

ORIGINAL ARTICLE

Crystal metamorphosis at stress extremes: how soft phonons turn into lattice defects

Xiaohui Liu^{1,2}, Jianfeng Gu¹, Yao Shen¹ and Ju Li^{2,3}

At 0 K, phonon instability controls the ideal strength and the ultrafast dynamics of defect nucleation in perfect crystals under high stress. However, how a soft phonon evolves into a lattice defect is still unclear. Here, we develop a full-Brillouin zone soft-phonon-searching algorithm that shows outstanding accuracy and efficiency for pinpointing general phonon instability within the joint material-reciprocal (\mathbf{x} - \mathbf{k}) spaces. By combining finite-element modeling with embedded phonon algorithm and atomistic simulation, we show how a zone-boundary soft phonon is first triggered in a simple metal (aluminum) under nanoindentation, subsequently leading to a transient new crystal phase and ensuing nucleation of a deformation twin with only one-half of the transformation strain of the conventional twin. We propose a two-stage mechanism governing the transformation of unstable short-wave phonons into lattice defects, which is fundamentally different from that initially triggered by soft long-wavelength phonons. The uncovered material dynamics at stress extremes reveal deep connections between delocalized phonons and localized defects trapped by the full nonlinear potential energy landscape and add to the rich repertoire of nonlinear dynamics found in nature. *NPG Asia Materials* (2016) 8, e320; doi:10.1038/am.2016.154; published online 21 October 2016

INTRODUCTION

The strength of solids at temperature $T=0$ K limits the attainable range of elastic strain engineering,¹ whereby finite elastic strain field^{2,3} $\epsilon(\mathbf{x})$ is tuned to yield better transistors,¹ solar cells,² superconductors⁴ and other devices. Phonon instability⁵ controls the ideal strength⁶ and influences stress-driven defect nucleation and/or phase transformation in an initially perfect crystalline lattice at 0 K. When a phonon frequency ω becomes imaginary, the harmonic oscillation will grow in amplitude and subsequently break lattice translational symmetry leading to defect nucleation.^{7,8}

Such processes are believed to occur in some low-temperature nanoindentation experiments where near-ideal strengths are experimentally measured.⁶ Previous simulation studies have focused on the long-wave phonon instability (elastic instability) with wave vector $\mathbf{k}\approx 0$ (Γ point) in the Brillouin zone (BZ) where an analytical formula can be derived based on the elastic constants and stress.^{7–12} However, the long-wave instability is just one special class of the general phonon instability in full BZ,^{13–23} which may be diagnosed by phonon calculation for homogeneously strained crystals under periodical boundary conditions (PBCs).⁵ While for crystals with inhomogeneous strain during nanoindentation, where BZ and PBC become ill-defined as a result of losing the lattice translational symmetry, it is reasonable to turn to examine the structural stability of each infinite lattice Θ that is homogeneously strained according to the local deformation at each

material point \mathbf{x} , if the material strain field is slowly varied. This is because the incipient instability volume $\tilde{\Omega}$ (centered at \mathbf{x}) in nanoindentation with a smooth indenter is usually localized inside the material without the direct contact with the external indenter,^{7,24,25} and its size increases with a decreasing local strain gradient $\nabla\epsilon$.²⁶ When $\nabla\epsilon$ is rather small, the atomic configuration in $\tilde{\Omega}$ approaches that of Θ , and the size of $\tilde{\Omega}$ would be significantly greater than the cutoff radius of the interatomic potential. It thus allows us, with diminishing error, to inspect the lattice stability of the infinite and homogeneously strained Θ instead of the inhomogeneous counterpart. To achieve this, development of new efficient algorithm is required, because in principle all the phonons in the full BZ (\mathbf{k} space) of Θ for every material point within \mathbf{x} space should be treated.

On the other hand, even if the soft phonons could be effectively pinpointed in crystals, it is still challenging to predict the final defect configuration. The unstable phonon mode indicates only the initial linear instability of atomic displacements, and as nonlinear effects develop, it becomes invalid as a descriptor at finite amplitude.¹⁵ In addition, a phonon is delocalized, and even a soft-phonon wave packet would span many atomic planes, whereas a defect such as a dislocation is localized down to one plane,²⁶ implying that there should be a localization process during the evolution of a soft-phonon pack prior to defect nucleation. For a long-wave soft phonon, the localization process into a dislocation loop has been demonstrated.^{7,8} Although the

¹School of Materials Science and Engineering, Shanghai Jiao Tong University, Shanghai, China; ²Department of Nuclear Science and Engineering, Massachusetts Institute of Technology, Cambridge, MA, USA and ³Department of Materials Science and Engineering, Massachusetts Institute of Technology, Cambridge, MA, USA
Correspondence: Professor J Gu, School of Materials Science and Engineering, Shanghai Jiao Tong University, 800 Dongchuan Road, Shanghai 200240, China.
E-mail: gujf@sjtu.edu.cn

or Professor J Li, Department of Nuclear Science and Engineering, Massachusetts Institute of Technology, 77 Massachusetts Avenue, Room 24-202, Cambridge, MA 02139, USA.

E-mail: liju@mit.edu

Received 23 April 2016; revised 5 July 2016; accepted 20 July 2016

softening of finite- \mathbf{k} phonons, particularly, the zone-boundary phonons, is quite common in shape-memory alloys,²⁷ ferroelectric compounds²⁸ and two-dimensional materials,^{18,20,23} their dynamical evolutions have not been previously reported.

In this paper, we develop a fast and robust full-BZ soft-phonon-searching algorithm we called *Phorion*. Combining finite-element modeling incorporating *Phorion* and atomistic simulations, we show how a zone-boundary phonon instability is first triggered in a simple metal (aluminum) under nanoindentation and how it metamorphoses into a deformation twin (DT). We propose a general mechanism that governs the dynamical transformation of the soft short-wave phonons.

