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Multiscale simulations of the hydration shells surrounding spherical Fe_3O_4 nanoparticles and effect on magnetic properties†

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Iron oxide magnetic nanoparticles (NPs) are excellent systems in catalysis and in nanomedicine, where they are mostly immersed in aqueous media. Even though the NP solvation by water is expected to play an active role, the detailed structural insight at the nanostructure oxide/water interface is still missing. Here, based on our previous efforts to obtain accurate models of dehydrated Fe_3O_4 NPs and of their magnetic properties and through multiscale molecular dynamics simulations combining the density functional tight binding method and force field, we unravel the atomistic details of the short range (chemical) and long range (physical) interfacial effects when magnetite nanoparticles are immersed in water. The influence of the first hydration shell on the structural, electronic and magnetic properties of Fe_3O_4 NPs is revealed by high-level hybrid density functional calculations. Hydrated Fe_3O_4 NPs possess larger magnetic moment than dehydrated ones. This work bridges the large gap between experimental studies on solvated Fe_3O_4 NPs and theoretical investigations on flat Fe_3O_4 surfaces covered with water and paves the way for further study of Fe_3O_4 NPs in biological environments.

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1. Introduction

Magnetite (Fe_3O_4) nanoparticles (NPs) are top-class materials for biomedical applications because of their excellent soft magnetism (high saturation magnetization and low coercive force), good biocompatibility and low cytotoxicity.^{1,2} They are used as new generation contrast agents for magnetic resonance imaging (MRI) and are ideal materials for targeted drug delivery, magnetic hyperthermia, bioseparation and biosensors.^{3–7} They are also widely used in catalysis.⁸ Fe_3O_4 NPs with variable sizes and different shapes including cubes, octahedra, rhombic dodecahedral, truncated octahedral and spheres have been successfully synthesized.^{9–16}

Because Fe_3O_4 NPs are often prepared from aqueous solutions^{14,15,17} and because in most of the potential applications mentioned above Fe_3O_4 NPs are supposed to act in an

aqueous environment, the surface/water interface as well as the solvation shells around the NP is expected to play a crucial role.¹⁸ However, the detailed atomistic description of the surface water chemistry and of the multilayer dynamical structures is not easy to be achieved, both experimentally and theoretically, and, in addition to that, it may largely depend on the NP shape and exposed facets.

Several studies have been conducted to explore the adsorption of water molecules on flat Fe_3O_4 surfaces.^{19–27} For instance, combining X-ray photoelectron spectroscopy (XPS) and density functional theory (DFT) calculations, Kendelewicz and co-workers proposed that, at low water vapor pressure and room temperature, water would not adsorb dissociatively on the $\text{Fe}_3\text{O}_4(001)$ surface, except on defect sites and that dissociation can be observed only between water vapor partial pressures of 10^{-4} and 10^{-2} Torr.^{19,21} At high water coverage on the $\text{Fe}_3\text{O}_4(001)$ surface, a mixed dissociated and undissociated adsorption mode was suggested by LEED,²² XPS^{21,25} and high-resolution electron energy loss spectroscopy (HREELS).²³ Similarly, on the $\text{Fe}_3\text{O}_4(111)$ surface, both dissociated water and undissociated water exist with increasing coverage, according to ultraviolet photoelectron spectroscopy (UPS) and thermal desorption spectroscopy (TDS).²⁴ On the theoretical side, DFT+U, hybrid density functional and density functional tight binding calculations indicate that, on the $\text{Fe}_3\text{O}_4(001)$ surface, isolated water molecules would not dissociate,

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whereas, at high water coverage, a mixed adsorption mode is favored.^{25–27} Water overlayers on the Fe₃O₄(001) surface were also investigated by multiscale molecular dynamics simulations.²⁷

However, as mentioned above, the interface between liquid water and Fe₃O₄ NPs is far different from considering water molecules on flat Fe₃O₄ surfaces. The detailed structure of the hydration shells for magnetite NPs in water was rarely investigated. By XPS and isosteric heat of adsorption, Tombácz proposed that a high density of hydroxyl groups exists at the surface of Fe₃O₄ NPs when exposed to water vapor.²⁸ By high-energy X-ray scattering, three distinct interatomic distance peaks (1.48, 1.95 and 2.39 Å) around faceted Fe₃O₄ NPs in water were found, which are attributed to hydrogen bonds between surface O atoms and H atoms in water, the Fe–O distance between Fe atoms on the surface and O atoms in molecular and dissociated water molecules.²⁹

To our knowledge, the theoretical study of magnetite NPs in water has not yet been performed for the main reason that the accurate simulation of magnetite NPs of realistic size in bulk water with quantum chemical methods is computationally very demanding. Magnetite is a complex magnetic material, which requires high-level quantum mechanical (QM) techniques, beyond standard density functional theory (DFT), for its correct description.^{26,30} Surrounding magnetite NPs with liquid water is, however, not feasible with DFT methods. Recently, through the development of new Fe–O interaction parameters, we managed to well describe magnetite by the Hubbard corrected self-consistent charge density functional tight-binding (SCC-DFTB+U; from now on, DFTB+U) method, which is a cheap and efficient quantum mechanical simulation method with comparable accuracy to DFT.³¹ Then, by combining DFTB+U and hybrid DFT methods, we finally succeeded in the quantum mechanical simulation of Fe₃O₄ NPs of realistic size (about 2.5 nm) and different shapes under vacuum.³²

In the present study, we have first investigated the adsorption behavior of one isolated water molecule on the Fe₃O₄ NP surface by putting one water molecule in contact with the bare roundish Fe₃O₄ NPs from our previous work.³² Then, we have adsorbed a water monolayer on a roundish NP and performed a global minimum structure search by a high temperature annealing process simulated by DFTB+U dynamics. At the end, by means of a multiscale approach, we surrounded the water-saturated NP with several water multilayers with the aim of simulating bulk water. The theoretical simulations are discussed in comparison with documented experimental data by high-energy X-ray scattering.²⁹

The extent of water dissociation on the surface of Fe₃O₄ NPs is obtained by performing DFTB+U dynamic simulations, whereas the influence of the first hydration shell on the structural, electronic and magnetic properties of Fe₃O₄ NPs is studied at the hybrid DFT level of theory. On top of that, by performing a quantum mechanics/molecular mechanics (QM/MM) molecular dynamics simulation of Fe₃O₄ NP in bulk

water, we obtain the atomistic insight into the structure of the solvation shells.

Our work fills the gap in theoretical simulations of magnetite NPs in water. It provides a clear picture of the chemistry at the NP surface and of the hydration shell structure around the NP of realistic size (about 1000 atoms) immersed in liquid water (more than 7000 water molecules). Understanding the fundamental properties of these types of interfaces is crucial for the development of nanoparticle-based applications in aqueous media.

