

SOLUBILITY AS A FUNCTION OF PROTEIN STRUCTURE AND SOLVENT COMPONENTS

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This review deals with ways of stabilizing proteins against aggregation and with methods to determine, predict, and increase solubility. Solvent additives (osmolytes) that stabilize proteins are listed with a description of their effects on proteins and on the solvation properties of water. Special attention is given to areas where solubility limitations pose major problems, as in the preparation of highly concentrated solutions of recombinant proteins for structural determination with NMR and X-ray crystallography, refolding of inclusion body proteins, studies of membrane protein dynamics, and in the formulation of proteins for pharmaceutical use. Structural factors relating to solubility and possibilities for protein engineering are analyzed.

t is generally known that proteins must be stored in an appropriate temperature and pH range to retain activity and prevent aggregation. This review discusses varying other solvent properties to maintain the stability and solubility of proteins.

Proteins are often most soluble in solution conditions mimicking their natural environment. Serum proteins are soluble in a pH and salt range where mature insulin, which is stored in acidic granules in the cell, precipitates¹. Bacterial proteins may prefer buffers containing glutamate or betaine, compounds that accumulate in response to high concentrations of Cl⁻ in the medium². Caseins and other Ca²⁺ associated proteins may require small amounts of this ion to maintain their native structure during purification⁴.⁵. The stability of lactase (β-galactosidase) is greatly increased in the presence of milk proteins⁶. But for most proteins, experimental determination of the solution properties can help in solvent design.

Low solubility in aqueous solvents is often regarded as an indication that a protein is "hydrophobic", as aggregation of integral membrane proteins after transfer to a hydrophilic environment is a well described phenomenon. But all proteins are to some extent hydrophobic, with tightly packed cores that exclude water. As native, properly folded structures aggregate less than unfolded, denatured ones, there is an intimate relationship between solubility and stability. The free energy of stabilization of proteins in aqueous solution is very low (ca. 12 kcal/mole at 30°C¹o); consequently proteins totter on the verge of denaturation¹o.¹¹. Protein stability can be increased by solvent additives or by alteration of the protein structure itself.

THE PROPERTIES OF PROTEINS IN SOLUTION

Defining solubility. The chemists definition of solubility, parts purified substance per 100 parts pure water, is not useful in a biological frame, as proteins in nature are never found in pure water. Blood and eukaryotic cytoplasm contain on the order of 0.15 M salt, with large quantities of trace metals, lipids and other proteins. The cytoplasm of bacteria is more variable, with salt concentration ranging from 0.3–0.6 M². The solubilizing effects of small molecules and even other proteins means that protein solubility does not correlate with purity¹².

Operationally, solubility is the maximum amount of protein in the presence of specified co-solutes that is not sedimented by 30,000 g centrifugation for 30 min¹³. An even stricter criterion, function retained after centrifuging for 1 h at 105,000×g, has been suggested for membrane proteins¹⁴. If one has a pure, lyophilized protein or a salt precipitate, one can determine solubility by adding increasing amounts of weighed solid, centrifuging, and measuring the protein content of the supernatant. Dissolved protein should reach a maximum (maximum solubility) and level off. (However, in the food industry, solubility is defined by sediment (in ml) remaining after centrifuging; the solubility index is thus inverse to the actual solubility¹⁵.)

The method described in the heading of Figure 1 allows definition of the solubility range of a protein in solution. A protein solution is diluted into a buffer series and the samples centrifuged in microconcentrators. As one can conveniently concentrate about 50 fold, a relatively small amount of protein is sufficient for the estimation.

Measuring stability. Methods for determining the thermodynamic stability of proteins use pH and temperature extremes or high concentrations of denaturants¹⁰. Although useful for discerning changes in the structural stability of mutant proteins that are not clear from activity data, they are not directly correlatable with the half life of proteins in solution. Since aggregation occurs at temperatures well below the Td for proteins, additives that stabilize proteins against aggregation may not necessarily affect the Td¹⁵.

The major problem with using thermodynamic measurements is their failure to account for the kinetic effects that lead to aggregation. Both the enthalpy (ΔH) and entropy (ΔS) of hydration vary greatly with temperature, but they cancel to give a relatively small measured free energy (ΔG) of hydration that seems to vary little with temperature. Most of the temperature dependent kinetic contribution, which is the more important in explaining hydrophobic effects, dissipates in alterations of the solvent structure around the protein and reversible deformation of the protein structure itself^{10,16}. Accurate determination of hydration shells can only be done from crystal structures. Clearly other methods of determining protein stability are needed.

