Complexes of Native Ubiquitin and Dodecyl Sulfate Illustrate the Nature of Hydrophobic and Electrostatic Interactions in the Binding of Proteins and Surfactants

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ABSTRACT: A previous study, using capillary electrophoresis (CE) [J. Am. Chem. Soc. 2008, 130, 17384–17393], reported that six discrete complexes of ubiquitin (UBI) and sodium dodecyl sulfate (SDS) form at different concentrations of SDS along the pathway to unfolding of UBI in solutions of SDS. One complex (which formed between 0.8 and 1.8 mM SDS) consisted of native UBI associated with approximately 11 molecules of SDS. The current study used CE and 15N/13C heteronuclear single quantum coherence (HSQC) NMR spectroscopy to identify residues in folded UBI that associate specifically with SDS at 0.8–1.8 mM SDS, and to correlate these associations with established biophysical and structural properties of this well-characterized protein. The ability of the surface charge and hydrophobicity of folded UBI to affect the association with SDS (at concentrations below the CMC) was studied, using CE, by converting lys-ε-NH3+ to lys-ε-NHCOCH3 groups. According to CE, the acetylation of lysine residues inhibited the binding of 11 SDS ([SDS] < 2 mM) and decreased the number of complexes of composition UBI-(NHAc)6·SDS, that formed on the pathway of unfolding of UBI-(NHAc)9 in SDS. A comparison of 15N-1H HSQC spectra at 0 mM and 1 mM SDS with calculated electrostatic surface potentials of folded UBI (e.g., solutions to the nonlinear Poisson–Boltzmann (PB) equation) suggested, however, that SDS binds preferentially to native UBI at hydrophobic residues that are formally neutral (i.e., Leu and Ile), but that have positive electrostatic surface potential (as predicted from solutions to nonlinear PB equations); SDS did not uniformly interact with residues that have formal positive charge (e.g., Lys or Arg). Cationic functional groups, therefore, promote the binding of SDS to folded UBI because these groups exert long-range effects on the positive electrostatic surface potential (which extend beyond their own van der Waals radii, as predicted from PB theory), and not because cationic groups are necessarily the site of ionic interactions with sulfate groups. Moreover, SDS associated with residues in native UBI without regard to their location in α-helix or β-sheet structure (although residues in hydrogen-bonded loops did not bind SDS). No correlation was observed between the association of an amino acid with SDS and the solvent accessibility of the residue or its rate of amide H/D exchange. This study establishes a few (of perhaps several) factors that control the simultaneous molecular recognition of multiple anionic amphiphiles by a folded cytosolic protein.

INTRODUCTION

This paper describes a study of the association of SDS with two small proteins—ubiquitin (UBI; also abbreviated (UBI-(NHAc))6) and its peracetylated derivative (UBI-(NHAc))9, in which all lysine groups and the amino terminus are acetylated. This acetylation neutralizes ~70% of the positively charged residues of UBI. The work has three objectives: (i) to determine if surface-accessible cationic residues on the surface of UBI are involved in specific electrostatic association with SDS, (ii) to clarify the relative importance of electrostatic and hydrophobic interactions in the interaction of an anionic surfactant with proteins, and (iii) to determine how the structural environment of residues in folded UBI (i.e., secondary structure and solvent accessibility) correlates with the interaction of those residues with SDS (at concentrations of SDS that do not unfold UBI).

The association between a protein and a charged surfactant involves both hydrophobic and electrostatic interactions, but the
relative importance of these interactions has not been defined. We believe that correlating the pathway of these associations (i.e., the number of surfactants that bind to the protein, as a function of the concentration of the surfactants) with the biophysical and physical-organic properties and characteristics of proteins (for example, net charge, electrostatic surface potential, hydrophobicity, secondary and tertiary structure, conformational flexibility, and fold family, to name a few) will help in understanding the interaction between surfactants and proteins, and the unfolding or aggregation of proteins that often results from this association. The interaction of lipids and proteins has, for example, been hypothesized to be a crucial step in promoting (or preventing) the aggregation of proteins and the pathogenesis of protein aggregation diseases.1–3

Ubiquitin Binds Up to 11 SDS Molecules without Unfolding. Using CE, we have previously demonstrated that UBI unfolds in SDS via an unexpectedly complicated pathway that involves the stepwise binding of discrete numbers of molecules of SDS and, therefore, the formation of distinct complexes along the unfolding pathway: UBI-SDS₁, UBI-SDS₂, and UBI-SDS₃,₃,₃ are especially stable. We sometimes refer to these complexes as intermediates, although some involve completely folded UBI (i.e., UBI-SDS₁,₁), and others completely unfolded UBI (i.e., UBI-SDS₃,₃,₃).

The complexes of UBI that form at low concentrations of SDS (<2 mM, i.e., UBI-SDS₁,₁) are remarkable for their conservation of native structure (as measured by circular dichroism (CD) and by rates of hydrogen–deuterium exchange). The complexes that form at higher concentrations of SDS (>2 mM; e.g., UBI-SDS₂,₂, UBI-SDS₃,₃, UBI-SDS₄,₂) show more α-helical structure than the native protein (as measured by CD) but retain very little tertiary structure, as demonstrated by the rapid rate of hydrogen–deuterium exchange. Above the critical micellar concentration (CMC), which for SDS is 3.4 mM (in 25 mM Tris, 192 mM glycine, pH 8.4, 25 °C), UBI is “unfolded” and saturated with approximately 42 molecules of SDS. This stoichiometry (UBI-SDS₄,₂) translates to approximately one SDS for every two amino acids.

Most of the proteins that we have studied with CE either do not form complexes of intermediate stoichiometry with SDS (e.g., most cannot bind SDS and remain natively folded) or do not form as many different complexes as does UBI. We have, so far, been unable to predict which proteins interact with SDS in ways that form stable aggregates of protein and surfactant, and that retain much or all of the tertiary structure of the protein. Nevertheless, because UBI is folded in some of the UBI-SDS complexes that form (i.e., UBI-SDS₁,₁) and because multiple interactions (as many as 11) between the folded protein and surfactant can be studied simultaneously, the UBI-SDS system could be particularly useful for acquiring general information on the interactions of folded proteins with surfactants (and perhaps with other small molecules in general).

Native Ubiquitin—Poly(dodecyl sulfate) Complexes Represent a Useful System in the Study of Molecular Recognition. The binding of SDS to folded UBI below the CMC does not involve the geometry of association common in many other protein—ligand systems—that is, the binding of a ligand inside a pocket. The binding of SDS to native UBI at 1 mM SDS, for example, appears to occur on the surface of the protein. It must be remembered, however, that the binding of SDS to the surface of UBI can be both strong and specific below the CMC: the complexes that form at ~1 mM SDS (e.g., UBI-SDS₁,₁) are equilibrium complexes. These complexes are characterized during CE by a single peak that is unimodal (and as sharp as the peak representing UBI-SDS₃,₃); this absence of peak broadening indicates that the interconversion of UBI-SDS₁,₁ between, for example, UBI-SDS₁₀, or UBI-SDS₁₄ is slower than the time scale of a CE experiment (~10–20 min). Native ubiquitin-poly surfactant complexes (prepared with SDS or with other surfactants) could, therefore, become useful systems for studying protein—ligand interactions, and systems that could yield new insights into the physical-organic chemistry that underlies both the hydrophobic effect and biomolecular recognition.

In this paper we have investigated, in particular, the importance of surface-accessible, cationic, lysine ε-NH₂ residues in the pathway of association of UBI with SDS, and we have identified residues in folded UBI that bind SDS at concentrations that are below its CMC. The two analytical tools used in the present study are CE and heteronuclear single quantum coherence (HSQC) NMR spectroscopy. Capillary electrophoresis provides information about the stoichiometry of protein—SDS interactions: the electrophoretic mobility of low- to medium-molecular-weight proteins (<60 kDa) in a bare capillary is sensitive to the binding of a single surfactant molecule. Because the native fold of UBI is retained in the first UBI-SDS complex that forms (at ~1 mM SDS), we can identify residues that interact with SDS below the CMC by shifts of amide N–H peaks in ¹³N–¹H HSQC and C–H peaks in ¹³C–¹H HSQC.

Our results show that the binding of SDS to folded UBI (and the pathway of unfolding of UBI) can be changed by altering the electrostatic surface potential and hydrophobicity of UBI. The results of our analysis of complexes of UBI and SDS with CE and HSQC also suggest, unexpectedly, that the negatively charged sulfate group of SDS does not, generally, associate selectively with the positively charged ammonium group of lysine (or the guanidinium group of arginine) below the CMC of SDS. The binding of SDS by UBI (below the CMC) is driven, instead, by an apparently balanced combination of electrostatic and hydrophobic forces: binding occurs at surfaces of the protein that both (i) are hydrophobic and (ii) have positive electrostatic surface potential (but not necessarily a formal positive charge).

### EXPERIMENTAL DESIGN

#### Selection of Ubiquitin.

We used UBI because it is a heat-stable, small protein (76 amino acids, 8565 Da), which has both α-helix and β-sheet elements, and which forms a variety of complexes with SDS that can be distinguished by CE. UBI is a tractable and widely used model protein for experimental (and sometimes theoretical) studies of protein folding. UBI contains seven lysines and one N-terminal methionine α-NH₂ group (all of these amino groups can be acetylated by acetic anhydride); one C-terminal ε-COO⁻ group; four arginines; one histidine; five aspartates and six glutamates; nine leucines; seven isoleucines; and three prolines. The theoretical isoelectric point of UBI is 6.56. The theoretical value of the net charge of UBI (Z_{SE}) at pH 8.4 was calculated, from its sequence, to be Z_{SE} = 0 (assuming that Z = +1 for the N-terminus; Z = −1 for the C-terminus; Z = −1 for Asp and Glu; Z = +1 for Arg and Lys).

The value of net charge of UBI that we estimated experimentally with CE (Z_{CE}) at pH 8.4 (25 mM Tris, 192 mM glycine) was Z_{CE} = −0.2 (in our previous study). The Supporting Information includes a discussion of the possible causes of this small difference between Z_{CE} and Z_{SE}.
Using Heteronuclear Single Quantum Coherence NMR Spectroscopy To Identify Electrostatic or Hydrophobic Interactions between Specific Residues in Folded UBI and SDS. Because a specific UBI-SDS complex is predominant at 0.8—1.2 mM SDS, and because the UBI proteins with up to 11 bound SDS have the same structure as UBI-SDS, we can use $^{15}$N—$^1$H HSQC NMR spectroscopy to identify the residues in UBI that associate with SDS at 0.8—1.2 mM.

