Examining the Mechanical Equilibrium of Microscopic Stresses in Molecular Simulations

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The microscopic stress field provides a unique connection between atomistic simulations and mechanics at the nanoscale. However, its definition remains ambiguous. Rather than a mere theoretical preoccupation, we show that this fact acutely manifests itself in local stress calculations of defective graphene, lipid bilayers, and fibrous proteins. We find that popular definitions of the microscopic stress violate the continuum statements of mechanical equilibrium, and we propose an unambiguous and physically sound definition.

The stress $\sigma(x)$ is a second-rank tensor field encoding the internal force distribution in a continuum system; when multiplied by a unit vector $n$, it provides the forces per unit area in the material across a surface passing through $x$ and perpendicular to $n$. The continuum stress field can be connected to the statistical mechanics of discrete particle systems, in what is called the microscopic stress. Today, the microscopic stress is increasingly used to recapitulate mechanical information contained in long molecular dynamics (MD) trajectories of nonuniform systems, or to connect molecular details with continuum physics at larger scales. Significant applications include defective bulk [1,2] and two-dimensional crystals [3,4], biomolecular assemblies such as lipid bilayers [5–7], membrane proteins [8,9], and even isolated molecules [10].

The average (or virial) stress of a periodic system can be uniquely defined and given a precise thermodynamical interpretation [11]. There are, however, multiple procedures to map a statistical mechanics ensemble into a stress field. This indeterminacy is expected since any divergence-free symmetric tensor field (self-equilibrated) can be added to the local stress without affecting the continuum statements of balance of linear and angular momentum for a system in equilibrium [12,13],

$$
\begin{align*}
\sigma_{ij,j} &= 0, \quad (1a) \\
\sigma_{ij} &= \sigma_{ji}, \quad (1b)
\end{align*}
$$

where we have ignored externally applied force and torque densities. The ambiguity in the microscopic stress is widely appreciated theoretically, but its practical consequences have been largely overlooked, partly because different definitions coincide for simple force fields [7]. As molecular simulations model increasingly complex systems, the issue of whether different definitions of the microscopic stress satisfy the fundamental requirements of mechanical equilibrium in Eq. (1) has not been systematically examined.

Here, we simulate the equilibrium behavior of three important condensed matter systems—defective graphene, lipid bilayers, and coiled-coil fibrous proteins—using MD simulations with realistic atomistic force fields, and study the significance of the definition of the microscopic stress. We consider the atomic virial stress [3,4,14] and several flavors of the Irving-Kirkwood stress [7,8,15–19]. Strikingly, we find that the atomic virial stress does not satisfy balance of linear momentum. Furthermore, we find that a widely used version of the Irving-Kirkwood stress [15] does not satisfy balance of angular momentum for systems with chiral constituents. Our results favor a recent canonical definition of the Irving-Kirkwood stress [16,17], which, however, is limited to multibody potentials with at most four-body interactions. Motivated by higher-body force fields used in materials science and biochemistry [20,21], we develop and test a new and general procedure [22] to uniquely obtain a physically meaningful microscopic stress.

Microscopic stress fields include a kinetic contribution $\sigma_K$, describing the flux of momentum due to internal vibrations, and a potential contribution $\sigma_V$, accounting for the interatomic forces. Since the difficulty in the microscopic stress definition stems from the potential part, we focus only on this contribution and refer to [23] for the expression of $\sigma_K$. For solid systems, where atoms stay relatively immobile, it is common to resort to the atomic virial stress definition [14], which is conceptually and computationally simpler than other definitions. In this method, the potential component is computed by distributing the virial of each potential contribution equally among the particles involved. More specifically, let the potential of the system be described additively as $V = \sum_{i=1}^M V_i$, with each contribution involving $n_i$ particles. Then, we have

$$
\sigma_V = \frac{1}{\Omega_\alpha} \sum_{i \in I^\alpha} \frac{1}{n_i} \left( \sum_{\beta} \frac{\partial V_i}{\partial r^\beta} \otimes r^\beta \right), \quad (2)
$$

where $r^\beta$ is the position of particle $\beta$, $I^\alpha$ collects the set of potential contributions involving particle $\alpha$, and $\Omega_\alpha$ is the volume of this particle.

