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A Martini coarse-grained model of the calcein fluorescent dye

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Abstract

Calcein leakage assays are a standard experimental set-up for probing the extent of damage induced by external agents on synthetic lipid vesicles. The fluorescence signal associated with calcein release from liposomes is the signature of vesicle disruption, transient pore formation or vesicle fusion. This type of assay is widely used to test the membrane disruptive effect of biological macromolecules, such as proteins, antimicrobial peptides and RNA and is also used on synthetic nanoparticles with a polymer, metal or oxide core. Little is known about the effect that calcein and other fluorescent dyes may have on the properties of lipid bilayers, potentially altering their structure and permeability. Here we develop a coarse-grained model of calcein that is compatible with the Martini force field for lipids. We validate the model by comparing its dimerization free energy, aggregation behavior at different concentrations and interaction with a 1-palmitoyl-2-oleoyl-*sn*-glycero-3-phosphocholine (POPC) membrane to those obtained at atomistic resolution. Our coarse-grained description of calcein makes it suitable for the simulation of large calcein-filled liposomes and of their interactions with external agents, allowing for a direct comparison between simulations and experimental liposome leakage assays.

Keywords: calcein, molecular dynamics, martini, coarse-graining, lipid membrane, computer simulations

S Supplementary material for this article is available online

(Some figures may appear in colour only in the online journal)

1. Introduction

Calcein is a slightly water soluble, green fluorescent dye that is widely used to study cell viability and as an indicator of lipid vesicle leakage by fluorescence microscopy imaging [1–3]. Calcein leakage assays are based on calcein self-quenching: at a concentration above 70 mM the fluorescence of the dye is quenched. In a typical leakage experiment, calcein is trapped into lipid vesicles or liposomes at a concentration above the self-quenching threshold. As calcein has a small permeability coefficient across a lipid bilayer [4], as long as the vesicles are intact the only fluorescence intensity that can be recorded is due to small leakage fluctuations. Varying the external

Original content from this work may be used under the terms of the Creative Commons Attribution 3.0 licence. Any further distribution of this work must maintain attribution to the author(s) and the title of the work, journal citation and DOI. conditions, such as adding an external agent to the solution, may lead to a change in the fluorescence intensity induced by calcein molecules that have leaked out from the vesicles to the surrounding environment, where calcein is dissolved at a smaller concentration. This is a signature of vesicle disruption, pore formation or vesicle fusion.

This type of assay is widely used to test the membrane disruptive effect of proteins [1], antimicrobial peptides [5], polycations [6], single strands of RNA [2] and nanoparticles [7–9] (NPs). Inorganic NPs, such as metal [10, 11] or oxide [12, 13] NPs, are nowadays considered as multivalent agents for biomedical applications, ranging from diagnostics to therapy. Many of these applications would require some degree of control on NP-membrane interactions: once the NPs have reached their cell target, non-disruptive membrane translocation or transient membrane damage can be effectively exploited to deliver drugs to the cytoplasm, while the disruption of significant portions of the plasma membrane can be lethal to the cell.

In this perspective, liposome leakage assays, possibly performed on synthetic vesicles, are fundamental to rationalize which physico-chemical factors drive the NP towards a gentle or disruptive type of interaction with the lipid bilayer. The role played by the NP charge, for example, has been investigated using calcein leakage assays by Goodman *et al* [8] who have found that cationic NPs induced transient poration of negatively charged liposomes, and that both cationic and anionic NPs can induce leakage from zwitterionic liposomes. Similarly, Liu *et al* [14] have shown the transient poration due to the interaction of zwitterionic liposomes with citratecapped Au NPs. Calcein leakage assays have also been used to test the effects of varying the NP core material [15], size, shape, and surface functionalization [16].

The computational approach, particularly that based on molecular dynamics (MD), can certainly be a useful complement to the experiments performed on synthetic vesicles. Simulations offer the unique opportunity to track NP-membrane interactions with atomistic or molecular details; although a fruitful comparison between experimental and *in silico* results also relies on the possibility of simulating systems whose composition is as close as possible to that of the experimental samples. Fluorescent probes then come into play.