The binding of a ligand (such as a surfactant) to residues in a protein can be conveniently detected by a shift in the amplitude $^{15}$N—$^1$H resonance of the interacting residue along the $^{15}$N or $^1$H axes (i.e., a chemical shift perturbation, denoted CSP or $\Delta \delta$), and in some cases, by a reduction of the intensity of the $^{15}$N—$^1$H resonance. Chemical shift perturbations have been widely used to characterize the interaction between proteins and charged ligands, including fatty acids (i.e., palmitic acid), anionic detergents, nucleic acids, oligonucleotides, lipids, carbohydrates, and small drug-like molecules. The specific residues whose $^{15}$N—$^1$H resonance shifts upon titration with a ligand in solution are often the same residues that are binding directly to the ligand according to X-ray crystallography—so long as no change in the conformation of the protein occurs (in solution) upon the binding of ligand.

We expect that electrostatic interaction between the ROSO$_3^-$ headgroup of SDS and any positively charged functional group in UBI (i.e., $\epsilon$-$\text{NH}_3^+$ or $\delta$-NH(CNH$_2$)$^+$) might be detected by movement in $^{15}$N—$^1$H resonance of the interacting residue along the $^{15}$N or $^1$H axes ($\Delta \delta$). We base these predictions on previously published reports of chemical shift perturbations of residues (in structurally well-characterized proteins such as UBI and nuclease) that engage in electrostatic interactions with negatively charged ligands (and micellar surfaces) that bear phosphate, carboxylic, and sulfate groups. The application of NMR spectroscopy to study the interaction of folded proteins with surfactants in detergent micelles (e.g., membrane proteins) has been reviewed by Wüthrich.

Approximating the Electrostatic Surface Potentials of Residues in Native Ubiquitin That Are Accessible to Solvent or Ligand. In order to determine if the sulfate group of SDS interacts with residues in UBI as a function of the local electrostatic environment of the residue (as opposed to, or in addition to, the formal charge of the residue), and in order to determine how the positive charge on some residues might be screened or dampened by solvation—and therefore not interact with R-OSO$_3^-$—we approximated the electrostatic surface potential of residues in folded UBI and mapped these potentials onto the solvent-accessible surface of UBI (based upon the X-ray crystal structure, PDB 1UBI). The electrostatic potential of residues at the solvent-accessible surface of UBI was approximated by solving the nonlinear Poisson—Boltzmann (PB) equation, using an implicit solvent model (e.g., treating solvent as a continuum with a uniform dielectric). A more detailed description of the PB implicit solvent model (e.g., the assumptions and weaknesses of the model) can be found in Supporting Information. One utility of numerical solutions to the PB equation is that they provide electrostatic information about the surface of a protein that is not obvious from an examination of the location of positively and negatively charged residues in the three-dimensional structure. The validity of these models for use in rationalizing the molecular recognition of charged ligands and proteins is largely empirical: proteins that bind DNA, for example, generally have patches of positive electrostatic surface potential (according to numerical solutions of PB equations) located at the site of binding for DNA. Phospholipases have patterns of electrostatic surface potential (at the site of lipid binding) that are thought to facilitate the binding of negatively charged lipids, many other examples of complementary electrostatic surface potentials between macromolecules that interact with one another, or with ligands, have been reviewed.

RESULTS AND DISCUSSION

Neutralizing Lys-$\epsilon$-$\text{NH}_3^+$ and $\alpha$-$\text{NH}_3^+$ of UBI with Acetylation Prevents the Formation of Folded Ubiquitin—Poly(dodecyl sulfate) Complexes. The electropherograms displayed in Figure 1 show the binding of different numbers of SDS molecules to UBI-$\text{(NH}_3^+)^8$ and UBI-(NHAc)$_8$ at different concentrations of SDS. For example, at 0 mM SDS, the electrophoretic mobility of UBI-$\text{(NH}_3^+)^8$ is similar to that of the neutral marker (dimethylformamide, DMF) because the net charge of UBI-$\text{(NH}_3^+)^8$ (denoted $Z_0$) is close to zero (at pH 8.4, $Z_0 = -0.2$). The peak for UBI-$\text{(NHAc)}_8$ shifts to a higher mobility as the concentration of SDS is increased from 0.0 to 0.2 and 0.4 mM because the binding of SDS increases the net negative charge of UBI-$\text{(NHAc)}_8$; the neutral marker peak remains at $\mu = 0 \text{ cm}^2 \text{v}^{-1} \text{s}^{-1}$.

The progressive binding of SDS to UBI-$\text{(NHAc)}_8$ is illustrated in the capillary electropherograms by a transition from the native protein (denoted “N”) to discrete complexes that we describe as groups “G” depending of their mobility. The first discrete and well-resolved complex (denoted G$_2$) forms via a set of complexes (denoted G$_1$, Figure 1a). Although G$_2$ represents the first discrete complex with a defined stoichiometry, G$_1$ represents, in fact, the first complex per se—actually a set of complexes. We estimate the net charge of the complex UBI-$\text{(NHAc)}_8$—SDS$_{1-1}$ to be $-10.1$ (assuming that each SDS molecule increases the negative charge by 0.9 unit$^{-1}$). The Supporting Information contains more details about the progression from N to G$_2$ (see Figure S1).

The electropherograms of UBI-(NHAc)$_8$ in different concentrations of SDS illustrate how SDS binds to UBI-(NHAc)$_8$ (Figure 1b). The mobility of UBI-(NHAc)$_8$ at 0 mM SDS ($\mu = 15 \text{ cm}^2 \text{v}^{-1} \text{s}^{-1}$) is significantly larger than for
Journal of the American Chemical Society

Figure 1. Studying the binding of SDS to UBI and peracetylated UBI with capillary electrophoresis. CE electropherograms of UBI-SDS complexes prepared by dialyzing UBI-(NH$_3^+$)$_8$ (a) and UBI-(NHAc)$_8$ (b) against Tris-glycine (25 mM Tris, 192 mM glycine) containing different concentrations of SDS (0–10 mM). In these electropherograms, UBI-(NH$_3^+$)$_8$ at 0 mM SDS is denoted “N”, and the successive complexes that form during the unfolding at higher concentrations of SDS (e.g., the UBI-SDS$_n$ complexes) are denoted “G$_n$”, where $n$ reflects the order in which the complex is formed along the unfolding pathway. (a) Because the net charge, Z$_n$ of UBI-(NH$_3^+$)$_8$ is so close to zero at pH 8.4 and 0 mM SDS (Z$_n = -0.2$), the mobility of UBI-(NH$_3^+$)$_8$ at 0 mM SDS (i.e., “N”) is similar to the mobility of the neutral marker DMF at 0 mM SDS. Therefore, the peak for DMF appears as a left-handed shoulder, at 0 mM SDS, that is unresolved from the peak for UBI-(NH$_3^+$)$_8$ at 0 mM SDS. The peak for UBI-(NH$_3^+$)$_8$ shifts to a higher mobility as the concentration of SDS is increased (as a result of the binding of SDS to UBI), but the neutral marker remains at mobility as the concentration of SDS is increased (as a result of the temperature = 25°C). (b) UBI-(NHAc)$_8$ has a mobility of $\mu$ = 15 cm$^2$ kV$^{-1}$ min$^{-1}$ at 0 mM SDS, and this mobility is constant up to $\sim$2.0 mM SDS, whereas the mobility of UBI-(NH$_3^+$)$_8$ in (a) was increasing from 0 to 2 mM SDS as a result of its binding with SDS. In both (a) and (b), electrophoresis was performed at 30 kV using a capillary made out of fused silica (total length of 60.2 cm, length to the detector of 50 cm, temperature = 25°C).

UBI-(NH$_3^+$)$_8$ at 0 mM SDS because UBI-(NHAc)$_8$ (Z = -7.4) is much more negatively charged than UBI-(NH$_3^+$)$_8$ (Z = -0.2).

The binding of SDS to UBI-(NHAc)$_8$ is quite different from that of UBI-(NH$_3^+$)$_8$. For example, UBI-(NHAc)$_8$ migrates during CE—at concentrations up to 1.8 mM SDS—as a single peak that has a mobility indistinguishable from that of UBI-(NHAc)$_8$ at 0 mM SDS (N’, $\mu \approx$ 15 cm$^2$ kV$^{-1}$ min$^{-1}$, Figure 1b). Therefore, SDS does not appear to be binding to UBI-(NHAc)$_8$ at concentrations of SDS below 1.8 mM. The inhibition of the recognition of SDS by UBI (at <1.8 mM SDS) by the neutralization of its lysine-$\varepsilon$-NH$_3^+$ groups (and N-terminus) is remarkable and demonstrates that electrostatic interactions are involved, somehow, in the binding of SDS with UBI—and that hydrophobic interactions are not so dominant that they can completely explain the association of SDS with UBI in the absence of electrostatic interactions.

The different capacities of UBI-(NHAc)$_8$ and UBI-(NH$_3^+$)$_8$ to bind SDS (at <1.8 mM SDS) caused us to hypothesize that the R-OSO$_3^-$ group of SDS formed specific electrostatic interactions with positively charged groups on the surface of UBI-(NH$_3^+$)$_8$. The neutralization of eight ammonium groups (seven lysines and the N-terminus) with acetate anhydride, however, prevented the binding of 11 molecules of SDS at <1.8 mM SDS, and not eight molecules of SDS, as might be expected if SDS molecules were associating selectively and specifically with Lys-$\varepsilon$-NH$_3^+$.$^4$ The four guanidinium groups of UBI-(NH$_3^+$)$_8$ remained positively charged in UBI-(NHAc)$_8$, after peracetylation (presumably), but these Arg residues did not bind SDS below a concentration of 1.8 mM. Therefore, there is not a 1:1 relationship between the number of positively charged functional groups in UBI-(NHAc)$_8$ (e.g., four guanidinium groups) and the number of SDS bound at 0–1.8 mM SDS (e.g., zero SDS). Moreover, there is not a 1:1 relationship between the number of amino groups in UBI-(NH$_3^+$)$_8$ and the maximum number of molecules of SDS (11) that bind at <1.8 mM. Thus, we conclude that the electrostatic interactions between UBI-(NH$_3^+$)$_8$ and 11 molecules of SDS (at <1.8 mM) are not occurring, exclusively, between the sulfate group of SDS and positively charged amino and guanidinium groups of the protein (although we cannot rule out the possibility that some lysine-$\varepsilon$-NH$_3^+$ groups are interacting with the SDS). We conclude, instead, that peracetylation inhibited the binding of SDS (at <1.8 mM) because it increased the net negative charge of UBI, or decreased the local positive electrostatic surface potential of regions on the surface of UBI that were binding SDS (possibly Ly residues or hydrophobic residues nearest the seven lysines or N-terminus).

