SYNTHETIC ANTIFREEZE PEPTIDE AND SYNTHETIC GENE CODING FOR ITS PRODUCTION

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Continuation of application No. 07/588,437, Sep. 25, 1990, abandoned.

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ABSTRACT
A synthetic antifreeze peptide and a synthetic gene coding for the antifreeze peptide have been produced. The antifreeze peptide has a greater number of repeating amino acid sequences than is present in the native antifreeze peptides from winter flounder upon which the synthetic antifreeze peptide was modeled. Each repeating amino acid sequence has two polar amino acid residues which are spaced a controlled distance apart so that the antifreeze peptide may inhibit ice formation. The synthetic gene has been expressed in E. coli. A synthetic insert fragment has been prepared which can be readily inserted into the synthetic gene to alter the number of repeating units and/or amino acid composition in the antifreeze peptide produced.

23 Claims, 10 Drawing Sheets
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(SEQ ID NO:1)

Asp—Thr—Ala—Ser—Asp—Ala—Ala—Ala—Ala—Ala—Leu—Thr—Ala—Ala—Asp—
Ala—Ala—Ala—Ala—Ala—Leu—Thr—Ala—Ala—Asp—
Ala—Ala—Ala—Ala—Ala—Ala—Ala—Thr—Ala—Ala—Xaa

FIG. 1

ANTIFREEZE PEPTIDE

HYDROGEN BOND

4.5 Å

a = AXIS

FIG. 2
(SEQ ID No:9)

5' - GCT GCG GCC GCG GCA GCC ACC GCA GCA ACT GCA - 3'
3' - A CGT CGA CGC CGG CGC CGT CGG TGG CGT CCA TG - 5'

(SEQ ID No:10)

- rAlaAlaAlaAlaAlaAlaAlaThrAlaAlaTh

(SEQ ID No:11)

FIG.5

(SEQ ID No:12)

5' - GCT GCG GCC GCG GCA GCC ACC GCA GCA ACT GCA GCT GCG GCC GCG GCA GCC ACC GCA GCA ACT GCA-3'
3' - A CGT CGA CGC CGG CGC CGT CGG TGG CGT CCA TGA CGT CGA CGC CGG CGC CGT CGG TGG CGT CCA TG-5'

FIG.6

(SEQ ID No:14)

5' - GCT GCG GCC GCG GCA GCC TGC GCA GCA ACT GCA -3'
3' - A CGT CGA CGC CGG CGC CGT CGG ACG CGT CCA TG - 5'

(SEQ ID No:15)

FIG.7
A. (SEQ ID No:16)

5' - CAGTGAATTCTAGATCTGTAAGGAGGTTATCGATGTCACCAGCTCTGGCCACGCCGCCTGGATCGGAGG

CiaI  

5' - GTGTGAATTCTAGATCTGTAAGGAGGTTATCGATGTCACCAGCTCTGGCCACGCCGCCTGGATCGGAGG

EcoRI  

3' - CACACTTAAAGCACATCTCCTCAAAATAGCTACAGGGTGGCCAGGCTGCGGCCCATGGCTCC

3' - CACACTTAAAGCACATCTCCTCAAAATAGCTACAGGGTGGCCAGGCTGCGGCCCATGGCTCC

FIG. 8A

B1 (SEQ ID No:19)

5' - GCCAGCAGCAGCTCCTACTCCTGCCACCG

PvuII

5' - GCCAGCAGCAGCTCCTACTCCTGCCACCG

3' - CGCCGGTGTCGTCGACGATGACGACGG

3' - CGCCGGTGTCGTCGACGATGACGACGG

(SEQ ID No:20)

FIG. 8B

B2. (SEQ ID No:22)

5' - CACCGCGCTGTCCGCGACCGCGACCGCGCTGCAGCTGCTGACCTGCGGAC

XmaII

5' - CACCGCGCTGTCCGCGACCGCGACCGCGCTGCAGCTGCTGACCTGCGGAC

(SEQ ID No:24)

3' - GCCGCTGTCGTCGACGATGACGACGG

3' - GCCGCTGTCGTCGACGATGACGACGG

FIG. 8C

B3. (SEQ ID No:25)

5' - GCCCGCACCGCGCTGCAGCTGCTGACCTGCGGAC

BacII

5' - GCCCGCACCGCGCTGCAGCTGCTGACCTGCGGAC

(SEQ ID No:27)

3' - GCCGCTGTCGTCGACGATGACGACGG

3' - GCCGCTGTCGTCGACGATGACGACGG

FIG. 8D

B4. (SEQ ID No:28)

