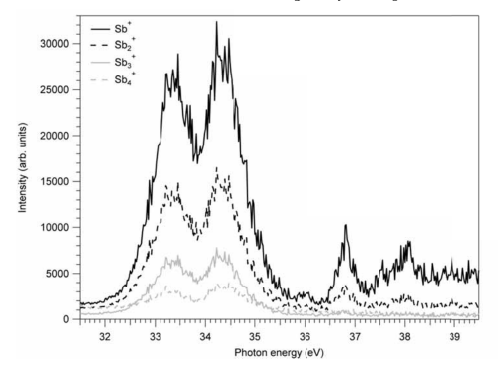


Figure 1. The PIY spectrum of Sb_4 clusters just below the 4d ionization threshold. The PIY spectrum shows clearly that the final states of the resonant Auger decay process are highly dissociative as only a very small amount of Sb_4 cations are present.

In order to further study the dissociation processes in more detail the PEPICO maps were recorded at the photon energy corresponding to the top of the first resonance feature. The electrons were collected in the binding energy region 7.6 eV to 18.75 eV corresponding to outer valence $(2e6t_25a_1)^{-1}$, inner valence $(5t_24a_1)^{-1}$ and shake-up and correlation satellite states $(2e6t_25a_1)^{-2}nl/n'l'$ and $(5t_24a_1)^{-2}nl/n'l'$ [1], where nl indicates and excited electronic state due to a shake-up process and n'l' an excited state due to correlation effects. The PEPICO recorded PEPICO map with false coincidence events subtracted is presented in Fig. 2.

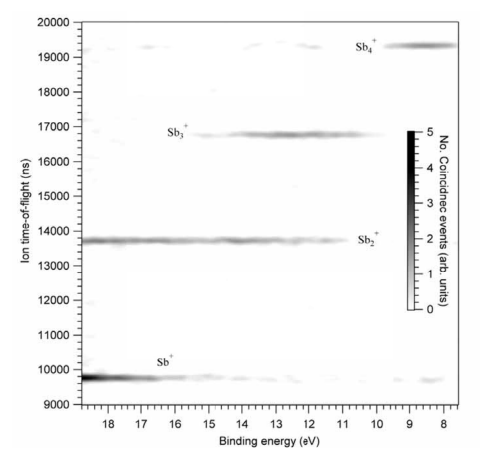


Figure 2. The PEPICO map from Sb4 clusters with false coincidence events subtracted.

The PEPICO map shows that the outer valence electrons do not contribute to the bonding of the cluster as no dissociation takes place after removing one electron from one of these orbitals. However, the removal of two electrons from the outer valence orbitals already starts to weaken the bonds resulting in fragmentation of the cluster. The removal of electrons from the inner valence orbitals always leads to a fragmentation of the cation .

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Spin Resolved Photoemission Spectroscopy at MAX; the detector upgrade and first data from a 1D topological surface

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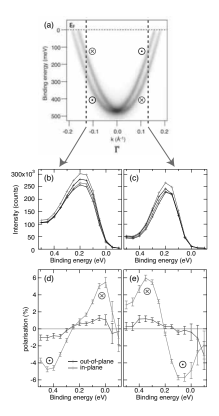


FIG. 1: The surface electronic structure of Au(111); panel (a) shows an overview of the Rashba split parabolic state, and the two green lines indicate the momenta at which EDCs were made using the MAX Mott detectors. Panels (b) and (c) show the raw 4-channel EDCs and (d) and (e) show the uncorrected polarisations extracted from the EDCs. The spin vector has a strong in-plane polarisation, and the spin direction (reversed between (d) and (e)) agrees with a simple Rashba model.

Detector test - Au(111): Following the replacement of the channeltrons at the Spin-resolving ARPES beamline I3 of MAX3 [1], a test was performed on the Rashba-split parabolic surface state of Au(111) (see Fig. 1). Panel (a) shows the spin-integrated photoemission intensity of the surface state (actual measurement from Ph. Hofmann [6], after reference [2]). From this overview two energy distribution curves (EDCs) are chosen and remeasured using the 4-channel Mott detectors at I3. The position of the two EDCs is indicated by the vertical green dashed lines, and the 4 spin signal EDCs are plotted as panels (b) and (c). From these 4-channel EDCs, it is straightforward to extract the uncorrected [7] polarisation as simply $P_{\parallel} = (I_{\rightarrow} - I_{\leftarrow})/(I_{\rightarrow} + I_{\leftarrow})$ and $P_{\perp} = (I_{\uparrow} - I_{||})/(I_{\uparrow} + I_{||})$ where $P_{||}$ and P_{\perp} are the in-plane and out-of-plane polarisations respectively, and $I_{\uparrow}, I_{\downarrow}, I_{\leftarrow}, I_{\rightarrow}$ are the intensities measured by the four spin-selective Mott detectors as plotted in panels (b) and (c). The resulting in-plane and out-of-plane polarisations are plotted in panels (d) and (e) of Fig. 1.

As expected, from previous studies in this surface [3], the Au(111) surface state shows a large in-plane reversal in polarisation, consistent with the Rasba spin splitting of the state. Between Fig 1 (d) and (e) the polarisation is reversed - consistent with the spin-orbit coupled nature of the state. The magnitude of the measured polarisation is ≈ 5 % and the raw EDC count rate is good [8], indicating that the machine is performing very well.

Vicinal surfaces of bismuth: Bismuth is one of the heaviest stable elements, and hence has unusually strong spin-orbit coupling associated with its electronic surface states [4]. By preparing strongly vicinal surfaces, we have previously demonstrated the 1dimensional (1D) surfaces can be engineered, and that they support strongly 1D spin-split surface states [5].

Here we have used a new vicinal surface Bi(441), which is also expected to have 1D character - a real-space trun-

cated bulk model of surface is depicted in Fig. 2(a). The 1D nature of the surface states is confirmed by recent ARPES measurements both at I3 and at ASTRID - an image through normal emission collected at I3 is shown as Fig. 2(b).

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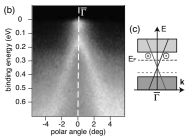


FIG. 2: (a) A truncated bulk model of Bi(441), showing the 1D nature of the surface. (b) ARPES measurement through normal emission, taken using the CCD detector at I3 and (c) a simplified model of the electronic structure showing the bulk bands in grey and the spin-split surface states in red and blue. The expected spin directions associated with these states are also shown.

the strong spin-orbit nature of bismuth surfaces, we expect this surface state to consist of two non-degenerate (i.e. single spin) bands in the projected bulk bandgap - a simplified picture of the expected structure of the surface state in shown in panel (c) of the figure.

Although the agreement between the measured surface electronic structure and the simple schematic representa-

From electron-counting arguments, and by considering

Although the agreement between the measured surface electronic structure and the simple schematic representation (Fig. 2(b) and (c)) is compelling, it is also necessary to measure the spin directions associated with these states in order to confirm that this non-degenerate single spin picture is correct. In order to do this, momentum dependent curves (MDCs) are made through this state. An MDC is performed at constant-binding energy whist the emission angle (and hence the in-plane momentum) is varied. Two such MDCs were made using the spin selective Mott detectors at I3; one MDC is made at the Fermi level, and one at a binding energy of 0.3 eV - in other words above and below the $\bar{\Gamma}$ point where the states cross and are hence degenerate.

In Fig. 3, the simplified picture of the surface electronic structure is repeated in the left hand panel, and the energies at which the two MDCs are made are indicated by horizontal green dashed lines. The middle panel shows a

simple model of the polarisation expected in the MDCs, and the right hand panel shows the corresponding measurements. As can be seen from the figure, the agreement is good, and hence the simplified picture of the spin-split surface electronic structure is verified.

Such a 1D spin-split surface state has significant implications; the odd number of Fermi level crossings between time-reversal invariant momentum points indicates that this state has metallic topological character. Furthermore, such 1D topological materials support a Spin Quantum Hall Effect - in other words, the strong coupling of an electron's spin its momentum means that carriers propagating in opposite directions will have opposite spins. - hence materials such as this are expected to have interesting spin-filtering transport properties.

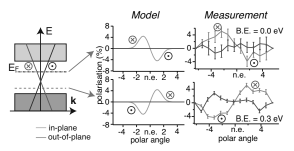


FIG. 3: The simplified surface electronic structure of Bi(441) together with the modelled and measured polarisations.

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 This has not been carried out here in order for the measured polarisation to be demonstrated, however a small normalising correction has been made such that the 4 detectors have the same collection efficiency
- [8] As shown in the raw EDCs in figure 1(b) and (c) each detector measures approximately 3×10^5 counts in the surface state. The integration time is 20 minutes per point.

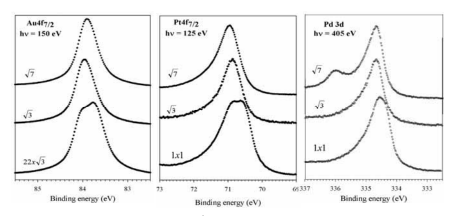
Surface structure, charge transfer and core level shifts on fcc(111)-I $\sqrt{3}$ and $\sqrt{7}$ surfaces

Sareh Ahmadi, Shun Yu, Mats Leandersson, Jonas Weissenrieder and Mats Göthelid Materialfysik, KTH and MAXLAB

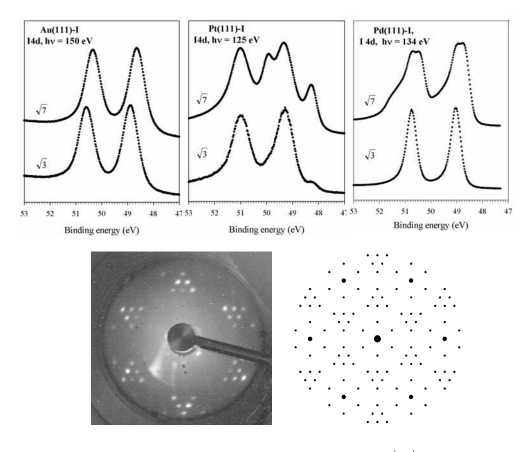
Iodine/iodide is used as redox couple in the Grätzel type solar cells. Charge is transferred from the metal electrode (usually a platinized oxide) by splitting of I_3^- into 3I. They donate charge and recombine into I_3^- on the dye sensitized TiO_2 electrode (the working electrode). Much effort has been put into understanding and improving the working electrode, but much less on the counter electrode. In order to improve the understanding of this side we have studied and compared some iodine induced structures on Au(111), Pt(111) and Pd(111) using valence band and core level spectroscopy at I4. One can question the relevance of such studies for real solar cell devices, but it has been shown by STM that iodine structures on Pd and Pt grown in solution share the same atomic geometry as those formed in UHV.

Halogens have been studied on several close packed fcc surfaces [1]. A commonly observed structure is the $(\sqrt{3}x\sqrt{3})R30^{\circ}$ phase at 1/3 ML halogen coverage. On Pd(111)-I it was found by normal incidence X-ray standing wavefield absorption that iodine occupies the fcc hollow [1]. The equivalent adsorption site was also proposed for iodine on Cu(111) [2], iodine on Pt(111) [3] and iodine on Rh(111) [4]. In addition to the 1/3 ML $(\sqrt{3}x\sqrt{3})R30^{\circ}$ structure on the Pt(111) surface, a $(\sqrt{7}x\sqrt{7})R19.1^{\circ}$ structure was observed at 0.43 ML and a (3x3) structure at higher coverage [5,6]. Both these latter structures were determined to be commensurate iodide adlayer structures [5].

On all three clean surfaces a core level shift (SCLS) to lower binding energy is observed. In all three cases this SCLS is removed when iodine adsorbs in the hollow site to form the $\sqrt{3}$ structure. Recent theory on Pt(111)-I [7] and Pd(111)-I [8] suggest that I is neutral in this position. Formation of the $\sqrt{7}$ structure induces rather different changes; on Au there is a weak broadening on the low binding energy side, on Pt a weak broadening instead appears on the high binding energy side and on Pd a new chemically shifted component due to iodide formation is observed.



I4d spectra are shown below. On all three $\sqrt{3}$ surfaces a single component is observed. The binding energies vary slightly between the surfaces. The widths are dramatically different; sharp on Pd(111), broad on Au(111) and broader on Pt(111). These differences are accentuated even more on the $\sqrt{7}$ structures; a moderate shift of the single component to lower binding energy on Au(111), a large chemical shift on Pt(111) and two new chemically shifted species on Pd(111). These differences in chemistry are accompanied by differences in electronic structure and the charge donating capacity of the different surfaces.



Experimental LEED patterns from Au(111)-I, together with a calculated $(\sqrt{7}x\sqrt{7})R19^{\circ}$ pattern.

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Photoemission from Graphene on SiC: asymmetry of replicas

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We have recorded angle resolved photoemission spectra for a nearly perfect graphene monolayer prepared on a SiC crystal. Other results from the characterization of sample preparation and characterization can be found in Ref. [1]. For the present measurements the polar detection angle is stepwise changed by 0.5° to record the emission of electrons from an approximately 1.2 Å-1 wide stripe of $k_{//}$ -values centered along the Gamma-K direction from -K to +K, this in an initial state energy range extending below the Dirac point. The sample was kept at around 100 K and the photon energy was 40 eV.

