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Human aB-crystallin discriminates between aggregation-prone and function-preserving variants of a client protein

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Abstract

Background—The eye lens crystallins are highly soluble proteins that are required to last the lifespan of an organism due to low protein turnover in the lens. Crystallin aggregation leads to formation of light-scattering aggregates known as cataract. The G18V mutation of human γ S-crystallin, which is associated with childhood-onset cataract, causes structural changes throughout the N-terminal domain and increases aggregation propensity. The holdase chaperone protein α B-crystallin does not interact with wild-type γ S-crystallin, but does bind its G18V variant (γ S-G18V). The specific molecular determinants of α B-crystallin binding to client proteins is incompletely charcterized. Here, a new variant of γ S, γ S-G18A, was created to test the limits of α B-crystallin selectivity.

Methods—Molecular dynamics simulations were used to investigate the structure and dynamics of γ S-G18A. The overall fold of γ S-G18A was assessed by circular dichroism (CD) spectroscopy and intrinsic tryptophan fluorescence. Its thermal unfolding temperature and aggregation propensity were characterized by CD and DLS, respectively. Solution-state NMR was used to characterize interactions between α B-crystallin and γ S-G18A.

Results— γ S-G18A exhibits minimal structural changes, but has compromised thermal stability relative to γ S-WT. The placement of alanine, rather than valine, at this highly conserved glycine position produces minor changes in hydrophobic surface exposure. However, human α B-crystallin does not bind the G18A variant, in contrast to previous observations for γ S-G18V, which aggregates at physiological temperature.

Conclusions—*a*B-crystallin is capable of distinguishing between aggregation-prone and function-preserving variants, and recognizing the transient unfolding or minor conformers that lead to aggregation in the disease-related variant.

General Significance—Human α B-crystallin distinguishes between highly similar variants of a structural crystallin, binding the cataract-related γ S-G18V variant, but not the function-preserving γ S-G18A variant, which is monomeric at physiological temperature.

Introduction

For many proteins, the solubility and oligomerization state are critical functional properties that are subject to alteration by mutation or post-translational modification. Protein aggregation is associated with many human pathologies, including Alzheimer's and Parkinson's diseases as well as chronic respiratory conditions, 1 transthyretin amyloidosis, 2 and sickle cell anemia. 3 Protein aggregates can cause disease via loss of function, as in α -antitrypsin deficiency: when α -antitrypsin aggregates, the normal function of this protein to inhibit proteolysis is lost, leading to tissue disruptions. 1 In other cases pathogenesis is due to direct cytotoxicity of the aggregates, as in transthyretin amyloidosis. 2

In the case of crystallins, their normal function is to maintain lens refractivity and transparency. These proteins are therefore exceptionally stable and soluble, even at high concentrations (above 400 mg/mL in humans). ^{4,5} In the eye lens there is a negligible amount of protein turnover, ⁶ meaning that the crystallins must remain soluble throughout the life of an organism, offering a unique model for the study of protein stability and solubility. With age, however, accumulated damage to the crystallins results in light-scattering aggregates causing an opacification of the lens, a condition known as cataract.

There are two main classes of lens crystallins, a- and $\beta\gamma$ -crystallins. The a-crystallins are small holdase chaperone proteins that form very large, multimeric complexes. ^{7,8} The $\beta\gamma$ -crystallins are highly refractive structural proteins that maintain the refractive index gradient of the eye lens. Post-translational modifications including deamidation, ⁹ glycation, ¹⁰ and UV-filter adduction ¹¹ have been shown to reduce crystallin solubility. Many mutations in both a- and $\beta\gamma$ -crystallins have been associated with early-onset cataract, including the R116C¹² and the R49C variants ¹³ of aA-crystallin, the R120G variant of aB-crystallin, ¹⁴ the S39C¹⁵ variant of γ S-crystallin, and the P23T^{16–18} variant of γ D-crystallin. In particular, the γ S-G18V variant of γ S-crystallin has been associated with childhood-onset cataract. ¹⁹

The structure of γ S-crystallin is typical of $\beta\gamma$ -crystallins in general, with two double Greek key domains. Although the sequences of these two domains are not identical, their structures are very similar, resulting in a highly symmetric protein. A theme that has emerged from investigation of cataract-associated variants of γ D- and γ S-crystallin is their structures are often not dramatically different from that of the wild-type protein. In the V24M variant, the core of the N-terminal domain is more closely packed, but the protein maintains the overall β -sheet structure. Similarly, the tertiary structures of the γ S-D26G²¹ and γ S-G18V²² variants are only subtly different from that of γ S-WT. The structure of γ S-crystallin with important features labeled is shown in Figure 1.

