The viscoelectric effect concerns the increase in viscosity of a polar liquid in an electric field due to its interaction with the dipolar molecules and was first determined for polar organic liquids more than 80 years ago. For the case of water, however, the most common polar liquid, direct measurement of the viscoelectric effect is challenging and has not to date been carried out, despite its importance in a wide range of electrokinetic and flow effects. In consequence, estimates of its magnitude for water vary by more than three orders of magnitude. Here, we measure the viscoelectric effect in water directly using a surface force balance by measuring the dynamic approach of two molecularly smooth surfaces with a controlled, uniform electric field between them across highly purified water. As the water is squeezed out of the gap between the approaching surfaces, viscous damping dominates the approach dynamics; this is modulated by the viscoelectric effect under the electric field across the water, enabling its magnitude to be directly determined as a function of the field. We measured a value for this magnitude, which differs by one and by two orders of magnitude, respectively, from its highest and lowest previously estimated values.

viscoelectric effect | electrokinetic phenomena | viscosity/electric field coupling | surface forces balance

The viscoelectric effect concerns the change in the viscosity of polar liquids in the presence of an electric field (1–3). It arises from the interaction of the field with the dipolar molecules, and while its molecular origins are still not well understood (4–6), it has considerable relevance in areas ranging from surface potential measurements (7–9) and boundary lubrication (10) to nanofluidics and its applications (11–13). Knowing the magnitude of the viscoelectric effect is thus of clear importance. It was first measured by Andrade and Dodd (1–3) for a range of polar organic liquids, by monitoring their flow in a narrow channel between metal electrodes across which a known electric field \( E \) was applied, and quantified via a viscoelectric coefficient \( f \) using an empirical relation based on their results:

\[
\eta(E) = \eta_0 (1 + f|E|^2),
\]

[1]

a simplified analysis leading to such a relation is given in Ref. (8). Here, \( \eta_0 \) is the unperturbed bulk liquid viscosity (i.e., in the absence of any field). For the case of water, however, the most ubiquitous and important polar liquid, measurement of its viscosity in the presence of a strong, uniform field presents a strong challenge (as discussed later in this section), and to our knowledge no such direct measurements have been reported. Over the past six decades, therefore, the magnitude of the viscoelectric effect in water has been only indirectly estimated by extrapolation from its values for organic liquids (8), from estimates of its effect on electrokinetic phenomena (11, 14–19), or by other approaches (7, 12, 20, 21). These estimated values, as expressed in the viscoelectric coefficient \( f \), vary over more than three orders of magnitude, ranging from \( f \approx 10^{-17} \) to \( 2.5 \times 10^{-14} \) (V/m)^2 \((S1 \text{ Appendix, Section 7)}\). For completeness, we note that results contradictory to the viscoelectric model have also been reported (22) (i.e., suggesting a decreased water viscosity in an electric field). The reasons for the large span of these estimated \( f \) values were attributed to various factors such as solid/liquid coupling, varying ionic sizes, and varying water permittivity (12, 19); however, while these factors may play some role, there is no evidence that they could lead to such large discrepancies.

We believe, rather, that the origin of the large variance in the estimated magnitude of the viscoelectric effect arises because none of the experimental studies on water to date in which the \( f \) values were estimated was direct, in the sense of probing how the water viscosity varied with field in a uniform electric field. In all cases, viscosity changes were presumed to occur only in the non-uniform, rapidly decaying electrostatic potential near charged surfaces immersed in water. Changes in electrophoretic mobility, electro-osmosis, or hydrodynamic dissipation or water mobility between similarly charged solid surfaces were then attributed to some mean viscosity increase in these thin surface-adjacent layers (7, 11, 12, 14–21). In practice, however, the effect on these electrokinetic phenomena of viscosity or water mobility changes in the thin layers where such nonuniform, rapidly decaying fields are present is not easy to quantify reliably, especially in the presence of salt ions (12). At the same time, measuring the viscosity of water in a uniform electric field between two surfaces at different potentials, as was done for the polar organic solvents (2, 3) and which would provide a direct determination of its viscoelectric effect, presents a considerable difficulty. This is due to two main factors and arises because, in contrast to organic solvents, water may self-dissociate. Firstly, the potential difference that