2. Computational details

2.1 DFT calculations

Hybrid DFT calculations (HSE06³³) were carried out using the CRYSTAL17 package^{34,35} for investigating the structural, electronic and magnetic properties of the Fe₃O₄ NP with one water monolayer adsorbed on it. In these calculations, the Kohn–Sham orbitals are expanded in Gaussian-type orbitals (the all-electron basis sets adopted for the elements in our model systems are H|5-11G*, O|8-411G* and Fe|8-6-411G*, according to the previous work on Fe₃O₄³⁰). The convergence criterion of 0.02 eV Å^{−1} for force was used for geometry optimization and the convergence criterion for total energy was set as 10^{−6} Hartree for all the calculations.

2.2 DFTB calculations

SCC-DFTB, which is a method cheaper and faster than DFT, can well describe Fe₃O₄ bulk, surface, NPs and the water/Fe₃O₄ interfaces.^{27,31,32} The SCC-DFTB method^{36–38} is an approximated DFT-based method that derives from the second-order expansion of the Kohn–Sham total energy in DFT with respect to the electron density fluctuations. The SCC-DFTB total energy can be defined as:

$$E_{\text{tot}} = \sum_i^{\text{occ}} \epsilon_i + \frac{1}{2} \sum_{\alpha, \beta}^N \gamma_{\alpha\beta} \Delta q_{\alpha} \Delta q_{\beta} + E_{\text{rep}}, \quad (1)$$

where the first term is the sum of the one-electron energies ϵ_i coming from the diagonalization of an approximated Hamiltonian matrix. Δq_{α} and Δq_{β} are the induced charges on the atoms α and β , respectively, and $\gamma_{\alpha\beta}$ is a coulombic-like interaction potential. E_{rep} is a short-range pairwise repulsive potential.

The SCC-DFTB calculations were carried out using the DFTB+ package.³⁹ The “trans3d-0-1” set of parameters⁴⁰ are used for the Fe–Fe and Fe–H interactions, and the “mio-1-1” set of parameters³⁶ are used for the O–O, H–O and H–H interactions. For the Fe–O interactions, we used the Slater–Koster files fitted by some of us previously,³¹ which can well describe Fe₃O₄ bulk, surface, NPs and the water/Fe₃O₄ interfaces.^{27,31,32} SCC-DFTB+U⁴¹ with an effective $U - J$ value of 3.5 eV was adopted to properly deal with the strong correlation effects among Fe 3d electrons, according to our previous work on magnetite.^{26,30,32} The convergence criterion of 10^{−4} a.u. for



force was used during geometry optimization and the convergence threshold on the self-consistent charge (SCC) procedure was set to be 10^{-5} a.u.

SCC-DFTB+U molecular dynamics was performed within the canonical ensemble (NVT) to obtain the global minimum structure of the Fe_3O_4 NP with one monolayer water adsorption. The time step was set as 1 fs and an Andersen thermostat⁴² was used to target the desired temperatures. To simulate the temperature annealing processes, the NP with a water monolayer was quickly heated up to 500 K (about 0.5 ps), kept at 500 K for 3.5 ps and then cooled down slowly to 100 K, when no further structural changes were expected. The total simulation time is 24.5 ps. To well describe the hydrogen bonds, a modified hydrogen bonding damping (HBD) function was introduced with a $\zeta = 4$ parameter⁴³ for the SCC-DFTB+U calculations.

2.3 QM/MM calculations

The QM/MM molecular dynamics in this work is carried out through the additive-coupling scheme,^{44,45} in which the total potential energy for the system is given by

$$E_{\text{TOTAL}} = E_{\text{QM}} + E_{\text{MM}} + E_{\text{QM/MM}}, \quad (2)$$

where E_{QM} and E_{MM} are the total potential energy of the QM and the MM sub-systems, respectively. $E_{\text{QM/MM}}$ stands for the total energy of the QM/MM coupling term, which accounts for electrostatic interactions and the vdW interactions between the QM and MM atoms. The Atomic Simulation Environment (ASE) interface⁴⁶ handles the coupling between the QM and the MM sub-systems through the electrostatic embedding QM/MM scheme.⁴⁷ We made use of the DFTB+ software³⁹ for the treatment of the QM part at the DFTB level, whereas for the MM part, composed only of classical water molecules, we employed the Amber16 code⁴⁸ with water described by the flexible q-SPC/Fw model.⁴⁹ For the QM part, the same setup as the DFTB calculations discussed above is adopted.

The coupling between the QM and MM regions consists of an electrostatic term calculated as the Coulomb interaction between the atomic charges of the QM part and the charges of the MM atoms, whereas all the non-Coulomb interactions (van der Waals) between the subsystems have been implemented as a Lennard-Jones (12–6) type potential,

$$U_{\text{vdW}}^{\text{QM/MM}} = \sum_{\text{A} \in \text{MM}} \sum_{\text{B} \in \text{QM}} \epsilon_{\text{AB}} \left[\left(\frac{\sigma_{\text{AB}}}{R_{\text{AB}}} \right)^{12} - 2 \left(\frac{\sigma_{\text{AB}}}{R_{\text{AB}}} \right)^6 \right], \quad (3)$$

where R is the distance between pair of atoms, ϵ is the depth of the attractive potential well, and σ is the distance at which the particle–particle potential energy U is zero. Values of ϵ and σ for oxygen and hydrogen atoms are taken from the SPC/Fw model,⁴⁹ and ϵ and σ for Fe atoms are taken from ref. 50. ϵ_{AB} and σ_{AB} cross-parameters are estimated using the Lorentz–Berthelot combining rules.⁵¹

To model the roundish Fe_3O_4 NP embedded in bulk water, we put the optimized hydrated NP into a water droplet with a diameter of 8 nm by using the PACKMOL program.⁵²

Molecular dynamics was performed within the NVT ensemble with a target temperature of 300 K maintained constant using the Berendsen thermostat. The Newton's equations of motion were integrated in time using the Velocity–Verlet algorithm⁵³ with a time step of 1 fs. The system has been equilibrated for 10 ps and then a production run has been performed for 20 ps.

