Atomistic shock Hugoniot simulation of single-crystal copper

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Planar shock waves in single-crystal copper were simulated using nonequilibrium molecular dynamics with a realistic embedded atom potential. The simulation results are in good agreement with new experimental data presented here, for the Hugoniot of single-crystal copper along \( \langle 100 \rangle \). Simulations were performed for Hugoniot pressures in the range 2 GPa – 800 GPa, up to well above the shock induced melting transition. Large anisotropies are found for shock propagation along \( \langle 100 \rangle \), \( \langle 110 \rangle \), and \( \langle 111 \rangle \), with quantitative differences from pair potentials results. Plastic deformation starts at \( U_p \approx 0.75 \text{ km/s} \), and melting occurs between 200 and 220 GPa, in agreement with the experimental melting pressure of polycrystalline copper. The Voigt and Reuss averages of our simulated Hugoniot do not compare well below melting with the experimental Hugoniot of polycrystalline copper. This is possibly due to experimental targets with preferential texturing and/or a much lower Hugoniot elastic limit. © 2004 American Institute of Physics.

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I. INTRODUCTION

Shock waves have long been used to study the equation of state of materials at extreme conditions.\(^1\)\(^-\)\(^3\) In addition, experiments on shock-induced plasticity, and fracture have provided useful insight into material deformation and failure.\(^4\) However, a number of issues are still not well understood. The constitutive equations are typically based on an equation of state that assumes isotropic material response, an assumption which is certainly not true for single crystals, and may even pose a problem for textured polycrystals. Furthermore, in the context of plasticity, a strong indication of limitations of our understanding is given by the classic Frost and Ashby deformation maps which show an “unexplored region” for deformation occurring at strain rates higher than \( 10^9/\text{s} \).\(^5\) Laser-induced shocks\(^6\) provide a new way to produce very high strain rates, up to \( \sim 10^{10}/\text{s} \) for experiments to be carried out in the near future at the National Ignition Facility (NIF). In addition, recent data for single crystals\(^7\)\(^,\)\(^8\) suggest that the plastic response in Cu occurs rapidly, but even with the fast dislocation motion in Cu, the kinetics are likely to be important on time scales that will soon be accessible in experiment (\( \sim 100 \text{ ps} \)).

There are current limitations on our abilities to model dynamic shock processes accurately, but advances in computing have greatly extended the capabilities of numerical simulations. In particular, molecular-dynamics (MD) simulations solve Newton’s equations of motion for a collection of interacting particles over a number of time steps.\(^9\) The size of the simulated system is limited by the number of available processors, and simulations of \( 10^7 – 10^{10} \) atoms are now possible using the largest parallel computers. MD simulations generally probe strain rates well above \( 10^6/\text{s} \) and, therefore, are a natural complement to understand atomic level mechanisms during shock compression. Linking to longer time and larger length scales could be accomplished within a multiscale framework, for instance, by dislocation dynamics informed by MD simulations and coupled to a finite elements mesh.\(^10\)

Most atomistic shock simulation studies have investigated single-crystal response to shocks along the \( \langle 100 \rangle \)
direction.\textsuperscript{11–13} On the other hand, nearly all experimental studies of shock waves in metals have been performed with polycrystalline samples.\textsuperscript{1–3} Clearly, in the majority of metals, the directional anisotropies in single crystals will give rise to direction-dependent Hugoniot relationships. Such anisotropies could be mapped by MD simulations and provide a guide to future and ongoing\textsuperscript{7,8} experimental efforts. Germann \textit{et al.}\textsuperscript{14} presented results for the Hugoniot of a Lennard Jones (LJ) fcc crystal showing a rich variety of behaviors depending on shock orientation.

A shock wave can be produced if a surface force, to which we will refer as a piston, is steadily applied to a material on one side. In the limit of zero piston pressure we expect to obtain a shock velocity equal to the longitudinal sound speed. The longitudinal/transverse sound speeds, in an isotropic elastic medium, can be calculated as \( c_{\text{elL}} = \sqrt{A_{\text{el}}/\rho} \), where \( K \) is the bulk modulus, \( \rho \) is the density of the material, and \( A_{\text{el}} = 3(1-\nu)/(1+\nu) ; A_{\text{el}} = 3(1-2\nu)/(1+\nu) \), with \( \nu \) the Poisson ratio. For nonzero piston pressure, the Hugoniot relationships,\textsuperscript{15} i.e., conservation of mass, momentum, and energy at the shock front, apply and give the shock speed \( U_s \) as \( U_s = U_p/e \), where \( U_p \) is the particle velocity and \( e = (1 - \rho_0/\rho) \) is the volumetric compressive strain. In the strong shock regime, when the plastic wave has overdriven the elastic wave, it is typically found that