Following the discovery that thin graphite films may be prepared by heating SiC [2] and that these give photoemission spectra with a quality on par with those obtained from other types of sample [3]. Graphene mono- and multilayer thick overlayers on SiC have been studied intensely in recent years when the transport properties of graphene have attracted a great deal of attention. Somewhat surprisingly the present measurements provide information not revealed in previous reports. The one observation discussed here concerns the replicas of the emission features characteristic of the wave vector range around the K point of the Brillouin zone. At initial state energies near the Dirac point one then obtains emission from states quite near the K point. In addition to this one notes the appearance of similar emission features, replicas, at other emission angles. This is explained by the existence of two spatial periods of the overlayer system, one for graphene and one for the surface layer of the substrate, which has the 6root3 structure. The dual periodicity will affect the electronic structure (new gaps, diffuse energy bands) as well as the photoemission process (diffraction of the photoexcited electrons). In the present case there is approximate registry but the reduced Brillouin zone is small. One might expect energy gaps at many boundaries but these will often be narrow and obscured by lifetime broadening of photoemission lines.

The present work has revealed several more of these replicas than noted in previous experiments. Fig.1a and 1b gives an overview of the results. Previously the hexagon of replicas around the K points have been noted and in another experiment two replicas closer to the origin have been observed in the Gamma-K direction, at 0.66~1/Å and 1.06~1/Å. For an initial energy somewhat above the Dirac point the emission intensity is asymmetric around the K-point and the asymmetry is reversed when the energy is below the Dirac point.

Together with the experimental points the positions expected due to diffraction are plotted. Since the reduced Brillouin zone is small many other replicas could be present. The ones that appear are understood in simple terms of a combination of the graphene and the unreconstructed SiC lattice. The simplest case is the given by the hexagon of replicas around the origin, which is obtained by diffraction involving only a reciprocal lattice vector of the substrate. One such

vector transfers electrons from the +K point by 2.36 1/Å to the left such that the electrons end up with $k_{//}$ = - 0.66 1/Å. One may note that as expected the diffracted spot preserves the asymmetry. We will not dwell on further details regarding the pattern shown in Fig.1 .

The high quality of the sample may explain the results obtained regarding the replicas. We conclude by pointing out that not all features appear as distinct as the replicas. There are regions in energy and k- space where bands are more diffuse than would be expected for say a 3D sample with high crystalline quality.

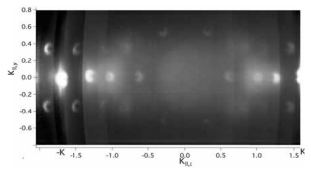


Fig. 1a: Experimental data at Fermi-level

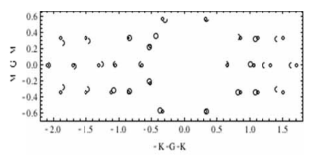


Fig 1b: Half and big circles: experimental data, small circles calculated.

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Electronic structure of the thallium induced 2×1 reconstruction on Si(001)

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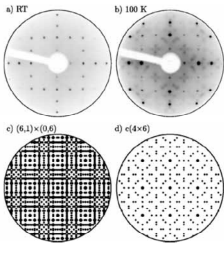
With a Tl coverage of one monolayer (ML), a 2×1 reconstruction is formed on the Si(001) surface at room temperature. In this study, low temperature angle resolved photoelectron spectroscopy (ARPES) data reveal four surface state bands associated with this Tl induced reconstruction. Calculated surface state band dispersions, obtained using the "pedestal + valley-bridge" model, are found to be similar to those obtained using ARPES. Inclusion of spin-orbit coupling is found to be important for the calculations to arrive at these results. A known effect of the strong spin-orbit coupling is the reluctance of the Tl $6s^2$ electrons to participate in the bonding, i.e., the inert pair effect. In the calculations, inclusion of spin-orbit coupling results in a ~ 5 eV downshift of the Tl $6s^2$ electrons, away from the dispersive surface states close to the Fermi level.

The atomic and electronic structures of reconstructions, induced by adsorption of group III metals on Si surfaces, have attracted a lot of attention through the years. These studies have mainly been focused on Al, Ga and In. Only recently, the attention has turned to Tl, the heaviest of the group III metals. Tl exhibits the peculiar inert pair effect, i.e., the $6s^2$ electrons tend to not participate in the bonding due to the strong spin-orbit coupling. As a consequence, Tl can exhibit a 1+ valence state, contrary to the other group III metals. By varying the temperature conditions, Tl was found to exhibit both the 1+ and the 3+ valence states in a study of the Si(111):Tl surface. Later, the valence state was found to be independent of coverage, as no evidence of the 3+ valence state was found for Tl coverages up to 1 monolayer (ML) in a core-level study of the same surface.

At sub ML Tl coverages, the Si(001) surface exhibits a series of 2×2 reconstructions, while a 2×1 reconstruction is formed at 1 ML. Of the group III metals, Tl is the only one that gives rise to a 1 ML 2×1 reconstruction. Total energy calculations and scanning tunneling microscopy (STM) have suggested that the "pedestal + valley-bridge" model is the likely atomic structure of this surface. The Si(001):Tl surface has been shown, using STM, to transform into a $(6,1)\times(0,6)$ periodicity. This is, however, not observed in low energy electron diffraction (LEED). The diffraction pattern resembles that of a $c(4\times6)$ periodicity. The variable valence state and the low temperature phase transitions make the Tl induced reconstructions interesting. Further, Tl/Si(001) has a possibility to show interesting spin splitting that originates from the Rashba effect, in similarity with Ti/Si(111) and Bi/Si(111).

P.E.J. Eriksson, K. Sakamoto, and R.I.G. Uhrberg, *Electronic structure of the thallium induced 2x1 reconstruction on Si(001)*, Phys. Rev. B <u>81</u>, 20422, 2010.

FIG. 1: LEED patterns obtained at a) RT using 111 eV electrons, and b) at 100 K using 94 eV electrons. c) and d) show simulated LEED patterns of two domain $(6,1)\times(0,6)$ and $c(4\times6)$ reconstructions, respectively. In d), integer order spots are recognized as they are slightly larger. The size of the simulated patterns matches the one in b).



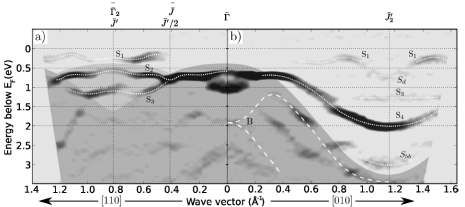


FIG. 2: Map showing features in 21.2 eV ARPES data obtained at 100 K along: a) the [110] direction and b) the [010] direction. Six identified surface state bands are marked by dotted curves. Four bands associated with the Tl induced 2×1 reconstruction are labeled S1-S4.

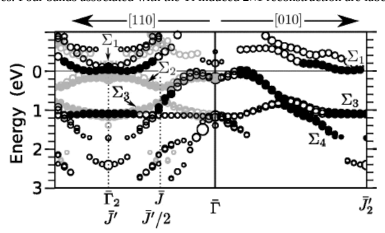


FIG. 3: Surface band dispersions calculated using the 1 ML Si(001):Tl 2×1 "pedestal + valley-bridge" model.

Photoemission study of (Zn,Co)O films with uniform Co distribution

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Diluted magnetic semiconductors (DMS) have been recently intensively studied for spintronic applications [1,2] and ZnCoO focuses a great part of an attention here [3,4]. Ferromagnetic ordering in (Zn,Co)O has been found and reported by several groups, but the origin of the ferromagnetism is still not clear. It has been related to defects in the ZnO lattice, formation of foreign phases like CoO and metal accumulations of the nanometers scale or uncompensated spins at the ZnCoO surface with higher Co content [5,6]. Dimensions of inclusions that contribute to room temperature ferromagnetism can be at the level of only few nm, so very difficult to detect.

There are only few resonant photoemission results on a (Zn, Co)O system reported to date [7, 8]. In each of these papers, experimental photoemission data were obtained for only one (Zn, Co)O sample and none of them related the electronic structure with the Co content and distribution. The aim of the photoemission measurements was to partially fill this gap.

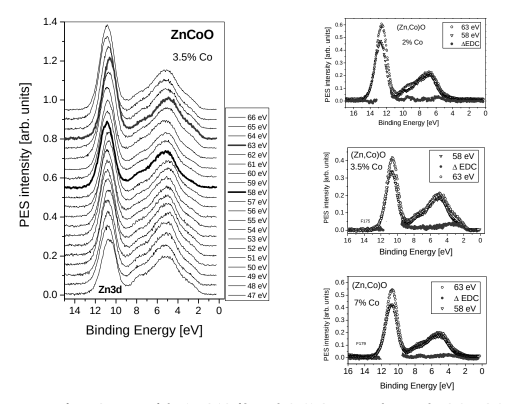


Figure 1. Left: EDC spectra of the (Zn,Co)O film with 3.5% Co measured across the Co3p→Co3d resonance. Thicker lines shows resonant (63 eV, red on-line) and anti-resonant (58 eV, orange on-line) spectra. Right: EDC spectra taken at resonance (63 eV, blue solid circles) and anti-resonance (58 eV, black open triangles) photon energy along with the calculate difference curve (ΔEDC, red open circles).

The investigated (Zn,Co)O films were grown by the Atomic Layer Deposition (ALD) method at the same temperature of 160° C and using the same zinc, cobalt and water precursors (dimethylzinc, Co(acac)₂ and water, respectively). We studied three samples with the Co content 2%, 3.5% and 7% as measured by SIMS and EDX.

The experiment was performed at the I4 beamline at MAXLAB. All the films were uniform and the EDX signal showed a homogeneous Co and Zn distribution in the 200 nm x 200 nm scale.

We used the resonant photoemission spectroscopy, which is as an effective tool for identification of the Co3d ion charge states in the valence band electronic structure of the (Zn,Co)O material. EDC's were taken for photon energies between 47 eV and 66 eV, e.g. across the $Co3p \rightarrow Co3d$ transition. In Fig. 1(left) we show the set of Energy Distribution Curves (EDC's) for (Zn,Co)O film with 3.5% of Co. For all three investigated films the maximum of the Fano resonance was observed at photon energy 63 eV and the minimum at 58 eV. In Fig. 1 (right) we present resonant and anti-resonant EDC spectra for films containing 2% Co, 3.5% Co and 7% Co. We also show the difference spectra (Δ EDC) that is related to the Co3d contribution to the valence band of the (Zn,Co)O system.

We noticed that resonant enhancement of the photoemission intensity from the Co3d shell does not scale with the cobalt content. The strongest photoemission response from the Co3d electron shell was observed for (Zn,Co)O film with 3.5% of cobalt. The resonant photoemission signal from the Co3d states for (Zn,Co)O films with 7% of Co was similar to that which was obtained for (Zn,Co)O film with 2% of cobalt.

Moreover, we also noticed that films with different content of cobalt give different Co3d contributions to the valence band electronic structure. For films containing 3.5% of Co we observe only one Co3d contribution, which is located between 2 eV and 4 eV below the Fermi level with the maximum at 3 eV (see Fig. 3b). For two remaining (Zn,Co)O films the additional Co3d contribution at higher binding energy is also present. In case of film with 2% of Co it is located 9 eV below the Fermi level (see Fig. 3a), while for (Zn,Co)O film with 7% of Co it is observed at 7.4 eV below the Fermi level (see Fig. 3c). The observed differences are tentatively ascribed to different hydrogen content, as we suppose that hydrogen promotes correlation of the *Co3d* electrons.

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Self-organized trimesic-acid nanomesh on Cu(100): adsorption geometry of the TMA molecules

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Self-organization of organic molecules into metallosupramolecular nanotemplates on metals and metal oxides is a promising avenue for bottom-up fabrication of functional surfaces. In particular, molecular species with functional groups for both intermolecular hydrogen bonding and covalent surface bonding readily facilitate self-assembly of highly organized and robust nanostructures. Potential applications range from enhanced surface corrosion resistance to covalent and specific immobilization of proteins in specific conformations and orientations for biosensor and implant technologies.

In the present study, the adsorption of trimesic acid (TMA) has been investigated on metallic Cu(100). While Cu is not widely used in bioapplications, its propensity to adopt a nanopatterned surface structure upon adsorption of molecules such as oxygen and nitrogen is highly useful for future studies in this field.

The Cu(100) surface was cleaned by cycles of Ar-ion bombardment at 500 K followed by annealing to 650 K. TMA was adsorbed on the surface at 300 K with physical vapor deposition in ultra high vacuum. Adlayers were investigated by Angle Resolved Ultraviolet Photoelectron Spectroscopy (ARUPS) at beamline I4 at MAX III. tunneling microscopy (STM) Scanning experiments were performed at TUT.

Figure 1 shows STM images from TMA covered Cu(100) at room temperature. As seen in Figures 1(a) and (b), at low coverages, TMA adsorbs preferentially at migrating Cu aggregates. As the coverage is increased, the TMA molecules begin to form a well-ordered nanomesh. This can be seen in Figures 1(c-f).

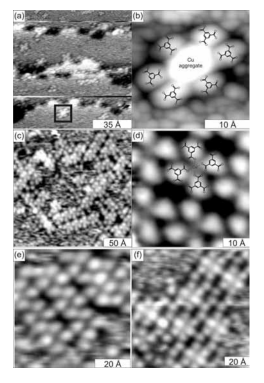


Figure 1: STM images of TMA/Cu(100) at the coverage of 2.8 Å (a and b) and 5.0 Å (c-f). The tunneling parameters: (a,b) the sample bias voltage $V_S = +0.42$ V, the tunneling current $I_T = 479$ pA; (c-e) $V_S = -1.60$ V, $I_T = 550$ pA; (f) $V_S = -1.40$ V, $I_T = 500$ pA.

ARUPS was employed in the investigation of the valence electronic structure of the TMA nanomesh and the orientation of TMA molecules.

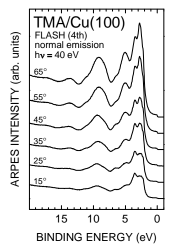


Figure 2: ARUPS data measured from TMA/Cu(100) at normal emission as a function of incident angle.

