Mechanochemistry of Protein 4.1’s Spectrin–Actin-Binding Domain: Ternary Complex Interactions, Membrane Binding, Network Integration, Structural Strengthening

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Abstract. Mechanical strength of the red cell membrane is dependent on ternary interactions among the skeletal proteins, spectrin, actin, and protein 4.1. Protein 4.1’s spectrin–actin-binding (SAB) domain is specified by an alternatively spliced exon encoding 21 amino acid (aa) and a constitutive exon encoding 59 aa. A series of truncated SAB peptides were engineered to define the sequences involved in spectrin–actin interactions, and also membrane strength. Analysis of in vitro supramolecular assemblies showed that gelation activity of SAB peptides correlates with their ability to recruit a critical amount of spectrin into the complex to cross-link actin filaments. Also, several SAB peptides appeared to exhibit a weak, cooperative actin-binding activity which mapped to the first 26 residues of the constitutive 59 aa. Fluorescence-imaged microdeformation was used to show SAB peptide integration into the elastic skeletal network of spectrin, actin, and protein 4.1. In situ membrane-binding and membrane-strengthening abilities of the SAB peptides correlated with their in vitro gelation activity. The findings imply that sites for strong spectrin binding include both the alternative 21-aa cassette and a conserved region near the middle of the 59 aa. However, it is shown that only weak SAB affinity is necessary for physiologically relevant action. Alternatively spliced exons can thus translate into strong modulation of specific protein interactions, economizing protein function in the cell without, in and of themselves, imparting unique function.

The erythrocyte is among the simplest and best characterized of cellular systems within which the chemistry of composition can be intricately related to structural function. Indeed, most of the red cell’s mechanical behavior, except for limited aspects of cell shape, appears to depend little on the complexities of intracellular metabolism (14). However, to withstand the forces of blood flow over a several-month lifespan, the erythrocyte’s membrane components must assemble into a highly durable structure. Physical limits to this durability are set by the system’s mechanochemistry, which is defined here as those molecular interactions that permit a pliable, yet resilient structure.

Spectrin, actin, and protein 4.1 are the principal structural proteins of the thin, cohesive skeletal network that underlies the red cell’s lipid bilayer (3, 5, 19). The network formed by these highly conserved and pervasive proteins confers resilience and durability to the red cell’s composite membrane (14, 16, 36, 44, 46). Mixtures of these network proteins also show that they can form very stable complexes in vitro (37) which at approximately micromolar protein concentrations can undergo a phase transition to a three-dimensional gel (7, 15, 49). In the formation of the basic ternary complex, protein 4.1 appears to play a crucial role in stabilizing spectrin cross-links between actin filaments. Consistent with cross-link reinforcement, red cells deficient in protein 4.1 possess membranes that fragment more readily than normal when exposed to high fluid shear (46, 51).

The crucial stabilizing function of protein 4.1 (~80 kD) localizes to a relatively small (~10-kD) spectrin–actin-binding domain (SAB) (10) whose function recently has been shown to be strongly modulated at the genetic level by alternative splicing. The SAB isoform expressed in the erythrocyte is encoded by two exons that translate to an alternative NH2-terminal 21-amino acid (aa) “cassette”

1. Abbreviations used in this paper: aa, amino acid; FL-PE, fluorescein-phosphatidyethanolamine; GPL, glycosyolphosphatidylinositil; GST, glutathione-S-transferase; IAF, iodoacetimido-5-fluorescein; SAB, spectrin-actin-binding domain.
and a constitutive 59-aa domain (8, 25). Studies with recombinant isoform constructs of the SAB (11, 24) demonstrated that inclusion of the 21-aa cassette is critical for spectrin binding and also for efficient, 4.1-like activity in both ternary complex formation and membrane strengthening. Whether the 21-aa peptide, by itself, completely specifies these biochemical and biophysical properties was not addressed. Neither was it determined whether SAB isoforms lacking the 21aa are completely inactive, or simply much less efficient in some functions.

To dissect methodically such issues of protein functionality, COOH-terminal deletants of the SAB were constructed (see Fig. 1) and their functions characterized. We found that the 21 aa cassette plus 43 residues from the downstream sequence (peptide designation 21.43) are required for strong, 4.1-like interactions with spectrin and actin. Ternary complex formation in vitro, however, does not strictly require the 21 aa, but does need at least the first 26 residues of the downstream sequence in tandem with either the NH2-terminal 21-aa cassette (to give a functional 21.26 peptide) or additional residues from the 59-aa domain (yielding a functional 0.59 peptide).

The physiological relevance of these in vitro results to both binding and mechanically strengthening intact membranes was also addressed. SAB peptide binding to normal membranes shows a highly specific, spectrin-dependent interaction. Integration of SAB peptide into a solidlike cytoskeletal network beneath the red cell bilayer is visually demonstrated by fluorescence imaged microdeformation (FIMD) (12). The strengthening effect of each SAB peptide on 4.1-deficient membranes is assessed in parallel with membrane binding; as found in vitro, the 21.43 is the most potent and effective at micromolar concentrations, while both the 21.26 and 0.59 are capable of strengthening membranes, but only at much higher peptide concentrations. The 21.20 is completely nonfunctional, even up to concentrations of ~250 μM. Since both the 21.26 and the 0.59, but not the 21.20 are functional, the first 26 residues appear to constitute an actin-binding site, and sequences flanking either side of this site seem essential for spectrin interactions. It therefore appears that actin binding in combination with even weak spectrin binding is sufficient, if not materially efficient, for network integration, stabilization, and structural strengthening.