Although the structures of γ S-G18V and γ S-WT γ S-crystallin are very similar, the holdase chaperone α B crystallin recognizes these subtle changes in structure, preferentially binding to the partially destabilized γ S-G18V variant but not the wild-type protein.²² The unfolding temperature of γ S-G18V is 6 °C higher than the aggregation temperature, indicating that the aggregation of the protein is not due to the protein simply unfolding;²³ instead γ S-G18V exhibits a slightly destabilized N-terminal domain characterized by an increase in

hydrophobic exposure on the surface of the protein. 24 Not all of the known cataract-associated variants of structural lens crystallins exhibit increased hydrophobic surface area or recognition by α B-crystallin. For instance, the cataract-associated I4F and V76D variants of γ D-crystallin do not bind to α B to any appreciable degree, even though these variants are noticeably destabilized relative to γ D-WT. 25

In order to investigate which local changes in structural crystallins enable recognition by α B-crystallin, we generated the novel G18A variant of γ S-crystallin (γ S-G18A). This variant was chosen to provide a minimal structural disruption that may or may not be recognized by α B-crystallin, as the mutated residue is located in a tight turn with backbone torsion angles that are canonically favorable only for glycine. We find that although γ S-G18A is thermally destabilized and somewhat aggregation-prone relative to γ S-WT, it exhibits minimal structural perturbations at physiological temperature and is not recognized by α B-crystallin.

Results and discussion

Experiments and molecular dynamics simulations suggest that the overall fold of γ S-G18A is similar to that of γ S-WT

The circular dichroism (CD) spectra of γ S-WT, γ S-G18A, and γ S-G18V were measured to probe changes in secondary structure. The spectrum of γ S-G18A exhibits a minimum at 218 nm, indicative of a primarily β -sheet structure (Fig. 2A). The spectrum of γ S-G18A is more similar to γ S-WT than γ S-G18V; it lacks the shoulder at 200–210 nm observed in the spectrum of γ S-G18V.^{19,23}

In vertebrate lens γ -crystallins, two buried tryptophans are ubiquitously conserved within each domain. ^{26,27} Each set acts independently as a FRET pair, enabling the γ -crystallins to quench UV radiation and act as a photoprotectant. ^{28,29} Minor changes to the chemical environment surrounding the tryptophans reduce their quenching efficiency, providing a sensitive measure to detect minor conformational changes. ^{30,31} Increased polarity of the tryptophan environment from solvent exposure results in red-shifting of the fluorescence spectrum. ³² This change can be observed for γ S-G18V, in which the fluorescence is broadened and the peak intensity is red-shifted to 328 nm (Fig. 2B). The emission spectrum of γ S-G18A is almost identical to that of γ S-WT, consistent with the absence of increased relative solvent accessibility (RSA) observed in the γ S-G18A Molecular Dynamics (MD) simulations (see below), and indicates that the chemical environments near the buried tryptophans are minimally perturbed (Supplementary Figure S1).

Anilinonaphthalene 8-sulfonate (ANS) is a small-molecule fluorophore often used to investigate hydrophobic surface exposure in proteins.^{33,34} Binding is mediated through both hydrophobic and electrostatic interactions.^{35,36} Increased hydrophobic surface area is observed in some cataract-related variants such as γ S-G18V,²⁴ γ S-V42M, and γ D-P23T, but not others, including γ S-D26G.^{20,21,34} ANS fluorescence increases moderately in the presence of γ S-G18A relative to γ S-WT, but the signal is dramatically weaker than for γ S-G18V (Fig. 2C). Similarly, a hypsochromic shift to 513 nm (from 525 nm) is observed for γ S-G18A compared to 495 nm for γ S-G18V. These changes indicate that γ S-G18A exhibits

a small increase in hydrophobic surface area relative to γ S-WT. We hypothesize that these changes are localized close to the mutation site; in our previous study of G18V the area around this loop opened up to expose more hydrophobic side-chains to solvent. ²⁴ Further, the strong hypsochromic shifting in conjunction with increased fluorescence suggests strong interactions with positively charged residues, such as Arg19 and Arg20.

MD simulations of γ S-WT and γ S-G18A were performed to investigate mutation-induced changes in protein structure and dynamics. No dramatic changes were observed in the Nterminal domain (NTD) or C-terminal domain (CTD) of \(\gamma \)S-WT or \(\gamma \)S-G18A as a whole based on C_{α} root mean square deviation (RMSD). Whole-protein changes are attributable to the flexible N-terminal tail extension and inter-domain linker regions of the protein (Fig. 3). At the residue level, \(\gamma \)S-G18A exhibits increased root mean-square fluctuations (RMSFs) within the a-helices and sequential loop sequences of the NTD (Supplementary Figure S2). Spatially, these regions are adjacent to the first β -hairpin, which contains residue 18. At the mutation site, no large changes in RMSF are observed. There is no increase in calculated solvent accessibility suggesting that the core of the protein would not be exposed and there would be no additional hydrophobic patches in 2S-G18A when compared to WT (Supplementary Figure S1). The ψ backbone angle of A18 is identical to that of G18, whereas the φ angle is slightly larger and more broadly distributed (Fig. 4). Although it is difficult to determine the exact significance of this change, alanine and valine exhibit a near identical distribution of ψ - φ torsion angles in lens β - and γ -crystallins (Supplementary Figure S3). Like V18 of γ S-G18V, the torsion angle of A18 is dissimilar to all native β - and y-crystallin alanines. Therefore, even though it is not as bulky or hydrophobic, alanine may be expected to cause similar changes in the protein backbone conformation at the mutation site. In \(\gamma \S-G18V \) the torsion angle changes results in altered backbone hydrogen bonding with the subsequent β -strand relative to wild-type. This change, however, is not observed for y-G18A. The absence of significant structural differences between yS-G18A and yS-WT as observed in the MD simulations does not necessarily preclude the possibility of increased aggregation propensity or α -crystallin binding, as it is well-established that structural changes to the lens y-crystallins are not necessary to alter protein stability and aggregation in the lens. 18,20,21,23