An alternative definition of the microscopic stress with a more solid statistical mechanics foundation was pioneered by Irving and Kirkwood [34,35] for two-body potentials. This approach, which we label as IK, defines the stress at
each point in space, $\sigma(x)$, rather than at the particles, and is thus more popular in fluid systems. The IK microscopic stress is identified by invoking Eq. (1a), which is therefore satisfied ab initio [34,35]. The potential component at position $x$ is given by

$$\sigma_V(x) = \left\langle \sum_{\alpha, \beta > \alpha} f_{\alpha\beta} @ r_{\alpha\beta} B(r', r, x) \right\rangle,$$

where $r_{\alpha\beta} = r_\beta - r_\alpha$, $B(r', r, x) = \int_0^1 \delta((1-s)r_\alpha + s r_\beta - x) ds$ is the bond function [35], and $\delta(r'-x)$ is the Dirac delta distribution centered at $x$. For two-body interactions $V_I = V_{\alpha\beta}(r_{\alpha\beta})$, we have $f_{\alpha\beta} = V_{\alpha\beta} r_{\alpha\beta} / r_{\alpha\beta}$, where $\rho_{\alpha\beta} = |r_{\alpha\beta}|$. In practice, space is subdivided into a grid and the microscopic stress is evaluated at discrete positions [7].

There have been a number of extensions of the IK framework to deal with multibody potentials [12,15,16,18,19]. Some of these approaches fully retain balance of linear momentum by construction [12,15,16,18], by defining $f_{\alpha\beta}$ as the terms of a pairwise force decomposition $F_{\alpha\beta} = \sum_{\beta} f_{\alpha\beta} \text{ satisfying } f_{\alpha\beta} = -f_{\beta\alpha}$, with $F_{\alpha\beta}$ being the total force acting on particle $\alpha$. The indeterminacy of the local stress tensor manifests itself in the IK definition because the pairwise force decomposition is not unique for potentials involving three or more particles. We consider initially the so-called central force decomposition (CFD) [13,16] and defer further discussion to later in this Letter.

We first consider a periodic graphene sheet with a Stone-Wales defect as a model system to compare the atomic virial and the IK stresses. The system is modeled with the force field described in [36] involving up to four-body interactions and simulated in an NVT ensemble at 300 K [23]. We compute microscopic stresses here and elsewhere in the Letter with a freely available implementation [7,37]. Figure 1(a) highlights the fundamental features of each stress definition. The atomic virial stress is defined discretely at the atoms, while the IK stress is continuous but exhibits marked concentrations along lines joining atomic positions, a signature of the force decomposition. It is often convenient to spatially average the discrete features in these fields with weighting functions such as Gaussian kernels [13,23,38]. Figure 1(b) shows that the spatially averaged fields $\sigma$ according to both notions of stress are qualitatively similar, although the atomic virial stress exhibits smaller magnitudes.

We analyze next whether these fields are in equilibrium, as physically expected. Importantly, it can be shown by a simple calculation that $\sigma_{ij}(x)$ is the spatial average of $\sigma_{ij}(y)$ [23]. Therefore, $\sigma_{ij}$ should be zero if the corresponding microscopic stress $\sigma_{ij}$ obeys Eq. (1a). Strikingly, the atomic virial stress is strongly out of equilibrium in the vicinity of the defect, Fig. 1(c). In contrast, the IK stress field exhibits nearly zero divergence (except from deviations due to discretization and limited statistical sampling). Thus, our results suggest that the atomic virial stress should be employed only for visualization purposes.