Significance

The viscosity of polar liquids increases in an electric field because of its interaction with the dipolar molecules. This viscoelectric effect was measured for organic liquids, but for water, the most important polar liquid, where it is crucial in areas from surface potential determination to nanofluidic applications, it is very challenging, and no direct measurements have been carried out to date. Consequently, estimates of its magnitude in water vary by more than a thousandfold. Here, we use a surface force balance to measure the dynamic approach of two molecularly smooth surfaces with a uniform, controlled electric field between them across water; this is modulated by the water viscosity and hence the viscoelectric effect, enabling its magnitude to be directly determined.

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may be applied between the surfaces across water is limited, if electrolysis is to be avoided (23, 24), and secondly, electrostatic screening implies that the field decays strongly (within a Debye screening length) away from the surfaces (25–27). Even in purified water with no added salt (as in the present study), the potential decays rapidly away from a charged surface (see, e.g., Fig. 1C), so that to measure viscosity in a uniform field between two surfaces, one would require flow channels of width of order some tens of nanometers or less, presenting a major challenge.

In the present study, we overcome this by directly probing the viscosity of purified water across which a uniform electric field acts while it is confined between two surfaces in a surface force balance (SFB). In our experiments, a molecularly smooth gold surface at a controlled (positive) surface potential approaches an atomically smooth mica surface at constant surface (negative) charge density, so that a known electric field acts across the water-filled gap of width $D$ between them; moreover, this field is very close to uniform at the most relevant surface separations ($D \leq 30$ nm, Fig. 1C). The dynamics of approach is strongly modulated by the viscous damping due to squeeze-out of the water as $D$ decreases, and hence by its viscosity in the uniform electric field; by monitoring the approach rate of the surfaces at high temporal (millisecond) and spatial (approximately angstrom) resolutions, we are able therefore to directly evaluate the magnitude of the viscoelectric effect (the value of $f$).

Results and Discussion

Electric Field Between Interacting Surfaces across Water. The SFB has been traditionally used to measure interactions between two single-crystal, atomically smooth mica surfaces as a function of their separation $D$, where, in water, each mica surface is negatively charged because of the dissociation of potassium ions (25, 26, 28), corresponding to a constant-charge surface with charge density $\sigma_{\text{mica}}$. Detailed descriptions of the SFB appear elsewhere (28, 29). More recently, by replacing one of the mica sheets with a template-stripped, molecularly smooth gold surface (30) (rms roughness $\sim 2$ Å, SI Appendix, Section 6; Fig. 2A) and controlling the potential $\psi_{\text{gold}}$ of the gold with a potentiostat in a piezo crystal, on which the top (gold) surface is mounted, at a speed $v_{\text{app}} < 10$ nm/s. This drives the gold surface toward the lower mica surface, which is mounted on a spring of constant $k$. Onset of the long-range electrostatic attraction is observed at surface separations $D \sim 100$ nm, and at lower $D$ values, the surfaces jump into contact because of an Euler-like spring-instability (33).

During the approach, the separation $D(t)$ is obtained from the interference fringes of equal chromatic order (FECO, Fig. 2B), using fast video recording ($\sim 10^3$ fps) at millisecond intervals. The typical noise in $D(t)$ is $\sim 2$ Å. The interaction between the surfaces as they approach arises from both conservative forces, i.e., electrostatic forces $F_{\text{PB}}$ described by the Poisson-Boltzmann (PB) equation and van der Waals (vdW) forces between the gold and mica surfaces across water, $F_{\text{vdw}}$, together with hydrodynamic forces due to the finite approach speed of the surfaces. The relative motion of the surfaces during the approach obeys the following equation, as previously described (33):

$$F_{\text{tot}}(D) = m \left( \frac{d^2 D}{dt^2} \right) - k(D_0 - D(t)) + v_{\text{app}} t$$

$$+ 6\pi R^2 \eta \left[ \frac{dD}{dt} / D(t) \right]$$  [2]

