

Water-induced correlation between single ions imaged at the solid-liquid interface.

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Abstract

When immersed into water, most solids develop a surface charge, which is neutralized by an accumulation of dissolved counterions at the interface. Although the density distribution of counterions perpendicular to the interface obeys well-established theories, little is known about counterions' lateral organisation at the surface of the solid. Here we show, using atomic force microscopy and computer simulations, that single hydrated metal ions can spontaneously form ordered structures at the surface of homogeneous solids in aqueous solutions. The structures are laterally stabilised only by water molecules with no need for specific interactions between the surface and the ions. The mechanism, studied here for several systems, is controlled by the hydration landscape of both the surface and the adsorbed ions. The existence of discrete ion domains could play an important role in interfacial phenomena such as charge transfer, crystal growth, nanoscale self-assembly and colloidal stability.

Introduction

The spatial arrangement of ions at solid-liquid interfaces is a problem central to many fields, ranging from electrochemistry¹ to colloidal science², heterogeneous catalysis³, surface science⁴ or biology⁵. When immersed into a liquid, most solids develop charges at their surface. In order to maintain electric neutrality, the counterions dissolved in the liquid tend to accumulate close to the solid's surface, forming an electrostatic double layer (EDL) that counterbalances the solid's surface charge. The precise structure and spatial distribution of ions in the EDL has been the subject of intense experimental and theoretical research over the last century. At the present time, the most widely used EDL model, the Gouy-Chapman-Stern model (GCS) (see e.g., Bard & Faulkner⁶) assumes a dense monolayer of counterions, the Stern or Helmholtz layer (hereafter called Stern layer) adsorbed onto the surface of the solid, followed by a diffuse layer of counterions whose density decays exponentially when moving away from the solid. Most EDL models, including GCS, rely on the so-called continuum assumption: ions are modelled as point charges in a continuous dielectric environment. Although routinely verified in the diffuse layer region, these models tend to fall short in the Stern layer region⁷ where the discrete size of ions can no longer be ignored. These difficulties are particularly obvious in highly charged systems^{8,9} and corrections have been proposed to better account for the complex interactions linking ions, liquid molecules and the surface structure of the solid¹⁰. Continuum models usually make the implicit assumption that the Stern layer is homogenous, with a uniform ion distribution. This assumption becomes problematic at the nanoscale where most solids exhibit some structure with localised charges and a well-defined hydration landscape that can determine the position and organisation of single ions at the interface. Such a complexity is key to many processes such as the folding and function of biomolecules¹¹, the accumulation and transfer of charges at the surface of extremophiles^{12,13} or at the electrodes of solar cells¹⁴, electro-friction¹⁵, the nucleation of nanoparticles on surfaces¹⁶, or the preferential adsorption of a particular type of ion onto

minerals¹⁷. Even at the macroscopic level, intense research is currently dedicated to the synthesis of colloid particles exhibiting non-homogenous charge distributions at their surface for better controlling the stability or ordering of colloidal solutions¹⁸.

Part of the difficulty in developing suitable models comes from the lack of experimental results. Probing the lateral organisation of ions within Stern layers is a challenging task and most results available to date stem either from computer simulations¹⁵ or theoretical developments^{19,20}. When available, experimental results are typically obtained indirectly¹⁰ or through techniques that require averaging over a large area of the interface²¹, hence implicitly relying on the assumption of a homogeneous Stern layer in order to provide a precise picture of the position of the ions and the liquid molecules.

These difficulties can in principle be overcome by local probe techniques such as atomic force microscopy (AFM). When operated in liquid, AFM is able to investigate solid-liquid interfaces locally, often with atomic- or molecular-level resolution²²⁻³¹ and in the presence of ions³²⁻³⁶. Studies allowing a clear and unequivocal identification of single ions within a Stern layer are however still sparse, with little information about the formation, stability and dynamics of adsorbed ions at the nanoscale.