Proteins with shorter half lives generally have larger subunit molecular weight, lower isoelectric points (pI), higher affinity for hydrophobic surfaces, and greater



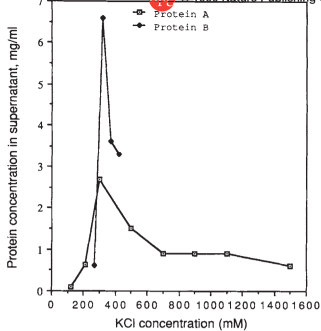


FIGURE 1 Solubility of T7 RNA polymerase (T7RP) as a function of salt concentration in 10 mM Na cacodylate buffer, pH 7.0, 1 mM DTT, and 0.02 mM PMSF. The polymerase solution (ca. 1 mg/ml in 0.1 M KCl, 20 mM tris pH 7.9, 5% glycerol) was diluted 1:10 with the indicated buffers and each sample was individually concentrated in 30 kD MW cutoff "Centricons" (Amicon). The protein in the supernatant (measured by the Coomassie blue assay) after concentration is indicated. The top curve ("Protein B") is from a second measurement using a finer salt gradient and more protein per sample.

susceptibility to proteases. Both of the latter characteristics can be used as the basis for determining enzyme stability in less extreme environments as well as the effect of additives on stability. As less stable proteins have a higher tendency to adsorb to surfaces¹⁷, resistance to mechanical shaking may be a useful indicator of solution half-life¹⁸. Trypsin digestion has been used to define the salt stabilization of hyalin⁵.

Determining surface charge. Isoelectric focusing gives the pI, the pH at which the protein shows no net charge in isoionic conditions. However, due to the binding of salt, one cannot assume that a protein in solution will be negatively charged at pH's above its pI (eg, acidic caseins bind Ca²⁺ and appear positively charged at pH 7⁴). At pH 7.5 and 50 mM salt, most proteins will bind to DEAE-coupled resins if they are negatively charged and to phospho- and other negatively charged resins if they are positively charged. The charge strength can be estimated from the salt concentration required for elution. Gel methods for following the changes in surface charge during protein folding and aggregation have also been developed¹⁹.

Generally, charged proteins can be "salted in" by counter ions. Binding of salts to proteins decreases bound water as well as the net charge at the surface. The solubility of lysozyme, a positively charged protein, was shown to vary more with the anion added than the cation; the anion dependence followed the Hofmeister series²⁰. The solubility of caseins with pI between pH 3 and 5 varies with the cation: sodium, potassium and ammonium caseinates are all more soluble than those prepared with calcium or aluminum^{4,13}.

Determining hydrophobicity. Binding to resins coupled with hydrophobic groups, like Phenylsepharose (Pharmacia) indicates the presence of hydrophobic resi-

dues at the protein surface. Proteins are applied in high salt (0.7–1 M ammonium sulfate), which furthers hydrophobic interactions, and then eluted with a decreasing salt gradient. Most proteins elute between 0.5 and 0.1 M salt; very hydrophobic proteins will not elute into low salt buffer unless the polarity is decreased by adding ethylene glycol. If a protein does not bind to Phenylsepharose, it either has a very hydrophilic surface (eg, RNase A) or it is aggregated.

One can determine the hydrophobicity of a purified protein or follow changes in exposure of hydrophobic groups during folding by measuring interaction with a hydrophobic dye or radioactive tracer (eg, 1-anilino-8-napthalenesulfonate²¹, or ¹²⁵I-TID, 3-(trifluoromethyl)-3-(m-(¹²⁵)iodophenyl)diazirine³).

Aggregation and precipitation. Precipitation via any agent can be: (1) Reversible, as after precipitation with salts or large organic molecules like polyethylene glycol (PEG). Because PEG molecules are excluded from the surface of the protein, a two phase system develops and the protein is concentrated into a smaller volume, where its chances of interacting with another protein molecule to form an aggregate are increased ("excluded volume" model^{22,23}). When the precipitant is removed, the water layer around the original molecule can reform and the protein molecules separate into soluble monomers.

The protein structure does not significantly change during reversible aggregation. A plot of protein in solution versus the concentration of the precipitant should look the same whether it is made with increasing precipitant (to precipitation) or decreasing precipitant (to solubility). Reversibility is assumed for most mathematical models of salting out 12 as well as some recent models of low salt aggregation phenomena 25.26.

(2) Partially reversible, a behaviour frequently seen in pH induced precipitation. Proteins precipitate around their pI and resolubilize as the pH is adjusted upward or downward. But during the pH adjustment, residues may change orientation. When the pH is readjusted, they may not be able to regain their former position and a mixture of structures (isomers) results. Even primary structure can change if a protein is held at acid pH for long periods of time, as for example the deamidation of asparagine 21 of insulin¹. A plot of protein in solution as a function of pH will depend on whether the protein has already precipitated. Kinetic modeling of pH dependent aggregation has been attempted by linear regression²⁷. Models could also use hysteresis equations.

(3) Irreversible, which is usually initiated by extreme changes in the solvent leading to protein denaturation. But some proteins (Fig. 1) also precipitate irreversibly when concentrated above their maximum solubility in a given buffer. Inactive flakes of protein form and remain insoluble even on redilution of the sample or transfer to a buffer of the correct salt concentration. The nature of this tight intermolecular binding is not easy to study, as the aggregates arise from many-body interactions potentially involving all parts of the protein. The initiation could be direct interaction of surface hydrophobic residues, or, as aggregation shows cooperativity, partial disturbance of the hydration sheath or unfolding of the protein structure allowing interaction between normally "buried" residues. Irreversible protein aggregation is not easily modeled. Thermal denaturation curves are done at very low protein concentration, to avoid aggregation terms in the equa-

BUFFER DESIGN FOR MAXIMIZING SOLUBILITY

The properties of water as a solvent. Water's high dielectric constant and its tendency to solvate ions makes it an active copartner in enzyme reactions. When NaCl

crystals are added to water, the atoms attract each other with only about 1/80th of the force in the dry state and the crystal dissolves. Analogously, dissolved proteins are coated with a "hydration shell" around charged and polar groups that prevents self-binding. This bound water does not freeze (some proteins are even efficient antifreezes²⁸) and has different properties from the surrounding solvent molecules²⁹. Bulk water molecules and the protein are in continual fluctuation, which leads to instability in the system^{11,30,31}.

