

CORRELATIONS BETWEEN INTERNAL MOBILITY AND STABILITY OF GLOBULAR PROTEINS

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ABSTRACT The recent work is surveyed which leads to the suggestions that the conformation of globular proteins in solution corresponds to a dynamic ensemble of rapidly interconverting spatial structures, that clusters of hydrophobic amino acid side chains have an important role in the architecture of protein molecules, and that mechanistic aspects of protein denaturation can be correlated with internal mobility seen in the native conformation. These conclusions resulted originally from high resolution ^1H nuclear magnetic resonance (NMR) studies of aromatic ring mobility, exchange of interior amide protons and thermal denaturation of the basic pancreatic trypsin inhibitor and a group of related proteins. Various new approaches to further characterize proteins in solution have now been taken and preliminary data are presented. These include computer graphics to outline hydrophobic clusters in globular protein structures, high resolution ^1H -NMR experiments at variable hydrostatic pressure and ^{13}C -NMR relaxation measurements. At the present early stage of these new investigations it appears that the hydrophobic cluster model for globular proteins is compatible with the data obtained.

INTRODUCTION

In a series of recent papers we described high resolution proton nuclear magnetic resonance (NMR) studies of the spatial structures in solution of a group of small globular proteins related to the basic pancreatic trypsin inhibitor (BPTI) either by chemical modification (1–4) or by homology (5, 6). For all the proteins considered, the average solution conformation coincided closely with the molecular structure of BPTI seen in single crystals (7). Thus we had a quite unique system for investigating the influence of local modifications of the covalent structure on the dynamic properties and the stability of a given spatial protein structure. The strategy used in these investigations is illustrated in Fig. 1. The NMR studies concentrated mainly on measurements of amide proton exchange rates, mobility of the aromatic rings and protein denaturation (8–13). Presently these earlier data are being complemented by various different experiments to investigate additional aspects of protein conformation in solution. This includes high resolution ^1H -NMR experiments at high hydrostatic pressure, computer graphics for detailed inspection of protein structures and measurements of nuclear spin relaxation times. Preliminary data obtained from these different approaches are discussed in the light of the previously suggested multi-state cluster description of globular proteins (9–11).

METHODS

^1H -NMR spectra at 360 MHz and ^{13}C -NMR spectra at 90.5 MHz were recorded on a Bruker HX 360 spectrometer. ^{13}C -NMR spectra at 25.1 MHz were obtained with a Varian XL-100 spectrometer. Experimental details on the sample preparation and the NMR techniques used were presented previously (1–6, 12, 13). High resolution 360 ^1H -NMR spectra at variable hydrostatic pressure were obtained with a home-built experimental device using a thick-walled glass capillary as a sample cell. A

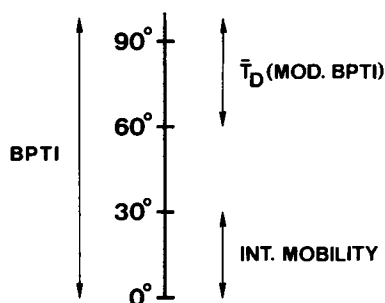


Figure 1 Strategy for the investigations of BPTI-related proteins. In aqueous solution the globular conformation of BPTI is stable over the temperature range from 0° to >95°C. Different, strictly localized modifications of the covalent structure yielded proteins with average denaturation temperature \bar{T}_D between 60° and 95°. It was then investigated whether the variations of T_D could be correlated with the internal flexibility of the globular protein structures manifested e.g. in the exchange of interior labile protons and in intramolecular rotational motions of the aromatic rings at temperatures far below \bar{T}_D .

detailed description of the high pressure apparatus, which was based on principles outlined by Yamada (14) and Volkl et al. (15), will be given elsewhere.¹

For the computer graphics studies we made use of the "Zentrum für Interaktives Rechnen" of the ETH. A DEC-10 computer was used with the program XRAY² to produce molecular models from a protein structure data bank. A PDP-11 computer with Evans and Sutherland picture system PS 2 was used with the program PROT² to orient the protein molecules to get the most informative views.

RESULTS

Amide Proton Exchange, Aromatic Ring Mobility, and Thermal Stability of Native and Chemically Modified BPTI

The structures of BPTI and a group of related proteins obtained by strictly localized chemical modifications are described in Fig. 2. Data on internal mobility and stability of these proteins are presented in Table I. Since the spin systems of all the eight aromatic rings (1–6) and the resonances of numerous amide protons (16, 17) were individually assigned, the data on internal rate processes could be correlated with specific locations in the protein structure. The following are some key observations made in these experiments.