Association of UBI-(NHAc)$_8$ with SDS is Reversible.

The formation of protein—protein aggregates (transient or irreversible) is a problem that complicates the study of interactions between proteins and chaotropic agents (e.g., protein folding). Protein aggregation is hypothesized to be the cause of numerous artifacts observed during studies of protein folding.$^{30}$ The aggregation of UBI, for example, has been hypothesized to cause its apparent three-state folding pathway, observed in guanidinium hydrochloride (a two-state pathway is observed in other studies).$^{31}$ We have previously demonstrated that the formation of complexes of SDS and UBI-(NH$_3^+$)$_8$ is reversible in tris-glycine buffer (at room temperature), and that very little protein is lost to aggregation during dialysis in different concentrations of SDS.$^4$ In order to prove that the association of SDS with UBI-(NHAc)$_8$ is also reversible, we prepared complexes of SDS and UBI-(NHAc)$_8$ from UBI-(NHAc)$_8$ that was initially unfolded, and also from UBI-(NHAc)$_8$ that was initially folded, and analyzed each with CE. Unfolded UBI-(NHAc)$_8$ was prepared by dialyzing solutions of UBI-(NHAc)$_8$ against SDS (up to 10 mM), as previously described for UBI-(NH$_3^+$)$_8$.$^4$ The two sets of electropherograms from these experiments are shown in Figure S2, and their similarity demonstrates that the unfolding of UBI-(NHAc)$_8$ is reversible in SDS (pH 8.4).

In order to determine whether the peracetylation of UBI significantly perturbed its structure or conformational stability, we also characterized UBI-(NHAc)$_8$ (in the absence of SDS) with CD (Figure S3), differential scanning calorimetry (DSC), and amide hydrogen—deuterium exchange (Figure S4). The similarity in the CD spectra of UBI-(NH$_3^+$)$_8$ and UBI-(NHAc)$_8$
more hydrophobic than UBI-(NHAc)₈·SDS₀ (because of the associated C₁₂H₂₅ chains) and UBI-(NH₃⁺)₈·(SDS)₁₁ might, therefore, interact with additional molecules of SDS more favorably (thermodynamically) than UBI-(NHAc)₈·SDS₀. Regardless of the explanation of this observation, G₄ intermediates, UBI-(NH₃⁺)₈·SDS₁₁ and UBI-(NHAc)₈·SDS₀ exist exclusively above 2.5 mM SDS (<3.5 mM), although they differ in charge by four units. We conclude that the presence of ϵ-NH₃⁺ groups is important for the formation of the first complex of UBI and SDS (e.g., UBI-SDS₁₁), but that ϵ-NH₃⁺ groups are not playing a major role thermodynamically during the intermediate stages of binding to SDS, and hydrophobic effects dominate electrostatics (although the latter are crucial in the latest stages of denaturation).

UBI-(NH₃⁺)₈·SDS₁₁ and UBI-(NHAc)₈ Unfold via the Binding of Indistinguishable Numbers of SDS Molecules. The mobilities of complexes of UBI-(NH₃⁺)₈ and UBI-(NHAc)₈ with SDS are approximately equal at concentration of SDS = 3.0 mM (μ ≈ 20 cm² kV⁻¹ min⁻¹) (Figure 1). The stoichiometries of the complexes at this concentration are, however, quite different: UBI-(NH₃⁺)₈·(SDS)₁₁ versus UBI-(NHAc)₈·(SDS)₁₄. The formation of G₄ complexes, therefore, involves binding 14 molecules of SDS by UBI-(NHAc)₈ and UBI-(NH₃⁺)₈·(SDS)₁₁.

The electrophoretic mobility of a complex of UBI and SDS can be approximated (eqs 1 and 2) as a function of its net charge (Z) and molecular weight (M). In eq 2, Z₀ is the net charge of the native UBI (Z₀(UBI-(NH₃⁺)₈) ≈ −0.2)₈, p is the number of acetylations, n is the number of equivalents of SDS bound to UBI-(NHAc)₈, (NH₃⁺)₈·(SDS)₁₁, ΔZ is the charge increment resulting from one acetylation or from the binding of one molecule of SDS, Mₐc is the increase in mass due to acetylation, MₛD is the molecular weight of dodecyl sulfate ions, Cₘ is a protein-specific constant, and α has a typical value of 2/3 for globular proteins. This value is consistent with the hydrodynamic drag being proportional to the surface area.

\[ μ = \frac{C_p Z}{M^\alpha} \]  

\[ μ = \frac{C_p}{(M_{UBI} + pM_{Ac} + nM_{SDS})^\alpha} \] 

We were able to estimate the stoichiometry of the complex making up G₄ for UBI-(NH₃⁺)₈ in our previous study (G₄ is UBI-(NH₃⁺)₈·(SDS)₁₄). The mobility of UBI-(NH₃⁺)₈·(SDS)₁₄ can be expressed using eq 1 (Z₀ = −0.2, p = 0, n = 25, ΔZ = −0.9, M_{UBI} = 8565 Da, and M_{SDS} = 265.4 Da) and can be set equal to the analogous expression for G₄ complexes of UBI-(NHAc)₈ (with p = 8, M_{Ac} = 42 Da), yielding n ≈ 14, and suggesting that 14 SDS molecules are bound to UBI-(NHAc)₈ at [SDS] = 3.0 mM. This number (n ≈ 14) is equal to the number of SDS molecules that we showed previously to bind to UBI-(NH₃⁺)₈ during the transition G₂→G₄. Complexes of UBI-(NHAc)₈ with SDS in G₄ therefore, have the composition UBI-(NH₃⁺)₈·(SDS)₁₄, while complexes of UBI-(NH₃⁺)₈ in G₄ have the composition UBI-(NH₃⁺)₈·(SDS)₁₁ during the transition G₂→G₄ and with UBI-(NHAc)₈ during the transition N→G₄ is unclear. The G₄ complex of UBI (e.g., UBI-(NH₃⁺)₈·(SDS)₁₁) is much more hydrophobic than UBI-(NHAc)₈ because the former has more aliphatic methylene groups on the surface of the protein.

Figure 2. SurfCE kinetic electropherograms of UBI-(NH₃⁺)₈ in the presence of SDS. UBI is injected in its native form (N) into a capillary previously filled with Tris-glycine and SDS, at a concentration of SDS labeled on each electropherogram. The ruler that is overlaid on the top of the electropherograms corresponds to the mass-calibrated mobilities labeled on each electropherogram. The ruler that is overlaid on the top of the electropherogram is reserved due to the equilibrium of bound and free SDS in the capillary.

indicated that the overall structures of UBI-(NH₃⁺)₈ and UBI-(NHAc)₈ were similar. Peracetylation did, however, increase the rate of amide hydrogen—deuterium exchange, and did lower the thermostability of the protein (Tₘ = 67 °C for UBI-(NHAc)₈; Tₘ > 90 °C for UBI-(NH₃⁺)₈), both DSC and hydrogen—deuterium exchange indicated that the peracetylated protein remained folded.

UBI-(NH₃⁺)₈·(SDS)₁₁ Unfolds at a Lower Concentration of SDS than Does UBI-(NHAc)₈. UBI-(NHAc)₈ begins to form a complex we denote “G₄” at a higher concentration of SDS (2.5 mM) than UBI-(NH₃⁺)₈, which begins to form the G₄ complex at concentrations as low as 1.4 mM SDS, although as a mixture with G₂ and G₃ intermediates (Figure 1). The G₄ complex of both UBI-(NH₃⁺)₈·(SDS)₁₁ and UBI-(NHAc)₈ exists up to 3.5–4.0 mM SDS, and is the only thermodynamically complex stable at 3.0 mM SDS. Our previous study⁴ of UBI-(NH₃⁺)₈ showed that the UBI-SDS complex G₄ is almost completely unfolded, based on the rate of H/D exchange at 3.0 mM SDS. The increase of α-helical content, measured by CD at 220 nm in both G₄ complexes, suggests they exist as mixed micelles with cylindrical shape. The lowered dielectric permittivity of the micelle surrounding the unfolded polypeptide increases the stability of H-bonds and promotes the formation of local folds (Figure S5). The rearrangement of the native β-sheet in UBI into an α-helix in mixed micelles is a slow and irreversible process, which yields complex G₄ upon extensive dialysis (170 h) with UBI with SDS. Electrophoresis of unmodified UBI-(NH₃⁺)₈ in the presence of SDS forms some G₄, but only as a mixture of complexes at higher concentrations of SDS (>5.0 mM) during a 10 min reaction in the capillary (Figure 2).

One explanation for the observation that UBI-(NH₃⁺)₈ forms the G₄ complex at a lower concentration of SDS than does UBI-(NHAc)₈ (by ~1 mM) is that UBI-(NH₃⁺)₈·(SDS)₁₁ is
Figure 3. Diagram summarizing the binding pathway of SDS to UBI-(NH₃⁺)₈ (blue) and peracetetylated UBI (red). The numbers of SDS molecules that are bound to each protein at different concentrations of SDS (0–10 mM SDS) are shown. The numbers of additional SDS that bind to each protein along the pathway of unfolding are also shown. Note: the structures of UBI-SDS complexes near the micellar regime are purely speculative and may (in actuality) not involve the types of configurations that are shown (i.e., the so-called “pearl necklace” or “bead on a string” configuration).

compared with UBI-(NHAc)₈, and this greater hydrophobicity of UBI-(NH₃⁺)₈(SDS)₋₁₁ should render it more favorable for interactions with SDS than UBI-(NHAc)₈. The UBI-(NH₃⁺)₈(SDS)₋₁₁ complex is also, however, more negatively charged than UBI-(NHAc)₈, and this greater negative charge could effectively cancel out any favorable hydrophobic interactions between UBI-SDS₁₁ and additional molecules of SDS.

