Molecular-scale Hydrophilicity Induced by Solute: Molecular-thick Charged Pancakes of Aqueous Salt Solution on Hydrophobic Carbon-based Surfaces

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We directly observed molecular-thick aqueous salt-solution pancakes on a hydrophobic graphite surface under ambient conditions employing atomic force microscopy. This observation indicates the unexpected molecular-scale hydrophilicity of the salt solution on graphite surfaces, which is different from the macroscopic wetting property of a droplet standing on the graphite surface. Interestingly, the pancakes spontaneously displayed strong positively charged behavior. Theoretical studies showed that the formation of such positively charged pancakes is attributed to cation–p interactions between Na+ ions in the aqueous solution and aromatic rings on the graphite surface, promoting the adsorption of water molecules together with cations onto the graphite surface; i.e., Na+ ions as a medium adsorbed to the graphite surface through cation–p interactions on one side while at the same time bonding to water molecules through hydration interaction on the other side at a molecular scale. These findings suggest that actual interactions regarding carbon-based graphitic surfaces including those of graphene, carbon nanotubes, and biochar may be significantly different from existing theory and they provide new insight into the control of surface wettability, interactions and related physical, chemical and biological processes.

Hydrophilic/hydrophobic interactions are among the most important driving forces of various physical phenomena1–25, such as wetting/dewetting2–5, the folding and native structure formation of protein6–9, drug/molecular delivery10,11, water purification/desalination12,13, molecular recognition14 and nanoparticle assembly/self-assembly in aqueous solution15,16. Conventionally the wetting property of a surface is determined by the macroscopic behavior of water, i.e., the contact angle of water droplets on the surface, and is widely used to analyze the interactions of the solid surface with other materials and dynamic properties at the interface. However, these interactions and dynamic properties are in fact dominated by the behavior of films of water molecules of molecular thickness on the surface. In recent years, molecular-thickness aqueous films have been observed on various hydrophilic surfaces1–4 and between two surfaces17. In contrast, on typical hydrophobic carbon-based surfaces, such as graphene/graphite surfaces, water films are known to only adsorb at extremely low temperature2.

Carbon-based surfaces widely exist as surfaces of both nanoscale and macroscopic materials26–34. Most of these carbon-based surfaces, such as graphite, graphene27,28, carbon nanotubes29,30, fullerenes31, biochar32,33 and activated carbon34, contain a graphitic surface composed of many aromatic rings, which are hexagonal carbon rings rich in p electrons. Moreover, graphitic surfaces rich in aromatic rings widely exist in biomolecules and organic molecules35, humus in the soil36, and polycyclic aromatic hydrocarbons in air pollutants37. It is well recognized that most of these graphitic surfaces rich in aromatic rings are hydrophobic and have wetting properties similar to those of graphite. However, some phenomena that only occur on hydrophilic surfaces have been recently observed on these graphitic carbon-based surfaces32–34,38–40. For example, biochar significantly increases water retention in a sandy soil even though it is well recognized as being hydrophobic32,33. The wetting and charge characteristics of graphene and graphitic surfaces are significantly impacted by the deposited substrates35–37 and
Results and discussion

We deposited a drop of NaCl solution or pure water of millimeter dimensions onto a freshly cleaved highly oriented pyrolytic graphite (HOPG) sheet. In both cases, large macroscopic droplets formed on the graphite surface. The static contact angles (SCAs) of the droplets of NaCl solution and pure water on the HOPG surface were respectively measured as $95^\circ \pm 4^\circ$ (see Fig. 1D) and $93^\circ \pm 2^\circ$ (see Fig. S1D). Aqueous droplets were then blown from the surface with air (see schematic drawings Fig. 1(A–C) for NaCl solution and Supporting Information for pure water). The systems were at room temperature and \( \sim 40\% \) relative humidity.

