Topological Defects in Solids with Odd Elasticity

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Crystallography typically studies collections of point particles whose interaction forces are the gradient of a potential. Lifting this assumption generically gives rise in the continuum limit to a form of elasticity with additional moduli known as odd elasticity. We show that such odd elastic moduli modify the strain induced by topological defects and their interactions, even reversing the stability of, otherwise, bound dislocation pairs. Beyond continuum theory, isolated dislocations can self propel via microscopic work cycles active at their cores that compete with conventional Peach-Koehler forces caused, for example, by an ambient torque density. We perform molecular dynamics simulations isolating active plastic processes and discuss their experimental relevance to solids composed of spinning particles, vortexlike objects, and robotic metamaterials.

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Topological defects are local singularities in an order parameter that have global consequences at large scales [1-9]. In active systems, topological defects exhibit distinctive properties such as self-propulsion or nonreciprocal interactions [10–31]. In the study of crystalline defects, it is often assumed that a potential energy governs the interactions between the constituent particles. This assumption, however, need not hold in driven and active solids. For example, Fig. 1(a) shows a nonconservative interaction force-one in which the work done between any two configurations depends on the path taken. Such microscopic forces generically give rise in the continuum limit to odd elasticity: additional moduli that break the major symmetry of the elastic modulus tensor [32,33]. Experimental signatures of odd elasticity have been reported in solids made of spinning embryos [34] and colloids [30] with hydrodynamic interactions, and robotic metamaterials [35,36]. Likewise, gyroscopic matter [37–46] and vortexlike objects [47–55], e.g., skyrmions [56–61], can exhibit a special case of odd elastic dynamics [see Supplemental Material (SM) [62]].

Crystallography without potentials.—A typical starting place in crystallography is a collection of point particles at positions $\underline{\mathbf{x}} = (\mathbf{x}^1, \mathbf{x}^2, ..., \mathbf{x}^N)$ interacting via forces that are the gradient of a potential

$$\mathbf{F}^{\alpha}(\underline{\mathbf{x}}) = -\frac{\partial V(\underline{\mathbf{x}})}{\partial \mathbf{x}^{\alpha}}.$$
 (1)

However, in general, Eq. (1) need not be valid. Experimentally relevant [29–31,34,72–77] counterexamples include pairwise forces of the form $\mathbf{F}^{\alpha}(\underline{\mathbf{x}}) = \sum_{\beta \neq \alpha} \mathbf{F}(\mathbf{x}^{\alpha} - \mathbf{x}^{\beta})$ where $\mathbf{F}(\mathbf{r})$ depends only on particle separation

$$\mathbf{F}(\mathbf{r}) = F^{\parallel}(r)\hat{\mathbf{r}} - F^{\perp}(r)\hat{\boldsymbol{\phi}}.$$
 (2)

Here, \mathbf{r} is the relative coordinate between two interacting particles, $\hat{\phi} = -\epsilon \cdot \hat{\mathbf{r}}$, and ϵ is the antisymmetric tensor. Henceforth, we will focus on first order dynamics $\dot{\mathbf{x}}^{\alpha} = \mathbf{F}^{\alpha}$, which arise in overdamped media, as well as in vortex [47–55] and gyroscopic [37–46] crystals in which the forces $\mathbf{F}^{\alpha} = \boldsymbol{\epsilon} \cdot (\partial V / \partial \mathbf{x}^{\alpha})$ are transverse to potential gradients, see SM [62], Sec. S1 A. Subject to first order dynamics, the quantity $P \equiv \dot{\mathbf{x}}^{\alpha} \cdot \mathbf{F}^{\alpha}(\underline{\mathbf{x}})$ is greater than zero for trajectories satisfying the equations of motion [78]. Of particular interest here are interparticle forces such that $W_{\mathcal{C}} = \oint_{\mathcal{C}} P dt \neq 0$ for closed contours \mathcal{C} [see Fig. 1(a)]. Notice that $\nabla \times \mathbf{F} = 0$ is equivalent to requiring that $W_{\mathcal{C}} = 0$ for all contractible loops C. In Newtonian mechanics, W_{C} has the interpretation of energy, and $\nabla \times \mathbf{F} \neq 0$ is equivalent to violating Maxwell-Betti reciprocity (MBR) [79], which means that the linear response matrix between force and displacement is no longer symmetric, see SM [62], Sec. S1 A.

