



Emerging investigator series: Curvature-induced hydrophobicity at imogolite-water interfaces

Journal:	Environmental Science: Nano
Manuscript ID	EN-ART-03-2020-000304.R2
Article Type:	Paper



Environmental Significance Statement

The wettability of mineral surfaces is a property controlling many environmental processes such us water transport in soils and geologic systems, or adsorption processes at mineral-water interfaces. Hygroscopicity of a mineral surface is often determined experimentally and interpreted by macroscopic approaches. Here, a microscopic view of two similar systems from the chemical point of view but with contrasting structures, show that the curvature of imogolite nanotubes induces a decreased hygroscopicity of its external convex surface with respect to the flat surface of gibbsite. This 'nanotube effect' could be relevant to other nanotubular systems with high curvatures, impacting their wettability properties, their colloidal stability and potentially enabling the formation of organo-nanotube complexes via hydrophobic interactions.

Journal Name

ARTICLE

Received 00th January 20xx, Accepted 00th January 20xx

DOI: 10.1039/x0xx00000x

www.rsc.org/

Emerging investigator series: Curvature-induced hydrophobicity at imogolite-water interfaces

Alejandro Fernandez-Martinez,*^a Jinhui Tao, ^b Adam F. Wallace, ^{*c} Ian C. Bourg, ^{*d} Mark R. Johnson, ^e James J. De Yoreo,^{b,f} Garrison Sposito, ^g Gabriel J. Cuello, ^e and Laurent Charlet ^a

Imogolite, a nanotubular aluminosilicate mineral, is commonly found in volcanic soils, where it exerts a control on carbon dynamics. Synthetic imogolites are used for the removal of contaminants from industrial effluents and are considered for a range of other applications including gas adsorption and functionalised heterogeneous catalysts. In spite of their environmental and industrial relevance, the properties of imogolite-water interfaces remain poorly understood. Here, an experimental and computational study is presented in which the structure and energetics of water are characterized on the curved external surface of imogolite and the hydrophilicity of this surface is contrasted with that of gibbsite, its planar counterpart. Atomic force spectroscopy experiments show that in spite of their identical surface structure, imogolite has a lower hygroscopicity than gibbsite. Molecular dynamics simulations provide an explanation for this observation: the curvature of imogolite prevents the formation of in-plane H-bonds along the directions of the nanotube circumference, lowering the enthalpy of adsorption of water molecules. The different arrangement of surface H-bonds and the resulting differences in hydration properties also affects the acidity constants of surface hydroxyl groups. This 'nanotube effect' may be relevant to other nanotubular systems with high curvatures, potentially impacting their wetting properties, their colloidal stability and their affinity towards hydrophobic organic moieties.

In spite of the importance of the phenomena outlined above,

the fundaments underlying the stability of adsorbed water on

mineral surfaces remain incompletely understood. One reason

for this lack of clarity is the scarcity of studies comparing the

atomistic-level structure and energetics of water on different

mineral surfaces. Studies based on interferometry,⁵

ellipsometry,⁶ and gravimetric adsorption measurements⁷ have

established a relatively extensive dataset on the adsorption of

water on a range of mineral surfaces, but these techniques

provide only a macroscopic, averaged view of the films.

Conversely, studies based on infrared spectroscopy,⁸⁻¹⁰

frequency-modulation atomic force microscopy,¹¹ or atomistic

simulations [i.e., molecular dynamics (MD),12 Monte Carlo

(MC),¹³ or density functional theory (DFT) simulations] have

produced detailed nanoscale views of adsorbed water on

mineral surfaces but have focused on a relatively small number

of surfaces, most predominantly the basal surface of muscovite

To date, only a few studies have compared water adsorption on

a range of different mineral surfaces using techniques that

provide atomistic-level information. Wang et al. (2006)¹⁴ used

MD simulations to compare the enthalpy and microstructural

details of adsorbed water films on the basal surfaces of five

Introduction

The affinity of water for mineral surfaces controls many fundamental processes in nature at different spatial and temporal scales. Examples include the multiphase flow of water and non-aqueous fluids (such as air, hydrocarbons, or supercritical CO₂) in geologic systems,¹ the nucleation and growth of biominerals in living organisms,² the formation of mineral-organic aggregates in soils,³ and the uptake of water vapour by mineral dust aerosols.⁴ In these situations and others, observed macroscopic-scale properties are fundamentally grounded in the hygroscopicity of the mineral surfaces and, in particular, in the affinity adsorbed water molecules and thin films for the mineral-air interface.

- ^{a.} Université Grenoble Alpes, Université Savoie Mont Blanc, CNRS, IRD, IFSTTAR, ISTerre, 38000 Grenoble, France
- ^{b.} Physical and Computational Sciences Directorate, Pacific Northwest National Laboratory, Richland, Washington 99352, United States
- ^c Department of Earth Sciences, University of Delaware, Newark, DE 19716, Unites States

^d Department of Civil and Environmental Engineering and Princeton Environmental Institute, Princeton University, Princeton, New Jersey 08544, United States

^{e.} Institut Laue Langevin, 71 Rue de Martyrs, 38000 Grenoble, France

^{f.} Department of Materials Science and Engineering, University of Washington, Seattle, Washington 98195, United States

- ⁹ Department of Environmental Science, Policy, and Management, University of
 - California, Berkeley, California 94720, United States
- Electronic Supplementary Information (ESI) available: [details of any supplementary information available should be included here]. See DOI: 10.1039/x0xx00000x

J. Name., 2013, 00, 1-3 | 1

mica.

60

ROYAL SOC OF CHEMIS

3

4

5

6

7

8

9

ARTICLE

adsorption on the basal surfaces of three uncharged layered minerals with a subtle structural difference: the presence and orientation of a hydroxyl group buried ~3 Å within the mineral (pyrophyllite, talc, and fluorotalc). These two studies of adsorption on minerals, combined with insights from studies of water adsorption on other surfaces such as metals¹⁶ or carbon nanotubes,¹⁷ reveal that mineral hygroscopicity is determined by multiple factors including the presence and localization of 10 surface charge sites and associated adsorbed counterions (that 11 can act as binding centres for water molecules), the presence 12 and density of hydroxyl groups (that can donate hydrogen 13 bonds to adsorbed water molecules), and the spatial 14 distribution of surface hydroxyls at the mineral-water interface 15 (that can facilitate or hinder interactions between adsorbed 16 water molecules). 17

18 A second reason for the lack of clarity outlined above is that 19 adsorbed water films are inherently two-dimensional phases: 20 their scale is atomistic in the direction normal to the surface but 21 macroscopic in other directions. Therefore, their stability is 22 sensitive to both specific short-range interactions with the 23 surface and collective long-range phenomena such the ability of 24 water molecules to collectively solvate adsorbed counterions 25 and capillary effects due to surface curvature and interfacial 26 energies. This sensitivity to both specific short-range and 27 collective long-range interactions is confirmed by recent 28 studies. However, it is not captured by the predominant 29 theoretical frameworks used to describe adsorbed water films, 30 as these assume the predominance either of short-range 31 interactions with the mineral surface (as in the case of the well-32 known Brunauer-Emmett-Teller theory) or of long-range 33 collective effects (as in the case of the Frenkel-Halsey-Hill, 34 Derjaguin-Landau-Verwey-Overbeek, and Köhler theories).4,7 35

In the present study, we approach the challenges outlined 36 37 above by characterizing and contrasting the hygroscopicities of two natural mineral surfaces: the external surface of imogolite 38 nanotubes and the basal surface of gibbsite nanosheets. These 39 two minerals present isostructural oxyhydroxide surfaces with 40 one key difference: the imogolite external surface is convex, 41 whereas the gibbsite basal surface is planar (Fig. 1). To our 42 knowledge, such a comparative examination of two surfaces of 43 unequal curvature has only been used in one previous study of 44 water adsorption (on a concave hydrophobic surface) which 45 showed results consistent with continuum scale descriptions 46 based on the Kelvin equation.¹⁸ It has not yet been applied to 47 either a convex or a hydrophilic surface and should provide 48 insight into the relative importance of short- vs. long-range 49 effects in controlling film stability. Our approach relies on a 50 combination of experimental and atomistic simulation 51 techniques that provide complementary atomistic-level views 52 of water adsorption, namely MD simulations and atomic force 53 spectroscopy measurements. We note that our study does not 54 directly examine macroscopic manifestations of hydrophilicity 55 such as the contact angles formed by water droplets (which 56 have a complex relationship to the microscopic properties of 57 interfacial water)¹⁵ although it does establish a basis for future 58 studies of these properties. 59