2.4 MM calculations

The potential energy form for the classical description of the systems is based on the well-established COMPASS force field⁵⁴ that estimates the vdW forces through a Lennard–Jones (6–9) potential while long-range electrostatics is modeled by a classical Coulomb potential, given by

$$E_{\text{vdW}} = \sum_{i,j} \epsilon_{ij} \left[2 \left(\frac{\sigma_{ij}}{r_{ij}} \right)^9 - 3 \left(\frac{\sigma_{ij}}{r_{ij}} \right)^6 \right], \quad (4)$$

$$E_{\text{elec}} = \frac{1}{4\pi\epsilon_0} \sum_{i,j} \frac{q_i q_j}{r_{ij}}. \quad (5)$$

Herein, σ_{ij} stands for the inter-atomic distance between a pair of atoms at which the potential energy reaches a minimum value, and ϵ_{ij} defines the potential well depth of the attractive component. q_i and q_j represent the partial atomic charges on the classical atoms i and j , separated by an inter-atomic distance r_{ij} . For unlike atoms, the sigma and epsilon LJ (6–9) cross-parameters are calculated by the 6th power combining rules⁵⁵ given by eqn (6) and (7), respectively:

$$\sigma_{ij} = \left(\frac{\sigma_i^6 + \sigma_j^6}{2} \right)^{1/6}, \quad (6)$$

$$\epsilon_{ij} = 2\sqrt{\epsilon_i \epsilon_j} \left(\frac{\sigma_i^3 \sigma_j^3}{\sigma_i^6 + \sigma_j^6} \right). \quad (7)$$

All MM-MD simulations were carried out with the LAMMPS program (version 7 Aug 2019).⁵⁶ The atomic coordinates of the Fe, O, and H atoms in the partially hydroxylated Fe_3O_4 NS are frozen at the DFTB-optimized geometry by zeroing the forces on these atoms every MM-MD simulation step. The partially hydrated NP is obtained by removing all the molecular water from the optimized fully hydrated NP. To solvate this system, we made use of the PACKMOL program⁵² to randomly put water molecules around the Fe_3O_4 NP within a simulation box of $100 \times 100 \times 100 \text{ \AA}^3$. Bonded and non-bonded FF parameters for the COMPASS-based three-site water model and the hydroxyl group are taken from the INTERFACE-FF.⁵⁷ The LJ (6–9) parameters for the Fe(II), Fe(III), and O(II) atom-types in the Fe_3O_4 NP are taken from ref. 50. For the Fe–O cross-interaction, we use an optimized set of LJ (6–9) parameters available in ref. 58. The partial-atomic charges on the magnetite atoms are derived from the DFT/HSE06 calculations in line with our previous work.²⁷

To vanish any atomic overlapping and minimize the total energy of the system before the equilibration phase, we carried out a minimization phase with a maximum number of 500 000



steps and a convergence tolerance of 10^{-7} for forces through the conjugate gradient algorithm. The equilibration phase was carried out for 10 ns in the isotherm-isobaric (NVT) ensemble until convergence of the bulk-water density at $T = 300$ K. The production phase explored 50 ns of the phase space in the NVT ensemble at $T = 300$ K. Electrostatic and LJ (6–9) potentials utilized a cut-off of 10 Å, and the Newton's equations of motion were solved using the Velocity–Verlet integrator⁵³ with a time step of 1.0 fs. The long-range solver particle–particle particle–mesh (PPPM)⁵⁹ handled the electrostatic interactions with a threshold of 10^{-6} for forces under Periodic Boundary Conditions (PBC).

3. Results and discussion

3.1 First hydration shell around the Fe_3O_4 NP: adsorption of a water monolayer

In our previous work, we have obtained the global minimum structure of a roundish Fe_3O_4 NP under vacuum ($(\text{Fe}_3\text{O}_4)_{136}(\text{H}_2\text{O})_{18}$) with a diameter of 2.5 nm.³² We expect that water adsorption might affect the curved surface structure. Before putting a whole water monolayer around the Fe_3O_4 NP, we first investigate the adsorption behavior of a single isolated water molecule on the various undercoordinated Fe sites present on the surface.³² We define two kinds of Fe ions in the NP, Fe_{Tet} and Fe_{Oct} , depending on the occupied type of bulk lattice site (tetrahedral and octahedral) and we keep this classification even when they become undercoordinated at the NP surface. Labels, including 3c, 4c, 5c and 6c, are then added to indicate the actual coordination number of the corresponding ions. For example, $\text{Fe}_{\text{Tet}-3c}$ and O_{3c} are a three-fold coordinated Fe ion at a tetrahedral site and a three-fold coordinated O ion, respectively.

According to previous studies,^{21–26} both molecular and dissociated (split into OH and H) water are present on flat Fe_3O_4 surfaces, with the water O atom chemically bound to surface Fe ions. Therefore, both molecular and dissociated adsorption modes are considered at different Fe sites on the NP surface and the resulting binding energy values for a single water molecule adsorbed on various sites are listed in Table 1. On

Table 1 The binding energies (E_b) for single water molecules adsorbed on different sites of the Fe_3O_4 NP

Adsorption site	Adsorption mode	E_b (eV)
$\text{Fe}_{\text{Tet}-3c}$	Molecular	−1.65 to −1.75
	Dissociated (H on O_{3c})	−0.91 to −1.00
	Dissociated (H on O_{2c})	−1.32 to −1.74
$\text{Fe}_{\text{Oct}-4c}$	Molecular	−1.82 to −1.95
	Dissociated (H on O_{3c})	−2.84 to −2.99
	Dissociated (H on O_{2c})	−3.17 to −3.90
$\text{Fe}_{\text{Oct}-5c}$	Molecular	−1.50 to −1.86
	Dissociated (H on O_{3c})	−1.11 to −1.24
	Dissociated (H on O_{2c})	−1.32 to −1.89

$\text{Fe}_{\text{Tet}-3c}$ sites, molecular adsorption is preferred, except for few cases where the H atom from dissociated water adsorbs on a two-fold coordinated surface oxygen atom (O_{2c}). This indicates that O_{2c} is far more reactive than O_{3c} , as one would expect. This agrees with previous results, which show that a single water molecule would not dissociate on $\text{Fe}_{\text{Tet}-3c}$ sites of the $\text{Fe}_3\text{O}_4(111)$ surface⁶⁰ because all the O atoms on the surface are three-fold coordinated. In contrast, on $\text{Fe}_{\text{Oct}-4c}$ sites, the dissociated mode is always favored with respect to the molecular mode, independent of the O site where H is adsorbed. On $\text{Fe}_{\text{Oct}-5c}$ sites, the situation is similar to that on $\text{Fe}_{\text{Tet}-3c}$ sites, *i.e.* molecular adsorption is preferred, except when the dissociated H atom binds to a surface O_{2c} site.

On the basis of the results above, we built an initial model of water-saturated Fe_3O_4 NPs by putting one undissociated water molecule on every $\text{Fe}_{\text{Tet}-3c}$ and $\text{Fe}_{\text{Oct}-5c}$ sites, one OH and one undissociated water molecule on every $\text{Fe}_{\text{Oct}-4c}$ site of the roundish NP cut from bulk Fe_3O_4 ($(\text{Fe}_3\text{O}_4)_{136}(\text{H}_2\text{O})_{18}$) from our previous work,³² as shown in Fig. 1. As a consequence, all the Fe ions in the NP become fully coordinated, leading to the molecular chemical formula of $(\text{Fe}_3\text{O}_4)_{136}(\text{H}_2\text{O})_{186}$. The H atoms from dissociated water are put preferentially on O_{2c} on the NP surface.