Continuum theory.—In the continuum, we describe the state of the solid via a continuous displacement field $\mathbf{u}(\mathbf{r})$, and we assume that the coarse-grained forces may be expressed as $f_j = \partial_i \sigma_{ij}$, where σ_{ij} is the Cauchy stress tensor (see Ref. [28] for a treatment of dislocations that lifts this assumption). We expand the Cauchy stress tensor σ_{ij} in terms of the displacement gradient u_{mn} to obtain $\sigma_{ij} = \sigma_{ij}^0 + C_{ijmn}u_{mn}$. Here, C_{ijmn} denotes the elastic modulus tensor and σ_{ij}^0 is the stress prior to deformation. In 2D isotropic crystals, the linearized stress-strain relationship is summarized by the pictorial equation [32]



FIG. 1. Odd elasticity modifies dislocation interactions and their stability. (a) A particle at point \mathbf{x}_1 exerts a force \mathbf{F} on a particle at point \mathbf{x}_2 . This force is nonconservative, so nonzero work is done along the closed cycle C. (b) A dislocation is defined by a Burgers vector **b** that represents the offset from what would, otherwise, be a closed loop γ . (c),(d) An orange dislocation is held stationary while a second antialigned dislocation is free to move along its glide plane subject to the Peach-Koehler force (red arrows). When |A/B| < 1, the free dislocation has two stable points located along rays forming an angle $\pi/4$ with the glide plane. The effective potential V_{eff} as a function of the horizontal (r_x) and vertical (r_y) distance between the dislocations. (e),(f) When A/B > 1, the rightmost stable equilibrium moves outward beyond $\pi/4$. (g),(h) When $A/B = \infty$, only one stable equilibrium position exists and the shaded region is an unstable basin.

$$\begin{pmatrix} \textcircled{\bullet} \\ \end{array} \\ \textcircled{\bullet} \\ \end{array} \\ \end{array}$$

See SM [62] Sec. S1 C for tensor notation. In Eq. (3), p_0 and τ_0 are mechanically interpreted in terms of the pressure and torque density associated with σ_{ij}^0 . The matrix in Eq. (3) corresponds to C_{ijmn} and has three diagonal components, the bulk *B*, shear μ , and rotational Γ moduli, and three off-diagonal moduli Λ , *A*, and K^o . The antisymmetric contributions to the matrix in Eq. (3) are what we refer to as odd elastic moduli [32]. The counterpart of $P = \dot{\mathbf{x}}^a \cdot \mathbf{F}^a$ in the continuum is $P = \int S_{ij} \dot{u}_{ij} d^2 r$, where S_{ij} is the first Piola-Kirchof tensor [80–84]. Writing $S_{ij} = S_{ij}^0 + \tilde{C}_{ijmn} u_{mn}$, we have

$$\tilde{C}_{ijmn} = C_{ijmn} + \sigma^0_{ij}\delta_{mn} - \sigma^0_{mj}\delta_{in}.$$
(4)