**Materials and Methods**

**Construction and Expression of SAB Peptides**

SAB domain deletion constructs (Fig. 1) were generated from a minigene encoding slightly more than the SAB domain of erythroid protein 4.1 (Glu180-Gly370) previously denoted 10k (11). Escherichia coli-expressed peptides were either affinity purified as glutathione-S-transferase (GST) fusion peptides or cleaved off the GST moiety by digestion with thrombin (20). Additional constructs to those in Fig. 1 included a 21.41, a 21.36, and a 21.31. A fusion peptide with the Xenopus laevis SAB (Mays, K. T., R. Winardi, L. Zon, N. Mohandas, and J. G. Conboy. 1992. Blood. 80:2725a.) was designated GST-XEN. Peptides were metabolically labeled by inducing the culture in a methionine-free minimal essential medium supplemented with [35S]methionine and Luria broth (15% vol/vol). Primary structures were partially confirmed by immunoblotting (11). NH2-terminal sequencing and mass spectrometry (Schischmanoff, P. O., unpublished data) further confirmed expression of the constructed sequences to within ~2 D. Peptide absorption spectra displayed the shapes expected (6) from the relative Tyr and Trp composition; spectrophotometry was therefore used to quantify protein concentration (31).

**Falling Ball Viscometry and Spectrin Composition of Networks**

The ability of recombinant peptides to form a supramolecular complex with spectrin and actin was assayed by falling ball viscometry (9). Purified spectrin tetramer (40, 50) and rabbit muscle F-actin (38) were mixed (at 1.8 and 14 μM, respectively) and the recombinant peptide was subsequently added (up to ~250 μM). The solution was drawn quickly into a 50-100-μl microcapillary and assayed by falling ball viscometry as described elsewhere (11). For composition of spectrin-actin peptide networks, protein mixtures were either prepared directly in ultramicrofuge tubes or transferred from a microcapillary. Centrifugation, sometimes through a sucrose cushion, at 80 k rpm, 22°C, for 7 min in a rotor (TLA-100; Beckman Instruments, Inc., Fullerton, CA) was followed by electrophoresis and densitometry (IS-1000; Alpha Innotech Corp., San Leandro, CA). Nearly all of the actin appeared in the pellets.

**Actin Filaments Cosedimentation Assay**

Fusion or thrombin-cleaved, 35S-labeled proteins were incubated in thick-walled, ultramicrofuge tubes with a mixture of spectrin (2 μM) and actin (7 μM) in parallel with a control tube having the same amount of peptide without spectrin-actin. The total volume was 50-100 μl, and included 50 μM BSA. After 1 h at 37°C, all tubes, up to 10 pairs, were centrifuged together as above. The supernatants and pellets were counted; the amount of bound peptide was typically determined from the difference in peptide concentrations in the supernatants of the sample and control tubes.

**Membrane-binding Method**

Typically, a small volume of in vitro translated 35S-labeled peptides was mixed with 400 μl of normal membranes. The membranes were obtained by simple hypotonic lysis (10 mM sodium phosphate, 4°C). The mixture of membranes plus peptides was incubated on ice for 20 min (47). To obtain resealed “ghosts” (cells with much reduced hemoglobin), salt was added (100 mM KCl, 1 mM MgCl2), and the membranes were incubated at 37°C for 30-60 min. After resealing, unbound peptide was washed away with isotonic PBS. The washed ghosts were then lysed a second time and rapidly washed in 10 vol of lysis buffer to extract internally free peptide. The pelleted membranes and aliquots from each wash were counted. All centrifugations were done at 15 k rpm for 3 min in a 4°C rotor (SS-34; DuPont Instruments, Sorvall Biomedical Division, Wilmington, DE).

**FIMD**

The micropipette aspiration technique is based on previous descriptions (13) with equipment differences and an image calibration scheme elaborated elsewhere (12). Details of fluorescent probes and membrane labeling are described elsewhere (12, 18). Labeling of skeletal F-actin was
accomplished with fluorescein or rhodamine phalloidin. Protein 4.1 was purified and labeled by previous methods with FITC (39, 48). Iodoacetimido-5-fluorescein (IAF) was conjugated to GST-21.43 and recombinant GST. Polyclonal IgGs to the α-II(44) domain of α-spectrin (S. Marchesi, Yale University, New Haven, CT) were proteolysed to antibody fragments and labeled with FITC. To emulate synthetically a lipid- or glycosphosphatidylinositol (GPI)-linked protein on the red cell's surface, Texas red-conjugated polyclonal IgGs against fluorescein were bound to fluorescein-phosphatidylethanolamine (FL-PE) labeled red cells.

Calculation of Network Conformation from Density Profiles

For an initial perspective of the analysis, deformation of a linear spring is characterized by one stretch ratio that indicates how the spring is stretched from an initial length L to a final length L'. For an axisymmetric elastic surface deformed to a second axisymmetric surface, the shape change and all geometric quantities are fully described by two stretch ratios, λ₁ and λ₂, which are continuous functions of position along orthogonal lines of curvature on the surface (14, and references therein). As an important application, if the surface density (mass per area) at a material point initially is ρ₀, then the final density ρ at that point is determined by how it is stretched: ρ = ρ₀/λ₁λ₂. A relative density can also be defined as ρ = ρ₀L'/L. The relative density of network components is obtained from FIMD (12) and the network stretches may be calculated by integration (Discher, D., unpublished calculations).

