# Structural and Thermodynamic Characterization of T4 Lysozyme Mutants and the Contribution of Internal Cavities to Pressure Denaturation<sup>†</sup>

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ABSTRACT: Using small-angle X-ray scattering (SAXS) and tryptophan fluorescence spectroscopy, we have identified multiple compact denatured states of a series of T4 lysozyme mutants that are stabilized by high pressures. Recent studies imply that the mechanism of pressure denaturation is the penetration of water into the protein rather than the transfer of hydrophobic residues into water. To investigate water penetration and the volume change associated with pressure denaturation, we studied the solution behavior of four T4 lysozyme mutants having different cavity volumes at low and neutral pH up to a pressure of 400 MPa (0.1 MPa = 0.9869 atm). At low pH, L99A T4 lysozyme expanded from a compact folded state to a partially unfolded state with a corresponding change in radius of gyration from 17 to 32 Å. The volume change upon denaturation correlated well with the total cavity volume, indicating that all of the molecule's major cavities are hydrated with pressure. As a direct comparison to high-pressure crystal structures of L99A T4 lysozyme solved at neutral pH [Collins, M. D., Hummer, G., Quillin, M. L., Matthews, B. W., and Gruner, S. M. (2005) Proc. Natl. Acad. Sci. U.S.A. 102, 16668-16671], pressure denaturation of L99A and the structurally similar L99G/E108V mutant was studied at neutral pH. The pressure-denatured state at neutral pH is even more compact than at low pH, and the small volume changes associated with denaturation suggest that the preferential filling of large cavities is responsible for the compactness of the pressure-denatured state. These results confirm that pressure denaturation is characteristically distinct from thermal or chemical denaturation.

The characterization of non-native states stabilized under a variety of conditions is important toward understanding how proteins fold into their biologically functional, native structures (2). It is widely believed that the dominant driving force in protein folding is the hydrophobic effect and that denaturation can be described as the transfer of hydrophobic residues to water. Although the hydrophobic compound transfer model (3-6) largely succeeds in explaining the thermodynamic stability of proteins as a function of temperature, it does not explain denaturation with pressure (7). In particular, this model fails to explain the magnitude and pressure dependence of the volume difference between the native and denatured states. Recent simulation studies and experimental work suggest that this failure is due to fundamental differences between the temperature- and pressure-denatured states of a protein (1, 8-12). These studies suggest that unlike thermally and chemically denatured states,

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The literature raises specific questions that require further study. Is pressure denaturation consistent with the water penetration model? How does the volume change upon denaturation correlate with cavity volume? Can other hydration mechanisms be distinguished from cavity filling? What constitutes a pressure-denatured state? We attempt to answer these questions by characterizing the pressure-denatured states of several mutants of the protein T4 lysozyme with varying cavity sizes.

T4 lysozyme is a small globular protein, 164 amino acid residues in length, with a molecular weight of 18.6 kDa (Figure 1 (f)) (13). Over 300 X-ray structures of T4 lysozyme mutants have been deposited in the Protein Data Bank, and the thermal stabilities of many of these mutants have been measured (14). Pressure-induced water filling of an enlarged hydrophobic cavity in the L99A mutant of the pseudo-wild-type T4 lysozyme (Figure 1 (b), cavity 6) has been observed by X-ray crystallography (1). Molecular dynamics simulations suggested that four water molecules cooperatively fill this cavity as the applied hydrostatic pressure is increased (1). No water molecules were

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FIGURE 1: T4 lysozyme structures shown from the same perspective. The external surface and buried cavities (shown in magenta) of (a) WT\* (1L63), (b) L99A (1L90), (c) L99G/E108V (1QUH), (d) A98L (1QS5), and (e) V149G (1G0P) identified with a 1.2 Å probe in MSMS (*34*) with all internal solvent molecules removed. Cavities are identified by numbers 1-10 (refer to Table 2). (f) Cartoon representation of WT\* T4 lysozyme. The C-terminal lobe is on the top side.

observed in the corresponding cavity of the pseudo-wild-type,  $WT^{*1}$  (Figure 1 (a), cavity 6) (1).

In this study, the solution behavior of L99A and three additional T4 lysozyme mutants (Figure 1) was compared using high-pressure small-angle X-ray scattering (SAXS) and tryptophan fluorescence spectroscopy. The three additional mutants were identified from approximately 300 T4 lysozyme mutants available as satisfying the criteria of having similar thermal stabilities as L99A, approximately 5 kcal/mol less than that of WT\* (13, 15-18), yet having differing internal cavity volumes. These features were important for this study in order observe complete denaturation transitions of all the mutants under identical solvent conditions and to determine the effect of internal cavity volume on the volume change of pressure denaturation. Two of the investigated mutants, L99A and L99G/E108V, are structurally very similar to WT\* except for differing volumes of the hydrophobic cavities at the mutation site where the 99th residue, leucine, L, has been replaced by alanine, A, or glycine, G (15, 16). The V149G mutation enlarges an existing solvent-bound polar cavity at the mutation site (17). A98L is destabilized relative to WT\* by a strain-inducing small-to-large amino acid substitution (18). SAXS was used to characterize the pressure-induced change in radius of gyration and overall shape of L99A T4 lysozyme. Thermodynamic analysis was performed on tryptophan fluorescence data taken at various solvent conditions on all mutants. A fluorescence quenching study of a selenomethionine variant of L99A provided additional information on global structural changes due to high pressure.

The results provide insight into the relationship between the thermodynamic and structural characteristics of T4 lysozyme. In particular, we characterize the contributions of cavity volume of the native state and electrostriction to the volume change of pressure denaturation. Low-resolution models of native and pressure-denatured solution structures have been determined. Multiple, compact pressure-denatured states are identified with differing degrees of hydration. These results confirm that pressure denaturation is characteristically distinct from thermal and chemical denaturation. It is shown that the study of pressure effects provides further understanding of the energy landscape and the structure—stability relationship of proteins.

### **EXPERIMENTAL PROCEDURES**

Sample and Buffer Preparation. In this work the cysteinefree pseudo-wild-type, C54T/C97A T4 lysozyme (13), is referred to as WT\*. All mutants used in this study contain the mutations C54T and C97A. The compositions of cell culture media used in these procedures are listed in supporting material 1 in the Supporting Information.

The L99A, L99G/E108V, and A98L mutants were expressed and purified by a modified version of the protocol described by Poteete et al. (19) (supporting material 2 in the Supporting Information). The V149G mutant was isolated using a modified version of the inclusion body protocol described by Vetter et al. (20) (supporting material 3 in the Supporting Information). A selenomethionine containing variant of the L99A mutant (Se-Met L99A) was prepared in a methionine deficient, selenomethionine rich growth medium using an adaptation of the procedure by Van Duyne et al. (21) using non methionine-auxotrophic cells (22) (see supporting material 4 in the Supporting Information).

High concentration buffers with low volume changes of ionization were chosen to stabilize the pH as a function of pressure (23, 24). At pH 3.0, 50 mM glycine buffer (Cat. 17-1323-01; GE Healthcare Bio-Sciences Corp., Piscataway, NJ) and at pH 7.0, 50 mM Tris-HCl buffer (T-3253; Sigma, St. Louis, MO) were used, unless otherwise noted. The NaCl (Cat. 7581; Mallinckrodt Baker, Phillipsburg, NJ) concentrations used were 20 and 100 mM. Buffers were sterilized with 0.22  $\mu$ m cellulose acetate filters (Catalog Number 431175; Corning Inc., Corning NY), stored at 4 °C, and used within two weeks of preparation.

High-Pressure Small Angle X-ray Scattering. SAXS samples were prepared up to 48 h in advance of experiments by dialyzing against buffers in microdialysis buttons (Cat. HR3-362; Hampton Research, Aliso Viejo, CA) closed with a 10 kDa molecular weight cutoff dialysis membrane (Cat. 68100; Pierce Biotech, Rockford, IL). Protein solution concentrations were adjusted to 4 to 25 g/L by UV absorption measurement (Nanodrop ND1000; Nanodrop Technologies, Wilmington, DE) and centrifugal reconcentrator; Millipore, Billerica, MA). The final dialysate solutions were used for sample dilutions and background scattering measurements. Samples were stored at 4 °C or on ice. Experiments were performed at room temperature.

