Forced Unfolding of Proteins Within Cells

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To identify cytoskeletal proteins that change conformation or assembly within stressed cells, in situ labeling of sterically shielded cysteines with fluorophores was analyzed by fluorescence imaging, quantitative mass spectrometry, and sequential two-dye labeling. Within red blood cells, shotgun labeling showed that shielded cysteines in the two isoforms of the cytoskeletal protein spectrin were increasingly labeled as a function of shear stress and time, indicative of forced unfolding of specific domains. Within mesenchymal stem cells—as a prototypical adherent cell—nonmuscle myosin IIa and vimentin are just two of the cytoskeletal proteins identified that show differential labeling in tensed versus drug-relaxed cells. Cysteine labeling of proteins within live cells can thus be used to fluorescently map out sites of molecular-scale deformation, and the results also suggest means to colocalize signaling events such as phosphorylation with forced unfolding.

Force-induced changes in protein conformation have long been postulated to contribute to the deformability of cells (1, 2). Likewise, in cell adhesion, forces of pico-Newton magnitude that result from cells pulling on matrix (3) are believed to induce conformational changes that initiate essential anchorase signals (4–8). Single-molecule measurements indeed show that domain unfolding occurs in reversible extension of purified cytoskeletal motor, and matrix adhesion proteins (9–12), and simulations of the molecular dynamics of protein extension have helped to clarify mechanisms (13–15). Direct cell-level evidence is lacking or even contrary to forced unfolding (16), although cytoskeletal association of a large and rare conformation-sensitive antibody has suggested extension of a proline-rich region in one protein within spread, fixed cells (17). The more broadly directed “shotgun” approach here is applied to live cells under physiological stresses and exploits small thiol-reactive probes that permanently label force-sensitive domains.

Cysteine (Cys) is a moderately hydrophobic amino acid that is frequently shielded by tertiary or quaternary protein structure. Labeling of cysteine’s SH moiety has been exploited in solution denaturation studies on a few small purified proteins (18, 19), as well as in an anemia-causing proline mutation in the red blood cell (RBC) protein spectrin (20). In addition, forced unfolding of single proteins with core-sequestered disulfides demonstrates reduction of the S-S within seconds by reactive thiols in the medium (21, 22). We show here, in intact cells, that force-induced changes in protein structure can also expose—for relatively rapid reaction—specific buried Cys (Fig. 1) in a number of key cytoskeletal proteins. Sequential n-dye-labeling with different color fluorophores (n = 2 here) proves to be a facile approach to amplifying signals from shielded sites relative to prelabelled surface sites (23). We illustrate the range of this in situ “Cys shotgun” approach, first, with the relatively simple human RBC, which allows for the most direct demonstration of forced unfolding in fluid-stressed cells, then, with human mesenchymal stem cells (MSCs) under cell-generated tension.

Red blood cells deform under the incessant stresses of blood flow, and the spectrin membrane cytoskeleton has proven central to cell deformability (24). Spectrin’s α and β chains interact to cross-link F-actin in this cell, and in single-molecule studies, spectrin’s tandem array of helical bundle domains (Fig. 1) are found to unfold at low forces (~picoNewton (25–27)) that could be generated by just a few myosin motor molecules. There are 20 Cys in α-spectrin and 15 Cys in β-spectrin, and some of these appear buried in crystal structures (Fig. 1) and homology models. To assess exposure of Cys in unfolding of spectrin and all of the other RBC membrane proteins, cells were reversely quenched to make hemoglobin-depleted pink “ghosts” that were resealed with entrapment of the Cys-reactive fluorescent dye IAEODANS. Dye-loaded cells were then either held static in suspension at various temperatures or else sheared over a physiological range of stresses with a standard fluid shearing device. After 5 min or more, cells were released, excess dye was quenched, and cells were imaged to assess membrane labeling (Fig. 2A).

