Role of the Conserved SRLFDQFFG Region of α-Crystallin, a Small Heat Shock Protein

EFFECT ON OLIGOMERIC SIZE, SUBUNIT EXCHANGE, AND CHAPERONE-LIKE ACTIVITY*

Small heat shock proteins (sHsps) are necessary for several cellular functions and in stress tolerance. Most sHsps are oligomers; intersubunit interactions leading to changes in oligomeric structure and exposure of specific regions may modulate their functioning. Many sHsps, including αA- and αB-crystallin, contain a well conserved SRLFDQFFG sequence motif in the N-terminal region. Sequence-based prediction shows that it exhibits helical propensity with amphipathic character, suggesting that it plays a critical role in the structure and function of α-crystallins. In order to investigate the role of this motif in the structure and function of sHsps, we have made constructs deleting this sequence from αA- and αB-crystallin, overexpressed, purified, and studied these engineered proteins. Circular dichroism spectroscopic studies show changes in tertiary and secondary structure on deletion of the sequence. Glycerol density gradient centrifugation and dynamic light scattering studies show that the multimeric size of the mutant proteins is significantly reduced, indicating a role for this motif in higher order organization of the subunits. Both deletion mutants exhibit similar oligomeric size and increased chaperone-like activity. Urea-induced denaturation study shows that the SRLFDQFFG sequence contributes significantly to the structural stability. Fluorescence resonance energy transfer studies show that the rate of exchange of the subunits in the αAdel-crystallin oligomer is higher compared with that in the αA-crystallin oligomer, suggesting that this region contributes to the oligomer dynamics in addition to the higher order assembly and structural stability. Thus, our study shows that the SRLFDQFFG sequence is one of the critical motifs in structure-function regulation of αA- and αB-crystallin.

α-Crystallin, a multimeric protein composed of two types of subunits, αA- and αB-crystallin, is abundantly present in the eye lens (1). Both αA- and αB-crystallin have subunit molecular masses of ~20 kDa each, share high sequence homology (2), can form homomultimers (3, 4), and are known to be present in other nonlenticular tissues as well (5, 6). Ingolia and Craig (7), while comparing the sequences of four small heat shock proteins from Drosophila, discovered the remarkable similarity between αB-crystallin and small heat shock proteins. Prompted by this sequence similarity, Klemenz et al. (8) investigated the expression of α-crystallin under heat shock and concluded that indeed α-crystallin is a heat shock protein. Horwitz (9) subsequently showed that α-crystallin prevents aggregation of other proteins like other heat shock proteins.

αB-crystallin and not αA-crystallin is stress-inducible. Its levels increase under stress such as heat shock, ischemia, oxidation, and infection and in various disease conditions (10–12). αA- and αB-crystallin, either in their hetero- or homo-oligomeric states, exhibit molecular chaperone-like properties in preventing protein aggregation (4, 9, 13, 14). α-Crystallin binds to partially unfolded, aggregation-prone intermediate states of proteins having molten globule-like properties via appropriately placed hydrophobic binding sites (15–18). Studies from our laboratory (13, 19, 20) and subsequently from other laboratories (21, 22) have shown that the chaperone-like activity of α-crystallins is temperature-dependent. The observed temperature-induced enhancement of activity involves a structural transition that enhances the activity by increasing or favorably reorganizing the hydrophobic substrate-binding surfaces (13, 14, 19–22). Its differential and reversible interaction with early unfolding, refolding-competent intermediates of target proteins decreases their partitioning into aggregation-prone late unfolding intermediates (18, 23). This appears to be important in the observed protection offered by α-crystallin to enzymes from heat-induced inactivation (23–26). α-Crystallins/sHsps2 are believed to serve as reservoirs to maintain partially unfolded target proteins in a folding-competent state, capable of subsequent refolding unassisted or by other chaperone systems (27).

Markedly decreased chaperone-like activity and altered structure (28, 29) due to the mutations of the conserved arginine residue, R120G in αB-crystallin and R116C in αA-crystallin, manifest in desmin-related myopathy and congenital cataract, respectively (30, 31). The R116C mutant of αA-crystallin offers reduced protection to apoptosis in lens epithelial cells subjected to UV stress (32). These and other similar observations suggest that the chaperone-like activity of α-crystallin is important in normal cellular function as well as in stress tolerance. Although growing evidence implicates the functional importance of α-crystallins in many cellular processes (27), knowledge about its structure-function relationship remains elusive largely due to the lack of an X-ray crystal structure or NMR solution structure.