The exchange rates of interior labile protons are correlated with the thermal stability of the proteins, i.e., at a given temperature the exchange rates were higher in the less stable proteins (Table I). In contrast, the mobility of the aromatic rings was not noticeably affected by localized chemical modifications (Fig. 2), unless the modification involved the immediate ring environment.

Different amide protons in the BPTI-related proteins exchanged with different rates (12). Overall, the protons of the β -sheet exchanged more slowly than those of the α -helix. Within the β -sheet the exchange was faster at both ends than in the central region. Similarly, different rates of the flipping motions about the C^β - C^γ bond were observed for the individual aromatic rings (18).

As a consequence of the chemical modifications in Fig. 2, the amide proton exchange rates were increased by several orders of magnitude (Table I). In contrast to these large differences

¹G. Wagner, manuscript submitted for publication.

²The programs XRAY and PROT were adapted from programs which were kindly provided to us by Richard M. Feldmann, National Institutes of Health, Bethesda, Md. 20205.

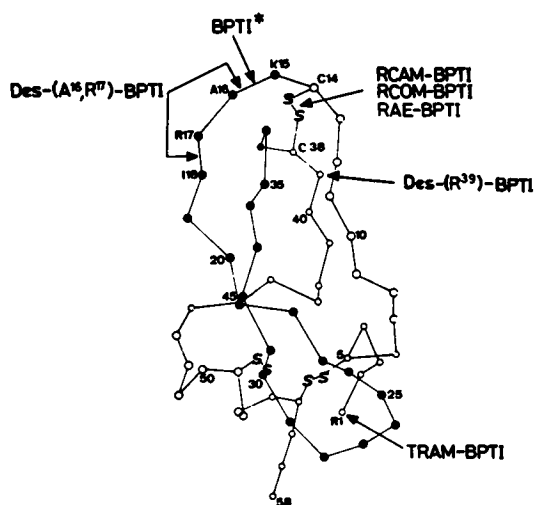


Figure 2 Projection of the α -carbon positions in the BPTI molecule obtained from a single crystal x-ray analysis (7). The structure contains three disulfide bonds, a twisted antiparallel β -sheet which extends through the entire length of the molecule, and a short α -helix near the C-terminus. The disulfide bonds are included in the drawing and the residues in the antiparallel β -sheet are identified by filled circles. The arrows indicate the locations of the chemical modifications discussed in this paper. Amino acid residues in the modification sites are identified by the IUPAC one-letter symbol and the position in the amino acid sequence. RCAM-BPTI, RCOM-BPTI and RAE-BPTI were obtained by reduction of the disulfide bond 14–38, with the cysteinyl residues protected by carboxamidomethylation, carboxymethylation or aminocethylation, respectively: BPTI* was obtained by cleavage of the peptide bond Lys 15—Ala 16, Des (A¹⁶, R¹⁷)-BPTI by cleavage of the peptide bond Lys 15—Ala 16 and removal of Ala 16 and Arg 17, TRAM-BPTI by transamination of the α -amino group of Arg 1, and Des-(R³⁹)-BPTI by cleavage of the peptide bond Arg 39—Ala 40 and removal of Arg 39. BPTI contains eight aromatic residues, i.e. four phenylalanines in the positions 4, 22, 33, and 45, and four tyrosines in the positions 10, 21, 23, and 35.

between the absolute rates, the rank order of the exchange of individual protons was with few exceptions identical in the different proteins. Thus, even though most of the chemical modifications were in peripheral locations in the molecular structure, e.g., at the *N*-terminus or at the reactive-site peptide bond, they caused an increase of the exchange rates throughout the entire protein. For each individual one of the interior amide protons, the exchange followed EX₂ kinetics (19) over the entire p²H range studied (12).

TABLE I
THERMAL STABILITY AND INTERNAL MOBILITY OF BPTI-RELATED
PROTEINS IN AQUEOUS SOLUTION

Protein	\bar{T}_D [°C] [p ² H 5.0]	k^{NH} [min ⁻¹] [p ² H 4.5, 36°C]	Phe-45 [p ² H 7.8, 4°C]	
			ν [s ⁻¹]	ΔG^\ddagger [kJ mol ⁻¹]
BPTI	>95	2×10^{-7}	30	60
TRAM-BPTI	[~95]	[$>2 \times 10^{-7}$]	30	60
BPTI*	85	5×10^{-5}	60	58
Des-(R ³⁹)-BPTI	85	3×10^{-4}	30	60
RCOM-BPTI	79	4×10^{-4}	30	60
RCAM-BPTI	76	1×10^{-4}	30	60
RAE-BPTI	70	6×10^{-4}	30	60
DES-(A ¹⁶ , R ¹⁷)-BPTI	>65	7×10^{-5}	60	58

The table lists the average denaturation temperature \bar{T}_D , the exchange rate constant, k^{NH} , for the most slowly exchanging amide proton, and the frequency ν and activation energy ΔG^\ddagger of the 180° flips of the Phe 45 ring about the C⁶-C' bond.

