Adhesive wear mechanisms in the presence of weak interfaces: Insights from an amorphous model system

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Engineering wear models are generally empirical and lack connections to the physical processes at the nano- to micro-scale. Here, we thus analyze wear particle formation for sliding interfaces in dry contact with full and reduced adhesion. Depending on the material and interface properties and the local slopes of the surfaces, we find that colliding surface asperities can either deform plastically, form wear particles, or slip along the contact junction surface without significant damage. We propose a mechanism map as a function of material properties and local geometry, and confirm it using quasi-2D and 3D molecular dynamics and finite-element simulations on an amorphous, silicon-like model material. The framework developed in the present paper conceptually ties the regimes of weak and strong adhesion contacts together and can explain that even unlubricated sliding contacts do not necessarily lead to catastrophic wear rates. A salient result of the present work is an analytical expression of a critical length scale, which incorporates interface properties and roughness parameters. Therefore, our findings provide a theoretical framework and a quantitative map to predict deformation mechanisms at individual contacts. In particular, contact junctions of sizes above the critical length scale contribute to the debris formation.

I. INTRODUCTION

Contacting solids in relative motion, i.e., they gradually lose volume. Such volume loss is inextricably linked to the roughness and the details of the contact between the two surfaces. In the case of adhesive wear, contacting surface asperities form a bond that leads to the detachment of wear particles during sliding, which are progressively evacuated from the system if they become loose. While such wear particles and the debris layer they form (collectively called the third body) were already observed experimentally a long time ago, the process of their creation is still under discussion. Such an understanding is essential, though, to establish predictive wear laws, which are currently still limited to empirical formulas that are not universally applicable.

Rabinowicz proposed a scenario in which particles become detached from the surfaces and might stick to the other [Fig. 1(a) right], possibly growing and becoming fully loose later on. Based on Griffith’s concept of a critical crack length for unstable crack propagation, one can define a minimum size a wear particle needs to have before it can break off by fracture or detach from a surface it is sticking to. This theoretical argument, though, contains no notion of the asperity collision process and consequently does not describe a shearing process as pictured in Fig. 1(a) left. Furthermore, it ignores the process proposed by Holm, in which plastic deformation of the asperities is responsible for wear. Recently, these theories were combined and extended for the initial particle formation [Fig. 1(a) left], by formulating a critical length scale in terms of a competition between plastic deformation of an asperity under load and its breaking off. In addition to predicting a minimum size of wear particles, it also provides a way to understand surface roughness evolution. Purely plastic deformation of asperities has been found to mostly flatten the surfaces or lead to welding, while the breaking off of wear particles seems to be the ingredient to reroughen worked surfaces, although surface kinks due to dislocation plasticity have also been suggested as sources of roughness. The resulting critical length scale is most generally defined as

$$d^* = \frac{f w_{\text{eff}}}{\overline{\tau}^2/2G},$$

where $w_{\text{eff}}$ is the effective fracture energy and $\overline{\tau}^2/2G$ the elastic energy density of an asperity with shear modulus $G$ when it is loaded to its elastic shear limit $\overline{\tau}$. The length $d$ refers to the diameter of spherical asperities, where asperities with $d < d^*$ deform plastically and those with $d > d^*$ form wear particles. The prefactor $f$ is a shape factor (see Appendix A), which includes information about the actual shape of the asperities and the geometry of the crack that leads to their detachment. The value of the effective fracture energy depends on the case. For the initial detachment [Fig. 1(a) left], two cracks in the material bulk are needed and thus $w_{\text{eff}} = w_b$, i.e., the bulk fracture energy. For the detachment of a sticking particle it is $w_{\text{eff}} = w_{\text{adh}}$, which is the adhesive energy. The details of $\overline{\tau}$ will be discussed in the course of the present work.

This model, though, neglects the fact that the interface formed between asperities is usually mechanically weaker than the bulk. This might be because the junction can effectively be thought of as a grain boundary, or due to surface contamination and chemical passivation, or even the presence of lubricants. In such a case, it would seem reasonable that any fracture occurs along the weaker interface [path 1 in Fig. 1(b)], as has been pointed out before. Since that would preclude the formation of wear particles, several improvements of this simple picture were proposed: If the junction interface has a well-defined yield stress $\overline{\tau}_{\text{adh}}$, it has been suggested that one simply needs to insert...
FIG. 1. Our current understanding of wear particle creation and evolution. (a) Two surface asperities adhere to each other and a wear particle starts to detach by cracks (red) propagating in the bulk material (left). A fully formed particle might be transferred to the opposite surface and get stuck there, based on the strength of adhesion (right). (b) Nevertheless, it is still unclear why the cracks should propagate in the bulk (path 2) instead of along path 1, given that the interface between two surfaces is usually weaker than the bulk. (c) One simple, possible solution is to require a local slope $\theta$ at the asperity contact, which is in fact expected for realistic contacts. Depending on $\theta$, path 2 could then become the preferred fracture path.

It into Eq. 1 instead of the bulk yield stress $\tilde{\tau}_b$, thereby just increasing $d^*$. An alternative solution is to take into account that the junction interface is inclined with respect to the sliding direction in most cases, which suggests that the resulting stress distribution might favor fracture in the bulk based on the inclination angle $\theta$ [Fig. 1(c)]. Indeed, some early work on macroscopic junction models supports the latter hypothesis, but is based on macroscale friction at the interface, which is not applicable on the relevant atomistic scales at junctions. Furthermore, the possibility of wear by atom-by-atom attrition has been found in some ultra-low wear conditions. It seems reasonable to assume that this process occurs when both wear particle formation and severe plastic deformation of the surface are suppressed and the surfaces just slip past each other, but this cannot be explained by the theory described above, since any adhesive contact in this framework either leads to wear particle formation or non-negligible plasticity. Ultra-low wear can only take place when both of these mechanisms are disabled.

In the present work, we therefore re-evaluate the elemental processes of adhesive wear, by now considering both the strength of the interface and its local inclination $\theta$, as well as add slip along the interface to the mechanism map. The competition between plastic deformation inside theasperities, wear particle formation, and slip can account for more realistic conditions with reduced wear, such as lubrication and surface passivation. We develop a simple stress-based model and test it using molecular dynamics (MD) and the finite-element method (FEM) in both plane-strain and full 3D simulations. In particular, we consider the question of the nature of interface slip: Is it plastic-like with a defined yield strength or is it crack-like and activated by stress concentrations?

II. THEORETICAL MODEL

We consider the case of the initial detachment of a wear particle [Fig. 1(a) left], for which we assume a simplified geometry of spherical asperities as shown in Fig. 2. The basic idea of the critical length scale model is that the surfaces keep sliding against each other, loading the asperities in shear, resulting in a shear stress $\tau$. At some point either the yield stress of the material is reached and the asperity deforms plastically, or enough elastic energy is stored in the asperity to open a crack and detach a wear particle. This picture needs to be extended, though, in order to be able to describe slip events at the junction interfaces, which differ from the plastic deformation of the asperities by causing little damage in the material, and which might occur in lubricated or ultra-low wear conditions. Furthermore, as shown in Fig. 1(b), it is not clear why fracture in the bulk should be preferred to slip along the weak interface, especially since the latter usually has a reduced adhesive energy.