Red Cell Membrane Fragmentation Assays

The strength of membranes in a shearing flow was qualitatively assessed in vitro as described elsewhere (4, 11). Typically, 150–250 μl of membranes was added to an equal volume of peptide. Normal and 4.1-deficient membranes incubated with lysis buffer instead of peptide served as positive and negative controls, respectively.

Results

Interactions with Spectrin–Actin In Vitro Show Minimal Functional Peptides Are the 21.26 and 0.59

The primary goal of this work was to elucidate the mechanism of SAB function in the red cell where the dominant interactions of protein 4.1's SAB occur in a ternary complex formed with F-actin and (limiting) amounts of spectrin at the membrane. The studies presented below therefore focus on this three-component system.

By analyzing the abilities of recombinant SAB peptides to induce gelation (Fig. 2 A) when mixed with F-actin (14 μM) and limiting spectrin (1.8 μM), we first identified peptides with no activity (21.10 and 21.20), weak activity (21.26 and 0.59), and strong activity (21.43 and 21.59) comparable to intact 4.1 (11, 15, also Schischmanoff, P. O., unpublished results). To characterize the mechanism of this gelation, spectrin incorporation into sedimentable, supramolecular assemblies was quantitated by densitometry after electrophoresis (vertical scale, Fig. 2 A). Peptide 21.43 was the most potent in stabilizing spectrin–actin interactions at low concentrations; its activity was again like that of purified 4.1. In comparison, both the 21.26 and 0.59 were between one and two orders of magnitude less active. Peptides shorter than the 21.26 had no effect on spectrin binding to F-actin. Despite the large differences in the activity of these peptides, the onset of gelation was consistently within a relatively narrow concentration range corresponding to 50–65% of spectrin recruitment into ternary complexes (light gray, horizontal band, Fig. 2 A). Thus, the mechanism of network assembly and material strengthening in vitro appears to be based on peptide-dependent stabilization of some critical number density of spectrin cross-links between actin filaments.

In related experiments, cosedimentation of 35S-labeled SAB peptides with F-actin (7 μM) and limiting spectrin (2 μM) revealed a two-phase binding process (Fig. 2 B). The 21.43 bound readily into the ternary complex until the spectrin appeared completely complexed with peptide and actin at a stoichiometry of ~1:1 peptide/spectrin (inset, Fig. 2 B). After this first, high affinity, spectrin-dependent phase, there seemed to be a weak, but cooperative, second phase of binding consistent with a binary interaction between peptide and F-actin. In support of this, the second phase of binding saturated at the total F-actin concentration of 7 μM. Decreasing the total F-actin concentration led to a decrease in the magnitude of saturation in this second phase. A weak, and apparently cooperative, interaction between actin and native, full-length 4.1 is consistent with previous reports (2, 36a).

In comparison to the 21.43, the 21.26 and 0.59 showed much less of the first, strong phase of spectrin-dependent binding, but clearly showed a weak, second phase of apparent actin binding. The minimal primary sequence in
common between these two constructs is comprised of the first 26 residues of the constitutive 59-aa domain. These 26 residues are thus implicated as the putative actin binding site in the SAB. Consistent with an important role for actin binding in structural functions like gelation, deletion peptides shorter than the 21.26 did not alter spectrin–actin interactions. Peptide inhibitors of $^{35}$S-21.43 binding included both unlabeled 21.43 and the *Xenopus laevis* SAB, but did not include the small F-actin–binding peptide rhodamine-phalloidin (data not shown).

**Ternary Complex Model Includes Cooperativity in Weak F-Actin Binding**

A previous analysis of ternary complex assembly (37) included both a spectrin–actin and a spectrin–4.1 association, but did not explicitly include any type of 4.1–actin interaction. The experiments presented in that previous study showed only a single, strong phase of spectrin binding to 10 μM actin in the presence of ~1 μM protein 4.1. This ternary interaction saturated at ~1:1 in 4.1/spectrin, had an apparent binary association constant of ~2 × 10$^6$ M$^{-1}$, and yielded a ternary association constant $K_a$ ~10$^{12}$ M$^{-2}$. While experiments did not extend beyond this first strong phase of ternary complex formation, the analysis assumed a very weak saturable second phase of 1:1 spectrin/actin interactions with a $K_a$ = 5 × 10$^5$ M$^{-1}$ estimated from a purely linear binding curve. The ratio of $K_a$ to $K_{sp}$ gave the thermodynamic contribution of 4.1 to ternary complex stabilization and since this ratio exceeded all prior measures of spectrin–4.1 association constants, the authors concluded that additional binding phenomena may contribute to ternary complex formation.

Based in part on this 1:1:1 stoichiometry in spectrin/actin/4.1 associations, our binding isotherms for the SAB constructs (Fig. 2) also suggest an initial ternary complex phase that uses up the limited spectrin. Our data further show a subsequent binary phase of weak, cooperative binding to F-actin. Cooperativity via an actin lattice is therefore proposed in an approximate ternary complex model (Fig. 3). Linear lattice models involving F-actin have been used before, for example, in matrix-based treatments (21) of cooperativity in troponin–tropomyosin binding to filaments (33). A comparable matrix-based analysis has also been performed on our data (Discher, D., unpublished calculations), but, in an effort to simplify a quantitative presentation, a rudimentary two-term expression is written below for the amount of bound (B) SAB peptide as a function of the amount of free (F) SAB peptide:

$$B = S_p(K_{sp} F)^n /[1 + (K_{sp} F)^n] + [A_{cto}] - S_p(K_{sp} F)^n /[1 + (K_{sp} F)^n](K_{sp} F)^n /[1 + (K_{sp} F)^n]$$  

(1)

The first term represents spectrin-dependent peptide–actin interactions in the ternary complex: peptide and spectrin are considered to interact while simultaneously bound to different sites on the same actin subunit. This spectrin-dependent component of peptide binding to F-actin is governed by an apparent binary association constant $K_{sp}$ (Fig. 3). The Hill exponent n represents cooperativity, in that whenever peptides are bound on neighboring actin subunits, they interact, perhaps allosterically. The second term represents purely peptide–actin interactions; a binary association constant $K_{sp}$ governs this binding.