When different separated resonance lines were observed through the thermal denaturation, differences of the order of up to 4° were found between the apparent denaturation temperatures for individual protons (20).

Since these resonances were previously individually assigned (21) the different denaturation behavior could be correlated with specific regions of the molecule. A general trend was that the α -protons of the polypeptide backbone showed lower apparent denaturation temperatures than the protons of hydrophobic side chains.³

High Resolution ¹H-NMR at High Hydrostatic Pressure

Since reaction volumes, ΔV , in chemical equilibria with equilibrium constant K are given by

$$\Delta V = -RT \frac{d \ln K}{dp}$$

and activation volumes, ΔV^\ddagger , in rate processes with rate constant k by

$$\Delta V^\ddagger = -RT \frac{d \ln k}{dp},$$

investigations at variable hydrostatic pressure, p , can provide essential data for the characterization of thermodynamic and mechanistic aspects of protein conformation in solution. We have started measurements of the dependence of the parameters of Table I on hydrostatic pressure in the range from 1 to 2,500 atm.

At present quantitative data were obtained for the rotational motions of the aromatic rings of Tyr 35 and Phe 45 in BPTI. The activation volumes, ΔV^\ddagger , for 180° flips of these two rings about the C ^{β} - C ^{γ} bond (8, 22) were found to be $63 \pm 20 \text{ \AA}^3$ and $50 \pm 10 \text{ \AA}^3$,³ respectively.¹ Positive activation volumes for rotational motions of interior aromatic rings in globular proteins appear not to be unique for BPTI, since similar results were obtained for three aromatic rings in ferricytochrome c.¹

¹³C-NMR Relaxation Studies

To complement the results on the internal mobility of BPTI-related proteins in Table I by measurements on a different time scale, the ¹³C-relaxation times T_1 and T_2 and the ¹³C [¹H]NOE (8) were studied at 25.1 MHz and 90.5 MHz (23).⁴

The main emphasis was on studies of backbone α -carbons, aromatic ring carbons, and methyl carbons of aliphatic amino acid side chains, which had previously been individually assigned (23, 24). An analysis of these data in terms of a "wobbling-in cone" model (23, 25)⁴ indicated that, in addition to the rapid methyl rotation, the methyl carbon relaxation times manifest librational motions of the side chains with correlation times of the order of 10⁻⁹ s and with angular displacements of 20°–50°.

Here we present an empirical comparison of relaxation data for BPTI and RCAM-BPTI. Table II shows that within the accuracy of these experiments no significant differences between the T_1 values of corresponding carbons in the two proteins were observed. Hence

³The same trend was since observed in several snake neurotoxins and cardiotoxins. In certain toxins the differences between the apparent denaturation temperatures for backbone α -protons and aromatic ring protons is considerably larger than 5°C. A. Chrzesczcyk, Ch. Moonen, J. Lauterwein, M. Lasdunski, W. Steinmetz, L. Visser, G. Wider, and K. Wüthrich, unpublished data.

⁴Richarz, R., K. Nagayama, K. and K. Wüthrich. Manuscript submitted for publication.

TABLE II
RELAXATION TIMES T_1 (msec) OF SELECTED ^{13}C -RESONANCE LINES IN BPTI
AND RCAM-BPTI AT 25.1 MHz, $\text{p}^2\text{H} = 4.1$, $T = 39^\circ\text{C}$.

Resonance Assignments	T_1	
	BPTI	RCAM-BPTI
α -carbons (average)	45 ± 3	48 ± 3
Tyr ϵ (average)	46 ± 3	45 ± 5
Ala 16 β	160 ± 20	180 ± 30
{ Ala 40 β		—
{ Ala 27 β		160 ± 20
Ala 48 β	150 ± 20	160 ± 20
Ala 58 β	~ 300	300 ± 30
Ile 18 γ^2	215 ± 30	170 ± 20
Ile 19 γ^2	215 ± 30	200 ± 20
{ Ile 18 δ	385 ± 30	360 ± 40
{ Ile 19 δ		
Met 52 ϵ	325 ± 30	330 ± 30

The relaxation times were measured with the inversion recovery method (26) in $2.5 \cdot 10^{-2}$ M solutions of the protein in $^2\text{H}_2\text{O}$.

there was no apparent correlation between internal segmental motions on the nanosecond time scale and the thermal stability of the globular proteins (Table I).

