Structural and Functional Analysis of p26, a Small Heat Shock Protein from 
*Artemia franciscana*

by

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for the degree of Doctor of Philosophy

at

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Dedicated to my parents, Maggie, and Rong...
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Abstract

The brine shrimp, *Artemia franciscana*, undergoes encystment and diapause, exhibiting a remarkable resistance to environmental stresses during this time. Encysted *Artemia* embryos contain an abundant small heat shock protein termed p26, a molecular chaperone contributing to development and survival. To investigate the role of p26 during *Artemia* development and upon exposure to stress, the structural and functional characteristics of wild type and modified p26 were determined. Wild type p26 resided predominantly in the cytoplasm of transiently transfected COS-1 cells, however mutated p26 occurred in both cytoplasm and nuclei. Translocation into nuclei was especially evident for variants missing the complete amino-terminal region, a single site mutant R114A, and a multiple-arginine-deleted mutant “R”. When produced in either prokaryotic or eukaryotic cells, modified p26 lacking the amino-terminal region occurred mainly as monomers and dimers, indicating the relevance of the amino-terminus for p26 oligomer formation. Assembly of higher order oligomers was strengthened by the carboxy-terminal extension, although removing the ten carboxy-terminal residues had only a marginal effect on oligomerization. A mechanism whereby oligomer disassembly assists translocation of p26 from cytoplasm into nuclei was suggested, this of importance because p26 translocates into *Artemia* embryo nuclei upon exposure to anoxia and thermal shock. However, when examined in *Artemia*, the results suggested a process more complex than just oligomer dissociation for regulation of p26 nuclear translocation. Bacterial thermostolerance and protein chaperoning assays demonstrated that modified p26 had decreased chaperoning capability, with truncated variants, mutant R114A and mutant “R” most dramatically affected. Oligomerization is necessary but not sufficient for full chaperone activity, since mutant R114A formed large oligomers while exhibiting limited chaperone capability. Circular dichroism revealed that secondary structures of most mutants, in comparison with wild type p26, possessed increased α-helices and decreased β-conformations. Intrinsic fluorescence and ANS-binding studies demonstrated altered aromatic residue microenvironments and reduced global hydrophobicity for p26 variants. Comparative modeling indicated the p26 α-crystallin domain is arranged in an immunoglobulin-like fold as a β-sandwich consisting of eight β-strands forming two antiparallel β-sheets. This work, which has important implications for understanding the relationship between p26 structural organization and functional mechanisms during *Artemia* development and stress, provides a model for study of other small heat shock proteins. As such, it may lead to a better appreciation of diseases, including ischemia/reperfusion injury, cataract, desmin-related myopathy and neurodegenerative disorders.
# List of Abbreviations and Symbols

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Definition</th>
</tr>
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<tbody>
<tr>
<td>ADCC</td>
<td>autosomal dominant congenital cataract</td>
</tr>
<tr>
<td>aTc</td>
<td>anhydrotetracycline</td>
</tr>
<tr>
<td>ATP</td>
<td>adenosine 5’-triphosphate</td>
</tr>
<tr>
<td>BEB</td>
<td>blot electrode buffer</td>
</tr>
<tr>
<td>Bis</td>
<td>N’, N’-methylenebisacrylamide</td>
</tr>
<tr>
<td>ANS</td>
<td>1-anilino-8-naphthalene-sulphonate</td>
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<tr>
<td>bp</td>
<td>base pairs</td>
</tr>
<tr>
<td>BSA</td>
<td>bovine serum albumin</td>
</tr>
<tr>
<td>cDNA</td>
<td>complementary deoxyribonucleic acid</td>
</tr>
<tr>
<td>CoA</td>
<td>coenzyme A</td>
</tr>
<tr>
<td>CS</td>
<td>citrate synthase</td>
</tr>
<tr>
<td>DAPI</td>
<td>4’, 6-diamidino-2-phenylindole, dihydrochloride</td>
</tr>
<tr>
<td>DEAE</td>
<td>diethylaminoethyl-Sephadex</td>
</tr>
<tr>
<td>DMEM</td>
<td>Dulbecco’s modified Eagle’s medium</td>
</tr>
<tr>
<td>DRM</td>
<td>desmin-related myopathy</td>
</tr>
<tr>
<td>DTNB</td>
<td>5, 5’-dithio-bis(2-nitrobenzoic acid)</td>
</tr>
<tr>
<td>DTT</td>
<td>dithiothreitol</td>
</tr>
<tr>
<td>ddH2O</td>
<td>double distilled water</td>
</tr>
<tr>
<td>ECL</td>
<td>enhanced chemiluminescence</td>
</tr>
<tr>
<td>EDTA</td>
<td>ethylene diaminetetraacetic acid</td>
</tr>
<tr>
<td>EGTA</td>
<td>ethylene glycol-bis (β-aminoethyl ether)-N, N, N’, N’-tetraacetic acid</td>
</tr>
<tr>
<td>FBS</td>
<td>fetal bovine serum</td>
</tr>
<tr>
<td>FITC</td>
<td>fluorescein isothiocyanate</td>
</tr>
<tr>
<td>g</td>
<td>gravity</td>
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<tr>
<td>h</td>
<td>hour</td>
</tr>
<tr>
<td>His</td>
<td>histidine</td>
</tr>
<tr>
<td>HRP</td>
<td>horse radish peroxidase</td>
</tr>
<tr>
<td>HSC</td>
<td>heat shock cognate protein</td>
</tr>
<tr>
<td>HSE</td>
<td>heat shock element</td>
</tr>
<tr>
<td>HSF</td>
<td>heat shock factor</td>
</tr>
<tr>
<td>HSP</td>
<td>heat shock protein</td>
</tr>
<tr>
<td>HST</td>
<td>high salt Tween 20</td>
</tr>
<tr>
<td>IgG</td>
<td>immunoglobulin G</td>
</tr>
<tr>
<td>IMAC</td>
<td>immobilized metal affinity chromatography</td>
</tr>
<tr>
<td>IPTG</td>
<td>Isopropyl-1-thio-β-D-galactopyranoside</td>
</tr>
<tr>
<td>kDa</td>
<td>kilodaltons</td>
</tr>
<tr>
<td>LB</td>
<td>Luria-Bertani</td>
</tr>
<tr>
<td>min</td>
<td>minute</td>
</tr>
<tr>
<td>PAGE</td>
<td>polyacrylamide gel electrophoresis</td>
</tr>
<tr>
<td>PBS</td>
<td>phosphate buffered saline</td>
</tr>
<tr>
<td>PCR</td>
<td>polymerase chain reaction</td>
</tr>
<tr>
<td>PIPES</td>
<td>piperazine-N, N’-bis (2-ethanesulfonic acid)</td>
</tr>
<tr>
<td>PMSF</td>
<td>phenylmethanesulfonyl fluoride</td>
</tr>
<tr>
<td>rpm</td>
<td>revolutions per minute</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Description</td>
</tr>
<tr>
<td>--------------</td>
<td>-------------</td>
</tr>
<tr>
<td>RT</td>
<td>room temperature</td>
</tr>
<tr>
<td>S1</td>
<td>cell-free extract of <em>Artemia</em></td>
</tr>
<tr>
<td>SDS</td>
<td>sodium dodecyl sulfate</td>
</tr>
<tr>
<td>sec</td>
<td>second</td>
</tr>
<tr>
<td>TAE</td>
<td>Tris, acetic acid, EDTA</td>
</tr>
<tr>
<td>TBS</td>
<td>Tris buffered saline</td>
</tr>
<tr>
<td>TEMED</td>
<td>N, N, N', N'-tetramethylethylenediamine</td>
</tr>
<tr>
<td>Tris</td>
<td>Tris-(hydroxymethyl)-aminomethane</td>
</tr>
<tr>
<td>Tween</td>
<td>polyoxyethylene sorbitan monolaurate</td>
</tr>
<tr>
<td>WT</td>
<td>wild type</td>
</tr>
<tr>
<td>X-Gal</td>
<td>5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside</td>
</tr>
</tbody>
</table>
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I. Introduction

1.1 Molecular chaperones

Over 40 years ago, an unusual gene expression profile was observed in the polytene chromosomes of salivary glands in heat shocked *Drosophila melanogaster* larvae (Ritossa, 1962). However, the products of these genes, the heat shock proteins (HSPs), were not identified and named until later, when a rapid and selective increase in the synthesis of specific proteins was demonstrated upon exposure of *Drosophila* larvae to higher than normal temperature (Tissières *et al.*, 1974). According to molecular weight, expression pattern, cell distribution and function, HSPs are categorized as families HSP100, HSP90, HSP70, HSP60, HSP40, and the small HSPs (sHSPs) (Liang and MacRae, 1997; Morimoto and Santoro, 1998; Tsan and Gao, 2004). Many HSPs are expressed constitutively at normal temperature and are indispensable in folding of nascent proteins and stabilizing their structure (Nover and Scharf, 1997; Morrow *et al.*, 2000; Pockley, 2003). Intracellular expression of HSPs is also induced by environmental stresses wherein HSPs function as molecular chaperones, maintaining protein structure and playing a critical role in cell metabolism, growth, differentiation and development (Ohtsuka and Hata, 2000).

The term, molecular chaperone, was first introduced to describe the role of nucleoplasmin, an abundant protein found in *Xenopus* oocytes, in the assembly of nucleosomes *in vitro*. Nucleoplasmin binds to purified histones and promotes nucleosome assembly by donating bound histones to assembling chromatin (Laskey *et al.*, 1978). The term was subsequently applied to define a larger family of unrelated proteins that mediate the correct folding of other polypeptides but are not themselves components of the final
functional protein (Hendrick and Hartl, 1993). Molecular chaperones inhibit nonproductive folding by binding specifically and non-covalently to interactive protein surfaces exposed transiently, with many able to refold partially denatured proteins in an ATP-dependent manner (Ellis, 1990; Ohtsuka and Hata, 2000).

1.2 A general view of sHSPs

1.2.1 sHSP Structure

Many, but not all molecular chaperones are HSPs, among which sHSPs compose a distinctive family characterized by a conserved “α-crystallin domain” of 80 to 100 amino acid residues. As a hallmark of the sHSP group, the α-crystallin domain is usually flanked by a nonconserved N-terminal region and a short flexible C-terminal extension (de Jong et al., 1998a; MacRae, 2000; Haslbeck, 2002; Hsu et al., 2003) (Fig. 1). sHSP monomers are usually 12-43 kDa in mass, and assemble into native oligomers of 9 to 40 subunits with mass up to 1 MDa (Muchowski et al., 1999; Fernando et al., 2003). sHSP oligomers undergo dynamic subunit exchange at elevated temperature usually by a rapid process involving ionic and hydrophobic interactions, this of importance in chaperone action (Bova et al., 1997; Friedrich et al., 2004; Haslbeck et al., 2004). sHSPs have a large binding capacity for denatured proteins which are held in a molten-globule folding-competent state until released and refolded by ATP-dependent chaperones such as HSP70/HSP60 upon restoration of physiologically permissive conditions (Fig. 2) (Lee and Vierling, 2000; Diamant et al., 2001; Mogk et al., 2003; Graf et al., 2004; Rekas et al., 2004). Unlike many other chaperones, sHSPs have limited ability to refold nonnative proteins (van Montfort et al., 2001a; Treweek et al., 2003).

Information on the quaternary structure of sHSPs is limited due to oligomer
Figure 1. Multiple sHSP sequence alignment

The amino acid sequences of selected sHSPs were analyzed by CLUSTAL W (1.82). HCRYAA, *Homo sapiens* αA-crystallin, P04289; HCRYAB, *H. sapiens* αB-crystallin, P02511; HHsp27, *H. sapiens* Hsp27, NP_001532; Ap26, *Artemia franciscana* p26, AAB87967; WHsp16.9, *Triticum aestivum* Hsp16.9, 1GME_A. (-) no amino acid residue; (*) identical residues; (:) conserved substitution; (.) semi-conserved substitution. sHSP domains are indicated above the alignment, while the secondary structure of *Triticum aestivum* Hsp16.9 is depicted under the alignment (van Montfort et al., 2001b).

Amino acid residues in red, small and hydrophobic; blue, acidic; magenta, basic; green, residues containing a hydroxyl or amine group. Residue number is indicated on the right.
Figure 1
Figure 2. sHSP oligomerization and chaperoning

Denaturation of native proteins upon exposure to stress leads to the formation of nonnative intermediates. Large oligomers of some sHSPs dissociate into smaller oligomers such as dimers (or trimers) which bind tightly to nonnative intermediates, thereby preventing irreversible aggregation. In some species, the large sHSP oligomers may bind to the intermediate proteins directly, without dissociation. Once physiologically permissive conditions are resumed, a limited number of bound intermediates may be released and refold spontaneously. The release of other nonnative intermediates and their reactivation, however, occur in the presence of ATP-dependent molecular chaperones such as HSP70 and HSP60. After release of substrate proteins, sHSP dimers (or trimers) reassociate into oligomers, perhaps as inactive storage forms (Treweek et al., 2003; Haslbeck et al., 2004; Stromer et al., 2004).
Figure 2

Protein aggregate → Oligomers

Dissociation → Dimers

Reassociation

Protein binding → Native protein

Denaturation

Renaturation

Spontaneous release/refolding

HSP70/60

Release

ADP + Pi

HSP70/60

Refolding

Dimer-substrates (Chaperoning)

Oligomer-substrates (Chaperoning)
polydispersity, with many sHSPs forming a continuum of structures composed of variable numbers of subunits (Haley et al., 2000). sHSPs form globular complexes as occurs for *Methanococcus jannaschii* (Mj) Hsp16.5 (Kim et al., 1998), mouse Hsp25 and human Hsp27 (Ehrnsperger et al., 1999; Rogalla et al., 1999), however, *Mycobacterium tuberculosis* (M) Hsp16.3 exists as nonameric triangles (Chang et al., 1996). Oligomer formation is usually a prerequisite for chaperone activity of most sHSPs, and the complexes may undergo distinct structural changes upon stress. For example, *Saccharomyces cerevisiae* Hsp42 functions as an active chaperone composed of large dynamic oligomers under either physiological or stressful conditions, while Hsp26 from the same organism disassembles reversibly into dimers for efficient interaction with unfolding proteins upon elevated temperatures (Haslbeck et al., 2004; Stromer et al., 2004).

Although the oligomers of most sHSPs show structural variability and are polydisperse, a few sHSPs have monodisperse oligomerization that is well characterized. Reported examples are the X-ray crystallographic structures of archaeal Mj Hsp16.5, a 24-subunit oligomer (Kim et al., 1998) (Fig. 3a), and wheat *Triticum aestivum* (Ta) Hsp16.9, a 12-subunit oligomer (van Montfort et al., 2001a) (Fig. 3b), which reveal common features although structurally different. Despite limited sequence identity, the α-crystallin domain in monomers of each crystallized sHSP oligomer exhibits a β-sandwich composed of two anti-parallel β-sheets (van Montfort et al., 2001b). The most extensive interface for both oligomers is between dimer subunits and the oligomers are stabilized by the interaction of hydrophobic residues in the C-terminal extension with a hydrophobic groove on the α-crystallin domain of a neighboring subunit dimer (Friedrich et al., 2004). The resolution
Figure 3. X-ray crystallographic structures of sHSPs

a. X-ray crystallographic structure of archaeal Mj Hsp16.5. The spherical 24-mer has an outer diameter of 12 nm and inner diameter of 6.5 nm; image resolution is 0.29 nm; b. X-ray crystallographic structure of wheat Ta Hsp16.9 arranged as dodecameric double disks which are approximately 9.5 nm wide and 5.5 nm high with a central cavity of about 2.5 nm wide; image resolution is 0.27 nm (Adapted from (Haley et al., 1998; Kim et al., 1998; van Montfort et al., 2001b)).
Figure 3
of the N-terminal arms from six subunits indicates wheat Ta Hsp16.9 oligomers are further stabilized by extensive N-terminal arm interactions, where essential contacts between dimers are formed by the first 40 N-terminal residues which generate pairs of interwining α-helical domains located inside the dodecamer (van Montfort et al., 2001b; Wintrode et al., 2003).

The dodecameric Ta Hsp16.9 consists of two disks, each comprising six α-crystallin domains organized in a trimer of dimers (Fig. 4a). The monomer folding and oligomer assembly are mutually interdependent processes wherein strand exchange, helix swapping, loose knots and hinged extension are involved (van Montfort et al., 2001b). The α-crystallin domain of each monomer consists of a β-sandwich composed of two antiparallel β-sheets encompassing a long loop, followed by a short C-terminal extension (Fig. 4d). Within a dimer, one monomer has a disordered N-terminal arm, while the other has a fully resolved N-terminal arm which is composed of helices connected by random coil. Residue R108 represents one of the most conserved sites among diverse sHSPs, and strengthens the dimer of Ta Hsp16.9 by forming an intermolecular salt bridge with E100 in the dimerization loop extending from the β-sandwich which engages in strand exchange to stabilize the partner monomer (Fig. 4c). The dimers of Ta Hsp16.9 form “eclipsed” tetramers whose bottom dimers appear hidden when viewed down the crystallographic three-fold axis, with β7 strands and part of the dimerization loops from the two monomers with ordered N-terminal arms, while the dimer interface is strengthened by two C-terminal extensions from monomers with disordered N-terminal arms (Fig. 4b). Residues 1147 and 1149 from β10 strand of a monomer are associated in a hydrophobic groove between strands β4 and β8 of the interacting monomer, forming a
Figure 4. *Ta* Hsp16.9 structure

a. An overview looking down the crystallographic three-fold axis of *Ta* Hsp16.9, which is perpendicular to three crystallographic two-fold axes. Dimers in the top disk are displayed in red, green and blue, dimers in the bottom disk in pink, sage and turquoise. The dimer-dimer interface that forms an “eclipsed” tetramer whose bottom dimers appear hidden when viewed down the crystallographic three-fold axis, is located between red-pink, blue-turquoise, and green-sage dimers; b. An outside view of one side of the *Ta* Hsp16.9 dodecamer, showing how dimers from the top and bottom disks are related by a two-fold axis to form an “eclipsed” tetramer. In the top dimer, the monomer with a disordered N-terminal arm is shown as a pink ribbon, while the monomer with an ordered arm appears as a solid molecular surface, with surface hydrophobic patches shown in blue. Monomers of the bottom dimer are represented by yellow and red ribbons. Residues I147 and I149 of the I/VX/V motif in the C-terminal extension from the bottom monomer (yellow) extending from one dimer to another cover hydrophobic grooves which are putative substrate binding sites in the α-crystallin domain of an adjacent dimer; c. The dimer of *Ta* Hsp16.9, composed of a complete monomer (red) and an N-terminal-arm-disordered monomer (pink). N- and C-terminals are labeled, as are E100 and R108, oppositely charged amino acid residues which form an intermolecular salt bridge in the dimerization loop; d. Ribbon diagram of *Ta* Hsp16.9 monomer with the ordered N-terminal arm shown in green, while the α-crystallin domain and the C-terminal extensions are in red (adapted from van Montfort *et al.*, 2001b).
binding motif (I/VXI/V) conserved among sHSPs and covering a putative substrate binding site in the α-crystallin domain of the monomer in the adjacent dimer (Fig. 4b) (van Montfort et al., 2001b).

