Structural plasticity of staphylococcal nuclease probed by perturbation with pressure and pH

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ABSTRACT

Despite significant progress in our understanding of protein folding and stability, important questions concerning the structural basis of folding cooperativity and the structural, thermodynamic, and dynamic origins of barriers between conformational states remain unanswered. This has hindered our understanding of the structural basis of important biochemical processes such as catalysis and energy transduction, which are governed partly by the response of proteins to changes in the charged state of H+ or e− binding centers. To examine this problem in detail, we applied NMR spectroscopy and SAXS at ambient and high-hydrostatic pressure to monitor the consequences of the ionization of Lys-66 in the V66K variant of staphylococcal nuclease (SNase).

The ionization of internal groups in proteins can trigger conformational change. Despite this being the structural basis of most biological energy transduction, these processes are poorly understood. Small angle X-ray scattering (SAXS) and nuclear magnetic resonance (NMR) spectroscopy experiments at ambient and high hydrostatic pressure were used to examine how the presence and ionization of Lys-66, buried in the hydrophobic core of a stabilized variant of staphylococcal nuclease, affect conformation and dynamics. NMR spectroscopy at atmospheric pressure showed previously that the neutral Lys-66 affects slow conformational fluctuations globally, whereas the effects of the charged form are localized to the region immediately surrounding position 66. Ab initio models from SAXS data suggest that when Lys-66 is charged the protein expands, which is consistent with results from NMR spectroscopy. The application of moderate pressure (<2 kbar) at pH values where Lys-66 is normally neutral at ambient pressure left most of the structure unperturbed but produced significant nonlinear changes in chemical shifts in the helix where Lys-66 is located. Above 2 kbar pressure at these pH values the protein with Lys-66 unfolded cooperatively adopting a relatively compact, albeit random structure according to Kratky analysis of the SAXS data. In contrast, at low pH and high pressure the unfolded state of the variant with Lys-66 is more expanded than that of the reference protein. The combined global and local view of the structural reorganization triggered by ionization of the internal Lys-66 reveals more detectable changes than were previously suggested by NMR spectroscopy at ambient pressure.


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Key words: protein electrostatics; conformational dynamics; pressure; NMR; SAXS.

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SNase is a rich model system for detailed examination of structural plasticity and structure-energy correlations. The thermodynamic stability of SNase can be modulated with mutagenesis and increased from 5.4 kcal mol\(^{-1}\) for the wild-type protein\(^1\) to nearly 12 kcal mol\(^{-1}\) for two highly stable variants known as NVIAGA\(^2\) and Δ+PHS\(^3\). The Δ+PHS protein, engineered by a deletion (44 to 49) and five substitutions (P117G, H124L and S128A, G50F and V15N), is of special interest because it has been shown to tolerate the presence and ionization of groups buried in its hydrophobic core.\(^3–10\) The majority of variants of Δ+PHS with internal ionizable groups retain a folded structure and cooperative unfolding profiles\(^4,8\) even when the internal groups are charged. The present study is focused on the V66K variant of Δ+PHS. Lys-66 titrates with a \(pK_a\) of 5.7, which is highly depressed relative to the normal \(pK_a\) of 10.4 for Lys in water.\(^3,6\) This depressed \(pK_a\) suggests that the interior of SNase is not as good a solvent as water; hence, the charged form of the amino group is destabilized relative to its neutral form.\(^6\)

Besides destabilizing the native state of a protein, the presence and ionization of a group buried in the hydrophobic core can trigger changes in conformation and dynamics, and could stabilize partially unfolded excited states in which the internal charge group can interact with water. This is precisely why internal ionizable groups can be useful for mapping the folding free energy landscape of proteins.\(^5,11\) The crystal structures of V66K variants of SNase are not particularly informative; they are indistinguishable from those of the reference Δ+PHS protein.\(^1,3,9\) In the structures obtained under conditions of pH where Lys-66 is neutral, the side chain of Lys-66 is buried deeply in the hydrophobic core of the protein, nearly 12 Å from the protein–water interface. In the structure obtained under conditions of pH where Lys-66 is charged, the backbone of the protein is unaffected but the side chain is disordered.\(^9\) Structure-based \(pK_a\) calculations with a variety of electrostatics models show that the behavior of Lys-66 is consistent with high-apparent polarizability in the hydrophobic interior of the protein;\(^6\) however, the crystal structures provide no clues into the origins of this high-apparent polarizability. Experiments and calculations suggest that it is related to conformational changes coupled to the ionization of the internal Lys-66.\(^5,8,9\) These are precisely the type of relaxation processes we wish to understand.