Beyond their utility to help resolve the fundamental questions outlined above, the systems investigated here have important implications in soil science (where gibbsite and imogolite are among the most important minerals in highly weathered soils such as boreal rainforest and volcanic ash soils)¹⁹ and in materials science (for example, in the synthesis of nanocomposite materials based on imogolite-like nanotubes).20,21

Materials and methods

Imogolite and Gibbsite

Imogolite (Al₂(OH)₃SiO₃OH) is an aluminosilicate mineral commonly found in the clay fractions of volcanic ash soils and spodosols.^{19,22–26} It has a high specific surface area (> 400 $m^2/g)^{27}$ and is known to strongly impact the fate and transfer of metals^{28–30} and natural organic matter in the soils in which it is found.³¹ It has a nanotubular geometry, with an internal face dominated by silanol (>SiOH⁰) functional groups and an external face composed by a curved gibbsite-like sheet (Al(OH)₃) with neutral aluminol functional groups (>Al₂OH⁰) (Fig. 1).^{32,33} The nanotubes have internal and external diameters on the order of 1 nm and 2.2 nm, respectively, and lengths of up to ~1000 nm.^{34,35} The presence of amphoteric surface sites and the lack of isomorphic substitutions make imogolite a mineral with a variable (pH-dependent) charge,³⁶ although some studies have reported that the external surface may carry a permanent negative charge resulting either from vacancies in the crystal structure or from a curvature-induced undersaturation of surface O atoms.^{27,37,38}

Gibbsite (Al(OH)₃) is a planar mineral present in bauxitic deposits and widespread in weathered soils. Its structure is composed of H-bonded dioctahedral layers, with the same stoichiometry as the external layer of imogolite.



Figure 1 Atomistic views of imogolite and gibbsite. The external surface of imogolite has a curved gibbsite-like (001) structure. The white spheres are H atoms in surface hydroxyl groups, which are drawn perpendicular to the surface in this representation. The orientations of gibbsite OH surface groups are the same as in bulk gibbsite as reported by Balan et al. (2006)³⁹.

Computational studies

Models of the gibbsite-water interface were constructed by cleaving a gibbsite structure along the (001) plane⁴⁰ to create a system containing two gibbsite layers in a $17.36 \times 15.23 \times 75$ Å $(a \times b \times c)$ simulation cell. For imogolite, a single nanotube was constructed in a 50 Å \times 50 Å \times 25.29 Å simulation cell. The

Journal Name

nanotube contained ten molecular $Al_2(OH)_3SiO_3OH$ units along the circumference, based on the reported structure of natural imogolite, with the 'zig-zag' configuration.^{32,33} A simulation cell of bulk liquid water was also simulated containing 256 water molecules (with no mineral). In total, the models of gibbsite and imogolite contained 72 and 180 surface OH groups, respectively.

The ClayFF model was used in all MD simulations. ClayFF is an interatomic potential model whose Hamiltonian includes Coulomb interactions between fixed-charge atoms and a 6-12 Lennard-Jones representation of Van der Waals interactions, a bond stretching term for surface and water OH bonds, and a three-body angular term to reproduce water angle bending modes.⁴¹ Gibbsite was one of the minerals used to parametrize ClayFF, and thus its structure is reproduced with high accuracy: the difference with the experimental lattice constant being less than 4.1%.¹⁴ The structure of imogolite is also reproduced with high fidelity.

The ClayFF model was originally developed for use with the SPC water model. This combination of models has been extensively used to evaluate water structure and properties and for the interaction of water with hydroxide mineral surfaces, including imogolite.^{14,41–45} In addition, the ClayFF model, when used with the slightly different SPC/E water model, has been repeatedly reported to give excellent predictions of the properties of water on mineral surfaces.^{46,47} Relative to the SPC/E model, the original SPC model produces somewhat less accurate prediction of water structure and dynamics,⁴³ but a more accurate prediction of the enthalpy and free energy of vaporization of liquid water.⁴⁸ As shown below, the SPC model successfully reproduces the two adsorption sites for water molecules at the gibbsite (001) surface reported by Veilly et al. (2008) using density functional theory methods.⁴⁹

Molecular dynamics simulations were performed in the canonical ensemble (*NVT*) using the program LAMMPS.⁵⁰ The structures of the dry surface models were relaxed during 0.5 ns equilibration runs. Each hydrated system was then prepared by loading the equilibrated dry surface model with a specified number of randomly placed water molecules. Following surface loading, the mineral structure was held stationary as the solvent was relaxed for 0.5 ns, after which the entire hydrated structure (mineral and solvent) was relaxed for an additional 1.0 ns. Simulations were performed on the NERSC supercomputer Franklin and on the University of Delaware's Caviness Cluster. Analyses of the trajectories were performed using custommade codes written in Fortran 95 and Python.

Helmholtz free energies of hydration (Δ F) were calculated from the enthalpy (Δ H) and entropic (Δ S) changes associated with water adsorption:

$$\Delta F(\phi) = \Delta H(\phi) - T\Delta S(\phi)$$
 eq. (1)

where the surface coverage, ϕ , is defined as the total number of water molecules normalized to the number of O atoms on the hydrated surface (e.g., the external surface of imogolite). Reported potential energies and entropies of hydration were derived using the following expression:⁵¹

$$\Delta X(\mathbf{\phi}) = \frac{\langle X(N) \rangle - \langle X(0) \rangle}{N} - X_b \{X = H, S\} \quad \text{eq. (2)}$$

where X(N) is the potential energy or entropy of the wet system, containing N water molecules, X(0) is the potential energy or entropy of the dry system and X_b is the potential energy or entropy of a water molecule in bulk water. The resulting values correspond to the free energy, enthalpy, and entropy of reaction associated with transferring ϕ monolayers of water from bulk liquid water to the dry mineral surface, normalized to the number of transferred water molecules. As reported by Smith (1998),⁵¹ the value of the enthalpic component can be approximated from the ensemble average of the system potential energy, which was computed here from 2.0 ns production runs. Addition of the kinetic energy to the enthalpic component does not result in significantly different results. Entropies were derived by analysing the vibrational density of states within the framework of the Two-Phase Thermodynamic Method (2PT)⁵² as implemented in an in-house code. The total system entropy was taken as the sum of the mineral and solvent entropies. For the solvent component, the translational, rotational, and internal vibrational states of each water molecule were considered in the determination of the 2PT entropy.⁵³ The entropy of the mineral component was derived by summing the translational entropies of each atom type contained within the mineral structure. Values of the specific free energy, enthalpy, and entropy of bulk water were calculated from a simulation cell containing 256 water molecules (with no mineral).

In addition to the MD simulations described above, DFT geometry optimizations of the hydrated structures (with one water layer) of imogolite and gibbsite were performed using the Vienna Ab-initio Simulation Package (VASP).54,55 The imogolite model used in these ab-initio calculations consisted of one imogolite unit cell with dimensions 35 Å \times 35 Å \times 8.43 Å. A vacuum gap of 15 Å was placed between the periodical images along the *a* and *b* directions, the tube being periodic along the c direction. A supercell of gibbsite containing 2×3 unit cells along the a and b directions (17.37 Å \times 15.23 Å \times 75 Å) and two layers of gibbsite (001) planes separated by a vacuum layer of 67 Å were used. The calculations were performed at the Gamma point (k = 0), using projector augmented wave (PAW) pseudopotentials (Kresse and Joubert, 1999) with a plane wave cutoff of 209 eV and the Perdew-Burke-Ernzerhof (PBE) functional of the generalized gradient approximation (GGA).56,57

Experimental sample description, synthesis, and preparation

A sample of gibbsite $[Al(OH)_3)]$ powder was kindly provided by Alcoa Corp (Aviles, Spain). Synthetic aluminosilicate imogolitelike nanotubes $[Al_2(OH)_3SiO_3OH]$ were synthesized following the procedure described by Denaix et al. (1999).⁵⁸ Aluminum nitrate (99.999%, Alfa Aesar) and Tetraethyl orthosilicate (99.999%, Alfa Aesar) were mixed in ultrapure water at an Al/Si ratio of 2 and aluminium concentration of 0.002 M. Dilute NaOH (0.1 M) was injected at a rate of 2 mL/min under vigorous stirring until

53

54

55

56

57

58

3

4

5

6

7

8

9

10

11

29

60

ARTICLE

reaching a hydrolysis ratio OH/Al of 1. After injection, the solution was stirred for 12 h to avoid the polymerization of Al hydroxides. The solutions were heated at 95°C during five days in closed Teflon bottles. To remove excess alcohol and Na⁺ and NO₃⁻ ions from solution, the solutions were dialyzed against ultrapure water for 15 days using a molecular weight cut-off membrane of 2000 Da (Spectra/Por[®] membranes, Carl Roth). After dialysis, part of the imogolite solution was freeze-dried and used for the experiments.