Protein stability in the solid state vs. solution. On the other hand, a protein completely stripped of its hydration shell is difficult to redissolve, as intermolecular hydrophobic forces must be broken. Lyophilization and other drying methods should thus be used with caution and osmotic stabilizers added where necessary to insure that the protein can be rehydrated. The water content of dry milk powder is a compromise between shelf life, which decreases with increasing content of water, and solubility, which increases with hydration index¹³.

Proteins in the solid state have different levels of reactivity depending on the water content. Dried protein with a water content below 22–25%, the minimum required for conformational flexibility and activity, is thermostable. Glycerol, which stabilizes proteins in solution, acts as a humectant on the powder and causes decomposition (as indicated by the Maillard browning reaction) at much lower water content (5–15%). Conversely, sorbitol competes for the hydration water of the protein and does not enhance denaturation³¹.

Solvent additives. There are many potential stabilizing co-solutes for proteins (Table 1). Buffers are described in several excellent reviews^{32,33} and will not be covered here. Table 1 is separated into groups of compounds that have varying effects on the solvation properties of water: the dielectric constant, chemical potential, viscosity, and the clathric tendency (surface tension). The first two qualities are related to protein polarity; the second two relate to the diffusion of the protein, its partial molar volume, and to hydrophobic hydration.

Osmolytic stabilizers. The first group of compounds are osmolytes, which are not strongly charged and have little effect on enzyme activity up to at least 1 M concentration³⁴. Their major effects are on the viscosity and surface tension of water, and hence on solvent ordering. Many of these compounds are used *in vivo* to control the osmotic pressure of eucaryotic and bacterial cells.

Osmolytes can be polyols, sugars, polysaccharides, neutral polymers, amino acids and their derivatives, and large dipolar molecules like trimethylamine N-oxide (TMAO). Glycerol is the most commonly used osmolyte, as it is easily removed by dialysis and does not interfere with ion-exchange chromatography. It does not alter the dielectric constant of the medium significantly and its stabilization effect on proteins seems to be due to its ability to enter into and strengthen the water lattice structure. High concentrations of glycerol decrease the diffusivity and the partial molar volume of proteins 15, thus lowering the rate of aggregate producing solute interactions.

Glycerol has major drawbacks, however, especially for large scale work, as it is an excellent substrate for bacteria. Xylitol, a potential substitute, is not degraded well by bacteria and can be recycled from buffers by alcohol precipitation. PEG can be added to *in vitro* systems for nucleic acid and protein synthesis, where sufficient molecular density but low ionic strength is needed.

Ionic stabilizers. Ionic compounds and salts can stabilize protein structure by shielding surface charges. Salts can also be considered as osmolytes and are used to some extent as such in vivo. E. coli transiently accumulated K⁺

and glutamate after osmotic shock, but within 30 min had switched to carbohydrates as osmoprotectants². Most ionic compounds will affect the dielectric constant and the chemical potential of the solvent and the protein at concentrations well below where they affect the other bulk properties of the solution. Normal bacterial and mammalian enzymes function at a rather low salt concentration and are inhibited by high salt. Halophilic organisms, which can accumulate as much as 7 molal K⁺ intracellularly, have adapted their enzymes to function in very high salt concentrations³⁴.

There is no general rule on salting in of proteins; models that work for one protein are not necessarily applicable to another¹². The salt concentration for maximum solubility frequently falls within a very narrow range. As shown in Figure 1, a 50 mM change in salt concentration gave as much as a 20-fold increase in dissolved T7 RNA polymerase. The solubilizing effect of ions are dependent on the size and charge distribution, but because polar groups on proteins are so diverse, it is hard to say a priori which ion will be best. Large ions are generally better at stabilizing proteins than small ones; in general, the more electronegative the ion, the more it interacts with and destabilizes protein structure.

The finest experimental work on the effect of salts on protein solubility (usually during salting out) has been done by crystallographers^{20,35,36}. The assumed mechanism for salting out by small molecules is that they compete for water molecules until the concentration is too low to maintain the hydration sheath around the protein³⁵.

Divalent cations can have extremely pronounced solubility effects at very low concentrations. Even 1 mM Ca²⁺ induces a conformational change characterized by insensitivity to trypsin in sea urchin hyalin, and Ca²⁺ and Mg²⁺ in the range of 1–20 mM encourage self-association⁵. Zn²⁺ aids in insulin solubilization as well as crystallization¹. As even tiny amounts of Cu, Zn, and Mn (among others) can also induce aggregation, chelators are often added to buffers.

Denaturants, chaotrophs, cryoprotectants, and other additives. One can solubilize almost any protein (usually at the expense of its activity) by chemical denaturation with perturbing ions. Urea stabilizes the unfolded states of proteins because essentially all protein parts, from the backbone to the tryptophan side chains, are more soluble in 6 M urea than water as evidenced by the free energy of transfer into this solvent³⁷. Another class of denaturants, "chaotrophs" like guanidinium, cetyltrimethyl ammonium salts, trichloroacetate, and thiocyanate ions disrupt hydrogen bond formation and disturb the hydration shell around proteins¹⁵. Detergents, amphiphilic compounds that lower the surface tension of water, bind to hydrophobic areas of proteins.

