Molecular Simulations of Surfactant Adsorption on Iron Oxide from Hydrocarbon Solvents

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Abstract

The performance of organic friction modifiers (OFMs) depends on their ability to adsorb onto surfaces and form protective monolayers. Understanding the relationship between OFM concentration in the base oil and the resulting surface coverage is important for improving lubricant formulations. Here, we use molecular dynamics (MD) simulations to study the adsorption of three OFMs: stearic acid (SA), glycerol monoostearate (GMS), and glycerol monooleate (GMO), onto a hematite surface from two hydrocarbon solvents: \( n \)-hexadecane and poly-\( \alpha \)-olefin (PAO). We calculate the potential of mean force of the adsorption process using the adaptive biasing force algorithm and the adsorption strength increases in the order: SA < GMS < GMO. We
estimate the minimum area occupied by OFM molecules on the surface using annealing MD simulations and obtained a similar hard-disk area for GMS and GMO, but a lower value for SA. Using the MD results, we determine the adsorption isotherms using the molecular thermodynamic theory (MTT), which agree well with one previous experimental data-set for SA on hematite. For two other experimental data-sets for SA, lateral interactions between surfactant molecules need to be accounted for within the MTT framework. SA forms monolayers with lower surface coverage than GMO and GMS at low concentrations, but also has the highest plateau coverage. We validate the adsorption energies from the MD simulations using high frequency reciprocating rig friction experiments with different concentrations of the OFMs in PAO. For OFMs with saturated tailgroups (SA and GMS), we obtain good agreement between the simulations and the experiments. The results deviate for OFMs containing Z-unsaturated tailgroups (GMO), due to the additional steric hindrance, which is not accounted for in the current simulation framework. This study demonstrates that MD simulations, alongside MTT, are an accurate and efficient tool to predict adsorption isotherms at solid-liquid interfaces.

**Introduction**

The adsorption and self-assembly of surfactants at solid-liquid interfaces plays a key role in a wide range of technological processes, from nanoparticle stabilisation to ore flotation. In tribology, the adsorption of polar lubricant additives from nonpolar base oils onto solid surfaces is critical to their effective operation. For example, organic friction modifiers (OFMs), such as stearic acid (SA), adsorb and form thin (∼2 nm) self-assembled monolayers (SAMs) on steel surfaces, which significantly reduce friction and wear. Similarly, inorganic friction modifiers, such as molybdenum dialkyldithiocarbamate (MoDTC), adsorb and decompose to form thin (∼1 nm), low-friction lamellar molybdenum disulphide films on steel. Moreover, antiwear additives, such as zinc dialkyldithiophosphate (ZDDP), adsorb, form SAMs, and
decompose on steel surfaces to form thick (∼100 nm) protective tribofilms, composed mostly of metal phosphates.\textsuperscript{6}

Modern lubricant formulations contain at least ten different types of additive dissolved in a blend of base oils.\textsuperscript{7} Combinations of different additives often behave antagonistically, that is the additive blend gives poorer performance than any of the individual additives alone at equal concentration.\textsuperscript{8} Prominent examples of antagonistic lubricant additive combinations include overbased calcium sulphonate detergents with ZDDP,\textsuperscript{9} succinimide dispersants with ZDDP,\textsuperscript{10} and MoDTC with ester OFMs.\textsuperscript{11} Many of these antagonistic interactions are due to competitive adsorption processes, whereby the most surface-active component occupies the majority of surface sites and inhibits the action of the other additives.\textsuperscript{8} Therefore, it is important to understand the relative adsorption strength of the various lubricant additives from base oils on steel surfaces.

OFMs are one of the earliest and most widely studied types of lubricant additive.\textsuperscript{12} Hardy and Doubleday\textsuperscript{13} were the first to suggest that OFM friction reduction was due to the formation of vertically-oriented surfactant monolayers on the sliding surfaces. They found that fatty acids with longer chain length (C\textsubscript{4}-C\textsubscript{12}) showed lower boundary friction and that fatty acids gave lower friction than fatty alcohols.\textsuperscript{13} Bowden and Leben\textsuperscript{14} showed that the friction of SA (C\textsubscript{18}) monolayers deposited on steel surfaces with the Langmuir-Blodgett technique was initially the same as that for SAMs formed from a hydrocarbon base oil. Fowkes\textsuperscript{15} showed that OFM adsorption strength from hydrocarbon solution onto steel varied in the order carboxylic acid > amine > alcohol, and Groszek\textsuperscript{16} showed that this correlated with antiwear performance. Rowe\textsuperscript{17} also found that adsorption strength and antiwear performance were correlated for OFMs, but with a slightly different order (amine > carboxylic acid > alcohol). Using these data, Rowe developed a mathematical model to calculate thermodynamic quantities describing adsorption from experimentally-obtained wear rates.\textsuperscript{17} Spikes
and Cameron\textsuperscript{18} showed that adsorption and lubricant failure both had a similar dependence on fatty amine concentration and temperature, suggesting that lubricant failure could be due to desorption of the OFM molecules. Jahanmir and Beltzer\textsuperscript{19} studied the adsorption and friction behaviour of a wide range of carboxylic acid, ester, alcohol, amine, and amide OFMs with different alkyl tailgroup lengths, branching, and unsaturation. They found that additives that adsorbed more strongly generally gave lower friction. \textsuperscript{19} Jahanmir and Beltzer\textsuperscript{20} also built upon the model due to Rowe\textsuperscript{17} to describe the relationship between the friction coefficient and adsorption energy for a two-component lubricant (i.e. base oil and one additive), assuming that the minimum friction coefficient observed at high OFM concentration was due to the formation of a close-packed SAM.\textsuperscript{20}

Recent depletion isotherm and polarised neutron reflectometry (PNR) experiments by Wood et al.\textsuperscript{21} for the adsorption from \textit{n}-dodecane onto iron oxide surfaces have shown that OFMs with linear, saturated tailgroups (SA) do form monolayers with a high surface coverage, similar to that expected for a close-packed monolayer, at high concentration. Conversely, OFMs with kinked, \textit{Z}-unsaturated tailgroups, such as oleic acid (OA), form monolayers with much lower surface coverage. This observation explained the higher friction coefficient for OA than SA reported in nanoscale atomic force microscopy (AFM) by Lundgren et al.\textsuperscript{22} and microscale surface forces apparatus (SFA) experiments by Ruths et al.\textsuperscript{23} The same observations were made at the macroscale by Campen et al.\textsuperscript{24} for OA and glycerol mono-oleate (GMO) compared to SA and glycerol mono-stearate (GMS) using steel substrates and an \textit{n}-hexadecane solvent. Using a range of OFMs with different headgroups and tailgroups, Fry et al.\textsuperscript{25} showed through a combination of quartz crystal microbalance (QCM), ellipsometry, and tribology experiments that OFMs which formed higher coverage monolayers gave lower initial friction coefficients.

In recent years, molecular simulations have provided additional insights into the nanoscale
behaviour of a wide range of lubricant additives, including OFMs.\textsuperscript{26,27} The adsorption energy of lubricant additives on metal oxide surfaces can be accurately determined with first principles methods such as density functional theory (DFT).\textsuperscript{28–30} However, due to its very high computational cost, DFT is typically performed in vacuum at zero temperature, and can only capture the enthalpy contributions to the adsorption free energy. For many surfactants at solid-liquid interfaces, entropy plays a central role in the adsorption process.\textsuperscript{31} These entropic contributions can be calculated using classical molecular dynamics (MD) simulations,\textsuperscript{32} which can be used to access much larger time and length scales than DFT. Nevertheless, capturing the adsorption and self-assembly processes from solution using realistic surfactant concentrations represents a considerable computational challenge.\textsuperscript{27}