Sequence comparison between several sHsps shows an evolutionarily conserved sequence of about 80–100 amino acids, 2

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§ The abbreviations used are: sHsps, small heat shock protein; FRET, fluorescence resonance energy transfer; DTT, dithiothreitol; AIAS, 4-acetamido-4′-(iodoacetyl)amino)stilbene-2,2′-disulfonic acid; LTI, lucifer yellow iodoacetamide; MOPS, 4-morpholinepropanesulfonic acid; bis-ANS, 1,1′-bi(4-anilino)naphthalenesulfonic acid.
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called the "α-crystallin domain" (33), flanked by the N-terminal domain and the C-terminal region (generally termed as C-terminal extension). The C-terminal region is largely unstructured and is believed to have a solubilizing role, since it contains many charged residues (34). Our recent study showed that besides its solubilizing role, it could also influence the structure and the chaperone-like activity significantly (35). The differences in the N-terminal domains and the C-terminal extensions of sHsps may result in their differing oligomeric assembly, size, dynamics, and functions.

Deletion of the first 63 residues of the N-terminal domain of αA-crystallin leads to a drastic reduction in the size of the protein to dimers or tetramers (36). However, truncation of the first 19 residues of the N-terminal domain of αA-crystallin does not alter the size significantly; the rate of subunit exchange also does not change significantly (37). Besides the α-crystallin domain, there are two notable regions (or motifs) that are conserved, especially among the mammalian sHsps: one in the N-terminal domain, RLFDQXFG, and another in the C-terminal extension, the LX(L/V) motif (33). However, the structural and functional significance of these regions is not understood. We therefore set out to investigate the role of the SRLFDQFFG sequence in the structure and chaperone-like activity of αA- and αB-crystallins by creating mutants that lack this sequence (αAΔ20–28 and αBΔ12–26), for simplicity termed αAdel- and αBdel-crystallin henceforth. Our results demonstrate that deletion of this sequence results in significant decrease in the oligomeric size and stability toward urea-induced denaturation, indicating that this sequence is one of the determinants of the quaternary structure of α-crystallins and contributes to the structural stability. Both the deletion mutants exhibit several-fold increased chaperone-like activity compared with that of αA- and αB-crystallin. Our results show that this region also contributes to the dynamics of subunit assembly as the αAdel-crystallin exhibits a significantly increased rate of subunit exchange compared with the wild type protein as monitored by fluorescence resonance energy transfer (FRET).

EXPERIMENTAL PROCEDURES

Materials—pET-21a (+), T7 promoter and terminator primers were obtained from Novagen (Madison, WI), pBSII/SK was from Stratagene, and insulin and bis-ANS were from Sigma. Sephacryl HR-300 was purchased from AP-Biotech. Dithiothreitol (DTT) was obtained from Sisco Research Laboratories (Mumbai, India). Fluorescence probes used in FRET experiments, namely the disodium salt of 4-acetamido-4'-((iodoacetyl)amino)stilbene-2,2'-disulfonic acid (AIAS) and diotassium salt of lucifer yellow iodocacetamide (LYI), were purchased from Molecular Probes, Inc. (Eugene, OR).

Creating Mutant αAdel- and αBdel-crystallin—Recombinant human αA- and αB-crystallin genes were cloned in pET-21a (+) as described earlier (28). These constructs were used as templates to generate the deletion mutants using PCR. For αAdel-crystallin, two independent PCRs were performed using T7 promoter primer and the mutagenic primer 5′-GGGTTGTAAGGCCCTTC-3′ as one primer pair and 5′-GAGGCCCCTTTTGGATGTG-3′ and T7 terminator primer as the second primer pair. Fragments of 157 and 588 bp of PCR products were digested using Ndel and HindIII, respectively, and cloned in the Ndel and HindIII sites of pET-21a (+) expression vector via a three-point ligation. The same strategy was employed for αBdel-crystallin using T7 promoter primer and the mutagenic primer 5′-GGGTTGTAAGGCCCTTC-3′ and T7 terminator primer to generate 140- and 532-bp fragments, respectively. These fragments were also three-point ligated into the Ndel and HindIII sites of pET-21a (+) expression vector. These constructs were verified by sequencing using a 3700 ABI automated DNA sequencer.