Another class of denaturants, organic solvents, lower the dielectric constant of water. The denaturing activity of hydrophobic solvents is due to a limited detergent effect and that they provide a competing interaction for the intramolecular hydrophobic interactions responsible for a stable tertiary structure. Some proteins are remarkably resistant to the denaturing effects of protic, hydrophilic organic solvents. The original method for isolation of insulin and human interferon- α from tissue and bacteria used extraction with acidic ethanol³⁸; crambin can be crystallized from 60% ethanol³⁹.

Two organic solvents frequently used as cryoprotectants, dimethylsulfoxide (DMSO), and ethyleneglycol, can also denature proteins. DMSO encourages unfolding by favoring peptide N-H-O=S solvent bonds over peptide N-H-O=C peptide bonds⁴⁰. Ethylene glycol, by reduc-

ing solvent polarity, weakens structural hydrophobic interactions.

Unless a protein is to be used *in vivo*, it is general practice to include protease inhibitors, sulfhydryl reductants, bacteriocides, and chelating agents in small amounts to all protein solvents. The more common additives are listed in Table 1c.

Concentrating proteins. Limits on the maximum protein concentration one can achieve are the structure of the protein, the buffer components and the purity of the protein preparation. Overloading the preparation on SDS acrylamide gels may not detect proteases that cause damage during concentration or storage. To minimize this contamination, during purification from bacterial extracts, the protein should completely change buffer at least three times. Suitable transfer methods are salt precipitation and dissolving in fresh buffer, binding to an affinity resin or HPLC column and elution, or gel filtration. Dialysis, flow-through affinity steps, and redissolving lyophilized samples do not count as buffer transfers. All purification buffers should be made with ultra-pure water and HPLC grade chemicals where possible, and sterilized to avoid the reintroduction of bacterial contaminants.

The most commonly used methods for concentration are salt precipitation, affinity chromatography, ultrafiltration, and occasionally, chromatofocusing, electrofocusing, and freeze condensation (for cryoresistant proteins). Very stable proteins and peptides can be lyophilized or spray dried and redissolved. One should get the preparation to as high a concentration as possible by judicious elution of the last affinity step.

The easiest method for concentrating proteins that cannot be lyophilized is ultrafiltration. Microconcentrators (Fig. 1) are useful for volumes up to 10 ml. Stirred pressure cells (Amicon, Millipore, or equivalent) are available for volumes between 10 and 500 ml, and membrane type can be selected according to the size and hydrophobicity of the protein. I was unable to use pressure cells for Mx protein or T7RP, however, as aggregation at the membrane surface was too high. The stir rate should be kept to a minimum as concentrated protein solutions are shear sensitive. For T7RP, losses were lowest with the Sartorius vacuum dialysis system, where I was able to concentrate to 40–50 mg/ml in 0.2 M ammonium sulfate buffer, pH 7.

Hollow fibers or parallel plate continuous flow systems can be scaled up to any size. The Minitan system from Millipore is a good intermediate size for lab use. Protein loss on the membranes is significantly higher than the maximum predicted by the manufacturers.

SITUATIONS WHERE PROTEIN SOLUBILITY BECOMES LIMITING

Refolding inclusion body (IB) proteins. IBs behave like protein that has been irreversibly precipitated. To obtain active protein, high concentrations of chaotrophic agents in the presence of sulfhydryl reducing agents are used to unfold the chains, which must then be refolded during removal of the denaturants. The primary refolding problem is aggregation of partially unfolded protein. In one study, the maximum protein concentration for efficient refolding was only 20 µg/ml41; for interleukin-2 the maximum was only 1 µg/ml⁴². Concentration by ultrafiltration after refolding is possible, but losses due to proteolysis, aggregation of isomers, and membrane binding are frequently very high. For tissue plasminogen activator (t-PA), the folding to intermediate states is rapid but the proper di-sulfide bonds form much more slowly. As the close to native folded forms are relatively soluble, timed addition of more unfolded protein concentrate (a

sort of "fed batch") can allow much higher final concentration of the extract^{41a}. Residual denaturant can also stabilize the native state of the protein; its optimal concentration in the final extract should also be determined. TMAO may be a useful osmolyte when refolding proteins from urea solution³⁴.

Every protein contaminant present during refolding increases the total dilution necessary to avoid aggregation. In addition, partially unfolded proteins are excellent protease substrates. Thus one of the major advances in inclusion body protein refolding has been the development of purification steps that can be used in the presence of the denaturant. These include gel filtration, certain types of affinity chromatography⁴³, and a new method based on the interaction between a poly-histidine peptide fused to the protein of interest and a nickel chelate column⁴⁴.

Alternate methods for refolding, such as binding denatured protein to thiol-sepharose columns or other affinity matrices and eluting with denaturant free buffers⁴⁵, are also being explored. It is possible that activated thiol sepharose mimics the structure of protein disulfide isomerase⁸⁴. Serine proteinases⁴⁶ and interleukin-4⁴⁷ refolding yields were greatly improved by pretreatment with glutathione. Interleukin-2 was renatured by dilution and autooxidation in the presence of Cu^{2+,42}.