\[
U_s = U_o + s_1 U_p, \tag{1}
\]

where \( s_1 \) is a constant in the range 0.5–2.5, and \( U_o \sim c_o \), the bulk sound speed. Of course, if there is a phase transition, the slope of the Hugoniot may change, and this has often been used as a diagnostic to detect such a transition. Using reasonable approximations for a model solid, it can be shown that the Grüneisen parameter \( \gamma_G \) is a function only of the compression, \( e \), and \( s_1 \).\textsuperscript{16} For the limit of \( e = 0 \), \( \gamma_G = 2s_1 - 1 \).\textsuperscript{15,16} On the other hand, the fact that \( s_1 \) is a constant for strong shocks implies that there is a limiting compression value, \( e = 1 - 1/s_1 \). The Grüneisen parameter at this compression limit is often given as \( \gamma_G = 2(s_1 - 1) \), which is smaller than the previous value by \( 1 \).\textsuperscript{16}

The relatively simple picture above is no longer true for an anisotropic solid. For instance, for propagation along \( \langle 100 \rangle \) in a cubic crystal, \( c_{\text{elL}}(\langle 100 \rangle) = \sqrt{c_{11} / \rho} \) and \( c_{\text{elT}}(\langle 100 \rangle) = \sqrt{c_{44} / \rho} \), where \( c_{ij} \) are the elastic constants of the cubic crystal. Experimentally, for Cu the asymmetry is large, \( c_{\text{elL}}(\langle 100 \rangle) / c_{\text{elT}}(\langle 100 \rangle) = 1.49 \), with the anisotropy ratio \( A = 2c_{44}/(c_{11} - c_{12}) = 3.21 \), compared to \( A = 1 \) for the isotropic case.\textsuperscript{17} The “isotropic” Grüneisen parameter is no longer applicable. One needs to calculate an anisotropic Grüneisen parameter and also needs a direction-dependent equation of state.\textsuperscript{18}

Typically, three regions may be identified in the \( U_s \) – \( U_p \) Hugoniot.\textsuperscript{14} For \( U_p < U_{\text{HEL}} \), only an elastic front is observed. At the Hugoniot elastic limit (HEL), a plastic wave appears, and this wave may be underdriven, moving slower than the elastic wave up to \( U_p < U_{\text{ps}} \). For \( U_p \geq U_{\text{ps}} \) the plastic wave overdrives the elastic front, which is not stable. In this last regime the velocity of the elastic front is not the same as the one of the plastic front, but lower. The region \( U_{\text{HEL}} \leq U_p < U_{\text{ps}} \) may be narrow or not exist at all, depending on the material. For any crystal direction, in the limit \( U_p \to 0, U_s \to c_{\text{elL}} > c_o \). On the other hand, in the strong shock regime Eq. (1) is valid. Since \( c_{\text{elL}} > c_o \), there must be a change in slope for velocities below the strong shock regime.

The dependence of the plastic wave speed on orientation may be understood in terms of both elastic and plastic anisotropy. Using the result of Drugan\textsuperscript{19} the steady-state shock behavior may be interpreted using the solution for a smooth wave. One such smooth wave solution for a rate-independent elastoplastic material, given by Lubliner,\textsuperscript{20} gives the wave speed as

\[
U_s = \sqrt{A_s / \rho}, \tag{2}
\]

with

\[
A_1 = K + (2/3)h - 2h^2/(3h + 6\mu), \tag{3}
\]

where \( \mu \) is the shear modulus, and \( h \) is the plastic modulus (hardening rate). For \( h \ll \mu \) (the usual case), this reduces to

\[
A_1 \sim K + (2/3)h. \tag{4}
\]

Thus the plastic wave speed depends in general upon bulk modulus, the shear modulus, and the plastic modulus. Although the bulk modulus of a material with cubic symmetry is independent of loading direction, both the shear and plastic response depend quite strongly on orientation. For copper, the shear modulus varies by over a factor of 3 depending on orientation\textsuperscript{21} while the strain hardening depends strongly on loading direction.\textsuperscript{22} As the shock strength increases the hardening response saturates resulting in less plastic anisotropy. Given these combined effects of elastic and plastic anisotropy, it is expected that the shock response of single crystals (and textured polycrystals) should be anisotropic.

