How clay delamination supports aseismic slip

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ABSTRACT

Aseismic slip is a stable fault slip, which allows strain to be relieved smoothly. Aseismic slip prevents the earthquake propagation, but it could nucleate an earthquake elsewhere. Understanding the mechanism of aseismic slip is promising in revealing the seismic cycle. Experimental evidence showed clay-rich fault gouge bears a low-friction strength, and the friction is strengthened with slip velocity (velocity-strengthening), which was thought to support aseismic slip. Clay minerals are comprised of platy crystalline layers with water intercalated between them, which may act as a lubricant. Sliding between clay layers was suspected to support aseismic slip but lacked a clarified mechanistic insight. We use non-equilibrium molecular dynamics simulations to show that shearinduced interlayer sliding is frictionally weak and velocity-strengthening, which evidences the role of clay minerals in aseismic slip. We find that interlayer water is a viscous fluid at most times, which explains the shear response of interlayer sliding. Depending on temperature and pressure conditions, intercalated water can be monolayer or bilayer, fluidic or ice like. Shear induces icelike water to transform into fluidic water, which happens as a stick-slip phenomenon reflecting a first-order transition. Increased pore fluid pressure leads to the transformation from monolayer to bilayer intercalated water, resulting in a lower friction strength and enhanced velocity-strengthening behavior. Our work suggests that disclosing the hydration state of a clay mineral is preliminary when studying fault mechanics.

Keywords: Smectite, interlayer, friction, aseismic slip

INTRODUCTION

Earthquakes arise as a consequence of frictional instabilities that cause stress accumulated over years to be relieved in sudden stick-slip events along the fault. Slip occurs rapidly as a rupture dynamically propagates over the fault surface, which generates seismic waves (Carlson et al. 1994; Scholz 1998). In contrast, stable creeping fault allows strain to be relieved smoothly and aseismically. Geodetic and seismological observations and geological records have shown a transition between aseismic slip and seismic rupture within the same fault zone (Avouac 2015; Bürgmann et al. 2000; Kawamura et al. 2012; Liu et al. 2020; Maurer and Johnson 2014; McGuire et al. 2012; Perfettini and Avouac 2014; Tarling et al. 2018). Aseismic slip or stable creep prevents the earthquake propagation along the fault, but the strain accumulation due to aseismic slip could nucleate an earthquake beyond the creeping fault zone. As aseismic slip is a common phenomenon in the earthquake preparatory phase, uncovering the underlying physics of this process is promising in earthquake forecasts (Collettini et al. 2019; Kawamura et al. 2012).

Whether a slip is stable or unstable depends on the velocity dependence of the steady-state friction. Sliding begins when shear stress reaches the Coulomb-Mohr failure criterion (Handin 1969). Once sliding, the Amontons' law characterizes the relationship between shear (τ) and effective normal stress ($P_{\sigma}-P_{f}$), in which P_{σ} is the normal stress and P_{f} is the pore fluid pressure):

$$\tau = \mu_{\rm d} \left(P_{\rm \sigma} - P_f \right) \tag{1}$$

where μ_d is the dynamic friction coefficient (Collettini et al. 2019). Materials that exhibit velocity-strengthening behavior, that is, μ_d in the steady-state sliding regime, increase with velocity and only produce stable creep; while velocity-weakening materials may exhibit either unstable stick-slip or conditionally stable behavior (Marone 1998; Scholz 1998). Based on the Dieterich-Ruina constitutive law, or the rate and state friction (RSF) law (Baumberger and Caroli 2006; Kawamura et al. 2012; Scholz 1998), the steady-state friction coefficient scales with the logarithm of sliding velocity ν :

$$\mu_{\rm d} = \mu_0 + (a - b) \ln(v/v_0). \tag{2}$$

If $a - b \ge 0$, the material is velocity-strengthening, otherwise it is velocity-weakening. Although the RSF law generally applies to rock, papers, wood, metals, and plastics (Heslot et al. 1994; Marone 1998), limitations of it have been reported, e.g., it no longer holds at high sliding velocity (Im et al. 2020; Kawamura et al. 2012). The microphysical foundation of the RSF law is that creep at asperity junctions between grains follows an Arrhenius activated rate process (Putelat et al. 2011; Rice et al. 2001).

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However, this foundation might be lost when grains are separated by melt or fluid (Di Toro et al. 2004; Hirose and Shimamoto 2005; Mizoguchi et al. 2006) and asperity junctions become rare. In such cases, the materials of fault gouge can be seen as an amorphous soft solid. The shearing flow of amorphous solid in the steady state manifests a power law dependence of shear stress with shear rate (Liu et al. 2021; Nicolas et al. 2018; Tsai et al. 2021), which is a regime distinct from that follows the RSF law.