Here we use AFM to image different hydrated monovalent metal ions adsorbed at the surface of mica in water. We show, with single ion resolution, that the metal ions do not adsorb randomly but form preferentially ordered structures such as rows or geometrical domains due to ion-ion attractive interactions controlled by the water at the interface. The results, confirmed by molecular dynamics (MD) simulations, show that water alone can provide a sufficient driving force to induce order within Stern layers, at least for chaotropic monovalent ions³⁷ (Throughout the paper we consistently use the concepts of "chaotropic" and "kosmotropic" as defined in reference 37). The key element is the interplay between the local solvation landscape of the solid at the interface, and the hydration structure of the ion. Electrostatic interactions induce the

adsorption of ions at the Stern layer but do not play any significant role in regulating interactions between monovalent ions within this region. This work presents, to the best of our knowledge, the first direct observation of this correlation mechanism. Significantly, the fact that it occurs between monovalent metal ions at interfaces suggests a common occurrence in natural and synthetic systems. Our findings are confirmed with different types of monovalent ions and also demonstrated on a polar self-assembled monolayer. The latter is chemically different than mica and, although only weakly charged, provides an appropriate hydration landscape that allows for the formation of ionic structures stable enough to be observed by AFM. The generality of the effect suggests that it could play an important role in wide range of systems, for example in controlling and enhancing charge transport at interfaces^{12,38}, in influencing crystal growth¹⁷ or the stability of colloidal solutions and emulsions¹⁸. Additionally, this effect could also be exploited in the design and self-assembly of nanomaterials³⁹ and used to template nanostructures¹⁶.

Results

Atomic Force Microscopy

Recently, we showed that amplitude-modulation AFM could derive quantitative information about the local free solvation energy of a solid fully immersed into a liquid, with sub-nanometre resolution³¹. This is possible when the vibrating AFM tip dissipates most of its energy within the interfacial liquid, without significant interaction between the tip and the solid^{30,31}. In this regime, hereafter referred to as “interfacial dissipation microscopy” (IDM), information about the local interfacial energy is contained in the phase image, acquired simultaneously with the topography image of the interface, and is related to the local dynamics of the liquid probed by the tip^{40,41}.

When ions adsorb at the surface of the solid, they induce a substantial perturbation of the local solvation environment, which can be detected at the single-ion level by IDM as the AFM tip forces the liquid around the imaged ion³⁵.

Single Rb^+ ions can be clearly identified in the topographic IDM images (Figure 1a). The adsorbed Rb^+ ions appear as distinct protrusions on the sites normally occupied by K^+ ions in bulk mica (hereafter called ‘binding sites’, Figure 1b). In the associated phase image the ions induce a sharp dark contrast, indicating an increase of the local tip energy dissipation (see also Supplementary Figures 1 and 2).

The IDM images reveal interesting peculiarities about the Rb^+ ions apparent dynamics and organisation at the surface of mica. First, the adsorbed Rb^+ ions appear surprisingly stable, with the same protrusion often visible in several consecutive line-scans. Theory⁴² and scanning polarisation microscopy measurements of ions mobility on mica⁴³ suggest that the residence time of adsorbed ions at a given binding site should not exceed microseconds, but our IDM measurements of the residence time of Rb^+ ions over a single binding site (Figure 1c) indicate an average time in the order of 100 milliseconds (see Supplementary Figure 3). The result is however likely to be limited by the time resolution capability of IDM measurements (~ 50 ms), which may probe ionic densities instead of single ions. Nonetheless, the protrusions unambiguously indicate the presence of Rb^+ ions at the corresponding sites, as evidenced when varying bulk ionic concentration (Figure 2b and Supplementary Figure 4).

Second, the ions do not adsorb randomly on the surface of mica, but rather appear to form small islands or rows (arrow in Figure 1a) instead of adopting a two-dimensional gas-like random arrangement. This behaviour is confirmed by a statistically significant excess of nearest neighbours for each adsorbed ion, when compared with the same number of ions arranged randomly ($p < 0.01$, Supplementary Methods Section 1). The result indicates attractive ion-ion interactions, something surprising given the equivalence of the binding sites and the expected

electrostatic repulsion between adjacent cations on the surface. This form of attractive interaction is responsible for the observed ion-ion correlation effects, and is at the centre of this study.

In order to better quantify this correlation, we performed IDM measurements of the apparent surface coverage of Rb^+ ions on mica, as a function of the RbCl bulk concentration in the aqueous solution (see Supplementary Methods Section 2 for the coverage analysis). These measurements provide Rb^+ adsorption isotherms from which it is possible to extract the adsorption free energy $\Delta G_{\text{Rb}^+}^0$ of the ions to the surface and the correlation energy E_c between adjacent ions, using appropriate models^{44,45}.