Because the \(pK_a\) of Lys-66 is very depressed, it is possible to use pH perturbation to exert electrostatic and conformational stress on the V66K protein. NMR spectroscopy at atmospheric pressure was used previously to show that at pH 7, where Lys-66 is buried and neutral,\(^5\) the V66K substitution affects the hydrogen exchange behavior throughout the protein, probably because the global thermodynamic stability of the protein is greatly diminished even when the internal Lys-66 is neutral. When Lys-66 is charged only the region immediately adjacent to the ionizable side chain is affected.\(^5\) Specifically, in \(^1\)H HSQC spectra, the resonances of backbone amides of residues 62–70 are broadened when these internal ionizable groups are charged; the rest of the protein appears to remain unaffected. This is consistent with an increase in local fluctuations concomitant with the ionization of Lys-66, but the exact nature of the fluctuations remains to be established. It is interesting that although burial and ionization of Lys-66 diminishes the thermodynamic stability of the protein significantly, the structural and dynamic consequences appear to be relatively minor.\(^8,10\)

In an attempt to improve our understanding of the effects of the ionization of Lys-66 on structure and dynamics over a wide range of timescales, we have monitored the effects of changes in pH concomitantly with nuclear magnetic resonance spectroscopy (NMR) and small angle X-ray scattering (SAXS). Because the pH sensitivity of this protein originates primarily from differences in hydration of the Lys residue when it is in water and when it is buried in a hydrophobic pocket, high-pressure was also used in attempts to perturb the native and unfolded state manifolds. Our interest in using high-hydrostatic pressure to examine the effects of internal polar and ionizable groups on conformational stability and dynamics stems from the premise of pressure as a useful tool for studying equilibrium processes governed by changes in water-inaccessible volume.\(^12,13\) The utility of pressure as a useful thermodynamic variable to examine processes affected by electrostatic hydration was already revealed by pressure-jump relaxation studies with variants of SNase with internal ionizable groups.\(^14,15\) Those studies showed that the hydration of the transition state ensemble can be affected dramatically by substitution with ionizable groups at positions that are normally part of the main hydrophobic core in the transition state ensemble of the wild-type protein. Although the effects of substitutions of hydrophobic positions with ionizable groups are subtle, the effects on the free-energy landscape can be very significant.\(^14,15\)

Application of hydrostatic pressure constitutes a unique and gentle thermodynamic perturbation to proteins. High-pressure selectively stabilizes conformations with smaller specific volumes. The working hypothesis,\(^16\) based on earlier studies on SNase\(^14–19\) and the c-Myb R2 domain\(^20\) is that the differences in volume between the various conformational states of proteins arise primarily from differences in their solvent excluded void volumes. Indeed, although the interior of proteins is generally quite well packed, packing is not perfect; hence there is unoccupied space inside proteins, some of which can be filled by water in locally unfolded states or on complete unfolding.\(^12,13\) The molar volume decreases as cavities become hydrated; therefore, the application of pressure is expected to stabilize progressively the lower volume conformers with higher degree of hydration in cavities. In principle, subtle conformational fluctuations that are unnoticed at ambient pressure can be enhanced with high pressure to levels.
detectable by NMR spectroscopy and SAXS. For these reasons, use of SAXS and NMR spectroscopy to monitor consequences of pressure perturbation coupled with pH perturbation (to control the charged state of the internal Lys) have promise to be useful to study subtle conformational reorganization coupled to the ionization of internal groups in proteins.

RESULTS

Effects of Lys-66 on conformational dynamics at ambient pressure

\(^{1}\)H-\(^{15}\)N HSQC spectra of the \(\Delta +\)PHS/V66K showed that when Lys-66 is charged, the resonances for backbone amides of residues surrounding Lys-66 are broadened beyond detection. A closer examination of the pH titration of the \(^{1}\)H-\(^{15}\)N HSQC spectrum of the \(\Delta +\)PHS/V66K protein (Fig. 1) revealed a strong pH-dependent decrease in the peak intensity, while the peak volumes remained constant, indicating conformational exchange on rather long timescales (>ms), coupled to the ionization of Lys-66. Certain residues [in blue in Fig 1(B,C)] exhibited a plateau across the \(pK_a\) of Lys-66, while others [in green in Fig 1(B,C)], particularly in the \(\beta\)-barrel near Lys-66, showed a minimum at the \(pK_a\). This latter behavior can be interpreted as arising either from a change in dynamics, or more likely, a decrease in the conformational heterogeneity for these residues once Lys-66 is completely protonated.

NMR relaxation techniques were used to compare the dynamics of the \(\Delta +\)PHS protein and its V66K variant at pH values above (6.5) and below (4.5) the \(pK_a\) of Lys-66. The rapid, picosecond dynamics of both proteins were rather constrained, with order parameters above 0.8 across the entire structure at both pH values (Supporting Information Fig. S1). The relaxation parameters (\(R_1\), \(R_2\), and heteronuclear NOE) exhibited no significant differ-
ences, demonstrating that insertion of a polar and ionizable residue into the hydrophobic core had very little effect on fast dynamics in the ps–ns range. CPMG relaxation dispersion measurements performed on the two proteins at pH 6.5 and 4.5 revealed no evidence for conformational exchange on the l to ms timescale at either pH value for the D1PHS protein, consistent with its very high stability. The same was true for the D1PHS/V66K variant at pH 6. However, when Lys-66 was positively charged at pH 4.5, conformational exchange was observed at 13 positions in the D1PHS/V66K variant [e.g., Fig. 2(A)], with exchange times in the range of 300 μs, and with four residues apparently exchanging on the millisecond timescale (Supporting Information Table I).

