Mechanism of Shiga Toxin Clustering on Membranes

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Supporting Information

ABSTRACT: The bacterial Shiga toxin interacts with its cellular receptor, the glycosphingolipid globotriaosylceramide (Gb3 or CD77), as a first step to entering target cells. Previous studies have shown that toxin molecules cluster on the plasma membrane, despite the apparent lack of direct interactions between them. The precise mechanism by which this clustering occurs remains poorly defined. Here, we used vesicle and cell systems and computer simulations to show that line tension due to curvature, height, or compositional mismatch, and lipid or solvent depletion cannot drive the clustering of Shiga toxin molecules. By contrast, in coarse-grained computer simulations, a correlation was found between clustering and toxin nanoparticle-driven suppression of membrane fluctuations, and experimentally we observed that clustering required the toxin molecules to be tightly bound to the membrane surface. The most likely interpretation of these findings is that a membrane fluctuation-induced force generates an effective attraction between toxin molecules. Such force would be of similar strength to the electrostatic force at separations around 1 nm, remain strong at distances up to the size of toxin molecules (several nanometers), and persist even beyond. This force is predicted to operate between manufactured nanoparticles providing they are sufficiently rigid and tightly bound to the plasma membrane, thereby suggesting a route for the targeting of nanoparticles to cells for biomedical applications.

KEYWORDS: Casimir force, fluctuation-induced force, endocytosis, invagination, membrane, clustering, glycosphingolipid, lectin

The bacterial Shiga-like toxins from Escherichia coli are responsible for pathological manifestations that can lead to hemolytic uremic syndrome, an endemic threat to human health.1 These toxins are composed of 2 subunits: a catalytic A-subunit, which modifies rRNA once translocated to the cytosol of target cells, leading to protein biosynthesis inhibition;2 and a homopentameric B-subunit, which mediates the toxin’s interaction with glycosphingolipids (GSLs) from the globoseries, which function as cellular toxin receptors.

The cell biology of the parental Shiga toxin from Shigella dysenteriae has been particularly well-studied.2,3 Its B-subunit (STxB), a rigid protein of around 7 nm in diameter, tightly binds up to 15 globotriaosylceramide (Gb3) GSL receptor molecules as the first step leading to toxin entry into cells.2 Current understanding suggests that the Shiga toxin-driven reorganization of membrane lipids endows the toxin−Gb3 complexes with curvature active properties that after clustering of toxin molecules enables the formation of endocytic membrane invaginations without a strict requirement for the cytosolic clathrin machinery (for a review, see Johannes et al.4). After endocytosis, Shiga toxin is then transported from early endosomes to the Golgi apparatus and the endoplasmic reticulum, from where the catalytic A-subunit is translocated to the cytosol to modify rRNA and thereby to inhibit protein biosynthesis.5

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Shiga toxin forms clusters on cell and model membranes despite the apparent absence of direct protein–protein interaction between toxin molecules based on hydrophobic or electrostatic effects, or mechanisms based on cholesterol-dependent phase separation. How this clustering occurs is still unknown. Possible mechanisms for membrane-associated particle clustering are based on acto-myosin-driven molecular focusing, which would operate at length scales in the range of tens to hundreds of nanometers, and membrane-mediated forces in which the particles perturb the membrane in a way that drives their aggregation. Such membrane-mediated forces include capillary (domain boundary) forces, which only arise under specific conditions, or lipid depletion forces, which are effective only in the subnanometric range. Membrane perturbations may also arise from generic properties of the particles, such as their size or shape, and have been studied by computer simulation. Transmembrane proteins whose height is mismatched with the width of the surrounding membrane have also been found to aggregate if they are sufficiently large. Membrane-bound STxB creates a small increment of curvature, which would be expected to yield a repulsive force between toxin molecules. However, our simulations and experiments show that reducing the rigidity of the bound STxB nanoparticles or displacing them from the membrane surface eliminates the clustering process. Strikingly, this effect is paralleled by a loss of STxB nanoparticle-mediated suppression of membrane fluctuations, strongly suggesting that fluctuation-induced forces generate an effective attraction between the STxB nanoparticles. This clustering mechanism is generic and should operate between any sufficiently rigid nanoparticles that are able to bind tightly to a fluid phase lipid bilayer, pointing to a route for the therapeutic delivery of nanoparticles into cells.