Practically any accurate IDM measurement requires first quantification of the effect of the IDM tip on the observed Rb^+ surface coverage. The tip is charged in water⁴⁶ and hence prone to electrostatic interactions with adsorbed ions. Additionally, the vibrating tip mechanically perturbs the Stern layer and may displace or remove adsorbed ions. Both these perturbations can artificially change the observed Rb^+ surface coverage. It is possible to quantify the energies E_e^{IDM} and E_m^{IDM} associated with respectively the electrostatic and mechanical perturbations induced by the IDM tip, and hence infer the properties of the unperturbed interface while retaining the IDM imaging resolution. This task is achieved by varying the mechanical perturbation in a quantifiable manner while keeping the electrostatic perturbation constant, using cantilevers with different stiffness but same tip design (Figure 2a, Supplementary Figure 5 and Supplementary Methods Section 3). Control experiments were also conducted with gold-coated tips (Supplementary Figure 6). Representative images of the IDM measurements for selected bulk RbCl concentrations are shown in Figure 2b.

Assuming that the large majority of Rb^+ ions are adsorbed in the inner-sphere hydration state at the surface of mica^{21,47} the IDM-derived isotherms can be fitted with two main models. The simplest model, the Langmuir isotherm⁴⁴, assumes a uniform and homogenous Stern layer with no interactions between adsorbing ions and therefore the correlation energy ($E_c \equiv 0$) is zero

(dashed line in Figure 2a). Alternatively, a modified Frumkin⁴⁵ isotherm model that takes into account both ion-ion correlations and contributions of the IDM tip perturbation, provides a substantially better fit to the data (solid line in Figure 2a, Supplementary Methods Section 3). The maximum coverage observed by IDM (10mM, Figure 2b) corresponds to all the binding sites being occupied and the isotherm is modelled accordingly. In reality, only half the binding sites should, on average, be occupied in order to ensure electro-neutrality at equilibrium⁴⁸. This suggests that at 10mM, the IDM images predominantly Rb^+ patches with empty area outside the imaging region. The existence of patches is consistent with correlations effects, and patches are likely to assemble in the vicinity of the negatively charged tip. The observed full surface coverage could also be explained by co-adsorption of Cl^- anions, known to occur for Cs^+ on mica⁴⁹. Co-adsorption is however unlikely to take place here^{48,50}, also in light of our simulation results (Supplementary Methods Section 4).

The main result from the isotherm analysis is the clear correlation between Rb^+ ions during the adsorption process. The associated correlation energy was found to be always attractive and similar, within uncertainty, for both types of cantilever $E_c = -7 \pm 3$ kJ/mol (Supplementary Methods Section 3). This value is consistent with the observed formation of Rb^+ rows and small domains at low salt concentration in the IDM images (Figure 1) and with statistical analysis of number of nearest neighbours (Supplementary Methods Section 1). Additionally, the fact that E_c is independent of the magnitude of the IDM perturbation indicates that the correlation is an intrinsic property of the interface. Time-dependent adsorption studies, such as that presented in Figure 1c, also showed preferential adsorption of ions on several adjacent sites simultaneously (Supplementary Figure 3), supporting the idea that $\text{Rb}^+ - \text{Rb}^+$ correlation effects stabilize the Stern layer.

This correlation can have two possible origins, either direct electrostatic interactions between adsorbed ions or hydration effects. Electrostatic correlation are known to play an important role

in systems with high charge density⁵¹ such as multivalent ions on DNA⁵², but they are unlikely to play a significant role here due to the low charge density of the ions involved. A simple estimate of the electrostatic energy gained by two isolated Rb^+ ions brought side by side on the surface of mica suggests that some other form of correlation must be taking place (Supplementary Discussion Section 1). We propose instead that the correlation is induced by a better coordination of the water molecules in the global hydration structure of neighbouring ions, compared to the hydration shell of single adsorbed ions. Under this hypothesis, although electrostatic interactions occur between solvent molecules, ions, and the surface, the correlation is effectively controlled by the structural organisation of water molecules at the interface. To test this hypothesis, we conducted molecular dynamics (MD) simulations so as to obtain a full molecular-level picture of the Rb^+ ions dynamics at the surface of mica in water.

