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Evidence for time dependent variation of glucagon secretion in mice.

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1	Evidence for diurnal variability of glucagon secretion
2	in mice
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26 **ABSTRACT**

27 Glucose metabolism is subjected to diurnal variation, which might be mediated by alterations in 28 the transcription pattern of clock genes and regulated by hormonal factors, as has been 29 demonstrated for insulin. However, whether also glucagon is involved in the diurnal variation of 30 glucose homeostasis is not known. We therefore examined glucagon secretion after meal 31 ingestion (meal tolerance test) and during hypoglycemia (hyperinsulinemic hypoglycemia clamp 32 at 2.5 mmol/l glucose) and in vitro from isolated islets at ZT3 versus ZT15 in normal C57BL/6J 33 mice and, furthermore, glucose levels and the insulin response to meal ingestion were also 34 examined at these time points in glucagon receptor knockout mice (GCGR-/-) and their wildtype 35 (wt) littermates.

36 We found in normal mice that whereas the glucagon response to meal ingestion was not 37 different between ZT3 and ZT15, the glucagon response to hypoglycemia was lower at ZT3 than 38 at ZT15 and glucagon secretion from isolated islets was higher at ZT3 than at ZT15. GCGR-/-39 mice displayed lower basal glucose, a lower insulin response to meal and a higher insulin 40 sensitivity than wt mice at ZT3 but not at ZT15. We conclude that glucagon secretion displays a 41 diurnal variability which is dependent both on intraislet and extraislet regulatory mechanisms in 42 normal mice and that the phenotype characteristics of a lower glucose and reduced insulin 43 response to meal in GCGR-/. mice are evident only during the light phase. These findings 44 suggest that glucagon signaling is a plausible contributor to the diurnal variation in glucose 45 homeostasis which may explain that the phenotype of the GCGR-/- mice is dependent on the 46 time of the day when it is examined.

48 Glucose metabolism displays circadian rhythm which is partially a result of dietary intake during 49 the active phase of the 24h period and maintenance of circulating glucose by hepatic glucose 50 production during the inactive phase (1). Glucose homeostasis is, however, also regulated by 51 the clock system, both by the clock genes in the suprachiasmatic nuclei in the hypothalamus and 52 by peripheral clock genes in many peripheral organs (1). Thus, each tissue contains its own 53 circadian clock-program that oscillates over the course of the 24 hour day and affects tissue-54 specific metabolic processes (2,3). Importantly, unlike the hypothalamus where the main time 55 giver (*zeitgeber*) is the light on the retina (4,5), food intake has been shown to be a stronger 56 zeitgeber in peripheral tissues (6). The intracellular signaling of clock genes consists of 57 interacting transcriptional positive and negative feedback limbs. The negative-feedback limb 58 involves three Period genes (Per1-3) and two Cryptochrome genes (Cry1 and 2) in the mouse, 59 whereas the positive-feedback arm involves the genes *Clock* and *Bmal1* (7). These genes 60 reciprocally regulate each other, establishing an oscillatory pattern of gene transcription. 61 The importance of the clock genes for glucose homeostasis is evident by findings that genetic 62 deletion of the clock transcription factor in the hypothalamus in mutant mice alters the diurnal 63 feeding pattern and results in overeating, obesity and a sign of metabolic syndrome 64 characterized with hyperglycemia and insulin deficiency (8). Furthermore, lesion in the 65 suprachiasmatic nuclei disrupts the circadian rhythm of glucose and insulin in mice (9). 66 Moreover, disruption in the transcription of *Clock* and *Bmal1* alters the expression of genes essential to beta cell function and leads to insulin deficiency and diabetes (10). The importance 67 68 of the clock system for glucose homeostasis and islet function is also emphasized by findings

69 that an autonomic rhythm exists within pancreatic beta cells (11,12) and that conditional 70 disruption of the clock in the pancreas results in impaired beta cell function and diabetes (13). 71 Recently, it was demonstrated that the pancreatic glucagon producing alpha cells is regulated 72 by the clock gene *Rev-erb alpha* such that silencing of this gene inhibits glucagon secretion 73 whereas a *Rev-erb alpha* agonist stimulates glucagon secretion (14). This would suggest that not 74 only insulin but also glucagon is the subjects of diurnal variation through clock regulation. This 75 would be of interest since glucagon stimulates hepatic glucose production which is a key 76 mechanism for preventing hypoglycemia during the inactive phase (1). Interestingly, it has also 77 been reported but not widely discussed, that the phenotype characteristic of a reduction in 78 circulating glucose in glucagon receptor knockout (GCGR-/-) mice is observed only in the 79 morning hours and vanishes later during the day (15), which may further indicate that glucagon 80 is involved in the diurnal variation of glucose homeostasis. 81 However, besides these studies there is little evidence linking glucagon signaling or glucagon

secretion to diurnal variation of glucose homeostasis. To gain further insight in the potential
involvement of glucagon in this respect, we compared the glucagon response to hypoglycemia
and meal test, and glucagon secretion from isolated islets between *zetigeber* time (ZT) 3 and
ZT15 in normal mice and compared glucose levels and insulin response to meal in GCGR-/- mice
and their wildtype littermates.