In order to model this, we keep the structure of Eq. 1,
but will derive a more complete expression for the critical stress \( \tau \). As can be seen in Fig. 2, slip events will be limited to the plane with normal vector \( \mathbf{n} \) in a slip direction \( \mathbf{m} \) perpendicular to it. Thus, we have to consider a resolved shear stress \( \tau_{RSS} \) on the interface, similar to a generalized Schmid factor.\(^{36}\) We assume that the asperities are loaded in shear by the sliding and that an additional normal stress \( p_N \) is applied. This normal stress is generally not equal to the macroscopically applied load and depends on the actual, local contact solution. The stress state in the junction is then assumed to be

\[
\sigma = \begin{bmatrix}
0 & 0 & \tau \\
0 & 0 & 0 \\
\tau & 0 & p_N \\
\end{bmatrix}.
\]

(Note that \( p_N \) is compressive for asperities in contact and thus negative.) With

\[
\mathbf{n} = \begin{pmatrix}
-\sin \theta \\
0 \\
\cos \theta \\
\end{pmatrix}, \quad \mathbf{m} = \begin{pmatrix}
\cos \theta \\
0 \\
\sin \theta \\
\end{pmatrix},
\]

this results in a resolved shear stress of

\[
\tau_{RSS} = (\sigma \cdot \mathbf{n}) \cdot \mathbf{m} = \tau \cos 2\theta + p_N \sin 2\theta.
\]

A negative \( \tau_{RSS} \) means that the asperities are driven to slide against the macroscopic sliding direction, i.e., the sliding mechanism is not active. This is because \( \tau \) increases with the sliding distance while the asperities interlock and will thus increasingly compensate for any “backsliding” caused by \( p_N \), which is assumed to be constant. For \( \theta \geq 45^\circ \), the slip mode is disabled and the asperities interlock. Here, only plastic deformation in the bulk is expected. In fact, the switch from slippage to interlock occurs at a critical angle \( \theta_c < 45^\circ \), which depends on the normal load and the material properties.

Of course, in a real contact, the normal vector of the junction is not constrained to the \( xz \) plane. Due to the imposed sliding direction, though, the slip direction \( \mathbf{m} \) is constrained in this manner. Hence, the effect on the resolved shear stress is relatively minor as long as the out-of-plane rotation is lower than \( 45^\circ \), as shown in Appendix B. Higher rotation angles should rather be interpreted as asperities passing by each other.

### A. The crossover angle

It stands to reason that the strength \( \tau_{adh} \) of the adhesive interface is smaller than or equal to the bulk yield strength \( \tau_b \). Thus, sliding will be preferred over bulk plasticity if \( \tau_{RSS} (\tau = \tau_b, \theta) > \tau_{adh} \). By solving Eq. 4 with \( \tau = \tau_b \) and \( \tau_{RSS} = \tau_{adh} \) we obtain the crossover angle

\[
\theta_c = \arctan \left( \frac{p_N / 2 \pm \sqrt{\tau_b^2 - \tau_{adh}^2 + p_N^2 / 4}}{\tau_b + \tau_{adh}} \right).
\]

Using the “plus” sign gives positive angles. We do not discuss \( \theta < 0 \) here, since asperity collisions during sliding are frontal. Such a case might nonetheless occur at the onset of sliding, or when the sliding direction changes, but then we expect the adhesive strength to dominate, especially since slip is aided by the normal load instead of suppressed.

### B. A refined critical length scale

For the moment, we will assume that a well-defined yield strength for the interface exists. As such, we can retain Eq. 1 by using a modified critical strength to obtain an angle and load-dependent critical length scale

\[
d^* (\theta, p_N) = \frac{w_{eff}}{\tilde{\tau} (\theta, p_N)^2 / 2G^*}, \quad \text{with}
\]

\[
\tilde{\tau} (\theta, p_N) = \begin{cases} 
\tilde{\tau}_b & \theta \geq \theta_c \\
\tilde{\tau}_{adh} - 0.5p_N \sin 2\theta / \cos 2\theta & \theta < \theta_c.
\end{cases}
\]

In some special cases \( \tilde{\tau} \) is constant and Eq. 1 is recovered:

For full adhesion (\( \tau_{adh} = \tau_b \), noting that \( p_N \leq 0 \)), we always obtain \( \theta_c = 0^\circ \). This means that the slip mechanism is disabled and we recover the originally proposed formula,\(^{46}\) where \( d^* \) is independent of \( \theta \) and \( p_N \). For \( \theta = 0^\circ \), i.e., grazing incident, there is also no dependence on \( p_N \) and the bulk shear strength is simply replaced by the interface shear strength.

### III. VERIFICATION USING MOLECULAR DYNAMICS SIMULATIONS

#### A. Methods and atomistic model

We used MD simulations in order to test and refine the proposed model. We chose an amorphous model material, a silicon-like glass, since worn surfaces often exhibit amorphous top layers\(^{37}\) and because we want to avoid the complications of crystal anisotropy for the present study. Additionally, glasses plastically deform by shear banding,\(^{38–41}\) which is known to have a well-defined yield strength that does not vary significantly in the presence of (flat) surfaces.\(^{42,43}\) Dislocation nucleation, on the other hand, is very sensitive to the nature of interfaces and to size effects,\(^{36}\) making an exact yield strength less accessible for the complex geometry of asperities.\(^{25}\)

To be able to study a large number of geometries, we need to reduce the computational requirements and thus need a brittle material (small \( d^* \)); in our case a silicon-like model material. There is a large choice of potentials for silicon, but most of them suffer from an unphysical ductility.\(^{44–47}\) State-of-the-art potentials fixing this problem exist in the form of screened bond-order potentials,\(^{48}\) but in our testing the computational cost is at least an order of magnitude higher than Stillinger–Weber\(^{49}\) or
Tersoff-type potentials. Instead, we opted for a modified Stillinger–Weber potential, in which the three-body parameter $\lambda_{SW}$ (which governs the bond directionality or stiffness of bond angles) has been doubled from 21 to 42, improving the description of fracture at the cost of other properties. While not suitable to accurately describe real silicon, the tunable brittleness of this model has proven useful to study fracture, as well as the plastic behavior and the vibrational properties of glasses.

All our simulations were performed using LAMMPS, using a timestep of 1 fs for the integration of the equations of motion. Glass samples were prepared following the procedure in Ref. 39: The system was quenched under full periodic boundary conditions from the melt at 3500 K down to 10 K at ambient pressure with a cooling rate of $10^{11}$ K/s, after which energy minimization was performed, yielding a system of size $20 \times 20 \times 20$ nm$^3$ with around 400,000 atoms. This initial preparation was performed with the Tersoff potential for silicon, since using the Stillinger–Weber potential during quenching is known to lead to a glass structure that is too liquid-like in comparison. To obtain the final sample, the system was equilibrated at 300 K and then a normal force was applied to a top and bottom layer (width 4 Å) of atoms in the direction to obtain the desired $p_N$ at the junction. A sliding velocity of 0.002 m/s was chosen and applied in the $z$ direction. Langevin thermostats at 300 K with a damping constant of 0.01 ps were applied next to these boundary layers over a width of 4 Å.

We used OVITO for the visualization and analysis of the simulations. The local von Mises shear invariant $\eta_i$ of the atomic shear strain was used to identify plastic events in the glass.