Solubilized cells were denatured, and all Cys that were not dye-labeled were alkylated with iodoacetamide (IAM). Separation of membrane proteins by one-dimensional (1D) SDS–polyacrylamide gel electrophoresis (SDS-PAGE), followed by densitometry, showed 50% more IAEODANS fluorescence in the bands of α- and β-spectrin from the shear samples (Fig. 2A: 15 runs at 37°C); sequential two-dye labeling magnifies this difference to >500%, as described below. Labeling under shear is only enhanced for spectrin: Labeling of the other major membrane proteins (ankyrin, protein 4.1, actin, etc.)

Fig. 1. Force-induced changes in protein structure within cells are hypothesized to expose novel binding sites for ligands. This example of a molecular dynamics simulation shows that Cys1167 in β-spectrin exposes 0 Å² surface area (of 224 Å²) until forced extension [e.g., (15)] exposes the –SH for reaction with a thiol-reactive fluorescent dye.
Liquid chromatography–coupled tandem MS (LC-MS/MS) was used to identify and quantify IAEDANS-modified Cys sites in spectrin bands after excision and trypsinization. Within α- and β-spectrin, respectively, 13 and 14 IAEDANS-Cys were detected (table S1). Because surface-exposed Cys are predominantly IAEDANS labeled and buried Cys are largely inaccessible until subsequent denaturation and IAM labeling, site-specific ratios of (IAEDANS/IAM) in ion chromatogram elution profiles (e.g., Fig. 2B) quantify Cys exposure. A majority of ratios are the same for shear and static samples, but at least six Cys sites are distinct with results for the ratio of dye-labeling rates over time $\phi(t) = ([\text{IAEDANS}/\text{IAM}]_\text{shear}/[\text{IAEDANS}/\text{IAM}]_\text{static})$ that are typically in the range of $\phi \approx 5$ to 10 at 60 min, but as high as 36 (Fig. 2C). Cleavage patterns of β-spectrin treated with NTCB (2-nitro-5-thiocyanobenzoic acid), which cleaves peptide bonds at unlabeled Cys (fig. S1A), are distinct for shear and static samples, consistent with qualitative differences in Cys exposure.

The structure of shear-sensitive β-R8-9 (28) shows that shear-labeled Cys$^{1167}$ is 100% buried or shielded within a tertiary fold—at least until force induces localized unfolding (Fig. 1). A recombinant multidomain construct β-R5-9 was, therefore, Cys-labeled at different temperatures for a fixed time and then analyzed by both fluorescence imaging and MS. Construct labeling is significant only at temperature $T \geq 25^\circ\text{C}$ (Fig. 2D), with evidence from MS of Cys$^{1167}$ labeling at 25°C. This apparent transition corresponds to the transition midpoint temperature ($T_m$) of β-R9 (29) and correlates with both the thermally induced loss of helicity and the loss of mechanical stability of R9 as determined by atomic force microscopy (AFM) forced extension. At higher temperatures, additional repeats are labeled as helicity is lost.

The shear-sensitive Cys$^{1203}$ was also studied in solution with a recombinant α-R12 domain for which modeling predicts partial shielding (Fig. 2E). At a given temperature, the ratio of dye-labeling rates $\Phi(T)$ for urea-denaturing conditions versus native conditions relates to the steric protection of a partially exposed cysteine (19) (fig. S2). The recombinant’s $\Phi(37^\circ\text{C}) \approx 4$ approximates the LC-MS/MS result for shear versus static samples’ $\Phi(37^\circ\text{C}) = 5$ (Fig. 2C). Normalization and rescaling to the folded state maximum $\Phi_{\text{max}} = 7$ (Fig. 4D) gives the fraction folded, and this decreases with $T$ in parallel again with the helicity loss upon heating (Fig. 2E and fig. S2).

