From dioxin to dioxin congeners: understanding the differences in hydrophobic aggregation in water and absorption into lipid membranes by means of atomistic simulations†

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Translocation of small molecules through a cell membrane barrier is a fundamental step to explain the response of cells to foreign molecules. Investigating the mechanisms through which this complex process takes place is especially important in the study of the adverse effects of toxicants. In this work, we start from the results of a previous simulation study of the mechanism of dioxin (2,3,7,8-tetrachlorodibenzo-p-dioxin) absorption into a model membrane, and extend it to four structural congeners of dioxin. The new molecules have been chosen taking into consideration the structural features that characterize dioxin: aromaticity, planarity, the presence of chlorine and oxygen atoms, and hydrophobicity. Our results for the absorption mechanism confirm our expectations based on the chemical structures, but also reveal some interesting differences in single-molecules and especially in cooperative actions underlying cluster absorption. The analysis of key parameters, such as free energies of transfer and translocation times, supports the idea that dioxin, more than its congeners investigated here, likely accumulates in cell membranes.

1 Introduction

The term “dioxin” is commonly used to refer to a family of toxic chemicals that share key structural features and cause harm through similar mechanisms of action. Overall, the dioxin family has more than two hundred members, based on polichlorinated dibenzo-p-dioxins (PCDDs), and polychlorinated dibenzofurans (PCDFs). Although not strictly dioxins, polychlorinated biphenyls (PCBs) also are included within this broad class. Only about thirty congeners of dioxin are of primary concern in risk assessment, with 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD, hereafter) recognized as the most toxic.

Dioxins are unintentionally produced as by-products in a number of human activities. These include combustion processes such as waste incineration, backyard trash burning, as well as some industrial processes such as chlorination processes, plastic and herbicide manufacturing. Once released in the environment, dioxins accumulate preferentially in sediments and soils, and, to a lesser extent, in air and groundwater. Depending on the exposure level, the toxic effects related to dioxin contamination in animals include immunological and hormonal dysfunctions, reproductive diseases, and teratogenesis. Studies of humans exposed to dioxins6,7 suggest that dioxins can cause chloracne, metabolic disorders, impaired reproduction, abnormal development, suppression of the immune system, and other systemic problems, including cancer.8–11

The toxic potency of dioxins is related to their chemical stability, and tendency to accumulate in fat tissues, where they can reach high concentrations prior to excretion. After the Seveso accident,12 the mode of action of TCDD and related dioxins has been quite extensively studied, and involves binding to the aryl hydrocarbon receptor (AhR).13,14 The AhR is a ligand-dependent transcription factor that induces expression of a number of genes encoding drug metabolizing enzymes.15 In the absence of a ligand the dioxin receptor is present in a latent conformation in the cytoplasmic compartment of the cell16 associated with the molecular chaperone hsp90.17 Binding of dioxins to the PAS-B domain of AhR triggers the translocation of AhR to the nucleus and its heterodimerization with the AhR nuclear translocator (Arnt).13,16,18 Finally, the AhR–Arnt complex binds xenobiotic response elements that encode the transcription of metabolizing enzymes such as Cyp1a1, Cyp1a2, and Cyp1b1.19

The very first step of this route is represented by the absorption and diffusion of the toxic molecules into cell membranes. Several
observations in both humans and animals have indicated that, although present in many tissues and organs, dioxins primarily accumulate in fat tissues because of their hydrophobic character.\textsuperscript{21,22} Phospholipids represent the major components of all cell membranes. Their lipophilic tails likely provide an ideal environment for dioxin storage and accumulation. Although not necessarily the final target of toxic action, cell membranes may behave as storage depots for dioxins, thus becoming an internal source of chronic exposure to these pollutants.\textsuperscript{21} Understanding dioxin absorption and distribution in cell membranes is therefore essential to elucidate dioxin toxicokinetics. The complexity of biological membranes and the lack of experimental data make this a challenging task.