MATERIALS AND METHODS

Full-BZ soft-phonon-searching algorithm: *Phorion*

We design *Phorion* to capture soft phonons in the joint material-reciprocal (\mathbf{x} - \mathbf{k}) spaces, in which the positioning of the ‘softest’ phonon is treated as a linearly constrained optimization problem:

$$\begin{aligned} \min_{\mathbf{x}, \mathbf{k}} f(\mathbf{x}, \mathbf{k}) &\equiv \omega^2/|\mathbf{k}|^2 \\ \text{s.t. } \mathbf{A}_1 \cdot \mathbf{x} &\leq \mathbf{b}_1 \\ \mathbf{A}_2 \cdot \mathbf{k} &\leq \mathbf{b}_2 \end{aligned} \quad (1)$$

where \mathbf{x} and \mathbf{k} are constrained within the sample volume and the first BZ, respectively; ω is calculated by diagonalizing the \mathbf{k} -dependent dynamical matrix and taking the lowest eigenvalue at each location \mathbf{x} with the finite strain. The objective function f is set as $\omega^2/|\mathbf{k}|^2$ rather than ω^2 to prevent the minimum $f(f_{\min})$ from being always pinned at the Γ point in the BZ during minimization. Obviously, once f_{\min} is detected to first hit zero at $(\mathbf{x}^*, \mathbf{k}^*)$ during the load ramp, the position \mathbf{x}^* and wave vector \mathbf{k}^* , as well as the polarization vector \mathbf{w}^*

of the soft phonon can be determined. Because there exist multiple local minima for the linearly constrained optimization problem, we implement an effective strategy to capture the global minimum reliably and efficiently by taking and utilizing the analytical gradients of Equation (1) (see Supplementary Sec. I for details).

Finite-element modeling

We seamlessly incorporate *Phorion* into the interatomic potential finite-element model (IPFEM),^{7,19} an extended local version of the quasicontinuum method.²⁹ We then use the IPFEM-*Phorion* approach to simulate a cylindrical (111) indentation of a single crystal Al plate and pinpoint the phonon instability within the joint \mathbf{x} - \mathbf{k} space at 0 K. IPFEM simulation of nanoindentation is performed using the commercial finite-element software ABAQUS (Dassault Systèmes, Vélizy-Villacoublay, France) by writing the UMAT subroutine, with the Zope–Mishin embedded-atom method (EAM) potential of Al³⁰ as the atomistic constitutive input. A single crystal Al plate, oriented as $x[11\bar{2}] - y[111] - z[1\bar{1}0]$ with a size of $2000 \times 1000 \text{ \AA}$ in the x - y plane, is indented on its top surface of the (111) plane at 0 K by an analytical rigid cylindrical indenter with a radius of 500 \AA lying along $z[1\bar{1}0]$. The finite-element model, divided into 130 592 linear plane-strain elements, is free on the top surface, and is fixed at the bottom and in the x direction on both sides. The indentation proceeds in a displacement control with an indentation depth of 0.1 \AA per step. *Phorion* is invoked at the end of each step.

Atomistic simulation

We use the LAMMPS code³¹ to perform the molecular dynamics (MD) simulation of nanoindentation. The size and boundary conditions of the indented Al plate in the x - y plane as well as the interatomic potential are the same as those in IPFEM simulation. PBC is applied in z , with half of the