We must note that the dissociation behavior depends not only on the adsorption sites but also on the extent of water coverage, as indicated by the studies of water on flat Fe_3O_4 surfaces.^{21–26} To obtain the global minimum structure of the water-saturated Fe_3O_4 NP, we then performed the MD run, with the DFTB+U method, simulating a temperature annealing process up to 500 K, whose temperature profile is shown in Fig. 1. After the annealing, the Fe_3O_4 NP was further fully optimized at the more accurate hybrid density functional HSE06 level of theory. We observe that, after the annealing, about half of water molecules on $\text{Fe}_{\text{Tet}-3c}$ dissociate into OH and H. Some of the H atoms from water dissociation go to O atoms on the NP surface and some of them react with OH on Fe_{Oct} sites forming molecular water again. The simultaneous dissociation and reforming water processes are due to the complex geomorphology of the NP surface, which cannot be found on flat surfaces. In the end, 43% water molecules on the NP surface are dissociated, indicating the strong ability of the NP surface to provoke this reaction. This agrees well with previous experimental work, in which a high density of hydroxyl groups at the surface of Fe_3O_4 NPs was found when exposed to water vapor.²⁸

By comparing the simulated extended X-ray absorption fine structure (EXAFS) of Fe_3O_4 bulk, of the NP under vacuum and of the water-saturated NP (with one adsorbed monolayer), as shown in Fig. S1 in the ESI,† we can conclude that water adsorption improves the degree of crystallinity of the NP due to the saturation of low-coordinated Fe ions on the surface. This agrees with a previous study on ZnS nanoparticles, which shows that the binding of water to the as-formed ZnS nanoparticles significantly reduces distortions of the surface.⁶¹ We expect that besides affecting the structure, water adsorption will have consequences also on the electronic and magnetic



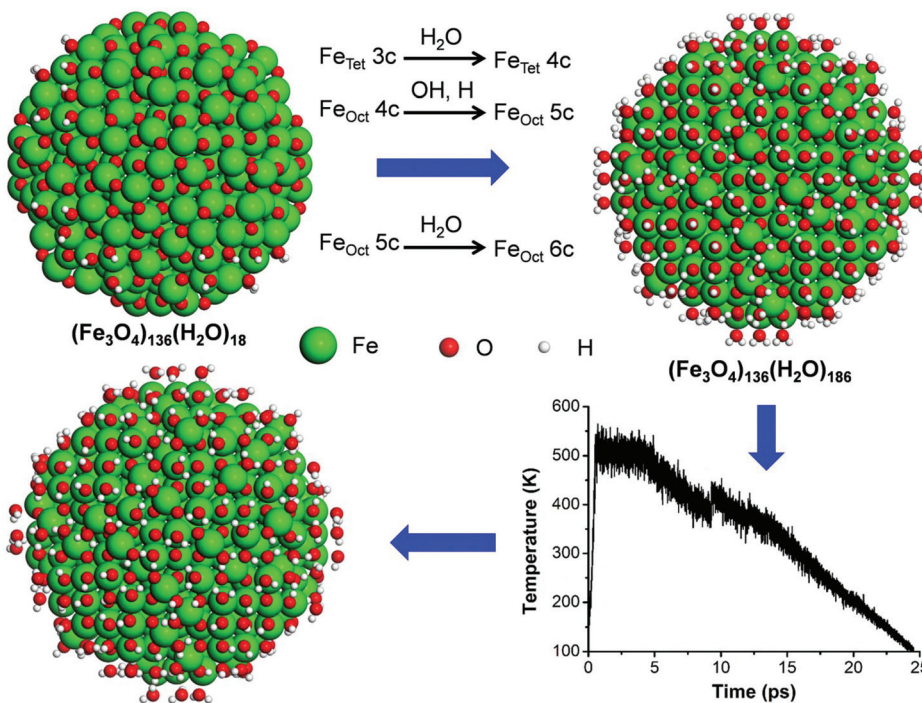


Fig. 1 The scheme of the saturation of the bare Fe_3O_4 NP with water molecules, including the simulated annealing temperature profile (bottom right) and global minimum structure of the magnetite NP saturated by 186 water molecules (bottom left). The color coding of atoms is given in the legend in the middle.

properties of Fe_3O_4 NPs. To prove this, we investigated the charge and spin distribution in the magnetite NP covered with a water monolayer using the HSE06 method, as shown in Fig. 2. Interestingly, we find that $\text{Fe}_{\text{Tet}}^{2+}$ ions exist not only on the surface but also inside the NP (Fig. 2a), which is in sharp contrast with the situation for the roundish Fe_3O_4 NP under vacuum, where $\text{Fe}_{\text{Tet}}^{2+}$ ions only exist on the surface. Detailed analysis of the charges of all the Fe ions in roundish Fe_3O_4 NP, under vacuum and under water covering, is shown in Fig. 2b. In sharp contrast with the NP under vacuum, the charge grouping in the water-saturated NP is more pronounced, similar to what has been observed in bulk magnetite.³⁰ Therefore, we can state that the effect of water adsorption is to make the NP more crystalline not only from the structural point of view but also from the charge ordering one.

The total magnetic moment (m_{tot}) of the water-saturated Fe_3O_4 NP is determined by a series of HSE06 full atomic relaxation calculations considering different m_{tot} values. As shown in Fig. S2 in the ESI,† the optimal m_{tot} of the NP after water saturation increases to $656\mu_{\text{B}}$ from $600\mu_{\text{B}}$ under vacuum, with high benefit for biomedical applications where NPs with high magnetic moments are more effective. The increment of the m_{tot} can be rationalized by the change in the charge distribution mentioned above. After water saturation, the number of $\text{Fe}_{\text{Tet}}^{2+}$ ions ($N(\text{Fe}_{\text{Tet}}^{2+})$) increases and the number of $\text{Fe}_{\text{Tet}}^{3+}$ ions ($N(\text{Fe}_{\text{Tet}}^{3+})$) decreases. In the meantime, the number of $\text{Fe}_{\text{Oct}}^{3+}$ ($N(\text{Fe}_{\text{Oct}}^{3+})$) increases and the number of $\text{Fe}_{\text{Oct}}^{2+}$ ($N(\text{Fe}_{\text{Oct}}^{2+})$) decreases in order to maintain the neutrality of the

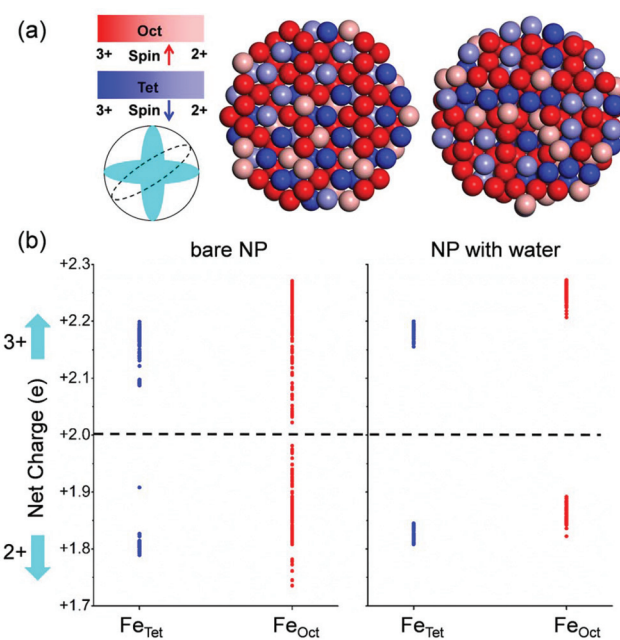


Fig. 2 (a) Selected dissected views showing the charge and spin distribution in the roundish magnetite NP covered with a water monolayer. Oxygen and hydrogen atoms are not shown. (b) The net charge distribution for the Fe ions at tetrahedral and octahedral sites in the optimized roundish magnetite NP without (on the left) and with (on the right) water adsorption. All the results are from HSE06 calculations.

