Summary of Ph.D thesis

Hsp70 chaperone homolog in *Synechocystis* PCC6803: the role of DnaK2 in defence against different stresses and in modulation of the physical state of tilakoid membranes

Written by Viktória Varvasovszki

Supervised by Dr. László Vigh

Institute of Biochemistry, Biological Research Center, Hungarian Academy of Sciences

Introduction

Living organisms have the capability to adapt to sudden changes in their environment. Since most of these stimuli (temperature up- and downshift, UV-B radiation, etc.) have detrimental effects on cells, they had to develop adequate protective systems. One of these general systems is the network of molecular chaperones, the so-called stress proteins. Molecular chaperones are specialized and universally conserved proteins that bind nonnative states of other proteins and assist them to reach a functional conformation, in most cases through the expenditure of ATP (Ellis, 1997). Chaperones function not only during the normal operation of cellular processes but in addition act to limit damage caused by stresses such as heat shock. DnaK proteins of prokaryotic cells belong to the Hsp70 chaperone family which is one of the highly conserved ubiquitous groups of heat shock proteins. The members of the Hsp70 chaperone machine are DnaK (70 kDa), DnaJ (40 kDa) and GrpE (20 kDa). These chaperones act together cooperatively with the Hsp60 chaperonin family during the process of assisted protein folding. The Hsp70 proteins play important role in both stress or non-stress conditions. Generally, they are involved in protein translocation across cytoplasmic membranes, protein degradation and induced by several different stress conditions, such as temperature and osmotic damages (Georgopoulos, 1993).

Our model system is the unicellular cianobacterium, *Synechocystis PCC6803* has been used extensively as a powerful model for studying the molecular mechanisms of stress response in photosynthetic organisms. Recently, the complete nucleotid sequence its genome and the potential protein coding regions have been determined (Kaneko, 1996). On the basis of this genomic map and in contrast to many other species, there are three dnaKs (as we designated them dnaK1, 2 and 3), four dnaJ genes and only a single copy of grpE in this organism.

It is noted, that three dnaK homologs were observed in the genome of another cyanobacterium, *Synechococcus PCC7942*, all exhibiting high similarity to each of the three *Synechocystis dnaKs*. Of these three homologues dnaK2 and dnaK3 were essential in *Synechococcus*, but only dnaK2 displayed a typical heat shock response (Nimura, 2001). DnaK1 and DnaK2 proteins were detectable mainly in the cytosolic fraction, and a significant amount of *Synechococcus* DnaK3 was localized to the thylakoid membrane (Nimura, 1996). A dnaJ homolog of *Synechococcus* is located
immediately downstream of dnaK3 (as in Synechocystis) and the two genes seemed to be co-transcribed. Moreover, since the DnaJ protein, which is essential for growth even under normal conditions, is also located mainly in the thylakoid of this cyanobacterium, a specific, photosynthesis-related function has been assigned to the membrane-colocalized DnaK3 and DnaJ, in which they presumably cooperate (Ouguchi, 1997). Pairing of the membrane-associated DnaJ-DnaK tandem with physiological relevance is further supported by the finding that the DnaJ-like DjlA, associated with the inner membrane, has been identified as a DnaK co-chaperone of E. coli (Genevaux, 2001).

The physical state of cell membranes is known to be a very sensitive monitor of the most diverse environmental changes. This feature was suggested to render cell membranes an ideal location for the primary temperature stress sensor (Vigh, 1993; Carratù, 1996). Strongly supporting this hypothesis, catalytic hydrogenation of unsaturated lipids in the surface membrane of cyanobacterial cells at constant temperature resulted in the activation of transcription of a desaturase gene (Vigh, 1993), which normally is induced by cold stress (Los, 1993). The rapid increase of the membrane fluidity induced by the direct physical effect of the temperature upshift during heat shock is well documented (Mejia, 1995; Dynlacht, 1992). On the other hand, all organisms examined to date produce the evolutionary conserved HSPs when they are exposed to a sudden, sublethal increase in the ambient temperature. If the primary heat shock sensor is membrane associated, one can suppose that any modification of membrane physical properties analogous to heat-induced perturbations also could lead to changes in the level of expression of heat shock genes. Further supporting the view that thermal stress is transduced into a cellular signal at the level of membrane, in parallel with the adaptive readjustment of membrane fluidity, the threshold induction temperature of HSPs was shown to be subject of temperature acclimation in various organisms (Piper, 1995; Dietz, 1992; Lehel, 1993).