The four interaction parameters for the data fits in Fig. 2 are tabulated in Table I. For all these fits, n = 2.25. By far the greatest variation among the fits of the 21.43, 21.26, and 0.59 appeared in the spectrin-dependent binding constant $K_{sp}$. For the 21.43, $K_{sp}$ was on average ~20-fold greater than that for the 21.26, and more than ~50-fold greater than that for the 0.59. These large differences in spectrin-dependent associations offer some explanation for the lack of detectable binary interactions in solution between spectrin and either the 0.59 (11) or the 21.26 (Schischmanoff, P. O., unpublished results). Finally, for an approximate ternary association constant $K_a$ (M$^{-1}$) ~2(2/$S_p$)*$K_{sp}$, the fits listed in Table I for the 21.43 give $K_a$ ~10$^{12}$ M$^{-2}$ which is in good agreement with previous determinations (38).

**SAB Peptide Binding to Normal Membranes Is Spectrin Dependent**

Various assays of our SAB peptides’ functions inside cells made use of a reversible, lytic incorporation technique. To first elucidate the process of incorporating into a near-normal system, low concentrations (less than or equal to nanomolar) of in vitro translated SAB peptides were loaded into normal cell ghosts during hypotonic lysis. The standard incorporation process (see Materials and Methods) involved a 20-min lysed state on ice during which hemoglobin can leave the cell and peptide can diffuse into the cell through a lytic pore(s) in the membrane (27). Re-sealed but fairly red ghosts were subsequently washed with PBS to remove radioactive peptide not trapped inside the ghosts (Fig. 4A); the membrane-binding ability of this unincorporated peptide was confirmed in subsequent as-

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<th>Table I. Protein 4.1 Peptides in Ternary Complex Formation</th>
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Model fit parameters for ternary complex sedimentation experiments in Fig. 2, A and B. Association constants are tabulated as their inverse, i.e., dissociation constant.

*Refers to binding of peptide in ternary complex with spectrin–actin.

1 Within 10% of measured concentrations.
Figure 4. Binding of SAB deletion peptides to normal red cell membranes. (A) Analysis of radioactive peptide distribution after incorporation by hypotonic lysis of normal cells and rescaling of the ghost membranes (see Materials and Methods). These incorporations were done with 8 μl of each in vitro translated peptide (12–18 nM initial) and 400 μl of membranes. The in vitro synthesis of SAB peptides provided an alternative to bacterial expression and allowed dilute, minimally perturbing tracer studies to be readily performed at the same small scale used in the rest of the work. Resealed ghosts were washed in 10 vol of PBS three times (P1 to P3). The membranes were then rescaled in 10 vol (L1), centrifuged within 4 min, and washed twice more with lysis buffer (L2, L3). 35S-peptide content of all the washes and the membrane pellets (M) was measured and normalized by the total amount of 35S-peptide added. (B) Membrane-bound peptide, as a homogeneous concentration, versus the sum of remaining peptide. The experiments as in A were repeated at various initial concentrations of peptide. The inset indicates the slope of the [35S]21.43 binding curves in the presence of various inhibitors: (i) GST-21.59 at ~25 μM, (ii) GST-XEN, an IgG against the 21-aa cassette (inset, Fig. 4 B), and (iii) preincubation with anti-21 aa IgG at ~1 μM, and (iv) without inhibitor present, the average slope ±SD. (C) Dependence of 21.59 bound over free on the ghost concentration. Pelleted ghosts were diluted in lysis buffer and treated as in A. Model fit of data is described in text. (D) Kinetics of 21.59 binding to normal membranes; experiment and model calculation. Experimental determination of binding kinetics followed A until 2, 6, or 10 min after the incubation on ice. At each time point, the membranes were diluted 10-fold with lysis buffer, centrifuged for 3 min, and the pellets and supernatants counted. For modeling the incorporation kinetics, it was assumed (19) that 2 × 10^5 molecules of endogenous 4.1 per cell (2 μM mean intracellular concentration) were in competition with the recombinant 21.59 (at <1 nM) for 2 × 10^5 skeletal binding sites (2 μM).

say, i.e., the free peptide is not inactive peptide. Parallel studies using fluorescent dextrans (10 kD) and quantitative fluorescence microscopy demonstrated that entrapped concentrations varied by <20% among all cells in a lysed sample. Unbound peptide trapped inside the ghosts by rescaling was released by a second lysis with washing before counting the membrane-bound peptide (Fig. 4 A).

