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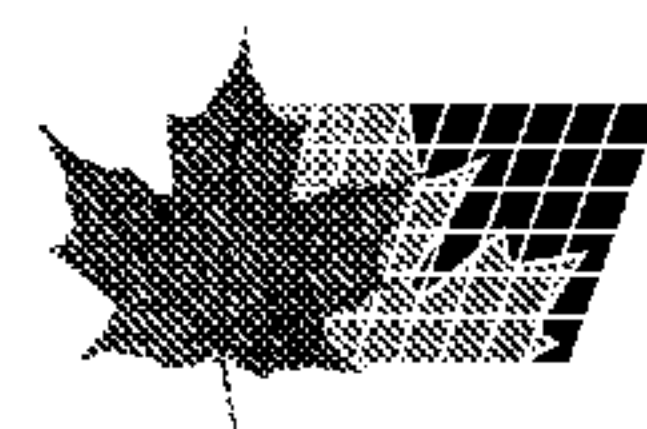
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(54) Title: TRANSGENIC ORGANISMS WITH LOWER GROWTH TEMPERATURE

(57) Abrégé/Abstract:

The present invention in general relates to the growth temperature of organisms, especially plants and microorganisms and the manipulation of the tolerable cultivation temperature. More specifically, the present invention relates to the expression of heterologous proteins in microorganisms, and especially to the heterologous expression of heat sensitive proteins in bacteria, either gram-negative or gram-positive. In a first aspect, the present invention provides a method for manipulation of cells and the resultant cells, characterized in that at least one gene from a psychrophilic micro organism coding for at least one chaperone or chaperonin is expressed. Such cells are selected among cultivated eukaryotic cells, i.e. animal and plant cells and entire plants, gram-negative and gram-positive bacteria, fungi and yeasts.



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(54) Title: TRANSGENIC ORGANISMS WITH LOWER GROWTH TEMPERATURE

(57) **Abstract:** The present invention in general relates to the growth temperature of organisms, especially plants and microorganisms and the manipulation of the tolerable cultivation temperature. More specifically, the present invention relates to the expression of heterologous proteins in microorganisms, and especially to the heterologous expression of heat sensitive proteins in bacteria, either gram-negative or gram-positive. In a first aspect, the present invention provides a method for manipulation of cells and the resultant cells, characterized in that at least one gene from a psychrophilic micro organism coding for at least one chaperone or chaperonin is expressed. Such cells are selected among cultivated eukaryotic cells, i.e. animal and plant cells and entire plants, gram-negative and gram-positive bacteria, fungi and yeasts.

Transgenic organisms with lower growth temperature

The present invention in general relates to the growth temperature of organisms, especially plants and microorganisms and the manipulation of the tolerable cultivation temperature. More specifically, the present invention relates to the expression of heterologous proteins in microorganisms, and especially to the heterologous expression of heat sensitive proteins in bacteria, either gram-negative or gram-positive.

It is generally known that the mesophilic host *E. coli* is suitable for expression of a wide range of heterologous proteins, both intracellular and secreted. When expression of proteins is induced on a large scale in *E. coli*, problems are often encountered due to the production of intracellularly agglomerated protein, which is enzymatically inactive. The reason for the inactivity of these agglomerates, also called inclusion bodies, is the misfolding of the polypeptide chains which are intensely synthesized after induction or which cannot attain their natural active conformation when expressed in *E. coli* or other hosts. Various attempts for the *in vitro* folding of purified agglomerated protein have been proposed and are used on an industrial scale. However, the folding *in vitro* requires numerous processing steps to produce enzymatically active protein, i.e. protein in its correct folding structure.

Feller, G. et al., Appl. Env. Microbiol, p. 1163-1165 (1998) describe the expression of the psychrophilic α -Amylase from the Antarctic psychrophile *Alteromonas haloplanktis* in *E. coli* by lowering the cultivation temperature of the transformed expression host to 18 °C. It was demonstrated that the expression of active enzyme could be increased over that of the wild type *Alteromonas haloplanktis* and furthermore that the recombinantly produced enzyme had the same kinetic parameters as the wild type enzyme produced at 4 °C. The authors therefore concluded that the psychrophilic enzyme is correctly folded when expressed recombinantly in *E. coli* at 18 °C.

A drawback of low cultivation temperatures of mesophilic host organisms is the dramatically reduced growth rate, and, consequently a reduced synthesis rate of the heterologous protein.

In view of the known state of art the present invention aims at providing organisms, especially micro organisms which are capable of growth at lower temperature, for instance at temperatures similar to the range of growth temperatures of psychrophiles. Furthermore, the present invention aims at providing an expression system for heterologous proteins in micro organisms which are capable of producing correctly folded protein, i.e. protein with a structure which retains the enzymatic or interactive activity of the native wild type protein.

In a first aspect, the present invention provides a method for manipulation of cells and the resultant cells, characterized in that at least one gene from a psychrophilic micro organism coding for at least one chaperone or chaperonin is expressed. Such cells are selected among cultivated eukaryotic cells, i.e. animal and plant cells and entire plants, gram-negative and gram-positive bacteria, fungi and yeasts.

In a second aspect, the present invention provides a method for producing heterologous proteins in micro organisms as well as the micro organisms themselves, i.e. animal and plant cells, gram-negative and gram-positive bacteria, fungi and yeasts, characterized in that at least one gene from a psychrophilic micro organism coding for at least one chaperone or chaperonin is expressed. The heterologous proteins to be expressed comprise gene-products from mesophilic as well as psychrophilic organisms.

In a third aspect, there is provided a method for in vitro folding of aggregated or misfolded protein, characterized in that at least one chaperone or chaperonin from a psychrophilic

micro-organism is contacted with the aggregated or misfolded protein in presence of necessary nucleotides.

In a fourth aspect, DNA and amino acid sequences are provided for native chaperonins Cpn10 and Cpn60 of *Oleispira antarctica* along with mutant chaperonins with altered characteristics as well as methodology and guidelines for cloning, expressing and adapting chaperones for enhancing the expression of heterologous proteins, and especially thermo-sensitive heterologous proteins, in their native conformation in host organisms, for adapting host organisms to lower growth temperatures and for re-folding, at low thermal conditions, denatured protein *in vitro*.

The invention is illustrated in relation to the chaperones from the psychrophilic bacterium *Oleispira antarctica*, which have been designated Cpn60 and Cpn10 and which are co-operative in their wild type forms.

The standard growth temperature of the widely used expression host bacterium *E. coli* is 37 °C with an experimental lower limit of approximately 15°C. The theoretical lower limit can be calculated to 7.5 °C by the square-root growth model of Ratkowsky et al., J. Bacteriol., 1222-1226 (1983).

It has now been discovered that the expression of chaperonin Cpn60 and its co-operating co-chaperonin Cpn10 in *E. coli* decreases the actual growth temperature to 0 to 7 °C with a theoretical minimum of –13.7 °C. The growth rate of these coldness-adapted *E. coli* reaches 0.28 /h at 8 °C and 0.22 /h at 4 °C.

When heterologous genes are expressed in *E. coli* which harbour both the chaperonin Cpn60 and its co-chaperonin Cpn10, then the expression can take place at significantly lower cultivation temperatures, i.e. 0 to 7 °C and thermo-sensitive protein can be produced by *E. coli* in its native conformation, e.g. enzymatically active.

It was shown that Cpn60 adapts its tertiary structure in a temperature-dependent manner. At the normal growth temperature of *O. antarctica* of 4 to 10 °C, the predominant tertiary structure of Cpn60 is a heptameric single ring of identical subunits along with lower molecular weight dissociation intermediates. In a temperature range of 12 to 24 °C, Cpn60 is

predominantly present as a 14-mer, consisting of two stacked rings, each comprised of seven identical subunits. However, the dissociation of the stacked heptameric rings at lower temperature is dependent on the presence of nucleotides of adenine, citidine, uridine or guanidine. It was shown that the activity of Cpn60 to refold denatured proteins is dependent on nucleotides, e.g. adenosine triphosphate (ATP).

From the following examples, analytical data of the chaperonins Cpn60 and Cpn10 of *Oleispira antarctica* will be apparent. As a consequence, the skilled persons will be enabled to identify and clone homologous genes encoding chaperones from either *Oleispira antarctica* itself or from other psychrophilic microorganisms, e.g. other eubacteria or archaeobacteria using the sequence information given for the chaperonins of *O. antarctica* and the cloning strategy below or other known procedures. As source organisms for chaperones with similar functional properties as and/or homology to those specifically disclosed herein, the following can be used: *Moraxella*, *Alteromonas haloplanktis*.

Homologous chaperones, derivatives or mutant forms of the chaperonins Cpn60 and Cpn10 of *Oleispira antarctica* which retain the functional properties in respect of the lowering of the growth temperature of a transformed mesophilic host organism and/or in respect of the chaperone activity to refold denatured proteins extracellularly, for example in vitro, are also accessible on the basis of the examples given below.

When screening other psychrophilic organisms for chaperonins homologous to those of *O. antarctica*, sequence alignment studies and comparisons can be employed, for example exploiting homologies of Cpn60 of *O. antarctica* to GroEL of *E. coli* and Cpn60 of *Paracoccus denitrificans* in order to identify residues with an influence on substrate specificity and/or conformation of the chaperonin. Variants can be constructed in accordance with the methodology presented below for producing mutant chaperonins of *O. antarctica*.