The material compositions of fault gouge primarily determine the slip stability. Gouges rich in phyllosilicate or clay minerals, which present crystallographic preferred orientation fabric (Fig. 1), were speculated to support stable creep (Avouac 2015; Carpenter et al. 2012; Collettini et al. 2019; French et al. 2015; Holdsworth 2004; Lockner et al. 2011). Clay minerals consist of platy crystalline layers, which weakly interacts with each other through electrostatic, van der Waals, or hydration forces (Smith et al. 2006; Whitley and Smith 2004). Local slip between water-lubricated smectite (a typical clay mineral) layers was suggested during shear (Aretusini et al. 2019). We speculate that some clay layers constitute "hot spots" (Amon et al. 2012; Pons et al. 2016). Local plastic events in hot spots, i.e., slips between layers, support the macroscopic plastic deformation, i.e., fault creep. However, the quantitative connection between fault stability and microscopic interlayer slips is still lacking. The RSF law is a macroscopic view of the shear deformation and it is phenomenological. It would be interesting to see whether the microscopic interlayer slip follows a similar form as the RSF law.

Slip frictions of lubricants (e.g., water film) confined between surfaces were widely investigated with shear experiments and non-equilibrium molecular dynamics (NEMD) simulations (de Wijn and Pettersson 2017; Dhopatkar et al. 2016; Diao and Espinosa-Marzal 2018; Ma et al. 2015; Xu and Leng 2018; Xu et al. 2020). With water as a lubricant, the interfacial structure and phase state or organization of water molecules have specific effects on friction between surfaces (Chen et al. 2015; de Wijn and Pettersson 2017; Dhopatkar et al. 2016; Jinesh and Frenken 2006; Ma et al. 2015; Ortiz-Young et al. 2013; Vilhena et al. 2016). We design NEMD simulations for the following goals: (1) to verify the possibility whether clay layers act as hot spots, which support the shear deformation of fault gouge; (2) to disclose whether the interlayer sliding is velocity-strengthening and explain why a clay mineral promotes aseismic slip; (3) to relate the shear behavior with the phase state of the clay interlayer water.

Temperature, normal stress, and pore fluid pressure are environmental variables that affect slip stability (Avouac 2015; French et al. 2015; Yang and Juanes 2018). These variables have influenced the interlayer hydration state of smectite (Smith et al. 2006; Whitley and Smith 2004; Zhou et al. 2020), which in turn affects lubrication between layers. Thus, first of all, the interlayer hydration state under different environmental conditions is determined in this study by combining equilibrium molecular dynamics (EMD) simulations and thermodynamic integrations (TI). Hence, through deriving the mechanical property of a hydration state under the same condition with NEMD simulations, the influences of those variables are clarified.

METHODS

Equilibrium molecular dynamics simulations

Arizona-type montmorillonite [unit-cell formula: (M^{x+})_{1/x}[Al₃Mg][Si₈] O20(OH)4·nH2O] was chosen as the smectite (clay mineral) model here, which bears only octahedral charges (Hensen et al. 2001; Tambach et al. 2004) and the intercalated cation M^{x+} is Na⁺ in this study (Fig. 2). The basic TOT layer (a layer consists of two tetrahedral and one octahedral sheets) of the model was based on the structure of pyrophyllite (Lee and Guggenheim 1981). We built a $7 \times 4 \times 4$ supercell of pyrophyllite. In each unit cell, there is a Mg atom replaced by an Al atom according to the formula of Arizona-type montmorillonite. Any two substitution sites are not adjacent. The range of water content (n) in this study is 0-18.92. The upper bound of the studied water content was determined so as to cover monolayer, bilayer, and trilayer hydration states (Zhou et al. 2020). Water molecules were inserted into the enlarged interlayer according to n. A total of 29 systems with different n were built. Three-dimensional periodic boundary conditions were applied. The electrostatic interaction was evaluated using the particlemesh Ewald method (Darden et al. 1993; Essmann et al. 1995). The cut-off radius for the short-range non-bonded interactions was set to 1.2 nm. The ClayFF force field (Cygan et al. 2004) was used to describe atoms in layers of smectite. Water molecules were described by the SPC/E model (Berendsen et al. 1987). The force field parameters of Na⁺ were adopted by that proposed by Smith and Dang (1994). Those parameters have long been used in conjunction with the ClayFF forcefield and verified to well deliver properties of Na+-intercalated smectite (Cygan et al. 2004; Zhou et al. 2020).

The studied temperatures were 300, 348, and 473 K, while pressures were 0.1, 60, and 120 MPa. Thus, there were altogether nine combinations of temperature and pressure conditions. The 300–473 K and 0.1–120 MPa ranges cover the most existence conditions of clay-rich gouge (Brown et al. 2017; Carpenter et al. 2012; Ikari et al. 2013; Lockner et al. 2011). Any temperature and pressure combination in this study is not referred to a specific geological setting but rather serves to show the generic influence of pressure or temperature variation.



FIGURE 1. Schematic of the relationship between plastic deformation of fault gouge and local slip between clay layers. (a) Fault gouge with large solid grains (gray particles), phyllosilicate or clay minerals (brown lines), and pore water (blue zone). (b) Local slip between clay layers. (c) Atomic model of a clay mineral. Si, Al, Mg, Na, O, and H atoms are in yellow, light purple, fluorescent green, purple, red, and white, respectively. (Color online.)