1.2.2 sHSP Function

sHSPs suppress aggregation of nonnative proteins thereby conferring stress resistance to cells and their functional relevance is widespread. Active α-crystallin, wherein αA-crystallin and αB-crystallin exist in a molar ratio of approximately 3:1, is essential for refracting light and maintaining lens transparency (Bera et al., 2002). In vertebrates, sHSPs interact directly with actin and intermediate filaments, potentially altering the filament assembly/disassembly equilibrium of cytoskeletal components (Perng et al., 1999). αB-crystallin negatively regulates TNF-α, DNA damage and differentiation-induced apoptosis, and protects cells against genomic instability (Kamracht et al., 2001; Kamracht et al., 2002; Bai et al., 2003). Overexpression of Hsp22 in the mitochondria of D. melanogaster motor neurons extends life span and increases resistance to oxidative injuries (Morrow et al., 2004). Several proteins are associated in vivo with Hsp16.6 from the photosynthetic cyanobacterium Synechocystis during heat treatment, revealing the potential for sHSPs to protect cellular functions as diverse as transcription, translation, cell signaling, and secondary metabolism (Basha et al., 2004). In bovine lens, helices in α-crystallin selectively recognize single-stranded and double-stranded DNA (Singh et al., 1998; Bloemendal et al., 1999), indicating potential relationships between sHSPs and nucleic acids in vivo. sHSPs may also be involved in the transient storage and protection of mRNA during stress-induced translation arrest (Scharf et al., 1998; Kedersha et al., 1999). Whether sHSPs bind RNA directly or indirectly via other proteins as intermediates
is uncertain. Studies on lens plasma membrane suggest that a subfraction of \(\alpha\)-crystallin becomes buried in the membrane bilayer (Cobb and Petrasch, 2000a), and the majority of heat-induced \textit{Synechocystis} Hsp16.6 is associated with thylakoid membranes (Hörúath \textit{et al.}, 1998; Lee \textit{et al.}, 2000), suggesting the involvement of these sHSPs in membrane fluidity regulation. It is thus reasonable to propose that upon stress, the cellular pool of sHSPs, at least in some organisms, is divided into two or more subfractions, including those responsible for protein chaperoning and those for membrane stability (Narbonhaus, 2002b).

1.3 Mutational analysis of sHSPs

Mutations to sHSPs may affect structural stability and compromise chaperone function, and this is the subject of intensive study. For example, internal deletion mutations of human \(\alpha\)A- and \(\alpha\)B-crystallin lacking residues SRLFDQFFG at positions 20-28 and 21-29 respectively, yield oligomers of significantly reduced size and decreased stability to urea-induced denaturation, indicating the motif is an important determinant of quaternary structure (Pasta \textit{et al.}, 2003). Removal of the N-terminus from \textit{M}t Hsp16.3 dissociates the nonameric oligomer into trimers, the building block of this sHSP (Koteiche and McHaourab, 2003). Deletion of the N-terminus of yeast Hsp26 reduces the twenty-four subunit complex to dimers and abolishes chaperone activity, indicating the structural and functional importance of the region (Stromer \textit{et al.}, 2004). The N-terminal region may be buried within sHSP oligomers, involved in monomer-monomer interactions, and of critical importance for the association of sHSPs with nonnative proteins and the formation of large oligomeric complexes (Kim \textit{et al.}, 1998; Stromer \textit{et al.}, 2004; Usui \textit{et al.}, 2004). For \textit{Mj} Hsp16.5, however, an N-terminal-deleted mutant behaves similarly to
its wild type counterpart, suggesting this part of the protein does not have a critical structural role, but is important in organizing monomers before final oligomer assembly (Kim et al., 2003b).

The $\alpha$-crystallin domain mediates subunit interactions and is responsible for the formation of sHSP dimers and trimers (Feil et al., 2001; Pasta et al., 2004). Mutations in this domain may have dramatic consequences. For instance, variations at F94 in the core $\alpha$-crystallin domain of HspH from the bacterium Bradyrhizobium japonicum disrupt function and dimerization, leading to monomer formation, whereas modification of G114, a conserved residue among sHSPs, impairs chaperone function without affecting oligomerization (Lentze et al., 2003). Mutant R120G of $\alpha$B-crystallin exhibits decreased $\beta$-sheet secondary structure, irregular tertiary and quaternary structure, the absence of a clear central cavity and a two-fold increase in oligomeric mass (Bova et al., 1999b). R120G loses the ability to inhibit proteolytic activation of caspase-3 which affects apoptosis regulation (Kamradt et al., 2002). Furthermore, R120G fails to stabilize the intermediate filament desmin and co-aggregates with this protein, leading to desmin-related myopathy (DRM) with symptoms similar to those caused by desmin mutations (Vicart et al., 1998; Goldfarb et al., 2004). The misfolding of R120G contributes to inclusion body formation, although their occurrence may be delayed or prevented by expression of wild type $\alpha$B-crystallin or other chaperones (Chavez-Zobel et al., 2003). Similarly, a mutation at the equivalent position in human $\alpha$A-crystallin, R116C, results in impaired chaperone activity, increased oligomeric polydispersity and enhanced membrane association (Cobb and Pettrash, 2000b), all potential influences in the pathogenesis of autosomal dominant congenital cataract (ADCC). R116C $\alpha$A-crystallin
has a diminished protective ability against UVA radiation-induced lens epithelial cell apoptosis (Andley et al., 2002). Missense mutations of K141 to either E or N in human HspB8 (Hsp22), a residue corresponding to R120 in αB-crystallin or R116 in αA-crystallin, lead to dysfunction of axonal transport and dysregulation of the cytoskeleton, causing motor neuron death in distal hereditary motor neuropathy (HMN) (Irobi et al., 2004). Together, the results indicate a single arginine or lysine mutation in the core α-crystallin domain has dramatic effects on the structural integrity and chaperoning function of human sHSPs. Substantial changes caused by these mutations are likely due to the fact that the arginine and lysine residues are buried and potentially form salt bridges with residues of opposite charge. The disruption or absence of such salt bridges could lead to protein conformational changes and loss of function (Kumar et al., 1999; Bera et al., 2002).

Besides acting primarily as a solubilizing agent for sHSPs and sHSP-substrate complexes formed during stress, the C-terminal extension is characterized by structural flexibility and sequence diversity. Being generally polar and hydrophilic, this extension plays a vital role in chaperone activity by maintaining sHSP solubility and binding to oppositely charged residues of unfolding proteins which are then held by the hydrophobic N-terminal region of sHSPs (Derham and Harding, 1999). The C-terminal extension of sHSPs is susceptible to cleavage, and deletion of the region results in modified tertiary structure, compromised solubility and decreased chaperone function (Takemoto et al., 1993; Andley et al., 1996). For instance, mouse Hsp25 lacking the C-terminal extension fails to stabilize α-lactalbumin against precipitation following reduction by dithiothreitol, and is less stable at elevated temperatures than the wild type, suggesting that the C-
terminal extension is indispensable for full chaperone activity (Lindner et al., 2000). Progressive deletion of the C-terminus from rat and human αA-crystallin leads to significant alteration of tertiary structure and dramatic decline in oligomer mass from approximately 550 kDa to 150 kDa (Thampi and Abraham, 2003), indicating a role in organization of higher order structures. Analysis of α-crystallin C-terminal truncations in hereditary cataractous rat lens demonstrated that a hydrophilic C-terminal extension is essential for chaperone activity, suggesting increased C-terminal truncation of α-crystallins in mutant lenses and reduction in chaperone function accelerates cataract formation (Takeuchi et al., 2004). Single site mutations to the C-terminal extension also affect sHSPs. As an example, alternation of isoleucine residues in the conserved I/VXI/V motif, which has a propensity to participate in inter-subunit interactions, results in oligomeric dissociation and chaperone activity decline for bacterial and plant sHSPs (Studer et al., 2002; Pasta et al., 2004).

1.4 The brine shrimp, *Artemia franciscana*

The crustacean, *Artemia franciscana*, has a remarkable ability to tolerate environmental hardships. The organism inhabits aquatic environments of high salinity, subject to temperature extremes, desiccation, anoxia, and variable food reserves (Drinkwater and Clegg, 1991; Clegg et al., 2000; MacRae, 2003). Two developmental pathways, namely ovoviviparous and oviparous, allow *Artemia* to survive these extremes (Fig. 5). Embryos developing via the ovoviviparous pathway are released from females as free-swimming nauplii or instar I larvae (MacRae, 2003). Alternatively, embryos undergoing oviparous development are arrested at the gastrula stage and then encyst. The cysts, composed of approximately 4000 cells enclosed in a shell permeable only to volatile molecules, enter a
Figure 5. Life history of *Artemia franciscana*

Fertilized *Artemia* oocytes may undergo ovoviviparous development where free-swimming nauplii are released (left). Alternatively, oviparous development occurs where development stops at late gastrulae and embryos encyst (top-left), enter diapause and are released from females (top-middle). Diapause continues until cysts are activated by environmental stimuli such as cold and desiccation (top-middle). Activated, post-diapause cysts develop into nauplii. However, if unfavorable conditions prevail, growth is arrested and cysts become quiescent until favorable conditions occur (Liang and MacRae, 1999).
Figure 5
dormant state known as diapause, where the metabolic rate is too low to be detected experimentally (Jackson and Clegg, 1996; Clegg et al., 2000; MacRae, 2003). Diapause continues, even under favorable conditions, until cysts are activated by the appropriate environmental stimulus, including desiccation and cold (Clegg et al., 2000; MacRae, 2003). Upon activation, cysts resume development, provided that appropriate temperature, water content and oxygen are available (Drinkwater and Clegg, 1991; Clegg et al., 1999). However, cysts remain quiescent if unfavorable conditions prevail.

Among organisms which experience diapause, encysted *Artemia* embryos possess an outstanding capability to tolerate environmental insults (Viner and Clegg, 2001). For example, post-diapause cysts will survive fully hydrated at physiological temperature in the absence of oxygen for four years, and the anoxic cysts maintain integrity even in the apparent absence of nuclear transcription, protein synthesis and ongoing energy metabolism (Hand, 1998; Clegg et al., 1999; Liang and MacRae, 1999; van Breukelen et al., 2000; Warner and Clegg, 2001). Interestingly, this appears contrary to the biological axiom that, upon normal hydration and temperature, cell maintenance requires a constant and substantial free energy flow (Warner and Clegg, 2001). Cysts have developed a mechanism to survive by which intracellular macromolecules and organelles are protected against irreversible denaturation due to water removal, heat, anoxia and reduced metabolism. The sHSPs, which function as molecular chaperones and may be developmentally regulated, are ideal candidates for this purpose.

1.5 p26, a small heat shock protein from *Artemia franciscana*

Diapause-destined embryos synthesize large amounts of a novel sHSP, namely p26, which is thought to act as a molecular chaperone during diapause, anoxia and other
stresses (MacRae, 2003). Composed of 20.8 kDa subunits, p26 forms oligomers as large as 34 subunits with a molecular mass of approximately 700 kDa (Liang et al., 1997b; MacRae, 2003). This protein makes up approximately 10-15% of total non-yolk embryonic protein and is most abundant in the encysted embryo stage of the life cycle (Clegg et al., 1994; Clegg et al., 1995). Produced mainly in oviparously developing Artemia, p26 peaks in encysted embryos and remains constant during post-gasttrula development until just before the emergence of nauplii from cysts (Liang and MacRae, 1999; MacRae, 2003). p26 is almost entirely missing from second instar larvae, and is neither detectable nor inducible in subsequent life stages (Clegg et al., 1999; Liang and MacRae, 1999). Instead of being heat-inducible, p26 is developmentally regulated, as occurs for Hsp12.6 from Caenorhabditis elegans, Hsp17.6 from sunflower and multiple sHSPs from D. melanogaster (Morrow et al., 2000; MacRae, 2003). During diapause, p26 exists in nuclei and cytoplasm (Jackson and Clegg, 1996; Clegg et al., 2000). Upon diapause termination, most p26 resides in cytoplasm, although about half re-enters nuclei if activated embryos experience anoxia or thermal stress (Clegg et al., 1994; Clegg et al., 1999). Cytoplasm-to-nuclei translocation of p26 also occurs in vitro when the pH of homogenates is reduced, and this is reversed by alkaline pH (Clegg et al., 1995; Clegg et al., 1999). The observation of p26 in discrete intranuclear compartments of Artemia cysts implies roles for the protein in specific processes (MacRae, 2003), perhaps including the inhibition of DNA replication which characterizes cyst development.

p26 exhibits molecular chaperone activity in vitro, protecting citrate synthase against heat induced inactivation and chaperoning Artemia tubulin which exposes hydrophobic amino acid residues as it ages (Liang et al., 1997b; Viner and Clegg, 2001; Day et al.,
2003). Additionally, p26 is thought to protect intracellular proteins of encysted *Artemia* from stress-induced denaturation (Liang and MacRae, 1999; MacRae, 2003). p26 also confers thermotolerance on transformed bacteria and enhances stress resistance in transgenic *D. melanogaster* (Crack *et al.*, 2002; Malaviarachchi, 2002), further indicating that p26 functions as a molecular chaperone *in vivo.*

As a member of the sHSP family, p26 possesses a conserved α-crystallin domain, with secondary structure predicted to consist predominantly of β-pleated sheets (Liang *et al.*, 1997a), although how these sheets are organized was not investigated. Previous data showed that the C-terminus of p26 is more important than the N-terminus for conferring thermotolerance on *Escherichia coli* and formation of large oligomers is not a prerequisite for the chaperoning function of p26 (Crack *et al.*, 2002). When compared with other sHSPs, p26 has an N-terminus distinctly enriched in both glycine and arginine (MacRae, 2003), and a serine/threonine-enriched patch in the unusually long, polar and hydrophilic C-terminus. Such specific variances between p26 and other sHSPs suggest these regions are structurally and functionally important. In addition to these unique characteristics, p26 possesses highly conserved sites within the core α-crystallin domain.

The purpose of the work described in this thesis was to analyze p26 structure and function. The results enhance our appreciation of *Artemia* development and have important implications for understanding the functional mechanisms of p26, while providing a model for the study of other sHSPs, many of which are involved in human diseases.
II. Materials and Methods

2.1 Site-directed mutagenesis of p26

Truncations, single amino acid substitutions and internal deletions of p26 cDNAs were generated by site-directed mutagenesis with the QuikChange™ Site-directed Mutagenesis kit (Stratagene, La Jolla, CA), using pRSET.C-p26-3-6-3 as template (Liang et al., 1997a), and designated primers (Tables 1 & 2). Fifty μl PCR reaction mixtures included 5 μl of 10 x reaction buffer, 1 μl (50 ng) of DNA extracted with the Wizard Plus SV Miniprep DNA purification kit (Promega, Madison, WI) following the manufacturer's instructions, 1 μl (125 ng) of sense primer, 1 μl (125 ng) of anti-sense primer, 1 μl of dNTP mix and 41 μl of ddH₂O. One μl of PfuTurbo DNA polymerase (2.5 U/μl) was added and sample reactions were overlayed with 30 μl of mineral oil. Mixtures were incubated for 30 sec at 95 °C prior to multiple cycles of 30 sec at 95 °C, 1 min at 55 °C, 8 min at 68 °C. To produce truncated or internally deleted p26 cDNAs, 18 cycles were employed, for single amino acid substitutions 12 cycles sufficed. A PTC-100 programmable thermal controller (MJ Research, Inc, Watertown, MA) was employed. After cycling, the reactions were placed on ice for 2 min, then 1 μl of the restriction enzyme Dpn I (10 U/μl) was added to each amplification reaction prior to incubation at 37 °C for 1 h. For transformation, 50 μl of gently thawed E. coli XL1-blue supercompetent cells were placed in prechilled Falcon 2059 polypropylene tubes, 1 μl of the Dpn I-treated DNAs was added, and the mixtures were incubated on ice for 30 min. The transformation mixtures were heated for 45 sec at 42 °C, placed on ice for 2 min, and 0.5 ml of NZY⁺ broth preheated to 42 °C was added. The transformation mixtures were
Table 1. Truncations of p26 cDNA

<table>
<thead>
<tr>
<th>p26 variants</th>
<th>Amino acid residues deleted</th>
<th>Primer sequences</th>
</tr>
</thead>
<tbody>
<tr>
<td>p26-NA36</td>
<td>1-36</td>
<td>5'-CTCGTGCCGATCAAAATGCCCTTCCGGAGAAG-3'</td>
</tr>
<tr>
<td></td>
<td></td>
<td>5'-CTTCTCCGGAAGGGCATTTTGATCGGCACGAG-3'</td>
</tr>
<tr>
<td>p26-NA60</td>
<td>1-60</td>
<td>5'-CTCGTGCCGATCAAAATGTCCTTGAGGGACACAGC-3'</td>
</tr>
<tr>
<td></td>
<td></td>
<td>5'-GCTGTGTCCCTCAAGGACATTTTGATCGGCACGAG-3'</td>
</tr>
<tr>
<td>p26-CA40</td>
<td>153-192</td>
<td>5'-GCTCAACCAACAGAACGTTAATCTGCAATTC-3'</td>
</tr>
<tr>
<td></td>
<td></td>
<td>5'-GAATGCAAGATTAACTGGTTGGAAGCAGCTAG-3'</td>
</tr>
<tr>
<td>p26-CA10</td>
<td>183-192</td>
<td>5'-GCTAGTTCAACACTCCATAATTCGCAATTC-3'</td>
</tr>
<tr>
<td></td>
<td></td>
<td>5'-GAATGCAAGATTATGGAGTTGAACAGCTAG-3'</td>
</tr>
<tr>
<td>p26-alpha</td>
<td>1-60</td>
<td>5'-CTCGTGCCGATCAAAATGTCCTTGAGGGACACAGC-3'</td>
</tr>
<tr>
<td></td>
<td></td>
<td>5'-GCTGTGTCCCTCAAGGACATTTTGATCGGCACGAG-3'</td>
</tr>
<tr>
<td></td>
<td>153-192</td>
<td>5'-GCTCAACCAACAGAACGTTAATCTGCAATTC-3'</td>
</tr>
<tr>
<td></td>
<td></td>
<td>5'-GAATGCAAGATTAACTGGTTGGAAGCAGCTAG-3'</td>
</tr>
</tbody>
</table>

Truncations of p26 cDNA were generated by site-directed mutagenesis using primers listed above in the right column. The name of each p26 variant, either cDNA or protein, is indicated in the left column and the positions of deleted amino acids are in the middle column.
Table 2. Single amino acid substitutions and internal deletions of p26 cDNA