Molecular Dynamics simulations

A ‘free’ interface (that is without the presence of an IDM probe) was used in the simulation in order to assess whether or not the IDM observations reflect the ‘unperturbed’ interface. Due to limited computational power, current simulations cannot directly replicate IDM experiments in terms of duration or system size. We therefore decided to compare the intrinsic stability of different geometrical configurations for adsorbed ions at the surface of mica in water. Three starting configurations were studied, progressively increasing the average number of direct neighbours between ions: (i) with the cations randomly distributed at the surface of mica (Figure 3a), (ii) with the cations forming a single row on mica (Figure 3b), and (iii) with the cations grouped in a single hexagonal domain on the surface (Fig 3c). Additionally, since hydration forces are expected to play an important role in the correlations effects, we also conducted the simulations with K^+ ions and Na^+ ions. The K^+ ions are generally seen as chaotropic³⁷ similarly to Rb^+ ions, but are naturally present in mica. In contrast, Na^+ ions are kosmotropic and exhibit a

smaller primary solvation shell with higher hydration energy than K^+ and Rb^+ . The simulations comprise ion-ion and ion-mica electrostatic interactions as well as hydration interactions between water molecules, ions and the mica. An ion located within 4 Å of the mica's hydroxyl oxygen is considered adsorbed (Supplementary Methods Section 4). The time-averaged interaction energies of the simulation runs are presented in Figure 3d for each ion type in the three different starting configurations, always with a similar number of ions.

A direct comparison between the simulation runs show that the overall free energy per ion is always higher when adsorbed in random configuration (Figure 3d). Decomposition of this free energy in its hydration (ion-water) and electrostatic (ion-ion and ion-mica) contributions shows that the energy dependence of the ion configuration is largely dominated by the hydration energy. The latter is about an order of magnitude larger than the ion-ion electrostatic potential, and always largest for the random configuration regardless of the type of ions considered. In the case of K^+ and Rb^+ , the hydration energy (and the total free energy) per ions decreases with the number of immediate neighbours. This trend is however not as clear for the Na^+ ions, mainly due to the particularly high mobility of Na^+ and the presence of multiple solvation states⁴⁷ (Supplementary Figures 7 and 8) leading to a rapid dissolution of the initial geometrical arrangements. This is not the case for K^+ and Rb^+ ions, which exist mainly in a single hydration state and tend to remain at the usual ion binding sites when adsorbed⁴⁷. They also exhibit little mobility throughout the simulation (Supplementary Figures 7 and 8), hence allowing for a more direct interpretation of the influence of their adsorption configuration on their respective free energy. One-way ANOVA analysis of the results demonstrated that the differences in total free energy between geometrical configurations are statistically meaningful for K^+ and Rb^+ ($p < 0.01$). Simulations also show that correlating ions have 10-15% less neighbouring water molecules over the entire length of the simulation compared to randomly adsorbed ions. This shows indirectly the favourability for the entire system to arrange ions on the surface in a

correlated way, further supporting hydration as the mechanism driving the local structuring of ions. In Figure 3(e) a simulation snapshot illustrating a possible mechanism is presented. Ions neighbouring each other can share coordinated water molecules and improve the hydration structure compared to a randomly adsorbed ion. Moreover, alternating positive (ions) and negative (water oxygen atoms) charges improves the stability of the structures as predicted by previous theoretical work⁵³.

Discussion

The IDM results consistently indicate that the Rb^+ and K^+ ions can form ordered structures when adsorbing at the surface of charged crystalline solids in solution. The attractive correlation between neighbouring ions is driven by a better coordination of the surrounding water molecules, as elucidated by the MD simulations. As a result, the effect should depend on the hydration structure of both the adsorbed ion and the surface of the solid. Furthermore, the simulations show that Rb^+ (and K^+ ions) prefer to form geometrical arrangements that favour nearest-neighbour configuration between adsorbed ions. Although only a limited number of configurations could be tested, the hexagonal arrangement (3.4 neighbours) appeared consistently more favourable than rows (2 neighbours), with both arrangements preferred over a random adsorption (0 neighbour). This 2D self-assembly process is driven by minimisation of the hydration water free energy. MD simulations show that two conditions are necessary for this ordering to take place: the ion's hydration energy has to dominate over ion-ion electrostatic interactions at the interface, and the hydration properties of the ions considered should allow the formation of an ordered monolayer, for example by favouring a single hydration state at the surface of the solid considered. The multiple solvation states of adsorbed Na^+ ions enable them to move rapidly between different locations not necessarily coinciding with a binding site and embed themselves deeper into the surface (Figure 4b and Supplementary Figure 7). The presence

