HAMLET interacts with lipid membranes and perturbs their structure and integrity.

Mossberg, Anki; Puchades, Maja; Halskau, Øyvind; Baumann, Anne; Lanekoff, Ingela; Chao, Yinxia; Martinez, Aurora; Svanborg, Catharina; Karlsson, Roger

Published in:
PLoS ONE

DOI:
10.1371/journal.pone.0009384

Published: 2010-01-01

Citation for published version (APA):
HAMLET Interacts with Lipid Membranes and Perturbs Their Structure and Integrity

Ann-Kristin Mossberg1*, Maja Puchades2*, Øyvind Halskau3, Anne Baumann3, Ingela Laneckoff2, Yinxia Chao4, Aurora Martinez2, Catharina Svanborg1,4, Roger Karlsson2

1 Section of Microbiology, Immunology, and Glycobiology, Institute of Laboratory Medicine, Lund University, Lund, Sweden, 2 Department of Chemistry, University of Gothenburg, Göteborg, Sweden, 3 Department of Biomedicine, University of Bergen, Bergen, Norway, 4 Singapore Immunology Network, A*STAR, Singapore, Singapore

Abstract

Background: Cell membrane interactions rely on lipid bilayer constituents and molecules inserted within the membrane, including specific receptors. HAMLET (human α-lactalbumin made lethal to tumor cells) is a tumoricidal complex of partially unfolded α-lactalbumin (HLA) and oleic acid that is internalized by tumor cells, suggesting that interactions with the phospholipid bilayer and/or specific receptors may be essential for the tumoricidal effect. This study examined whether HAMLET interacts with artificial membranes and alters membrane structure.

Methodology/Principal Findings: We show by surface plasmon resonance that HAMLET binds with high affinity to surface adherent, unilamellar vesicles of lipids with varying acyl chain composition and net charge. Fluorescence imaging revealed that HAMLET accumulates in membranes of vesicles and perturbs their structure, resulting in increased membrane fluidity. Furthermore, HAMLET disrupted membrane integrity at neutral pH and physiological conditions, as shown by fluorophore leakage experiments. These effects did not occur with either native HLA or a constitutively unfolded Cys-Ala HLA mutant (rHLAall-Ala). HAMLET also bound to plasma membrane vesicles formed from intact tumor cells, with accumulation in certain membrane areas, but the complex was not internalized by these vesicles or by the synthetic membrane vesicles.

Conclusions/Significance: The results illustrate the difference in membrane affinity between the fatty acid bound and fatty acid free forms of partially unfolded HLA and suggest that HAMLET engages membranes by a mechanism requiring both the protein and the fatty acid. Furthermore, HAMLET binding alters the morphology of the membrane and compromises its integrity, suggesting that membrane perturbation could be an initial step in inducing cell death.

Introduction

Lipid bilayers maintain cellular integrity by forming interactive barriers against the environment. They delineate the extracellular and intracellular spaces and play a critical role in the regulated uptake and export of essential components. Molecular interactions with this barrier in vivo may rely on lipid membrane constituents per se, or on specialized lipid-rich membrane domains, such as lipid rafts [1]. In addition, molecular recognition mechanisms often require specific receptors, which are embedded in lipid membranes, and structural modification of these receptors leads to the activation of highly regulated signaling pathways.

Membrane interactions are also essential in controlling cell survival and proliferation, and alterations in membrane composition and function accompany cancer development [2]. Molecules with selectivity for tumor cell membranes are important, to better understand the properties of tumor cells and to design agonists that attack and kill such cells with better selectivity. HAMLET (human α-lactalbumin made lethal to tumor cells) is a complex of partially unfolded α-lactalbumin (HLA) and oleic acid with broad tumoricidal activity. The complex interacts with tumor cell membranes, becoming rapidly internalized and translocated to the cell nuclei [3]. It appears that HAMLET triggers cell death by activating several signaling pathways and by interacting with cellular organelles, including proteasomes and mitochondria [4,5]. HAMLET is also medically relevant, as demonstrated in clinical studies of human skin papillomas and bladder cancer [6,7].