<table>
<thead>
<tr>
<th>Regions of p26</th>
<th>Mutations</th>
<th>Primer sequences</th>
</tr>
</thead>
<tbody>
<tr>
<td>N-terminal region</td>
<td>W17R 5'-GGTATGACTGACCCAAAGGTCATTCATCGATTTGG-3' 5'-CCAAATGAGTCACCTTGGGTATCGAACC-3'</td>
<td></td>
</tr>
<tr>
<td></td>
<td>D19A 5'-GACCCATGTCCTCCTCATTGGATTGGG-3' 5'-GCAAATCCAAATTGGGACGACATCG-3'</td>
<td></td>
</tr>
<tr>
<td></td>
<td>P20A 5'-GCCAACAGCTGATTGTCATGGATTGGGAGCTG-3' 5'-GACCAAAACTCCAGACGATAATCAGG-3'</td>
<td></td>
</tr>
<tr>
<td></td>
<td>F21A 5'-GACCCATTGCCATCGCTGTCGATTTGGGAGCTG-3' 5'-GACCAAAAATTCAGCTGATGCAATCCACC-3'</td>
<td></td>
</tr>
<tr>
<td></td>
<td>G22A 5'-GAGTCATGTCCTCATTGGCGATTTGGGAGCTG-3' 5'-GCCACAAATGCAATTGGGATCACG-3'</td>
<td></td>
</tr>
<tr>
<td></td>
<td>&quot;G&quot; deletion 5'-GCCAACATTCCATGATGGGACCTGGATATTGAC-3' 5'-GCCAACATGCAATTGGGATCACG-3'</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(8-29:22) 5'-GTCATCTAACATTGGATCTGCATGGGACCTGGATATTGAC-3' 5'-GTCATCTAACATTGGATCTG-3'</td>
<td></td>
</tr>
<tr>
<td></td>
<td>&quot;R&quot; deletion 5'-GGACCTTGTATGGACCTGACGACTACC-3' 5'-GGTATCTGGGACCCATATACTGCGACG-3'</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(36-45:10) 5'-GGACCTTGTATGGACCTGACGACTACC-3' 5'-GGTATCTGGGACCCATATACTGCGACG-3'</td>
<td></td>
</tr>
</tbody>
</table>

| α-crystallin domain | D67A 5'-GGGACACAGCTGCTGAAATTCCAGTATACC-3' 5'-CTGAACATGGAAATTGACGCTGTCCTTCC-3' | |
|                    | F69R 5'-GGGACACAGCTGCTGAAATTCCAGTATACC-3' 5'-CTGAACATGGAAATTGACGCTGTCCTTCC-3' | |
|                    | R110G 5'-GGGACACAGCTGCTGAAATTCCAGTATACC-3' 5'-CTGAACATGGAAATTGACGCTGTCCTTCC-3' | |
|                    | F112R 5'-GGGACACAGCTGCTGAAATTCCAGTATACC-3' 5'-CTGAACATGGAAATTGACGCTGTCCTTCC-3' | |
|                    | R114A 5'-GGGACACAGCTGCTGAAATTCCAGTATACC-3' 5'-CTGAACATGGAAATTGACGCTGTCCTTCC-3' | |
|                    | Y116D 5'-GGGACACAGCTGCTGAAATTCCAGTATACC-3' 5'-CTGAACATGGAAATTGACGCTGTCCTTCC-3' | |
|                    | G135A 5'-GGGACACAGCTGCTGAAATTCCAGTATACC-3' 5'-CTGAACATGGAAATTGACGCTGTCCTTCC-3' | |
|                    | L137R 5'-GGGACACAGCTGCTGAAATTCCAGTATACC-3' 5'-CTGAACATGGAAATTGACGCTGTCCTTCC-3' | |
|                    | T138A 5'-GGGACACAGCTGCTGAAATTCCAGTATACC-3' 5'-CTGAACATGGAAATTGACGCTGTCCTTCC-3' | |

C-terminal extension

| "TS" deletion | 5'-GGTAATGCAAGGGGGACGACGACGAGCTGTCCTTCC-3' 5'-GCTCCACCCAGCAGACGAGCTGTCCTTCC-3' | |
| (169-186:18) | "GGAA" deletion 5'-GCTCCACCCAGCAGACGAGCTGTCCTTCC-3' 5'-GCTCCACCCAGCAGACGAGCTGTCCTTCC-3' | |
| "AA" deletion | 5'-GCTCCACCCAGCAGACGAGCTGTCCTTCC-3' 5'-GCTCCACCCAGCAGACGAGCTGTCCTTCC-3' | |

Single amino acid substitutions and internal deletions of p26 cDNA were generated by site-directed mutagenesis using the listed primers. The p26 regions which encompass mutations are indicated in the left column. Single amino acid substitutions are named
starting with the mutated amino acid, followed by its position, and then the replacing amino acid. “G” is the abbreviation for the modification where amino acid residues G8 to G29 are deleted; “R” is the internal deletion where residues R36 to R45 are lost; “TS” represents the internal deletion where residues T169 to T186 are missing; “GGA” is the deletion where the last four C-terminal residues of p26 are removed; “AA” refers to the deletion which lacks the last two residues of p26. In later work, only four single amino acid substitutions generated in the α-crystallin domain including R110G, F112R, R114A, Y116D, and three internal deletions “G”, “R”, “TS” of p26 were selected for further studies.
incubated at 37 °C for 1 h with shaking at 225-250 rpm, after which 250 μl of each transformation mixture was spread on agar plates containing 50 μg/ml ampicillin (Sigma, St. Louis, MO) and incubated at 37 °C. Bacterial clones were selected randomly and propagated for DNA mini-preparation, with the identity of each mutant verified by DNA sequencing (DNA Sequencing Facility, Center for Applied Genomics, Hospital for Sick Children, Toronto, ON).

2.2 Cloning of p26 cDNAs in eukaryotic and prokaryotic expression vectors

For cloning in eukaryotic and prokaryotic expression vectors, p26 cDNA-containing pRSET.C plasmids were digested with Bam HI and Xho I, and electrophoresed in agarose gels from which inserts were recovered with the GFX™ PCR DNA and Gel Band purification kit (Amersham Biosciences, Piscataway, NJ). p26 cDNA, either full length or containing single amino acid substitutions and internal deletions was inserted into pcDNA4/TO/myc-His.A (Invitrogen, San Diego, CA), a eukaryotic expression vector which was pre-linearized with the same restriction enzymes, with T4 DNA ligase (MBI Fermentas, Hanover, MD) and constructs were transformed into E. coli strain DH5α (Invitrogen, Carlsbad, CA). These p26 cDNAs were also subcloned into pPROTet.E233 (Clontech Laboratories, Inc, Palo Alto, CA), a His-tag containing prokaryotic expression vector, between the Bam HI and Xba I sites. Full length and truncated forms of p26 cDNAs were cloned in pRSET.C and recovered by digestion with Bam HI and Xho I, before subsequently cloned into pSecCMV (Invitrogen, San Diego, CA) for expression in mammalian cells. p26-full which contained the full length p26 polypeptide, p26-NΔ60 where residues 1-60 were removed, and p26-alpha which comprised only the α-crystallin domain of p26 were also excised with Bam HI and Xba I and cloned into the eukaryotic
vector pcDNA4/His.A (Invitrogen, San Diego, CA) for expression as His-tagged fusion proteins. The vectors used in these experiments are listed in Table 3. After cloning, all inserts were sized by restriction digestion followed by electrophoresis in 1% agarose gels, and PCR fidelity was verified by DNA sequencing.

2.3 p26 cDNA expression in transiently transfected COS-1 cells

COS-1 cells were maintained at 37 °C under 5% CO2 in Dulbecco's Modified Eagles Medium (DMEM) (Gibco, Burlington, ON) containing 10% fetal bovine serum (FBS, Gibco) and 2% penicillin/streptomycin (Gibco). For small scale transfections, 60 μl of DMEM lacking serum and antibiotics, 1 μg of purified plasmid DNA and 3 μl of SuperFect™ (Qiagen, Mississauga, ON) were mixed in 5 ml polypropylene tubes. The transfection mixtures were incubated at room temperature for 15 min, placed in individual wells of 6-well plates (Becton Dickinson Labware, Franklin Lakes, NJ) containing COS-1 cells grown to 60% confluency and incubated overnight at 37 °C. For large scale transfection, T75 culture flasks (Becton Dickinson Labware) were employed, with the amount of each transfection reagent 6 times that used for 6-well plates. Cells were trypsinized with trypsin-EDTA (Gibco) 24 h after transfection, centrifuged at 1,500 x g for 5 min and washed with 1 ml of phosphate buffered saline (PBS) (140 mM NaCl, 2.7 mM KCl, 8.0 mM Na2HPO4, 1.5 mM KH2PO4, pH 7.4). Cells were collected by centrifugation at 1,500 x g for 5 min, resuspended in 50 μl of lysis buffer consisting of 50 mM Tris-HCl, pH 7.8, 150 mM NaCl, 0.5% Nonidet P-40, 1 mM phenylmethylsulfonyl fluoride (PMSF) (Sigma), 1 μg/ml pepstatin A (Sigma) and 1 μg/ml leupeptin (Sigma) and incubated on ice for 20 min. Cell lysates were centrifuged at
Table 3. Vectors used for cloning and expression of p26 cDNA

<table>
<thead>
<tr>
<th>Types of p26 cDNA mutations</th>
<th>Prokaryotic expression vectors and restriction sites (His-tagged)</th>
<th>Eukaryotic expression vectors and restriction sites</th>
</tr>
</thead>
<tbody>
<tr>
<td>Truncations</td>
<td>pRSET.C</td>
<td>pSecCMV</td>
</tr>
<tr>
<td></td>
<td><em>Bam HI</em> and <em>Xho I</em></td>
<td><em>Bam HI</em> and <em>Xba I</em></td>
</tr>
<tr>
<td>Single amino acid substitutions</td>
<td>pPROTet.E233</td>
<td>pcDNA4/TO/myc-His.A</td>
</tr>
<tr>
<td></td>
<td><em>Bam HI</em> and <em>Xba I</em></td>
<td><em>Bam HI</em> and <em>Xho I</em></td>
</tr>
<tr>
<td>Internal deletions</td>
<td>pPROTet.E233</td>
<td>pcDNA4/TO/myc-His.A</td>
</tr>
<tr>
<td></td>
<td><em>Bam HI</em> and <em>Xba I</em></td>
<td><em>Bam HI</em> and <em>Xho I</em></td>
</tr>
</tbody>
</table>

The vectors used for cloning and expression of p26 cDNAs in prokaryotic and eukaryotic cells are listed. The types of p26 cDNA variants are indicated in the left column. Vector names and restriction sites used in cloning are indicated in the remaining columns. "His-tagged" means the vector contains either a 6×His or a 6×HN tag.
10,000 × g for 10 min, supernatants were transferred to fresh tubes and protein concentrations determined by the Bradford assay (Bio-Rad, Hercules, CA).

2.4 SDS-polyacrylamide gel electrophoresis (SDS-PAGE), Western blotting and immunodetection of proteins

Equal amounts of cell-free extracts from COS-1 cells expressing p26 variants were electrophoresed in 12.5% SDS-polyacrylamide gels which were either stained with Coomassie Brilliant Blue R-250 (Sigma) or the proteins were transferred to nitrocellulose membranes (Bio-Rad, Hercules, CA). The membranes were rinsed briefly with TBS, and then stained with 2% Ponceau-S (Sigma) in 3% trichloroacetic acid (TCA) to check transfer efficiency. The membranes were incubated in 5% low fat Carnation milk powder in TBS-Tween for 45 min, followed by incubation for 30 min at room temperature with either anti-p26 antibody (Liang et al., 1997a) diluted 1:10000 in HST (10 mM Tris, pH 7.4, 1 M NaCl, 0.5% (v/v) Tween 20) or Omni-probe (Santa Cruz Biotechnology, Inc., Santa Cruz, CA), an antibody recognizing the (His)6/Xpress epitope region encoded by pRSET.C and diluted 1:2000 in HST buffer. The blots were washed twice for 3 min in TBS-Tween (10 mM Tris, pH 7.4, 0.14 M NaCl, 0.1% (v/v) Tween 20), twice for 3 min in HST, and once for 3 min in TBS-Tween, before incubation for 30 min with horseradish peroxidase (HRP)-conjugated goat anti-rabbit IgG antibody diluted 1:10000 in HST or HRP-conjugated goat anti-mouse IgG antibody (Jackson ImmunoResearch) diluted 1:5000 in HST. The immunoconjugates were detected with the Western Lightning Enhanced Chemiluminescence (ECL) Reagent Plus (PerkinElmer Life Sciences, Boston, MA) using Fuji medical X-ray film (Fuji Photo Film Co., Ltd, Tokyo).
2.5 Immunolocalization of p26 in transfected COS-1 cells

Twenty-four h after transfection, cells cultured on cover-slips were fixed for 20 min at room temperature in 4% paraformaldehyde, rinsed twice with PBS and permeabilized for 5 min in PBS containing 0.2% (v/v) Triton X-100 (PBS-Triton). Each cover-slip received 35 μl of anti-p26 antibody diluted 1:500 in PBS-Triton and was incubated for 30 min. The cells were washed three times with PBS-Triton, then incubated for 20 min at room temperature with 35 μl of fluorescein isothiocyanate (FITC)-conjugated goat anti-rabbit IgG antibody (Jackson ImmunoResearch) diluted 1:500 in PBS-Triton. The cells were washed three times with PBS-Triton, incubated with RNaseA (Amersham Biosciences, Baie d’Urfe, Quebec) at 5 mg/ml for 10 min and stained with propidium iodide (Sigma) at 0.4 mg/ml for 2 min. Cover-slips were rinsed with PBS-Triton, PBS, and distilled water before being placed on slides with Vectashield™ mounting medium (Vector Laboratories, Burlingame, CA). Cells expressing His-tagged p26 fusion proteins were stained with Omni-probe diluted in PBS-Triton, followed by FITC-conjugated goat anti-mouse IgG antibody (Jackson ImmunoResearch). Immunofluorescently stained cells were examined with a Wild Leitz Aristoplan microscope (Wild Leitz Ltd, Heerbrugg, Switzerland) and a Zeiss 410 inverted confocal laser scanning microscope (Carl Zeiss, Thornwood, NY).

2.6 p26 oligomer formation in COS-1 cells

Transfected COS-1 cells grown to confluency in T75 culture flasks and collected as described above were incubated on ice for 20 min in lysis buffer (50 mM Tris-HCl, pH 7.8, 150 mM NaCl, 1% (v/v) Nonidet P-40,) containing 1 mM PMSF (Sigma), 1 μg/ml pepstatin A (Sigma) and 1 μg/ml leupeptin (Sigma), before centrifugation at 10,000 × g
for 10 min. Sucrose gradients of 10-50% were prepared by layering 5 ml of 10% (w/v) sucrose on 5 ml of 50% (w/v) sucrose in 0.1 M Tris/glycine buffer, pH 7.4, in 12 ml tubes and centrifuging at 200,000 × g for 3 h at 15 °C. Equal amounts of protein extract from transfected COS-1 cells were applied to individual gradients and centrifuged at 200,000 × g for 21 h at 4 °C followed by collection of 0.8 ml fractions. Fifteen µl from each fraction was electrophoresed in 12.5% SDS polyacrylamide gels, blotted to nitrocellulose, and probed with either anti-p26 antibody or Omni-probe as described previously. Molecular mass standards, namely alpha-lactalbumin (14.2 kDa), carbonic anhydrase (29 kDa), bovine serum albumin (66 kDa), alcohol dehydrogenase (150 kDa), apoferritin (443 kDa), and thyroglobulin (669 kDa) (Sigma) were centrifuged separately and their locations determined by measuring the A₂₈₀ of gradient fractions.

2.7 Expression of p26 cDNA in E. coli

pRSET.C vectors containing full length and truncated p26 cDNAs were transferred into E. coli BL21(DE3)pLysS (Novagen, Madison, WI). For expression of p26, 2 ml of LB medium containing 50 µg/ml ampicillin was inoculated with a single E. coli colony and incubated overnight with shaking at 37 °C. The cultures were added to 50 ml of LB medium containing 50 µg/ml ampicillin in 125 ml Erlenmeyer flasks and incubated with shaking at 37°C until the A₆₀₀ of cultures reached 0.5-0.7. Isopropyl-1-thio-β-D-galactopyranoside (IPTG) (Sigma) was then added to a final concentration of 1 mM and incubation continued for 6 h. The cultures were cooled on ice for 5 min, and centrifuged at 5,000 × g for 5 min at 4 °C. The cell pellets were washed once with 1 × Extraction/Wash buffer (50 mM Na₃PO₄, 300 mM NaCl, pH 7.5) and resuspended in 4 ml of the same buffer, prior to addition of 100 µg/ml lysozyme (Sigma), 1 mM PMSF
(Sigma), 1 μg/ml pepstatin A (Sigma) and 1 μg/ml leupeptin (Sigma). After multiple freeze-thaw cycles, the mixtures were incubated at 30 °C for 15 min, cooled on ice and sonicated three times for 10 sec using a Branson Sonifier™ 150 (Branson Ultrasonics, Danbury, CT) at medium setting with intermittent cooling on ice for 30 sec. The sonicated cells were centrifuged at 12,000 × g for 20 min at 4 °C, the supernatants recovered and protein concentrations ascertained with the Bradford assay kit (Bio-Rad, Hercules, CA).

For the expression of p26 cDNA containing single amino acid substitutions and internal deletions, recombinant vectors constructed with pPROTe.E233 were transformed into E. coli BL21PRO (Clontech Laboratories, Inc., Mississauga, ON). Single colonies were picked from LB plates containing 50 μg/ml spectinomycin (Sigma) and 34 μg/ml chloramphenicol (Sigma), inoculated in 2 ml of LB medium and incubated overnight at 37 °C. Fifty ml of LB medium containing spectinomycin/chloramphenicol were inoculated with each overnight culture, and incubated with shaking at 37 °C until the A₆₀₀ reached 0.5-0.7. Anhydrotetracycline (aTc) (Clontech Laboratories) was added to a final concentration of 100 ng/ml, and the incubation continued for 6 h. Cell-free extracts were prepared as just described.

2.8 Purification of p26

Five ml aliquots of thoroughly resuspended BD TALON resin (BD Biosciences Clontech, Mississauga, ON) were transferred to sterile 50 ml tubes, centrifuged at 700 × g for 2 min, and the pellets were resuspended in 10 bed volumes of 1 × Extraction/Wash buffer (50 mM Na₂PO₄, 300 mM NaCl, 10 mM Imidazole, pH 7.5) prior to brief agitation. After re-centrifugation at 700 × g for 2 min, the resin was pelleted, and the
wash was repeated. Bacterial cell-free extracts were added to the resin, and the mixtures were agitated gently at room temperature for 20 min on a platform shaker, before centrifugation at 700 × g for 5 min. The supernatants were removed carefully, the resin was washed with gentle agitation in 10 bed volumes of 1 × Extraction/Wash buffer for 10 min and centrifuged at 700 × g for 5 min, after which the wash was repeated. One bed volume of the 1 × Extraction/Wash buffer was added, the resuspended resin was transferred to 2 ml BD TALON™ affinity gravity-flow columns (BD Biosciences Clontech) and allowed to settle out of suspension. The buffer was drained and columns were washed once with 5 bed volumes of 1 × Extraction/Wash buffer. The polyhistidine-tagged proteins were eluted by adding 5 bed volumes of 1 × Elution buffer (50 mM Na₃PO₄, 300 mM NaCl, 150 mM Imidazole, pH 7.5) to columns, 500-μl fractions were collected and the A₂₈₀ of each fraction measured. The purified samples were dialyzed for 4 h at room temperature with one change of buffer, then overnight at 4 °C against 10 mM NaH₂PO₄, pH 7.1. Dialyzed samples were concentrated in CentriprepYM-10 centrifugal filter devices (Amicon Bioseparations, Billerica, MA) following manufacturer’s instructions.

2.9 Oligomerization of p26

Half-ml samples of p26-containing cell-free extracts from either transfected COS-1 cells or transformed bacteria, and purified samples of p26, were applied to continuous 10-50% (w/v) sucrose gradients, which were prepared with 5 ml of 10% (w/v) sucrose and 5 ml of 50% (w/v) sucrose in 0.1 M Tris/Glycine buffer, pH 7.4, using a gradient maker (MRA, Boston, MA). The gradients were centrifuged at 200,000 × g for 12 h at 4 °C in an SW41-TI Beckman rotor. Tube bottoms were punctured with a 21-gauge needle after
centrifugation, 0.8 ml fractions were collected and 15 μl from each fraction was
electrophoresed in 12.5% SDS-polyacrylamide gels before blotting to nitrocellulose for
immunodetection. For purified truncated versions of p26, the A_{280} of each gradient
fraction was measured with a SPECTRAmax PLUS microplate spectrophotometer
(Molecular Devices, Sunnyvale, CA). The number of p26 monomers per oligomer was
calculated using a p26 molecular mass of 20.8 kDa as determined by GENERUNNER
(version 3.05, Hastings Software, Inc.) with corrections for protein modifications as
necessary. Alpha-lactalbumin (14.2 kDa), carbonic anhydrase (29 kDa), bovine serum
albumin (66 kDa), alcohol dehydrogenase (150 kDa), apoferritin (443 kDa), and
thyroglobulin (669 kDa) (Sigma) were centrifuged separately and their locations
determined in sucrose gradients by measuring the A_{280} of each gradient fraction.