87

88 2. Materials and methods

89 2.1 Animals and anesthesia

90 Female C57BL/6J mice were obtained from Taconic (Skensved, Denmark) and housed on arrival

91 at 22° in a 12h light-dark cycle (6 am to 6 pm). The generation of GCGR-/- mice and their

92 wildtype littermates has been described previously (15). A standard research diet R34

93 (Lantmännen, Stockholm, Sweden) and water was provided *ad lib*. Mice were anesthetized prior

94 to all experiments using an intraperitoneal injection of midazolam (18 mg/kg animal, Dormicum,

95 Hoffman-La Roche, Basel, Switzerland) and Fluanisone/Fentanyl (41/9 mg/kg animal

96 respectably, Hypnorm, Janssen, Beerse, Belgium). All experimental procedures were performed

97 in agreement with the Animal Ethics Committee in Lund, Sweden. The experiments were

98 performed at ZT 3 (9 am) and ZT15 (9 pm) in regard to glucose homeostasis after meal challenge

and during hypoglycemia. Some data were also collected at ZT9 (3 pm) and ZT21 (3 am).

100 *2.2 Mixed meal tolerance test (MTT)*

101 The MTT was performed following 5 h of fasting. A 60/20/20E% Glucose/Protein/Lipid mixed 102 meal solution was administered as a 500 μL gavage as previously described (16). Blood samples 103 were collected from the retrobulbar intraorbital capillary plexus before (0 min) and at 15, 30, 45 104 and 60 min in the experimental series for measurements of insulin or at 5, 10 and 20 min in the 105 experimental series for measurements of glucagon following oral gavage. Plasma samples for 106 glucose and hormone determination were stored at -20° awaiting analysis.

107 2.3 Hypoglycemic hyperinsulinemic clamp

108 The hypoglycemic clamp was performed following 5 h of fasting. Surgery and clamp 109 experiments were performed as previously described (17) with the protocol modification of 110 returning of red blood cells (18). Briefly, the right jugular vein and the left carotid artery were 111 catheterized using catheters filled with heparinized saline (100 U/mL). The mice remained 112 anesthetized to reduce variation in the blood glucose concentrations due to stress. Following 113 baseline sampling, synthetic human insulin (Actrapid[®], Novo Nordisk, Bagsvaerd, Denmark) was 114 infused as a continuous infusion (15 mU/kg animal/min) at a pace of $2 \mu L/min$ for 90 minutes. Blood glucose in ~5 µL whole blood was determined every 10 minutes with an Accu-Chek Aviva 115 116 blood glucose monitor (Hoffman-LaRoche). A variable amount of a 10% glucose (Sigma-Aldrich, 117 MO, USA) solution was infused to maintain blood glucose levels at 2.5 mmol/L. Glucose 118 requirement to maintain target glucose was represented by the glucose infusion rate (GIR) during the final 30 min steady state of the clamp. 119

120 *2.4 Islet experiments*

121 Pancreatic islets were isolated at ZT3 and ZT15 by collagenase digestion and handpicked under 122 the microscope. Batches of freshly isolated islets were pre-incubated in HEPES balanced salt 123 solution containing 125 mmol/L NaCl, 5.9 mmol/l KCL, 1.28 mmol/L CaCl₂, 1.2 mmol/L MgCl₂, 124 25 mmol/L HEPES (pH 7.4), 5.6 mmol/L glucose and 0.1% fatty acid free BSA (Boehringer 125 Mannheim, Mannheim, Germany) at 37°C during 60 min. Thereafter, islets in groups of three 126 were incubated in 200 µl of the above described buffer but with 2.8 and 11.1 mM glucose without 127 or with addition of arginine (10 mM) at 37°C during 60 min. Aliquots of the buffer were 128 collected and stored at -20°C until analysis of insulin levels.