<sup>&</sup>lt;sup>1</sup> Abbreviations: SAXS, small-angle X-ray scattering; SVD, singular value decomposition; EOM, the SAXS program Ensemble Optimization Method; WT\*, a cysteine-free pseudo-wild-type mutant of T4 lysozyme; L99A, a leucine-99 to alanine mutant of WT\*; L99G/E108V, a mutant of WT\* with the mutations, leucine-99 to glycine and glutamic acid-108 to valine; A98L, an alanine-98 to leucine mutant of WT\*; V149G, a valine-149 to glycine mutant of WT\*; Se-Met L99A, L99A with methionines replaced by selenomethionines.

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sample	condition	$\langle \lambda \rangle_{\rm N} \; ({\rm nm})^b$	$\langle \lambda \rangle_{\rm D} \ ({\rm nm})^b$	$\Delta G^{\circ}$ (kcal/mol)	$\Delta V^{\circ}$ (Å <sup>3</sup> )	$P_{\rm m}~({\rm MPa})^c$
			pH 3.0, 16 °C			
L99G/E108V	20 mM NaCl	$338.4 \pm 1.0$	$360.3 \pm 0.3$	$1.75 \pm 0.24$	$-235.8 \pm 25.0$	51.7
	100 mM NaCl	$339.0 \pm 0.5$	$359.3 \pm 0.2$	$2.21 \pm 0.17$	$-212.8 \pm 13.8$	72.3
L99A	20 mM NaCl	$339.2 \pm 0.4$	$360.6 \pm 0.2$	$2.54 \pm 0.15$	$-187.3 \pm 10.0$	94.5
	100 mM NaCl	$339.7 \pm 0.2$	$358.5\pm0.2$	$2.75 \pm 0.14$	$-161.4 \pm 7.5$	118.7
A98L	20 mM NaCl	$341.6 \pm 0.2$	$357.4 \pm 0.3$	$2.49 \pm 0.13$	$-127.2 \pm 6.8$	136.0
	100 mM NaCl	$341.8 \pm 0.4$	$353.7\pm0.2$	$2.07\pm0.22$	$-107.7 \pm 10.0$	133.7
V149G	20 mM NaCl	$343.3 \pm 0.4$	$358.7\pm0.2$	$1.30 \pm 0.10$	$-134.1 \pm 7.3$	67.4
	100 mM NaCl	$342.4\pm1.0$	$352.3\pm0.2$	$0.58\pm0.17$	$-99.2\pm10.0$	40.8
			pH 7.0, 24 °C			
L99G/E108V	20 mM NaCl	$340.3\pm0.2$	$355.6\pm0.3$	$2.56\pm0.17$	$-102.8 \pm 7.0$	172.8
L99A	20 mM NaCl	$340.9\pm0.2$	$356.1\pm0.5$	$3.19\pm0.26$	$-93.1\pm8.2$	238.1

<sup>*a*</sup> Results of fitting Equation 8 (Experimental Procedures) to data. <sup>*p*</sup> Subscripts "N" and "D" refer to the native and denatured states, respectively. <sup>*c*</sup>  $P_m$  is the midpoint pressure of denaturation.

Table 2: Cavity Volumes of T4 Lysozyme Mutants <sup>a</sup>							
cav. #	WT* (1L63)	WT* $(MC)^b$	L99A (1L90)	L99G/E108V (1QUH)	A98L (1QS5)	V149G (1G0P)	comments
0	21044.0	21113.6	21103.3	20866.9	20919.7	21312.7	protein Surface
1	50.3 (0)	44.9 (0)	51.9 (0)	47.5 (0)	46.8 (0)	49.2 (0)	Sol 171, 179
2	4.6				10.4		
3	25.3 (0)	22.2 (0)	27.6 (0)	23.6 (0)	37.8 (0)	$106.3 (0)^{c}$	Sol 208, Trp138
4	25.8 (0)	29.3 (4.2)	26.7 (0)			27.5 (0)	Sol 175
5	7.1	11.9	9.2		7.7	9.6	
6	39.2	39.1	161.1	$222.5(87.2)^d$	53.7	40.7	
7	24.6	23.9	28.9	24.3	24.3	25.0	Trp126
8				5.0 (0)	7.1 (0)		Sol 213
9				5.5	8.5		
10					12.1		

<sup>*a*</sup> Volumes in Å<sup>3</sup> of molecular surfaces found in crystallographic structures (Experimental Procedures) of T4 lysozyme mutants with internal solvent molecules removed. Volumes of solvent-containing cavity determined with internal solvent molecules kept in the structures are shown inside the parentheses. The molecular surfaces were found using a 1.2 Å probe in MSMS (*34*). Molecular surface 0 is the outer surface of the proteins, while 1–10 were cavities isolated from the external solvent. Surfaces detected by MSMS but found to intersect with the external surface of the protein are not shown. <sup>*b*</sup> WT\* structure was provided by Marcus Collins (private communication). <sup>*c*</sup> Cavity 3 in V149G contained Sol 323 (alternative site 423) and 324 according to the crystal structure. <sup>*d*</sup> Similarly, cavity 6 in L99G/E108V contains Sol 401 and 402.

The custom built high-pressure SAXS cell, equipment, and data reduction procedures were previously described (25). X-ray experiments were performed at the G1 station at the Cornell High Energy Synchrotron Source (CHESS, Cornell University, Ithaca, NY). Data on 10 g/L L99A T4 lysozyme in 50 mM glycine, 100 mM NaCl, pH 3.0 (q range of 0.019-0.328 Å<sup>-1</sup>) and 16 g/L WT\* T4 lysozyme in 50 mM glycine, 150 mM NaCl, pH 3.0 (q range of 0.025-0.295  $Å^{-1}$ ) were collected with a detector distance of 1.25 m and a 12 keV 250  $\mu$ m square beam. The salt concentration was adjusted at each protein concentration to minimize protein interactions (25). The momentum transfer, q, is defined as q $= 4\pi/\lambda \sin \theta$ , where  $2\theta$  is the scattering angle and  $\lambda$  is the X-ray wavelength. The samples were contained in custom acrylic inner cells (ALine Inc., Redondo Beach, CA) (25) with 7.5  $\mu$ m Kapton film windows (Spectromembrane No. 3022; Chemplex Industries, Palm City, FL). Data were collected on a home-built  $1024 \times 1024$  pixel CCD detector, and the transmission intensity was measured with a homemade PIN diode beamstop. At each pressure, protein samples were equilibrated for 10 to 60 min. Buffer scattering was measured at each pressure. Measurements were repeated on fresh samples to identify radiation damage. Exposures exhibiting radiation damage were not used for analysis.

The scattering profile of the protein, I(q), was produced by subtracting the transmission-normalized scattering profile of the buffer taken at the sample pressure from that of the protein solution without further scaling. A Guinier approximation can be applied to the low q region of the scattering profiles.

$$I(q) \approx I(0) \exp(-q^2 R_g^2/3)$$
 (1)

The radius of gyration,  $R_g$ , and zero-angle scattering intensity, I(0), are determined from a linear fit to the Guinier plot,  $\log(I(q))$  vs q, in the approximate region where the condition  $qR_g < 1.3$  holds (26). I(0) is a function of the electron density contrast between the protein and solvent,  $\Delta \rho$ , the excluded volume of the protein,  $V_p$ , and the structure factor, S(q), which is a function of protein—protein interactions (27).

$$I(0) \propto S(0) (\Delta \rho \cdot V_{\rm p})^2 \tag{2}$$

The radius of gyration can also be determined from the pair distance distribution function, P(r) (27).

$$R_{g}^{2} = \frac{\int_{0}^{D_{\text{max}}} P(r)r^{2} \,\mathrm{d}r}{2\int_{0}^{D_{\text{max}}} P(r) \,\mathrm{d}r}$$
(3)

P(r) was calculated with program GNOM, which implements the indirect Fourier transform method (27, 28) using the experimental I(q) as the input. This method is sensitive to low q data and, therefore, the choice of the maximum dimension of the protein,  $D_{\text{max}}$ , where P(r) approaches zero. Thus, it was only applied to data where  $D_{\text{max}}$  could be unambiguously determined. Low-resolution models of protein structures were generated from the I(q) produced by GNOM using the *ab initio* reconstruction program GASBOR (29). The scattering profile of the denatured state was also analyzed with the Ensemble Optimization Method (EOM) package developed for modeling flexible proteins in solution (*30*).

*High-Pressure Fluorescence Spectroscopy.* Three tryptophan residues are present in the C-terminal lobe of T4 lysozyme (Trp126, Trp138, and Trp158). Samples for tryptophan fluorescence spectroscopy were prepared up to 48 h in advance of experiments by dialyzing 100–200  $\mu$ L protein solutions in microdialysis floats (Cat. PI-69560; Thermo Scientific Pierce Slide-A-Lyzer, Thermo Fisher Scientific, Waltham, MA). The dialyzed protein solutions were further diluted to a final volume of 2.4–2.5 mL, resulting in protein concentrations of <3  $\mu$ M. No contaminants from the dialysis step were detected spectroscopically.