Membranes are dynamic assemblies of molecules,\textsuperscript{23} characterized by the capability of self-ordering in a water environment. The absence of covalent bonds between molecules allows subtle structural variations in response to the environmental changes. Atomistic molecular dynamics (MD) approaches have become standard numerical tools to study these variations at the molecular level.\textsuperscript{24–26} So far, these methods have been successfully applied to passive membrane transport,\textsuperscript{27–34} with the aim of characterizing the dynamics of the absorption process,\textsuperscript{33,34} as well as the effect of solute molecules on membrane properties.\textsuperscript{30,32} Current MD simulations cannot deal with the tremendous complexity of real cell membranes.\textsuperscript{23} Nonetheless, the use of simplified model membranes, essentially consisting of phospholipid bilayers, provides a good starting point to answer some fundamental questions and can help in the development of more refined models. Recently,\textsuperscript{35} we have applied MD to the study of TCDD translocation from water into a 1,2-dipalmitoyl-sn-glycero-3-phosphocholine (DPPC) bilayer membrane. Through the calculation of the free energy of transfer, we have shown that TCDD absorption is a spontaneous process, with a free energy minimum at about \(-40\) kJ mol\(^{-1}\) with respect to the aqueous phase. Our simulations have also revealed the tendency of TCDD clusters to quickly penetrate the membrane, and soon disaggregate once the absorption process completes. Although the concentration of TCDD in our simulations was higher than that usually observed in experimental data, this modeling approach can provide insights into the mechanisms of dioxin absorption and distribution in cell membranes.

To ensure consistency with our previous work, we used the same 1,2-dipalmitoyl-sn-glycero-3-phosphocholine (DPPC) model membrane. Besides, the model system was improved by replacing pure water with a physiological (0.15 M NaCl) solution. MD simulations were performed for both single molecules and their aggregates.

### 1.1 Simulated systems

Fig. 1 shows the molecular sketches of TCDD and the four congeners chosen for the present investigation. These are anthracene (ANTH), tetrahydrodibenzo-p-dioxin (THDD), 3,5,3\textsuperscript{'},5\textsuperscript{'}-tetrachloro-biphenyl (TCBP), and 1,2-dihydroxytetrahydrodibenzo-p-dioxin (THDO). The structure of TCDD is characterized by the presence of aromatic rings. The ether linkages between the two benzene rings formally make the compound not completely aromatic because the central ring does not conform to the Hückel rule. However, the two oxygen atoms participate in the conjugation of the benzene rings, increasing the electron availability. In addition, the four chlorine atoms contribute to enhance the benzene activity. These heteroatoms also make the compound polar and able to form hydrogen bonds with hydrogen donors. The molecule is fundamentally flat and does not possess rotatable dihedral bonds.

In order to study the importance of these structural features with respect to the interaction with the membrane, we chose four congeners whose structures can be seen as variations of the TCDD structure. ANTH is a polycyclic aromatic hydrocarbon (PAH). Compared to TCDD it can be considered its parent hydrocarbon structure. Due to its hydrophobic character and the absence of heteroatoms, ANTH is structurally the most diverse compound from TCDD. The first step from this structure to the TCDD one is the substitution of the two central CH groups with oxygen atoms. This leads to THDD, which has the same structure as that of TCDD, but no chlorine atoms. It is also hydrophobic but can accept H-bonds from water molecules. If we consider the possibility of structural variations introduced by cell metabolism, we can hypothesize the introduction of two hydroxyl groups in the 1,2 positions of the tricyclic moiety on the THDD structure, thus generating THDO. This is the simplest metabolic transformation of aromatic compounds made by oxidative enzymes, and its all-hydrocarbon version is a well known metabolite of PAHs.\textsuperscript{38} THDO is still hydrophobic, but it can accept and donate H-bonds both intramolecularly and intermolecularly. Thus, it can be expected to interact with water molecules and with DPPC headgroups more strongly than THDD and TCDD.