Little is known about the effects that the many different fluorescent dyes used in the experiments, sometimes at very large concentration, may have on the properties of lipid bilayers or on the interactions between them and the NPs. Calcein is not an exception: at physiological pH the molecule is expected to bring four negatively charged COO⁻ groups, which may lead to a favorable interaction with the polar lipid head region. At large concentrations (>100 mM) this interaction may have effects on the structural and mechanical properties of the bilayer, in turn affecting bilayer permeability and its interaction between external agents or NPs. For these reasons, it is of interest to study the calceinmembrane interactions using a computational approach.

Here we present the development of a coarse-grained (CG) model of calcein that is compatible with the popular Martini force field for lipids [17–19]. The model resolution makes it suitable for the simulation of calcein-filled liposomes and of their interactions with external agents. We base our model on the structural properties of calcein, as obtained from atomistic simulations, and validate it by comparing the aggregation properties of the CG calcein in water to those obtained at atomistic level.

As calcein is charged at physiological pH, it is interesting to evaluate the performance of different versions of the Martini force field, which treat the electrostatic interactions differently. We compare three versions of Martini: the first is the standard Martini force field [17] (STD). In STD Martini, electrostatic interactions are described as short-range interactions and the water polarizability is implicitly taken into account with a uniform dielectric constant $\varepsilon_r = 15$. The long-range contribution to the electrostatic interactions can be added to the STD model via the particle-mesh Ewald (PME) method, an approach that has been shown [20] to improve the performance of the model when dealing with highly charged molecules. Here, we will always include the PME energy term in our STD simulations. The second model we use explicitly includes water polarizability [18] (PW). The model uses PME and a uniform dielectric constant $\varepsilon_r = 2.5$. We also test the polarizable water model (refPW) developed by Michalowsky *et al* [19], in which water dielectric properties are refined with respect to the original PW model.

In the methods section we provide all the details about the force field parameters, MD run set-up and system composition; in the results section we present the development of the CG Martini model of calcein, and its validation in terms of dimerization free energy, aggregation behavior at different concentrations and interaction with a 1-palmitoyl-2-oleoyl-*sn*-glycero-3-phosphocholine (POPC) membrane. Eventually, the different CG model performances and perspective used are discussed.

2. Methods

Both atomistic and CG simulations, as well as the analysis, are performed using the Gromacs 2016 MD software [21].

2.1. Atomistic simulations

Atomistic simulations are performed with the OPLS unitedatom (OPLS-UA) force field [22], the TIP3P water model [23] and the Berger parameters for lipids [24]. The time step is set to 2 fs. All simulations are performed in the NpT ensemble: the temperature is kept constant at 310K by the velocity rescale thermostat [25] ($\tau_T = 2$ ps); the pressure is kept constant at 1 bar by the Berendsen barostat ($\tau_p = 1$ ps) for the equilibration runs and by the Parrinello–Rahman algorithm [26] ($\tau_p = 1$ ps) for the production runs. The cut-off of the Van der Waals interactions is set to 1 nm, while the electrostatic interaction is treated via the PME method with a grid spacing of 0.12 nm.

The simulation of a single calcein in water is made in a cubic box of $5 \times 5 \times 5 \text{ nm}^3$ filled with water and four Na⁺ counterions. The unbiased simulations with calcein at different concentrations are made in a cubic box of $10 \times 10 \times 10$ nm³ containing water, salt at physiological concentration and Na⁺ counterions. At 30 mM calcein concentration, 20 calcein molecules are solvated by about 32000 water molecules; at 150 mM calcein concentration, 100 calcein molecules are solvated by about 30000 water molecules. The simulations for the dimerization process are made with two calcein molecules solvated by about 33000 water molecules and counterions in a dodecahedron box (radius of the inscribed sphere about 5.5 nm). To study the interaction between calcein and a model zwitterionic lipid membrane we set-up a cubic box of $6 \times 6 \times 10 \text{ nm}^3$ with a symmetric membrane composed of 114 POPC lipids, 7100 water molecules, salt at physiological concentration and counterions. Simulations are initialized with the calcein molecule placed on top of the membrane, in the water phase.