The term on the left is the total conservative surface force $F_{\text{tot}} = F_{\text{vdw}} + F_{\text{PB}}$ (SI Appendix, section 1), which is balanced by the inertial term (first term on right), the force due to distortion of the spring from its unperturbed state at separation $D_0$ at $t = 0$ (second term on right), and the hydrodynamic resistance (third term on right), where $R$ is the mean radius of curvature of the curved mica and gold surfaces and $\eta$ is the water viscosity in the gap, which depends on the electric field through the viscoelectric effect (Eq. 1). The inertial term (where $m \sim 2$ gm is the mass of the lower lens) is readily shown to be negligible compared to the other terms (33). In this study, the viscoelastic effect (34–37),

Purified water with no added salt (Materials and Methods) was added to the SFB bath immersing the mica and gold surfaces. When a positive potential is applied to the gold, a long-range electrostatic attraction acts between the surfaces, as previously observed (28, 32). The dynamics of approach was initiated (at time $t = 0$, at a separation $D = D_0 > 300$ nm, larger than the range of any surface interactions) by imposing a steady extension on the spring from its unperturbed state at separation $D_0$, i.e., electrostatic forces $F_{\text{PB}}$ described by the Poisson-Boltzmann (PB) equation and van der Waals (vdW) forces between the gold and mica surfaces across water, $F_{\text{vdw}}$, together with hydrodynamic forces due to the finite approach speed of the surfaces. The relative motion of the surfaces during the approach obeys the following equation, as previously described (33):

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The unperturbed viscosity of water. The smooth curve is the predicted curve on the proton concentration in the immediate vicinity of the gold surface. The working electrode is the gold surface coating a cylindrical lens, mounted on the piezoelectric tube (PZT), facing the back-silvered mica surface glued on a similar lens, in a crossed-cylinder configuration, supported by the spring \( k \). Images of FECO formed by white light interference between the gold and the silver layers were recorded at high frame rates (up to 1,000 fps) during the approach of the surfaces. Separation (D) was evaluated with subnanometer precision from the wavelengths of the fringe tips obtained from a double Gaussian fit to the pixel intensity profile (FECO image shown here taken at 100 fps for better contrast). An electric potential (controlled by a potentiostat) is applied to the gold surface via a three-electrode configuration. The counter (C) and reference (R) electrodes were constructed with platinum wires. The working (W) electrode is the gold surface. \( \psi_{\text{applied}} \) is imposed by the potentiostat.

which describes the resistance to the flow of the counterions in the diffuse double layers adjacent to the gold and mica surfaces (and should not be confused with the viscoelectric effect, which is the subject of our study), is negligible because of the low ion concentration and may therefore be safely ignored in the hydrodynamic resistance term (SI Appendix, Section 4). The electric field in the gap \( E(D) \) can be evaluated by the PB equation for given \( \psi_{\text{gold}} \) and \( \sigma_{\text{mica}} \). \( \psi_{\text{gold}} \) is unknown a priori from the applied potential on the gold because of a pseudo electrode (Fig. 2C) but may be evaluated from the force versus distance profile \( F(D) \) as the surfaces approach (32); \( \sigma_{\text{mica}} \) is controlled by the equilibrium of surface cation adsorption, mostly dependent on the proton concentration in the immediate vicinity of the mica (26), and may likewise be determined from the \( F(D) \) profile (32).