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FIGURE 2. Schematic of water exchange between smectite interlayer and pore fluid. The TOT layer contains SiO_4 tetrahedron (yellow), AlO_6 octahedron (fluorescent green). Water molecules [O atoms (red), H atoms (white)] and Na⁺ ions (purple) are intercalated between layers. (Color online.)

GROMACS 5.1 package (Hess et al. 2008) was used to perform simulations. The equation of motion was integrated with the leapfrog algorithm with a time step of 1.0 fs. A 7 ns isothermal-isobaric (NPT) simulation was performed with each dimension of the box scaled independently. Simulated annealing from 500 K to target temperature was performed during the first 2 ns, so as to well equilibrate structure. Berendsen thermostat and barostat (Berendsen et al. 1984) were used in this stage. Then, an NP_aT simulation, during which only the *z*-dimension of the box was scaled, was run for 70–150 ns. The Nosé-Hoover thermostat (Parrinello and Rahman 1985; Hoover 1985; Nose 1984) and Parrinello-Rahman barostat (Parrinello and Rahman 1981) were used in this stage. Data were saved every 0.1 ps in the equilibrium NP_aT simulations. The coupled pressure in the *z* dimension (0.1, 60, and 120 MPa) is normal stress P_a on the basal plane of smectite.

Bulk phase with 2165 water molecules, which mimic the pore fluid phase, was also simulated. An NPT simulation was performed for 18 ns. The same three temperatures (300, 348, and 473 K) were considered. At 300 K, the investigated pressures ranged from -120 to 120 MPa with a gradient of 10 MPa, except that 0 MPa was replaced by 0.1 MPa. At 348 K, because the bulk water phase blows up at -120 MPa, the pressure range was narrowed to be from -110 to 120 MPa. At 473 K, the pressure range was from -40 to 120 MPa. Simulated annealing from 500 K to target temperature was performed during the first 3 ns of an NPT simulation. Data were collected every 0.1 ps in the equilibrium NPT simulations. It is noted that water simulated here at all conditions is in the liquid state, as shown by the molecular volume of water (Online Materials¹ S1), which is consistent with the liquid state description (Lopez-Lemus et al. 2008). The simulated water at some conditions (i.e., 473 K and negative pressure) is superheated liquid (Gallo et al. 2016), or in other words, in a metastable state. The stable state at such conditions is a vapor state or a vapor/liquid coexistence one. As we try to provide a generic insight on how the variation of temperature or pressure affects the coupling between pore and interlayer water, we do not simulate the stable state but rather the metastable one, which maintains the liquid behavior. The coupled pressure of the bulk phase represents the pore fluid pressure P_f. In this study, we do not explicitly consider the influence of solvated ions on the effective pore fluid pressure or osmotic pressure (Kohns et al. 2016; Koop et al. 2000). In principle, the chemical potential of the bulk phase simulated here is equal to that of the real pore fluid with the same osmotic pressure P_{f} .

Determination of stable hydration state of smectite

For water in smectite coupled with pore fluid (Fig. 2), we assumed no water flow between the two phases, i.e., a static condition. The static condition implied:

$$\mu_i = \mu_f \tag{3}$$

$$\Pi = P_{\sigma} - P_{L}$$

$$(4)$$

where μ_i and μ_f are chemical potentials of intercalated and pore fluid water, respectively, P_L is the lateral stress of water intercalated in smectite, and Π is the disjoining pressure which is the inherent property of intercalated water (Shchukin et al. 2001). If both smectite and pore fluid are under the same hydrostatic pressure, Equations 4 and 5 are derived to be $P_a = P_L = P_f$ and $\Pi = 0$, as assumed in our previous study (Chen et al. 2020a). However, here we are interested in the mechanical non-equilibrium state with $P_{\sigma} \neq P_{f}$. We express the hydration Gibbs free energy G_{hyd} as a function of the interlayer spacing *l*:

$$G_{\text{hyd}}(l) = -A \int_{0}^{l} \Pi(l) dl' + (P_{\sigma} - P_{f}) \cdot A \cdot l$$
(6)

in which the former term accounts for the work done by intercalated water, while the latter accounts for that by environment. A is the area of a basal surface. The minimum of $G_{hyd}(l)$ corresponds to the static state. Alternatively, G_{hyd} can be seen as a function of the number of intercalated water molecules (N):

$$G_{\rm hyd}(N) = \int_{0}^{N} [\mu_{\rm i}(N, P_{\sigma}) + \mu_{\rm f}(P_{f})] dN'$$
(7)

 μ_i and μ_f at certain temperature and pressure conditions are derived with TI through decoupling interactions between a water molecule and the surrounding environment, which can be found in our previous studies (Chen et al. 2020a, 2020b; Zhou et al. 2020) and briefly introduced in the Online Materials¹ S2.