NMR chemical shifts of γ S-G18A indicate only mild structural perturbations

The ^{1}H - ^{15}N -HSQC 37 spectrum of γ S-G18A (Fig. 5A) was measured in order to compare the NMR chemical shifts, a sensitive probe of the local chemical environment, to those of γ S-WT and γ S-G18V, which were measured previously. Chemical shift assignments were determined by comparison to the γ S-WT and γ S-G18V assignments from the BioMagResBank (BMRB) (Entries 17576 and 17582), in which 72 % of shifts were assigned. As a whole, the chemical shift perturbations (CSP) for γ S-G18A indicate that the structure is much more similar to γ S-WT than γ S-G18V (Fig. 5B,C). The most pronounced differences are observed in the first 50 residues of the N-terminal domain. Even for these residues, the γ S-G18A chemical shifts are still more similar to γ S-WT than to γ S-G18V. Residues 17-20 were not assignable, probably due to local structural rearrangement.

γ S-G18A is of intermediate stability and aggregation propensity, between γ S-WT and γ S-G18V

The stability of γ S-G18A relative to γ S-WT and γ S-G18V was measured in response to chemical stress. Chemical denaturation was performed using guanidine hydrochloride and assessed using the 355/325 nm fluorescence ratio (Fig. 6A). \(\gamma \S-\text{WT} \) and \(\gamma \S-\text{G18A} \) follow a two-state unfolding model with near identical unfolding midpoints at 2.1 M. In 2S-G18V, three state-unfolding is observed with an unfolding intermediate between ~1 M and 2 M. The unfolding midpoint leading up to the intermediate is at 0.9 M and leading up to complete unfolding is 2.3 M. We hypothesize that γ S-G18A does not exhibit an intermediate unfolding state because the minimal structural changes near the mutation site are not sufficient to catalyze unfolding of the N-terminal domain. This is corroborated by the "m" parameter for each protein, which is the dependance of G on concentration of denaturant. For the first transition observed in γ S-G18V, m is higher than any slope in the second transition suggesting that some part of the protein is more suseptible to denaturation by guanidine hydrochloride. The δG_{app} of unfolding (Table 1) for each protein was calculated by using a linear approximation of a two-state unfolding curve as previously described.³⁹ yS-G18V was separated into two, two-state unfolding curves to generate its thermodynamic properties. G_{app} of unfolding was estimated by extrapolating a line of best fit for each data set to 0^{39} (Supplementary Figure S4).

Thermal denaturation as measured by circular dichroism (CD) as a function of temperature (Fig. 6B). As previously observed, γ S-WT is highly stable, with a unfolding midpoint at 77.67±0.03 °C whereas γ S-G18V has the lowest unfolding midpoint at 66.74±0.04 °C. γ S-G18A is of intermediate stability, with a unfolding midpoint at 69.87±0.05 °C. Although γ S-G18A is nearly as robust as γ S-WT with respect to chemical denaturation, its thermal stability of γ S-G18A is noticably depressed.

Although γ S-crystallin has a high thermal unfolding midpoint, thermal aggregation occurs at much lower temperatures, probably due to transiently unfolded states. Using dynamic light scattering (DLS), we measured protein aggregation as a function of temperature to investigate the behavior of γ S-G18A (Fig. 6C). Between 44 °C and 47 °C, aggregates 10 nm in diameter are observable. These aggregates exceed 20 nm by 53 °C, after which large, insoluble aggregates 1000 nm are observed. The formation of intermediate size aggregates of γ S-G18A is similar to that of γ S-G18V, but at elevated temperature. In contrast, γ S-WT does not exhibit any aggregation until 51–53 °C, when large, insoluble aggregates rapidly form. The similarity in thermal unfolding and aggregation of γ S-G18A to γ S-G18V suggests that γ S-G18A has a similar aggregation pathway to γ S-G18V, but shifted such that its onset occurs above physiological temperature.