Derivatives or mutant forms of *O. antarctica* can readily be obtained by genetically engineering the DNA sequence of the genes encoding Cpn60 and/or Cpn10. Such mutants may have altered substrate binding specificities, altered nucleotide binding properties or an altered secondary or tertiary structure or altered interaction parameters of a chaperonin with its co-chaperonin, e.g. of mutant Cpn60 with mutant Cpn10. It can be expected that mutations introduced at sites responsible for substrate binding change the substrate specificity,

mutations at sites responsible for the association of subunits to the single ring conformation change the single ring structure, and mutations at sites responsible for the interaction of ring structures with one another to stacked rings change the conformation and consequently the stability and the temperature-dependent association of rings and ultimately their activity and/or substrate specificity.

Short description of the figures:

Figure 1 shows the amino acid sequences of native Cpn10 (SEQ ID No 1) and Cpn60 (SEQ ID No 2) of *O. antarctica*.

Figure 2 shows the DNA sequences of native Cpn10 and Cpn60 (SEQ ID No 3) of *O. antarctica*.

Figure 3 shows the amino acid sequence of esterase (SEQ ID No 4) of *O. antarctica*.

Figure 4 shows the DNA sequence of esterase (SEQ ID No 5) of *O. antarctica*.

Figure 5 shows the amino acid sequences of native Cpn10 (SEQ ID No 6) and Cpn60 (SEQ ID No 7) of *O. antarctica* and esterase (SEQ ID No 8) of *O. antarctica*.

Figure 6 shows the DNA sequence of the expression cassette of native Cpn10 and Cpn60 and of *O. antarctica* with the esterase (SEQ ID No 9) of *O. antarctica*.

Figure 7 shows the amino acid sequences expressed from the expression vector coding for the co-expression of native Cpn10 (SEQ ID No 10) and the stabilized single ring mutant Glu461Ala/Ser463Ala/Val464Ala of Cpn60 (SEQ ID No 11) of *O. antarctica* with the esterase (SEQ ID No 12) of *O. antarctica*.

Figure 8 shows the DNA sequence of the expression vector SEQ ID No 13) coding for the co-expression of native Cpn10 (SEQ ID No 10) and the stabilized single ring mutant Cpn60 (SEQ ID No 11) of *O. antarctica* with the esterase (SEQ ID No 12) of *O. antarctica*.

Figure 9 shows the amino acid sequences of native Cpn10 (SEQ ID No 14) and of the stabilized single ring mutant Glu461Ala/Ser463Ala/Val464Ala of Cpn60 (SEQ ID No 15) of *O. antarctica*.

Figure 10 shows the DNA sequence of the stabilized single ring mutant Glu461Ala/Ser463Ala/Val464Ala of Cpn60 (SEQ ID No 16) of *O. antarctica*.

The examples below describe the methodology of cloning the chaperonin genes Cpn60 and Cpn10 from *O. antarctica* as well as generating mutant and variant chaperones. Therefore, the skilled person will be instructed on how to influence stability and activity parameters of chaperones having similar functional parameters as those described specifically.

Example 1: Cultivation of *Oleispira antarctica* and isolation of Cpn60 and Cpn10:

O. antarcitca RB-8 were cultivated at 4 °C in 400 mL ONR7a medium (Dyksterhouse et al., I. J. Sys. Bacteriol. 116-123 (1995)) supplemented with 0.2 vol % Tween 80 (Sigma Chemicals) to an optical density of 0.7 to 0.8 at 600 nm, harvested by centrifugation (4500 x g, 30 min, 4 °C) and frozen at -20 °C. Thawed cells were suspended in two-fold volume of buffer containing 50 mM Tris-HCl, pH 7.0, 10 mM MgCl₂, 10 mM KCl, 2 mM EDTA, 1 mM DTT, 1 tablet protease inhibitor cocktail (Roche) and Dnase I grade II, then homogenized in a French press at 68.95 bar (1000 psi), centrifuged (35,000 x g, 35 min, 4 °C) and the supernatant concentrated by ultrafiltration by centrifugation against a membrane with a cut-off at 10 kDa (Centricon, Amicon Inc.) to 2 mL.

The purification was by elution from a Mono-Q HR 10/10 ion exchange column, equilibrated with 50 mM Tris-HCl, pH 7.0, 10 mM MgCl₂, 10 mM KCl, with a 0 – 1 M NaCl gradient in the same buffer for 200 min at 2.0 mL/min. Fractions containing Cpn60 and Cpn10 were identified by SDS-PAGE with subsequent blotting and immunodetection with a polyclonal antibody directed against the N-termini of both Cpn10 and Cpn60 as well as by an activity test of the refolding activity using chemically denatured rhodanese as the substrate. Active fractions were eluted at 0.30 – 0.45 M NaCl, pooled, dialyzed against 50 mM Tris-HCl, pH 7.0, 10 mM MgCl₂, 10 mM KCl, 150 mM NaCl and concentrated by centrifugation against a membrane with a cut-off at 10 kDa (Centricon, Amicon Inc.). The concentrated pooled fractions were purified by gel-filtration on a Superdex 200 16/60 column, equilibrated in 50

mM Tris-HCl, pH 7.0, 10 mM MgCl₂, 10 mM KCl, 150 mM NaCl at 4 °C at a flow rate of 1 mL/min. Fractions were pooled, dialyzed against 50 mM Tris-HCl, pH 7.0, 10 mM MgCl₂, 10 mM KCl, concentrated by ultrafiltration by centrifugation (10 kD cut-off membrane), then further purified by ion-exchange chromatography on a Mono-Q HR 5/5 column at 0.5 mL/min of a linear gradient (20 mL) of 0 – 1 M NaCl and fractions pooled.

Example 2: Cloning and characterization of chaperonin Cpn60 and Cpn10 of *Oleispira antarctica*.

Starting from *Oleispira antarctica* RB-8, available as DSMZ accession No 14852, a comprehensive genomic library comprising 5 x 10⁸ phage particles /µL, total of 8 mL with an average insert size of 7.5 kb was created using the ZAP Express Kit of Stratagene according to the manufacturers instructions. Briefly 7.5 kb fragments of genomic DNA from *O. antarctica* were cloned into a plasmid using the well known procedure of the ZAP Express kit (Stratagene).

Degenerate forward primer 5'-GCI GCI GGI ATG AAY CCI ATG G (Seq ID No 17) and reverse primer 5'-CCI CCI CCI GCI ACI ACI CCY TC (Seq ID No 18) were designed on the basis of the sequences analyzed from purified chaperonin fragments SVAAAGMNPMDLQR (Seq ID No 19) and VEEGVVAGGGVAAL-LR (Seq ID No 20), respectively. These primers were used for PCR amplification of the genomic DNA of strain RB-8 (Smits et al., *Environ. Microbiol.* 307-317 (1999)). The amplified fragment of approximately 930 bp was cloned into plasmid pCR2.1 (Invitrogen). Briefly, the fragment was purified from an agarose gel and ligated into the pCR2.1 plasmid. Subsequent sequencing of the cloned PCR product revealed a high similarity of its deducible amino acid sequence to the Cpn60/Hsp60 family.

The cloned PCR fragment was then excised from vector pCR2.1, labelled with digoxigenin (DIG DNA Labelling Kit, Roche Diagnostics) and used as a hybridisation probe to screen the lambda phage genomic library of RB-8. From phage plaques that hybridized and identified using the immuno-detection of digoxigenin (Roche Diagnostics), the cloned DNA fragments were rescued with the infection of helper phage f1 to give plasmids pBK-CMV. The inserted DNA fragments of pBK-CMV was sequenced. The amino acid sequence of Cpn60 and Cpn10 translated from the DNA sequence (shown in Figure 2) are given below in Figure 1.

Example 3: Simultaneous expression of Cpn60 and Cpn10 of *Oleispira antarctica* in *E. coli*.

The expression of Cpn60 and Cpn10 was induced from the vector pPST26, originating from the lambda clone No 26 that hybridized with the DNA probe for cpn60, designated pBK26, which was deleted upstream by a restriction with PstI. The expression vector carries both genes in the orientation that enables their expression from the P_{lac} -promoter. For overexpression, *E. coli* cells XL-1 Blue MRF' were transformed with pPST26, grown in LB medium to an optical density at 600 nm of 0.6 to 0.8 and induced by the addition of IPTG (isopropyl- β -D-thiogalactopyranoside) to a final concentration of 2 mM. Cells were harvested by centrifugation 2 to 3 hours after induction, resuspended in 2 mM EDTA, 1 mM dithiothreitol (DTT), 1 tablet protease inhibitor cocktail (Roche) and Dnase I grade II, incubated on ice for 30-45 min and sonicated for 4 min total time. The cell lysate was centrifuged at 10,000 x g, 30 min, 4 °C and the soluble supernatant was dialyzed overnight against 50 mM Tris-HCl, pH 7.0, 10 mM MgCl₂, 10 mM KCl, then concentrated by ultrafiltration by centrifugation against a membrane with a cut-off at 10 kDa (Centricon, Amicon Inc.) to 2 mL.