Within RBCs, the labeling kinetics for IAEDANS provides a basis for dramatically increasing the shear/static–labeling ratio. Surface-exposed Cys in spectrin label to saturation within 30 to 60 min with a first-order rate constant (0.13 min$^{-1}$) (Fig. 3A), whereas in the shear sample, the ~50% enhanced labeling is slower (0.04 min$^{-1}$), consistent with a second population of shear-exposed Cys sites. In sequential two-dye labeling, static-accessible sites are labeled to near saturation with IAEDANS (blue), and then shear is applied with labeling by a second (green) Cys-reactive fluorophore (Fig. 3B). N-(4,4-Difluoro-1,3,5,7-tetramethyl-4-bora-3a,4a-diaza-s-indacene-2-yl) iodoacetamide (BODIPY-IA) labeling of spectrin within RBC's increases with both time and stress, yielding results for the shear/static ratio of ~5 to 6.5 after 60 min at the highest shears of $\sigma = 0.8$ to 1.7 Pa. Site-specific MS gave $\Phi \approx 10$ (Fig. 2C).

The effective labeling rate $k$ for the secondary BODIPY-IA–labeling reaction is 30 times as fast in maximally sheared cells as in static cells (Fig. 3B) and fits to a force ($f$)-dependent Linderstrom-Lang (fLL) scheme in which shear stress, proportional to $f$, shifts the conformational equilibrium.

\[
N \times \frac{k_{\text{ff}}e^{2f/(\kappa a)T}}{k_{\text{ff}}e^{2f/(\kappa a)T} + k_{\text{ff}}e^{2f/(\kappa a)T}} \rightarrow U \frac{k_2}{k_1e^{2f/(\kappa a)T}} P
\]

Similar to forced unfolding in the cited single-molecule studies, the folded state $N$ shifts toward unfolded state $U$ with a force bias on the forward and reverse reactions, and the activated transition state $I^*$ along the reaction coordinate $x$ is at the respective distances $\Delta x > 0$, $N \rightarrow I^*$ and $\Delta x' < 0$, $U \rightarrow I^*$. Here, the $U$ state’s exposed Cys can be labeled by dye at concentration $c$ and rate $k_2$ to yield measured
product $P$ (table S1 and fig. S1B). The analytical solution to the fLL scheme (SOM text) fits well to the spectrin in-cell experiments here when we use parameter values similar to those obtained from single-molecule studies on spectrin, such as $k_{1f} \approx 0.002 \text{ min}^{-1}$ (25). The fits are also consistent with expectations that (i) the refolding rate is only important in the static sample and at low stress, (ii) labeling is limiting at very high stress, and (iii) labeling in the intermediate stress regime is sufficiently rapid that the rate of exposure of sequestered Cys is exponential with respect to stress.

Consistent with past speculations that spectrin unfolding occurs to some extent within RBCs (1, 30), about 40% of spectrin’s domains studied as isolated domains unfolded with $T_m \approx 37^\circ \text{C}$ (29). Within static cells, IAEDANS labeling of spectrin Cys occurs slowly at low temperature as expected (fig. S3A), and, although site-selective labeling rates will be needed for an accurate assessment of in-cell melting of spectrin, 60 min of labeling yields an apparent melting curve (fig. S3B) in close agreement with the summed melting profile calculated from individual recombinant spectrin domains (29). Moreover, at all temperatures up to thermal disruption of cells (~45°C), shear enhances labeling of spectrin but not actin (fig. S3).