A high-pressure fluorescence cell (HP Cell; ISS, Champaign, IL) equipped with sapphire windows was used with a scanning fluorimeter (Chronos; ISS, Champaign, IL). The pressurizing medium in the high-pressure cell, isopropanol, was separated from the pressurizing medium (Fluorinert FC-77; 3M, St. Paul, MN) in the high-pressure pump (Cat. No. 37-6.75-60; High Pressure Equipment Co., Erie, PA) by a piston in a home-built stainless steel high-pressure reservoir. Samples were contained in a square bottle with a cylindrical neck formed by fusing a short 3 mm inner diameter quartz tube to the top of a 10 mm path length square quartz cell (Cat. No. CQ110; VitroCom, Mountain Lakes, NJ). The sample-filled bottle was sealed with a soft bulb made from polyolefin heat shrink tubing (Cat. No. 03F3745; Newark Electronics, Chicago, IL) and positioned in the high-pressure cell with a homemade Delrin cuvette holder. The large flat windows of this cuvette minimize scattering of the incident beam, while the soft bulb closure acts as a deformable pressure-transmitting membrane.

Samples were excited at 295 nm with a bandwidth of 8 nm to minimize tyrosine excitation, and emission from 310 to 550 nm was collected at 90° with a bandwidth of 4 nm. Vinci software (ISS, Champaign, IL) was used for instrument control and data acquisition. Protein solutions were equilibrated at each pressure for 20 to 90 min. Buffer reference spectra were pressure independent and featureless, with the exception of a small pressure independent water Raman peak around 330 nm. For this reason only one representative reference spectrum was taken at 50 MPa for each protein sample. Sample temperature was maintained with a water circulation pump (Neslab, Thermo Scientific, Waltham, MA).

Data were analyzed with MATLAB (The MathWorks, Natick, MA). The collected emission spectra were background-subtracted. Intensity corrections were performed with calibration data collected by ISS (Champaign, IL) with 4 nm excitation and emission bandwidths. The center of spectral mass,  $\langle \lambda \rangle$ , of each spectrum was calculated in the region 310 to 550 nm with the following equation (*31*):

$$\langle \nu \rangle = \sum_{i=1}^{N} \nu_i I_i / \sum_{i=1}^{N} I_i$$
(4)

where  $I_i$  is the fluorescence intensity at the wavenumber,  $\nu_i$ , and N is the number of data points collected in the spectrum. The weighted average  $\langle \nu \rangle$  was then converted to wavelength,  $\langle \lambda \rangle$ . For two-state denaturation, the thermodynamic stability of the protein is the free energy difference between the native (N) and denatured (D) states.

$$\Delta G = G_{\rm D} - G_{\rm N} \tag{5}$$

Assuming that the protein solution is an ideal solution,  $\Delta G$  at a given pressure can be expressed as a function of the observable quantity, the center of spectral mass.

$$\Delta G = -RT \ln K_{\rm eq} = -RT \ln \left( \frac{\langle \lambda \rangle - \langle \lambda \rangle_{\rm N}}{\langle \lambda \rangle_{\rm D} - \langle \lambda \rangle} \right) \tag{6}$$

where  $K_{eq}$  is the equilibrium constant, *R* is the gas constant (83.1447 mL-bar/mol-K), *T* is the experimental temperature, and  $\langle \lambda \rangle_N$  and  $\langle \lambda \rangle_D$  are the spectral centers of mass of the native and denatured states.  $\langle \lambda \rangle_N$  and  $\langle \lambda \rangle_D$  were inferred by fitting experimental data. No baseline corrections to the spectral centers of mass were necessary.

The dominant contribution to the pressure-induced change in protein stability is the volume change (32, 33). Thus, a first-order approximation was used to model the denaturation of T4 lysozyme. The pressure stability with respect to a reference pressure,  $P_0$ , can be expressed by

$$\Delta G = \Delta G^{\circ} + \Delta V^{\circ} (P - P_0) \tag{7}$$

where  $\Delta G^{\circ}$  and  $\Delta V^{\circ}$  are the free energy difference and volume change of denaturation at standard pressure  $P_0$ , 1 atm, and a constant experimental temperature, *T*. Combining eqs 6 and 7, we arrive at the following model for our data:

$$\langle \lambda \rangle = \frac{\langle \lambda \rangle_{\rm N} + \langle \lambda \rangle_{\rm D} \exp\{-[\Delta G^{\circ} + \Delta V^{\circ} (P - P_0)]/RT\}}{1 + \exp\{-[\Delta G^{\circ} + \Delta V^{\circ} (P - P_0)]/RT\}}$$
(8)

where  $\langle \lambda \rangle_{\rm N}$ ,  $\langle \lambda \rangle_{\rm D}$ ,  $\Delta G^{\circ}$ , and  $\Delta V^{\circ}$  are free parameters.

*Cavity Volume Calculation and Visualization.* Cavity calculations were performed on the atomic structures of L99A (PDB accession code 1L90), L99G/E108V (1QUH), A98L (1QS5), V149G (1G0P), and WT\* (1L63). An additional atomic structure of WT\* was provided by Marcus Collins (private communication). Cavities were identified using the program MSMS (*34*) with a 1.2 Å probe. The identified molecular surfaces were viewed using the program PyMol (Delano Scientific LLC, Palo Alto, CA) with a script provided by Warren DeLano (private communication).

#### RESULTS

Thermodynamic Analysis of Fluorescence Measurements. The fluorescence of tryptophan is sensitive to the polarity of its local environment and can be used to monitor protein conformational changes. Both the solvent and protein contribute to environmental polarity, such that protein denaturation is generally accompanied by a red-shift in tryptophan fluorescence (35). Fluorescence measurements were made on the T4 lysozyme mutants at pH 3.0 and 7.0. Thermodynamic analysis was performed on the pressureinduced change in the center of spectral mass (defined in Experimental Procedures).

All samples exhibited two-state behavior (native versus denatured) under pressure, consistent with observations made in other denaturation studies (36, 37). The peak intensities of the mutants were normalized to that of WT\*, which did not denature under our conditions, to correct for pressure-

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induced changes in fluorescence quenching and solvent transmission. An isosbestic point was evident in the pressurecorrected spectra of each denaturation series (Figure 6), indicative of two-state denaturation. Singular value decomposition (SVD) analysis determined that the data could be adequately described as linear combinations of two independent SVD states. For each data set, denaturation curves reconstructed from the two significant SVD states fit the experimental spectra with a goodness of fit parameter,  $R^2 \ge$ 0.9986. The reversibility calculated from the center of spectral mass upon decompression to ambient pressure was 80-93%. The low end of this range applied to samples maintained at high pressure for several hours. The effects of nonreversibility were apparent only at low pressure, and therefore, low-pressure data were collected first in order of increasing pressure. Thermodynamic fits (eq 8) performed on the centers of spectral mass yielded goodness of fit parameters,  $R^2 \ge 0.9990$ .

Fluorescence measurements were made on L99G, L99G/ E108V, A98L, V149G, and WT\* at pH 3.0. The thermal denaturation of T4 lysozyme has been extensively characterized at pH 3.0 (15, 16, 18, 38), and therefore this pH was chosen as a major test condition for this work. The low pH also enabled pressure denaturation of all the mutants to complete below 400 MPa, the pressure limit of our apparatus. At pH 7, these mutants are more stable, and not all of them denature at 400 MPa. The temperature was set to 16 °C, where the folded fractions of the least stable mutants, L99G/ E108V and V149G were nearly 1 at ambient pressure. The centers of spectral mass of all mutants showed sigmoidal dependence on pressure, indicative of conformational changes (Figure 2), while that of WT\* did not change up to 300 MPa (not shown). The centers of spectral mass of the native and denatured states were similar for L99A and L99G/E108V at each solvent condition. This observation is consistent with the structural similarity of L99A and L99G/E108V compared with the other two mutants. The steepness of the denaturation curves is a function of the magnitude of the volume change of denaturation,  $\Delta V^{\circ}$ , while the pressure at which the transition begins is indicative of the stability at ambient pressure,  $\Delta G^{\circ}$ . The results from the thermodynamic fits are presented in Table 1. At pH 3.0, A98L and V149G exhibited similar volume changes although V149G was less stable, denaturing at a lower pressure. Compared to A98L and V149G, L99A and L99G/E108V showed large volume changes. The magnitude of  $\Delta V^{\circ}$  was also dependent on the ionic strength of the solvent. An increase from 20 to 100 mM NaCl resulted in a reduction by approximately 25 Å<sup>3</sup> in the magnitudes of  $\Delta V^{\circ}$  for all mutants. At both NaCl concentrations, the volume change of L99G/E108V was roughly 50 Å<sup>3</sup> greater than that of L99A.

