Virtual melting as a new mechanism of stress relaxation under high strain rate loading

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Generation and motion of dislocations and twinning are the main mechanisms of plastic deformation. A new mechanism of plastic deformation and stress relaxation at high strain rates (109-1012 s-1) is proposed, under which virtual melting occurs at temperatures much below the melting temperature. Virtual melting is predicted using a developed, advanced thermodynamic approach and confirmed by large-scale molecular dynamics simulations of shockwave propagation and quasi-isentropic compression in both single and defective crystals. The work and energy of nonhydrostatic stresses at the shock front drastically increase the driving force for melting from the uniaxially compressed solid state, reducing the melting temperature by 80% or 4,000 K. After melting, the relaxation of nonhydrostatic stresses leads to an undercooled and unstable liquid, which recrystallizes in picosecond time scales to a hydrostatically loaded crystal. Characteristic parameters for virtual melting are determined from molecular dynamics simulations of Cu shocked/compressed along the $\langle 110 \rangle$ and $\langle 111 \rangle$ directions and Al shocked/compressed along the (110) direction.

high strain-rate plasticity | relaxation of non-hydrostatic stresses | thermodynamics under uniaxial straining

eneration and motion of dislocations, twinning, and crystal-Generation and motion of dissociations, and the main mechanisms of plastic deformation and relaxation of nonhydrostatic stresses that are reflected in the deformation-mechanism maps (1, 2). Recent large-scale nonequilibrium molecular dynamics (NEMD) simulations of shockwave propagation in fcc metallic single crystals have revealed that for wave propagation along the $\langle 110 \rangle$ and $\langle 111 \rangle$ directions, melting occurs at temperatures below the equilibrium melt temperature $T_m(p)$ at the corresponding shock pressure p—e.g., for Cu by 20% in ref. 3 and by 7–8% (\pm 4%) in ref. 4. However, for shockwaves along the $\langle 001 \rangle$ direction, superheating is observed. A possible mechanism for the observed melting below $T_m(p)$ has been attributed to solid-state disordering due to high defect densities. Indeed, a small reduction in the melt temperature due to defects has been reported (5). It has also been argued that anomalous plastic flow rather than bulk melt can explain the experimentally observed low melt temperatures of Ta under pressure (6). Here, we propose a new deformation mechanism, under which melting can occur at temperatures much below $T_m(p)$ in materials subjected to high deviatoric stresses (such as those produced under shock loading and isentropic deformation) and compete with defect nucleation mechanisms, as will be shown here in the case of metals. We have developed a thermomechanical theory of melting under uniaxial deformation that predicts extremely large driving forces. The thermodynamic driving force for melting is due to the energy and work of nonhydrostatic stresses. The theory is quite general and not materialspecific, and it is most applicable at deformation rates sufficiently high, where melting may proceed at rates faster than traditional mechanisms of plasticity. As a proof of concept, we have systematically studied melting under high strain rates in two fcc metals, Cu and Al, employing molecular dynamics (MD) methods that simulate quasi-isentropic and shockwave loading conditions. These simulations confirm for the first time that melting can occur at temperatures much below (by 80% or 4,000 K) $T_m(p)$. After melting, the nonhydrostatic stresses relax, leading to an undercooled and unstable liquid, which for the metals studied here recrystallizes in ps time scales. Because the conditions for melting (nonhydrostatic stresses) disappear upon melting, and because of the short time scale involved, we have termed this new deformation mechanism virtual melting (VM). Because fcc metals can be easily deformed by traditional dislocation and twinning mechanisms, the strain rates at which VM is observed and confirmed via our MD simulations of Cu and Al crystals are necessarily high (approximately $10^9 - 10^{12} \text{ s}^{-1}$) and represent an upper bound in the regime of strain rates to study and confirm our thermodynamic predictions. For materials with suppressed plasticity (such as Si, Ge, high-strength materials, alloys, or complex organic molecules), VM may very well be observed at much lower strain rates. Because the thermodynamic VM formalism is quite general, in what follows we will primarily discuss our results for Cu and Al shocked or quasi-isentropically compressed along the (110) direction and Cu shocked or quasi-isentropically compressed along the (111) direction. Simulation details are provided in SI Appendix. We find that the reduction of the melt temperature correlates with the magnitude of the nonhydrostatic (shear) stress at the shock front, which is consistent with thermodynamics.

Results and Discussion

Development of Thermodynamics of Melting Under Nonhydrostatic Conditions (see details in SI Appendix). Fig. 1A shows, by way of example, the stress σ_1 -uniaxial strain curve for a Cu single crystal loaded in the $\langle 110 \rangle$ direction. The data was obtained from quasiisentropic MD simulations employing a well-tested embedded atom method (EAM) potential for Cu (7). Because in a planar shock, the two lateral strains are zero, the uniaxial strain $\varepsilon = l/l_0 - 1$ is equal to the volumetric strain, where l_0 and lare the initial and the current lengths of the sample, respectively. Melting starts at strain $\varepsilon = \varepsilon^*$ and stress $\sigma_1 = \sigma_1^m$, and it ends at strain $\varepsilon = \varepsilon^m$ and at the same stress σ_1 . Because the transformation proceeds under a variable stress tensor, classical thermodynamic approaches based on thermodynamic potential differences and thermodynamic equilibrium across an interface (8-10) are not applicable. In fact, the reduction of the thermodynamic melt temperature of the stress-free solid, $T_m(0)$, caused by deviatoric stresses was estimated to be just 1 K (8, 10), which is why it was not considered as a possible reason for the reduction in $T_m(0)$ in shockwaves. We here will consider a homogeneous transformation process in a finite volume, expanding on our advanced thermodynamic approach to crystal-crystal transformations (11, 12). Because melting is assumed to be the only

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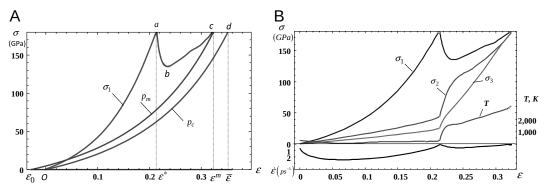


Fig. 1. (A) Stress-uniaxial strain curve of a Cu crystal until melting (σ_1 , the same as in B) and equations of state of molten (ρ_m) and crystalline (ρ_c) phases. Area between curves {Oabcd} and {Od} is the additional driving force for melting due to nonhydrostatic loading. (B) Variation of normal stresses σ_i , temperature, and prescribed strain rate vs. uniaxial strain obtained using MD simulations for $\langle 110 \rangle$ shock loading of Cu.

dissipative process, the thermodynamic driving force per unit unloaded volume for the melting is equal to the total dissipation, as follows

$$X_{m} = \int_{\varepsilon^{*}}^{\varepsilon^{m}} \sigma_{1}(\varepsilon) d\varepsilon + \Delta s (T - T_{m}(0))$$

$$+ (\psi_{c}^{h}(\varepsilon^{*}) - \psi_{m}^{h}(\varepsilon^{m})) + \psi_{c}^{\text{dev}}(\varepsilon^{*}).$$
[1]