Example 4: Mutants of Cpn60 with increased stability of the tetradecameric structure

Mutants of Cpn60 were constructed which show an increased stability of the two stacked heptameric ring conformations, forming the tetradecamer of Cpn60. Mutations were introduced by site-directed mutagenesis using primers in PCR amplifications which carry the desired nucleotide exchanges to yield a different codon, as it is generally known to the skilled person. For introducing the mutations into the wild type gene of Cpn60 from *O. antarctica*, the following oligonucleotides were used in PCR:

Lys468Thr: 5'-GGT GGT CAG TGG TTG TTG ATA CAG TGA AAT CTG GCA CAG-3' (Seq ID No 21) and 5'-CCT GTG CCA GAT TTC ACT GTA TCA ACA ACC ACT GAC C-3' (Seq ID No 22)

Ser471Gly: 5'-GGT GAT AAA GTG AAA GGT GGC ACA GGT AGC-3' (Seq ID No 23) and 5'-GCT ACC TGT GCC ACC TTT CAC TTT ATC AAC-3' (Seq ID No 24)

Lys471Thr: 5'-GGT CAG TGG TTG TTG ATA CAG TGA AAG GTG GCA CAG **GTA**
GCT TTG G-3' (Seq ID No 25) and 5'-CCA AAG CTA CCT GTG CCA CCT TTC **ACT**
GTA TCA ACA ACC ACT GAC C-3' (Seq ID No 26)

GLU461ALA/SER463ALA/VAL464ALA: 5'-CCT AAC GCA GGT GCT GCA **GGG** GCA
GCG GTT GTT GAT AAA GTG-3' (Seq ID No 27) and 5'-CTC TTT ATC AAC AA**C** CGC
TGC CCC TGC AGC ACC TGC GTT ACC-3' (Seq ID No 28).

Firstly, Lysin 468 was exchanged for a Threonin, and secondly, Serin 471 was exchanged for a Glycine and thirdly, a double mutant Lys468Thr/Ser471Gly was produced. These plasmids were expressed in *E. coli* strain XLOR as described in Example 3 with the appropriate antibiotic kanamycin added. All three mutants demonstrated a more stable association of the heptameric rings to the tetradecameric stacked ring structure during native gel electrophoresis (7.5 % PAGE, poly acryl amide gel electrophoresis according to Laemmli).

Example 5: Mutant of Cpn60 with decreased stability of the tetradecameric structure

As a fourth variant, a mutant with three amino acid substitutions was produced as above, introducing the mutations Glu461Ala, Ser463Ala and Val464Ala. This mutant was shown in native PAGE to have a single ring heptameric conformation with an apparent mass of approximately 400 kDa, which corresponds to the wild-type single heptameric ring conformation.

The above described mutants were purified as described in Example 1. The analysis of the mutant proteins by circular dichroism demonstrated that the triple mutant Glu461Ala/Ser463Ala/Val464Ala as well as the double mutant Lys468Thr/Ser471Gly were not destabilized in their respective overall secondary conformations in comparison to the wild-type Cpn60.

Using the measurement of peptide ellipticity at 220 nm to monitor the loss of secondary structure due to increasing temperature, it could be demonstrated that the stabilized double ring mutant Lys468Thr/Ser471Gly has an increased temperature stability at 45 – 55 °C and a rate for ATP hydrolysis 1.3 to 1.6 times higher than the wild type and the stabilized single

ring mutant Glu461Ala/Ser463Ala/Val464Ala, the latter having temperature stability up to 24 – 28 °C.

Similar results for an increased temperature stability at increased hydrolysis rates for ATP are obtained for mutants of Cpn60 of *O. antarctica* Leu468Thr and Ser471Gly, which each show an increased stability of the double ring structure compared to the wild type, at least in presence of ATP.

With a view to adapting chaperones to the cultivation temperature, it can therefore be concluded that single ring variants of chaperonins, especially of Cpn60 of *O. antarctica* RB-8, are essential for producing protein in its correct conformation at low temperatures, e.g. at 0 – 8 °C, whereas variants of chaperonins, especially of Cpn60 of *O. antarctica* RB-8, are essential for producing protein in its correct conformation at higher temperatures, e.g. at above 10 - 12 °C.

Example 6: Chaperone activity of Cpn60 and its mutants *in vitro*

The wild type Cpn60 from *O. antarctica*, without its co-chaperonin Cpn10 has refolding activity under both physiological and temperature stress conditions of 0 – 30 °C, which is in correlation with the range of the growth temperature of the source organism.

When used in *in vitro* folding procedures, isolated wild type Cpn60 refolds denatured protein as a single ring at 4 to 8 °C, whereas at >12 °C, the predominant conformation of the active form is the double ring complex.

When using chemically denatured mtMDH (mitochondrial malate dehydrogenase) (Nielsen et al., Mol. Cell 93-99 (1995)) as the substrate for *in vitro* refolding procedures, it was found that spontaneous refolding occurred at about 16 – 24 % at a temperature range of 4 to 30 °C in 50 mM Tris-HCl, pH 7.0, 50 mM MgCl₂, 50 mM KCl.

When testing the refolding activity of the wild type Cpn60, the stabilized single ring mutant Glu461Ala/Ser463Ala/Val464Ala, and the stabilized double ring mutant Lys468Thr/Ser471Gly under the same conditions with added Cpn10 and ATP (1 mM), it was found that the stabilized single ring mutant catalysed refolding at 4 to 8 °C at 70 – 80%, but

was inactive for refolding at above 10 °C. The behaviour of the stabilized single ring mutant led to the conclusion that at the low temperature, the co-chaperonin Cpn10 could bind even to the heptameric conformation, whereas at elevated temperature presumably Cpn10 could not be released from this heptameric single ring.

In contrast to the stabilized single ring mutant, the stabilized double ring mutant Lys468Thr/Ser471Gly yields a low refolding effect (20%) at low temperature of 4 °C, approximately four times lower than the stabilized single ring mutant (80%).

Wild type Cpn60 was active from 4 to 20 °C, showing a higher refolding activity at lower temperatures.

Although the wild type Cpn60 as well as the stabilized single ring mutant could not catalyse refolding at temperatures of above 25 – 30 °C, the stabilized double ring mutant was active, i.e. at 28 °C the refolding yield was more than ten times that of the wild type Cpn60 and that of the stabilized single ring mutant, with activity up to 36 °C.

From these results it can be inferred that the temperature range, in which a chaperonin is active for refolding denatured protein can be influenced by its conformation as a stabilized single or stabilized double ring variant. Furthermore, at least the amino acids in homologous positions as Lys468 and Ser 471, respectively, in Cpn60 of *O. antarctica* are responsible for this temperature range of chaperone activity.

Furthermore it is demonstrated that even as a single ring structure, Cpn60 catalyzes the refolding of denatured protein when in combination with its co-chaperonin. Further experiments using a competition assay of radio-labelled Cpn10 and non-labelled Cpn10 at 4 °C and 20 °C demonstrated that the co-chaperonin is released from the stabilized single ring mutant of Cpn60 only at the lower temperature and in presence of denatured protein substrate (denatured mtMDH). At the higher temperature, the bound co-chaperonin was not released from this mutant Cpn60 when denatured protein substrate was added. When testing the stabilized double ring mutant of Cpn60 under the same conditions, the release of the co-chaperonin was independent from the temperature and lower by a factor of 4 – 5 compared to the stabilized single ring mutant.

Example 7: Influence of variant chaperones on the growth of transformed host organisms

The effect of the presence of a gene product coding for the wild type chaperone from a psychrophilic organism as well as of variant chaperones thereof have been assessed for the growth of *E. coli* at varying temperatures. *E. coli* have been transformed with a plasmid bearing, under the control of an IPTG inducible lac promoter the gene for wild type chaperonin Cpn60 and its co-chaperonin Cpn10 of *O. antarctica*, the stabilized single ring mutant Glu461Ala/Ser463Ala/Val464Ala, and the stabilized double ring mutant Lys468Thr/Ser471Gly, respectively. As shown in Figure 9, *E. coli* without heterologous chaperone grew at 15 °C only to some extent (OD_{600} after 48 h incubation $0.74 +/- 0.24$), at 4 °C, no growth was observed. Only *E. coli* expressing the wild type chaperonin Cpn60 and Cpn10 or the stabilized single ring mutant Glu461Ala/Ser463Ala/Val464Ala grew at 4 °C up to an $OD_{600} = 1.5 +/- 0.14$ after 48 h.. However, at 15 °C, the stabilized single ring mutant Glu461Ala/Ser463Ala/Val464Ala ($OD_{600} = 0.75 +/- 0.10$) did not enhance viability, but wild type Cpn60 and Cpn10 ($OD_{600} = 1.45 +/- 0.12$) and the stabilized double ring mutant Lys468Thr/Ser471Gly ($OD_{600} = 1.63 +/- 0.24$) allowed for an enhanced growth.

From these findings it is clear that when screening for or constructing variant chaperonins it has to be considered that at lower temperatures for growth of host cells and for refolding of denatured protein to their native conformation the effectiveness, i.e. activity of the chaperonin is dependent on its tertiary conformation. As has been detailed for chaperonin Cpn60 and Cpn10 of *O. antarctica*, the active conformation of Cpn60 at low temperatures, i.e. below 12 °C is the single heptameric ring structure, whereas at higher temperatures, i.e. above 12 °C, the active conformation of Cpn60 is the double ring structure.