of small kosmotropic ions sitting between the main binding sites has already been reported in previous AFM studies³⁴ and RAXR experiments have also shown that several kosmotropic ions can adsorb in different hydration states simultaneously⁴⁷. The surface of the solid plays an important role in the correlation effects by imposing a set distance between adjacent binding sites.

In order to verify experimentally the MD predictions for K^+ and Na^+ ions, we obtained IDM images of the surface of mica in KCl and NaCl solutions (Figure 4a-d). The results are fully consistent with the simulations: the K^+ ions appear similar to Rb^+ ions, forming domains and rows on the surface and allowing imaging conditions almost as stable as in RbCl (see also Supplementary Figure 9 and Supplementary Methods Section 4). In contrast, Na^+ ions appear poorly defined and show no clear structural organisation on the surface. Furthermore the high cations mobility on the surface and the presence of multiple hydration states prevented stable IDM imaging.

Interestingly, the phase shift observed for images of K^+ and Na^+ is different despite a same cantilever/tip and identical imaging condition. This difference reflects that of the two ions' hydration shells, suggesting that IDM can even differentiate between adsorbed ions through their specific modification of the interfacial hydration structure. Differences between ions are likely to be enhanced in IDM measurements due to the momentary confinement of the interface by the IDM tip pushing water around the ions' solvation shell. Previous studies have shown that the hydration structure of a particular ion often dominates its interfacial behaviour under confinement⁵⁴. In IDM measurements, although confinement effects are limited by the small surface of the tip effectively imaging and the slow confining speed ($<50 \mu\text{m/s}$), they may nonetheless enhance the observations of ion-ion correlations that would otherwise be difficult to capture. Indeed, previous RAXR measurements were unable to detect correlations between Rb^+ ions within experimental uncertainty⁴⁸. Present MD simulations of the 'free' (that is unconfined)

interface show that correlations effects constitute an intrinsic property of the system, which we believe is, at best, enhanced by the IDM measurements (Supplementary Discussion Sections 2, 3 and 4).

To test the generality of our findings we assessed a final experiment on a different solid: a self-assembled monolayer (SAM) exposing polar but neutral alcohol headgroups (mercaptohexanol, inset Figure 5). The SAM-functionalised gold surface develops a surface potential in aqueous solution. Previous reports indicate zeta-potentials ranging between -40 and -50mV for gold nanoparticles coated with alcohol-terminated alkanes in an aqueous solution at a similar pH^{55,56}. The SAM surface potential is weaker than that of mica. Significantly, the SAM surface in direct contact with the liquid is not charged and the polar alcohol groups interact with the liquid mainly through the formation of hydrogen bonds. The surface potential of the SAM can hence be seen as a diffused background creating an incentive for cations to form an EDL close to the SAM's surface. The molecular packing of the SAM creates a hydration landscape similar to that of mica at the nanoscale^{26,57} and ion-ion correlations can be expected to occur. Figure 5 compares high-resolution IDM images acquired in ultrapure water and in 10mM RbCl. In presence of the salt Rb⁺ clusters can be identified at the surface of the SAM, yielding a phase contrast consistent with the results on mica. The clusters are however less clearly defined than on mica and can easily be removed by the tip. This can be explained by the relatively weak surface potential of the SAM that limits the incentive for adsorbed ions to remain at the surface. Nonetheless, the results confirm that in the presence of an appropriate hydration landscape, correlated ion clusters can form solely through hydration forces, and that the effect is not limited to mica.