Here, ψ^h and ψ^{dev} are the elastic energy of hydrostatic and deviatoric stresses (strains), Δs is the jump in entropy, T is the temperature at which the melting starts at $\varepsilon = \varepsilon^*$, and subscripts c and m refer to crystalline and molten states, respectively. An increase in temperature during the deformation increases X_m and can be taken into account through an effective temperature (11). For simplicity, we neglect it and underestimate X_m . Let $p_c(\varepsilon)$ and $p_m(\varepsilon)$ be the pressure-volumetric strain equation of state of the crystalline and molten phases, respectively (Fig. 1A), with $\varepsilon_c(p)$ and $\varepsilon_m(p)$ being their inverse equations. The equation of state for the melt starts at point ($\varepsilon = \varepsilon_0$; $\sigma_1 = 0$), where $\varepsilon_0 < 0$ is the volumetric transformation strain at melting at pressure p = 0. After some manipulations, the condition $X_m = 0$ provides the equilibrium melt temperature under uniaxial straining T_m^{nh} :

$$T_m^{nh} = T_m(\sigma_1) - \left(\int_0^{\varepsilon^m} \sigma_1 d\varepsilon - \int_0^{\bar{\varepsilon}} p_c d\varepsilon + \sigma_1(\bar{\varepsilon} - \varepsilon^m) \right) / \Delta s,$$
[2]

where $\bar{\epsilon} = \varepsilon_c(\sigma_1^m)$ and $\varepsilon^m = \varepsilon_m(\sigma_1^m)$ (Fig. 1A). Geometrically, the magnitude of the negative mechanical part of the driving force for melting under hydrostatic conditions is equal to the area $\{O\varepsilon_0cd\}$ between the equation of state for melt and crystal. This area characterizes the increase in the melt temperature under hydrostatic

loading from $T_m(0) = 1,357$ K to 5,087 K (at 179.1 GPa) (see Table S1 in SI Appendix). The difference between the areas under the stress-strain curve $\sigma_1(\varepsilon)$ {Oabcd} and {Od}, which represents the equation of state $p_c(\varepsilon)$ for the single crystal, provides an additional driving force for melting due to non-hydrostatic loading (the term in parentheses in Eq. 2). This difference is almost three times larger than the area {O $\varepsilon_0 cd$ } and produces close to a threefold reduction in $T_m(p)$ when compared with the increase associated with the hydrostatic pressure. All parameters for Cu and Al melting obtained from NEMD simulations of shock loading along the $\langle 110 \rangle$ direction and for Cu along the $\langle 111 \rangle$ direction can be found in Table S1 in SI Appendix.

In Fig. 1B, melting commences at a peak stress $\sigma^m =$ 179.1 GPa, corresponding to $\varepsilon^* = 0.214$, and ends at the same stress $\sigma_1 = 179.1$ GPa but at $\varepsilon^m = 0.324$. For this case, the reduction in the thermodynamic melt temperature calculated with the help of Eq. 2 is enormous (Fig. 2A): $\Delta T_m = 10074\,$ K—i.e., from $T_m(179.1) = 5,087 \text{ K to } T_m^{nh} = -5,013 \text{ K. Reductions of similar}$ orders of magnitude were found for Al, and for other loading directions (Fig. 2B and Table S1 in SI Appendix). It should be noted that a negative T_m^{nh} does not imply the existence of a negative temperature: At this temperature the calculated thermodynamic driving force for melting due to nonhydrostatic stresses is zero. In order for melting to proceed within ps time scales, some overheating—which cannot be estimated using thermodynamics—is necessary. Furthermore, MD simulations (see below) have shown that melting can occur at least at $0.2T_m(p)$. The calculated reduction in the melt temperature ΔT_m vs. strain ε^* and corresponding stress $\sigma_1(\varepsilon^*)$ can be approximated by $\Delta T_m = A_{\sigma} + B_{\sigma}\sigma_1 + C_{\sigma}\sigma_1^2 =$ $A_{\varepsilon} + B_{\varepsilon} \varepsilon^* + C_{\varepsilon} \varepsilon^{*2}$. All constants for Cu and Al are listed in Table S3 in SI Appendix.

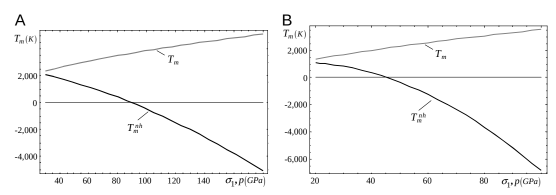


Fig. 2. Calculated melting temperature under hydrostatic T_m and nonhydrostatic T_m^{nh} conditions vs. pressure (uniaxial load) for shock loading of Cu (A) and Al (B) along the $\langle 110 \rangle$ direction.

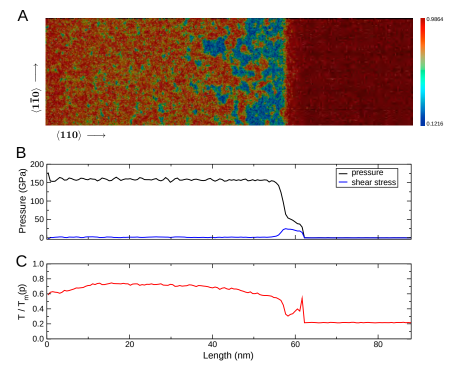


Fig. 3. (A) Atomic configuration of Cu slab shocked to a pressure of 160 GPa (red atoms are solid, blue are liquid); (B and C) pressure and temperature profiles, respectively, along the shock direction. The temperature is normalized by the equilibrium melt temperature $T_m(p)$ at the corresponding shock pressure along the profile.

MD Simulations. Fig. 3A shows a thin slice (about 20-Å thick) of the atomic configuration from an NEMD simulation of Cu shocked along the (110) direction to a pressure of 160 GPa (see also Movie S1). Liquid and solid atoms were identified with a local order parameter q_6 , which measures the symmetry correlation of an atom with its neighbors (13, 14). Solid atoms (red) are well correlated with their neighbors, while liquid atoms (blue) are not. The material at the shock front melts at a temperature $T \simeq 0.3 T_m(p)$ at the corresponding pressure along the profile (Fig. 3 A-C) and recrystallizes behind the shock front at $T \simeq 0.8 T_m(p)$ and at a rate that increases with increase in $T_m(p) - T$. This is consistent with the behavior of an over-cooled liquid rather than amorphous solid (see SI Appendix). No evidence of dislocation or extended defects was found.