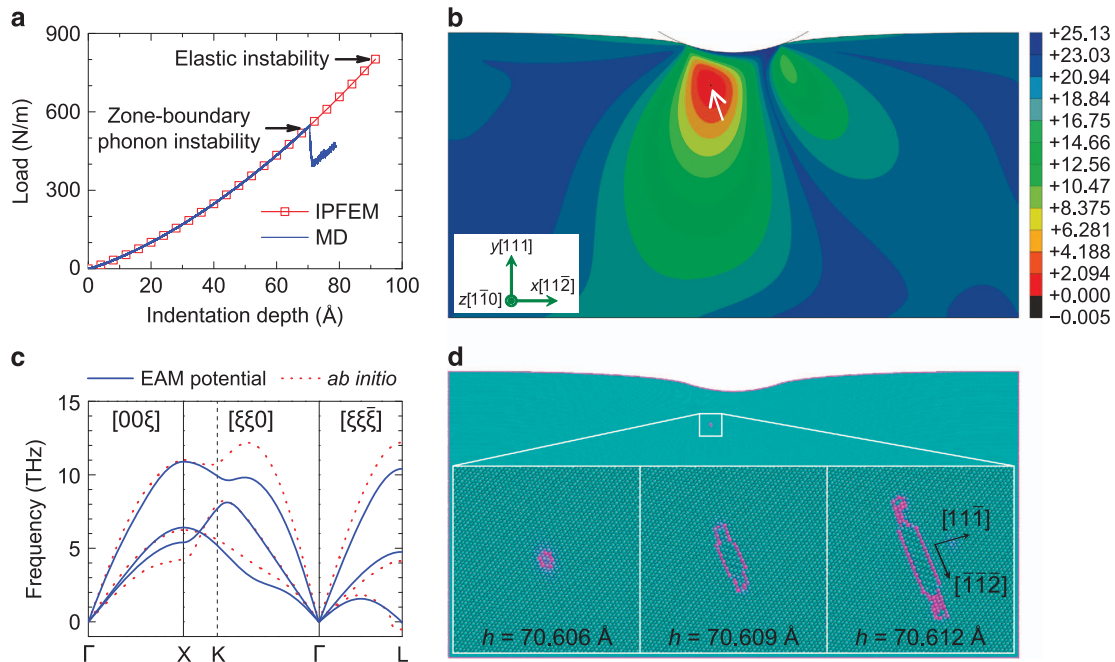


Figure 1 IPFEM-*Phorion* and MD simulations of (111) cylindrical nanoindentation of Al. The size of the Al plate in x - y plane is $2000 \times 1000 \text{ \AA}$ and the indenter radius is 500 \AA . (a) Load (P) versus indentation depth (h) by IPFEM and MD simulations. *Phorion* and the acoustic tensor formalism embedded in IPFEM predict a zone-boundary phonon instability and an elastic instability occurring at $h=69.7$ and 91.5 \AA , respectively. (b) Contour of the objective function f in Equation (1) at $h=69.7 \text{ \AA}$ by IPFEM-*Phorion* simulation, in which the wave vector \mathbf{k} is fixed at $[11\bar{1}]/2$. Homogeneous lattice instability site is indicated by the arrow, where f vanishes first at 184.6 \AA beneath the top surface and 79.0 \AA away from the central y axis. (c) Phonon spectra of the Al crystal lattice strained according to the local deformation gradient of the predicted lattice instability site shown in (b), based on the Zope–Mishin Al EAM potential (blue solid) and *ab initio* calculation using the Goedecker–Teter–Hutter Al pseudopotential (red dot). Both predict an acoustic phonon branch softening at $L = [11\bar{1}]/2$, with a polarization displacement vector $\mathbf{w}^*/[1\bar{1}0]$. (d) Atomic configurations at the early stages of DT nucleation by MD simulation. Insets show magnified snapshots of how the DT is homogeneously nucleated. DT embryo center is positioned at 186.9 \AA beneath the top surface and 80.7 \AA displaced from the central y axis. Atoms are colored according to the central symmetric parameter.²⁴ IPFEM, interatomic potential finite-element model; MD, molecular dynamics.

thickness greater than the cutoff radius given by the Zope–Mishin potential. About two million atoms are contained in the simulation box. Following ref. 24, nanoindentation is implemented by introducing an external repulsive potential acting upon the atoms on the top surface with a force constant of $10 \text{ eV } \text{Å}^{-3}$; this virtual indenter proceeds at a speed of 1 m s^{-1} . The time step is 1 fs , and the temperature is controlled at 1 K using an NVT ensemble with the Nosé–Hoover thermostat.^{32,33}

Ab initio calculation

To verify that the soft phonon predicted by IPFEM-*Phorion* based on the Zope–Mishin potential is real, we use the ABINIT code³⁴ to perform an *ab initio* calculation of the phonon spectra of the homogeneously strained lattice at the predicted instability site. The Goedecker–Teter–Hutter Al pseudopotential is adopted within the local density approximation^{35–37} with a 20 Ha energy cutoff. Marzari’s ‘cold-smearing’ method³⁸ is used for BZ integrations with a smearing parameter of 0.04 Ha . The lattice constant a_0 of Al is first obtained using a $12 \times 12 \times 12$ Monkhorst–Pack³⁹ k -point grid and is found to be 3.99 Å , in good agreement with the experimental 4.03 Å ⁴⁰ value and the 4.05 Å value given by the Zope–Mishin potential. The Al lattice with the calculated a_0 is then subjected to a homogeneous strain according to the local deformation gradient at the IPFEM-*Phorion*-predicted instability site; the phonon frequencies of which are calculated using the density functional perturbation theory⁴¹ with a $20 \times 20 \times 20$ Monkhorst–Pack k -point grid.

RESULTS AND DISCUSSION

Figure 1a shows the indentation load (P) versus the indentation depth (h) response from the IPFEM simulation. With remarkable computational efficiency (Supplementary Sec. I), *Phorion* predicts that a zone-boundary phonon instability occurs inside the crystal at $h = 69.7 \text{ Å}$ (Figure 1b), with a local Mises shear stress as large as 12.1 GPa . Examination of the phonon dispersion at the instability site based on the Zope–Mishin potential (Figure 1c) shows that a transverse acoustic phonon branch softens at L with a wave vector \mathbf{k}^* of $[11\bar{1}]/2$ and polarization vector $\mathbf{w}^*/[\bar{1}10]$. This soft mode is also verified by the *ab initio* calculation. To compare, we also utilize the acoustic tensor formalism^{42,43} (equivalent to the Λ criterion⁷) within IPFEM to predict the onset of elastic (long-wave) instability. As shown in Figure 1a, the critical h is predicted to be 91.5 Å , occurring much later than the zone-boundary phonon instability. Moreover, the instability position and the mode are also different (Supplementary Figure S4).

To verify the bi-continuum prediction (joint $\mathbf{x}-\mathbf{k}$ space), we have carried out direct MD simulation using the same empirical potential.

As shown in Figure 1a, the $P-h$ curves obtained by MD and IPFEM simulations agree very well before the first load drop in MD, and the critical h of 70.6 Å is very close to that predicted by *Phorion*, whereas the prediction by the $\mathbf{k} \approx 0$ acoustic tensor criterion lags far behind. In MD, the load drop is found to correspond to the homogeneous nucleation of a DT, and the position of the DT embryo center agrees nearly perfectly with *Phorion*’s prediction (Figure 1d). The observed twinning plane of $(11\bar{1})$ is consistent with \mathbf{k}^* provided by *Phorion*. However, the DT shear strain direction, which is $[\bar{1}12]$, differs from the phonon polarization direction $\mathbf{w}^*/[\bar{1}10]$; indeed, the two are perpendicular. This deviates greatly from the previous understanding for long-wave soft phonons.⁸ We also note a curious fact that the phonon-predicted (\mathbf{k}^* , \mathbf{w}^*) shear between adjacent atomic planes has zero Schmid factor in Figure 1b, so no mechanical work can be conducted during the initial bifurcation instability. It is then intriguing to connect the predicted soft-phonon mode with the homogeneous nucleation of the DT embryo.