129 *2.5 Analysis*

Plasma glucose during the MTT was measured with the glucose oxidase method. Plasma and
medium insulin was analysed with sandwich immunoassay technique (ELISA; Mercodia, Uppsala,
Sweden) using double monoclonal antibodies according to manufacturer's protocol. Plasma
glucagon was analyzed with ELISA (Mercodia), using double monoclonal antibodies, according to
manufacturer's protocol.

135 *2.6 Calculations and statistics*

136 All data are presented as mean ± S.E.M. Basal insulin sensitivity during MTT was determined 137 with the quantitative insulin sensitivity check index (QUICKI) which has been well validated in 138 mice (19). Clamp insulin sensitivity (SI_{Clamp}) and glucose clearance per unit of insulin (Cl_{Clamp}) was 139 calculated as previously described (20). Comparisons between groups were performed using a 140 two-tailed Student's t-test (paired when applicable) or a 2-way ANOVA with a Holm-Sidak's 141 multiple comparison test post hoc. Comparisons within groups between time points were 142 performed using repeated measure ANOVA and difference from time point 0 min was calculated 143 post hoc using Holm-Sidak's multiple comparison test. Incremental area under the curve (iAUC) 144 was calculated using the trapezoidal rule.

145

146 **3. Results**

147 *3.1 Glucagon response to meal ingestion in normal mice*

Whereas baseline blood glucose did not differ between ZT3 and ZT15 (Fig. 1A), glucose
excursion after MTT was lower at ZT3 compared to ZT15 at 10 min (Fig. 1A). In contrast, there
was no significant difference in the glucagon response to MTT between ZT3 and ZT15 (Figs. 1B
and 1C).

152 *3.2 Glucagon response to hypoglycemia in normal mice*

153 To study the glucagon response to hypoglycemia, hyperinsulinemic hypoglycemic clamp at 2.5 154 mmol/L was undertaken at ZT3 and ZT15 in normal mice; at this glucose level a robust glucagon 155 response is provoked (18). Basal blood glucose or blood glucose during the clamp did not differ 156 between ZT3 and ZT15 (Fig. 2A) but the GIR needed to maintain target blood glucose of 2.5 157 mmol/L was significantly lower at ZT3 compared to ZT15 (Figs. 2B and 2C). Consequently, 158 insulin sensitivity (SI_{Clamp}) was higher at ZT15 than at ZT3 (4.8±0.9 vs 1.5±0.2 L/kg x min, 159 p=0.003, Fig. 2E) and so was glucose clearance per unit of insulin (2.1 \pm 0.5 vs 0.6 \pm 0.1 L²/kg x min 160 x mmol, p=0.006; Fig. 2F). The glucagon response to hypoglycemia was significantly higher at 161 ZT15 than at ZT3, both when measured in absolute concentrations $(7.5\pm1.2 \text{ vs } 3.3\pm1.5 \text{ pmol/L})$ 162 p=0.019; Fig. 2G) and when estimated as fold change over basal (4.8±1.2 vs 1.9±0.54, p=0.035;

163 Fig. 2H).

164 3.3 Glucagon secretion from isolated islets

165 Glucagon secretion from isolated islets from normal mice at 2.8 or 11.1 mmol/L was not

166 different at ZT3 versus ZT15. However, glucagon secretion in response to 10 mmol/L

- arginine was higher at ZT3 than at ZT15 both at 2.8 mmol/L and 11.1 mmol/L glucose (both
 p<0.001).
- 169 3.4 GCGR knockout alters the circadian rhythm of metabolism
- 170 GCGR-/- mice had lower circulating glucose than their wt littermates at ZT3 (4.2±0.2 versus
- 171 7.4±0.3 mmol/L, p<0.001) and ZT9 (5.5±0.2 versus 7.6 ±0.2 mmol/L, p=0.0002) but not at ZT15
- 172 (5.2±0.2 versus 5.4±0.1 mmol/L; Figs. 3A-C). GCGR-/- mice had also a lower insulin response to
- meal than wt mice at ZT3 and ZT9 but not at ZT15 (Figs. 3D-G). Insulin sensitivity, measured as
- 174 QUICKI after meal ingestion, was lower in GCGR-/- than in wt mice at ZT3 and ZT9, but not at
- 175 ZT15 (Fig. 3H).