It was an intriguing finding of our workgroup that the heat shock proteins GroESL of Escherichia coli and Hsp17 of Synechocystis proved to be thylakoid-associated during heat shock (Horváth, 1998; Török, 2001). We also have shown in a model using lipid membranes and active GroESL oligimers that the chaperone binding to lipids is governed by the composition and physical state of the host membranes. Chaperones, which associated with unilamellar vesicles and stabilized the membranes at high temperature by
increasing its microviscosity, retained their capability to assist protein folding (Török, 1997). The small heat shock protein Hsp17 of *Synechocystis*, inducible by heat and ethanol, also was found to be peripherically associated with the membrane in *Leuconostoc oenos* (Jobin, 1997).

We assume that the ability of HSPs to alter membrane organization and physical parameters is a rapid, reversible and powerful tool of cellular adaptation. It may antagonize the heat-induced lipid disorganization of the membrane and thus might serve to preserve membrane structure and function during heat stress (Vigh, 1998). Furthermore, the association of HSPs with membranes likely causes inactivation of the membrane perturbation signal induced by heat, thereby turning off the heat shock genes in a feedback loop. The modulation of membrane physical order may repress transcription of heat shock genes in the heat-modified-state, explaining the known temporality of induction of the stress response. Therefore, such proposed “cross-talk” between the membrane located sensor and the heat shock response suggests the existence of a feedback mechanism of heat shock gene regulation (Vigh and Maresca, 2002).

**Specific aims**

Previously, our laboratory has been studying the Hsp60 (chaperonin) family with regard to the function of these proteins in the defence of biological membranes during heat stress. Setting out from that generally accepted view that in the diversified world of stress proteins, the role of the chaperone families are closely related to each other, the subject of this thesis was the investigation of the members of another universal group of chaperones, the Hsp70-family in *Synechocystis* PCC6803.

In this direction our specific aim was to

1. Test whether the *hsp70*-*, hsp40*- and *hsp20* homolog ORFs found on the genetic map of *Synechocystis* PCC6803 have transcriptional activity on basal level and during heat stress.
2. Observe that the protein products of the heat-inducible chaperone genes localize within parts of the cell and whether they are bound to the membranes.

3. Clone viable *hsp70*-deficient mutant *Synechocystis* strains.

4. Show if the inactivation (even partial) of a *hsp70* homolog gene in *Synechocystis* has influence on the transcriptional or translational expression of other chaperone family members.

5. Clarify the nature of physiological changes caused by different stresses (as heat and cold stress, photoinhibition or UV-B radiation) and their combination in the membrane-related photosynthetic processes.

6. Reveal whether the amount of the Hsp70 homolog proteins in *Synechocystis* cells has any influence on the physical composition or state of the thylakoid membranes.

**Experimental procedures**

**Culture conditions**

*Synechocystis* PCC 6803 cells were grown photoautotrophically to mid log phase at 30 °C in BG-11 medium at a constant photon flux density of 70 µE m⁻² s⁻¹ as described previously (Lehel et al, 1992). The *dnaK2* strain was cultured and maintained in BG-11 medium supplemented with 300 µg/ml kanamycin. Cells were heat-shocked for different time intervals, as designated.

**RT-PCR and Northern analysis**

Total RNA was purified according to Mohamed and Jansson (1989). For RT-PCR, 3 µg of total RNA was reverse-transcribed by using the SuperScript Preamplification System for First Strand cDNA Synthesis (Gibco BRL). PCR reactions (94 °C 30 s, 65 °C 1 min, 72 °C 1 min, 35 cycles) were performed with ORF-specific primers. 20 µl of each PCR products was analyzed on a 0.6% agarose gel. Denaturing agarose gel electrophoresis and Northern hybridizations with the different chaperone probes were performed as in Horváth et al. (1998).
**Primer extension**

Synthetic oligonucleotids were used to determine the transcriptional startpoint of the *dnaK2* gene. Primers were labelled with T4 polynucleotide kinase (USB) as recommended by the manufacturer. 5 pmol of labelled oligonucleotides were coprecipitated with 25 µg of total RNA. After denaturation at 95°C for 2 min, annealing was carried out at room temperature for 20 min. Extension reactions were performed at 42°C in a total volume of 15 µl containing 50mM Tris-HCl pH 8.3, 8 mM MgCl₂, 50mM NaCl, 5 mM DTT, 1 mM each of dNTPs, 20U Rnasin (Promega) and 12U AMV reverse transcriptase (USB). Reactions were stopped after 30 min, and loaded onto a standard sequencing gel.