Fig. 4A shows the shear stress $\tau = 1/2(\sigma_1 - 1/2(\sigma_2 + \sigma_3))$ as a function of strain rate for Cu compressed quasi-isentropically. The shear stress at $\dot{\varepsilon} = 0$ corresponds to the elastic limit, which is characteristically high for defect-free crystals. The elastic limit can be lowered by introducing pre-existing dislocations (PED) (15). Data shown are for samples with defect densities approximately 10^{12} cm⁻². Fig. 4B clearly shows a strong reduction in the melt temperature (even by 4,000 K) for quasi-isentropic compression with increasing shear stress. The final strain was fixed at 0.34, while the strain rate was varied and the temperature was controlled to a value that would yield the same maximum fraction of liquid atoms (25%) for all simulations. In shock compression, where the temperatures are higher, the fraction of pre-melted atoms behind the shock is significantly higher than 25% (Fig. 3A).

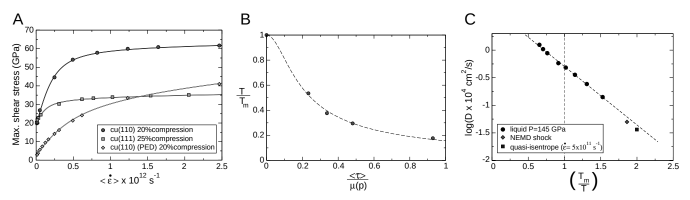


Fig. 4. (A) Maximum shear stress vs. average strain rate in Cu defect-free and with pre-existing defects (PED), quasi-isentropically and uniaxially compressed to 0.20 and 0.25 along the (110) and (111) directions. The PED crystal is nominally oriented along the (110) direction. (B) Melting temperature (normalized by $T_m(160) = 4816$ K) vs. average shear stress (normalized by the shear modulus at p = 160 GPa) for defective Cu quasi-isentropically compressed to a final strain of 0.34 at different strain rates. (C) Diffusion coefficient of liquid Cu at a pressure of 145 GPa (circles) evaluated from the atomic mean-squared displacements. Also shown is the diffusion coefficient associated with a region behind the shock front for Cu shocked along (111) to a pressure of 145 GPa (diamond) as well as Cu isentropically compressed along $\langle 111 \rangle$ to the same pressure at a strain rate of $5 \times 10^{11} \text{ s}^{-1}$ (square). The dashed line is a fit to the liquid data, extrapolated to lower temperatures.

Solid-state disordering has been proposed as the main mechanism of pre-melting in shock along the $\langle 110 \rangle$ and $\langle 111 \rangle$ orientations in Cu single crystals (4). In order to distinguish between a hot, amorphous solid or a supercooled liquid, the self-diffusion coefficient of the liquid region behind the shock front was computed from the mean-squared displacement. Fig. 4C shows the diffusion coefficient as a function of temperature at a pressure of 145 GPa for the liquid state of the EAM model Cu used in this work. Also shown is the diffusion coefficient of a region behind the shock front in a Cu slab shocked along the (111) direction to the same pressure. Because all points belong to the same Arrhenius line, these results confirmed that the region behind the shock front is a supercooled liquid rather than an amorphous solid. Note that VM was observed for isothermal (111) loading of Cu even at 300 K (Table S1 in SI Appendix)-i.e., at $T \simeq 0.055 T_m(p)$. However, recrystallization is so fast that the diffusion coefficient could not be determined.

Conclusions

At high strain rates ($\dot{\varepsilon} \sim 10^9 - 10^{12} \text{ s}^{-1}$ in metals) and high shear stresses, VM can compete with traditional defect nucleation mechanisms and therefore should be incorporated in the deformation-mechanism maps (1, 2). This prediction is based on a new thermomechanical theory of melting under uniaxial straining that is quite general and that can be applied to any material subjected to strain rates sufficiently large in order to allow melting to proceed faster than traditional mechanisms of plasticity. Virtual melting has been confirmed for the first time by MD simulations for the most critical case of fcc metals. For materials with suppressed plasticity, VM may be observed at lower strain rates and stresses than the ones sampled here [e.g., in a recent MD study (16) deformation of organic α-HMX crystal is shown to occur through generation and motion of dislocations for shock strengths up to 14 GPa and via formation of amorphous shear nanobands at higher pressure, which may be related to VM]. Our thermodynamic approach can be extended for amorphization and

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sublimation as well as for arbitrary 3D loading, in particular for pressure and shear experiments. The concept of the VM was introduced in refs. (17-20) as an intermediate step in crystal-crystal, crystal-amorphous, and crystal-gas transformations. Here, however, VM was for the first time directly observed in MD simulations of plastic straining. The driving force for VM in phase transformations in refs. (17–20) is due to the relaxation of the deviatoric stresses and also disappears upon melting, as in the case of VM here. This suggests that our results here indirectly support the plausibility of VM in phase transformation as well. Finally, while the sampled compression rates in our simulations of Cu and Al were necessarily high, they are close to what can be achieved in laser-driven shock experiments (21). However, due to the short time scales additional efforts are required for experimental verification of VM, particularly in metals. Such regimes are also relevant for nuclear explosion and meteorite impact as well as for planned experiments in large laser facilities such as the National Ignition Facilities at the Lawrence Livermore National Laboratory in the United States and the Laboratoire pour l'Utilisation des Lasers Intenses in France.

Materials and Methods

We have performed multimillion atom NEMD simulations of shockwave compression as well as smaller, Hugoniostat-method (22) and MD simulations of quasi-isentropic compression (15). The NEMD simulations comprised up to 12 million atoms arranged in a rectangular slab with periodic boundary conditions in the transverse directions. The procedure used to initiate a shockwave of a given strength is detailed in ref. 23. The systems studied were Cu and Al single crystals as well as Cu crystals with pre-existing defects. A detailed description of the MD simulations and methods is available in SI Appendix.

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Supporting Information

Virtual Melting: A New Mechanism of Stress Relaxation Under High Strain Rate Loading

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1. Methods of Advanced Thermodynamic Calculations

Since melting is considered the only dissipative process, the thermodynamic driving force per unit unloaded volume for the melting for isothermal loading is equal to the total dissipation [1, 2]:

$$X_m = W + \Delta s(T - T_m(0)) + (\psi_c^h(\varepsilon^*) - \psi_m^h(\varepsilon^m)) + \psi_c^{dev}(\varepsilon^*). \tag{1}$$

Here, $W = \int_{\varepsilon^*}^{\varepsilon^m} \sigma_1(\varepsilon) d\varepsilon$ is the transformation work, which takes into account the actual stress-strain curve $\sigma_1(\varepsilon)$ during the melting; ψ^h and ψ^{dev} are the elastic energy of hydrostatic and deviatoric stresses (strains); Δs is the jump in entropy; T is the temperature at which the melting is started (at $\varepsilon = \varepsilon^*$); $T_m(0)$ is the thermodynamic melting temperature for stress-free solid; and subscripts c and m are for crystalline and molten states, respectively.