DISCUSSION

Multi-State Hydrophobic Cluster Model for Globular Proteins

Fig. 3 shows a scheme of fundamental structural aspects of a dynamic multistate model for globular proteins which was suggested on the basis of the data in Table I and additional observations (9–11, 27). The fundamental point in Fig. 3 is that the protein molecule consists of hydrophobic clusters which are loosely connected by covalent bonds and held in fixed

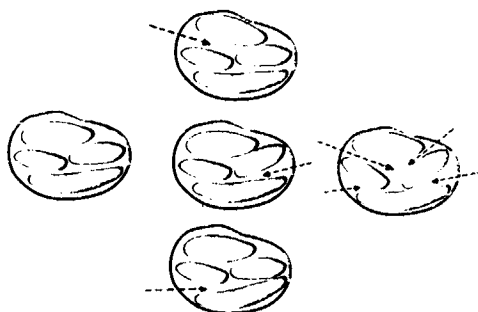


Figure 3 Schematic two-dimensional representation of a dynamic multistate model for globular proteins. Clusters formed by hydrophobic side chains are the pillars of the molecular structure. The individual clusters are loosely linked together by the polypeptide backbone and interact primarily via polar groups located on their surfaces, e.g., by hydrogen bonding. In the compact average structure on the left, interior amide protons in the interfaces between different clusters are shielded from the solvent. The three species in the center represent distorted structures contained in the molecular conformation at temperatures far below the denaturation point. Through variations of the relative spatial orientations of the intact clusters, individual interior amide protons are exposed to the solvent, as indicated by the arrows. The drawing on the right represents a more strongly distorted open structure which would be characteristic for the species prevailing near denaturing conditions.

spatial orientations mainly by interactions between polar groups on the cluster surfaces, i.e., primarily hydrogen bonded secondary structures. Transitions between the molecular structures contained in the protein conformation are by two different types of intramolecular fluctuations which promote, respectively, exchange of interior amide protons or rotational motions of the aromatic rings: (a) Translational and rotational motions of the intact hydrophobic clusters relative to each other primarily expose labile protons on the interior surface of the clusters to the solvent, thus enabling exchange to take place. These are "global" fluctuations of the protein structure in the sense that they are related to the thermal stability and that a localized variation of the covalent structure affects nearly identically the exchange rates of all amide protons throughout the protein (13). (b) Structure fluctuations within the hydrophobic clusters determine the mobility of the aromatic rings, since these are located in the interior of the clusters. The experiments showed that the fluctuations manifested in the ring flips were strictly localized, i.e. not correlated with the stability of the protein and not susceptible to long range effects from chemical modifications (9–11). The observations of different apparent denaturation temperatures for backbone fragments on the surface of the clusters and for different hydrophobic amino acid side chains located in different clusters could be explained with the assumption that cooperative unfolding in the structure of Fig. 3 extends only over the individual clusters. In a first phase of the process of unfolding the regions of contact between the different clusters would be altered, which would be followed by cooperative decay of individual clusters (20). The new experiments described in this paper were designed to provide additional criteria for distinguishing between different types of internal fluctuations of protein structures and for characterization of the hydrophobic clusters in Fig. 3.

Measurements of the activation volumes, ΔV^\ddagger , for the aromatic ring flips and the amide proton exchange appear to provide additional evidence that these two rate processes are correlated with different types of internal fluctuations of the protein structure. While we measured positive ΔV^\ddagger s for the ring flips, Carter et al. (28) reported small positive and negative ΔV^\ddagger for proton exchange in aqueous solutions of different proteins. ^1H -NMR measurements of ΔV^\ddagger for the exchange of individual amide protons in BPTI, where the same pressure range will be covered as for the studies of the ring flips,¹ are in progress in our laboratory. A reliable check of the above conclusion should thus soon be available.

Measurements of ΔV^\ddagger are also a potential source of information on the mechanism of the aromatic ring flips (22, 29). Positive values for ΔV^\ddagger show that the ring flips occur when the protein adopts a larger than average volume. It is then interesting that the ΔV^\ddagger s measured for Tyr 35 and Phe 45 correspond approximately to the volume of the atoms near the rings which stick into the sphere occupied by the rotating ring (22). This may indicate that the "viscosity" of the ring environment, as defined in (29), does not have a dominant influence during the actual process of ring rotation.