The mechanism of HAMLET uptake by tumor cells is unclear, however. HAMLET uptake has previously been quantified by radioactive labeling, quantitative confocal imaging and flow cytometry [4,8]. Attempts have also been made to calculate the equivalents of membrane receptor sites. Such techniques are not fully applicable to HAMLET and tumor cells, however, due to rapid uptake of the complex and loss of membrane integrity as a result of the cell death process. Artificial membranes may thus be useful to specifically address how HAMLET interacts with lipid...
HAMLET Interacts with Membrane

Materials and Methods

Materials

1,2-Dioleoylphosphatidylglycerol (DOPG), porcine brain phosphatidylserine (PBPS), soybean lecithin (polar extract) and egg yolk phosphatidylcholine (EYPC) lipids were from Avanti Polar Lipids (Alabaster, USA). Chemicals were from Merck (Darmstadt, Germany) and Sigma-Aldrich (St. Louis, MO, USA). 8-Aminonaphthalene-1,3,6-trisulfonic acid, disodium salt (ANTS), and p-xylene-bis-N-pyrimidinum bromide (DPX) used as a fluorophore and quencher, respectively, in the leakage assay were from Sigma-Aldrich. Chromatographic matrices were from Amersham Biosciences (Uppsala, Sweden) (HIC, DEAE-Sepha-cell) and DEAE-Trisacryl M from BioSephra (Villeneuville, France).

HAMLET

The HAMLET complex was formed from partially unfolded HLA and oleic acid, as described [16]. Recombinant HLA Alb-Ala was expressed in BL21* Escherichia coli using the T7-polymerase-based vector pAE92 (kindly donated by Peter S. Kim). Expression was induced by IPTG (Saveen & Werner AB; Limhamn, Sweden), rHLA Alb-Ala was purified according to Kim et al [19] with slight modifications [14]. For fluorescence imaging, HAMLET, HLA and rHLA Alb-Ala were labeled with AlexaFluor 568 (Molecular Probes, Carlsbad, CA, USA) according to the manufacturer’s instructions.

Cellular Assays

A549 lung carcinoma (ATCC CCL-185) cells were cultured in RPMI with NEAA, sodium pyruvate and 5% fetal calf serum (FCS) and Pheochromocytoma (PC12) cells in RPMI with donor horse serum (10%) and FCS (5%) in collagen IV coated flasks (VWR International, USA). To study HAMLET uptake, A549 cells were grown overnight on poly-L-lysine chamber slides (Nunc, Thermofisher, Roskilde, Denmark) and treated with AlexaFluor 568-labeled HAMLET, native HLA or rHLA Alb-Ala in medium without FCS (35 μM, 1 hour at 37°C in 5% CO2). Morphological changes and labeled protein uptake was recorded by real-time cellular imaging and confocal microscopy of fixed cells (LSM510 META confocal system, Carl Zeiss, Jena, Germany).

To quantify cell death, cells in RPMI without FCS were seeded into 24 well plates and incubated with different concentrations of each agonist at 37°C in 5% CO2. ATP measurements used ATPlite Luminescence ATP detection assay system (Perkin-Elmer, Waltham, MA, USA) and Trypan blue exclusion was recorded by interference contrast microscopy.

Lipid Vesicles

Large Unilamellar Vesicles (LUVs) were produced from EYPC, PBPS and DOPG, as described [17,18]. The prerequisite amounts of chloroform-dissolved lipids (1:1 mixes of DOPG:EYPC and PBPS:EYPC, and EYPC alone) were transferred to Kimble glass tubes wrapped in aluminum foil. The chloroform solutions were dried to lipid films under dry N2-pressure, and residual chloroform was removed under vacuum. The samples were vortexed with 10 mM citric acid/20 mM Na2HPO4 buffer pH 7.0 and allowed to hydrate overnight. For liposome preparation, the solutions were subjected to seven freeze-thaw cycles using liquid N2 and a water bath at 55°C [19]. Hydrated multilamellar structures were extruded (LIPEX extruder, Northern Lipids, Burnaby, BC, Canada), forced through two Millipore filters (pore size 200 nm) 10 times using 12-bar N2 pressures and transferred to clean Kimble tubes for immediate usage. In the ANTS/DPX leakage experiments, the lipid film hydrating buffer contained 12.5 mM ANTS and 45 mM DPX. The free fluorophores and quenchers were removed by gel-filtration on a Sephadex G-75 exchange column.