aB-crystallin does not bind √S-G18A

Our characterization of γ S-G18A indicates that protein stability is reduced compared to γ S-WT, however, not as dramatically as it is in γ S-G18V. Similarly, the aggregation propensity is intermediate, with an onset temperature around 45 °C. Moreover, γ S-G18A does not appear to undergo large structural changes and does not exhibit a large increase in hydrophobic surface exposure. We next investigated how γ S-G18A would interact with α B-

crystallin to learn more about how mutated, but essentially structurally unmodified, γ crystallins behave as client proteins. Specifically, the objective was to determine whether the
holdase chaperone α B-crystallin could discriminate between γ S-G18A and γ S-G18V. To
this end, 1 H- 15 N-HSQC experiments were performed using mixed samples of 15 N-labeled γ S-G18A and natural abundance human α B-crystallin.

aB-crystallin forms high-molecular weight complexes that enable it to keep bound client proteins soluble. Therefore, if aB-crystallin were able to recognize and bind γS-G18A, then the NMR spectra of the latter should display an increase in line-width and a change in chemical shift for residues that participate in binding to aB-crystallin, as previously observed for γS-G18V.²² The HSQC spectra (Fig. 7A) of ¹⁵N-labeled γS-G18A in the presence of natural abundance aB do not exhibit these changes, instead showing only modest chemical shift perturbations (Fig. 7B) compared with γS-G18A alone, even as temperature increases. The vast majority of the residues have a CSP of less than 0.05 ppm at both 25 (Fig. 7C) and 37 °C (Fig. 7D). Using a threshold of 2*RMS, the CSP of only 3 residues at 25 °C and only 4 residues at 37 °C are large enough to suggest significant conformational change. In both cases these residues with the highest CSP are in the NTD suggesting that there might be some very weak and transient interaction between the NTD of γS-G18A and αB-crystallin.

When comparing the NMR solved structure of γ S-WT to the NMR structure of γ S-G18V and the predicted structure of γ S-G18A there are some key structural differences that could account for the changes in solubility and α B-crystallin selectivity (Fig. 8). It has been previously shown that the binding interface for α B-crystallin includes the 3 cysteine residues on γ S-G18V.²⁴ In this regard γ S-G18A is much more similar to γ S-WT (Fig. 8A), so it is possible that the conformation of this region of γ S-G18A does not promote interactions with α B-crystallin, while the altered conformation seen in γ S-G18V does (Fig. 8B).

Conclusion

CD spectra and NMR chemical shifts suggest that the structure of γ S-G18A is much more similar to γ S-WT than to γ S-G18V. However, the protein is thermally sensitive and there was a noticeable increase in aggregation propensity upon heating. Although it might be expected that a protein of intermediate aggregation potential between γ S-WT and γ S-G18V would also bind α B-crystallin to some degree, this is not the case for γ S-G18A. This indicates that α B-crystallin is capable of discriminating very subtle structural changes in its client proteins, binding to the aggregation-prone G18V variant while leaving the function-preserving G18A variant untouched.

Materials and Methods

Protein expression and purification

The G18A variant of human γ S-crystallin was generated using standard site directed mutagenesis. The cDNA encoding each protein and containing an N-terminal 6x-His tag was ligated into a pET28a(+) vector (Novagen, Darmstadt, Germany) and transformed into Rosetta *E. coli* for expression. All natural abundance proteins were expressed via β -D-1-

thiogalactopyranoside induction. A culture of Lysogeny broth (LB) was an inoculated from a starter culture at OD_{600} of ~0.05 and grown to an OD_{600} of ~1. The growth culture was grown at 37 °C to an OD_{600} of ~1.2 and then induced with 0.5 mM IPTG. After 24 hours at 25 °C the cells were harvested via centrifugation at 4000 rpm. The cells were lysed by sonication and subsequently centrifuged. The lysate supernatant was purified by nickel affinity chromatography and digested using TEV protease (produced in house) to remove the 6x His tag. The TEV protease and 6x His tag were removed by a second round of nickel affinity chromatography. Finally, size exclusion chromatography using a Superdex-75 SEC column was run to confirm purity. Protein masses were further confirmed using a Waters Xevo XS-QTOF.

For 15 N-labeled protein expression, a culture of *E. coli* in natural abundance LB was grown to an OD₆₀₀ of ~1.2. The culture was spun down and the pelleted cells were resuspended in an equivalent volume of M9 minimal media. The culture was allowed to grow for 1 h at 16 °C and before being induced with β -D-1-thiogalactopyranoside at a final concentration of 0.5 mM. The cells were then grown at 18 °C for 36 hours. Purification was performed using the same procedure as the unlabeled abundance protein.

ANS Fluorescence

 γ S-WT, γ S-G18V, and γ S-G18A samples at 1 mg/mL (10 mM sodium phosphate, pH 6.9) were incubated with 750 μ M of 1-Anilinonaphthalene-8-Sulfonic Acid (ANS) for 1 hour at room temperature. Spectra from 450–500 nm were collected on a Cary Eclipse Fluorescence Spectrophotometer (Agilent Technology Inc., Santa Clara California, USA) using a 390 nm excitation and 5 nm slit widths.