Figure 2 shows ARUPS data measured from TMA/Cu(100) at normal emission. Emission from molecular orbitals of the phenyl ring (at around 5 eV, 9.4 eV, 11.1 eV and 14 eV) indicate that the ring remains intact upon adsorption.

Emission at around 5 eV originates from the π states of the phenyl ring, whereas the transition seen at 9.4 eV is a superposition of π and σ states. Due to the exclusion rules of electronic transitions, the relative emission rate from orbitals with π and σ symmetry depends on the polarization of the incoming light.

By changing the angle-of-incidence of the linearly polarized light, we have changed the polarization from p- to s-type. Should the phenyl ring orientation be parallel to the surface or perpendicular to it, we would expect to see strong polarization dependence

in the relative intensities of π and σ states. However, the observed polarization dependence is weak, thus indicating that plane of the phenyl ring is most likely tilted. This result is in good agreement with our earlier XAS results from D1011.

The most likely explanation for the apparent tilt of the phenyl ring is that the preferred adsorption site is in the proximity of Cu adatom clusters seen in Figure 1. TMA bonds strongly to a Cu cluster via its OH groups and, since the Cu cluster resides on top of the surface, the optimal TMA adsorption geometry requires tilting of the phenyl ring away from the surface plane.

Combined experimental and theoretical study of surface core-level shifts on Ge(111)-c(2×8)

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The number and atomic origin of components in core-level photoemission from the clean Ge(111)- $c(2\times8)$ surface so far remain an open issue. Typically, two surface core-level shifts (SCLS) have been found in 3d spectra taken with the total-energy resolution of 0.2-0.3 eV [1-6]. A third SCLS has been identified in spectra taken with higher resolution (80 meV) [7]. The assignment of these SCLS to individual atoms is still debated, which is hindered by the large size of the $c(2\times8)$ unit cell and the lack of *ab initio* core-level calculations.

In this study, we demonstrate the presence of four SCLS in the 3d spectra of Ge(111)-c(2×8) measured with improved resolution (48-58 meV), and interpret these data on the basis of density functional theory (DFT) calculations. The measurements were performed on beamline I4 at MAX-III by using a SPECS Phoibos 100 electron analyzer. The spectra were taken at 100 K with photon energies hv = 40-90 eV and emission angles $\theta_e = 0$ -80°. The acceptance angle was $\pm 8^\circ$. The Ge(111)-c(2×8) surface was prepared by repeated cycles of 700–1000-eV Ar⁺ sputtering at 775 K and subsequent annealing at 900 K until a sharp low-energy electron diffraction pattern was observed. The calculations were carried out by using Vienna *ab initio* simulation package (VASP).

Figs. 1 and 2 show Ge 3d spectra measured at various experimental conditions and their decompositions, respectively. The spectra are fitted by five spin-orbit-split Voigt functions which are the bulk (B) and surface (S_1 - S_4) components. The binding energies of S_1 - S_4 are respectively -0.290, -0.755, +0.150, and -0.125 eV relative to the bulk. The spin-orbit splitting and Lorentzian width are respectively 0.588 and 0.150 eV for all the components. The branching ratio is 0.658 \pm 10% at θ_e = 0° and 0.663 \pm 10% at θ_e = 60°. The Gaussian width is found to vary between 0.161 and 0.250 eV for different components. Fig. 3 illustrates residuals obtained in three fitting schemes: the fit with four surface components (Fig. 2) and two other fits with three and five surface components (not shown). It is seen that three surface components are not enough to reproduce the 3d spectrum around the relative binding energies of -0.2 eV and 0.35 eV (note that in this range, the S_4 is identified in Fig. 2). An introducing of the fifth surface component, in contrast, does not lead to any noticeable improvement of the fit quality. Thus, we propose the fitting scheme with the four SCLS (S_1 - S_4) which describes the 3d spectra adequately.

In order to interpret the measured SCLS, DFT calculations were performed by VASP. The slab with the full-periodicity $c(2\times8)$ surface unit was used. The SCLS were calculated for two different fully optimized atomic configurations: the structure with the adatoms at T4 sites, which is energetically favorable, and the structure with the adatoms at H3 sites, which is slightly less stable. In Fig. 4, the calculation results, i.e. the initial state and complete screening DFT SCLS and the differences between these values (SCLS^{IS}, SCLS^{CS}, and Δ_{SCLS}), are shown for the two structures. The analysis reveals that the Ge 3d line shape can be reproduced well within the initial state model, and that the measured shifts agree fairly with the T4 structure. On this basis, the following assignments are made: the atomic origin of S_2 is the rest atoms, and the origin of S_3 is the adatoms and third-layer atoms which are below the adatoms. The S_1 and S_4 are contributed by the first- and second-layer atoms, with the core-level binding-energy splitting of these atoms being too small to resolve in experiment.

Thus, the four surface components are resolved in the 3d emission from the Ge(111)-c(2×8) surface. The atomic origins of these components are interpreted on the basis of DFT calculations, providing with more detailed understanding of 3d photoemission line shape from this reconstruction.

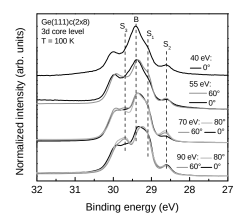


Fig. 1. Raw 3d spectra of Ge(111)-c(2×8) at various hv and θ_e .

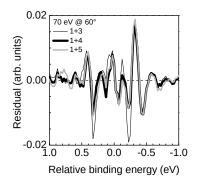


Fig. 3. Residuals obtained in the fitting schemes with three, four, and five surface components.

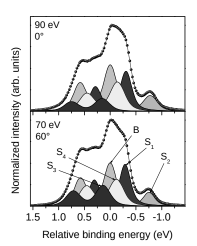


Fig. 2. Decomposition of spectra at $(hv, \theta_e) = (70 \text{ eV}, 60^\circ)$ and $(90 \text{ eV}, 0^\circ)$.

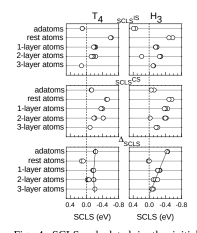


Fig. 4. SCLS calculated in the initial state and complete screening models, and their differences for the T4 and H3 structures.

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Unoccupied electronic structure of graphite probed by Angle-resolved Photoemission spectroscopy

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Graphite valence band structure has been intensively studied both theoretically and experimentally [1–4]. The interest in graphite has been renewed recently because of the discovery of Graphene out of Graphite by peeling off the top layer [5]. Graphene has become the most promising material for future electronics due to its exceptional electronic structure around the Fermi energy [6]. Graphite can be regarded as a prototype two dimensional material with full valence bands structure characterization along parallel to the surface and strong localization perpendicular to the surface.

Here, we present the angle-resolved photoemission spectroscopy (ARPES) results up to the range of secondary electron on single crystal graphite. In addition to the valence band structure which is well known and reported widely, we find some anomalous bands which appear as a constant kinetic energy features at different photon energies. This band are neither reported in ARPES of graphite nor predicted by the bulk band structure calculations. The experimental results are discussed in relation to the VLEED experiments performed by Strocov *et al.* [7], which essentially probes the photoemission final states, showing that the VLEED information can be found in the ARPES experiments. This makes the ARPES a versatile experimental tool for mapping both the occupied and unoccupied part of band structure with full k resolution [8]. The experiments on single crystal graphite were conducted at I4 beamline of MAX-Lab synchrotron in Sweden. The graphite single crystals were cleaved in-situ using the aluminum post in preparation chamber.

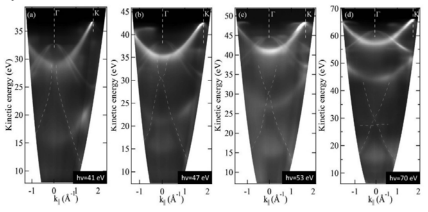


Figure 1: ARPES spectra of single crystalline graphite using (a) 41 eV photon energy along ΓK symmetry direction, (b) 47 eV photon energy along ΓK direction, (c) 53 eV photon energy along ΓK direction and (d) 70 eV photon energy along ΓK direction. Anomalous bands are shown with yellow dashed line.

In order to study the origin of these anomalous bands, we have performed ARPES measurements at different photon energy in synchrotron. The same anomalous band can be seen in ARPES spectra of different photon energy as shown in Figure 1. Apart from the anomalous band seen in 41 eV photon energy ARPES spectra, we can see there are two more bands visible at 70 eV photon energy. There is a constant energy (non-dispersive) band near kinetic energy 28.5 eV and an upper dispersing band ranging from 30 eV to 42 eV kinetic energy. These other two anomalous bands were not visible or partly visible

in lower photon energies as those were masked by the other valence bands of the graphite. At 70 eV photon energy, the anomalous bands are in a region free of any other bands and can be distinguished very clearly. All these three anomalous bands are dispersed in constant kinetic energy for different photon energy. The dispersions along the ΓK and ΓM symmetry directions are very similar (not shown).

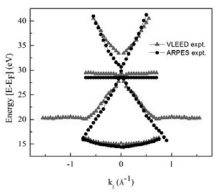


Figure 2: Comparison of the energy position and dispersion of anomalous bands in ARPES spectra with the VLEED results reported by Strocov *et al* [7] of graphite.

Analyzing the different photon energy data and seeing the constant kinetic energy nature of these three anomalous bands, we propose that these bands are coming from the unoccupied states or photoelectron final state bands of graphite due to secondary electron emission process. Since the VLEED experiment maps the unoccupied valence band structure of the solid, we compare the VLEED data reported by Strocov *et al.* [7] with our anomalous bands accomplished by ARPES experiment as shown

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in Figure 2. We establish the equivalence of VLEED and secondary electron emission process in the connection of mapping the unoccupied bands of graphite. In particular the experimental ARPES spectrum on graphite supports the idea of mapping the unoccupied bands or photoelectron final states by performing ARPES up to low kinetic energy (secondary electron energy) region. As a consequence, the ambiguity about the ${\bf k}$ can be regained and absolute band structure can be mapped by performing a single ARPES measurement.

In conclusion, we found the presence of the unoccupied bands or photoelectron final states in the ARPES spectrum of single crystal graphite. The unoccupied bands were mapped by the secondary electron emission process produced by the cascaded scattering of the photoelectron ejecting from the valence band. We confirm the equivalence of VLEED and secondary electron emission process as shown by Strocov *et al.* [8] in the connection of mapping the unoccupied bands of graphite. In particular the experimental ARPES spectra on graphite single crystal supports the idea of mapping the unoccupied bands or photoelectron final states by performing ARPES up to low kinetic energy (secondary electron energy) region.

This work was supported by the DST-facility for the synchrotron utilization, Department of Science and Technology, Govt. of India for experiments at MAX-Lab, Sweden. We acknowledge the Micro-Nano initiative programme of Department of Atomic Energy (DAE), Govt. of India for generous funding and support.

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Software development for analysis of 2-D ARPES data at BL I4

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In recent years, angle resolved photoelectron spectroscopy (ARPES) has been revolutionized by the introduction of electron analyzers with 2-D detectors. Data using such analyzers can be recorded as a 2-D image with one energy axis and one angle axis as illustrated in Fig. 1 (top row). The images show the distribution of the emitted electrons on the two dimensional energy-angle plane of the detector. The energy axis can be converted into a binding energy (E) axis and the angle axis into a k-parallel axis. When BL I4 was built up, the Phoibos 100 analyzer from Specs was chosen. However, at that time there was no commercial software available that could process the data in a way that met the demands of the users. We therefore decided to develop software with the aim to incorporate various new features when required. The new software is a result of many discussions and communications with the beam line management and other users.

The new program, "Blender.ipf", was written as a macro (procedure) of the Igor software package. The data files obtained by the software of the Specs analyzer are saved in certain formats such as *.sp2 and *.xy. The manufacturer of the analyzer provided a macro to read and load the data as a matrix which we used in a raw or a modified form in our own macro.

The analyzer at BL I4 can be rotated around the horizontal lens axis. This allows for two basic modes of operation. In the vertical mode (V-mode) the energy dispersive plane of the analyzer is vertical and in the H-mode it is horizontal. The V-mode is used to obtain angle scans over a wide range along an azimuthal direction that is horizontal. However, the angle range within one image is limited and depends on what lens mode is used. To visualize the data over a wide angle range a set of E- θ images obtained at different emission angles, θ_e , (different polar angles of the vertically mounted manipulator) need to be stitched. The first main function of the program ("blending") is to stitch images separated by an angle $\Delta\theta_e$. In Figure 1, top row, there are 3 images collected in the V-mode. The images represent frames collected along a selected azimuthal direction at emission angles θ_e of 12, 22 and 32°, respectively (the angle between the surface normal and the lens axis of the analyzer). The angular widths of the images are determined by the angular range of the lens mode. A smaller angle width can be selected in order to cut out irregularities at the edges of the images. The width of each image in this example is 21° which means that neighboring images overlap by 11°. The result of the stitching is presented in the lower panel of Fig. 1, where the angle axis has been converted into a k-parallel axis. The software has algorithms for finding the proper angle and energy shifts in order to avoid artificial kinks in the dispersions going from one image to the next. There is also a procedure that accounts for the difference in intensity between images, resulting in a seamless stitching.

The H-mode is used to obtain E- θ images for various polar angles of the manipulator. The images in this mode represent a set of vertical slices through the Brillouin zone. One slice for each emission angle, θ_e . The second function of the software ("build 3D") visualizes the set of images spaced equally by some small angle $\Delta\theta_e$ (typically <0.5°). The set of the images collected in the H-mode can be visualized in various ways. Users usually have to deal with at least 100 such images which all together form a 3D object with one energy axis and two angular axes. Such a set can be visualized by cuts along a particular plane through the data set. So far tools to visualize the data along planes normal to all the axes are implemented. Two examples of cuts perpendicular to the energy axis (constant energy plane) are shown in Fig. 2. These images are already transformed to show the energy contours on a k_x , k_y plane.