A mix of multilamellar and unilamellar giant unilamellar vesicles (GUVs) were created from soybean lecithin (Polar extract, 45% phosphatidylcholine and 22% phosphatidylethanolamine; 18% phosphatidylinositol and 7% phosphatidic acid) or EYPC (72% phosphatidylcholine, 11% phosphatidylethanolamine and 16% other neutral lipids). Lipids were dissolved in chloroform (3 mg in 200 μl of chloroform), dried and then swelled in PBS, pH 7.2 (3 ml) overnight at 8°C and dispersed (1 mg lipid per ml) 4000–6000 RU. The proteins (concentration range of 0–80 mM and dissolved in either buffer at pH 7.4 or pH 5.0) were mixed over the immobilized LUVs (flow rate of 10 μl/min) containing functional membranes with functional ion channels, used for patch-clamping [20]. The lipid film was allowed to swell on a glass bottom dish for microscopy (no. 1 glass thickness) for 5 minutes whereby surface-attached vesicles were formed.

Surface Plasmon Resonance

The Biacore 3000 biosensor system (Biacore AB, Uppsala, Sweden) was used at 25°C, as described [21]. L1 sensor chip surfaces were cleaned with a 2-min injection of isopropanol: 50 mM NaOH (1:1) at a flow rate of 20 μl/min, followed by 30 min washing with running buffer (HBSS-N buffer: 10 mM HEPES, pH 7.4, 150 mM NaCl). LUV solutions were diluted with running buffer (pH 7.4) or acetate buffer (pH 5.0, 0.6–1 mM) and injected (flow rate of 10 μl/min), resulting in a deposition yielding measurements of 4000–6000 RU. The proteins (concentration range of 0–80 mM and dissolved in either buffer at pH 7.4 or pH 5.0, to match the buffer used in the previous step) were injected over the immobilized LUVs (flow rate of 100 μl/min). The sensor chip surface was regenerated by injecting isopropanol:50 mM NaOH (40:60, v/v) and running buffer. Non-specific binding in the reference cell was subtracted. Binding affinity was estimated by fitting 3-parameter sigmoidal function to all concentration-RU data pairs, by nonlinear least-squares analysis, yielding S0.5-values reflecting the protein concentration necessary for a half-maximal response [21]. The fitting procedure also provided error estimates of the S0.5-values. The BIAl evaluation program, version 3.2 (Biacore AS) was used for analysis of the sensorgrams. Curve fits were performed using SigmaPlot version 9.01 (Systat Software, Inc.).
Fluorescence-Monitored Leakage Assays