Expression and Purification of the Recombinant Wild Type and Mutant Proteins—The wild type and the mutant recombinant proteins were overexpressed in Escherichia coli BL21(DE3) cells. All the purification processes were performed using 50 mM Tris-HCl buffer (pH 7.4) containing 100 mM NaCl, 1 mM EDTA with 0.02% sodium azide (TNE). Bacterial cells were lysed in TNE buffer using lysozyme and then sonicated to shear the genomic DNA. The purification procedure for the wild type and mutant αA- and αB-crystallin was essentially as described by Sun et al. (4). The wild type and mutant proteins were precipitated with 30–50% saturated ammonium sulfate. The protein pellet was dissolved in TNE buffer and loaded on to a Sephacryl HR-300 gel filtration column (1.8 × 130 cm). The fractions containing α-crystallin were pooled and subjected to ion exchange chromatography using Mono Q. The purified proteins were dialyzed against the TNE buffer and concentrated by ultrafiltration. The purity of the wild type and deletion mutant proteins was checked by SDS-polyacrylamide gel electrophoresis and found to be homogeneous. The concentrations of the protein samples were determined by the method described by Pace et al. (38).

Chaperone Assays—The chaperone-like activity of the wild type and mutant proteins was studied using DTT-induced aggregation of insulin, catalase (232 kDa), and aldolase (158 kDa) were used as standards for concentration of the probes at 37 °C. Protein samples were incubated with 100 mM NaCl in the absence or the presence of chaperones (at the different concentrations mentioned in the figure legends to obtain different chaperone to target protein ratios) was incubated at 37 °C. Aggregation was initiated by adding DTT to a final concentration of 20 mM. The aggregation of the target protein was monitored as right angle light scattering using a Hitachi F-4500 fluorescence spectrophotometer. The samples were placed in the thermostated cuvette holder, and light scattering was measured as a function of time by setting the excitation and emission monochromators at 485 nm with the excitation and emission band passes set at 3 nm.

Fluorescence Studies—All fluorescence spectra were recorded using a Hitachi F-4500 fluorescence spectrophotometer with excitation and emission band passes set at 5 and 3 nm, respectively. All spectra were recorded in corrected spectrum mode. Intrinsic tryptophan fluorescence spectra of the wild type and the mutant α-crystallins (0.15 mg/ml) in 10 mM sodium phosphate buffer (pH 7.4) containing 100 mM NaCl were recorded with exciting the sample with 285-nm light. Urea-induced denaturation of the wild type and deletion mutants of α-crystallin was studied by monitoring the changes in the tryptophan fluorescence. Proteins at 0.2 mg/ml in 50 mM Tris-HCl buffer (pH 7.2) in the presence of various concentrations of urea were incubated at room temperature (25 °C) for 2 h, and the tryptophan fluorescence spectrum of these samples was recorded with the excitation and emission band passes of 280 and 340 nm. Fluorescence spectra of the samples were recorded from 400 to 600 nm with the excitation wavelength set at 390 nm.

Circular Dichroism Studies—Near- and far-UV CD spectra of the wild type and the mutant α-crystallins were recorded using a JASCO J-715 spectropolarimeter. Experiments were performed with 1.0 mg/ml protein in TNE buffer using a 1-cm path length cell for the near-UV region and a 0.01-cm path length cell for far-UV region. All spectra reported are the average of five accumulations and are blank corrected. CD spectra were recorded through Grazing-Grating Gradient—Optical Pathlength Correlation. The experiment was carried out essentially as described by Lambert et al. (39). αA-crystallin and the deletion mutants (1 mg) in TNE buffer (pH 7.4) were loaded on top of a 12-mL linear gradient of glycerol (10–40%) made in the same buffer. The tubes were centrifuged for 18 h at 30,000 rpm in a Beckman SW41 rotor at 4 °C. Fractions (0.3 ml) were withdrawn from the top using a Haake-Buchler Auto Densi-Flow IIC gradient former/ remover, and optical density at 280 nm of the fractions was measured using a Shimadzu UV-1601 spectrophotometer. Thryoglobin (669 kDa), catalase (232 kDa), and aldolase (158 kDa) were used as standards for estimating the molecular masses of the α-crystallin samples.