Let $p_c(\varepsilon)$ and $p_m(\varepsilon)$ be the equations of state of the crystalline and molten phases (Fig. 1). The equation of state for melt starts at point ($\varepsilon = \varepsilon_0$ and $\sigma_1 = 0$), where $\varepsilon_0 < 0$ is the volumetric transformation strain at melting at pressure p = 0. Then,

$$\psi_c^h(\varepsilon^*) = \int_0^{\varepsilon^*} p_c d\varepsilon; \quad \psi_c^{dev}(\varepsilon^*) + \psi_c^h(\varepsilon^*) = \int_0^{\varepsilon^*} \sigma_1 d\varepsilon; \quad \psi_m^h(\varepsilon^m) = \int_{\varepsilon_0}^{\varepsilon_m} p_m d\varepsilon, \tag{2}$$

and

$$X_m = \int_0^{\varepsilon^m} \sigma_1 d\varepsilon - \int_{\varepsilon_0}^{\varepsilon_m} p_m d\varepsilon + \Delta s (T - T_m(0)). \tag{3}$$

Geometrically, the mechanical driving force for melting is equal to the difference between the areas under the stress-strain curve for $\sigma_1(\varepsilon)$ { $Oabc\varepsilon^m$ } and for $p_m(\varepsilon)$ { $\varepsilon_0c\varepsilon^m$ } (Fig. 1). For melting under constant temperature and hydrostatic pressure $p = \sigma_1$, the condition for thermodynamic equilibrium is

$$X^{h} = \sigma_{1}(\varepsilon_{m} - \bar{\varepsilon}) + \Delta s(T_{m}(\sigma_{1}) - T_{m}(0)) + (\psi_{c}^{h}(\bar{\varepsilon}) - \psi_{m}^{h}(\varepsilon_{m})) = 0, \tag{4}$$

where $T_m(\sigma_1)$ is the equilibrium melting temperature under hydrostatic pressure $p = \sigma_1$, $\bar{\varepsilon} = \varepsilon_c(\sigma_1)$, $\varepsilon_c(p)$ and $\varepsilon_m(p)$ are the inverse equations of state for crystal and melt. The magnitude of the negative mechanical part of X_h is $\{O\varepsilon_0cd\}$ (Fig. 1). It follows from Eq. (4):

$$T_m(\sigma_1) = T_m(0) + \left(\sigma_1(\bar{\varepsilon} - \varepsilon_m) - (\psi_c^h(\bar{\varepsilon}) - \psi_m^h(\varepsilon_m))\right) / \Delta s. \tag{5}$$

Expressing $\Delta s T_m(0)$ from Eq. (4) and substituting it in Eq. (1), we obtain

$$X_m = W + \sigma_1(\bar{\varepsilon} - \varepsilon_m) + \Delta s(T - T_m(\sigma_1)) + (\psi_c^h(\varepsilon^*) - \psi_c^h(\bar{\varepsilon})) + \psi_c^{dev}(\varepsilon^*).$$
 (6)

Then, the condition $X_m = 0$ results in

$$T_m^{nh} = T_m(\sigma_1) - \left(\psi_c^{dev}(\varepsilon^*) + W + \sigma_1(\bar{\varepsilon} - \varepsilon^m) - (\psi_c^h(\bar{\varepsilon}) - \psi_c^h(\varepsilon^*))\right) / \Delta s =$$

$$T_m(\sigma_1) - \left(\int_0^{\varepsilon^m} \sigma_1 d\varepsilon - \int_0^{\bar{\varepsilon}} p_c d\varepsilon + \sigma_1(\bar{\varepsilon} - \varepsilon^m)\right) / \Delta s. \tag{7}$$

Geometrically, additional driving force for melting due to nonhydrostatic loading (the term in parentheses in Eq. (7)) is equal to the difference between the areas under the stress-strain curve $\sigma_1(\varepsilon)$ {Oabcd} and the equation of state $p_c(\varepsilon)$ for crystal {Od}. All parameters for melting of Cu and Al for loading in the <110> direction and for Cu loading in the <111> direction are given in Table 1S.

Table 1S. Parameters for crystal loading (ε^* , ε^m , $\overline{\varepsilon}$, and σ_1^m , see Figs. 1 and 4S), maximum strain rate $\dot{\varepsilon}_{\max}$, initial T^* and final T_f temperatures, as well as melting temperature under hydrostatic conditions $T_m(\sigma_1^m)$ and decrease in melting temperature due to nonhydrostatic loading ΔT_m .

	£*	$\boldsymbol{\mathcal{E}}^m$	$\overline{\mathcal{E}}$	$\dot{\mathcal{E}}_{ ext{max}}$	$\sigma_{\scriptscriptstyle 1}^{\scriptscriptstyle \mathrm{m}}$	<i>T</i> *	T_f	$T_{\scriptscriptstyle m}(\sigma_{\scriptscriptstyle 1}^{\scriptscriptstyle m m})$	ΔT_{m}
				ps^{-1}	GPa	K	K	K	K
Cu <110>	0.214	0.324	0.351	2.48	179.1	325	3084	5087	10074
Cu <111>	0.297	0.360	0.369	4.3	204.9	300	300	5434	9197
Cu <111>	0.301	0.353	0.375	3.97	215.1	197	2604	5565	8022
Al <110>	0.227	0.356	0.386	3.87	98.17	305	2711	3544	9945

Table 2S. Parameters in Murnaghan's (8) and Simon's (11) equations in Supplementary Materials.

	p_0	b	T_0	α	β
	GPa		K	GPa	
Cu	39.096	3.98	1327	15.3691	0.529458
Al	33.576	2.81	324	2.56529	0.647822

Table 3S. Parameters in Eq.(14) in Supplementary Materials for the reduction in melting temperature due to nonhydrostatic loading.

	T_f	A_{σ}	B_{σ}	C_{σ}	$A_{arepsilon}$	$B_{arepsilon}$	$C_{arepsilon}$
	K	K	K/GPa	K/GPa ²	K	K	K
Cu <110>	3084	-1128	44.67	0.1008	4174	-82665	517956
Cu <111>	300	-1845	64.52	0.0518	-2182	20997	59602
Cu <111>	2604	-1137	42.24	0.0020	-299	-2684	101753
Al <110>	2711	-441	15.26	0.9295	5162	-96069	493689

The Murnaghan's equation of state for crystal, $p_c(\varepsilon)$, its inverse form, and the corresponding expressions for ψ_c^h and ψ_c^{dev} are:

$$p_c = p_0((1-\varepsilon)^{-b} - 1); \quad \varepsilon = 1 - \left(1 + \frac{p_c}{p_0}\right)^{-1/b},$$
 (8)

$$\psi_c^h = \int_0^{\varepsilon} p d\varepsilon = \frac{p_0}{b-1} \left((1-\varepsilon)^{b-1} - \varepsilon(b-1) - 1 \right), \quad \psi_c^{dev} = \int_0^{\varepsilon^*} \sigma_1 d\varepsilon - \psi_c^h(\varepsilon^*), \tag{9}$$

where parameters p_0 and b for Cu and Al are given in Table 2S.