In the continuum, the Maxwell-Betti reciprocity theorem states the internal forces must be nonconservative if $\tilde{C}_{ijmn} \neq \tilde{C}_{mnij}$ [79]. Odd elasticity $(C_{ijkl} \neq C_{klij})$ coincides with broken MBR $(\tilde{C}_{ijkl} \neq \tilde{C}_{klij})$ when no ambient stress is present $(\sigma_{ij}^0 = 0)$. In terms of the moduli in Eq. (3), the condition for MBR, $\tilde{C}_{ijmn} = \tilde{C}_{mnij}$, reads

$$2K^o = A - \Lambda = 2\tau_0. \tag{5}$$

Notice from Eq. (5) that odd elasticity can arise even when MBR holds (i.e., the microscopic forces are conservative) provided that τ_0 is nonvanishing. For instance, the transverse microscopic force $F^{\perp}(r) \propto (1/r)$ is curl free, i.e., conservative and, nonetheless, gives rise to A and K^o (SM [62], Sec. S1 E). In this case, A and K^o can be detected from static stress-strain measurements, but the work they generate during strain cycles must be cancelled by τ_0 . The distinction between C_{ijkl} and \tilde{C}_{ijkl} vanishes when no ambient stress σ_{ij}^0 is present. See SM [62], Sec. S1 E for how crystals with purely transverse interactions, such as lattices of vortexlike objects [47–61], or gyroscopes [37–46], can be mathematically cast as a special case of odd elasticity with $B = \mu = 0$.

Microscopics.—To relate the moduli to the microscopic transverse forces, we linearize Eq. (2) about the lattice spacing *a*: $F^{\perp}(r) = F_0^{\perp} - k^a(r-a)$. The resulting odd elastic moduli for a hexagonal lattice read

$$A \approx \frac{\sqrt{3}}{2} \left(k^a + \frac{F_0^{\perp}}{a} \right), \qquad K^o \approx \frac{\sqrt{3}}{4} \left(k^a - \frac{3F_0^{\perp}}{a} \right), \quad (6)$$

along with an ambient torque density $\tau_0 = \sqrt{3}F_0^{\perp}/a$, see SM, Sec. S1D and Refs. [28,32]. Additionally, the modulus A arises whenever the full torque density $\tau = \epsilon_{ij}\sigma_{ij}/2$ couples to local dilation $\partial_i u_i = -\delta\rho/\rho_0$, namely, $A = (d\tau/d\rho)\rho_0$. The forces in Eq. (2) depend only on r and, therefore, cannot contribute to Γ and Λ which couple to solid body rotations. However, Γ and Λ can arise in response to external fields or interactions with a substrate [8], see SM [62], Sec. S1 E for examples. Henceforth, we set $\Lambda = \Gamma = 0$, see S.M. [62], Sec. S2 C for a general treatment. *Continuum solutions.*—Topological defects are singularities where $u_i(\mathbf{r})$ becomes multivalued, e.g., the dislocation in Fig. 1(b) is defined by the Burgers vector b_i

$$b_j = \oint_{\gamma} \partial_i u_j dr_i. \tag{7}$$

where γ is a counterclockwise contour around the dislocation. We solve $\partial_i \sigma_{ij}(\mathbf{r}) = 0$ together with Eq. (7) to obtain static solutions of the dislocation displacement field (SM [62], Sec. S2 C)

$$\mathbf{u}_{\text{disl}} = \frac{1}{2\pi} \left\{ \boldsymbol{\phi} \mathbf{b} + \frac{1-\nu}{2} \log(r) \boldsymbol{\epsilon} \cdot \mathbf{b} - \frac{1+\nu}{2} (\hat{\mathbf{r}} \cdot \mathbf{b}) \hat{\boldsymbol{\phi}} - \nu^{o} [\log(r) \mathbf{b} + (\hat{\boldsymbol{\phi}} \cdot \mathbf{b}) \hat{\boldsymbol{\phi}}] \right\}.$$
(8)

where *r* and ϕ are polar coordinates about the defect. The elastic properties enter only through (i) a modified Poisson's ratio, ν , (SM [62], Sec. S2 C) and (ii) a purely nonreciprocal odd ratio [32]

$$\nu^{o} = \frac{BK^{o} - A\mu}{\mu(B + \mu) + K^{o}(A + K^{o})}.$$
(9)

The effect of ν^{o} in Eq. (8) is to globally rotate the local shear axis by an angle $\delta \alpha$

$$\delta \alpha = -\frac{1}{2} \arctan\left(\frac{2\nu^o}{1+\nu}\right). \tag{10}$$

See SM [62], Figs. S5 and S6 for an illustration and numerical validation. Sec. S2 B in the SM [62], provides similar results for point defects and isolated disclinations, which have recently been observed in experiments [34] of spinning embryos interacting via transverse forces, cf. Eq. (2).