Dynamic Light Scattering Studies — The hydrodynamic radii of the wild type and mutant α-crystallins were measured using dynamic light scattering measurements using a DynaPro MSX Dynamic Light Scattering Instrument from Protein Solutions Inc. The protein samples were diluted through a 0.22-μm filter, and 50 μl of 2 mg/ml α-crystallin sample was used. The cell holder was thermostated at 22 °C. Measurements were made at a fixed angle of 90° using an incident laser beam of 532 nm. The measurements were made with a 1-s time constant at 10 s for each measurement at the sensitivity of 10%. The data were analyzed using graphical size analysis software, Dynamics, provided with the instrument.

Subunit Exchange Studies—The cysteine residues in αA- and αAdel-crystallin were covalently labeled with the fluorescence probes, AEG and fluorescein isothiocyanate (FITC) and AlexaFluor 647, respectively, separately by incubating the protein with the probes in 20 mM MOPS buffer (pH 7.9) containing 100 mM NaCl with a 250 μM concentration of the probes at 37 °C for 18 h. The unlabeled probes were removed by passing the samples through a desalting column (PD10), eluted using 50 mM sodium phosphate buffer (pH 7.5) containing 100 mM NaCl. The samples were diluted with 20 mM MOPS buffer (pH 7.9) containing 100 mM NaCl, and 50 μl of 2 mg/ml α-crystallin sample was used. The cell holder was thermostated at 22 °C. Measurements were made at a fixed angle of 90° using an incident laser beam of 532 nm. The measurements were made with a 1-s time constant at 10 s for each measurement at the sensitivity of 10%. The data were analyzed using graphical size analysis software, Dynamics, provided with the instrument.
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Fig. 1. A, conservation of the RLFDQXFG motif in the N-terminal domains of various small heat shock proteins. B, sequence-based prediction of the secondary structural propensity of the N-terminal domains of αA- and αB-crystallin by various programs using different algorithms: UCSC Computational Biology (58), Sspro8 (59), Homology (60–62), APSSP2 (64), PSIPRED (65), GOR4 (66), and Chou-Fasman (67). The full-length sequences were subjected to prediction methods, and the results of only the N-terminal domains are shown. H, e, t, and g, helix, strand, turn, and 3_10 helix, respectively. Dashed line, random coil. C, helical wheel representation of the sequence SRLFDQFF showing spatial dispositions of the hydrophobic and hydrophilic residues.

RESULTS AND DISCUSSION

Earlier studies indicated that the N-terminal region and the α-crystallin domains are involved in subunit interactions (40, 41). However, it is not clear which specific sequence(s) in the N-terminal domain is crucial for the quaternary structure of α-crystallins. In order to identify the peptide regions that might be involved in the subunit interactions, we have earlier screened proteolytic fragments of α-crystallin for possible interaction with the α-crystallin heteroaggregate (42). One of the fragments that showed significant interaction is a 27-residue peptide from the N-terminal region. Further analysis showed that this fragment contains RLFDQXFG, a well conserved motif in the N-terminal region, among the mammalian sHsps (33). Conservation of this motif in many sHsps (Fig. 1A) suggests that it plays a specific role in the structure and function of these proteins. This motif, SRLFDQFFG, is identical in both αA- and αB-crystallin. Sequence-based structural prediction with different algorithms shows that this sequence has some propensity to form a helical structure (Fig. 1B). The helical wheel analysis on the side chain disposition of the residues in the sequence shows that the putative helical segment can exhibit amphiphatic character (Fig. 1C). It is important to note that an analogous sequence (residues 7–11) in wheat Hsp16.9, SNSFD, has been found to make contact with residues Trp46, Arg109-Phe110 in the α-crystallin domain inter- and intramolecularly (43). It is possible that the corresponding SR-LFDQXFG motif in αA- and αB-crystallin may have such intramolecular interactions. Thus, this region in αA- and αB-crystallin may be an important determinant of structure and function. In order to investigate the role of this conserved motif, we have created deletion mutants of αA- and αB-crystallin, which lack the sequence SRLFDQFFG, αAdel- or αBdel-crystallin, and studied the structure and chaperone-like activity with respect to their wild type counterparts.