The jump in entropy Δs versus pressure p along the melting line $T_m(p)$ is found using the Clausius-Clapeyron relation

$$\Delta s = -\varepsilon_v / (dT_m(p)/dp), \tag{10}$$

where ε_v is the volumetric strain during the melting under the pressure (equal to ε_0 for p=0). Since we consider cubic crystals with isotropic thermal expansion, the effect of deviatoric stresses on entropy will be neglected. The melting temperature versus pressure is found according to the Simon equation:

$$T_m = T_0 (1 + p/\alpha)^{\beta}, \qquad (11)$$

with parameters T_0 , α , and β given in Table 2S for Cu and Al. The volume change during the melting $\varepsilon_v < 0$ along the melting line was calculated using MD simulations at constant temperature and pressure and approximated by the functions

$$|\varepsilon_v| = 0.008915 + 0.95820/p - 4.4612/p^2$$
 (12)

for Al and

$$|\varepsilon_v| = 0.013712 + 1.14436/p - 19.3125/p^2 + 105.80245/p^3$$
 (13)

for Cu. Substitution of Eqs. (11), (12), or (13) in Eq. (10) gives us jump in entropy. Functions $|\varepsilon_v|$ and Δs vs. pressure are shown in Figs. 1S and 2S for Cu and Al, respectively. While the

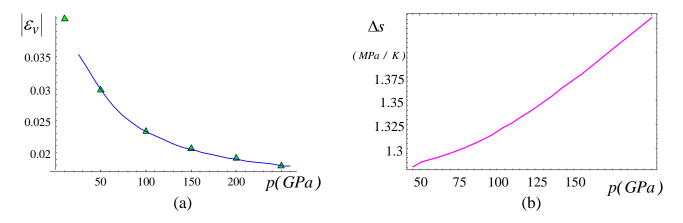


Figure 1S. (a) Jump in volumetric strain vs. pressure calculated using MD constant pressure and temperature simulations for Cu. (b) Jump in entropy during melting vs. pressure calculated using Clausius-Clapeyron equation (10) for Cu.

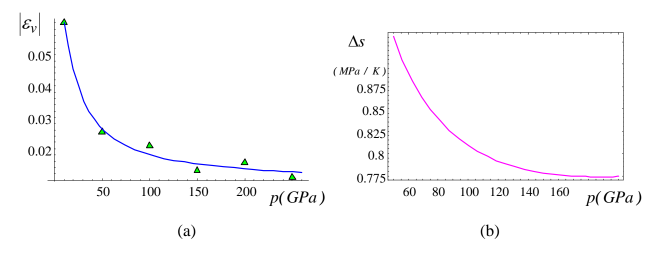


Figure 2S. (a) Jump in volumetric strain vs. pressure calculated using MD constant pressure and temperature simulations for Al. (b) Jump in entropy during melting vs. pressure calculated using Clausius-Clapeyron equation (10) for Al.

pressure dependence of Δs is very weak, it is interesting to note that it has different slopes for Cu and Al.

To calculate the melting temperature decrease for lower strains ε^* and corresponding stress $\sigma_1(\varepsilon^*)$, one needs to know the stress-strain curve $\sigma_1(\varepsilon)$ for elastic uniaxial loading (we will take it from Fig. 1, but it can be found phenomenologically if pressure dependence of elastic constants is known), the equation of state for solid $p(\varepsilon_c)$, and the stress strain curve $\sigma_1(\varepsilon)$ during melting, which has to be determined by MD simulation for each ε^* separately. To avoid additional MD simulations, we take into account that the reduction in σ_1 during the melting (line abc in Fig. 1) results in multiplication of work under constant stress $\sigma_1(\bar{\varepsilon}-\varepsilon^*)$ by a factor h=0.8831 for Al (or h=0.8847 for Cu) for <110> loading, which we used for all smaller ε^* . For <111> loading of Cu, we obtain h=0.9504 for $T_f=300K$ and h=0.8311 for $T_f=2604K$. The calculated melting temperature decrease (Eq. (7)) for lower strains ε^* and corresponding stress $\sigma_1(\varepsilon^*)$ can be approximated in the form

$$\Delta T_m = T_m(\sigma_1) - T_m^{nh}(\sigma_1) = A_\sigma + B_\sigma \sigma_1 + C_\sigma \sigma_1^2 = A_\varepsilon + B_\varepsilon \varepsilon^* + C_\varepsilon \varepsilon^{*2}, \tag{14}$$

where all constants are listed in Table 3S.

2. Methods of Molecular Dynamics Simulations

We performed multi-million atom non-equilibrium molecular dynamics (NEMD) simulations of shock wave compression, as well as smaller, Hugoniostat, and molecular dynamics (MD) simulations of quasi-isentropic compression to explored the dynamical evolution of the shocked state as a function of shock compression and crystallographic direction. The NEMD simulations comprised up to 12 million atoms arranged in a rectangular slab with periodic boundary conditions in the transverse directions. We employed the high-performance parallel molecular dynamics code

SPaSM [3] for the multi-million NEMD simulations. The procedure used to initiate a shock wave of a given strength is detailed in Ref. [4]: the crystal impacts an infinitely massive piston with a velocity $-u_p$, which produces a shock wave that propagates away from the piston with velocity $u_s - u_p$, u_s being the shock velocity. The NEMD sampled time scales were nominally of the order of 20-40 ps.

These simulations were augmented by constant-stress Hugoniostat simulations comprising 10⁵ atoms arranged approximately in a cube. Details of the method can be found in Ref. [5]. We also carried out quasi-isentropic compression simulations, in which the system was uniaxially and homogeneously compressed to a final uniaxial strain using a specified time-profile for the strain-rate. Details of this method are given in [6].

The systems we chose to study were copper and aluminum single crystals shocked or quasiisentropically compressed along the $\langle 110 \rangle$ direction and aluminum single crystal along the $\langle 111 \rangle$ direction. These crystallographic directions are known to produce large shear stresses with strain. The atomic interactions in the Cu and Al simulations were described using the welltested embedded-atom method (EAM) models for Cu [7] and Al [8].