Dislocation interactions.—The modified stress field alters dislocation interactions. First, consider the work done in quasistatically moving a test dislocation by δX_i through a preexisting stress field $\sigma_{ij}^{(\text{pre})}$ obeying $\partial_i \sigma_{ij}^{(\text{pre})} = 0$. Regardless of the constitutive properties of the material, the work done by $\sigma_{ij}^{(\text{pre})}$ is given by the Peach-Koehler (PK) formula $\delta W = f_m^{\text{PK}} \delta X_m$, where $f_m^{\text{PK}} = \epsilon_{mi} \sigma_{ij}^{(\text{pre})} b_j$ (SM [62], Sec. S3 A). The continuum interaction between two dislocations is the PK force experienced by the test dislocation as a result of the stress field generated by the other. In Figs. 1(c)–1(h), we examine the interaction between two antiparallel dislocations in the presence of odd elastic moduli. The modulus A provides a nontrivial modification

$$f^{\rm PK}(r_x) = \frac{(1-\nu)b_1b_2}{\pi r^4} (r_x^2 - r_y^2) (Br_x + Ar_y), \qquad (11)$$

where f^{PK} is the PK force projected onto the glide plane of the dislocation. When A = 0, the dislocations obey the classic result: their separation vector forms an angle $\pi/4$ with respect to their glide planes [2,85]. When 0 < A/B < 1, the mechanically stable positions remain the same while their basins of attraction change. When A/B > 1, the right equilibrium point moves out beyond the $\pi/4$ angle. When $A/B \rightarrow \infty$, the rightmost basin becomes unstable. When the Burgers vectors are not parallel, the dislocation interactions are nonreciprocal in the sense of being nonmutual: their forces are not equal and opposite (SM [62], Sec. S3 B).

Dislocation motion from PK forces.—While the continuum theory provides insights at long length scales, whether and in what directions dislocations actually move depends on microscopic details. To illustrate this, we perform overdamped molecular dynamics simulations of particles interacting with a radial force $F^{\parallel}(r)$ given by a Lennard-Jones (LJ) force and three different realizations of the transverse force $F^{\perp}(r)$ in Eq. (2), see Figs. 2(a) and 2(b) and Movie S1 in the SM [62].

First, consider a transverse interaction $F_{LI}^{\perp}(r)$, which like F^{\parallel} , is an LJ force: a dislocation introduced to the center remains stationary as in a passive crystal. The reason is that the total force on any particle is simply a rotation of the force due to F^{\parallel} . Hence, the static configuration guaranteed by energy minimization for F^{\parallel} remains static when F_{LJ}^{\perp} is introduced. Next, we introduce F_{Lub}^{\perp} , which is a monotonically decreasing function of a single sign shown in Fig. 2(a), generically representative of hydrodynamic, lubrication, and frictional forces between self-spinning particles [29,34,72,76,77]. For F_{Lub}^{\perp} , the dislocation travels at a near constant speed to the left. Since F_{Lub}^{\perp} is nonzero at the first neighbor shell, it produces an ambient torque density τ_0 . Upon setting $\sigma_{ij}^{(\text{pre})} = \tau_0 \epsilon_{ij}$, the direction of dislocation motion follows the standard PK force expression $f_i^{\text{PK}} = \epsilon_{ij}\sigma_{jk}^{(\text{pre})}b_k = -\tau_0 b_i$, in agreement with the experiments and analysis of spinning-colloid crystals in Refs. [28,30].