Since this temperature dependence of the chaperonin activity has been elucidated for Cpn60 of *O. antarctica*, the skilled persons will be able to apply these findings to homologous chaperones from psychrophiles and to mutant and variant forms thereof using standard methodology. In detail, the influence of the amino acids Lys468 and Ser471 on the stabilisation of the association of the heptameric rings of Cpn60 has been demonstrated. Accordingly, the influence on stability of the tertiary conformation of their functionally equivalent residues in homologous chaperonins is evident and can be used to manipulate the tertiary structure and, as a consequence the temperature dependence of homologous and variant chaperones.

As one mutant form of a homologous chaperonin, the GroEL from *E. coli* was mutated doubly by Thr468Lys and Gly471Ser to arrive at a chaperonin with altered activity at temperatures below that of unmodified GroEL. The sequence of GroEL of *E. coli* is available as accession No P06139 at the Swissprot databank.

Example 8: Chaperone activity of Cpn60 and its mutants *in vivo*

The influence of chaperones from psychrophilic organisms on the expression of protein in its native, i.e. non-denatured conformation is demonstrated on the example of the temperature sensitive esterase from *O. antarctica* in *E. coli* with and without presence of the heterologous chaperonin Cpn60 from *O. antarctica*.

The esterase gene from *O. antarctica* RB-8 (DSMZ No. 14852T) was cloned from the genomic lambda library described in example 2. Detection of clones expressing active esterase was after infection of *E. coli* XL1-Blue MRF' and incubation by overlay with an aqueous solution containing per mL 60 µL naphthyl acetate (20 mg/mL acetone), 0.25 mM IPTG and 16 µL of Fast Blue RR (80mg/mL dimethyl sulfoxide). Positive clones exhibited a brown halo after about 2 h incubation and were isolated after consequent phage particle dilution, infection of *E. coli* and halo detection. The inserted DNA sequence from positive clones was rescued by helper phage infection and sequenced. The amino acid sequence and the DNA sequence of the esterase from *O. antarctica* are given in Figures 3 and 4, respectively.

For expression experiments of the thermo sensitive esterase as cloned above, the esterase gene (est) was cloned into an *E. coli* expression vector under the control of a lac-promoter, alternatively in combination with an expression cassette under a lac-promoter of the wild type chaperonin Cpn60 and its co-chaperonin Cpn10 (cpn10::cpn60::est, see Figures 5 and 6 for amino acid and DNA sequences, respectively) and in combination with the stabilized single ring mutant Glu461Ala/Ser463Ala/Val464Ala and Cpn10 (cpn10::stabilized single ring mutant Glu461Ala/Ser463Ala/Val464Ala::est, see Figures 7 and 8 for amino acid and DNA sequences, respectively), under control from a lac-promoter as well. Standard PCR-cloning procedures with primers designed according to the established gene sequences were used.

E. coli cells TOP10 (Invitrogen) were transformed with the above plasmids by electroporation and incubated in LB broth containing the appropriate antibiotic kanamycin at 4, 8, 10, 15, 20, 30, and 37 °C each. Induction was done with 1 mM IPTG. Once the cultures reached maximal esterase activities, the cells were harvested by centrifugation, resuspended in 50 mM Tris-HCl, pH 7.0, 50 mM MgCl₂, 50 mM KCl containing one protease inhibitor tablet (Roche Diagnostics) and Dnase I grade II, incubated on ice for 30 – 45 min and sonicated for a total of 4 min. After centrifugation at 10,000 x g for 30 min at 30 , the supernatant was dialysed overnight against 50 mM Tris-HCl, pH 7.0, 50 mM MgCl₂, 50 mM KCl and concentrated by ultrafiltration with a 10 kDa cut-off membrane (Centricon, Millipore).

Purification was by ion-exchange chromatography on Mono-Q HR 10/10, equilibrated with 50 mM Tris-HCl, pH 7.0, 50 mM MgCl₂, 50 mM KCl, elution with a 0 – 1 M NaCl gradient in the same buffer for 200 min at 0.5 mL/min. Esterase containing fractions were eluted at about 0.3 M NaCl and pooled. After changing the buffer by dialysis and ultrafiltration, the pool was loaded onto a Resource 15PHE hydrophobic chromatography column, previously equilibrated with 50 mM Tris-HCl, pH 7.0, 1 M (NH₄)₂SO₄, washed with a decreasing 1.0 – 0 linear (NH₄)₂SO₄ gradient in 10 mM Tris-HCl, pH 7.0. The fractions active for esterase were pooled, dialysed against 10 mM Tris-HCl, pH 7.0, 150 mM NaCl and concentrated as before. Finally, gel filtration was performed on a Superose 12 HR 10/30 in 10 mM Tris-HCl, pH 7.0, 150 mM NaCl at 4 °C and 0.4 mL/min. N-terminal sequencing was employed to corroborate the identity of the enzyme.

The results are given in table 1 below.

Table 1:

Growth temperature [°C]	Without additional chaperonin		cpn10::cpn60::est		cpn10::stabilized single ring mutant Glu46Ala/Ser463Ala/Val464Ala::est	
	Protein expression ¹	Esterase activity ²	Protein expression ¹	Esterase activity ²	Protein expression ¹	Esterase activity ²
37	2 - 5	12	2 - 5	12	2 - 5	12
30	2 - 5	127	2 - 5	127	2 - 5	127
20	<2	504	2 - 5	768	<2	528
15	<1	1560 ³	2 - 5	2040 ⁴	<1	1400 ³
10	n.d.	n.d.	2 - 5	2304 ⁵	2 - 5	2304 ⁶
8	n.d.	n.d.	2 - 5	2400 ⁵	2 - 5	2400 ⁶
4	n.d.	n.d.	2 - 5	2400 ⁵	2 - 5	2400 ⁶

¹ Expression as % by weight of total cell protein² Whole cell activity in µmol tributyrin per min per g cell lyophilisate, measured in Tris-HCl buffer, pH 8.5 at 20 °C³ Growth rate of 0.15 /h, late-exponential phase reached in 60 h⁴ Growth rate of 0.46 /h, late-exponential phase reached in 24 h⁵ Growth rate of 0.22 – 0.28 /h, late-exponential phase reached in 30 h⁶ Growth rate of 0.5 /h, late-exponential phase reached in 20 h

n.d. No growth observed or growth rate below 0.01 /h

When analysing the heterologously expressed cloned esterase from *O. antarctica* in *E. coli* under identical conditions except for the presence of IPTG inducible chaperonin genes it was found that without additional chaperonins in *E. coli*, at 37 °C the hydrolytic activity was very low, i.e. 190 µmol/min/g. When reducing the cultivation temperature, higher specific activity of the esterase was observed. However, the higher specific activity correlated with a dramatically reduced expression level at lower temperatures.

In *E. coli* also expressing the chaperonin from *O. antarctica* in its wild type and stabilized single ring mutant forms, respectively, esterase was expressed at much higher level, with the wild type chaperonin reaching the late-exponential growth phase after 30 h and the stabilized single ring mutant chaperoning after only 20 h.

When comparing the different chaperonins expressed, it becomes clear that their structure greatly influences their activity at different temperatures. In detail, the stabilized single ring mutant Glu461Ala/Ser463Ala/Val464Ala was only efficient for production of active esterase at below 10 °C.

At temperatures above 20 °C, the esterase activity was significantly lower for all transformants and it is assumed that this is due to the instability of the esterase at these temperatures. However, when analysing the fluorescence intensity of esterase obtained from cultures at 4 °C and 37 °C for both chaperonin transformants, i.e. wild type cpn10::cpn60::est and cpn10::stabilized single ring mutant Glu461Ala/Ser463Ala/Val464Ala::est, the fluorescence intensity of esterase for each transformant measured for the 4 °C culture were five times higher than those for the 37 °C culture. Therefore, misfolding of the thermo-sensitive esterase due to its expression at 37 °C can practically ruled out but higher fluorescence values for the esterase expressed at 4 °C indicate a better folding state, correlating with a higher specific esterase activity.

Claims:

1. Microorganism, characterized by the presence of a DNA sequence encoding a functional chaperone of a psychrophilic bacterium.
2. Microorganism according to claim 1, characterized in the DNA sequence encoding a functional chaperonin of a psychrophilic bacterium.
3. Microorganism according to claim 1, characterized in the DNA sequence encoding the chaperonin Cpn60 and/or Cpn 10 (SEQ ID No 1 and/or 2) of *Oleispira antarctica*.
4. Microorganism according to claim 1, characterized in the DNA sequence encoding a functional homolog of the chaperonin Cpn60 and/or Cpn10 of *Oleispira antarctica* (Seq ID Nr. 1 and/or 2) from a psychrophilic bacterium.
5. Microorganism according to claim 4, characterized in that the psychrophilic bacterium is selected from the group consisting of *Moraxella*, and *Alteromonas haloplanktis*.
6. Microorganism according to claim 1, characterized in the DNA sequence encoding a functional mutant of the chaperonin Cpn60 and/or Cpn 10 (Seq ID No 1 and/or 2) of *Oleispira antarctica*.
7. Microorganism according to claim 1, characterized in the DNA sequence encoding the stabilized single ring mutant chaperonin Glu461Ala/Ser463Ala/Val464Ala of Cpn60 (Seq ID No 11) or the mutant chaperonin Lys468Thr/Ser471Gly and/or Cpn 10.
8. Microorganism according to one of the preceding claims, which is selected among animal cell lines, plant cell lines, gram-positive or gram-negative bacteria, fungi and yeasts.
9. Microorganism according to one of the preceding claims, characterized in that the heterologous protein has enzymatic activity or hormonal activity in its native conformation.
10. Microorganism according to one of the preceding claims, characterized in that the DNA sequence encoding a functional chaperone is located chromosomally, extra-chromosomally, or mitochondrially, or in chloroplasts of plants.
11. Process for producing a protein by heterologous expression in a host microorganism containing a gene sequence encoding the heterologous protein, characterized in that a microorganism according to one of the preceding claims is used.
12. Process according to claim 11, characterized in that the host organism is cultivated at a temperature below 25 °C, preferably 4 to 15 °C.

