Hydration layers are formed on hydrophilic crystalline surfaces immersed in water. Their existence has also been predicted for hydrophobic surfaces, yet the experimental evidence is controversial. Using 3D-AFM imaging, we probed the interfacial water structure of hydrophobic and hydrophilic surfaces with atomic-scale spatial resolution. We demonstrate that the atomic-scale structure of interfacial water on crystalline surfaces presents two antagonistic arrangements. On mica, a common hydrophilic crystalline surface, the interface is characterized by the formation of 2 to 3 hydration layers separated by approximately 0.3 nm. On hydrophobic surfaces such as graphite or hexagonal boron nitride (h-BN), the interface is characterized by the formation of 2 to 4 layers separated by about 0.5 nm. The latter interlayer distance indicates that water molecules are expelled from the vicinity of the surface and replaced by hydrocarbon molecules. This creates a new 1.5–2 nm thick interface between the hydrophobic surface and the bulk water. Molecular dynamics simulations reproduced the experimental data and confirmed the above interfacial water structures.

**Introduction**

The interface of an aqueous solution with a solid surface is ubiquitous in chemistry, biology and technology. It participates in many processes from heterogeneous catalysis and electrochemistry to protein folding and cell signaling. Yet the molecular-scale understanding of the interactions between aqueous solutions and solid interfaces remains unsatisfactory.1,2

Hydration layers are key components of the structure of interfacial water. Our understanding of hydration layers comes from the surface force apparatus,3,4 X-ray methods,5,6 high-resolution AFM images,7–12 molecular dynamics (MD) and other theoretical methods.1,13–19 The paradigm emerging from those results is that the attractive interaction of the water molecules with the atoms of the crystalline surface leads to an alternation of high and low density water layers.

The spacing between hydration layers measured on hydrophilic surfaces such as mica and calcite surfaces is about 0.3 nm,7 this is, very close to the van der Waals diameter of a water molecule. The same type of layering structure and spacing has been predicted by MD simulations on graphitic and other hydrophobic surfaces.14–16 Yet the experimental evidence is either inconclusive or controversial.17–20 Explanations about the structure and composition of interfacial water on a hydrophobic surface range from the formation of ice-like water21 to the adsorption of air-borne molecules22–24 or the layering of condensed gas molecules.25,26,29–31

Here we have applied a newly developed AFM mode that provides atomic resolution 3D images of the organization of water on mica, graphite and h-BN to explain the structure of interfacial water on solid surfaces. Fig. 1a shows a volume AFM image of water on mica. Different 2D maps and images of the interface can be extracted from the volume data, such as maps perpendicular to the surface (Fig. 1b) or images parallel to it at different heights (Fig. 1c). The 2D maps perpendicular to the surface show the hydration layers, while the 2D images parallel to the surface reveal the mica lattice, the adsorbed ions or the organization of water molecules in the hydration layers. The data can be processed to obtain the dependence of the force as a function of the distance (FDC) to the surface (Fig. 1d).21,32–34

**Methods**

**Experimental Methods**

**AFM Imaging**

Atomic force microscopy (AFM) images were obtained using a Dimension Icon atomic force microscope (Bruker) equipped with a Nanoscope V controller and an RTESPA AFM head, with a silicon cantilever (NanoSensors) with a spring constant of 2 N m\(^{-1}\) and a resonance frequency of 300 kHz. The AFM images were recorded in the contact mode with a tip-sample force of 2 nN and a scan rate of 1 Hz. The force-distance curves were obtained through the tapping mode with a tip-sample force of 1 nN and a scan rate of 1 Hz.