2.10 p26-induced thermotolerance in E. coli

To examine thermotolerance induction in E. coli expressing p26, 2 ml of LB medium
containing either ampicillin/IPTG or spectinomycin/chloramphenicol/anhidrotetracycline
was inoculated with single bacterial colonies and incubated overnight at 37 °C with
shaking. Upon 1:10 dilution in Falcon 2059 polypropylene tubes (Becton Dickinson
Labware, Franklin Lakes, NJ), cultures were transferred to a 54 °C water bath and 100 μl
samples were removed at 0, 15, 30, 45 and 60 min. Samples were diluted in cold LB
medium and kept on ice before plated on LB agar and incubated at 37 °C overnight. All
colony-forming units were counted, and data from three separate heat shock experiments
were pooled in order to assess survival. Between data groups, two-sample t-tests were
performed at a confidence level of 95.0% with a statistical software MINITAB 14.12.0
(Minitab Inc., State College, PA) to evaluate the significance of difference which was
accepted at the $P < 0.05$ level. To ensure that bacteria contained p26, small amounts of
cells were collected before heating, homogenized, and electrophoresed in 12.5% SDS-
polyacrylamide gels, followed by transfer to nitrocellulose membranes and
immunodetection of p26.

2.11 In vitro chaperone activity of p26

2.11.1 Citrate synthase aggregation

A 0.178 mM stock solution of dimeric citrate synthase (Sigma) was diluted with 40
mM HEPES/KOH buffer, pH 7.5 to a final concentration of 150 nM in 1.0 ml cuvettes
and heated at 43 °C with 37.5 nM, 75 nM, 150 nM, 300 nM, 600 nM, 1.2 μM and 2.4 μM
of purified, bacterially-produced p26. p26 molarity in reaction mixtures was based on
monomeric molecular masses of p26 variants. Heat induced aggregation of citrate
synthase was monitored by measuring solution turbidity at 360 nm with a SPECTRAmax
PLUS spectrophotometer (Molecular Devices) every 2 min for 1 h. Bovine serum
albumin (BSA) (Sigma) and immunoglobulin G (IgG) (Sigma) were used at final
concentrations of 600 nM or 1.2 μM to evaluate non-specific protection for citrate
synthase.

2.11.2 Citrate synthase inactivation

Citrate synthase was incubated in the presence and absence of p26 at 43 °C, small
aliquots of the enzyme were withdrawn at various time points during heat denaturation
and enzyme activity was measured. Reaction mixtures contained 940 μl of TE (50 mM
Tris-HCl, pH 7.5, 2 mM EDTA), 10 μl of 10 mM oxaloacetic acid (OAA) (Sigma), 10 μl
of 10 mM 5,5'-Dithiobis (2-nitrobenzoic acid) (DTNB) (Sigma) and 30 μl of 5 mM
acetyl-CoA (Sigma). The reaction, initiated by adding 10 μl of 150 nM citrate synthase,
was monitored at 25 °C as an increase in absorption at 412 nm which was determined with a SPECTRAmax PLUS spectrophotometer (Molecular Devices).

2.11.3 Insulin aggregation

In 300 µl reaction systems (10 mM phosphate buffer, 100 mM NaCl, pH 7.4) prepared in a 96-well plate, insulin (Sigma) was dissolved to a final concentration of 0.023 mg/ml (4.0 µM), while p26 was added to final concentrations of 0.05 µM, 0.1 µM, 0.2 µM, 0.4 µM, 0.8 µM, 1.6 µM and 3.2 µM. Dithiothreitol (DTT) (Sigma) was added to a final concentration of 20 mM in each well and the aggregation extent was measured at 25 °C by monitoring light scattering at 400 nm in a SPECTRAmax PLUS spectrophotometer (Molecular Devices) for 30 min at 2 min intervals.

2.12 Far-UV circular dichroism measurements

Far-UV circular dichroism (CD) spectra were recorded over 180-260 nm in a JASCO J-810/A001960750 spectropolarimeter (Japan Spectroscopic, Tokyo, Japan). At 25 °C, a 0.1 cm path length quartz cuvette containing purified p26 at 0.2 mg/ml in 10 mM NaH₂PO₄, pH 7.1, was employed, and an average of 3 scans was taken for each spectrum with a bandwidth of 2 nm, with all scans corrected for buffer and smoothed to eliminate background noise. Secondary structure parameters were calculated with the CDNN v2.1 deconvolution program (Martin-Luther-Universität Halle-Wittenberg, Germany).

2.13 Intrinsic tryptophan fluorescence and ANS-binding measurements

Purified p26 was diluted in 10 mM NaH₂PO₄, pH 7.1 to a final concentration of 0.06 mg/ml and fluorescence spectra were measured at 25 °C in 96-well plates with a SPECTRAmax GEMINIXS fluorescence spectrophotometer (Molecular Devices). The emission wavelength was set at 340 nm with a 2 nm band pass, and fluorescence
excitation was detected over a range of 250-310 nm to determine the maximal absorbance wavelength. Then, the excitation wavelength was set to 280 nm with a 2 nm band pass, and fluorescence emission was detected over a range of 310-400 nm. Each spectrum was recorded in duplicate using two independent sample preparations.

Two µl of 8.0 mM ANS (Molecular Probes, Eugene, OR) stock prepared in 10 mM NaH₂PO₄, pH 7.1 were added to 198 µl of each protein sample, and the mixtures were incubated for 5 min at either 25 °C or 43 °C. The excitation wavelength was set to 388 nm with a band pass of 8 nm, and the emission wavelength was set to 473 nm with a band pass of 8 nm. Measurements were taken with an AMINCO Bowman series z luminescence spectrometer (AMINCO, Rochester, NY) equipped with a thermostated circular water-bath, and with all conditions chosen to minimize inner filter effects. A standard curve was constructed with a series of ANS concentrations from 0.025 to 1.0 µM for future reference.

2.14 p26 purification from Artemia cysts

2.14.1 Preparation of Artemia cell-free extracts

Encysted Artemia embryos (cysts) (INVE Aquaculture, Inc., Ogden, UT) were hydrated in distilled water at 4 °C for 3 h, collected on a Buchner funnel, washed with cold distilled water and homogenized with a Retsch motorized mortar and pestle (Brinkman Instruments Canada, Rexdale, ON) in PIPES buffer (100 mM Pipes, 1 mM EGTA, 1 mM MgCl₂, pH 6.5). The homogenates were centrifuged at 16,000 × g for 10 min at 4 °C and the supernatants were passed through two layers of Miracloth (Calbiochem, La Jolla, CA) before centrifugation at 40,000 × g for 30 min at 4 °C. The upper two thirds of each
supernatant was transferred to a fresh tube and re-centrifuged for 20 min. The resulting supernatants were used immediately or frozen at −80 °C.

2.14.2 Purification of IgG from rabbit serum and p26 from Artemia cell-free extracts

Preparation of IgG from anti-p26 serum, coupling of purified IgG to protein A Sepharose™ CL-4B (Amersham Biosciences, Baie d’Urfe, Quebec), and purification of p26 were performed as described previously (Liang et al., 1997b).

2.15 Effect of pH on p26 oligomers

Cell-free extracts of encysted Artemia embryos were prepared as just described, except that the PIPES buffer for homogenization was at either pH 6.5 or 7.0. Either immediately after preparation, or following incubation at room temperature for 30 min, supernatants were centrifuged at 200,000 × g for 12 h at 4 °C in 10-50% continuous sucrose gradients at the pH used for homogenization. The gradients were fractionated and 10 μl of each sample was electrophoresed in SDS-polyacrylamide gels, blotted to nitrocellulose and probed with anti-p26 antibody as described above.

2.16 Effect of heat shock on p26 oligomers

Ten g of Artemia cysts in 500 ml of distilled H₂O were brought to 22 °C, then heated to 50 °C over 1 h with vigorous aeration in a Programmable/Digital Immersion Circulator (VWR, Mississauga, ON). Heat shocked and control cysts, the latter incubated on ice for 30 min, were homogenized separately by hand for 2 min in a chilled mortar and pestle in 35 ml of cold HPC (1 mM CaCl₂, 0.05 mM PIPES, 6.4% Hexylene glycol, pH 7.0) (Liang et al., 1997a), followed by one passage in a Dounce homogenizer, filtration through Miracloth, and centrifugation at 2,000 × g for 10 min at 4 °C to pellet nuclei. Supernatants were centrifuged at 40,000 × g for 30 min at 4 °C, and the upper two thirds
was placed in a fresh tube and re-centrifuged for 20 min. Supernatants were centrifuged in 10-50% continuous sucrose gradients, fractionated and analyzed on western blots with anti-p26 antibody after SDS-PAGE.

The nuclei-containing pellets from above were rinsed twice with 40 ml of HPC, resuspended in 17.5 ml of the same buffer, applied to 25 ml cushions of 75% (v/v) Percoll (Sigma) in 150 mM NaCl, 14 mM MgCl₂, 16 mM Tris, pH 7.0, and centrifuged at 16,000 × g for 30 min at 4 °C in a Beckman JS-13.1 swinging bucket rotor. Cloudy layers were transferred to fresh tubes, brought to 15 ml with HPC, placed on a 25 ml Percoll cushion and centrifuged as above. Nuclei were recovered, mixed with an equal volume of HPC, centrifuged at 16,000 x g for 30 min at 4 °C, and the pellets were suspended in 1.0 ml of HPC. Samples were stained with 0.4% DAPI (Molecular Probes, Eugene, OR) and examined microscopically. Nuclei from control and heat shocked embryos were lysed in treatment buffer, electrophoresed in 12.5% SDS polyacrylamide gels, blotted to nitrocellulose and probed with anti-p26 antibody. For sucrose density gradient centrifugation, extracts were prepared by suspending purified nuclei in 1.0 ml of ice cold extraction buffer (20 mM HEPES/KOH, 2.5 mM MgCl₂, 0.42 M NaCl, 25% (v/v) glycerol, 0.2 mM EDTA, 0.5 mM DTT, pH 7.9) containing 1 mM PMSF, 1 µg/ml pepstatin A and 1 µg/ml leupeptin, and sonicating three times for 10 sec using a Branson Sonifier™ 150 at medium setting with intermittent cooling on ice for 30 sec. The sonicated samples were incubated on ice for 30 min, and then centrifuged at 25,000 × g for 30 min at 4 °C with supernatants collected. Nuclear extracts were centrifuged in 10-50% continuous sucrose gradients at 200,000 × g for 12 h at 4 °C. The gradients were
fractionated and samples were electrophoresed in SDS polyacrylamide gels, blotted to nitrocellulose and probed with antibody to localize p26.

2.17 Modeling of p26 structure

To model the three-dimensional structure of p26, the Swiss-Model Protein Modeling Server (version 36.0003, Biozentrum University Basel, Basel and Swiss institute of Bioinformatics, Geneva, Switzerland and GlaxoSmithKline R&D S.A., Raleigh, NC, US) (Guex and Peitsch, 1997; Guex et al., 1999; Schwede et al., 2003) was applied. Wheat Ta Hsp16.9 (ExPDB entry code: 1GME) was used as a template for sequence alignment when the “alignment interface” function of Swiss-Model was employed, with the resulting alignment as a guide for modeling. The structure of p26 was modeled encompassing A2–P158 corresponding to S2–G151 of Ta Hsp16.9, and was shown in a monomer, with partial sequence of p26 C-terminal (A159–A192) deleted for alignment optimization. The returned three-dimensional model was improved with the “improve fit” function of Swiss-PdbViewer (version 3.7, GlaxoSmithKline R&D, Geneva, Switzerland) and sent for a second round modeling.
III. Results

3.1 p26 truncations

3.1.1 Generation and cloning of p26 cDNAs

Five truncated p26 cDNAs were generated by site-directed mutagenesis, with pRSET.C-p26-3-6-3 (Liang et al., 1997), a prokaryotic expression vector containing full length p26 cDNA as template. Six different cDNAs were obtained, including p26-full-His, encoding the full length p26 polypeptide; p26-NA36-His, residues 1-36 deleted; p26-NA60-His, residues 1-60 removed; p26-CΔ40-His, final forty residues eliminated; p26-CΔ10-His, lacking the last ten residues; p26-alpha-His, missing residues 1-60 and 153-192, thereby consisting only of the α-crystallin domain (Fig. 6). All p26 cDNAs generated by site-directed mutagenesis contained, in addition to the p26 coding sequence, an amino-terminal peptide of 13 residues encoded by p26-3-6-3, the His-tag and the Xpress epitope region. The presence of the His-tag is indicated by “−His” in the name of the p26 cDNA or protein. To examine the size of each cloned cDNA, minipreped pRSET.C plasmids as p26 constructs were double digested with Bam HI and Xho I, and the digestion products were electrophoresed in 1% agarose gels, revealing bands of insert DNAs at the expected positions (not shown). p26 constructs derived from pRSET.C-p26-3-6-3 were used for expression in prokaryotic systems. The p26 cDNAs recovered from previously used storage vector pCRII (Crack et al., 2002) were subcloned into pSecCMV for expression in COS-1 cells. p26-full, p26-NA60 and p26-alpha were also subcloned into pcDNA4/His.A for expression as His-tagged fusion proteins. All cloned p26 cDNA fragments were sized by electrophoresis in 1% agarose gels after restriction digestion of recombinant plasmids, with PCR fidelity confirmed by DNA sequencing (not shown).
Figure 6. Schematic representation and sequence of p26 cDNAs

a, truncations of p26 cDNAs generated by site-directed mutagenesis were cloned in the prokaryotic expression vector pRSET.C and the eukaryotic expression vectors pSecCMV and pcDNA4/His.A. The primers used for site-directed mutagenesis are listed in Table 1. The box labeled 6xHis is a His-tag-encoding sequence present in the vectors pRSET.C and pcDNA4/His.A; b, amino acid sequence of p26 (NCBI accession number AAB87967). The amino acid residues removed to generate p26-NΔ60 and p26-CΔ40 are shaded and those for p26-NΔ36 and p26-CΔ10 are shaded and bold.
Figure 6
3.1.2 p26 expression in bacteria

IPTG induction of transformed *E. coli* BL21(DE3)pLysS yielded polypeptides on western blots of the appropriate sizes that reacted with Omni-probe, although bands corresponding to p26 were not readily visible in Coomassie blue stained gels (Figs. 7a, b). With the exception of p26-NΔ60-His and p26-alpha-His, all of the polypeptides were recognized by anti-p26 antibody (not shown). Subsequent to purification on TALON™ affinity columns, single bands of the expected size appeared on Coomassie blue stained gels (Fig. 7c). These polypeptides reacted strongly with Omni-probe (Fig. 7d) indicating they were His-tagged p26, although purified p26-NΔ60-His and p26-alpha-His again failed to react with anti-p26 antibody. Protein bands were not detected on blots containing extract from *E. coli* transformed with vector lacking p26 cDNA (Figs. 7a, b, lane 7).

3.1.3 p26 oligomer formation

The molecular mass of oligomers assembled from purified p26-full-His synthesized in *E. coli* BL21(DE3)pLysS ranged from 25.8 kDa to 600 kDa, with the peak at about 150 kDa, or 6 monomers per oligomer (Fig. 8, Table 4). The oligomers assembled with full length p26 were larger than those obtained with truncated proteins, although oligomers of p26-CΔ10-His were similar in mass. At the other extreme, p26-NΔ60-His and p26-alpha-His migrated predominately as monomers and dimers, although larger complexes occurred. Oligomers assembled from other p26 variants were intermediate in mass to those just described.
Figure 7. Purification of p26 synthesized in E. coli

Cell free extracts from transformed E. coli induced with IPTG were electrophoresed in SDS polyacrylamide gels and either stained with Coomassie blue (a) or blotted to nitrocellulose and reacted with Omni-probe (b). Proteins purified by affinity chromatography were electrophoresed in SDS polyacrylamide gels and either stained with Coomassie blue (c), or blotted to nitrocellulose and reacted with Omni-probe (d). The lanes contained: 1, p26-NΔ36-His; 2, p26-NΔ60-His; 3, p26-alpha-His; 4, p26-CΔ40-His; 5, p26-CΔ10-His; 6, p26-full-His; 7, vector lacking p26 cDNA. All lanes received 10 μl of sample. M, molecular mass markers of 97, 66, 45, 31, 21 and 14 kDa.
Figure 8. Oligomer formation by purified p26

Bacterially produced p26 purified to apparent homogeneity by affinity chromatography was centrifuged at 200,000 x g for 12 h at 4 °C on 10-50% sucrose gradients. The gradients were fractionated and the A$_{280}$ of each fraction was plotted against fraction number. The top of each gradient is to the right and fractions are numbered across the bottom. The molecular mass markers, alpha-lactalbumin, 14.2 kDa; carbonic anhydrase, 29 kDa; bovine serum albumin, 66 kDa; alcohol dehydrogenase, 150 kDa; apoferritin, 443 kDa; and thyroglobulin, 669 kDa are indicated by numbered arrows.
Figure 8
Table 4. Oligomerization of p26 purified from *E. coli* BL21(DE3)pLysS

<table>
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<th>p26 variant</th>
<th>Monomer mass (kDa)</th>
<th>Oligomer Mass range (kDa)</th>
<th>Oligomer Mass peak (kDa)</th>
<th>Monomer number (peak)</th>
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</table>

The molecular mass of oligomers produced by p26 purified from transformed *E. coli* BL21(DE3)pLysS was determined by reading the $A_{280}$ of samples obtained by fractionation of sucrose density gradients. Monomer mass refers to the molecular mass of p26 polypeptides. Oligomer mass range represents the smallest to largest oligomers observed while oligomer mass peak refers to the mass of the most prevalent oligomer. Monomer number peak refers to the number of subunits in the most prevalent oligomer.
3.1.4 Chaperone activity

Purified p26-full-His possessed the greatest chaperone activity and p26-alpha-His the least, as determined by heat-induced denaturation of citrate synthase, although all variants provided some protection (Fig. 9). At a chaperone to target molar ratio of 4:1 (p26 monomer to citrate synthase dimer), or 1:1.5 if the peak oligomer size of p26-full-His is used for comparison, citrate synthase denaturation was inhibited almost completely after 1 h at 43 °C (Fig. 9a). At a 1:4 molar ratio of p26-full-His to citrate synthase, or 1:24 (oligomer to dimer), purified p26 reduced the heat-induced turbidity increase by approximately 25% (Fig. 9e). In contrast, p26-alpha-His almost completely lost chaperone activity even at molar ratios of 4:1 (Fig. 9). p26-NAΔ60-His was marginally better than p26-alpha-His as a chaperone, followed by p26-NAΔ36-His and p26-CAΔ40-His which were similar to one another. p26-CAΔ10-His approached p26-full-His in chaperone potency especially at high concentrations. In contrast, BSA and IgG at 600 nM provided almost no protection at all upon heating of citrate synthase (Fig. 9f).