Appropriate choice of buffer during the refolding step can also improve yields at higher concentrations of protein¹¹. As optimal refolding conditions vary with the protein, one should either dilute the denatured sample into or dialyze it against many different buffers, and measure active or soluble protein after centrifugation.

Solubilization and reconstitution of membrane enzyme systems. Difficulties in solubilizing proteins from membranes have greatly limited structure and function studies⁴⁸. Membrane proteins function in an amphiphilic environment and fold differently from cytoplasmic proteins: they turn their hydrophobic sites outward rather than inward. This probably accounts for why computer programs developed from soluble proteins predict the opposite of the known X-ray structures for membrane proteins⁴⁹. This structural difference also accounts for the failure of detergents to solubilize IB proteins.

The only way to isolate most integral membrane proteins is to extract them from their lipid environment with bulky detergents (typically Triton X-100 or Emulphogen BC-720). The protein is integrated into a detergent micelle with detergent replacing phospholipids or proteins that were previously in contact with the hydrophobic surfaces¹⁴. Even if the protein is not inactivated by this treatment, low critical micelle concentration (CMC) detergents interfere with protein concentration (by giving a gel), functional assays, and further purification steps (as the detergent's properties dominate the protein's).

Thus proteins are transferred after the initial extraction to less harsh detergents forming smaller micelles¹⁸ via gel filtration. For detergents with CMC's too low to allow for efficient dilution into monomers, one may need to use highly polar micelle dispersing agents like ethanediol or bile salts⁵⁰. NMR structural studies of small membrane proteins in micelles⁵¹ is possible.

A major advance in membrane protein crystallization is the use of "small amphiphiles" to replace detergents binding to the face of the protein⁴⁸. One can thus prevent some of the problems caused by phase separation at higher salt and protein precipitation as the detergent in the micelles becomes too concentrated.

Osmolytic stabilizers (20% glycerol) or high salt (0.3–0.4 M KCl) added before the detergent may stabilize the tertiary structure of the protein during extraction and

dilution of the protein into proteoliposomes⁵². Glycerol or PEG is needed for efficient elution of membrane proteins from chromatofocusing columns⁵³. *In vitro* assays of transport systems from bacteria^{24,52}, signal peptidase from yeast⁵⁴, and the tamoxifen binding protein from a breast cancer cell line⁵⁵ were only possible by judicious control of the salt concentration during detergent extraction of the membrane.

As there is some evidence that high salt concentrations can stabilize secondary structural elements even during tertiary structure disruption²¹, the need for osmolytic stabilization may indicate that membrane proteins can undergo a transition phase, "molten globule" state during solubilization. This state is defined for soluble proteins as an intermediate during reversible unfolding which retains compact structure and CD spectrum similar to the native state, but shows other evidence (eg, increased binding of a hydrophobic dye) of a non-native tertiary structure.

Very concentrated protein solutions and NMR work. As growth factors and enzymes are so active, one generally works with solutions containing less than 1 μg/ml. But much more concentrated solutions are required for microinjection into cells, for clinical trials of drugs and for analytical studies of protein structure. There are many references on preparing proteins for X-ray studies^{1,35}. As it has only recently been shown to be a general method for protein structure determination⁵⁶, less has been written on preparing proteins for NMR. The major requirement for good spectra is absolutely pure protein at high (1–20 mM) concentration.

Most structure determination by $^1\text{H-NMR}$ used solutions in D_2O and H_2O at acid pH. Acid conditions encourage aggregation and protein unfolding, which shortens sample life. Solvent protons can significantly obscure regions of interest in the protein spectrum (C^{α} protons), so buffers are usually phosphate or deuterated Tris. Some groups prefer to work without ionic stabilizers, as they can blur peak profiles and cause excessive heat-up of the sample during measurements. These stringent requirements obviously limit the proteins that can be studied by the technique to small, stable ones.

Assuming the solubility requirements are met, structure analysis for up to 80 amino acid proteins is almost routine⁵⁷. The recent descriptions of well resolved (but very complex) 2-D NOESY and COSY spectra for urokinase (54 kD; solution was 1.5 mM in D₂O at pH 4.5)⁵⁸, as well as the interaction of pepsin (35 kD) with its ¹⁵N-labeled inhibitor⁵⁹ show that investigation of even larger proteins is possible. Isotope-edited NMR spectroscopy, which selectively detects only protons bound to isotopically labeled (¹⁵N, ¹³C) nuclei, allows larger proteins to be analyzed and widens the choice of non-interfering buffers^{33,59-61}.

The solution should be stable during the measurement, which for larger proteins means addition of some salt. Staphylococcal nuclease was solubilized in 0.3 M NaCl at pH 7.6, ovomucoid domains (55 amino acids) were soluble to 12–15 mM in 0.2 M KCl at pH 8⁶², and yeast phosphoglycerate kinase substrate binding was studied in 0.1 M Na ²H-acetate buffer at pH 7.1 (unspecified enzyme concentration)⁶³. Narrowest line widths were obtained for a solution of thrombin (35 kD) concentrated to 0.5 mM in 0.2 M KCl at neutral pH. Significant line broadening was seen if the protein concentration was increased or at lower salt concentrations at the same pH (Gerhard Wagner, personal communication).