3. Results of additional MD simulations

1. The role of high shear stresses on the lowering of the melt temperature can be studied in greater detail via quasi-isentropic compression simulations, where the final compressive strain can be fixed, while the uniaxial strain-rate is varied. Using these types of simulations, we carried out a systematic study of virtual melting in both single and defective Cu crystals. We employed approximately cubic samples with 1.2×10^5 to 1.5×10^6 atoms and periodic boundary conditions. The final strain was fixed, and the strain rate was varied over 3 decades (from 2.5×10^9 to 2.5×10^{12} s⁻¹). As shown in Fig. 4 for Cu, along <110>, the shear stress increases rapidly with

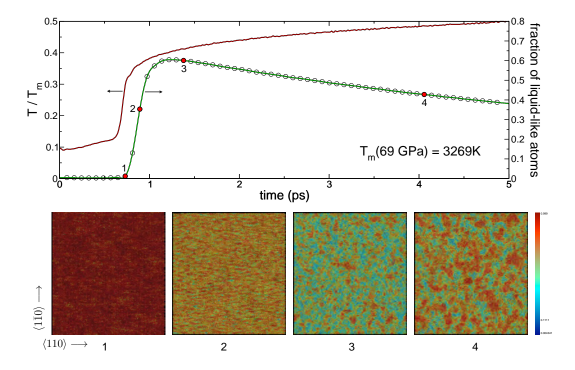
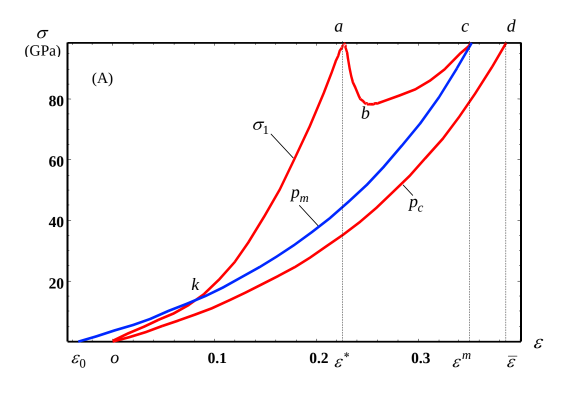


Figure 3S. 1.35-million-atom simulation of Cu quasi-isentropically compressed along <110> at an average strain rate of 2.5×10^{11} s⁻¹ to a final strain of 0.20. The temperature profile (upper line) has been normalized by the equilibrium melt temperature at the final pressure (69 GPa). The fraction of liquid-like atoms (lower line) was computed from the order parameter q_6 . The atomic profiles numbered at the bottom corresponds to the times numbered along the path. Atoms are colored according to q_6 : solid (red) and liquid (blue).

compression. Under quasi-isentropic compression and at sufficiently high strain rates, we find that the system can melt at $T \simeq 0.35 T_m(p)$ (Fig. 3S). Figure 3S also shows a sequence of snapshots of the atomic configurations in a 1.35-million-atom single crystal Cu simulation in which the system was quasi-isentropically compressed along the <110> orientation at an average strain rate of $2.5 \times 10^{11} \ \mathrm{s^{-1}}$ to a final strain of 0.20. Also shown is the temperature (red line) normalized by the equilibrium melt temperature at the equilibrium pressure (69 GPa). The fraction of liquid atoms (green line) as a function of time was computed by following the evolution of the local order parameter q_6 and evaluating the fraction of atoms with a value less than 0.6. It takes about 0.8 ps to compress the sample to strain of 0.20 at $\dot{\varepsilon} = 0.25 \text{ ps}^{-1}$. At the point labeled 1 in Fig. 3S, the atoms are still in an elastic and uniaxially compressed state, with a very high elastic energy corresponding to the highest shear stress (45 GPa) along the path. From point 1 to point 3, the system transforms from uniaxially compressed elastic solid to an isotropic state due to virtual melting. The virtual melting is partial at this strain and strain rate, and the liquid regions (blue atoms) coarsened rapidly. At the point labeled 3, the material is more isotropic, the elastic energy has been reduced significantly, and the temperature has increased due to the enthalpy difference between the isotropic liquid and the uniaxially compressed elastic solid. The time scale for this conversion is less than 1 ps. The decrease of the fraction of liquid atoms with time is caused by re-solidification from the melt (red atoms, point 4 along the path) followed by further reduction in the shear stress and an increase in temperature. From the enthalpy differences between the uniaxially compressed solid, supercooled liquid, and isotropic solid, the temperature profile as a function of time can be evaluated with excellent agreement with the simulations.

2. In addition, crystallization kinetics unambiguously suggests that the disordering represents the virtual melt rather than amorphization. Increase in temperature increases thermal fluctuations and promotes kinetics of crystallization from both melt and glass. However, the driving force for crystallization of the melt and glass is the increasing function of $T_m - T$ and $T - T_g > 0$, respectively (where T_g is the glass transition temperature); thus temperature has opposite effect on these driving forces. We assume that T_g is below temperature at the end of our calculations, otherwise, there is no sense to talk about the crystallization of glass. For glass, the driving force for crystallization increases with temperature growth and crystallization rate has to be faster with increasing temperature. For melt, the driving force reduces with temperature increase and (if thermodynamic effect exceeds the kinetic one) crystallization rate may decrease with growing temperature. Since in all our MD simulations the crystallization rate decreased with the growing temperature, we can conclude that the disordered state is the virtual melt rather than glass.

- 3. For strain rate smaller than some threshold, melting occurs heterogeneously rather than in the entire volume (Fig. 3S). The volume fraction of disordered regions decreases with the decrease in strain rate. Note that if the remaining crystals are surrounded by melt and are under hydrostatic conditions, the total energy will be smaller than for complete melting, and the driving force will be even larger. This is because the energy of hydrostatically loaded crystal is much smaller than the energy of the melt below melting temperature. Crystallization time drastically depends on the existence of residual crystalline clusters, which can serve as nucleation cites. The higher are the shock strength (or strain rate) and temperature, the less probability there is of finding residual crystalline clusters in melt. Much more time is required for homogeneous nucleation of the crystal phase.
- 4. Fig. 4S shows the stresses and temperature evolution, and prescribed strain rate vs. uniaxial strain obtained using MD simulations for <110> shock loading of Al, similar to Fig. 1 for Cu. A well-tested EAM potential for Cu [8] was used.



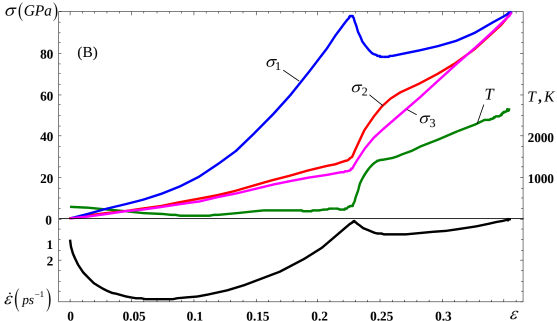


Figure 4S. (A) Stress-uniaxial strain curve of an Al crystal until melting (σ_1 , the same as in Fig. 4S, B) and equations of state of molten (p_m) and crystalline (p_c) phases. Area between curves {Oabcd} and {Od} is the additional driving force for melting due to nonhydrostatic loading. (B) Variation of normal stresses σ_i , temperature, and prescribed strain rate vs. uniaxial strain obtained using MD simulations for <110> shock loading of Al.

5. Supporting movie

Cu single crystal compressed quasi-isentropically along the <111> direction (out of plane) at a strain rate of $3 \times 10^{11} s^{-1}$ to a final uniaxial strain of 33% (σ_1 =174 GPa) is shown. Axis x is along the <1 $\bar{1}$ 0> direction and axis y is along the <11 $\bar{2}$ > direction. The cross section of atomic configuration in the left panel is 10 nm × 10 nm. Only atoms within a 1.7 nm thick slice are shown for clarity. Atoms are colored according to q_6 [9, 10] (red atoms are solid, green atoms are liquid). The right panel shows the time evolution of the temperature (normalized by the equilibrium melt temperature at the corresponding pressure), pressure (p) and shear stress (τ).