3.1.5 Synthesis and oligomerization of p26 in transfected mammalian cells

Except for p26-NAΔ60 and p26-alpha, which reacted exclusively with Omni-probe, protein extracts from COS-1 cells transfected with p26 cDNA-containing expression vectors yielded polypeptides of the expected size on western blots when probed with anti-p26 antibody (Fig. 10). Neither primary antibody gave protein bands on blots with extracts from COS-1 cells transfected with vectors lacking p26 cDNA (not shown). Full length p26 synthesized in COS-1 cells produced oligomers as large as 512 kDa and composed of up to 21 monomers, with monomer number essentially the same in the presence and absence of His-tag (Figs. 11a, b). Oligomers assembled with p26-CAΔ10
Figure 9. p26 chaperone activity

Bacterially produced p26 purified to apparent homogeneity was heated at 43 °C for 1 h with 150 nm citrate synthase and solution turbidity was measured at A_{360}. p26 was at final concentrations of: a, 600 nM; b, 300 nM; c, 150 nM; d, 75 nM; e, 37.5 nM. The curves are: 1, No p26; 2, p26-alpha-His; 3, p26-NA60-His; 4, p26-NA36-His; 5, p26-CA40-His; 6, p26-CA10-His; 7, p26-full-His and they occupy the same relative positions in panels a-e. Panel f contains 150 nM citrate synthase incubated in the absence of other proteins (1) but with either 600 nM BSA (2) or 600 nM IgG (3).
Figure 10. p26 synthesis in transfected COS-1 cells

Extracts prepared from COS-1 cells transfected with the pSecCMV eukaryotic expression vector containing p26 cDNAs were electrophoresed in SDS polyacrylamide gels and either stained with Coomassie blue (a) or transferred to nitrocellulose and stained with anti-p26 antibody followed by HRP-conjugated goat anti-rabbit IgG antibody (b). Each lane received 10 μl of extract from cells transfected with 1, p26-NΔ36; 2, p26-NΔ60; 3, p26-alpha; 4, p26-CΔ40; 5, p26-CΔ10; 6, p26-full. M, molecular mass markers of 97, 66, 45, 31, 21 and 14 kDa. Western blots containing extracts from COS-1 cells transfected with the pcDNA4/His.A expression vector containing p26 cDNAs and prepared as just described were stained with either anti-p26 antibody followed by HRP-conjugated goat anti-rabbit IgG antibody (c) or Omni-probe followed by HRP-conjugated goat anti-mouse IgG (d). Each lane in c and d received 10 μl of extract from cells transfected with 1, p26-full-His; 2, p26-NΔ60-His; p26-alpha-His.
Figure 11. p26 oligomer formation in transfected COS-1 cells

Extracts prepared from COS-1 cells transiently transfected with p26-containing vectors were centrifuged in 10-50% sucrose gradients and 15 μl samples from each fraction were electrophoresed in SDS polyacrylamide gels, transferred to nitrocellulose, and probed with either anti-p26 antibody followed by HRP-conjugated goat anti-rabbit IgG (a, c, d, e) or Omni-probe followed by HRP-conjugated goat anti-mouse IgG (b, f, g). The gradients contained, a, p26-full; b, p26-full-His; c, p26-Δ10; d, p26-Δ40; e, p26-Δ36; f, p26-Δ60-His; g, p26-alpha-His. The top of each gradient is to the right and fractions are numbered across the top. The positions of molecular mass markers, α-lactalbumin, 14.2 kDa; carbonic anhydrase, 29 kDa; bovine serum albumin, 66 kDa; alcohol dehydrogenase, 150 kDa; apoferritin, 443 kDa; and thyroglobulin, 669 kDa are indicated by numbered arrows.
were close in size to those produced from full length p26 (Fig. 11c), but removal of the complete C-terminal extension gave oligomers of a narrower size range and smaller mass (Fig. 11d). Oligomer mass became progressively smaller upon sequential removal of the N-terminus (Figs. 11e, f) with polypeptides encoded by p26-NΔ60-His present as monomers and dimers, as was true for p26-alpha-His (Fig. 11g), although the latter was more enriched in monomers. The properties of p26 oligomers produced in COS-1 cells are summarized in Table 5.

3.1.6 Intracellular localization of truncated p26 in COS-1 cells

In order to monitor p26 synthesis, transiently transfected COS-1 cells were stained with anti-p26 antibody or Omni-probe and propidium iodide, revealing that polypeptides encoded by p26-full and p26-full-His (Figs. 12a, b) and the C-terminal truncations, p26-CΔ40 (Fig. 12c) and p26-CΔ10 (Fig. 12d), localized predominantly to the cytoplasm. In contrast, upon N-terminal truncation p26 occurred in cytoplasm and nuclei, the latter shown by yellow staining (overlapping of red and green). p26-NΔ36 encoded polypeptides entered the nuclei of only some transfected cells (Figs. 12e, f), but polypeptides p26-NΔ60/p26-NΔ60-His lacking the N-terminus (Figs. 12g, h), or p26-alpha/p26-alpha-His composed of the α-crystallin domain (Figs. 12i, j), resided in the nuclei of all transfected cells, indicating that the His-tag had no effect on p26 localization. The results, in concert with analysis of oligomer formation, suggested that oligomer disassembly is responsible for p26 movement into nuclei of COS-1 cells, and by extrapolation, the nuclei of encysted Artemia embryos.
Table 5. Oligomerization of p26 synthesized in transfected COS-1 cells

<table>
<thead>
<tr>
<th>p26 variant</th>
<th>Monomer mass (kDa)</th>
<th>Oligomer Mass range (kDa)</th>
<th>peak (kDa)</th>
<th>Monomer number peak</th>
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<tr>
<td>p26-CΔ10</td>
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<td>20-438</td>
<td>100</td>
<td>5</td>
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</table>

The molecular mass of p26 oligomers was determined by immunopробing of samples electrophoresed in SDS polyacrylamide gels and transferred to western blots after sucrose gradient centrifugation. Monomer mass refers to the molecular mass of p26 polypeptides. Oligomer mass range represents the smallest to largest oligomers observed while oligomer mass peak refers to the mass of the most prevalent oligomer. Monomer number peak refers to the number of subunits in the most prevalent oligomer.
Figure 12. p26 intracellular localization in transfected COS-1 cells

COS-1 cells transfected with the p26 cDNA-containing vector pSecCMV were incubated with antibody to p26 followed by FITC-conjugated goat anti-rabbit IgG antibody (green) (a, c, d, e, f). Cells transfected with the p26 cDNA-containing vector pcDNA4/His.A were exposed to anti-p26 antibody followed by FITC-conjugated goat anti-rabbit IgG (green) (g,i) or to Omni-probe followed by FITC-conjugated goat anti-mouse IgG (green) (b, h, j). Nuclei were stained with propidium iodide (red) and samples were examined by confocal microscopy. a, p26-full; b, p26-full-His; c, p26-CAΔ40; d, p26-CAΔ10; e and f, p26-NAΔ36; g, p26-NAΔ60; h, p26-NAΔ60-His; i, p26-alpha; j, p26-alpha-His. The scale bar represents 50 μm and all figures are the same magnification.
3.1.7 Effects of pH and heat shock on oligomer size of p26 from *Artemia* embryos

p26 moves into the nuclei of *Artemia* embryos upon exposure to reduced pH *in vitro* and upon heat shock *in vivo* by an unknown mechanism that, as suggested by the previous results, entails oligomer mass reduction. However, the p26 oligomers obtained from *Artemia* cysts homogenized at either pH 6.5 or 7.0 were identical in mass, whether or not they were incubated for 30 min at RT before gradient centrifugation (Fig. 13a-d). p26 migrated into the nuclei of heat shocked *Artemia* embryos under the conditions used in this work (Figs. 14a, b), but the size of p26 oligomers in extracts from cysts (Figs. 14c, d) and nuclei (Figs. 14e, f) remained constantly similar. The p26 in nuclear extracts tended to smear upon electrophoresis and blotting, probably due to the presence of DNA.

3.2 Single amino acid substitutions of the p26 α-crystallin domain

3.2.1 Mutation and cloning of p26 cDNAs

Four single amino acid substitutions in the α-crystallin domain of p26, namely R110G, F112R, R114A and Y116D, were generated by site-directed mutagenesis using the template pRSET.C-p26-3-6-3 (Liang *et al.*, 1997). The mutated cDNAs, along with wild type (WT) cDNA were then cloned into the eukaryotic expression vector pcDNA4/TO/myc-His.A (in this thesis, WT is equivalent to full length). The p26 cDNA-containing pcDNA4/TO/myc-His.A plasmids were digested with *Bam* HI and *Xba* I, with the inserts ligated into the His-tag containing prokaryotic expression vector pPROTet.E233. The sizes of insert cDNAs in eukaryotic and prokaryotic expression constructs were examined by restriction digestion, and the identity of each cDNA was confirmed by automated DNA sequencing (not shown).
Figure 13. p26 oligomer size is unaffected by pH

Protein extracts obtained from *Artemia* cysts prepared at either pH 7.0 (a, c) or pH 6.5 (b, d), were centrifuged in 10-50% continuous sucrose gradients either immediately after preparation (a, b), or following incubation at room temperature for 30 min (c, d). Fifteen μl fractions from each gradient were electrophoresed in 12.5% SDS polyacrylamide gels, transferred to nitrocellulose, and probed with anti-p26 antibody followed by HRP-conjugated goat anti-rabbit IgG. The top of each gradient is to the right and fractions are numbered across the top. The positions of molecular mass markers, alpha-lactalbumin, 14.2 kDa; carbonic anhydrase, 29 kDa; bovine serum albumin, 66 kDa; alcohol dehydrogenase, 150 kDa; apoferritin, 443 kDa; and thyroglobulin, 669 kDa are indicated by numbered arrows.
Figure 14. p26 oligomer size is unaffected by heat shock

Extracts prepared from heat shocked *Artemia* cysts and their nuclei were electrophoresed in SDS polyacrylamide gels and either stained with Coomassie blue (a) or blotted to nitrocellulose and stained with anti-p26 antibody (b). The lanes contained: 1, total cyst extract, heat shocked; 2, total cyst extract, control; 3, nuclear extract, heat shocked; 4, nuclear extract, control. Total extract from heated (c) and control (d) cysts and nuclear extracts from heated (e) and control (f) cysts were centrifuged in continuous 10-50% sucrose gradients and 15 μl fractions from each gradient were electrophoresed in SDS polyacrylamide gels, transferred to nitrocellulose, and probed with anti-p26 antibody followed by HRP-conjugated goat anti-rabbit IgG. The top of each gradient is to the right and fractions are numbered across the top. The positions of molecular mass markers, alpha-lactalbumin, 14.2 kDa; carbonic anhydrase, 29 kDa; bovine serum albumin, 66 kDa; alcohol dehydrogenase, 150 kDa; apoferritin, 443 kDa; and thyroglobulin, 669 kDa are indicated by numbered arrows.
3.2.2 Purification of p26 produced in bacteria

The pPROTet.E233 constructs were transformed into *E. coli* strain BL21PRO, and p26 expression was induced with anhydrotetracycline (aTc). After electrophoresis, light bands corresponding in expected size to p26 variants were visible in Coomassie blue stained SDS polyacrylamide gels, and these polypeptides reacted with antibody to p26 (Figs. 15a, b). Extract from *E. coli* transformed with vector without p26 cDNA lacked anti-p26 antibody reactive polypeptides (Figs. 15a, b, lane 6). After purification on TALON™ affinity columns, single bands of the expected size were seen on Coomassie blue stained gels (Fig. 15c) and these polypeptides reacted strongly with antibody to p26 (Fig. 15d). Each fusion protein of recombinant p26 expressed in *E. coli* BL21PRO contained, in addition to the p26 sequence (20.8 kDa), a 6×HN-tag followed by 14 amino acids prior to the multiple cloning site (3.2 kDa) and an amino-terminal peptide of 13 residues encoded by p26-3-6-3 but not part of the p26 sequence (1.5 kDa).

3.2.3 Oligomer formation by bacterially produced p26

Oligomers formed by each p26 mutant were similar in mass before and after purification from bacterial extracts (Figs. 16a, b). The molecular mass of oligomers assembled from WT p26 ranged from approximately 29 kDa to 669 kDa and this range was greater than for any p26 containing single amino acid substitutions. Among the four mutants, the range in oligomer mass was greatest for R114A, narrowest for F112R, and intermediate for R110G and Y116D. WT p26 exhibited the largest oligomers at 669 kDa, whereas for the modified p26 variants, R114A had the largest oligomers, although they were still smaller than for WT, and the maximum oligomer mass was smallest for F112R. These results are summarized in Table 6, with the median oligomer mass and monomer
Figure 15. Purification of p26 synthesized in *E. coli* BL21PRO

Cell free extracts from transformed *E. coli* BL21PRO induced with aTc were electrophoresed in SDS polyacrylamide gels and either stained with Coomassie blue (a) or blotted to nitrocellulose and reacted with antibody to p26 (b). Proteins purified by affinity chromatography were electrophoresed in SDS polyacrylamide gels and either stained with Coomassie blue (c), or blotted to nitrocellulose and reacted with antibody to p26 (d). The lanes contained: 1, R110G; 2, F112R; 3, R114A; 4, Y116D; 5, WT; 6, vector lacking p26 cDNA. Arrow, p26. All lanes received 10 µl of sample. M, molecular mass markers of 97, 66, 45, 31, 21 and 14 kDa.
**Figure 16. Oligomer formation by p26 synthesized in *E. coli* BL21PRO**

Bacterially produced p26 either before (a) or after purification to apparent homogeneity by affinity chromatography (b) was centrifuged at 200,000 x g for 12 h at 4 °C on 10-50% continuous sucrose gradients. The gradients were fractionated and 15-μl samples from each fraction were electrophoresed in SDS polyacrylamide gels, blotted to nitrocellulose and reacted with antibody to p26 followed by HRP-conjugated goat anti-rabbit IgG. The top of each gradient is to the right and fractions are numbered across the top. The molecular mass markers, alpha-lactalbumin, 14.2 kDa; carbonic anhydrase, 29 kDa; bovine serum albumin, 66 kDa; alcohol dehydrogenase, 150 kDa; apoferritin, 443 kDa; and thyroglobulin, 669 kDa are indicated by numbered arrows.
Figure 16
Table 6. Oligomerization of p26 produced in *E. coli* BL21PRO

<table>
<thead>
<tr>
<th>p26 variant</th>
<th>Monomer mass (kDa)</th>
<th>Oligomer Mass Range (kDa)</th>
<th>Median Oligomer Mass (kDa)</th>
<th>Median Monomer number</th>
</tr>
</thead>
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<td>R110G</td>
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</tr>
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<td>F112R</td>
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<td>R114A</td>
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<td>Y116D</td>
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</tr>
<tr>
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The molecular mass of p26 oligomers produced by transformed *E. coli* BL21PRO was determined by sucrose density gradient centrifugation. Monomer mass refers to the molecular mass of p26 polypeptides. Oligomer mass range represents the smallest to largest oligomers observed while median oligomer mass refers to the mass of the median size oligomer. Median monomer number refers to the number of subunits in the median size oligomer.
number given for each p26 variant. R114A is unusual within the latter comparison in that there was a greater number of oligomers above the median position than below, thus the median values may represent underestimations.

3.2.4 p26 induced bacterial thermotolerance

E. coli BL21PRO expressing all variants of recombinant p26 examined in this study were significantly more thermotolerant than bacteria containing only the pPROTet.E233 vector which did not survive heat shock, and of these, bacteria producing WT p26 were the most resistant to heat stress (Fig. 17). The levels of thermotolerance induced by expression of R110G, F112R and Y116D were similar to each other (P > 0.05) and significantly higher than the thermotolerance conferred by R114A (P < 0.05).

3.2.5 Chaperone activity of WT and modified p26

3.2.5.1 p26 protects citrate synthase against heat-induced aggregation

Purified WT p26 possessed the greatest chaperone activity and R114A the least, as determined by heat-induced aggregation of citrate synthase, although all variants provided protection (Fig. 18). At final p26 concentrations of 1200 nM and 600 nM when chaperone to target molar ratio is 8:1 and 4:1 respectively (p26 monomer to citrate synthase dimer), or 1:1 and 0.5:1 respectively if the median oligomer size of WT p26 is used for comparison, citrate synthase aggregation was inhibited almost completely for 1 h at 43 °C (Figs. 18a, b). At a final p26 concentration of 37.5 nM, wherein the molar ratio of WT p26 to citrate synthase is 1:4, or 1:32 if the median oligomer size of p26 is used for comparison, purified WT p26 reduced heat-induced turbidity by 46% (Fig. 18f). At each concentration from 37.5 nM to 1200 nM, mutants R110G, F112R and Y116D possessed similar levels of chaperone activity, significantly better than R114A, but less
Figure 17. Thermotolerance of transformed bacteria
Transformed \textit{E. coli} BL21PRO were incubated at 54 °C for periods up to 1 h, diluted and plated in duplicate on LB agar followed by incubation at 37 °C for 16 h. Colonies were counted and the $\log_{10}$ values of colony forming units (CFU) per ml were plotted against the length of heat shock in min. Bacteria containing the pPROTet.E233 vector lacking a p26 cDNA insert did not survive the 60 min heat shock. Results are averaged from three independent experiments.
Figure 18. p26 chaperone activity on citrate synthase

Bacterially produced p26 purified to apparent homogeneity was heated at 43 °C for 1 h with 150 nM citrate synthase, and solution turbidity was measured at 360 nm. The final concentrations of p26 based on monomer molecular mass were: a, 1200 nM; b, 600 nM; c, 300 nM; d, 150 nM; e, 75 nM; f, 37.5 nM. The curves are: 1, No p26; 2, R114A; 3, F112R; 4, Y116D; 5, R110G; 6, WT, and they occupy the same relative positions in a-f.
Figure 18
than WT (Fig. 18a-f). R110G, F112R and Y116D almost completely inhibited citrate synthase aggregation at 1200 nM (Fig. 18a), while R114A only reached this chaperone level at 2400 nM (not shown), twice the concentration required for other p26 variants. However, R114A still offered a significant amount of protection to citrate synthase even at lower ratios to substrate. BSA and IgG at 1200 nM provided almost no protection upon heating of citrate synthase when compared to the solution containing only enzyme (not shown).

3.2.5.2 p26 prevents citrate synthase from heat-induced inactivation

WT p26 protected citrate synthase against heat induced inactivation more effectively than any of the modified variants (Fig. 19). Of the tested samples, R114A exhibited the least protection at each chaperone concentration, this in contrast to R110G, F112R and Y116D which were more effective although not as good as WT. When the concentration of WT p26 reached 1200 nM, only limited enzyme inactivation occurred, with the detected enzyme activity almost the same as those determined for samples containing citrate synthase before heat shock (not shown).