12 The resulting powder was characterized using transmission 13 electron microscopy (TEM) and Fourier transform infrared 14 spectroscopy (FTIR) and showed good agreement with the 15 expected features of synthetic imogolite (see Supporting 16 Information Figures S1 and S2). TEM investigations were carried 17 out on a Philips CM 200 microscope operating at 200 kV. Prior 18 to observation, the powders were immersed in an aqueous 19 chloroform solution. A droplet of this solution was deposited on 20 a copper grid. TEM observations were made after evaporation 21 of the solvent. FTIR spectra of the imogolite freeze-dried 22 powders were taken to characterize the product obtained after 23 the synthesis. Transmission FTIR spectra were obtained using a 24 Bruker HYPERION 3000 FTIR microscope in transmission mode 25 on KBr pellets (1 mg of imogolite in 100 mg of KBr). One hundred 26 scans taken using a resolution of 2 cm⁻¹ were averaged to obtain 27 spectra in the range 4000–370 cm⁻¹. 28

Force spectroscopy

30 Force spectroscopy measurements were performed using SiOH-31 terminated tips under ultrapure water to directly probe the 32 interaction between SiOH and the interfacial water or solid 33 surface. Based on previous studies, the free energy measured 34 by DFS probes the sum of SiOH-interfacial water hydrogen bond 35 strength decrease and interfacial water entropy increase during 36 H-bond rupture by pulling the silicon tip away from the gibbsite 37 or imogolite surface.^{59,60} Although we do not expect to directly 38 measure the free energy of water adsorption on these two 39 minerals due to the chemical difference between SiOH and 40 HOH, this method gives the most straightforward way to 41 evaluate the relative affinity between water and mineral with 42 surface specificity, which is proportional to hydrophilicity. The 43 main assumption made here is thus that the interaction of a 44 hydroxylated AFM tip with an oxide surface is a reasonable first-45 order proxy for the attachment of a water molecule at the 46 surface. $^{\rm 59,61-65}$ To that end, aliquots of synthetic imogolite and 47 gibbsite suspensions were deposited onto muscovite mica disks 48 and dried under N_2 . The disks were mounted on a fluid cell for 49 analyses. Tip functionalization (hydroxylation) was performed 50 by plasma cleaning of Si tips on Si₃N₄ AFM cantilevers (Bruker, 51 SNL-10, CA) for 2 min (see Fig. 1a). Measurements were made 52 with the MFP-3D Atomic Force Microscope (Asylum Research, 53 Santa Barbara, CA). To account for any surface heterogeneity, a 54 custom routine randomly sampled points on the surface to give 55 a representative average. A constant approach velocity of 1.04 56 μ m/s was used for every pulling speed studied. An 8 nm 57 deflection trigger was used to contact the surface and dwell for 58 1 s before pulling away. The AFM tip was vertically retracted 59

from the substrate at five different pulling rates: 651 nm/s, 1040 nm/s, 1740 nm/s, 2600 nm/s and 3910 nm/s. The value of the cantilever spring constant was determined by deflection sensitivity and thermal fluctuation measurements.⁶⁶ The collected force versus separation curves typically exhibited a characteristic single rupture event. Curves showing more than one bond breaking event were discarded. We cannot exclude that multiple bond-breaking events are confounded within a single curve. This remains a limitation of the technique.

The approximate analytical expression for the mean rupture force for a single bond is: 60,67,68

$$\langle f \rangle = \frac{\Delta G}{\nu x_t} \left\{ 1 - \left[\frac{k_B T}{\Delta G} ln \frac{k_{u}^{\delta G} + \gamma}{\frac{k_{u} e^{k_B T} + \gamma}{\frac{k_{u} e f}{k_B T dt}}} \right]^{\nu} \right\} \quad \text{eq. (3)}$$

where $\langle f \rangle$ is the mean rupture force for a single bond at any loading rate; ΔG is the equilibrium free energy of binding $\Delta G =$ $G_u - G_b$ of the bonded system, G_b , relative to the free cantilever G_u ; x_t is the distance between the free-energy minimum and the transition state; ν depends on the shape of the potential energy well ($\nu = 2/3$ for a linear-cubic potential, $\nu = 1/2$ for a harmonic potential, and $\nu = 1$ recovers the phenomenological Bell theory); k_B is the Boltzmann constant; T is absolute temperature; k_u^0 is the intrinsic rate coefficient for the system in absence of applied force; $\gamma = 0.577$ is the Euler-Mascheroni constant; and df/dt is the loading rate. The energy ΔG of the tip-substrate interaction can be calculated from data on $\langle f \rangle$ vs. df/dt.

Results and discussion

Hydration energies: force spectroscopy and molecular dynamics

Fig 2 shows results of the force spectroscopy experiments. Fig. 2b shows the rupture force between the hydroxylated Si (SiOH) tip and the gibbsite or imogolite surface as a function of loading rate. The solid curve in Fig. 2 is a fit to the data according to the theoretical dependence of the bond rupture force on the loading rate (eq. 3). Equilibrium free energies of binding for the SiOH-gibbsite (001) and SiOH-imogolite systems are reported in Table 1. The energy range of the bonds formed between the hydroxylated tip and the surface falls within the expected range for water H-bonds.^{69,70} The Δ G value for imogolite is larger than that for gibbsite by 4.41 k_BT (10.92 kJ/mol), which implies that the gibbsite (001) face forms more stable interactions with SiOH groups (and hence potentially also with water molecules). Representative examples of force vs. distance curves are shown in Fig. S3.

In order to understand the origin of this difference between the two systems, a series of molecular dynamics simulations were performed with different levels of hydration. Values of the average free energy of hydration per water molecule are shown in Fig. 3. All the values of the average free energy of hydration are negative, indicating that the dry mineral surface is more hygroscopic than bulk liquid water. In other words, both surfaces should carry an adsorbed water film when exposed to ambient water-saturated air. As expected, all curves approach zero (i.e., the energetics of water in the film become more bulk-

Journal Name

liquid-like) with increasing film thickness. The lower affinity of water for the curved imogolite surface as compared to the planar gibbsite surface is confirmed by the simulations: almost all the points of the free energy curves, in particular those at low surface coverages, are above the values for gibbsite. Interestingly, a minimum value is observed for both surfaces in the average enthalpy of hydration, but this minimum is positioned at different ϕ values for imogolite ($\phi \sim 1$) and gibbsite (φ ~ 0.6). The origin of these differences is further explored below through detailed analyses of the interactions between surface hydroxyl groups and interfacial water molecules (Fig. 4). We note here that the analysis of the adsorption energies provided above reflect the stability of the entire adsorbed water films (i.e., reported energetics are a collective property emerging from the network of H-bonds at the interfaces). Analyses of adsorption energies of individual water molecules are beyond the scope of this manuscript.

Table 1 Fitted parameters of the interaction model between the SiOH tip and the
gibbsite (001) and imogolite surfaces. Results were obtained using the harmonic
potential approximation to the surface-tip interaction (i.e., v = 1/2 in Eq. 4).

Surface	Gibbsite (001)	Imogolite external
$k_u^0 (s^{-1})$	$\textbf{3.78} \pm \textbf{1.43}$	$\textbf{74.43} \pm \textbf{13.37}$
$x_t (nm)$	$\textbf{0.13}\pm\textbf{0.02}$	$\textbf{0.10}\pm\textbf{0.02}$
$\Delta G (k_B T)$	$\textbf{-7.88} \pm \textbf{0.14}$	$\textbf{-3.47}\pm0.27$



Figure 2 Determination of the equilibrium Gibbs energy of binding of the SiOH tip to the gibbsite and imogolite surfaces. (a) Schematic view illustrating the SiOH tip placed directly on the solid surface. (b) Dynamic force spectra for the rupture of bonds between the tip and the gibbsite (001) or imogolite surfaces (spring constant of 43.30 pN/nm). The solid curves are fit to the data using Eq. 3. (c-d) Histograms of bond rupture forces measured from repeated force–distance trajectories for gibbsite (c) and imogolite (d). Normalized histograms for increasing loading rate (28.1, 45.0, 75.3, 112.6 and 169.3 nN/s) are offset for clarity. The vertical range is 50% for (c) and 60% for (d).