RESULTS

FCS Setup To Measure STxB Clustering. Membrane-bound STxB was studied in a micropipette aspiration setup that provides control over membrane tension in a model membrane system, giant unilamellar vesicles (GUVs), while simultaneously allowing the analysis of clustering of membrane constituents by fluorescence correlation spectroscopy (FCS) (Figure 1A). GUVs (95 mol % 1,2-dioleoyl-sn-glycero-3-phosphocholine (DOPC)) were prepared by electroformation with 5 mol % C22:1 Gb3 species, which have previously been shown to favor STxB-driven membrane bending. Stretching the membrane to high values of approximately $10^{-3}$ N/m provided us with two advantages in precisely testing the extent of STxB clustering. First, STxB-induced formation of tubular membrane invaginations was prevented, which allowed us to compare the different lipid species used in our experiments, independent of their ability to support tubulation. Second, stretching the membrane suppressed high amplitude membrane fluctuations that would otherwise have resulted in non-STxB diffusion related intensity fluctuations in the FCS measurements.
fluorescence intensity fluctuations observed in our experiments were predominantly due to diffusion of STxB-Alexa488 molecules in and out of the cross section of the membrane within the focal FCS observation spot. Diffusion time distributions were measured by fitting the autocorrelation curves using the maximum entropy method-based fitting routine (MEMFCS). This allowed us to deduce the size distribution of the diffusing STxB clusters independent of the dimensionality of the equation used in the fitting process (Supporting Information, Figure S115) and the photophysics of the dye (Supporting Information, Figure S116). We found that the peak of the distribution of the diffusion times shifts with time to higher values and the distribution broadens (expressed as $\tau_p$ in Figure 1C,D). These observations suggest that the peak size of the STxB clusters as well as the size heterogeneity grew with time.

**Line Tension Is Not Required for STxB Clustering.** Next we exploited the above experimental setup to test whether commonly understood membrane-mediated forces can explain STxB clustering.

Lipid chain length mismatch can be very effective for driving lipid phase segregation. By creating a locally distinct lipid phase environment, STxB could aggregate through a capillary effect. This possibility was addressed in our setup by using C18:1 Gb3 species whose chain length matched that of the bulk membrane lipid DOPC. Even under these conditions, STxB still clustered efficiently (Figure 1D), demonstrating that lipid chain length mismatch is not strictly required.

Compositional mismatch is another mechanism for line-tension driven clustering. We therefore tested GUVs with a bulk Gb3 concentration of 30 mol %, mimicking the estimated Gb3 concentration bound under STxB molecules (see Supporting Information for further details). STxB still formed invaginations at 37 °C, above the phase transition temperature for 30% Gb3 and 70% DOPC (Figure 1B), indicating that clustering is maintained and that lipid compositional mismatch is not required. These results lead us to conclude that conventional line tension-based mechanisms are not an obligatory requirement for STxB clustering.

**No Impact of STxB on the Hydrophobic Bulk of the Bilayer.** Since STxB, via its receptor, Gb3, is connected to the hydrophobic region of the bilayer, one might suspect that the bound proteins modify the lipid chain entropy thereby driving their clustering. To address this possibility, we reanalyzed trajectories from our previously published work and observed no major differences in the lipid chain order parameter (Supporting Information, Figure S110) nor the rotational correlation function of DOPC lipids under STxB (Supporting Information, Figure S111), when compared to DOPC lipids far from STxB. Also, when the chain order parameter of Gb3 bound to STxB was compared to that of unbound Gb3, no evidence for STxB-induced ordering could be observed (see Supporting Information, Figure S112). Therefore, an entropic effect of STxB on the hydrophobic region of the bilayer as a source of clustering appears highly unlikely.