Typical force versus separation approach profiles are shown in Fig. 3A and B (normalized as \( F_{\text{applied}}/R \) in the Derjaguin approximation), evaluated from the experimentally measured \( D(t) \) via Eq. 2 on the assumption that the viscosity is at its unperturbed (i.e., zero field) bulk water value \( \eta_0 \) throughout. This is a good approximation at separations \( D > 40 \text{ nm} \), where the mean electric field between the mica and gold surfaces is so low (see Fig. 1B) that differences between \( \eta \) and \( \eta_0 \) are negligible at all \( D > 40 \text{ nm} \). This allows the unknown parameters \( \sigma_{\text{mica}}, \psi_{\text{gold}}, \) and bulk concentration \( \psi_0 \) to be extracted through comparison with solutions of the PB equation (together with vdW forces) for \( D > 40 \text{ nm} \) (details in Materials and Methods and SI Appendix, Section 1). The resulting theoretical fits are shown in Fig. 3 A and B; the deviation at \( D < \sim 40 \text{ nm} \) between the experimental curve deduced from Eq. 2 and the theoretical force fit based on the PB and vdW equations is due entirely to the fact that the viscoelectric effect is not taken into account in Eq. 2, as confirmed in the next section. Inset in Fig. 3B is shown for comparison data from an earlier study (32) of interactions between mica and gold at the same applied potentials on the gold (0.2 V), in which \( \sigma_{\text{mica}}, \psi_{\text{gold}}, \) and bulk concentration \( \psi_0 \) were independently determined and were evaluated from Eq. 2 ignoring the viscoelectric effect. These data show, as expected, close similarity to the previous study at \( D > \sim 40 \text{ nm} \) (the slight differences at lower \( D \) values are likely due to the much lower video acquisition rate and thus the time resolution in the earlier work).

Using these parameters obtained from the force profiles in Fig. 3, we may evaluate the potential on the mica surface \( \psi_{\text{mica}} \), the charge density on the gold surface \( \sigma_{\text{gold}} \), and the potential in the gap \( \psi(d) (0 < d < D) \) by numerically solving the nonlinearized PB equation (SI Appendix, Section 1) at varying separations \( D \), as shown in Fig. 1.

Fig. 4A shows that as the separation \( D \) decreases, \( \sigma_{\text{gold}} \) approaches \( \sigma_{\text{mica}} \) and \( \psi_{\text{mica}} \) approaches \( \psi_{\text{gold}} \) as earlier discussed (28). Ultimately the two surfaces mutually neutralize when in contact. The mean electric field \( E = |\psi_{\text{gold}} - \psi_{\text{mica}}|/D \) is shown in Fig. 1B. Although \( \psi_{\text{mica}} \) becomes less negative as \( D \) decreases, \( E \) increases because of the hyperbolically increasing factor \( 1/D \) and eventually plateaus as \( D \) approaches zero. For our analysis with the viscoelectric model (Eq. 1), \( E = E' \) is an excellent approximation at small separations as \( \psi(d) \) becomes more linear in trend (Fig. 1C) so that the field is essentially uniform for \( D < \sim 40 \text{ nm} \). At large separations, our fits to Eq. 2 are seen to be justified by the small magnitude of \( E \); at an order of 0.004 V/nm or less, water viscosity is enhanced by \( \sim 1.6\% \) or less, assuming a viscoelectric coefficient of order \( 10^{-15} \text{ m}^4/\text{N}^2 \) (see next section), which, as noted earlier, has negligible impact on the approach dynamics at \( D > 40 \text{ nm} \).

Fig. 3. (A, B) Representative force versus surface separation profiles evaluated from the experimentally measured \( D(t) \) variation via Eq. 2, assuming unperturbed viscosity of water. The smooth curve is the predicted \( F_{\text{applied}} + F_{\text{vdW}} \) fitted to the region \( D > \sim 40 \text{ nm} \) and used to extract values of \( \sigma_{\text{mica}}, \psi_0, \) and \( \psi \) (see SI Appendix for details). The inset in B compares typical force profiles in our study (gray and red curves) with results from an earlier study by Tivony et al. (32) (blue dots) between mica and gold surfaces at the same applied gold potential (0.2 V), showing close correspondence for \( D > \sim 40 \text{ nm} \).
Magnitude of Viscoelectric Effect Determined from Surface Approach Dynamics. To analyze the effect of the viscoelectric effect on the dynamics of approach, we evaluated $\frac{dD}{dt}(D)$, Eq. 3—obtained by rearranging Eq. 2—from the $D(t)$ variation determined directly from the video recording of the approach:

$$\frac{dD}{dt} = \left( F_{\text{tot}} D + kDv_D D - kD^2 + kv_{\text{app}} Dr \right) \frac{1}{6\pi R^2 \eta (1 + f E^2)^2}.$$  \[3\]

The damping term now manifests the viscoelectric effect in the enhanced value of $\eta = \eta_0 (1 + f \cdot E^2)$. The advantage of using $dD/dt(D)$ as the metric as opposed to $F(D)$ is that, given that $D(t)$ is the primary measurable through the FECO position (Fig. 3B), the error propagated from $D$ is minimized by allowing only one linear operation on the parameter.