We have compared the tryptophan fluorescence spectra of the wild type proteins and deletion mutants of αA- and αB-crystallin and found that the spectra of both αAdel- and αBdel-crystallin differ only marginally from those of the respective wild type proteins (data not shown). This result shows that the microenvironment of the tryptophan residues of αA- and αB-crystallin does not get altered significantly upon deletion of the sequence SRLFDQFFG. However, this result does not rule out the possibility of tertiary structural alterations in other parts of the molecules. Fig. 2A shows that the near-UV CD spectra of αA- and αAdel-crystallin do not overlap with each other and differ in the region between 270 and 300 nm. The difference between near-UV CD spectra of αB- and αBdel-crystallin is more pronounced (Fig. 2B). Drastic changes are observed in the region between 270 and 290 nm.
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The effect of the deletion of the specific conserved sequence, SRLFDQFFG, on the secondary structure of the proteins has been investigated by far-UV CD spectra. In Fig. 2, panels C and D compare the far-UV CD spectra of the mutant and the wild type αA- and αB-crystallin, respectively. α-Crystallins exhibit CD spectra indicative of mainly β-sheet structure with minor content of helical structure (44, 45). As mentioned earlier, sequence-based prediction shows that the deleted specific sequence shows some propensity to form helical structure. It is therefore expected that deletion of this sequence would result in decreased helical content, and since helical structure contributes more toward the observed ellipticity, the far-UV CD spectra of the deletion mutants should exhibit decreased ellipticity. However, the far-UV CD spectra of both αAdel- and αBdel-crystallin show increased ellipticity compared with that of the wild type proteins. This result suggests induction of some secondary structural elements in some regions of the sequence, which could probably arise due to packing alterations upon deletion of the specific sequence.

As mentioned earlier, the N-terminal region and the "α-crystallin domain" are involved in subunit interactions (40, 41). Deletion of the first 56 or 63 residues results in a drastic decrease in the size, whereas truncation of the first 19 residues does not result in significant alteration in the size of αA-crystallin (36, 37). It therefore appears that the sequence between residues 19 and 56 may contribute significantly to the subunit interactions and hence the oligomeric size of the protein. However, whether some specific sequence(s) within this large segment or the segment as a whole contribute to such intersubunit interactions is not clear. In order to understand the role of the conserved SRLFDQFFG sequence in αA- and αB-crystallin in the oligomerization, we have performed gel filtration and dynamic light scattering studies on the wild type and the mutant proteins. Fig. 3 compares the sedimentation profiles of the wild type and deletion mutants on a 10–40% glycerol density gradient. αA-crystallin (Fig. 3A) sediments faster than αB-crystallin (Fig. 3B), and the molecular masses of these proteins have been estimated to be ~550 and 480-kDa, respectively (see "Experimental Procedures" for details). From the profiles in Fig. 3, A and B, it is evident that the deletion mutants, αAdel- and αBdel-crystallin, sediment much slower compared with the respective wild type proteins, indicating that the deletion mutants exhibit lower molecular mass. It is also to be noted that both αAdel and αBdel show a similar profile and peak at the same position, the estimated molecular mass of these mutants being ~300 kDa.

Table I shows the hydrodynamic radii ($R_h$) and polydispersity of wild type αA- and αB-crystallin as well as those of the deletion mutants as measured by dynamic light scattering. The hydrodynamic radius of αA-crystallin is larger (8.88 nm) than that of αB-crystallin (7.85 nm). It is seen from Table I that the polydispersity of αA-crystallin is higher than that of αB-crystallin. It is evident from Table I that deletion of the conserved SRLFDQFFG sequence results in a decrease in $R_h$ value as well as the polydispersity in both αA- and αB-crystallin.

Thus, our sedimentation and dynamic light scattering studies on the wild type proteins and the deletion mutants clearly show that the conserved SRLFDQFFG sequence in αA- and αB-crystallin contributes to the quaternary structure of the protein, especially in higher order subunit assembly. It is important to note that in earlier studies, where 56 or 63 residues were deleted from the N terminus, a drastic decrease in the size leading to dimers and tetramers was observed. Deletion of SRLFDQFFG sequence in αA- and αB-crystallin alters the oligomer formation but does not lead to dimers and trimers.
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Fig. 3. Sedimentation of the wild type α-crystallins and the deletion mutants through a linear glycerol gradient (10–40%). A, "A"-crystallin; B, "B"-crystallin; C, αBdel-crystallin. The positions of proteins used for standard molecular masses are also indicated: aldolase (a) (198 kDa), catalase (b) (292 kDa), and thyroglobulin (c) (669 kDa). See "Experimental Procedures" for details.