Figure 3 Hydration free energy of imogolite and gibbsite (blue circles and orange squares) as a function of surface coverage (left panel) and its breakdown into enthalpic and entropic components (center and right panels). The results quantify the average (per water molecule) energy associated with transferring ϕ water monolayers from bulk liquid water to the dry mineral surface. The results show that enthalpy favours liquid water uptake on the dry mineral surface, whereas entropic effects inhibit this uptake.





ARTICLE

Figure 4 Snapshots of the MD simulations showing H-bonded water molecules in both systems at monolayer coverage (φ=1). Left: view of the gibbsite-water interface, showing two water molecules (highlighted in yellow) giving (left) and receiving (right) a H-bond to/from surface hydroxyls. Right: view of the imogolite-water interface with two water molecules (yellow) receiving (left) and giving (right) H-bonds from/to surface hydroxyls.

Orientation of surface hydroxyls. The projections of the O and H atoms from surface hydroxyl groups onto the (001) surface of gibbsite and onto the external surface of imogolite are shown in Fig. 5, together with histograms of the angle β formed between the **OH** vector of each hydroxyl and the vector normal to the surface for the dry (ϕ =0) and hydrated surfaces (ϕ =4). Histograms for other hydration states are shown in Fig. S4.



Figure 5 Top: Projections of surface OH groups onto the (001) gibbsite surface (left) and the external imogolite surface (right). The orange circles represent the positions of surface O atoms. The blue regions indicate the more probable positions of hydroxyl H atoms. Red arrows indicate OH groups perpendicular to the surface. Green arrows indicate OH groups parallel to the surface. The position of the empty octahedral site is placed between three O atoms, at the end of the green arrows in the left figure. Bottom: histograms showing the distribution of the β angle formed between the OH vector and the normal to the surface for ϕ =0 (left) and ϕ =4 (right).

The histograms show a strong effect of water on the orientations of the hydroxyl groups. The presence of adsorbed water induces an orientation of the surface hydroxyls which is qualitatively similar on both surfaces. The histogram of the β angle for ϕ =4 shows two peaks placed at ~20° and ~85°, i.e., almost 'normal' and almost 'parallel' to the mineral surfaces. A difference between the two surfaces is observed, with a significant higher number of 'normal' hydroxyls in the case of imogolite. The projections in Fig. 5 show differences in the orientations along the xy directions (plane of the mineral surfaces): For gibbsite, the OH groups tend to form H-bonds with water or with neighbouring O atoms (towards the empty octahedral site characteristic of its dioctahedral surface structure). On the external imogolite surface, however, the curvature of the nanotube prevents the formation of H-bonds between surface OH groups along its circumference. Thus, only ~1/3 of the OHs are parallel to the surface, with their direction lying along the c axis, where there is no curvature.

Structure of adsorbed water at the mineral-water interface. The normalized density profiles of water computed at the gibbsiteand imogolite-water interfaces are shown in Fig. S5. A similar profile, with minima and maxima positioned at similar distances, is observed on both surfaces, indicating that the two surfaces exert similar influence over the structure of adsorbed water. A difference in the width of the first water peak, broader in the case of imogolite, indicates a less ordered first layer, as expected for a less hydrophilic surface.⁷¹ The simplest explanation for this difference is the geometric effect caused by the curvature of the imogolite surface, for example the impact of curvature on the orientation of surface OH groups noted above.

Examination of the angular distributions of water molecules at the surface (Figs. 6 and 7) further suggests that the orientation of surface OH groups strongly influences the properties of adsorbed water. In particular, our calculations reveal the existence of two conformations of adsorbed water on both surfaces. Two angles are needed to describe the orientation of a water molecule. The angle θ is defined between the water dipole vector and the vector normal to the mineral surface, whereas α is the angle between the vector joining the two water hydrogen atoms and a vector normal to the mineral surface. The occurrence of two adsorbed water conformations is most obvious in the distribution of $\boldsymbol{\theta}$ angles at low water coverage, where two peaks can be distinguished in the first water monolayer (d \approx 2.5 Å) of all the histograms: one centred at $\theta \approx 122^{\circ}$ and the other at $\theta \approx 75^{\circ}$. In the case of gibbsite, the peak position of the more acute angle shifts towards lower values as the water coverage increases. The distribution of the α angle is similarly bimodal, particularly at low water coverage, as shown in Fig. 7.

Further examination of the simulation trajectories in light of the angular distributions noted above shows that the first orientation (type A: $\theta \approx 120^\circ$, $\alpha \approx 45^\circ$), corresponds to a water molecule with an OH pointing towards the surface and donating a hydrogen bond to a surface oxygen. The second orientation (type B: $\theta \approx 55^{\circ}$, $\alpha \approx 90^{\circ}$) corresponds to a water molecule that accepts a hydrogen bond from the surface (Fig. 4). These two types of adsorbed water were described by Wang et al. (2006) and Veilly et al. (2008) in their investigations of water adsorption on gibbsite.14,49 Orientation B was shown by Veilly et al. (2008) to be stabilized when two hydroxyl groups participate in the binding. However, although these authors mention the strong hydrogen bonding of orientation A, no comparison is made with the B orientation. On gibbsite, the intensity corresponding to the A orientation ($\theta \approx 120^\circ$) does not increase with increasing water coverage from ϕ = 0.5 to 1.0, meaning that additional first-layer water molecules adsorb on the surface with the B orientation. On imogolite, the same water

orientations are observed,⁴³ but A orientation does not become saturated until $\phi = 1.0$ as shown in the evolution of the peak at $\theta \approx 120^{\circ}$ between $\phi = 0.5$ to $\phi = 1$ in Fig. 6. A previous study reported the occurrence of A type water on imogolite, but did not discuss the B type.⁴³ Instead, a third type was described in which the water dipole vector points away from the surface, and that may occur in positive charged planar regions of the imogolite surface as predicted by the vibrational breathing mode.⁴³

Journal Name

The second layer of water molecules on the two mineral surfaces (d \approx 5 Å) show much less orientational preference than the first layer. On gibbsite, angular distribution maps reveal a minor preference for water molecules orienting one of their OH groups towards the surface ($\theta \approx 130^\circ$, $\alpha \approx 90^\circ$). Again, the effect

of curvature is seen in the angular distribution of water molecules in the second hydration shell of imogolite, which is less structured than that of gibbsite.

Water H-bonds with the surface. An analysis of the hydrogen bonding of surface OH groups with water molecules provides further insight into the mechanisms of water adsorption. Histograms of the number of H-bonds donated or accepted by surface OH groups to/from water molecules are shown in Fig. 8. The criteria used to define a hydrogen bond are the following: a water molecule was considered H-bonded if it was located within the first peak of the O-H pair correlation function, which extends to 2.2 Å in the case of a receiving H-bond from water and to 2.3 Å for H-bonds donated to water, similarly to the distances found in a previous study on metal oxides.⁷²



Figure 6 Histograms of the θ orientations of water molecules at the surfaces of imogolite (top) and gibbsite (bottom) as a function of water coverage (numbers at the top). Blue color indicates a low probability of finding a certain orientation, and red represents the maximum probability.



4

5

6



Figure 8 Percentage of surface hydroxyl groups donating or accepting 1 hydrogen bond from water molecules, for both imogolite (blue circles) and gibbsite (orange squares).



Figure 9 Percentage of surface OH groups forming 0, 1 or 2 H-bonds with water molecules. About a 15% of the sites are able to form two H-bonds with water molecules, in comparison with a 2% in gibbsite. At any instant, a significant number of surface OH groups do not H-bond with water. Squares: gibbsite; Circles: imogolite.

Our results on the number of H-bonds donated and received by surface OH groups are consistent with the orientational distributions shown in Figs. 6-7: the number of hydrogen bond acceptors (water molecules of type A) becomes saturated at $\phi \approx 0.6$ on gibbsite and $\phi \approx 1.0$ on imogolite. On each surface, the water coverage where the type A water population saturates coincides with that where the enthalpy of hydration is most favourable (Fig. 3), suggesting that the surface hydroxyls that are oriented parallel to the surface (Fig. 5) may provide high

energy sites for the adsorption of type-A water molecules, and that the greater tendency of surface OH groups to lie parallel to the surface of gibbsite (particularly at low degrees of hydration) may cause the greater affinity of water molecules for this surface.

The percentage of surface hydroxyls forming 0, 1 or 2 H-bonds with water is shown in Fig. 9. It is interesting to note that, although most of the surface OH sites are forming H-bonds, all curves reach a plateau, and that a significant number of surface sites remain 'dry' even at high surface coverage values.