Fig. 4 shows several typical $dD/dt(D)$ profiles, taken at different extracted values of the surface parameters, all showing these common features: at large separations, $dD/dt(D)$ is equal to the driving speed $v_{\text{app}}$ of the piezo crystal ($<10$ nm/s), while at smaller separations, $D < \sim 80$ nm, the approach velocity increases under the increasing electrostatic attraction while being modulated by the viscosity-dependent hydrodynamic stress as in Eq. 3. At $D = D_{\text{min}} \sim 10$ to 20 nm, a local minimum in $dD/dt(D)$ is seen due to the decreasing driving term $F_{\text{tot}} \cdot D$ at small distances. Physically, this may be understood as follows: The electrostatic component of the attractive force $F_{\text{tot}}$ (Eq. 2) arises largely because of the expulsion of counterion pairs between the oppositely charged surfaces (38, 39), and its magnitude, for surface separations $D$ of order the Debye screening length or less for our sphere-on-flat geometry, scales approximately as $(1/\eta D)^2$ (38), while the damping term continues to increase as $(1/D)$ (39). This is similar to the $D$ variation of the damping term in Eq. 2, so that the surfaces initially accelerate toward each other under the increasing attractive force. For $D < \sim 30$ nm in the conditions of our experiments, however, solution of the PB equation shows that the attraction varies more weakly than $(1/D)$ (38), while the damping term continues to increase as $(1/D)$, leading to the deceleration and eventual decrease in $dD/dt$ (SI Appendix, section 5). The two pairs of figure panels, Fig. 4A and B and C and D show $dD/dt(D)$ profiles measured in two independent experiments but at the same contact point in each experiment (i.e., identical values of $c_p$ and $\sigma_{\text{nica}}$ for each pair of figures, different for each pair), with only the potential applied on the gold surface being varied. A higher applied potential on gold, from 0.07 V in Fig. 4A to 0.1 V in Fig. 4B and from 0.05 V in Fig. 4C to 0.08 V in Fig. 4D, results in higher jump-in speed at all distances as expected and $\sim 70$ nm/s and 85 nm/s differences, respectively.
at the \( \frac{dD}{dt}(D) \) local minima of Fig. 4 B and D compared with Fig. 4 A and C.

To evaluate the viscoelectric coefficient \( f \), first we evaluated \( D(t) \) and \( \frac{dD}{dt}(D) \) with \( f = 0 \) from Eq. 3 using the theoretically predicted \( F_{\text{dip}} \) with values of \( \eta_{\text{mica}}, \psi_{\text{gold}}, \) and \( \eta_b \) extracted directly from the large \( D > 40 \text{ nm} \) force profile as described previously. We then derive

\[
\eta = \frac{dD/dt}{d\psi/d\psi} = g_{\text{exp}} \quad \text{where} \\
g = \left( F_{\text{net}}D + k_D D - k_D^2 + k_{\text{app}}D \right),
\]