The binding of SDS to UBI-(NHAc)₈ and UBI-(NH₃⁺)₈ is similar above the CMC; the shape and mobilities of the electropherograms are almost indistinguishable for UBI-(NH₃⁺)₈ and UBI-(NHAc)₈ (Figure 1, [SDS] > 3.5 mM). Calculating the number of SDS molecules that are bound to UBI-(NHAc)₈ and UBI-(NH₃⁺)₈ at >3.5 mM (with eq 2) demonstrated, however, that UBI-(NHAc)₈ has ~11 fewer SDS molecules bound than UBI-(NH₃⁺)₈ under denaturing conditions (∼CMC).

A Kinetically Stable Ubiquitin-Poly(dodecyl sulfate) Complex (e.g., UBI-(NH₃⁺)₈(SDS)₇) Forms on the Path from N to G₂. The CE experiments that we have done have involved complexes of UBI and SDS that are formed at equilibrium, by prolonged exposure of UBI to solutions of SDS. We are also interested, however, in studying the kinetics of binding of SDS to UBI. We investigated the kinetics of binding of SDS to UBI-(NH₃⁺)₈ in the transitions N → G₁ → G₂ by using a technique we refer to as surfactant capillary electrophoresis (SurfCE). SurfCE is a technique that makes it possible to observe both the rate and stoichiometry of association of SDS with proteins, in situ (e.g., during capillary electrophoresis). In SurfCE, the protein is injected in its native, folded form into a capillary that is filled with a solution of SDS at a selected concentration; the reaction time therefore corresponds to the time of electrophoresis (e.g., ∼7 min with a 60-cm capillary). Rapid association of proteins with SDS is reflected in an immediate conversion of the peak corresponding to N into a new peak with a higher mobility. The slow association of SDS to proteins or a slow equilibration between multiple aggregates with different compositions will result in a broad peak.

Analysis of the interaction of UBI-(NH₃⁺)₈ with SDS by SurfCE (Figure 2) revealed several additional complexes with mobilities between the peaks N and G₂ (i.e., 1.5 < μ < 14 cm² kV⁻¹ min⁻¹); these ubiquitin-poly(dodecyl sulfate) complexes are not present in the equilibrium electropherograms for UBI-(NH₃⁺)₈ (Figure 1). The peak at mobility μ ≈ 11.5 cm² kV⁻¹ min⁻¹ is present over a narrow range of SDS concentrations (1.2 < [SDS] < 3.0 mM). The constant mobility of this peak suggests that a particular species with a discrete stoichiometry is formed along the path from N to G₂. Since this peak is absent in electropherograms of equilibrated mixtures (Figure 1a), we infer that this complex is kinetically labile, and forms along the pathway from N to G₂. Comparison of the mobility of this peak with the mass-calibrated mobilities of the rungs of UBI charge ladder (rulers overlaid on top of the electropherograms in Figure 2) indicates that this complex has the composition UBI-(NH₃⁺)₈(SDS)₇.

Monitoring the Binding of SDS to Residues in UBI-(NH₃⁺)₈ with ¹⁵N–¹H HSQC. Figure 3 summarizes the thermodynamic pathway of association of SDS with UBI-(NH₃⁺)₈ and UBI-(NHAc)₈. The complex UBI-(NH₃⁺)₈(SDS)₇ described in the previous section is an unstable intermediate on the pathway of formation of the first stable complex UBI-(NH₃⁺)₈(SDS)₁₁. In order to determine which residues in UBI-(NH₃⁺)₈ interact with SDS at concentrations below the CMC, we studied UBI-SDS complexes with ¹⁵N–¹H HSQC NMR. We focused on interactions between SDS and UBI-(NH₃⁺)₈ at 1.0 mM SDS, because the protein retains its fold at this concentration and forms only a single complex, UBI-(NH₃⁺)₈(SDS)₁₁, according to CE experiments.

With ¹⁵N–¹H HSQC, we were able to monitor the association of SDS to amino acid residues of UBI-(NH₃⁺)₈ in 1 mM SDS by measuring the movement in the ¹⁵N–¹H correlation signal (Δδ) of each residue (Figures 6 and 7). The value Δδ represents the Euclidian distance (adjusted for the characteristic frequency of precession of N and H) by which each ¹⁵N–¹H correlation signal was shifted, calculated by eq 3:

$$
\Delta \delta = \sqrt{(\Delta ^{15}N / S)^2 + \Delta ^{13}H^2}
$$

One important limitation of ¹⁵N–¹H HSQC NMR as a tool for studying SDS binding to UBI at 1 mM is that there is an interaction between a specific residue and surfactant will not be detected if the association occurs at a rate faster than the rate of the ¹⁵N–¹H HSQC scan, which occurs on a millisecond time scale. For example, a portion of the dodecyl chain of an SDS molecule might be mobile (to some degree) when bound to the surface of UBI, and different gauche rotamers might interact with different residues. If dynamic rotamer interactions are occurring, and a particular SDS molecule is populating multiple rotamers with different chemical shifts, then we would only be able to detect those rotamer interactions that are populated longer than milliseconds. The majority of ¹⁵N–¹H resonances were
Lys6, Leu67, and Val70. Residue labels are positioned next to the 

The correlation signal of Lys6 was not shifted by the presence of 1 mM SDS, indicating that Lys6 did not interact with SDS; Leu67 and Val70 do interact with SDS, as demonstrated by the shifting of their signals from 0 to 1 mM SDS. (b) Expanded version of full spectra shown in (a); residue labels are positioned next to the 

The 15N\(^{-1}\)H correlation signal for two residues (Glu24, Ala46) were too weak to be detected at 0 or 1 mM SDS; the chemical shifts for three residues (Arg42, Gln49, and Arg72) could not be determined at 1 mM SDS because the 15N\(^{-1}\)H correlation signal in 1 mM SDS disappeared entirely, or could not be assigned to a residue with confidence. In addition, the three proline residues (Pro19, Pro37, and Pro38) were also not observable, because proline does not have a backbone amide N–H. 15N\(^{-1}\)H correlation signals for four other residues in UBI are unassigned at 0 mM SDS and pH 7.4: Met1, Thr9, Gly10, and Arg74.

The values of Δ\(\delta\) for several residues of UBI-(NH\(^3\))\(_8\) at 0–1 mM SDS were close to zero (e.g., 0.00 < Δ\(\delta\) < 0.02 ppm; Lys 6, Thr12, Glu16, Val17, Glu18, Ser20, Asp21, Thr22, Asp39, Gln40, Leu56; Tyr59, and Asn60; Figure 5); we conclude that these residues did not bind SDS, or only bound SDS weakly. The 15N\(^{-1}\)H correlation signals for other residues of UBI, however, shifted considerably on going from 0 to 1 mM SDS, with 0.15 < Δ\(\delta\) < 0.30 ppm, and we conclude that these residues specifically bind SDS (i.e., Val5, Leu8, Lys11, Ile13, Thr14, Leu43, Ile44, Leu50, His58, Val70, and Leu71; Figure 5). Table 1 (below) lists the residues of UBI-(NH\(^3\))\(_8\) with the largest and smallest values of Δ\(\delta\) in 0–1 mM SDS. The Supporting Information contains a complete list for all 63 residues.

A comparison of the values of Δ\(\delta\) with a map of the secondary structure of UBI (Figure 5) revealed that the residues contained in hydrogen-bonded loops in UBI were least affected by SDS (in comparison to residues in the β-sheet or α-helix); this observation suggests that SDS does not interact specifically with residues in loops (e.g., residues 18–22, 39–40, and 56–59; residues 39–40 and 56–59 have also been described by some as a 310-helix).

Most of the residues in UBI-(NH\(^3\))\(_8\) with the largest values of Δ\(\delta\) are hydrophobic residues with no formal charge (Figure 5, Table 1). Some residues that have a formal positive charge (and

**Figure 4.** Studying the binding of SDS to UBI at 1 mM SDS with 15N\(^{-1}\)H HSQC NMR. (a) HSQC spectra of human ubiquitin (labeled with 15N) in the presence of 0 and 1 mM SDS (pH 7.53, 25 mM Tris); the concentration of ubiquitin was 150 \(\mu\)M. An expanded view shows Lys6, Leu67, and Val70. Residue labels are positioned next to the 15N\(^{-1}\)H correlation signal for each residue at 0 mM SDS (blue). The correlation signal of Lys6 was not shifted by the presence of 1 mM SDS, indicating that Lys6 did not interact with SDS; Leu67 and Val70 do interact with SDS, as demonstrated by the shifting of their signals from 0 to 1 mM SDS. (b) Expanded version of full spectra shown in (a); residue labels are positioned next to the 15N\(^{-1}\)H correlation signal for each residue at 0 mM SDS (blue).
are expected to be positively charged in UBI) did exhibit large changes in chemical shift in the presence of 1 mM SDS (i.e., Lys11 and Lys29), and we conclude that these residues are specifically interacting with SDS; several residues with a formal negative charge showed little or no perturbation (i.e., Glu18, Asp21, and Asp39). This correlation, however, was not uniformly observed for ionic residues: the least perturbed residue in UBI-(NH$_3$$^+$)$_8$ at 1 mM SDS, for example, was Lys6 (e.g., $\Delta\delta = 0.00180$), and we conclude that Lys6 did not associate with SDS, whereas a large change in chemical shift was observed for Asp32 from 0 to 1 mM SDS (e.g., $\Delta\delta = 0.14069$). An analysis of the structure of UBI obtained by either NMR or X-ray crystallography, indicates that Lys6 is exposed to solvent and that the $\varepsilon$-NH$_3^+$ is not hydrogen-bonded with any nearby residues (Figure S6). The reasons why SDS does not associate with this solvent-exposed, positively charged residue are unknown. We do expect that an electrostatic interaction between the $\varepsilon$-NH$_3^+$ of Lys6 and the ROSO$_3^-$ of SDS can be detected with $^1$H-$^1$N HSQC. Flynn and co-workers have demonstrated, in studies that aimed to mimic molecular confinement and crowding, that Lys6 of UBI has the fifth-largest value of $\Delta\delta$ (of any residue) when the reverse micelles of sodium bis(2-ethylhexyl)sulfosuccinate are decreased in size (by varying the amount of water in the organic–aqueous–micellar mixture). We hypothesize that the large value of $\Delta\delta$ for Lys6 in Flynn’s study is likely to be the result of a direct interaction between R-OSO$_3^-$ and Lys-$\varepsilon$-NH$_3^+$.