Recombinant peptides 21.59 and 21.43 bound strongly to red cell membranes with a normal content of endogenous protein 4.1 (Fig. 4, A and B). The peptide 21.26, functional in ternary complex assays, failed to show significant binding to these membranes even though the 21.26 contains the putative actin-binding site. Consistent with the revelation of a strong spectrin-dependent binding ability, previous results showed that peptides lacking the 21 aa, including the 0.59, could not bind either spectrin in vitro, or normal membranes in situ (11). Peptides 21.36 and 21.31 were found to bind normal membranes only slightly better than the 21.26 (data not shown), suggesting that these intermediate peptides also interact with spectrin only weakly.

As a further demonstration of membrane-binding specificity, potent inhibitors of the 35S-21.43 were shown to include unlabeled GST-21.59, GST-XEN, an IgG against the 21-aa cassette (inset, Fig. 4 B), and fluorescently labeled GST-21.43. GST protein alone did not inhibit membrane binding (data not shown). In accordance with the proposed spectrin dependence, as more spectrin was added in the form of ghost membranes (ghost volume fraction ϕ), membrane binding of the 35S-21.59 also increased (Fig. 4 C).

To characterize membrane binding kinetics and verify that an equilibrium could be reached within the 20 min on ice, the time-course for incorporation of the 21.59 was studied systematically. Ghosts lysed for 10 min before the addition of peptide showed the same amount of membrane binding as the typical 20-min reaction on ice. By then adding the 21.59 for preset intervals <10 min and not rescaling at all, it was then found that 10 min was necessary to reach equilibrium (Fig. 4 D). Since similar binding was obtained in the absence of rescaling, the typical 37°C rescaling step does not appear to dramatically perturb the 0°C equilibrium. Rescaling was highly effective, however, since preincubating ghosts at 37°C for 10 or 30 min before the addition of peptide, reduced membrane binding by >90%.

Spectrin-dependent Competition with Endogenous Protein 4.1

Binding specificity and the correlations with a strong spectrin-dependent association, particularly the lack of membrane interactions of the 21.26, along with the use of compositionally complex normal membranes (wherein [4.1]_{tot} = [Sp]_{tot}) all suggest that membrane binding is a process involving a competition between the endogenous protein 4.1 and the exogenous SAB for spectrin-dependent binding. Hence, the equilibrium and kinetics of binding (Fig. 4) are analyzed as peptide, P, competing with endogenous 4.1 for binding to skeleton, Sk:

\[
P + \text{Sk} \leftrightarrow \text{P*Sk} \quad K = \frac{k_f}{k_r} \quad (2a)\]

\[
4.1 + \text{Sk} \leftrightarrow 4.1*\text{Sk} \quad K_e = \alpha K \quad (2b)\]

The association constants \(K\) and \(K_e\) are allowed to differ by an accelerating factor \(\alpha\). We expect the skeletal sites to be spectrin-dependent binding sites. Also, because the spectrin–actin-binding activities of the 21.43 and 4.1 appeared equivalent in vitro, the factor \(\alpha\) is intended primarily to capture the effect of protein 4.1's additional interactions with the cytoplasmic domains of both band 3 and the glycoporphins (29, 17). These interactions tend to concentrate the endogenous 4.1 to the red cell's periphery in the region of the skeleton, accelerating protein 4.1's interactions with skeletal components.

The fractional saturation of the total sites \(S_0\), \(S = \frac{[\text{Sk}]_{tot}}{[\text{Sk}]_{tot}}\), by the bound peptide, B: B/S0 = KF/[1 + KF + αKI] (21). The total amount of peptide is much smaller than the total concentration of skeletal binding sites, i.e.,
Membrane Localization of Fluorescent SAB Peptide

In Situ

As a means of explicitly verifying membrane localization and proper integration into network structure, fluorescently labeled molecules in resealed ghosts were imaged after aspirating a ghost into a glass micropipette (Fig. 5). The phospholipid analogue FL-PE readily intercalates into the bilayer and gives a definitive edge-brightness to the image of the sphere outside the pipette (Fig. 5 B). Fluorescein-labeled antibody fragment against spectrin (Fig. 5 A) which includes resealing with salt at 37°C and may therefore give the coarser measure, we get $K = 5 \times 10^6 M^{-1}$, $\alpha' < 0.2$.

The incorporation kinetics data were fit (Fig. 4 D) by the solution of two coupled ordinary differential equations based on the competitive binding model (Eqs. 2a and 2b) and using $K = 0.9 \times 10^6 M^{-1}$, $\alpha' < 0.04$ to 0.2, and $K_t = 0.5 \times 10^5 M^{-1}s^{-1}$. Thus, endogenous protein 4.1 has a slight kinetic advantage (fivefold) in binding to the skeleton from its vantage point on integral membrane proteins.

Another reason for using SAB is to an apparent binary association constant, $K_{ps}$ (M$^{-1}$), for the first phase of spectrin-limited peptide binding to actin in a ternary-complex. As in the ternary-complex analysis, $K_{sp} \approx (S_0/2)K_p$; hence a $K_{sp}$ $\sim 10^4 M^{-1}$ is estimated from $K_j \approx 10^{12} M^{-2}$ and $S_0 \approx 2 \mu M$.