Circular dichroism (CD) and thermal unfolding

The circular dichroism of all proteins were collected using a J-810 spectropolarimeter (JASCO, Easton MD). Samples were prepared at 0.1 mg/mL (10 mM sodium phosphate, pH 6.9) and measured over the 190–250 nm wavelength range at RT. Thermal denaturing of proteins was assessed via the absorbance at 218.0 nm. A temperature ramp of 2 °C/min was applied over the temperature range of 25–85 °C using 5 second equilibrations and 1 °C increments. Thermal denaturation experiments were performed at a sample concentration of 0.25 mg/mL (10 mM sodium phosphate, 150 mM NaCl, 1 mM DTT, pH 6.9). The 50% unfolding point was calculated by fitting each curve to a two-state unfolding model as described previously.²³

Tryptophan fluorescence and chemical unfolding

The intrinsic tryptophan fluorescence of all proteins was measured using a SpectraMax GeminiEM spectrometer (Molecular Devices, USA). Samples were excited at 295 nm and emission intensity was recorded from 300–400 nm. All samples were prepared at 0.1 mg/mL in 10 mM sodium phosphate, 50 mM sodium chloride, 0.05% sodium azide, pH 6.9.

The chemical unfolding via guanidine hydrochloride were assessed by 355/325 fluorescence ratio. Protein samples at 0.1 mg/mL were prepared in increasing concentrations of guanidine hydrochloride up to 6 M. The reaction mixtures were equilibrated at room temperature for

48 hours. Unfolding curves were generated by calculating the fluorescence ratio of 355 nm to 325 nm.

Dynamic light scattering

Protein thermal aggregation was measured via dynamic light scattering (DLS) using a Zetasizer Nano-ZS (Malvern Analytical, Malvern, United Kingdom). The number mean was used to assess particle sizes across a temperature range of 25–80 °C. Protein was diluted to 1 mg/mL in 10 mM sodium phosphate, 0.05% sodium azide, pH 6.9.

Solution-state NMR

 $^{15}\text{N-HSQC}$ experiments were conducted using an $^{1}\text{H-}^{13}\text{C-}^{15}\text{N}$ 5 mm tri-axis PFG triple resonance probe attached to a Varian <code>textitUnityINOVA</code> spectrometer (Agilent Technologies) operating at 800 MHz (Oxford Instruments). Proteins were concentrated to 35 mg/mL and then supplemented with 2mM TMSP and 10% D₂O. ^{1}H chemical shifts were referenced to TMSP and ^{15}N shifts were referenced indirectly to TMSP. NMR data were processed using NMRPipe 40 and analyzed using NMRFAM-Sparky. 41

aB-crystallin binding assay

Lyophilized αB -crystallin was resuspended in 10 mM sodium phosphate at pH 6.9 and mixed in a 2:1 ratio with the ^{15}N -labeled γS -G18A variant of γS -crystallin at a concentration of 1.5 mM. A series of ^{15}N -HSQC experiments were then performed at 25, 27, 30, 32, 35, and 37 °C. The sample was incubated at the experimental temperature for 1 hour and the ^{15}N -HSQC experiment was run for 2 hours at each temperature. The chemical shifts collected were compared to a sample of a ^{15}N -labeled γS -G18A variant of γS -crystallin at a concentration of 1.5 mM. Chemical shift perturbations (CSP) were calculated using the following formula:

$$\Delta \delta_{avg} = \sqrt{\frac{\left(\Delta \delta_N / 5\right)^2 + \left(\Delta \delta_H\right)^2}{2}}$$

Molecular dynamics simulations

The top configuration of the human γ S-crystallin solution-state NMR structure²² (PDB ID 2M3T) was used as the initial protein configuration in the γ S-WT simulation and in the γ S-G18A simulation after performing the Gly to Ala substitution at residue 18. The γ S-WT simulation system consisted of a single protein chain, 11,493 water molecules and a single sodium counterion to neutralize the protein net charge for a total of 37,347 atoms. The initial simulation cell size was $63.573 \times 81.480 \times 77.315 \text{ Å}^3$. The γ S-G18A simulation system consisted of a single protein chain, 12,860 water molecules and a single sodium counterion for a total of 41,451 atoms. The initial simulation cell size was $81.52 \times 67.71 \times 80.34 \text{ Å}^3$.

The simulations were performed with NAMD 2.9.⁴² Each simulation system was subjected to 10,000 steps of conjugate-gradient energy minimization followed by a 1-ns MD run at constant pressure (1 bar) and a constant temperature (300 K) over which the protein was released from its initial configuration in a stepwise manner using harmonic restraints on all

backbone heavy atoms. The simulations were then run for 518 ns (γ S-WT) and 1048 ns (γ S-G18A) at constant temperature (300 K) and pressure (1 bar). The CHARMM36 force field^{43–45} was used for proteins and ions and the TIP3P⁴⁶ model was used for waters. The smooth particle mesh Ewald method^{47,48} was used to calculate electrostatic interactions. Short-range, real-space interactions were cut off at 12 Å by means of a switching function. A reversible, multiple time-step algorithm⁴⁹ was used to integrate the equations of motion with a time step of 4 fs for electrostatic forces, 2 fs for short-range nonbonded forces. The integration time step for bonded forces was 1 fs in the γ S-WT simulation an 2 fs in the γ S-G18A simulations . All bond lengths involving hydrogen atoms were held fixed using the SHAKE⁵⁰ and SETTLE⁵¹ algorithms. A Langevin dynamics scheme was used for temperature control, and a Nosé-Hoover-Langevin piston was used for pressure control.^{52,53}