I would like to express my gratitude to Prof. Roger Uhrberg, who initiated the project, and helped me with all the theoretical aspects of the project and contributed with new ideas and improvements. I am also thankful to Dr. Balasubramanian Thiagarajan, the beam line manager, who welcomed and encouraged our initiative, and who made important suggestions which helped to improve the macro. I would also like to thank all who had an impact on the project, especially Dr Johan Adell for general software discussions and Hafiz Muhammad Sohail for testing the macro and finding various "program bugs".

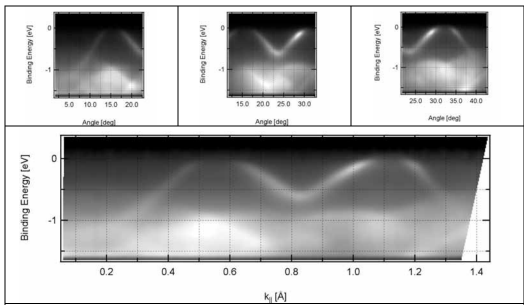


Fig. 1 Top row. Energy versus angle images obtained at an emission angle of 12, 22, and 32\mathbb{Q} respectively, with the analyzer in the V-mode. Lower panel. Wide range representation of the data after stitching. The angle scale has been converted into a k-parallel axis.

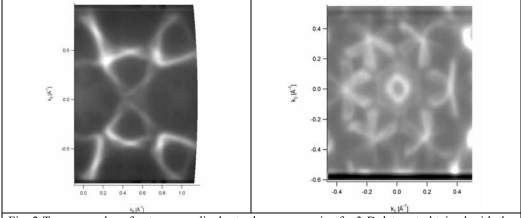


Fig. 2 Two examples of cuts perpendicular to the energy axis of a 3-D data set obtained with the analyzer in the H-mode. The images show constant energy curves in a k_x, k_y plane.

Chemical Doping of Epitaxial Graphene on SiC

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Chemical modification of graphene is a topic of current interest, especially functionalisation of the edges. The goal is the introduction of functional groups giving new and interesting properties to the graphene sheet; supply means of attaching entities with an attractive functionality or enable further processing control of the graphene layer.

We have launched a systematic study where we aim to introduce functionality at graphene edges, either in a gas-phase based approach using ozone under oxidizing conditions, or a liquid based approach where we expose the graphene to concentrated HCl which is under net non-oxidizing conditions. Both methods are selectively targeting the edges of graphene to mainly introduce oxygen and hydrogen containing functional groups. The question arises whether these treatments are electronically doping the graphene, i.e. are the functional groups adding or withdrawing charge to or from the π -system of graphene in any significant amount.

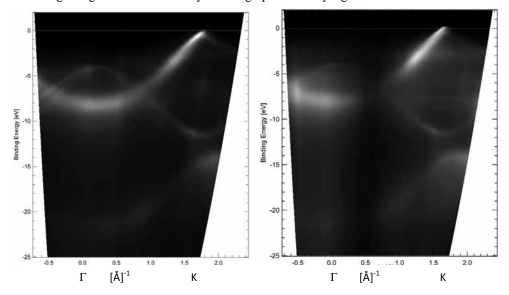


Figure 1: ARPES maps in the \(\Gamma \)-K direction of few layer graphene on SiC(0001) (left) and after HCl treatment (right), hv = 60 eV. The samples have been annealed at 550 K and 710K respectively.

In the present experiment, we investigate the effect on doping by performing Angle Resolved Photoelectron Spectroscopy (ARPES) at I4, MAX-Lab. The electronic structure of graphene is peculiar owing to the 2-dimensionaltity of the material and as a result, the π -electrons form bands with linear dispersion in the K-points of the surface Brillouin zone. The points where the

500 MAX-lab ACTIVITY REPORT 2010 filled and empty bands meet are known as the Dirac points. The effect of doping this electronic system can be easily seen in the band maps, as the Dirac points move up or down with the emptying and filling of electrons. Figure 1 above presents band maps along the Γ -K direction in the surface Brillouin zone, the photon energy is 60eV. The left panel shows the band structure of graphene on SiC after degassing the sample to 550 K while the right panel shows the HCl treated sample after heat treatment at 710 K. A more detailed view of the band structure at the K-point is presented in Figure 2 below. The Dirac point is found slightly below the Fermi level as is expected in the graphene on SiC case¹. There might be a small doping effect from adding functional groups on the surface. The Dirac point is found at about 0.2 eV binding energy in the present experiment while for un-doped samples it is found at 0.4 eV binding energy. The attack by HCl is mainly on the edges of the graphene layer as we cannot see any significant change in the Fermi level after exposure.

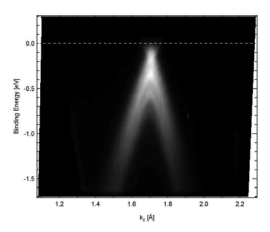


Figure 2: The Dirac cone at the K-point, hv = 60 eV.

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High-resolution core-level photoelectron spectroscopy of the Bi/Si(110)-(2x3) surface

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Heavy element adsorbed light element substrates are systems where exotic physical phenomena occur. One of such phenomena is the Rashba-Bychkov (RB) (or simply Rashba) effect [1] that originates from the spin-orbit coupling and the broken space-inversion symmetry at the surface. The RB effect, which has been observed on clean surfaces of noble metals [2] and heavy group V elements [3], are enhanced in these heavy element adsorbed light element systems [4]. In our former studies, we have investigated the RB effect on heavy elements Tl [5] and Bi [6] adsorbed Si(111) surfaces, and discuss the relation between the RB effect and the symmetry of the surface. In the case of Tl/Si(111)-(1x1), the surface state showed a "curious splitting" around the \overline{K} point. By performing spin-resolved ARPES measurements and first-principles electronic structure calculation, we concluded this curious splitting, which has not been observed on any other 2D or 3D system so far, to originate from the "C₃ symmetry" of the \overline{K} point of this system. On Bi/Si(111)-($\sqrt{3}x\sqrt{3}$), we found another peculiar Rashba effect at the \overline{K} point of this surface. That is, although the time-reversal symmetry was believed to be a necessary condition for the Rashba splitting and there is no time-reversal symmetry at the \overline{K} point, the "C_{3V} symmetry" originates a RB effect at a point without time reversal.

Since our former studies indicate that the 2D symmetry of the surface plays an important role to the RB effect, and because a proper understanding about the atomic structure of the surface is indispensable to comprehend a peculiar RB effect, we have performed core-level photoemission measurements on the Bi/Si(110)-(2x3) surface. The symmetry of the Bi/Si(110)-(2x3) surface is different from those of the systems used in our previous studies and thus this surface is a candidate to show new peculiar RB effects. The LEED pattern of the Bi/Si(110)-(2x3) surface shown in Fig. 1 indicates the high quality of the sample. In Figs. 2 and 3, we show the Bi 5d and Si 2p core-levels, respectively. The simple Bi 5d core-level spectrum shown in Fig. 2 and the observation of only one surface component in the Si 2p core-level spectra of Fig. 3 indicates that the atomic structure of this surface would be rather simple.

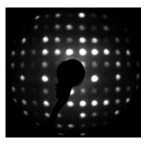


Fig. 1: LEED pattern of the Bi/Si(110)-(2x3) surface obtained at 100 eV.

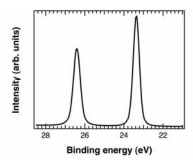


Fig. 2: Bi 5d core-level spectrum measured at hv=80 eV.

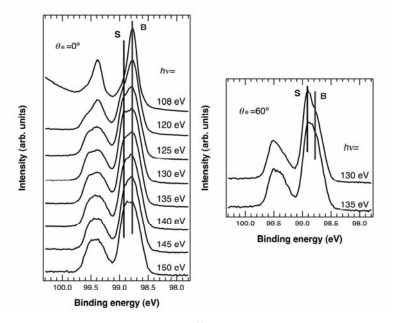


Fig. 3: Si 2p core-level spectra measured at different photon energies and emission angles

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Electronic structure of Ce₂TSi₃ systems; magnetic Kondo lattice (T=Rh) and a Fermi liquid (T=Co)

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1. Introduction

The project is devoted to investigations of Ce_2CoSi_3 and Ce_2RhSi_3 compounds, which exhibit peculiar electronic properties originating from the behaviour of Cerium f-electrons. Ce_2RhSi_3 is an antiferromagnetic Kondo system, thus exhibiting a competition between RKKY interaction and a Kondo effect [1]. Ce_2CoSi_3 is a moderate heavy fermion compound with the specific heat coefficient $\gamma = 160$ mJ/(mole K^2), what signifies that the hybridization between Ce f-electrons and conducting band electrons is much stronger than for its counterpart containing Rhodium. It was reported that a quantum phase transition appears if Co is gradually substituted for Rh [2]. Hence, the studies of these systems gain a new dimension as the two investigated compounds are located on both sides of a quantum critical point.

Present knowledge of the electronic structure of Ce_2TSi_3 systems (T=Rh, Co) is very limited. The first results concerning the angle integrated photoemission spectroscopy of Ce_2CoSi_3 valence band are known [3]. Therefore, the general purpose of the research is to determine the band structure topography and the Fermi surface. The second, but more important goal concerns the behaviour of f-electrons and their role in the physical phenomena observed in these systems.

2. Experimental

Single crystals of Ce₂CoSi₃ and Ce₂RhSi₃ were grown by Czochralski method at the Institute of Low Temperature and Structural Research in Wrocław. Studies of resonant photoelectron spectroscopy (RESPES) and angle resolved photoelectron spectroscopy (ARPES) were carried out at the beamline I4 of the MAX III storage ring with the PHOIBOS 100 electron analyzer from SPECS and photon energy between 36 eV and 150 eV.

The single crystals were cleaved in a vacuum of 1*10⁻¹⁰ mbar, cooled to the temperature of 100 K and subjected to the photoemission studies.

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3. Results

The valence bands of Ce_2RhSi_3 and Ce_2CoSi_3 were scanned for the incident photon energies between hv=88 eV and hv=150 eV. The energies were selected to exploit the effect of Ce 4d – Ce 4f resonance, which is observed near 120 eV. The exemplary results of angle intergrated spectra for Ce_2RhSi_3 are shown in the Figure 1. Spectral intensity related to Ce f-electrons increases when the photon energy approaches the resonance energy. This fact was used to determine the contribution from f-electrons, which is characterized by major peaks at the Fermi energy (Kondo resonance) and at 0.25 eV.

ARPES measurements were performed on Ce₂RhSi₃ and Ce₂CoSi₃ crystals as well. However, despite many attempts no clear dispersion was detected as the investigated compounds are very demanding systems.

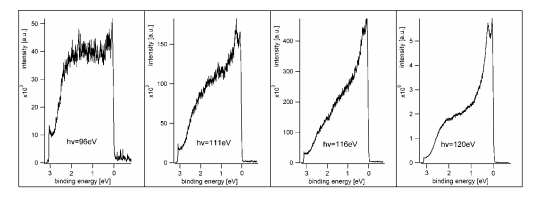


Figure 1. The valence band of Ce₂RhSi₃ measured with photon energies of 96 eV, 111 eV, 116 eV and 120 eV.

4. Summary

The main achievement of the studies carried out at the beamline I4 is a set of the valence band spectra obtained for both Ce_2RhSi_3 and Ce_2CoSi_3 with a series of photon energies, which correspond to a crossing through the $Ce\ 4d\ -4f$ resonance. The data are used to separate the contribution from $Ce\-f$ states to the valence band and will be compared to band structure calculations.

5. Acknowledgments

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Li intercalation on epitaxial graphene on 6H-SiC

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We have demonstrated an intercalation process of Li in monolayer graphene on SiC(0001). Our results show that Li atoms penetrate through the graphene as well as the carbon buffer layer and intercalate at the SiC/buffer layer interface. The process starts immediately after deposition, see Fig. 1. The Li atoms bond to Si atoms in the uppermost Si-C bilayer and create a dipole layer at the interface that induces a ca. 2 eV shift of the SiC core levels. The Li atoms do at the same time transform the carbon buffer layer into a second graphene layer. After Li deposition three π -bands are observed at the K-point, see Fig. 2. After annealing when a more even Li distribution and intercalation is obtained two distinct π -bands appear at the K-point. Li gives rise to electron doping and lowers the Dirac point (E_D) by 0.25-0.5 eV depending on coverage [1,2]

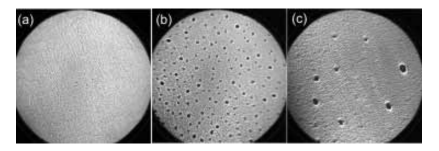


Fig. 1. (a) LEEM image of a monolayer graphene grown on SiC (0001); the field of view (FOV) is 50 μ m and the electron energy is E_{vac} -0.5 eV. (b) LEEM image illustrating the graphene surface after Li deposition. (c) LEEM image recorded after annealing the sample to 320°C.

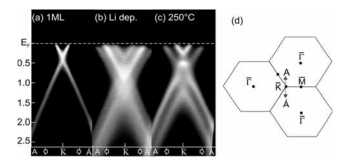


Fig. 2. The π band around the K point recorded from (a) monolayer graphene, (b) after Li deposition and (c) after annealing at 250°C. (d) Schematic drawing of the 2D Brillouin zone of graphene and the direction of scan.