177 **4.** Discussion

178 As most species, both mice and humans exhibit oscillatory patterns in behavior and 179 physiological functions over the course of the day (1). Central and peripheral gene clocks 180 regulate this and they are in turn regulated by the effect of light on the retina of the eye (4,5), 181 by food intake (21,22) as well as by specific metabolic hormones (23). In this study, we have 182 explored the potential role of glucagon in this respect by examining the glucagon secretion 183 during hypoglycemia and after meal ingestion as well as in vitro at ZT3 versus ZT15 in normal 184 C57BL/6J mice and basal and postprandial glucose levels were also examined at these time 185 points in GCGR-/- mice and their wildetype (wt) littermates. 186 A main general novel finding of this study is that there indeed is a diurnal variability in glucagon 187 secretion in normal mice. The detail of this variability is, however, dependent on the 188 experimental condition. Thus, whereas the glucagon counterregulation to hypoglycemia is lower 189 at ZT3 than at ZT15, arginine-stimulated glucagon secretion from isolated islets shows the 190 opposite pattern, being higher at ZT3 than at ZT15, and glucose-dependent glucagon secretion 191 from islets and the glucagon secretion to meal ingestion is the same at ZT3 and ZT15. Therefore, 192 the diurnal variability in glucagon secretion is complex and regulated both by islet and extraislet 193 mechanisms since many factors regulate glucagon secretion besides the capacity in the islet 194 alpha cells.

To study glucagon secretion during hypoglycemia we used our recently developed hypoglycemic clamp in mice, where we demonstrated a clear glucagon response when glucose levels were reduced (18). The glucagon response under this condition is complexly regulated by secretory

198 capability from the alpha cells when glucose levels are reduced in combination with stimulation 199 by other counter-regulatory hormones, such as epinephrine released from the adrenals, and the 200 autonomic nerves (24). Since we did not observe any diurnal variability in the effect of low 201 glucose on glucagon secretion from isolated islets between ZT3 and ZT15, our conclusion is that 202 the variability during hypoglycemia is not dependent on different glucose sensitivity in alpha 203 cells. Instead, the difference between the light and dark phase in glucagon response to 204 hypoglycemia may rather be caused by a diurnal variation in the other counterregulatory 205 hormones. The lower glucagon at ZT3 compared to ZT15 during hypoglycemia coincided with a 206 lower insulin sensitivity as judged by a lower glucose infusion rate to maintain the target 207 hypoglycemic glucose level during the clamp at ZT3. This may suggest a cross-talk between 208 insulin sensitivity and glucagon secretion such that when insulin sensitivity is lower, the 209 requirement for glucagon to restore hypoglycemia is more limited.

210

211 To examine the potential of glucagon variability during another physiological condition we used 212 a mixed meal test, by applying a recently developed model when a mixture of glucose, fat and 213 protein resembling a mixed meal was administered to mice (16). In this model, glucagon 214 secretion is stimulated, which is mainly achieved by a combination of fatty acids and amino 215 acids derived from the meal constituents. We found that there was no difference in the 216 glucagon response to meal ingestion when we compared ZT3 and ZT15, suggesting that in 217 contrast to the glucagon counterregulation to hypoglycemia, there is no evidence of a diurnal 218 variability in the glucagon response to meal ingestion. It was therefore a surprise when we documented a clear diurnal variability in glucagon secretion from islets in response to arginine, 219

both at low and high glucose, with a higher glucagon secretion at ZT3 than at ZT15. This shows
an interesting diurnal variability in the capacity to secrete glucagon, which is not reflected in a
similar difference in vivo with a more modest stimulation. The mechanism and potential
contribution of this diurnal variability needs now to be examined in more detail.

224 To examine whether the diurnal variability in glucagon secretion is important for glucose 225 homeostasis, we assessed the hormonal response to a meal ingestion in GCGR-/- mice and wt 226 controls. As reported previously, a characteristic phenotype in GCGR-/- mice is lower baseline 227 glucose, impaired insulin secretion after arginine stimulation and enhanced insulin sensitivity 228 (15,25). We confirm here that these mice, compared to their wt littermates, have reduced 229 baseline glucose, reduced insulin response to meal ingestion and increased insulin sensitivity. 230 However, the main finding in this respect is that these phenotype characteristics were evident 231 only at ZT3 and not seen at ZT15. This further suggests a potential contribution of glucagon to 232 diurnal variability of glucose homeostasis. However, the mechanisms explaining these 233 discrepancies between ZT3 and ZT15 in GCGR-/- mice remain to be established. Due to hyperproduction of pro-glucagon, the GCGR -/- animals have increased levels of both glucagon and 234 235 glucagon-like peptide-1 (GLP-1) (25), which might contribute to the phenotype in these animals. 236 However, although recent studies have suggested that GLP-1 has a circadian rhythm (26), it 237 remains yet to be shown whether it affects peripheral gene clocks. It would be interesting to 238 test this, as a recent paper did (27), in a double knockout model. Nevertheless, a consequence 239 of our findings is that the well known phenotype of these mice with reduction of glucose levels 240 and impaired insulin secretion (15,25) depends on the time of the day when it is measured. In a 241 broader perspective, this raises questions on results derived from other hormone-altering