13. Process according to claim 11 or 12, characterized in that the heterologous protein is selected from the group consisting of mammalian proteins, psychrophilic mammalian or bacterial proteins, mesophilic bacterial, fungal or yeast proteins, and mutant or fusion variants thereof.
14. Process for changing the conformation of denatured proteins into their native and/or active conformation, characterized by the step of contacting the denatured protein with a functional chaperone of a psychrophilic bacterium.
15. Process according to claim 14, characterized in that the chaperone is the chaperonin Cpn60 and/or Cpn 10 (Seq ID No 1 and/or 2) of *Oleispira antarctica* in presence of at least one nucleotide, preferably adenosine triphosphate.
16. Process according to claim 11, characterized in that the chaperone is a functional homolog of the chaperonin Cpn60 and/or Cpn 10 (Seq ID No 1 and/or 2) from a psychrophilic bacterium or a functional mutant of the chaperonin Cpn60 and/or Cpn 10 (Seq ID No 1 and/or 2) of *Oleispira antarctica*.
17. Process according to one of claims 11 to 16, characterized in that the contacting is performed extracellularly or *in vitro*.
18. Process according to claim 17, characterized in that the contacting uses at least one immobilized chaperone.
19. Plant, characterized in that it can grow at lower ambient temperatures due to the presence of a DNA sequence encoding a cold-active functional chaperone of a psychrophilic bacterium or plant.
20. Plant according to claim 19, characterized in the DNA sequence encoding a functional chaperonin selected from the group consisting of Cpn60 and/or Cpn 10 (SEQ ID No 1 and/or 2) of *Oleispira antarctica*, a functional homolog thereof, and the stabilized single ring mutant chaperonin Glu461Ala/Ser463Ala/Val464Ala of Cpn60 (Seq ID No 11).

Figure 1:

Amino acid sequences of Cpn60 and Cpn10:

SEQ ID No 1: Cpn10 (encoded by nucleotides pos. 458-751 of Figure 2):

MKIRPLHDRIVVRRKEEETATAGGIILPGAAAEEKPNQGVVISVGTGRILDNGSVQALA
VNEGDDVVVFGKYSGQNTIDIDGEELLILNESDIYGVLEA

SEQ ID No 2: Cpn60 (encoded by nucleotides pos. 800-2446 of Figure 2):

MAAKDVLFGDSARAKMLVGVNILADAVRVTLGPKGKNVIEKSFGAPIITKDGVSVA
AREIELKDKFENMGAQMVKEVASQANDQAGDGTTATVLAQAIISEGLKSVAAGMN
PMIDLKRGIDKATAAVVAAIKEQAQPCLDTKAIAQVGTISANADETVGRLIAEAMEKV
GKEGVITVEEGKGLEDELDVVEGMQFDRGYLSPYFINNQEKMVTEMENPLILLVDKK
IDNLQELLPILENVAKSGRPLLIVAEDVEGQALATLVVNNLRGTFKVAAVKAPGFGD
RRKAMLQDLAILTGGQVISEELGMSLETADPSSLGTASKVVIDKENTVIVDGAGTEAS
VNTRVDQIRAEIESSTSVDYDIEKLQERVAKLAGGVAVIKVGAGSEMEMKEKKDRVD
DALHATRAAVEEGVVAGGGVALIRALSSVTVVGDNEDQNVGIALALRAMEAPIRQI
AGNAGAEGSVVVDKVKSGTGSFGFNASTGEYGDMIAMGILDPAKVTRSSLQAAASI
AGLMITTEAMVADAPVEEGAGGMPDMGGMGGMGGMPGMM

Figure 2:

SEQ ID No 3: DNA coding for Cpn60 and Cpn10:

Cpn10, pos. 458-751

Cpn60, pos. 800-2446

atcaaaaaatgcagcaaggacagattcctgccaagaatttagcagaaggttctgttagcactggccggcgcttattattaacgccgg
gttttgtcactgatgcgcgtgggtttacattactcgtccccgcacgcgtaaagcggtgtccataaggtgattgcatttattaccctc
gcatgatgactgcaagcagcttcaagcgcacggtagtttcaggaaggctcgtaaagatgtacattgcacactgactcgcaaagca
gtcatgaaaaaatcacaattgaaggcgaatataccaaagacgataagttaggtattttcggctagccgttcaaattcctagtaaaagccc

Figure 3:

SEQ ID No 4: Amino acid sequence of esterase cloned from *Oleispira antarctica* (EstRB8):

EstRB8 (encoded by nucleotides 1145 to 2143 Frame 2 of Figure 4) 333 aa

MKNTLKSSSRFSLKQLGTGALISSLFFGGCTTQQDNL YTGVMSLARDSAGLEVKTA
 SAGDVNLTYMERQGSDKDNAESVILLHGFSADKDNWILFTKEFDEKYHIVIAVDLAG
 HGDSEQLLTDYGLIKQAERLDIFLSGLGVNSFHIAGNNSMGGAIASIYSLSHPEKVKS
 TLIDAAGVDGDTESSEYYKVLAEGKNPLIATDEASFEYRMGFTMTQPPFLPWPLRPSLL
 RKTЛАRAEINNKIFSMLKTKERLGMTNFQQKIEVKMAQHPLPTLIMWGKEDRVLD
 VSAAAFFKKIIPQATVHIFPEVGHLPMEIPSESAKVYEEFLSSIK

Figure 4:

SEQ ID No 5: DNA fragment from plasmid pBK1Est coding for esterase of *Oleispira antarctica* (EstRB8):

Nucleotide positions 1-100 correspond to reverse complement of positions 1196-1121 and 3799-3939 correspond to reverse complement of 1043-952 of pBK-CMV vector (Stratagene).

Positions 101-105 are *Bam*H – *Sau*3A1 fusion and positions 3795-3798 are *Sau*3A1-*Bam*H-fusion.

acaggaaacagctatgaccattgattacgccaagctcgaaattaaccctactaaaggaaacaaagctggagctcgccgcctgcag
 gtcgacactagtggatcaacggcgttcatggtaactggctgagttcagcgtcataatgccatgcgatactggccgtcatgactgagttact
 tcttctgctagcaccgattttctaatacgccagctctttatattctgaacggcaactgatgttagttttactaaccggcttttaggcatgg
 taaaactctcgatattcaaattattactgttcatattacaatcatagtgatgttagttttactaaccggcttttaggcatgg
 taagcattattacactcatcgccgttattaaattgtctaaataaaaaatccctgtagcggaaaaattcagcaaatgcacatttttattatttc
 ttggcaataccaaagaattcatcgattttagatgatgacattaaggcaggcaacttggctattaaactacagtcaaaatgcaatttttagacat
 ctcattcaagcaactgacgaacactatggcttagcgtaaagaccttgcactgtcgagcgtagaacaccttcaggtattcacaatagcagtct
 tattttatttaccctcgactaaagactgaattcaataacactacatattgccttaagtcgacatattcaagataaaagatgccttcactgacatca
 gtcaccaacaatcaaatccaaatcgcaaaaactcataaaactagccatccaaatcccaaaagcggtcaaaaatgaa
 acgagcacgtcacaaaaatcaatttatacgtaacgaaaccaggtaacttatcgtttttagcgtttttccactaatgaaagaga
 aaagtcttaattcactggctttggcgatccgcacccatcataagaaatttagtaatggcatgctactggctttaaaaagaatcagttaatt
 gaagaaacctcgcttatctcagccattaccgctgtagccgaatttgcgttatcctcagccatgatccaaactgacgccaattataagac
 atactaattaaactcccttaattgagaagaataatgaaaaacacactcaaattcctcatcagttttactctgaaacaactcggcaccggc
 gctctgattatctccagttttgcgttatcctcagccatgatccaaactgacgccaattataagac
 gctggctagaagttaaaacagcctctgcccgtgacgtcaatcttacttatggaacgccaaggcagtgacaaagataatgccgaaag