2.2. Coarse-grained simulations

The CG simulations are performed with the Martini force field [17]. The time step is set to 20 fs. All simulations are



Figure 1. Left panel: chemical structure (top) and atomistic visualization (bottom). In the top image, colored circles indicate the mapping of atoms onto CG beads. Right panel: CG description, mapping showing bead types, their numbering and the bonds between them. Thinner lines represent harmonic bonds, while thicker ones are constrained bonds. Bottom-right: visualization of a CG calcein molecule.

performed in the NpT ensemble with temperature and pressure set to 310K and 1 bar, respectively. The velocity rescale thermostat [25] ($\tau_{\rm T} = 1 \, {\rm ps}$), Berendsen ($\tau_{\rm p} = 4 \, {\rm ps}$) and Parrinello–Rahman [26] ($\tau_{\rm p} = 12 \, {\rm ps}$) barostat are used. For the simulations with the STD model, the cut-off of the Van der Waals interactions is set to 1.1 nm and the dielectric constant is $\varepsilon_{\rm r} = 15$. The simulations performed with the PW model have a Van der Waals cut-off set to 1.2 nm and a dielectric constant $\varepsilon_{\rm r} = 2.5$. For the refPW model the Van de Waals cut-off is 1.1 nm. In all cases, the Verlet scheme for the neighbor list update and the PME method for the electrostatic interactions, with a grid spacing of 0.12 nm, are used.

The simulation boxes are set-up in the same way as in the atomistic case with the same box dimensions and number of calcein molecules. A factor of 4 to 1 is considered to rescale the number of water molecules from atomistic to CG. Counterions are always present and ions at physiological concentration are present as in the atomistic case.

2.3. Umbrella sampling simulations

The analyses of the umbrella sampling [27] simulations are performed with the WHAM [28] algorithm using the 'gmx wham' tool [29] of the Gromacs suite. The order parameter for the umbrella sampling of the dimerization process of two calcein molecules is the distance between their centers of mass (COMs). We sample, both at atomistic and CG level, 25 windows: from a distance of 0.4 nm to 5 nm. In the atomistic case, each window is equilibrated for 10 ns followed by a production run of 50 ns. In the CG case, the equilibration runs are 50 ns long while the production runs are 500 ns long. The total simulated time is about 1.5 μ s (atomistic) and 13.7 μ s (CG).

With regards to the umbrella sampling simulations of the desorption process of one calcein molecule from the POPC membrane we use, as an order parameter, the projection along the *z*-axis of the distance between the COM of the membrane and the COM of the calcein molecule. In this case 17 windows are sampled: from 1.6 nm to 4 nm. Each window is initialized with a frame extracted from the corresponding unbiased simulation. At about 1.6 nm the calcein molecule is partially embedded in the membrane head region while, at a distance larger than 5 nm, it is in the water phase. Each window of equilibration and production runs lasts 15 ns + 75 ns (atomistic) and 50 ns + 800 ns (CG). The total simulated time is about 1.5 μ s (atomistic) and 14.5 μ s (CG).

2.4. Contacts and aggregates analysis

The contacts analysis is performed with the 'mindist' Gromacs tool. Two atoms or beads are considered in contact if their distance is lower than a threshold of 0.8 nm. The analysis of the aggregates is performed with the 'g_aggregate' tool, a Gromacs compatible tool developed by Barnoud *et al* [30]. Two calcein molecules are assigned to the same cluster if the minimum distance between their atoms or beads is lower than a cut-off value which we set to 1.2 nm.

3. Results and discussion

3.1. Atomistic model

The parameterization of the atomistic model of calcein is based on the OPLS-UA force field developed by Jorgensen *et al* [22]. The full details of the topology are available in our open online repository [31].

In the left panel of figure 1 the chemical structure of calcein and its atomistic UA representation are shown. We have a central hydrophobic region that is highly rigid and planar and the 1-phthalanone perpendicular to the central region. The atom type assignment, the bonded and non-bonded interaction



Figure 2. Distributions of the dihedral angle between beads 5–6–8–9 (left panel) and beads 9–12–10–11 (right panel) (see figure 1 for bead numbering). Blue represents the atomistic model (OPLS), green is the coarse-grained standard Martini model (STD) and orange is the Martini model with water polarizability (PW).

parameters, as well as the partial charge assignment, are derived from the OPLS-UA parameters for hydrocarbons and proteins [22, 32, 33] and from the AMBER parameters for nucleic acid and proteins [34, 35].