**Local Curvature Induced by STxB Does Not Drive Clustering.** Shiga toxin induces negative local membrane curvature, since it results in macroscopic membrane invagination on GUVs (see Römer et al.) and membrane bending in silico. The question thus arises whether such a curvature imprint can generate an attractive force. Indeed, for highly curved particles, an attractive curvature-induced force has been described previously. Later, the same authors performed a more precise evaluation of the range of the curvature imprint for which the curvature-induced force is attractive. Notably, it was found that for contact angles up to 45°, the curvature-induced force is described well by the linear approximation and is predicted to be repulsive. STxB clearly falls into this category with a low contact angle around 7°, resulting from a

![Figure 2. DPD simulations of particles on a fluctuating membrane.](image)
spontaneous curvature radius that is 5-fold bigger than the protein’s size.\textsuperscript{14} We can therefore exclude induced membrane curvature as the driving force for STxB clustering.

\textbf{Coarse-Grained Simulations of Clustering.} The experimental results so far suggest that the clustering of STxB molecules is independent of the precise molecular structure of the membrane, and cannot be explained by known membrane-mediated clustering mechanisms. To proceed further, we used a coarse-grained simulation technique, dissipative particle dynamics (DPD), to study the clustering of nanoparticles (NP) that mimic the physical properties (but not the molecular details) of Shiga toxin on an amphiphilic bilayer.

Figure 3. STxB clustering on GUVs containing Gb3 species with flexible linkers. (A) Molecular structure of the different lipids used in this study. (B) Binding of 200 nM STxB-Alexa488 to vesicles prepared with 5% mentioned Gb3 species and 95% DOPC. The yellow rectangle depicts the range within which FCS measurements were taken. (C) The percentage of vesicles displaying tubulation decreases with increasing linker length (black circles). The extent of tubulation (red boxes), that is, the sum of length of all the invaginations for a given vesicle divided by the circumference in that particular cross section (schematic in panel F), also decreases with increasing linker length. (D) An example snapshot of vesicles containing C22:1 Gb3 showing extensive tubulation. (E) An example snapshot of vesicles prepared with C22:1 Gb3\_EG, showing almost no tubulation. (F) Schematic describing the measurement of extent of tubulation (see panel C). Scale bars for panels D and E, 10 μm.
Previous studies have shown that DPD simulations of lipid bilayers faithfully reproduce membrane physical properties, such as area per lipid and membrane elasticity for single \(^\text{19}\) and multicomponent bilayers.\(^\text{20,21}\) After confirming that our simulations replicated these properties (not shown), we added nanoparticles to the simulation whose shape (pentagonal), size (7 nm cross-sectional size), and material properties (high stiffness) mimic those of STxB. The detailed procedure is described in the Supporting Information, including how NP stiffness is quantified by a “rigidity parameter” \(k\). The STxB binding to Gb3 was modeled as a nonspecific attraction between the lipid head groups and the nanoparticles. The simulated STxB nanoparticles are large enough that atomic features of their binding, including H-bonding, need not be explicitly represented, which makes DPD a suitable simulation technique. Our results are therefore robust against the details of the force field used and depend only on generic properties of the particles that are independent of their detailed molecular structure. We eliminated the possibility that clusters arise from direct binding between nanoparticles by giving them a short-range (smaller than the particle radius) repulsive interfacial interaction. Furthermore, height and compositional mismatch and curvature were excluded by choosing conditions in which, respectively, all lipid tails are of the same length, the NPs are flat, and the same molecular lipid species are used throughout (Supporting Information, Figure SI8B,C).