The $^{15}$N-$^1$H HSQC experiments provided incomplete data on the association of arginine residues with SDS. One of the four arginines (Arg 74) could not be assigned at 0 mM SDS, and correlation signals for two others (Arg43, Arg72) could not be assigned at 1 mM SDS, suggesting that these amides interact with surfactant on an intermediate time scale. A fourth arginine, however (Arg54), was observed at 0 and 1 mM SDS, and the value of $\Delta\delta$ was small ($\Delta\delta = 0.02433$); we infer that Arg54 did not associate with SDS, or at least did not associate with SDS as often, or as tightly, as did other residues in UBI with positive formal charge (e.g., Lys11; $\Delta\delta = 0.17593$).

We did not analyze UBI-(NH$_2$Ac)$_8$ with HSQC because the acetylation of lysine residues (and the amino terminus) will shift the $^{15}$N-$^1$H correlation signals of residues in unpredictable directions, so that most $^{15}$N-$^1$H correlation signals would need to be completely reassigned to each residue (the experimental work required to do these assignments is beyond the scope of this study). Nevertheless, our analysis of UBI-(NH$_3^+$)$_8$ at 0–1 mM SDS with HSQC shows that three solvent-exposed, cationic residues—Lys6, Lys48, and Arg54—have a small value of $\Delta\delta$ (i.e., $\Delta\delta < 0.03$), indicating that these residues do not specifically bind SDS. Therefore, the results of our analysis of UBI-(NH$_3^+$)$_8$ with HSQC do not support the hypothesis—based on the stoichiometry inferred from CE experiments (Figure 1)—that the negatively charged sulfate groups of SDS associate uniformly with positively charged functional groups (lys-$\varepsilon$-NH$_3^+$) more strongly than Arg-NHC($\equiv$NH$_2^+$)NH$_2^+$ on the surface of UBI-(NH$_3^+$)$_8$ in approximately 1:1 stoichiometry at 0.8–1.4 mM SDS. The results of HSQC do, however, suggest that some cationic residues (e.g., Lys11 and Lys29) are associating, in some fashion, with SDS. A comparison of $^{15}$N-$^1$H HSQC data with electrostatic surface potentials (described below) offers an explanation for this unexpected lack of correlation between $\Delta\delta$ and formal charge for each residue in UBI.

Monitoring the Interaction of Side Chains of UBI and SDS with $^{13}$C-$^1$H HSQC. We carried out $^{13}$C-$^1$H HSQC experiments to supplement the results of $^{15}$N-$^1$H HSQC experiments that could potentially miss interactions between UBI and SDS. For example, $^{13}$C-$^1$H HSQC suggests that SDS is not interacting with Lys6; the backbone amide of Lys6 could, however, possibly be unperturbed by electrostatic interactions between the R-OSO$_3^-$ and $\varepsilon$-NH$_3^+$ groups. Figure 8 shows the overlay of $^{13}$C-$^1$H HSQC spectra of UBI with 0 and 1 mM SDS. This plot of $^{13}$C-$^1$H HSQC spectra reveals a striking attenuation of the intensities of the majority of resonances in the presence of 1 mM SDS. The resonances corresponding to the majority of $\alpha$-CH and $\beta$-CH peaks are completely attenuated in the presence of 1 mM SDS, while the remaining few include the $\alpha$-CH of Ile23 and $\beta$-CH of Ala28 and Ala46. The resonance of the methyl group of Ala28 is virtually stationary, while those of Ala46 and $\alpha$-CH of Ile23 are significantly shifted in the presence of 1 mM SDS. These results suggest that Ala46 and Ile23 are involved in specific interactions with SDS, but Ala28 is not. Correspondingly, the amide chemical shift of I23 ($\Delta\delta = 0.11$) is significantly greater than that of A28 ($\Delta\delta = 0.02$). Similarly, for a pair of valines, movement of any $^{13}$C-$^1$H resonances correlates with greater amide shifts: in Figure 8, $\gamma$-CH resonances of V17 and V26 move only slightly ($^{15}$N-$^1$H $\Delta\delta = 0.012$, 0.02), while $\gamma$-CH of V70 shifts dramatically and corresponds to a greater amide shift ($\Delta\delta = 0.15$).

Several of the few remaining resonances in the $^{13}$C-$^1$H HSQC spectra of UBI in 1 mM SDS correspond to the methylenes of lysines; $\varepsilon$-CH, $\delta$-CH, and $\gamma$-CH. Although the latter do not exhibit significant chemical shifts, we can infer that underlying interactions with SDS result in bound states with lifetimes longer than milliseconds. The existence of such resonances (e.g., $\gamma$-CH’s of K29, K33, and K63) correlates with the larger $\Delta\delta$ of amide HSQC. The resonances of HSQC signals for K6, however, are neither shifted or significantly attenuated in the presence of 1 mM SDS: we easily resolved the $\delta$-CH, $\gamma$-CH, and $\varepsilon$-CH of K6. This lack of perturbation or attenuation of K6 when studied by either $^{13}$C-$^1$H or $^{15}$N-$^1$H HSQC provides support for the conclusion that SDS does not interact with this highly solvent-exposed lysine residue. Other lysines, such as K27, also produced very small changes in amide chemical shifts, but the $\varepsilon$-CH was completely attenuated. The alkyl protons of SDS contribute to spin–lattice relaxation of aliphatic protons of UBI. The similar energy levels of these protons on the protein and SDS allow for undesirable transfers of magnetization, but these require proximity. The loss of the vast majority of resonances, including those from the hydrophobic core of UBI in the presence of 1 mM SDS, indicates that the backbone is entirely exposed to interactions with the n-alkyl tail. This result suggests that the n-alkyl tails of the surfactant transiently penetrate the hydrophobic core of the protein, pushing the strands of the backbone of UBI apart (e.g., the protein-SDS complex undergoes “breathing” motions).

The greater effect of SDS on relaxation of $^{13}$C-$^1$H magnetization than on that of $^{15}$N-$^1$H suggests that the alkyl tails make a unique contribution to relaxation of aliphatic protons. This relaxation must come from the undesirable mode of relaxation of magnetization—spin–spin or transverse (T2 relaxation)—rather than the typically faster spin–lattice (longitudinal or T1 relaxation). The longitudinal relaxation, which is strongly dependent on the size of the protein, would affect both carbon and nitrogen equally. The $^{13}$C-$^1$H spectrum at 1 mM SDS shows resonances for methyl moieties from valine, leucine, isoleucine, and threonine residues, as well as a number of methylene groups.
These groups are known to have slow rates of T1 relaxation, because of free and rapid rotation, so we conclude that dynamic, low-affinity (nonspecific) interactions of SDS contribute to spin–spin relaxation of aliphatic protons in UBI. This dynamic interaction of 1 mM SDS affects both Hα and side chains but does not disrupt the H-bonding network or the secondary structure of UBI according to CD and hydrogen/deuterium exchange.4

These observations suggest that the effects of SDS on the side chains of UBI fall in two classes: weak, transient interactions with aliphatic methyl and methylene protons, and stable interactions with sub-millimolar affinity and millisecond lifetimes.

SDS (1 mM) Interacts Preferentially with Hydrophobic Residues in UBI-(NH₃⁺) That Have a Positive Electrostatic Surface Potential. We hypothesized that a better understanding might emerge as to why SDS associates with surface residues in UBI-(NH₃⁺)₈ that have no formal charge (i.e., Ile13 or Leu43), and not with other formally neutral residues or residues with formal positive charge (i.e., Lys6, Val17, and Thr22)—and why the acetylation of lysine inhibits the binding of SDS to UBI—when we took into consideration the local electrostatic potential at the surface of a folded protein. We used a nonlinear PB calculation to approximate the electrostatic potential of the solvent-accessible surface of UBI. Figure 6 shows a comparison of the electrostatic surface potential of the solvent-accessible surface of UBI-(NH₃⁺)₈ with the value of Δδ for each residue during HSQC experiments at 0 and 1 mM SDS. Red represents residues with small chemical shift perturbations; white, intermediate chemical shift perturbations; blue, large chemical shift perturbations. Gray indicates residues that could not be studied (i.e., proline) or residues that could not be assigned a ^15N–^1H correlation signal. Surface-exposed lysine (K) residues and the single histidine residue (H) are labeled. The surface of UBI that was most affected by 1 mM SDS was made up of nonpolar, hydrophobic residues that existed in a positively charged environment: Leu71, Leu43, Leu8, Val70, Ile44, and Gly47. The surface of UBI that was least affected by SDS was a polar region, containing multiple glutamate and aspartate residues (and one valine), that was negatively charged: Glu16, Val17, Glu18, Asp21, and Thr22. The lone arginine that was observable in HSQC experiments (Arg54) had a small value of Δδ (0.02433 ppm) and is electrostatically neutral, but is also flanked on three sides by residues with a negative electrostatic surface potential. Remarkably, Lys6 and Lys48 also have a small value of Δδ (Lys6 has the smallest Δδ of any residue at 1 mM SDS), but both residues have a positive electrostatic surface potential and are surrounded by other positively charged residues.

![Image](https://dx.doi.org/10.1021/ja205735q)
(PDB code 1UBI) for the electrostatic calculations and for making a color-coded surface rendering of UBI that illustrates the magnitude of \( \Delta \delta \) for each surface residue of UBI at 0–1 mM SDS.