We focus next on the various extensions of the IK stress for multibody interactions. A generalization of the method of planes has been proposed by Heinz, Paul, and Binder (HPB) [18,19], which is consistent with the IK stress for two-body potentials and also recovers the global virial stress. We find that this method does not exactly satisfy balance of linear momentum, albeit with a smaller error than the atomic virial stress [23]. We examine next other IK generalizations satisfying Eq. (1a) by construction. A natural force decomposition satisfying $f_{\alpha\beta} = -f_{\beta\alpha}$ was proposed by Goetz and Lipowsky [15],

$$f_{GLD}^{\alpha\beta} = \sum_{I=1}^M \frac{1}{n_I} \left( \frac{\partial V_I}{\partial r} - \frac{\partial V_I}{\partial r^\prime} \right).$$

This Goetz-Lipowsky decomposition (GLD) has been widely employed to analyze MD simulations ([5,7,8] and references therein), yet it produces noncentral forces; i.e., in general $f_{GLD}^{\alpha\beta}$ is not parallel to $r_{\alpha\beta}$. In principle, this could lead to nonsymmetric stresses [12,16]. Recently, a central force decomposition has been proposed by Admal and Tadmor [13,16], which produces a symmetric stress tensor by construction because $f_{\alpha\beta}$ is defined such that it is always parallel to $r_{\alpha\beta}$, see Eq. (3). By invariance with respect to rigid body transformations, the additive potential contributions can be represented in terms of interatomic distances, $V_I(r_{12}^{\prime}, \ldots, r_{n_I}^{\prime}) = V_I(r_1, \ldots, r_n)$, leading to the CFD pairwise forces

$$f_{CFD}^{\alpha\beta} = \sum_{I=1}^M \frac{\partial V_I}{\partial r} r_{\alpha\beta}^\prime.$$

The CFD and GLD force decompositions result in pairwise forces with large differences in magnitude and direction [23]. To explore the features of each force decomposition, we consider a lipid bilayer system of fluid DPPC.
(1,2-dipalmitoyl-sn-glycero-3-phosphocholine), which is isotropic in the membrane plane \((x - y)\) at the simulated temperature \((323 \, \text{K})\). In addition to the conventional profiles of all stress components across the thickness, we adopt an unconventional but intuitive method to visualize stress through the traction vector, \(\mathbf{t} = \sigma \cdot \mathbf{n}\), on a given internal surface with unit normal \(\mathbf{n}\) (Fig. 2). The traction can be decomposed into a normal and a tangential component, \(\mathbf{t} = t_n \mathbf{n} + \mathbf{t}_\tau\). Here, we consider a test cylinder, and represent the normal traction \(t_n\) as a color map and the tangential component \(\mathbf{t}_\tau\) using arrows.

In agreement with the symmetries of this system and the fluidity of the bilayer, the CFD stress is diagonal with equal lateral components \((\sigma_{xx} = \sigma_{yy})\) and a normal constant component across the bilayer \(\sigma_{zz}(z) = \text{const.}\) as dictated by Eq. (1a) [Fig. 2(a)]. We note that for a bilayer in the gel phase, the off-diagonal components could be nonzero but should nevertheless respect the symmetry of \(\sigma\). In sharp contrast, the GLD stress exhibits nonzero in-plane off-diagonal components, which are antisymmetric \((\sigma_{xy} = -\sigma_{yx})\) and of significant magnitude [Fig. 2(b)], hence violating Eq. (1b). Furthermore, CFD and GLD produce significantly different lateral components, and, hence, normal tractions \((t_n = \sigma_{xx} = \sigma_{yy})\) [Fig. 2(c)].