Flat molecules can have a privileged interaction with phospholipid assemblies because, if properly oriented, they can be expected to more easily pass through the phospholipid

![Fig. 1](image-url) Molecular structures of the selected compounds. The arrows connect closely-related chemical structures.
barrier. TCBP is an example of non-flat compounds belonging to the PCB class. This toxic molecule has four chlorine atoms (as TCDD), but no oxygen atom. TCBP is characterized by the presence of a single rotatable C–C bond between the two benzene rings. This feature makes the rotation around the biphenyl C–C bond possible, introducing a new degree of freedom. The free rotation of this bond may hinder membrane permeation. TCBP is also hydrophobic, but cannot form stable hydrogen bonds.

1.2 Numerical methods

We have performed both conventional and $z$-constrained MD simulations. All simulations were carried out using the GROMACS-5.0 program suite. Pressure and temperature were maintained at 1 atm and 325 K. For the DPPC molecules, we adopted the force field by Ulmschneider et al., whereas the TIP3P one was used for water molecules. Suitable OPLS-AA parameters were used to develop the force fields for the solute molecules. Plain simulations were carried out for systems containing 1 (N1) or 10 (N10) solute molecules. In all cases, the MD input was assembled starting from a pre-equilibrated bilayer containing 128 DPPC molecules in water (3655 molecules). In order to give more room to the solute molecules, and prevent the bilayer to interact with its own top and bottom periodic images, the size of the simulation box along the $z$ axis was doubled and filled with water molecules. Na$^+$ and Cl$^-$ were inserted afterwards in order to obtain a physiological (0.15 M NaCl) solution. The simulation box was equilibrated within the $NPT$ ensemble at 325 K, for 20 ns, in order to adjust the box size and correct the density. The final box size was about $6.5 \times 6.5 \times 11.0$ nm$^3$. A program written by our group was then used to insert the contaminants in the aqueous phase. This was accomplished by removing the minimum number of water molecules surrounding them.

After this step, $NPT$ production runs were performed for each system. For N1 systems, 3 to 5 independent $NPT$ simulations were carried out in order to investigate the absorption dynamics. The resulting trajectories were used both to calculate structural and orientational parameters, and to generate the configurations to be used in $z$-constrained simulations (see below). For N10 systems, 3 production runs were performed to assess the dynamics of cluster formation and absorption and calculate the relevant descriptors. The overall duration of these simulations was between 50 and 200 ns, although in some cases (TCBP), runs longer than 500 ns were carried out.

Beside plain MD simulations, we performed $z$-constrained simulations by constraining the distance separating the solute and the bilayer center-of-mass along the $z$-coordinate. Five independent sets were assembled for each species, each made up by twenty-one equally spaced points (from 0 to 4 nm) along the $z$ axis. For each position within each set, a 20 ns $NPT$ simulation was performed to calculate the mean force, $F(z)$, acting on the solute molecule. In order to improve the statistical convergence of the force estimates, two strategies were adopted. First, the simulation time of some “critical points” located in the polar headgroup region (between 1.8 and 2.4 nm) was extended from 20 to 40 ns in all the constrained runs. Additionally, the first 10 ns of each constrained run were considered as equilibration and neglected in the calculation of the average force and all related quantities. One additional test was also performed to verify that our $z$-constrained runs were not biased by the existence of poorly sampled regions along a coordinate orthogonal to the constrained one (see ESI† for details). The free energy of transfer ($\Delta G(z)$) from the bulk water ($z = 4$ nm) to a given position along the bilayer normal ($z$) was evaluated as potential of the mean force (PMF): \[
\Delta G(z) = -\int_0^4 F(z')dz'.
\]

The local diffusion coefficient was evaluated according to the force autocorrelation function method. To improve integral evaluations, both the average force and the local diffusion coefficient were fitted by cubic Bèzier splines. An analogous fitting process was performed on the free energies of transfer. For each set of simulations, the permeability coefficients were obtained from the solute resistance profiles, according to the inhomogeneous solubility-diffusion model. The final values of free energies of transfer, local diffusion coefficients, and solute resistances reported below were obtained, for each solute molecule, averaging over the five independent sets. The same was done for permeability coefficients and translocation times. A detailed discussion about the choice of the simulation parameters can be found in the ESI† together with the results of some numerical tests aimed at validating our calculations.