LUV integrity was examined according to Ellens et al [22]. LUVs with ANTS/DPX encapsulation were diluted to 1 mM lipid concentration using citrate/Na₂HPO₄ at pH 5.0 or 7.0. Half-milliliter volumes were added to a 10 mm Hellma GmbH & Co quartz cuvette, and fluorescence was measured with a Perkin-Elmer LS5 Luminescence Spectrometer. Volumes of protein and oleic acid (OA) stock solution were added stepwise directly to the cuvette with gentle mixing (cumulative protein concentration 1, 5, 10, 20, 40, 60 μM). Fluorescence was recorded for each concentration (355 nm excitation, 450–550 nm emission, scan speed of 200 nm/ min, 3 and 5 nm slit widths for the excitation and emission pathways), 1 min after addition of the protein 25°C. Spectra were dominated by ANTS fluorescence (λ_{max} at 510 nm). The emission intensity at this wavelength depends on ANTS/DPX proximity, and an increased intensity indicates loss of liposome integrity as the average distance between ANTS and DPX increases. Controls were measured and subtracted. Triton X-100 was added to the cuvette to a final concentration of 2 mM and fluorescence at 510 nm was arbitrarily set to 100%, i.e., complete LUV breakdown. Fluorescence at 510 nm prior to addition of stock solution was arbitrarily set to 0%, i.e., no leakage. This way of benchmarking the resulting change in fluorescence is convenient for comparing leakage responses for liposomes of a given lipid mix. The values represent the means of two experiments (<5% variation was obtained). The leakage assay is a well-established technique that has been optimized to make variations in quantum yield negligible [22], as recently reviewed and validated by Almeida and Pokorny [23]. Controls were performed to confirm that increases in fluorescence are not caused by changes in the molecular environment at the end-point of the experiments (i.e. by the presence of HAMLET, HLA and Triton X-100 at defined concentrations), and are uniquely associated to the leakage from liposomes. Briefly, controls consisted of measuring the effect of HAMLET or HLA addition, in the absence or presence of Triton X-100, on the fluorescence at 510 nm (355 nm excitation) of samples of ANTS and ANTS/DPX at different concentrations. The additions did not cause differences in the fluorescence intensity of ANTS.

Imaging of HAMLET Interactions with Artificial Lipid Membranes

Alexa-labeled HAMLET or controls were added to GUVs in glass bottom dishes (0.5 mg/ml with 10% fluorescently labeled protein). Unbound protein was removed after 20 minutes using precision syringe pumping (15 minutes 2 ml/min). Microphotographs were taken before and after the labeled proteins were added, using confocal laser scanning microscopy (Leica TCS SP2 RS, Wetzlar, Germany), with a PL APO CS 63×/1.2 W corr objective for image acquisition. Brightness and contrast was adjusted in the Leica software. For intensity profile measurements, line scans were obtained with Image J software (http://rsbweb.nih.gov/ij/). An inverted fluorescence microscope (Leica DMI 6000B) was used to record morphological changes (bright-field, Leica DFC350FX camera system).

Plasma Membrane Vesicles

Pheochromocytoma (PC12) cells were cultured for 2–4 days and plasma membrane vesicle (PMV) formation was induced as described by Bauer et al [15]. PC12 cells with surface-attached PMVs and free vesicles were then rinsed twice in washing buffer and transferred to a glass bottom dish (Wilco well) for incubation with Alexa-labeled proteins. PMVs from lung carcinoma cells (A549) were prepared by the same technique.

Results

1. HAMLET Interacts with Neutral and Negatively Charged Membranes at Physiological pH

To understand the membrane affinity of HAMLET, we examined its interaction with biologically relevant membranes in vitro, using surface plasmon resonance with LUVs immobilized on a Biacore L1 sensor chip.

The composition of the LUVs chosen was based on previous studies of HLA interacting with and penetrating into such bilayers. Briefly, binding was optimal for lipid mixtures with an overall negative charge of 50% [24] achieved by 1:1 mixtures containing either DOPG or PBPS as the negative component, and EYPC as the zwitterionic component. Liposomes prepared with DOPG or PBPS alone would contribute 100% negative charge, while EYPC-only liposomes would have 0%; thus, a 1:1 mixture yields 50% negative charge. As fluidity also defines the membrane quality of potential importance for HLA-membrane interactions [Agasoster, 2003 #165], three types of LUVs (DOPG:EYPC, PBPS:EYPC and EYPC lipid mixtures) representing different states of fluidity and charge were used to investigate HAMLET binding (Fig. 1A).

The DOPG:EYPC liposomes are in the fluid lipid state, while the two latter species are close to their transition temperature, T_c [24,25]. Since 18:1 oleic acid acts as the HAMLET co-factor, the amount of 18:1 fatty acid 18:1 acyl chains in the liposomes is of interest when interpreting the binding results. The DOPG:EYPC lipid mix contains 66% 18:1 acyl chains, while the PBPS:EYPC and EYPC mixtures contain approximately 33%.