cgttatttttattacacggttctctgctgataaagataactggattctttaccaaagaattcgataaaaatcatgttatcgctgtcgatttgcgggacatggcgattcagaacaattattaacgactgattacggctcataaaacaagccgagcgtagatcttcttatctggcttagggtaactcattcacatcgccgtaattcaatgggggggctatcagcgcaatctacagttgagtcacccagagaaagttaaaagtcttacattgatcgatgcagcagggtcgatggcgatactgaaagcgaatactacaaagtttggcagaaggtaagaatccttaattgcaactgatgaagcaagttgaataccgcatgggttaccatgactcagcctccttacctggccactaagaccttattacgtaaaacgctagccgtgccgagatcaataacaaaattttccgatatgctgaaaaccaaagaacgttttaggaatgactaacttcaacacagaaaattgaagtaaaaatggctcaacatccattgccaacactgattatgtgggcaaagaagatcgcttgcgtatccgcagcagcggcctcaaaaaataattccacaagcaactgttcatattttcctgaagtaggccacctatggtagaaattccttagtgaagacgctaaagttaatggcagagtttgccttattaaataagagcacataatcatgactgacttataaacagccaagcattttaaatgctggctttatttaatggccaaattattcaacgaccaagctctgcccggaaaatcgcagtgggttcttgcattcaacagcaacaaacgtgaaatccccgtaatcgattttctgattatcaaaaatacatactttccaccagcatattaactcaactttaaactcgccgcccaccttataacactggcagtcaattcgacaatggtacctgcccggaaacaggatgctttaaaatcgattcgactcactgctgacggttacgatgcttgcgagaaaaacgagtcgtgcgtcaataaaagaaacccatccatccactgcattgcaggccaccgaataacgtatcatgatgattttgtctctggaaataccgctttagaaatagtggtttgatacgcgcttcgctgcgaataatatcttctctgctaagagttgcggatggcatacataactcgcttgcattaaagattaataataatagttAACAGTATATTGAACCTGAGGGTCTGAAGAACTCTAACCTCTGAAGAACCTGAGGCCGCTAGAGAGAAAAGACCATGATAATTTCATCTGCCATGAGAGCCTTATCATGAAAGCCTGTGCTTAAATCAATCATTATTTATTTCATCTTAAATTGAAATAACCAATTTCATATATAATTTCACACTACCCTATCTCACTAGACTTCCC CGCAGCGCAAACAAATCAACGCAAGTTACAATAAGCGTTGC

Figure 5:

Amino acid sequences expressed from vector pBK1CpnEst: - the co-expression of fragments encoding native chaperonines with the esterase gene (EstRB8), all from *Oleispira antarctica*

SEQ ID No 6: cpn10 (nucleotides 113 to 403: Frame 2 of Figure 6) 97 aa:

MKIRPLHDIVVRRKEEETATAGGIILPGAAAEEKPNQGVVISVGTGRILDNGSVQALA
VNEGDVVVFGKYSGQNTIDIDGEELLILNESDIYGVLEA

SEQ ID No 7: cpn60 (nucleotides 455 to 2098: Frame 2 of Figure 6) 548 aa:

MAAKDVLFGDSARAKMLVGVNILADA VRVTLGPKG RNVVIEKSFG APIITKDGVSV
AREIELKDKFENMGAQMVK EVASQANDQAGDGTTATVLAQAIIS EGLKSVAAGMN
PM DLKRGIDKATAAVVAAIKEQAQPCLDTKAIAQVGTISANADETVGRLIAEAMEKV
GKEGVITVEEGKGLEDELDV VEGMQFD RGYLSPYFINNQE KMTVEMENPLILLVDKK
IDNLQELLPILENAVKSGRPLLIV AEDVEGQALATLVVNNLRGTFKVA AVKAPGFGD
RRKAMLQDLAILTGGQVISEELGMSLETADPSSLGTASKV VVIDKENTVIVDGAGTEAS
VNTRVDQIRAEIESSTS DYDIEKLQERVAKLAGGVAVIKVGAGSEM MEMKEKKDRVD
DALHATRAAVEEGVVAGGGVALIRALSSVTVVGDNEDQNVGIALALRAMEAPIRQI
AGNAGAEGSVVVDKVKSGTGSFGFNASTGEY GDMIAMGILDPAK VTRSSLQAAASI
AGLMITTEAMVADAPVEEGAGGMPDMGGMGGMGGMPGMM

SEQ ID No 8: estRB8 (nucleotides 2579 to 3577: Frame 2 of Figure 6) 333 aa:

MKNTLKSSSRFSLKQLGTGALISSLFFGGCTTQQDNLYTGVM SLARD SAGLEV KTA
SAGDVNLTYMERQGSDKD NAESVILLHGFSADKDNWILFTKEFDE KYH VIAV DLAG
HGDSEQLLTTDYGLIKQAERLDIFLSGLGVNSFH IAGNSMGG AISAIYSL SHPEKVKSL
TLIDAAGV DGDTESEYYKV LAEGKNPLI ATDEASFEYRMGFTMTQPPFLPWPLRPSLL
RKTLARA EINNK IFSDMLKT KERLGMTNFQQKIEVKMAQHPLPTLIMWGKEDRVLD
VSAAA AFKKIIPQATV HIFPEV GHLP MVEIPS ESAK VYEEFLSSI K

Figure 6:

SEQ ID No 9: pBK1CpnEst: - the fusion of native chaperonine-coding fragments with esterase of *Oleispira antarctica* (EstRB8)

The DNA fragment coding for Cpn10 and Cpn60 is flanked by *SacI* site (pos. 69-75) and *SalI* site (encoded by pos. 2138-2143 of Figure 7):

Nucleotide positions 1-75 correspond to reverse complement of positions 1196-1121 and positions 5233-5273 correspond to reverse complement of 1043-952 of pBK-CMV vector (Stratagene)

Small letters – the Cpn10-Cpn60 encoding fragment,

Capital italics – fragments of vector pBK-CMV

Capital letters – fragment coding for EstRB8 from plasmid pBK1Est

gttgggtgcgggtctgaaatggaaatgaaagagaagaaaagaccgtgtgacgatgcacttcattgcaactcgccagcggtaagaag
gtgttgtgcgggtgggtgtgcttgattcgccactcttcattgtaaccgtgtggtgataacgaagatcaaaacgtcggtattgcat
tggcacttcgtgcgtggaaagctctatccgtcaaattcgccggtaacgcagggtctaagggtcagtgggtgtgataaagtgaaatctg
gcacaggtagcttggtttaacgccacaggtagtggcgatattgcgtatgggtatttagaccctgcaaaagtacgcgttc
atcttacaagccgcggcgttatcgccagggttatgtatcacaaccgaagccatggtgccgtatgcgtatgggtatgcgtatccgt
gtatgcctgatatggcgcatgggtggatggcggtatgcctggcatgtatcactttgtgattcattgtcctgatctgcgttaccgt
GTCGACATATTCAAGATAAAGATGCCTCACTGACATCAGTCACCAACAATCAAT
CAAACACCAATACCAATCGCAAAAACACTCATAAAACCTAGCCGATCACCAAAATCCC
AAAAGCGTTCAAAAATGAAACGAGCACGTACACACAAAATCAATTATACGCTAA
CGAACCCAGGTCAAACCTATCGTTTTGAGCACGTTGTTCCACTAATGAAAGA
GAAAAGTCGTTAACCTACTGGCTTTGGCGTATCCGCACCTTCACATAGAAATT
GTAATGGCATGCTACTGGCCTTAAAAAGAATCAGTTAACATGAAGAACCTCGCT
TATCTCAGCCATTACCGCTGTAGCCGAATTGCGCTTATCCTCAGCCATGATTAAA
CTGACGCCAATTAATATAAGACATACTAATTAATAACTCCCTTAATTGAGAAGAA
TAATGAAAAACACACTCAAATCCTCATCACGTTTAGTCTGAAACAACTCGGCAC
CGCGCTCTGATTATCTCCAGTTGTTCTCGGTGGTGCACCACAAACACAAG
ATAATTATACACAGGGTTATGCTCTTGCGAGAGACAGCGCTGGCCTAGAAGT
TAAAACAGCCTCTGCCGGTACGTCAATCTTACTTATATGGAACGCCAAGGCAGT
GACAAAGATAATGCCGAAAGCGTTATTTTACACGGTTCTGCTGATAAAAG
ATAACTGGATTCTTTACCAAAGAATTGATGAAAAATATCATGTTATCGCTGTC
GATTAGCGGGACATGGCGATTAGAACAAATTATTAACGACTGATTACGGCTCA
TAAAACAAGCCAGCGTTAGATATCTCTTATCTGGCTTAGGGTTAACTCATTT
CACATGCCGGTAATTCAATGGGGGGGCTATCAGCGCAATCTACAGTTGAGTC
ACCCAGAGAAAGTTAAAAGTCTTACATTGATCGATGCAGCAGGTGTCGATGGCG
ATACTGAAAGCGAATACTACAAAGTTGGCAGAAGGTAAGAACCTTTAATTGC
AACTGATGAAGCAAGTTGAATAACCGCATGGTTCAACCAGTCACTCAGCCTCCT
TTCCTACCTTGGCCACTAAGACCTTCTTATTACGTAACCGCTAGCCGTGCCGA
GATCAATAACAAAATTTCGATATGCTGAAACCAAGAACGTTAGGAATG
ACTAACTTCAACAGAAAATTGAAGTGAAGGCTAACATCCATTGCCAACAC
TGATTATGTGGGGCAAAGAAGATCGCGTTCTGACGTATCCGCAGCAGCGGCCTT
CAAAAAAAATAATTCCACAAAGCAACTGTCATATTTCCTGAAGTAGGCCACCTA
CCTATGGTAGAAATTCTAGTGAAAGCGCTAAAGTTATGAAGAGTTTGTCCCT
CTATTAAATAAGAGCACATAATCATGACTGACTTATAAACAGCCAAGCATTAAA
ATGCTTGGCTGTTATTAAATGCCAAATTATTCAACGACCAAGCTCTGCGGTAA