In this study, we derive $\mu_i(N)$ and μ_f at conditions ($P_\sigma = 0.1$ MPa, T = 300 K), ($P_\sigma = 60$ MPa, T = 348 K), and ($P_\sigma = 60$ MPa, T = 473 K) through decoupling interactions. $\mu_i(N)$ under the same T but different P_σ conditions can be derived by the following integration:

$$\mu_{i}\left(N,P_{\sigma,j},T\right) - \mu_{i}\left(N,P_{\sigma,i},T\right) = \int_{P_{\sigma,j}}^{P_{\sigma,j}} \left(\frac{\partial V}{\partial N}\right)_{N,P_{\sigma}',T} dP_{\sigma}'$$
(8)

in which

$$\left(\frac{\partial V}{\partial N}\right)_{N,P'_{\sigma},T}$$

is the partial derivative of volume with N, as derived with equilibrium configurations with various N (Chen et al. 2020a). If $P_{\alpha,j}$ is close to $P_{\alpha,i}$ and the variation of

$$\left(\frac{\partial V}{\partial N}\right)_{N,P'_{\sigma},T}$$

with P'_{σ} is small, the chemical potential difference can be approximated as:

$$\mu_{i}\left(N,P_{\sigma,j},T\right) - \mu_{i}\left(N,P_{\sigma,i},T\right) \approx \frac{P_{\sigma,j} - P_{\sigma,i}}{2} \cdot \left[\left(\frac{\partial V}{\partial N}\right)_{N,P_{\sigma,j},T} + \left(\frac{\partial V}{\partial N}\right)_{N,P_{\sigma,j},T} \right].$$

$$(9)$$

With Equation 9, we derive $\mu_i(N)$ under all the studied pressure conditions. As to μ_f the dependence of it on pressure is similarly:

$$\mu_{\rm f}(P_{f,j},T) - \mu_{\rm f}(P_{f,j},T) = \int_{P_{f,j}}^{r_f} V_f(P_f',T) dP_f'$$
(10)

where V_f is the molecular volume of pore fluid water (bulk water here). Equilibrium data of V_f at discrete P'_f conditions are recorded (Online Materials¹ S1). Through fitting the data with a third-order polynomial equation, we derive the $V_f(P'_f)$ function.



FIGURE 3. Schematic of NEMD simulation model: (**a**) side view; (**b**) top view. Color representations are the same as those in Figure 2. (Color online.)

Substituting the function into Equation 10, with the known $\mu_t(P_{f,i},T)$ derived with TI, $\mu_f(P_{f,j},T)$ is achieved. With μ_i and μ_f and Equation 7, we get $G_{hyd}(N)$. $G_{hyd}(N)$ is transformed into $G_{hyd}(n)$, in which *n* is water content in the chemical formula of smectite. *n* at the minimum of G_{hyd} corresponds to the stable hydration state of smectite under conditions characterized by P_o, P_f , and *T*.

Non-equilibrium molecular dynamics simulations

The equilibrium configuration of a stable hydration state at a condition (P_o, P_f , and T) was used to initiate a NEMD simulation in an NP_oT ensemble. The two TOT layers and water intercalated between them in the middle of the smectite model are seen as one group, referred to as a pulled group (Fig. 3a). A linear potential parallel with the basal plane was added to the pulled group. Thus, the pulled group was in a field like a gravitational field and, therefore, a constant force (F) was at the center of mass of the group. F does not vary with the number of atoms in the pulled group. A large enough F would lead to dislocation between layers. In a steady state, the friction force is equal to F. The shear stress is expressed as:



FIGURE 4. Hydration Gibbs free energy G_{hyd} normalized by basal surface area A as a function of water content *n*. Results at different temperatures and normal stresses are shown in different panels. In each panel, different free energy curves refer to cases with different pore fluid pressures. The global free energy minimum is denoted by a red symbol. (Color online.)



FIGURE 5. The stable water content (*n*) of smectite as a function of pore fluid pressure (P_f) in cases at different temperatures (*T*) and normal stresses (P_o) (**a**–**c**). Snapshots of monolayer (1W) and bilayer (2W) hydration states are shown (**d** and **e**). Color representations are the same as those in Figure 1. (Color online.)

 $\tau = F/2A \tag{11}$

where A is the area of a basal surface. Because the pulled group was in contact with two basal surfaces, the denominator in Equation 11 was 2A. In this study, F along directions x, -x, y, and -y were adopted (Fig. 3b). Thus, we denote τ as τ_x , τ_{xx} , τ_y , and τ_{-y} , whose subscripts represent pull directions.

A NEMD simulation lasted for 1-6 ns. The simulation time depended on how long it took for the pulled group to reach a steady state. The topmost and downmost TOT layers and water molecules between them across the periodic boundary are seen as the reference group (Fig. 3a), whose center of mass was kept fixed during a simulation. At a given condition, a series of NEMD simulations with different *F* on the pulled group were performed. The displacement of the pulled group with time is recorded.