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Band Structures of Ag films on $Sn/Si(111)\sqrt{3}\times\sqrt{3}$

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Ultrathin layer systems, which may show discrete quantum-well states (QWS) due to spatial confinement of electrons, have attracted a lot of attention in the fields of the surface and solid-state physics. Over the past decades, a large number of studies have been devoted to metal thin films on metal substrates. However, the observation of quantum-well states has been restricted to a few systems. In semiconductor layer systems, on the other hand, these effects are well known and some of them have been used in electronic devices. Recently several studies have reported thin metal films (Ag or Pb) grown on clean Ge or Si surfaces at reduced sample temperature [1,2]. These films show an energy quantization effect in their valence band states. Inspired by these findings, we have investigated the electronic structure of Ag films on the Sn/Si(111) $\sqrt{3} \times \sqrt{3}$ surface by angle-resolved valence band and core-level photoelectron spectroscopy.

A series of coverages (8-14 monolayers (ML)) of Ag were deposited on the Sn/Si(111)- $\sqrt{3}\times\sqrt{3}$ surface at low temperature (120 K). Figure 1 shows a gray-scale valence band image recorded with a photon energy of 21.2 eV from a thin Ag film at 100 K. The new analyzer (Phoibos 100), equipped with a CCD camera, was set with a wide angle mode (angular resolution 0.4°), which covers an angular range of ± 13 . The pass energy was set 10 eV, together with 0.2 mm entrance and 0.5 mm exit slits in the detector, which gives an energy resolution of 17.5 meV. For the current system with a thickness in the nanometer range, the valence electrons in Ag with energies within the fundamental gap of Si will be partially confined. The resulting quantum-well states disperse with in-plane momentum to form subbands. As illustrated in Fig. 1, the valence band of the Ag film clearly shows dispersions of its quantum-well states. There is a strong emission just located at the point near the Fermi-level. A parabolic-like metallic band (n=1) starts from k_/=±0.5 at the Fermi-level, and disperses steeply downwards to -1.5 eV at the point. Interestingly, this band seems to break at -0.5 eV and forms a very flat band near the point. Another parabolic-like band (n=2) starts from k_/=±0.9 at the Fermi-level, and disperses steeply downwards to -3.0 eV near the $\overline{\Gamma}$ point. The second band has a similar behavior as the first one. They also break down at -1.0 eV. These breaks in the subbands form wiggles and kinks that resemble the concave curves observed in the Aq/Ge(111) system [1]. They originate from the top valence state band of the substrate. This is a clear indication of an interaction between the Ag film and the underlying Si substrate. But the most interesting finding in this study was a very strong band at $k_f = \pm 1.3$ near the Fermi-level, which has not been observed before.

As shown in Fig. 2, the strong metallic band has an extremely steep dispersion, which seems folded at the M point of the £1 surface Brillouin zone (SBZ). By comparing its shape and dispersion with the theoretical calculation [3], this band should be a free-electron-like band with an sp character. The reason that it has not been observed in earlier studies is because this band disperses too quick to be "seen" in a normal spectrum with a limit angular resolution.

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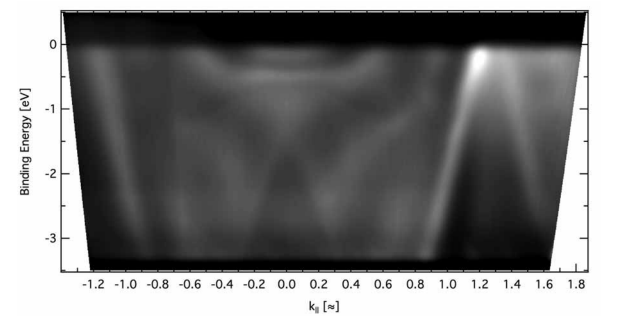


Fig. 1 Dispersions of band structure from a thin Ag film on the Sn/Si(111) $\sqrt{3}\times\sqrt{3}$ substrate along the Γ M direction, recorded by the CCD camera with a photon energy of 21.2 eV at 100 K. The wide angle range was obtained by adding ten partially overlapping frames, each covering $\pm 13^{\circ}$. The individual frames were stitched together using the software provided by Dr. Jacek Osiecki.

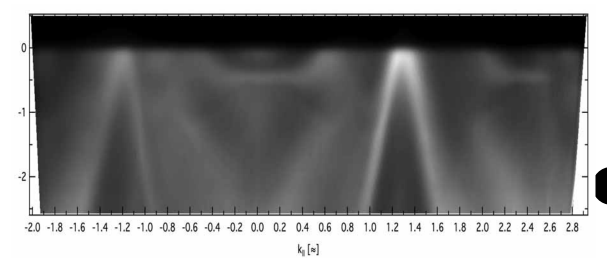


Fig. 2 Dispersions of band structure from a thin Ag film on the Sn/Si(111) $\sqrt{3}\times\sqrt{3}$ substrate along the Γ M direction, recorded by the CCD camera with a photon energy of 40 eV at 100 K. The wide angle range was obtained by adding ten partially overlapping frames, each covering $\pm 13^{\circ}$.

ENERGETIC ELECTRONS



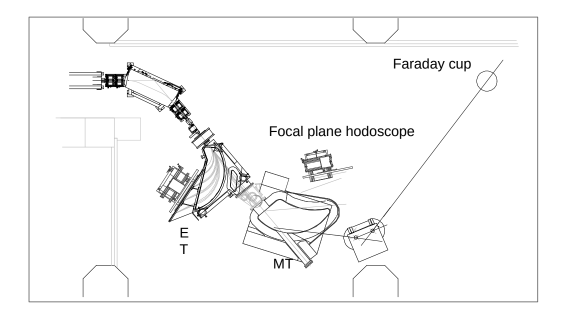
A gathering of nuclear scientists at a coffee break during the Annual User Meeting, 8 November, 2010. Around the table; Lennart Isaksson, John Annand, Kevin Fissum, Ken Andersen, Grant O'Rielly, Kenneth Livingston, and David Hamlton.

Photo: Annika Nyberg

Energetic Electrons

During 2010 the available beam time for the nuclear physics program at MAX-lab was the same as in 2009: 18 weeks. A detailed account of the program during these weeks may be found on the homepage of MAX-lab.

The tagged photon facility consists of an injector, two linear accelerators equipped with SLED cavities, operating at 10 Hz providing an electron pulse width of about 150 ns. This pulsed beam is injected into the pulse stretcher ring MAX I, and slowly extracted over 100 ms resulting in an almost continuous electron beam in the nuclear physics area. For an average current in the ring of typically 20 mA; the stretched beam current, measured in a Faraday cup, is about 20 nA. The electron beam impinges on a thin Al-foil and bremsstrahlung is produced. The residual (final) electron is momentum analyzed in one of two magnetic spectrometers. The non-interacting part of the electron beam is dumped in a shielded Faraday cup. The extracted beam contains an initial, short burst of electrons which are not contained in the ring. An abort kicker is used to dump these electrons before they reach the experimental cave.



The two magnetic spectrometers are the main tagger (MT) which covers photon energies corresponding to 10 – 70 per cent of the incoming electron energy, and the end point tagger (ET) which allows the use of tagged photons as close as 10 MeV from the endpoint of the bremsstrahlung spectrum. Full energy coverage implies that the focal plane hodoscope has to be shifted along the focal plane. This applies to both tagging magnets. The focal plane hodoscope was provided, along with the tagging spectrometers, by the Saskatchewan Accelerator Laboratory in Saskatoon. The SAL hodoscope consists of 63 overlapping scintillators (3 mm thick, 50 mm high and 25 mm wide) providing 62 coincidence channels. The same hodoscope is used at both tagging magnets.

A new focal plane hodoscope with 160 scintillators has been designed and has been funded by the Knut and Alice Wallenberg Foundation. The construction is well underway and the hodoscope is expected to be commissioned in 2011.

The bremsstrahlung beam produced enters into the experimental cave (from the top left in the figure) through a collimator (with diameter 4, 12 or 19 mm). The energy of a photon interacting in a nuclear target is obtained from a time coincidence between a reaction product, measured in the detection system, and an overlap channel in the focal plane hodoscope. The photon energy is obtained from the energy difference between the incoming and residual electrons. During 2010 two electron beam energies were used, 165 MeV and 193 MeV. The count rate in the focal plane hodoscope is typically 1 – 1.5 MHz per MeV.

The normal mode of operation of the MAX I as a stretcher ring is the following. The beam is available Tuesday morning until Monday morning the following week. There are normally three interruptions per day to allow filling the two storage rings MAX II and MAX III. In a normal week some 120 hours of stretched beam is delivered to the nuclear physics area. Students operate the accelerator during evening, night and weekend shifts. This system has worked well over the 18 weeks in 2010.

The Nuclear Physics Users at MAX-lab met on November 9-10 at a combined PAC and User meeting in conjunction with the MAX-lab Annual User Meeting. Four new proposals and letters of intent were presented, as well as status reports from ongoing experiments, facility-related reports and talks from invited speakers.

A summary of the activities in the beam time periods during 2010

Period	Electron energy Photon energy range	Projects
25; 2010-01-25 – 02-05	164.7 MeV; 12.9 – 68.5 MeV	PWO-crystal measurements
- " -	164.7 MeV; 6.7 – 66.4 MeV	Beam test
- " -	164.7 MeV; 63.7 – 102.4 MeV	Compton scattering on ¹⁶ O
26; 2010-03-01 – 03-28	164.7 MeV; 63.7 – 102.4 MeV	Beam diagnostics and rate-dependence measurements
- " -	192.9 MeV; 167.5 – 179.7 MeV	(γ,π^*) CH ₂ , C
27; 2010-06-07 – 07-05	192.9 MeV; 167.5 – 179.7 MeV	(γ,π^*) CH $_{2'}$ CD $_{2'}$ C, ⁶ Li
28; 2010-09-27 – 10-31	164.7 MeV; 81.6 – 115.8 MeV	Compton scattering on ² H, C
29; 2010-11-15 – 12-05	164.7 MeV; 81.6 – 115.8 MeV	Compton scattering on C
- " -	192.9 MeV; 167.4 – 179.5 MeV	(γ,π*) Та

The PhD thesis of Luke Myers, based on work carried out at MAX-lab, was defended in 2010.

During 2010 the following students carried out their project work for the BSc or the MSc at MAX-lab: Colleen Allen, Mohamed Luis El-Sheikh, Ulrika Forsberg and Matty Litwack.

REPORTS FROM NUCLEAR PHYSICS AT MAX-Iab

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MAX-lab ACTIVITY REPORT 2010 515

Total Photo-absorption Cross Section of 6,7 Li Below π -threshold

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The study of few body systems via electromagnetic probes is of particular interest to the field of nuclear physics. The electromagnetic interaction of the photon (both real and virtual) with the nucleus is well known and due to progress in theoretical techniques the final state of reactions involving these systems can now be described exactly. This allows the different components of the theoretical descriptions of such reactions, like the choice of nucleon-nucleon (NN) potential used, to be tested without having to resort to using approximations, as is necessary for heavier nuclei.

The Li nucleus is of particular interest as calculations which employ the Lorentz Integral Transform method [1] exist for both the ⁶Li and ⁷Li isotopes [2, 3] with which experimental measurements can be compared. Results of these calculations for the total photo-absorption cross section of ⁶Li and ⁷Li can be seen in figure 1; the ⁷Li calculations are compared to data from a previous measurement [4]. As can be seen in figure 1a the use of different *NN*-potentials yields different results for the calculated cross section. With a measurement of high enough precision these different potentials can be compared to see which gives the best description of the data.

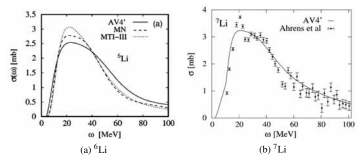
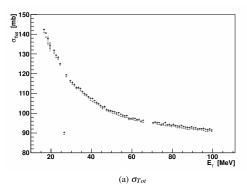


Figure 1: Theoretical calculations for the total photo-absorption cross section for ⁶Li [2] and ⁷Li [3]. The data in figure 1b are from a previous measurement [4].

The aim of this experiment is to measure the total photo-absorption cross section of the ⁶Li and ⁷Li isotopes. The cross section will be measured using the photon flux attenuation method. The equation

$$N(E_{\gamma}) = N_0(E_{\gamma}) \exp\{-n\sigma_{tot}(E_{\gamma})\}$$
 (1)

gives the flux of photons $N(E_{\gamma})$ at a photon energy E_{γ} that pass through a piece of matter of number of nuclei n and total photo-absorption cross section $\sigma_{tot}(E_{\gamma})$ for an incident photon flux of $N_0(E_{\gamma})$. The cross section is determined



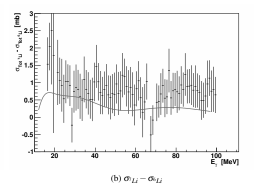


Figure 2: Figure 2a shows the measured total cross section for the two Li targets, red points are ⁶Li and black points are ⁷Li. Figure 2b shows the difference between the two measured Li total cross sections compared to a theoretical calculation [5] of the difference between the two Li nuclear cross sections as shown by the red line.

by taking the ratio of $N(E_{\gamma})/N_0(E_{\gamma})$ where $N(E_{\gamma})$ and $N_0(E_{\gamma})$ are determined by measuring the photon flux with and without the material in the incident photon beam. The total cross section σ_{tot} consists of two parts

$$\sigma_{tot} = \sigma_a + \sigma_n \tag{2}$$

where σ_n and σ_a are the total nuclear and atomic cross sections respectively. The atomic cross section is about two orders of magnitude greater than the nuclear cross section and will be subtracted from the measured total cross section leaving the nuclear cross section. The atomic cross section is well known and will be determined using a Monte Carlo simulation of the experimental set-up.