The overall picture emerging from IDM measurements and MD simulations highlights the importance of the interplay between the hydration properties of adsorbed ions and of that of the surface for ordering the ions. In the case of mica, chaotropic ions such as K⁺ or Rb⁺ appear mostly located within the first hydration layer with limited effect over the distal layers

(Supplementary Figures 7 and 10). This minimal disruption of the mica's interfacial water structure allows the formation of stable ionic structures guided by the particular hydration landscape of mica. The self-assembly takes place within mica's first hydration layer and is guided by a minimisation of the overall hydration energy of both the mica and the adsorbed ions. The resulting correlation effect does not require any specific interaction between the ions and the surface and can, in principle, occur at any interface which exhibiting an appropriate hydration landscape. Additionally, the short Bjerrum length of hydrated monovalent ions (~1 nm, Supplementary Discussion Section 1) makes it possible for correlation effects to take place at the surface of solids with lower surface charge density than mica, as illustrated for the SAM. Finally, given the ubiquity of K^+ ion in biology, this type of water-induced correlation may be commonplace at the surface of biomembranes where gel-phase phospholipid domains tend to adopt an arrangement with a similar periodicity as the mica lattice or the SAM studied here. Biological interfaces are however more complex, due to their strong and local dependence on the surrounding pH, and the conformational flexibility of the molecules. Correlated ionic networks are however likely to play key roles in charge transfer^{12,38} and membrane shaping⁵⁸, and both experiments and simulations are already underway to examine these possibilities.

In this paper, we have shown that single metal ions can form ordered structures at solid-liquid interfaces through water-induced correlation effects. The effect, observed experimentally at the single-ion level by IDM, is driven by the interfacial water, and depends on the hydration properties of the ions considered. MD simulations of different types of monovalent ions at the interface with mica highlighted the delicate interplay between the hydration structure of the solid and that of the adsorbing ions, and correctly predicted the IDM observations. The simplicity of the system and the generality of the effect suggest that it may spontaneously occur in a large variety of systems.

Methods

Interfacial Dissipation Microscope

All the AFM data were acquired on a Multimode Nanoscope IIIa (Digital Instruments) operated in the IDM regime (see text). The sample and the scanning tip were fully immersed into the imaging solution. Before each experiment, the AFM liquid cell was thoroughly washed by consecutive sonication in isopropanol and ultrapure water and then dried under nitrogen flow. A cantilever was then mounted on the cell and engaged over clean mica in ultrapure water. After the AFM had reached thermal equilibrium, the ultrapure water was removed and the desired solution was injected into the cell. We used standard silicon nitride cantilevers (Olympus RC800 PSA, Olympus, Tokyo) with two different lever length but identical tip design. The nominal spring constant of the levers were $k_s=0.1$ N/m (softer lever) or $k_h=0.76$ N/m (stiffer lever). Cantilevers from a same wafer were used to allow for better comparison of the results. The cantilevers were calibrated using their thermal spectra⁵⁹ after each experiment and found to have typical stiffness and Q-factors of $k_h \sim 0.5$ N/m and $Q_h \sim 4.4$ (stiffer lever), and $k_s \sim 0.1$ N/m and $Q_s \sim 2.5$ (softer lever). The cantilever was driven acoustically in a liquid cell, close to its resonance frequency. Typical working amplitudes A were kept between 0.8 nm and 1.5 nm with the setpoint ration A/A_0 as large as possible where A_0 is the free vibration amplitude of the lever in the liquid far from the mica surface. When operated in these conditions, atomic-level resolution topographic and phase images of the type shown in Figure 1, 2 and 4 could be routinely achieved. We found that too small amplitudes (<0.5 nm) resulted in no apparent surface coverage by the ions due to the tip ‘sweeping’ the ions sideways.

Generally, since the IDM tip can mechanically remove ions from the surface (see Figure 2), we always carefully balanced the imaging setpoint so as to keep it as large as possible. Nonetheless, minute setpoint variations were sufficient to introduce variations in the apparent surface

coverage; this is the main source of error in our statistical analysis (see Supplementary Methods Section 1 and 2).

Sample preparation

High quality muscovite mica (grade V) was purchased from SPI supply (West Chester, PA, USA) and used in all the experiments. The mica was freshly cleaved with adhesive tape and rinsed copiously with ultrapure water (18.2 M Ω , Merck Millipore, Billerica, MA, USA) before use.