Focusing on the tangential traction, we note that \(\mathbf{t}_\tau\) is parallel to the bilayer plane with sense and magnitude given by \(\mathbf{t}_\tau = \mathbf{n} \times \mathbf{r}\). As expected, for CFD \(\mathbf{t}_\tau\) is nearly zero. For GLD, however, it is clear from Fig. 2(c) that the non-symmetry of the stress tensor introduces distributed torques of opposite sign in each leaflet of the bilayer, since \(\sigma_{xy}(z) = -\sigma_{yx}(z)\). We hypothesize that such behavior may be due to the internal structure of each lipid, since the headgroup portion of DPPC contains a chiral carbon. We test this hypothesis by comparing the stress tensors for three systems with different mixtures of the two DPPC enantiomers (L-DPPC and D-DPPC). Consistent with this hypothesis, the torques induced in each monolayer according to GLD adopt the same sign for a system with one monolayer composed solely of L-DPPC and the second monolayer composed solely of D-DPPC [Fig. 2(d)]. Mixing equal numbers of each chiral lipid in both monolayers results in nearly zero distributed torques according to the GLD stress. Thus, the off-diagonal components of the GLD stress tensor reflect the average chirality of the molecular composition. In contrast, we find that the CFD stress tensor is essentially unchanged by the lipid chirality [23].

To physically interpret the GLD distributed torques, it is necessary to resort to an extended theory of continuum mechanics. In micropolar continuum theories, these torques can be balanced locally by invoking a couple stress field \(\mathbf{m}\), which in equilibrium satisfies \(\sigma_{ij,j} = \nabla m^k\), where \(\epsilon_{ijk}\) is the Levi-Civita symbol [12,39–41]. In our situation, however, there is no compelling physical justification for this field since the primary objects of our model are achiral point particles [41] and there is no apparent external source for \(\mathbf{m}\). Thus, although the connection between the non-symmetry of the IK-GLD stress and molecular chirality is very appealing, this example undermines its mechanical interpretation. The HPB stress [18,19] produces nonsymmetric stresses very similar to GLD for this system [23].

The microscopic stress tensor not only serves as a tool to explore the local distribution of forces, but can also provide important material properties. For instance, the Gaussian curvature elastic modulus of lipid bilayers can be computed as \(\kappa = \int (\sigma_{xx} + \sigma_{yy})/2 - \sigma_{zz} z^2 dz\), which is highly sensitive to the features of the stress profile (see [6] and references therein). For the three bilayer systems with different chiralities in Fig. 2, we obtain \(\kappa_{\text{CFD}} = (-6.4, -6.7, -6.1) \times 10^{20} \, \text{J}\), in agreement with the common estimates of \(\kappa\) in the order of the negative of the bending modulus \(-5–15 \times 10^{20} \, \text{J}\) [42]. Strikingly, we find \(\kappa_{\text{GLD}} = (0.91, 0.57, 1.3) \times 10^{20} \, \text{J}\), with the wrong sign—suggesting that a DPPC bilayer would be unstable [43]—and widely varying magnitudes.

Taken together, these results show that the choice of microscopic stress definition is not a mere theoretical preoccupation. Our results strongly favor the IK-CFD definition, which, unlike the atomic virial or the IK-GLD stresses, identically satisfies Eq. (1) for a system in equilibrium. However, CFD is not uniquely defined when \(n_1 > 4\). The geometric reason behind this ambiguity is that the \(n_1(n_1 + 1)/2\) interatomic distances \((r_{12}, \ldots, r_{n_1^{-1}, n_1})\)
involved in a given multibody potential $V_I$ cannot be arbitrarily chosen in $D_I = \mathbb{R}^{n_I^2/2}_{I}$. There are geometric conditions that make these distances realizable by a system of $n_I$ particles, which define the so-called shape space $S_I \subset D_I$. When $n_I > 4$, the dimension of the manifold $S_I$ is smaller than $n_I(n_I + 1)/2$, and, therefore, the differential calculus involved in Eq. (5) needs to be carefully considered [44,45]. More practically, when $n_I > 4$ there are infinitely many different ways to express the potential, $V_I$, in terms of interatomic distances, each resulting in a different force decomposition and microscopic stress [13,16].