We added improper dihedrals to impose the planarity of the four ethanoate groups ($-CH_2-COO^-$), that of the three rings of the central region, that of the γ -butyrolactone and that of the benzene rings. In the case of the two sp³ N atoms, we added an improper dihedral to keep the tetrahedral geometry.

3.2. Martini model development

3.2.1. Mapping. In the Martini force field, each particle (or bead) represents a chemical building block. The CG bead interactions are parameterized so as to reproduce the free energy of transfer from water to oil of the chemical group they represent. Two bead sizes are comprised within the model: normal beads have a Lennard–Jones σ parameter of 0.47 nm and typically represent groups of four heavy atoms; small beads have a $\sigma = 0.43$ nm and are used to represent both groups of three or two atoms [36, 37]. For the calcein molecule we adopt both normal and small bead types.

In figure 1 the chemical structure (left panel) with the related mapping scheme and the Martini CG model (right panel) are shown. The reader is referred to our online repository for the atomistic and CG topology files [31].

Since we are interested in the negatively charged $(4e^{-})$ form of the calcein molecule, all four carboxyl groups are deprotonated and mapped onto four Qa beads [37] that carry a negative unit charge each. The other Martini types are chosen based on the logP of the chemical group they represent, or in analogy with the Martini models of other compounds, as provided in detail in the supplementary material.

We remark that the choice of the Martini bead types does not change in the STD, PW and refPW models of calcein. 3.2.2. Bonded interactions. The parameterization of the CG bonded interactions is based on the reproduction of the target atomistic distributions of distances and angles. Atomistic distributions are obtained from a trajectory of a single calcein molecule in water [38]. In figures S1 and S2 (stacks. iop.org/JPhysD/51/384002/mmedia) of the supplementary material, bond and angle distributions are shown, respectively. The equilibrium distances or angles and the force constants of the harmonic (or cosine-harmonic for angles) functions are chosen so as to reproduce the peak position and the width of the target distributions. Bond constraints are used whenever the force constant needed to reproduce the atomistic distribution is larger than 12 000 kJ mol⁻¹ nm⁻². The parameterization of the benzene ring is that proposed by De Jong *et al* [39].

In order to keep the central region of the molecule planar, we used two improper dihedrals (between beads 4-6-5-8 and 7-8-5-6, see figure 1) with equilibrium angles and force constants chosen to fit the atomistic distributions, as shown in figure S3. To keep bead 9 in the same plane of the central region we added an improper dihedral between beads 5-6-8-9 and the corresponding angle distribution is shown in the left panel of figure 2. The position of the 1-phthalanone is kept perpendicular to the central region by an improper dihedral (beads 9-12-10-11, equilibrium angle of 180° and force constant of 80 kJ mol⁻¹ rad⁻²) and by an angle interaction between beads 5-9-10. At CG level, beads 9-12-10-11 are coplanar, as shown in the right panel of figure 2.

3.3. Validation

In calcein release experiments the calcein solution inside the liposome, at high concentration (>70 mM), is quenched by both static and dynamic quenching mechanisms [40, 41]. We thus validate our CG models based on dimerization free



Figure 3. The PMF of the dimerization process of two calcein molecules as a function of their distance, *d*. The PMF profiles are shifted to have a value of 0 kJ mol⁻¹ in the water phase. Each profile is also shifted by the factor $2k_BT \ln(d)$ to take into account the Jacobian of the transformation from Cartesian to spherical coordinates [42].



Figure 4. A snapshot of the dimer states. In panel A, the atomistic and CG dimer state in which the distance between the calcein COMs is about 0.5 nm and we observe the staking of the central regions. In panel B, the CG dimer in which we observe the stacking of the benzene ring of the 1-phtalanone and the distance between the COMs is about 0.8 nm.

energies and aggregation behavior of calcein at different concentrations.

3.3.1 Dimerization process. We calculate the potential of mean force (PMF) of the dimerization of two calcein molecules in a water solution. The PMF is obtained from umbrella sampling simulations in which the order parameter is the distance between the COM of the two molecules. For details of the simulation set-up see section 2.