Ramachandran analysis

The observed phi and psi angles for glycine, alanine, and valine were determined by collecting all available native vertebrate lens $\beta\gamma$ -crystallins from the PDB (40 structures). Angles were calculated by using the DSSP package and processed using R. ^{54,55}

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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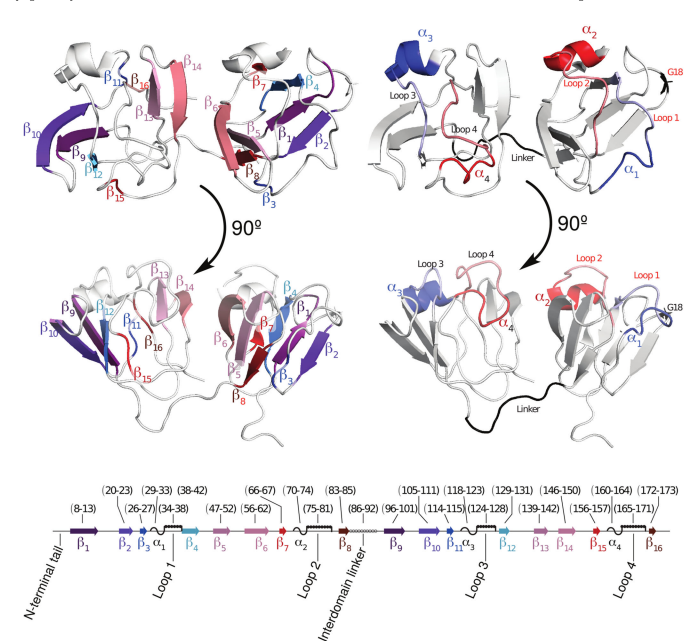


Figure 1: A schematic view of human γ S-WT (PDB ID: 2M3T), highlighting key sequence features, including β -strands (Left) and α -helices or loops (Right) on the structure. The mutated residue (G18) is labeled as well. The span of each helix, strand, or loop is laid out linearly (Bottom) to indicate how the sequence of γ S-WT relates to the structure.

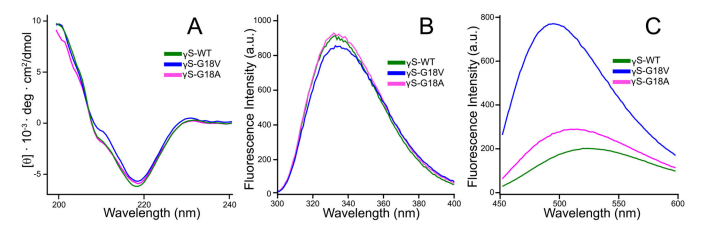


Figure 2: CD spectra indicate that the overall fold of γ S-G18A is similar to that of γ S-WT. (**A**) CD spectra of γ S-WT, γ S-G18A, and γ S-G18V. The spectra of all three proteins are similar overall, however γ S-G18V has a shoulder at 200-210 nm not observed in the others. (**B**) Intrinsic tryptophan fluorescence spectra of γ S-WT, γ S-G18A, and γ S-G18V after excitation at 280 nm. The spectrum of the γ S-G18V variant exhibits a slight red shift relative to the other two. (**C**) ANS binding assay for γ S-WT, γ S-G18A, and γ S-G18V. The γ S-G18V variant has the highest intensity, indicating that the γ S-G18V variant has the largest hydrophobic surface area exposed to solvent.

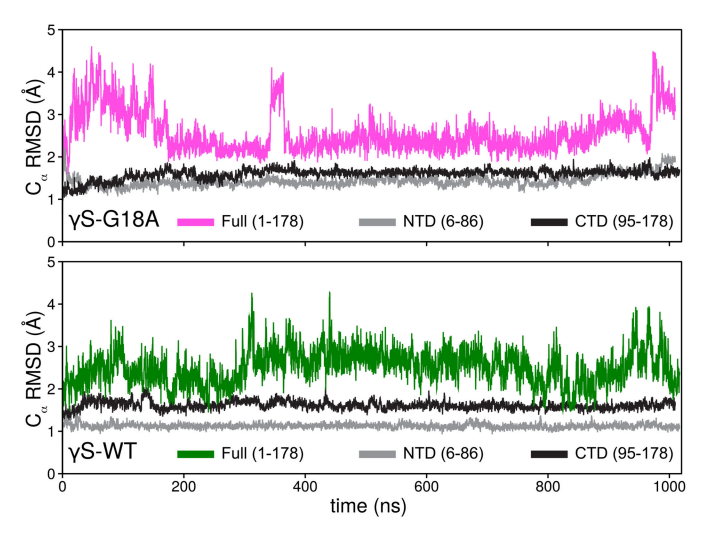


Figure 3: C_a RMSD from the initial configuration. The time evolution of γ S-G18A predicts a well-folded protein much like γ S-WT. Full = RMSD of all α -carbons. CTD = RMSD of residues 95-178. NTD = RMSD of residues 6-86.