NP. As a consequence, according to the formula for estimating the m_{tot} of Fe_3O_4 NPs, proposed in our previous work,³² *i.e.* $m_{\text{tot}} = 5 \times (N(\text{Fe}_{\text{Oct}}^{3+}) - N(\text{Fe}_{\text{Tet}}^{3+})) + 4 \times (N(\text{Fe}_{\text{Oct}}^{2+}) - N(\text{Fe}_{\text{Tet}}^{2+}))$, the m_{tot} increases. Finally, we must note that the main features of the electronic structure of the NP do not change much after water saturation, as indicated by comparison of the density of states shown in Fig. S3 in the ESI.†

3.2 Immersion of the hydrated Fe_3O_4 NP in bulk water

As mentioned in the introduction, to gain a detailed structural insight into the water solvation shells around the roundish Fe_3O_4 NP, we built a realistic model by immersing the optimized fully hydrated Fe_3O_4 NP from the previous section into a water droplet with a diameter of 8 nm, as shown in Fig. 3. Then a multiscale QM/MM molecular dynamics simulation has been performed at 300 K for 30 ps, where the hydrated Fe_3O_4 NP is treated at the DFTB level, whereas the water molecules around are treated at the MM level.

First, we focus on the innermost (first) hydration shell, *i.e.*, the direct bonds between surface Fe ions and OH and H_2O , which is treated at the DFTB level of theory. The bond length distribution of the Fe–O at the NP/water interface is plotted in Fig. 4. There is a prominent peak centered at 1.99 Å with a shoulder at 1.93 Å, which can well explain the experimentally observed peak at 1.95 Å.²⁹ By projecting the total Fe–O bond length distribution for different Fe ions (Fe_{Tet} and Fe_{Oct}) with different O ions (O in OH and H_2O), the peak can be rationalized in detail. The peak at 1.99 Å is mainly contributed by

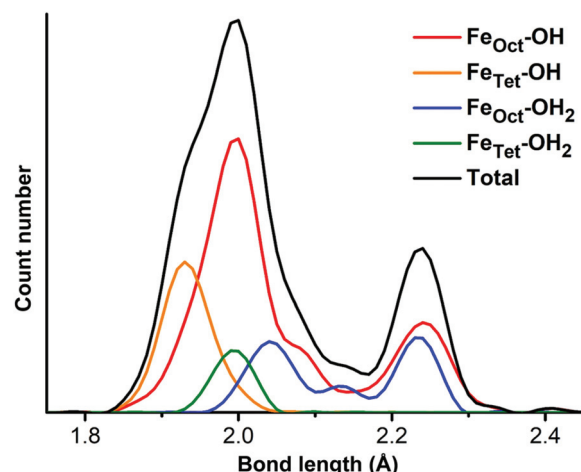


Fig. 4 Fe–O bond length distribution at the Fe_3O_4 NP/water interface.

bonds between Fe_{Oct} ions and OH (red line in Fig. 4) and the shoulder at 1.93 Å is due to the bonds between Fe_{Tet} and OH (orange line). The bond length of $\text{Fe}_{\text{Tet}}\text{–OH}_2$ is also around 1.99 Å (green line). The bond lengths of $\text{Fe}_{\text{Oct}}\text{–OH}_2$ are longer than those of $\text{Fe}_{\text{Tet}}\text{–OH}_2$ and are widely distributed in the range from 1.99 Å to 2.26 Å (blue line). Another peak centered at 2.24 Å (black line) is very close to the experimentally observed peak at 2.39 Å,²⁹ which is contributed by both $\text{Fe}_{\text{Oct}}\text{–OH}$ and $\text{Fe}_{\text{Oct}}\text{–OH}_2$ bonds. Note that the $\text{Fe}_{\text{Oct}}\text{–OH}$ bond length depends on the coordination of the O atom of the OH group. When one OH is bonded to one Fe ion, the bond length is shorter (peak at 1.99 Å), whereas when one OH is bonded with two Fe ions, the bond length is longer (peak at 2.24 Å).

Then, we analyze all the hydration shells. We extract from the QM/MM molecular dynamics runs the radial distribution function (RDF), $g(d)$, of O in water molecules with respect to the NP center. To gain a deeper insight into the Fe_3O_4 NP/water interface, a decomposed RDF based on the O-type is presented in Fig. 5, *i.e.*, O from molecular water molecules in the MM part (black line), O from molecular water molecules in the QM part (red line) and O from the dissociated OH in the QM part (green line) are shown, respectively. The RDF is normalized by the density of bulk water. In addition, to clearly set the position of the NP surface, the RDFs of the O and Fe atoms in the NP, with meaningless y-values, are also plotted (blue and gray lines, respectively). According to the RDF (blue, gray and green lines), the NP surface atoms are set in a window range of distances between 12 Å and 14 Å, because the NP is not perfectly spherical. Consequently, the water O atoms present a wide distribution of d values (green and red lines).

The distribution of O in water in the MM part (black line in Fig. 5) presents two clear peaks, one centered at 15.5 Å and the other at 16.5 Å. Through a detailed check of the structures of MM water molecules during the dynamics in the range between 14.2 Å and 15.7 Å (about 2 to 3 Å from the NP surface), we could unravel the nature of the interactions in this

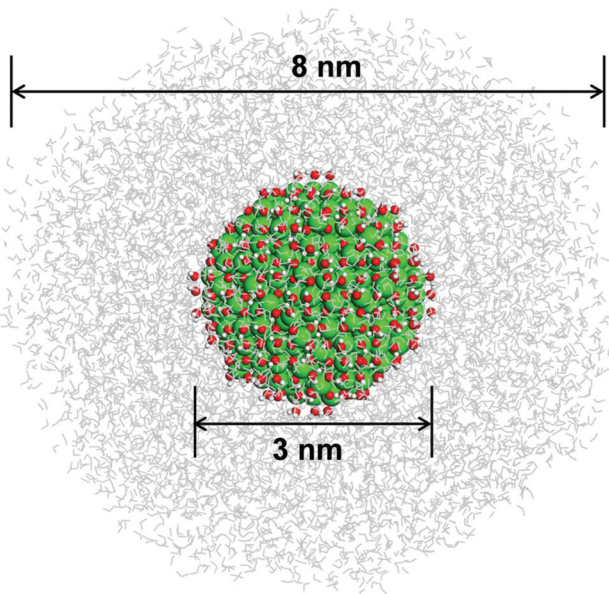


Fig. 3 Atomic structure of the hydrated Fe_3O_4 NP embedded in a water droplet with an overall stoichiometry of $(\text{Fe}_3\text{O}_4)_{136}(\text{H}_2\text{O})_{7379}$. The water droplet is 8 nm in diameter and the hydrated Fe_3O_4 NP is about 3 nm in diameter. Fe, O, and H atoms of the hydrated NP (treated at the DFTB level of theory) are shown as green, red and white balls, respectively. Molecular water molecules of the water droplet (treated at the MM level of theory) are represented by gray lines.