A comparison of the PMF between the atomistic and CG models is shown in figure 3. As we can see, all models agree that the bound state is more stable than the dissolved state. The free energy difference between the unbound and bound state is about 17 kJ mol⁻¹ for the atomistic model. The STD and refPW model overestimates it, with a free energy difference of ~29 kJ mol⁻¹ and ~20 kJ mol⁻¹, respectively, while

the PW model slightly underestimates it, with a difference of ~11 kJ mol⁻¹. The atomistic, PW and refPW models predict a small barrier for aggregation, that is not captured by the STD model. In all cases, the free energy barriers for dimerization and separation are of a few $k_{B}T$, allowing for the dynamic formation and disruption of small aggregates on experimentally relevant time scales. The width of the dimer state well is not the same in the atomistic and CG models: the latter (particularly the PW model) shows two dimer states that slightly differ in energy. In figure 4 the two states are shown. The CG dimer that corresponds to a distance between the calcein molecules of about 0.5 nm (left panel of figure 4, state A) is the same as that observed in the atomistic case. State A is more populated in the CG simulations with the STD model. This state is characterized by the flat contact between the central planar regions of calcein, with the 1-phthalanone ring pointing to the outer



Figure 5. The total number of aggregates as a function of the time. Data are smoothed for clarity of visualization; the raw data are shown in figure S4. Left panel: a system with calcein at 30 mM. Right panel: a system with calcein at 150 mM. In the system with calcein at 150 mM we have also tested the refined Martini polarizable water model (refPW) shown in red.



Figure 6. Contacts between the atoms or beads of the benzene ring of the 1-phthalanone in the unbiased simulation of calcein at 150 mM concentration.

direction. The other dimer state, at about 0.8 nm (right panel of figure 4, state B), is the one in which the benzene rings are facing each other in a sandwich π - π stacking. This state is the most populated in the CG simulations with the PW and refPW models and is not observed in atomistic simulations.

3.3.2. Aggregation behavior. The self-quenching concentration threshold for calcein is about 70 mM. We set-up atomistic and CG simulations at two different calcein concentrations, well above and well below the threshold: 30 mM and 150 mM. We use the atomistic system as a reference and compare it to the STD and PW systems at both concentrations; for the system at 150 mM we also test the refPW. Figure 5 shows the number of aggregates as a function of time as obtained with the different models.

At both concentrations, but with different kinetics, the atomistic simulations predict that the calcein molecules aggregate into one single cluster. The STD model agrees with the atomistic result at both concentrations. On the contrary, the CG models with water polarizability are not able



Figure 7. Snapshots of the membrane-bound configuration at atomistic (left) and CG Martini standard (right) level. The head region of the lipid membrane is shown as a grey surface. The hydrophobic lipid tails are shown as thin grey sticks. The water molecules and ions, are not shown for clarity.

to reproduce this effect in any of our simulations. By visualizing the PW or refPW trajectories, we observe a dynamic formation-destruction of micelles of calcein molecules that do not tend to form a stable cluster. The time scales of the simulations are certainly too short to exclude the possibility of disaggregation events for the atomistic and STD models: indeed, at large calcein concentration, both the atomistic and the STD CG model predict transient disaggregation events, on time scales of microseconds.

With regards to the structure of the aggregates, we also observed that the PW model has the tendency to stabilize micellar aggregates in which the benzene rings of the 1-phthalanone face the interior of the micelle. On the contrary, visual inspection of the atomistic runs suggests that most of the benzene rings are in the water phase. We calculated the number of benzene-benzene contacts, as shown in figure 6. We see that the CG models give a number of benzene-benzene contacts larger than the atomistic model by a factor of 1.5–2. We conclude that the STD model is, among the CG models tested here, the one that best reproduces the atomistic aggregation behavior, characterized by the formation of large aggregates with possible detachments on time scales of microseconds; as for the structure of the aggregates, all the CG models tested share the limitation of an overestimation of the benzene-benzene contacts.

3.3.3. Calcein-membrane interactions. To test the calceinmembrane interactions at atomistic and CG level, we ran a set of unbiased simulations (5.7 μ s at atomistic level and 20 μ s at CG level, for each model) comprising a POPC membrane (114 lipids) and a single calcein molecule, initially placed far on top of the membrane surface.