Dislocation self-propulsion from microscopic work cycles at their cores.-Now, we show that not all mechanisms of dislocation propulsion can be captured by continuum considerations. The continuum PK force is derived from a coarse-grained approximation to the work done during dislocation motion. However, when a potential is not well defined, contributions to δW from short length scales need not average out during dislocation motion. In some cases, they can even overcome continuum predictions. To illustrate this, we use, as a probe, the force $F_{\delta}^{\perp}(r)$ narrowly peaked at a tunable interparticle distance $r = \delta$ and with the same sign as F_{Lub}^{\perp} . However, when the peak δ lies half way between the first and second neighbor shells, the dislocation now travels to the right, the opposite direction of F_{Lub}^{\perp} [see Figs. 2(a) and 2(b)]. Since the force F_{δ}^{\perp} is negligible at the first and second neighbor shells, the odd moduli A and K^o , as well as the ambient torque density τ_0 , are vanishingly small. This suggests that the underlying mechanism of dislocation motility evades the standard continuum explanation in terms of PK forces provided in the previous paragraph.

As we now show, this dislocation motility is a form of self-propulsion associated with microscopic work cycles acting at dislocation cores. First, we highlight all the bonds



FIG. 2. Dislocations self propel via active work cycles at their cores. (a) Three transverse interactions F_{LJ}^{\perp} (purple), F_{Lub}^{\perp} (teal), and F_{δ}^{\perp} (orange), with the neighbor shells highlighted by grey lines. Inset: A hexagonal lattice with first and second neighbor shells highlighted. (b) Particles are arranged in a free floating circular cluster with a single dislocation located at the center, and the dislocation position is tracked as a function of time. Simulations are performed with clusters of radius R = 50(dashed) and R = 100 (solid). See Supplemental Material (SM) [62], Movie S1. (c) Bonds crossing the glide plane of a dislocation are highlighted. Hue indicates the position of the bond in real space (blue: left, red: right). Opacity indicates the length of the bond (nearest neighbors darkest). (d) The highlighted bonds are plotted with their bases aligned. As the dislocation moves one unit cell to the right, the tops of the bonds trace out a contour C(black dashed). The gray arrows depict the interaction force field. See SM [62], Movie S2. (e) The interaction F_{δ}^{\perp} is varied by changing the location δ of its peak (pink: smaller δ , green: larger δ). For each value of δ , the position of the dislocation is tracked as a function of time. (f) The magnitude of the Peach-Koehler force $f^{\rm PK}$ and the active core force $f^{\rm core}$ as a function of the peak position δ . The vertical lines represent the values of δ used in the simulation. The direction change of the dislocation motion coincides with the crossover between f^{core} and f^{PK} .

that straddle the glide plane [Fig. 2(c)] and align their bases so that they are viewed in the space of their relative coordinates (r_x, r_y) [Fig. 2(d)]. Crucially, as the dislocation moves by a single unit cell, each highlighted bond assumes the position of its neighbor to the right. Next, we concatenate all the individual bond trajectories into a single contour C (dashed line) that begins at $r_x = -\infty$ and ends at $r_x = \infty$. The total work done when each of the bonds moves a short distance is then equivalent to that of a single bond traveling the entire contour, cf. Fig. 1(a). Notice that, if the force falls off faster than 1/r, then the contour may be closed in the upper half plane. Similar to the single-bond cycle shown in Fig. 1(a), the corresponding work W_{glide} reads

$$W_{\text{glide}} \approx \oint_{\mathcal{C}} \mathbf{F} \cdot d\mathbf{r} = \int_{\mathcal{A}} \nabla \times \mathbf{F} \, d^2 r,$$
 (12)

where A is the upper half plane enclosed by C (SM [62], Sec. S4 and Movie S2).