In principle, if one is only interested in the Hugoniot curve for polycrystals, one can obtain very good agreement with experiments by calculating an accurate bulk modulus as a function of the pressure and temperature,\textsuperscript{23} and any potential giving that functional form will also suffice. This is because the constant \( c_0 \) is equal to the bulk sound velocity at zero pressure (\( \sqrt{K/\rho} \)).

With this information one can obtain the equilibrium states that form the Hugoniot. However, this approach, or the use of equilibrium MD calculations,\textsuperscript{24} does not provide any information on when will plastic behavior start, what kind of plastic behavior will be found, etc. This is where nonequilibrium MD simulations play an important role, with the disadvantage that they are computer intensive. Constrained techniques, like the “Hugoniostat,”\textsuperscript{13} can bridge these two approaches. In this work, we present a comparison of experiments and simulation for the (100) shock Hugoniot of solid Cu, and simulation results for the Hugoniot along other crystalline directions, finally averaging these results to compare with experiments on polycrystals. The overall behavior for an fcc LJ solid presented by Germann \textit{et al.}\textsuperscript{14} is qualitatively simi-
lar to the one seen here for embedded-atom method (EAM) potentials, but important quantitative differences arise as discussed below.

A. Experiments

Plate impact experiments [(one-dimensional 1D) strain] were performed on single-crystal copper using the 35 mm light gas gun at Lawrence Livermore National Laboratory. The flyer plate (impactor) of polycrystalline copper was 1.5 mm thick. The target plate was 5 mm thick. The experiments were designed to look at the spall behavior of copper and measured free surface velocity using laser interferometry.\textsuperscript{25} Hugoniot measurements were backed out of the data using an impedance matching technique. At the flyer-target interface, pressure, and particle velocity are identical in both samples. Using the Hugoniot relation \( P = \rho_0 U_p U_s \), where \( P \) is the Hugoniot pressure and \( \rho_0 \) is the initial density, allows one to calculate the shock velocity of the target assuming that one knows the Hugoniot of the polycrystalline copper flyer [a standard - \( U_s = 3.94 \text{ km/s} + 1.489 U_p(\text{km/s}) \)] and the particle velocity of the target which is taken to be one half the free surface velocity.

B. MD Simulations

The simulations were performed with the mdcask code,\textsuperscript{26} adapted to simulate shock waves.\textsuperscript{27} A box elongated along the \( z \) direction was equilibrated during several picoseconds at 1.5 K, using periodic boundaries only along the \( x \) and \( y \) directions. A square pulse shock wave was applied along \( \tilde{z} \) by adding an external force to few planes of atoms on one of the free surfaces. The applied force was kept constant throughout these simulations, although a time dependent profile could be applied. Velocity profiles were analyzed at subsequent times to calculate both \( U_p \) and \( U_s \). Following a transient stage, typically around 0.5 ps, the elastic shock profiles reached a steady state, allowing a determination of \( U_p \) and \( U_s \) with errors generally smaller than 5%. Both plastic and elastic fronts were seen in the simulations above a “plastic threshold”, as for the LJ simulations.\textsuperscript{14} Our simulations, employing a planar, but fully 3D geometry, are quite different from “equilibrium” simulations where the Hugoniot is extracted from a “hydrostatic” compression at relatively small system sizes.\textsuperscript{24} Thus, we can easily capture the evolution of the shock without the assumption of local thermal equilibrium, together with the detailed dynamics of the deformation originating at the front.

Most “nonequilibrium” shock simulation studies of fcc solids have used pair potentials, including LJ,\textsuperscript{11–14} Morse,\textsuperscript{28} exponential-6 (Ref. 29), etc. Taylor and Dodson recently presented results on the Hugoniot of EAM Cu along (100) using an EAM potential, for \( U_p \in (0, 2.5) \) and a target with \( 4 \times 4 \times 30 \) unit cells.\textsuperscript{28} Ryazanov et al.\textsuperscript{30} have also presented some simulations of point defect formation by multiple weak shock waves in EAM Cu, and Kum\textsuperscript{31} has shown results for shock waves at a single \( U_p \) above the plastic limit for EAM and Morse Ni along \( \langle 100 \rangle \), \( \langle 110 \rangle \) and \( \langle 111 \rangle \). None of these simulations have been compared with experimental data on single crystals.