To unravel this continuum-atomic connection, we perform a focused analysis of the evolution of *Phorion*-pinpointed soft phonon by direct MD simulation. First, an Al supercell under PBC is created with a homogeneous strain according to the local deformation gradient tensor at the instability site (Figure 1b), as shown in Figure 2a. It is oriented such that its x and z axes are along the soft phonon’s $\mathbf{w}^*/[\bar{1}10]$ and $\mathbf{k}^*/(11\bar{1})$ plane normal, respectively. A low-amplitude (perturbational), high-wave vector atomic displacement wave according to the soft phonon’s polarization is then injected into the supercell (Figure 2a). Its ultrafast evolution dynamics are then traced directly with MD. Figure 2b shows the evolution of the perturbational wave prior to defect nucleation. It can be observed that, instead of continuous growth, the wave experiences an amplitude oscillation while keeping its original shape up to $\sim 19.8 \text{ ps}$ (Figure 2b I–IV). Then its upper and lower envelopes bend into sinusoidal shapes, and their amplitudes continue to grow until $\sim 23.5 \text{ ps}$ (Figure 2b V–VI). Finally, the envelope curves evolve into a defect embryo residing in only a couple of atomic layers and forms a conspicuous displacement shuffling along $x[\bar{1}10]$ within the packet (Figure 2b VII–VIII).

To understand the zone-boundary phonon driven process above, we analyze the temporal evolution of the displacement in x (u_x) of each $(11\bar{1})$ atomic layer. As shown in Figure 2c, we may divide the odd- and even-numbered $(11\bar{1})$ atomic layers into A_1 and A_2 groups,

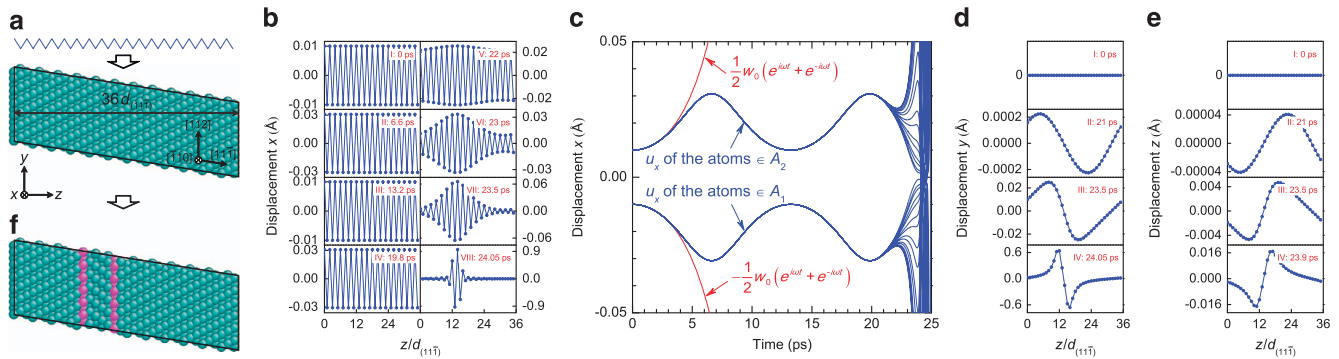


Figure 2 Structural metamorphosis of a perturbed Al supercell. (a) Al supercell under PBCs is subjected to the same strain as that at the lattice instability site predicted by *Phorion* shown in Figure 1b. It is perturbed by a plane wave according to the instability mode of the soft L phonon shown in Figure 1c, with an initial amplitude w_0 of 0.01 Å . (b, d and e) Spatial evolutions of the displacements of the 36 $(11\bar{1})$ atomic layers along x , y and z , respectively. (c) Temporal evolution of the displacements x of the $(11\bar{1})$ atomic layers (in blue), which are initially bunched together in two groups, A_1 and A_2 . They are compared with the linear exponential growth curves (in red), in which the L phonon’s imaginary circular frequency $\omega = 0.406i \text{ THz}$. (f) Atomic configuration of the supercell at 24.05 ps , in which a deformation twin is nucleated. Atoms are colored according to the central symmetry parameter, with red atoms constituting two twinning planes. PBC, periodical boundary condition.

in which their u_x are initially bunched together, thereby being grouped into the initial lower- and upper-straight envelope lines. Such bunching is maintained until ~ 19.8 ps, accounting for the undistortion of the wave profile shown in Figure 2b I–IV. During this stage, u_x of A_1 and A_2 groups first follow linear exponential growth as $u_x(t) = \mp \frac{1}{2} w_0 (e^{i\omega t} + e^{-i\omega t})$ with an imaginary ω within the harmonic approximation, where w_0 is the initial amplitude of the perturbational wave. However, they then deviate from this pure amplitude growth and start to oscillate. Such a response is attributed to the anharmonic terms in the total potential energy expansion $\Delta\Psi = c_2 u_A^2 + c_4 u_A^4 + \dots$, where u_A is the phonon displacement. The lack of the cubic term is owing to the symmetry of the zone-boundary phonon mode (see Supplementary Sec. III for the analytical expressions of $\Delta\Psi$). Figure 3 compares $\Delta\Psi - u_A$ of the harmonic expansion with that including the fourth-order anharmonic term. In contrast to a downward parabola of the harmonic expansion, the significant anharmonic effect yields a double-well potential, which in principle could arrest the perturbational wave and suppress the amplitude