The SAM of mercaptohexanol where created on gold substrates. The substrates, commercially available (PHASIS, Geneva, Switzerland) consisted of gold evaporated on mica packaged in a sealed nitrogen atmosphere. Upon opening the substrates were immediately placed in vials containing 10 ml of ethanolic solution of 6-mercaptohexan-1-ol (mercaptohexanol, MH) (TCI America, Portland, Oregon, USA). The thiol concentration was 0.01 mM. The substrates were left in the solution in dark at room temperature for a period of 7 days, at which point they were rinsed with ethanol seven times and dried under a gentle argon-flow.

All the solutions were prepared in ultrapure water with 99.9% pure salts purchased from Sigma-Aldrich and used as received.

Molecular Dynamics simulations

Based on the previously reported crystal structure of muscovite⁶⁰ we constructed for the molecular dynamics (MD) simulations a crystal measuring approximately 41×55×40 Å with the (001) plane set parallel to the xy-plane of the simulation referential. The exposed plane naturally contains potassium ions and depending on the required starting configuration all, none or a selected number of these ions were removed or replaced with a different ion species, resulting in either a random, row or hexagon configuration for the surface ions. In all cases we ensured

complete system charge neutrality. This fully atomistic model of muscovite is centred in a periodic simulation box and solvated in the direction perpendicular to the exposed plane, ensuring a water thickness on top of the mica of at least 5 nm.

To describe the interactions between all atoms, two force fields are used. For the muscovite mica interactions and all ion interactions we used the CLAYFF force field⁶¹, which is a general force field suitable for the simulation of hydrated and multicomponent mineral systems and their interfaces with aqueous solutions. The interactions between water molecules were described by the TIP3P model of the CHARMM force field⁶². Recently it has been shown that the CLAYFF and CHARMM force fields are compatible and can be used together⁶³⁻⁶⁶. All simulations have been performed using the MD code NAMD⁶⁷. For visual inspection VMD⁶⁸ was used and most of the analysis was performed using the Python library MDAnalysis⁶⁹.

Three different surface arrangements were examined for each type of ion: (i) a row of ions along the (100) direction, (ii). a hexagon and (iii) ions randomly placed. In the case of the row and the random distributions, 8 ions were placed at the surface, while 7 ions (a full hexagon) were used in the hexagonal configuration. Examples of these arrangements are depicted in Figure 3a-c.

For each of the nine different MD simulations a similar protocol is followed. This includes an initial minimisation of the energy associated to the water only (allowing bad contacts to be removed), followed by a short dynamics run of 5 ps to equilibrate the entire system at ambient conditions, and finally the production run of 1 ns. Only the latter is used for analysis. The current simulation set-up and protocol is similar to the setup used previously to study the interaction between calcite, water and ions³⁵, but in the current work all simulations are performed in duplicate to improve statistical accuracy and to account for individual aberrations.

In order to verify our muscovite model we computed basic properties such as water residence time near the muscovite surface and around the ions in solution, as well as the hydration numbers for these ions. We found all of the measured properties to be in good agreement with previously

reported data^{47,49,70}. For instance, the hydration numbers computed from our simulations are 5.8, 7.1 and 7.9 for Na⁺, K⁺ and Rb⁺, respectively in bulk water. As rubidium is a less common ion to encounter in mineral force fields like CLAYFF or organic force fields like CHARMM, the properties for this ion were checked more carefully and found comparable to experimental values reported^{21,48,50}.

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Authors contributions

KV designed the experiments. MR and KV conducted the IDM experiments and data analysis. PS carried out all the simulations and analysed the resulting data. The manuscript was written by KV with contributions from MR and PS. All the authors commented on the manuscript and discussed the data.

Competing financial interests

The authors declare no competing financial interests

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Figure captions

Figure 1: IDM imaging of Rb^+ ions adsorbed at the surface of mica in an aqueous solution. The solution contains 3 mM RbCl . Individual ions can be distinguished both in topography (a) and in phase (b, arrow), often forming geometrical domains or rows. Occasionally, a bright ‘tail’ is visible adjacent to the dark spots attributed to adsorbed ions. This tail is induced by a momentary increase of the tip vibration amplitude as the feedback loop completes its correction, and is due to the particularly gentle imaging conditions. The atomic structure of mica evidences the position of the ion binding sites, which are occupied by K^+ ions in the bulk crystal (b). Repeatedly scanning over a same binding site (c), yields substantial height variations in time as single Rb^+ ions adsorb and desorb from the site (see also Supplementary Figure 3). The scale bar is 3nm (a) and the colour bars represent 1nm (topography) and 15° (phase).