Table I

| Protein          | $R_h$ | Polydispersity | Polydispersity |
|------------------|-------|----------------|----------------|
| αA-crystallin    | 8.88  | 2.85           | 22.17          |
| αAdel-crystallin | 7.20  | 1.05           | 14.4           |
| αB-crystallin    | 7.85  | 1.43           | 18.2           |
| αBdel-crystallin | 7.39  | 1.24           | 16.6           |

* The values are the averages of three different experimental values. Each experiment value is an average of 20 data accumulations.

Thus, it appears that the rest of the residues in the segment 19–56 contribute other interacting sites in the intersubunit interactions.

In order to find out whether these structural alterations lead to changes in the hydrophobic surfaces on the proteins, we have probed the hydrophobic surfaces using the fluorescent probe, bis-ANS. Upon binding to apolar surfaces, its fluorescence intensity enhances with a blue shift in the emission maximum of bis-ANS. Upon binding to apolar surfaces, its fluorescence intensity in arbitrary units. Fig. 4 shows that the deletion mutants are more susceptible to urea-induced denaturation than their respective wild type proteins. This result indicates that the SRLFDQFFG sequence contributes significantly to the overall conformational stability of α-crystallins.

In order to investigate whether structural and stability alterations lead to changes in function, we have investigated the chaperone-like activity of the wild type and the mutant α-crystallins toward DTT-induced aggregation of insulin at 37 °C (Fig. 6A). It is seen from the figure that the chaperone-like activity of αB-crystallin is higher than that of αA-crystallin, consistent with the earlier reports on the chaperone-like activity of these proteins (4, 14, 17, 18); a 1:1 weight ratio of αA-crystallin to insulin is required to obtain about 80% protection, whereas only a 0.4:1 ratio of αB-crystallin to insulin is required to obtain similar protection. Interestingly, both the mutants, αAdel- and αBdel-crystallin, exhibit enhanced chaperone-like activity compared with that of the wild type proteins. αBdel-crystallin, however, is more active compared with αAdel-crystallin (Fig. 6A). The concentrations or the weight ratios of the chaperones to target protein required to get a comparable extent of protection differ quite significantly and also vary with temperature. Therefore, deriving a parameter using the chaperone activity data at different ratios under a given set of experimental conditions would be useful in order to make a meaningful comparison. The ratio of target protein concentration to the chaperone concentration would reflect the efficiency of the chaperone. We have therefore compared the weight ratios of the chaperones to target protein required to get a comparable extent of protection differ quite significantly and also vary with temperature. Therefore, deriving a parameter using the chaperone activity data at different ratios under a given set of experimental conditions would be useful in order to make a meaningful comparison. The ratio of target protein concentration to the chaperone concentration would reflect the efficiency of the chaperone. We have therefore compared the weight ratios of the chaperones to target protein to the chaperone at 50% protection of the aggregation. Fig. 6B compares the chaperone efficiencies of the wild type and the deletion mutant proteins. From the figure it is evident that the chaperone efficiency of αA- and αB-crystallin are increased severalfold upon deletion of the SRLFDQFFG sequence; the chaperone efficiency of αAdel-crystallin is 8.4-fold higher than that of αA-crystallin, and that of αBdel-crystallin is 6.0-fold higher than that of αB-crystallin. Thus, the fold increase in the chaperone-like activity of αA-crystallin upon deletion of the SRLFDQFFG sequence is higher than that observed for αB-crystallin.
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As mentioned earlier, α-crystallins exhibit temperature-dependent chaperone-like activity (13, 19, 22). At 25 °C, both αA- and αB-crystallin exhibit less chaperone-like activity; at a 1:1 weight ratio of α-crystallins to insulin, the percentage of protection was found to be 9.2 and 16.3% in the case of αA- and αB-crystallin, respectively. However, even at a 0.5:1 weight ratio, the percentage of protection observed in the cases of αAdel- and αBdel-crystallin was 47.5 and 95.7, respectively. Thus, these results show that the deletion mutants exhibit enhanced activity even at the lower temperature (25 °C).