Until recently, a certain surface coverage of surfactant molecules on the solid surfaces was usually assumed in MD simulations of OFMs.\textsuperscript{26,27} However, Chia et al.\textsuperscript{33} and Jaishankar et al.\textsuperscript{34} used MD simulations to study the adsorption of surfactants on iron oxide surfaces from hydrocarbon solvents. Chia et al.\textsuperscript{33} studied the competitive adsorption of the fuel additive 2-[(3-(dimethylamino)propyl)amino]methyl-4-dodecyl-6-methyl-phenol and ethanol from \textit{i}-octane onto the \(\alpha\)-Fe\(_2\)O\(_3\)(0001) surface at different temperatures. They showed that, while the free energy of adsorption of the additive was approximately two times larger than that for a single ethanol molecule, a single additive needs to displace up to five ethanol molecules to strongly adsorb onto the surface. Jaishankar et al.\textsuperscript{34} studied SA adsorption from \textit{n}-heptane and \textit{n}-hexadecane onto Fe\(_3\)O\(_4\)(111), Fe\(_3\)O\(_4\)(001), and hydroxylated Fe\(_3\)O\(_4\)(001) surfaces at room temperature. They found that adsorption was stronger on the hydroxylated surface and adsorption from \textit{n}-hexadecane was much slower than from \textit{n}-heptane due to stronger alignment of the molecules on the surface. They used a combined approach using MD and molecular thermodynamic theory (MTT) to calculate adsorption isotherms, which agreed well with experimental data obtained with QCM.\textsuperscript{34} The theoretical MTT model for the adsorption of non-ionic surfactants, originally developed by Nikas et
al.,\textsuperscript{35} was applied in combination with MD for the first time by Sresht et al.\textsuperscript{36} to study surface tensions at a water-air interface. Since then, it has been used to study adsorption in many systems, such as sodium dodecyl sulphate (SDS) on graphene from water,\textsuperscript{37} SDS on $\alpha$-Al$_2$O$_3$ nanoparticles from water,\textsuperscript{38} and organomolybdenum friction modifiers on boron-doped diamond-like-carbon (DLC) from $n$-heptane.\textsuperscript{39}

Chia et al.\textsuperscript{33} and Jaishankar et al.\textsuperscript{34} extracted the potential of mean force (PMF) along the reaction coordinate using umbrella sampling (US) and the weighted histogram analysis method (WHAM).\textsuperscript{40,41} Even using modern simulation software and high performance computing (HPC) architectures, this method requires hundreds of nanoseconds of simulation time to obtain converged PMFs. For example, Jaishankar et al.\textsuperscript{34} used an overall simulation time of more than 1 $\mu$s to obtain the PMF describing the adsorption of SA onto hematite from $n$-hexadecane. A potentially more efficient method to obtain the PMF for surfactants at solid-liquid interfaces is the adaptive biasing force (ABF) algorithm.\textsuperscript{42,43} The ABF method has been used to study adsorption at solid-liquid interfaces for a range of surfactants from water, such as SDS on graphene,\textsuperscript{37} chlorogenic acid on polydimethylsiloxane,\textsuperscript{44} lipoic acid on silver,\textsuperscript{45} peptides on gold,\textsuperscript{46} proteins on hydroxyapatite\textsuperscript{47} and graphene,\textsuperscript{48} ethylene glycols on calcite\textsuperscript{49} and hematite,\textsuperscript{50} and a wide range of organic molecules on carbon nanotubes,\textsuperscript{51} graphene,\textsuperscript{52} and graphene oxide.\textsuperscript{53}

In this study, we compare the adsorption of different OFMs from hydrocarbon solvents onto iron oxide surfaces. More specifically, ABF-MD is used to study the adsorption of SA, GMS and GMO from $n$-hexadecane and hydrogenated 1-decene trimer onto the $\alpha$-Fe$_2$O$_3$(100) surface. We show that the OFM headgroup and tailgroup structure, as well as the solvent, significantly affect the adsorption behaviour. Once the adsorption PMF is obtained, the hard-disk area occupied by the molecules on the surface due to steric effects is estimated using separate annealing MD simulations. Subsequently, the relation between the bulk con-
centration of the OFMs in the solvent and the surface coverage, i.e. the adsorption isotherm, is determined using the MTT model.\(^{34,35,54}\) Finally, the Jahanmir and Beltzer model,\(^{20}\) together with High Frequency Reciprocating Rig (HFRR) friction measurements, are used to experimentally validate the calculated adsorption energies.\(^{20,34}\)

**Methods**

**System Setup**

The studied OFMs are SA, a carboxylic acid containing a saturated \(C_{18}\) tailgroup, GMS, a glyceride mono-ester containing a saturated \(C_{18}\) tailgroup, and GMO, a glyceride ester containing a \(C_{18}\) tailgroup with a \(Z\)-double bond in the middle of the chain. While SA is commonly used as a prototypical OFM, it is no longer used in commercial formulations due to its acidic nature, and has been replaced with amines, amides, and esters, such as GMS and GMO.\(^{12}\) The molecular structures of the OFMs are shown in Fig. 1a.

There has been some debate in the literature as to whether GMO and GMS adsorb and reduce friction as intact glyceride esters or are hydrolysed to form the corresponding carboxylic acid that adsorbs, self-assembles, and reduces friction.\(^{12,55–57}\) Recent experiments have shown that the initial friction of GMO solutions is much lower than OA on silica surfaces.\(^{25}\) Therefore, in this study, we consider the adsorption of intact GMO and GMS molecules.

For the hydrocarbon solvent, \(n\)-hexadecane was compared to hydrogenated 1-decene trimer. While \(n\)-hexadecane is commonly used as a model base oil in adsorption and boundary friction experiments,\(^{24,25}\) hydrogenated 1-decene trimer is the primary constituent of commercial PAO4 base oil, a poly-\(\alpha\)-olefin with a kinematic viscosity of 4 cSt at 100 °C.\(^{58}\) The molecular structures of the base oils are shown in Fig. 1b.
Figure 1: a) OFM molecular structures, from left to right: GMO, GMS and SA. b) Solvent molecular structures, from left to right: \( n \)-hexadecane and hydrogenated 1-decene trimer (PAO4).
All of the systems for the PMF adsorption calculation were constructed using the Materials and Processes Simulations (MAPS) platform from Scienomics SARL, whereas the systems to estimate the hard-disk area on the surface were constructed using proprietary scripting.

Steel is used for the majority of lubricated engineering components. When exposed to air, steel forms passivating iron oxide surfaces, such as hematite ($\alpha$-Fe$_2$O$_3$). Two slabs of $\alpha$-Fe$_2$O$_3$ with dimensions ($x \times y \times z$) of approximately $5.5 \text{ nm} \times 5.5 \text{ nm} \times 12 \text{ nm}$ were used as the substrates, with the (100) plane exposed to the liquid phase. We selected the $\alpha$-Fe$_2$O$_3$(100) surface. The surface energy of this facet from DFT calculations is approximately $1.4 \text{ J m}^{-2}$. Using the classical force field, we obtain a surface energy of $3.2 \text{ J m}^{-2}$. Although this is an overestimation, it is similar to that obtained previously ($2.3 \text{ J m}^{-2}$) using a different interatomic potential. Periodic boundary conditions were imposed in the $x$- and $y$-directions. Initially, the two hematite slabs were separated by 12 nm of vacuum space. We do not consider surface hydroxylation since this is not expected to have a large effect on the relative adsorption strength between the OFMs. Previous MD simulations showed that SA adsorption from $n$-heptane was slightly stronger ($\sim 5 \text{ kJ mol}^{-1}$) on hydroxylated Fe$_3$O$_4$(001) than bare Fe$_3$O$_4$(001), which was similar to the difference observed between Fe$_3$O$_4$(001) and Fe$_3$O$_4$(111).

For the adsorption calculations, a single OFM molecule was placed at the centre of the simulation cell, and either 700 $n$-hexadecane or 373 PAO molecules were randomly distributed between the slabs using the amorphous builder plugin in MAPS, see Figure 2. This gave a similar total number of C atoms ($\sim 11200$) for both systems. Two systems were used for the hard-disk area calculations (see Supporting Information). In the first set-up (dimerisation simulations), two OFM molecules were placed close to the surface of one of the hematite slabs, using the same system size and number of solvent molecules as in the adsorption calculations. In the second set-up (annealing simulations), one hematite slab was covered with a pre-formed monolayer of OFM molecules with different surface coverages.
The number of OFM molecules included varied from 106 to 198, yielding initial surface coverages, $\Gamma = 3.5-6.5$ molecules/nm$^2$.

![VMD visualization of a representative system used for the adsorption simulations (GMO in $n$-hexadecane) on $\alpha$-Fe$_2$O$_3$(100). OFM atoms shown as beads: O atoms in red, H atoms in white, C atoms from OFM in blue and Fe atoms in gray. Solvent molecules shown as lines, C atoms shown in brown, H atoms not shown. The dark blue box depicts the simulation cell, exhibiting periodic boundary conditions in the $x$ and $y$ directions (i.e. parallel to the surfaces).](image)

The base oil and OFM molecules were modeled using the all-atom long chain-optimised potential for liquid simulations (L-OPLS-AA) force field. This is an updated version of OPLS-AA in which some of the parameters for C and H atoms are adjusted to accurately
reproduce the experimental density and viscosity of long chain alkanes.\textsuperscript{64} Parameters for the other atoms remain unchanged from the original OPLS-AA force field.\textsuperscript{63} The Lennard-Jones parameters and partial charges for the Fe and O atoms in the $\alpha$-$\text{Fe}_2\text{O}_3$ surface were taken from Berro et al.\textsuperscript{65} Moreover, harmonic bonds with a force constant of 130 kcal mol$^{-1}$ Å$^{-1}$ were added between atoms within 0.3 nm of each other in their lattice positions.\textsuperscript{65} Bond lengths were taken from X-ray diffraction experiments by Blake et al.\textsuperscript{66} This combination of force fields has already been successfully used by Ewen et al.\textsuperscript{67} in nonequilibrium molecular dynamics (NEMD) simulations of OFM friction.