3.2.5.3 p26 protects insulin against DTT-induced aggregation

WT p26 was the most effective in preventing DTT-induced insulin aggregation (Fig. 20), with R114A the least effective, and R110G, F112R and Y116D moderately better than R114A. At 0.1 μM, WT p26 inhibited insulin aggregation by 39% after 30 min, and the aggregation was almost completely suppressed at 1.6 μM. At this concentration, the molar ratio of chaperone to target is 0.4:1 (monomer to monomer), or 0.05:1 if the median oligomer size of WT p26 is used for comparison. BSA and IgG at 1.6 μM did not inhibit insulin aggregation (not shown).
Figure 19. Citrate synthase inactivation

Citrate synthase at 150 nM was heated at 43 °C for 1 h in either the absence or the presence of p26 and the enzyme activity remaining was determined. p26 concentrations were at 1200 nM, 600 nM, 300 nM, 150 nM, 75 nM, 37.5 nM. The p26 variants tested were: 1, R110G; 2, F112R; 3, R114A; 4, Y116D; 5, WT; 6, No p26, and they are in the same order at each concentration tested.
Figure 20. Insulin aggregation

Bacterially produced p26 purified to apparent homogeneity was incubated with insulin for 30 min in the presence of DTT and solution turbidity was measured at 400 nm. The p26 variants tested were: 1, R110G; 2, F112R; 3, R114A; 4, Y116D; 5, WT; 6, No p26, and they are in the same order in each histogram group.
3.2.6 p26 localization in transfected COS-1 cells

Transiently transfected COS-1 cells were immunofluorescently stained and examined by confocal laser scanning microscopy. WT p26 was localized predominantly in the cytoplasm of transfected cells (Fig. 21a). Mutants R110G, F112R and Y116D occurred in the cytoplasm but not in all nuclei of transfected cells (Figs. 21b, c, d). R114A was found in the cytoplasm and nuclei of all transfected COS-1 cells (Fig. 21e).

3.2.7 Oligomerization of p26 in COS-1 cells

The molecular mass of WT p26 oligomers assembled in COS-1 cells ranged from approximately 14.2 kDa to 500 kDa which was similar to that of R114A (Fig. 22). Among the four mutants, F112R exhibited the narrowest range in oligomer size, while R110G and Y116D were intermediate. The oligomeric pattern of each sample from COS-1 was generally similar to that from E. coli (Fig. 16), except that maximal oligomer mass for WT p26 was reduced. The results are summarized in Table 7.

3.2.8 The secondary structure of p26 was modified upon mutation

Far-UV CD spectra of WT p26 purified from transformed bacteria had a negative shoulder near 214 nm and a positive shoulder near 194 nm (Fig. 23-1), these characteristic of β-sheets. p26 from Artemia had similar negative and positive peaks (Fig. 23-2). Most of the α-crystallin domain mutants possessed a wider negative shoulder encompassing approximately 220 nm to 208 nm, with each mutant showing a positive shoulder near 194 nm (Figs. 23-3-6). Among them, R114A gave the greatest peaks for both the negative and positive shoulders, with higher intensities than other mutants and WT (Fig. 23-5). The calculated secondary structure elements of each sample are listed in Table 8, indicating decreased β-structure and increased α-helical constituents for the
Figure 21. p26 localization in transfected COS-1 cells

COS-1 cells transiently transfected with the p26 cDNA-containing vector pcDNA/4/TO/myc-His.A were incubated with antibody to p26 followed by FITC-conjugated goat anti-rabbit IgG antibody (green). Nuclei were stained with propidium iodide (red). a, WT; b, R110G; c, F112R; d, Y116D; e, R114A. The scale bar represents 100 μm and all figures are the same magnification.
Figure 22. Oligomer formation of p26 in transfected COS-1 cells

Extracts prepared from COS-1 cells transiently transfected with p26-containing vectors were centrifuged in 10-50% continuous sucrose gradients. The gradients were fractionated and 15-μl samples from each fraction were electrophoresed in SDS polyacrylamide gels, blotted to nitrocellulose and probed with antibody to p26 followed by HRP-conjugated goat anti-rabbit IgG antibody. The top of each gradient is to the right and fractions are numbered across the top. The molecular mass markers, alpha-lactalbumin, 14.2 kDa; carbonic anhydrase, 29 kDa; bovine serum albumin, 66 kDa; alcohol dehydrogenase, 150 kDa; apoferritin, 443 kDa; and thyroglobulin, 669 kDa are indicated by numbered arrows.
Figure 22
Table 7. Oligomerization of p26 synthesized in transfected COS-1 cells

<table>
<thead>
<tr>
<th>p26 variant</th>
<th>Monomer mass (kDa)</th>
<th>Oligomer Mass Range (kDa)</th>
<th>Median (kDa)</th>
<th>Median monomer number</th>
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<tbody>
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<td>R110G</td>
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<td>80</td>
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The molecular mass of p26 oligomers produced by COS-1 cells was determined by immunoprobing samples electrophoresed in SDS-polyacrylamide gels and transferred to western blots after sucrose gradient centrifugation. Monomer mass refers to the molecular mass of p26 polypeptides. Oligomer mass range represents the smallest to largest oligomers observed while median oligomer mass refers to the mass of the median size oligomer. Median monomer number refers to the number of subunits in the median size oligomer.
Figure 23. Far-UV circular dichroism spectra of p26

Each spectrum represents the average of 3 scans of purified p26 dissolved in 10 mM NaH₂PO₄, pH 7.1, at a final concentration of 0.2 mg/ml. The absorption data were expressed as molar ellipticity in degrees cm²/dmol (m deg). 1, WT from bacteria; 2, WT from Artemia; 3, R110G; 4, F112R; 5, R114A; 6, Y116D.
Table 8. Secondary structure elements of p26 calculated from CD spectra

<table>
<thead>
<tr>
<th>Elements</th>
<th>R110G (%)</th>
<th>F112R (%)</th>
<th>R114A (%)</th>
<th>Y116D (%)</th>
<th>WT (E. coli) (%)</th>
<th>WT (Artemia) (%)</th>
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Secondary structure elements of p26 were calculated with the CDNN v2.1 deconvolution program, with the percentage of α-helix, β-antiparallel, β-parallel, β-turn and random coil indicated for each sample.
variants, with these observations most pronounced for R114A.

3.2.9 Fluorescence studies of p26 revealed altered tertiary structure and reduced surface hydrophobicity

Fluorescence emission of each sample was detected over a range of 310-400 nm with an excitation wavelength at 280 nm. WT p26 from bacteria gave a maximal emission peak at 344 nm, while WT from *Artemia* was at 352 nm, with the former thus blue-shifted over the latter, although their emission intensities were similar (Fig. 24-1, 2). In comparison, all mutants exhibited reduced emission intensities, with the peaks of R110G, F112R, Y116D at 344 nm and the peak of R114A at 358 nm (Fig. 24-3-6). The intensity of R114A was the lowest with a red-shift in comparison with other samples (Fig. 24-5), indicating altered microenvironments and mobilities for aromatic amino acid residues.

ANS-binding experiments were carried out at an excitation wavelength of 388 nm with an emission wavelength at 473 nm. WT p26 from bacteria and *Artemia* gave similar fluorescence intensities (Fig. 25), but all four mutants exhibited decreased ANS-binding capacity when compared with WT at either 25 ºC or 43 ºC. Among them R114A possessed the lowest ability to bind ANS. When tested at 25 ºC, the fluorescence intensity of R114A was 45.5% that of WT from bacteria, and at 43 ºC, it was 40.1%. With elevated temperature, the fluorescence intensity of each sample increased. For instance, the emission of p26 from *Artemia* at 43 ºC was 73.9% higher than at 25 ºC. However, the relative increases in fluorescence intensity of mutants when temperature changed from 25 ºC to 43 ºC were less remarkable than that of WT. As an example, the increase of R110G was 19.4%, in contrast to that of WT from *E. coli* as 72.4%.
Figure 24. Intrinsic fluorescence of p26

Purified p26 was diluted in 10 mM NaH₂PO₄, pH 7.1 to a final concentration of 0.06 mg/ml. The excitation wavelength was set to 280 nm with a 2 nm band pass and fluorescence emission was detected over a range of 310-400 nm. Each spectra was recorded in duplicate using two independent sample preparations. 1, WT from bacteria; 2, WT from Artemia; 3, R110G; 4, F112R; 5, R114A; 6, Y116D.
Figure 24
Figure 25. Surface hydrophobicity of p26

Purified p26 solutions at 0.06 mg/ml in 10 mM NaH₂PO₄, pH 7.1 were oversaturated with ANS and the fluorescence was measured with an excitation wavelength at 388 nm, a band pass of 8 nm, and an emission wavelength set to 473 nm with a band pass of 8 nm. Measurements were carried out at either 25 °C (shaded columns) or 43 °C (filled columns). The fluorescence generated by buffer containing ANS but no p26 was subtracted from the readings for each sample.
Figure 25
3.3 Internal deletions of p26

3.3.1 Mutation and cloning of p26 cDNAs

Three internal deletions of p26, namely “G” (multiple glycine deletion), “R” (multiple arginine deletion), and “TS” (multiple serine/threonine deletion) were generated by site-directed mutagenesis using pRSET.C-p26-3-6-3 as template. Extracted plasmids for the mutants were digested with Bam HI and Xho I and the products were electrophoresed in 1% agarose gels, revealing DNA inserts at the expected positions. The identity of each insert cDNA was confirmed by DNA sequencing (not shown). The modified and WT cDNAs were then subcloned between the Bam HI and Xho I restriction sites of the eukaryotic expression vector pcDNA4/TO/myc-His.A. These p26 cDNA containing constructs were digested with Bam HI and Xba I and the resulting DNA fragments were ligated into the prokaryotic expression vector pPROtet.E233. The sizes of cDNAs inserted into eukaryotic and prokaryotic expression vectors were examined by restriction digestion, and their identities confirmed by automated DNA sequencing (not shown).

3.3.2 Purification of p26 synthesized in bacteria

The pPROtet.E233 constructs were transformed into E. coli strain BL21PRO, and expression was induced with anhydrotetracycline. Cell-free extracts containing expressed polypeptides were electrophoresed in SDS polyacrylamide gels, blotted to nitrocellulose and reacted with antibody to p26. Bands corresponding in size to p26 variants were visible in Coomassie blue stained gels and these polypeptides reacted with antibody to p26 (Figs. 26a, b). The “TS” construct gave the lowest yield and protein bands were not detected on blots containing extract from E. coli transformed with vector only. Upon purification with BD TALON™ IMAC system, single bands of the expected size
Figure 26. Purification of p26 synthesized in E. coli BL21PRO

Cell free extracts from transformed E. coli BL21PRO induced with aTc were electrophoresed in SDS polyacrylamide gels and either stained with Coomassie blue (a) or blotted to nitrocellulose and reacted with antibody to p26 (b). Proteins purified by affinity chromatography were electrophoresed in SDS polyacrylamide gels and either stained with Coomassie blue (c), or blotted to nitrocellulose and reacted with antibody to p26 (d). The lanes contained: 1, “G”; 2, “R”; 3, “TS”; 4, WT; 5, vector lacking p26 cDNA. Arrow, p26. All lanes received 10 µl of sample. M, molecular mass markers of 97, 66, 45, 31, 21 and 14 kDa.
appeared on Coomassie blue stained gels and these polypeptides reacted with antibody to p26 on western blots (Figs. 26c, d).

3.3.3 p26 oligomer formation

The oligomer size of each bacterially expressed p26 variant was similar before and after purification (Figs. 27a, b). The molecular mass of oligomers assembled from purified WT p26 synthesized in *E. coli* ranged from 29 kDa to 669 kDa, whereas the range was 14 kDa to 443 kDa for “G” and 14 kDa to 300 kDa for “R” and “TS”. The median mass of WT oligomers was greater than for any of the mutants. The results are summarized in Table 9.

3.3.4 Bacterial thermotolerance

*E. coli* BL21PRO expressing all variants of recombinant p26 examined in this study were significantly more thermotolerant than bacteria containing only the pPROTet.E233 vector which did not survive heat shock, and of these, bacteria producing WT p26 were the most resistant to heat stress (Fig. 28). The levels of thermotolerance induced by expression of “G” and “TS” were similar to each other (*P* > 0.05) and significantly higher than the thermotolerance conferred by variant “R” (*P* < 0.05).

3.3.5 Chaperone activity

3.3.5.1 p26 protects citrate synthase against heat-induced aggregation

Purified WT p26 possessed the greatest chaperone activity, while mutant “R” exhibited the least, as determined by heat-induced aggregation of citrate synthase, although all mutants provided some protection (Fig. 29). At final WT p26 concentrations of 1200 nM and 600 nM when chaperone to target molar ratio is 8:1 and 4:1 respectively (p26 monomer to citrate synthase dimer), or 1:1 and 0.5:1 respectively if the median oligomer
Figure 27. Oligomer formation by p26 synthesized in *E. coli* BL21PRO

Bacterially produced p26 either before (a) or after purification to apparent homogeneity (b) was centrifuged at 200,000 x g for 12 h at 4 °C on 10-50% continuous sucrose gradients. The gradients were fractionated and 15-µl samples from each fraction were electrophoresed in SDS polyacrylamide gels, blotted to nitrocellulose and reacted with antibody to p26 followed by HRP-conjugated goat anti-rabbit IgG. The top of each gradient is to the right and fractions are numbered across the top. The molecular mass markers, alpha-lactalbumin, 14.2 kDa; carbonic anhydrase, 29 kDa; bovine serum albumin, 66 kDa; alcohol dehydrogenase, 150 kDa; apoferritin, 443 kDa; and thyroglobulin, 669 kDa are indicated by numbered arrows.
Figure 27
Table 9. Oligomerization of p26 produced in *E. coli* BL21PRO

<table>
<thead>
<tr>
<th>p26 variant</th>
<th>Monomer mass (kDa)</th>
<th>Oligomer Mass Range (kDa)</th>
<th>Median (kDa)</th>
<th>Median monomer number</th>
</tr>
</thead>
<tbody>
<tr>
<td>&quot;G&quot;</td>
<td>23.7</td>
<td>14-443</td>
<td>95</td>
<td>4</td>
</tr>
<tr>
<td>&quot;R&quot;</td>
<td>24.1</td>
<td>14-300</td>
<td>66</td>
<td>3</td>
</tr>
<tr>
<td>&quot;TS&quot;</td>
<td>23.8</td>
<td>14-300</td>
<td>66</td>
<td>3</td>
</tr>
<tr>
<td>WT</td>
<td>25.5</td>
<td>29-669</td>
<td>200</td>
<td>8</td>
</tr>
</tbody>
</table>

The molecular mass of p26 oligomers produced by transformed *E. coli* BL21PRO was determined by sucrose density gradient centrifugation. Monomer mass refers to the molecular mass of p26 polypeptides. Oligomer mass range represents the smallest to largest oligomers observed while median oligomer mass refers to the mass of the median size oligomer. Median monomer number refers to the number of subunits in the median size oligomer.
Figure 28. Thermotolerance of transformed bacteria

Transformed *E. coli* BL21PRO were incubated at 54 °C for periods up to 1 h, diluted and plated in duplicate on LB agar followed by incubation at 37 °C for 16 h. Colonies were counted and the Log$_{10}$ values of colony forming units (CFU) per ml were plotted against the length of heat shock in min. Bacteria containing the pPROTet.E233 vector lacking a p26 cDNA insert did not survive the 60 min heat shock. Results are averaged from three independent experiments.
Figure 28
Figure 29. p26 chaperone activity on citrate synthase

Bacterially produced p26 purified to apparent homogeneity was heated at 43 °C for 1 h with 150 nM citrate synthase, and solution turbidity was measured at 360 nm. The final concentrations of p26 based on monomer molecular mass were: a, 1200 nM; b, 600 nM; c, 300 nM; d, 150 nM; e, 75 nM; f, 37.5 nM. The curves are: 1, No p26; 2, “R”; 3, “TS”; 4, “G”; 5, WT, and they occupy the same relative positions in a-f.
Figure 29
size of WT p26 is used for comparison, citrate synthase aggregation was inhibited almost completely for 1 h at 43 °C (Figs. 29a, b). At a final p26 concentration of 37.5 nM, where the molar ratio of WT p26 to citrate synthase is 1:4, or 1:32 if the median oligomer size of p26 is used for comparison, purified WT p26 reduced heat-induced turbidity by 46% after 1 h at 43 °C (Fig. 29f). WT p26 at each concentration from 37.5 nM to 1200 nM gave the best protection, mutants “G”, “TS” and “R” followed in decreasing order of activity (Figs. 29a-f). None of the three mutants with internal deletions completely inhibited the aggregation of citrate synthase at 1200 nM (Fig. 29a), nor were higher concentrations more effective (not shown). However, “R”, the least effective chaperone, still offered a significant amount of protection to citrate synthase even at the lowest ratio to substrate tested. BSA and IgG at 1200 nM provided almost no protection when heated with citrate synthase (not shown).

3.3.5.2 p26 prevents citrate synthase from heat-induced inactivation

WT p26 was the most effective in protecting citrate synthase against heat induced inactivation, followed by “G” and “TS” which were very similar and then “R” which exhibited the least protection at each chaperone concentration (Fig. 30). When the concentration of WT p26 was 1200 nM, the enzyme activity remaining after 1 h at 43 °C was almost the same as in samples containing citrate synthase before heat shock (not shown).

3.3.5.3 p26 protects insulin against DTT-induced aggregation

Purified WT p26 was the most effective in preventing DTT-induced denaturation of insulin at room temperature (Fig. 31), whereas mutant “R” had the least ability to protect insulin. Mutants “G” and “TS” provided an intermediate level of protection. At 0.1 μM,
Figure 30. Citrate synthase inactivation

Citrate synthase at 150 nM was heated at 43 °C for 1 h in either the absence or the presence of p26 and the enzyme activity remaining was determined. p26 concentrations were at 1200 nM, 600 nM, 300 nM, 150 nM, 75 nM, 37.5 nM. The p26 variants tested were: 1, "G"; 2, "R"; 3, "TS"; 4, WT; 5, No p26, and they are in the same order at each concentration tested.
Figure 30
Figure 31. Insulin aggregation

Bacterially produced p26 purified to apparent homogeneity was incubated with insulin for 30 min in the presence of DTT and solution turbidity was measured at 400 nm. The p26 variants tested were: 1, “G”; 2, “R”; 3, “TS”; 4, WT; 5, No p26, and they are in the same order in each histogram group.
Figure 31
WT p26 inhibited insulin aggregation by 39% after 30 min, and the aggregation was almost completely suppressed at 1.6 μM. At this concentration, the molar ratio of chaperone to target is 0.4:1 (monomer to monomer), or 0.05:1 if the median oligomer size of WT p26 is used for comparison. BSA and IgG at 1.6 μM did not inhibit insulin aggregation (not shown).

3.3.6 Immunolocalization of p26 in COS-1 cells

Transiently transfected COS-1 cells were immunofluorescently stained and examined by confocal laser scanning microscopy. WT p26 was localized predominantly in the cytoplasm of transfected cells (Fig. 32a). Mutants “G” and “TS” occurred in the cytoplasm of all transfected cells and the nuclei of some (Figs. 32b, c), whereas “R” was found in the cytoplasm and nuclei of all transfected COS-1 cells (Fig. 32d).

3.3.7 Expression and oligomerization of p26 in COS-1 cells

The molecular mass of WT p26 oligomers assembled in COS-1 cells ranged from approximately 14 kDa to 500 kDa which was broader than for the mutants (Fig. 33). Among the four mutants, “G” exhibited slightly broader range in oligomer size than “R” and “TS”. The oligomeric pattern of each sample from COS-1 was generally similar to that from E. coli (Fig. 27), except that maximal oligomer mass for WT p26 was reduced. The results are summarized in Table 10.