Because how the thermostat is applied could affect NEMD simulation results (de Wijn and Pettersson 2017), we tried two thermostating ways. (I) Separate thermostat: we separated two thermostating groups, one was the reference group, and the other was the pulled group and the two water layers beneath or above it. These two groups were separately thermostated, so that heat generated by frictional sliding would be removed. (II) Partial thermostat: only the reference group was thermostated, so it allowed local temperature increase due to heat generated by frictional sliding. With these two thermostatic methods, whether the frictional heat has an influence on sliding will be shown.

RESULTS

Hydration state of smectite at different P_{σ} , P_{f} , and T conditions

At a condition characterized by P_o , P_f , and T, the global minimum of $G_{hyd}(n)$ is referred to as the stable hydration state of smectite (Fig. 4). At conditions when P_f is high enough, there is no global free energy minimum in $G_{hyd}(n)$. In these cases, when n is larger than some value, $G_{hyd}(n)$ decreases monotonically as n increases. Thus, layer-by-layer stacking smectite is not available under these conditions, and exfoliation of smectite is anticipated.

We compile *n* referring to global minima at different conditions and show them in Figure 5. There are roughly two hydration states of smectite, one with n < 6.0, and the other with *n* larger than 8.0. Water in the former state roughly exhibits a monolayer form (Fig. 5d), while water in the latter appears as a bilayer (Fig. 5e). These water arrangements are consistent with previous studies (Mazo et al. 2008; Zhang et al. 2014; Zhou et al. 2020). We refer to the two states as 1W and 2W, respectively, consistent with the previous designation (Ferrage 2016). At a P_{σ} and T condition, the increase of P_f generally induces the transformation from a 1W state to a 2W one. At the same temperature, when P_{σ} is higher, it requires a higher P_f to transform smectite from a 1W state to a 2W one (Figs. 5a–5c). At the condition T = 300 K and $P_{\sigma} = 0.1$ MPa, only a 2W state is available under the studied range of pore fluid pressure (Figs. 4a and 5a), the same as that at the condition T = 473 K and $P_{\sigma} = 0.1$ MPa (Figs. 4g and 5c).

Simulated interlayer sliding of smectite

With stable hydration states of smectite at different P_o , P_f , and T conditions, NEMD simulations were performed to show sliding between layers. The displacement of the pulled layers with time is shown in Figure 6. When F is too small, the pulled group is stationary in the steady state (Figs. 6a and 6d). As Fincreases, the pulled group exhibits a stepwise movement (Figs. 6b and 6e). When F is large, a linear displacement (s) with time (t) is observed (Figs. 6c and 6f). Through linear fitting of s(t) in the steady state (Fig. 6), we derive the slope of s(t), which is the velocity component along x, -x, y, and -y directions. The direction of v is not necessarily along the pull direction, as a small non-zero velocity component perpendicular to it is observed (Online Materials¹ Figs. S2–S7). In the following, we just refer to the magnitude of v.

In the steady state, the velocity of water next to the pulled layer is statistically the same as the steady-state velocity of that layer. At the same time, the water next to the reference layer



FIGURE 6. Displacement (*s*) of the pulled group as a function of time (*t*). *x* and *y* denote displacement components along *x* and *y* coordinates, respectively. A constant force *F*, which is along *x* or -x direction is exerted on the pulled group. Linear fit of the displacement component along the *x* coordinate in the steady state is shown. "II" denotes that partial thermostat was used. (Color online.)

almost does not flow. An example of the distribution of the steady-state water velocity is shown in Figure 7.

$$\ln \tau = k \ln v + \ln \tau_{\rm i} \tag{12}$$

Shear response of smectite interlayer

The relationships between shear stress and velocity magnitude are shown in Figure 8 (results with partial thermostat) and in Online Materials¹ Figure S8 (results with separate thermostat). For steady slidings, τ increases with steady-state velocity *v*, corresponding to a velocity-strengthening behavior. However, we find that τ does not scale with ln*v*, i.e., the interlayer sliding does not follow the form of the RSF law. Instead, a log-log scaling between τ and *v* is found. $\tau(v)$ can be phenomenologically fitted with the following equation:



FIGURE 7. Velocity profile of water in the steady state. The blue dash lines mark velocities of the pulled and stationary layers. The green dash lines show the linear fits to the velocity profiles of water. (Color online.)