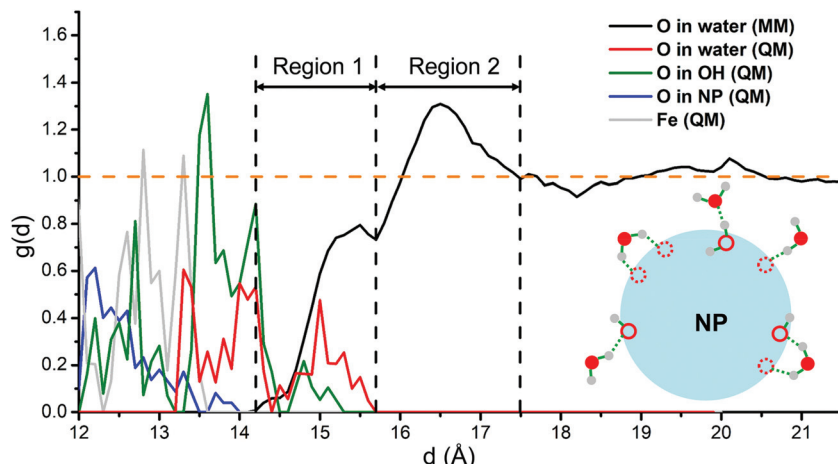


Fig. 5 RDF of O in water molecules as extracted from the QM/MM molecular dynamics runs of the hydrated Fe_3O_4 NP enclosed in a water droplet. The zero reference is the center of the NP. The distribution function of O and Fe in Fe_3O_4 NP are also shown (blue and gray lines) just to indicate the position of the surface of the NP. Inset schematically shows the location of water molecules in region 1 in the MM part and the interaction between these water molecules and the NP surface. The solid red balls represent O of water in region 1 in the MM part. The hollow red balls are O of water or OH directly bonded with Fe ions. The dotted green lines represent hydrogen bonds.

region 1, which are schematically shown in Fig. 5. Some of the water molecules in region 1 interact with the surface O of the NP through hydrogen bonds and some of them interact with the innermost hydration shell through hydrogen bonds.

Based on the analysis above, therefore, we can classify the water molecules and OH directly bonded with Fe ions as the first hydration shell, whereas the water molecules in region 1 as the second hydration shell. All the water molecules in region 2 of Fig. 5 do not interact directly with the NP but interact with the second hydration shell through hydrogen bonds and, therefore, can be classified as the third hydration shell. The RDF quickly converges to 1.0 after 17.5 Å, indicating that the water above 5 Å from the NP surface is essentially bulk water. Similarly, previous MM molecular dynamics simulations

show that water close to the surfaces of the hematite nanoparticles also forms several ordered layers.⁶²

Note that during the QM/MM molecular dynamics, atoms in the QM part are frozen to their optimized position due to some technical issues. Therefore, the absolute value of the RDF intensity for the O in the QM part (green and red lines) is not precise. However, we believe that atoms in the first hydration shell would not move so much during the dynamics because they are tightly bonded to the NP surface atoms.

To exclude any effect of freezing water molecules during the QM/MM molecular dynamics, we also performed MM molecular dynamics where all the water molecules are free to move for a longer simulation run (60 ns in total) and in a larger box (10 nm × 10 nm × 10 nm). In this model, the partially hydrated

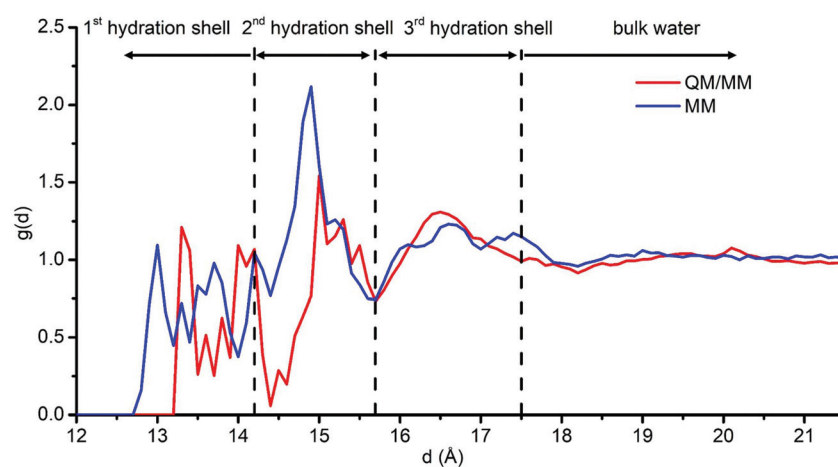


Fig. 6 Radial distribution function $g(d)$ of O in water molecules as extracted from the QM/MM (the red line) and MM (the blue line) molecular dynamics runs of the hydrated Fe_3O_4 NP enclosed in a water droplet, respectively. The zero reference is the center of the NP.



NP, obtained by removing all the molecular water from the optimized hydrated NP, is fully optimized by DFTB+U and then put in a large water box. The RDF for the O atoms in the water molecules based on the last 50 ns of the MM molecular dynamics run is shown in Fig. 6 (blue line). To compare the MM results with those from the QM/MM simulation, we must sum up the black and red lines (not the pink one) in Fig. 5 and display the resulting merged red line in Fig. 6, because the RDF extracted from the MM calculations is only for the O atoms belonging to molecular (undissociated) water molecules, which does not include the contributions of dissociated OH groups fixed on the NP surface. In general, the curves from QM/MM and MM agree quite well one with the other, indicating three hydration shells. However, there is a shift of peaks (about 0.3 Å) in the first hydration shell. This shift is because the optimized force field parameters used in the MM simulation tend to maximize the number of hydrogen bonds established by water molecules with the nanoparticle surface atoms, resulting in a shorter distance between O in water molecules and the center of the NP, as discussed in our previous paper.⁵⁸ The good agreement between QM/MM and MM data not only confirms the reliability of our results but also suggests that with a chemically stable and fully optimized Fe_3O_4 NP structure (by hybrid density functional methods) and with reasonable force field parameters⁵⁸ MM molecular dynamics can be an extremely efficient way to explore Fe_3O_4 NP/water interfaces and even more complex systems, such as functionalized Fe_3O_4 NPs in water for biomedical applications.