PROTEIN ENGINEERING TO INCREASE SOLUBILITY

Amino acid solubility and water affinity. Individual

amino acids vary greatly in solubility and affinity for water (Table 2). Protein solubility is based on the ability of soluble, polar residues to interact with water in such a way that the rest of the protein can maintain an active structure. According to the "hydrophobic collapse" model of protein folding, the driving force for folding is hydrophobic amino acid clustering to avoid water, with the eventual secondary and tertiary structure further stabilized by hydrogen bonding and electrostatic interactions^{8,9}. The distribution of polarity toward the surface is so typical that it has been used as a criterion for protein design⁶⁴.

The data in Table 2 shows that the tendency of residues to be "buried" (definitions range from less than 5% of the residue surface exposed to solvent⁶⁵ to up to 30%⁹) in a protein agrees with these generalizations. Most positively charged and amide side chain residues (His, Lys, Arg, Gln, Asn) were on the surfaces of the proteins studied, and the interiors were primarily composed of the aliphatics Gly, Ala, Ile, Leu, Val and the aromatic Phe. But only 23% of the Trp residues and 13% of the Tyr in the structures were non-accessible to solvent, similar to that of the negative polar residues Glu (20%) and Asp (14.5%). One could agree that the large volume that Trp and Tyr residues occupy makes them difficult to completely bury in a small protein, but more likely the aqueous affinity of the tryptophan imidazole ring and the hydroxyl group of tyrosine were underestimated by early hydrophobicity measurements.

Peptide solubility. There is also a great difference in solubility in secondary structural elements, as illustrated by peptides designed to adopt one conformation or another. For peptides of more than 8 amino acids, sequences favoring α -helix/random coil structures are more soluble in polar solvents than those forming β -sheet structures. The sum of the Chou-Fasman coil index (Table 2) for individual amino acids correlated with the solubility of a series of peptides. The tendency of peptides to form β -sheets could be significantly reduced by the strategic positioning of tertiary peptide bonds (protected residues or prolines) at intervals in the sequence⁴⁰.

The small membrane interacting protein melittin has a positively charged end, which makes it soluble in water, but the protein spontaneously forms a tetramer through interactions at the hydrophobic end²⁶. For other peptides, insertion of arg-NO₂ residues or replacement of hydrophobic residues improved solubility and lowered aggregation tendencies⁶⁶.

Primary structure alterations. Small changes in the protein's primary structure can have drastic effects on stability and solubility. Replacement of the hydrophobic (-EGNFFGKIIDYIKLMFHHWFG) carboxy terminal amino acids of $E.\ coli$ penicillin binding protein 5 with a shorter hydrophilic sequence (-IRRPAAKLE) made the protein water soluble and allowed crystallization⁶⁷. A 13 residue deletion (EVLNENLLRFFVA) in α -casein makes the molecule more soluble⁴. Note that both of the deleted sequences contained FF. Phenylalanine residues are likely to self-interact and are frequently found at subunit interfaces⁶⁸.

A series of point mutations altered the stability and solubility of insulin without significantly affecting the biological activity. In particular, it was possible to replace the asparagine at position 21, which deamidates in acid solution and leads to dimer formation, with Gly, Ser, Thr, Asp, His, and Arg. Similarly, the tendency of yeast cytochrome c to autoreduction and dimerization was eliminated by substituting a Thr for Cys- 107^{69} . A hybrid interferon- α protein precipitated at low salt, unlike either of the parent molecules⁷⁰.

The fragility of protein structure is the major limitation



on the industrial use of enzymes. Thus the question of | lysine and glutamic acid with arginine and aspartic acid, what makes proteins stable at high temperatures and in organic solvents and whether the two correlate is not purely academic. Specific sequence changes in proteins

and a preference for the hydrophobic amino acids Phe, Val and Ile over Leu, Ala and Met71. Most of these changes occur in α -helical regions and increase the net from thermophilic organisms show a tendency to replace | hydrophobicity of the residue 72. Crambin, a plant toxin