Table I. Parameters for the potentials discussed in the text. “\( r_{\text{cut}} \) shell” indicates the last shell of neighbors included by the potential cutoff. \( d_{\text{SF}} \) is the equilibrium separation between SF at \( P = 0 \).

<table>
<thead>
<tr>
<th>Potential</th>
<th>LJ*</th>
<th>EAM1</th>
<th>EAM2</th>
<th>Expt.</th>
</tr>
</thead>
<tbody>
<tr>
<td>( r_{\text{cut}} ) shell</td>
<td>Second</td>
<td>Third</td>
<td>Fourth</td>
<td>( \cdots )</td>
</tr>
<tr>
<td>( \gamma_\text{SF}(\text{mJ/m}^2) )</td>
<td>0</td>
<td>11.4</td>
<td>44.0</td>
<td>45.0^* – 78.0^*</td>
</tr>
<tr>
<td>( d_{\text{SF}}(\text{nm}) )</td>
<td>( \infty )</td>
<td>( -8 )</td>
<td>( -2 )</td>
<td>( -2 ) – ( -1 )</td>
</tr>
</tbody>
</table>

\textsuperscript{*Reference 14.} \textsuperscript{34}^b Reference 34. \textsuperscript{35}^c Reference 35.

Here we consider two many body potentials, of the embedded atom form, EAM1 (Ref. 32) and EAM2.\textsuperscript{23} For shock waves along (100) most simulations were performed with sample sizes of \( 50 \times 50 \times 200 \) fcc cells, i.e., \( 2 \times 10^6 \) atoms. Using the lattice parameter for Cu, \( a_0 = 3.615 \text{ Å} \), the size of the sample was \( 18.1 \times 18.1 \times 72.3 \text{ nm}^3 \). Several of the results for EAM2 were obtained for smaller samples with a cross section of \( 25 \times 25 \) cells, and results for these two sample sizes were indistinguishable for the EAM2 potential. Near the threshold for plasticity, plastic shock waves appeared few picoseconds after the shock was applied and it took them few additional picoseconds to reach a steady profile, stressing the need to simulate very large samples (more than 100 fcc cells long) for reliable calculation of \( U_s, \text{plastic} \) with this scheme. For the other two crystalline directions similar sample sizes were used. The following velocities will be given in km/s, unless noted otherwise.

For any pair potential, \( c_{12} = c_{44} \), which fixes the anisotropy. This relationship does not hold experimentally for metals and many body potentials, such as EAM, are needed to solve this discrepancy. In addition, it is important to point out that any potential with range \( r_{\text{cut}} \) less than the third nearest neighbor distance will have a stable stacking fault (SF) energy equal to zero, \( \gamma_{\text{SF}} = 0 \). These potentials may have a nonzero, small, unstable SF energy which will provide a barrier for dislocation nucleation. However, once partial dislocations are nucleated, only \( \gamma_{\text{SF}} > 0 \) results in a finite separation between partial dislocations in the fcc crystal, \( d_{\text{SF}} \). Therefore the behavior of dislocations cannot be accurately simulated for short range potentials. Table I shows some relevant properties for the potentials discussed in the text. \( d_{\text{SF}} \) is given at zero pressure and may increase under pressure. At a minimum, the lateral size of an MD target must accommodate \( d_{\text{SF}} \). Using the zero-pressure value for \( d_{\text{SF}} \) at least six unit cells are required in the lateral direction for the EAM2 potential and 22 for the EAM1 potential. EAM2 was fitted to the \( ab \text{ initio} \) cold curve of Cu, making it particularly attractive for shock simulations.