For the analysis of the ^{13}C -relaxation data in Table II it is essential that previous studies showed the average solution structures of BPTI and RCAM-BPTI to be closely similar. Hence the correlation times for overall rotational motions should be nearly identical and different relaxation times would thus manifest different librational mobility for corresponding groups in the two proteins. Similar to the aromatic ring flips (Table I) the librational motions of the backbone and the aliphatic amino acid side chains thus appear to manifest internal fluctuations of the protein structure which are not correlated with the thermal stability of the globular structure. Interestingly most of the methyl carbon resonances in Table II correspond to amino acid side chains located near the aromatic rings in the hydrophobic clusters.

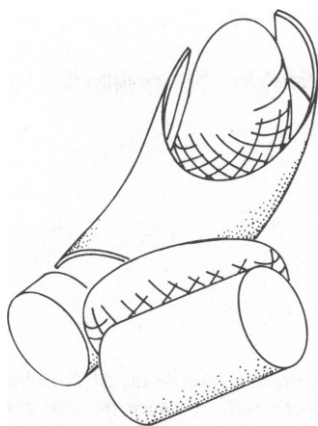


Figure 4 Schematic presentation of the hydrophobic cluster structure of BPTI. Two hydrophobic clusters, which are characterized in more detail in Fig. 5, a hydrophilic layer, which is described in detail in Fig. 6, and two helices are distinguished. A hydrophobic cluster which contains the side chains of Phe 23, Phe 33, and Tyr 35 is surrounded by the hydrophilic layer. The second cluster between the hydrophilic layer and the C-terminal helix includes the side chains of Phe 4, Tyr 21, Tyr 23, and Phe 45. The short *N*-terminal helix has direct H-bond connections to the hydrophilic layer.

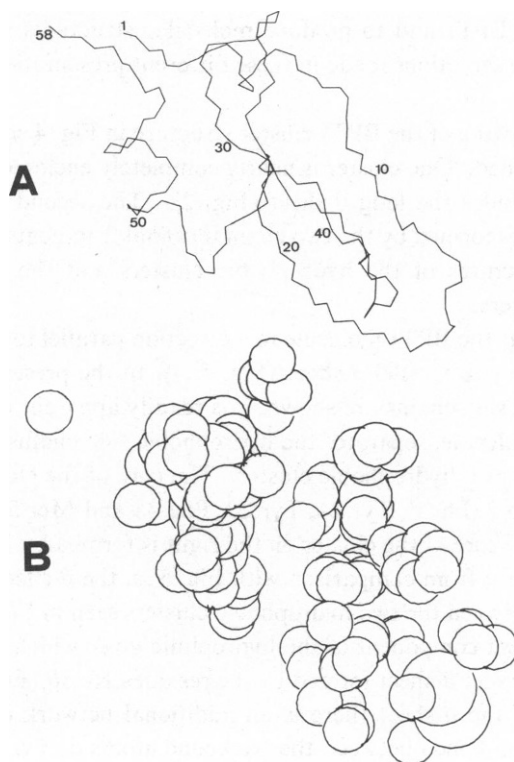


Figure 5 Computer drawing of hydrophobic clusters in BPTI. The orientation of the molecule resulted from rotations by 40°, 22°, and 100°, respectively, about the fixed *x*-, *y*-, and *z*-axes used by Deisenhofer and Steigeman (7). Trace *A* shows a line drawing of the polypeptide backbone from this point of view. *B* presents a space filling drawing of the side chains of Ala, Val, Leu, Ile, Pro, Met, Cys, Phe, and Tyr. No backbone atoms are shown except for the α -carbons and the amide nitrogens of Pro. Atomic radii of 1.5, 1.65, and 2.5 Å were used for C, N, and S, hydrogen and oxygen atoms are not shown.

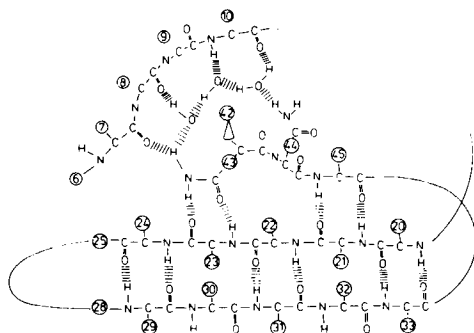


Figure 6 Planar projection of the hydrophilic layer in the BPTI molecule (Fig. 4). This layer includes the triple-stranded β -sheet and hydrogen bonds between the side chains of Asn 43 and Asn 44, the backbone fragment 7-10 and three internal water molecules (7). The α -carbons are identified by the position of the residues in the amino acid sequence (5-7). The α -protons are not shown. Residues 8 and 9 are prolines. Only the central region of the β -sheet is shown (see Fig. 2). The peptide bond 41-42 points out of the plane of projection. The missing protons of the water molecules are located above the projection plane.