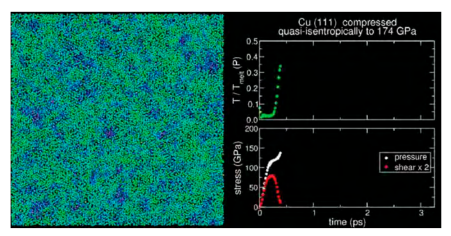
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Supporting Information

Levitas and Ravelo 10.1073/pnas.1203285109



Movie S1. Cu single crystal compressed quasi-isentropically along the $\langle 111 \rangle$ direction (out of plane) at a strain rate of 3×10^{11} s⁻¹ to a final uniaxial strain of 33% ($\sigma_1 = 174$ GPa) is shown. Axis x is along the $\langle 110 \rangle$ direction and axis y is along the $\langle 112 \rangle$ direction. The cross section of atomic configuration in the left panel is 10 nm \times 10 nm. Only atoms within a 1.7 nm thick slice are shown for clarity. Atoms are colored according to q_6 (1, 2) (red atoms are solid, green atoms are liquid). The right panel shows the time evolution of the temperature (normalized by the equilibrium melt temperature at the corresponding pressure), pressure (ρ) and shear stress (τ).

1 Ten Wolde PR, Ruiz-Montero MJ, Frankel D (1995) Numerical evidence for bcc ordering at the surface of a critical fcc nucleus. Phys Rev Lett 75:2714–2717.

² Steinhardt PJ, Nelson DR, Ronchetti M (1983) Bond-orientational order in liquids and glasses. *Phys Rev B Condens Matter Mater Phys* 28:784–805. Movie S1 (MOV)

tissue on implantation (Fig. 1b), making the composition and architecture of the vessels more physiological. In this respect, it would be interesting to see if, on seeding the engineered matrices of Miller and coworkers with smooth muscle cells, the cells would, on perfusion, migrate towards the endothelialized channels *in vitro*, just as they do *in vivo*¹⁰.

Still, even if complex thick tissues can eventually be built *in vitro*, it is unlikely that faithful copies of real organs will ever be fabricated. Yet, do we really need to reproduce physiological organs with minute details? For clinical purposes it will probably be sufficient to make

autologous tissue that functions just as well, if not better, than the original tissue to be replaced. Perhaps in the not-so-distant future a patient could just walk into a specialized facility, shed some cells, and have a functionally correct tissue or organ built from scratch. As the work of Miller and colleagues hints at the possibility that such engineered tissues could be maintained *in vitro*, the notion of on-demand off-the-shelf organs does not seem like science fiction anymore.

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SHOCK RELIEF

Like us, materials respond to stress in many ways. Like us, some materials simply break. Metals may relieve stress by spawning dislocations or other defects that permit plastic flow. Other solids, such as ice and silica, can undergo amorphization when compressed, or may be transformed to different crystal phases. It now seems that melting is another option for crystals seeking to relax under stress.

For example, simulations of copper subjected to shock waves — that is, to non-hydrostatic stress — suggest that, along some crystal axes, melting can occur a little below the equilibrium melting temperature $T_{\rm m}$ (ref. 1). It's not been clear how this happens. Is the liquid state produced by massive accumulation of stressinduced defects? Can we be sure that this apparent melting is not something that looks superficially like it, such as a special kind of plastic flow? Levitas and Ravelo have now explained how melting can indeed happen at very high strain rates in response to shockwave propagation in crystals, even at temperatures far (by as much as 80 per cent, or 4,000 K) below $T_{\rm m}$ (ref. 2).

The relationships between defect formation, disordering, amorphization and melting have long been a topic of inquiry and controversy, and it is fair to say that they are still not fully understood. True melting, as opposed to solid-state disordering,

requires a thermodynamic driving force. Levitas and co-workers have previously argued that a solid—solid phase transition can induce transient local melting at the moving interface between the phases, more than 100 K below $T_{\rm m}$, to relieve stresses there and allow the elastic energy to relax³; they have very recently reported experimental evidence of such effects in a crystalline nanofibre⁴. Such transient melting might also facilitate high-pressure amorphization as much as 1,000 K below $T_{\rm m}$ (ref. 5).

The curious thing about this kind of melting is that, once it has happened, it negates its own driving force, so that recrystallization happens almost at once. For this reason, the researchers called the phenomenon virtual melting.

Levitas and Ravelo now propose that virtual melting can also account for the response of metals like copper to high non-hydrostatic stresses like those induced by shockwaves. Their simulations of copper subjected to a shock pressure of around 160 GPa reveal a molten state just in front of the wave, with a corresponding large drop in nonhydrostatic stress there, even though the local temperature can be just a fifth of $T_{\rm m}$. The metal recrystallizes in a matter of picoseconds, but a genuine (supercooled) liquid state can still be distinguished from a hot



PHILIP BALL

amorphous one by calculating the atomic self-diffusion rates.

Spotting this phenomenon experimentally will be challenging, given the short timescales. But ultrafast electron diffraction has already been used to study comparably ephemeral effects in surface phase transitions⁶, so it's not a vain hope. And the conditions considered by Levitas and Ravelo are by no means unrealistic, being relevant for example to meteorite impacts, nuclear explosions and the sort of laser-induced compression explored in nuclear fusion.

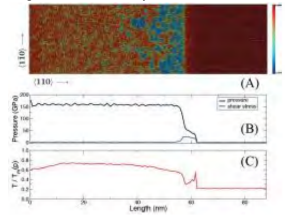
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Crystals take a chill pill: A thermomechanical theory of lowtemperature melting

August 21st, 2012 in Physics / Condensed Matter



Enlarge

(A) Atomic configuration of Cu slab shocked to a pressure of 160 GPa (red atoms are solid, blue are liquid); (B and C) pressure and temperature profiles, respectively, along the shock direction. The temperature is normalized by the equilibrium melt temperature Tmopp at the corresponding shock pressure along the profile. Copyright © PNAS, doi:10.1073/pnas.1203285109

(Phys.org) -- Virtual melting is a phase transition phenomenon associated with solid-solid phase transformation and relaxation of nonhydrostatic stresses and other effects in HMX explosives, as well as with crystal-crystal and crystal-amorphous phase transformation under high pressure. Since its initial formulation by Iowa State University Professor Valery I. Levitas and coworkers, virtual melting has been estimated to occur at 100 - 1000 °K below the material's equilibrium melt temperature. Recently, however, Levitas and University of Texas Associate Professor Ramon Ravelo proposed a new deformation mechanism on which melting can occur at temperatures 4000 °K lower than the equilibrium melt temperature in materials subjected to high deviatoric stresses (where stress components vary by direction, and which control the degree of body distortion) in a shock wave. In addition, they've developed a novel thermomechanical theory of melting that predicts extremely large reduction in melting temperature.