Usually, many salt particles and aggregates were observed remaining on the graphite if the salt concentration was too high, and the surface was observed to be clean if the concentration was too low. However, at a salt concentration of \( \sim 20 \text{mM} \), we observed many thin films of the salt solution (which we termed pancakes) on the HOPG surface approximately 10 min after treatment with the salt solution in most cases by tapping-mode atomic force microscopy (AFM) imaging as shown in Fig. 1(E) (see details in Supporting Information). The pancakes had apparent height of about 0.6 nm (Fig. 1(F)) in most cases and about 0.3 nm in some cases. The lateral scale of the pancakes spanned from a few hundred nanometers to several micrometers, which is about 3–4 orders of magnitude greater than the height. In one experiment, we determined the probability distribution of the areas of 217 pancakes that we observed (Fig. 1(K)). From this distribution, we computed the average area as \( 0.20 \mu m^2 \). We found that pancakes formed at relative humidity ranging from 30%–70% and that no pancakes formed on surfaces treated only with pure water (see Supporting Information).

The pancakes appeared to move across the graphite surface over time. The AFM images of the surfaces acquired after 1 hour differ from those acquired 10 min after treatment with the salt solution (Fig. 1(F) vs. Fig. 1E). In Fig. 1(F), for example, the pancake has gradually enlarged and there is a visible bulge at the edge. The dis-
tance between pancakes decreased and the pancakes gradually coalesced with one another. These motion behaviors of pancakes enlarging and coalescing reduce the charge distribution and surface energy (see details in Supporting Information).

We also employed non-contact-mode vibrating scanning polarization force microscopy in imaging the HOPG surface treated by the salt solution to minimize the effect of AFM tips. As shown in Fig. S3 in the Supporting Information, the pancakes are again clearly seen to move. The motion of the pancakes was further demonstrated by the repair of a partially damaged pancake and the behavior of a thoroughly removed pancake (see Supporting Information). These observations suggest that the pancakes are liquid.

Remarkably, the salt-solution pancakes showed behavior that they were not charge neutral. The electrostatic force microscopy image obtained with a tip voltage of +3 V (Fig. 2(A)) showed clear and bright pancakes, and these pancakes darkened when the voltage was −3 V (Fig. 2(B)).

From Kelvin probe force microscopy images, we obtained the potentials on the pancakes relative to the substrate. We measured the probability distribution of the electric potentials of 60 pancakes and determined the average pancake potential to be ∼27 mV (Fig. 2, C).

The observed liquid pancakes on HOPG were unexpected since the HOPG surface is hydrophobic and the salt solution would be expected to form droplets (Fig. 1D). As the NaCl droplets dry, the left salt should form particles or aggregates of small particles on HOPG at 40% relative humidity, which is far below the deliquescence humidity of NaCl (around 75% relative humidity at room temperature). This clearly indicates that the hydrophobic graphite surfaces had "apparent" and strong molecular-scale hydrophilicity with respect to the salt solution under ambient conditions.

The key mechanism of the physics of these unexpected charged solution pancakes is the cation−π interactions. The graphite surface adsorbed ions, and water molecules were then adsorbed onto the surface with the help of the ions. To further demonstrate the hydrophobic/hydrophilic transition due to the Na+ adsorption onto graphite, we performed molecular dynamics (MD) simulations with a modification of the cation−π interaction44 of the NaCl solution on graphite. We mimic these behaviors by applying additional acceleration a along the line between the x and z directions on a NaCl drop solution (comprising 810 water molecules and 45 sodium ions and 45 chloride ions) on the graphite surface. For each system, MD simulations were performed for five parallel samples with different initial configurations, with each simulation running for 2 ns. In a typical case shown in Fig. 1(G–I) where a = 0.1 nm/ps², at t = 0.96 ns, some of the solution moved upward and separated from the remaining solution (Fig. 1H) (see the movie in Supporting Information). Interestingly, there was a Cl− ion in the part of the solution departing the surface. The solution pancake remaining on the solid surface (see Fig. 1I) was thus positively charged because of the greater number of Na+ ions. A variation of this charge behavior was observed by Martinez-Martín et al.41 in the adsorption of atmospheric contaminants, such as a polycyclic aromatic hydrocarbon and its isomers, on graphitic surfaces.