| Compounds: | Mode of action | Amount used |
|--------------------------------------------------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-------------------------------------------|
| Osmolytic stabilizers. These generally have little dire | ct interaction with proteins but affect the bulk solution pr | operties of water. |
| Polyols and sugars | | |
| glycerol, erythritol, arabitol, sorbitol, mannitol, xylitol, mannisidomannitol, glucosylglycerol, glucose, fructose, sucrose, trehalose, isofluoroside | These stabilize the lattice structure of water, thus increasing surface tension and viscosity. They stabilize hydration shells and protect against aggregation by increasing the molecular density of the solution without changing the dielectric constant. | 10–40% |
| Polymers | | |
| dextrans, levans, polyethylene glycol | Polymers increase molecular density and solvent viscosity, thus lowering protein aggregation in a single phase system. At high polymer concentration, a two phase system develops and the protein aggregates in the phase where its concentration is the highest. | 1–15% |
| Amino acids and derivatives thereof | | |
| glycine, alanine, proline, taurine, betaine, octopine, glutamate, sarcosine, γ-aminobutyric acid, trimethylamine N-oxide (TMAO) | Small amino acids with no net charge, like gly and ala, have weak electrostatic interactions with proteins. Octopine is a derivative of Arg that is less denaturing to proteins. TMAO stabilizes proteins even in the presence of denaturants like urea. Most of these compounds increase the surface tension of water. | 20-500 mM |
| onic compounds. These affect enzyme reactions and range than the above compounds. | their stabilizing effects on protein occur in a much narro | wer concentration |
| Stabilizing | | |
| citrate, sulfates, acetate, phosphates, quaternary amines | Larger anions shield charges and can stabilize proteins at low concentrations. At high concentrations they lead to precipitation by competing for water molecules. | 20-400 mM |
| Destabilizing | | |
| hlorides, nitrates, thiocyanates | These are generally less stabilizing than large ions but are also useful for charge shielding at lower concentrations. | 20-400 mM ¹ |
| Denaturing (chaotrophs) | | |
| urea, guanidinium salts, trichloroacetates, cetylmethylammonium salts, organic solvents | Denaturants either stabilize the unfolded state of proteins (urea) or perturb protein structure by interfering with hydrogen bonding or disturbing the hydration shell. | 0.2-8 M |
| Other common additives (mostly non-physiological) the buffer but do not change the bulk solvent physical | hat either interact directly with proteins or specifically aff Il properties. | ect impurities in |
| 2-mercaptoethanol, dithiothreitol (DTT), glutathione | Reductants, protect free sulfhydryls from oxidation; prevent intermolecular sulfydryl crosslinking | 1-5 mM 0.1-1 mM ² 1-4 mM |
| phenylmethylsulfonyl fluoride (PMSF), benzamidine | Inhibit serine proteases by reacting with the active site serine hydroxyl group | 0.02-0.05 mM 1 mM |
| eupeptin, peptides | Protect from protease attack by serving as alternate substrates | <1 mM |
| ethylene diamine tetra-acetic acid (EDTA), ethylene-bis(oxyethylenenitrilo)tetracetic acid EGTA) | Chelate divalent metal ions which may react with proteins; inhibit metalloproteases; EDTA at 5-20 mM aids in the lysis of G ⁻ bacteria by lysozyme | 0.01-0.1 mM for buffers |
| catechols, phenolics NaN3 | Bacteriocides | <0.05% <0.1% |

¹Higher concentrations may be used when working with enzymes from halophilic organisms.

²DTT is a potential denaturant of proteins at higher temperatures and has limited solubility in high salt. The concentrations indicated should not be exceeded.

³Dissolve PMSF to 20 mM in isopropanol. The indicated concentrations represent the maximum solubility in aqueous buffers.

that is extremely stable in polar organic solvents, contains no Met and has a higher content of Phe, Val, and Ile residues than the hydrophilic plant toxins to which it is related39.

Post-isolation alterations. One can alter the solubility of isolated proteins in vitro by coupling to polyethylene glycol. Such modifications have been shown to significantly increase the activity, aqueous solubility, and in vivo half-life of interleukin-276. Native lipase M from Candida rugosa, which acts on non-polar substrates, is soluble up to 12% in water but insoluble in benzene. PEG 5000-lipase dissolved rapidly and was active in benzene, toluene, chloroform, and trichloroethane⁷⁷.

Designer proteins. As site-directed mutagenesis is relatively straightforward for recombinant proteins, one might simply replace surface hydrophobic amino acids with acidic residues when aggregation problems arise. But what residues are at the surface and what will the changes do to the tertiary structure and the enzymatic activity? Obviously, the problem of designing soluble proteins is greatly dependent on the ability to predict protein structure.

The Chou-Fasman rules, like most programs used to predict secondary structure from primary sequence data, are based on the study of known structures and the pattern of amino acid usage discerned from them⁷⁸. The learning capabilities of neural networks may be the basis for the next generation of predictive programs⁷⁹. Although the determined "code" can predict where ahelices are likely to occur, β-sheets and turns are less easy to locate. Faster computing techniques have allowed the development of local energy minimization of conformations to predict stable structures80, but the problem of dealing with solvent energies remains. Further, many intermediate secondary structures disappear before the native state is reached81, and no program in use today correctly predicts tertiary structures. Thus mutation is still guesswork.

Mutation of proteins like T4 lysozyme⁶⁹ or RNase A (this institute) may aid in structure based stability design. For example, conversion of a single Thr residue near the carboxy terminus of T4 lysozyme to Ile, Gln, Ser, Arg, or His lowered the stability of the molecule compared to the wild type⁶⁹. Such mutations are rather easy to produce but

TABLE 2 The aqueous solubility and affinity of the amino acids, their relative tendency to exist in a coil conformation (Pc) and accessibility to solvent in protein crystal structures (percent buried). The calculations in the last two columns are based on Table II of reference 65. The amino acid names are followed by the one letter code in parentheses.