It is noteworthy that recent DFT calculations have shown that strong molecule-surface interactions can occur for OFMs on iron oxide surfaces in the absence of solvent molecules.\textsuperscript{29} DFT calculations have also been used to parametrize classical force fields to more accurately represent amide-surface chemisorption.\textsuperscript{68} However, this task has not yet been executed for the OFM molecules of interest in this study.

**Simulation procedure**

MD simulations were performed using the Large-scale Atomic/Molecular Massively Parallel Simulator (LAMMPS) software.\textsuperscript{69} The velocity-Verlet algorithm\textsuperscript{70} was used to integrate the MD equations of motion. A time step of 1 fs was used for all of the simulations and all bonds involving H atoms were fixed using the SHAKE algorithm.\textsuperscript{71} Long-range electrostatic interactions were calculated using the slab implementation of the particle-particle, particle-mesh (PPPM) algorithm with a relative force accuracy of $10^{-4}$.\textsuperscript{72} The PMFs were extracted using the ABF method\textsuperscript{42,43} with the collective variables (colvars) module implemented in LAMMPS.\textsuperscript{73}

Before starting the PMF production runs, the systems were energy minimized using the conjugate gradient algorithm, with all of the hematite atoms being fixed in place. Subsequently, the system was equilibrated at 300 K and 101 kPa for 2 ns. The outer layer of atoms in the bottom wall of hematite were fixed in all directions, whereas the outer layer of the top
wall was fixed only in the $x$- and $y$-directions, but allowed to move in the $z$-direction during the equilibration phase. These outer layers consisted of all of the Fe and O atoms that were 0.1 nm or less from the bottom and the top of the simulation box respectively (440 atoms in both cases).

The pressure was controlled during the equilibration phase by applying a constant force to the outer layer of atoms in the top wall. After equilibration, the outer layers of both hematite slabs were fixed in place for the subsequent PMF production simulations. At the start of the equilibration phase, the velocities of the solvent, OFM and non-fixed hematite atoms were initialized using with a Gaussian distribution of velocities corresponding to a temperature of 300 K. The temperature was then controlled using the Nosé-Hoover\textsuperscript{74,75} thermostat with a time relaxation constant of 0.1 ps. The thermostat was applied to all of the non-fixed atoms.

The PMF was calculated using the ABF method, an importance sampling algorithm developed by Pohorille and Darve.\textsuperscript{42,76} The algorithm is based on the computation of the mean force along a certain collective variable (colvar),\textsuperscript{73} which is divided into small bins. The mean force is then cancelled out by an equal and opposite biasing force. This allows the system to move freely along that colvar, overcoming energy barriers and recovering ergodicity. The mean force profiles converge once the relevant range of the colvar has been explored and sufficient samples have been accumulated within each bin. The convergence of this method is subservient to the efficient exploration of the phase space orthogonal to the chosen colvar. In our case, this corresponds to the different orientations and conformations of the OFM molecules for each distance to the surface. Finally, the PMF is calculated from the mean force profile, since the average force exerted by the system onto the colvar is the gradient of the free energy profile along the colvar with a sign change:

$$\frac{dA}{d\xi}(\xi_0) = -\langle F(\xi_0) \rangle,$$  \hspace{1cm} (1)

where $A$ is the free energy profile as a function of the colvar $\xi$ and $F$ is the force exerted by the system onto the colvar.\textsuperscript{42,76} In principle, $\xi$ is a continuous variable and equation 1 applies
for every value $\xi_0$. In practice, the colvar is divided into small bins of width $\Delta \xi$ and samples are acquired for each bin during the simulation to compute the mean force in the interval $(\xi_0, \xi_0 + \Delta \xi)$. Then, the mean value of those samples for each bin is used to compute the gradient and, through integration, the PMF. The review by Comer et al.\textsuperscript{77} provides a more detailed description of the ABF method, together with recommendations and good practices.

**Adsorption PMF**

To estimate the PMF that captures the adsorption process, the colvar\textsuperscript{73} was defined as the projection onto the z-axis of the distance between the center of mass (COM) of the OFM polar headgroup and the COM of the topmost layer of atoms of the bottom hematite surface. To accelerate convergence, the relevant range of the collective variable can be divided into several regions or ”windows” to which the ABF algorithm is applied simultaneously. This approach is known as the stratification scheme.\textsuperscript{78} The length of these windows should not be too small, as restricting the range of the colvar too much in a single simulation may lead to a slowdown in convergence or even a loss of ergodicity.\textsuperscript{77} In other words, the simulation should visit the same configurations and in the same proportion as it would do using a much larger window, so that the artificial limit to the colvar explored in each window does not introduce a bias in the sampling.

In this case, the window closest to the surface should be large enough so as to avoid that the tail remains intermingled in the first maximum of the solvent layering while the colvar is being explored. Otherwise all the samples acquired for the first window would correspond to the same tail position, biasing the exploration of the phase space. Due to these considerations, the calculations for SA and GMS were run with three windows: from 0.18/0.23 to 2.0 nm, 2.0 to 3.0 nm and 3.0 to 4.0 nm, since their saturated linear tailgroups are more extended than the kinked Z-unsaturated tailgroups in GMO. Therefore, four windows were used for GMO: 0.23 to 1.0 nm, 1.0 to 2.0 nm, 2.0 to 3.0 nm and 3.0 to 4.0 nm. The initial
configurations for each window were obtained using steered molecular dynamics (SMD), as implemented in the LAMMPS colvars package. The force constant of the applied harmonic biasing potential was 10 kcal mol$^{-1}$Å$^{-2}$, and the equilibrium distance was moved at a velocity of 4 m s$^{-1}$ from the initial position of the molecule towards the surface. One initial condition was saved for each window and was equilibrated for 2 ns before applying the ABF algorithm to each window. The chosen width of the colvar bins was $\Delta \xi = 0.005$ nm.

The ABF algorithm was applied to each window until convergence was reached. As for other importance sampling techniques, the assessment of convergence is not trivial. Several necessary conditions for convergence were monitored to determine a reasonable trade-off between computational cost and precision. These included the uniform increase of the samples across bins, the absence of further changes to the PMF, and the continuity of the mean force between different windows (see the Supporting Information). The mean force converged more slowly in the windows that were close to the surface due to the existence of several orientations and conformations that correspond to different values of the mean force. The total simulation times required to obtain converged PMFs varied between 165 ns and 750 ns for windows adjacent to the surface and between 75 ns and 150 ns for the others. The calculations using $n$-hexadecane converged more quickly than those with PAO4. This is consistent with the observations of Jaishankar et al., who found that calculations converged more quickly for $n$-heptane than $n$-hexadecane, due to shorter relaxation times of the former.

**Hard-disk area**

To obtain a simulated adsorption isotherm using MTT, it is also necessary to estimate the area that each adsorbed molecule occupies on the surface. As explained by Nikas et al., several difficulties arise concerning the estimation of the hard-disk area of each molecule on the surface due to steric effects. For instance, this area depends on the molecular conforma-
tions accessible to the adsorbed surfactant, which will depend on the surface coverage. As a result, the effective cross-sectional area will be coverage-dependent. In practice, it is usually sufficient to estimate the hard-disk area at high concentration and thus high coverage, since this is where the steric repulsion of adsorbed molecules limits further increases to the surface coverage.