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Conventional microscopy was used for most of the imaging since it provides a sensitivity advantage, but optical sectioning (Fig. 6 A, inset) was also used to clearly verify edge enrichment. Quantitatively, for a spherical ghost cytoplasmic vol of $\sim 150 \mu m^3$ filled by unbound label and a range in skeletal thickness between 5 and 100 nm (estimated from either actin monomer dimensions or spectrin dimer contour length; 5) binding sites at the membrane would be enriched relative to the cellular concentration by between 200- and 12-fold, respectively. For protein 4.1, these numbers imply that its apparent concentration at the membrane ranges between 36 and 600 $\mu M$, substantially higher than its mean cellular concentration of 3 $\mu M$. As further evidence, edge brightness with rhodamine-phalloidin was measured as a function of added phalloidin (Fig. 6 A); a fit of the phalloidin-binding data with a simple bimolecular model indicated a high-affinity membrane association constant that was within a factor of 7 of in vitro mea-
Figure 6. (A) Specificity of rhodamine-phalloidin labeling of a normal ghost membrane. Left inset image is a rhodamine-phalloidin–labeled ghost imaged with a confocal microscope; right image is taken with a conventional fluorescence microscope. The curve indicates edge intensity of images using conventional microscopy and measured as a function of total phalloidin added. Rhodamine-phalloidin was serially diluted in lysis buffer, after measuring the highest concentration using a dye/phalloidin ratio of 1:1 and a peak absorptivity of 76,000 M⁻¹cm⁻¹. The fitted curve used 2 × 10⁻⁷ M for the concentration of F-actin binding sites as calculated from 4 × 10⁵ actin molecules per membrane (19) and the volume of cells lysed. The apparent association constant is 0.33 × 10⁻⁷ M⁻¹ which is slightly weaker than in vitro assays. Average ±SD are shown for at least nine ghosts labeled at each concentration. (B) Localized network stretching conformations and SAB peptide integration. Plot of a typical network density profile, ρ, on an aspirated membrane projection along with calculated stretch ratios: axial stretch, λₐ, and circumferential stretch, λ₂ (see Materials and Methods). The abscissa is in the shape of the aspirated projection, and shows the axial distance z scaled by the pipette radius Rp. To the left of the projection is illustrated a square piece of the cytoskeletal meshwork before deformation. This meshwork is dilated at the tip of the projection, and about halfway down the cylinder the meshwork is in a rectangular state of pure shear. The principal directions of stretch and the deformation of an initial square are also sketched on the projection.

measurements for phalloidin binding to F-actin. These coarse calculations together with the confocal image (Fig. 6 A, inset) provide more explicit illustrations of the membrane-concentrating effect and demonstrate that physiological concentrations can be much higher than the micromolar range.

In Situ Integration into the Solidlike Cytoskeletal Network

In cells other than erythrocytes, actin-binding proteins localize to stress fibers, focal contacts, and a number of other structures (e.g., 34). If one were to deform reversibly such cells and redistribute the structures, one would still expect to see a colocalization of the actin and actin-binding proteins to the same redistributed structures. In the aspirated images of Fig. 5 then, specificity of localization and structural connectivity are indicated by the fluorescence patterns obtained after deforming the uniformly labeled red cell into the micropipette (12). In short, FL-PE shows that the lipid bilayer conveys like a fluid sheet to cover uniformly (Fig. 5 B) a strong gradient in density of the underlying skeleton as labeled with either antiserpectrin, rhodamine-phalloidin, FITC-4.1, or IAF-GST-21.43 (Fig. 5, C, D, E, and F, respectively). Formation of this density gradient is much like the very tangible process of pushing a finger into the palm of a rubber glove: looking towards the extreme tip of the finger, one sees that the material becomes progressively more dilated. Important in contrast, is the finding that lipid-linked proteins, both lipid-bound antibody (Fig. 5 H) and GPI-anchored protein CD-59 (12), which presumably have no means for connection to the underlying skeleton, give a response in deformation nearly opposite to the network probes (12, for clarification).

Quantitative “fingerprints” of these component distributions are given by the intensity sections along the projections shown in small insets below each cell (Fig. 5, insets). For lipid probe, the intensity is nearly uniform, but for spectrin, actin, 4.1, and recombinant SAB peptide, a strong gradient in density appears. Particularly note that these insets show the network density is nonvanishing at the tip of the projection. This gradient is stable (up to 30 min) and reversible upon release from aspiration, indicating a solidlike skeletal network and consistent with a resilient and durable membrane. The two-dimensional nature of the network permits calculation (see Materials and Methods) of how an initially small square of cytoskeletal meshwork can, depending on its location, be transformed to various rectangular conformations down the length of the projection (Fig. 6 B). At the very tip or cap of the projection, the initially square bit of meshwork is simply dilated in the deformation. Toward the pipette entrance, the material is highly stretched in the axial direction, but compressed in the lateral direction; this conformational state is one of extensive shearing and is a sustainable state for a solidlike but not a labile, fluidlike network. The fact that the SAB peptide gives the same apparent density field along the projection as the native skeletal components, explicitly shows integration of the peptide into the same elastic network structure whether this structure is sheared, dilated, or compressed reversibly. This suggests that both the SAB peptide and protein 4.1 should have similar effects on network integrity or strength.

Red Cell Membrane Strength Is Determined by SAB Peptides’ Ability to Stabilize Spectrin Cross-links of Actin In Situ

Red cells deficient in protein 4.1 possess membranes that are more fragmentable than normal membranes (46, 51). However, both purified, exogenous 4.1 (47) and recombinant GST-21.59 (11) can, at approximately micromolar cellular concentrations, increase membrane strength to normal. To test this crucial aspect of in situ cellular function of our SAB constructs, mutant red cells with deficiencies in protein 4.1 were lysed in the presence of each peptide. After resealing and suspending the ghosts in a viscous dextran, each sample was subjected to a membrane-fragmentation assay.