where τ_i and k are defined as initial shear stress and velocitystrengthening constant here. In results with either thermostat, Equation 12 well fits most data (Fig. 8; Online Materials¹ Fig. S8). The log-log plots in Online Materials¹ Figures S9 and S10 clearly show the general well fittings. However, in some cases (1W state with T = 300 K and $P_{\sigma} = 60$ or 120 MPa; 2W state with T = 300 K and $P_{\sigma} = 120$ MPa), a discontinuous jump of v with τ is observed (Figs. 8a, 8b, and 8h; Online Materials¹ Figs. S8a, S8b, and S8h). This is reminiscent of the stick-slip regime when a surface is driven with a spring at a constant velocity, which probably implies a dynamic phase transition (Drummond and Israelachvili 2001; Urbakh et al. 2004). Here, the pulled group remains stationary until τ reaches a threshold. Equilibrium trajectories of the stationary state show that an intercalated water molecule vibrates but almost does not diffuse (Online Materials¹ Videos). It is an ice-like structure (Chen et al. 2020a; Leng and Cummings 2006). When τ is smaller than the threshold, a snapshot shows that a hydrogen bond (HB) network is well developed among water molecules (Fig. 9a). When τ is larger than the threshold, the HB network is largely broken (Fig. 9b), corresponding to a first-order phase transition. We calculate the number of HBs per water molecule forms $(N_{\rm HB})$ with the HB criterion: the donor-acceptor distance, the hydrogenacceptor distance, and the hydrogen-donor-acceptor angle are <0.35 nm, 0.245 nm, and 30°, respectively (Laage and Hynes 2006, 2008). As τ increases, we observe a discontinuous drop of $N_{\rm HB}$ (Fig. 9c). The radial distribution function [g(r)] between water O atoms also shows a discontinuous structural variation (Fig. 9d). They evidence a shear-induced ice-like to liquid-like

phase transition. In other cases where *v* continuously increases with τ (Fig. 8; Online Materials¹ Fig. S8), no distinct structural variation happens as shown by N_{HB} (Fig. 9c) and g(r) (Fig. 9d). Good fitting with Equation 12 implies even a small non-zero τ could initiate an interlayer slip given that simulation time was as long as possible. In addition, for the 1W state at 473 K with $P_{\sigma} = 60$ or 120 MPa, Equation 12 well fits low-velocity data, but it deviates from high-velocity ones (Figs. 8o and 8p; Online Materials¹ Figs. S8o and S8p). This state is characterized by a relatively lower water content (n = 3.9). The deviation from a simple log-log scaling probably implies a multiple-stage sliding dynamics but without phase transition. Based on the log-log plots in Online Materials¹ Figures S9 and S10, when v is as small as <10 nm/ns, a deviation from the log-log plot is observed. This can also be explained as multiple-stage sliding dynamics. Nevertheless, the deviation of the magnitude of τ from the loglog plot is just small in such a low-velocity regime.

DISCUSSION

Comparison between simulations and laboratory friction experiments

The shear velocity magnitude reported here is of the order of nm/ns (i.e., m/s), but that reported in laboratory friction experiment is much smaller and of the order of μ m/s (Carpenter et al. 2012; Diao and Espinosa-Marzal 2018; Ikari et al. 2013). In most cases we studied here, the log-log scaling between τ and v (Fig. 8) implies that even a small finite τ can induce an interlayer slip. Given that simulation time was as long as possible, a stepwise movement of the pulled layer under a finite τ like those in



FIGURE 8. Shear stress (τ) as a function of velocity magnitude (v) of the pulled group at 300 K (left), 348 K (middle), and 473 K (right). Subscripts (x, -x, y, and -y) denote directions of the shear stress. Fittings with Equation 12 are shown. "II" denotes that partial thermostat was used. (Color online.)



FIGURE 9. Structure of water in the interlayer of smectite. Panels **a** and **b** show top-view configurations of water before and after a shear-induced phase transition, respectively. Panel **c** shows number of hydrogen bonds ($N_{\rm HB}$) per water molecule forms as a function of shear stress. Panel **d** shows the radial distribution function between water O atoms. These results were derived with partial thermostat. (Color online.)

Figures 6b and 6e is anticipated. However, due to the limitation of the current computational powers, reproducing such a low shear velocity as those in laboratory experiments is impossible. Nevertheless, in cases without dynamic phase transition, the log-log scaling between τ and ν could be extrapolated to the lower velocity regime. It provides an estimate of the relationship between τ and ν . In cases with dynamic phase transitions, the threshold value of τ , which induces a transition, and the initiate shear velocity are hard to be determined as it takes long enough simulations to guarantee the reliability. In those cases, the shear dislocation between clay layers is harder to be induced.

Diao and Espinosa-Marzal (2018) slide a silica tip along a calcite surface in aqueous solution with atomic force microscopy (AFM) to produce a shear rate of the order of μ m/s with a shear stress of the order of Pa. However, the shear experiment on fault gouge produces the same order of a shear rate of the amorphous solid with a shear stress of a much higher order of MPa (Collettini

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et al. 2019). This discrepancy reflects that, the macroscopic shear displacement is different from the microscopic one. If the AFM tip was under a shear stress of the same order as in shear experiments on fault gouge, a shear rate of a much higher order is anticipated. We use the concept of hot spots (Amon et al. 2012; Pons et al. 2016) to refer to sites in amorphous solid which are more susceptible to plastic deformations. Those sites could be grain boundaries intercalated by water or clay interlayers. Under an applied macroscopic shear stress, the local shear rates in hot spots are higher than the macroscopic one. Because shear sliding between clay layers is velocity-strengthening, the sliding would stop, and the stress in hot spots would be released and propagated to environmental sites. In this study, we show that under a shear stress of the order of MPa (the same order as in shear experiments on fault gouge) the shear rate between clay layers is of the high order of m/s, which evidences the role of clay layers as hot spots.