Earlier studies showed that the V66K substitution affected the H/D exchange properties of the protein globally, even when Lys-66 is neutral.5 The ratio of the protection factors measured here between the two proteins at pH 6.5 and 4.5 confirmed the global destabilization of the V66K variant relative to the Δ+PHS background, with decreases in the protection factor of up to four orders of magnitude at both pH values for residues distributed throughout the protein [Figs. 3(A,B) and Supporting Information Figs. S2(A–D) and S3(A–B)]. The residues exhibiting the largest decrease in protection factor reside in helix-1, where Lys-66 is found, and in the β-barrel. Overall, the effects of the V66K substitution on the conformational dynamics of the protein probed by H/D exchange indicate that, while both the global fluctuation and the local fluctuation are increased by the substitution, the global fluctuations are enhanced by protonation of Lys-66.

Small-angle X-ray scattering (SAXS)27,28 experiments at atmospheric pressure were performed with Δ+PHS and its V66K variant to probe the global structural consequences of the V66K substitution and of the ionization of Lys-66, whereas at pH above 5.7, the Δ+PHS protein and its V66K variant exhibited similar Rg values (16.9 ± 0.4 and 16.6 ± 0.8 Å) and pair distribution functions, at pH 4.5, where Lys-66 is charged, the V66K variant exhibited a significantly broader pair-distance distribution function and a larger Rg value (18.1 ± 0.5 Å) (Supporting Information Fig. S4(A–C), Fig. 4(A,B), Table I). The Δ+PHS background also showed some expansion at pH 4.5 (17.6 ± 0.3 Å). The Kratky plots [Fig 5(A,B)] regardless of variant or pH were all very similar, although not identical, with a strong maximum consistent with structured, globular proteins.

Ab initio modeling of the SAXS data for Δ+PHS and its V66K variant at atmospheric pressure and pH 6 was

Table I

<table>
<thead>
<tr>
<th>Variant</th>
<th>Experimental conditions</th>
<th>Rg/Å</th>
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<tbody>
<tr>
<td>Δ+PHS</td>
<td>1 bar, pH 7.0</td>
<td>16.9 ± 0.4</td>
</tr>
<tr>
<td>Δ+PHS</td>
<td>1 bar, pH 4.5</td>
<td>17.6 ± 0.3</td>
</tr>
<tr>
<td>Δ+PHS + V66K</td>
<td>1 bar, pH 6.0</td>
<td>16.8 ± 0.8</td>
</tr>
<tr>
<td>Δ+PHS + V66K</td>
<td>1 bar, pH 4.5</td>
<td>18.1 ± 0.5</td>
</tr>
<tr>
<td>Δ+PHS + V66K</td>
<td>3 kbar, pH 6.0</td>
<td>19.8 ± 2.0</td>
</tr>
<tr>
<td>Δ+PHS + V66K</td>
<td>1 kbar, pH 4.5</td>
<td>22.8 ± 0.7</td>
</tr>
<tr>
<td>Δ+PHS + V66K</td>
<td>2 kbar, pH 4.5</td>
<td>32.5 ± 2.0</td>
</tr>
<tr>
<td>Δ+PHS + V66K</td>
<td>3 kbar, pH 4.5</td>
<td>30.2 ± 2.0</td>
</tr>
</tbody>
</table>
performed with the program DAMMIF.\textsuperscript{29} The most probable model from among 16 separate calculations for the Δ+PHS protein revealed a generally globular structure, with a circular indentation [Fig. 6(A,B)]. Superposition of the crystal coordinates of the Δ+PHS structure (PDB accession code 3bdc.pdb) with the residues that exhibit faster exchange rates, as compared with the Δ+PHS background colored in pink to red from 10-fold to 10,000-fold faster. Blue residues are exchanged 10-fold to 100-fold more slowly. NMR acquisitions were carried out at 20°C. Under these conditions, the exchange occurs in the EX2 limit.

**Figure 3**

H/D exchange measurements of the Δ+PHS protein and its V66K variant. (A) ratio of the exchange rates of the amide protons of Δ+PHS SNase with respect to those measured for the Δ+PHS/V66K variant at pH 6.5; (B) ribbon diagram of Δ+PHS/V66K with the residues that exhibit faster exchange rates, as compared with the Δ+PHS background colored in pink to red from 10-fold to 10,000-fold faster. Blue residues are exchanged 10-fold to 100-fold more slowly. NMR acquisitions were carried out at 20°C. Under these conditions, the exchange occurs in the EX2 limit.

**Figure 4**

Normalized pair-distance distribution functions calculated from the SAXS data on Δ+PHS and its V66K variant. (A) Δ+PHS at pH 7 and atmospheric pressure ( ), 1 kbar ( ), 2 kbar ( ) and 3 kbar ( ); (B) Δ+PHS/V66K at pH 4.5 ( ) and pH 6.0 ( ) at atmospheric pressure; (C) Δ+PHS/V66K at pH 6.0 and atmospheric pressure ( ) and 3 kbar ( ).

experimental scattering from the disordered C-terminal residues 141–149, although a contribution from the disordered N-terminal residues 1–7 cannot be excluded. We...
note that such *ab initio* modeling, while appropriate for stable globular structures, does not yield reliable envelopes for dynamically disordered regions, and that, therefore, this protrusion in the model should not be interpreted as a true structural envelope.