Hydrophobic Clusters in the Crystal Structure of BPTI

The atomic coordinates of Deisenhofer and Steigemann (7) were used to construct a "Labquip" model of BPTI and to produce molecular structures with a computer graphics system. The major observations made in these different presentations of BPTI are illustrated in Figs. 4-6.

The schematic drawing of the BPTI cluster structure in Fig. 4 shows that two hydrophobic clusters can be discerned. One cluster is nearly completely enclosed by a layer of hydrophilic fragments which includes the long β -sheet (Fig. 2). The second cluster is enclosed by the β -sheet and the helices formed by the two chain terminal fragments. Figs. 5 and 6 describe in more detail the structures of the hydrophobic clusters and the hydrophilic layer located between the two clusters.

In Fig. 5 we look at the BPTI molecule in a direction parallel to the hydrogen bonds of the central portion of the antiparallel β -sheet (Fig. 5 A). In the presentation of Fig. 5 B, where only the hydrophobic side chains are shown, it is readily apparent that a cleft, which extends through the entire molecule, separates the hydrophobic side chains into two groups. Each of the two groups contains a hydrophobic cluster. The core of the cluster on the left consists of the side chains of Pro 2, Phe 4, Tyr 21, Tyr 23, Phe 45 and Met 52, and the disulfide bonds 5-55 and 30-51. The core of the cluster on the right is formed by Pro 9, Phe 22, Phe 33, and Tyr 35. As can be seen from comparison with Fig. 5 A, the β -sheet of the BPTI structure is located in the cleft between the two hydrophobic clusters seen in Fig. 5 B.

The most prominent component of the hydrophilic layer which separates the two clusters (Fig. 4) is the antiparallel β -sheet formed by the residues 16-36 (Fig. 2). Fig. 6 shows that in the central region of the β -sheet there is an additional network of hydrogen bonded polar groups. The hydrogen bonds between the backbone atoms of Tyr 21 and Phe 45 produce a short stretch of triple stranded β -sheet, and the side chain of Asn 43 is hydrogen bonded to the backbone atoms of Tyr 23 in the β -sheet. The side chains of Asn 43 and Asn 44 are then, via three interior water molecules, connected to the backbone fragment 7-10 and to the short *N*-terminal helix (Fig. 4). This hydrophilic layer essentially divides the molecule into two parts. In its shape it resembles a half barrel which encloses one of the hydrophobic clusters (Fig. 4).

Overall, inspection of the crystal structure of BPTI thus revealed features which appear to be compatible with the hydrophobic cluster architecture suggested by the solution studies (Fig. 3). Fig. 4 would in particular provide an explanation for the thermal denaturation studies, which so far has allowed us to distinguish three different apparent denaturation temperatures for individual polypeptide fragments (20). On the basis of the individual resonance assignments the three different denaturation temperatures could be correlated with the two clusters and the contact area between the clusters.

Use of the facilities at the Zentrum für Interaktives Rechnen (ZIR) of the ETH is gratefully acknowledged. Financial support was obtained from the Schweizerischer Nationalfonds (project. 3.0046.76). W. Braun was the recipient of a fellowship from the Schweizerische Kommission für Molekularbiologie (SKMB). We would like to thank H. Roder and M. Meier for the preparation of the large quantities of RCAM-BPTI needed for the ^{13}C -NMR experiments.

Received for publication 29 November 1979.

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DISCUSSION

Session Chairman: Hans Frauenfelder Scribe: Carolyn Ritz-Gold

FRAUENFELDER: We have an extended comment from Joseph Rosa.

ROSA: We have observed the exchange behavior of the S-protein of RNase S. The results extend some of the findings described here for BPTI. We looked at exchange rates as a function of pH, temperature and S-peptide binding using a tritium exchange method that yields average rates for known local regions of the protein. The details of the method have been published by Rosa and Richards (1979, *J. Mol. Biol.*, 133:399-416). The portions of S-protein whose exchange rates we monitored included regions of α -helix, β -sheet and the C-terminal tetrapeptide.