AATCGCAGTGGGTTCTTGTTCATCAACAGCAACAAACGTGAAATACCCCGTA
ATCGCATTCTGATTATCAAAATACATACTTCCACCAGCATATTAACCTCAAC
TTTAAACTCGCCGCCCTACCTCTATAAACACTGGCAGTCAATTGACAATGGTAC
CTGCGGGAACAGGATGCTAAAATCGATTGACTGCTGACGGTTACGATGCT
TTGTCGAGAAAAACGAGTCGCTGCAATAAAAGAAACCTCATCCATCCACTGCATT
GCAGTGCCACCGAATAACGTATCATGATGATTGTTGTCTGGAAATACCGCTT
AGAAAATAGTGGTTTGATACGCGCTTCGCTGCGCAATAATATCTTCTGCTAA
GAGTTGCGGATGGCATACATAACTCGCTGATTAAGATTAATAATAAGTTA
ACAGTATATTGAAC TGAGGGTCTGAAGAACTCTAACCTCTGAAGAACTTGAG
GCCGCTAGAGAGAAAAGACCA GTGATAATATTGATCTTGCCATGAGAGCTTATC
ATGAAAGCCTGTGCTAAAATCAATCATTATATTGATCTTTAATTGAAATAA
TACCAATATATTGATATATAATTCAACACTACCCCTATCTCACTAGACTCCC
GCATAGGCGCAAACAATCAACGCAAGTTACAATAAGCGGTCGCTGCAACAC
ATGCCCTAGCGTCTAAAGTAGCACGCACAACACTGGCCAGTCGTACTAGCCC
TGCGATTGTCAGACGAGCAACAAGCGCTATTAAACTTACCTAAATTCTAAC
ACCACCATGGTCTTCCACAAACTCAAAAAACTCGTCAAATCCGCTGCAATT
TAAACGCGATGACATAGATCTAACGATTATCAAACCCGCATTCAAGCGCTCATT
AAAAACGCACCACTGGCAAGAAGTTCTACCTGCACTGACCAATATGCAAGCGGC
GGCGGAAGAGCTGCCTTGATCGATCAAGAAGAAGGGAGCAGCAAAGAGGAAA
ACAATCAAAAAGAGGAGAGCAATCAAATAAAACGAGTTATTGAGGATTAAAT
TTTAAAACAGGTATTTAATACCTCTCGTAGTAAACAATGACTGTATTACAC
AAAAATAAAATAGAGGTATACCATGTCAAACATCTGGTTGAAGTACCAAAGATTG
AAGTATTAAACCGTCAAATGGAAAATACTGCCTGCAGCAACTAGGCATTCAAAT
TACAGAAATTGGCGATGATTATATCACTGGCACAAATGCCAGCAGATGCACGTACC
TTCCAGCCAATGGACTGATTGATGGCGCTCAAATGTATTGCTGGCAGAAACAC
TGGCAGCATGGCAGCTAACTGCTGTATTAAATTGCTCAAGAATATTGTGTTGG
CCAAGAAATTACGCCAACACATACGCGGTGTTCGTCCGGCATAGTACTGGC
ACAGCAACGCTAGTACACAAAGGAAGAACCTCCCAGATTGGAAATTGCGATC
GTTAACGATCCAAAGAATTCAAAGCTTCTCGAGAGTACTTCTAGAGCGGCCGCGGG
CCCATCGATTCCACCCGGTGGGTACCAGGTAAGTGTACCCATTGCCCTATAGT
GAGTCGTATTACAATTCACTGGCCGTCGTTTAC

Figure 7:

Amino acid sequences expressed from vector pBK1CpnSREst: - the co-expression of the stabilized single ring mutant chaperonin with the esterase gene (EstRB8) from *Oleispira antarctica* (cpn10::stabilized single ring mutant Glu461Ala/Ser463Ala/Val464Ala::est)

SEQ ID No 10: cpn10 (nucleotides 113 to 403: Frame 2 of Figure 8) 97 aa:

MKIRPLHDRIVRRKEEETATAGGIILPGAAAEEKPNQGVVISVGTGRILDNGSVQALA
VNEGDVVVFGKYSGQNTIDIDGEELLILNESDIYGVLEA

Below – ***Capital bold letters*** are the mutations introduced

SEQ ID No 11: stabilized single ring mutant of cpn60 (nucleotides 455 to 2098: Frame 2 of Figure 8) 548 aa:

MAAKDVLFGDSARAKMLGVNILADAVRVTLGPKGKGRNVVIEKSFGAPIITKDGVS
AREIELKDKFENMGAQMVKVASQANDQAGDGTTATVLAQAIISSEGKSVAAAGMN
PMIDLKRGIDKATAAVVAIAIKEQAQPCLDTKAIAQVGTISANADETVGRLIAEAMEKV
GKEGVITVEEGKGLEDELDVVEGMQFDRGYLSPYFINNQEKMVTMEMENPLILLVDKK
IDNLQELLPILENAKSGRPLLIVAEDEVGQALATLVVNNLRGTFKVAAVKAPGFGD
RRKAMLQDLAILTGGQVISEELGMSLETADPSSLGTASKVVIDKENTVIVDGAGTEAS
VNTRVDQIRAEIESSTDYDIEKLQERVAKLAGGVAVIKVGAGSEMEMKEKKDRVD
DALHATRAAVEEGVVAGGGVALIRALSSVTVVGDNEDQNVGIALALRAMEAPIRQI
AGNAGAAGAAVVDKVKSGTGSFGFNASTGEYGDMIAMGILDPAKVTRSSLQAAASI
AGLMITTEAMVADAPVEEGAGGMPDMGGMGGMGGMPGMM

SEQ ID No 12: EstRB8 (nucleotides 2579 to 3577: Frame 2 of Figure 8) 333 aa:

MKNTLKSSSRFLSLKQLGTGALISSLFFGGCTTQQDNLYTGVMMSLARDSAGLEVKT
SAGDVNLTYMERQGSDKDNAESVILLHGFSADKDNWILFTKEFDEKYHVIAVDLAG
HGDSEQLLTTDYGLIKQAERLDIFLSGLGVNSFHAGNSMGGAIYSLSHPEVKSL
TLIDAAGVGDGDESEYYKVLAEKNPLIATDEASFYRMGFTMTQPPFLPWPLRPSLL

RKTLARAЕINNKIFSMLKTKERLGMTNFQQKIEVKMAQHPLPTLIMWGKEDRVLD
VSAAAAFKKIIPQATVHFPEVGHLPMVEIPSESAKVYEEFLSSIK

Figure 8:

SEQ ID No 13: DNA sequence of vector pBK1CpnSREst: the expression cassette for the co-expression of the stabilized single ring mutant chaperonin with the esterase gene (EstRB8) from *Oleispira antarctica* (cpn10::stabilized single ring mutant Glu461Ala/Ser463Ala/Val464Ala::est)

Nucleotide positions 1-75 correspond to reverse complement of positions 1196-1121 and positions 5233-5273 correspond to reverse complement of 1043-952 of pBK-CMV vector (Stratagene)

DNA fragment coding for Cpn10 and Cpn60 is flanked by *SacI* site (pos. 69-75) and *Sall* site (pos. 2138-2143).

In the DNA sequence:

Small letters – the Cpn10-Cpn60 coding fragment,

Capital italics – fragments of vector

Capital letters – fragment coding for EstRB8 from plasmid pBK1Est

Capital bold letters = introduced mutations

gatacaaaagcaatcgctcaggtagggacaatctctgccaatgccatgaaacgggtgcgttaattgctgaagcgatggaaaaagt
cggttaagaagggtgattaccgtgaagaaggcaaaggccgtgaagacgagcttgcgttgcgttgcgttgcgttgcgttgcgttgcgtt
acttgtctccgtacttcatcaacaaccaagaaaaaatgaccgtagaaatggaaaatccattaattctattgggtgataagaaaattgataaac
cttcaagagctgtgccaattctgaaaacgtcgctaaatcaggtcgtccattattgatcggtgcgtgaagatgtgaaggccaagcactagc
aacattggtagtaaacaactgcgcggcacattcaagggtgcagcggtaaagcccctggcgatcgctgtaaagcgatgtgca
agatcttgcacatcttgcgggttcaggttattctgaagagctaggatgtcttagaaactgcggatcctcttgggtacggcaa
gcaagggttatcgataaagaaaacaccgtgattgtatggcgaggactgaagcaagcgttaatactcggttgcaccagatccgt
ctgaaatcgaaagctgacttctgattacgacatcgaaaagttacaagaacgcgttgcgttgcggcggttgcgttgcgttgcgttgcgttgcgt
gttgggtcgggttctgaaatggaaatgaaagagaagaaagaccgtgttgcgttgcacttcatgcaactcgccagcgggtgaagaag
gtgttgcgggtgggtgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgt
tggcacttcgtgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgt
ctggcacaggttagcttggtttaacgccagcacaggtgagtatggcgatgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgt
gttcatctacaagccgcggcgttatcgcagggttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgt
ggtggatgcgttatggcgcatgggtggatggcggtatgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgttgcgt
ccgtGTCGACATATTCAAGATAAAGATGCCTTCACTGACATCAGTCACCAACAATC
AATCAAACACCAATACCAATCGAAAAACTCATAAAATAGCCGATCACCAAAT
CCCCAAAGCGTTCAAAATGAAACGAGCACGTACACACAAATCAATTATACGC
TAACGAACCAGGTCAAAACTATCGTTTTGAGCACGTTGTTCCACTAATGAAA
GAGAAAAGTCGTTAATTCACTGGCTTGGCGTATCCGCACCTCACATAGAAAAT
TAGTAATGGCATGCTACTGGCCTTAAAAAGAATCAGTTAATTGAAGAACCTCG
CTTATCTCAGCCATTACCGCTGTAGCCGAATTGCGCTTATCCTCAGCCATGATTA
AACTGACGCCAATTAAATATAAGACATAACTAATTAAACTCCCTTAATTGAGAAG
AATAATGAAAAACACACTCAAATCCTCATCACGTTTAGTCTGAAACAAACTCGGC
ACCGGCGCTTGATTATCTCCAGTTGTTCTCGGTGGTGCACCAACACAAACA
AGATAATTATACACAGGGTTATGTCTCTGCGAGAGACAGCGCTGGCCTAGAA
GTTAAAACAGCCTCTGCCGGTGACGTCAATCTTACTTATATGGAACGCCAAGGCA
GTGACAAAGATAATGCCGAAAGCGTTATTATTACACGGTTCTGCTGCTGATAA
AGATAACTGGATTCTTTACCAAAGAATCGATGAAAAATATCATGTTATCGCT
GTCGATTAGCGGGACATGGCGATTAGAACAATTATTAACGACTGATTACGGTC
TCATAAAACAAGCCGAGCGTTAGATATCTTCTGCTTAGGGCTTAGGGTTAACTC
ATTCACATCGCCGGTAATTCAATGGGGGGGCTATCGCAATCTACAGTTG
AGTCACCCAGAGAAAGTTAAAAGTCTTACATTGATCGATGCAGCAGGTGTCGATG
GCGATACTGAAAGCGAATACTACAAAGTTGGCAGAAGGTAAGAATCCTTAAAT
TGCAACTGATGAAGCAAGTTGAAATACCGCATGGGTTACCATGACTCAGCCT