**Molecular Dynamics Simulations**

Molecular dynamics simulations were performed on a graphene surface using the GROMACS software package.29 The simulations were performed using the Amber99SB force field and the TIP3P water model. The systems were equilibrated at 300 K for 100 ps and then run for 1000 ps at constant temperature and pressure. The simulation box was 2 × 2 × 2 nm\(^3\) with 64 water molecules per cell. The time step was 2 fs and the long-range electrostatic interactions were calculated using the Ewald summation method.
Results

AFM experiments

The FDCs measured on mica, graphite and h-BN surfaces immersed in liquid water provide quantitative information on the interfacial water (Fig. 2a–d). The force–distance curves show an oscillatory behaviour that alternates attractive and repulsive regions until the tip establishes mechanical contact with the surface. From then on, the force becomes increasingly repulsive. The peaks observed in the FDC were associated with peaks in the liquid density although they are not located at the same distance from the solid surface.35–37

The shape of the FDCs measured on mica (Fig. 2a), graphite (Fig. 2b) and h-BN (Fig. 2c) are quite similar, however, the interlayer separations are different. The spacing between the layers measured on graphite (0.51 nm) and h-BN (0.43 nm) were larger than the interlayer separations measured on mica (0.3 nm). A variety of experimental results obtained on hydrophilic surfaces7 and MD simulations [see below] on both hydrophilic and hydrophobic surfaces13–17 indicates that the layering structure of water has a periodicity of 0.30 ± 0.03 nm. Therefore, the layering observed on graphite and h-BN is incompatible with the presence of water. Interestingly, the interlayer distances observed on graphite agree with the values measured in interfaces formed by organic solvents on a variety of crystalline surfaces.19,38–40 In addition, the values measured on graphite and h-BN immersed in purified water are similar to separations measured on graphite immersed in alkanes (Fig. 2d).

The experiments reported here have been performed with purified water characterized by a resistivity of 18.2 MΩ cm⁻¹. It is known that purified water carries trace amounts of hydrocarbon molecules (1–10 ppb).41 It is likely that these hydrocarbons originate from ambient air.28,42–44 They might be still

Fig. 1 3D-AFM image of a solid–water interface. (a) Mica–water interface. The 3D image can be split up into different 2D maps and images. An image of the mica lattice is shown at the bottom. (b) 2D maps of the mica–water interface taken at different orientations with respect to the mica basal plane. The maps represent the tip’s phase shift as a function of the tip’s position (xy, z). (c) 2D images parallel to the mica surface taken at different z distances. Lateral ordering of the water is seen, e.g. in the image taken at z = 0.35 nm. (d) Force distance curve obtained by processing the bottom panel of (b).

Fig. 2 Force distance curves (FDC). (a) FDC obtained on mica immersed in purified water. (b) FDC obtained on HOPG immersed in purified water. (c) FDC obtained on h-BN immersed in purified water. (d) FDC obtained on HOPG immersed in hexane.
biasing force method. Although the tip radius is expected to
tance from the aforementioned interfaces using the adaptive
lated the average force on this model tip as a function of dis-
graphite
Fig. 3a near a mica
have constructed a model AFM tip, which is illustrated in
during the experiments.

Furthermore, the FDCs for h-BN and graphite surfaces
experimental FDC for graphite immersed in hexane (Fig. 2d).

This value is similar to the distance between maxima in the
experimental FDC for graphite immersed in hexane (Fig. 2d).

The distance between force
peaks is well reproduced. Likewise, the computational FDC for
graphite–hexane interface matches well with the experimental
FDC for that interface (Fig. 3f). On the other hand, the
calculations using a model of a pristine graphite–water interface
do not even exhibit qualitative agreement with the graph-

The computational force profile for the mica–water inter-
face agrees remarkably well with the corresponding experi-
mental FDC (Fig. 3d). In particular, the distance between force
peaks is well reproduced. Likewise, the computational FDC for
the graphite–hexane interface matches well with the experimental
FDC for that interface (Fig. 3f). On the other hand, the
calculations using a model of a pristine graphite–water interface
do not even exhibit qualitative agreement with the graph-

The hydrocarbons adopt a more ordered structure near the
graphite than in bulk. As illustrated in Fig. S9a, hexane
molecules at the graphite surface tend to lie flat on the surface
and align with each other. Subsequent layers show similar, but
weaker, tendencies to align parallel to the surface. To quantify
this tendency, we have calculated the orientational order par-

ter of hexane C–C bonds as function of distance from
graphite and mica surfaces (Fig. S9b). The hexane molecules
tend to align parallel to the surface at distances where the
hexane density is high and perpendicular to the surface in
between, where the density is low. Hence, the diffusion of a

present in the liquid after the application of stringent purifi-
cation processes. We propose that either airborne or liquid-
borne hydrocarbon molecules might accumulate on the graph-
ite (h-BN)–liquid water interface while being absent from the
mica–water interface.