From the rate of exchange of the isolated S-protein, we found that at pH 2.8 all these fragments had at least one site exchanging relatively slowly. Based on the increase in (OH^-) , the exchange rates in going from pH 2.8 to pH 5.35 would be predicted to be ~ 100 times greater than the rates at pH 2.8. Depending on the fragment, the rates of exchange at pH 5.35 were, however, observed to be only 2-4 times greater. Similarly, the rates of exchange at pH 7 would be predicted to be 35-45 times greater than rates at pH 5.35, but the observed rates at pH 7 were only 5-20 times greater.

This apparent violation of first-order dependence on pH is undoubtedly due to the dramatic increase in the thermal stability of the S-protein at pH 7 compared to pH 2.8. These results therefore point to a similar correlation of exchange rates with thermal stability as that observed for BPTI. The relative exchange rates are presumably linked to the relative thermal stability by intramolecular "vibrational" modes. They are probably not normal modes in the usual sense and may or may not be present in the type of picosecond simulation presented here by McCammon and Karplus.

This pH dependence and correlation with thermal stability also correlates well with Jim Matthew's electrostatic calculations. This kind of argument linking exchange rates to thermal stability via either a distribution of microstates or oscillatory modes is also consistent with the temperature dependence of the exchange rates. If activation energy is plotted vs. temperature, we see from the activation energy at pH 2.8, e.g., that even at low temperature, many degrees of freedom would be populated at the level of kT . We would also predict that exchange would be mediated by a low energy process.

If the thermal stability of the protein is increased, e.g. by going to pH 5.35, we would predict that the electrostatic interactions presumably contributing to the thermal stability and tightening the protein will have local interactions and effectively increase the force constants of the "springs" contributing to these degrees of freedom. We would

therefore predict the loss of certain vibrational modes. The remaining modes are still apparently able to permit exchange via a low-energy process, although the deformations affecting each site may well be different.

As the temperature is increased further, more energetic modes become populated and the activation energy will increase. If the stability of the protein is increased sufficiently, e.g., by binding S-peptide, the thermally available vibrational modes might not be sufficient to permit exchange. This is what we actually observe; in fact, at low temperature, many S protein sites do not exchange at a measureable rate. In contrast to BPTI, however, these effects are not equally distributed among the various fragments.

MCCAMMON: In your paper, it was suggested that the apparent activation volume for ring rotation can be interpreted in terms of formation of a spherical "cavity" with sufficient volume to permit relatively unhindered rotation. One alternative, or complementary, rationale or physical model for this apparent activation volume is based on the collisions between the rings and the atoms of the surrounding protein during rotation. At low pressure these collisions are apparently too infrequent to produce significant slowing of the rotational motion. At hundreds or thousands of atmospheres pressure, however, the collision frequency will probably be increased to the point where frictional slowing of rotation might occur—a well-known phenomenon in simple liquids. This slowing of rotation with increased pressure would, of course, appear as an apparent activation volume. This mechanism is suggested to be complementary since the trajectory calculations indicate that both types of processes occur. In addition, the observation that some of these trajectories pass through regions where the density of the atoms around the ring is somewhat lower than in the x-ray structure indicates that density fluctuations do play a role in these processes. My point is that collisional effects probably do also.

KARPLUS: The simple molecule butane in CCl_4 has been studied from this point of view by Montgomery, Chandler, and Berne. They compared the activation volume from the difference in the size of the trans and gauche forms of the molecule as it isomerizes relative to the viscosity effect. For reasonable pressures, the viscosity effect is significantly larger than the "pure" activation volume effect. Obviously, the details of this effect depend on the density of the system and the coupling. My point is that it is important to have these additional measurements to relate to the more complex system.

ROSENBERG: Your concept of global fluctuation, in contrast to, e.g., smaller local fluctuations, to explain hydrogen exchange from amides was based on the correlation of exchange with thermal stability and an adequate enthalpy of activation for the exchanging hydrogens. We have tried to observe these global fluctuations by studying the somewhat larger proteins lysozyme and myoglobin. In a plot of the activation energy for lysozyme as a function of hydrogens remaining (i.e., going from outside to inside) with the heat of ionization of water subtracted, we find that the activation energy is close to 15 kcal/mol^{-1} over a wide range. If now the energy for the exchange reaction itself is also subtracted ($6\text{--}9 \text{ kcal/mol}^{-1}$), the actual enthalpy for exchange is $<5 \text{ kcal/mol}^{-1}$.