HAMLET bound strongly and in a concentration-dependent manner to all three membranes at near physiological conditions (pH 7.4, 150 mM NaCl), with only small differences in absolute binding and S0.5 values (S0.5-values ranging from 6.99–7.94, with standard errors in the range of 0.07–0.40; Figs. 1A and B). HAMLET binding was not influenced by membrane fluidity, charge or 18:1 fatty acid acyl chain composition. HLA did not bind to any of the LUVs under physiological conditions (Fig. 1A), confirming previous studies where a pH<5 was optimal for HLA binding to bilayers with 50% negatively charged phospholipids. The results show that HAMLET binds broadly to membranes under physiological conditions, in contrast to HLA.

2. HAMLET Interacts with Giant Unilamellar Vesicle Membranes

The membrane interactions of HAMLET were further investigated by confocal microscopy imaging of GUVs composed of EYPC or soybean lipids at pH 7. LUVs were not suitable for confocal imaging, as they do not form stable unilamellar vesicles under these conditions. The GUVs, prepared by dehydration-rehydration, yielded a mixture of unilamellar (single layer) and multilamellar (several layers) phospholipid membranes. The membrane properties of the respective vesicles were the same, however, regardless of size and structure. Conclusions were based on pair-wise analysis within each preparation.

Membranes of EYPC lipid vesicles were clearly visualized by binding of fluorescently labelled HAMLET, whereas HLA and free Alexa 568 dye associated more weakly to the vesicles, as shown by the low fluorescence signal (Fig. 2A). To quantify the membrane association of HAMLET in unilamellar vesicles, we used intensity profile measurements with ImageJ software and a median filter radius of 1.0 pixels. There was a clear HAMLET association with both EYPC and soybean membranes, but only a weak signal inside the unilamellar vesicle, suggesting no substantial internalization of the HAMLET complex (Fig. 2B).
The results show that HAMLET binds more strongly than native HLA to both neutral and slightly negatively charged artificial membranes.

3. Membrane Perturbations by HAMLET

To test if membrane integrity was affected by HAMLET, LUVs were loaded with ANTS fluorophore and DPX quencher, and leakage into the extravesicular space was quantified by measuring ANTS fluorescence. Native α-lactalbumin disrupts liposomes at pH 5, but has no effect at neutral pH or on zwitterionic liposomes at acidic pHs [24]. HAMLET disrupted negatively charged (DOPG:EYPC) and PBPS:EYPC after one minute at neutral pH (Fig. 3B,C). Fluorophore leakage was higher for DOPG:EYPC, followed by PBPS:EYPC (Fig. 3B and C), indicating that the degree to which HAMLET can alter membrane structure is influenced by the membrane’s lipid composition and physical characteristics. HAMLET caused a maximal leakage, 100–120% relative to that induced by the Triton X-100 control, used for benchmarking the leakage response within each experimental series on a given liposome lipid mix. Such high relative permeability suggests that HAMLET is a very effective disruptor of negatively charged membranes at neutral pH. HAMLET bound zwitterionic vesicles (EYPC) (Fig. 1A) and although no disruption was observed at neutral conditions (Fig. 3A), leakage was measured at pH 5.0 (Fig. 3E). Free oleic acid had no detectable effect on LUVs independent of phospholipidic composition (Fig. 3). Thus, HAMLET, but not HLA or oleic acid, permeabilizes negatively charged lipid bilayers under physiological conditions. Moreover, HAMLET is a very efficient disruptor of DOPG:EYPC and PBPS:EYPC vesicles, with the former, enriched in oleoyl-sidechains, being even more susceptible to disruption by HAMLET than by Triton X-100.

We also documented the effect of HAMLET on membranes by light microscopy. Vesicles made of EYPC in PBS buffer at pH 7 remained morphologically unchanged for one hour, but the addition of HAMLET affected the shape of both multi- and unilamellar vesicles. We observed a change from round and defined to elongated, heterogeneous morphology after 40 minutes (Fig. 4). In addition, continuous monitoring revealed an increase in membrane movement, with a subsequent increase in the number of unilamellar structures (data not shown). The contribution of protein unfolding to this change in membrane morphology was examined by comparing HAMLET to native HLA and to the rHLA<sup>1-44</sup> mutant, which lacks disulphide bonds and is therefore unable to fold into the native state. Morphological changes of membrane vesicles were not observed after treatment with either native HLA or rHLA<sup>1-44</sup>. The results show that HAMLET, but not HLA or rHLA<sup>1-44</sup>, alters the morphology of membrane vesicles.