The DPD simulations showed (see Supporting Information, Movie 1) that two rigid toxin NPs initially placed apart on the membrane approached each other whereupon they adhered dependent on the stiffness and the size of the toxin NPs, as described below. Repeated simulations confirmed that, once bound, NPs did not separate for at least 9.6 \(\mu\)s (Figure 2A, \(r_0\) condition, blue trace/box where \(r_0\) is the unit of length in the DPD simulations and is defined in the Supporting Information). For comparison, the NPs diffuse a distance equal to their diameter in approximately 10 \(\mu\)s (Supporting Information, Figure SI8-A) indicating that the toxin NPs have overcome the entropic separation tendency and adhere. Strikingly, toxin NPs dampened the membrane’s thermal fluctuations in our DPD simulations (Supporting Information, Figure SI7A, \(k = 64\), blue dot; Figure SI7B for lipid L-02; see Supporting Information for details on the method to calculate the suppression of membrane fluctuations). This effect on fluctuations will be followed up throughout all subsequent experiments and its relevance to the clustering process addressed in the Discussion.

Why Do Not All Proteins Cluster? The above clustering mechanism might be thought to operate among all membrane-associated proteins, which would contradict the experimental observation that many membrane proteins do not cluster.\(^\text{22–24}\) The clustering mechanism must therefore depend on properties that are not shared by all membrane proteins but that are present when the NPs in the DPD simulations bind to the membrane, for example, NP size, geometry, stiffness, and coupling to the membrane.

In a first set of simulations, we found that NPs with sizes similar to that of the NP from Supporting Information, Movie 1, but with different geometries (square, hexagonal, and disk-like), aggregated on tense or tensionless bilayers (not shown). These results demonstrate that the membrane tension and NP shape do not remove the clustering although the force may be altered. By contrast, we found that clustering disappeared when the nanoparticle diameter was less than about 2.5 nm (Figure 2A, red trace; Supporting Information, Movies 2–4).

We next modified the internal stiffness of the NPs to explore the effects of their rigidity and found that clustering vanished for values of the rigidity parameter \(k = 16\) or below (Figure 2B). This result is in agreement with that found in Monte Carlo simulations of inclusions on a fluctuating surface by Weikl.\(^\text{25}\) The suppression of membrane fluctuations was no longer observed in conditions under which a loss of rigidity led to a loss of clustering (Supporting Information, Figure SI7A). We also note that solvent depletion forces are still present for highly flexible NPs, demonstrating that these were not sufficient for clustering.

These results showed that clustering in DPD simulations is strongly dependent on the NP size and internal rigidity. Unfortunately, neither of these properties can be modified experimentally in the STxB system. However, the mechanical coupling of a NP onto the bilayer is experimentally accessible and was therefore addressed next.

STxB Clustering Requires Tight Mechanical Coupling to the Membrane. To test the importance of tight mechanical coupling between NPs and membrane, we added flexible linker segments (gray beads in the schematics of Figure 2C) of increasing length (L-12 or L-22 carry 1 or 2 linker beads, respectively) to the lipid head groups (red beads). In this DPD simulation, distal regions (green beads) remained attractive to the toxin NPs, which therefore adsorbed to the membrane in all cases. When toxin NPs were forced to aggregate, they separated again when linkers were longer than 1 nm (Figure 2C, L-12 and L-22 conditions; black and red traces/boxes, respectively; Supporting Information, Movies 5 and 6), whereas they remained bound for up to 4.8 \(\mu\)s in all simulations in the absence of any flexible linker segments (Figure 2C, L-02 condition, blue trace/box). Importantly, the loss of clustering with increasing linker length again correlated with the loss of suppression of membrane fluctuations (Supporting Information, Figure SI7B), pointing to the possibility of a causal link between fluctuations and clustering (see Discussion).