CCTTCCTACCTGGCCACTAAGACCTTCTTATTACGTAAAACGCTAGCCCGTGC
CGAGATCAATAACAAAATTTTCCGATATGCTGAAAACCAAAGAACGTTAGGA
ATGACTAACTTCAACAGAAAATTGAAGTGAAAATGGCTAACATCCATTGCCAA
CACTGATTATGTGGGGCAAAGAAGATCGCGTTCTGACGTATCCGCAGCAGCGGC
CTTCAAAAAAATAATTCCACAAGCAACTGTTCATATTTCCTGAAGTAGGCCAC
CTACCTATGGTAGAAATTCTAGTGAAAGCGCTAAAGTTATGAAGAGTTTGT
CCTCTATTAAATAAGAGCACATAATCATGACTGACTTATAAACAGCCAAGCATT
AAAATGCTGGCTGTTATTAAATGCCAAATTATTCAACGACCAAGCTCTGCG
GTAAAATCGCAGTGGTTCTGTTCATCAACAGCAACAAACGTGAAATACCC
CGTAATCGCATTTCCTGATTATCAAAATACATACTTCCACCAGCATATTAACTT
CAACTTTAAACTCGTCCGCCCTACCTCTATAACACTGGCAGTCATTGACAATG
GTACCTGCAGGAACAGGATGCTTAAATCGATTGACTGCTGACGGTTACGA
TGCTTGTCGAGAAAAACGAGTCGCTGCAATAAAAGAAACCTCATCCATCCACTG
CATTGCAGTGCCACCGAATAACGTATCATGATGATTGTTGTCTCTGGAAATACC
GCTTAGAAATAGGTTTGATACGCGCTTCGCTGCGCAATAATATCTTCTCT
GCTAAGAGTTGCGGATGGCATACATAAAACTCGCTTGATTAAGATTAATAATAAAT
AGTTAACAGTATATTGAAC TGAGGGTCTGAAGAACTCTAATACCTCTGAAGAACT
TTGAGGCCGCTAGAGAGAAAAGACCAGTGATAATATTCATCTGCCATGAGAGC
TTATCATGAAAGCCTGTGCTTAAATCAATCATTATATTATTATCTTAAATTGA
AATAATACCAATATATTCATATATAATTACACTACCCTATCTCACTAGACTT
CCCGCGCATAGGCGCAAACAATCAACGCAAGTTCACAATAAGCGGTTCGCTGC
AACACATGCCCTAGCGTCTAAAGTAGCACGCACAACACTGGCCAGTCGTACTAGC
CCCTTGCATTGCGACGAGCAACAGCGCTATTAAACTTACCTAAATTTC
TAACCACCAATTGGTTCTTCCACAAACTCAAAAAACTCGTCAAATCCGCTTG
CAATTAAACGCGATGACATAGATCTAATCGATTATCAAACCCGATTCAAGCGC
TCATTAAAAACGCACCACTGGCAAGAAGTTCTACCTGCACTGACCAATATGCAAG
CGGGCGGAGAGCTGCCATTGATCGATCAAGAAGAAGGGAGCAGCAAAGAGG
AAAACAATCAAAAAGAGGAGAGCAATCAAATAAAACGAGTTATTGAGGATT
AATTAAACAGGTATATTAAATACCTCTCGTAGTAAACAATGACTGTATT
CACAAAAATAAATAGAGGTATACCATGTCAAACATCTGGTTGAAGTACCAAAG
ATTGAAGTATTAAACCGTCAAATGGAAAATCTGCAGCAACTAGGCATT
AAATTACAGAAATTGGCGATGATTATACACTGGCACAATGCCAGCAGATGCACG
TACCTTCCAGCCAATGGACTGATTCAATGGGGCTCAAATGTATTGCTGGCAGAA
ACACTGGGCAGCATGGCAGCTAACTGCTGTATTAAATTGTCTCAAGAATATTGTG

TTGGCCAAGAAATTAACGCCAACCATACGCGGTTCGTTCCGGCATAGTGAC
 TGGCACAGCAACGCTAGTACACAAAGGAAGAACCTCCCAGATTGGGAAATTG
 CATCGTTAACGATCAAAGAATTCAAAAAGCTCTCGAGAGTACTCTAGAGCGGCCG
CGGGCCATCGATTTCCACCCGGGTGGGGTACCAGGTAAAGTGTACCCAATTGCCCT
ATAGTGAGTCGTATTACAATTCACTGGCCGTCGTTTAC

Figure 9:

Amino acid sequence of the stabilized single ring mutant Glu461Ala/Ser463Ala/Val464Ala of Cpn60:

SEQ ID No 14: Cpn10 (nucleotides 458-751 of Figure 10):

MKIRPLHDRIVVRRKEEETATAGGIILPGAAAEEKPNQGVVISVGTGRILDNGSVQALA
 VNEGDRVVFVFGKYSGQNTIDIDGEELLILNESDIYGVLEA

SEQ ID No 15: Cpn60 (nucleotides 458-751 of Figure 10):

MAAKDVLFGDSARAKMLVGVNILADAVRVTLGPGRNVVIEKSFGAPIITKDGVSV
 AREIELDKFENMGAQMVKEVASQANDQAGDGTTATVLAQAIISEGLKSVAAGMN
 PMDLKRKGIDKATAAVVAAIKEQAQPCLDTKAIAQVGTISANADETVGRLIAEAMEKV
 GKEGVITVEEGKGLEDELDVVEGMQFDRGYLSFYFINNQEKMVTLEMENPLILLVDKK
 IDNLQELLPILENVAKSGRPLLIVAEDVEGQALATLVVNNLRGTFKVAAVKAPGFD
 RRKAMLQDLAILTGGQVISEELGMSLETADPSSLGTASKVVIDKENTVIVDGAGTEAS
 VNTRVDQIRAEIESSTDYDIEKLQERVAKLAGGVAVIKVGAGSEMEMKEKKDRVD
 DALHATRAAVEEGVVAGGGVALIRALSSVTVVGDNEDQNVGIALALRAMEAPIRQI
 AGNAGAAGAAVVDKVKSGTGSFGFNASTGEYGDMIAMGILDPAKVTRSSLQAAASI
 AGLMITTEAMVADAPVEEGAGGMPDMGGMGGMGGMPGMM

Figure 10:

SEQ ID No 16: DNA sequence of the stabilized single ring mutant

Glu461Ala/Ser463Ala/Val464Ala:

In the DNA sequence:

Small letters – the Cpn10-Cpn60 coding fragment,

Big bold letters = introduced mutations

ggtaacgcagggtctg**C**aggg**G**cag**C**ggtttgtataaagtgaaatctggcacaggttagcttggtttaacgccagcacaggt
agtatggcgatatgattgcgatgggtatttttagaccctgcaaaagtacgcgttcatctctacaagccgcggcgtctatcgcagggttgat
gatcacaaccgaagccatgggtgcggatgcgcctgttgaagaaggcgctgggtatgcctgatatggcggcatgggtggaatggg
cggtatgcctggcatgatgtaatcacttgtgattcattgcctgatctgcttaccgtgtaaaaagatcaggctaaggctgtctctataaaa
agccgtatctttagtggatgtttgtctgtgaaaacgacattttggagtgccggctttttgattttggtcataaaattcagaatattgtgt
attttatgttaactagctggcctataatgtttagttcctctgggtggcatgatctcatggtacttcacttaagcctgattcactgcg
gctttaacagtaaaataaacgcaacgtagaacataataagcgtatggcattaatgaagacggctgcatttaattcagatc