B. Results in plane strain

In a first step, we performed plane-strain simulations on cylindrical asperities in a quasi-2D geometry with a thickness of 20 nm [see Fig. 4(a)]. While the critical length scale as defined by Eq. 6 contains no fit parameters, it was found in an earlier 2D study that the shape factor $f$ is closer to 1.5 than to the predicted value of $8/\pi \approx 2.5$. In the interface in a layer of 1 nm thickness, leading to a disturbed short-range order as it would occur in a shear band. We investigated the behavior of these interfaces by a shear test on a bulk sample under periodic boundary conditions with a shear rate of $4 \times 10^7$ /s. Figure 3 shows the resulting stress–strain curves and snapshots are provided in Fig. S.1. The bulk material deforms by shear banding, as expected, connected to the typical, rate-dependent stress overshoot. Reduction of the adhesive energy leads to a complete loss of shear strength at the onset of sliding, since the energetic cost of plasticity in the neighboring bulk is higher than for separation of the interfaces during slip (evidenced by a reduced coordination number of silicon atoms at the interface and a lack of plastic events in the bulk), leading to a relatively “brittle” stick-to-slip transition. The damaged layer, however, is close in its behavior to the unmodified bulk material after the onset of plasticity and shows only a slight softening. Since slip along interfaces can be thought of as shear banding, these shear curves can be used as an accurate description of the material behavior in the junctions of ourasperity-contact geometries.
Therefore, we started by performing simulations with full adhesion for different \( \theta \). According to Eq. 6, we expect no influence of the contact angle, since \( \tilde{\tau}_{\text{adh}} = \tilde{\tau}_b \) and thus \( \theta_c = 0^\circ \), meaning that no slip is expected. Indeed, as plotted in Fig. 5(a), we found for all \( \theta \) that asperities smaller than \( d^* \approx 7 \text{ nm} \) deform plastically, while larger asperities detach to form wear particles. Since the fracture of the material is close to perfectly brittle [cf. Fig. 4(d)], we assume that \( w_b \approx 2 \gamma_s \). Together with the other material parameters and \( d^* \), and by using Eq. 6, we derive a value of \( f \approx 1.5 \), consistent with the earlier findings.

Subsequent simulations with reduced adhesion at the interface between asperities exhibit the three predicted mechanisms of slip, plastic deformation of the asperities, and detachment of wear particles [see Fig. 4(b)–(d)]. Slip occurs, as expected, without significant plasticity inside the two sliding bodies [Fig. 4(b)]. In the case of plasticity, only one of the asperities is damaged, while the other stays mostly intact. This was observed before in simulations of asperity collisions in diamond-like carbon\(^{64}\) and is a consequence of the shear softening and strain localization behavior of the material. In strain-hardening materials, a more even distribution of plastic strain would be expected.

Figure 5 shows that the observed mechanisms and the transitions between them are in accord with the model predictions of Eq. 6 for different angles, normal loads, and adhesion potentials. We differentiate between slip and plasticity inside the asperities by calculating the atomic shear strain \( \eta_i \) and checking if plasticity is constrained to the interface or if it occurs in the bulk. As a criterion, we only consider significant, irreversible shape changes of the asperities to indicate bulk plasticity [compare Fig. 4(b) with Fig. 4(c), for example]. With this, we find that the critical angle \( \theta_c \) is correctly predicted.

Note that in order to accurately compare the results to the model, we measured the inclination angle \( \theta \) of the interface just before any non-elastic event occurred. We found that \( \theta \) can be reduced by up to \( 10^\circ \) during the elastic deformation, although the large change is a consequence of the large elastic limit of our model material.

In several cases, we observe that the mechanism can switch, e.g., when a wear particle starts detaching, but the interface rotates to a low-angle plane and starts slipping before the crack propagation can fully unload the contacting asperities (Fig. S.2 in\(^59\)). Nevertheless, the angle at which the switch occurs is in accord with our theoretical model. This hints that the detachment of such asperities could be a fatigue-like process, since the pre-crack that is formed is not expected to heal completely and can possibly grow again more easily upon a following asperity collision.

Moreover, because we control \( p_N \) by a constant force, the actual stress at the junction increases during slip. Therefore, we observe increased plasticity at the end of slip, see Fig. 4(b) and Fig. S.3 in\(^59\). In cases close to \( \theta_c \), this plasticity can be significant. This emphasizes that
the evolution of contacting surfaces is a highly complex and dynamic process, which is not fully predictable from the initial contact configuration. Our mechanism map, though, demonstrates that a simplified picture is not too far from reality and is useful to categorize and understand the asperity-level mechanisms of wear.

C. Results for the 3D geometry

While the plane-strain simulations are useful to explore the possible mechanisms of adhesive wear and suggest that our model is predictive, they do not fully represent all aspects of real contact geometries. Thus, we extended the simulations to a full 3D geometry by considering hemispherical asperities [see Fig. 6(a)]. We again started from simulations with full adhesion and found $d^* \approx 18 \text{ nm}$, which, together with the material parameters and Eq. 6, corresponds to a shape factor of $f \approx 3.5$. This shape factor is in fact the expected factor for overlapping, hemispherical asperities (see Appendix A for the derivation), which means that Eq. 6 is fully predictive for wear particle formation in the 3D case under full adhesion conditions without any fit parameter.

We then extended the simulations to junction interfaces with either reduced adhesion or with a damaged layer. As for the plane-strain simulations, the three mechanisms of slip [Fig. 6(b)–(c)], plasticity [Fig. 6(d)], and particle detachment [Fig. 6(e)–(f)] were observed. As shown in Fig. 7, the mechanism map partly agrees with the model prediction, but we unexpectedly found that interfaces with either reduced adhesion or a damaged interface layer have a tendency for slip at small $\theta$, even for large asperity sizes $d \gg d^*$ (light gray areas on the graphs). We thus repeated one of the simulations displaying slip while outputting data in smaller time increments. Figure 8 indicates that the start of slip occurs at edges lateral to the sliding direction, hinting at slip nucleation by a stress concentration.

In order to elucidate the reason for this difference in behavior between quasi-2D and full 3D simulations, we turned to finite-element modelling of our geometry using dynamically inserted cohesive elements. A detailed description of the setup and results is presented in Appendix C. While this model does not account for plasticity, it is useful to study the transition from interface slip to detachment of wear particles. We find that a shear stress concentration indeed necessarily arises in the same position as in the MD simulation, which leads to slip along the interface, even for $d \gg d^*$. Given the existence of a stress concentration, this is not surprising if the fracture toughness of the junction is reduced compared to the bulk material: The competition between particle detachment and slip is not governed by an energetic criterion as in...
FIG. 6. Snapshots of asperity interactions in the 3D simulations. (a) An example setup, the colors serve to differentiate the two bodies and highlight the interface. The slip mechanism is depicted both for the reduced adhesion (b) and the damaged (c) interface. (d) Plastic deformation of an asperity. (e)–(f) Formation of wear particles. The reduced adhesion interface (e) exhibits less plastic activity than the damaged interface (f), neither interface starts slipping, though. The snapshots (b)–(f) show zooms on the junctions and are colored according to the atomic shear strain $\eta_i$.

FIG. 7. Mechanism map for the 3D simulations. The data points represent MD simulations and indicate the observed mechanisms. Simulations were conducted with (a) full adhesion at the interface between asperities, (b) reduced adhesion (green curve in Fig. 3), and (c) a damaged interface layer (purple curve in Fig. 3). In all figures, the solid lines show the prediction for $d^*$ of Eq. 6: Contacting asperities bigger than $d^*$ are expected to form wear particles (red). Additionally, the dotted lines indicate $\theta_c$, i.e., the transition from significant bulk plasticity at $\theta > \theta_c$ (yellow) to slip at $\theta < \theta_c$ (blue) for smaller asperities. The data show that the predictions are reproduced well, except for the gray areas in (b) and (c). Here, slip can occur in a regime where wear particle formation is expected. For the damaged layer, the predicted behavior can be recovered by blunting the interface (half-filled data points), while the interface with reduced adhesion is not influenced by the blunting. As in Fig. 5, data points connected by arrows indicate a switch from an incomplete particle detachment to slip along the interface. This can occur due to the rotation of the interface while the particle is still forming. The simulation marked with an asterisk (*) exhibited a transition first from particle detachment to slip and then from slip to plasticity, since the effective normal load $p_N$ increases during sliding (cf. Fig. 5). The simulation marked with a dagger (†) exhibited a small slip event along the interface that triggered a shear band and plastic deformation even slightly above $d^*$. 