Molecular dynamics simulations

To test the above hypothesis and to understand the free energy
factors that drive the formation of interfacial water structures,
we performed several sets of MD simulations. First, simula-
tions of mica–solvent and graphite–solvent interfaces pre-
dicted that the structure of the solvent overlayers are mostly
determined by the chemical nature of the solvent. Notably, as
shown in Fig. S7 (ESI), the distance between the primary and
secondary density maxima at mica–water and graphite–water
interfaces (0.28 nm) is less than that at mica–hexane and
graphite–hexane interfaces (0.44 nm). In fact, the density pro-
files of all straight-chain alkanes considered (from pentane to
decane) show a well-defined characteristic length of 0.44 nm.

Furthermore, the FDCs for h-BN and graphite surfaces
immersed in water (Fig. 2b and c) show similar characteristic
lengths, supporting the hypothesis that alkanes or molecules
with prominent alkane moieties are present at the interfaces
during the experiments.

To estimate the forces measured in the experiments, we
have constructed a model AFM tip, which is illustrated in
Fig. 3a near a mica–water interface, and in Fig. 3b and c near
graphite–water and graphite–hexane interfaces. We have calcu-
lated the average force on this model tip as a function of dis-
tance from the aforementioned interfaces using the adaptive
biasing force method. Although the tip radius is expected to
be several nanometers, the magnitude of the measured forces
suggests that contact with the solvation layers or surface is
made by some small asperity; hence, our model has a contact
area of only a few atoms. Tests with various alternative models
showed that the qualitative results depended little on the
details of the tip asperity model.

The computational force profile for the mica–water inter-
face agrees remarkably well with the corresponding experi-
mental FDC (Fig. 3d). In particular, the distance between force
peaks is well reproduced. Likewise, the computational FDC for
the graphite–hexane interface matches well with the experimental
FDC for that interface (Fig. 3f). On the other hand, the
calculations using a model of a pristine graphite–water interface
do not even exhibit qualitative agreement with the graph-
ite–water FDC (Fig. 3e). This disagreement is not surprising
given that we hypothesize that a pristine graphite–water inter-
face is a poor model of the interface. Indeed, the simulation
system including the graphite–hexane interface (Fig. 3e)
appears to be a much better model of the experiment for
graphite immersed in water than the system with the pristine
graphite–water interface (Fig. 3b). The comparison between
experimental and computational FDC requires some consider-
ations about the origin of the z-coordinate (see ESI†).
hexane molecule from one solvation layer to another requires its reorientation, making diffusion near the surface significantly slower than in bulk hexane (Fig. S9c†).

**Time evolution of the interfacial water structure**

To understand the origin of the hydrocarbon layers formed at a graphite–water interface we studied the evolution of the interfacial water structure as a function of the time passed from the preparation of the crystalline surface. We prepared two types of surfaces, fresh and exposed. A fresh surface, either from mica, graphite or h-BN, was prepared by cleavage with the resulting surface immediately immersed in purified water (<2 s). An exposed surface was prepared by cleavage with the resulting surface being exposed to ambient air for 30 minutes, then immersed in purified water. The experiments reported in Fig. 2 were performed on exposed surfaces.

Fig. 4 shows the interlayer distances measured on fresh and exposed mica and graphite surfaces. On mica, we did not observe any significant differences between the structure of the interfacial water on fresh and exposed surfaces. On the other hand, the interfacial water structure observed on graphite showed a marked difference between fresh and exposed surfaces. On exposed graphite surfaces, the interlayer distances were centered at 0.45 (1st), 0.55 (2nd) and 0.52 nm (3rd). Those values were very close to the values obtained by MD on a graphite surface immersed in an alkane solvent. On fresh surfaces, we observed a decrease of the mean values, in particular for the 1st layer (~0.35 nm). This value was close to 0.32 nm which characterizes hydration layers. It suggested that water molecules might form hydration layers on graphite whenever hydrocarbons are not adsorbed on the surface. In addition, the difference observed between the mean and median values indicated that the interfacial water structure on a fresh graphite surface was somehow unstable. A low amount of hydrocarbon species adsorbed on graphite is likely to generate a competition between water and hydrocarbon molecules which might lead to a changing interfacial water structure.