In the spirit of [46–48], we propose an alternative thermodynamic derivation of the IK microscopic stress, which naturally and unambiguously extends CFD to multibody potentials. In analogy to the Doyle-Ericksen equation of continuum mechanics [49,50], the stress tensor can be defined from covariance arguments as

$$
\sigma(x) = \frac{2}{\sqrt{|g(x)|}} \frac{\delta A}{\delta g(x)},
$$

where $g(x)$ is the Jacobian determinant of the metric and $\delta A$ is the variation of the canonical free energy with respect to an infinitesimal change of metric $\delta g(x)$ resulting from a change of coordinates. As fully detailed in [22], this variational formalism identifies the covariant central force decomposition (cCFD)

$$
f_{\text{cCFD}} = \sum_{I=1}^{M} (\nabla S_I \hat{V}_I)_{\text{cCFD}} r_{\text{cCFD}},
$$

where $\nabla S_I \hat{V}_I$ is the covariant derivative of the potential along the shape space $S_I$. For four- or fewer-body potentials, cCFD and CFD in Eq. (5) coincide. However, cCFD circumvents the main limitation of CFD by providing a unique expression for potentials with any number of particles. In practice, $(\nabla S_I \hat{V}_I)_{\text{cCFD}}$ can be computed by projecting $\partial \hat{V}_I / \partial r_{\text{cCFD}}$ for an arbitrary extension onto $S_I$ [23]. Through a different rationale, the projection of the CFD onto the shape space has been recently discussed in [51]. Our assumption that the potential is additively decomposed into a many-body expansion is appropriate for most classical force fields. For semiempirical methods based on density functional theory concepts, such as the embedded-atom model [20], this additive structure is not apparent. We refer the interested reader to [22,51] for further discussion.

We test the cCFD microscopic stress by considering a coiled-coil structural protein, composed of two identical α-helical chains that wrap around each other to form a super-helix. The coiled-coil structure is a double “zipper,” with an inner core of intercalating hydrophobic amino acids that are flanked by opposing negatively and positively charged amino acids, Fig. 3(a). We model this system as an infinitely long periodic molecule with a widely used protein force field (CHARMM22/CMAP) [21,52] involving up to five-body interactions. We compare the tractions on the surface of the coiled-coil protein, essentially exerted by the solvent, calculated with GLD, cCFD, and another seemingly reasonable way to fix the indeterminacy of CFD (by minimizing the norm of the force decomposition) that we call nCFD [23]. We find that for GLD and cCFD, $t_n$ exhibits a similar pattern that follows the left-handed helical structure, Figs. 3(b) and 3(c). The zippered interface between the two chains is dominated by outward tractions (red), which transition to inward tractions (blue) at the periphery of the protein. In contrast, nCFD produces spurious maps of $t_n$ [Fig. 3(d)], highlighting the need for a physically meaningful method to fix the gauge freedom of CFD [40]. Focusing on $\tau$, cCFD presents tractions that locally equilibrate and do not produce net forces. GLD, however, produces a predominantly leftward traction field following the twist of the protein as a result of the non-symmetric components of $\sigma$. Quantitatively, the GLD tractions produce a net force per unit protein length along its axis of 34 mN/m, whereas for cCFD we have 0.35 mN/m. Thus, the GLD stress again contains information about chirality but produces tractions that cannot be physically balanced in our periodic system, which does not undergo any translation or rotation during the course of the simulation.

In summary, the ambiguity of the microscopic stress acutely emerges when analyzing complex materials and biomolecular assemblies. Strikingly, the widely used atomic virial stress does not satisfy balance of linear momentum, while a popular version of the Irving-Kirkwood stress does not satisfy balance of angular momentum as a result of molecular chirality. In contrast, Irving-Kirkwood stress fields based on a covariant central force decomposition can be
rationally derived irrespective of the many-body nature of the potential and satisfy by construction mechanical equilibrium in Eq. (1). This definition of microscopic stress thus provides a solid footing to systematically connect MD simulations and the mechanical behavior of materials at the nanoscale [6,53].

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