Since W_{glide} is associated with motion through one lattice spacing, the corresponding force on the dislocation reads $f^{\text{core}} = (1/a)W_{\text{glide}}$, and it is directed along the glide plane. In principle, the detailed shape of C depends on protocol and dynamics. However, a useful first approximation is to take C to be the line at $r_y = (\sqrt{3}/2)a$, giving

$$f^{\text{core}} = \frac{1}{a} W_{\text{glide}} \approx \frac{1}{a} \int_{-\infty}^{\infty} F_x^{\perp} dr_x \bigg|_{r_y = \frac{\sqrt{3}}{2}a}.$$
 (13)

In Fig. 2(e), we perform simulations with F_{δ}^{\perp} , but we vary the parameter δ , which sets the location of the central peak. At small δ , there is significant overlap between F_{δ}^{\perp} and the first neighbor shell, giving rise to a large ambient torque density $\tau_0 \approx \sqrt{3}F^{\perp}(a)/a$ and the corresponding PK force f^{PK} . Figure 2(f) shows the relative magnitudes of f^{core} and f^{PK} as a function of δ . The crossover in dominant force coincides with the sign reversal in dislocation speed corroborating our theoretical derivation of f^{core} . While the sign reversal is a dramatic effect that occurs under specific conditions, f^{core} is generically present for all nonconservative microscopic interaction forces. Solids whose microscopic interactions violate Newton's third law can also display spontaneous dislocation motion [28].

Active plasticity.-Finally, we examine the effects of odd elasticity on plastic deformation. In Fig. 3(a), we perform a simple uniaxial compression of a solid interacting via a transverse lubrication force (see, also, SM [62], Movies S3 and S4). Before the first dislocation nucleates, odd elasticity biases the stress distribution [Figs. 3(a) and 3(d)]. At the end of the compression, the permanently deformed shape of the beam breaks all mirror symmetries [Figs. 3(b) and 3(e)]. The change in final shape arises because the biased stress distribution favors dislocation nucleation from the upper-right- and lower-left-hand corners [Figs. 3(c) and 3(f)]. Empirically, we find that introducing transverse forces generally lowers the plastic yield strain at which the first dislocation nucleates [Fig. 3(g)]. In Fig. 3(h), we consider a single dislocation in the center of a disk. In a passive medium, a uniform compression induces an isotropic stress $-B(\delta \rho / \rho) \delta_{ij}$ with an associated $f_i^{\text{PK}} =$ $-B(\delta\rho/\rho)\epsilon_{ij}b_j$ in the climb direction. This typically results



FIG. 3. Active plasticity in odd elastic media. (a)–(c) The compression of a solid beam with a standard LJ interaction of strength e^{\parallel} . (d)–(f) The same numerical experiment with the addition of transverse lubrication forces of magnitude e^{\perp} . See SM [62], Movies S3–S4. (a),(d) show the per-particle stress, resolved on the [11] and [11] glide planes and summed, immediately prior to the first dislocation nucleation. (b),(e) After significant plastic deformation, we color particles by the cumulative number of neighbor changes in their first coordination shell (N_{swap}). (c),(f) N_{swap} averaged in the vicinity of the lower-left- and lower-right-hand corners as a function of strain. The strain is $\Delta h/h$ where *h* is the height of the beam. (g) The critical strain at first nucleation for the bottom left-hand (green) and righthand (pink) corners. (h) A disk of particles interacting via radial and transverse LJ forces is subject to compression (negative dilation). At higher compression, a torque density is induced throughout the cluster which drives dislocation motion via the PK force. See the SM [62], Sec. S5 for additional simulation details.

in no motion or defect splitting. The odd elastic solid in Fig. 3(h) features a F_{LJ}^{\perp} which induces no dislocation motion in the absence of external stresses [recall Fig. 2(b)]. However, due to the odd modulus *A*, an area change gives rise to a torque density $\tau = A(\delta \rho / \rho)$, which, in turn, promotes motion along the glide plane via $f_i^{PK} = -\tau b_i$.

To summarize, we studied how defect strains, interactions, and motility are modified in systems for which the interactions are more general than standard pairwise, potential forces.

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