Despite the similarity in their $R_g$ values, the *ab initio* model for the $\Delta+$PHS/V66K variant [Fig. 6(D,E)] appears quite different from that of the $\Delta+$PHS reference protein. In particular, the DAMMIF modeling of the $\Delta+$PHS/V66K variant produced two extensions that, as in the case of the single protrusion calculated for the reference protein, may also arise from disordered regions, which are more important in the $\Delta+$PHS/V66K protein than in the reference protein. At pH 6, close to the $pK_a$ of 5.7 of Lys-66, the SAXS data might be reporting on the superposition of multiple conformations. A slow dynamic disordering of helix 1 is supported by the H/D exchange data, which revealed substantial increases in the helix 1 exchange rates for $\Delta+$PHS/V66K compared with the $\Delta+$PHS reference protein. The *ab initio* model of the V66K variant at pH 4.5 shows a very elongated structure [Fig. 6(F)], suggesting that under conditions where Lys-66 is charged, the protein populates a state in which part or all of helix-1 is separated from the core of the structure. This is consistent with the NMR results that show even faster H/D exchange, the loss of the resonances for residues 62–66 and the broadening of resonances in the

**Figure 5**
Kratky plots of (A) $\Delta+$PHS/V66K SNase at: (□) pH 6 and atmospheric pressure, (○) pH 6 and 3 kbar, (△) pH 4.5 and atmospheric pressure, (+) pH 4.5 and 1 kbar, (×) pH 4.5 and 2 kbar, (•) pH 4.5 and 3 kbar; (B) $\Delta+$PHS SNase at: (□) pH 7 and atmospheric pressure, (○) pH 7 and 1 kbar, (△) pH 7 and 2 kbar, (×) pH 7 and 3 kbar.

**Figure 6**
Results of *ab initio* modeling of the $p(r)$ plots for $\Delta+$PHS SNase and its V66K variants under a variety of conditions. (A) and (B) *ab initio* models of $\Delta+$PHS at pH 7 and atmospheric pressure in two 90° orientations, (C) the crystal structure of $\Delta+$PHS modeled into the *ab initio* sphere model, with $\alpha$-helices colored in red, $\beta$-strands in yellow, loops and turns in orange, position 66 in green, and the C-terminal 3-10 helix in blue, (D) and (E) *ab initio* models of $\Delta+$PHS/V66K at atmospheric pressure and pH 6 in two 90° orientations, (F) *ab initio* model of $\Delta+$PHS/V66K at atmospheric pressure and pH 4.5, (G) *ab initio* model of $\Delta+$PHS at pH 7 and 3 kbar, (H) *ab initio* model of $\Delta+$PHS/V66K at pH 6 and 3 kbar, (I) *ab initio* model of $\Delta+$PHS/V66K at pH 4.5 and 1 kbar.
Effects of the neutral Lys-66 on conformational dynamics monitored with variable pressure NMR spectroscopy

Variable pressure NMR spectroscopy was used as an additional means of assessing the effect of the V66K substitution on the malleability of the protein at pH 6.3. Under these conditions, Lys-66 is neutral and no significant differences in the protein's ps–ms dynamics were observed at atmospheric pressure as compared with the Δ+PHS reference protein. Pressure induced significant shifts in both the $^1$H and $^{15}$N frequencies in the $^1$H–$^{15}$N HSQC spectrum for Δ+PHS/V66K at pH 6.3 [Figs. 7(A,B)] at all sites, reflecting average changes in hydrogen bond distances and in $\phi$, $\psi$, $\chi$ torsion angles.$^{21-23}$ The pressure-dependent $^1$H–$^{15}$N HSQC spectra of the protein when Lys-66 is charged.

Figure 7
Pressure-dependent $^1$H–$^{15}$N HSQC spectra of Δ+PHS/V66K SNase at pH 6.3. NMR acquisition was carried out at 21 °C. (A) Complete HSQC spectra between 30 bar and 3 kbar as indicated. (B) Synthetic HSQC revealing the amide chemical shift variations of the folded state with pressure. Peaks shift black to yellow to purple to brown with increasing pressure between atmospheric pressure and 3 kbar.
shifts in the HSQC resonances of the reference protein were generally smaller than those observed for the V66K variant (spectra not shown). Variations in the shifts over the entire chain reveal heterogeneous conformational fluctuation. Continuous shifts without loss of signal intensity or broadening of signals indicate that these fluctuations are rapid on the NMR time scale (fluctuations faster than ~ms). Although the majority of the shifts are linear, some are quite significantly nonlinear. The linear shifts are thought to represent the pressure-induced conformational fluctuation within the folded conformation, whereas the nonlinear shifts are attributed primarily to the increase in the population of alternative or low-lying excited state conformations.21–23 Fitting both $^1$H and $^{15}$N shifts for the $\Delta$+PHS and $\Delta$+PHS/V66K proteins to a second-order polynomial [Supporting Information Fig. S6(A–D)] revealed that in both the proton and nitrogen dimensions, the nonlinear coefficients for the $\Delta$+PHS background are quite small, on the order of the average observed for BPTI, one of the least pressure sensitive proteins tested to date.21,23,30 The response of the $\Delta$+PHS/V66K variant to pressure was similar to that of the reference protein over most of its structure. However, significant nonlinear pressure responses were observed in the $\Delta$+PHS/V66K variant clustered near Lys-66 in helix-1 and in the $\beta$-barrel (Fig. 8 B and D). These results suggest that the V66K substitution increased significantly the pressure-induced conformational fluctuations (<ms) into low-lying excited states and that these fluctuations involve a change in conformation in helix-1 and the $\beta$-barrel.