A first beam time took place in September 2009 and analysis of these data is underway. The experimental set-up consisted of a target frame with four targets, an empty one (to determine $N_0(E_\gamma)$), the two Li targets and a 12 C target for normalisation and monitoring purposes. A BaF₂ calorimeter detector was used for detection of photons that traversed the target material with a veto detector placed in front of this to remove charged particles produced in the target. Two incident photon energy ranges were used for the measurement, $17 \le E_\gamma \le 70$ MeV and $60 \le E_\gamma \le 100$ MeV.

The measured total cross sections for the two Li targets are shown in figure 2a. An accidental subtraction has been applied as well as a correction for background events in the experimental hall. Further corrections are necessary and are expected to shift these results a bit.

Figure 2b shows the difference between the two measured cross sections of figure 2a ($\sigma_{^7Li} - \sigma_{^6Li}$). This difference roughly represents the difference in nuclear cross sections as the atomic cross section should be very similar for both nuclei. The data are compared to the calculated difference in nuclear cross sections [5] shown by the red line where the Argonne V4' NN potential has been used. The theoretical difference is smaller than the experimental one for the whole incident photon range covered. The discrepancy could be due to different things being compared in the figure and this may improve after the analysis of the measured data is completed.

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Helium Photodisintegration using the Gas-Scintillator Active Target

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The measurement of photodisintegration reactions on few-body nuclei, from breakup threshold to pion production threshold, is a major component of the MAX-lab nuclear physics programme. The main motivation is to test the latest ab initio methods to describe few-body nuclear properties, based either on the best phenomenological nucleon-nucleon interaction potentials, or alternatively on potentials derived using Chiral Perturbation Theory. Photonuclear observables are sensitive to the input to these theoretical calculations [1, 2], which are in urgent need of experimental verification. The present series of experiments is designed to measure total and partial cross sections for ⁴He photodisintegration, using a gas-scintillator Active Target (AT), and extends previous measurements at MAX-lab [3].

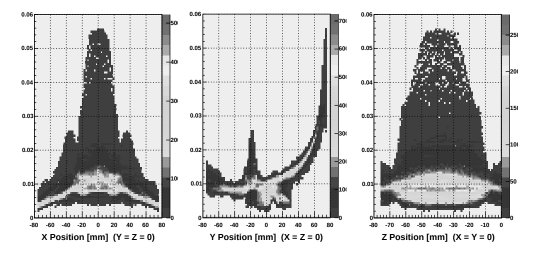


Figure 1: False-colour plot of Monte Carlo calculations of the probability of a scintillation photon (y-axis of plot) reaching the photocathode of a single photomultiplier. Photons have been generated along the X-axis (horizontal), Y-axis (vertical) and Z-axis (along beam axis) within a single AT cell. For events produced along the central axis of the target cell there is around 1% probability of a photon reaching the PMT.

The experiment was performed at MAX-lab in 2009 and the AT operated at a pressure 20 bar, running comfortably at the maximum available tagged-photon beam intensity. An electron beam energy of 164.8 MeV was employed and the broad-band tagging spectrometer produced photons of known energy in the range 11.5 - 67.4 MeV. Data taking was performed with different trigger conditions to help to determine the effects of AT detection efficiency in the energy range close to breakup threshold.

In addition to the AT, 2, 10-inch NaI(Tl) counters were employed to detect coincident, Compton-scattered photons from $^4\text{He}(\gamma, \gamma'$ $^4\text{He})$, and an array of 12 liquid scintillators (Nordball) detected neutrons from $^4\text{He}(\gamma, Xn)$ reactions. Details of the experiment can be viewed at the *Electronic Logbooks for Nuclear Physics at MAX-lab* (http://np1.maxlab.lu.se:7713/). The first analysis objective, to obtain the inclusive γ + 4 He cross section, is making good progress. It will be followed by analyses of partial reaction channels, producing

coincident photons or neutrons in the final state, which are not stopped in the robust walls of the AT. This will provide a feasibility study of future experiments on Compton scattering from He and also a measurement of ${}^4\text{He}\,(\gamma,n)$. The AT signal gives the time reference to determine neutron momentum by time-of-flight and in this case the random-coincidence background is largely unaffected by the complicated stretched-beam time structure.

The analysis to obtain the inclusive reaction yield as a function of incident photon energy has proved relatively fast, once procedures to extract the AT-Tagger coincidence signal from a strongly fluctuating background was perfected. The background arises from a variety of periodic processes within the MAX-I stretcher ring. However the conversion from a yield to a cross section requires an evaluation of the AT detection efficiency and also of background processes within the AT. Thus much of the data analysis effort has gone into the development of a Monte Carlo model of the AT, based on the CERN toolkit Geant-4. This model is vital to understand the effect of trigger thresholds and background on the shape of the excitation function, as close to the photodisintegration energy threshold the final-state, particle energies are small.

The Monte Carlo calculations have been split into 2 sections.

- 1. The calculation of the scintillation light transport through the target to the photomultipliers: this has been done to model the position dependence of the signal amplitude from the target. Optical photons were generated from known points inside the target and the numbers reaching the photocathode of a PMT calculated. The energy spectrum of those photons followed that of N₂ fluorescent emission. In total 9 × 10⁹ photons were generated and used to build a 3-D map of the position dependence of the light-collection efficiency within a target cell. A sample of results showing the variation of photon collection efficiency along the x, y and z AT coordinate axes is shown in Fig.1. These calculations are being compared to pulse-height calibration measurements made with an α particle source placed inside the target cell.
- 2. The modeling of the photo reaction processes within the target: this includes not only the γ + 4 He interactions of interest, but also background processes such as interactions in the Be and Al-mylar windows. Atomic interactions such as Compton scattering and pair production are also considered. The energy and angle dependence of 4 He(γ , X) reactions was taken from previous measurements, theoretical calculations or modeled with phase-space distributions were this information is not available. The energy loss of charged reaction products was calculated and weighted, using the light-collection efficiency map described above, to produce a simulated pulse height. From this the detection efficiency for a given pulse-height threshold is being calculated.

The scintillation transport calculations are now complete and work is in progress to evaluate the model dependence of efficiency and background corrections to the inclusive excitation function. This work is being performed as part of the Glasgow Ph.D. thesis of S. Al Jebali, which will be submitted in 2011. When this is complete an analysis of the exclusive channels, involving detection of an external photon or neutron will be made. A new measurement of ${}^4\text{He}(\gamma, n\, {}^3\text{He})$ [5] is scheduled for 2011 and will be performed in collaboration with the Karkov and Lund groups. This will use linearly polarised photons, produced by coherent bremsstrahlung on a diamond radiator, to measure the Σ beam asymmetry. After this a series of measurements on ${}^3\text{He}$ is planned.

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The new focal plane hodoscope at the tagged photon facility

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The Main Tagger (MT) at MAX-lab was moved from the Saskatchewan Accelerator Laboratory (SAL), Canada, to MAX-lab, in connection with the upgrade of the tagged photon facility. In addition, a hodoscope consisting of 63 overlapping plastic scintillators became available. The energy resolution depends on the incoming electron energy E_0 and the magnetic field of the tagging magnet. The central momentum of the MT, $P_0 = 198.6 \cdot E_0 / S$, where S is a number related to the exit path of the direct electron beam towards the beam dump. We use S = 345 or 475. The focal plane of the MT is linear from $0.8 \ P_0$ to $1.4 \ P_0$ and slightly curved over the range $0.6 \ P_0$ to $0.8 \ P_0$. The length of the focal plane is about 1240 mm. It is possible to use also parts of the focal plane that are outside the range $0.8 \ to 1.4 \ P_0$, however, the characteristics of this part has to determined for each setting using a NaI spectrometer in the direct tagged photon beam. The energy resolution is typically about 500 keV.

A new hodoscope is under construction funded by the Knut and Alice Wallenberg Foundation and by MAX-lab. The hodoscope consists of 160 scintillators, each 10.5 mm wide and 3 mm thick. The trajectories of the electrons form a 45° angle with respect to the focal plane and will pass the scintillators at 90°, see figure 1. The dimensions are chosen to obtain as high an energy resolution as possible using the smallest available PM tubes (Hamamatsu R1635). The total length of the new focal plane hodoscope is 1200 mm. The light guides from the scintillators to the PM tubes are designed to allow for two rows of PM tubes, every second PM tube is along one row. Figure 1 shows one of these rows. In reality the row is displayed with respect to the scintillators. The distance between the rows is determined by the size of the mymetal shield (diameter 15 mm) around the tubes.



Figure 1

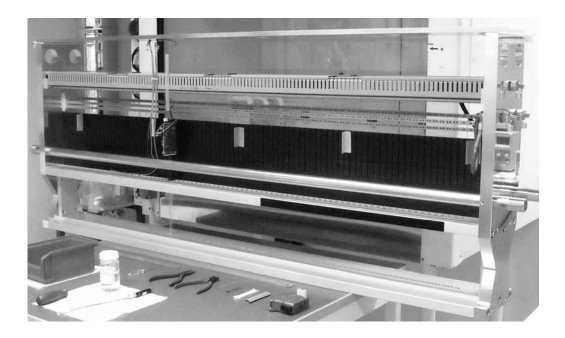
Each PM tube is mounted on a separate HV/discriminator card similar to the Glasgow design used at the tagging system at MAMI in Mainz. Each card has an analogue signal and two ECL signals, the three signals will be transported about 30 m to the counting rooms outside the shielded experimental area. The cards make it possible to require an overlap coincidence between two adjacent scintillators to improve the energy resolution, however, the cards can also be used in a stand-alone mode.

Examples of energy ranges and energy resolutions expected at the hodoscope are given in Table 1 for the MT with two values of S. The ET will provide tagged photon energies as close as 10 MeV from the endpoint of the bremsstrahlung spectrum.

Table 1

Electron energy (MeV)	Photon energy range (MeV)	Photon energy resolution (MeV)	Tagging system
140	12 - 75	0.39	MT (345)
165	68.4 - 123.6	0.35	MT (475)
190	78.8 - 142.3	0.40	MT (475)
190	15.0 - 102.5	0.55	MT (345)

All parts of the new hodoscope are now available and the assembly is in progress. Figure 2 shows the frame that will be used to hold scintillators, discriminator cards, cables etc.



RESPONSE OF THE PANDA ELECTROMAGNETIC CALORIMETER TO PHOTONS IN THE ENERGY RANGE 10 – 120 MeV.

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The physics programme at the PANDA experiment at FAIR (Facility for Antiproton and Ion Research) relies heavily on the capability to measure photons with good energy, position and timing resolution over a wide dynamic range starting at a few MeV up to several GeV energy. The identification of a single photon as well as the coincident detection of the decay photons from multi-meson exit channels requires a high and homogenous efficiency. The invariant mass reconstruction requires also a precise determination of the opening angles between decay photons. This can only be achieved by a highly granular calorimeter. A compact scintillator material with a short Moliére radius will provide a minimum occupancy of the detector modules. These requirements makes the electromagnetic calorimeter one of the most difficult and expensive part of the whole PANDA detector. The chosen detector material is PbWO₄ (PWO). Very encouraging results have been obtained by operating such crystals at low temperatures down to -25° C where the light yield increases by a factor of 2.5 as compared to room temperature. A new material composition for the PWO crystals (PWO-II) has given an additional increase of the light output by a factor more than 2. This will give a substantially increased performance with respect to the PWO crystals used at the CMS experiment at LHC. The electromagnetic calorimeter of the target spectrometer consists of a barrel part with 11 360 crystals of 11 different shapes, a forward end cap with 3600 crystals and a backward end cap with 592 crystals. Avalanche Photo Diodes (APDs) will be used for the barrel part and Vacuum Photo Triods (VPTs) are anticipated for the forward end cap.

Four Swedish groups are members of the PANDA collaboration, the nuclear physics groups at KTH, Lund University, Stockholm University and Uppsala University. The Swedish groups have jointly decided to focus on the electromagnetic calorimeter with special emphasis of the forward end cap as the Swedish hardware contribution to PANDA. By a grant from the Swedish Research Council approximately half of the crystals for the forward end cap have been purchased.

A good understanding of the EMC response to low-energy photons is very important in order to reach the desired resolution in PANDA. MAX-lab is a unique facility for the investigation of the PWO response to low-energy photons, i.e. linearity and energy resolution. The response at higher energies will be measured at MAMI, Mainz and ELSA, Bonn. In a series of experiments the response of PWO crystals to photons in region 10 - 120 MeV is measured at the tagged photon facility at MAX-lab. In March of 2009, a 5 x 5 matrix of crystals for the forward end cap of PANDA, equipped with photo-multiplier tubes, was inserted in a collimated photon beam. An unexpectedly low resolution ($\sigma \approx 2.6$ MeV) for photon energies around 25 MeV compared to the resolution at higher and lower energies ($\sigma \approx 2.4$ MeV at 32 MeV and $\sigma \approx 1.7$ MeV at 13 MeV) was observed. We concluded from the data that this effect

was most probably due bad resolution of the tagging facility for electrons with momenta 40% larger than the central momentum in the tagging magnet, rather than a property of the PWO matrix..