OFMs are known to form vertically-orientated SAMs on iron oxide surfaces. Therefore, the surfactant headgroup size will mostly determine the hard-disk area of the adsorbed molecule. Two approaches were used to estimate the hard-disk area occupied by the OFMs on the hematite surface. In the first approach, the hard-disk radius was estimated using the ABF method to calculate the PMF for the lateral interactions between OFMs. Two OFMs were adsorbed on the hematite surface and the projection onto the $x$-$y$ plane of the distance between the headgroups was used as colvar. Subsequently, the cross-sectional area occupied by each OFM on the hematite surface was estimated using the position of the PMF minimum, following the approach used by Jaishankar et al. for SA at the $n$-heptane-iron oxide interface and Sresht et al. for polyethylene glycol at the air-water interface. The results obtained using this first method can be found in the Supporting Information. In the second approach, annealing simulations of preformed OFM monolayers were performed. A similar approach to that described by Zhang et al. was used here. Different numbers of OFM molecules were placed close to the surface to form monolayers with a range of initial surface coverages. A subsequent equilibration of 2 ns allowed the molecules to adsorb onto the surface. Then the temperature was increased instantaneously from 300 K to 400 K and the system was equilibrated for 16 ns, which allowed the unbound molecules to desorb and escape the surfactant monolayer. Subsequently the temperature was reduced back to 300 K for another 2 ns. The final coverage was then estimated by determining the number of adsorbed and desorbed molecules, based on the distances of the hydroxyl oxygen atom of the headgroup to the hematite surface. The threshold distance (0.4 nm) was chosen such that for low concentrations, the number of adsorbed molecules at the end was equal to the number
of initially placed molecules. Varying the number of the initially placed molecules allowed us to determine the maximum coverage and thus the cross-sectional area of each molecule. No base oil molecules were included in the annealing simulations since tests showed that their presence did not significantly affect the results. The same observation was made in previous MD simulations of SA monolayer stability on iron oxide surfaces.\textsuperscript{82}

**MTT model**

Once the adsorption PMF and the area per molecule were determined, the MTT model for surfactant adsorption developed by Nikas et al.\textsuperscript{35} was used to construct adsorption isotherms for the OFMs on hematite from PAO4. The model starts with a surface equation of state based on the following assumptions: (1) single monolayer adsorption, (2) the surfactants in the monolayer are modelled as hard-disks of radius $r$ with short-ranged attractive interactions, (3) exclusively non-ionic surfactants are considered, and (4) the headgroups are located on the Gibbs dividing surface (minimum of the PMF) while the tailgroups are pointing towards the liquid phase. From this equation of state, the surface chemical potential, $\mu^\sigma$, is derived.

The surfactants in the bulk are considered to be below the critical micelle concentration (CMC),\textsuperscript{83} thus the bulk chemical potential of the surfactants $\mu^b$ is that of an ideal solution with no contributions coming from the interactions between surfactants. Imposing thermodynamic equilibrium $\mu^\sigma = \mu^b$, the surface coverage $\Gamma$ and the bulk molar fraction $X$ are related in a transcendental equation that can be solved using the bisection or Newton method. For more details, see the original publication\textsuperscript{35} or the derivation from Sresht et al.\textsuperscript{54} The final result reads:

$$\ln X = \frac{\Delta \mu^0}{k_B T} + \ln \left( \frac{\Gamma a}{1 - \Gamma a} \right) + \Gamma a \frac{3 - 2 \Gamma a}{(1 - \Gamma a)^2} + 2B \Gamma$$

(2)
where $X$ is the bulk mole fraction of monomer surfactant (assumed here to be the total concentration of surfactant without distinguishing between monomers, dimers, or micelles), $\Gamma$ is the surface coverage, $a$ is the hard-disk area, and $\Delta \mu^0 \equiv \mu^\sigma - \mu^b$ is the free energy of adsorption at infinite dilution, i.e. with only one molecule. And $B$ is the coefficient associated to attractive pairwise interactions between surfactant molecules, which won’t be taken into account in our computational results. $\Delta \mu^0$ is calculated from the transfer free energy or difference in free energy between the bulk region and the adsorbed state. If the interface region is broad and not well defined, then the minimum of the PMF can be used directly for $\Delta \mu^0$, considering that the surfactants are located at the Gibbs dividing surface, as proposed by Sresht et al.\textsuperscript{54} Here, however, we follow the approach used by Jaishankar et al.,\textsuperscript{34} where the surfactants are not considered to be in a 2D plane above the hematite surface, but inside a 3D interface, where the different adsorbed states are taken into account. In this case, the expression for $\Delta \mu^0$ reads:

$$
\Delta \mu^0 = -k_B T \ln \left[ \frac{aL}{v_s(z_1 - z_0)} \int_{z_0}^{z_1} e^{-\Delta G(z) / k_B T} dz \right]
$$

where $L = z_2 - z_1$, $z_2$ marks the point at which the PMF deviates from the bulk value, $z_1$ is the upper bound of the free energy well and $z_0$ is the lower bound of the free energy well. $v_s$ is the molecular volume of the solvent and $\Delta G(z)$ is the adsorption PMF.

Jaishankar et al.\textsuperscript{34} considered an additional term that takes into account the entropic penalty of increasing the coverage as the chains are not free to rotate. This contribution has been analytically estimated and increases the potential energy of the surfactants at the surface\textsuperscript{84} and is included here as well in the MTT model:

$$
\ln X = \frac{\Delta \mu^0}{k_B T} + \ln \left( \frac{\Gamma a}{1 - \Gamma a} \right) + \Gamma a \frac{3 - 2\Gamma a}{(1 - \Gamma a)^2} + 2B\Gamma + \frac{\pi^2}{8} N_c L_c^3 \Gamma^2.
$$

Here $N_c = (n_c + 1)/3.6$ is what Nagarajan et al.\textsuperscript{84} consider as an orientationally free segment.
formed by 3.6 methylene groups, with \( n_c - 1 \) being the number of methylene groups \( \text{CH}_2 \) in the tail and the final methyl group counting as two methylenes. Each group of 3.6 methylenes is considered to be a free-orientation segment of length \( L_c = 0.46 \). For a more detailed explanation of the MTT model, see Refs. 34–36.

In hydrocarbon solvents, carboxylic acids such as SA\textsuperscript{85} dimerise, while glycerol esters, such as GMO\textsuperscript{86} and GMS\textsuperscript{87} can form reverse micelles. Previous MD simulations have shown that SA monomers and dimers exhibit equal affinity for iron oxide surfaces.\textsuperscript{34} As micellation is not taken into account here, the results will be representative of systems below the CMC of the surfactants in the hydrocarbon solvent.\textsuperscript{83} To our knowledge, the CMC has not been quantified for the systems studied here. Future studies could use the same techniques adopted here to study the adsorption of OFM reverse micelles on solid surfaces from hydrocarbon solvents.\textsuperscript{56,88}

**Experiments**

To validate the calculations of the adsorption free energy, friction was used as a proxy for surface coverage using the Jahanmir and Beltzer model.\textsuperscript{20} The obtained surface coverage can then be related to the adsorption energy. Friction measurements were performed using a high-frequency reciprocating rig (HFRR) from PCS Instruments (Acton, UK).\textsuperscript{89} The standard specimens were used, namely a 6 mm diameter ball (upper) loaded on a 10 mm diameter disk (lower). Both of the specimens were made from AISI E-52100 steel. A 2 N load was applied to the upper specimen, which followed a reciprocating movement of 10 Hz with a stroke length of 0.1 mm, giving a maximum velocity of 2 mm s\textsuperscript{-1}. The corresponding maximum Hertzian pressure was 0.8 GPa, whereas the mean was 0.5 GPa. High pressure and low speed were used to ensure that boundary lubrication conditions were maintained over most of the stroke. This is important since it is here that the friction behaviour depends strongly on the OFM concentration and consequently the coverage.\textsuperscript{12,24} Ambient temperature (300 K) was used and each test was conducted for 1 h to reach steady state friction. The friction coefficient
was measured for pure base oil and for OFM concentrations between $10^{-6}$ M and 0.02 M, for a total number of 7 HFRR tests. At the beginning of each test and once the contact was flooded, the ball and disc were held stationary and no pressure was applied during 5 minutes to allow for the formation of the OFM film, which is similar to the procedure used by Fry et al.\textsuperscript{25} The two specimens were cleaned prior to each test with acetone and toluene. The base oil used was PAO4 from ExxonMobil Chemical (Houston, USA) and the OFMs were GMO and GMS supplied by Sigma-Aldrich (Madrid, Spain) with a high purity grade (> 99%). The steady-state friction coefficient was determined averaging over the last 10 min of the experiment. Three repetitions were used for each OFM and concentration to reduce the noise in the HFRR data.