$\tilde{\tau}_{adh} = \tilde{\tau}_b$  \hspace{1cm} $\rho_N = 0.1\tilde{\tau}_b$

$\tilde{\tau}_{adh} = 0.7\tilde{\tau}_b$  \hspace{1cm} $\rho_N = 0.1\tilde{\tau}_b$

$\tilde{\tau}_{adh} = 0.6\tilde{\tau}_b$  \hspace{1cm} $\rho_N = 0.1\tilde{\tau}_b$

$\tilde{\tau}_b$  \hspace{1cm} $\rho_N = 0.1\tilde{\tau}_b$

$\tilde{\tau}_b$  \hspace{1cm} $\rho_N = 0.1\tilde{\tau}_b$

$\tilde{\tau}_b$  \hspace{1cm} $\rho_N = 0.1\tilde{\tau}_b$
FIG. 8. Evidence of a lateral stress concentration in an MD simulation, which is responsible for the nucleation of slip. The snapshot shows a slice through the junction, with atoms colored according to the atomic shear strain $\eta_i$. The arrow on the junction indicates the sliding direction, and the onset of slip at the lateral corners can be seen. (The flat surface was made darker to make the asperity easier to discern.)

FIG. 9. Snapshots of a blunted interface in a 3D simulation with a damaged interface layer. The asperity has a diameter of $d = 50\,\text{nm} > d^*$ and consequently forms a wear particle as predicted by Eq. 6, in contrast to the same simulation with a sharp interface, which exhibits slip along the interface [cf. Fig. 7(c) at $\theta = 0^\circ$].

Eq. 6 anymore, but by a competition between stress intensity factors. If the junction has full adhesive strength, on the other hand, the mode I crack at the base of the asperity is always preferred over the mode II interface slip (cf. Fig. S.5 in 59). This also explains why the damaged layer in the MD simulations has a higher resistance towards interface slip: It does not weaken catastrophically upon slip and can therefore accommodate small deformations around the stress concentrator. In other words, it has a larger plastic zone size and therefore a higher fracture toughness than the interfaces with reduced adhesion potential (compare the area under the stress–strain curves in Fig. 3).

As such, the competition between slip and wear particle detachment depends on the details of fracture toughness and relative stress intensity factors of the two processes. Of course, the simulations considered until now always possessed sharp interfaces, while contacts might also exhibit blunted corners at the junction (e.g., due to diffusion). To explore if we can recover the original model formulation by reducing the stress concentration, we repeated several MD simulations with blunted corners around the junction as depicted in Fig. 9. We restricted ourselves to simulations around critical points in the mechanism map in order to reduce the computational demands. In the case of the reduced adhesion potential at the interface [Fig. 7(b)], no change of mechanism due to blunting could be observed for $\theta < \theta_c$. This is most likely because the stress concentration is only reduced, but not removed by blunting the interface. Due to the rather abrupt stick-to-slip transition of this interface model, any onset of slip will be catastrophic. For the damaged layer, though, a well-defined yield strength and $d^*$ are recovered for all $\theta$ [Fig. 7(c)]. The value of $d^*$ for blunted interfaces with small inclination angles is slightly overestimated. It should be noted, though, that the damaged interface’s strength is subject to some uncertainty, since the exact damage depends on the somewhat random local short-range order of the glass.

A switch from wear particle detachment to slip due to rotation of the slip plane, as observed in the plane-strain case discussed above, is also present if the interface is “brittle” (reduced adhesion), but not if it is shear-band-like (damaged layer). This emphasizes again that these processes are ultimately controlled by the exact stress distributions and process zones that arise. A purely yield-strength-based argument as laid out in Section II can only apply if the junction interface behaves bulk-like, which most likely applies to mature junctions. Additionally, nanoscale roughness of the junction interface might also help to suppress the “brittle” slip behavior. If that does not apply, a competition between a mode II crack along the interface and a mode I crack that detaches the particle has to be considered for $\theta < \theta_c$ and $d > d^*$ (gray areas in Fig. 7).

Finally, for the derivation of our theoretical model in Section II, we assumed that a deviation of the sliding direction by an angle $\phi$ from the $xz$ plane sketched in Fig. 2 does not change the mechanism map. In Appendix B, we show that this is reasonable for $|\phi| \leq 45^\circ$ under the assumption that the slip of the junction will be constrained by the macroscopic sliding direction. In order to verify this, we repeated the simulations with a damaged interface, initial angle $\theta = 35^\circ$, and $d = 10\,\text{nm}$, 20 nm, and 30 nm and rotated the sliding direction by $\phi = 30^\circ$. As expected, no change in mechanism was observed (Fig. S.6 in 59). This means that Eq. 6 is robust even for asperity collisions that are not perfectly frontal.
IV. DISCUSSION

The mechanism maps outlined above generally agree with the limited available literature. Simulations of asperity collisions in diamond-like carbon were interpreted in terms of the overlap of the asperities, transitioning from localized shear banding at the contact spot to a large plastic zone with increasing overlap. Since the asperities were cylindrical and in general a slightly reduced adhesion due to lattice mismatch can be assumed, the overlap can be reinterpreted as an angle of contact, in agreement with our results of transition from slip to bulk plasticity.

Beyond the nanoscale, the present work also has consequences for our understanding of wear at the engineering scale. First, large angles between the interface and the sliding direction lead to an increased probability of wear particle formation, meaning that a high root mean square roughness of the surface leads to more asperity interlock and more wear. Consequently, though, the roughness should reduce during running-in (by wearing off high and thin asperities), while the real contact area and the size of the individual contact spots increases and the wear rate decreases. Without the slope-dependent slip mechanism, this effect cannot be explained, since increased contact spot sizes should lead to the formation of more and larger wear particles. Our results provide an answer: The local slopes will be flatter due to the change of roughness and thus the probability for slip increases, especially in lubricated conditions. Second, in a previous study, it was found that the wear coefficient cannot simply be derived only from the assumption that contact spots larger than a critical value form wear particles, while smaller ones deform plastically. Indeed, this view misses important complexities, such as interactions between contact spots, plastic deformations due to the normal load, and as the present work indicates, slip. This slip depends on the strength of the adhesive bond and the sharpness of the interface, and as such is subject to time-dependent phenomena such as aging and interface creep. With appropriate knowledge about the parameters of the interface, wear debris formation, surface deformation due to plasticity, and slip can now be treated quantitatively using our framework.