Origin of the shear response of smectite interlayer

Diao and Espinosa-Marzal (2018) found a log-log scaling between friction force and velocity in the AFM experiment. The log-log scaling between τ and ν might be common when an aqueous layer is confined between sliding surfaces. Such a scaling is reminiscent of shear flow of non-Newtonian fluid, which can be described by the Ostwald de Waele equation (Chhabra 2010):

$$\tau = m(\nu/h)^k \tag{13}$$

where *m* and *k* are constants and *h* is the fluid thickness. Equation 13 can be transformed into Equation 12 with $\ln \tau_i = \ln m - k \ln h$. Thus, the log-log scaling between τ and *v* is in fact derived from the Ostwald de Waele equation, which describes shear fluid flow. The shear flow is manifested by an approximately linear distribution of water velocity with distance (Fig. 7). Thus, the shear rate (*v*/*h*) under a shear stress (τ) is controlled by viscosity (η) of interlayer water (Landau and Lifschitz 1987):

$$\nu/h = \tau/\eta. \tag{14}$$

Combining Equations 13 and 14, η can be derived with the following equation:

$$\eta = m(\nu/h)^{k-1}.$$
 (15)

We show the relationship between η and v/h in Figure 10 and Online Materials¹ Figure S11. The function line of Equation 15 is also presented according to $m = \exp(\ln\tau_i + k\ln h)$ and with the fitted values of τ_i and k (Fig. 8; Online Materials¹ Fig. S8). Figure 10 shows Equation 15 generally explains the log-log scaling between η and ν/h . η decreases with shear rate, which implies interlayer water is a shear-thinning fluid (Hartkamp et al. 2013; Leng and Cummings 2005; Thompson et al. 1995). Shear thinning appears when the structural relaxation rate of fluid is smaller than the shear rate (Leng and Cummings 2005).

The viscosity response to shear rate is the intrinsic property of interlayer water, which explains log-log scaling between τ and v and the velocity-strengthening behavior. Different viscosities of water in 1W and 2W states and variations with shear rate give rise to different shear responses. The phenomenological RSF law with a form of a $\tau \sim \ln v$ scaling (Baumberger and Caroli 2006; Kawamura et al. 2012; Scholz 1998) is not applicable for the viscous shear between clay layers. For sliding between solid surfaces with water as a lubricant, experiments have found a $\tau \sim \ln v$ or a $\ln \tau \sim \ln v$ scaling (Diao and Espinosa-Marzal 2018; Ma et al. 2015). The $\tau \sim \ln v$ relation was found in the low-hydration regime. In this regime, hydrated ions at the surface act as asperity contacts, so that dislocation between surfaces is an activated process requiring the crossing of an energy barrier. However, even for the 1W hydration state studied here, shear between layers follows a $\ln \tau \sim \ln v$ relation. Snapshots show ions are not present as asperities (Figs. 5d and 5e) but flow with water. The viscous flow of water supports the $\ln \tau \sim \ln v$ relation. As for the interlayer with ice-like water, a slide only begins after a dynamic phase transition, which is an activated process. Thus, if a statistical analysis was on a clay matrix that consists of many interlayers with ice-like water, the deformation of the matrix may follow the $\tau \sim \ln v$ relation. This scenario appears at a high pressure but close to ambienttemperature conditions.



FIGURE 10. The effective viscosity (η) as a function of shear rate (ν/h) at 348 K with separate (I) (**a**, **b**, **c**) and partial (II) thermostats (**d**, **e**, **f**). Solid lines are function lines of Equation 15 derived with fitted values of τ_i and *k*. Black and yellow lines correspond to results of the 1W and 2W states, respectively. (Color online.)

Effects of P_{σ} , P_{f} , and T on shear strength and its velocity dependence

Through fitting $\tau(v)$ (Fig. 8), we derive τ_i and k, which are dynamic characteristics of interlayer water with a fixed content under conditions of T and P_{σ} . τ_i is the shear stress at a low-sliding velocity. It is noted that τ_i is not the yield shear stress because when interlayer water is in a fluidic state, steady sliding is always available only if τ is non-zero. When interlayer water is in an ice-like state under low-shear stress, the fitted τ_i is smaller than the threshold stress, which induces a phase transition. As shown in the thermodynamic analysis, the variation of effective normal stress ($\Delta P = P_{\sigma} - P_{f}$) can lead to alternation of a hydration state, which in turn changes τ_i . We show τ_i as a function of ΔP in Figure 11. The error bar of ΔP refers to the range of P_f in which the hydration state of smectite does not change (Figs. 5a-5c). According to Equation 1, with τ_i and ΔP , friction coefficient μ_d can be derived in principle. However, the variation of τ_i with ΔP in an isothermal condition is stepwise (Fig. 11). When interlayer water is in a fluidic state and ΔP is increased and the hydration state is not changed, the variation of τ_i is almost negligible. The alternation of a hydration state due to increased ΔP leads to a sharp increase of τ_i . Due to the nonlinear relation between τ_i and ΔP , the Amontons' law breaks down here. Linear fitting of $\tau_i(\Delta P)$ would result in an apparent friction coefficient not larger than 0.3, as guided by the $\tau_i = 0.3\Delta P$ line in Figure 11. Laboratory experimental studies have shown that clay-rich samples exhibit a macroscopic friction as low as $0.1 < \mu_d < 0.3$ (Collettini et al. 2019). Thus, this study here discloses that the low macroscopic friction strength can be attributed to the shear displacement between smectite layers.