3.3.8 The secondary structure of p26 was altered by mutation

Far-UV CD spectra of WT p26 purified from either transformed bacteria or Artemia cysts had a negative shoulder near 214 nm and a positive shoulder near 194 nm, these characteristic of β-sheets (Fig. 34). The mutants with internal deletions possessed wider negative shoulders encompassing approximately 220 nm to 208 nm, with each mutant
Figure 32. p26 localization in transfected COS-1 cells

COS-1 cells transiently transfected with the p26 cDNA-containing vector pcDNA/4/TO/myc-His.A were incubated with antibody to p26 followed by FITC-conjugated goat anti-rabbit IgG antibody (green). Nuclei were stained with propidium iodide (red). a, WT; b, “G”; c, “TS”; d, “R”. The scale bar represents 100 μm and all figures are the same magnification.
Figure 33. Oligomer formation of p26 in transiently transfected COS-1 cells

Extracts prepared from COS-1 cells transiently transfected with p26-containing vectors were centrifuged in 10-50\% continuous sucrose gradients. The gradients were fractionated and 15-\mu l samples from each fraction were electrophoresed in SDS polyacrylamide gels, blotted to nitrocellulose and probed with antibody to p26 followed by HRP-conjugated goat anti-rabbit IgG antibody. The top of each gradient is to the right and fractions are numbered across the top. The molecular mass markers, alpha-lactalbumin, 14.2 kDa; carbonic anhydrase, 29 kDa; bovine serum albumin, 66 kDa; alcohol dehydrogenase, 150 kDa; apoferritin, 443 kDa; and thyroglobulin, 669 kDa are indicated by numbered arrows.
### Table 10. Oligomerization of p26 synthesized in transfected COS-1 cells

<table>
<thead>
<tr>
<th>p26 variant</th>
<th>Monomer mass (kDa)</th>
<th>Oligomer Mass Range (kDa)</th>
<th>Median (kDa)</th>
<th>Median monomer number</th>
</tr>
</thead>
<tbody>
<tr>
<td>“G”</td>
<td>18.7</td>
<td>14-443</td>
<td>80</td>
<td>4</td>
</tr>
<tr>
<td>“R”</td>
<td>19.4</td>
<td>14-300</td>
<td>66</td>
<td>3</td>
</tr>
<tr>
<td>“TS”</td>
<td>19.1</td>
<td>14-300</td>
<td>66</td>
<td>3</td>
</tr>
<tr>
<td>WT</td>
<td>20.8</td>
<td>14-500</td>
<td>100</td>
<td>5</td>
</tr>
</tbody>
</table>

The molecular mass of p26 oligomers produced by COS-1 cells was determined by immunoprophobing samples electrophoresed in SDS-polyacrylamide gels and transferred to membranes for Western blots after sucrose gradient centrifugation. Monomer mass refers to the molecular mass of p26 polypeptides. Oligomer mass range represents the smallest to largest oligomers observed while median oligomer mass refers to the mass of the median size oligomer. Median monomer number refers to the number of subunits in the median size oligomer.
Figure 34. Far-UV circular dichroism spectra of p26

Each spectrum represents the average of 3 scans of purified p26 dissolved in 10 mM NaH₂PO₄, pH 7.1, at a final concentration of 0.2 mg/ml. The absorption data were expressed as molar ellipticity in degrees cm²/dmol (m deg). Tested samples are: 1, WT from bacteria; 2, WT from Artemia; 3, “G”; 4, “R”; 5, “TS”.
showing a positive shoulder near 194 nm. Among all p26 variants examined, "R" gave
the greatest peak for both the negative and positive shoulders. The calculated secondary
structure elements of each sample are listed in Table 11, indicating decreased β-structure
and increased α-helical constituents for the variants, with these characteristics most
pronounced for "R".

3.3.9 Tertiary structure and surface hydrophobicity were affected by mutation

Fluorescence emission of each sample was detected over a range of 310-400 nm with
an excitation wavelength at 280 nm. WT p26 from bacteria gave a maximal emission
peak at 344 nm, while WT from Artemia was at 352 nm, with the former thus blue-
shifted over the latter, although their emission intensities were similar (Fig. 35). In
comparison, all mutants exhibited reduced emission intensities, with the peak of "G" and
"TS" at 348 nm and "R" at 360 nm. The fluorescence intensity of "R" was the lowest
with a red-shift in comparison with other samples. The results indicate altered
microenvironments and mobilities for aromatic residues.

ANS-binding experiments were carried out at an excitation wavelength of 388 nm with
an emission wavelength at 473 nm. WT p26 from bacteria and Artemia gave similar
fluorescence intensities (Fig. 36), but all three mutants exhibited decreased ANS-binding
capacity when compared with WT at either 25 °C or 43 °C. Among them "R" possessed
the lowest ability to bind ANS. When tested at 25 °C, the fluorescence intensity of "R"
was 38.5% that of WT from bacteria, and at 43 °C, it was only 28.7%. With elevated
temperature, the fluorescence of each sample increased. For instance, the emission of p26
from Artemia at 43 °C was 73.9% higher than at 25 °C. However, the relative increases in
fluorescence intensity of mutants when temperature changed from 25 °C to 43 °C were
Table 11. Secondary structure elements of p26 calculated from CD spectra

<table>
<thead>
<tr>
<th>Elements</th>
<th>&quot;G&quot; (%)</th>
<th>&quot;R&quot; (%)</th>
<th>&quot;TS&quot; (%)</th>
<th>WT (E. coli) (%)</th>
<th>WT (Artemia) (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>α-helix</td>
<td>21.8</td>
<td>24.9</td>
<td>22.2</td>
<td>17.7</td>
<td>17.7</td>
</tr>
<tr>
<td>β-antiparallel</td>
<td>17.8</td>
<td>15.6</td>
<td>16.4</td>
<td>21.6</td>
<td>23.0</td>
</tr>
<tr>
<td>β-parallel</td>
<td>9.8</td>
<td>9.4</td>
<td>9.5</td>
<td>10.3</td>
<td>10.0</td>
</tr>
<tr>
<td>β-turn</td>
<td>17.1</td>
<td>17.4</td>
<td>17.5</td>
<td>16.5</td>
<td>16.8</td>
</tr>
<tr>
<td>Random coil</td>
<td>33.5</td>
<td>32.7</td>
<td>34.4</td>
<td>34.0</td>
<td>32.5</td>
</tr>
</tbody>
</table>

Secondary structure elements of p26 were calculated with the CDNN v2.1 deconvolution program, with the percentage of α-helix, β-antiparallel, β-parallel, β-turn and random coil indicated for each sample.
Figure 35. Intrinsic fluorescence of p26

Purified p26 was diluted in 10 mM NaH₂PO₄, pH 7.1 to a final concentration of 0.06 mg/ml. The excitation wavelength was set to 280 nm with a 2 nm band pass and fluorescence emission was detected over a range of 310-400 nm. Each spectrum was recorded in duplicate using two independent sample preparations. 1, WT from bacteria; 2, WT from Artemia; 3, “G”; 4, “R”; 5, “TS”.

Figure 35
Figure 36. Surface hydrophobicity of p26

Purified p26 at 0.06 mg/ml in 10 mM NaH$_2$PO$_4$, pH 7.1 was oversaturated with ANS and the fluorescence was measured with an excitation wavelength at 388 nm, a band pass of 8 nm, and an emission wavelength set to 473 nm with a band pass of 8 nm. Measurements were carried out at either 25 °C (shaded columns) or 43 °C (filled columns). Tested samples are: 1, “G”; 2, “R”; 3, “TS”; 4, WT from bacteria; 5, WT from Artemia. The fluorescence generated by buffer containing ANS but no p26 was subtracted from the readings for each sample.
Figure 36
less remarkable than for WT. As an example, the increase for “R” was 27.9%, in contrast to WT from *E. coli* which was 72.4%.

3.4 Comparative modeling of p26 structure

The 29% sequence identity between α-crystallin domains of p26 and wheat *Ta* Hsp16.9 allowed generation of a p26 model based on the crystal structure of *Ta* Hsp16.9 (Fig. 37). The overall sequence identity between these two sHsps was 20%, and the lack of significant sequence identity between p26 and *Ta* Hsp16.9 C-terminal extensions precluded modeling this portion of p26. Thus, residues A159 to A192 were deleted from the p26 sequence for a better alignment interface. The similarity between the proteins indicates each p26 monomer contains a buried N-terminal region, a partially exposed α-crystallin domain and a solvent-accessible C-terminal extension, as initially shown for *Ta* Hsp16.9 (van Montfort *et al.*, 2001b). The α-crystallin domain of p26 forms an immunoglobulin-like fold composed of eight β-strands organized into two anti-parallel β-sheets. In the p26 model, loop 1 composed of residues 22FGFGFGGMDL32 is located in the center of the N-terminal region, whereas loop 2 composed of residues 58TPGLSR63 is in the region connecting the N-terminal region and α-crystallin domain. Loop 3 (102SGEYGHVQRE111) protrudes from β-sheets of the α-crystallin domain and corresponds to the loop encompassing the β6 strand in *Ta* Hsp16.9. Loop 4 (132SSDGV136) equates to a sequence connecting β8 and β9 of *Ta* Hsp16.9. Residues 153VIPTP158 form the partial C-terminal extension of p26, and this sequence corresponds to a stretch close to the β10 strand in the C-terminal extension of *Ta* Hsp16.9. Hydrophobic residues I147 and I149 of *Ta* Hsp16.9 bind in a hydrophobic groove between strands β4 and β8 of a neighboring monomer in intact oligomers.
Figure 37. Comparative modeling of p26

Wheat *Ta* Hsp16.9 was used as a template for sequence alignment with the “alignment interface” program of Swiss-Model. The p26 residues A2-P158, corresponding to S2-G151 of *Ta* Hsp16.9 were used for modeling and the result is shown as a monomer, with C-terminal residues A159-A192 deleted for alignment optimization. The returned three-dimensional model was enhanced with the “improve fit” function of Swiss-PdbViewer and sent for second round modeling. Internal deletions “G” and “R” are indicated by labeled arrows; the location of the four α-crystallin domain single amino acid substitutions in strands β6-β7 (arrow labeled) are hidden from view by overlaying strands β2-β3 (arrow labeled) (a). The arrow-labeled A2 and P158 position the first and last amino acid residues, respectively, in the p26 model. The N-terminal region, α-crystallin domain and C-terminal extension are indicated by labeled arrows. Loops within the p26 monomer are indicated in b, and the large loop of the wheat *Ta* Hsp16.9 monomer corresponding to p26 loop 3 is indicated in c.
Figure 37
IV. Discussion

p26 exists in large amounts in encysted *Artemia* embryos, and is thought to play an important role in protecting embryos against environmental hardships. As the only crustacean sHSP described in detail so far, p26 has the ability to effectively prevent proteins from irreversible aggregation during *Artemia* development. p26 structure and function were characterized by site-directed mutagenesis in this work, which allows a further appreciation of p26 relevance during embryo diapause, desiccation and other stress, thereby also shedding light on the functional mechanisms of sHSPs in human diseases.

4.1 Sequence analysis of p26

Like most sHSPs, p26 is composed of three regions, with the conserved "α-crystallin domain" as the hallmark of this protein family. p26 was predicted to lack α-helical constituents in the α-crystallin domain, which consists predominantly of β-sheets (Liang *et al.*, 1997a). p26 forms oligomers and exhibits chaperone activity *in vivo* and *in vitro* (Liang *et al.*, 1997b; Crack *et al.*, 2002), characteristics potentially dominated by conserved sequences. The p26 α-crystallin domain is flanked by the 60-residue N-terminus and the 40-residue C-terminus, both of limited similarity to corresponding regions in other sHSPs. The nonconserved N-terminal region and C-terminal extension may determine the distinct properties of p26. As compared with other sHSPs, a unique characteristic of the p26 N-terminal region is the presence of 10 glycines in the first 29 amino acid residues, which are in a 22-residue peptide. Other sHSPs usually have only 3 or 4 glycines in the equivalent region. Seven arginines occur between residues 36-51 of p26, with 6 distributed in a 10-residue group. The first 60 residues of p26 share limited
sequence identity with other sHSPs, but exhibit a weak similarity in terms of the
enrichment in glycine, arginine and phenylalanine. p26 shares many highly conserved
residues with other sHSPs in the central α-crystallin domain, especially in the sequence
110REFRRRY116. Mutations to the arginine residues equivalent to R114 of p26 in αA-
and αB-crystallin result in autosomal dominant congenital cataract and desmin-related
myopathy, respectively (Litt et al., 1998; Vicart et al., 1998).

The sHSP C-terminal extension is variable in length and sequence from species to
species, being absent from C. elegans Hsp12.6, while reaching 49 residues in Drosophila
Hsp27 and 50 residues in Artemia p26 (Leroux et al., 1997a; Liang et al., 1997a; de Jong
et al., 1998b). Composed predominantly of polar and charged amino acid residues, the C-
terminal extension solubilizes either sHSP oligomers or the complexes formed between
sHSPs and substrates (Lindner et al., 2000). Removal of this extension will likely
decrease protein solubility and compromise chaperone function. The p26 C-terminal
extension encompasses a serine/threonine-enriched stretch, wherein 13 serines or
threonines exist within 24 residues. The structural and functional relevance of this motif
was unexplored prior to this work.

4.2 Oligomerization of p26

Most sHSPs tend to assemble into large oligomeric complexes as a prerequisite for
chaperone activity. Native p26 extracted from Artemia cysts forms polydisperse
oligomers with a maximum mass of 669 kDa composed of up to 32 subunits, as is also
ture for recombinant p26 from bacteria although there are more smaller oligomers (Crack
et al., 2002). p26 oligomers in COS-1 cells are from 20.8 kDa to 500 kDa. p26 appears as
particles about 15 nm in diameter as determined by electron microscopy, and they are
similar in size to human αB-crystallin and Hsp27, native bovine α-crystallin, and yeast Hsp26 (Clegg et al., 1999; Haslbeck et al., 1999; Haley et al., 2000). The oligomeric polydispersity of p26 contrasts to *Mj* Hsp16.5 which consists of 24 monomers organized into a spherical complex, *Ta* Hsp16.9 existing as a dodecamer assembled into double disks and *Mt* Hsp16.3 which forms a triangular structure as a nonamer composed of three trimers (Kim et al., 1998; van Montfort et al., 2001b; Abulimiti et al., 2003).

The 17WSDPF21 motif in p26 is conserved and corresponds to 15SWEPF19 in Chinese hamster Hsp27, a sequence important in oligomer formation and chaperone activity (Theriault et al., 2004). This motif was missing in truncations p26-NΔ36 and p26-NΔ60, and the absence of higher order oligomers in both cases suggests the involvement of this sequence in oligomer assembly, a proposal currently under study. Mutated *Mj* Hsp16.5 lacking the first N-terminal 12 amino acid residues fails to assemble into spherical oligomers, indicating the region is required for oligomer organization (Kim et al., 2003a). Other N-terminal hydrophobic elements may contribute to p26 structural organization, with the deleted residues 1-36 and 1-60 possessing 72.2% and 61.7% hydrophobicity, respectively, both higher than the 51% hydrophobicity of the p26 full-length sequence.

Truncated p26 consisting of only the α-crystallin domain existed predominantly as dimers and monomers. The α-crystallin domains of many sHSPs assemble into dimers rather than larger oligomers, while oligomer assembly and dynamics are controlled by the N-terminal region (Koteiche and McHaourab, 2002; Stromer et al., 2004). For example, analysis of yeast Hsp26 revealed that the N-terminal region is critical for formation of Hsp26 large oligomers (Stromer et al., 2004). This is consistent with data presented in
this thesis and corroborates the importance of the p26 N-terminal region in oligomer assembly. An interesting point is that sHSP oligomerization is insensitive to sequence extension at the N-terminus (Leroux et al., 1997b; Muchowski et al., 1997). For example, adding a tail of 4 kDa to the C. elegans Hsp16-2 N-terminal did not disturb oligomerization, and after fusion of the 42 kDa maltose-binding protein to the N-terminus of αB-crystallin, the recombinant protein remained as high molecular mass oligomers displaying full chaperone function (Leroux et al., 1997b; Muchowski et al., 1997). In this work, an elongation of approximately 5 kDa to the WT p26 N-terminus did not interfere with oligomer assembly dramatically.

Removal of 10 C-terminal residues had little effect on p26. However, deleting the entire C-terminal extension including the conserved I/V-X-I/V (as VPI for p26) reduced oligomerization more dramatically, and a similar case occurred for B. japonicum HspH (Studer et al., 2002). The p26 C-terminal extension influences oligomer assembly less than the N-terminal region. In comparison, pea Hsp17.7 variants lacking 3, 5, or 10 amino acid residues of the C-terminal fail to form full size oligomers (Kirschner et al., 2000), while C-terminal truncated B. japonicum HspB and HspC without the conserved IXI motif exist as small oligomers (Narberhaus, 2002). Deletion of the last 11 or more C-terminal residues of αA-crystallin significantly decreases oligomeric mass, although removing up to 10 residues causes little change (Thampi and Abraham, 2003). High resolution crystal structures of Ta Hsp16.9 reveal inter-dimer interactions between hydrophobic residues around the C-terminal extension β10 strand with a hydrophobic patch on the surface of α-crystallin β4, β5 and β8 strands, indicating the involvement of
C-terminal sequence encompassing the conserved IXI motif in sHSP oligomerization (van Montfort et al., 2001b; Giese and Vierling, 2002; Kim et al., 2003a).

Single site mutations to the conserved α-crystallin domain of p26 resulted in generally smaller oligomers than for WT p26, with F112R oligomers exhibiting a much narrower range than WT. Intriguingly, the largest oligomers obtained for mutant R114A were 500 KDa, contrasting to R116C αA-crystallin and R120G αB-crystallin, mutants with molecular mass of >2 MDa and 1.4 MDa, respectively (Bova et al., 1999a; Andley et al., 2002). Clearly, modification of this highly conserved arginine residue had less effect on p26 oligomer formation than for the α-crystallins.

4.3 Chaperone activity of p26

sHSPs are effective molecular chaperones, recognizing and interacting with nonnative rather than native structures. The chaperone activity of p26 towards heat-induced citrate synthase aggregation and inactivation and DTT-induced insulin aggregation was determined. Citrate synthase, a model protein for the characterization of chaperone activity has been used extensively for studies of molecular chaperones including GroEL, HSP90 and sHSPs (Muchowski and Clark, 1998). Citrate synthase is inactivated at 43 °C, a temperature relevant to physiological heat shock conditions, and it subsequently forms aggregation-prone intermediates, thereby allowing examination of sHSP chaperone activity (Buchner et al., 1998; Rajaraman et al., 2001). Insulin is another ideal substrate for investigation of sHSP chaperone function. The β-chain of insulin aggregates upon reduction of the disulfide bond between the α- and β-chains, increasing solution turbidity. sHSPs bind to the unfolding β-chain and reduce aggregation (Haslbeck et al., 1999). This assay allows examination of p26 substrate aggregation kinetics in the absence
of heat shock, and mimics non-thermal stressful conditions that *Artemia* cysts experience during development.

In contrast to GroEL, which accommodates one or two substrate molecules per 14 subunits (Houry *et al.*, 1999), sHSPs usually have a much higher binding capacity. A 2:1 molar ratio of chaperone to substrate (monomer to monomer) often prevents most protein aggregation (Studer and Narberhaus, 2000). Native p26 from *Artemia* and recombinant p26 from bacteria efficiently prevented thermally induced citrate synthase aggregation and chemically induced insulin aggregation, with a molar ratio of p26 to substrate of 4:1 for the former (monomer to dimer) and 0.4:1 for the latter. The former contrasts a molar ratio of 7:1 for *C. elegans* Hsp16-2 and 8:1 for *Mt* Hsp16.3 to citrate synthase (Chang *et al.*, 1996; Leroux *et al.*, 1997b), and a molar ratio of 3:1 for αA- and αB-crystallins to the same substrate (Santhoshkumar and Sharma, 2001). The relative binding capacity of sHSPs for DTT-sensitive proteins ranges from 2:1 (sHSPs to insulin or α-lactalbumin) to 10:1 (sHSPs to ovotransferrin) (Lindner *et al.*, 1998). It is reasonable to speculate that p26 binding capability in *Artemia* cysts equals values observed *in vitro*, and considerable amounts of protein can be stored by interacting with p26. Due to the abundance of p26 in encysted embryos, this provides a large reservoir of proteins for renaturation under permissive conditions as *Artemia* resumes development.