5' - CCACCAGCAGCAGCGCTGCAGCTACCAGCAGCTGCTGCGGAC

PstI

5' - CCACCAGCAGCAGCGCTGCAGCTACCAGCAGCTGCTGCGGAC

(SEQ ID No:29)

3' - GTGTCGTCTGTCGACCGGAGCCGGCTGATGGCCGGCCGCTGGACC

3' - GTGTCGTCTGTCGACCGGAGCCGGCTGATGGCCGGCCGCTGGACC

FIG. 8E
C. (SEQ ID No:31)

\[
\begin{align*}
5' - & \text{GCTAAAGCTGCGGCTCTGCAACGCGACTGCTGCGGCGGACTGCTGCGGCGGACTGCTGCGGCAGCGGCTCGT} \\
& \quad \text{AlaLysAlaAlaAlaLeuThrAlaAlaAsnAlaAlaAlaAlaAlaThrAlaAlaAlaAlaAlaAlaAla} \\
& \quad \text{Arg} \\
3' - & \text{ACGTCGATTCCGACGAGACTGACGACGTTCGGCGGCAGCGCGCCGCTGAGCGCAGCGGCGTCGCCGAGCA} \\
& \quad \text{PstI} \\
\end{align*}
\]

(SEQ ID No:33)

\[
\begin{align*}
\text{BamHI} \\
& \quad \text{GGTTGATAAG-3'} \\
& \quad \text{Gly****} \\
& \quad \text{CCAACATTCCTAG-5'} \\
\end{align*}
\]

FIG. 8F

CΔI (SEQ ID No:34)

\[
\begin{align*}
5' - & \text{GCTAAACCTGCGGCTCTGCAACGCAGCGACTGCTGCGGCGGACTGCTGCGGCGGACTGCTGCGGCGGCTCGT} \\
& \quad \text{AlaLysAlaAlaAlaLeuThrAlaAlaAsnAlaAlaAlaAlaAlaThrAlaAlaAlaAlaAlaAlaAla} \\
& \quad \text{Arg} \\
3' - & \text{ACGTCGATTCCGACGAGACTGACGACGTTCGGCGGCAGCGCGCCGCTGAGCGCAGCGGCGTCGCCGAGCA} \\
& \quad \text{PstI} \\
\end{align*}
\]

\[
\begin{align*}
\text{BamHI} \\
& \quad \text{GGTTGATAAG-3'} \\
& \quad \text{GlyCysLysAspP} \\
& \quad \text{CCAACATTCCTAG-5'} \\
\end{align*}
\]

(SEQ ID No:35) BamHI

FIG. 8G
(SEQ ID NO:37) \text{EcoRI} \quad \text{SstII} \\
5'-\text{AATTCAGATCTGCAGAATCGCCCTCCGACGCCG}-3' \\
(SEQ ID NO:39) \hspace{1cm} \text{Asn Ser Asp Leu Arg Asp Thr Ala Ser Asp Ala Ala} \\
3'-\text{GTCTAGACGCTCTATGGCGAAGGCTGCCG} - 5'(SEQ ID NO:38)

FIG. 10
SYNTHETIC ANTIFREEZE PEPTIDE AND SYNTHETIC GENE CODING FOR ITS PRODUCTION

This application is a continuation of Ser. No. 07/588,437, filed Sep. 25, 1990, now abandoned.

BACKGROUND OF THE INVENTION

1. Field of the Invention

The present invention generally relates to compositions used to prevent ice formation and, more particularly, to a synthetic polypeptide which lowers the freezing point of water and a synthetic gene for producing the polypeptide in bacteria.

2. Description of the Prior Art

Ice formation is an ever present problem in today's world. Ice on roadways and bridges is believed to be the cause of numerous automobile accidents. Ice on aircraft wings is known to decrease lift and increase the weight of the aircraft and is believed to be responsible for causing at least one jumbo jet to crash land. Ice continually forms on the decks of boats which sail in arctic and antarctic waters, thereby making operations aboard the boats extremely hazardous. Ice caused during severe weather can also result in millions of dollars worth of damage to crops such as grapes and oranges.

Ice prevention in the roadway, bridge, and boat environments typically involves applying large amounts of toxic chemicals such as glycols and salts to ice formed on the surfaces of the roadway, bridge or boat. Application of glycols and salts is reactive in nature since the conditions for ice formation must all ready be present when the glycols and salts are applied. The chemicals used for ice prevention can be harmful to the environment and tend to cause rust formation on automobiles which travel over the salt covered roadways. Ice prevention in aircraft usually involves onboard inflatable members and heating elements which break and melt the ice after it forms. Such ice protection schemes require energy and are only suitable for specific applications. Presently, there is no generally accepted ice prevention scheme used in agriculture.