This problem was studied in January 2010. The energy resolution (and the absolute energies) of the tagged photon beam was independently measured using a large sodium iodide detector (CATS). Preliminary results are presented in Figure 1. The bad resolution of the tagger is clearly seen in the energy range 16-33 MeV. For the absolute energies we obtained a result deviating by 7% from the nominal energies

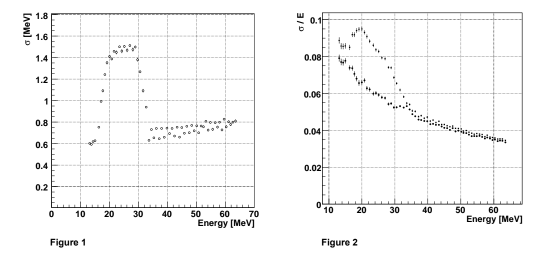


Figure 1: The resolution (σ = FWHM/2.35) as a function of photon energy for the CATS detector. In the analysis of the response the contributions from the central part and the peripheral segments were added.

In addition, a new prototype for the forward end cap was tested. The prototype houses 64 crystals but was in a first stage equipped with 25 crystals in a 5 x 5 matrix configuration using photomultiplier tubes for read-out. The response was measured at -25 $^{\circ}$ C. Two different pulse-handling electronics were used. In the first preamplifiers, shaping amplifiers and peak-sensing ADCs were used. In the second, the anode signals were fed directly to flash-ADCs. In figure 2 we show the result for the relative resolution, before and after correcting for the inherent tagger resolution.

Figure 2: The relative resolution σ/E ($\sigma=FWHM/2.35$) as a function of photon energy for the 5x5 crystal forward end cap prototype at -25 ^{o}C , measured (filled squares) and corrected for the energy resolution of the tagger (filled circles).

Detection of positively charged pions with a CsI/SSD telescope

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Pion photoproduction experiments are one of the few ways to explicitly test predictions made at low energies by QCD based effective theories. The research program within the Nuclear Physics group at MAX-lab focuses strongly on this area.

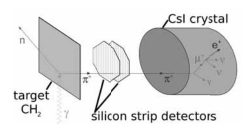


Figure 1: Diagram of a π^+ produced by a photon interacting with a proton in a CH₂ target, which is then detected by the CsI/SSD telescope.

One of the detector setups used in the detection of positively charged pions is a ΔE - ΔE -E telescope formed by two thin silicon strip detectors (SSD) and a CsI(Tl) crystal. The two SSDs are 0.5 mm thick, 64 mm wide, octagonal in shape and are comprised of 64, 1 mm wide strips. The strips are read out in groups of two, which, for a separation distance between the SSDs of 15 mm, gives an angular resolution of approximately 15°. The CsI(Tl) crystal is cylindrical in shape, 4 inches in length with a 5 inch diameter, read out by a 5 inch photomultiplier attached to the end. The setup was arranged with the first SSD placed at either 90° or 60° to the photon beam, approximately 10 cm from a 1 mm thick CH₂ target lying in the photon beam path, and the CsI(Tl) crystal placed 15 mm behind the second SSD, as depicted in Figure 1. In this arrangement, the solid angle of the telescope is set by the second SSD,

and is approximately 330 msr. The separation of particles of different mass is accomplished with the standard Δ E-E method, by examining the energy deposited in the SSDs compared to that deposited in the CsI(Tl). An example of this is shown in Figure 2-a, where clear separations between different particles are seen. This method however cannot be used to distinguish between positively and negatively charged pions. To separate pions with opposite charges from each other, the decay signature of the π^+ was searched for. A π^- entering the CsI(Tl) detector is quickly captured by a nucleus, while a π^+ is repelled by a nucleus and decays according to the following two step process:

$$\begin{split} \pi^+ \to \mu^+ + \nu & \left(\tau_{\pi^+} = 26.0 \, ns, \, E_{\mu^+} = 4.12 \, MeV \right) \\ \mu^+ \to e^+ + 2\nu & \left(\tau_{\mu^+} = 2.197 \, \mu s, \, E_{e^+} = 5 \to 50 \, MeV \right) \end{split}$$

The use of a 20 MHz Flash ADC attached to the CsI(Tl) detector makes it is possible to look for the presence of this process. An example of events recorded in the Flash ADC is shown in Figure 3. Choosing events such as these, with certain peak separations and heights, it is possible to better separate positively charged pions from other particles, as was done in constructing Figure 2-b.

An experimental beam-time dedicated to the photoproduction of positively charged pions took place over 4 weeks during June and July of 2010. During this time, various types of targets were used in conjunction with the CsI/SSD telescope, with the emphasis being on ${\rm CH_2}$, of which approximately 180 hours of data were recorded. The aim of this experiment was to study the photoproduction of pions from the hydrogen nucleus, therefore background measurements with a pure Carbon target were also performed, 40 hours of such data were recorded. Photons striking the target were produced via bremsstrahlung using electrons with energies of 193 MeV.

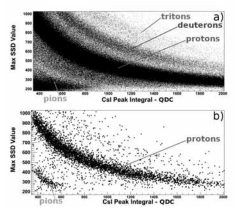


Figure 2: Energy deposits (in arbitrary units) in the first SSD versus in the CsI(Tl) crystal, a) without any event selection, and b) with cuts on the pulses seen in the CsI(Tl) using the Flash ADC. Pions are clearly separated from protons and from the background in b).

With the detector at 90° to the photon beam path and a 1 mm thick CH_2 target in place, Figure 4 shows the time coincidences between the focal plane and experimental detectors, with events in the coincidence spike and the random regions having been used to produce 5. The time separation between the two Flash ADC pulses for selected events can be seen in Figure 6, which has a distribution of that which is expected for the second pulse to have come from μ^+ decay. Figures 4 and 6 together are strong evidence that tagged pions have been detected.

Having tagged pions means that individual photons can be associated with the pions they produced through interactions with the target. A full analysis of the data is still in progress, however preliminary results appear promising with approximately 2700 tagged pions being detected from all targets and detector angles by the CsI/SSD telescope.

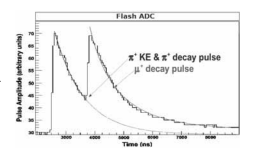


Figure 3: Readout of the CsI(Tl) detector as a function of time for one event using the Flash ADC.

The amount of data existing world wide for the process $\gamma + p \to \pi^+ n$ is lacking at energies close to pion production threshold. Eventually, differential cross sections will be calculated for this process at various angels and energies, in order to fill-in the gaps in the world data set. These measurements will then be compared with various competing theoretical predictions.

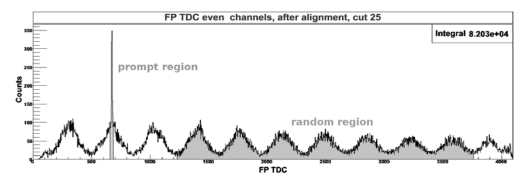


Figure 4: Time separation between the electrons hitting the focal plane detector and reaction products striking the telescope. The large peak on the left is from true coincident events, the rest are random events. The highlighted random region is used to subtract the random contribution that is present under the coincidence peak.

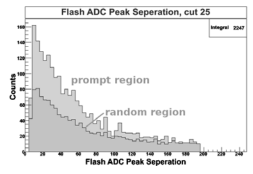


Figure 5: Time separation between the two pulses in the Flash ADC for both random and prompt events where the random contribution has been scaled down. This is data for a particular target and detector angle.

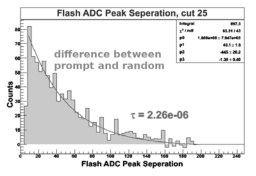


Figure 6: The difference between the prompt and random distributions shown in Figure 5. The fit to the distribution gives strong support to these events being the result of a π^+ decay, which decays into a μ^+ , which in turn has a lifetime of 2.197 μ s.

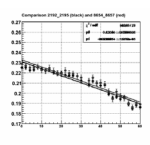
International Research Experience for Students (IRES)

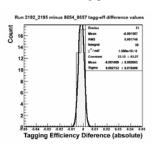
H. Caceres ^a, K. DiBenedetto ^a, K. England ^b, D. Kelleher ^b, M. Litwack ^c, and B. Smith ^a, W.J. Briscoe ^a, N. Benmouna ^d, K.G. Fissum ^e and G.V. O'Rielly ^b

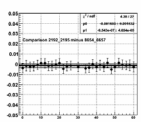
a) The George Washington University, Washington DC 22201, USA
b) University of Massachusetts Dartmouth, North Dartmouth MA, 02740, USA
c) Georgia Institute of Technology, Atlanta, GA 3033, USA
d) Montgomery College, Rockville, MD 20850, USA
e) Lund University & MAX-lab, Lund SE-221 00, Sweden

Supported by the US National Science Foundation's Office of International Science and Engineering through an *International Research Experience for Students (IRES)* grant (Award 0853760), five undergraduate students from three US institutions participated in the pion photoproduction experiment NP-014 being performed by the PIONS@MAXLAB Collaboration in June 2010. Mr. H. Caceres, Ms. Kristina DiBenedetto and Mr. Brian Smith from The George Washington University, Ms. Khayla England and Mr. Daniel Kelleher, both from the University of Massachusetts Dartmouth, and Mr. Matty Litwack from Georgia Institute of Technology were integral members of the collaboration and participated in the detector and target set-up, took data acquisition shifts monitoring the apparatus and data quality, and were each responsible for a research project involving system calibration or event analysis. The work performed by the students was crucial for the success of both their projects and the measurements.

Mr. Caceres and Mr. Litwack each had a project which involved the photon tagging efficiency, which determines the fraction of bremsstrahlung photons which make it to the target. Mr. Litwack



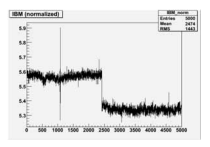


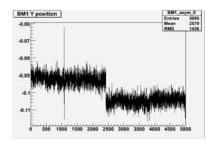


The top left panel shows the tagging efficiency for each tagger channel with (red) and without (black) a background subtraction made. The lower left panel shows the channel-by-channel difference for the tagging efficiency compared with base-line values. These differences are plotted as a histogram in the top right panel.

worked on developing a GEANT-4 monte-carlo simulation of MAX-lab electron photon tagging system in order to understand fully this property of the tagged photon beam. Mr. Caceres was responsible for the daily dedicated calibration runs to measure the tagging efficiency. The resulting data were analyzed to determine the tagging efficiency for the individual tagger channels. By comparing each day's value with previous measurements is was possible to create an estimate of the overall systematic uncertainty in the tagging efficiency. Knowledge of this quantity is essential to the determination of the absolute cross sections from these measurements.

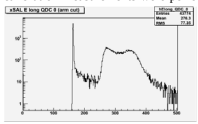
Ms. DiBenedetto's analysis project involved incorporating new photon beam monitoring hardware (X-Y beam position monitors and an in-beam photon flux monitor) into the measurement data acquisition system and online display. These beam monitors were essential for maintaining the quality and behaviour of the photon beam, quickly signaling even small changes in the beam parameters.





The left panel shows the in-beam flux monitor (IBM) rate normalized to the tagger focal plane rate. The sudden drop in this ratio was due to a change in the current to one of the MAX-I ring magnets. This change caused a shift in the steering of the electron beam. This mis-steering was also seen in the photon beam vertical (Y) position shown in the right panel.

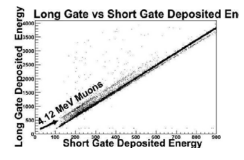
Ms. England worked on the gain measurements for the xSAL counters. These important calibration measurements were performed each day during the NP014 run period. The gamma-

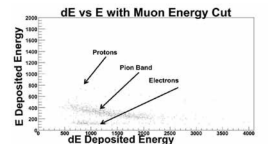


ray spectrum produced by a Th-C γ -ray source was measured and recorded for analysis. By fitting this spectrum to extract the pedestal value and the position of the Compton edge, a two-point fit to these points was made to determine the energy calibration for each of the xSAL E-counters. These calibration measurements were used to monitor for gain changes in the xSAL detectors. This is necessary in order to accurately determine the pion energies from the $\gamma p \to n \pi^+$ reaction.

The γ -ray spectrum recorded for one of the xSAL counters (located at 90°). The pedestal (near channel 160) and the Compton edge (near channel 360) are clearly seen. This data will be fitted with a gaussian peak and a smooth background to best determine the Compton edge position.

Mr. Kelleher and Mr. Smith worked on the data analysis code to optimize the event identification procedure used to separate the pion events from the very large proton and electron background. This event analysis was based on identifying the 4.12 MeV muons from the $\pi^+ \rightarrow \mu^+$ decay from the energy deposited in a long-gated versus short-gated QDC.





Elastic Compton Scattering on Deuterium

Compton@MAX-lab Collaboration

The Compton collaboration performed several runs in the past year. First of all, in March 2010, a week-long test run was done in order to investigate rate-dependent effects in the tagger. A simulation of the tagger had been developed which enabled such corrections to be modeled, but the simulation needed to have some benchmark data with which to validate its results. The test run used plastic scintillator paddles to generate x-triggers and a wide range of tagger rates was investigated. The data from this test run effectively served to "calibrate" the tagger simulation so that it could be used reliably to calculate correction factors (for effects such as stolen coincidences and ghosts) for our major runs on deuterium Compton scattering.

The next big production run on deuterium occurred in the October-November running period. This involved a 4-week run for more deuterium production data in the tagger energy range E_{γ} = 90-115 MeV, as well as a 2-week run on carbon as a further crosscheck on our normalization procedures. The carbon running was done under various conditions to mock up the actual run conditions of earlier production runs, including variations due to shaker frequency, tagger discriminator output width, and tagger rate. The tagger rates that were used for the carbon running covered the range from high-rate production running (~1 MHz) down to lower rates (200 kHz) as part of our investigation of rate-dependent effects.

The October-November runs utilized an electron beam energy of 165 MeV, giving a tagged energy range of 90-115 MeV. As part of this run, the multi-hit TDC's were used extensively, and also were compared to the output of the single-hit TDC's. The multi-hit units are not affected by stolen coincidences due to high rates, since they do not have a single stop, and so this comparison aided in the study of rate-dependent effects in the data. The beam duty factor was carefully monitored using the recently installed "duty-factor circuit" and so an online relative measure of the beam time structure was available as a diagnostic tool.