For the atomistic and STD models the calcein molecule binds to the membrane and unbinds spontaneously. Binding and unbinding events can be monitored by tracking the distance between calcein and the COM of phosphate groups, as shown in figures S6 and S8. Two metastable states can be distinguished in which the calcein molecule interacts with the membrane. In the first state, or bound state, the hydrophobic benzene ring of calcein interacts with the glycerol groups of the lipids. The correlation between the calcein-COM of membrane distance and the number of contacts between the benzene ring of calcein and the lipid glycerol groups is shown in figure S7. A snapshot of this membrane-bound configuration is shown in figure 7 for both the atomistic and STD resolutions. In CG simulations, in this configuration the calcein molecule is at a distance of about 0.1 nm from the phosphate groups; in atomistic simulations the distance is about 0.2 nm. In the second metastable state, or adsorbed state, the calcein molecule only interacts with the phosphate groups, and can be found at a distance of 1.2 nm (STD) and 0.7 nm (atomistic). On the contrary, the simulation with the PW model does not sample any bound or adsorbed state in the simulation time. The refPW model still predicts that the water phase is the most favorable state, but with small probability compared to the STD and atomistic models, it also predicts a bound metastable state at a distance of 0.25 nm from the phosphate group (see figure S8).

To be more quantitative, we calculated the PMF of the binding process of one calcein from the water phase to the membrane-bound state, showed in figure 8. The energy profile reveals two metastable states for both the atomistic and the STD model. The refPW model predicts one metastable bound state, but its most favorable configuration is in the water phase. The PMF confirms the absence of any stable state near the membrane for the PW model. The STD energy barrier to go from the adsorbed state to the bound state is about 3.5 kJ mol⁻¹, while the adsorbed state is more stable than the bound state by ~1 kJ mol⁻¹. The barrier for the atomistic model is higher, and it sets to 5.4 kJ mol⁻¹. The energy barrier connecting the dissolved state to the bound state for the refPW model is 6.4 kJ mol⁻¹.

3.4. Discussion

We can now summarize and discuss the similarities and the differences between the three CG models of calcein we have tested. All the models predict that the dimerization process



Figure 8. The PMF for the binding process. It is obtained from umbrella sampling simulations in which the order parameter is the z distance between the COM of the calcein molecule and the COM of the membrane. The PMF is plotted as a function of the z distance of the calcein-COM from the phosphate groups, for clarity. The PMF profiles are shifted to zero in the water phase.

is thermodynamically favored (figure 3), in agreement with the prediction of the atomistic force field. Only the polarizable water models (PW and refPW) are able to reproduce the energy barriers of the binding and unbinding processes. The STD model overestimates the population of the dimer state over the dissolved one and does not describe the small barrier for aggregation that is foreseen by the PW, refPW and atomistic models. The behavior of the STD model, which we have employed here with the addition of long-range electrostatic interactions, is likely to be attributed to an overestimation of hydrophobic interactions that has previously been reported for other aromatic compounds [43, 44]. A more detailed inspection of the CG dimers reveals the existence of a metastable minimum in which the benzene rings of the 1-phthalanone are in π - π sandwich stacking, a configuration that is not sampled during the atomistic umbrella sampling calculations.