Initially, the deformation response for all the samples is about the same value (Fig. 7), indicating that all ghosts start as comparably stretched ellipsoids. After a very short interval of sustained deformation, however, the 4.1-deficient membranes (~40% deficiency) show a transition to steady fragmentation as indicated by the decay in the deformation index. Normal membranes fragment much later. Addition of the peptide 21.20 to the 4.1-deficient membranes did not alter the membrane fragmentation process (Fig. 7 A). The 21.43, in contrast, was able to strengthen the 4.1-deficient membranes to a level comparable to nor-
membranes could be strengthened by peptides 21.43, 21.26, and 0.59, but not 21.20 at high concentrations. Thrombin-cleaved, 35S-labeled peptides were added to ghosts prepared from partially (~40%) 4.1-deficient red cells. Peptide added equals the amount of total peptide divided by the volume after addition to membranes. The relative strength of the resealed membranes was assessed by membrane fragmentation under fluid shear and is plotted as a time-dependent decline in deformation index. Responses of normal (solid line) and 4.1(-) membranes (large dashed line) defined the range of membrane strengthening effect for each construct. (A) Peptide 21.20, even at very high concentrations, did not alter the fragmentation rate. (B) Peptide 21.43 restored membrane strength at as low as 1.5 μM, consistent with published results for either purified protein 4.1 or GST-21.59, while (C) increasing the concentration of 21.26 resulted in a decreased rate of fragmentation, hence increased membrane strength. (D) Entrapment of 0.59 at 37 μM and, to a lesser extent, at 13 μM showed a positive effect.

A, B, C, D

Figure 7. Membranes can be strengthened by peptides 21.43, 21.26, and 0.59, but not 21.20 at high concentrations. Thrombin-cleaved, 35S-labeled peptides were added to ghosts prepared from partially (~40%) 4.1-deficient red cells. Peptide added equals the amount of total peptide divided by the volume after addition to membranes. The relative strength of the resealed membranes was assessed by membrane fragmentation under fluid shear and is plotted as a time-dependent decline in deformation index. Responses of normal (solid line) and 4.1(-) membranes (large dashed line) defined the range of membrane strengthening effect for each construct. (A) Peptide 21.20, even at very high concentrations, did not alter the fragmentation rate. (B) Peptide 21.43 restored membrane strength at as low as 1.5 μM, consistent with published results for either purified protein 4.1 or GST-21.59, while (C) increasing the concentration of 21.26 resulted in a decreased rate of fragmentation, hence increased membrane strength. (D) Entrapment of 0.59 at 37 μM and, to a lesser extent, at 13 μM showed a positive effect.

For a quantitative approach to the incorporation chemistry, we used 35S-labeled, thrombin-cleaved peptides in the fragmentation assays. Measurements of the entrapped peptide concentrations showed that the concentration of the inactive peptide 21.20 inside ghosts was linear over the full range of added peptide, extending to ~50 μM entrapped. In contrast, the amounts of 21.43, 21.26, and 0.59 entrapped at a given concentration of added peptide were consistently higher than the 21.20, and appeared as an additional, specific membrane-binding component. The amount of specific binding (Fig. 8 A) was therefore obtained by subtracting the baseline amount of 21.20 physically entrapped by rescaling. In addition, for the 21.43 at two concentrations, the amount of membrane-bound peptide after a second lysis was directly measured (as per Fig. 4 A), and was found to coincide with the calculation based on subtraction of the 21.20 baseline. When the ternary complex model was fit through the membrane-binding data (Fig. 8 A), we used a value for available spectrin (0.8 μM = 40% of 2 μM) consistent with an ~40% (32) deficiency in endogenous protein 4.1 (normally 2 μM). The actin available for binding used in fitting the results (2.4 μM; Table II) appears to be slightly less than an estimate (2.9 μM) based on the total actin content (4.1 μM minus the endogenous 4.1 (60% of 2 μM). Importantly, the trends in spectrin-dependent binding constants obtained in vitro (Table I) are substantially replicated in the fits of binding to membrane structures in situ (Table II). Competition with endogenous 4.1 has been ignored in this membrane-binding analysis, since such competition is likely to have an effect only at high 21.43 concentrations after the 40% of skeletal
Table II. Interaction of Protein 4.1 Peptides with the Membrane

<table>
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<th>21.43</th>
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<tr>
<td>$s_\beta, \mu M$</td>
<td>2.4</td>
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Model fit parameters for binding of peptides to membranes (Fig. 8 A) used in fragmentation assay.

Table II. Interaction of Protein 4.1 Peptides with the Membrane

sites left unoccupied by 4.1 become largely filled by the 21.43.

Full membrane-strengthening effect (Fig. 8 A, solid points) by the 21.43 occurs with much lower binding to the membrane (>0.3 μM) than required for the 0.59 (~2 μM). The underlying mechanism that gives rise to this approximately sevenfold difference in effectiveness is significantly clarified, however, by calculating for each SAB construct the peptide-facilitated increase in spectrin–actin associations in the membrane. This spectrin-dependent phase of the binding of each peptide, the first term in Eq. 1, is plotted as the predicted fraction of membrane spectrin that binds to actin. A background shading scheme also illustrates the effect on membrane strength (Fig. 8 B). As found in vitro (Fig. 2 B), spectrin–actin stabilization is very closely associated with material strengthening.