4. Interaction of HAMLET with PMVs from Intact Tumor Cells

To compare the effect of HAMLET on artificial protein-free membranes versus tumor cell membranes (containing membrane receptors), we prepared PMVs from the tumor cell lines PC12 [15]. Free PMVs and PMVs (blebs) still attached to cells were exposed to Alexa-labeled HAMLET at concentrations corresponding to the LD<sub>50</sub> cell death value (0.5 mg/ml). Alexa-labeled HLA and free Alexa dye were used as controls. HAMLET bound strongly to tumor cell PMVs (Fig. 5A, I & II). In contrast to the artificial protein-free membranes, the HAMLET distribution on PMVs was patchy, indicating that specific membrane regions may have higher affinity for the complex. There was no clear evidence of translocation across the membrane into the interior of the PMVs, but the remainder of the tumor cells from which the blebs had been derived fluoresced strongly, showing that Alexa-labeled HAMLET had been taken up. Alexa-labeled HLA bound weakly to the PMVs and cell remnants (Fig. 5A, III). Alexa dye did not interact with the PMVs per se, but cells fluoresced weakly, possibly due to non-specific uptake through the ruptured plasma membrane (Fig. 5A, IV). To confirm our results, we tested in parallel PMVs from another tumor cell line, A549 lung carcinoma cells, and used a different visualization marker. The vesicles also bound HAMLET with a patchy distribution, as visualized with Nile red detection (Fig. 5B I–III). After 20 minutes, a change in membrane morphology was observed, with an accumulation of Nile red in the membranes accompanied by elongated, convoluted shapes.
Oleic Acid Binds PMVs but Does Not Disrupt LUVs or GUVs

In parallel with HAMLET and HLA, the different vesicle preparations (LUVs, GUVs, and PMVs) were exposed to OA, at a concentration corresponding to the stoichiometry of 4–6 OA residues per HLA in the HAMLET complex. In ANTS/DPX-loaded vesicles, OA alone did not cause an increase in extravesicular fluorescence, indicating that the membranes remained intact (Fig. 3). In the membrane morphology studies, OA alone did not cause any detectable change in vesicle shape or flexibility (data not shown). OA did incorporate into PMV membranes. However, OA became associated with the PMVs, resulting in weak fluorescence compared

Figure 2. Confocal imaging of HAMLET binding to lipid membranes in GUVs. A. Differential interference contrast images and corresponding fluorescence confocal microscopy images. EYPC lipid vesicle membranes were visualized after application of Alexa-labeled HAMLET (I), HLA (II) or free Alexa 568 dye (III). HAMLET bound to multiple layers of lipid membranes in the multilamellar vesicles. Vesicles after application of Alexa-labeled HLA or free Alexa 568 dye indicated weak fluorescence association. B. Membrane detection of LUVs by fluorescence confocal microscopy, and corresponding intensity profile measurements, using ImageJ software and a median filter radius of 1.0 pixels to process the images before the profile measurements. I & II) Binding of HAMLET to EYPC vesicles; confocal micrograph and corresponding DIC microscopy images. White arrowheads illustrate unilamellar structures. III) Intensity profile measurement across the line shown in I, confirming the strong association of HAMLET to the membrane. IV) Binding of Alexa-labeled HAMLET to soybean phospholipid choline vesicles. V) Corresponding DIC micrograph. Black arrowheads illustrate unilamellar structures and the black arrow indicates a multilamellar structure. VI) Intensity profile measurement of the line shown in IV. HAMLET bound strongly to soybean vesicle membranes.
doi:10.1371/journal.pone.0009384.g002
HAMLET, but concentrated in spots on the membrane (arrows in Fig. 5C I–III) indicating an accumulation in membrane dots. There was no change in PMV morphology comparable to that seen with HAMLET.