We returned to the GUV/FCS setup to test experimentally how soft linkers influence STxB clustering. For this, we synthesized three artificial Gb3 species: C22:1 Gb3\(_{\text{EG}_{\text{a}}\text{r}}\), in which the globotriose sugar headgroup was separated from the ceramide backbone by 1, 3, or 7 ethylene glycol (EG) units (Figure 3A), corresponding to 1.0, 1.75, or 3.25 nm of extended linker length. Details of the chemical synthesis can be found in the Supporting Information. GUVs were prepared with these artificial Gb3 species, as described above. The binding of STxB was similar in all cases (Figure 3B). By contrast, there was a marked difference in the ability to form inward tubules (Figure 3C, black circles, D, and E), and the extent of tubulation (Figure 3F) was clearly reduced with increasing linker length (Figure 3C, red boxes).

Using FCS on micropipette aspirated GUVs for which tubulation was suppressed by tensing the membranes allowed us to precisely compare the extent of STxB clustering for all lipids. Compared to the natural C22:1 Gb3 species, STxB clustering kinetics was already slightly reduced on C22:1 Gb3\(_{\text{EG}_{\text{r}}\text{r}}\) (Figure 4). For Gb3\(_{\text{EG}_{\text{r}}\text{r}}\) and Gb3\(_{\text{EG}_{\text{a}}\text{r}}\), no significant aggregation was observed as \(r_0\) did not extend beyond 20 ms and the width of \(r_0\) distribution remained comparable with time (Figure 4). Whereas clusters formed from natural Gb3 have hydrodynamic radii of 180 nm, estimated as described in Supporting Information, those...
formed from Gb3_EG3 and Gb3_EG7 have a hydrodynamic radius that remains below 10 nm, which is comparable to the size of a single STxB particle. These results closely mirror the ones made with soft linkers in DPD simulations (Figure 2C) and suggest a systematic decrease in the degree of clustering, related to the ethylene glycol linker lengths between the head and tail groups of the corresponding Gb3 species. Since the density of toxins at the cell surface is similar in all conditions, our findings further support the absence of direct protein–protein interactions between STxB molecules. These experiments provide evidence that tight mechanical coupling onto the membrane is essential for STxB clustering.

To examine if flexible EG linkers also affected STxB clustering on cellular membranes, we incorporated synthetic C18:1 Gb3 or C18:1 Gb3_EG7 species into GSL-deficient GM95 cells. We first showed that the binding of the toxin on the cells incorporated with these lipids was similar (Figure 5A,B). Consistent with the experiments on GUVs, we found using FCS measurements that STxB clustering was strongly compromised on cells with C18:1 Gb3_EG7 species (Figure 5C,D,E).

We finally exploited phosphine-based quenching to differentiate internalized STxB from plasma membrane-accessible extracellular STxB. Using this approach, we found that the endocytic uptake of STxB was strongly reduced on cells that were reconstituted with C18:1 Gb3_EG7 species, when compared to cells that were reconstituted with natural C18:1 Gb3 (Figure 6A,B). Both the number of STxB-positive endosomes per cell and the breadth of STxB fluorescence intensity distribution were strongly reduced in the C18:1 Gb3_EG7 condition.

Thus, the mechanical coupling of STxB to the plasma membrane and its capacity to cluster were of critical importance to its efficient uptake into cells.

**DISCUSSION**

Clustering of the bacterial Shiga toxin on the cellular plasma membrane is required for infection of the cell. Several
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We estimate from FCS experiments on highly tensed vesicles (for which invaginations cannot form) that clusters can grow into domains that are hundreds of nanometers in size (Supporting Information), indicating their phase separation into toxin-rich domains embedded in the surrounding toxin-poor membrane. When this phase separation occurs on membranes with low values of membrane tension, the small increment of curvature induced by each toxin particle, which has been predicted from atomistic Molecular Dynamics simulations to be 0.035 nm$^{-1}$, creates a spontaneous curvature in the toxin-rich domains that subsequently drives their tubular invagination, as seen in Figure 1B.