Neither BSA nor IgG significantly affected the aggregation or inactivation kinetics of target proteins, indicating that effects observed with p26 are due to its chaperone activity instead of non-specific protein interactions. However, chaperoning by modified p26 was compromised. p26 variants either composed of only the α-crystallin domain or without the N-terminal region were relatively poor chaperones in comparison with WT, and
activity decreased as truncation increased. The single site mutation R114A led to a considerable loss of p26 chaperone ability. Incubation of R114A p26 with thermally denatured citrate synthase did not inhibit aggregation as well as WT, nor did R114A p26 suppress citrate synthase aggregation effectively until its concentration was increased to 1200 nM when the molar ratio of chaperone to substrate was 8:1. Even at concentrations of R114A p26 higher than 1200 nM, substrate aggregation remained the same, indicating the mutant had reached its maximum ability to chaperone citrate synthase at this concentration. The same results were found for the inhibition of DTT-induced insulin aggregation, where R114A p26 concentrations of 1.6 μM or higher were required for effective chaperoning. That R114A lost substantial in vitro chaperone capability against thermal or chemical stress suggests that the substrate-binding site is impaired, preventing effective contact between the chaperone and substrates.

Among three internal deletions, “R” was most detrimental and had the lowest capability to suppress protein aggregation. Although both R114A and “R” exhibited reduced chaperone activity, they functioned as chaperones to a certain extent, in contrast to either R116C αA-crystallin whose chaperone activity is almost completely abolished, or R120G αB-crystallin which enhances the kinetics and extent of substrate aggregation (Kumar et al., 1999; Chavez-Zobel et al., 2003). Although α-crystallins usually function for the entire lifespan of mammalian, a single site mutation may bring highly pathological effects. By comparison, p26 seems more resistant to modification of the conserved arginine residue than its mammalian counterparts, and our data suggest p26 may be more stable than other sHSPs.
4.4 Secondary structure of p26

WT p26 from *Artemia* and bacteria possess similar secondary structure elements, while most p26 variants exhibit increased α-helical conformations and decreased β-structure. This was especially remarkable for R114A, whose α-helical percentage increased by 8%. Residue R114 of p26 is in the β7 strand which is buried in the α-crystallin domain, and it has no apparent access to aqueous environments as indicated in Fig. 37. Mutation of R114A may disrupt the organization of the β-strand-enriched α-crystallin domain and decrease β-strand amounts, as occurred when αB-crystallin was modified in the same fashion (Bova et al., 1999a). Charged, buried amino acid residues are usually paired with a residue of opposite charge by forming salt bridges, this of structural importance, and addition or removal of these residues leads to protein conformational changes (Shroff et al., 2000). For example, T4 lysozyme is destabilized upon positioning of an unpaired, charged residue in its hydrophobic core (Xu et al., 1998). R114 of p26 is a charged, bulky residue which was replaced by the small uncharged alanine, potentially leading to destabilization of p26 structure. Although p26 oligomerization remained largely unchanged upon mutation of R114A, the secondary structure was modified and chaperone activity was compromised.

Deletion of the arginine-enriched stretch 36RPFRRRRMMRR45 in the N-terminal region led to similar consequences, within the α-helical percentage increased by 7.2%. This stretch is located in a random coil of the p26 N-terminal region, and deleting the sequence may produce new α-helices, thereby altering p26 secondary structure composition.
4.5 Intrinsic p26 fluorescence and surface hydrophobicity

Fluorescence emission is observed when electrons return from the excited state to the ground state after proteins absorb light, and is sensitive to microenvironment changes of chromophores, such as side chains of aromatic residues especially tryptophans. Protein conformational perturbations may lead to large changes in fluorescence emission, with shifts in wavelength and alterations in intensity detected upon protein unfolding and refolding. Protein fluorescence is thus an excellent marker for investigation of protein conformational changes. All p26 variants had a reduction in fluorescence intensity as compared to WT, implying altered microenvironments for tryptophan residues. This indicates that mutations introduced changes to p26 tertiary structure. A significant decrease in fluorescence intensity occurred for mutants R114A and “R”, suggesting aromatic residues especially tryptophans in each protein are in more flexible environments with increased mobility than for WT p26, as was reported for R116C αA-crystallin and R114A αB-crystallin (Kumar et al., 1999).

Surface hydrophobicity is an important parameter for chaperone-substrate interaction, although not the sole determinant (Reddy et al., 2000). ANS is a hydrophobic molecule that becomes highly fluorescent upon binding to protein hydrophobic sites or patches, providing a sensitive measure of protein hydrophobicity. ANS binding showed considerably reduced surface hydrophobicity for R114A, with changes to other α-crystallin domain mutants more moderate. This is consistent with changes in the intrinsic fluorescence spectra for mutant R114A, both indicating that this mutation caused a stronger conformational change in p26 than did other α-crystallin domain mutations. Internal deletion of multiple arginine residues had consequences similar to those caused
by the R114A mutation, although the deletion was in the N-terminal region. Arginine residues are positively charged, thus a decrease in surface hydrophobicity upon deletion of an arginine-enriched stretch, as occurred in the “R” mutant, implies reorganization of protein tertiary structure.

4.6 Nuclear migration of p26

Some sHSPs enter nuclei upon stress, although their mission within the organelle remains elusive. For instance, a subpopulation of Hsp20 migrates into the nuclei of cultured rat neonatal cardiac myocytes during heat stress, indicating a role for the protein in myocyte nuclei (van de Klundert and de Jong, 1999). αB-crystallin and Hsp27 are nuclear speckle components in unstressed, transcriptionally active cells, and Hsp27 is found in the nucleolus, implying these sHsps have similar but not completely overlapping roles in the nucleus (van den IJssel et al., 2003). Mutation R120G in αB-crystallin inhibits nuclear speckle localization, potentially affecting the nuclear functions of αB-crystallin and the RNA splicing machinery, and contributing to development of cardiomyopathy and cataract phenotypes (van den IJssel et al., 2003).

p26 is found in Artemia nuclei during diapause and upon stress including heat shock, anoxia in vivo, and low pH in vitro (Willsie and Clegg, 2001). The movement of p26 into nuclei implies association with specific nuclear components, such as protein assemblies required for DNA replication, transcription, RNA splicing and genome maintenance. The majority of nuclear p26 and a lamin share similar distributions in Artemia nuclei and are associated with the nuclear matrix, implying that p26 stabilizes nuclear matrix proteins (Willsie and Clegg, 2002). p26 sequence analysis failed to reveal a typical nuclear localization signal (NLS) consisting of either a short stretch of lysines (PKKKRKV) or a
bipartite signal where a spacer region of 10 to 20 amino acid residues separates a pair of basic clusters (KRPAATKKAGQAKKKK) (Liang et al., 1997a). In contrast, the variable p26 N-terminal region is uniquely enriched in arginine, which potentially exists as an unusual cellular localization signal, the absence of which may disrupt p26 intracellular distribution. However, removal of these arginines resulted in the strong tendency of modified p26 to translocate into transiently transfected COS-1 nuclei, indicating the arginine residues are not required for translocation. The single site substitution R114A had similar consequences for nuclear migration. This is in contrast to mutation R116C of αA-crystallin, which is equivalent to R114A of p26, but does not modify the cytoplasmic localization of αA-crystallin (Andley et al., 2002).

Interestingly, a novel R49C αA-crystallin variant is abnormally localized to the nucleus of human lens epithelial cells (HLE B-3) and fails to protect cells from staurosporine-induced apoptosis. R49C is the first dominant nuclear cataract mutation located outside the phylogenetically conserved sHSP α-crystallin domain (Mackay et al., 2003).

How p26 is translocated into nuclei remains unknown. One proposal is that decreased oligomer assembly permits p26 movement through nuclear pore complexes by simple diffusion. However, modified p26 with reduced oligomer size due to C-terminal truncation remained in the cytoplasm, and mutant R114A existing as oligomers of similar size to that of WT readily entered nuclei, suggesting that simple diffusion does not occur. In addition, p26 oligomers in heat shocked and control cysts including those from nuclei were similar, even though their mass was maintained in reduced pH, a condition promoting p26 translocation into Artemia nuclei in vivo and in vitro. Thus, the migration of p26 from cytoplasm into nuclei depends on a mechanism other than oligomer mass
reduction, although the transient dissociation into small oligomers followed by rapid reassociation within nuclei is possible.

4.7 Comparative modeling of p26

According to the model generated in this study, the p26 monomer likely has a buried N-terminal region, a partially exposed α-crystallin domain and a solvent-accessible C-terminal extension, this a general pattern of diverse sHsps (Treweek et al., 2003). Because of its hydrophobic character, the p26 N-terminal region may be involved in oligomer formation. The α-crystallin domain is an immunoglobulin-like fold as a β-sandwich, composed of eight β-strands organized into two antiparallel β-sheets. The C-terminal extension is polar, flexible and thought to maintain p26 oligomer solubility and stability, thereby minimizing protein precipitation.

There are deviations in the number and size of loops in the polypeptide backbone between Ta Hsp16.5 and the modeled p26. The most obvious difference is the loop, designated as 3 for p26, which connects β5 and β7 strands and is smaller for p26 than for Ta Hsp16.9. Ta Hsp16.9 monomers are stabilized at the dimer interface by interactions among residues of the long loop extending from the α-crystallin β-strands (van Montfort et al., 2001b). By analogy, loop 3 of p26 may be involved in structural stability and dimer formation, an interesting possibility that deserves further study.

Small oligomers such as dimers are the main building blocks of sHsps and subunit-subunit interactions are strongest along the dimerization interface (Sobott et al., 2002). Ta Hsp16.9 dimers form tetramers using β7 strands and the dimerization loops from two monomers with ordered N-terminal arms (van Montfort et al., 2001b). In this work, the single point mutation F112R which substituted the conserved hydrophobic phenylalanine
with a positively charged arginine, interfered with oligomerization and resulted in relatively small but active p26 complexes. In a similar case, the substitution of F94 of *B. japonicum* HspH, a residue corresponding to F112 of p26 and potentially involved in subunit-subunit interactions, resulted in oligomers of dramatically reduced size (Lentze et al., 2003). Thus, F112, a residue located in the $\beta$7 strand of p26 and close to loop 3, is possibly involved in inter-subunit interaction. Internal deletion “G” which lacks loop 1 did not disturb p26 oligomerization dramatically, and this variant exhibited relatively good chaperone activity. This suggests loop 1 has limited involvement in protein stability and function. The structural and functional importance of loop 2 and loop 4 in p26 remain to be determined.

4.8 Conclusions

Site-directed mutagenesis is a powerful approach to explore relationships between protein structure and function. Truncation of either the N- or C-terminus of p26 affected oligomerization and chaperone activity, although to different extents, with the N-terminal region of critical relevance for assembly of large oligomers. For single site mutations, dramatic effects were observed when the highly conserved, positively charged residue R114, located in the central $\alpha$-crystallin domain of p26, was replaced by alanine. There are several possibilities for the involvement of this conserved residue in p26 structure and function. A salt bridge formed between R114 and another residue in the protein may be important for structural integrity. Disruption of such an ionic interaction within p26 explains the altered secondary structure, disturbed tertiary structures, reduced surface hydrophobicity, and compromised chaperone activity. Alternatively, interference of ionic interactions between p26 and denaturing substrates is another possibility. Even though it
seems that binding of substrates mainly depends on hydrophobic interactions, contributions of electrostatic forces cannot be excluded.

The oligomeric range of mutant F112R was narrower than for R114A, but its chaperone activity was higher, demonstrating the absence of a positively proportional relationship between oligomerization and chaperone function. As the least effective mutants in this study, R114A and “R” exhibited significantly decreased chaperone function in vivo and in vitro, but their protection capabilities remained relatively high. This indicates the robust nature of p26 and that it can effectively accommodate critical single site mutations and multiple deletions.

p26 functions as an effective sHSP not only protecting substrate against aggregation in vitro but conferring thermotolerance on bacteria cells in vivo. It is thus reasonable to speculate that in oviparously developing Artemia cysts p26 prevents irreversible denaturation of intracellular proteins, which may be refolded by other molecular chaperones in an ATP-dependent manner once favorable conditions are resumed. However, p26 chaperone efficiency is still low in comparison with WT α-crystallins which account for approximately one-third of the total soluble lens protein and maintain refractive power (Santhoshkumar and Sharma, 2001). Of multicellular eukaryotes, Artemia cysts are among the most resistant to environmental insults, surviving years of continuous anoxia, desiccation and temperature extremes. This undoubtedly necessitates molecular chaperone function and the sHSP, p26, which is produced in large amounts in oviparous embryos, is very likely an important component of this impressive stress resistance.
Appendix I. Solutions and Recipes

Solutions for SDS-Polyacrylamide Gel Electrophoresis

1A: 39 g acrylamide
    1 g bis
    20 ml glycerol
    Add ddH₂O to 100 ml.

1B: 18.3 g Tris
    2.5 ml 20% SDS stock
    Add ddH₂O to 100 ml, pH 8.8.

1C: 0.1 ml TEMED
    Add ddH₂O to 50 ml.

2B: 6.0 g Tris
    2.5 ml 20% SDS stock
    Add ddH₂O to 100 ml, pH 6.8.

2C: 0.2 ml TEMED
    Add ddH₂O to 10 ml.

1D: 50 mg Ammonium persulfate
    Add ddH₂O to 10 ml.

Running Buffer (4L)

12.0 g Tris
57.6 g glycine
8.0 ml 20% SDS stock
Add ddH₂O to 4L. Adjust to pH 8.1-8.3.

Coomassie Brilliant Blue Staining Solution (2L)

10 g Coomassie Brilliant Blue R-250
140 ml acetic acid
800 ml methanol
Add ddH₂O to 2L. Stir 1 h, filter through miracloth.

Destaining Solution (2L)

400 ml methanol
140 ml acetic acid
100 ml glycerol
Add ddH₂O to 2L. Filter through miracloth.

4×Treatment Buffer (20ml)

0.6 g Tris
1.6 g SDS
8.0 ml glycerol
4.0 ml 2-mercaptoethanol
Add ddH₂O to 20 ml. Adjust to pH 6.8 with HCl.

Solutions for Western Blotting and Immunodetection

144
Blot Electrode Buffer (4L)
12.0 g Tris
57.6 g glycine
800 ml methanol
Add ddH₂O to 4L.

TBS Buffer (1L)
1.21 g Tris
8.18 NaCl
Add ddH₂O to 1L. Adjust pH to 7.4 with HCl.

TBS-Tween Buffer (1L)
999ml TBS buffer
1ml Tween 20

HST Buffer (1L)
1.21 g Tris
58.4 g NaCl
5.0 ml Tween 20
Add ddH₂O to 1L. Adjust pH to 7.4 with HCl.

2% Ponceau S Solution
2.0 g Ponceau S
30 g Trichloroacetic acid (TCA)
Add ddH₂O to 100 ml. Dilute 1:9 in ddH₂O for staining blots.

Developer
100 ml Developer
Add ddH₂O to 500 ml.

Fixer
124.5 ml Solution A
13.7 ml Solution B
Add ddH₂O to 500 ml.

Media for Bacteria Culture
LB medium
10 g Tryptone
5 g Yeast extract
10 NaCl
Add ddH₂O to 1L, adjust pH to 7.0 with NaOH.
1.5% agar for solid broth. Autoclave at 121 °C for 15 min.

NZY⁺ medium
10g NZ Amine (casein hydrolysate)
5 g Yeast extract
5 g NaCl
Add ddH₂O to 1L, adjust pH to 7.5 with NaOH. Autoclave at 121 °C for 15 min.

**Solutions for DNA Gel Electrophoresis**

6×DNA Loading Buffer
0.25% Bromophenol blue
0.25% Xylene cynol
15% Ficoll (type 400)
Add ddH₂O.

50×TAE
242 g Tris
57.1 ml Glacial acetic acid
100 ml 0.5 M EDTA, pH 8.0
Add ddH₂O to 1L.

**DNA Storage Solution**

TE Buffer
10 mM Tris
1 mM EDTA
Add ddH₂O. Adjust pH to 8.0 with HCl.

**Solutions for Immunofluorescence Staining of COS-1 Cells**

5×PBS
40 g NaCl
1.0 g KCl
5.75 Na₂HPO₄
1 g KH₂PO₄
Add ddH₂O to 1L, pH 7.3-7.4 as mixed.

Triton X-100 PBS
100 ml PBS
200 µl Triton X-100

Paraformaldehyde Fixative
25 ml PBS
1g paraformaldehyde.

**Solutions for Artemia Nuclei Isolation**

10×HPC Buffer (500 ml)
0.735 g CaCl₂·2H₂O
0.075 g PIPES
Add ddH₂O to 500 ml. Adjust pH to 6.5 with NaOH. Autoclave.

1×HPC Solution
50 ml  10×HPC
32 ml  Hexylene glycol
418 ml ddH₂O

10×NaCl/MgCl₂/Tris Solution (100 ml)
8.7 g  NaCl
2.8 g  MgCl₂-6H₂O
1.9 g  Tris
Add ddH₂O to 100 ml. Adjust pH to 6.5 with HCl. Autoclave.

Percoll Solution
188 ml  Percoll
25 ml  10× NaCl/MgCl₂/Tris Solution
37 ml  autoclaved ddH₂O

Solutions for Protein Purification
1×Extraction/Wash Buffer
50 mM  Na₃PO₄
300 mM NaCl
Add ddH₂O. Adjust pH to 7.5.

1×Elution Buffer
50 mM  Na₃PO₄
300 mM NaCl
150 mM Imidazole
Add ddH₂O. Adjust pH to 7.5.

Protease Inhibitor Cocktail
Solution A (5 ml)
5 mg  Leupeptin
5 mg  Soybean trypsin inhibitor
Dissolve in PIPES buffer.

Solution B (5 ml)
5 mg  Pepstatin A
10 mg  PMSF
Dissolve in methanol.
Appendix II. Prokaryotic Expression Vectors Applied in p26 cDNA Cloning
Appendix III. Eukaryotic Expression Vectors Applied in p26 cDNA Cloning

![Diagram of expression vectors](image-url)
Appendix IV. Schematic Representation of Single Amino Acid Substitutions and Internal Deletions of p26

N-terminal region  α-crystallin domain  C-terminal extension

1  60  153  192

R110G/F112R/R114A/Y116D

N-terminal region  α-crystallin domain  C-terminal extension

1  60  153  192

“G” Deletion  “R” Deletion  “TS” Deletion
Appendix V. Sequence Alignment of *Ta* Hsp16.9 and p26

The amino acid sequences of *Ta* Hsp16.9 and p26 were analyzed by CLUSTAL W (1.82). WHsp16.9, *Triticum aestivum* Hsp16.9, 1GME_A; Ap26, *Artemia franciscana* p26, AAB87967. (-) no amino acid residue; (*) identical residues; (;) conserved substitution; (.) semi-conserved substitution. The secondary structure of *Triticum aestivum* Hsp16.9 is depicted above the alignment (van Montfort *et al.*, 2001b), while the predicted secondary structure of p26 is below the alignment. Amino acid residues in red, small and hydrophobic; blue, acidic; magenta, basic; green, residues containing a hydroxyl or amine group. Residue number is indicated on the right.
References


Lee, S., Owen, H. A., Prochaska, D. J. and Barnum, S. R. 2000. HSP16.6 is involved in the development of thermotolerance and thylakoid stability in the