which can be evaluated for the range of \( E \) values in the most relevant distances \( 8 \text{ nm} < D < 30 \text{ nm} \). A plot of \( n/\eta_0 \) versus \( E^2 \) is shown in Fig. 4 E, using values extracted from the \( dD/dt \) curves with least noise (Fig. 4 A and B) for \( (n/\eta_0) \) and from the PB equation (for \( E^2 \), Fig. 1 B). Data were binned at equally spaced intervals of \( E^2 \) and averaged. From Eq. 1, the slope of this plot is simply the viscoelectric coefficient \( f \), and its best fit value is \( f = 9.9 \pm 2.8 \times 10^{-16} \text{ m}^2/\text{V}^2 \). To confirm this, \( dD/dt(D) \) was solved with this value of \( f \) for each of the \( dD/dt \) curves in Fig. 4 A–D, showing excellent agreement (as also for additional \( dD/dt(D) \) profiles in SI Appendix, Section 2). In particular, this is seen comparing the fits of Eq. 3 to the data in Fig. 4 A and B and C and D, with each pair taken at the same contact point (in independent experiments) but at different \( \psi_{\text{gold}} \); crucially, the best fits to the experimental \( dD/dt(D) \) data are associated with the same \( f \) value for all four \( dD/dt(D) \) profiles, even though the uniform fields across the gap at \( D < \sim 40 \text{ nm} \) are significantly different. For comparison, the predicted curves according to Eq. 3 but using \( f \) values corresponding to the highest and lowest literature estimates to date (SI Appendix, Section 7) are also shown in Fig. 4 A, showing clearly that they do not fit the data [similar misfits of these extremal \( f \) values to the data apply to the other \( dD/dt(D) \) profiles but are omitted for clarity]. Finally, in Fig. 4 F, we revisit the force profiles \( F/R \) versus \( D, \psi \), for which the discrepancy between theory and the forces evaluated from Eq. 2 for \( D < \sim 40 \text{ nm} \) arose from ignoring the viscoelectric effect (i.e., setting \( f = 0 \) in Eq. 2): We see that once the directly determined value of \( f \) is inserted into Eq. 2, there is close agreement (within the scatter) between the evaluated forces and the theoretical prediction, further supporting our determination of the viscoelectric coefficient.

Conclusions

We have measured the viscoelectric effect in water by directly determining the changes in water viscosity under uniform electric fields through its modulation of the hydrodynamic damping between two approaching surfaces. Our result differs by up to one to two orders of magnitude from many of the earlier indirect estimations (7, 11, 12, 14–21), which were in the range \( 10^{-14} – 10^{-17} \) (SI Appendix, Section 7) (though some of these indirect estimates, e.g., Refs. 7 and 8, were comparable to our measured one). Molecular dynamics simulation results suggest lower \( f \) values at fields much higher than in our study (4, 5, 40) and also that the relation of Eq. 1 may not apply at such fields, while our results (Fig. 4 E) are consistent with Eq. 1. We do not know the reason for these discrepancies between experiment and MD simulations, though we should note that the fields used in the simulations could not be implemented over the length scales of water thickness in our experiments (10 to 30 nm), as electrolysis would occur (41); for this reason, comparison may not be appropriate. The main difference between our direct determination and previous estimates is that the latter were based on leveraging the electric field from the rapidly decaying potential gradient of double layers formed at a surface (or the field between two similar surfaces, which switches sign at the midplane due to symmetry). The analysis in those cases was based on an averaged electric field in the thin interfacial layer or in the intersurface gap (within which it varied very strongly), while in our study, the intersurface potential variation was very closely linear and its gradient—the electric field—therefore uniform across the water. We believe therefore that our measured value may be viewed as a direct measurement of the viscoelectric effect in water, conceptually similar to the direct measurements of this effect in polar organic liquids commenced over 80 y ago (1–3) but differing from all previous estimates of this effect in water, none of which involved measuring viscosity changes in uniform fields. It thus provides a benchmark of the viscoelectric effect, a basic phenomenon associated with coupling of the electric field to the dipole of water molecules, after decades of estimates covering a range of values differing by over a thousandfold. Improved characterization of the viscoelectric effect may shed light on many electrokinetic problems and help interpret boundary effects in aqueous environment, which has significant impact in real-life applications, as well as underpinning attempts at a better understanding of its molecular origins (4–6).