The thermal Casimir force depends on the ability of membrane-adsorbed particles to suppress the thermal membrane fluctuations beneath them because this restricts the fluctuation spectrum of the free membrane regions and gives rise to the attractive force between the particles. It was therefore of interest to note that STx B nanoparticles dampened the membrane’s thermal fluctuations in our DPD simulations. Importantly, this suppression of membrane fluctuations was not observed under the same conditions in which loss of rigidity or the presence of long flexible linkers led to a loss of clustering. The thermal Casimir effect is also expected to increase with nanoparticle size and rigidity, which agrees with our simulations.

One unintuitive aspect of the Casimir force is that it is nonadditive as pointed out in several previous studies. The total Casimir force between any pair of particles in a many body system is still attractive and can be larger than the sum of the pairwise forces.

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Probing the thermal Casimir force in membranes experimentally is a challenging task. It is not possible to use the temperature or membrane tension as control parameters
because membranes undergo phase changes or rupture events within a relatively small range of temperatures or values of membrane tension. The relevant short-distance behavior caused by the Casimir-like force is practically insensitive to changes within this experimentally accessible regime of temperatures or membrane tension values.

For experimental testing, we therefore exploited the fact that the thermal Casimir force arises only when membrane inclusions locally suppress membrane fluctuations as validated in our simulations. Using Gb3 species for which flexible linkers separate the carbohydrate part from the ceramide backbone, thus removing the suppression of the membrane fluctuations at the binding sites of the B-subunit of Shiga toxin (STxB), we could experimentally show that a tight mechanical coupling of STxB onto the membrane was required for toxin clustering. Therefore, unlike the capillary forces mentioned above, whose removal one by one did not prevent the clustering, when the conditions for the thermal Casimir force are removed (by displacing the toxin particles with the linker) the clustering is eliminated, indicating that the thermal Casimir force is essential for clustering to occur.

The thermal Casimir force hypothesis provides a fresh view of clustering processes that operate in concert on biological membranes. It is also versatile compared to mechanisms such as capillary attraction because the only constraint on the (fluid) membrane composition is the presence of an appropriate binding partner (e.g., the GSL Gb3 for STxB). We hypothesize that bacterial and viral evolution proceeding only by modifying the protein sequence of the infectious disease particles has harnessed a generic, fundamental force to initiate cellular entry. Many pathogens and pathogenic factors bind to GSLs for their entry into cells, indicating that a Casimir-like force-driven clustering mechanism may apply also to these. Furthermore, thermal Casimir interactions should occur for manufactured rigid nanoparticles that tightly adsorb onto the plasma membrane. The clustering mechanism that we describe here is therefore expected to contribute to cellular, pathogenic, and pharmaceutical drug delivery processes.

**MATERIALS AND METHODS**

**Preparation of GUVs.** 1,2-Dioleoyl-sn-glycero-3-phosphocholine (DOPC) was purchased from Avanti Polar Lipids (Alabaster, Alabama, USA). GUVs were prepared by electroformation. One microliter of DOPC/Chol (X = C22:1, C22:1 Gb3_EG, C22:1 Gb3_EG1, or C22:1 Gb3_EG2) at 95:5 ratio at 1 mg/mL in chloroform was spread on two indium tin oxide-coated glass plate electrodes that were spaced 4 mm apart. The electrodes with the lipid films were immersed in a chamber containing 320 mM sucrose solution and were connected to a power generator. Electroformation was performed at 2 V and 10 Hz for 1 h at 65 °C. The GUVs were released from the electrodes by changing the frequency to 2 Hz for 30 min, and transferred into a chamber with equimolar PBS.

**Micropipette Aspiration.** Glass micropipettes were prepared using a Sutter P-2000 micropipette puller (Novato, USA). The glass micropipettes were passivated with β-casein at 5 mg/mL. GUVs were placed in a homemade chamber consisting of two coverslips separated by 1 mm, aspirated with a glass micropipette, and membrane tension was adjusted by changing the difference of hydrostatic pressure. For each GUV used for FCS measurements, the membrane was stressed to approximately 10−3 N/m to minimize intensity variations contributed by the fluctuations of the membrane along the long axis of the focal spot.