For direct comparison with the room-temperature highpressure crystal structures of L99A and WT\* acquired by Collins et al. (1), fluorescence measurements were also made at pH 7.0, 24 °C. L99G/E108V was also studied because of its structural similarity to L99A. We observed sigmoidal transitions for L99A and L99G/E108V as a function of pressure, while no transition was detected for WT\* up to 350 MPa (Figure 3). The denaturation curves of the two mutants are similarly shaped with a gradual transition except that of L99G/E108V is shifted by about 65 MPa to a lower pressure with respect to that of L99A. In contrast to the result



FIGURE 2: Pressure denaturation of T4 lysozyme mutants in pH 3.0 50 mM glycine 20 mM NaCl (diamond) and 100 mM NaCl (circle) buffers monitored at 16 °C by tryptophan fluorescence spectroscopy. (a) L99G/E108V (open) and L99A (closed). (b) V149G (open) and A98L (closed).



FIGURE 3: Pressure denaturation of L99G/E108V (diamond), L99A (circle), and WT\* (plus) T4 lysozyme in pH 7.0 50 mM Tris HCl 20 mM NaCl buffer monitored at 24 °C by tryptophan fluorescence spectroscopy.

at pH 3.0, the volume changes of denaturation of the two mutants at pH 7.0 were similar and small in magnitude (approximately  $-100 \text{ Å}^3$ ). The results of thermodynamic fits are given in Table 1.

Structural Information from SAXS Measurements. Smallangle X-ray scattering is sensitive to electron density distributions with inhomogeneities on the length scales of 10 to 100 Å and thereby provides structural information for proteins in solution. At pH 3.0 where the SAXS measurements were made, T4 lysozyme is highly positively charged. To minimize the effect of interparticle interference on the intensity profile, a specific NaCl concentration was required at a given protein concentration. These interaction-free



FIGURE 4: (a) Radius of gyration,  $R_g$ , as a function of pressure for 10 g/l L99A in 50 mM glycine 100 mM NaCl pH 3.0 buffer at room temperature.  $R_g$  was determined by a Guinier fit to the  $qR_g$ < 1.3 region of scattering profiles taken at each pressure. The error bars are larger at high pressure because only one exposure was taken at each pressure. At lower pressure, the sample was less susceptible to radiation damage-induced aggregation and multiple exposures were taken at each pressure, which enabled averaging of images. A two-state thermodynamic fit is shown (solid line) to guide the eye. (b) The zero-angle scattering intensity of L99A (circle) and the WT\* (diamond) at pH 3.0. WT\* does not denature below 300 MPa.

conditions were first identified at ambient pressure (25). SAXS profiles were measured at pressures ranging from 28 to 300 MPa. Due to the exposure limit set by radiation damage, the data presented here were taken on two aliquots of the same sample. The small-angle X-ray scattering of L99A T4 lysozyme exhibited pressure-reversibility. SAXS was not attempted at pH 7.0 both to limit the scope of the required beam time and because not all the mutants denature within the accessible pressure range at this pH.

Guinier analysis was performed on the low q region of the SAXS profiles. The results of Guinier fits to the data are shown in Figure 4. We previously reported a radius of gyration of  $16.5 \pm 0.3$  Å for L99A at ambient pressure (25), which is in agreement with that calculated from the crystal structure of a folded monomer, 16.4 Å. At 28 MPa, the radius of gyration determined over q = 0.024-0.076 Å<sup>-1</sup> was 17.1  $\pm$  0.1 Å, indicating that L99A is mostly folded at this pressure. As the applied pressure was increased, the radius of gyration increased to  $31.6 \pm 1.7$  Å (determined over q =0.024-0.038 Å<sup>-1</sup>) at 300 MPa. As the radius of gyration ( $R_g$ ) as determined by Guinier analysis is only accurate for homogeneous samples, we do not place strong meaning on the specific value of  $R_g$  at intermediate pressures where L99A was a mixture of folded and denatured forms. Pair distance distribution functions were computed using GNOM (28) from data taken at 28 and 300 MPa with a maximum size,  $D_{\text{max}}$ , of 54 and 134 Å, respectively (Figure 5 (a)). Radii of gyration determined from the pair distance distribution functions were 17.0 and 34.5 Å, in good agreement to the Guinier fits. The  $R_{\rm g}$  of the pressure-denatured state was quite small in comparison to 40.7 Å, the predicted radius of gyration of a fully unfolded polypeptide of the same length (39). WT\* did not show change up to 300 MPa (25). The linearity of the Guinier plots at low q indicated that aggregates were not detected. The 28 and 300 MPa scattering profiles are shown in the inset of Figure 5 (a) as Kratky plots, i.e.  $Iq^2$  vs q. Krakty plots emphasize the I(q) decay in the intermediate q region, which is sensitive to the shape of the protein (40). The 28 MPa profile shows a peak around 0.1  $\text{\AA}^{-1}$ , indicating that L99A is compact and globular at this pressure. At 300 MPa, this peak is greatly reduced and no an additional peak at lower q is evident, indicating that L99A is extended and largely monomeric at high pressure.

The zero-angle scattering intensity, I(0), determined by Guinier analysis is shown in Figure 4 (b). I(0) is a function of the electron density contrast between the hydrated protein and solvent, the excluded volume of the protein, molecular weight, and interprotein interactions. We have previously shown that the electron density contrast of WT\* decreases with pressure because of the greater compressibility of the solvent compared to the protein (25). Relative to WT\*, L99A shows an increase in I(0). As L99A is highly charged at pH 3.0 and sensitive to the ionic strength, we attribute the increase in I(0) to changes in hydration or protein interactions that accompany denaturation (41-43). Although I(0) is also a function of the protein volume, the volume changes due to denaturation are on the order of 1% of the volume of the protein and therefore do not significantly affect I(0).

For further structural characterization, low-resolution models of L99A were produced using the ab initio reconstruction program GASBOR (29) and the denatured ensemble was modeled with the EOM package (30). Five GASBOR reconstructions were performed on the small X-ray scattering data taken at 28 and 300 MPa on the assumption that L99A is a monomer at both pressures. The results of this reconstruction are shown in Figure 5 (c) and (d). The low-pressure reconstructions resembled the crystal structure of L99A in exterior shape and size. The high-pressure reconstructions do not represent actual structures found in solution, as the denatured state is an ensemble without a unique conformation, but rather, they are models that closely match the experimental scattering profile. While the low-pressure structures are compact, the high-pressure reconstructions are noticeably more extended. The small-angle X-ray scattering profile taken at 300 MPa was also examined with EOM. EOM generates a large pool of randomly generated conformations available for a polypeptide of a certain sequence (44) from which an optimization algorithm is used to select the subset that best fits the experimental scattering profile. Two starting pools were generated. The first contained 10,000 random coil conformations with an average  $R_g$  of 38.0 Å, and the second contained 10,000 unfolded conformations with residual structure with an average  $R_{\rm g}$  of 29.7 Å. The ensemble selected from the pool with residual structure had an average  $R_{\rm g}$  of 31.2 Å, in agreement with Guinier analysis, and better described our experimental data than the selected



FIGURE 5: (a) Pair distance distribution functions of native (28 MPa, solid line) and denatured (300 MPa, dotted line) L99A T4 lysozyme at pH 3.0 obtained with GNOM (28). Inset: Kratky representation of the 28 MPa (1) and 300 MPa data (2). (b) The scattering profile of L99A at 300 MPa (black) was examined with the Ensemble Optimization Method (30). An ensemble of unfolded conformers with residual structure (red scattering profile) better described the experimental data compared to an ensemble of random coil conformers (blue scattering profile). The distribution of  $R_g$  in the first ensemble is shown in the inset. (c) Side and top views of the crystal structure of L99A (right) and a representative low-resolution structure at 28 MPa obtained with GASBOR (29) (left) show good agreement. (d) GASBOR models that fit well to 300 MPa data were extended.