Mechanically induced changes in protein structure within other cells are likely to be of importance not only to cell deformability but also to various mechano-signaling processes. MSCs, for example, are contractile (31) and strain the underlying matrix in differentiation (32). Traction force measurements on soft matrices show that MSCs, like many tissue cells (4), generate stresses (~kPa) that are ~1000 times fluid shear stresses imposed on RBCs. Inhibition of non-muscle myosin II (NMM II) with the drug blebbistatin (33) relaxes and softens MSCs (31) and also blocks differentiation. For molecular insight into structural changes, as well as a proteomic characterization of MSCs, the Cys sites were studied (fig. S3). Immunofluorescence imaging shows cytoskeletal labeling in the tensed state of MSCs for which polymerization does not greatly disturb MSCs (34). Cell lysates for the tensed cells and blebbistatin-relaxed cells were Cys-quenched, and, once again, 1D SDS-PAGE was followed by densitometry and then LC/MS/MS analyses of all prominently labeled bands (~40 kD) (fig. 4B). The cytoskeletal proteome of MSCs (table S2) appears typical of nucleated cells in consisting of actin, myosin, the actin-binding proteins filamin and nonerythroid spectrin, as well as tubulin and the mesenchymal intermediate filament protein vimentin. Site-specific differences in Cys labeling with blebbistatin are seen in several proteins, and two prove illustrative.

Nonmuscle myosin IIA shows greater mBBBr-labeling in the tensed state of MSCs for which immunofluorescence imaging shows cytoskeletal striation (Fig. 4C); in contrast, blebbistatin disorganizes NMM II, consistent with blocking it in a low-affinity state for actin (34). MS analysis identifies Cys 90 as a differentially labeled site, and homology models of NMM II show this site is buried between the head and rod domains. Transient allosteric changes that occur in stress generation during NMM II’s normal ATP hydrolysis likely expose Cys 90, whereas blebbistatin

![Fig. 3](image)

**Fig. 3.** Stress-dependent kinetics and sequential two-dye labeling for amplification. (A) Sheared ($\sigma = 0.93 \text{ Pa}$) versus static RBCs show distinct labeling kinetics for spectrin. IAEDANS labeling at 2.5 mM of the static-labeled, surface-exposed spectrin sites fit a first-order rate of 0.13 min$^{-1}$, the shear-exposed sites (66% more from fit) label more slowly at 0.04 min$^{-1}$. (B) Sequential labeling entails labeling of the surface Cys (blue) for 30 min, after which BODIPY-IA (green) is added at $>10$-fold excess (6 mM). Half of the cells are then exposed to shear with timed aliquots again quenched by IAM and analyzed by densitometry of the green fluorescent spectrin. BODIPY-spectrin at 60 min increases with $\sigma$ (top) as does the effective rate of labeling (bottom). Error bars (±SD) indicate three or more measurements (six for $\sigma = 0$). The data are fit by the exact solution of the fLL reaction for $P(t)$, either with or without refolding (SOM text). The intermediate regime of stress (dotted black line) is dominated by unfolding and increases exponentially with stress, exhibiting a characteristic stress of $\sigma_c = 0.5 \text{ Pa}$ and giving $k_{1f}^{unfolding} = 0.004 \text{ min}^{-1}$. (Inset) The effect of force on computed kinetics using the fit parameters.

![Fig. 4](image)

**Fig. 4.** In-cell labeling of human stem cells either tensed or relaxed. (A) The membrane-permeable Cys-reactive fluorophore, mBBBr, was added to 1-week cultures of MSCs with active myosins and tensed cytoskeletons and also to MSCs treated with myosin-inhibiting blebbistatin for 1 day to relax the cells. Imaging shows homogeneous labeling with 0.5 mM mBBBr for 40 min. (B) SDS-PAGE and densitometry of samples (±SD, three experiments) that were either blebbistatin treated (relaxed) or untreated (tensed) show several protein bands in which the fluorescence intensities are different (normalized to protein load). Lysates were quenched with $\beta$-mercaptoethanol (50 mM) before analysis. (C) Immunofluorescence imaging of NMM IIA in tensed cells (top) and relaxed cells (bottom) indicates a spatial redistribution of myosin with drug. Scale, 5 $\mu$m. MS analyses of excised myosin bands detected labeling of Cys 90, which appears buried within the fold of NMM IIA homology models. (D) Vimentin labeling in monomeric, and polymeric forms display different degrees of fluorescence (error bars from two experiments), which indicates that polymerization sterically blocks Cys 327 for labeling.
eliminates force-generation and keeps this site buried.