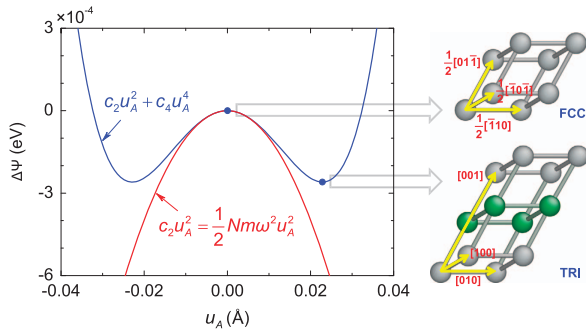


Figure 3 Response of the potential energy change ($\Delta\Psi$) of the Al supercell to the phonon displacement (u_A) of the perturbational wave. $\Delta\Psi$ can be fitted as $c_2 u_A^2 + c_4 u_A^4$ (in blue), where $c_2 = -0.993 \text{ eV } \text{Å}^{-2}$ and $c_4 = -949.7 \text{ eV } \text{Å}^{-4}$, compared with the harmonic-approximation response following $\Delta\Psi = c_2 u_A^2 = \frac{1}{2} Nm \omega^2 u_A^2$ (in red), where N is the number of atoms in the supercell and m is the atomic mass of Al. Primitive cells of the FCC and TRI structures, corresponding to the center and the bottom of the double wells, respectively, are shown on the right, and they have $d_{[001]}^{\text{TRI}} = 2d_{[11\bar{1}]}^{\text{FCC}}$. Two basis atoms in the TRI primitive cell are colored with gray and green, respectively. TRI, triclinic.

growth. The behavior of the injected wave in stages I–IV in Figure 2b could thus be understood. Indeed, using the Landau polynomials language of phase transitions, what Figure 3 describes is a phase transformation that should occur after the L -phonon ‘spinodal’ instability (in displacement, instead of chemical concentration) to a new crystal structure. In Figure 3, adjacent $(11\bar{1})$ atomic layers undergo displacement shuffling along $[\bar{1}10]$ on the path from the center to one of the double wells, which is actually accompanied by a phase transformation from the original face-centered cubic (FCC) phase to a new triclinic (TRI) phase. As shown in the primitive cell of the TRI structure, there exist two basis atoms occupying two sites (denoted as B_1 and B_2) on adjacent $(11\bar{1})$ atomic layers that exactly belong to the A_1 and A_2 groups, respectively.

However, this phase transformation, triggered by the zone-boundary soft phonon, does not significantly relax the strain energy, and the TRI phase is still subjected to high stress. Therefore, it is still necessary to keep inspecting its structural instability. If the TRI structure were mechanically stable, the injected perturbational and subsequently grown wave could be dissipated by phonon scattering⁴⁴ and permanently arrested by the double-well potential, which would lead to a polymorphic phase transition such as those observed in some indentation experiments.^{45,46} Otherwise, the second-generation lattice instability within the TRI structure would affect the wave growth and may drive it to escape from the TRI phase in a different direction not reflected in the Landau polynomial plot of Figure 3.

In fact, second-generation phonon spectra analysis shows that the TRI structure is elastically unstable for a long wave with $\mathbf{k} // [001]^{\text{TRI}}$ (Figure 4a), where $[001]^{\text{TRI}}$ is a reciprocal vector to the real-space lattice vectors of TRI shown in Figure 3. For the present supercell with a limited size along z , only a soft phonon with \mathbf{k} of $[001]^{\text{TRI}}/18$ is allowed to be excited and is expected to generate two independent unstable long waves denoted as β_1 and β_2 that involve the collective movement of the B_1 - and B_2 -sited (or A_1 - and A_2 -grouped) atoms, respectively. Eigenmode analysis shows that each of the long waves can be decomposed into three polarization components including two transverse waves β_i^x and β_i^y oscillating along x and y , respectively, and a longitudinal wave β_i^z oscillating along z ($i=1, 2$), with their normalized amplitudes v and phase φ shown in Supplementary Table S3. As the displacement shuffling along $[\bar{1}10]$ breaks the central symmetry of B_1 - and B_2 -sited atoms in the TRI structure, these

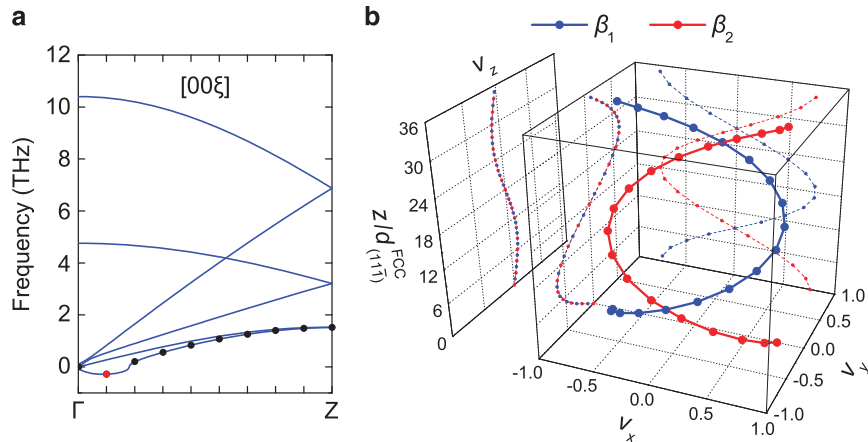


Figure 4 Near- Γ phonon instability of the TRI structure. (a) Phonon spectra of the TRI structure along the Γ -Z path in the BZ. For the present supercell with $l_z = 18d_{[001]}^{\text{TRI}}$, there are nine k points allowable distributed uniformly between (Γ, Z) , of which the frequencies on the softened phonon branch are denoted as solid dots. Among these, only the k point at $[001]^{\text{TRI}}/18$ has an imaginary frequency, denoted as the red dot. (b) Wave profiles with normalized amplitudes of two unstable long waves β_1 and β_2 excited by the soft phonon denoted in (a). Two long waves spatially constitute a double helix, with their polarization components shown on the projective planes. BZ, Brillouin zone; TRI, triclinic.

polarization components have different phases, yielding a double-helix profile for the β_1 – β_2 pair (Figure 4b). For the β_1^x and β_2^x waves, owing to a phase difference of π , their superposition upon the original injected wave would bend its lower and upper envelope lines into a pair of sinusoidal curves oscillating oppositely and coherently, and drive the system to escape from the double-well potential; this explains the profile of the injected wave shown in Figure 2b V. The continuous amplitude growth of the sinusoidal envelope curves (Figure 2b VI) followed by progressive localization (Figure 2b VII–VIII) agrees well with the localization behavior of an unstable elastic wave,⁸ even though ‘wave-front steepening’ that breaks the positive strain–negative strain symmetry is not observed, resulting from the vanishing shear stress component $\tau_{zx}=0$ (zero Schmid factor) here.