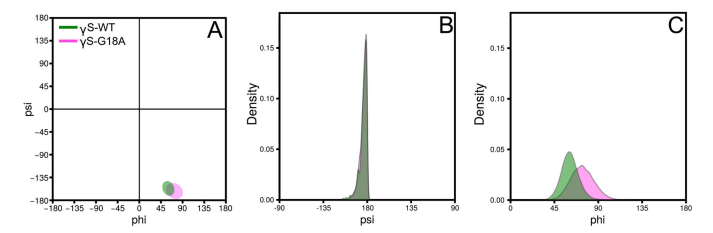


Figure 4: Backbone torsion angles for lens γ S-crystallin residue 18. (**A**) The ψ - φ populations of γ S-G18A are similar to those of γ S-WT. (**B**) The ψ torsion angle distributions are almost identical, while (**C**) the γ S-G18A φ torsional angle distribution is broader that in γ S-WT.

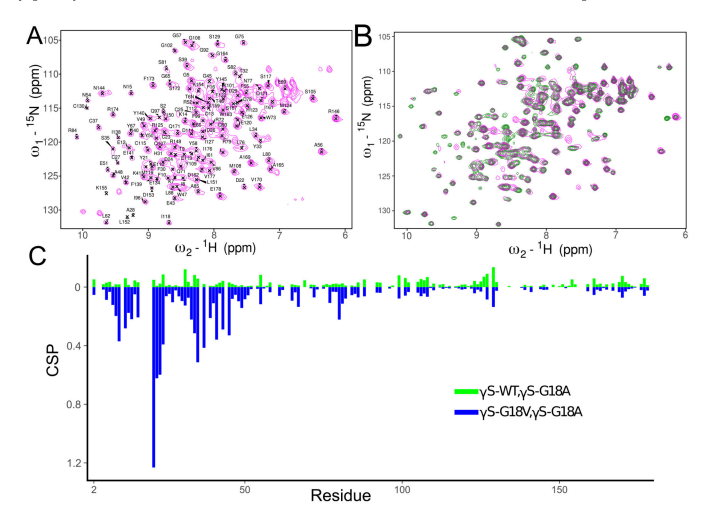


Figure 5: (A) $^{1}\text{H}^{-15}\text{N}$ HSQC spectrum of γ S-G18A. Peaks were assigned by comparing the spectrum of γ S-G18A to the previously assigned spectra of γ S-WT and γ S-G18V. (B) Overlaid $^{1}\text{H}^{-15}\text{N}$ HSQC spectra of γ S-WT (green) and γ S-G18A (magenta). The two spectra overlay very well but some peaks are in slightly different positions. (C) Comparison of the chemical shift differences between γ S-G18A and γ S-WT (green) and γ S-G18A to γ S-G18V (blue).

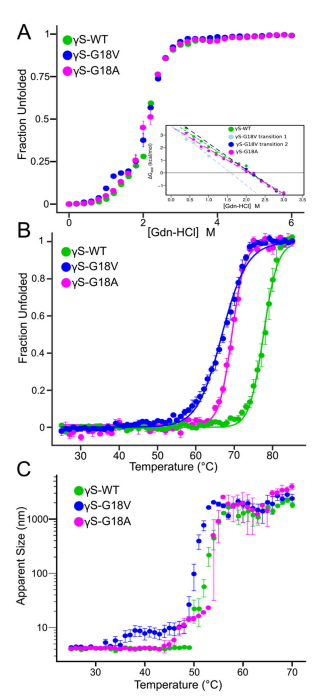


Figure 6: The stability and aggregation propensity of γ S-G18A are more similar to γ S-WT than to γ S-G18V, although decreased stability is observed. (A) Chemical stability of γ S-WT, γ S-G18A, and γ S-G18V was measured by taking the ratio of 355/325 nm after incubation in increasing concentrations of guanidine hydrochloride. Both γ S-WT and γ S-G18A show very similar results. However, γ S-G18V is the least stable protein, exhibiting a three-stage unfolding pattern. The inset shows the linear extrapolation used to calculate the parameters in Table 1. (B) Thermal unfolding curves were generated by monitoring the CD intensity at

218 nm as a function of temperature for γ S-WT (green), γ S-G18A (magenta), and γ S-G18V (blue). γ S-WT is the most stable protein with a thermal unfolding midpoint temperature (T_m) of 77.67±0.03 °C. The γ S-G18A variant is of intermediate stability, with a T_m of 69.87±0.05 °C, while the γ S-G18V variant is the least stable with a T_m of 66.74±0.04 °C. (C) Thermal aggregation propensity was measured by DLS. Both γ S-G18A and γ S-G18V are aggregation-prone relative to wild-type, however, γ S-G18V forms small aggregates at physiological temperature, where γ S-G18A remains monomeric. Each experiment was repeated three times.