We have also colored the surface of UBI according to the magnitude of the chemical shift \( \Delta \delta \) at 0–1 mM SDS. Figure 6 shows the correlation between the local electrostatic surface potential of native UBI and the magnitude of \( \Delta \delta \) at 0–1 mM SDS. In general, a residue contained within patches of UBI-(NH\(_3^+\))\(_8\) with positive electrostatic potential interacted with SDS more tightly (or strongly) than did a residue with a negative electrostatic surface potential—irrespective of the formal charge of the residue (Figure 6a,b). It is worth noting that UBI-(NH\(_3^+\))\(_8\) is, electrostatically speaking, a Janus-faced protein.\(^{37,38}\) The electrostatic potential of surface residues of UBI-(NH\(_3^+\))\(_8\) seems to reveal why several residues, including hydrophobic residues with no formal charge, have larger values of \( \Delta \delta \) than other residues, such as lysine or arginine, which have a positive formal charge. The nine hydrophobic residues of UBI-(NH\(_3^+\))\(_8\) that have the largest values of \( \Delta \delta \) in 1 mM SDS (e.g., Ile13, Leu43, Leu71, Thr14, Leu50, Leu8, Gly47, Val70, Ile44; \( 0.15967 < \Delta \delta < 0.31160 \)) are located in regions that have a local positive electrostatic potential (Figure 6a)—that is, in regions where positive charge is not screened by solvent. In fact, Leu8, Val70, Leu71, and Ile44 are all clustered together in the same hydrophobic face of UBI-(NH\(_3^+\))\(_8\) with a positive electrostatic potential (Figure 6a, top left image). We note that this hydrophobic face (i.e., the “Ile44 face”) of UBI comprises a \( \beta \)-sheet and has been identified, previously, to be the site of recognition of UBI by multiple families of proteins, including UBA\(_3\) (ubiquitin-associated domains), UIMs\(^{40}\) (ubiquitin-interacting motifs), CUB\(^{41}\) (coupling of ubiquitin to ER degradation), and NZFs\(^{42}\) (N1p14 zinc fingers), as well as sites of UBI-UBI interactions of poly-ubiquitin chains.\(^{43}\)

In contrast, residues that have the lowest values of \( \Delta \delta \) at 0–1 mM SDS are located on the opposite face of UBI, and in a region with negative electrostatic potential (e.g., Val17, Thr22, Tyr59, and Asn60; \( 0.01098 < \Delta \delta < 0.01468 \)). We also point out that the only arginine residue for which we have data, Arg54, has a small value of \( \Delta \delta \) (\( \Delta \delta = 0.02433 \)), and that moreover, Arg54 does not have a positive electrostatic surface potential (according to the PB calculation). The approximately zero electrostatic surface potential of Arg54 (Figure 6a), calculated from the PB equation, is caused by (according to the model and its approximations) a high degree of solvation of the guanidino group of Arg54; the highly solvated nature of Arg54 might explain why SDS does not have a specific electrostatic interaction with Arg54. Moreover, Arg54 is flanked on three sides by residues with a negative electrostatic potential, which will contribute to the neutralization of the surface potential of Arg54 (via screened Coulombic interactions). A previous study by Flynn and co-workers showed that Arg54 had the second lowest chemical shift perturbation of any residue in UBI, when folded UBI was confined inside reverse micelles of sodium bis(2-ethylhexyl) sulfosuccinate (in this study, \( \Delta \delta < 0.005 \) for Arg54). We also rationalize these previous observations in terms of the zero electrostatic surface potential of Arg54 and its surrounding by anionic residues.

The major finding of Figures 6–8—that SDS interacts preferentially (below the CMC) with hydrophobic residues that have positive electrostatic surface potential—is consistent with the result shown in Figure 1, that the acetylation of seven lysine-\( \epsilon \)-NH\(_3^+\) (and one \( \alpha \)-NH\(_3^+\)) inhibits the binding of 11 SDS below the CMC. We conclude that the peracetylation of UBI-(NH\(_3^+\))\(_8\) inhibits the binding of SDS to (primarily) hydrophobic residues on the surface of UBI, and to a lesser degree cationic or polar residues, because peracetylation affects the electrostatic surface potential of these hydrophobic residues, and not because peracetylation neutralized specific R-NH\(_3^+\) sites that could have been points of contact with the sulfate group of SDS. The amino groups of lysine are, therefore, important for the binding of SDS to UBI—not necessarily because they represent cationic sites where the sulfate group can bind, but because the cationic groups are (according to nonlinear PB methods) a source of positive electrostatic surface potential that extends beyond their own van der Waals radii.

The electrostatic surface potential of amino acid residues in UBI cannot, however, entirely explain the degree with which these residues associate with SDS: Thr12 and Lys6, for example, both have positively charged surfaces but have low values of \( \Delta \delta \) (Lys6 = 0.01109 and Thr12 = 0.00180); Asp32 has one of the largest values of \( \Delta \delta \) for any residue (0.14069) but has a negative electrostatic potential, and is surrounded by many negatively charged residues (Figure 6). We have no explanation for these observations involving Lys6, Thr12, and Asp32.

The general observation that hydrophobic residues in UBI-(NH\(_3^+\))\(_8\) have larger values of \( \Delta \delta \) than cationic residues (at 0–1 mM SDS) demonstrates that hydrophobic interactions between the dodecyl chains of SDS are a large driving force for the specific binding of SDS to UBI. In order to begin to quantify how the hydrophobicity of a residue in UBI-(NH\(_3^+\))\(_8\) affects its association with SDS, we plotted the average value of \( \Delta \delta \) for each type of amino acid (i.e., lysines, leucines, aspartates, etc.) as a function of their hydrophobicity (Figure 7). The values of

\[
R^2 = 0.7552 \\
m = 0.0162 
\]

\[
R^2 = 0.7693 \\
m = 0.0200 
\]
The chemical shifts of amides and modeling the surface potential suggest that the highest affinity sides are hydrophobic residues in regions of positive electrostatic potential, such as those involved in formation of the first stable intermediate, UBI-(NH$_3^+$)$_8$(SDS)$_{11}$.

A recent study by Xiao et al. has used $^1$H–$^{15}$N HSQC NMR spectroscopy to monitor the binding of sodium perfluorooctanoate (SPOF, an anionic surfactant) to UBI (0.5 mM, pH 5.7), below and above the CMC (the CMC of SPOF was reported to be 31.0 mM).$^{15}$ This study reported that the structure of UBI is largely unaffected by concentrations of SPOF < 3 mM. Even though the Xiao et al. study involved a surfactant with a perfluorinated tail and studied UBI below its isoelectric point, we find that our observations are qualitatively similar. The previous study by Xiao et al. supports, for example, our observation that Lys6 and Arg54 are not associated with SDS at 1 mM SDS: the $^1$H–$^{15}$N correlation signals for Lys6 and Arg54 were not affected—either in intensity or in position—by the presence of 2.9 mM SPOF.

The Conformational Flexibility and Solvent Accessibility of a Residue in UBI-(NH$_3^+$)$_8$ and Its Association with SDS. We also suspected that other biophysical properties of residues in UBI, such as the conformational flexibility or solvent accessibility, might also contribute to the binding of SDS with UBI below the CMC. Because UBI is a model protein that has been studied in depth by structural biologists, there are biophysical and structural data available for most amino acids of the folded protein. In order to determine how the structural properties of residues of UBI-(NH$_3^+$)$_8$ might influence their association with SDS—and perhaps explain the $\Delta$ values of some residues (i.e., Lys6, Thr12, Asp32)—we compared the values of $\Delta$ from $^{15}$N–$^1$H HSQC at 0–1 mM SDS with some of these available data, including (i) the solvent accessibility of the side chain ($\text{SA}_{\text{side}}$) and main chain ($\text{SA}_{\text{main}}$) of each residue, as estimated from the crystal structure of UBI, and (ii) the rate of amide hydrogen/deuterium (H/D) exchange for each residue in UBI (measured with NMR). Figure S7 (Supporting Information) shows a comparison of $\Delta$ values of each residue in 0–1 mM SDS, and the static solvent accessibility of the side chain and main chain of each residue in folded UBI. A comparison of the $\Delta$ of each residue in 0–1 mM SDS and the rate of amide hydrogen—deuterium exchange of each residue is also shown in Figure S7. We did not observe any general correlation between the $\Delta$ of a residue, its accessibility to solvent, or the rate of amide hydrogen—deuterium exchange of that residue. A detailed discussion of these results is included in the Supporting Information.

Characteristics of the Binding Sites of SDS on the Surface of Folded Ubiquitin. Our analysis of UBI at 0 and 1 mM SDS with $^{15}$N–$^1$H and $^{13}$C–$^1$H HSQC, and a comparison of these results with the electrostatic surface potential, secondary structure, hydrophobicity, solvent accessibility, and rate of H/D exchange of residues in UBI, has shown that the first stable intermediate with SDS stably bound to UBI, such as occurs at 1 mM SDS, largely involves regions where hydrophobic side chains are adjacent to positively charged residues. The association of SDS to specific amino acids in folded UBI is not strongly influenced by type of secondary structure, whether located in a $\beta$-sheet or an $\alpha$-helix; however, the loops are characterized by small chemical shifts (Figure 6). Presumably, the loops provide a less-stable docking platform for SDS than either a $\beta$-sheet or an $\alpha$-helix. The results of this analysis are summarized in Table 1, which lists some of the structural, biophysical, and physical-organic properties of the 12 residues that were most affected by their interaction with SDS. Several peaks remain and are unchanged in 1 mM SDS (i.e., $\varepsilon$-CH and $\gamma$-CH of several Lys, and $\delta$-CH of Arg54 and Arg72), indicating no interaction with SDS. Several $\gamma$-CH and $\delta$-CH of Ile and Leu remain but are significantly shifted (denoted with curved arrows), indicating a specific interaction with SDS. Notably, the $\gamma$-CH, $\delta$-CH, and $\varepsilon$-CH of Lys6 (a solvent-exposed lysine residue) are not attenuated or shifted by the presence of 1 mM SDS. The resonance labeled with an asterisk is likely the $\alpha$-CH of Gly76 on the freely rotating C-terminus of UBI.

The hydrocarbon chain of SDS is linear and saturated and is therefore similar, structurally and chemically, to the alkyl chain of octanol (with the exception of four additional methylene groups).$^{44}$

We observed a significant correlation between the interaction of well-represented amino acids in UBI with 1 mM SDS and their hydrophobicity (Figure 7). This plot shows that the $^{15}$N–$^1$H resonance signals of hydrophobic amino acids are more perturbed by 1 mM SDS than hydrophilic residues. A linear least-squares regression of each data set in Figure 7 yielded $R^2 \approx 0.75$. We infer from this correlation that the SDS associates preferentially with hydrophobic side chains, not hydrophilic or cationic amino acids.