Levitas and Ravelo began their investigation when Levitas gave a talk on virtual melting in 2005 at Los Alamos National Laboratory, and there met Ravelo. "Ramon was very critical and asked a lot of questions," Levitas recounts to Phys.org. "He noted that in atomistic simulations of shock wave propagation in defect-free crystals, it had been observed that disordering occurred at the shock front at temperatures much below the melt temperature at the corresponding shock pressure along some directions – but along others, melting occurred at or above the melting temperature."

Moreover, Levitas adds, it was common wisdom at the time that amorphization occurs due to the high stresses generated at the shock front. "This led us to ask, is the pre-melting observed in atomistic simulations due to mechanical instabilities, such as strain-induced amorphization, which are not related to melting? Or it is indeed virtual melting? Which loading parameters control the melting? What is the lowest temperature at which melting can occur? These were tough questions," Levitas

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acknowledges, "and we then started our collaboration, which after seven years has resulted in the current paper."

Levitas relates the main challenges they faced. "I needed to develop a thermodynamic theory of melting under uniaxial straining typical for shock waves." At the time, known theories for melting under nonhydrostatic conditions – that is, when loading is different in different directions – considered thermodynamic equilibrium to be between melt and nonhydrostatically stressed solid. Reduction in melting temperature due to nonhydrostatic stresses was estimated to be just 1 °K.

"This definitely did not sound promising," Levitas continues. "In contrast, melting in our case represents a deformation process and thermodynamic theory for processes was not developed. I did my best to develop a general formal theory and apply it to the specific processes, which Ramon observed in molecular dynamics simulations. "Since Ramon and I have different backgrounds and consider phenomena from completely different positions, one of the most challenging problems was to understand each other and to make our concepts consistent. A few times I misunderstood how Ramon verbally describes results of atomistic simulations, and developed a theory for the wrong scenarios. For example," Levitas illustrates, "one time I used wrong the video player to play his movie with atomistic simulations and mistakenly observed that melting occurs at the surface of the sample only. I found this very exciting and developed a theory for this scenario, since it sounds reasonable that if uniaxially loaded sample melts along all the surfaces, it would become hydrostatically loaded and further melting would be impossible."

Their next step, Levitas explains, was determining material parameters for a developed theory and application it to the specific melting process: While thermodynamic theory is quite universal and is applicable to any material, for some materials and loading conditions reduction in melting temperature can be small and for others very large. "This study was done through our synergistic collaboration: Ramon gave me parameters determined from his atomistic simulations for loading of copper and aluminum in several directions. "I then made thermodynamic predictions of the reduction in melting temperature for these specific materials."

There were also significant challenges in confirming virtual melting by large-scale molecular dynamics simulations. "Thermodynamic theory predicted how much melting temperature can be reduced due to uniaxial versus hydrostatic loading" Levitas explains. "For very high pressure this reduction was drastic: about 10,000 °K. The thermodynamic approach has important advantages in that it's universal – for example, it's not limited to specific materials and atomic structure. However, thermodynamics never says that some process will occur, but only that it can occur – that it's thermodynamically admissible."

In contrast, Levitas notes, molecular dynamics simulations are performed for a specific material and atomic structure and interaction, but they reproduce all actual physical processes. "It may happen that while melting is possible from the thermodynamic point of view, it does not occur due to kinetic reasons or because other processes – such as dislocation plasticity or twinning – more rapidly release elastic stresses. Thus, one of the challenges in molecular dynamics simulations was to conceptually prove the existence of the virtual melting, which was done for the first time. Another challenge was to prove that the observed disordered state is indeed melt rather than an amorphous solid. Next, we had to investigate which parameters promote the melting, how actual melting temperature depends on them, and identify the lowest melting temperature."

Addressing these challenges required a range of insights, innovations and techniques. "I believe that the main insights and innovations were in our development of a combined thermodynamic and molecular

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dynamics approach, as well as in the proof of virtual melting existing in a shock wave several thousand degrees Kelvin below the melting temperature," Levitas says. "Also, our results further support the idea that the virtual melting is a general <u>phenomenon</u> with various realizations." While they previously suggested virtual melting as the mechanism of crystal-crystal <u>phase transformation</u>, amorphization and sublimation, here the scientists found that it also can serve as a mechanism of plastic deformation. "Lastly, since the thermodynamic features of the virtual melting here and in previous papers on phase transformations are quite similar, our first molecular dynamics confirmation of the virtual melting indirectly supports the plausibility of the virtual melting as an intermediate stage for phase transformations."

Levitas outlines how their findings impact nuclear explosions, meteorite impacts, and planned experiments in large laser facilities. "Virtual melting can compete with traditional mechanisms of plastic flow only at very high strain rates. Such conditions can be satisfied during nuclear explosions and meteorite impacts, which is why our results may be utilized for simulation of these phenomena." Introducing new and completely unexpected deformation mechanisms, he adds, may also lead to essential progress in their understanding and predictive modeling.

"The importance of such high strain rate regimes," Levitas continues, "is supported by the fact that several laboratories around the world are now developing corresponding facilities for their experimental studies – and our results may find experimental confirmation in such studies. Finally," Levitas notes, "due to the complexity of interpreting experimental results under such extreme conditions, it's always good to have an idea about which phenomena could be found – and our virtual melting process is one of them."

Levitas also describes other areas of research and technology that might benefit from their findings. "In the current paper, virtual melting was conceptually confirmed by molecular dynamics simulations for the most unexpected case of metals, like copper and aluminum." In these metals, traditional plasticity is very pronounced, which is why extremely high strain rates are required to activate an alternative mechanism like virtual melting. Since thermodynamic consideration is quite generic, this mechanism is expected in many other materials. "For materials with suppressed plasticity – for example, ceramics, high-strength alloys, or complex organic compounds – much lower strain rates may be required," Levitas points out. "Then the virtual melting may play part in more traditional high-strain rate fields, like penetration of the projectile in a target in armor ceramic, propagation of shock waves, behavior of explosive materials, and deformation in shear bands."

Regarding next steps in their research, notes Levitas, "We plan to extend our thermodynamic approach for arbitrary 3D loading – in particular for deformation under high pressure and shear strains. It can also be extended for amorphization and sublimation, which can be considered as mechanisms of stress <u>relaxation</u>. Applying a phase field approach to the phenomena discussed is another important task. Finally," Levitas concludes, "in molecular dynamic simulations we'll study polycrystalline metals and materials with suppressed plasticity."

More information: Virtual melting as a new mechanism of stress relaxation under high strain rate loading, PNAS August 14, 2012 vol. 109 no. 33 13204-13207, doi:10.1073/pnas.1203285109

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"Crystals take a chill pill: A thermomechanical theory of low-temperature melting." August 21st, 2012. http://phys.org/news/2012-08-crystals-chill-pill-thermomechanical-theory.html

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