V. CONCLUSION

In conclusion, we derived a revised criterion for the formation of wear particles in the adhesive wear regime by considering not only the properties of the two bulk materials in contact, but by investigating the interface in more detail. For blunted interfaces with sufficient toughness—as would be expected at junctions that are sufficiently mature for diffusive processes to have acted—a well-defined interface shear stress exists. The junction strength in this case can be derived by a competition between the critical resolved interface shear stress (comparable to the concept of the Schmid factor) and the bulk shear strength. Weak interfaces, on the other hand, are very sensitive to stress concentrators, and will easily slip in preference to other mechanisms. This means that only high-angle asperity collisions can lead to wear particle formation. Finally, when the interface strength approaches the bulk strength, no slip occurs anymore and only plastic deformation and wear particle detachment are expected. As a result, flat surfaces are in general more amenable to slip and low wear, as long as they are well lubricated and as long as the contact patches do not grow above a critical size, which is expected to be very large compared to the typical contact junction sizes in lubricated conditions. As soon as lubrication is lost and adhesion increases, the critical size for wear particle formation will drop drastically and severe adhesive wear is expected to set in. The present paper provides a theoretical framework and a quantitative map to predict these transitions.

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Appendix A: Shape factor

The shape factor $f$ in Eqs. 1 and 6 contains information about the actual shape of the asperities and the geometry of the cracks needed to detach a wear particle. In the case of a contact between two asperities, each of volume $V$, the maximal stored elastic energy is $2V\bar{\tau}^2/2G$. Two cracks with area $A$ are needed to form a particle, which requires a work of $2Aw_b$. Often, the assumption of hemispherical asperities with diameter $d$ and with a circular fracture area at their base is made: $A = \pi d^2/4$ and $V = \pi d^3/12$. The balance of the two energies gives $d^* = 3w_b/(\bar{\tau}^2/2G)$ and thus $f = 3$, as shown in earlier work.

In our simulations, the asperities are no longer perfectly hemispherical. Thus, in order to derive the exact shape factor $f$ for our 3D geometries, we first consider the volume of the asperities. Since they are overlapping hemispheres, each asperity can be described as a spherical segment (cf. Fig. 2). A spherical segment is a solid bounded by a radius of $h$, and a volume of $\pi h(3a^2 + 3b^2 + h^2)/6$. Since one cut goes through the sphere’s center, $a = d/2$. The second cut represents the junction, for which we chose a radius of $b = j/2 = 3d/8$. Together with the relation $d^2/4 = a^2 + ((a^2 - b^2 - h^2)/2h)^2$, we obtain a volume of

$$V = \frac{41\sqrt{7\pi}}{1536}d^3 \quad (A1)$$
per asperity. We keep the assumption of a relatively flat crack path, corresponding to the circular base of the asperities. As above, we solve for the critical length by the equating potential energy with the work for crack propagation and obtain

$$d^* = \frac{384\sqrt{\pi}}{287} \frac{w_b}{\tau_0^2/2G} \approx 3.54 \frac{w_b}{\tau_0^2/2G}. \quad (A2)$$

The result of \( f \approx 3.54 \) agrees with the data in Section III C without a fit parameter.

Finally, one can also attempt to define a critical size for a hemispherical wear particle sticking to one of the surfaces. Using the same arguments as before, we obtain a maximal stored elastic energy of \( V\tau^2/2G \) and a total adhesive energy of \( A\tau_0^2 \), also resulting in \( f = 3 \). This mechanism, though, is not as well defined: Rabinowicz assumed that the detachment is due to residual compressive stress and thus used yield strength in tension and Young’s modulus instead of shear strength and shear modulus.\(^5\) But in this case the residual tensile stress is most likely smaller than the yield strength. Rabinowicz thus made a back-of-the-envelope estimation that the residual stress is around 10% of the yield strength and thereby arrives at \( f = 30 \), but did not justify this choice any further. Therefore, this mechanism is not yet fully understood, neither quantitatively nor qualitatively.

**Appendix B: Rotation of the junction interface normal vector out of the \( xz \) plane**

In a 3D geometry, the derivation of resolved shear stress (Eqs. 2–4) is only valid if the normal vector of the slip plane remains in the \( xz \) plane. We will show below that a rotation out of this plane by an angle \( \phi \) does not significantly change the resolved stress, as long as the slip direction remains in the \( xz \) plane. The latter is a reasonable assumption, since the sliding direction is macroscopically imposed.

The additional rotation can be included in the normal vector \( n \):

$$n’ = \begin{pmatrix} -\sin \theta \cos \phi \\ +\sin \theta \sin \phi \\ \cos \theta \end{pmatrix} \quad m = \begin{pmatrix} \cos \theta \\ 0 \\ \sin \theta \end{pmatrix}, \quad (B1)$$

resulting in a resolved shear stress of

$$\tau_{\text{RSS}}’ = (\hat{n} n’) \cdot m = \tau(\cos^2 \theta - \sin^2 \theta \cos \phi) + p_N \frac{\sin 2\theta}{2}. \quad (B2)$$

The term relating to \( p_N \) is unchanged. Figure 10 shows that the change of \( \tau_{\text{RSS}}’ \) with \( \phi \leq 45^\circ \) is minor and can mostly be ignored. As discussed in Section III C and shown in Fig. S.6,\(^59\) this is confirmed by simulations with \( \phi = 30^\circ \), which exhibit the same mechanisms as the simulations with \( \phi = 0^\circ \). For \( \phi > 45^\circ \), slip is preferred, which should be interpreted as asperities passing by each other.

**Appendix C: FEM simulations of slip at the junction**

In order to elucidate the reason for the difference in the slip-to-fracture transition between quasi-2D and full 3D simulations, we turned to simplified finite-element modelling of our geometry. We used the software Akantu\(^80,81\) to model a homogeneous, isotropic, linear elastic material with large deformations. Fracture was modeled using cohesive elements with the linear irreversible law by Shozzi and Molinari.\(^82\) The simulations were performed with an explicit time-integration scheme, where the cohesive elements were inserted dynamically.\(^81\) The geometry and the sliding velocity of 20 m/s match the MD simulations at \( \theta = 0^\circ \). Visualization was performed with ParaView.\(^83\)

We chose parameters to match the model glass and the interface with reduced adhesion (\( \tau_{\text{ad}} = 0.77\tau_0 \)) in the MD simulations: For the linear elastic, isotropic bulk material, a density of 2.17 g/cm\(^3\), a Young’s modulus of 149 GPa, and a Poisson’s ratio of 0.2447 were used.

The cohesive elements inserted in the bulk of the material had a critical insertion stress \( \sigma_c = 7.9 \) GPa (the shear yield strength of the glass) with a critical effective opening displacement of \( \delta_c = 0.664 \pm 0.45 \) nm at which cohesion is lost. This corresponds to a fracture energy of 2.62 J/m\(^2\), which was the same for mode I and mode II opening.

![FIG. 10. Influence of out-of-plane rotation \( \phi \). With increasing \( \phi \), the resolved shear stress in the junction interface increases, thereby promoting interface slip. For \( \phi \leq 45^\circ \) and low \( \theta \), though, this increase is very small. Values are plotted for \( p_N = 0 \), since the influence of the normal load is independent of \( \phi \). The inset shows the definition of \( \phi \), sketched as a top view of an asperity contact: The blue asperity is connected to the blue bottom surface, sliding in direction \( \xi \), while the red asperity belongs to the top surface (not pictured), which slides in direction \( -\zeta \).](attachment:image.png)
FIG. 11. FEM analysis of the difference of stress states between the plane strain and the 3D geometries for junctions with reduced adhesion. (a) In the 2D case of an asperity with \( d < d^* \), the interface starts slipping in the junction center, where the shear stress \( \tau_{xz} \) is highest. The bulk is colored according to the shear stress, while the cohesive elements are colored according to the damage parameter. (b) After the onset of slip, the cracks at the base do not open and wear particle formation is suppressed. (c) For \( d > d^* \), no slip events can be observed and crack propagation leads to particle formation. (d) For the 3D case, only cohesive elements and a slice of the junction are shown. The blue circle represents the base of the bottom asperity, while the junction is colored according to the shear stress. Stress concentrations lateral to the sliding direction are visible and act as nucleation sites for slip (e)–(f). In contrast to the predictions of Eq. 6, such slip occurs even for sizes \( d \gg d^* \).