II. RESULTS

In Fig. 1 we have plotted the simulated \( \langle 100 \rangle \) Hugoniot along with recent single-crystal Cu gas-gun data. The EAM2 potential is in good agreement with the experimental results, while the EAM1 potential is shifted slightly to higher values of \( U_s \). Results for both EAM1 and EAM2 are quite close, indicating that the long range part of the potential (beyond third nearest neighbor) does not play a key role in the \( \langle 100 \rangle \).
Hugoniot below melting. The different values of $\gamma_{EF}$ do not make a significant difference for the elastic Hugoniot, but do impact the plastic deformation at a given $U_p$, giving different stacking fault densities for the two potentials at the same piston velocity. The fit to the strong shock regime for a LJ crystal is also included, and shows a large deviation with respect to the experimental data and the EAM results. The LJ potential gives $U_{s} = c_{ol,(100)} + 1.92 U_{p}$, where the value of $c_{ol,(100)}$ could be fit to the experimental data by choosing the appropriate LJ parameters. The EAM2 potential gives $U_{s} = (4.1 \pm 0.1) + (1.3 \pm 0.1) U_{p}$ for the interval $U_{p} = (0, 1.5) \ldots$ and it is extremely close to the experimental data. The Morse potential provides a good way to compare results for many body and pair potentials, since a LJ potential is very close to body and pair potentials, since a LJ potential is very close to

As expected, when $U_p = 0$, $U_s - c_{ol} = 4.3$. Indeed, Fig. 1 shows a small plateau, where the shock velocity stays constant within our error bars as $U_p = 0$. Plastic response appears at $(U_p/c_{ol}) \sim 0.14$, that corresponds to a compression of about 14%. This is lower than the value for short-range LJ, which was $(U_p/c_{ol}) \sim 0.2$, but happens at about the same compression. The pressure for the HEL is therefore $32 \pm 2$ GPa, much higher than experimental values. This is not unexpected, since even “perfect” crystals used in the experiments have a pre-existing density of defects that act as nucleation sites for plasticity to begin at lower stress value. Holiann and Lomdahl showed that a nonplanar piston leads to a reduced HEL, and preliminary simulations we have carried out including extended defects in our perfect crystal do lower the HEL significantly.

The shear stress behind the shock front is only a small fraction of the applied stress, while the hydrostatic pressure is comparable to the applied stress. Therefore, the net shear stress near the plastic threshold is only few GPa, similar to the ideal yield stress of single crystals. It is difficult to calculate the HEL for uniaxial compression, but estimates assuming a perfect elastoplastic solid without strain rate effects are often used.

The varied behavior seen in Fig. 2 translates into widely different Hugoniot relationships along different orientations. Figure 4 shows the MD results using the potential EAM2 for the shock Hugoniot along $\langle 100 \rangle$, $\langle 110 \rangle$ and $\langle 111 \rangle$ directions.

\begin{align}
\text{HEL} &= P_H + \frac{2}{3} Y_0 \\
&= (1 - \nu ) Y_0 \\
&= \left( \frac{K}{2\mu} + \frac{2}{3} \right) Y_0,
\end{align}

where $Y_0$ is the yield strength of the material, and all quantities are evaluated at the appropriate hydrostatic pressure $P_H$ which is not known \textit{a priori}. Using zero pressure values Eq. (5c) gives $-4.6$ GPa, and employing values of $K$ and $\mu$ at $-35$ GPa does not give a significant increase in this estimate. The large difference between the HEL from MD and this estimate indicates the limitations of using simplified models for shock behavior in solids.

Figure 2 shows snapshots of the velocity profile along the $z$ direction 3 ps after the shock was applied, for a piston pressure close to 100 GPa. In all cases there is an elastic and plastic wave. For the $\langle 100 \rangle$ direction both the plastic and elastic wave move at roughly the same speed, and can only be separated examining the structure of the sample behind the shock. For the $\langle 111 \rangle$ direction, an elastic precursor is seen, followed by the plastic wave.