**Generation of *dnaK2* mutant *Synechocystis* strain**

A 1.7 kb PCR fragment of the *Synechocystis dnaK2* gene was amplified with specific primers. *Synechocystis* genome being used as a template. The resulting fragment was cloned into the *HincII* site of pUC18. The integral *HincII* fragment of the *dnaK2* was replaced with the *SmaI* fragment of pUC4-KIXX vector (Pharmacia) containing the Tn5 Kan^R^ cartridge. *Synechocystis* cells were transformed with this construct according to Williams (1988). Mutant colonies were grown on BG-11 plates containing increasing concentrations of kanamycin (up to 300 µg/ml).

**Isolation of genomic DNA and Southern hybridization**

In order to isolate genomic DNA, 200 ml of *Synechocystis* culture (OD_{800}=1.6-1.8) was harvested by centrifugation at 9 000 rpm for 10 min, resuspended in 1 ml of 10 mM Tris/1 mM EDTA (TE) solution, and spun down at 12 000 rpm for 2 min The pellet was resuspended in 100 µl of 500 mM EDTA, and 900 µl of protoplast solution (10mg/ml lysozyme, 500 mM mannitol in TE) was then added. After incubation at 37 °C for 90 min, the pellet was collected by spinning at 14 000 rpm for 2 min next washed with 1 ml of TE containing 500 mM mannitol, centrifuged as before, and then resuspended in 1.2 ml of TE. The sample was divided into two portions and an equal volume of phenol was added. Following a 5-min incubation on ice, the tubes were spun at 12 000 rpm at 4 °C for 5 min The supernatants were vortexed with 600 µl of chloroform each, and the phases were separated by a short centrifugation. Incubation with 20 µl of 10 mg/ml RNAse was carried out at 37 °C for 20 min, and the samples were then treated with phenol and chloroform as above. Finally, DNA was precipitated with isopropanol at −20 °C for 2 hours. The pellet was washed with ethanol, dried and resuspended in sterile distilled water. For genomic Southern analysis, 4 µg of total DNA was digested with the indicated enzymes, and then run on
0.8% agarose gel. The DNA was next blotted onto Zetaprobe Genomic Tested membrane as recommended by the manufacturer (BioRad). Hybridization was performed with a radioactively labeled (Multiprime DNA labelling system, Amersham) dnaK2-specific probe according to the manufacturer’s instructions.

**Western analysis**
Mid log phase cultures were incubated under light (70 \( \mu E \ m^{-2} \ s^{-1} \)) at designated temperatures for 3 hours. The cells were next harvested and resuspended in SDS sample buffer. Equal amounts of proteins were separated on 8-15% SDS-PAGE, electroblotted to PVDF membrane (Millipore, Bedford, Ma) and reacted with antiserum against *E. coli* DnaK (SPA-880, Stressgene), *Synechococcus* DnaK3 (a kind gift of H. Yoshikawa), *Synechocystis* HSP17 (a kind gift of E. Vierling) and *Synechocystis* GroEL (made in our laboratory). Peroxidase-conjugated anti-mouse (DnaK, DnaK3) or anti-rabbit IgG (Sigma) was used as secondary antibody. Immunodetection was carried out by the enhanced chemiluminescence (ECL) method (Amersham, UK).

**2-Dimensional gel electrophoresis**
The isoelectronic focusing were carried out according to O’Farell (1975) and Laemli (1970).

**Determination of photosynthetic activity**
Heat inactivation of photosynthesis was measured by means of oxygen exchange with a Clarke-type oxygen electrode (Hansatech) as in Lehel et al. (1993). Control cells (grown at 30 °C) or heat-preadapted cells (exposed to 44 °C for 3 hours) were heat-shocked at 49 °C in darkness for different lengths of time, and the evolution of oxygen was then measured at the growth temperature (30°C).

**Photoinhibition assay**
*Synechocystis* cells grown at different temperatures were harvested by centrifugation (10 min at 7000 rpm) and then set to the same chlorophyll concentration (6.5 \( \mu \text{gChl/ml} \)). Volumes of 100 ml of the cells were set for treatments in special containers so that they yield a layer 10mm thick. During the photoinhibition treatment the light intensity was constant at 1200 \( \mu E \ m^{-2} \ s^{-1} \) provided by a set of 6 halogen spot-lights and the temperature was constant at the value required by the experimental protocol. Recovery of the cells was performed using 120 \( \mu E \ m^{-2} \ s^{-1} \) light intensity. During the course of the experiment 2ml samples were taken at 30min intervals for measuring the oxigene evolution in the presence of 2,5 (dimethyl) p-
benzoquinone as electron acceptor. The values recorded were plotted as a function of time using Microcal Origin Professional 5.0 software.