One of the characteristic features that small heat shock proteins exhibit is that subunits of one sHsp can exchange between the multimers of the same protein or with subunits of other related sHsps. Although the exact role of this dynamic property is not clear, it appears to be of some functional importance, since αB-crystallin and Hsp25/27 form heteromultimers with one another in vivo (47–50). We therefore studied the dynamic property of subunit exchange of the wild type and the deletion mutants. Bova et al. (48) have shown that the subunit exchange of rat αA-crystallin can be monitored using FRET by labeling the single cysteine residue by AIAS as donor and LYI as acceptor fluorescent probes. The human αA-crystallin has two cysteine residues at positions 131 and 142, whereas αB-crystallin does not have any cysteine residues in its sequence. We have therefore studied the subunit exchange only in the wild type and the mutant αA-crystallin. We have labeled one batch of the proteins with AIAS and the other with LYI and mixed these two labeled αA-crystallin or mutant in a 1:1 weight ratio at the indicated temperature and monitored the subunit exchange by FRET at 37 °C. As shown in Fig. 7, while the exchange progresses, the fluorescence intensity of donor decreases with a concomitant increase in the acceptor fluorescence upon exciting the sample with the donor absorption maximum, 332 nm. The ratio of the fluorescence intensity at the emission maximum of the acceptor LYI (520 nm) to that of the donor AIAS (410 nm) monitored as a function of incubation time would thus reflect on the rate of subunit exchange. It is seen from Fig. 7 (A, B, and D) that the rate of subunit exchange between the homomultimers of αAdel-crystallin is significantly higher than that between homomultimers of αA-crystallin. Thus, it is evident from our results that the SRLFDQFFG sequence not only determines the higher order assembly but also contributes to a significant extent to the stability or the architecture of the oligomeric assembly, since the deletion of this region results in increased dynamic nature of the assembly. We have also investigated whether the wild type and the mutant αA-crystallin can exchange their subunits with each other. Fig. 7 (C and D) shows clearly that both αA- and αAdel-crystallin exchange their subunits to form a heteroassembly. From the initial slope of the curves in Fig. 7D, it appears that...
the rate of exchange of subunits between the wild type and mutant protein lies between that of the individual homomultimers of α- or αAdel-crystallin.

We have also investigated the subunit exchange at 25 °C. As seen from Fig. 8, subunit exchange in α-crystallin is not significant at 25 °C, whereas it is quite significant in the case of αAdel-crystallin, although it is slower compared with that at 37 °C. It is evident from earlier studies that temperature-induced increase in the rate of subunit exchange in α-crystallin (47, 48, 50) parallels the temperature-induced increase in the chaperone-like activity (13). The dynamic properties of subunit assembly in sHsps are important for their activity (51, 52). Our results of enhanced chaperone-like activity of deletion mutant of αA-crystallin correlate well with the increased rate of subunit exchange of the multimeric assembly.

Masking and unmasking the chaperone sites under certain conditions might be a mode of regulation of the activities of heat shock proteins. Destabilization of the multimeric assembly by temperature and other post-translational modifications such as phosphorylation appears to alter the structure and the activity of several molecular chaperones and heat shock proteins (43, 53–57). It is therefore important to understand the local structures or sequences that are critical in determining the structure and function of the molecule. Our study shows that the SRLFDQFFG sequence may be at least one such critical motif present in small heat shock proteins, particularly in αA- and αB-crystallin, and changes around this region such as phosphorylation may be regulatory in nature. It is important to note that one of the phosphorylatable serines is the residue before the SRLFDQFFG sequence in αB-crystallin. It is possible that phosphorylation of this residue may change the interaction of the sequence with other parts of the molecule and hence modulate the structure and the chaperone-like activity of the protein.

We conclude that the putative amphipathic helical sequence, SRLFDQFFG, at the N-terminal domain of α-crystallins contributes to the higher order assembly of their subunits as well as structural stability. Interestingly, the mutants exhibit increased chaperone-like activity, increased exposed hydrophobic surfaces, and increased rate of subunit exchange compared with the wild type proteins. It appears that loosening of subunit organization leading to more dynamic properties enhances the available chaperone sites for the target proteins.

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