At higher temperatures, the activation energy for thermal unfolding can be calculated from the thermal equilibrium data of Vale and Privalov. We have stabilized lysozyme by adding glycerol and destabilized it by adding denaturants to vary its thermal stability and melting temperature over a wide range. We have found practically no corresponding change in hydrogen exchange at 25°C . We also reversibly adsorbed myoglobin to silica beads and studied its thermal stability by scanning calorimetry. We found that thermal unfolding, as such, was totally abolished. Its stability was lowered; it evidently folded over a large temperature range with successive segmental motions. However, the pattern of hydrogen exchange for the majority of protons was changed by no more than 5–10%. These results make it difficult for me to believe that global fluctuations—if operationally defined as being correlated to thermal unfolding and having an adequate enthalpy—are a major cause of hydrogen exchange at lower temperatures in globular proteins. Is it possible that such fluctuations are characteristic of the family of proteins having a very high thermal stability.

WÜTHRICH: Your observations disagree with ours for all the proteins we have studied thus far. In addition to the series of chemical modifications of BPTI, we have observed groups of homologous inhibitors, snake toxins and scorpion toxins, as well as cytochromes C. For all these proteins we found a clear-cut qualitative correlation between the amide proton exchange rates and the stability. We also studied the effect of denaturants on BPTI and on one of its reduced forms at low temperatures (2°C) and found an increased rate of exchange when the stability of the protein was lowered by addition of guanidinium HCl, ethanol, or urea.

LUMRY: The activation volume for reversible thermal unfolding near room temperature is extremely high, whereas the change in overall volume is very small, so one needs to be careful in mixing rate constants and equilibrium constants when dealing with the effects of pressure on stability and unfolding.

TAINER: You find 2 hydrophobic clusters separated by an antiparallel β -sheet. I have three related questions: What can you say about the differential stability of the two hydrophobic clusters? How different are their respective temperatures of denaturation? How do these differ from that of the β -sheet backbone?

WÜTHRICH: We measured apparent denaturation temperatures for groups in the two clusters and for several α -protons located at the interface. In the case of BPTI, modified BPTI, and the snake toxins, the α -protons showed consistently lower apparent denaturation temperatures than the groups within the clusters. Indeed, this observation was fundamental to the proposed structure and to the concept of relative cluster movements, since changes in the orientation of one cluster with respect to the others would be expected to change the spectral properties of the backbone protons at the interface but not those of groups in the cluster interiors. The difference between the apparent denaturation temperatures of the α -protons and the groups in the most stable cluster is $\sim 5^\circ\text{C}$, and the difference between the two clusters is $\sim 2^\circ\text{C}$. For one of the snake toxins, the difference between the apparent denaturation temperature of the α -protons and that of aromatic residues was $\sim 10^\circ\text{C}$.

TAINER: Your thermal denaturation study suggests that hydrophobic clusters are important in stabilizing molecular architecture; however, it can be argued that hydrophobic forces increase as a function of temperature. Have you tried varying other parameters to assess the importance of hydrophobic clusters in structural stability?

WÜTHRICH: In a review paper, Timasheff reports that presence of trichloroethanol decreases the apparent denaturation temperature of the aromatic groups in lysozyme with respect to that of the backbone groups. This was observed using CD spectroscopy. We have not yet found a denaturant that would lower the apparent denaturation temperature of the aromatic groups of BPTI below that of the backbone groups. We have so far tried various alcohols, urea, and guanidinium HCl.

HENDRICKSON: Can you obtain specific information about the motion of the aromatic rings in the clusters, for example, the r.m.s. rotational amplitudes about the $\sim C_\alpha-C_\beta$ and $C_\beta-C_\gamma$ bonds?

WÜTHRICH: The averaged resonances of pairs of protons in symmetrical positions in the rings fall essentially in the center of the two resonances seen in the anisotropic environment provided by the protein. This indicates that, aside from the flipping about the $C_\beta-C_\gamma$ bond, other motions have only low amplitudes. Additional evidence comes from the temperature factors for the aromatic rings in the BPTI crystal structure. They are smaller than the average for the entire protein structure. This temperature factor would reflect the rapid small amplitude motions, but not the ring flips because they are too rare. So, current evidence seems to indicate that all motions other than ring flips are of small amplitude.

HENDRICKSON: Can that amplitude be determined from the NMR data?

WÜTHRICH: Not exactly. All we can say is that these additional small amplitude motions are manifested in the ^{13}C relaxation data. These motions can, however, be estimated theoretically.

LEVY: We are looking at the effect of picosecond motional averaging, which would tend uniformly to increase the T_1 s. In considering relative effects, in general the aromatic rings look much like the backbone carbons: they are highly hindered in their picosecond motions, but some of the aliphatic side chains can have substantial averaging leading to increases in T_1 s by factors of 2 or 3 over the backbone α -carbons.