**UV-B treatment procedure**

UV-B irradiation was performed in open, flat glass containers in which 330 ml of cell suspension (30 µgChl/ml) of 15mm depth was continuously stirred at room temperature. Before UV-B treatment, the cells were incubated under visible light of physiological intensity and were kept at room temperature under continuous stirring. Cell cultures grown at 30°C were subjected to 22°C and treated with UV-B radiation with 312 nm maximal wavelength and 60 µE m⁻² s⁻¹ intensity. During the recovery process, samples were incubated under physiological light conditions. The photosynthetic oxygen evolution was measured by Hansatech DW2 oxygen electrode.

**Fluidity measurement of thylakoid membranes**

3 µl of a 0.2 mM solution of 1,6-diphenyl-1,3,5-hexatriene (DPH) in terahydrofuran was added to 3 ml suspension of thylakoids (isolated as in Kovács et al. 1994) containing 3 µg of Chl a, and the suspension was incubated for 40 min at 4°C. Steady-state fluorescence anisotropy was determined as in (Török et al, 1997).

**Fatty acid analysis**

The extracted lipids Sato and Murata, 1988) were methylated by using HCl/methanol (5:95, wt/wt) at 80°C for 2 hours. The esterified fatty acids were analyzed by gas chromatography (HP3396A, Hewlett-Packard) on a SP2330 column (Supelco) as in (Vigh et al., 1993).

**Summary of the results**

We examined the expression and the function of the Hsp70 homolog DnaK chaperone family in the photoautotrophic cyanobacterium, *Synechocystis PCC6803*.

1. We have established that, unlike in *Synechococcus*, **only one of the three dnaK genes, dnaK2, is transcribed and it exhibits a typical heat stress response in Synechocystis PCC6803**. Northern experiments, primer extension reactions and RT-PCR analysis were carried out with specific probes for ORFs, but we failed to detect any transcripts corresponding to either dnaK1 or dnaK3, under either normal or heat-shock conditions. To our surprise, **together with grpE, all dnaJ homolog genes are**
expressed constitutively and appear to be uninducible by high-temperature stress.

2. According to the Western analysis, we have shown that at normal temperature, the DnaK2 protein is located both in the cytoplasm and the thylakoid membrane fraction. Under heat shock conditions, the expression of this chaperone is significant and a well-defined amount of DnaK2 becomes membrane-bound.

3. We have prepared a partially dnaK2-deficient merodiploid Synechocystis strain using the method of site-directed insertional mutagenesis, since we could not disrupt all copies of dnaK2 in Synechocystis, which strongly suggests that this gene is essential for growth under normal conditions.

4. Following the photosynthetic oxigene evolution of the different strains, we have proved that the partially DnaK2-deficient cells displayed temperature sensitive phenotype, but were able to acquire thermoderotolerance.

5. The study of the photoinhibition of the mutant and wild type Synechocystis cultures adapted different growth temperatures has given the following results:
   - In case of the cells grown at 30°C, the high light treatment alone did not cause considerable changes in the photosynthetic activity of the different cultures. There was a significant decrease in this parameter in case of both strains when they were subjected to 22°C.
   - Both cultures adapted to 36°C have shown the photoinhibition effect even at their growth temperature.
   - Strains cultured at 22°C tolerated the high light treatment less at their growth temperature than at 30°C. They both showed decreased oxigene-evolution when they were returned to normal light conditions.

6. According to the analysis of UV-B tolerance, we have established that the photosynthetic oxigen-evolution of the dnaK2 mutant cells dropped to the 40% of the normal level and it was decreasing also during the recovery process and the mutant cells are finally died. In contrast, the wild type strain have been almost completely recovered.
7. The result of the experiments concerning the study of the changes in the membrane physical state using the method of fluorescence-anisotropy showed that thylakoid membranes derived from the dnaK2 mutant cells are always more rigid, than these of the wild type strain, regardless the temperature preadaptation.

8. The analysis of the fatty acid composition of the thylakoid membranes has revealed a significant increase in the level of oleic acid (18:1) in the lipids of the mutant cells in comparison with the wild type strain, while the amount of the linolic acid and the $\gamma$-linoleic acid is diminished.

In conclusion, as the photosynthesis of the dnaK2 mutant is more sensitive to high temperature, but is able to acquire thermotolerance, this might suggest that a specific composition and specific levels of stress proteins are a prerequisite of effective stress defense, but the heat-adaptation process is affected less strongly. Additional factors, such as remodeling the composition and physical state of the membranes are likely to act to explain the remaining capacity of partial dnaK mutants to develop thermotolerance by preadaptation.

According to the results of this thesis, we could prove at the first time that an organism deficient in a basic heat shock protein, DnaK, is a membrane-mutant at the same time.
Publications

*In: Genes and their products for tolerance to physical stresses in plants.* (Leone, A., Grillo, S. Eds.), Springer-Verlag, Heidelberg, pp. 21-29.