The 3D case, though, shows a deviation from our initial assumptions. In Fig. 11(d), a shear stress concentration is visible at the junction, lateral to the sliding direction. Despite \( d \gg d^* \), interface slip starts at the stress concentration and propagates inwards. This matches the MD simulation with high time resolution shown in Fig. 8. When the modeled junction interface has the same properties as the bulk material, on the other hand, a wear particle detaches by fracture at the base of the asperities (cf. Fig. S.5 in 59).

An analysis of the stress state in the junction shows the principal difference between the 2D and 3D case: In the \( xz \) plane, the free surfaces of the asperities require

(see Ref. 82 for details). For cohesive elements inserted along the junction interface, we reduced the toughness by setting \( \sigma_c = 5.5 \text{ GPa} \) and \( \delta_c = 0.2 \text{ nm} \) for a fracture energy of 0.55 J/m².

While this model cannot account for plasticity, it is useful to study the transition from interface slip to detachment of wear particles.

Figures 11(a) and (b) show a 2D plane-strain simulation with \( d < d^* \). Clearly, the shear stress is maximal in the center of the contact junction, where a mode II crack nucleates and propagates, leading to slip. The cracks at the bottom of the asperity never fully form (damage much smaller than 1). For \( d > d^* \) [Fig. 11(c)], on the other hand, the interface never damages and a wear particle is detached.
that

\[ \begin{bmatrix} \sigma_{xx} & \tau_{xy} & \tau_{xz} \\ \tau_{xy} & \sigma_{yy} & \tau_{yz} \\ \tau_{xz} & \tau_{yz} & \sigma_{zz} \end{bmatrix} \begin{bmatrix} \cos \theta \\ 0 \\ \sin \theta \end{bmatrix} = \begin{bmatrix} \sigma_{xx} \cos \theta + \tau_{xz} \sin \theta \\ \tau_{xy} \cos \theta + \tau_{yz} \sin \theta \\ \tau_{xz} \cos \theta + \sigma_{zz} \sin \theta \end{bmatrix} = 0. \]  

(C1)

Since we can assume that \( \sigma_{xx} \approx 0 \), it follows that \( \tau_{xz} \) has to be zero. In the case of \( \theta = 0^\circ \), even this assumption is not needed and \( \tau_{xz} \) is required to be exactly zero.

In the \( yz \) plane, though, no constraint is placed on \( \tau_{xz} \):

\[ \begin{bmatrix} \sigma_{xx} & \tau_{xy} & \tau_{xz} \\ \tau_{xy} & \sigma_{yy} & \tau_{yz} \\ \tau_{xz} & \tau_{yz} & \sigma_{zz} \end{bmatrix} \begin{bmatrix} 0 \\ 0 \\ 1 \end{bmatrix} = \begin{bmatrix} \tau_{xy} \\ \sigma_{yy} \\ \tau_{yz} \end{bmatrix} = 0. \]  

(C2)

Therefore, the sharp neck at the interface leads to a stress concentration. Here, the slip mode cannot be treated like plasticity with a defined shear strength, but slippage is crack-like. As demonstrated in Section III C, a blunt interface can reduce the stress concentration and disable this crack-like mechanism in some cases.