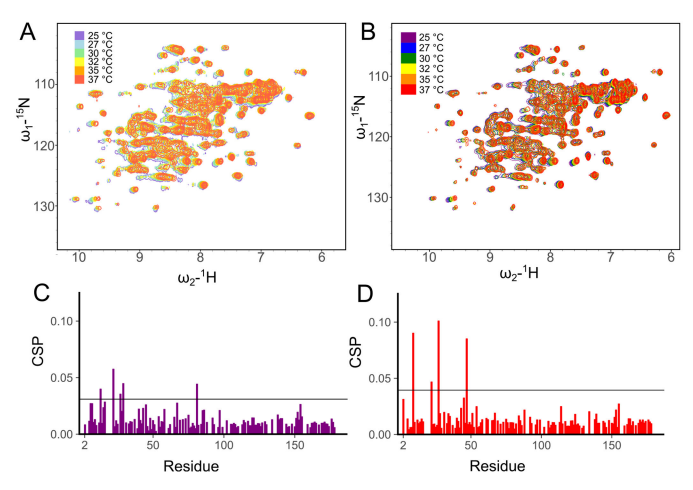


Figure 7: α B-crystallin does not specifically bind γ S-G18A. (A) 15 N-HSQC spectra of 15 N-labeled γ S-G18A as a function of temperature. (B) 15 N-HSQC spectra of 15 N-labeled γ S-G18A mixed with natural abundance α B-crystallin. The spectra are almost identical, suggesting little interaction between γ S-G18A and α B. (C) Observed CSPs for γ S-G18A compared to γ S-G18A+ α B at 25 °C indicate very little change in conformation, suggesting minimal interaction. Only four residues have a CSP greater than 2 * *RMS*, residues 13, 22, 29, and 81. (D) a similar observation is made at 37 °C where only four residues show a CSP greater than 2 * *RMS*, these residues are 9, 22, 27, and 47.

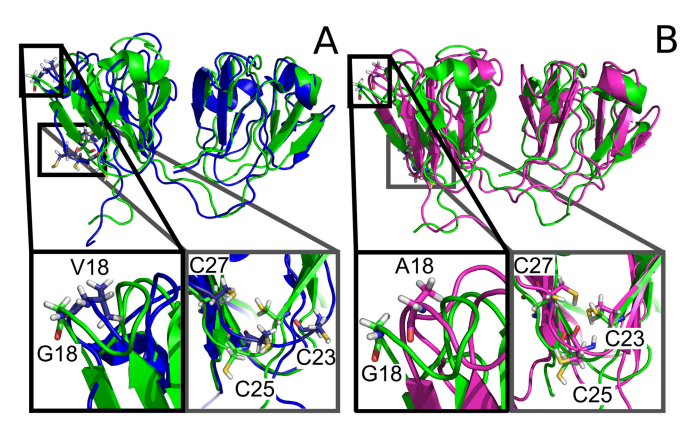


Figure 8:

The predicted structure of γ S-G18A is very similar to that of γ S-WT. (**A**) The overall structures of γ S-WT (PDB ID: 2M3T) and γ S-G18V (PDBID: 2M3U) are very similar but there are some key differences. The loop region around the mutation site exhibits an altered conformation suggesting altered salt-bridging interactions with this loop region. The cysteine residues in γ S-G18V are pointed out into the solvent and not turned into the protein like in γ S-WT. (**B**) Comparing a predicted structure of γ S-G18A generated from molecular dynamics simulations to the NMR structure of γ S-WT shows some important similarities. Although γ S-G18A has a slightly altered conformation in the loop region around the mutation site, the disturbances are not sufficient enough to alter the conformation of the loop containing cysteines 23, 25 and 27, in contrast to γ S-G18V, where these cysteines are more exposed to solvent.

Table 1:

Thermodynamic parameters as calculated from chemical unfolding in Gnd-HCl. The midpoint of unfolding is similar for all three proteins in the final transition, but γ S-G18V shows a stable intermediate. The parameters were calculated from the lines shown in Supplementary Figure S4 where "m" represents the slope of each line. The midpoint of unfolding was read at the x-intercept for each line and G_{app} was read at the y-intercept

Transition 1			Transition 2			
Protein	Midpoint [Gnd- HCl]	G _{app} (kcal/ mol)	m (kcal·mol ⁻¹ ·M	Midpoint [Gnd-HCl]	G _{app} (kcal/mol)	m (kcal·mol ⁻¹ ·M ⁻¹)
WT				2.14±0.01 M	4.07±0.01	-1.93±0.01
G18V	1.58±0.01 M	3.73±0.1	-2.36 ± 0.01	2.3 M±0.01	4.43±0.01	-2.14 ± 0.01
G18A				2.11±0.01 M	3.68±0.01	-1.75 ± 0.01