FIGURE 6: Tryptophan fluorescence spectra of L99A (color) and Se-Met L99A (black) in 50 mM glycine 20 mM NaCl pH 3.0 16 °C with increasing pressure (direction indicated by arrow). The emission intensities were normalized at high pressure (200 MPa) to emphasize the pressure-induced intensity increase of Se-Met L99A compared with L99A. Inset: Spectra of L99A (red) and Se-Met L99A (black) at low pressure (25 MPa) scaled by a constant factor.

random coil ensemble, which had an average  $R_g$  of 36.1 Å (Figure 5 (b)).

Structural Information from Fluorescence Quenching Measurements. Additional information on global conformational changes was derived from fluorescence quenching measurements. T4 lysozyme has five methionine residues in close proximity to the three tryptophan residues present in the C-terminal lobe. The sulfur atom in methionine quenches tryptophan emission and provides a probe of the methionine-tryptophan separation. Denaturation of T4 lysozyme is usually accompanied by an increase in emission intensity (45), consistent with an increased methionine–tryptophan distance. In the selenomethionine variant of L99A (Se-Met L99A), the methionines are replaced with selenomethionines. As selenium and sulfur have differing quenching strengths (46), a comparison of L99A and Se-Met L99A yields additional information on the spatial compactness of the unfolded state of L99A.

Tryptophan fluorescence measurements were made on Se-Met L99A in pH 3.0, 50 mM glycine, 20 mM NaCl and pH 7.0, 50 mM Tris, 20 mM NaCl buffers and compared to those of L99A under the same conditions. The intensity increase accompanying denaturation was more pronounced for Se-Met L99A than for L99A, consistent with the stronger quenching ability of selenium (Figure 6). Figure 7 (a) shows that at each pressure, the centers of spectral mass were similar for L99A and Se-Met L99A in the same solvent, indicating that these two mutants are structurally similar at any given pressure and that the introduction of selenomethionines does not significantly distort the shape of the fluorescence spectra or change the denaturation behavior.

Pressure-induced changes to the fluorescence yield of tryptophan or transmission through the solvent were factored out in the ratio of the emission intensities of L99A and Se-Met-L99A,  $I_{L99A}$  and  $I_{Se-MetL99A}$ , at the same pressure. The ratios were calculated as follows at each pressure:

$$\left\langle \frac{I_{\text{Se-MetL99A}}}{I_{\text{L99A}}} \right\rangle = \frac{1}{N} \sum_{i=1}^{N} \frac{I_{\text{Se-MetL99A}}(\lambda_i)}{I_{\text{L99A}}(\lambda_i)} \tag{9}$$

The sum was taken over points in the wavelength range 310-400 nm where the fluorescence signal is strong. *N* is the number of ratios in the sum (91, in this case). Figure 7 (b) shows a comparison of these ratios at pH 3.0, 50 mM glycine, 20 mM NaCl and pH 7.0, 50 mM Tris, 20 mM NaCl as a function of denatured fraction. They are normalized at 0.1 MPa. We see that as denaturation progresses, the ratio is greater at pH 3.0 than at pH 7.0. This result suggests that the denatured state at pH 7.0 is more compact than at pH 3.0.



FIGURE 7: Pressure denaturation of L99A and Se-Met L99A in 50 mM glycine 20 mM NaCl pH 3.0 16 °C and 50 mM Tris HCl 20 mM NaCl pH 7.0 24 °C. (a) Centers of spectral mass of L99A (open) and Se-Met L99A (closed) at pH 3.0 (diamond) and pH 7.0 (circle). L99A and Se-Met L99A show similar denaturation curves, indicative of structural similarity at each pressure. (b) Normalized intensity ratios of Se-Met L99A and L99A emission at pH 3.0 (closed diamond) and pH 7.0 (closed diamond) and pH 7.0 (open circle). At pH 7.0, the ratio shows little change with pressure, while at pH 3.0, there is a large increase. This indicates a greater separation between tryptophan and selenomethionine residues in the denatured state at pH 3.0 compared with pH 7.0, suggesting that the pH 3.0 denatured state is more unfolded than at pH 7.0.

Cavity Volume Calculations and Internal Waters. The volumes contained by the external surface and the volumes of internal cavities present in the crystal structures of the T4 lysozyme mutants were calculated to determine if a correlation exists between the volume changes of denaturation extracted from thermodynamic analysis and the volume of buried cavities. In order to identify and measure buried cavities, crystal structures (PDB accession codes 1L90 for L99A, 1QUH for L99G/E108V, 1QS5 for A98L, 1G0P for V149G, and 1L63 for WT\*) were analyzed with a 1.2 Å probe using MSMS (34). MSMS implements a rolling probe to determine reduced surfaces, from which the solvent excluded surface can be computed. Two sets of molecular surfaces were produced for each structure. In the first set of calculations, all crystallographically determined solvent particles were manually removed from the PDB files prior to analysis to eliminate false surface cavities caused by surface water, and to enable detection of hydrated internal cavities. The external surface and six to nine cavities were identified for each mutant (Figure 1). In the second set, internal water molecules were retained to determine changes in cavity size due to hydration. Four water molecules, Sol 171, 179, 175, and 208, are considered conserved internal solvent molecules in T4 lysozyme (47). Using a 1.2 Å probe in MSMS (34) to define the molecular surfaces, Sol 213 was also observed within the external surface of all mutants. Sol 175 was found either in a solvent-exposed pocket on the surface or within the external surface. As Sol 175 was considered to fully occupy its cavity, its exact location did not contribute to the total cavity volume. Both L99G/E108V and V149G structures contain two additional poorly ordered water molecules in their respective mutation-enlarged cavities. The L99G/E108V structure contains Sol 401 and 402 in the hydrophobic cavity near the L99G mutation (Figure 1 (c), cavity 6). The V149G mutation enlarges the polar cavity binding Sol 208 (Figure 1 (e), cavity 3) and introduces Sol 323 (alternative site 423) and 324. Molecular surfaces that intersected or were located outside the external surface were not considered internal cavities. For comparison, a second crystal structure of WT\* (provided by Marcus Collins, private communication) was also analyzed. The variability of individual cavity volumes in the two WT\* structures was within 5.4 Å<sup>3</sup> (3% of the total cavity volume of WT\*). Table 2 summarizes the results.

#### DISCUSSION

Structural Characterization of the Native and Pressure-Denatured States in Solution. The low-resolution solution structures of native L99A T4 lysozyme at pH 3.0 obtained from SAXS showed good agreement with the crystal structure solved at neutral pH. This supports the reasonable assumption that the shapes and sizes of the internal cavities in the crystal structure are representative of the solution structure. For the pressure-denatured state of L99A at pH 3.0, SAXS results suggest that the average overall conformation is extended while local regions may have residual structure. The radius of gyration of the pressure-denatured state was smaller than that predicted for the chemically denatured state (39), consistent with the view that the widely accepted model of protein denaturation, i.e. the transfer of core residues to water, does not appropriately describe pressure denaturation. Fluorescence quenching measurements suggest that the pressure-denatured state of L99A at pH 7.0 is even more compact than at pH 3.0.

Unfolding at pH 3.0. The volume change accompanying denaturation is contributed by volume changes of the protein atoms, the solvent-excluded cavities, and the hydration of solvent-exposed residues. As the atoms are the least compressible component, it is thought that the major contributions to the volume change of denaturation are the elimination of solvent-excluded cavities and the hydration of solventexposed residues (32, 48). Around charged residues, electrostriction of water molecules is known to occur (32). However, the volume properties associated with the hydration of hydrophobic residues are still under investigation, particularly with respect to pressure (12, 32, 49, 50). It is therefore difficult to probe the contributions of the hydration changes. By comparing mutants with various volumes of internal cavities, however, we can investigate how these volumes correlate with the volume change of denaturation.