As indicated above, the TRI structure would also generate polarization components oscillating along y and z , which contribute to the DT nucleation as well. For the β_1^y and β_2^y waves with the same ν and φ (Supplementary Table S3), they are expected to be merged into a single long wave β^y with phase $\pi/2$ ahead of β_1^x , involving collective displacements along y of all the A_1 - and A_2 -grouped atoms. Such a prediction is verified by tracing the evolution of u_y for all the $(11\bar{1})$ atomic layers, shown in Figure 2d. It can be observed that a single long u_y wave is excited, followed by the amplitude growth (Figure 2d I–II), wave-front steepening (Figure 2d III), and shrinking into a couple of atomic layers and thus producing a uniform shear along $[\bar{1}12]$ (Figure 2d IV); this wave actually behaves as an unstable elastic wave.⁸ Simultaneously, a single u_z wave also develops owing to crystallographic shear-tension coupling⁸ and experiences the dynamics similar to those of the u_y wave (Figure 2e), albeit with much smaller amplitude.

Therefore, eventually, the TRI phase turns out to be just an intermediate product, decomposing within tens of picoseconds by the second-stage, near- Γ phonon instability to evolve into a lattice defect–DT, as shown in Figure 2f. Its atomic structure is verified to be consistent with that in Figure 1d, proving the continuum-atomic linkage and the validity of our bi-continuum prediction (joint $\mathbf{x}-\mathbf{k}$ space) as well as focal study of nucleation dynamics with PBCs. Unexpectedly, and quite distinct from a conventional DT in which all atomic planes slip along the same direction (same Shockley partial Burgers vector) thereby producing a uniform shear strain, the DTs in Figure 1d and 2f have two different slip vectors alternating along $[12\bar{1}]$ and $[\bar{2}1\bar{1}]$ on every successive $(11\bar{1})$ planes. In fact, our DT embryo is mainly achieved by the superposition of the displacement shuffling along $[\bar{1}10]$ resulting from the localization of the β_1^x and β_2^x waves, and the shear strain along $[\bar{1}12]$ contributed by the accumulated deformation before the L -phonon softening plus the localization of the β^y wave (β^z contribution can be neglected owing to a small amplitude). The two slip vectors are produced through $\pm\frac{1}{4}[1\bar{1}0] + \frac{1}{12}[\bar{1}12] = \frac{1}{6}[12\bar{1}]$ and $\frac{1}{6}[\bar{2}1\bar{1}]$ for adjacent $(11\bar{1})$ planes, and the stacking order is thus changed from the original ‘ $\dots ABCABCAB\dots$ ’, for example, to ‘ $\dots A[\bar{B}ACBAC]A\dots$ ’. In other words, a DT with a width of $5d_{(11\bar{1})}$ nucleates homogeneously in the lattice (see Supplementary Figure S5 for the schematic of how the stacking order is changed). If viewed from $[1\bar{1}0]$, the DT has an averaged transformation shear along $[\bar{1}12]$, where the average relative slip vector between two adjacent $(11\bar{1})$ planes is $\frac{1}{12}[\bar{1}12]$, only 50% of that of the conventional textbook DT (Supplementary Figure S6), leading to a significantly smaller transformation strain and smaller elastic energy of embedding. Interestingly, our DT embryo shows a crude resemblance to the DT growing from a grain boundary following a dislocation–reaction and cross-slip mechanism recently proposed by Zhu *et al.*,^{47,48} which also incorporates two different slip vectors or alternating Shockley partial

Burgers vectors. To be revealed as the product of two degenerate soft phonons is quite a surprise, though.

CONCLUSIONS

In summary, we have developed a robust full-BZ soft-phonon-searching algorithm, which can be combined with other multiscale simulation methods or *ab initio* calculation tools to predict the lattice instability and the ideal strength of materials that enable elastic strain engineering.¹ We have revealed how a continuum soft phonon evolves into a discrete atomic defect within tens of picoseconds in an indented perfect crystal at stress extreme. This involves four stages: (1) linear growth of an unstable short wave triggered by a softened L phonon, well described by the harmonic-approximation phonon theory; (2) arrest of the short wave by a double-well potential in the form of an anharmonic Landau polynomial, indicating a polymorphic phase transformation from FCC to TRI; (3) excitation of two independent unstable long waves by a soft near- Γ phonon in the TRI lattice; and (4) growth and localization of the two long waves until trapped in a couple of atomic layers, where an unconventional DT embryo with staggered slip Burgers vectors and 50% of the total transformation strain of the textbook FCC DT is formed. In this two-stage soft-phonon cascade, the Landau polynomial has a critical role, with the anharmonic energy terms arresting the dynamics of an L phonon in FCC followed by a near- Γ phonon in TRI. In Supplementary Sec. IV, we show an analytical scaling theory that explains why an unstable long elastic wave is much more difficult to be trapped in a polynomial energy landscape.

The Figure 3 scenario is not unlike the well-known temperature-driven phase transformation within a double-well potential⁴⁹ in phonon coordinate. However, a salient feature of the stress-driven martensitic transformation owing to a softened zone-boundary phonon is that the new phase still withstands high stress, and it is thus highly likely to be unstable as well and prone to the next-stage evolution. This is because a first-generation transition yields little transformation strain, as the zone-boundary phonon has too short wavelength, unlike a long-wavelength phonon that can significantly relax the strain in a local region. In view of this, we believe that the proposed two-stage transformation mechanism can generally describe the dynamics of the homogeneous defect nucleation induced by a softened zone-boundary (or nearby) phonon at stress extremes. As we show in Supplementary Sec. V, it can also fully account for the homogeneous nucleation of a dislocation. The general utility of IPFEM-*Phorion* bi-continuum stability analysis has been repeatedly verified by our direct atomistic simulations. The uncovered material dynamics at stress extremes reveal deep connections between delocalized phonons in \mathbf{x} and \mathbf{k} continua—a wave phenomenon controlled by the harmonic part of potential—and localized atomic defects trapped by the full nonlinear interatomic potential energy landscape, which adds to the rich repertoire of nonlinear dynamics found in nature.^{50,51}

CONFLICT OF INTEREST

The authors declare no conflict of interest.