The above data indicates that airborne hydrocarbons are the constituents of the interfacial layers observed on hydrophobic surfaces. The adsorption of hydrocarbons might have two different pathways. They could be directly adsorbed from the air-solid surface interface or from an air–liquid interface.

**Comparison with hydrophilic surfaces**

Further evidence of the accumulation of liquid-borne molecules on graphite and h-BN comes from the data extracted from 3D-AFM images. Fig. 5a shows a 2D xz map of the mica–water interface and two xy images taken, respectively, at the mica surface (z = 0.00 nm) and 0.35 nm above the surface. At z = 0.35 nm, the image shows a honeycomb structure formed by water molecules that surround the K⁺ ions that sit on top of the mica trigonal cavities. Hydrocarbon molecules are not present in this interface.

Fig. 5b shows a 2D xz map and two xy images of the graphite–water interface. Three hydrophobic layers are resolved (top panel). The middle panel reveals a ripple structure with a periodicity of 4.8 nm. A finer orthogonal structure with a periodicity of 0.5 nm is also shown. The latter value matches the interlayer distance observed in the 2D xz panel and the periodicity of alkane monolayers formed on graphite and h-BN.46,47 By breaking into this layer we could recover the graphite lattice (bottom panel).48 Fig. 5c shows the 2D xz maps and xy images for h-BN immersed in water. The results coincide with those reported for graphite (Fig. 5b)).
Discussion

The formation of gas layers (N₂) was suggested to explain the observed interlayer distances.²⁹,³⁰ Our data do not support that interpretation. To further rule out the gas adsorption model we have performed additional experiments and simulations. Experiments performed by using degassed water did not show any changes in the interlayer distances. A comprehensive MD simulation was performed to study the adsorption of N₂ on graphite. Fig. 6 shows simulations of a model tip for graphite–hexane and graphite–N₂ + water interfaces. The experimental data obtained on exposed graphite immersed in purified water agrees with the simulation of a graphite-alkane interface. The force predicted for a graphite-N₂ gas + water interface agreed neither with the shape nor with the intermaxima distances of the experimental FDCs. Other MD simulations performed on hydrophobic surfaces⁴⁹ favored the quick formation of gas nanobubbles from an adsorbed layer. This result also contradicted the gas adsorption hypothesis.

The structure of the interface is determined by a competition between water and hydrocarbon molecules from the environment. On mica, water experiences strong electrostatic interactions with the surface, overwhelming relatively weak surface-hydrocarbon interactions and resulting in a net repulsive thermodynamic force on hydrocarbon molecules. On hydrophobic surfaces, particularly graphite and h-BN, water
molecules are expelled from the interface by hydrocarbons, reducing the overall area of water–hydrophobic contact and resulting in more favorable water–water interaction (Fig. S8†).

Our model of interfacial water on crystalline hydrophobic surfaces challenges the prevalent paradigm in the surface science community.1 It questions the interpretation of some recent experimental data22,24 or the use of hydrophobic nanopores to confine water.50 At the same time, it provides a comprehensive understanding of the interfacial water and support the use of passivating layers to prevent contamination.51 Let’s illustrate one of the implications. The dielectric constant $\varepsilon$ of water confined between two atomically flat hydrophobic walls was measured down to 1 nm separations.24 A value of $\varepsilon = 2$ was measured for very small separations (1–2 nm), this is, far from the value of bulk water $\varepsilon = 80$. The results were explained by the confinement of the water molecules which inhibits the rotational freedom of the dipole moments.24 However, the existence of a 1–2 nm layer of hydrocarbon molecules provides a different explanation because alkane molecules have already a very low dielectric constant ($\varepsilon = 2$ at $T = 295$ K).