2 Results and discussion

2.1 Single-molecule simulations

We describe the results of our MD calculations, starting from the results of conventional simulations, e.g. unrestrained single-molecule simulations (N1, hereafter). All molecules were absorbed by the DPPC membrane with capture times varying between 10 and 80 ns. The capture times were different from one run to another. In most cases, the molecules were observed to repeatedly approach the membrane surface before absorption occurred. Once started, the absorption process took about 1–2 ns to complete, regardless of the molecule considered. Afterwards, the molecules were found to reach stable positions within the bilayer, according to their free energy minima (see below). Some simulations were also extended to check for the possibility of membrane crossing. This process, however, was never observed within the maximum simulation time considered in this work (about 100 ns). Numerical estimates of the translocation time will be discussed below. The trajectories extracted from these runs were used to calculate structural as well as orientational parameters, namely the membrane thickness, the area-per-lipid, the solute orientation with respect to the bilayer normal, and the distance between the solute and the bilayer center-of-mass. Regardless of the species considered, the membrane thickness was found to vary between 4.2 and 4.4 nm during the simulations. This result well compares with
those published in the literature for pure DPPC membranes.30
No significant variation of this parameter was observed upon
solute absorption. Similarly, the area-per-lipid was not signifi-
cantly affected by solute permeation. Its values ranged between
0.5 and 0.7 nm² per molecule. Fig. S1–S5 in the ESI† provide an
example of the evolution of these descriptors over time (one NPT
run for each solute molecule). These plots also display the evolu-
tion of the tilt angle (θ) between the solutes and the bilayer surface
normal. In all cases, the molecules entering the membrane were
preferentially oriented at relatively small θ. On the basis of these
observations, we conclude that the differences in the structure of
the solute molecules did not appreciably affect the mechanism of
the absorption process.

### 2.1.1 Free energies of transfer.

In order to gain more information about the absorption process, we calculated the
free energies of transfer from the water phase to the membrane
interior at different z-positions along the bilayer normal. These
were obtained as PMFs, by integrating the average force acting
on the solute molecules along the bilayer normal (see eqn (1)).
Fig. 2 shows the corresponding profiles for all the species
considered. The standard deviations, reported in the ESI†
(Fig. S11), and here omitted for clarity, were of the order of
±3 kJ mol⁻¹ for the molecules in the membrane, except for
TCBP characterized by larger deviations.

The profiles confirm that absorption is spontaneous for all
the species considered. Crossing the polar headgroups was a
barrier-free process, always followed by a steep gain in energy,
demonstrating that the molecules preferred the hydrophobic
part of the membrane to the water environment. As shown in
Table 1, each molecule has its own energy minimum at slightly
different positions inside the membrane.

The free energy minimum shows the ascending order: TCDD <
TCBP < THDD < ANTH < THDO. These results indicate that the
free energy change on transferring these molecules from water
was higher for moieties containing chlorine (TCDD and TCBP),

| Molecule | ΔG(ζ_min) [kJ mol⁻¹] | ζ_min [nm] |
|----------|---------------------|-----------|
| TCDD     | −51.58 (±2.59)      | 1.19      |
| TCBP     | −41.17 (±5.26)      | 1.08      |
| THDD     | −36.55 (±1.61)      | 1.06      |
| ANTH     | −32.84 (±2.48)      | 1.10      |
| THDO     | −29.96 (±3.17)      | 1.17      |

a Standard errors are reported in parentheses.

than for the less polar hydrocarbons ANTH and THDD. The result
for TCDD agrees with our previous estimate (−43 ± 1.8 kJ mol⁻¹),
obtained for the TCDD/DPPC/water system, in the absence of
NaCl. For THDO, the result is consistent with the presence of two
hydroxyl groups. Once inside the membrane the molecules move
more or less easily crossing the membrane centre. The energy
barriers for this process show small magnitudes for all species
(about 5 kJ mol⁻¹), except for THDO (about 20 kJ mol⁻¹). In fact,
for this molecule crossing of the membrane centre was never
observed during the simulations.