Effect of the curvature on the hydroxyl acidity. To gain additional insight into the impact of surface curvature on surface hydration, we used a bond-valence model approach to characterize the coordination of O atoms on hydrated gibbsite and imogolite surfaces. The bond valence model approach was initially developed to help explain the structures of crystalline minerals⁷³ and has been extended to help predict the acidity constants of surface OH groups, a key parameter influencing the reactivity of (oxyhydr)oxide surfaces including those examined here.^{74,75,76,77} For example, the well-known MUSIC model predicts the pK_a values of surface OH groups using the expression:⁷⁵

$$-pK_a = logK_a = -A(\sum_j s_j + V) \quad \text{eq. (4)}$$

where V is the valence of oxygen (-2), $\sum_{j} s_{j}$ is the sum of the bond valences s_i associated with all the bonds *j* reaching the oxygen atom, and A = 19.8 is a phenomenological constant. In the case of gibbsite and imogolite, the bond valence sum $\sum_i s_i$ for surface O atoms in >Al₂OH₀ functional groups includes the bond valences of the two Al-O bonds and the O-H covalent bond formed by a surface O atom, as well as the bond valence of all H-bonds received by the O atom. The MUSIC model uses a fixed value of s = 0.8 valence units (v.u.) for each covalent O-H bond and s = 0.2 v.u. for each H-bond accepted from a water molecule,75 such that each additional H-bond received by a surface O atom increases its acidity by 4 pH units. In previous applications of this model to imogolite and gibbsite, Hiemstra and co-workers concluded that >Al₂OH₀ functional groups on the gibbsite (001) surface have pK_a values of 0.0 and 11.9, i.e., they are not reactive in the pH range of environmental relevance,^{78,79} in agreement with other estimates.^{80,81} A previous study of imogolite assumed that the number of Hbonds received by surface O atoms is the same as on gibbsite, such that the acidity constants of both surfaces were predicted to be identical.27

Here, instead of assigning a fixed bond valence of 0.2 to each Hbond, we used an alternative approach whereby the bond valence associated with an individual bond decreases with the length of interatomic distance following the relation:⁷³ Journal Name

1 2

3

4

5

6

7

8

9

10

11

12

13

14

15

16

17

18

19

20

21 22

23

24

25

26

27

28

29

30

31

32

33

34

35

36

37

38

39

40

41

42

43

44

45

46

47

48

49

50

51

60

$$s_j = exp\left(\frac{r_0 - r_j}{B}\right)$$
 eq. (5)

where r_j is the bond length (including H-bonded water molecules) and r_0 and B are fitted parameters specific of each anion-cation pair. For O-H bonds, values of r_0 and B determined by Brown are shown in Table 2. Equation 5 yields a value of s_{O-H} that tends asymptotically to zero for large r_{O-H} values. This allows the bond valence sum $\sum_j s_j$ to be evaluated over all water molecules around a surface site, in contrast to the analysis presented in Figs. 8-9 (where the number of H-bonds was quantified using a distance cut-off value as a criterion). The inclusion of all the water molecules in the bond-valence sum was suggested by Brown (2002), who observed that interactions of water molecules in bulk water placed at long distances from each other should be considered if proper valence sums were to be maintained.^{82,83}

Table 2Parameters used to calculate the bond-valence of surface OH bonds, O...H and $O...H_2O$ hydrogen bonds, using eq. 5.

r _{о-н} (Å)	<i>r</i> ₀ (Å)	В
< 1.05	0.907	0.28
$1.05 \leq r_{O-H} \leq 1.70$	0.569	0.94
> 1.70	0.99	0.59

Bond valences corresponding to hydrogen bonds formed between surface oxygen atoms and water molecules or surface hydroxyls are reported in Fig. 10 as a function of the surface coverage. The differences between the two systems reflect the influence of the curvature. In Fig. 10, the blue symbols show the bond valence shared by a surface O atom with neighbouring structural H atoms: 0.88 v.u. for gibbsite and 0.83 v.u. for imogolite. This value equals the sum of the bond valence associated with the covalent O-H bond (0.72 v.u. according to Eq. (5) with the parameters from Table 2) and the bond valence associated with H-bonds between surface hydroxyls (0.16 v.u. for gibbsite, 0.11 v.u. for imogolite). In short, surface O atoms on the flat gibbsite surface receive approximated 50% more bond valence from intra-surface H-bonds than O atoms on the curved imogolite surface, in agreement with the greater fraction of OH groups pointing perpendicularly to the surface on imogolite vs. gibbsite.

As shown by the red symbols in Fig. 10, at water coverages greater than one monolayer, the greater tendency of surface O atoms on gibbsite (vs. imogolite) to receive H-bonds from other surface OH groups is partly compensated by their smaller tendency to receive H-bonds from water molecules. At submonolayer coverages, however, gibbsite surface O atoms share more bond valence than imogolite surface O atoms with both neighbouring OH groups and with adsorbed water molecules.

In addition to the degree of H-bonding of surface O atoms, 52 values of the protonation constants of the neutral >Al₂OH₀ sites 53 calculated by the MUSIC model are very sensitive to small 54 variations of the metal-oxygen bond lengths. In order to obtain 55 these protonation constants using Eq. (4), values for the Al-O 56 bond lengths need to be determined. Different experimental 57 techniques can be used to obtain this information (e.g., Pair 58 Distribution Function, X-ray Absorption Spectroscopy). 59

This journal is © The Royal Society of Chemistry 20xx



Figure 10 Bond valences of surface oxygen sites resulting from H-bonding with water molecules (red) and with other surface OH groups, plus direct OH bonding (blue). The top (balck) curve indicates the total bond valence received by surface O atoms. Squares: gibbsite; Circles: imogolite.

However, determining the local information about the bonds with surface oxygen atoms remains an experimental challenge. For these reasons, DFT calculations of gibbsite and imogolite–water interfaces were used to obtain an estimation of the average Al-O bond lengths. It is worth noting that the inplane H-bonding configuration along the axis of the imogolite tube observed for surface hydroxyls groups in classical MD simulations is reproduced by the more accurate DFT calculations (see Fig. S6). In Fig. 10, the Al-O bond lengths (r_{Al-O}) have been plotted as the independent variable, and the s_{Al-O} bond valences have been calculated using equation (5) with B = 0.37 and $r_0 = 1.651$ Å.

Both pK_{a1} values follow a trend towards lower values as the Al-O distances increase. The r_{Al-O} bond lengths obtained from DFT calculations give an estimation of the pK_{a1} for each surface. For imogolite we obtained an average value of $\langle r_{Al-O} \rangle = 1.95$ Å which yields a pK_{a1-imogolite} = -0.92 ± 0.24. The value obtained for the gibbsite groups is of $\langle r_{Al-O} \rangle = 1.94$ Å, yielding a pK_{a1-gibbsite} = -0.05 ± 0.50. The acidity constant for imogolite doubly coordinated neutral groups is shifted towards more acidic values by a $\Delta pK_{a1} = 0.87 \pm 0.75$. The value for gibbsite is in close agreement with the value reported by Hiemstra et al. (1996) using the CD-MUSIC model (pK_{a1} = 0.0), and in disagreement with other studies, which proposed values of pK_{a1} between 5.2 and 5.9.^{80,81}



3

4

5

6

7

8

9

10

11

12

13

14

15

16

17

18

19

20

21

22

23

ARTICLE

Figure 11 Acidity constants for the surface >Al₂OH⁰ functional groups calculated using Eq. (5) as a function of Al-O bond length. The scale in the top abscissa indicates the total bond valence due to the two Al-O bonds. The error bars (gray and pink areas) have been calculated from the uncertainties in the s_{0-H} bond valences. An estimation of the absolute acidity constant has been done using <r_{Al-O}> distances obtained from DFT-optimized structures.

Conclusions

The effect of curvature on surface wetting and acidity has been explored using two mineral surfaces with the same surface chemistry, the external curved surface of imogolite and the planar (001) surface of gibbsite. The difference in measured free energy values (7.88 k_BT and 3.47 k_BT) for gibbsite vs imogolite is consistent with our MD simulation predictions of the difference in hydrophilicity between the two minerals. The larger values obtained by AFM vs. MD simulation may reflect a formation of stronger H-bonds by SiOH vs. HOH or the existence of multiple bonds ruptures during the pulling of our AFM tips. The consistency of the experimental and simulation results give us confidence that this methodology is adequate to explore molecular interactions at the mineral-water interface.