Figure 8. Studying the binding of SDS to UBI at 1 mM SDS with $^{13}$C–$^1$H HSQC NMR. HSQC spectra of human ubiquitin (uniformly labeled with $^{13}$C) in the presence of 0 and 1 mM SDS (pH 7.53, 25 mM Tris). The majority of $^{13}$C–$^1$H signals are attenuated by the interaction with SDS. Several peaks remain and are unchanged in 1 mM SDS (i.e., $\varepsilon$-CH and $\gamma$-CH of several Lys, and $\delta$-CH of Arg54 and Arg72), indicating no interaction with SDS. Several $\gamma$-CH and $\delta$-CH of Ile and Leu remain but are significantly shifted (denoted with curved arrows), indicating a specific interaction with SDS.

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**Characteristics of the Binding Sites of SDS on the Surface of Folded Ubiquitin.** Our analysis of UBI at 0 and 1 mM SDS with $^{15}$N–$^1$H and $^{13}$C–$^1$H HSQC, and a comparison of these results with the electrostatic surface potential, secondary structure, hydrophobicity, solvent accessibility, and rate of H/D exchange of residues in UBI, has shown that the first stable intermediate with SDS stably bound to UBI, such as occurs at 1 mM SDS, largely involves regions where hydrophobic side chains are adjacent to positively charged residues. The association of SDS to specific amino acids in folded UBI is not strongly influenced by type of secondary structure, whether located in a $\beta$-sheet or an $\alpha$-helix; however, the loops are characterized by small chemical shifts (Figure 6). Presumably, the loops provide a less-stable docking platform for SDS than either a $\beta$-sheet or an $\alpha$-helix. The results of this analysis are summarized in Table 1, which lists some of the structural, biophysical, and physical-organic properties of the 12 residues that were most affected by their interaction with SDS. Several peaks remain and are unchanged in 1 mM SDS (i.e., $\varepsilon$-CH and $\gamma$-CH of several Lys, and $\delta$-CH of Arg54 and Arg72), indicating no interaction with SDS. Several $\gamma$-CH and $\delta$-CH of Ile and Leu remain but are significantly shifted (denoted with curved arrows), indicating a specific interaction with SDS. Notably, the $\gamma$-CH, $\delta$-CH, and $\varepsilon$-CH of Lys6 (a solvent-exposed lysine residue) are not attenuated or shifted by the presence of 1 mM SDS. The resonance labeled with an asterisk is likely the $\alpha$-CH of Gly76 on the freely rotating C-terminus of UBI.
1 mM SDS (e.g., $\Delta \delta > 0.14$), and the properties of the 12 residues that were the least affected by 1 mM SDS (e.g., $\Delta \delta < 0.02$).

There are few structures available in the Protein Data Bank of SDS coordinated to folded proteins. The only example that we could find is the recent X-ray data of lysozyme$^{45}$ obtained from protein crystals that were grown in the presence of 2 mM SDS: a single dodecyl sulfate molecule was observed to be bound at the surface of lysozyme.$^{45}$ Details of this X-ray crystal structure, and details of X-ray crystal structures of other proteins coordinated with sulfatide, are included in the Supporting Information.

Predicting the structure of SDS molecules that are bound to the surface of UBI (i.e., at 1 mM SDS) with computational docking methods—for example, predicting where the sulfate group binds, and the conformation of the alkyl tail—is likely to be difficult because the binding of each additional SDS molecule will alter both the electrostatic surface potential of UBI and the hydrophobicity. From considering the results of Figure 7—that hydrophobic functional groups bind SDS more favorably than ionic or polar groups—we find no reason to exclude the possibility that the alkyl chains of different SDS molecules, which are bound to the surface of folded UBI, are interacting with one another.

**Electrostatic and Hydrophobic Interactions between SDS and UBI Occur Cooperatively.** Does UBI recognize the hydrophobic tail of SDS more than the anionic head at concentrations of 1 mM SDS? Our CE and HSQC results demonstrate that native UBI-(NH$^3_3$)$_8$ provides a type of surface (hydrophobic, but with a positive electrostatic surface potential) that allows electrostatic and hydrophobic interactions between SDS and UBI to occur cooperatively. The residues in UBI that were found to have the largest values of $\Delta \delta$ (at 1 mM SDS) were hydrophobic residues that had the greatest magnitude of positive electrostatic surface potential (i.e., Ile13, Leu 43; Table 1). The majority of the residues with the least interaction with SDS were hydrophobic residues that had the greatest magnitude of negative electrostatic surface potential (i.e., Val17, Thr22; Table 1). We infer from the HSQC analysis that salt-bridges are not forming between the sulfate group of SDS and positively charged groups of lysine and arginine (although salt bridges between some lysine or arginine residues might be forming). This absence of a general interaction between SDS and Lys or Arg is not so surprising when the electrostatic surface potential of UBI (Figure 6) is taken into account: lysine and arginine residues in UBI do not always have the largest positive electrostatic surface potentials (e.g., Arg54 $\approx$ 4 eV; Ile13 $\approx$ 4 eV) because, presumably, solvent is screening the charge of these residues. Therefore, the location of a salt-bridge—if one were to form between UBI and SDS—would be with a residue bearing a poorly solvated charge.

**CONCLUSIONS**

The hydrophobicity and surface potential (not formal charge) of an amino acid residue in folded UBI appear to be more important than secondary structure and solvent accessibility in explaining its association with SDS below the CMC. The strength of association of residues in UBI and SDS at sub-denaturing concentrations (<1.0 mM) seems to be equally governed by the residue’s hydrophobicity and its local electrostatic environment, but neither property alone is sufficient to explain the binding of SDS to folded UBI. A third property (the $2^\alpha$ structure of the residue) appears to play a lesser role (Table 1): residues in hydrogen-bonded loops appear to be protected from interacting with SDS; however, SDS did not prefer residues in $\alpha$-helical structure over those in $\beta$-sheet structure. We infer that—at least for UBI—the presence of (formally) cationic functional groups is important for the binding of SDS because cationic sites provide positive electrostatic surface potential that extends beyond the van der Waals radius of the functional group (toward formally neutral hydrophobic regions), and not necessarily because the cationic site forms an ionic bond (“saltbridge”) with the sulfate group (although these interactions might occur for some cationic residues, such as Lys11, but not for others, such as Lys6 and Arg54). In summary: (i) surfactants bind preferentially to hydrophobic regions with positive

<table>
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* $^a$ eV$_{local}$ refers to electrostatic potential in Figure 6; $^2^\alpha$ refers to $\alpha$-helical or $\beta$-sheet secondary structure; $\cap$ denotes loop structure; SA refers to solvent accessibility of the side chain or main chain, as approximated from the X-ray crystal structure; HydroP refers to hydrophobicity of each residue; $^{44} k_{1/2}$ is the previously measured rate of amide hydrogen deuterium exchange (s$^{-1}$).$^{45}$ The changes in chemical shift of each residue in the HSQC spectra at 0 and 1 mM SDS are derived from data in Figures 4 and 5 and are denoted $\Delta \delta$. (Table 1). We infer from the HSQC analysis that salt-bridges are not forming between the sulfate group of SDS and positively charged groups of lysine and arginine (although salt bridges between some lysine or arginine residues might be forming). This absence of a general interaction between SDS and Lys or Arg is not so surprising when the electrostatic surface potential of UBI (Figure 6) is taken into account: lysine and arginine residues in UBI do not always have the largest positive electrostatic surface potentials (e.g., Arg54 $\approx$ 4 eV; Ile13 $\approx$ 4 eV) because, presumably, solvent is screening the charge of these residues. Therefore, the location of a salt-bridge—if one were to form between UBI and SDS—would be with a residue bearing a poorly solvated charge.

The hydrophobicity and surface potential (not formal charge) of an amino acid residue in folded UBI appear to be more important than secondary structure and solvent accessibility in explaining its association with SDS below the CMC. The strength of association of residues in UBI and SDS at sub-denaturing concentrations (<1.0 mM) seems to be equally governed by the residue’s hydrophobicity and its local electrostatic environment, but neither property alone is sufficient to explain the binding of SDS to folded UBI. A third property (the $2^\alpha$ structure of the residue) appears to play a lesser role (Table 1): residues in hydrogen-bonded loops appear to be protected from interacting with SDS; however, SDS did not prefer residues in $\alpha$-helical structure over those in $\beta$-sheet structure. We infer that—at least for UBI—the presence of (formally) cationic functional groups is important for the binding of SDS because cationic sites provide positive electrostatic surface potential that extends beyond the van der Waals radius of the functional group (toward formally neutral hydrophobic regions), and not necessarily because the cationic site forms an ionic bond (“saltbridge”) with the sulfate group (although these interactions might occur for some cationic residues, such as Lys11, but not for others, such as Lys6 and Arg54). In summary: (i) surfactants bind preferentially to hydrophobic regions with positive
electrostatic surface potential (this location may or may not be at cationic residues, depending upon solvation), and (ii) salt bridges do not necessarily form between the sulfate group and cationic groups in the protein. We conclude that the acetylation of lysine residues in UBI inhibits the binding of the initial ∼11 molecules of SDS, below the CMC of SDS, primarily because acetylation increases the negative surface potential of UBI, and not because acetylation eliminates amino functionalities that might have been sites of binding for the sulfate group of SDS. In general, the two amino residues in UBI that interacted most favorably with SDS, below the CMC, were the branched hydrophobic residues leucine and isoleucine, so long as they were adjacent to areas of positive electrostatic surface potential and were not part of a hydrogen-bonded loop. Specific interactions of SDS with aliphatic protons of side chains resulted in shifting of several $^{13}$C−$^{1}H$ resonances, but the majority of the side chains and all of the Hα’s (with the exception of the flexible N-terminus) were attenuated by dynamic broadening of resonances resulting from rapid kinetics of binding.

Previous work has suggested that $β$-stranded proteins are, in general, more kinetically resistant to unfolding by SDS than are $α$–$β$ proteins, but we did not find with UBI that similar amino acids residues bind SDS differently in $β$-strands and $α$-helices. Moreover, the solvent accessibility of the side chain of a residue, and the conformational flexibility of the residue, do not seem to correlate with its association with SDS at 1 mM SDS, although Table 1 suggests that peptide backbones that are solvent exposed interact with SDS slightly more than backbones that are buried (Table 1).