4. Conclusions

In this work, we presented a multistep investigation of the interaction of roundish Fe_3O_4 NPs with water, based on hybrid DFT, QM(DFTB)/MM and MM calculations.

At the hybrid DFT level of theory, we first investigated single water molecule adsorption modes on various types of uncoordinated Fe sites present on a realistic curved nanoparticle. Then, by decorating all the adsorption sites, we studied a full water monolayer coverage and observed the effect of the first hydration shell on the structural and electronic properties of Fe_3O_4 NPs: they become more crystalline and the total magnetic moment increases, which has a positive impact on their nanomedical applications. We found out that 43% of water molecules in the first hydration shell dissociate, suggesting a high tendency of roundish Fe_3O_4 NPs to become hydroxylated.

By a multi-scale approach combining the density functional tight-binding method and classical force field calculations (QM(DFTB)/MM), we investigated the atomistic structure of the hydration shells around the roundish Fe_3O_4 NP immersed in bulk water. We identified three hydration shells and determined that above 5 Å away from the NP surface water holds essentially its bulk properties. The bond length distribution of the Fe–O at the NP/water interface successfully explains the experimental observations. Interactions between the first and

the second hydration shells as well as between the second and the third hydration shells are mainly through H-bonds. Parallel MM calculations lead to very similar results, which suggest that the very cheap MM approach could be used to obtain reliable information on the dynamical behavior of magnetite NPs in aqueous media.

Our work fills the gap in theoretical simulations of magnetite NPs in water and paves the way for further studies of more complex functionalized Fe_3O_4 NPs in biological environments for nanomedical applications.

Conflicts of interest

The authors declare no conflict of interest.

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References

- 1 J. M. Perez, L. Josephson, T. O'Loughlin, D. Högemann and R. Weissleder, *Nat. Biotechnol.*, 2002, **20**, 816–820.
- 2 J. Liu, Z. Sun, Y. Deng, Y. Zou, C. Li, X. Guo, L. Xiong, Y. Gao, F. Li and D. Zhao, *Angew. Chem., Int. Ed.*, 2009, **48**, 5875–5879.
- 3 W. Wu, Z. Wu, T. Yu, C. Jiang and W. Kim, *Sci. Technol. Adv. Mater.*, 2015, **16**, 023501.
- 4 Q. A. Pankhurst, N. T. K. Thanh, S. K. Jones and J. Dobson, *J. Phys. D: Appl. Phys.*, 2009, **42**, 224001.
- 5 C. Sun, J. S. Lee and M. Zhang, *Adv. Drug Delivery Rev.*, 2008, **60**, 1252–1265.
- 6 S. Laurent, D. Forge, M. Port, A. Roch, C. Robic, L. V. Elst and R. N. Muller, *Chem. Rev.*, 2008, **108**, 2064–2110.
- 7 M. Colombo, S. Carregal-Romero, M. F. Casula, L. Gutiérrez, M. P. Morales, I. B. Böhm, J. T. Heverhagen, D. Prosperi and W. J. Parak, *Chem. Soc. Rev.*, 2012, **41**, 4306–4334.
- 8 M. B. Gawande, P. S. Brancoa and R. S. Varma, *Chem. Soc. Rev.*, 2013, **42**, 3371.
- 9 S. Sun and H. Zeng, *J. Am. Chem. Soc.*, 2002, **124**, 8204–8205.
- 10 Y. Hou, J. Yu and S. Gao, *J. Mater. Chem.*, 2003, **13**, 1983–1987.
- 11 J. Park, K. An, Y. Hwang, J. Park, H. Noh, J. Kim, J. Park, N. Hwang and T. Hyeon, *Nat. Mater.*, 2004, **3**, 891–895.
- 12 M. V. Kovalenko, M. I. Bodnarchuk, R. T. Lechner, G. Hesser, F. Schäffler and W. Heiss, *J. Am. Chem. Soc.*, 2007, **129**, 6352–6353.
- 13 L. Zhao and L. Duan, *Eur. J. Inorg. Chem.*, 2010, 5635–5639.

14 X. Li, D. Liu, S. Song, X. Wang, X. Ge and H. Zhang, *CrystEngComm*, 2011, **13**, 6017–6020.

15 L. Zhao, H. Zhang, Y. Xing, S. Song, S. Yu, W. Shi, X. Guo, J. Yang, Y. Lei and F. Cao, *Chem. Mater.*, 2008, **20**, 198–204.

16 A. G. Roca, M. P. Morales and C. J. Serna, *IEEE Trans. Magn.*, 2006, **42**, 3025–3029.

17 X. Cheng, J. Jiang, D. Jiang and Z. Zhao, *J. Phys. Chem. C*, 2014, **118**, 12588–12598.

18 A. E. Nel, L. Mädler, D. Velegol, T. Xia, E. M. V. Hoek, P. Somasundaran, F. Klaessig, V. Castranova and M. Thompson, *Nat. Mater.*, 2009, **8**, 543–557.

19 T. Kendelewicz, P. Liu, C. S. Doyle, G. E. Brown, E. J. Nelson and S. A. Chambers, *Surf. Sci.*, 2000, **453**, 32–46.

20 G. S. Parkinson, Z. Novotný, P. Jacobson, M. Schmid and U. Diebold, *J. Am. Chem. Soc.*, 2011, **133**, 12650–12655.

21 T. Kendelewicz, S. Kaya, J. T. Newberg, H. Bluhm, N. Mulakaluri, W. Moritz, M. Scheffler, A. Nilsson, R. Pentcheva and G. E. Brown, *J. Phys. Chem. C*, 2013, **117**, 2719–2733.

22 N. Mulakaluri, R. Pentcheva, M. Wieland, W. Moritz and M. Scheffler, *Phys. Rev. Lett.*, 2009, **103**, 176102.

23 S. Liu, S. Wang, W. Li, J. Guo and Q. Guo, *J. Phys. Chem. C*, 2013, **117**, 14070–14074.

24 Y. Joseph, C. Kuhrs, W. Ranke and W. Weiss, *Surf. Sci.*, 1999, **433–435**, 114–118.

25 M. Meier, J. Hulva, Z. Jakub, J. Pavleč, M. Setvin, R. Bliem, M. Schmid, U. Diebold, C. Franchini and G. S. Parkinson, *Proc. Natl. Acad. Sci. U. S. A.*, 2018, **115**, E5642–E5650.