Differential labeling in MSCs is also found at the single Cys site of vimentin (Fig. 4D): Cys\(^{127}\) is in the central coil that mediates coiled-coil polymerization (35). Quaternary structure changes in solution have been exploited previously to understand hemoglobin tetramerization [e.g., (36)], as well as actin-binding interfaces [e.g., (37)], and here blebbistatin-induced depolymerization of vimentin with Cys exposure in the MSCs appears consistent with blebbistatin-induced softening of the cells (32). Purified vimentin confirms enhanced labeling as monomer (~50% more) versus polymer (Fig. 4D), and Western blot analysis plus immunofluorescence imaging of labeled proteins within cells [e.g., (38)] indicates no differences in overall expression. In-cell labeling thus identifies structural changes within tensed cell but not relaxing cells. Nucleated cells typhified by MSCs have a complex intracellular force map when inferred from cell tractions in the surrounding matrix (4, 32, 38), and even simple cells such as RBCs sustain stresses at the molecular level in unknown ways (1, 29). The overall Cys-shotgun methodology here not only provides evidence of force-induced changes that propagate in both tertiary and quaternary structures within cells but also, through LC-MS/MS, provides useful proteomic information, as well as new opportunities for fluorescence imaging. In addition, whereas fluorescence resonance energy-transfer imaging of labeled proteins within cells [e.g., (39)] might allow imaging of force-induced conformational changes in real time, the time-integrated analyses here are complementary and also readily extended to engineered sites in wild-type and mutant proteins for in-cell assessments of perturbations. This idea seems very likely to extend to coincidence detection of Cys labeling with posttranslational events such as phosphorylation so as to precisely colocalize conformational changes with signaling events.

References and Notes

23. Materials and methods are available as supporting material on Science Online.
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Supporting Online Material

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Materials and Methods
SOM Text
Figs. S1 to S4
Tables S1 and S2
References and Notes
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Monitoring of Blood Vessels and Tissues by a Population of Monocytes with Patrolling Behavior

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The cellular immune response to tissue damage and infection requires the recruitment of blood leukocytes. This process is mediated through a classical multistep mechanism, which involves transient rolling on the endothelium and recognition of inflammation followed by extravasation. We have shown, by direct examination of blood monocyte functions in vivo, that a subset of monocytes patrols healthy tissues through long-range crawling on the resting endothelium. This patrolling behavior depended on the integrin LFA-1 and the chemokine receptor CXCR1 and was required for rapid tissue invasion at the site of an infection by this “resident” monocyte population, which initiated an early immune response and differentiated into macrophages.

Mammalian monocytes consist of two main subsets of immune cells (1, 2), which arise from a common hematopoietic progenitor, the macrophage and dendritic cell (DC) precursor (MDP, or monoblast) that also gives rise to conventional resident spleen DCs (cDCs) and several tissue macrophage subsets (3, 4). So-called “inflammatory” monocytes express the cell surface protein Ly6c (Gr1\(^\text{+}\)), the chemokine receptor CCR2, and the adhesion molecule L-selectin and are selectively recruited to inflamed tissues and lymph nodes (1, 5). They are able to differentiate into inflammatory DC (1, 6, 7) and can replenish resident cell compartments in the skin, digestive tract, and lung (1, 8). The second subset of monocytes has been termed “resident” in mice (1, 2) because they were found in both resting and inflamed tissues, although their functions are still unknown. This subset is defined by a smaller size, high expression of the chemokine receptor CXCR1 and LFA-1 integrin, and by the lack of expression of Ly6c (Gr1\(^\text{+}\)), CCR2, and L-selectin (1, 2). Two monocyte subsets can also be identified in humans (9), with CD14\(^{+}\)CD16\(^{-}\) monocytes resembling mouse inflammatory monocytes and CD14\(^{+}\)CD16\(^{+}\) monocytes sharing a phenotype similar to that of mouse resident monocytes (1). Resident and