Global unfolding of $\Delta$+PHS/V66K monitored by NMR spectroscopy and SAXS at high pressure

Besides the nonlinear effect of pressure on some chemical shifts, at pH 6.3 the $\Delta$+PHS/V66K variant underwent major spectral changes in the $^1$H-$^{15}$N HSQC as pressure increased to 3 kbar [Fig 7(A)]. The well-dispersed resonances of the folded state reversibly decreased in intensity while those less resolved central peaks corresponding to the unfolded protein increased in intensity, clearly showing that the entire polypeptide chain became unfolded, in complete agreement with fluorescence experiments under similar conditions.14 The observation of separate signals for the folded and unfolded conformers reveals that global unfolding is slow on the NMR time scale (>ms). It was possible to follow the loss of peak intensity over the entire pressure range for 47 of the 138 residues of the protein. Plots of peak intensity versus pressure for these residues were analyzed (Supporting Information Fig. S7) to obtain residue specific values of the apparent unfolding free energy ($\Delta G_u$) and volume change on unfolding ($\Delta V_u$). The average $\Delta V_u$ calculated from the values of all the residues was $-67.7 \pm 16$ mL mol$^{-1}$, and the average $\Delta G_u$ was $3.4 \pm 0.8$ kcal mol$^{-1}$, in agreement the values of $\Delta V_u$ ($-60 \pm 6$ mL mol$^{-1}$) and $\Delta G_u$ (3.0 ± 0.3 kcal mol$^{-1}$) obtained previously for this variant at pH 6.0 from analysis of pressure-induced fluorescence changes.14

The effect of pressure on the global structure of the reference $\Delta$+PHS protein and its V66K variant was also

![Figure 8](image-url)
characterized by SAXS measurements at high pressure [Supporting Information Fig. S4(B)]. Ab initio modeling of the \( p(r) \) functions of the reference protein [Fig. 4(A)] revealed a very limited change in shape at 3 kbar, which modeled only as an attenuation of the circular indention [Fig. 6(G)] in the globular structure. In contrast, the major change in conformation of the \( \Delta + \)PHS/V66K variant, observed by high-pressure NMR and fluorescence, was confirmed by SAXS. The \( p(r) \) for this variant at pH 6 [Fig. 4(C)] broadened and the \( R_g \) increased from 16.6 ± 0.4 to 19.8 ± 2.0 Å (Table I), indicating expansion of the chain relative to the native state. However, this value for the \( R_g \) of the pressure-unfolded \( \Delta + \)PHS/V66K variant is much smaller than the value of 35 Å obtained previously for the high-pressure unfolding of the true wild-type form of SNase\(^{17}\) under similar conditions (pH 5.5). Hence, even with the V66K substitution, the pressure unfolded state of the \( \Delta + \)PHS variant remained much less expanded than that of the wild-type protein. The Kratky plot obtained from the scattering data on the V66K variant at 3 kbar and at pH 6 [Fig. 5(A)] exhibited a small maximum reflecting some residual globular structure in this unfolded state. Ab initio modeling of the scattering curves under these conditions [Fig. 6(H)] was consistent with a significantly expanded structure relative to atmospheric pressure. Furthermore, the SAXS data show that ionization of Lys-66 (at pH 4.5) shifted the unfolding transition to lower pressure which was complete by 2 kbar, as observed previously by fluorescence.\(^{14}\)

The radius of gyration at pH 4.5 and 2 kbar (≈32 Å, Table I), was close to the value of 35 Å\(^{17}\) observed for pressure denatured WT SNase, yet it was not as large as for chemically denatured wild-type SNase (\( R_g = 37.2 \pm 1.2 \) Å)\(^{31}\) or for the temperature denatured WT protein (\( R_g = 45 \pm 2 \) Å).\(^{17}\) Kratky plots [Fig. 5(A)] of the high-pressure SAXS profiles (2 and 3 kbar) at pH 4.5 showed no maximum, indicating that under these conditions the protein behaved like a random polymer. Likewise, the ab initio model [Fig. 6(I)] was consistent with a highly expanded chain. Because Lys-66 in the \( \Delta + \)PHS/V66K variant is charged in the unfolded state at pH 6 as well as at pH 4.5, the observed expansion of the pressure unfolded state at low pH must arise from general destabilization of residual interactions arising from increased electrostatic repulsion in the unfolded state as carboxylic groups begin to become neutralized.