To quantify cavity volumes, structural changes caused by a mutation must be identified, and the solvent occupancy of each cavity must be quantified. In a manner similar to the estimation of a mutation-induced change in stability ( $\Delta\Delta G$ ) by solvent transfer free energies, the change in cavity volume

Table 3: Denaturation Volume Changes Predicted by Side-Chain Volumes  $^{a}$ 

	$\Delta V^{\circ}_{\text{predict}}$ (Å <sup>3</sup> )	$\delta \Delta V (\text{\AA}^3)$
L99G/E108V	$206.2\pm10.0$	$29.6\pm26.9$
A98L	$73.1 \pm 10.0$	$54.1 \pm 12.1$
V149G	$187.1\pm10.0$	$-53.0\pm12.4$

<sup>*a*</sup> The experimental denaturation volume change,  $\Delta V^{\circ}$ , of L99A at pH 3.0 20 mM NaCl was used as a reference.  $\delta \Delta V = \Delta V^{\circ}_{\text{experiment}} - \Delta V^{\circ}_{\text{predict}}$ . Volume changes of other mutants were predicted with van der Waals volumes of side chains (52) of exchanged amino acids.

as a result of a single buried mutation can-as a first approximation-be estimated as the difference between the van der Waals volumes of the exchanged side chains. This analysis has been performed on staphylococcal nuclease and ribonuclease A mutants to predict the mutation-induced change in cavity volumes (8, 51). Particularly in the case of large-to-small amino acid substitutions in staphylococcal nuclease, the cavity volumes predicted by this method successfully correlated with the  $\Delta V^{\circ}$  of pressure denaturation, lending support to the hypothesis that internal hydration of cavities is implicated in pressure denaturation. However, in both studies, atomic resolution structures were not available for most of the mutants to support the assumption that the structural changes were localized at the mutation sites. The same side-chain volume analysis, when applied to T4 lysozyme, fails to predict mutation-induced changes in cavity volumes that correlate with  $\Delta V^{\circ}$ . Using the side chain volumes of Leu, Ala, Val, and Gly reported by Richards et al. (52) and the experimental  $\Delta V^{\circ}$  of L99A at pH 3.0, 50 mM glycine, 20 mM NaCl as a reference,  $\Delta V^{\circ}$  was predicted for L99G/E108V, A98L, and V149G (8). Note that the mutation, E108V, was not included in this analysis because the 108th residue is on the surface of the protein. We therefore do not expect differences in the hydration of Glu (E) and Val (V) to contribute differently to the volume change of denaturation. The predicted results in Table 3 show, in particular, the failure to account for the observed behavior of A98L and V149G. The  $\Delta V^{\circ}$  of V149G was predicted to be as large in magnitude as L99A, while that of A98L was predicted to be very small in magnitude, roughly half of the experimentally observed value.

The failure of this method to estimate mutation-induced changes in cavity volumes is not surprising. Previous crystallographic studies of T4 lysozyme have also shown that structural changes due to a mutation cannot be easily predicted. At some buried sites, a large-to-small amino acid substitution in T4 lysozyme created or enlarged a cavity, while at other sites, a similar mutation caused a rearrangement of the protein that filled the cavity (53). Moreover, for mutations that permitted the creation or enlargement of cavities, the new cavity volume could not be simply predicted by a side-chain analysis using the WT\* structure and the van der Waals volumes of the exchanged amino acids. In the case of L99A, the change in cavity size is slightly smaller than expected. An extreme case was L99G T4 lysozyme. Unlike L99A, the L99G mutation exposes the large cavity to the solvent. A second mutation, E108V, far from the cavity and on the surface of the protein was required to close the opening of the molecule's surface created by the L99G mutation (16). The structural changes caused by a small-tolarge amino acid substitution such as A98L are less predictable (18).

Crystal structures of T4 lysozyme were thus necessary to quantify cavity volumes. The internal cavities of the mutants identified with a 1.2 Å probe are shown in Figure 1 (a)–(e), and the corresponding volumes are presented in Table 2. An inspection of the crystal structures explains the large discrepancies between the experimental  $\Delta V^{\circ}$  and that predicted by side-chain volume analysis for A98L and V149G (Table 3). The A98L mutation does not fill the cavity at the mutation site, but instead, causes the strain-induced formation of several small cavities throughout the protein (Figure 1 (d)). Therefore the total cavity volume is greater than that predicted by side-chain volume analysis. In the case of V149G, the mutation not only enlarges a cavity (Figure 1 (e)) but this cavity accommodates two additional solventbinding sites. Thus, the effective volume of this cavity is likely smaller than that predicted by side-chain volume analysis.

The volume of a single water molecule in bulk is approximately 30  $Å^3$ , which is roughly the size of a small cavity in T4 lysozyme. Therefore, the solvent occupancy of each cavity can have large effects on its effective volume. Cavity 1 (Figure 1, Table 2) is a conserved solvent-binding site in T4 lysozyme (47) and contains Sol 171 and 179. Similarly, cavity 3 binds another conserved internal water molecule, Sol 208. Cavities 4 and 8 contain Sol 175 and 213, respectively. Crystal structures enable the identification of internal water molecules that otherwise may not be identified. However, the average occupancy of internal water is not easily determined by conventional model-based crystallographic refinement methods (54). Sol 171, 175, 179, and 208 have been observed in a large number of mutant structures solved over many independent experiments, and several of these water molecules show evidence of hydrogen bonding stabilization (47). Sol 213 was also observed in all of the mutants studied in this work. We therefore made the reasonable assumption that these water molecules have full occupancy in all of the studied mutants.

The occupancies of poorly ordered water molecules introduced by V149G and L99G/E108V were more difficult to assign. In the crystal structures, the enlarged cavity at the V149G mutation site (Figure 1 (e), cavity 3) contains two additional water molecules (Sol 323 and 324) in addition to Sol 208, while that of L99G/E108V (Figure 1 (c), cavity 6) contains Sol 401 and 402. We first assume that both of these cavities are fully occupied. Under this assumption, however, no correlation is apparent in the relationship between  $\Delta V^{\circ}$ at pH 3.0 with the calculated total cavity volumes (Figure 8 (a)). As discussed earlier, the  $\Delta V^{\circ}$  of V149G predicted by side-chain volume analysis was much larger than the experimentally determined value because solvent binding of the mutation-enlarged cavity was not accounted for. As this mutation site is the location of the only buried polar network in T4 lysozyme (17), we believe that it is reasonable that Sol 323 and 324 occupy this cavity at least partially under the solvent conditions used in this study. This may explain why the tryptophan fluorescence spectrum of V149G in its folded state was the most red-shifted of all the mutants (Figure 2). It is possible that the tryptophan residue lining this polar cavity (Trp 138) reports the increased hydration of V149G.

In contrast, we speculate that the enlarged hydrophobic cavity in L99G/E108V is empty at pH 3.0. The enlarged



FIGURE 8: The volume changes of denaturation (see Table 1) of L99G/E108V, L99A, V149G, and A98L T4 lysozyme in pH 3.0 buffer, 16 °C at 20 mM NaCl (open diamond) and 100 mM NaCl (closed circle) compared to the total cavity volumes calculated from crystal structures using two methods. (a) No correlation is apparent when full occupancy of all crystallographically determined internal solvent-biding sites was assumed in the calculation of cavity volumes. (b) The large cavity of L99G/E108V was assumed to be empty, while full occupancy was assumed for all other internal solvent molecules. The volume changes of denaturation correlate with the total cavity volumes.

cavity of L99A is believed to be empty (1, 15), and the crystal structures of L99G/E108V and L99A are very similar except for the differing volumes of the enlarged cavities (16). At pH 3.0, the magnitude of the volume change upon pressure denaturation was roughly 50 Å<sup>3</sup> greater for L99G/ E108V compared to L99A. This value is similar to the difference in volumes of the enlarged cavities (Figure 1 (b) and (c), cavity 6) in their empty states (61.4  $Å^3$ ), which are the dominant contributors to the total cavity volumes of the two mutants. This suggests that the occupancies of the two cavities are similar at pH 3.0. The total cavity volumes were recalculated with the new assumption that the enlarged cavity in L99G/E108V is empty and that all other crystallographically determined solvent-binding sites are fully occupied. A correlation between denaturation volume changes and total cavity volume is now observed (Figure 8 (b)). The proportionality of the two quantities suggests that all the cavities are hydrated under pressure denaturation at this pH. The slope of the linear fits was dependent on the probe size used to calculate the cavity volumes. Using a 1.4 Å probe gave a smaller slope as fewer small cavities could be detected, reducing the estimated total cavity volume of small cavity mutants by a greater fraction than that of the large cavity mutants.

The magnitude of the volume change of denaturation was also dependent on the ionic strength, reflected by the  $\sim 40$  Å<sup>3</sup> change in vertical positions of linear fits in Figure 8 (b).

We consider two possible mechanisms of hydration changes around solvent-exposed residues to explain this ionic strength dependence. At pH 3.0, T4 lysozyme is highly positively charged. Most of the charged residues are on the surface of T4 lysozyme, and no change in the hydration of these residues is expected from denaturation. However, there are several buried and semiburied salt bridges in T4 lysozyme. Crystal structures of T4 lysozyme solved at various ionic strengths demonstrate that a change in ionic strength does not affect buried salt bridges in the native state (55). These salt bridges are likely exposed in the pressure-denatured state, and the presence of counterions around the dissociated salt pairs would reduce the effect of electrostriction on the denaturation volume change. The presence of counterions may also affect the compactness of the denatured state, and thus change the number of solvent-exposed residues. Refolding of an acid-denatured protein has been observed at high anion concentrations (56). This is thought to be due to reduced charge repulsions in the protein. A similar behavior can be expected to occur to T4 lysozyme at pH 3.0. This may be the basis for the apparent blue-shift in the tryptophan fluorescence of the denatured state with increased NaCl concentration. Regardless of mechanism, our data indicate that under the conditions investigated at low pH, all of the internal cavities are hydrated while the ionic strength of the solvent affects the volume contribution of hydration around solvent-exposed residues.