**Fluorescence Correlation Spectroscopy.** FCS was performed using a Picoquant accessory FCS unit on a Nikon confocal scanning microscope. STxB was labeled with Alexa488 (Life Technologies), according to the manufacturers instructions. The 488 nm beam was focused on the bottom pole of the membrane, and the emission was collected from 510 to 560 nm using a dichroic mirror and a filter in front of the avalanche photodiodes. Data were collected using Symphotime64 software and exported in ASCII format for independent data analysis. Every 6 min, data were collected in run lengths of 30 s for a total duration of 30 min and then grouped. The autocorrelation curve for every recording was averaged and used for MEMFCS analysis (Supporting Information, Figure S113).

**Data Analysis.** The FCS data was analyzed according to the maximum entropy method using:

\[
G(\tau) = \sum_{i=1}^{n} \left( \frac{1}{1 + \tau/T_{0,i}} \right) \left( \frac{1}{1 + \frac{2}{N_{0,i}}} \right)^{1/2}
\]

MEMFCS minimizes \(\chi^2\) as well as maximizes the entropic quantity, defined by \(S = \sum p_i \ln p_i\), which then detects a distribution of diffusion times, the amplitude of which is related as \(p_i = \frac{1}{\sum q_i}\) for each event. \(G(\tau)\) corresponds to the sum of all amplitudes. By normalizing the amplitudes, the logarithmic distribution of diffusion times represents the probability that a specific noninteracting species has a particular diffusion time. Typically, the distribution maximizes at the most probable diffusion time for a single distribution. Fits were obtained by setting the number of species to 150, with the diffusion times ranging from \(1 \times 10^{-3}\) to 1000 ms. The fits were deemed good by the residuals for all the curves and the apparent diffusion times corresponding to photophysical processes and diffusion were verified independently by a membrane binding protein, MinD (53 kDa), labeled with Alexa488. The fact that the time scales of the photophysical processes are below 0.01 ms as expected and diffusion times of MinD peak at a few milliseconds verifies the MEMFCS fitting routine and confirms the diffusion time of monomeric STxB (Supporting Information, Figure S116).

**Incorporation of Lipids into Cell Membranes.** For incorporation of lipids, GM95 cells were incubated for 48 h in serum free medium with C18:1 Gb3 or C18:1 Gb3_EG7. To minimize the amount of Gb3 nonspecifically attached to cells and the glass surface, cells were washed 3 times with medium containing 10% FCS. Subsequently cells were detached with accutase, collected by centrifugation, and replated in fibronectin-coated glass bottom dishes for 60 min. Cells were incubated with 0.5 μM STxB-Alexa488 for 20 min at 37 °C, washed to remove unbound STxB, and imaged at RT. C18:1 Gb3_EG7 incorporated into the plasma membrane less efficiently than C18:1 Gb3. To achieve comparable STxB-Alexa488 intensities on the plasma membrane, lipid and toxin concentration were adjusted: C18:1 Gb3 was incorporated at 3.3 μM, and C18:1 Gb3_EG7 at 50 μM.

**Endocytosis Assay.** STxB-Cy5 was allowed to bind on ice to GM95 cells incubated with C18:1 Gb3 or C18:1 Gb3_EG7 for 15 min in DMEM buffer supplemented with HEPES at 25 mM. Cells were shifted for 10 min to 37 °C and fixed with 4% PFA in phosphate buffer. To quench the fluorescence of membrane-bound, non-intenalized STxB-Cy5, samples were imaged in a buffer containing 0.2 M Tris at pH 9, 100 mM TCEP, 5% glucose, 1 mM ascorbic acid, and an oxygen scavenging system consisting of 0.5 mg/mL glucose oxidase, 40 μg/mL catalase, and 5% glucose.11 Exposing to 633 nm laser quenched all extracellular STxB-Cy5 and allowed distinguished the fusion time. Typically, the distribution maximizes at the most probable diffusion time for a single distribution. Fits were obtained by setting the number of species to 150, with the diffusion times ranging from \(1 \times 10^{-3}\) to 1000 ms. The fits were deemed good by the residuals for all the curves and the apparent diffusion times corresponding to photophysical processes and diffusion were verified independently by a membrane binding protein, MinD (53 kDa), labeled with Alexa488. The fact that the time scales of the photophysical processes are below 0.01 ms as expected and diffusion times of MinD peak at a few milliseconds verifies the MEMFCS fitting routine and confirms the diffusion time of monomeric STxB (Supporting Information, Figure S116).