In the LJ simulations a number of elastic precursors was seen at zero temperature for shock waves along $\langle 100 \rangle$ and for shock waves along $\langle 110 \rangle$ when the initial temperature of the sample was below $T_{melt}/10$. We observe elastic precursors at finite temperature below the plastic limit for $\langle 100 \rangle$, and for all simulated piston pressures along $\langle 110 \rangle$. For instance, Fig. 3 shows one snapshot 4 ps after the shock for $P_H \sim 50$ GPa. The kinetic energy map shows the layering of planes, alternating low and high kinetic energy regions, due to plane-plane collisions. A potential energy map shows the complimentary effect, as in a “harmonic oscillator,” with lowest potential energy corresponding to highest kinetic energy and vice versa.
The experimental data for polycrystalline Cu for $U_p < 4$ can be reasonably well fit by $U_s = 3.96 + 1.5 U_p$ for the pressure range simulated, and this fit is also shown in Fig. 4. Melting was detected by both the jump of the Hugoniot in the pressure-temperature plane and the pair correlation function of our shocked sample giving a liquid structure. Below melting, the $k_{100}$ and $k_{110}$ curves have slopes much lower than those obtained from the LJ potential simulations. The slope of the Hugoniot does not change noticeably due to melting, which occurs in the range 200–220 GPa (2.0–2.2 Mbar), in agreement with experiments on polycrystalline Cu, with equilibrium “hydrostatic” MD simulations, and with a recent shock-release model of melting. The higher values for the $k_{110}$ Hugoniot are related to the plane-plane collisions that propagate the shock faster than along the other two directions. Using our simulation data and $g_{ijk} = 2 s_{ijk} - 1$, we have calculated the limiting value of the directional Grüneisen parameters. These values are 1.6, 3.4, and 3.1 for $k_{100}$, $k_{110}$ and $k_{111}$. For shock data of polycrystalline Cu, the experimental isotropic Grüneisen parameter, $\gamma_{ij}^{\text{iso}}$, has been reported to be 1.99 and 1.98, in Refs. 45 and 15, respectively.

How can we relate our simulations to the many experimental results on polycrystalline Cu? There have been a number of studies using numerical continuum models of shock propagation through rectangular or spherical grains, but unfortunately, different values of the shock velocities were not taken into account. A model numerical simulation of shock propagation through a polycrystalline slab with anisotropic shock velocities shows large deviations from the isotropic case, even for small anisotropies. As an initial step, we have attempted to provide limiting bounds for polycrystalline Cu using the Voigt ($\overline{U}_{\text{V}}$) and Reuss ($\overline{U}_{\text{R}}$) averages of our single-crystal results. This analysis assumes that grain boundary effects can be neglected and that the grain distribution along directions other than
tatively the results for pressures below melting agree with results from LJ solids, with several quantitative differences.

We observed that the shock velocity has a plateau, with $U_p = c_{dl}$, in the elastic region at low $U_p$. The HEL occurs at 32±2 GPa for all studied directions, at a compression of ~14% and a ratio $(U_p/c_{dl}) = 0.14–0.2$, similar to the LJ potential results. Since simulations were carried out for perfect single crystals, the HEL from MD is much higher than the experimental value and estimates based on a simple elastoplastic model of the solid. Melting is found to happen in the range 200–220 GPa, in agreement with both experiments and equilibrium MD simulations. Melting happens at $(U_p/U_s) = 0.375$, i.e., $(U_p/c_{dl}) = 0.5–0.75$, while for the LJ crystal $(U_p/c_{dl}) = 1$ at melting.

The slopes of the Hugoniot along (100) and (111) are much softer for EAM Cu than for the LJ system. The (110) and (111) results which were relatively close to each other for LJ in the strong shock regime below melting, now show large differences. As shock pressure increases, the Hugoniot curves for all directions do converge to the polycrystalline result, as expected, but the convergence happens near the melting pressure for the plastic front and at even higher pressures for the elastic precursor.

We also presented a simplified analysis of the average shock velocity for polycrystalline Cu, and find that the value of the average Hugoniot is up to 20% larger than the experimental Hugoniot for polycrystals. This difference decreases if one assumes a dominant (100) texture. This is consistent with the observation that a large contribution from (100) texturing is required in polycrystalline experiments to account for the experimental value of the limiting Grüneisen parameter. The calculation of directional Grüneisen parameters for both pair and many body potentials would allow a direct comparison of our limiting compression factors, and would pave the way for future anisotropic equations of state.

We have carried out some preliminary Hugoniot calculations for crystals including defects which decrease the HEL, and they seem to indicate that the elastic Hugoniot is not changed, while the plastic Hugoniot moves closer to the polycrystalline Hugoniot data. Even well annealed Cu single crystals have a dislocation density $10^5–10^6$/cm$^2$, leading to a low experimental HEL. Therefore, dislocation sources, and not texture, could be an alternative explanation to the departure of the calculated single crystal Hugoniot with respect to the polycrystal Hugoniot, even after averaging over different directions.

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**III. SUMMARY**

We have presented MD simulations of the Hugoniot relationship for Cu, for pressures in the range of $2–800$ GPa (20 kbar–8 MBar), which includes the melting transition. Our simulation results agree well with new experimental data for single crystal shocks along (100) and $U_p = 1$. Qualitatively the results for pressures below melting agree with results from LJ solids, with several quantitative differences.

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