To validate the calculation of the adsorption energy at infinite dilution, the Jahanmir and Beltzer model was used.\textsuperscript{20} This model relates the measured friction from the HFRR experiments with the fractional coverage. The model assumes that the contact is in boundary lubrication conditions (so friction is given by asperity contact and not the viscosity of the base oil) and that the friction coefficient is a linear function of the fractional coverage:

$$f = f_b(1 - \theta) + f_a \theta,$$  \hspace{1cm} (5)

where $f$ is the measured friction coefficient, $\theta \equiv \Gamma/\Gamma_{max}$ is the fractional coverage, $f_a$ is the friction coefficient at maximum additive coverage and $f_b$ is the friction coefficient with no additive present. Although we considered $\theta = 1$ to determine $f_a$ from the HFRR measurements, complete coverage is not achieved in practice according to our computational adsorption isotherm results. This observation is in agreement with previous MTT-MD studies.\textsuperscript{34}

Once $f_a$ and $f_b$ are determined, the Jahanmir and Beltzer model can be applied to the friction data to obtain a relation between the OFM bulk concentration and the fractional surface coverage, to which an adsorption isotherm model can be fitted. Jaishankar et al.\textsuperscript{34} used the Langmuir model for this purpose. However, according to Jahanmir and Beltzer,\textsuperscript{20}
the Temkin isotherm model is better suited for these OFMs than the Langmuir isotherm model since it is able to account for inhomogeneous surfaces and lateral interactions. In the Temkin isotherm model the adsorption energy increases linearly with coverage (making the adsorption process less favorable):

$$\Delta G_{ads} = \Delta G_0 + \alpha \theta,$$  \hspace{1cm} (6)

where $\Delta G_0$ is the adsorption energy at $\theta = 0$, $\alpha$ is a positive constant depending on the interactions of the molecules on the surface, and $\theta$ refers to the fractional coverage. As explained by Jahanmir and Beltzer\textsuperscript{20} the complete form of the Temkin adsorption isotherm is complex and trying to fit several parameters to the HFRR data would result in several possible combinations of $\alpha$ and $\Delta G_0$. However, at intermediate fractional coverages ($0.8 \geq \theta \geq 0.2$) the isotherm takes a simpler form:

$$\theta = (RT/\alpha) \ln(X/K_0) , \ K_0 = e^{\Delta G_0/RT}.$$  \hspace{1cm} (7)

Representing the fractional coverage as a function of $\ln(X)$, $\alpha$ can be found from the slope $m$, whereas $\Delta G_0$ is found from the intercept $n$ as shown in equations 8 and 9.

$$\alpha = \frac{RT}{m} \hspace{1cm} (8)$$
$$\Delta G_{ads} = -\alpha n \hspace{1cm} (9)$$
Results and discussion

Adsorption

For both solvents, the bulk density in the centre of the film was in good agreement with experimental values. We obtained approximately $742 \pm 5$ g/L for $n$-hexadecane and $778 \pm 20$ g/L for PAO4, which is below the experimental values by 2% and 4%, respectively. At the end of the equilibration phase, the separation between the two walls was 12 nm for $n$-hexadecane and 11 nm for PAO4. Pronounced layering of the solvent molecules can be observed close to the surfaces in the mass density profiles (Fig. 3). This layering is caused by van der Waals and electrostatic interactions between the base oil molecules and the hematite surfaces. The layering next to the wall is stronger for linear $n$-hexadecane than branched PAO4, which agrees with observations from previous MD simulations. The longer period of the density oscillations in Fig. 3 shows that each molecular layer is slightly thicker for PAO4 than $n$-hexadecane, which is a result of its branched structure.

Figure 3: Mass density profiles for the PAO4 and $n$-hexadecane solvent molecules on $\alpha$-Fe$_2$O$_3$(100) at the end of the equilibration phase (left) and corresponding image from VMD for $n$-hexadecane (right). Atom colors the same as in Fig. 2.

The PMF is calculated by integrating the negative of the mean force and taking its value in the bulk region as reference (i.e. zero). The bulk solvent is defined as the region where the
interactions of both the OFM and the solvent with the surface are negligible (3-4 nm from the surface). For the molecules considered here, the mean force $F(\xi)$ close to the surface and thus the changes of the PMF in that region are mainly determined by the interaction of the headgroup with the surface, whereas the mean force in the transition from the bulk to the surface proximity is mainly determined by the tailgroup-solvent interactions, since the interactions of the polar headgroups with the surface are still weak. As we have chosen as reference the free energy within the bulk to be zero, the depth of the minimum will be determined by the interactions of the OFM molecule with both the surface and the solvent. For all the OFMs studied here, the most stable conformation (PMF minima) involves the surfactant molecule lying parallel to the hematite surface. This observation is consistent with previous DFT calculations that showed that, at low surface coverage, such conformations maximise the strength of the OFM-surface interactions. The surfactant molecules only lie flat on the surface for GMS, while for SA and GMO there is a trapped solvent layer separating the tailgroup and the surface. This is attributed to the additional flexibility in GMS from the ester headgroup and the fully saturated tailgroup.

All calculated adsorption PMFs (Figs. 4a, 4b, 4c) exhibit at least one minimum close to the surface. The PMF profiles also show energy barriers that correspond to the interaction of the tail with the layering of the solvent molecules close to the surface. There are only minor differences between PMFs from $n$-hexadecane and PAO4. The simulation time required to reach convergence of the PMF profile is much greater for PAO4 than $n$-hexadecane, due to the longer relaxation time of the solvent molecules. The results for SA are in reasonable agreement with the ones from Jaishankar et al. for the adsorption of SA from $n$-hexadecane onto Fe$_3$O$_4$ (111). As the SA molecule approaches the $\alpha$-Fe$_2$O$_3$ (100) surface, the free energy oscillates with a period ($\sim$0.5 nm) that corresponds to the thickness of the solvent layer (Fig. 3). At OFM-surface distances of around 2 nm, 1.5 nm, 1.0 nm, and 0.5 nm, there are free energy barriers that correspond to the OFM moving through the
Figure 4: Adsorption PMF for the SA (a), GMO (b), and GMS (c) onto the $\alpha$-Fe$_2$O$_3$(100) surface obtained using the ABF algorithm.\textsuperscript{42,76} PAO4 and $n$-hexadecane were used for SA and GMO while only PAO4 was used for GMS. VMD snapshots\textsuperscript{61} show the molecular configurations at different points during the adsorption process from PAO4 (solvent molecules not shown for clarity). Atom colors the same as in Fig. 2.
solvent layers. The tailgroup is preferentially located in the maxima of the solvent mass
density profile, and a small energy barrier has to be crossed for the tailgroup to move be-
tween these maxima. The height of these free energy barriers increases as the OFM gets
closer to the surface, because the solvent layering is more pronounced. The final barrier,
corresponding to moving through the solvent layer closest to the surface, is approximately
10 kJ mol\(^{-1}\) for SA in PAO4 and approximately 5 kJ mol\(^{-1}\) for the other OFM-solvent
combinations. For SA, the barrier height is larger for PAO4 than n-hexadecane, despite the
stronger layering for the latter (Fig. 3). This implies that the solvent layers are more difficult
for SA to penetrate for branched alkanes solvents than linear ones. In practical terms, this
means that SA adsorption will be slower from PAO4 than n-hexadecane at 300 K.\(^{34}\) The
improved ability for the saturated linear OFM molecule to penetrate the n-alkane of similar
chain length could be the molecular mechanism of the chain matching phenomena observed
experimentally.\(^{91}\) For GMO, the free energy barrier heights are similar for both solvents. In
between these barriers, there are local minima in the free energy profiles corresponding to
outer-sphere complexes. These have also been observed in previous MD simulations of oxalic
acid adsorption on rutile from water\(^ {92}\) and napthenic acid adsorption on calcite from water.\(^ {93}\)

For SA, the headgroup was defined as the carboxyl group \(-\text{COOH}\). Only one inner-
sphere adsorption mode was observed in the PMF at 0.23 nm above the surface, see Fig. 4a.
The \(-\text{OH}\) group of the carboxylic acid was oriented towards either one oxygen atom or a
bridge between two oxygen atoms of the hematite surface with a OH···O distance of \(\sim 0.16\)
nm, while the \(=\text{O}\) atom of the carboxylic group was oriented towards a Fe atom of the surface
with a \(=\text{O}···\text{Fe}\) distance of \(\sim 0.25\) nm. This configuration gave an adsorption energy of -27.8
kJ mol\(^{-1}\) for SA on \(\alpha\)-Fe\(_2\)O\(_3\)(100) from PAO4. The SA adsorption is slightly stronger (-32.4
kJ mol\(^{-1}\)) from n-hexadecane, because the tailgroup is only able to lie flat to the surface in
the latter case. This value is in good agreement with that obtained by Jaishankar et al.\(^ {34}\)
for SA adsorption from n-hexadecane onto Fe\(_3\)O\(_4\)(111) (-29.1 kJ mol\(^{-1}\)).
For both GMO and GMS, the headgroup was defined as the carboxyl and glycerol groups \(-\text{COO} - \text{C}_3\text{H}_5\text{O}_2\text{H}_2\). Both of these surfactants have several polar groups capable of interacting with the oxygen or iron atoms in the surface. They are also large, flexible molecules that can adopt many different possible conformations at the solid-liquid interface. Consequently, two distinct inner-sphere adsorption modes are present for GMS and GMO, as can be seen from the corresponding PMFs. The different situations and adsorption modes depicted in Fig. 4b and 4c are associated to changes in the derivative of the PMF, i.e. changes in the mean force experienced by the colvar at that distance.\(^{73}\) The barriers between 0.6 nm and the bulk are once again due to the interaction between the tail and the solvent molecules.