Our results also reveal a few points regarding the unfolding of UBI in SDS. Importantly, UBI-(NH$_3^+$)$_8$ binds 11 more molecules of SDS than UBI-(NHAc)$_8$ at concentrations of SDS that are below and above the CMC; this observation suggests that the 11 SDS molecules bound to UBI-(NH$_3^+$)$_8$ in the folded state remain bound in the unfolded state. Nevertheless, the pathway of association of SDS with UBI in the unfolding regime is not influenced by the initial number of cationic residues: according to CD spectroscopy both UBI-(NH$_3^+$)$_8$ and UBI-(NHAc)$_8$ unfold via the binding of indistinguishable numbers of SDS molecules (i.e., 14 SDS; see the CD spectra in Figure S5).

We have previously used capillary electrophoresis to study the unfolding of several proteins in SDS, but UBI (e.g., UBI-(NH$_3^+$)$_8$) is unique among the 18 that we studied in that UBI binds multiple equivalents of SDS below the CMC without unfolding. Most proteins that we have studied do not associate with large numbers of SDS below the CMC; most proteins do not bind SDS without unfolding, and most do not follow a complex pathway of unfolding that involves multiple stable, distinct complexes with defined stoichiometry. We still do not understand why UBI forms a stable intermediate with SDS below the CMC, without unfolding, and why UBI forms more distinct complexes with SDS and populates more intermediates during unfolding than other proteins that we have studied. We point out that UBI is one of the least negatively charged proteins that we have studied ($Z = -0.2$ at pH 8.4), and our previous investigations into the unfolding of proteins in SDS with CE are biased against proteins that have a net positive charge (studying SDS binding to positively charged proteins with a bare silica capillary is difficult because of absorption onto the negatively charged capillary). This study has shown, however, that the peracylation of UBI causes the protein to unfold in a manner that is similar to most of the other proteins that we have studied: peracylation eliminates the ability of the protein to coordinate SDS without unfolding and causes UBI to unfold in SDS via a two-state transition.

In studying of a subject—the association of proteins with surfactants in particular, and molecular recognition in general—that is important, but very complicated, this work represents a step toward a more detailed understanding. The combination of capillary electrophoresis, HSQC NMR, and protein charge ladders is certainly a useful, new technique to investigate quickly (and reproducibly) various hypotheses concerning important interactions in these processes.

### Materials and Methods

#### Chemicals.

Ubiquitin from bovine erythrocytes (all experiments with the same lot no. 075K740S, Tris-NH$_3$ glycine, sodium dodecyl sulfate, acetic anhydride, dioxane, N,N-dimethylformamide, and N-(2-hydroxyethyl)piperazine-N’-(4-butanesulfonic acid) (HEPES) were purchased from Sigma-Aldrich. Dialysis microtubes (Slide-A-Lyzer MINI, MWCO 3.5k) were purchased from Pierce and rinsed several times with ultrapure water prior to use. SDS was recrystallized three times from hot ethanol, and the absence of any alcohol precursor (dodecanol) was checked by NMR. Tris-glycine buffer (25 mM Tris, 192 mM glycine) was prepared from salts purchased from Aldrich and filtered through a 0.22-μm membrane prior to use. Preparation of UBI in solutions of SDS by dialysis was carried out as described in a previous study.

#### Synthesis of Peracylated UBI.

The acylation of UBI (20 μM) with acetic anhydride was carried out in 100 mM HEPES at pH 9.0. Neat acetic anhydride was diluted 1:10 into dioxane, 100 stoichiometric equivalents (with respect to lysine and N-terminus) were added to buffered solutions of UBI, and the reaction was allowed to proceed overnight at room temperature. The concentration of dioxane in the aqueous solution of UBI did not exceed 5%. Dioxane was eliminated by extensive dialysis against Tris-glycine. The peracetylated protein was then concentrated using Centricron centrifugal filtration devices (Millipore, 5 kDa MWCO) to a concentration of 50 μM.

#### Capillary Electrophoresis (CE).

We used a Beckman Coulter P/ACE electrophoresis apparatus, with a capillary length of 60.2 cm (50 cm to the detector), and composed of fused silica (inner diameter of 50 μm; Polymicro Technologies TSP050375, Part No. 2000017, lot LHQ02A). Buffers were composed of Tris-glycine (pH 8.4) containing concentrations of SDS from 0 to 10 mM. CE was performed at an applied voltage of 30 kV for 10 min. The capillary was equilibrated before each electrophoresis by a sequential rinsing with (i) methanol, (ii) HCl (1 M), (iii) NaOH (0.1 M), (iv) ultrapure water (Millipore Advantage Ultrapure Water System, conductivity of 18.2 MΩ·cm and total organic content not exceeding 4 ppb), and (v) the running buffer for the electrophoresis (3 min at 20 psi prior to each injection of protein). The solutions containing UBI-SDS complexes (∼50 μM; we used an extinction coefficient for UBI of ε(280) = 1490 L/(mol·cm)) were injected in the capillary at 0.5 psi for 20 s. Samples were injected at a temperature of 4 °C, and the temperature of the capillary was maintained at 25 °C during electrophoresis. Each series of experiments was carried out with the same capillary, equilibrated after each set of 20 runs.

The mobility of a UBI-SDS complex was calculated according to eq 4, where $\mu$ is the mobility of the migrating species (in cm$^2$·kV$^{-1}$·min$^{-1}$), $L_T$ the total length of the capillary, $L_p$ the length up to the detector, $V$ the applied voltage, $t_{no}$ the time of the neutral marker, and $t$ the time of the migrating species:

$$\mu = \frac{L_P \cdot L_T}{V \left( \frac{1}{t_{no}} - \frac{1}{t} \right)} \tag{4}$$
Circular Dichroism (CD) Spectroscopy. CD spectra were collected on a spectropolarimeter (Jasco Inc.) equipped with a quartz cuvette with a 1-mm path length. For far-UV measurements (205–260 nm), the concentration of proteins was 50 μM at (25 °C). Each sample was scanned 10 times at 20 nm/min with a step size of 0.5 nm. The molar ellipticity \( \theta (\text{deg} \ M^{-1} \ cm^{-1}) \) was obtained by using eq. (5) with \( L_1 \) and \( L_2 \) (the path length of the cuvette used for the measurement of the absorbance at 220 nm is \( L_1 = 1 \ cm \); for the cuvette used to collect CD spectra, \( L_2 = 0.1 \ cm \)).

\[
\theta (\text{deg} \ M^{-1} \ cm^{-1}) = \frac{\theta_{\text{meas}} (\text{deg} \ M^{-1} \ cm^{-1})}{A_{220} L_1} \]

15N−1H and 13C−1H HSQC NMR. The NMR experiments were carried out on a Bruker 600 MHz spectrometer equipped with a cryo-probe. All experiments were conducted using 25 mM (DOCD)2CND2 (or D(11)-Tris) in 1:9 H2O:D2O at pH 7.53, with the samples at room temperature (20 °C). The deuterated form of Tris buffer was obtained from Cambridge Isotope Labs (Cambridge, MA), as well as 15N- and 13C-1H-labeled human UBI (76 amino acids), 15N−1H HSQC spectra of uniformly 15N-labeled UBI (100 μM) were collected for 45 min, while those in the presence of 1 mM SDS were collected over a period of 12 h and then averaged. 13C HSQC spectra of uniformly 13C- and 15N-labeled UBI (150 μM) in the absence of SDS were collected for 2 h, and those in 1 mM SDS were collected over a period of 4.5 h, maintaining the samples at room temperature. The 15N−1H HSQC experiments were performed using solutions with 150 μM UBI-(NH3+)8. We chose this concentration of UBI (low for a typical protein15N−1H HSQC experiment) so that the UBI:SDS ratio would be as comparable with CE experiments as possible (CE experiments were carried out at 50 μM UBI-(NH3+)8). We assigned residues to UBI at pH 7.53 and processed images using Cara (v 1.8.4). We used Sparky software to analyze the chemical shifts (\( \delta \)).

**ASSOCIATED CONTENT**

Supporting Information. Circular dichroism spectra of UBI-(NH3+)8 and UBI-(NHAc)8, high-resolution CE electrophorograms of the transition N→G1→G2, and electrophorograms of BCA dialyzed against SDS. This material is available free of charge via the Internet at http://pubs.acs.org.

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**ACKNOWLEDGMENT**

This research was funded by NIH award GM 051559. B.F.S. was supported by a NIH Ruth L. Kirchstein NRSA postdoctoral fellowship (GM081055), and G.F.S. was supported by a “Lavoisier Générale” postdoctoral fellowship (Ministère des Affaires Etrangères Français). Molecular graphics images of UBI were produced using the UCSF Chimera package from the Resource for Biocomputing, Visualization, and Informatics at the University of California, San Francisco (supported by NIH P41 RR-01081). The authors thank Elizabeth V. Shaw for taking the photographs used to create Table of Contents and Abstract graphics.

**REFERENCES**

6. This point is supported by the absence of clefs or pockets on the surface of folded UBI, and the observation that no pockets appear to form in UBI upon its association with <1.4 mM SDS (i.e., previous work has shown that the secondary and tertiary structure of UBI does not change upon the binding of the first 11 SDS).
33. The Hansch hydrophobic parameter for RSO4− is −0.476.
34. This rationalization assumes that the 11 RSDS molecules bound to UBI-(NH3+)8 in its G2 state (e.g., UBI-(NH3+)8(SDS)11) render the surface of the protein more hydrophobic than the NHCOCH3 groups introduced onto the surface of UBI-(NHAc)8 by peracetylation (the binding of SDS also renders the surface more zwitterionic). The increase in surface hydrophobicity due to the acetylation of lysines in UBI-(NHAc)8—that is, the addition of eight -COCH3 groups—is certainly small compared to the increase in surface hydrophobicity reflected by the association of ~130 methylene units resulting from the binding of 11 molecules of SDS.
(38) The positively charged side of UBI is composed of an entire face of a \(\beta\)-sheet and is hydrophobic. The negatively charged side contains a loop structure and also contains the single \(\alpha\)-helix; this \(\alpha\)-helix contains three cationic residues (all lysine) but also contains three anionic residues and is close to being overall electrically neutral.