ACKNOWLEDGEMENTS

This work was supported by the Natural Science Foundation of China under Grant Nos. 50971090, 51071102, 51201101 and 51471107. JL acknowledges the support of NSF DMR-1410636. XL acknowledges helpful discussion with M.H. Wu about the *ab initio* calculation of phonon spectra.

- 1 Li, J., Shan, Z. W. & Ma, E. Elastic strain engineering for unprecedented materials properties. *MRS Bull.* **39**, 108–117 (2014).
- 2 Feng, J., Qian, X., Huang, C.-W. & Li, J. Strain-engineered artificial atom as a broad-spectrum solar energy funnel. *Nat. Photonics* **6**, 866–872 (2012).
- 3 Li, H., Contryman, A. W., Qian, X., Ardakani, S. M., Gong, Y., Wang, X., Weisse, J. M., Lee, C. H., Zhao, J., Ajayan, P. M., Li, J., Manoharan, H. C. & Zheng, X. Optoelectronic crystal of artificial atoms in strain-textured molybdenum disulphide. *Nat. Commun.* **6**, 7381 (2015).
- 4 Hao, S., Cui, L., Wang, H., Jiang, D., Liu, Y., Yan, J., Ren, Y., Han, X., Brown, D. E. & Li, J. Retaining large and adjustable elastic strains of kilogram-scale Nb nanowires. *ACS Appl. Mat. Interfaces* **8**, 2917–2922 (2016).
- 5 Liu, F., Ming, P. M. & Li, J. *Ab initio* calculation of ideal strength and phonon instability of graphene under tension. *Phys. Rev. B* **76**, 064120 (2007).
- 6 Zhu, T. & Li, J. Ultra-strength materials. *Prog. Mater. Sci.* **55**, 710–757 (2010).
- 7 Li, J., Van Vliet, K. J., Zhu, T., Yip, S. & Suresh, S. Atomistic mechanisms governing elastic limit and incipient plasticity in crystals. *Nature* **418**, 307–310 (2002).
- 8 Li, J., Ngan, A. H. W. & Gumbsch, P. Atomistic modeling of mechanical behavior. *Acta Mater.* **51**, 5711–5742 (2003).
- 9 Van Vliet, K. J., Li, J., Zhu, T., Yip, S. & Suresh, S. Quantifying the early stages of plasticity through nanoscale experiments and simulations. *Phys. Rev. B* **67**, 104105 (2003).
- 10 Zhu, T., Li, J., Van Vliet, K. J., Ogata, S., Yip, S. & Suresh, S. Predictive modeling of nanoindentation-induced homogeneous dislocation nucleation in copper. *J. Mech. Phys. Solids* **52**, 691–724 (2004).
- 11 Fago, M., Hayes, R. L., Carter, E. A. & Ortiz, M. Density-functional-theory-based local quasicontinuum method: prediction of dislocation nucleation. *Phys. Rev. B* **70**, 100102 (2004).
- 12 Liu, X. H., Gu, J. F., Shen, Y. & Chen, C. F. Anisotropy in homogeneous dislocation nucleation by nanoindentation of single crystal Cu. *Scripta Mater.* **58**, 564–567 (2008).
- 13 Einarsdóttir, K., Sadigh, B., Grimvall, G. & Ozoliņš, V. Phonon instabilities in fcc and bcc Tungsten. *Phys. Rev. Lett.* **79**, 2073–2076 (1997).
- 14 Li, J. & Yip, S. Atomistic measures of materials strength. *Comput. Model. Eng. Sci.* **3**, 219–227 (2002).
- 15 Clatterbuck, D. M., Krenn, C. R., Cohen, M. L. & Morris, J. W. Phonon instabilities and the ideal strength of aluminum. *Phys. Rev. Lett.* **91**, 135501 (2003).
- 16 Dmitriev, S. V., Li, J., Yoshikawa, N. & Shibutani, Y. Theoretical strength of 2D hexagonal crystals: application to bubble raft indentation. *Philos. Mag.* **85**, 2177–2195 (2005).
- 17 Dubois, S. M. M., Rignanese, G. M., Pardo, T. & Charlier, J. C. Ideal strength of silicon: an *ab initio* study. *Phys. Rev. B* **74**, 235203 (2006).
- 18 Marianetti, C. A. & Yevick, H. G. Failure mechanisms of graphene under tension. *Phys. Rev. Lett.* **105**, 245502 (2010).
- 19 Liu, X. H., Gu, J. F., Shen, Y., Li, J. & Chen, C. F. Lattice dynamical finite-element method. *Acta Mater.* **58**, 510–523 (2010).
- 20 Li, T. Ideal strength and phonon instability in single-layer MoS₂. *Phys. Rev. B* **85**, 235407 (2012).
- 21 Grimvall, G., Magyari-Köpe, B., Ozoliņš, V. & Persson, K. A. Lattice instabilities in metallic elements. *Rev. Mod. Phys.* **84**, 945–986 (2012).
- 22 Zhou, W., Zhang, Y., Sun, H. & Chen, C. Ideal strength and structural instability of aluminum at finite temperatures. *Phys. Rev. B* **86**, 054118 (2012).
- 23 Isaacs, E. B. & Marianetti, C. A. Ideal strength and phonon instability of strained monolayer materials. *Phys. Rev. B* **89**, 184111 (2014).
- 24 Kelchner, C. L., Plimpton, S. J. & Hamilton, J. C. Dislocation nucleation and defect structure during surface indentation. *Phys. Rev. B* **58**, 11085–11088 (1998).
- 25 Gouldstone, A., Van Vliet, K. J. & Suresh, S. Nanoindentation—simulation of defect nucleation in a crystal. *Nature* **411**, 656–656 (2001).