At \(\xi \sim 0.5\) nm, the first of the two hydroxyl groups interacts with the oxygen atoms of the hematite surface with a \(\text{OH} \cdots \text{O}\) distance of \(\sim 0.16\) nm. At \(\xi \sim 0.4\) nm, both hydroxyl groups are interacting with the oxygen atoms of the surface, and the PMF exhibits a local minimum. \(\xi \sim 0.3\) nm corresponds to the global adsorption minimum, where the \(=\text{O}\) atom of the headgroup also interacts with the surface with a \(=\text{O} \cdots \text{Fe}\) distance of \(\sim 0.26\) nm. There is a small energy barrier between the two adsorption modes related to the energy required to further approach the headgroup center of mass to the hematite surface. In this transition, the tailgroup goes from being located at the second maximum of the solvent mass density profile at 0.7-0.8 nm to being partially or completely located at the first maximum at 0.3 nm from the surface. In the case of GMS, the tailgroup at the global minimum lies flat to the surface, whereas for GMO, the half of the C atoms lie flat to the surface, with the remaining half pointing into the solvent layer due to the presence of the \(Z\)-alkene group.

The adsorption of the alkene group provides a significant increase in adsorption strength for GMO (\(-41.8\) kJ mol\(^{-1}\)) than GMS (\(-34.2\) kJ mol\(^{-1}\)) on \(\alpha\)-\(\text{Fe}_2\text{O}_3\)(100) from PAO4. This observation is in agreement with previous experiments of fatty acids on steel surfaces\(^{94}\) as well as MD simulations of fatty acids on DLC surfaces.\(^{95}\) The adsorption energy for GMO is slightly larger from branched PAO4 than linear \(n\)-hexadecane (\(-39.0\) kJ mol\(^{-1}\)), which is
opposite to the trend observed for SA.

**Hard-disk area**

The hard-disk area was determined using annealing MD simulations. Various monolayers of different coverages were created for the three OFMs, namely from 3.5 molecules/nm$^2$ to 6.5 molecules/nm$^2$ for SA and from 3.5 molecules/nm$^2$ to 5 molecules/nm$^2$ for GMO and GMS. The final coverage, or alternatively the final area per molecule, can be obtained by counting the number of molecules adsorbed at the end of the simulation. The threshold distance to consider a molecule as desorbed was 0.4 nm, measured from the oxygen of the hydroxyl group to the COM of the first layer of surface atoms. Increasing the threshold distance above 0.35 nm did not change the results significantly, whereas decreasing it below this value underestimated the number of adsorbed molecules at low coverages, where all the molecules are assumed to be adsorbed, as shown in the Supporting Information.

Fig. 5 shows that the number of adsorbed molecules at the end of the simulation is equal to the number of molecules initially placed on the surface up to a certain value of the initial coverage. Beyond this critical value, a plateau is reached and the final coverage does not increase further regardless of the initial coverage. This maximum number of adsorbed molecules is 170 for SA, 121 for GMO and 120 for GMS. Consequently, the area per molecule at maximum coverage is 0.1796 nm$^2$ for SA, 0.2523 nm$^2$ for GMO and 0.2544 nm$^2$ for GMS.

It is important to note that these results only give information regarding the minimum area due to steric effects that can be occupied by molecules adsorbed on the surface in a thermodynamically stable monolayer. It is not necessarily a realistic depiction of the maximum coverage that can be kinetically accessed in realistic timescales, since the OFM monolayers are preformed. Jaishankar et al.\textsuperscript{34} suggested that the maximum accessible coverage can be illustrated both using the MTT and the random sequential adsorption (RSA) model.\textsuperscript{96} The RSA model describes the irreversible adsorption of hard-disks on a uniform surface, which predicts a maximum coverage below the closed-packed maximum one, in which all of the
Figure 5: Comparison of the initial and final surface coverage for the three OFMs following the annealing simulations on $\alpha$-$\text{Fe}_2\text{O}_3(100)$ using a threshold OFM-surface distance of 0.4 nm. Colored dashed lines are guides for the eye. VMD snapshots show the molecular configurations after the annealing process. Atom colors the same as in Fig. 2.

surface would be covered. This is a result of irreversible adsorption and absence of restructuring between adsorption events, creating a jamming limit that was in agreement with the results by Jaishankar et al.\textsuperscript{34}

In reality the value of the effective hard-disk area occupied by each molecule on the surface depends on the coverage, in particular if the molecule has access to several adsorption modes with different conformations. The hard-disk area parameter used in the MTT model should correspond to the case of high coverage, where steric effects become important as explained by Nikas et al.\textsuperscript{35} From the literature, one can find several values for different OFMs. One should distinguish between area per molecule (total area divided by the number of molecules) and effective hard-disk area (due to steric effects, serving as input for the MTT model). The former will always be greater than or equal than the later. Davidson et al.\textsuperscript{55} calculated the area per molecule for 1-monoctyl glycerol at high coverage with MD, finding an average
area per molecule of 0.225 nm\(^2\). Kawaguchi et al.\(^{97}\) studied the surface properties of mono-, di- and triglycerol monostearate monolayers at the air-water interface, finding a limiting area per GMS molecule of 0.26 nm\(^2\). They explain that the area per molecule of GMS is larger than the area per molecule of SA (0.2 nm\(^2\)\(^{98}\) or 0.22 nm\(^2\)\(^{99}\)) due to the larger headgroup size.

The translation from area per molecule to hard-disk area requires the disk packing efficiency to be estimated. The packing efficiency would be 0.906 for hexagonal packing (hp) or 0.785 for square packing (sp). The packing efficiency of the OFMs is likely to be tighter if there are stronger interactions between them (e.g. hydrogen bonds or van der Waals interactions between tailgroups).

Table 1: Values of the hard-disk area obtained from the annealing MD simulations on \(\alpha{-}\text{Fe}_2\text{O}_3\)(100). \(A_{\text{sim}}\) refers to the area per molecule at maximum coverage, \(A_{\text{hp}}\) refers to the effective hard-disk area per molecule assuming hexagonal packing.

<table>
<thead>
<tr>
<th>OFM</th>
<th>(A_{\text{sim}}) [nm(^2)]</th>
<th>(A_{\text{hp}}) [nm(^2)]</th>
</tr>
</thead>
<tbody>
<tr>
<td>SA</td>
<td>0.180</td>
<td>0.163</td>
</tr>
<tr>
<td>GMO</td>
<td>0.252</td>
<td>0.228</td>
</tr>
<tr>
<td>GMS</td>
<td>0.254</td>
<td>0.230</td>
</tr>
</tbody>
</table>

A summary of all the values for the area per molecule and the effective hard-disk area per molecule can be found in Table 1. As hexagonal packing is the most efficient way to pack disks on a surface, it is reasonable to assume at most a ratio of 0.906 between the raw area per molecule and the hard-disk area. In addition, hexagonal packing can be observed at the end of the annealing MD simulations (see Supporting information). Consequently, the results of the annealing simulations assuming hexagonal packing (\(A_{\text{hp}}\) in Table 1) are then used as input for the MTT model. The values in Table 1 are in good agreement with those available in the literature for similar molecules.\(^{55,97-99}\) Interestingly, the estimated hard-disk area for both GMO and GMS is very similar. This is in contrast with studies from Wood et al.,\(^{21}\) who measured a lower surface coverage for unsaturated OFMs with respect to their saturated counterparts. They explained these findings in terms of a reduced surface packing efficiency due to the presence of the kinked Z-alkene. However, in the annealing
simulations presented here no significant difference is observed between both, suggesting that the observed differences in surface coverage by Wood et al.\textsuperscript{21} are instead due to kinetic effects due to steric hindrance, rather than surface packing efficiency.

**Adsorption isotherms from MTT**

The main input parameters for the MTT model are the adsorption energy, the hard-disk area, and the number of orientationally-free carbon atoms $n_c$ in the tail. The value of the hard-disk area for each OFM is taken as $A_{hp}$ from Table 1. The adsorption energy is calculated for each OFM using equation 3 and the PMF, with the solvent volume $v_s$ for PAO4 being 0.8525 nm$^3$ and 0.488 nm$^3$ for $n$-hexadecane. For SA the adsorption energy was calculated using $z_0 = 0.18$ nm, $z_1 = 0.365$ nm and $z_2 = 1.8$ nm for PAO4 and yielded $\Delta \mu_{sa}^0 = -27.8$ kJ mol$^{-1}$. While for the $n$-hexadecane results $z_0 = 0.18$ nm, $z_1 = 0.4$ nm and $z_2 = 1.8$ which yielded $\Delta \mu_{sa}^0 = -32.4$ kJ mol$^{-1}$. For the adsorption of GMO from PAO4, we used $z_0 = 0.22$ nm, $z_1 = 0.59$ nm and $z_2 = 1.4$ nm and obtained an adsorption energy $\Delta \mu_{gmo}^0 = -41.8$ kJ mol$^{-1}$. Whereas for the corresponding $n$-hexadecane results $z_0 = 0.25$ nm, $z_1 = 0.57$ nm and $z_2 = 1.63$ which yielded $\Delta \mu_{gmo}^0 = -39.0$ kJ mol$^{-1}$. For GMS the same values of $z_0$, $z_1$ and $z_2$ are used as in GMO-PAO4 and the adsorption energy is $\Delta \mu_{gms}^0 = -34.2$ kJ mol$^{-1}$. The value of $n_c$ was taken as 17 for SA and GMS and 15 for GMO, since the C atoms involved in the alkene group are not counted.