Efforts have been made to find less toxic alternatives for ice prevention which are suitable for ice prevention/suppression in a wide variety of applications. U.S. Pat. Nos. 4,045,910 and 4,161,084 to Amy et al. disclose protecting plants from frost damage by applying non-ice nucleating bacteria to the plants before the onset of freezing cold. The non-ice nucleating bacteria are supposed to compete with native ice nucleating bacteria and prevent ice formation by reducing the number of potential "triggers" to crystallization. One drawback of the Amy et al. ice prevention method is that it involves the release of genetically modified bacteria into the environment. U.S. Pat. No. 4,484,409 to Caple et al. discloses chemically synthesizing polymeric ice nucleation inhibitors via free radical polymerization. The polymers produced in Caple et al. have a tightly controlled spacing of about 15 Angstroms (Å) between the hydrophilic and hydrophobic groups. The polymers are sprayed on the plants and are designed to inhibit ice formation. U.S. Pat. No. 4,834,899 to Kleveland discloses applying a bactericide to plants to prevent frost damage by killing the ice nucleating bacteria. U.S. Pat. No. 4,601,842 to Caple et al. discloses applying a proteinaceous material obtained from cold weather plants to growing crops for protection from frost damage.

Recently, naturally produced antifreeze materials that are present in cold water fish have been investigated for their utility as ice suppressors (see, for example, Picket et al., Eur J. Biochem. 143:35–38 (1984)). These materials are either peptides or glycopeptides and have been designated as AFPs for "antifreeze peptides" and AFGPs for "antifreeze glycopeptides", respectively. The AFPs and AFGPs are produced by the fish to prevent the formation of ice in their body fluids so that they may survive in water temperatures below freezing. There are several subclassifications of AFPs and AFGPs, with the simplest being the alanine rich peptides found in several species of flounder.

FIG. 1 shows the primary amino acid sequence of a native AFP isolated from the winter flounder Pseudopleuronectes americanus (see, DeVries, Phil. Trans. of the Roy. Soc. of London (series B) 304:575–588 (1984)). This is a class 1 or alanine rich AFP. There are 38 amino acids (SEQ ID NO:1) in the AFP with the last 33 amino acids comprising essentially 3 repeats of 11 amino acids (i.e., there are slight differences in the repeating groups where, for example, leucine (leu) is substituted for alanine (ala) and another polar amino acid (xxa) can be substituted for aspartic acid (asp)). The non-polar amino acids, i.e., ala and leu, are used primarily as "spacers" which position the polar amino acids, i.e., thr and asp, approximately 4.5 angstroms (Å) apart. The AFP of SEQ ID NO:1 is but one example of a wide variety of AFPs. Most naturally occurring class 1 AFPs are approximately 70% ala and include three to five repeats of a sequence which includes two amino acids capable of hydrogen binding, i.e., thr and asp, spaced approximately 4.5 Å apart. Chakrabartty et al., in Journal of Biological Chemistry 264:11307–11312 (1989) and 264:11313–11316 (1989), disclosed the direct chemical synthesis of AFPs. It was determined in Chakrabartty et al. that the minimum size for activity of an AFP is 3 repeats of the sequence. There are no known naturally produced AFPs that include greater than 5 repeats of the sequence.

FIG. 2, which is taken from the above cited DeVries article, shows an AFP interacting with an ice crystal during ice formation. As shown in FIG. 2, the secondary structure of a class 1 AFP is an alpha helix and the tertiary structure is a straight rod. The polar amino acids, thr and asp, are positioned on one side of the rod and form hydrogen bonds with the water molecules along the α-axis of the ice crystal. The 4.5 Å spacing of the polar amino acids in the AFP is precisely the spacing of water molecules in a forming ice crystal.

FIGS. 3a and 3b, which are also taken from the above-cited DeVries article, shows that ice crystals form in two directions. The crystals grow outward along the α-axis and at right angles to this plane along the c-axis. By binding to the forming prism faces in the c-axis, the AFPs disrupt the formation of the step, thereby preventing smooth formation of the face, and resulting in curved fronts. Site 10 shows an area where two AFP molecules have bound closely together which has resulted in an even more effective blockage of ice crystal growth. The same extra blockage effect would occur with a longer molecule that can block more of the step.