#### 2.1.2 Local diffusion coefficients, solute resistances, per-
meability coefficients, and translocation times in N1 systems.

Beside free energies of transfer, the average force values were
also used to compare the diffusion and permeation of single
molecules across the DPPC membrane. To this end, we calculated
the diffusion coefficient, solute resistance, permeability and
average translocation times for all species.

Fig. 3 compares the local diffusion coefficients, D(ζ), calculated
for all species. The differences among the different species are
limited. In bulk water, diffusion coefficients of the order of
2.5 × 10⁻⁵ cm² s⁻¹ were obtained. Approaching the membrane
heads the diffusion coefficients decreased, reaching the mini-
mum value inside the membrane, with an order of magnitude
(10⁻⁶ cm² s⁻¹) consistent with the results of calculations on
similar systems.32 The figure inset shows the small variations
near the membrane centre: all the compounds show greater

![Fig. 2](image1.png)

**Fig. 2** Plot of the free energies of transfer along the bilayer normal,
with ζ = 0 corresponding to the bilayer center-of-mass. Standard errors
are omitted.

![Fig. 3](image2.png)

**Fig. 3** Plot of the diffusion coefficients along the bilayer normal. The inset
magnifies the region between 0 and 2 nm.
$D(z)$ in the membrane center, where the alkyl tails are less dense. In addition, ANTH and THDD, the most lipophilic compounds, have slightly greater values than the other molecules.

Fig. 4 shows the solute resistance profiles $R(z)$ along the bilayer normal, calculated by combining the free energy profiles and the local diffusion coefficients:

$$R(z) = \frac{e^{\Delta G(z)/k_B T}}{D(z)} \, dz. \quad (2)$$

According to the above equation, $R(z)$ can be expected to show correlations with the free energy of transfer and the local diffusion coefficient. In the bulk water phase, where $e^{\Delta G(z)/k_B T}$ approaches unity and $D(z)$ is large, $R(z)$ shows small but non-negligible values for all molecules. Approaching the membrane heads, $R(z)$ steeply increases to reach a maximum in the membrane head region. Two points are worth noting. First, the maximum of TCDD is not aligned with that of the other molecules. Its location, in the water phase, is due to the steep decrease in $\Delta G(z)$ upon approaching the polar headgroups. Second, for TCBP, $R(z)$ peaks at a value two/three times larger than that observed for the other molecules (0.115 cm s$^{-1}$). Also in this case, the outcome is related to the free energy barrier upon crossing the barrier heads.

On going from the membrane heads to the interior, $e^{\Delta G(z)/k_B T}$ decreases and so does $R(z)$. As expected, the minima of $R(z)$ are very close to those of $\Delta G(z)$. Substantial differences in $R(z)$ values can be noticed across the different species, TCDD showing a minimum value about two orders of magnitude smaller than its congeners. Approaching the membrane center, $R(z)$ moderately increases for all compounds except THDO, for which a thousand-fold increase was observed.

The permeability coefficient of a membrane, hereafter abbreviated as $P$, is generally defined as the proportionality factor linking the steady-state flux of molecules per unit area ($J$) to the concentration difference between its two sides: $J = P(C_{\text{out}} - C_{\text{in}})$. Using this equation to estimate $P$ in atomistic simulations is unfeasible, due to the limited time scale accessible to MD. According to the inhomogeneous solubility-diffusion model,40,55–57 $P$ can be more conveniently obtained from $R(z)$ as

$$P = \left[ \frac{1}{\int_{-4}^{4} R(z) \, dz} \right]^{-1}. \quad (3)$$

The limits of integration correspond to the bulk aqueous phase on the two sides of the membrane. Table 2 collects the numerical values calculated using the above equation (see the ESI† for details).

| Molecule | $P$ [cm s$^{-1}$] |
|----------|------------------|
| TCBP     | 12.4 ± 4.3       |
| ANTH     | 17.0 ± 3.6       |
| THDD     | 23.8 ± 4.2       |
| THDO     | 26.8 ± 5.3       |
| TCDD     | 29.6 ± 3.5       |

Table 2 Permeability coefficients of dioxin congeners$^a$

$^a$ Standard errors are reported in parentheses.