**Conclusions**

The prevalent paradigm about hydration layers was derived from experiments performed on hydrophilic surfaces. Assuming a pristine and uncontaminated interface, MD simulations supported their existence on hydrophobic surfaces. However, ambient air and purified water have a trace of hydrocarbon molecules ($\leq 10$ ppb in water). These molecules, either by direct adsorption from the air-solid interface or by diffusion from the air–liquid interface, might control the organization of water on hydrophobic surfaces. Until the present work, the presence of trace hydrocarbons was commonly ignored in MD simulations.

Atomic resolution volume images reveal a dual interfacial water organization. The structure and composition of the solid–water interface depends on the hydrophilic/hydrophobic character of the surface. On mica, the interfacial water is formed by hydration layers while on graphite and h-BN, the interfacial layer is formed by airborne and waterborne hydrocarbon molecules.

These results illustrate the complexity of real water interfaces where a trace amount of hydrocarbons might dominate the intrinsic properties of the interface. In addition, the data illustrate how advanced high-resolution methods are needed to reach a fundamental understanding of the interactions of real liquid water with surfaces, with implications in molecular biology, energy storage and environmental engineering, among others.

**Experimental**

**Crystalline surfaces**

Flakes of hexagonal-boron nitride [h-BN, purchased from HQ Graphene, Netherlands], were mechanically exfoliated with adhesive tape and transferred onto clean Si/SiO$_2$ substrates (275 nm SiO$_2$, thermally oxidized) using a polydimethylsiloxane (PDMS) stamp. For cleaning, the substrates were sequentially ultrasonicated in acetone (99.6%, Acros Organics), ethanol ($\geq 99.8\%$, Sigma Aldrich), and ultrapure water. After drying the substrates with a flow of nitrogen gas, they were then exposed to oxygen plasma for 15 min (Diener Electronic, Germany).

Highly oriented pyrolytic graphite (HOPG, grade ZYB) was purchased from Bruker (USA) and cleaved with adhesive tape before the experiment. Muscovite mica (Grade V-1) was purchased from SPI supplies (USA). The mica was freshly cleaved with adhesive tape and rinsed copiously with ultrapure water before the experiments.

**Solvents**

Ultrapure water was freshly obtained before the experiments (ELGA Maxima, 18.2 M$\Omega$ cm$^{-1}$). The water’s pH value reached a value of 5.6 a few minutes after obtaining it from the machine (Hanna Instruments HI 9024). Some experiments on h-BN and HOPG were performed in $n$-hexane ($\geq 99\%$, Scharlab), $n$-octane ($\geq 99\%$, Sigma-Aldrich) and $n$-pentadecane ($\geq 99\%$, Sigma-Aldrich).

**AFM imaging**

We used a home-made three-dimensional AFM.11,27 The 3D-AFM was implemented on a Cypher VRS platform. Three-dimensional AFM imaging is performed in the amplitude modulation mode23 by exciting the microcantilever at its first eigenmode. At the same time that the cantilever oscillates with respect to its equilibrium position, a sinusoidal signal is applied to the $z$-piezo to modify the relative $z$-distance between the sample and the tip. We have used $z$-piezo displacements with amplitudes between a 1 and 2.5 nm and a period (fre-
frequency) of 10 ms (100 Hz). This z-range was chosen because we did not observe features in the liquid structure above \( \geq 1.5 \) nm. The z-piezo signal is synchronized with the xy-displacements in such a way that for each xy-position on the surface of the material, the tip performs a single and complete z-cycle.

The z-data is read out every 10.24 \( \mu \)s and stored in 512 pixels (256 pixels half cycle). Each xy-plane of the 3D map contains 80 \( \times \) 64 pixels. Hence, the total time to acquire such a 3D-AFM image is 52 s.