| Amino acid | Solubility ^a | Water affini | Water affinity ^b | | | |
|--------------------------------------------------------------------------------------------------------|---------------------------------------|-----------------------------------------------------------|--------------------------------------------------------------------------|-----------------------------------------------------|--------------------------------------------------------------------------|-----------------------------------------------|
| | | F: | W: | P _c ^c | Percent buried ^d | $V_R(A^s)^e$ |
| Aliphatics | | | | | | |
| glycine (G) alanine (A) isoleucine (I) leucine (L) valine (V) | 25.0 16.7 4.1 2.4 8.9 | 0 0.31 1.8 1.7 1.22 | 2.39 1.94 2.15 2.28 1.99 | 1.5 0.7 0.66 0.68 0.62 | 37% (10) 38% (12) 65% (12) 41% (10) 56% (15) | 66 92 169 168 142 |
| Aromatics | | | | | | |
| phenylalanine (F) tryptophan (W) tyrosine (Y) | 2.97 1.14 0.045 | 1.79 2.25 0.96 | $ \begin{array}{r} -0.76 \\ -5.88 \\ -6.11 \end{array} $ | 0.71 0.75 1.06 | 48% (5) 23% (1.5) 13% (2.2) | 203 240 203 |
| Hydroxy/Sulfur | | | | | | |
| serine (S) threonine (T) methionine (M) cystine cysteine (C) | 5.0 s 3.4 0.01 s | -0.004 0.26 1.23 | -5.06 -4.88 -1.48 | 1.32 1.07 0.58 1.18 | 24% (8) 25% (5.5) 50% (2) 47% (3) | 99 122 171 106 |
| Proline | | | | | | |
| proline (P) hydroxy-L-proline | (160) 36.1 | 0.72 | NA | 1.59 | 24% (3) | 129 |
| Charged/Amides | | | | | | |
| aspartic acid (D) glutamic acid (E) asparagine (N) glutamine (Q) histidine (H) lysine (K) arginine (R) | 0.5 0.86 3.1 3.6 4.2 s | -0.77 -0.64 -0.6 -0.22 0.13 -0.99 -1.01 | -10.95 -10.20 -9.68 -9.38 -10.27 -9.52 -19.92 | 1.2 0.83 1.35 0.86 1.06 0.98 1.04 | 14.5% (3) 20% (2) 10% (2) 6.3% (2.2) 19% (1.2) 4.2% (0.1) | 125 155 135 161 167 171 225 |

^aSolubility of the amino acids in g/100 g water at 25°C. Source: CRC Handbook of Chemistry and Physics, 68th Edition (1987–88) and Langs Handbook of Chemistry (12th edition). Sigma L-proline was not soluble at more than 1g/ml, even at 40°C (my

measurement). s = freely soluble.

bTwo different scales are shown. F is the hydrophobicity scale of Fauchère et al.⁷³ which is based on the partition coefficient of the Nα-acetyl-amino acid amides in octanol/water relative to glycine. W is the hydration potential (water affinity) of the amino acid side that a scalculated from the free energy of transfer of the side chain (eg, methane for A) from the vapor phase to water (see ref. 74 for details). Note that both of these scales differ from the frequently used Nozaki and Tanford⁷⁵, which assigns values only to residues considered hydrophobic (A: 0.5; I:1.8; L:1.8; V:1.22; M:1.3; C:0.5; F:2.5; Y:2.3; W:3.4; all other amino acids:0). Coil conformation parameter based on Chou-Fasman data⁶⁰. The parameter is based on the frequency with which a residue is

present in a coil relative to its overall occurrence in the 29 proteins studied.

This column represents the tendency of an amino acid to be buried (less than 5% of residue available to solvent) in the interior of a protein, and is based on the structures of 9 proteins (total of about 2000 individual residues studied, with 587 of these (29%) buried). The first number indicates how often each amino acid was found buried, relative to the number of residues of this amino acid found in the proteins. The number in parentheses indicates the number of buried residues of this amino acid found relative to all buried residues. For other calculation methods with similar results, see 9 and 74a.

Average volume of buried residues, calculated from the surface area of the side chain (ref. 29, 64).



time-consuming to characterize; even selective mutagenesis may deplete the graduate student supply long before all the possibilities are exhausted.

It may be easier to design a soluble protein from scratch, and make "designer proteins" from designer genes82. The potential usefulness of this approach was recently demonstrated by the production in E. coli of an α -helical protein designed from "first principles". The tetramer was soluble in the bacteria and seems to be both αhelical (by its CD spectrum) and very stable (-22 kcal/ mol). Betabellin, a predominantly β-sheet engineered protein which is being made synthetically, may also be coming into solution83.

These proteins show that although the folding language is not understood, a primitive but internally consistent translation is available. If this subcode really works, the next molecules should be stable, soluble, and active.

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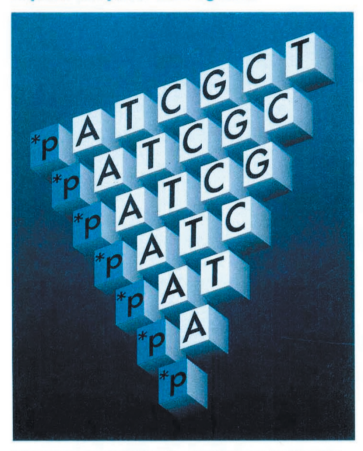
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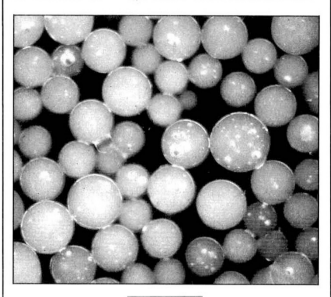
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