Previous MD simulations have shown that the adsorption energy can be influenced by the type of iron oxide, the crystallographic orientation, surface hydroxylation, and the solvent type.\textsuperscript{34} The adsorption isotherm for SA calculated with MTT (Eq. (4))\textsuperscript{35} with parameters obtained from the MD simulations\textsuperscript{34,36} on $\alpha$-Fe$_2$O$_3$(100) is compared to data from previous iron oxide powder adsorption experiments in Fig. 6. The adsorption isotherm without accounting for lateral interactions between adsorbed surfactants ($B = 0$), a simplification suggested by Jaishankar et al.,\textsuperscript{34} is shown as a solid line. Using this approach, the MTT results are in good agreement with the experimental results reported by Jaishankar et al.\textsuperscript{34}
onto Fe$_3$O$_4$ from $n$-hexadecane. However, the surface coverage at intermediate concentrations is underestimated compared to the experimental results reported by Wood et al.$^{21}$ onto $\alpha$-Fe$_2$O$_3$ from $n$-dodecane and Studt$^{100}$ onto iron from $n$-hexadecane. The calculated adsorption free energy from the PMF of SA on $\alpha$-Fe$_2$O$_3$(100) from $n$-hexadecane (-32.4 kJ mol$^{-1}$) agrees well with previous estimates (-29.1 kJ mol$^{-1}$) on Fe$_3$O$_4$(111) from $n$-hexadecane.$^{34}$ Moreover, the hard-disk area (Table 1) is also consistent with previous estimates of the molecular area of SA.$^{98,99}$

Consequently, we suggest that the rate at which coverage increases with OFM concentration is underestimated at intermediate coverage compared to some experiments due to the absence of lateral interactions in this version of the MTT model. To account for these interactions, one should estimate the dispersion forces between adsorbed molecules on the surface and then calculate the $B$ parameter of Eq. (4), as explained in Refs. 35,36. One possibility would be to calculate the PMF between two adsorbed OFMs using the distance between headgroups as a colvar,$^{73}$ as suggested by Sresht et al.$^{36}$ These calculations are reported for our systems in the Supporting Information. Our results yield an overestimation of the $B$ parameter, which we attribute to issues with convergence of the PMF profiles. To obtain satisfactory convergence for these systems would require several $\mu$s of simulation time. However, to show that including these lateral interactions would approach our results to the experimental ones, we have used a non-linear least squares fitting of Eq. (4) to the experimental data, using the same values for the hard-disk area and the adsorption energy as before, while optimizing $B$ (see Fig. 7). The resulting MTT adsorption isotherms closely match the experimental data where $B = 0.114$ nm$^2$ for Jaishankar et al.,$^{34}$ $B = -0.453$ nm$^2$ for Wood et al.,$^{21}$ and $B = -0.847$ nm$^2$ for Studt.$^{100}$ A negative $B$ value is expected due to the attractive nature of the interactions. The values obtained to fit the experimental data are similar in magnitude to those obtained previously for C$_{10}$E$_4$ polyethylene glycol surfactants at the air-water interface.$^{36}$ The accurate determination of lateral interactions will be the focus of future MD simulations. This will likely require the use of a smaller hydrocarbon.
solvent to accelerate convergence and modification of the colvar to accelerate the exploration of the different degrees of freedom, such as the tailgroup positions with respect to the headgroups. The inclusion of $B$ in the MTT fitting is physically justified. Although the van der Waals interactions between the tailgroup are similar in strength to the tailgroup-solvent interactions, there will also be an entropic benefit of displacing adsorbed solvent molecules.\textsuperscript{32} Moreover, hydrogen bonding between the headgroups of adsorbed OFMs will provide an additional attractive component to the lateral interactions.\textsuperscript{67}

At relatively high concentrations (molar fraction $> 10^{-5}$-$10^{-3}$), the surface coverage of SA on hematite exceeds the jamming limit from the RSA model (green dashed line in Fig. 6)\textsuperscript{96} for both our MD data and two of the experimental data sets.\textsuperscript{21,100} This suggests that RSA model, as proposed by Jaishankar et al.,\textsuperscript{34} should be used with care for these systems.
Figure 6: Adsorption isotherm obtained using the MTT model (Eq. (4))\textsuperscript{35} for SA on $\alpha$-Fe$_2$O$_3$(100) from $n$-hexadecane, with the parameters calculated from the MD simulations (solid line). The results are compared to previously obtained experimental data (points) for SA adsorption on iron oxide powder from $n$-dodecane (Wood et al.\textsuperscript{21}) and $n$-hexadecane (Jaishankar et al.\textsuperscript{34} and Studt\textsuperscript{100}). Fits of the MTT model including lateral interactions ($B$ parameter in Eq. (4)) are also included for each experimental data-set for SA (dashed lines). The maximum coverage predicted by the RSA model (0.547/$a$)\textsuperscript{96} and the theoretical maximum coverage assuming hexagonal packing (0.906/$a$) are shown as green and blue dashed horizontal lines, respectively.

Fig. 7 shows adsorption isotherms obtained using the MTT model (Eq. (4))\textsuperscript{35} for the three OFMs on $\alpha$-Fe$_2$O$_3$(100) from PAO4, with the parameters calculated from the MD simulations, without lateral interactions ($B = 0$). Of the three OFMs studied, SA yields the highest maximum coverage (3.6 molecules/nm$^2$) due to its smaller headgroup size compared to GMO and GMS. This is in agreement with the maximum coverage of SA obtained experimentally by Wood et al.\textsuperscript{21} and Studt,\textsuperscript{100} validating the value used for the hard-disk area. Since both GMO and GMS have the same headgroup, they yield a very similar maximum coverage at high concentrations, where the surface coverage is mostly determined by the packing efficiency of the molecules. GMO yields the highest coverage of the three OFMs
at low concentration, which is due to its larger adsorption energy. GMS has a coverage between GMO and SA at low concentration because of its intermediate adsorption energy. At relatively high concentrations (molar fraction \( > 10^{-3} \)), the surface coverage of both GMS and GMO on hematite exceed the jamming limit from the RSA model (green dashed line in Fig. 7).^96

![Figure 7: Adsorption isotherms obtained using the MTT model (Eq. (4))^35 for the three OFMs on \( \alpha-\text{Fe}_2\text{O}_3(100) \) from PAO4, with the parameters calculated from the MD simulations, without lateral interactions \( (B = 0) \). The maximum coverage predicted by the RSA model \( (0.547/a)^96 \) and the theoretical maximum coverage assuming hexagonal packing \( (0.906/a) \) for GMS and GMO are plotted in green and blue dashed horizontal lines, respectively.

Experimentally, Wood et al.^21 observed a higher surface coverage for fatty acids with fewer \( Z \)-unsaturated groups (stearic > oleic > linoleic) at equal concentration. Campen et al.^24 showed that unsaturated OFMs gave higher friction that was independent of the sliding speed, which can be explained in terms of a low surface coverage. At equal surface coverage Ewen et al.^67 found similar friction for both saturated and unsaturated fatty acids.
in NEMD simulations, ascribing the differences found experimentally to the superior ability of saturated OFMs to form high-coverage monolayers with respect to their unsaturated counterparts. This difference between saturated and unsaturated OFMs is not reproduced with the present methodology. The form of the MTT model that has been used here only depends on the hard-disk area and the adsorption energy. The former is very similar for both GMS and GMO, since it is mainly determined by the headgroup, while the latter is more negative for GMO, due to its additional Z-alkene adsorption site.

The version of the MTT model used here does not appear to fully account for the differences in tailgroup saturation observed experimentally. The original MTT model included an extra term (with another parameter) that took into account the van der Waals interactions between surfactants on the surface. This parameter might be able to capture the differences in tailgroup saturation.

In order to differentiate GMO and GMS (and more generally saturated and unsaturated OFMs) in the current MD-MTT framework, a slightly different approach is required. We suggest that the PMF for OFM adsorption should be obtained on iron oxide surfaces partially covered by preformed monolayers with different surface coverages. This type of simulation has been performed previously for alkyl phosphonate monolayers on aluminium oxide surfaces\textsuperscript{101} This way, the steric barriers to the formation of high-coverage monolayers could be uncovered. This, however, is likely to require extensive computational resources (several µs of simulation time) and is beyond the scope of this current study.