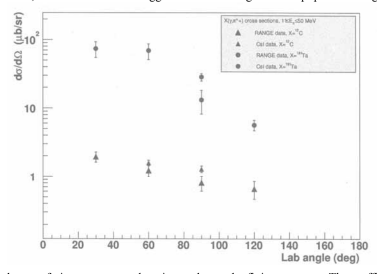
The NaI detectors BUNI, CATS and DIANA were placed at angles of 60°, 120° and 150°, respectively. This adds to our data set from November 2009 in which we measured 60°, 90° and 150°. These current runs (2010) were taken with essentially the same beam and tagger conditions as last year (2009), as related to the shaker frequency and the tagger discriminator width. These data from 2009 and 2010 constitute the Ph.D. thesis data of Khayrullo Shonyozov, who is a student at the University of Kentucky (thesis supervisor Mike Kovash).

Near-threshold π^+ production in photonuclear reactions with heavy nuclei

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Pion photoproduction cross sections on heavy nuclei close to threshold provide clean information on in-medium pion-nucleon interaction and pion propagation through nuclear matter, in contrast to (multi)hadron induced reactions where the hadronic cascade destroys this information grossly. (γ,π) experiments at higher energies have been performed at a few international facilities but in the near-threshold region data is scarce. Non-tagged data of this kind was however recorded at MAX-lab in 2007 and presented in Ref. [1] in 2008. Results from reactions in fairly light target nuclei (up to 27 AI) indicated that a deep (-20 MeV) optical potential is needed to explain the pion cross sections by cascade models.

The first experiment with tagged photons followed in 2009 where a heavy monoisotopic target nucleus, 181 Ta, was introduced. The tagger was here arranged to accept photon energies of 172 - 182 MeV.



Two kinds of detector systems were used to register pions, RANGE telescopes [1] utilizing the fast πu decay as signum for π^+ and a CSI telescope [2] that uses the 2.2 µs, µe decay to identify π^+ . The figure shows differential, $d\sigma/d\Omega$, cross sections of pions with energy > 11 MeV, coming from reactions in Ta and C. The registered number of pions has been corrected for decay-inflight, reactions in the detector material

losses of pions, muons and positrons due to the finite geometry. These efficiencies have been obtained via GEANT4 simulations. Since protons (22-108~MeV) could be registered without bias this gives an extra possibility to check the absolute normalization by comparing to existing (γ,p) data. The most important message is that both detector systems measure cross sections that are quite similar in the three overlapping points shown in the figure.

This data is under theoretical analysis. In particular we use the RELDIS cascade model for comparison [3]. Preliminary calculations indicate that the optical potential may be somewhat less deep than reported before and especially that its depth does not increase very much between Al and Ta.

In a (γ,π) run, dedicated for heavy targets (experiment NP08-02) in November 2010 we improved the statistics on pions produced in Ta reactions by some 10 times and extended the measurements to larger angles as well. The tagger was again set up to accept 172-182 MeV photons but due to some mis-match with energy measurements in (γ,d) reactions this acceptance is going to be controlled once more.

The analysis of the new data has just started and here we can only report that π^+ identification works well. Because of the improved statistics a serious attempt will be made to determine also π^- cross sections by subtracting the corrected number of π^+ from the total number of charged pions obtained from $\Delta E - E$ identification.

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MAX-lab ACTIVITY REPORT 2010

Positive Pion Photoproduction from the Proton Experiment NP-014

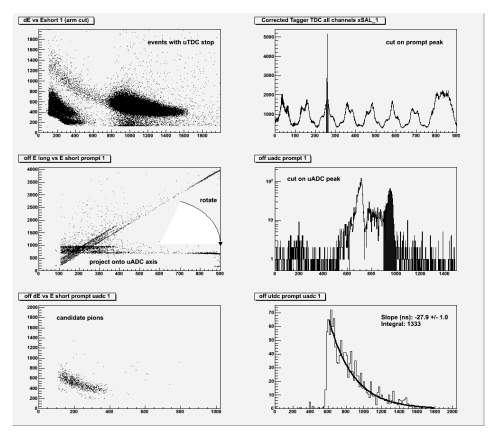
G.V. O'Rielly
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PIONS@MAX-lab Collaboration

One of the current challenges in nuclear science is to connect the observed properties of the nucleon with the theoretical framework provided by QCD [1]. One approach to address this question is to compare measurement results with theoretical predictions for those reactions where both can be done well. One such reaction is pion photoproduction from the nucleon in the nuclear regime. The development of modern descriptions of the strong interaction based on QCD can now provide detailed predictions for this reaction. Comparison of these predictions with the results from accurate measurements of these reactions serves as a stringent test of these QCD-based models. In addition, measurements of the differential cross sections for the $\gamma p \to n\pi^+$ reaction channel are needed to extract the p-wave contribution for comparison with the predictions from Chiral Perturbation Theory calculations. These measurements are also essential for the partial-wave analyses performed by the MAID and SAID groups. Despite this broad interest, there are few modern data sets for the $\gamma p \to n\pi^+$ reaction at energies below the Δ -resonance, with extensive angular distributions only at threshold and above 200 MeV, and no tagged photon data in the energy range between 155 and 185 MeV.

To address this need for new, high-quality data on the charged pion channels at energies below the Δ -resonance, the PIONS@MAXLAB collaboration is performing measurements of the $\gamma p \to n \pi^+$ reaction using the MAX-lab tagged photon facility. To detect the out-going pion, several different detector systems are being used. This report discusses the results obtained by the xSAL detectors, which are a set of four monolithic plastic scintillator counters configured as ΔE -E telescopes. The stopping counters have dimensions chosen to stop the highest energy pions produced by the tagged photon beam at MAX-lab. Complementing the measurements with the xSAL telescopes are measurements made using other detector systems with different pion identification methods; the RANGE and CsI detectors. The use of several independent detector systems will provide important cross-checks throughout the event analysis and for the final measurement results.

A dedicated four-week production run took place during June 2010. The xSAL detectors and trigger electronics were configured in the same manner as was used during the June 2009 run period. Some modifications were made to the experimental set-up in order to increase the event rate, including a larger detector acceptance and a faster DAQ system. The four xSAL telescopes were placed at laboratory angles of 90°, 135° and two at 45°, these angles have been identified as having the maximum sensitivity to differences between the existing MAID and SAID partial-wave analysis solutions [2]. Data were collected from C and CH₂ targets. Data for the $\gamma p \rightarrow n\pi^+$ reaction will come from the CH₂ target, with the C target used for background subtraction.

Due to the very low event rate for the (γ, π^+) reaction, standard particle identification techniques based on ΔE vs. E *stopping power* are insufficient to separate pions from the much larger electron and proton background. To reliably identify pion events, analysis methods based on detecting the $\pi^+ \to \mu^+$ decay in the xSAL E-counter were used. This technique identifies the 4.12 MeV muon based on the energy deposited in a short- versus long-gated QDC.



(top left) The ΔE vs. E stopping power for all events recorded in one of the xSAL counters. The pion events are swamped by background electrons and protons. (top right) The tagger TDC spectrum with the prompt coincidence peak identified in red. (middle left) The energy deposited in the long- vs. short-gated QDC. The candidate pion events are located above the equal energy band produced by the background events. This plot is rotated so that these event bands are horizontal. (middle right) The projection of the long vs. short QDC bands with the candidate pion events identified in blue. (bottom left) The ΔE vs. E stopping power for candidate pion events. The pion events are now clearly seen and well separated from the remaining background events. (bottom right) The time distribution of these candidate pion events exhibits a decay time of (27.9 \pm 1.0) ns, consistent with the known $\pi \to \mu$ lifetime.

This use of both energy and timing information from the π^+ decay in the xSAL E-counter enables the reliable identification of pion events with a well-understood detection efficiency. The analysis to determine final pion yields from each xSAL telescope is currently underway. Additional work is required to determine corrections to account for the misidentification and loss fractions due to the analysis cuts, and to determine the absolute π^+ -detection efficiency. Monte-Carlo simulations, using the *GEANT-4* package, of the experiment and detector systems are currently underway to determine the acceptance for each xSAL telescope. This will be combined with information from the photon tagger to determine the final differential cross sections for the $\gamma p \to n \pi^+$ reaction.

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Investigation of the MAX I photon beam using a pair spectrometer

A. Thelin

As for today the energy of the photons in the MAX I beam is determined by the energy of the electrons which produces them through Bremsstrahlung. To further investigate characteristics of the beam, it would be of interest to measure the energy of the actual photons. It was suggested that this could be done using a pair spectrometer. The whole system can be seen in Figure 1 and works as follows:

- The photon produced by Bremsstrahlung enters a converter.
- Photon is occasionally converted into an e⁻ e⁺ pair.
- e and e are separated by a magnetic field.
- Position sensitive detectors are hit by the particles.
- A second pair of detectors is placed behind the position sensitive detectors and act as triggers to determine if two particles hit the detectors simultaneous.
- The signal from the detectors are registered in a DAQ and analyzed.

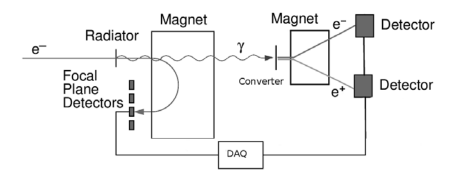


Figure 1. A simplified picture of the experimental setup.

Simulations made in Geant4 of the system, indicated that a relation between position of the e⁻ e⁺ pair and incident electron energy could be made. The results from the simulations were produced under near perfect conditions and real-life experiment were expected to produce more uncertain results, due to showers from shielding and a broad photon beam to name a few examples.

In the real experiment, a total of three runs were made, all with slightly different settings in order for maximum amount of information to be attained. The output signals from the two position sensitive detectors, registering the e⁻ and e⁺ respectively, are shown in Figure 2.

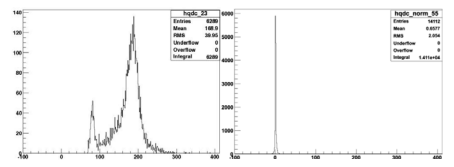


Figure 2. Left: Signal from detector A Right: Signal from detector B.

The two signals in Figure 2 are typical for all QDC channels. The output from detector A is believed to be caused by a reflection in the detector, possibly between the scintillating fibers and the PM tube in the position sensitive detectors. The abnormal signal from detector A makes an analysis difficult and no relation between incident photon energy and e e position could be established. However a more in depth analysis of the focal plane detectors, Figure 3, revealed that the detectors might have been placed too close center of the beam. This could also be a reason for the unexpected result.

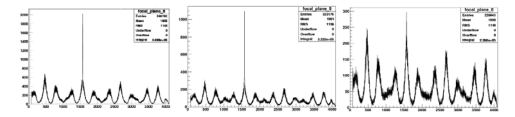


Figure 3. Output from one of the focal plane detectors for all three runs made in the experiment. This indicates that the detectors were placed too close the center of the beam.

Since expected results could not be attained from the experiment, and as simulations indicate that it should be possible, it is recommended that studies continue in this field, perhaps as another master thesis. It would be advisable to start by making sure that both position sensitive detectors function properly before continuing with the experiment.



Emelie Hilner, Department of Synchrotron Radiation Research at Lund University, preparing her presentation at the Annual User Meeting, 8 November 2010. Photo: Annika Nyberg

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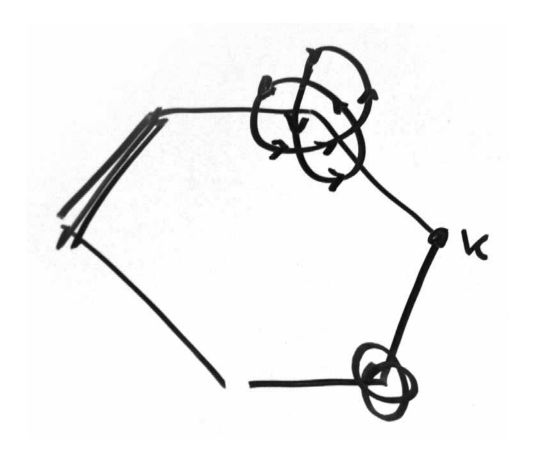
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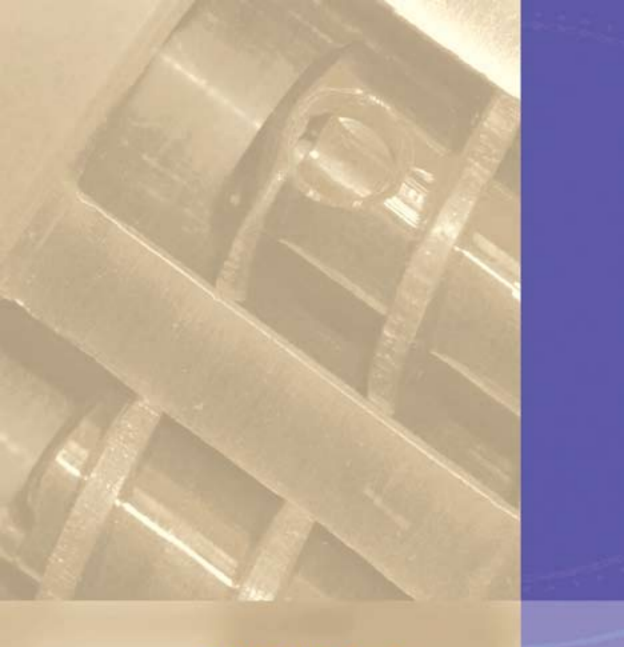
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