**DISCUSSION AND CONCLUSIONS**

Previous studies of the \( \Delta + \)PHS protein and its V66K variant with NMR spectroscopy have shown that the response of the protein to the ionization of Lys-66 is limited and localized to the vicinity of the side chain of the lysine.\(^5\) Because structural and dynamic consequences of the presence and ionization of the internal Lys-66 are affected by the volume differences between Val and Lys, and by the differences in hydration of Lys in water and Lys inside a protein, the application of high-hydrostatic pressure was expected to shed insight into the structural determinants of the properties of the internal Lys-66 and the response of the protein to the presence of a charged group in the mid of its hydrophobic core.

The present studies at atmospheric pressure show that at pH values close to the \( pK_a \) value of Lys-66, the protein exists in dynamic equilibrium on relatively slow timescales between two or more conformations. The maximal loss of intensity at the \( pK_a \) of Lys-66 suggests that a few residues in the barrel are exchanged broadened on a rather slow timescale (\( > \)ms) owing to a mixed population of protonated and unprotonated Lys-66. A dynamic equilibrium of the side chain between a buried position, where it is neutral and one less buried, where it is charged and more hydrated, is compatible with this slow timescale, but so is an equilibrium between ordered and disordered backbone for the segment 62 to 67. Interestingly, the fast (ps–ns) relaxation of the protein was unperturbed by the V66K substitution or by the ionization of Lys-66, supporting the former interpretation. Ab initio modeling of SAXS data also suggested the coexistence of several conformations near the \( pK_a \), one native, and one or more conformations in dynamic equilibrium, probably involving some or all of helix-1, which becomes more exposed to solvent and maybe also more disorganized on complete protonation of Lys-66 at pH values below 5.7.

Application of moderate pressures to the \( \Delta + \)PHS/V66K variant at pH just above the \( pK_a \) of Lys-66 and below the pressure unfolding limit allowed observation of the effect of pressure on the \( ^{15} \)N and \( ^{1} \)H resonances arising from the folded state manifold. These experiments revealed that only residues in the region adjacent to Lys-66 (helix 1 and β-barrel) showed significant nonlinear pressure profiles of their chemical shifts, indicative of the pressure-induced population of alternate, more open conformations with a smaller specific volume due to increased hydration and loss of cavities. It is of interest that the residues exhibiting the nonlinear pressure effect are found in the same region of the protein, around Lys-66, as those that are affected on protonation of this residue. Although the ensemble of low-lying excited conformations populated by protonation of Lys-66 are surely distinct from those accessed by application of pressure to this variant above the \( pK_a \) of Lys-66, they are likely to share some of the same features, and clearly involve dynamic disordering of the same regions. The internal cavity where Lys-66 resides is smaller in the V66K variant than in the true wild-type or \( \Delta + \)PHS reference protein, then, the variant should be less pressure sensitive. In contrast, hydration of the charged lysine residue should decrease the molar volume; this could account for the pressure-induced conformational changes. The two different types of perturbation that were applied (pressure and pH), operate through fundamentally different physical
principles, but both reveal local plasticity in the V66K variant that were not apparent under standard conditions.

Application of hydrostatic pressures larger than 2 kbar results in the unfolding of the Δ+PHS/V66K protein. The NMR data are consistent with a concerted two state global unfolding. Interestingly, the SAXS data revealed that the pressure-unfolded form of the protein is significantly more compact than that of wild-type SNase under comparable conditions of high pressure, pH and temperature, despite the fact that the calculated volume changes and tryptophan fluorescence properties are nearly identical. The more condensed chain at high pressure for the unfolded state of the Δ+PHS/V66K variant could arise from residual stabilizing interactions in the hydrophobic core that are inherent to the Δ+PHS reference protein (as compared with the true wild-type form of SNase). The differences in compactness between the unfolded states of the variants of SNase reported here and previously may be less apparent using chemical denaturants because the physical basis for chemical denaturation involves interaction of the denaturant with the polypeptide chain, and, hence, denaturants, unlike pressure, intrinsically favor expanded structures. In contrast, pressure acts simply to reduce volume, and, hence, residual interactions that do not contribute to volume effects are often preserved at high pressure.32–35 Apparently, further disruption of the core in the case of wild-type SNase compared with the compact denatured state populated by the Δ+PHS/V66K protein is not associated with a further decrease in volume. The differences observed in chain compaction in the pressure unfolded ensemble for the V66K variant compared with wild-type SNase suggests that pressure may reveal intrinsic sequence-dependent characteristics of the unfolded state that are obscured using harsher, chemical perturbation.