The overestimation of the hydrophobic interactions by the CG models is also confirmed by the unbiased simulations of calcein aggregation. The number of ring-ring contacts is overestimated by all CG models, particularly by the STD model. However, in terms of aggregation, the agreement between the PW models and the atomistic model is not as satisfying as one could expect based on the dimerization free energy profile. Both the CG models that include water polarizability (PW and refPW) tend to form a highly dynamic number of calcein micelles that do not aggregate in a stable cluster. These micelles are formed by a variable number of calcein molecules, depending on the calcein concentration, as shown in figure S5 in which the maximum size of the micelles is plotted as a function of the simulation time. A common characteristic of these micelles is that the packing of the molecules is driven by the 1-phthalanone rings that tend to stick to each other in the inner part of the micelle. On the contrary, at atomistic resolution the packing of the molecules is driven by the stacking of the central plane of calcein. During the model optimization we explored alternative mappings with the aim to improve the performance of the PW models in this respect. We have tried to have a less hydrophobic benzene with the introduction of a new Martini bead type that is equal to the SC5 bead but has a decreased self-interaction (-6%) and an increased interaction with the water beads (+6.5%). The improvement is little and does not justify the introduction of a new bead type. The final PW model uses an Nda type to describe the central C-O-C group of calcein. A straightforward choice based on standard Martini mappings would suggest to use the Na type, which has only hydrogen acceptor character. However, the Nda-Nda interactions are stronger than the Na-Na interactions $(4.5 \text{ kJ mol}^{-1} \text{ versus } 4 \text{ kJ mol}^{-1})$, and thus provide a better aggregation behavior. We have also tried to further increase the self-interaction of the beads of the central plane (beads 4, 5 and 7, up to +11%). Again, this fine tuning did not significantly improve the aggregation behavior of the PW or refPW models. Overall, and particularly in terms of aggregation at high concentration, the behavior of the STD (with long-range electrostatics) model seems in qualitative agreement with the atomistic model.

Experimentally, fluorescence lifetime measurements [41] suggest that calcein's self-quenching method is based on both static quenching and dynamic quenching, the latter prevailing over the former. The results obtained with the atomistic and CG STD model would suggest that the static quenching deriving from calcein-calcein aggregation (over time scales longer than the fluorescence lifetime, 4 ns) is the main mechanism of fluorescence quenching; on the other hand, PW models show an

aggregation behavior which is more consistent with a dynamic quenching mechanism.

Our simulations show a good agreement between the STD model predictions and the atomistic ones in terms of interaction with the lipid membrane. Both the STD and atomistic models indicate the existence of two metastable states in which the calcein is in close contact with the membrane (the bound state is shown in figure 7). On the contrary, the PW model shows no interaction at all between the negatively charged calcein and the membrane surface (figure 8). The refPW model slightly improves the situation, but it does not predict stable binding either (figure 8). This latter behavior is at odds with the general experimental observation of stable interactions between phosphatidylcholine membranes and anionic macromolecules [45] and nanoparticles [9, 46]. A possible explanation for this behavior could be the much larger hydration of the lipid head groups that is provided by the PW model. We calculated the density of water as a function of the z component of the distance from the COM of a POPC membrane, in the absence of calcein, with the OPLS-UA, STD, PW and refPW models. Table S1 contains the water density recorded in the lipid phosphate plane and shows that the STD underestimates the atomistic water density by 4%, while PW and refPW overestimate it by 11% and 14%, respectively. Head group hydration could screen the favorable interaction of the negatively charged groups of calcein with the membrane choline groups. These observations lead us to the conclusion that the use of the STD model of calcein could be preferred over the use of the PW model, particularly when looking at calcein in the presence of a lipid membrane.

We anticipate that our CG model of calcein will be useful to the interpretation of vesicle and liposome leakage assays. Different external agents, such as cell penetrating peptides, polymer and inorganic NPs, can damage the lipid bilayer via different mechanisms and to a different extent. In order to achieve a molecular interpretation of the experimental data, the possibility for inclusion of the leaking dye in the simulations may help to clarify the role played by the dye itself during membrane poration.

Calcein is also used as a model metabolite to study the permeability of gap junction channels. Zonta *et al* [47], for example, used calcein to probe the permeability of the human connexin 26 channel protein to negatively charged molecules bearing different total charges. They found that while the neutral form of calcein is able to cross the channel and the transition rate is comparable to that obtained experimentally, the channel is impermeable to the fully deprotonated $(-4e^-)$ calcein. With our CG model it would be relatively easy to change the protonation state of the dye and to assess its role during translocation [48], accessing longer time scales than with a pure atomistic approach.

4. Conclusions

In this paper we have developed and tested a CG model of calcein: a fluorescent dye routinely used in liposome leakage assays. The model is compatible with the Martini force field, and as such is suitable for the simulation, via MD, of the interaction between synthetic nanoparticles and large lipid bilayers (e.g. calcein-containing liposomes with diameters of tens of nm). The availability of a CG model of calcein is a step forward in the direction of a better integration of experimental and computational data, as the model allows us to perform simulations in which the effect of the dye during the membrane poration process is explicitly taken into account and with molecular resolution.

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