Discussion

COOH-terminal deletion constructs and an NH2-terminal isoform of protein 4.1's SAB domain have been assayed for their ability to enter into ternary associations with spectrin–actin in vitro and also to specifically bind, integrate into, and mechanically strengthen red cell membranes in situ. Highly efficient, membrane-strengthening interactions involving both spectrin and actin require both the NH2-terminal 21 aa, and also the 43 residues of the constitutive 59-aa domain. However, a reduced yet adequate, membrane-strengthening activity arises from either the 21 aa together with the first ~26 residues of the 59 aa or just the 59 aa alone. Combined with similar findings in ternary complex formation in vitro, a simplistic structural model of SAB function emerges (Fig. 9): two sites for interaction flank the ~26 residue actin-binding site. Exactly half of the residues of the putative actin-binding site are charged, consistent with localization to the protein surface. This sequence also contains Lys487 which, through deletion of this conserved 20th of 26 residues, has recently been implicated in both defective red cell membranes and an inability to bind spectrin–actin (30). The two putative spectrin-binding sites are largely conserved among species, consistent with our results for X. laevis 21.59 (Fig. 4 B). Structural artifacts arising from peptide construction could contribute to the results, but preliminary CD spectra suggest that small if any changes in global peptide conformation result from our deletions in primary structure (Schischmanoff, P. O., unpublished data). In further support of two spectrin sites, a tyrosine kinase phosphorylates at the 40th residue of the 59 aa (45); both phosphoadducts of the SAB have substantially reduced spectrin–actin-binding activity (28, 45). Thus, in addition to the very strong genetic modulation of activity provided by splicing in or out the alternative NH2-terminal exon, the split in the spectrin-binding interaction establishes two physiological pathways for strong modulation via post translational modification.

Because the constitutive 59-aa domain on its own is found functional, albeit weakly, in both gelation and membrane-strengthening assays, expression of the alternatively spliced 21 aa may be interpreted as a biologically efficient (42), rather than physically necessary means of achieving the red cell membrane’s required strength. It should be emphasized that the mean intracellular concentration of 0.59 sufficient for fully normal strengthening (~12 μM, Fig. 8) is substantially below the minimum estimate of protein 4.1 concentration at the membrane (>36 μM).

After controlled, reversible aspiration of ghosts, spectrin, actin, 4.1, and the recombinant 21.43 appeared to have moved collectively as an integrated structural unit. Fragmentation of 4.1-deficient red cell membranes under high fluid shear was subsequently shown to be modulated by the SAB peptides’ abilities to bind spectrin–actin. The rapid, fluid-forced fragmentation of the red cell membrane seems likely to be a process of catastrophic failure at the weakest point in the structure. In the absence of protein 4.1, the membrane’s weak point would appear to be the bare spectrin–actin interaction. Addition of enough of a recombinant SAB peptide, even an inefficient stabilizer like the 0.59, ultimately shifts the membrane's weak point to another set of strength limiting interactions. Below this strength limit, concomitant increases in spectrin–actin stabilization and material strength, either membrane strength (Fig. 8 B) or gel strength (Fig. 2 A), appear roughly peptide independent. This suggests that, for all the functional SAB peptides, it is primarily the stabilization of spectrin cross-links of F-actin that solidifies and strengthens the structure.

Like protein 4.1, the spectrin–actin-stabilizing phosphoprotein adducin seems to bind F-actin cooperatively (35). Adducin’s binary interaction with actin can cooperatively drive spectrin into a spectrin–actin–adducin ternary complex, similar perhaps to the SAB peptides 21.26 and 0.59, which interact with spectrin only weakly (Fig. 2 A). β-adducin is also alternatively spliced, and occurs in a number of tissues as an isoform lacking portions of the spectrin–actin binding COOH-terminus (Gilligan, D. M., and V. Bennett. 1994. Mol. Biol. Cell. 5:421a; Hughes, C. A., and V. Bennett. 1994. Mol. Biol. Cell. 5:421a).

The present data reinforce the finding (11) that protein
4.1’s role in membrane strength, as assessed by the fluid shear-fragmentation assay, is essentially determined by its SAB interactions, independent of further associations of protein 4.1 with glycoporphins, p55, band 3, or phosphatidylserine. A role for glycoporphin C as a site for linking the bilayer and network was suggested by studies showing glycoporphin C-deficient red cells are more easily fragmented in fluid shear than normal membranes (43). However, additional studies of the glycoporphin C deficiency have noted a coexisting deficiency in protein 4.1 (1), and we have recently found that addition of either purified protein 4.1 or just an SAB peptide to red cells completely deficient in protein 4.1 with glycophorins, p55, band 3, or phosphatidylserine, strengthened these membranes to normal levels (43). Interactions at the membrane clearly have a concentrating effect (e.g., Fig. 6 A, inset). For protein 4.1, peripheral interactions with bilayer structures would tend to accelerate its interactions with network spectrin and actin. Quantitative evidence for this hypothesis was provided, to a limited extent, by the competitive binding of SAB to normal membranes: multifunctional 4.1 appears to have a fivefold kinetic advantage in competition for membrane localization. The net result of this mechanism for membrane stabilization may be an evolved effort to economize with a fixed and minimal amount of protein. Splicing in the 21-aa cassette has a similar effect as it tremendously increases SAB activity and thereby reduces the amount of free protein. Both of these mechanisms for the enhanced binding of 4.1 to spectrin–actin in situ thus act to minimize the total amount of protein needed for network stabilization. The studies presented here on the mechanism for SAB interactions with the membrane skeletal protein 4.1’s SAB domain therefore support the idea that, besides the possibilities of contributing directly to cell mechanical properties, multiplicity in structural protein interactions can play an important role in cell protein economics.

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