26 H. Liu and C. Di Valentin, *Nanoscale*, 2018, **10**, 11021.

27 H. Liu, E. Bianchetti, P. Siani and C. Di Valentin, *J. Chem. Phys.*, 2020, **152**, 124711.

28 E. Tombácz, A. Hajdú, E. Illés, K. László, G. Garberoglio and P. Jedlovszky, *Langmuir*, 2009, **25**, 13007–13014.

29 S. L. J. Thomä, S. W. Krauss, M. Eckardt, P. Chater and M. Zobel, *Nat. Commun.*, 2019, **10**, 995.

30 H. Liu and C. Di Valentin, *J. Phys. Chem. C*, 2017, **121**, 25736.

31 H. Liu, G. Seifert and C. Di Valentin, *J. Chem. Phys.*, 2019, **150**, 094703.

32 H. Liu and C. Di Valentin, *Phys. Rev. Lett.*, 2019, **123**, 186101.

33 A. V. Kruckau, O. A. Vydrov, A. F. Izmaylov and G. E. Scuseria, *J. Chem. Phys.*, 2006, **125**, 224106.

34 R. Dovesi, A. Erba, R. Orlando, C. M. Zicovich-Wilson, B. Civalleri, L. Maschio, M. Rerat, S. Casassa, J. Baima, S. Salustro and B. Kirtman, *Wiley Interdiscip. Rev.: Comput. Mol. Sci.*, 2018, **8**, e1360.

35 R. Dovesi, V. R. Saunders, C. Roetti, R. Orlando, C. M. Zicovich-Wilson, F. Pascale, B. Civalleri, K. Doll, N. M. Harrison, I. J. Bush, P. D'Arco, M. Llunell, M. Causà, Y. Noël, L. Maschio, A. Erba, M. Rerat and S. Casassa, *CRYSTAL17 User's Manual*, University of Torino, Torino, Italy, 2017.

36 M. Elstner, D. Porezag, G. Jungnickel, J. Elsner, M. Haugk, T. Frauenheim, S. Suhai and G. Seifert, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1998, **58**, 7260.

37 G. Seifert and J. Joswig, *Wiley Interdiscip. Rev.: Comput. Mol. Sci.*, 2012, **2**, 456.

38 M. Elstner and G. Seifert, *Philos. Trans. R. Soc., A*, 2014, **372**, 20120483.

39 B. Aradi, B. Hourahine and T. Frauenheim, *J. Phys. Chem. A*, 2007, **111**, 5678.

40 G. Zheng, H. A. Witek, P. Bobadova-Parvanova, S. Irle, D. G. Musaev, R. Prabhakar and K. Morokuma, *Theory Comput.*, 2007, **3**, 1349.

41 B. Hourahine, S. Sanna, B. Aradi, C. Köhler, Th. Niehaus and T. Frauenheim, *J. Phys. Chem. A*, 2007, **111**, 5671.

42 H. C. Andersen, *J. Chem. Phys.*, 1980, **72**, 2384.

43 H. Hu, Z. Lu, M. Elstner, J. Hermans and W. Yang, *J. Phys. Chem. A*, 2007, **111**, 5685–5691.

44 H. Lin and D. G. Truhlar, *Theor. Chem. Acc.*, 2007, **117**, 185.

45 W. Han, M. Elstner, K. J. Jalkanen, T. Frauenheim and S. Suhai, *Int. J. Quantum Chem.*, 2000, **78**, 459–479.

46 A. H. Larsen, J. J. Mortensen, J. Blomqvist, I. E. Castelli, R. Christensen, M. Dułak, J. Friis, M. N. Groves, B. Hammer, C. Hargus, E. D. Hermes, P. C. Jennings, P. B. Jensen, J. Kermode, J. R. Kitchin, E. L. Kolsbørg, J. Kubal, K. Kaasbørg, S. Lysgaard, J. B. Maronsson, T. Maxson, T. Olsen, L. Pastewka, A. Peterson, C. Rostgaard, J. Schiøtz, O. Schütt, M. Strange, K. S. Thygesen, T. Vegge, L. Vilhelmsen, M. Walter, Z. Zeng and K. W. Jacobsen, *J. Phys.: Condens. Matter*, 2017, **29**, 273002.

47 A. O. Dohn, E. O. Jónsson, G. Levi, J. J. Mortensen, O. Lopez-Acevedo, K. S. Thygesen, K. W. Jacobsen, J. Ulstrup, N. E. Henriksen and K. B. Møller, *J. Chem. Theory Comput.*, 2017, **13**, 6010.

48 D. A. Case, R. M. Betz, D. S. Cerutti, T. E. Cheatham, III, T. A. Darden, R. E. Duke, T. J. Giese, H. Gohlke, A. W. Goetz, N. Homeyer, S. Izadi, P. Janowski, J. Kaus, A. Kovalenko, T. S. Lee, S. LeGrand, P. Li, C. Lin, T. Luchko, R. Luo, B. Madej, D. Mermelstein, K. M. Merz, G. Monard, H. Nguyen, H. T. Nguyen, I. Omelyan, A. Onufriev, D. R. Roe, A. Roitberg, C. Sagué, C. L. Simmerling, W. M. Botello-Smith, J. Swails, R. C. Walker, J. Wang, R. M. Wolf, X. Wu, L. Xiao and P. A. Kollman, *AMBER 2016*, University of California, San Francisco, 2016.

49 F. Paesani, W. Zhang, D. A. Case, T. E. Cheatham and G. A. Voth, *J. Chem. Phys.*, 2006, **125**, 184507.

50 L. Zhao, L. Liu and H. Sun, *J. Phys. Chem. C*, 2007, **111**, 10610–10617.

51 A. L. Parrill and K. B. Lipkowitz, *Reviews in Computational Chemistry, Volume 29*, John Wiley & Sons, Inc, Hoboken, NJ, 2016.

52 L. Martínez, R. Andrade, E. G. Birgin and J. M. Martínez, *J. Comput. Chem.*, 2009, **30**, 2157.

53 L. Verlet, *Phys. Rev.*, 1967, **159**, 98.

54 H. Sun, *J. Phys. Chem. B*, 1998, **102**, 7338–7364.

55 M. Waldman and A. T. Hagler, *J. Comput. Chem.*, 1993, **14**, 1077–1084.



56 S. Plimpton, *J. Comput. Phys.*, 1995, **117**, 1–19.

57 H. Heinz, T. J. Lin, R. Kishore Mishra and F. S. Emami, *Langmuir*, 2013, **29**, 1754.

58 P. Siani, E. Bianchetti, H. Liu and C. Di Valentin, *J. Chem. Phys.*, 2021, **154**, 034702.

59 R. W. Hockney and J. W. Eastwood, *Computer Simulation Using Particles*, CRC Press, 1988.

60 P. Dementyev, K. Dostert, F. Ivars-Barceló, C. P. O'Brien, F. Mirabella, S. Schauermann, X. Li, J. Paier, J. Sauer and H. Freund, *Angew. Chem., Int. Ed.*, 2015, **54**, 13942–13946.

61 H. Zhang, B. Gilbert, F. Huang and J. F. Banfield, *Nature*, 2003, **424**, 1025–1029.

62 D. Spagnoli, B. Gilbert, G. A. Waychunas and J. F. Banfield, *Geochim. Cosmochim. Acta*, 2009, **73**, 4023–4033.