Materials and Methods

SFB. As illustrated in Fig. 2 A, two plano-cylindrical glass lenses are arranged in a cross-cylinder configuration, with interaction geometry equivalent to a sphere approaching a flat surface. The upper lens is coated with a gold surface prepared through template stripping (30). The lower lens is covered by a single-crystal, back-silvered mica sheet. The upper lens is driven by the piezo crystal in a steady motion enabled by a function generator at a speed \( \nu_{\text{app}} < 10 \text{ mm/s} \) toward the lower lens, mounted on a spring (constant \( k = 89 \text{ mN/mm} \)), which bends in response to the total force between the surfaces. Normal force measurements were acquired from the bending of the spring \( k \) and by analyzing the hydrodynamics (Eq. 2). The three-electrode configuration (Fig. 2 C) enables positive potentials to be imposed onto the gold surface through a potentiostat when the platinum electrodes are positioned in the water immersing the surfaces, contained in a quartz bath to prevent short to ground. Further details of the technique are provided elsewhere (28, 32).

Preparation of Template-Striped Gold Surface. The gold surfaces were prepared by evaporating 99.999% pure gold pellet (Kurt J. Lesker and Materion) \( \sim\) leaving a few nm of mica, \( \sim\) 30 nm. A plot of \( R \) versus \( D \) for each of the \( dD/dt \) curves in Fig. 4 A–D, showing excellent agreement (as also for additional \( dD/dt(D) \) profiles in SI Appendix, Section 2). In particular, this is seen comparing the fits of Eq. 3 to the data in Fig. 4 A and B and C and D, with each pair taken at the same contact point (in independent experiments) but at different \( \psi_{\text{gold}} \); crucially, the best fits to the experimental \( dD/dt(D) \) data are associated with the same \( f \) value for all four \( dD/dt(D) \) profiles, even though the uniform fields across the gap at \( D < \sim 40 \text{ nm} \) are significantly different. For comparison, the predicted curves according to Eq. 3 but using \( f \) values corresponding to the highest and lowest literature estimates to date (SI Appendix, Section 7) are also shown in Fig. 4 A, showing clearly that they do not fit the data [similar misfits of these extremal \( f \) values to the data apply to the other \( dD/dt(D) \) profiles but are omitted for clarity]. Finally, in Fig. 4 F, we revisit the force profiles \( F/R \) versus \( D, \psi \), for which the discrepancy between theory and the forces evaluated from Eq. 2 for \( D < \sim 40 \text{ nm} \) arose from ignoring the viscoelectric effect (i.e., setting \( f = 0 \) in Eq. 2): We see that once the directly determined value of \( f \) is inserted into Eq. 2, there is close agreement (within the scatter) between the evaluated forces and the theoretical prediction, further supporting our determination of the viscoelectric coefficient.

Water Purification. Water was purified with the Barnstead GenPure Pro Water Purification System, using high-performance ultraviolet assembly to reduce total organic content (TOC) to ultralow levels. The resistivity of the purified water is 18.2 MΩ · cm at 25°C with TOC < 1 ppb.

Image Acquisition and Analysis. A Hamamatsu Fusion C14440-20UP camera was set up to capture image sequence of FECO directly from the spectrometer without any lens in between. A calibration image was captured with a mercury (neon) calibration lamp, from which a linear relation between the pixel position and the wavelength was fitted. The image resolution is ~0.4 Å in wavelength per pixel, and the best achieved resolution in surface-to-surface separation per pixel is 2.3 Å with mica leaflets thinner than 3 μm. During the jump-in events, images were acquired at 993 fps. The pixel position of the fringe tip, corresponding to the minimal distance between the lenses, was evaluated by fitting a double Gaussian function to the pixel intensity of the fringe doublet (Fig. 2B). The separation \( D \) was evaluated using the multilayer matrix method, as described in previous works (30, 32, 42, 43).

Post-Data-Acquisition Processing. To reduce the noise in the separation \( D(t) \) trajectory, multiple recordings were acquired consecutively at the same contact point for each applied gold surface potential. \( D(t) \) was smoothed with a moving average filter at 0.02 s intervals, chosen carefully to avoid artifacts from smoothing. \( D(t) / dt \) was computed and averaged over repeated trajectories to reduce noise (SI Appendix, Section 3). Note that, as demonstrated
previously, charge inversion on the gold surface alone neutralizes the mica surface charge when in contact. Therefore, hysteresis observed in consecutive jump-in events of mica–mica systems due to unequivilibrated charge release (44) is not observed in the gold–mica system (28).