Figure 3A shows the total numbers of Cl− and Na+ ions in the solution separating from the graphite surface at different accelerations. There are fewer Na+ ions than Cl− ions in the separated solution and, generally speaking, there are more Na+ and Cl− ions in the separated solution with increasing acceleration. This positive charge behavior of the pancake on the graphite surface clearly results from the interactions between Cl− ions and the graphite surface being much weaker than the interactions between Na+ ions and the graphite surface (i.e., the hydrated Cl−−π interaction (∼1.8 kcal/mol) is only about 1/10 of the hydrated Na+−π interaction (∼16.4 kcal/mol)44).

To study the behavior of the solution remaining on the graphite surface, we further performed MD simulations on the remaining NaCl solution for 4 ns after blowing away the separated solution. From the distribution probability of oxygen atoms along the z direction shown in Fig. 3B for each simulation, we see that there are three peaks at ZO = ∼0.32, ∼0.63 and ∼0.86 nm. The first two peaks are consistent with the experimental observations of the height of the pancake structures of ∼0.3 and ∼0.6 nm. As a increases, the peak at ZO = −0.86 nm reduces and even disappears. Only three and two simulations show clear peaks at a = 0.2 and 0.3 nm/ps², respectively. Thus, the heights of the resulting pancakes depend on the acceleration a. We note that the existence of charged pancakes is robust as can be clearly seen at all four accelerations (a = 0.10, 0.15, 0.20 and 0.30 nm/ps²) in our simulations. This further demonstrates the consistency between the MD simulations and experimental observations.

We also investigated the main reason for the inconsistency in the wetting property between molecular and macroscopic scales. The flat graphite surface on the macroscopic scale is made up of a large number of graphene flakes stacking on a molecular surface. The macroscopic flat graphite surface is actually formed by a large number of molecular-level graphene layers, and there are many steps between the layers (see Fig. 1(E, F) and Fig. 2(A, B)). Theoretical calculations based on density functional theory (see details in Supporting Information) show that the Na+ ions diffuse easily on the graphene flakes (barrier energies of only ∼3 kcal/mol, Fig. S5B) but with more difficulty across the steps between layers (barrier energies exceeding 100 kcal/mol, Fig. S5B). Thus, Na+ ions bind at the aromatic rings on the graphite surface retaining their hydration water molecules, resulting in the formation of molecular-thick pancakes of aqueous salt solution on the molecular-level graphene layers (see Fig. 1(E, F)), yet the drop of salt solution remains intact because there are many molecular-scale-layer steps hindering the diffusion of Na+ ions on the macroscopic surface.

Moreover, the many hydrophobic steps reduce the surface energy of the graphite surface (Fig. S6), resulting in the macroscopic graphite surface being more hydrophobic than the molecular-scale graphene flakes (see details in Supporting Information). It is much
scale53, which also affects the wetting property at different (macroscopic and molecular) scales. In summary, the different properties of the graphitic surface and Na⁺ ion distributions between molecular and macroscopic scales produce the inconsistency in the wetting property between molecular (hydrophilicity) and macroscopic (hydrophobicity) scales.

Conclusion

We conclude that, counter to intuition, molecular-thick films of aqueous salt solution can stably exist on a hydrophobic carbon-based surface under ambient conditions. Experimental and theoretical results show the unexpected molecular-scale hydrophilicity on hydrophobic surfaces. The cation–π interactions and the different properties of the graphitic surface and Na⁺ ion distributions between molecular and macroscopic scales generate an inconsistency in the wetting property between molecular (hydrophilicity) and macroscopic (hydrophobicity) scales. Interestingly, the formed pancakes spontaneously display positively charged behavior. Considering that the key ingredient in the production of molecular-scale pancakes, the aromatic rings, is commonly found in biomolecules, the findings may clarify the actual interactions of biomolecules. The underlying mechanism should be helpful in understanding and controlling the functional characteristics of carbon-based materials for various applications such as drug delivery, water purification, employing carbon nanotubes, ion filtration employing graphene pores, hydrogen storage employing graphene/graphite, and other applications of carbon-based nanomaterials.