The detailed examination of the structural and dynamic consequences of the presence and ionization of groups in the hydrophobic core of a protein has revealed that the response of the protein can be rather subtle. The comparison of site-specific structural and dynamic information obtained from NMR with the global information about protein shape and chain condensation available from SAXS has confirmed that there is a change, that the change is localized to the region in the vicinity of Lys-66, but that, surprisingly, most of the protein remains intact after the ionization of the internal group. By using coupled changes in hydrostatic pressure and pH to perturb the system, we were able to examine the effect of the substitution on the local malleability of the structure, as well as the surprising robustness of the folded ensemble. These important observations confirm that the apparent dielectric constant needed to reproduce the pKₐ of Lys-66 with continuum electrostatics methods is high because the subtle, dynamic structural reorganization coupled to the ionization of the internal Lys needs to be accounted for implicitly in this dielectric constant.5,8,9 New microscopic methods that account explicitly for conformational reorganization coupled to the ionization of internal groups will be required for accurate calculation of the pKₐ of Lys-66 and for understanding the structural basis of function in proteins that depend on internal ionizable groups to perform energy transduction.36–38

MATERIALS AND METHODS

Protein preparation

The Δ+PHS variant of SNase and the Δ+PHS/V66K were prepared as described previously.7 Uniform 15N labeling was obtained from over-expression of recombinant protein in E. coli grown in M9 medium containing 15NH₄Cl as the sole nitrogen source. After purification, proteins were dialyzed against 1M KCl, then against water before lyophilization.

NMR relaxation dispersion-HD exchange and pH titrations

Protein samples were dissolved at 1mM concentration in 600 µl of Tris 10 mM buffer (for data at pH 6.5) or in Acetate 25 mM buffer with 150 mM NaCl (for data at pH 4.5). 5% of D₂O was then added for the lock procedure. Temperature was carefully adjusted to 20°C using a calibration sample (80% glycol in d₆-DMSO). For pH titration adjustments of pH values to 7.1, 6.9, 6.6, 6.0, 5.6, 5.1, 4.6, 4.3, and 4.0 was done by small additions of HCl and NaOH solutions. 1H and 15N chemical shift variations were measured on HSQC spectra recorded at 600 MHz on Δ+PHS and Δ+PHS/V66K 15N-labeled samples, on a Bruker Avance III spectrometer.

Heteronuclear relaxation parameters were obtained with Δ+PHS and Δ+PHS/V66K samples uniformly enriched in 15N at pH 4.5 on Bruker Avance III spectrometers operating at three different magnetic fields corresponding to proton frequencies of 400, 500, and 600 MHz. Heteronuclear longitudinal relaxation rates (R₁) measurements were performed at these three fields using 10 inversion-recovery experiments with a relaxation delay of 18 to 1206 ms. Heteronuclear transversal relaxation rates (R₂) were obtained at 500 MHz using 10 CPMG experiments with a relaxation delay of 16 to 192 ms. Heteronuclear NOE values were calculated from two experiments performed at 500 MHz, one collected with and one without proton presaturation. The final five relaxation rates measured for each individual amide vectors were converted into five spectral density values following the fast spectral density mapping strategy.39 The corresponding spectral density functions were then fitted by a “model free” approach, using the DYNA MOF software, allowing calculation of the order parameter and specific correlation times for each residue.

CPMG dispersion relaxation experiments were performed at 500 MHz using 10 frequencies from 101.874 to 2267.274 s⁻¹ and a τCPMG delay of 40 ms. The sample
protection factors were calculated from the observed exchange limit expression (Meiboom-Gill model).

H/D exchange

For H/D exchange experiments, samples were dissolved in D2O at a concentration of 1 mM after lyophilization. [^1H,^15N]-HSQC spectra were recorded at either 500 or 600 MHz. The time series used for each set of experiments ranged from 5 mn to 120 h (Δ+PHS pH 6.5), 5 mn to 136 h (Δ+PHS/V66K pH 6.5), 20 mn to 148 h (Δ+PHS pH 4.5) and 20 mn to 96 h (Δ+PHS/V66K pH 4.5). Amide proton protection factors were calculated from the observed exchange rates (kex) obtained from the time dependence of the peaks intensities using an exponential decay.

High pressure NMR

For the high pressure NMR experiments, protein samples were prepared with 20 mM d-Tris-HCl buffer. Protein concentration and sample pH were 1.2 mM and pH 6.3, respectively. NMR data was collected at 21 °C. Spectral assignments of 1H and 15N cross-peaks for the proteins have been done previously. High-pressure NMR experiments were performed on a Bruker DRX 800 spectrometer combined with the on-line high pressure NMR cell technique as previously described. At all pressures, 1H chemical shifts were directly referenced to the methyl resonance of DSS, while 15N chemical shifts were referenced indirectly to the absolute frequency ratios \( \frac{\text{15N}}{\text{1H}} = 0.101329118 \).

Under physiological conditions, the protein can be considered to be nearly fully folded in conformer N, giving NMR signal intensity \( I_N \) for site \( i \). When with increasing pressure either local or global unfolding takes the protein into another conformer U, the original signal intensity \( I_N \) for site \( i \) decreases to a final value, \( I_p \), which at each measured pressure is \( I_p \). The decrease in the intensity, \( I_N - I_p \), divided by the intensity \( I_p \) represents the equilibrium constant \( [U]/[N] \) at site \( i \):

\[
K_i(p) = \frac{[U]}{[N]} = \frac{(I_N - I_p)}{(I_N - I_p)}
\]

In actual application, \( I_p \) represents individual cross peak intensities in two-dimensional HSQC spectra. The Gibbs free energy difference \( \Delta G(p) \) between the two conformers is expressed as a function of pressure \( p \) by Eq. (2), under the assumption of negligible changes of differential compressibility with pressure between the two conformers.