The values of $P$ follow the ascending order: TCBP < ANTH < THDD < THDO < TCDD. For TCBP the result is dominated by the high solute resistance in the polar headgroup region. For the remaining molecules, the differences in $R(z)$ are less relevant; ANTH and THDD have similar values. The position of THDO, intermediate between THDD and TCDD, can be rationalized considering the presence of two hydroxyl groups. In an apolar environment, such as the lipophilic tails of DPPC, the two hydroxyl groups make an intramolecular H-bond that decreases the lipophobicity of the compound; this effect is well-known in the literature.32 It should be noted that, especially for THDO and TCDD, significant differences in $P$ were observed on going from one set to another, as testified by the standard deviations. In any case, our results suggest that, among the compounds considered, TCDD is the one with the strongest interactions with the membrane.

As mentioned above, no species was observed to cross the DPPC bilayer within the time scale explored by our simulations. In order to get a measure of the time required for this process to occur, we used the average force to estimate the translocation time ($\tau$) assuming Brownian motion. It should be noted that the assumption of Brownian motion not necessarily hold in cell membranes, and therefore, this calculation only provides an order-of-magnitude estimate of $\tau$. Table 3 collects the translocation times for all species, calculated according to the following equation:

$$\tau = \frac{1}{D} \int_{-4}^{4} e^{\Delta G(z)/k_B T} dy \int_{-4}^{y} e^{-\Delta G(z)/k_B T} \, dz. \quad (4)$$

Table 3 Translocation times

| Molecule | $\tau$ [s] |
|----------|-----------|
| TCDD     | 6.21      |
| TCBP     | 0.29      |
| THDD     | 1.38 × 10$^{-2}$ |
| ANTH     | 2.97 × 10$^{-3}$ |
| THDO     | 1.44 × 10$^{-3}$ |

Fig. 4 Logarithmic plot of solute resistance profiles along the bilayer normal.
where \( P \) represents the average force, while \( D \) the average value of \( D(z) \), calculated taking into account the location of the molecules after the translocation process (see the ESI† for details).

TCDD and TCBP show the highest \( \tau \) values, and therefore, tend to reside within the membrane longer than the other species considered. This is consistent with the fact these species are more stabilized by the interactions with the membrane tails (see the free energy minima in Fig. 2). On going from these molecules to the chlorine-free ones, the translocation times become two or three orders of magnitude smaller. For THDO, the translocation time is the smallest. This apparently contrasts with the increase of \( R(z) \) approaching the membrane centre. At the same time, however, the THDO resistance maximum is lower than the other molecules, finally resulting in a faster transport. We should also note that the translocation time (1440 \( \mu \)s) is anyhow far beyond the time scale that we reached with unconstrained MD simulations.

### 2.2 Many-molecule simulations

In our previous paper, we reported the results of MD studies performed on TCDD clusters made up of 1, 2, 5, and 10 molecules. In all cases, we observed TCDD aggregation to form molecular clusters. The smaller clusters could form and break up many times during an \( NPT \) simulation. Clusters consisting of 10 molecules were long lasting and very stable. In addition, we found that the absorption of one molecule belonging to the cluster immediately resulted in the sequential absorption of all other molecules.

Taking into account these outcomes, in the present study we considered N10 systems, namely systems consisting of 10 solute molecules in a physiological solution with the DPPC membrane. We performed three independent \( NPT \) runs for each species, including TCDD. The solute molecules were initially placed far apart, in order to test the possibility of spontaneous aggregation. Since, except for THDO, all molecules were hydrophobic, we expected the clusters to show the same behavior as TCDD ones, in terms of cluster formation and cooperative absorption. The results we obtained were partly unexpected.