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1 F. P. Bowden and D. Tabor, “The area of contact between stationary and moving surfaces,” Proc. R. Soc. Lond. A 169, 391–413 (1939).
2 E. Rabinowicz and D. Tabor, “Metallic transfer between sliding metals: an autoradiographic study,” Proc. R. Soc. Lond. A 208, 455–475 (1951).
3 J. T. Burwell and C. D. Strang, “On the empirical law of adhesive wear,” J. Appl. Phys. 23, 18–28 (1952).
4 J. F. Archard, “Contact and rubbing of flat surfaces,” J. Appl. Phys. 24, 981–988 (1953).
5 E. Rabinowicz, “The effect of size on the looseness of wear fragments,” Wear 2, 4–8 (1958).
6 M. Godet, “The third-body approach: A mechanical view of wear,” Wear 100, 437–452 (1984).
7 M. Godet, “Third-bodies in tribology,” Wear 136, 29–45 (1990).
8 R. Aghababaei, D. H. Warner, and J.-F. Molinari, “On the debris-level origins of adhesive wear,” Proc. Natl. Acad. Sci. U.S.A. 114, 7935–7940 (2017).
9 L. Frérot, R. Aghababaei, and J.-F. Molinari, “A mechanical understanding of the wear coefficient: From single to multiple asperities contact,” J. Mech. Phys. Solids 114, 172–184 (2018).
10 J.-F. Molinari, R. Aghababaei, T. Brink, L. Frérot, and E. Milanese, “Adhesive wear mechanisms uncovered by atomistic simulations,” Friction 6, 245–259 (2018).
11 H. C. Meng and K. C. Ludema, “Wear models and predictive equations: their form and content,” Wear 181–183, 443–457 (1995).
12 E. Rabinowicz, “Practical uses of the surface energy criterion,” Wear 7, 9–22 (1964).
13 E. Rabinowicz, Friction and Wear of Materials, 2nd ed. (John Wiley & Sons, New York, USA, 1995).
14 A. A. Griffith, “VI. The phenomena of rupture and flow in solids,” Phil. Trans. R. Soc. Lond. A 221, 163–198 (1921).
15 R. Holm, Electric Contacts: Theory and Application, 3rd ed. (Springer, Berlin, 1967).
16 R. Aghababaei, D. H. Warner, and J.-F. Molinari, “Critical length scale controls adhesive wear mechanisms,” Nat. Commun. 7, 11816 (2016).
17 E. Milanese, T. Brink, R. Aghababaei, and J.-F. Molinari, “Emergence of self-affine surfaces during adhesive wear,” submitted (2018).
18 M. R. Sørensen, K. W. Jacobsen, and P. Stoltze, “Simulations of atomic-scale sliding friction,” Phys. Rev. B 53, 2101–2113 (1996).
19 P. Spijker, G. Anciaux, and J.-F. Molinari, “Dry sliding contact between rough surfaces at the atomistic scale,” Tribol. Lett. 44, 279 (2011).
20 L. Pastewka, S. Moser, P. Gumbsch, and M. Moseler, “Anisotropic mechanical amorphization drives wear in diamond,” Nat. Mater. 10, 34–38 (2011).
21 J. Zhong, R. Shakiba, and J. B. Adams, “Molecular dynamics simulation of severe adhesive wear on a rough aluminum substrate,” J. Phys. D: Appl. Phys. 46, 055307 (2013).
22 Z.-D. Sha, V. Sorkin, P. S. Brancicio, Q.-X. Pei, Y.-W. Zhang, and D. J. Srolovitz, “Large-scale molecular dynamics simulations of wear in diamond-like carbon at the nanoscale,” Appl. Phys. Lett. 103, 095118 (2013).
23 P. Stoyanov, P. A. Romero, R. Merz, M. Kopnarski, M. Stricker, P. Stemmer, M. Dienwiebel, and M. Moseler, “Nanoscale sliding friction phenomena at the interface of diamond-like carbon and tungsten,” Acta Mater. 67, 395–408 (2014).
24 M. I. De Barros Bouchet, C. Matta, B. Vacher, Th. LeMogne, J. M. Martin, J. von Lautz, T. Ma, L. Pastewka, J. Otschik, P. Gumbsch, and M. Moseler, “Energy filtering transmission electron microscopy and atomistic simulations of tribo-induced hybridization change of nanocrystalline diamond coating,” Carbon 87, 317–329 (2015).
25 S. P. Venugopalan and L. Nicola, “Indentation of a plastically deforming metal crystal with a self-affine rigid surface: a dislocation dynamics study,” Acta Mater. (2018), 10.1016/j.actamat.2018.10.020.
26 K. Komvopoulos, N. Saka, and N. P. Suh, “The mechanism of friction in boundary lubrication,” J. Tribol. 107, 452–462 (1985).
27 I.-M. Feng, “Metal transfer and wear,” J. Appl. Phys. 23, 1011–1019 (1952).
28 A. P. Green, “Friction between unlubricated metals: a theoretical analysis of the junction model,” Proc. R. Soc. Lond. A 228, 191 (1955).
29 J. A. Greenwood and D. Tabor, “Deformation properties of friction junctions,” Proc. Phys. Soc. London, Sect. B 68, 609 (1955).
30 A. Erdemir and C. Donnet, “Tribology of diamond-like carbon films: recent progress and future prospects,” J. Phys. D: Appl. Phys. 39, R311 (2006).
31 T. D. B. Jacobs, B. Gotsmann, M. A. Lantz, and R. W. Carpick, “On the application of transition state theory to
atomic-scale wear,” Tribol. Lett. 39, 257–271 (2010).
32. H. Bhaskaran, B. Gotsmann, A. Sebastian, U. Dreck-
scher, M. A. Lantz, M. Despont, P. Jaroenpibul, R. W. Carpick, Y. Chen, and K. Sridharan, “Ultralow nanoscale wear through atom-by-atom attrition in silicon-containing diamond-like carbon,” Nat. Nanotechnol. 5, 181–185 (2010).
33. T. W. Scharf and S. V. Prasad, “Solid lubricants: a review,” J. Mater. Sci. 48, 511–531 (2013).
34. G. Zeng, C.-K. Tan, N. Tansu, and B. A. Krick, “Ultralow wear of gallium nitride,” Appl. Phys. Lett. 109, 051602 (2016).
35. J. F. Curry, T. F. Babuska, T. A. Furnish, P. Lu, D. P. Adams, A. B. Kustas, B. L. Nation, M. T. Dugger, M. Chand-
dross, B. G. Clark, B. L. Boyce, C. A. Schuh, and N. Argibay, “Achieving ultralow wear with stable nanocrystals-
talline metals,” Adv. Mater. 30, 1802026 (2018).
36. G. Gottstein, Physical Foundations of Materials Science, 1st ed. (Springer, Berlin, Germany, 2004).
37. M. Scherge, D. Shakhvorostov, and K. Pühlmann, “Fundamental wear mechanism of metals,” Wear 255, 395–400 (2003).
38. M. Talati, T. Albaret, and A. Tanguy, “Atomic simulations of elastic and plastic properties in amorphous silicon,” Europhys. Lett. 86, 66005 (2009).
39. C. Fusco, T. Albaret, and A. Tanguy, “Role of local order in the small-scale plasticity of model amorphous materials,” Phys. Rev. E 82, 066116 (2010).
40. M. L. Falk and J. S. Langer, “Deformation and failure of amorphous, solidlike materials,” Annu. Rev. Condens. Matter Phys. 2, 353–373 (2011).
41. A. L. Greer, Y. Q. Cheng, and E. Ma, “Shear bands in metallic glasses,” Mat. Sci. Eng. R 74, 71–132 (2013).
42. Y. Q. Cheng and E. Ma, “Intrinsic shear strength of metallic glass,” Acta Mater. 59, 1800–1807 (2011).
43. K. Albe, Y. Ritter, and D. Šopu, “Enhancing the plasticity of metallic glasses: Shear band formation, nanocompos-
ites and nanoglasses investigated by molecular dynamics simulations,” Mech. Mater. 67, 94–103 (2013).
44. D. Holland and M. Marder, “Ideal brittle fracture of silicon studied with molecular dynamics,” Phys. Rev. Lett. 80, 746–749 (1998).
45. D. Holland and M. Marder, “Erratum: Ideal brittle fracture of silicon studied with molecular dynamics [Phys. Rev. Lett. 80, 746 (1998)],” Phys. Rev. Lett. 81, 4029–4029 (1998).
46. L. Pastewka, P. Pou, R. Pérez, P. Gumbsch, and M. Moseler, “Describing bond-breaking processes by re-
active potentials: Importance of an environment-dependent interaction range,” Phys. Rev. B 78, 161402 (2008).
47. L. Pastewka, M. Mrovec, M. Moseler, and P. Gumbsch, “Bond order potentials for fracture, wear, and plasticity,” MRS Bull. 37, 493–503 (2012).
48. L. Pastewka, A. Klemenz, P. Gumbsch, and M. Moseler, “Screened empirical bond-order potentials for Si-C,” Phys. Rev. B 87, 205410 (2013).
49. F. H. Stillinger and T. A. Weber, “Computer simulation of local order in condensed phases of silicon,” Phys. Rev. B 31, 5262–5271 (1985).
50. J. Tersoff, “Empirical interatomic potential for silicon with improved elastic properties,” Phys. Rev. B 38, 9902–9905 (1988).
51. J. A. Hauch, D. Holland, M. P. Marder, and H. L. Swinney, “Dynamic fracture in single crystal silicon,” Phys. Rev. Lett. 82, 3823–3826 (1999).
52. D. Holland and M. Marder, “Cracks and atoms,” Adv. Mater. 11, 793–806 (1999).
53. C. Fusco, T. Albaret, and A. Tanguy, “Rheological properties vs. local dynamics in model disordered materials at low temperature,” Eur. Phys. J. E 37, 43 (2014).
54. Y. Shi, J. Luo, F. Yuan, and L. Huang, “Intrinsic ductility of glassy solids,” J. Appl. Phys. 115, 045328 (2014).
55. Y. M. Beltukov, C. Fusco, D. A. Parshin, and A. Tanguy, “Boson peak and Ioffe-Regel criterion in amorphous silic-
on-like materials: The effect of bond directionality,” Phys. Rev. E 93, 023006 (2016).
56. S. Plimpton, “Fast parallel algorithms for short-range molecular dynamics,” J. Comp. Phys. 117, 1–19 (1995), https://lammps.sandia.gov/.
57. M. Ishimaru, S. Munetoh, and T. Motooka, “Generation of amorphous silicon structures by rapid quenching: A molecular-dynamics study,” Phys. Rev. B 56, 15133–15138 (1997).
58. M. J. Demkovicz and A. S. Argon, “Liquidielofractive environments act as plasticity carriers in amorphous silicon,” Phys. Rev. B 72, 245205 (2005).
59. See Supplemental Material at [URL will be inserted by publisher] for additional simulation snapshots.
60. Y. Ritter and K. Albe, “Chemical and topological order in shear bands of Cu64Zr48 and Cu60Zr64 glasses,” J. Appl. Phys. 111, 103527 (2012).
61. C. A. Schuh, T. C. Hufnagel, and U. Ramamurthy, “Mech-
anical behavior of amorphous alloys,” Acta Mater. 55, 4067–4109 (2007).
62. A. Stukowski, “Visualization and analysis of atomistic simulation data with OVITO – the open visualization tool,” Modell. Simul. Mater. Sci. Eng. 18, 015012 (2010), https://ovito.org/.
63. F. Shimizu, S. Ogata, and J. Li, “Theory of shear banding in metallic glasses and molecular dynamics calculations,” Mater. Trans. 48, 2923–2927 (2007).
64. J. von Lautz, L. Pastewka, P. Gumbsch, and M. Moseler, “Molecular dynamic simulation of collision-induced third-
body formation in hydrogen-free diamond-like carbon as-
perities,” Tribol. Lett. 63, 26 (2016).
65. P. J. Blau, “On the nature of running-in,” Tribol. Int. 38, 1007–1012 (2005).
66. R. Aghababaei, T. Brink, and J.-F. Molinari, “Asperity-
level origins of transition from mild to severe wear,” Phys. Rev. Lett. 120, 186105 (2018).
67. B. N. J. Persson, “Elastoplastic contact between randomly rough surfaces,” Phys. Rev. Lett. 87, 116101 (2001).
68. L. Pei, S. Hyun, J.-F. Molinari, and Mark O. Robbins, “Finite element modeling of elasto-plastic contact between rough surfaces,” J. Mech. Phys. Solids 53, 2385–2409 (2005).
69. S. Solhjoo and A. I. Vakis, “Continuum mechanics at the atomic scale: Insights into non-adhesive contacts us-
ing molecular dynamics simulations,” J. Appl. Phys. 120, 215102 (2016).
70. L. Frérot, M. Bonnet, J.-F. Molinari, and G. Anciaux, “A Fourier-accelerated volume integral method for elastoplastic contact,” arXiv:1811.11558 [cond-mat.soft] (2018).
71. Z. Li and I. Szufarska, “Multiphysics model of chemical aging in frictional contacts,” Phys. Rev. Materials 2, 063602 (2018).
72. Z. Li, L. Pastewka, and I. Szufarska, “Chemical aging of large-scale randomly rough frictional contacts,” Phys. Rev. E 98, 023001 (2018).
73. S. Dillavou and S. M. Rubinstein, “Nonmonotonic aging and memory in a frictional interface,” Phys. Rev. Lett. 120,
R. L. Coble, “A model for boundary diffusion controlled creep in polycrystalline materials,” J. Appl. Phys. 34, 1679–1682 (1963).