- 26 Miller, R. E. & Rodney, D. On the nonlocal nature of dislocation nucleation during nanoindentation. *J. Mech. Phys. Solids* **56**, 1203–1223 (2008).
- 27 Otsuka, K. & Ren, X. Physical metallurgy of Ti-Ni-based shape memory alloys. *Prog. Mater. Sci.* **50**, 511–678 (2005).
- 28 Blinc, R. & Zeks, B. *Soft Modes in Ferroelectrics and Antiferroelectrics* (Elsevier, New York, NY, USA, 1974).
- 29 Tadmor, E. B., Ortiz, M. & Phillips, R. Quasicontinuum analysis of defects in solids. *Philos. Mag. A* **73**, 1529–1563 (1996).
- 30 Zope, R. R. & Mishin, Y. Interatomic potentials for atomistic simulations of the Ti-Al system. *Phys. Rev. B* **68**, 241021–2410214 (2003).
- 31 Plimpton, S. Fast parallel algorithms for short-range molecular dynamics. *J. Comput. Phys.* **117**, 1–19 (1995).
- 32 Nosé, S. A unified formulation of the constant temperature molecular dynamics methods. *J. Chem. Phys.* **81**, 511–519 (1984).
- 33 Hoover, W. G. Canonical dynamics: equilibrium phase-space distributions. *Phys. Rev. A* **31**, 1695–1697 (1985).
- 34 Gonze, X., Beuken, J.-M., Caracas, R., Detraux, F., Fuchs, M., Rignanese, G.-M., Sindic, L., Verstraete, M., Zerah, G., Jollet, F., Torrent, M., Roy, A., Mikami, M., Ghosez, Ph., Raty, J.-Y. & Allan, D. C. First-principles computation of material properties: the ABINIT software project. *Comput. Mater. Sci.* **25**, 478–492 (2002).
- 35 Ceperley, D. M. & Alder, B. J. Ground state of the electron gas by a stochastic method. *Phys. Rev. Lett.* **45**, 566–569 (1980).
- 36 Perdew, J. P. & Wang, Y. Accurate and simple analytic representation of the electron-gas correlation energy. *Phys. Rev. B* **45**, 13244–13249 (1992).
- 37 Goedecker, S., Teter, M. & Hutter, J. Separable dual-space Gaussian pseudopotentials. *Phys. Rev. B* **54**, 1703–1710 (1996).
- 38 Marzari, N. *Ab-initio Molecular Dynamics for Metallic Systems* (University of Cambridge, 1996).
- 39 Monkhorst, H. J. & Pack, J. D. Special points for Brillouin-zone integrations. *Phys. Rev. B* **13**, 5188–5192 (1976).
- 40 Eisenmann, B., Schäfer, H., Landolt-Börnstein. in *Structure Data of Elements and Intermetallic Phases. Elements, Borides, Carbides, Hydrides* (eds Hellwege K.-H. & Hellwege A. M.) (Springer-Verlag, New York, NY, USA, 1988).
- 41 Baroni, S., de Gironcoli, S., Dal Corso, A. & Giannozzi, P. Phonons and related crystal properties from density-functional perturbation theory. *Rev. Mod. Phys.* **73**, 515 (2001).
- 42 Hill, R. Acceleration waves in solids. *J. Mech. Phys. Solids* **10**, 1–16 (1962).
- 43 Rice, J. R. in *Theoretical and Applied Mechanics* (ed. Koiter W. T.) Vol. 1, 207–220 (North-Holland, Amsterdam, The Netherlands, 1976).
- 44 Li, J., Porter, L. & Yip, S. Atomistic modeling of finite-temperature properties of crystalline β -SiC: II. Thermal conductivity and effects of point defects. *J. Nucl. Mater.* **255**, 139–152 (1998).
- 45 Pharr, G. M., Oliver, W. C. & Harding, D. S. New evidence for a pressure-induced phase-transformation during the indentation of silicon. *J. Mater. Res.* **6**, 1129–1130 (1991).
- 46 Jang, J. I., Lance, M. J., Wen, S. Q. & Pharr, G. M. Evidence for nanoindentation-induced phase transformations in germanium. *Appl. Phys. Lett.* **86**, 131907 (2005).
- 47 Zhu, Y. T., Wu, X. L., Liao, X. Z., Narayan, J., Mathadhu, S. N. & Kecskés, L. J. Twinning partial multiplication at grain boundary in nanocrystalline fcc metals. *Appl. Phys. Lett.* **95**, 031909 (2009).
- 48 Zhu, Y. T., Liao, X. Z. & Wu, X. L. Deformation twinning in nanocrystalline materials. *Prog. Mater. Sci.* **57**, 1–62 (2012).
- 49 Landau, L. D. & Lifshitz, E. M. *Statistical Physics* (Elsevier Science, 2013).
- 50 Thompson, J. M. T. & Stewart, H. B. *Nonlinear Dynamics and Chaos* (Wiley, Chichester, UK, 2002).
- 51 Strogatz, S. H. *Nonlinear Dynamics and Chaos: With Applications to Physics, Biology, Chemistry, and Engineering* (Westview Press, Boulder, CO, USA, 2014).



This work is licensed under a Creative Commons Attribution 4.0 International License. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder to reproduce the material. To view a copy of this license, visit <http://creativecommons.org/licenses/by/4.0/>

© The Author(s) 2016

Supplementary Information accompanies the paper on the NPG Asia Materials website (<http://www.nature.com/am>)