Cavity Filling at pH 7.0. Fluorescence measurements were made at pH 7.0, 24 °C as a direct comparison to the highpressure, room-temperature crystal structures of L99A and WT\* T4 lysozyme (1). According to the high-pressure crystal structures acquired by Collins et al., cooperative water filling of the large hydrophobic cavity in L99A (Figure 1 (b), cavity 6) is favorable under pressure even without unfolding (1). We emphasize that, in the crystal state, large conformational changes such as unfolding do not occur. Only the large cavity of L99A showed increased internal electron density with pressure (PDB structures 2B6T, 2B6W, 2B6X, 2B6Y, 2B6Z, 2B70, 2B72, 2B73, 2B74, 2B75). No other pressure-induced changes in internal hydration were observed in the crystal structures. WT\*, which has a smaller corresponding cavity (Figure 1 (a), cavity 6), showed no water filling up to 200 MPa (PDB structures 20E4, 20E7 are at 100 MPa; 20E9, 20EA at 200 MPa). Molecular dynamics simulations performed on the high-pressure structures of L99A suggested cooperative filling of four water molecules in the large cavity (1).

In solution, L99A underwent a pressure-induced change in fluorescence corresponding to a denaturation volume change of  $\Delta V^{\circ} = -93.4 \pm 8.2$  Å<sup>3</sup>, nearly a factor of 2 smaller than at pH 3.0, with a pressure midpoint of roughly 240 MPa. Structural information obtained from comparative fluorescence quenching of L99A and Se-Met L99A indicated that the pressure-denatured state of L99A at pH 7.0 is even more compact than that at pH 3.0. While our methods do not provide direct evidence of preferential water filling of the large cavity in L99A in solution at this pH, we explored this is a possible mechanism to explain the small volume change and compactness of the denatured state. Assuming that the volume change of transferring each water molecule from the bulk to the large cavity is 30 Å<sup>3</sup>, this mechanism would suggest that in solution, the large cavity is filled by



FIGURE 9: Comparison of change in water occupancy of L99A at neutral pH determined by X-ray crystallography (diamond; reprinted with permission from ref *I*. Copyright 2005 The National Academy of Sciences of the USA) and fluorescence spectroscopy (circle). A two-state (0 or 3 water molecules) thermodynamic fit to the data is shown on each data set with a fixed volume change of 90 Å<sup>3</sup> (volume of three water molecules in bulk). The fit to the crystallography data is shifted to lower pressure by 82 MPa relative to the fluorescence data. This stability difference may be due to subtle differences between structure and dynamics of the T4 lysozyme molecule in solution and in the crystal. The change in water occupancy of WT\* determined by fluorescence is shown for reference (cross).

three water molecules (93.4 Å<sup>3</sup> ~ 3 × 30 Å<sup>3</sup>). This is in reasonable agreement with the predictions made by the molecular dynamics simulations (1). For comparison with the change in water occupancy of the large cavity determined experimentally by integrated electron densities in the crystal structures (1), the centers of spectral mass of L99A were converted to change in water occupancy under the assumption that under pressure, the large cavity is filled by three water molecules (Figure 9). A two-state thermodynamic fit to the converted fluorescence data with a fixed volume change of three bulk water molecules shows good agreement with the crystallographic results. This suggests that preferential water filling of the large cavity is a plausible mechanism to explain the small volume change and size of the denatured state at this pH.

As a comparison to L99A, fluorescence measurements were also made on the structurally similar mutant, L99G/ E108V. Although the large cavity of L99G/E108V is roughly 61 Å<sup>3</sup> greater in volume than the corresponding cavity in L99A, the denaturation volume changes were the same to within error (approximately 10 Å<sup>3</sup>). As mentioned earlier, the crystal structure of L99G/E108V shows two poorly ordered water molecules in the large cavity (Sol 401 and 402). We interpret this to mean that at neutral pH, where the crystal structure was solved, the large cavity of L99G/E108V is partially filled with a maximum occupancy of two water molecules. By taking two water molecules into consideration, the difference in volume changes of cavity-filling for L99A and L99G/E108V can be as small as 1 Å<sup>3</sup> (61 Å<sup>3</sup> ~ 2 × 30 Å<sup>3</sup>).

One of the paradoxes of pressure denaturation has been the small magnitudes of observed volume changes. It has been thought that a positive volume contribution must exist, possibly due to the hydration of hydrophobic residues (32). While this is plausible, it cannot be assumed that all the existing cavities and packing defects of a protein are hydrated or eliminated with the application of pressure. The highpressure crystallography study of L99A showed that water penetration of the large cavity and not other, smaller cavities in the native structure of L99A was possible in part due to the cooperativity (1). Only the hydrophobic cavity large enough to accommodate multiple waters interacting with hydrogen bonds was filled in the folded structure.

## CONCLUSIONS

Our results imply the existence of multiple pressuredenatured states of the T4 lysozyme family with differing levels of internal hydration and unfolding that were dependent on the solvent conditions and particularly pH. The results were thus consistent with water penetration of the protein rather than transfer of hydrophobic residues from the core of the protein to the water as the mechanism of pressure denaturation. At pH 3.0, the pressure-denatured state was extended but more compact than the predicted size of a chemically denatured protein of the same length (39). We were able to relate the volume change of denaturation to cavity volume and hydration changes by comparing multiple mutants of T4 lysozyme with known crystal structures and varying the ionic strength of the solvent. The magnitudes of the volume changes of denaturation for L99A, L99G/E108V, A98L, and V149G T4 lysozyme positively correlated with the total cavity volume. These results suggest that at pH 3.0, all the cavities were hydrated in the pressure-denatured state. Increasing the ionic strength reduced the magnitude of the volume change likely as a result of counterion interactions with salt pairs exposed in the denatured state. At pH 7.0, the denaturation thermodynamics of L99A in solution was in good agreement with the preferential cavity filling observed by high-pressure crystallography and MD simulations (1). The pressure-denatured state at pH 7.0 was more compact in size than at pH 3.0. These results suggest that at pH 7.0, not all the cavities are filled. We speculate several mechanisms for the pH-dependence on the hydration of the denatured state. pH may affect the polarity of internal cavities. It may also affect the dynamics of the protein and therefore affect the water-penetration pathways. Finally, it is also possible that the charge state of the protein affects the compactness of the denatured state, which in turn may lead to preferential cavity filling as is the case for folded structures (1).

We showed that the problem of pressure denaturation requires understanding of the change not only in water occupancy as a function of pressure but also in the volume properties of the protein and interacting waters. The availability of crystal structures was necessary for the quantification of cavity volumes and solvent occupancy of the T4 lysozyme mutants investigated in our work. However, we recognize that a crystal structure represents a static picture of the average conformation of a protein under a specific condition. As a protein is a fluctuating system, the cavities and internal solvent molecules are not static. The calculation of a cavity volume from an atomic structure also depends on the probe size and the van der Waals radii of the amino acids used for identifying molecular surfaces. For the T4 lysozyme mutants used in this study, a 1.2 Å probe was preferred over a 1.4 Å probe. Small cavities that were found in all mutants by a 1.2 Å probe were not consistently identified with a 1.4 Å probe. The magnitudes of the

experimentally observed volume changes of denaturation were roughly the size of several water molecules. The interpretation of volume changes from native structures, therefore, requires certainty in the occupancy of internal water molecules in the native state and its dependence on the solvent condition.

In this work, we investigated the contribution of cavity volumes and electrostriction to the volume change of pressure denaturation. The volume properties of hydration waters, particularly around hydrophobic residues as a function of pressure, remain to be established. Finally, it was assumed in this study that the volume change accompanying denaturation was pressure-independent because the magnitude and sign of isothermal compressibility change with denaturation, i.e. the pressure dependence of the denaturation volume change, are still under debate. Direct measurements of the isothermal compressibility change associated with pressure denaturation have not yet been made (57). With the exception of the study by Seemann et al. (33), the isothermal compressibility change has only been determined from fitting a thermodynamic model with a second-order expansion of  $\Delta G$  to denaturation data (57). It is noted that the sign of the isothermal compressibility is defined inconsistently in the literature, causing confusion on the sign of the second-order term of  $\Delta G$  (32, 48, 57–60). A second-order fit also increases the number of free fitting parameters such that overfitting of data is also a concern. Further investigations of the pressure-dependence of the volume change accompanying denaturation are suggested.