24 A mechanistic explanation of the observed differences in 25 wettability is obtained from analyses of the MD results. These 26 reveal differences in the geometry of water molecules adsorbed 27 within the first hydration layer of the mineral-water interface: 28 whereas gibbsite is able to accept hydrogen bonds from water 29 molecules forming a highly structured first hydration layer (A-30 type), the curvature of imogolite prevents the formation of 31 some of the H-bonds between surface hydroxyl groups, making 32 the adsorption of water molecules less energetically favourable 33 at sub-monolayer coverages. The different stereo-chemical 34 conditions of water adsorbed on the two systems are further 35 translated into different acidities for surface >Al₂OH⁰ sites. The 36 results show more acidic pK_{a1} values for doubly-coordinated 37 neutral sites on imogolite than on gibbsite, by $\Delta p K_{a1} = 0.87 \pm$ 38 0.75. The absolute values of the protonation constants reported 39 in this work coincide with those reported by Hiemstra and Van 40 Riemsdijk (1996),⁷⁵ who used the same CD-MUSIC model. 41

42 The lower hygroscopicity observed for imogolite may be related 43 to several properties that have been classically described as 44 specific to this mineral. Of these, the formation of stable 45 organo-mineral complexes has been largely reported in the soil 46 science literature as a characteristic of soils where imogolite 47 (and other allophanic clays) are present.^{31,84–88} The formation of 48 organo-mineral complexes may be facilitated by the high 49 density of functional groups at the surface of imogolite. 50 However, prior to specific interactions with functional groups, 51 the organic molecule has to approach the imogolite surface. The 52 less hydrophilic character of imogolite would be beneficial for 53 this first step, as water is less structured on imogolite surfaces 54 than on other minerals.¹⁴ We note here however that the 55 complexity of soil organic matter and the multiple faces of 56 mineral exposed to the soil solution make organo-mineral 57 interactions complex.⁸⁹ The wetting properties studied here are 58 therefore only a part of the puzzle. Further investigations of this 59 property should be made in the future to correctly assess the 60

retention of organic matter by imogolite. Our simulations shed light on another important property of imogolite: its tendency to form bundles. The free energy profile of aggregation of two mineral particles has been shown to depend upon energy barriers related to highly ordered layers of water between the mineral surfaces.^{90,91} In the case of imogolite this shell of adsorbed water is less structured, and a lower energy barrier is expected to occur, thus potentially facilitating their aggregation.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

Barry Bickmore and Kideok K. Kwon are gratefully acknowledged for enriching discussions. Loïc Vidal (Inst. Sciences des Matériaux de Mulhouse, France) is thanked for TEM analyses. A.F.-M. acknowledges the Région Rhône-Alpes for the financial support received through an Explora'DOC fellowship for a 6-months stay at the University of California, Berkeley. 2PT analysis was performed by A.F.W. with support of the U.S. Department of Energy (DOE), Office of Science, Office of Basic Energy Sciences (BES), Chemical Sciences, Geosciences, & Biosciences (CSGB) Division, under Award Number DE-SC0018439. I.C.B. was supported by the U.S. Department of Energy, Office of Science, Office of Basic Energy Sciences, Geosciences Program under Award DE-SC0018419. Molecular dynamics simulations were performed using resources of the National Energy Research Scientific Computing Center (NERSC), which is supported by the U.S. Department of Energy, Office of Science, under Award DE-AC02-05CH11231.

References

- E. Shahraeeni and D. Or, Pore-scale analysis of evaporation and condensation dynamics in porous media, *Langmuir*, 2010, **26**, 13924–13936.
- S. Elhadj, J. J. De Yoreo, J. R. Hoyer and P. M. Dove, Role of molecular charge and hydrophilicity in regulating the kinetics of crystal growth, *Proc. Natl. Acad. Sci. U. S. A.*, 2006, **103**, 19237–19242.
- 3 C. J. Bronick and R. Lal, Soil structure and management: A review, *Geoderma*, 2005, **124**, 3–22.
 - M. Tang, D. J. Cziczo and V. H. Grassian, Interactions of Water with Mineral Dust Aerosol: Water Adsorption, Hygroscopicity, Cloud Condensation, and Ice Nucleation, *Chem. Rev.*, 2016, **116**, 4205–4259.
- 5 T. E. Balmer, H. K. Christenson, N. D. Spencer and M. Heuberger, The Effect of Surface Ions on Water Adsorption to Mica, *Langmuir*, 2008, **24**, 1566–1569.
 - D. Beaglehole and H. K. Christenson, Vapor adsorption on mica and silicon Entropy effects, layering and surface forces, *J. Phys. Chem.*, 1992, **96**, 3395–3403.
 - R. M. Pashley, Multilayer adsorption of water on silica: An

4

6

ו ר	3001110			
2		analysis of experimental results 1 Colloid Interface Sci	24	R
3		1980 78 246–248	24	ch
4 5	8	D B Asay A L Barnette and S H Kim Effects of surface		fr
5	0	chemistry on structure and thermodynamics of water		41
0		layers at solid-vapor interfaces I Phys Chem C 2009 113	25	<u>-</u> -
/		2128–2133	25	Δ
0	9	W Cantrell and G E Ewing Thin film water on muscovite		St
9 10	5	mica / Phys Chem B 2001 105 5434–5439		n
10	10	W Cheng K Hanna and L E Boily Water Vanor Binding on		18
11	10	Organic Matter-Coated Minerals Environ Sci Technol	26	Δ
12		2019 53 1252–1257	20	V
13	11	T. Arai, K. Sato, A. Jida and M. Tomitori, Ouasi-stabilized		Co
15		hydration layers on muscovite mica under a thin water film	27	J.
15		grown from humid air. Sci. Rep., 2017. 7 , 4054.		C
10	12	D. Argyris, T. Ho. D. R. Cole and A. Striolo, Molecular	28	A.
12		dynamics studies of interfacial water at the alumina		zi
10		surface, J. Phys. Chem. C. 2011, 115 , 2038–2046.		ex
20	13	R. Debbarma and A. Malani. Comparative Study of Water	29	C.
20	10	Adsorption on a H+ and K+ Ion Exposed Mica Surface:		0
21		Monte Carlo Simulation Study, Lanamuir, 2016, 32 , 1034–		R
22		1046.		ar
23	14	J. W. Wang, A. G. Kalinichev and R. J. Kirkpatrick. Effects of		73
24		substrate structure and composition on the structure.	30	S.
25		dynamics, and energetics of water at mineral surfaces: A		Di
20		molecular dynamics modeling study. <i>Geochim. Cosmochim.</i>		in
27		Acta, 2006, 70 , 562–582.		89
20	15	B. Rotenberg, A. J. Patel and D. Chandler, Molecular	31	M
30	-	explanation for why talc surfaces can be both hydrophilic	-	ar
31		and hydrophobic. J. Am. Chem. Soc., 2011. 133, 20521-7.		st
37	16	A. Hodgson and S. Hag. Water adsorption and the wetting	32	Ρ.
32		of metal surfaces. Surf. Sci. Rep., 2009. 64, 381–451.		М
34	17	A. Striolo, A. A. Chialvo, K. E. Gubbins and P. T. Cummings,		si
35		Water in carbon nanotubes: Adsorption isotherms and		24
36		thermodynamic properties from molecular simulation, J.	33	G
37		Chem. Phys., 2005, 122 , 234712.		D.
38	18	E. Xi, V. Venkateshwaran, L. Li, N. Rego, A. J. Patel, S. Garde		St
39		and M. L. Klein, Hydrophobicity of proteins and		st
40		nanostructured solutes is governed by topographical and	34	К.
41		chemical context, Proc. Natl. Acad. Sci. U. S. A., 2017, 114,		Μ
42		13345–13350.	35	C.
43	19	K. Wada, in Minerals in soil environments, eds. J. B. Dixon		D
44		and S. B. Weed, 1989, pp. 1051–1087.		Fi
45	20	E. Paineau, Imogolite Nanotubes: A Flexible Nanoplatform		ar
46		with Multipurpose Applications, Appl. Sci., 2018, 8, 1921.		N
47	21	G. Monet, E. Paineau, Z. Chai, M. S. Amara, A. Orecchini, M.	36	A.
48		Jimenéz-Ruiz, A. Ruiz-Caridad, L. Fine, S. Rouzière, LM.		Pi
49		Liu, G. Teobaldi, S. Rols and P. Launois, Solid wetting-layers	37	F.
50		in inorganic nano-reactors: the water in imogolite		ar
51		nanotube case, <i>Nanoscale Adv.</i> , 2020, 2 , 1869–1877.		al
52	22	F. C. Ugolini and R. A. Dahlgren, Weathering environments		Pł
53		and occurrence of imogolite and allophane in selected	38	L.
54		andisols and spodosols, Soil Sci. Soc. Am. J., 1991, 55,		D
55		1166–1171.		el
56	23	R. A. Dahlgren and F. C. Ugolini, Formation and stability of		36
57		imogolite in a tephritic spodosol, Cascade Range,	39	E.
58		Washington, USA, Geochim. Cosmochim. Acta, 1989, 53,		st
59		1897–1904.		Μ
60				

Journal Name

- R. A. Dahlgren and F. C. Ugolini, Distribution and characterization of short-range-order minerals in spodosols from the Washington cascades, *Geoderma*, 1991, **48**, 391– 413.
- C. Levard, E. Doelsch, I. Basile-Doelsch, Z. Abidin, H. Miche,
 A. Masion, J. Rose, D. Borschneck and J.-Y. Bottero,
 Structure and distribution of allophanes, imogolite and
 proto-imogolite in volcanic soils, *Geoderma*, 2012, 183–
 184, 100–108.