**HAMLET Internalization and Cell Death Require Both Unfolded HLA and Oleic Acid**

To further examine membrane binding of intact tumor cells, A549 lung carcinoma cells were exposed to Alexa-labeled HAMLET or HLA. Membrane interactions and effects on tumor cell morphology were first recorded by live-cell imaging (Fig. 6A). HAMLET rapidly became associated with the cell membrane, and internalization occurred immediately (Fig. 6A).

Over a six-hour period, HAMLET-treated cells changed morphologically. HAMLET became membrane-associated and internalized after 30 minutes, and after three hours the cells rounded up and formed blebs. After six hours, membrane integrity was lost and all cells had internalized HAMLET (Fig. 6A). There was no binding, uptake or morphological change after cell exposure to the protein component of HAMLET alone. Neither fatty acid free HLA nor the unfolded mutant, rHLAAl-Ala, entered or killed cells (Fig. 6B). In HAMLET-treated cells, tumor cell death coincided with HAMLET internalization, and was detected by trypan blue exclusion and ATP measurements after one- and three-hour exposures, (Fig. 6C).

The results show that membrane binding and internalization by tumor cells require partially unfolded HLA and oleic acid. Importantly, unfolding alone did not enable HLA to bind to cell membranes, become internalized or kill carcinoma cells.

**Discussion**

The tumoricidal HAMLET complex interacts with the tumor cell membrane, is internalized and translocates to the cell nucleus [3]. This study used artificial membranes to define whether HAMLET can interact with phospholipid bilayers in the absence of specific tumor cell membrane constituents. We showed that HAMLET bound membranes at physiological pH, and as a result, perturbed the membrane integrity and caused leakage of vesicular contents to the exterior. In addition, the morphology of vesicles composed of natural lipid mixtures was altered. The effects were quite dramatic, differing markedly from those of natively folded HLA, which failed to bind membranes at neutral pH. Significant HLA binding has only been recorded at pH<5, where the proton gradient near the negatively charged membrane is thought to be sufficient to remove negative charges at key residues, and thus drive protein unfolding necessary for membrane binding [11,26,27]. To determine whether...
For native HLA, at least, membrane curvature is of secondary importance for binding relative to membrane charge and the protein’s membrane-bound folding state [24,26,29]. That HAMLET has high affinity for a range of liposomes examined here (neutral, charged, LUVs and GUVs) suggests that curvature is also less important for the partially folded protein. A previous study [25] of SUVs prepared with similar lipid mixtures suggested that at room temperature the mixtures are either in the lα-state (DOPG:EYPC) or within the transition temperature interval (EYPC, PBPS:EYPC). Moreover, HLA increases the membrane Tc upon binding. Whether this latter feature is true of HAMLET is an open question, but we interpreted this as indicating that the degree to which the native protein penetrates the membrane and the physical state of the membrane were coupled.

Thermodynamic access to intermediately folded protein states has been proposed to drive membrane binding of proteins, including HLA [27,30,31,32]. Membrane curvature is a known contributor in many systems. However, for HLA interacting with negatively charged membranes, this factor is less important than bulk conditions and membrane charge [24,28]. For HAMLET, unfolding of HLA alone was not sufficient to create membrane affinity and to permeabilize the membrane, emphasizing that the role of oleic acid, in addition to kinetically trapping HAMLET in its active conformation, might be to enhance membrane affinity. Increased hydrophobicity through the fatty acid acyl-chain, or modulation of the local charges on the protein through shielding and hydrogen bonding are possibilities. It remains unclear if short-range forces could consolidate the interaction with the subset of conformations available that are most favorable for binding the membrane.