\[
\Delta G(p) = -RT \ln K = \Delta G^0 + \Delta V^0 (p - p^0)
\]

Here, \( R \) is the gas constant, \( T \) is the absolute temperature, \( K(p) \) is the equilibrium constant, and \( \Delta G(p) \) and \( \Delta V^0 \) are the Gibbs free energy differences at pressure \( p \) and \( p_0 (=1 \text{ bar}) \), respectively. \( \Delta V^0 \) is the partial molar volume difference between the two conformers at pressure \( p_0 (=1 \text{ bar}) \). The experimentally measured values of \( I_p \) were fitted to Eqs. (1) and (2), for the parameters, \( \Delta G^0 \) and \( \Delta V^0 \) and the asymptotic values \( I_0 \) and \( I_f \) using the program, Bioeqs.43,44

SAXS sample preparation

For the SAXS experiments, protein solutions were prepared immediately before the measurements were performed. To avoid pressure-induced changes of solution pH, 25 mM Tris buffer was used in the pH range of 6–7, respectively. At pH 4.5, these buffers do not keep the pH constant. Thus, a 15 mM Na acetate buffer solution was used instead. All buffers were prepared with deionized water. The pH was adjusted using concentrated 6N HCl. The protein concentration in the solutions was 3.5–5.0 mg mL\(^{-1}\). At this concentration, the solution is diluted enough so that the single scattering approximation is valid.46 However, the scattering signal is still intense enough.

SAXS experiments

SAXS experiments were performed at beamline BL9 of DELTA (TU Dortmund, Germany) and at beamline BW4 of HASYLAB (DESY, Hamburg, Germany). To investigate the SAXS signal of the protein solutions under high pressure, a special sample cell using two flat diamond windows was used.49 A range of 1 bar to 3 kbar was covered in the experiments. Owing to the different brilliances at the various beamlines of the synchrotron facilities used, the exposure time of the protein solution with X-rays was adapted to the experimental conditions. At BW4 (\( \lambda = 1.3808 \) A\(^{-1}\)), using the Pilatus 300 K detector, a \( q \)-range 0.2 to 2.8 nm\(^{-1}\) could be covered. The exposure time was 20 min. A quite similar exposure time was used at BL9 by using a MAR345 image plate detector. A wavelength of \( \lambda = 1.239 \) A\(^{-1}\) allowed covering a \( q \)-range of 0.3 to 3.5 nm\(^{-1}\).

SAXS data analysis

In the case of highly diluted protein solutions, the SAXS signal \( I(q) \) is the spherically averaged squared modulus of the Fourier transform of the protein electron density, \( \rho(\mathbf{r}) \):

\[
I(q) = n \cdot P(q) = n \cdot \left( \int_{\Omega} \rho(\mathbf{r}) \cdot e^{i\mathbf{q} \cdot \mathbf{r}} d^3r \right)^2
\]

where \( n \) denotes the protein’s number density, \( P(q) \) the protein’s form factor and \( \langle \cdot \rangle_\Omega \) denotes the spherical average. The wavevector transfer \( q = (4\pi/\lambda) \sin \theta \) is a function of the scattering angle 2\( \theta \) and the incident
wavelength $\lambda$. For small $q$ values, the scattered intensity can be expressed by the so-called Guinier approximation:

$$I(q) = I(0) \cdot e^{-R_g^2 q^2/3}. \quad (2)$$

The radius of gyration $R_g$, describing the characteristic size of a scattering object, is given as: \text{50}

$$R_g^2 = \frac{\int \rho(r) \cdot r^2 \, dr}{\int \rho(r) \, dr} = \frac{\int_0^{D_{max}} p(r) r^2 \, dr}{2 \int_0^{D_{max}} p(r) \, dr}, \quad (3)$$

where $D_{max}$ is the maximum dimension of the protein and $p(r)$ is the so-called pair-distance distribution function, characterizing the shape and size of the particle. The Guinier approximation is valid for $q$-values up to $q_{\text{max}} \leq 1.3/R_g$. \text{27}

After azimuthally averaging of the SANS spectra, the scattering signals were normalized and corrected for solvent and background scattering by subtracting the scattering signals of the corresponding pure buffer solution. To obtain the radii of gyration, the low $q$-range showing a linear curve shape in the $I(q)$ versus $q^2$ representation was fitted according to Guinier’s approximation. The pair-distance distribution functions were calculated by fitting the scattering data using the program GNOM.\text{51} To reveal if the proteins are folded or denatured, SANS curves are shown in the so-called Kratky plot, that is, $q$ versus $q^2 \cdot I(q)$. In the case of a globular, folded protein, the scattering curve exhibits a distinct maximum, whereas in the case of an unfolded polypeptide, the curve displays a flat plateau.\text{51} For each structure, 16 independent ab initio calculations were started using the program DAMMIF.\text{29} The resulting 16 ab initio structures were aligned by using DAMAVER to build the “most probable” model for each protein and the crystal structures were fit into the ab initio models using the same program.\text{52,53}

REFERENCES