 A. G. Jongmans, P. Verburg, A. Nieuwenhuyse and F.
 Vanoort, Allophane, imogolite and gibbsite in coatings in a Costa-Rican andisol, *Geoderma*, 1995, 64, 327–342.

- J. P. Gustafsson, The surface chemistry of imogolite, *Clays Clay Miner.*, 2001, **49**, 73–80.
- 28 A. Abdelfattah and K. Wada, Adsorption of lead, copper, zinc, cobalt and cadmium by soils that differ in cationexchange materials, J. Soil Sci., 1981, **32**, 271–283.
- C. Levard, E. Doelsch, J. Rose, A. Masion, I. Basile-Doelsch,
 O. Proux, J. L. Hazemann, D. Borschneck and J. Y. Bottero,
 Role of natural nanoparticles on the speciation of Ni in
 andosols of Ia Reunion, *Geochim. Cosmochim. Acta*, 2009,
 73, 4750–4760.
- S. Rahman, H. Takaki, M. Tamai and Y. Nagatomo,
 Distribution of zinc, manganese, copper, cobalt, and nickel in Andosols profiles, *Soil Sci. Plant Nutr.*, 1996, **42**, 881– 891.
- M. S. Torn, S. E. Trumbore, O. A. Chadwick, P. M. Vitousek and D. M. Hendricks, Mineral control of soil organic carbon storage and turnover, *Nature*, 1997, **389**, 170–173.
- P. D. Cradwick, K. Wada, J. D. Russell, Yoshinag.N, C. R.
 Masson and V. C. Farmer, Imogolite, a hydrated aluminum silicate of tubular structure, *Nature-Physical Sci.*, 1972, 240, 187-.
- G. Monet, M. S. Amara, S. Rouzière, E. Paineau, Z. Chai, J.
 D. Elliott, E. Poli, L.-M. Liu, G. Teobaldi and P. Launois, Structural resolution of inorganic nanotubes with complex stoichiometry, *Nat. Commun.*, 2018, **9**, 2033.
- K. Wada and N. Yoshinaga, Structure of imogolite, Am.
 Mineral., 1969, 54, 50–71.
- C. Levard, A. Masion, J. Rose, E. Doelsch, D. Borschneck, C. Dominici, F. Ziarelli and J. Y. Bottero, Synthesis of Imogolite Fibers from Decimolar Concentration at Low Temperature and Ambient Pressure: A Promising Route for Inexpensive Nanotubes, J. Am. Chem. Soc., 2009, **131**, 17080-+.
- 6 A. Fernandez-Martinez and L. J. Michot, *Physicochemical Properties of Imogolite*, 2016, vol. 7.
- F. Alvarez-Ramirez, Ab initio simulation of the structural and electronic properties of aluminosilicate and aluminogermanate natotubes with imogolite-like structure, *Phys. Rev. B*, 2007, **76**, 125421.
- L. Guimaraes, A. N. Enyashin, J. Frenzel, T. Heine, H. A.
 Duarte and G. Seifert, Imogolite nanotubes: Stability, electronic, and mechanical properties, ACS Nano, 2007, 1, 362–368.
- E. Balan, M. Lazzeri, G. Morin and F. Mauri, First-principles study of the OH-stretching modes of gibbsite, *Am. Mineral.*, 2006, **91**, 115–119.

1	ARTI	CLE		Journal Name
2	40	H Scalfold and M Woddo, Polinoment of the crystal	56	G Krosse and D. Jouhart. From ultrasoft proudenations
3	40	structure of gibbsite Al(OH)3 Zeitschrift für Krist 1974	50	d. Riesse and D. Joubert, From unrason pseudopotentials
4 5		139 , 129–135.		1999. 59 . 1758–1775.
6	41	R. T. Cygan, J. J. Liang and A. G. Kalinichev, Molecular	57	J. P. Perdew, K. Burke and M. Ernzerhof, Generalized
7		models of hydroxide, oxyhydroxide, and clay phases and		gradient approximation made simple, Phys. Rev. Lett.,
8		the development of a general force field, J. Phys. Chem. B,		1996, 77 , 3865–3868.
9		2004, 108 , 1255–1266.	58	L. Denaix, I. Lamy and J. Y. Bottero, Structure and affinity
10	42	B. Creton, D. Bougeard, K. S. Smirnov, J. Guilment and O.		towards Cd2+, Cu2+, Pb2+ of synthetic colloidal
11		Poncelet, Molecular dynamics study of hydrated imogolite.		amorphous aluminosilicates and their precursors, Colloids
12		1. Vibrational dynamics of the nanotube, J. Phys. Chem. C,		Surfaces a-Physicochemical Eng. Asp., 1999, 158 , 315–325.
13		2008, 112 , 10013–10020.	59	A. Noy, D. V. Vezenov and C. M. Lieber, Chemical force
14	43	B. Creton, D. Bougeard, K. S. Smirnov, J. Guilment and O.		microscopy, Annu. Rev. Mater. Sci., 1997, 27 , 381–421.
15		Poncelet, Molecular dynamics study of hydrated imogolite	60	R. W. Friddle, A. Noy and J. J. De Yoreo, Interpreting the
16		- 2. Structure and dynamics of confined water, <i>Phys. Chem.</i>		widespread nonlinear force spectra of intermolecular
17	4.4	Chem. Phys., 2008, 10, 4879–4888.	61	bonds., <i>Proc. Natl. Acad. Sci. U. S. A.</i> , 2012, 109 , 13573–8.
18	44	B. Creton, D. Bougeard, K. S. Smirnov, J. Guiment and O.	01	S. A. Gazze, I. Hallin, G. Quilli, E. Dudley, G. P. Matthews, R. Roos, G. Van Koulon, S. H. Doorr and L. W. Francis
19		of allophane I Phys Chem C 2008 112 358–364		Organic matter identifies the nano-mechanical properties
20	45	R T Cygan I A Greathouse H Heinz and A G Kalinichev		of native soil aggregates Nanoscale 2018 10 520–525
21	43	Molecular models and simulations of layered materials. <i>J.</i>	62	T. Hassenkam, J. J. Skovbierg and S. J. S. Stipp, Probing the
22		Mater. Chem., 2009, 19 , 2470–2481.		intrinsically oil-wet surfaces of pores in north sea chalk at
23	46	I. C. Bourg, S. S. Lee, P. Fenter and C. Tournassat, Stern		subpore resolution, Proc. Natl. Acad. Sci. U. S. A., 2009,
25		Layer Structure and Energetics at Mica-Water Interfaces, J.		106 , 6071–6076.
26		Phys. Chem. C, 2017, 121 , 9402–9412.	63	P. D. Ashby and C. M. Lieber, Ultra-sensitive imaging and
27	47	E. Ferrage, B. A. Sakharov, L. J. Michot, A. Delville, A. Bauer,		interfacial analysis of patterned hydrophilic SAM surfaces
28		B. Lanson, S. Grangeon, G. Frapper, M. Jiménez-Ruiz and G.		using energy dissipation chemical force microscopy, J. Am.
29		J. Cuello, Hydration properties and interlayer organization		Chem. Soc., 2005, 127 , 6814–6818.
30		of water and ions in synthetic na-smectite with tetrahedral	64	W. Fu and W. Zhang, Measurement of the surface
31		layer charge. Part 2. Toward a precise coupling between		hydrophobicity of engineered nanoparticles using an
32		molecular simulations and diffraction data, J. Phys. Chem.		atomic force microscope, Phys. Chem. Chem. Phys., 2018,
33	40	<i>C</i> , 2011, 115 , 1867–1881.	CF	20 , 24434–24443.
34	48	H. Zhang and S. J. Singer, Analysis of the subcritical carbon	65	P. Stock, J. I. Monroe, T. Utzig, D. J. Smith, M. S. Shell and
35		6296		Molecular Scale Using Force Spectroscopy and Molecular
36 27	49	E Veilly I Roques M C Iodin-Caumon B Humbert B		Dynamics Simulations ACS Nano 2017 11 2586–2597
20 20	45	Drot and F. Simoni. Uranyl interaction with the hydrated	66	O. Dudko, G. Hummer and A. Szabo, Intrinsic Bates and
20		(001) basal face of gibbsite: A combined theoretical and	00	Activation Free Energies from Single-Molecule Pulling
40		spectroscopic study, J. Chem. Phys., 2008, 129 , 244704.		Experiments, <i>Phys. Rev. Lett.</i> , 2006, 96 , 108101.
41	50	S. Plimpton, Fast parallel algorithms for short-range	67	J. L. Hutter and J. Bechhoefer, Calibration of atomic-force
42		molecular dynamics, J. Comput. Phys., 1995, 117, 1–19.		microscope tips, <i>Rev. Sci. Instrum.</i> , 1993, 64 , 1868–1873.
43	51	D. E. Smith, Molecular computer simulations of the	68	R. W. Friddle, K. Battle, V. Trubetskoy, J. Tao, E. A. Salter, J.
44		swelling properties and interlayer structure of cesium		Moradian-Oldak, J. J. De Yoreo and A. Wierzbicki, Single-
45		montmorillonite, <i>Langmuir</i> , 1998, 14 , 5959–5967.		molecule determination of the face-specific adsorption of
46	52	ST. Lin, M. Blanco and W. Goddard, The two-phase model		Amelogenin's C-terminus on hydroxyapatite., Angew.
47		for calculating thermodynamic properties of liquids from		Chem. Int. Ed. Engl., 2011, 50 , 7541–5.
48		molecular dynamics: Validation for the phase diagram of	69	T. D. Kühne and R. Z. Khaliullin, Electronic signature of the
49		Lennard-Jones fluids, J. Chem. Phys., 2003, 119 , 11792.		instantaneous asymmetry in the first coordination shell of
50	53	SI. Lin, P. K. Maiti and W. Goddard, Two-phase	70	liquid water, Nat. Commun., 2013, 4 , 1–7.
51		entropy of water from molecular dynamics simulations.	70	G. Desiraju and T. Steiner, The Weak Hydrogen Bona in
52		Phys. Chem. B 2010, 114, 8191–8		Online Oxford LIK 2010
53	54	G Kresse and L Hafner, Ab-initio molecular dynamics	71	D T Limmer A P Willard P Madden and D Chandler
54 55	51	simulation of the liquid-metal amorphous semiconductor		Hydration of metal surfaces can be dynamically
56		transition in germanium, <i>Phys. Rev. B</i> , 1994, 49 , 14251–		heterogeneous and hydrophobic, Proc. Natl. Acad. Sci. U. S.
57		14269.		A., 2013, 110 , 4200–4205.
58	55	G. Kresse and J. Hafner, Ab-initio molecular dynamics for	72	M. L. Machesky, M. Predota, D. J. Wesolowski, L. Vlcek, P.
59		liquid metals, Phys. Rev. B, 1993, 47, 558–561.		T. Cummings, J. Rosenqvist, M. K. Ridley, J. D. Kubicki, A. V
60				