Fitting Parameters in PB Equation. The nonlinearized PB equation with constant potential on the gold surface and constant charge on the mica surface was solved numerically for more than 2,000 combinations of $\psi_{\text{mica}}(D) = \psi_{\text{mica}}(D_0)$, and bulk concentration $c_b$, to sample $\psi_{\text{mica}}(D) = \psi_{\text{mica}}(D_0)$ at 0.01 V intervals, $\psi_{\text{gold}}(D)$ at 0.01 V intervals, and bulk concentration $c_b$ from $5 \times 10^{-7}$ to $8 \times 10^{-3}$ M at 1 × $10^{-6}$ M intervals. Each PB solution was augmented by van der Waals force $F_{\text{vdW}}(D)$ with Hamaker constant $A_H = 9.0 \times 10^{-20}$ J (28, 32). The total force $F_{\text{tot}}$ was compared to the normal force profile, i.e., the right-hand side of Eq. 2 ($|\text{f} - \text{r}|$), which was evaluated using the smoothed $|\text{f}|$ as described previously, averaged over repeated profiles, and smoothed with a 0.05 s interval moving average filter. Best fit was selected using the least mean-square deviation at large separations ($D > 40$ nm) where the viscoelastic effect is small, subject only to the constraint $\psi = 0$. For each selected set of parameters, $\psi_{\text{mica}}(D)$ and $\psi_{\text{gold}}(D)$ are evaluated for demonstrating charge inversion and for estimating the electric field $E(D)$ at varying separations (Fig. 1 A and C, and SI Appendix, section 1). Uncertainties in these parameters manifest as uncertainties in the evaluated electric field and are indicated as error bars in the data of Fig. 4E.

Numerical Solution to the Equation of Motion. Eq. 3 was made dimensionless by introducing

$$D = D^* L, \quad \psi_{\text{app}} = \psi_{\text{app}} U, \quad F_{\text{tot}} = F_{\text{tot}} D^*, \quad k = k^* F, \quad t = t^* L^* D^*,$$

which gives

$$\frac{dD^*}{dt^*} = N_0 \left( F_{\text{tot}} D^* + k^* D^*_0 D^* - k^* D^* - k^* \psi_{\text{app}} D^* \right)^\tau - \frac{F_L}{6\pi \eta L_0 (1 + \tau^2)}$$

[5]

where the dimensionless number $N_0$ indicates the relative scale of the surface forces compared to the viscous force. Eq. 5 was solved numerically using the second-order Crank-Nicolson method (45), taking $F_{\text{Tot}}(D)$ and the corresponding $E(D)$ as inputs. $\psi_{\text{app}}$ was evaluated by linear fits to the experimentally measured $D(t)$ at large separations $D > 200$ nm, where $F_{\text{Tot}}$ and spring force are mostly absent. Parameters used were spring constant $k = 89$ N/m, lens radius $R = 0.01$ m, bulk water viscosity at room temperature $\eta_0 = 8.9 \times 10^{-4}$ Pa s, and initial separation $D_0 = 500$ nm. To avoid spurious computational errors due to the exponentially growing $|\psi_{\text{Tot}}|$ as $D$ decreases, the dimensionless time step $\Delta t$, at each discrete step $n$ was adjusted as

$$\Delta t_n = \frac{\text{l}_n}{\text{Un}_n} \times 10^{-5} s^{-1},$$

and by updating the force scale $F$, the length scale $L$, and the velocity scale $U$ such that

$$L_n = D^*_n \times L_0, \quad U_n = \frac{D^*_n - D^*_0}{\Delta t_n} \times \frac{U_0}{\Delta t_0}, \quad F_n = F_{\text{Tot}}(D^*_n, L_n)$$

where the initial values are $L_0 = D_0$, $U_0 = \psi_{\text{app}}$, and $F_0 = F_{\text{Tot}}(D_0)$.

Data Availability. All study data are included in the article and/or supporting information.

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