Both TCDD and TCBP rapidly formed clusters. Once formed, the clusters were stable and we did not observe any occurrence of cluster breakdown. The clusters entered the membrane at different times. For TCDD the dynamics of cluster absorption was similar to that observed in our previous work: the molecules belonging to the cluster were absorbed sequentially, within a relatively short time (about 60 ns in two runs). TCBP clusters, in contrast, required much more time to enter the membrane. In fact, the TCBP cluster was absorbed by the membrane only in one simulation. In both remaining runs, each extended well beyond 500 ns, we never observed TCBP absorption. The solute resistance profile of the isolated molecule provides a qualitative rationale for this outcome, although it does not account for the interactions among TCBP molecules. An analysis of the geometrical properties of TCBP clusters has evidenced the tendency of TCBP clusters to keep an average distance from the membrane heads larger than that of the other molecules. We shall return to this point below.

ANTH, THDD, and THDO also formed clusters. These, however, were characterized by a highly variable composition. As a consequence, the molecules entered the membrane at different times, often isolated, or sequentially like TCDD. ANTH and THDD had similar behaviors. We observed the clusters to form and separate repeatedly during the simulations. Often, the molecules were absorbed individually, sometimes leaving the cluster in proximity of the membrane. As mentioned above, THDO can form H-bonds, and has strong interactions with the membrane headgroups. Once formed, the whole THDO cluster was observed to spend a long time at the bilayer surface, before entering the membrane. Fig. 5 compares THDO (top) and TCDD absorption (bottom).

Regardless of the species considered, all clusters rapidly broke down into separate molecules after absorption. Fig. 6 gives an example of the final distribution of the molecules for ANTH (left), and THDO (right). ANTH molecules were widely distributed all over the membrane interior. The same was also observed for TCDD, THDD and TCBP. THDO molecules, conversely, were preferentially located close to the membrane polar heads. In this figure we note that THDO molecules entered the membrane from both sides due to system periodicity. However, as for the N1 system, crossing of the membrane centre was never observed during the simulations.
To better understand the different behaviour of the clusters in the water phase, we performed an analysis of their stability. In order to qualitatively judge cluster stability, we calculated the Coulomb and Lennard-Jones intermolecular energies associated with the removal of one molecule from the largest cluster. In order to simulate the effect of this process, for a cluster comprising \( N \) molecules, we calculated the difference between two intermolecular energies, that of the cluster, namely \( E_N \), and that obtained by averaging the energies of the \( N \) sub-clusters \( (E_{N-1}, \text{hereafter}) \), containing \( N-1 \) molecules. The sub-clusters were obtained by deleting in turn each molecule from the cluster. DPPC, water, and ions, were not considered in the calculation. The calculation was performed on 500 MD frames extracted from the simulations, where the clusters were present as whole aggregates (10 molecules), with the exception of ANTH where the maximum cluster size was 9. See the ESI for details of the selection algorithm. For each species, and each MD frame, the energy difference \( \Delta E_N \) was calculated as

\[
\Delta E_N = \frac{1}{N} \sum E_N - \frac{1}{N-1} \sum E_{N-1}.
\]

Table 4 collects the \( \Delta E_N \) values, as averages over all frames. For comparison purposes, we reported the Lennard-Jones (L) and the electrostatic (C) contributions to the total interaction energy (T). It should be noted that, since the interactions of the molecules with all the other system components were neglected, \( \Delta E_N \) cannot be expected to provide a quantitative measure of cluster stability.

Lennard-Jones interactions always increased cluster stability. The stability order according to \( \Delta E_N \) is: TCDD < TCBP < THDO < THDD < ANTH. This order is consistent with the outcome of the MD simulations: TCBP and TCDD formed stable clusters. THDO exhibits a similar behaviour, however, the interactions with water and with the polar headgroups promoted cluster break-down. ANTH and THDD clusters are far less appreciable.

Even considering that the calculation is not a quantitative measure of the cluster stability, these results well agree with the inspection of the MD trajectories. Hydrophobicity is not the sole factor behind cluster formation. Intermolecular interactions between the clustered molecules also play an important role. To propose a rationale to the difficulty encountered by TCBP clusters in entering the membrane, we analyzed the values of the distance between the cluster geometrical centre and the bilayer centre. As a representative for this distance we considered the distance of each molecule from the membrane center along the \( z \) coordinate. For each species, 500 MD frames were considered, the same used in the calculation of the stabilization energies (see above). As mentioned above, in all these frames, the clusters were present as whole aggregates and were located in the water phase.