The magnitude of the volume change accompanying pressure denaturation of a protein is generally less than 1% of the protein volume and on the order of several bulk water molecules in volume. Our results support the growing view that the pressure-denatured state of a protein is one in which the penetration by a few water molecules is favored and emphasizes the importance of the role of water in protein folding and biological processes.

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#### SUPPORTING INFORMATION AVAILABLE

Additional experimental details. This material is available free of charge via the Internet at http://pubs.acs.org.

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# Supporting Online Material

# Supporting Material 1: Composition of Cell Culture Broths

Proteins used in this work were expressed in bacteria grown in media similar in composition, but not identical to standard Luria-Bertani broth or M9a media. The media referred to as modified Luria-Bertani broth contains 12 g tryptone, 10 g NaCl, 1 g glucose and 5 g yeast extract per liter of broth. LBH broth contains 10 g tryptone, 5 g NaCl, 1 ml of 1 N NaOH and 5 g yeast extract per liter of broth. The version of M9a media used for these experiments contains 7 g Na<sub>2</sub>HPO<sub>4</sub>· 7H<sub>2</sub>O, 3 g KH<sub>2</sub>PO<sub>4</sub>, 1 g NH<sub>4</sub>Cl, 0.5 g NaCl mixed with one liter of water and autoclaved, followed by sterile addition of 10 mL 20% glucose, 0.4 mL 0.25 M CaCl<sub>2</sub>, 1 mL 1 M MgSO<sub>4</sub>, and 2 mL 0.5 mg/mL thiamine.

## Supporting Material 2: Production of L99A, L99G/E108V, and A98L mutants

The L99A, L99G/E108V, and A98L mutants were expressed and purified by a modified version of the protocol described by Poteete, *et al.* (1). For clarity, the entire procedure is listed here. *E. coli* strain RR1 containing lysozyme-producing plasmids were streaked from frozen cultures onto modified LB-agar plates containing 100  $\mu$ g/mL ampicillin. Single colonies were used to inoculate 100 mL cultures of LBH broth (Supporting material 1) containing 200  $\mu$ g/mL ampicillin, and were incubated overnight at 32° C. The 100 mL culture was diluted into four 1 L cultures of LBH broth in Fernbach flasks with aeration. Protein expression was induced by an IPTG concentration of 180 mg/L at an optical density of 0.6. After 90 minutes the cultures were pelleted and resuspended in 100 mL of 50 mM Tris-HCl, pH 7.5, 10 mM Na<sub>3</sub>HEDTA, 0.1% triton X-100 buffer with 1 protease inhibitor tablet (Complete Mini, EDTA-free Protease Inhibitor Cocktail Tablets,

Roche Applied Science, Indianapolis, IN, USA). The viscosity due to the bacterial genomic DNA was reduced by DNase I treatment, sonication or the use of a French press. The products from the DNase I treatment were removed by dialysis (Spectra/Por 10 kDa MWCO, Spectrum Laboratories, Inc., Rancho Dominguez, CA, USA). The dialyzed solution was then loaded onto a 2.5 x 5 cm CM-sepharose column, pre-equilibrated with 50 mM Tris-HCl, 1 mM sodium EDTA, pH 7.25 and gradient eluted using a 0 to 0.3 M NaCl gradient in the same buffer. Proteins were dialyzed into 0.025 M NaPO<sub>4</sub>, pH 5.8, then concentrated using SP sephadex and stepped off with 0.55 M NaCl, 0.1 M NaPO<sub>4</sub>, 1 mM sodium EDTA, 0.01% sodium azide, pH 6.6, and stored in the same buffer (*1*).

# Supporting Material 3: Production of Seleno-methionine Containing Mutants

The V149G mutant was isolated using a modified version of the inclusion body protocol described by Vetter *et al.* (2). Protein expression was induced at 37 °C. Following expression, the culture was pelleted by centrifugation at 4,700 x g. The bacterial pellet was resuspended in a buffer composed of 50 mM Tris-HCl, 10 mM sodium EDTA, pH 8.0 plus 1 protease inhibitor tablet (Roche) and then sonicated. The resuspended mixture was then centrifuged for 30 minutes at 27,200 x g to pellet the cellular debris. The pellet was resuspended in 50 mM Tris-HCl, 10 mM sodium EDTA, 50 mM NaCl, 1 mM phenyl methane sulfonyl fluoride, 2.5 mM benzamide, 0.1 mM DTT, pH 8.0 (2). One tenth of the volume of the resuspension buffer of 2% Triton X-100 in 50 mM Tris, 10 mM sodium EDTA, pH 8.0 was added to the resuspended bacterial pellet solution (2). The suspension was stirred overnight at 4 °C, and then centrifuged at 27,200 x g for 30 minutes. Ten volumes of 2.5% octyl- $\beta$ -D-glucopyranoside in 50 mM Tris, 10 mM sodium EDTA, pH 8.0, per apparent volume of pellet were added and the suspension was stirred at 4 °C for 4 hours. Following centrifugation at 27,200 x g for 30 minutes, the pellet was washed with double-deionized water. The pellet was then suspended in freshly made 4 M urea (unbuffered, pH 6 to 7). The pH of the urea solution was lowered to 3-3.5 by the addition of 10 mM glycine, 2 N phosphoric acid, then mixed well for several minutes, and centrifuged at 12,000 x g for 15 minutes. The supernatant, containing the protein, was dialyzed against a 50 mM citrate, 10% glycerol, pH 3.0 buffer at 4 °C overnight, and then dialyzed against a 50 mM citric acid buffer, 10% glycerol, pH adjusted to 5.5 using NaOH at 4 °C for 8-16 hours. The solution was centrifuged at 12,000 x g for 25 minutes. The procedure yields at least 40 mg of protein, estimated to be approximately 99 % pure.

## Supporting Material 4: Production of Seleno-methionine Containing Mutants

A seleno-methionine containing variant of the L99A mutant (Se-Met L99A) was prepared in a methionine deficient, seleno-methionine rich growth medium using an adaptation of the procedure by Van Duyne *et al.* (*3*). Frozen stocks of *E. coli* strain RR1 (*4*) containing an L99A expressing plasmid were streaked onto modified LB-ampicillin agar plates and grown overnight. Single colonies were used to inoculate 200 mL modified LB cultures containing 200 µg/mL ampicillin. The 200 mL cultures were grown for approximately 8 hours at 37 °C. Each 200 mL culture was used to inoculate a 1.25 L culture of M9a media (7 g Na<sub>2</sub>HPO<sub>4</sub>· 7H<sub>2</sub>O, 3 g KH<sub>2</sub>PO<sub>4</sub>, 1 g NH<sub>4</sub>Cl, 0.5 g NaCl mixed with one liter of water and autoclaved, followed by sterile addition of 10 mL 20% glucose, 0.4 mL 0.25 M CaCl<sub>2</sub>, 1 mL 1 M MgSO<sub>4</sub>, and 2 mL 0.5 mg/mL thiamine). Each 1.25 L culture was shaken at 250 rpm at 37 °C overnight. The next day, each 1.25 L culture was transferred to a 4 °C cold room for 30 minutes. Methionine biosynthesis was halted by adding 50 mL of M9a media containing an amino acid mixture of 100 mg lysine hydrochloride, 100 mg threonine, 100 mg phenylalanine, 50 mg leucine, 50 mg isoleucine and 50 mg valine to each 1.25 L culture. The 1.3 L cultures were shaken at 30 °C for approximately 30 minutes at 150 rpm. Next 50 mg/L of L-(+)-selenomethionine (Anatrace, Maunee, OH) was added and protein expression was induced by addition of 45 mg of IPTG per 1.3 L culture. Induction was allowed to proceed for 4 hours.

Following protein expression the cell cultures were pelleted at 5,000 x g for 10 minutes. The cell pellet was resuspended in 50 mM NaCl, 50 mM Tris pH 7.5 buffer with 15 mM methionine to prevent oxidation, and lysed by sonication for 7 minutes. The cellular debris was pelleted by centrifugation at 17,000 x g for 20 minutes. The supernatant was loaded onto a column containing a 2.5 cm bed of CM sepharose. This procedure yields approximately 70 mg of Se-Met L99A per 4 L of culture.

# Supporting References

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