Free fatty acids are used by all cells as an energy source and as building blocks for hormones, glycolipids and lipoproteins and are constantly taken up by cells. Although the uptake mechanisms are not fully understood, a range of receptors and transport systems are thought to be involved [33]. Fatty acids also cross lipid bilayers by receptor-independent mechanisms; this has been ascribed to their hydrophobicity [34]. We have confirmed elsewhere that tumor cells internalize free oleic acid as well as the HAMLET complex, suggesting that the primary membrane interactions may depend on oleic acid as well as the protein (manuscript in preparation). In the present study, neither HLA nor oleic acid alone could alter membrane structure or disrupt vesicles, suggesting that both the unfolded protein and the lipid are necessary bind to and disrupt membranes.

The difference in membrane affinity between HAMLET and HLA was confirmed using tumor cell PMVs, suggesting that the artificial membrane systems might reflect important properties of intact cell membranes. Previous studies using microscopy and proteomic analysis of cytosolic proteins have shown that the PMV interior is filled with cytosolic molecules but does not contain cytoskeletal elements such as actin or microtubules [35]. The lipid composition of the PMVs is representative of the donor cell plasma membrane, with phospholipid/cholesterol ratios of 2:1 [36] and PMVs display phase separation of proteins and lipids [37]. HAMLET was distributed in patches on PMV membranes, indicating that specific membrane regions bound HAMLET more strongly than others. Thus, whereas lipid bilayers and their constituents are sufficient for HAMLET binding, specific receptors for HAMLET or oleic acid could also contribute in tumor cell interaction. This is consistent with preliminary studies indicating that HAMLET internalization is influenced by specific signaling pathways and by the metabolic state of tumor cells (P. Storms et al manuscript in preparation). The membrane interactions described here may allow the complex to reach specific membrane-associated receptor structures, thus enabling HAMLET to activate the complex signaling pathways involved in cell death.
Figure 5. Interaction of HAMLET with tumor cell PMVs. Pheochromocytoma cell (PC12) PMVs on glass-bottom dishes were exposed to Alexa-labeled HAMLET, Alexa-labeled HLA or free Alexa 568 dye. Membrane fluorescence was observed by confocal fluorescence microscopy and morphology by differential interference contrast microscopy. A) Detection of Alexa-HAMLET on PMVs (I, white arrowheads). Cell remnants are also highly stained (I, right panel). Black arrowheads indicate the PMV membrane (II). Weak detection of PMVs and cell remnants after application of Alexa-labeled HLA (III). Free Alexa dye served as a negative control (IV). B) PMVs from A549 cells exposed to HAMLET (35 μM, 0, 10, 20 min) and visualized with Nile red. An increase in fluorescence intensity and change of shape was observed over time. Internal convoluted membrane structures developed (arrows). C) PMVs from A549 cells exposed to OA and visualized with Nile red (0, 10, 20 min) showed a gradual increase in membrane binding at distinct membrane spots (arrows).

doi:10.1371/journal.pone.0009384.g005
Acknowledgments
We would like to thank Dr. Ming Ying for expert help with surface plasmon resonance measurements, Brigitte Bauer and the Centre for Cellular Imaging at the Sahlgrenska Academy, University of Gothenburg, and Susan Basiri and Pontus Linden for preparing HAMLET and α-lactalbumin variants.

Author Contributions
Conceived and designed the experiments: AKM MP ØH IL AM CS RK. Performed the experiments: AKM MP ØH AB IL YC RK. Analyzed the data: AKM MP ØH AB IL YC AM CS RK. Contributed reagents/materials/analysis tools: AKM MP ØH IL YC AM CS RK. Wrote the paper: AKM MP ØH IL AM CS RK.

References

HAMLET Interacts with Membrane


Ellens H, Beut J, Szoka FC (1984) pH-induced destabilization of phosphati-


Agasoster AV, Halskau O, Fuglebakk E, Froystein NA, Muga A, et al. (2003) The interaction of peripheral proteins and membranes studied with alpha-


van der Goot FG, Gonzalez-Manas JM, Lakey JH, Pattus F (1991) A 'molten-


van der Goot FG, Gonzalez-Manan JM, Lakey JH, Fattus F (1991) A 'molten-


Kleinfeld AM (2000) The interaction of peripheral proteins and membranes studied with alpha-