Conventional high-resolution AFM imaging was performed on a modified commercial microscope (Cypher VRS, Asylum Research, Oxford Instruments). Before each experiment the AFM holder was cleaned with ethanol and distilled water. After the cleaning procedure, the samples were mounted onto the microscope sample stage. Then, a small droplet of imaging solution was placed on the surface. Immediately after that, the cantilever was immersed into the liquid and the liquid cell was sealed. Within the cell, the temperature was held constant at (28.0 \pm 0.1) \(^\circ\)C.

Silicon cantilevers with silicon tips were used for 3D-AFM imaging (ArrowUHF AuD and ArrowUHF Al, NanoAndMore, Germany). Cantilevers were cleaned first in a mixture (50 : 50 in volume) of isopropanol (99.6\%, Acros Organics) and ultrapure water, rinsed with ultrapure water and then placed in a UV-Ozone cleaner (PSD-UV3, Novascan Technologies, USA) for 1 h. A contactless method was used to calibrate the microcantilevers.

**Molecular dynamics protocols**

The molecular dynamics simulations followed protocols validated against experiment in our previous work.\(^{53,54}\) All simulations were performed with NAMD 2.13. Lennard-Jones interactions were calculated with a smooth 1.0–1.2 nm cutoff. For all simulations, the pressure was maintained at 1.01325 bar using the Langevin piston method and the temperature was maintained at 295 K using a Langevin thermostat with a damping parameter of 1 ps\(^{-1}\). All simulations were performed with mass repartitioning of non-water hydrogen atoms (with the mass increased by a factor of 3). The equations of motion were integrated with a 3 fs time step. Electrostatic interactions were calculated using the particle-mesh Ewald method with a grid spacing <0.12 nm. Water molecules were represented by the TIP3P model of the CHARMM force field and kept rigid using the SETTLE method. The lengths of other covalent bonds involving hydrogen were constrained.

For organic molecules and graphitic carbon, interatomic interactions were represented using the CHARMM General Force Field (CGenFF), version 4.3. We have previously demonstrated that this force field agrees well with experiment for the thermodynamics of adsorption of small organic molecules on graphitic carbon in aqueous solution\(^{53,54}\) as well as in organic solvents.\(^{55}\) Muscovite mica was represented using CLAYFF. Previous studies\(^{56}\) have validated the combination of CLAYFF with the CHARMM force field framework. Mean forces on the model tip asperity as a function of distance from the surfaces were calculated by the adaptive biasing force method\(^{45}\) as implemented in the Colvars module\(^{57}\) using a 0.005 nm grid. To mimic the fact that the AFM tip is attached to a relatively rigid assembly, a restraint was applied to maintain the orientation of the tip asperity also using the Colvars.

The graphite systems included two rectangular graphene sheets stacked atop one another and aligned perpendicular to the z-axis, with mean dimensions of 2.935 \( \times \) 2.966 nm\(^2\). As in previous work\(^{53,54}\) the atoms of the lower sheet were harmonically restrained to their initial z-position. All calculations of surface interactions were performed relative to the upper sheet, to which no restraints were applied. The muscovite mica surface was created by a 2 \( \times \) 2 replication of the final structure given by Wang et al.\(^\text{58}\) to produce a bilayer mica sheet with mean dimensions of 3.583 \( \times \) 4.140 nm\(^2\). The bottom plane of tetrahedral aluminium atoms was harmonically restrained to their initial z-position. Again, all calculations of surface interactions were performed relative to the unrestrained upper layer of mica. Water molecules were added using VMD to create mica–water and graphite–water systems with mean z dimensions of 5.66 and 5.19 nm, respectively. Simulation systems including organic solvents were constructed using the program PackMol to yield systems with z-dimensions of 5.37 nm\(^3\). All systems underwent 2000 steps of energy minimization and 150 ps of equilibration before beginning production simulations.

**Author contributions**

M. R. U. and S. B. performed the experiments. M. R. U., S. B. and R. G. designed and analyzed the experiments. J. C. and R. G. planned the MD simulations. J. C. and R. T. designed, performed and analyzed the MD simulations. R. G. coordinated the activity and wrote the manuscript and SI with contributions from J. C., M. R. U. and S. B. All the authors discussed the results.

**Conflicts of interest**

There are no conflicts to declare.

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