Figure 2: Adsorption isotherm of Rb^+ ions at the surface of mica measured by IDM in solution. The isotherms (a) comprise measurements conducted with two types of cantilevers that exhibit different spring constants ($k_s \sim 0.1$ N/m for the softer lever, and $k_s \sim 0.5$ N/m for the stiffer lever) but a same tip design. The solid lines represent data fits with the Frumkin isotherm, which allows for ion-ion correlations. The dashed lines provide fits without correlations (Langmuir isotherm) (see Supplementary Methods Section 3 for details). The apparent surface coverage derived from IDM measurements with the softer cantilever is larger than for the stiffer cantilever, consistent with the expected smaller mechanical perturbation (and hence smaller E_m^{IDM}) of the Stern layer by the softer cantilever. The results indicate a relatively small mechanical perturbation energy by the measuring IDM tip ($E_m^{IDM} < 0.4$ kJ/mol for the softer lever, Supplementary Figure 11) but a substantially larger electrostatic perturbation of $E_e^{IDM} \sim 7$ kJ/mol for both cantilevers. Control experiments with gold-coated tips confirm the dominance of the electrostatic perturbation

(Supplementary Figure 5). Representative topographic images of the IDM measurements are shown in (b) for the stiffer lever. (see also Supplementary Figures 4 and Supplementary Methods Section 3 for a full set of data). The scale bar is 5nm in (b).

Figure 3: Data from the molecular dynamics simulations of mica in aqueous solution. Three different starting configurations have been tested: with the cations randomly adsorbed (a), forming a row on mica (b) or a hexagon (c). The average hydration (open square), electrostatic (open triangles) and total (open circles) free energy of the adsorbed ions is given for each type of ion an each starting configuration (d). The data in (d) is extracted from the evolution of the hydration energy per adsorbed ion throughout the simulation. The error bars in (d) are two standard deviations. In (e) a schematic depiction of the water surrounding adsorbed K^+ . A clear difference is visible between isolated ions or ions forming geometrical patterns where the location of shared water molecules is well defined. From MD-simulations, the density of oxygen atoms of the water molecules is also shown. In the case of single ion the value of the oxygen density is the radial average around the K^+ ion. In the case of row of K^+ the average oxygen density is obtained by considering a rectangle of 0.5 nm width aligned with the row of the ions and centred on them.

Figure 4: Adsorption of K^+ and Na^+ ions at the surface of mica seen by simulation and IDM. In order to allow for a direct comparison, images in K^+ (a), Na^+ (b) and water (c) are presented side by side and on a same scale. In each case, a simulation snapshot with semi-transparent water molecules is shown (top) together with the corresponding IDM image (bottom). All the IDM data were acquired with a similar cantilever/tip and in identical operating conditions. The KCl

and NaCl bulk concentration are 1mM and 0.5mM so as to ensure only partial coverage of the mica in each case. The adsorbed ions consistently induce a darker contrast in the phase (as for Rb^+) although with a magnitude specific to each ion type. The contrast is less pronounced for Na^+ ions due to their positions more “embedded” in the mica. The Na^+ ions sitting further from the surface are only occasionally imaged by the tip. This is also reflected in the smoother topographic image achieved in NaCl. The scale bar is 3nm for all images. The colour scale bars are 1nm (a) and 0.5nm (b, c) for topography and 30° (a-c) for phase.

Figure 5: IDM images of a mercaptohexanol SAMs in water and in a 10mM RbCl solution. In both cases, the SAM lattice can be identified. In the presence of the salt, Rb^+ ions can be seen clustering at the SAM surface (arrows), creating a consistent darker contrast in the phase image. The ion clusters are however easily removed by the scanning tip, making them appear less clearly than on mica due to a weaker attractive interactions with the surface (weaker surface potential). The chemical structure of a mercaptohexanol molecule is illustrated in the inset. The scale bars are 4nm (water) and 3nm (RbCl) and colour scale bars are 0.7nm for topography and 20° for phase.