This journal is © The Royal Society of Chemistry 20xx

2		
3		Bandura, N. Kumar and J. O. Sofo, Surface Protonation at
<u>л</u>		the Rutile (110) Interface: Explicit Incorporation of
5		Solvation Structure within the Refined MUSIC Model
5		Framework, <i>Janamuir</i> , 2008, 24 , 12331–12339,
7	73	L D Brown and D Altermatt Bond-valence parameters
/	/5	obtained from a systematic analysis of the Inorganic
0		Crystal-Stucture Database Acta Crystalloar Sect B-
9		Structural Sci. 1085 41 244 247
10	74	Siluciului Sci., 1985, 41 , 244–247.
11	74	G. Sposito, The surjuce chemistry of hatarda 2004
12		Oxford University Press, New York, 1st edn., 2004.
13	75	I. Hiemstra and W. H. VanRiemsdijk, A surface structural
14		approach to ion adsorption: The charge distribution (CD)
15		model, J. Colloid Interface Sci., 1996, 179 , 488–508.
16	76	Y. Arai, M. McBeath, J. R. Bargar, J. Joye and J. A. Davis,
17		Uranyl adsorption and surface speciation at the imogolite-
18		water interface: Self-consistent spectroscopic and surface
19		complexation models, Geochim. Cosmochim. Acta, 2006,
20		70 , 2492–2509.
21	77	D. A. Sverjensky, Prediction of surface charge on oxides in
22		salt solutions: Revisions for 1 : 1 (M+L-) electrolytes,
23		Geochim. Cosmochim. Acta, 2005, 69, 225–257.
24	78	T. Hiemstra, P. Venema and W. H. VanRiemsdijk, Intrinsic
25		proton affinity of reactive surface groups of metal
25		(hydr)oxides: The bond valence principle, J. Colloid
20		Interface Sci 1996 184 680–692
27	79	T Hiemstra H Yong and W H Van Riemsdijk Interfacial
20	15	charging phenomena of aluminum (Hydr)oxides <i>Langmuir</i>
29		1000 15 50/2_5055
30	<u>80</u>	Y Cap and C V Franks Charging behavior of the Cibbsite
31	80	hasal (001) surface in NaCl colution investigated by AEM
32		basar (001) surface in Naci solution investigated by AFIV
33	04	Colloidal probe technique, Langmuir, 2006, 22 , 6087–6092.
34	81	B. R. Bickmore, C. J. Tadanier, K. M. Rosso, W. D. Wonn and
35		D. L. Eggett, Bond-valence methods for pka prediction:
36		critical reanalysis and a new approach, Geochim.
37		Cosmochim. Acta, 2004, 68 , 2025–2042.
38	82	I. D. Brown, The Chemical Bond in Inorganic Chemistry: The
39		Bond Valence Model, Oxford University Press, New York,
40		2002.
41	83	B. R. Bickmore, K. M. Rosso, I. D. Brown and S. Kerisit,
42		Bond-Valence Constraints on Liquid Water Structure, J.
43		Phys. Chem. A, 2009, 113 , 1847–1857.
44	84	I. Basile-Doelsch, R. Amundson, W. E. E. Stone, C. A.
45		Masiello, J. Y. Bottero, F. Colin, F. Masin, D. Borschneck and
46		J. D. Meunier, Mineralogical control of organic carbon
47		dynamics in a volcanic ash soil on La Reunion, Eur. J. Soil
48		Sci., 2005, 56 , 689–703.
49	85	T. Chevallier, T. Woignier, J. Toucet, E. Blanchart and P.
50		Dieudonne, in 14th International Sol-Gel Conference.
51		Montpellier, FRANCE, 2007, pp. 231–238.
51	86	M Egli M Nater A Mirabella S Raimondi M Ploetze and
52	00	L Alioth Clay minerals oxybydroxide formation element
55		leaching and humus development in volcanic soils
54		Cooderma 2008 142 101 114
55	07	B K C Thong and C D Vien Nanonerticles in the Self
56	0/	D. N. G. Theng and G. D. Yuan, Nanoparticles in the Soll
57	00	Environment, <i>Elements</i> , 2008, 4 , 395–399.
58	88	IVI. A. WIISON, N. H. Iran, A. S. Millev, G. S. K. Kannangara,
59		H. VOIK and G. Q. M. Lu, Nanomaterials in soils, Geoderma,
60		

Journal Name

1

2008, **146**, 291–302.

- C. Rasmussen, K. Heckman, W. R. Wieder, M. Keiluweit, C. R. Lawrence, A. A. Berhe, J. C. Blankinship, S. E. Crow, J. L. Druhan, C. E. Hicks Pries, E. Marin-Spiotta, A. F. Plante, C. Schädel, J. P. Schimel, C. A. Sierra, A. Thompson and R. Wagai, Beyond clay: towards an improved set of variables for predicting soil organic matter content, *Biogeochemistry*, 2018, **137**, 297–306.
- D. Spagnoli, J. F. Banfield and S. C. Parker, Free energy change of aggregation of nanoparticles, *J. Phys. Chem. C*, 2008, **112**, 14731–14736.
- D. Li, M. H. Nielsen, J. R. I. Lee, C. Frandsen, J. F. Banfield and J. J. De Yoreo, Direction-specific interactions control crystal growth by oriented attachment, *Science (80-.).*, 2012, 336, 1014–1018.