It is noted that even for samples rich in phyllosilicate minerals without intercalated water, the macroscopic friction still fulfills the range $0.1 < \mu_d < 0.3$ (Collettini et al. 2019). Thus, the low friction coefficient is not exclusively determined by intercalated water. If the variation of ΔP is too small to induce a transition between different hydration states of smectite, the alternation of τ_i is almost negligible (Fig. 11) and μ_d would be approaching 0. Thus, the major mechanical difference between clay minerals in different hydration states is not μ_d but τ_i , which quantifies the ability as hot spots. τ_i for the 2W state is generally smaller than that for the 1W one (Fig. 11), as viscosity of interlayer water in the 2W state is smaller (Fig. 10).

As temperature increases, for the same hydration state, τ_i generally decreases due to a decrease of viscosity. When the temperature is constant, if the hydration state does not change, increasing ΔP generally does not lead to a considerable variation of τ_i , except in case increased ΔP leads to a phase transition, e.g., 2W state at 300 K transforms from a fluidic state to an ice-like one from 60 to 120 MPa. However, τ_i of the ice-like state is meaningless as it is much smaller than the yield stress.

At the same temperature, k for the 2W state is generally larger than that for the 1W state (Fig. 12), implying velocitystrengthening friction is elevated. It can be explained by the viscosity of interlayer water in the 2W state, which decreases less with shear rate (Fig. 10). For the 2W state, temperature increase generally leads to the enlargement of k, but it is not clear for the 1W state, as also explained by the viscosity variation. When temperature is unchanged, variation of ΔP without changing the hydration state does not lead to a considerable variation of k.

We provide the first mechanistic insight on how delamination between clay layers supports aseismic slip. The interlayer water lubricant gives rise to a velocity-strengthening behavior, as interlayer water is a viscous fluid. Interlayer water is a shear-



FIGURE 11. The initial shear stress (τ_i) as a function of the effective normal stress (ΔP). Panels **a** and **b** show results with separate (I) and partial (II) thermostats, respectively. The number marked next to the symbol is the P_{σ} value whose unit is MPa. The function line of $\tau_i = 0.3\Delta P$ is shown to guide the upper bound of the friction coefficient. (Color online.)



FIGURE 12. The velocity-strengthening constant (k) as a function of the effective normal stress (ΔP). Panels **a** and **b** show results with separate (I) and partial (II) thermostats, respectively. The number marked next to the symbol is the P_{σ} value whose unit is MPa. (Color online.)

thinning fluid whose viscosity response to shear rate determines the relationship between shear stress and velocity. The variation of temperature, normal stress, or pore fluid pressure alters hydration state of an interlayer, and thus it causes changes in dynamic characteristics. Viscosity of the 2W hydration state is lower than that of the 1W one, which results in the lower frictional resistance. The velocity-strengthening behavior is more significant for the 2W state due to less decrement of viscosity with shear rate. Thus, clay minerals in the 2W hydration state are more prone to act as hot spots.

We suggest that a separate thermostat gives rise to a more accurate estimation of τ_i , but with a partial thermostat, k is better estimated. With a partial thermostat, frictional sliding accompanies heat generation, which implies energy dissipation. As a result, the increment of v requires a larger increment of τ than in the case with a separate thermostat. Thus, k for the 2W state simulated with partial thermostat is lower (Fig. 12b). As heat generation is also anticipated when sliding natural samples, the partial thermostat better estimates k. On the other hand, when sliding has not happened or sliding is very weak, energy dissipation into environment is fast. Thus, the separate thermostat is more reasonable as the temperature is consistent with environment. τ_i is the stress at a low shear rate so that it is better derived with the separate thermostat. Thus, Figures 11a and 12b are recommended estimations of initial shear stress and velocity-strengthening ability, respectively.

IMPLICATIONS

Properties of minerals primarily determine fault mechanics. The viscous deformation of serpentine antigorite at high pressure impedes stress buildup and supports interseismic creep (Hilairet et al. 2007). Similarly, the viscous shear between clay layers shown here supports aseismic slip. Shear behavior of smectite does not monotonically vary with pressure or temperature, but the alternation of a hydration state with the environment plays a key role. We suggest that disclosing the hydration state of a clay mineral is preliminary. Understanding local slip between clay layers paves the way for uncovering the constitutive law to describe fault gouge deformation. In this work, we verify the role of clay minerals as hot spots, which support aseismic slip, but future numerical studies would be needed to connect the local plastic deformation in clay layers and macroscopic fault dynamics.

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