M. F. Ashby, “A first report on deformation-mechanism maps,” Acta Metall. 20, 887–897 (1972).

E. Arzt, M. F. Ashby, and R. A. Verrall, “Interface controlled diffusional creep,” Acta Metall. 31, 1977–1989 (1983).

C. Kalcher, T. Brink, J. Rohrer, A. Stukowski, and K. Albe, “Interface-controlled creep in metallic glass composites,” Acta Mater. 141, 251–260 (2017).

Please note that in the original publication, the factor 2 in front of $G$ was moved into $f$ and $w$ included a factor 2 to account for the fact that two cracks are needed for particle formation. These differences cancel out and we arrive at the same value of $f$.

Note that while asperities with $\theta \neq 0^\circ$ are not spherical segments, they have the same volume as long as the plane for cutting the junction is rotated around the sphere’s center and as long as the cuts do not intersect. Both conditions are true for all geometries we used.

N. Richart and J.-F. Molinari, “Implementation of a parallel finite-element library: Test case on a non-local continuum damage model,” Finite Elem. Anal. Des. 100, 41–46 (2015), https://akantu.ch.

M. Vocialta, N. Richart, and J.-F. Molinari, “3D dynamic fragmentation with parallel dynamic insertion of cohesive elements,” Int. J. Numer. Meth. Engng. 109, 1655–1678 (2017).

L. Snozzi and J.-F. Molinari, “A cohesive element model for mixed mode loading with frictional contact capability,” Int. J. Numer. Methods Eng. 93, 510–526 (2013).

J. Ahrens, B. Geveci, and C. Law, “ParaView: An end-user tool for large-data visualization,” in Visualization Handbook, edited by Charles D. Hansen and Chris R. Johnson (Elsevier Butterworth–Heinemann, 2005) Chap. 36, pp. 717–731, https://www.paraview.org/.
Supplemental Material

Adhesive wear mechanisms in the presence of weak interfaces:
Insights from an amorphous model system

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Fig. S.1: Snapshots of bulk shear tests with periodic boundary conditions. All simulation boxes have a size of roughly 20 × 20 × 20 nm$^3$. (a) Homogeneous bulk glass at a macroscopic shear strain of $\gamma = 24\%$, color coded according to the local atomic shear strain $\eta_i$. A single, dominant shear band is visible. (b) Simulation with a damaged layer (arrows) of width 1 nm at $\gamma = 12\%$. The shear band is forced onto the plane of the damaged layer, but some diffuse plasticity can be observed around the shear band. (c)–(d) The simulations with interfaces were modeled by two identical material blocks with a reduced adhesion potential between them. Due to the periodic boundary conditions, this results in two interface planes (arrows). Slip is completely confined to one of these two planes and no significant plasticity is observed next to the interface. The snapshots were taken at shear strains of 12\% (c) and 4\% (d), respectively.
Fig. S.2: Slip after partial wear particle detachment due to decreasing inclination. Snapshots from a simulation with interface adhesion strength of $\tilde{\tau}_{\text{adh}} = 0.7 \tilde{\tau}_b$, $d = 10$ nm, $p_N = 0.1 \tilde{\tau}_b$, and initial $\theta = 30^\circ$. At the onset of fracture, $\theta = 25^\circ$. As soon as the interface rotates to $\theta \approx 15^\circ$, slip sets in.

Fig. S.3: Plastification during slip due to increasing normal pressure. Snapshots from a simulation with interface adhesion strength of $\tilde{\tau}_{\text{adh}} = 0.15 \tilde{\tau}_b$, $d = 5$ nm, $p_N = 0.1 \tilde{\tau}_b$, and initial $\theta = 45^\circ$. The junction area reduces after the onset of slip, leading to an increase of $p_N$ and plastic flow.

Fig. S.4: Sensitivity of the interface to stress concentrations. Simulations with interface adhesion strength of $\tilde{\tau}_{\text{adh}} = 0.15 \tilde{\tau}_b$, $d = 30$ nm, $p_N = 0.1 \tilde{\tau}_b$, and initial $\theta = 45^\circ$. (a) If the inclination $\theta$ is high, its edge can come close to the stress concentration at the corner of the asperity. This seems to lead to the nucleation of a slip front even in cases where our model predicts particle detachment. (b) By putting the asperities on small "sockets" (height $\approx d/10$), the stress concentrator is moved away from the junction interface and a wear particle detaches, as expected. The latter procedure was performed for the simulations with initial $\theta \geq 45^\circ$ in Fig. 5(c) and (d).

Fig. S.5: FEM simulation of a contact junction without weakened interface. The left image shows only the solid, while the right image includes the cohesive elements. The cracks appear, as expected, at the bottom of the asperities and lead to the detachment of a wear particle.
Fig. S.6: Simulations with a sliding direction deviating from the xz plane. It can be seen here—as expected from the derivation in Appendix B—that the mechanism is unchanged for a rotation of the sliding direction by $\phi = 30^\circ$: Asperities with $d \leq 20 \text{ nm}$ plastify (a)–(d), while asperities with $d = 30 \text{ nm}$ form wear particles (e)–(f), independent of $\phi$. 