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**Simulation Methods.** The details of the dissipative particle dynamics simulations and the model for proteins in these simulations, as well as the force field parameters in which proteins suppress membrane fluctuations, can be found in the Supporting Information.
ASSOCIATED CONTENT

Supporting Information

In the associated Supporting Information we provide movies that show the results of the DPD and MD simulations, and detailed descriptions of our experimental and theoretical work as described below. The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsnano.6b05706.

Description of the thermal Casimir force between membrane inclusions as presented in the literature and our calculations using the proximity force approximation of the thermal Casimir force between a pair of polygonal particles at separations that are small compared to their size, the dissipative particle dynamics simulations of nanoparticles adsorbed to a fluctuating membrane, showing that only when the nanoparticles suppress the membrane fluctuations do they experience a clustering force, reanalysis of trajectories from our previously published molecular dynamics (MD) simulations, showing that the binding of STxB to the membrane does not change the lipids’ chain order parameter nor their rotational diffusion, experimental protocol for adding the STxB toxin particles to the vesicles and cells, estimation of the membrane coverage due to STxB binding to Gb3 lipids, FCS experiments that are used to measure the decrease in diffusion as the toxin nanoparticles aggregate, and complete organic synthesis reactions for the PEG-modified Gb3 lipids.

Movie 1, flat pentagonal NPs with radius $R = 5.5$ nm attract after adhesion to the bilayer surface and remain attached to each other until the end of the simulation (AVI)

Movie 2, flat pentagonal NPs with radius $R = 2.4$ nm are forced to aggregate and after the force is removed (starting configuration of the system) start to separate immediately (AVI)

Movie 3, flat pentagonal NPs with radius $R = 3.5$ nm are forced to aggregate and after the force is removed (starting configuration of the system) are still attached to each other for about 4 μs (AVI)

Movie 4, flat pentagonal NPs with radius $R = 5.5$ nm are forced to aggregate and after the force is removed (starting configuration of the system) are still attached to each other for at least 9.6 μs (AVI)

Movie 5, flat pentagonal NPs with radius $R = 5.5$ nm adhere to a bilayer composed of the lipid with structure L-12, are forced to aggregate, and after the force is removed (starting configuration of the system) are attached to each other for about 2.5 μs (AVI)

Movie 6, flat pentagonal NPs with radius $R = 5.5$ nm adhere to a bilayer composed of the lipid with structure L-22, are forced to aggregate, and after the force is removed (starting configuration of the simulation) start to separate immediately (AVI)

Movie 7, spherical NPs with pentagonal base adhered to a bilayer composed of the lipid with shown in Figure SI2-1 are forced to aggregate and after the force is removed (starting configuration of the simulation) start to separate immediately showing that curvature energy does not mediate an attractive force (AVI)

Movie 8, spherical NPs with pentagonal base adhered to a bilayer composed of the lipid with shown in Figure SI2-1 are forced to aggregate and after the force is removed (starting configuration of the simulation) start to separate immediately showing that curvature energy does not mediate an attractive force (AVI)

Movie 9, 200 ns all-atom MD simulation of a STxB protein bound to a membrane from a trajectory of our previously published paper (AVI)

REFERENCES


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Formation. Induced by Shiga Toxin B-Subunit: from Molecular Structure to Tube