**Friction experiments**

The experimental HFRR results are obtained calculating the steady-state friction coefficient from three repetitions of the HFRR experiments at each OFM concentration. The results for GMO and GMS are shown in Figure 8. The highest friction coefficient (0.263) is observed for the pure PAO4 base oil (not depicted in Figure 8). The lowest friction coefficients are observed for the highest OFM concentrations in PAO4, corresponding to 0.02 M for
GMO (0.0103 in molar fraction) or 0.01 M for GMS (0.005 molar fraction). These concentrations correspond to the high-coverage regime (∼3 molecules/nm²) according to the calculated MTT adsorption isotherms (see Fig. 7). This observation is in contrast to the results of Jaishankar et al. for SA. Our results suggest that, on steel surfaces, SA starts to reduce friction when the surface coverage reaches ∼2 molecules/nm² and minimum friction is reached at ∼4 molecules/nm², which is approaching a close-packed monolayer. This is the same surface coverage at which minimum friction was observed on silica surfaces for octadecylamine monolayers formed from n-hexadecane. GMO and GMS also begin to reduce friction at ∼2 molecules/nm², while minimum friction is achieved at a lower coverage of ∼3 molecules/nm².

Figure 8: Steady state friction coefficient from the HFRR experiments as a function of molar concentration for GMO (left) and GMS (right). The error bars represent the standard deviation of the data used to calculate the steady-state friction coefficient.

Subsequently, the Jahanmir and Beltzer model is applied to the friction coefficient results to obtain the coverage as a function of the logarithm of the molar concentration. Fitting the Temkin adsorption isotherm to intermediate concentrations for each OFM, we obtain an experimental estimate of the adsorption energy, see Figure 9. This approach yields an adsorption energy of -24.9 kJ mol⁻¹ for GMO (α = 13.1) and -33.2 kJ mol⁻¹ for GMS (α = 15.5). These values are the same order of magnitude as the computational results calculated through the adsorption PMF of each OFM, see Table 2.
Table 2: Adsorption energy of OFMs from PAO4 or n-hexadecane on ABF-MD ($\Delta \mu_{MD}$) and experimental values for GMO and GMS estimated using fits of the Temkin isotherm to HFRR friction data on steel in PAO4 ($\Delta \mu_{exp}$), * experimental data for SA taken from HFRR experiments in PAO6 from Ref. 34 using the Langmuir isotherm.

<table>
<thead>
<tr>
<th>OFM</th>
<th>SA</th>
<th>GMS</th>
<th>GMO</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\Delta \mu_{MD}$ [kJ mol$^{-1}$] (n-hexadecane)</td>
<td>-32.4</td>
<td>-</td>
<td>-39.0</td>
</tr>
<tr>
<td>$\Delta \mu_{MD}$ [kJ mol$^{-1}$] (PAO)</td>
<td>-27.8</td>
<td>-34.2</td>
<td>-41.8</td>
</tr>
<tr>
<td>$\Delta \mu_{exp}$ [kJ mol$^{-1}$] (PAO)</td>
<td>-26.4*</td>
<td>-33.2</td>
<td>-24.9</td>
</tr>
</tbody>
</table>

The agreement between the computational and experimental results is much better for GMS than for GMO and the $\alpha$ value found for both OFMs is the same. One would expect to find different $\alpha$ values if different chain packing efficiencies lead to reduced dispersion forces between adsorbed molecules in the case of GMO and thus lower coverage. This was the case with SA and iso-stearic acid reported by Jahanmir and Beltzer,$^{20}$ in which the presence of the methyl side group decreases packing efficiency and thus dispersion forces. They found a different slope $m$ and therefore a different $\alpha$ for both surfactants. The same authors, however, ascribed the differences in coverage between SA, elaidic acid and oleic acid to changes in chain dispersion interaction even though they found similar values of $\alpha$ for the three surfactants.

Overall, the differences between our computational adsorption energy (with a single molecule) and the experimental estimation fitting the Temkin adsorption isotherm for GMO can be understood through the steric hindrance for molecules adsorbing into partially formed monolayers. The kink in the $Z$-unsaturated tailgroup of GMO will lead to larger kinetic barriers for molecules to pass through the film and adsorb on the hematite surface. As well as affecting the kinetics of adsorption, this may also influence the thermodynamics. The larger steric barrier could result in an effective reduction in the adsorption energy for GMO (as predicted from the HFRR experiments), because the molecules become trapped in the monolayer before reaching the hematite surface.$^{102}$ Another factor worth considering is that
the PMF calculation is completely agnostic to the interaction between the OFM molecules in solution. It is possible that reverse micelle formation is more favourable for GMO than GMS in PAO4,\textsuperscript{86,87} thus reducing the availability of molecules to adsorb on the hematite surface.

![Figure 9: Fractional surface coverage as a function of the logarithm of the molar fraction with GMO (left) and GMS (right). Error bars are calculated from the propagation of the standard deviation of the friction measures.](image)

For the adsorption energy of SA, several references can be found in the literature. Jahanmir and Beltzer\textsuperscript{20} used their model and the Temkin adsorption isotherm as done here to estimate the adsorption energy at full coverage ($\Delta G_{ads}(\theta = 1)$) obtaining -23 kJ mol$^{-1}$. Jais-hankar et al.\textsuperscript{34} estimated the adsorption energy of SA at infinite dilution computationally, as done here through the PMF of adsorption ($\Delta G_{0} = -26.4$ kJ mol$^{-1}$) and experimentally with powder adsorption experiments, quartz crystal microbalance and HFRR as done here, obtaining estimates between -26.4 kJ mol$^{-1}$ and -29.1 kJ mol$^{-1}$. These values are in excellent agreement with the computational results presented here, see Table 2.

**Conclusions**

In this work, we have used MD simulations together with the MTT model to predict the adsorption isotherm of three OFMs on an iron oxide surface from hydrocarbon solvents.
First, we calculated the adsorption PMF for the three OFMs using MD and the ABF method, which proved to be more computationally efficient than other comparable enhanced sampling approaches. The adsorption strength increased in the order: SA < GMS < GMO, suggesting that the adsorption energy increases with the number of functional groups that interact with the surface. The stronger adsorption energy for GMO than GMS is due to the additional Z-alkene functional group in the tailgroup. Adsorption from both n-hexadecane and PAO4 give similar results, with the calculations using PAO4 taking longer to converge, due to slower transitions of the OFM tailgroup between solvent layers. Subsequently, the hard-disk surface area has been estimated using annealing simulations of pre-formed monolayer films at different surface coverages. The results show that the hard-disk area increases in the order: SA < GMS ∼ GMO.

The adsorption energy and hard-disk area from the ABF-MD simulations are used to calculate the adsorption isotherm using the MTT model. The surface coverage at low concentrations is mainly controlled by the adsorption energy and the maximum coverage by the hard-disk area. Lateral interactions between adsorbed molecules control how fast the adsorption isotherm reaches its asymptotic value and are necessary to better describe some of the literature experimental results. The adsorption isotherm for GMS and GMO are similar, in contrast with previous experimental results regarding the packing of saturated and Z-unsaturated surfactants. Since both molecules have very similar hard-disk areas, we propose that these experimental differences in coverage are mainly due to differences in the kinetic barriers to the formation of high-coverage monolayers. This factor could be investigated in future MD simulations of adsorption onto pre-formed monolayers.

Finally, the calculated adsorption energies from the PMFs have been validated through HFRR experiments. For all of the OFMS, the friction coefficient starts to reduce at a coverage of ∼2 molecules/nm², reaching a minimum value of ∼4 molecules/nm² for SA and ∼4 molecules/nm² for GMS and GMO. These values are well above the jamming limit predicted by the RSA model. Using the Jahanmir and Beltzer model, a good agreement is
obtained between the ABF-MD results and the experimental estimation of the adsorption energy for the OFMs (SA and GMS) with saturated tailgroups. However, the adsorption energy is overestimated by around 10 KJ mol$^{-1}$ for the OFM with a Z-unsaturated tailgroup (GMO). We expect that the differences for the unsaturated OFMs could be accounted for by incorporating the decrease in adsorption energy on already partially formed monolayers in future MD simulations. Overall, this study shows that it is possible to predict the adsorption isotherm of saturated OFMs in silico.

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**Supporting Information Available**

Additional figures and discussion on the convergence of the ABF calculations, results of the hard-disk area calculations, and the effects of various parameters on the MTT fits.

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Graphical TOC Entry

Adsorption energy

Hard-disk surface area

MTT Adsorption isotherm

- RA
- CXB
- G2B

Molar fraction