Fig. 7 shows the distribution of distances for each compound. It is easy to note that the minimum distance of TCBP from the membrane center rarely falls below 3 nm. As discussed above, both \( \Delta G(z) \) and \( R(z) \) show that the interaction between the molecules and the membrane just takes place starting at \( \sim 2.8 \) nm. Hence, the TCBP cluster has statistically less chances to penetrate the membrane.

### 3 Summary and conclusions

In this work, we used molecular dynamics to compare the absorption dynamics of TCDD in DPPC bilayers with a group of four structurally similar congeners. These were obtained by modifications of the TCDD structure with respect to its key structural features: aromatic character, planarity, presence of chlorine atoms, and hydrophobicity.

The results of single-molecule simulations have evidenced some aspects of the interaction between the solute molecules and the membrane that can be rationalized on the basis of the structural properties. We have shown that TCDD and its congeners share some features: they spontaneously enter the membrane, their free energies of transfer indicate the presence of energy minima inside the membrane. Once inside the membrane, the molecules settle preferentially at 1 nm from the membrane center, although wide displacements may occasionally occur,
leading to membrane center crossing. Beside the analogies, some important differences also emerge from our results. ANTH and THDD are hydrophobic molecules that similarly interact with the membrane. The presence of chlorine atoms has a different impact on the absorption behavior. Both TCDD and TCBP show deep energy minima and freely move inside the membrane, in agreement with their translocation times. However, the other calculated variables discriminate the two compounds: TCDD has the highest permeability coefficient and TCBP the lowest. TCBP has the highest solute resistance upon crossing the polar headgroups. These differences are likely related to the rotational freedom about the biphenyl dihedral.

Due to the presence of two hydroxyl groups, THDO is the least lipophilic compound, the sole compound that can accept and donate hydrogen bonds, which can stabilize intermolecular interactions. This is confirmed by the significant interactions with the membrane heads and the difficulty in crossing the membrane centre. Although not affecting the absorption mechanism, the presence of hydroxyl groups increases the permeability and decreases the translocation time.

The results obtained for clusters revealed some unexpected outcomes, often related to cluster formation in bulk water. Despite the hydrophobicity of all compounds, only TCDD and TCBP formed large and stable clusters made up of all solute molecules. Consistently with our previous study, TCDD clusters were absorbed quickly. By contrast, the absorption of TCBP clusters required longer times and was less frequently observed. The clusters formed by the other species, especially ANTH, were characterized by highly variable compositions. The behaviour in the water phase had important consequences on the absorption dynamics, since poorly clustered molecules were often individually absorbed. The importance of intermolecular interactions in the cluster formation and absorption was also evidenced by the analysis of cluster stability.

Eventually, compared to its congeners, TCDD emerged as the species with the highest tendency to penetrate the DPPC bilayer and accumulate in its interior. Due to the high complexity of real biological membranes in comparison with our model system, this result is only a first step toward a better understanding of the absorption and transport mechanism of TCDD into cells that represent a fundamental requisite for its toxic action.

It is clear that the compounds considered in this study only cover a fraction of the chemical space spanned by the structural moieties of TCDD. A thorough study on the effect of some functional groups on membrane absorption would have required more congeners. In this respect, however, we note that the calculation of the free energy of transfer and the related properties via z-constrained MD simulations was very CPU demanding (about 300 000 hours on a Tier-1 supercomputer), despite the limited number of compounds considered. Besides, z-constrained MD offers some advantages with respect to other methods, in that it has been validated and successfully applied to quite many solute/membrane systems. In view of the extension to more complex systems, we are currently testing other approaches, such as meta-dynamics and adaptive biasing force. The inclusion of polarization effects would additionally be considered as a major step toward a more realistic description of the molecular interactions.

**Acknowledgements**

G. S. received partial funding through computational time at CINECA (Iscrac project MASCRADE) and the Università degli Studi di Milano.

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