242 mouse models where a diurnal variation in glucose homeostasis might exaggerate or occlude 243 the phenotype and it highlights the importance of time dependent effects in metabolic 244 phenotyping. Furthermore, the true circadian rhythm in glucose homeostasis with particular 245 attention to glucose needs now to be tested over the entire 24 hr period in at least two cycles. 246 In conclusion, glucagon secretion displays a diurnal variability which is dependent both on 247 intraislet and extraislet regulatory mechanisms in normal mice and, furthermore, the phenotype 248 characteristics of a lower glucose, reduced insulin response to meal and lower insulin sensitivity 249 in GCGR-/- mice are evident only during the light phase. Our results therefore suggest that there 250 is a link between glucagon signaling and the diurnal variation of glucose homeostasis and that 251 the phenotype of the GCGR-/- mice is dependent on the time of the day when it is examined.

ARTICLE INFORMATION

255	Author contributions. S.M designed study, performed experiments, analyzed data, drafted and			
256	wrote manuscript and the final version of manuscript. B.A. designed study, analyzed data, wrote			
257	and revised manuscript and approved final version. B.A is the guarantor of this work and, as			
258	such, had full access to all the data in the study and takes responsibility for the integrity of the			
259	data and the accuracy of the data analysis.			
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265	Duality of Interest. The authors have nothing to disclose in relation to this study.			

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339 **FIGURE LEGENDS**

Figure 1—Plasma glucose (A) and glucagon levels (B), and fold change increase in glucagon
compared to basal (0 min; C) during a MTT in female C57BL/6J mice at ZT3 (open circle) or at
ZT15 (square). Mean±SEM are shown, n=7 for each group, *p<0.05 paired comparison between
groups.

Figure 2—Blood glucose levels (A), cumulated glucose infusion (B) and steady state glucose
infusion rate (GIR;C) during a hyperinsulinemic hypoglycemic clamp in female C57BL/6J mice at
ZT3 (open circle/white bars) or at ZT15 (square/black bars). Steady state glucose is obtained
during the last 30 minutes of the experiment. Insulin levels (D), insulin sensitivity index (SI; E),
glucose clearance per unit of insulin (CI; F), glucagon levels (G) and fold change increase in
glucagon compared to basal (H) during hyperinsulinaemic hypoglycaemic at ZT3 (open
circle/white bars) or at ZT15 (square/black bars). Mean±SEM are shown, n=8 for each group,

- 351 *p<0.05, **p<0.01, ***p<0.001, *p<0.05 comparison between groups, #p<0.05 compared to
 352 basal (0 min) for each group.
- **Figure 3**—Plasma glucose levels (A-C), iAUC of insulin levels (D), plasma insulin levels (E-G) and
- insulin sensitivity measured through QUICKI (H) during a MTT in female C57BL/6J mice at ZT3 (A
- and E), ZT9 (B and F) and ZT15 (C and G) in GCGR-/- (square/striped bar) and wt mice
- 356 (circle/white bar). Mean±SEM are shown, n=18-20 for each group, *p<0.05, **p<0.01,
- 357 ***p<0.001, *p<0.05 comparison between groups, [#]p<0.05 compared to ZT15 minutes for each
- 358 group.
- 359

360 Table 1 Glucagon secretion from isolated islets from C57BL/6J mice after incubation for 1 hr in

361 glucose at 2.8 mmol/L or 11.1 mmol/L without or with addition of aarginine at 10 mmol/L at ZT3

362 or ZT15. A total of 24 incubations with 3 islets in each from 3 mice were performed. Means \pm

- 363 S.E.M. are shown.
- 364

	Glucagon ZT3	Glucagon ZT15
	(pg/islet/hr)	(pg/islet/hr)
2.8 mmol/L glucose	3.9±0.7	3.6±0.7
2.8 mmol/L glucose +10 mmol/L arginine	9.2±1.0	6.3±0.9
11.1 mmol/L glucose	3.0±0.4	3.1±1.0
11.1 mmol/L glucose + 10 mmol/L arginine	5.3±0.7	2.4±0.3

365 366

FIGURES

369 Figure 1









373 Figure 3374