Methods

Experimental Section. Materials. NaCl (crystal purity 99.99%) was purchased from Sinopharm Chemical Reagent Co., Ltd., and was dissolved in ELGA laboratory water to a final concentration of 20 mM. The highly oriented pyrolytic graphite (HOPG) was provided by Molecular Devices and Tools for Nano Technology Co. Zelenograd, Moscow, Russia. The electric conductive adhesive (DAD-40) was purchased from the Shanghai Research Institute of Synthetic Resins.

Sample Preparation. The HOPG fragment was fixed to the sample holder using electric conductive adhesive with the working side up, and was freshly cleaved using a single-crystal diamond knife. The NaCl sample was prepared according to a process previously reported for the observation of liquid nanodroplets of KOH by scanning polarization force microscopy on graphite. Briefly, a drop (~20 μl) of NaCl solution (20 mM) was deposited on the HOPG substrate, and the droplet was then dried with a stream of air at room temperature and 40%–60% relative humidity. The relative humidity and temperature of a sample chamber (SDH-01N, Shanghai Jianheng Instrument Co.) were controlled with accuracy of 5% and 0.1°C, respectively. The as-prepared sample was placed in this chamber for a given amount of time.

Atomic Force Microscope (AFM) Imaging. Experiments were performed on a commercial AFM (Nanoscope IIIa, Veeco/Digital Instruments, Santa Barbara, CA, USA) equipped with a J scanner (100 μm × 100 μm) and E scanner (15 μm × 15 μm). Silicon etched probes (NSC18/Ti-Pt, MikroMasch Co., length: 230 μm, width: 40 μm, thickness: 3 μm, nominated spring constant: 3.5 N/m, resonant frequency: 60–90 kHz) were used in other experiments. The Ti-Pt coating comprised a 10-nm Pt layer on a 20-nm Ti sublayer, which provided greater adhesion and electromigration firmness than if using Pt alone. The Ti-Pt coating formed on both the tip and reflective side of the cantilever. The resulting tip radius with the coating was 40 nm. The morphological features of the AFM images, namely the height and width, were analyzed using AFM accessory software (ver. 7.30). All AFM images were adequately flattened using the software to correct the distortion at a micrometer scale, but no other digital operation was carried out. All AFM data were obtained at room temperature, whereas relative humidity was measured by a hygrometer with accuracy of 5% (SDH-01N, Shanghai Jianheng Instrument Co.).

SCA Measurement. The SCA measurement was made on an Attension Theta system (KSV Instruments Ltd., Finland). The volume of each droplet of NaCl solution or pure water was ~5 μl and each droplet was carefully touched to the sample surface. A digital camera was used to take images of all droplets, and the values of SCA were automatically computed by the supplied calculation software. Each HOPG sample was measured at three different points and the average value was reported.

Computational Methods. The cation–π interactions between Na⁺ and the graphite surface are represented by the model potential

\[
V = \varepsilon \left( z_1 - z_2 \right)^2 - \frac{2 \varepsilon z_1 z_2}{z_1 + z_2},
\]

where the parameters \( \varepsilon \) and \( z_0 \) are the adsorption energy and balance position (the vertical distance between the Na⁺ ion and the surface) of Na⁺ relative to the graphite surface and \( z \) is the vertical distance between the Na⁺ ion and the surface; these are the main potential parameters describing the cation–π interaction. Their values are \( \varepsilon = 3.8 \text{ Å} \) and \( z_0 = 16.4 \text{ kcal/mol} \). Molecular dynamics simulations were carried out using the program NAMD2/VMD,9 packages;60 with the CHARM force field,61 at time steps of 2 fs and with the O-H bonds and C atoms held fixed (see detail in Supporting Information).

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Author contributions

G.S. and T.L. performed molecular dynamics simulations. H.F. and G.S. carried out most of the theoretical analysis. Y.S. and J.H. designed and observed the experimental investigation. C.W. and B.S. carried out some theoretical analysis. Y.W. carried out some experimental analysis. H.F., J.H. and G.S. contributed most of the ideas and wrote the paper. All authors discussed the results and commented on the manuscript.

Additional information

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