The mechanism of ice crystal growth suppression by AFPs illustrated in FIGS. 2 and 3b falls under the "adsorption inhibition hypothesis" proposed by DeVries. As illustrated, the ice crystal can only grow in unblocked regions because the presence of the AFP ties up the potential insertion sites for new water to come into the lattice. In theory, since all water adds to the lattice by hydrogen bonding too, if all possible active sites are bound up, the AFPs must by moved out of the way before new water can be added. Attachment of the AFP to the ice crystal increases the ratio of molecular volume to surface area, and in order
for freezing to occur, more energy must be removed from
the system than would be required in the absence of the AFPs.
Hence, the freezing point of water is lowered by the binding
action of AFPs. The mechanism of action of AFPs is
somewhat analogous to “poisoning” the growth of a crystal
by the presence of an impurity where the AFP acts as the
impurity. Melting point, however, is not lowered, and this
phenomenon, which has been known for many years, is
referred to as “thermal hysteresis”. Freezing point depres-
sion is not a colligative reaction, i.e., very low concentra-
tions of AFP in pure solutions are known to have approxi-
mately five hundred times greater depression than
colligative processes would predict.

Ideally, AFPs could be used to suppress ice formation in
a wide variety of environments. AFPs have the advantage
that they can be applied to a road surface or to an agricultural
plant ahead of time so that they would interact with ice
during formation and, further, they can be applied after the
onset of ice formation and serve to prevent continued ice
crystal formation. In addition, since AFPs are simply
polypeptide chains, they pose no hazard to the environment.
In winter flounder, the concentration of AFPs range from
1.0% to 3.0% depending on the species and the season;
therefore, AFPs are not produced in large enough quantities
in fish for the fish to be harvested as a source for an ice
preventing agent. Moreover, AFPs and AFGPs are typically
only produced in the fish during the winter months. As noted
above, Chakrabartty et al. have shown it is possible to
synthesize AFP using direct chemical processes; however,
these processes can be expensive and time consuming.
Peters et al., in Protein Engineering 3:145–151 (1989),
disclosed producing a semisynthetic winter flounder AFP in
Escherichia coli (E. coli). In Peters et al., a gene constructed
of a fused synthetic deoxyribonucleic acid (DNA) fragment
and a DNA fragment derived from a full length winter
flounder clone was inserted into a plasmid and the plasmid
was placed in the E. coli for production of a fusion protein.
The biosynthetic fusion protein produced contained part of
a pro-AFP and part of a β-galactosidase peptide and had
limited antifreeze activity after cleavage from
β-galactosidase. What is needed is a synthetic polypeptide
which has greater freezing point depression capability than
those AFPs which occur in nature and which can be eco-
nomically produced by biosynthetic processes.

SUMMARY OF THE INVENTION

It is therefore an object of the present invention to provide
a new AFP modeled after the native AFP found in the winter
flounder, Pseudopleuronectes americanus, but which is
longer than the native AFP and should have greater ability
to suppress ice formation than that which can be achieved by
solutions of the native AFP.

It is another object of this invention to provide a gene
coding for the new AFP which can be expressed in a
bacterial host.

It is yet another object of this invention to provide a
deoxyribonucleic acid (DNA) insert sequence which can be
easily placed in the gene coding for the new AFP which
ultimately will extend the length of new AFP produced by
the bacterial host and may provide for the crosslinking of
separate polypeptide chains.

According to the invention, an artificial peptide with a
specific sequence of amino acids modeled after an AFP
found in the winter flounder, Pseudopleuronectes americanus,
has been designed. The gene necessary to produce the artificial AFP has been synthesized and inserted
into E. coli, and the gene has been expressed in the E. coli.

In addition, a DNA insert sequence has been synthesized
which can be readily inserted into the gene for producing the
artificial AFP and serves the function of extending the length
of the AFP ultimately produced by the bacterial host and
may provide for the crosslinking of separate polypeptide
chains.

BRIEF DESCRIPTION OF THE DRAWINGS

The foregoing and other objects, aspects and advantages
will be better understood from the following detailed
description of a preferred embodiment of the invention with
reference to the drawings, in which:

FIG. 1 is the primary amino acid sequence of an AFP
found in the winter flounder Pseudopleuronectes americ-
ana;

FIG. 2 is a representational drawing of a forming ice
crystal and an antifreeze peptide showing how the spacing
allows hydrogen bonding to the face of the forming ice
crystal;

FIGS. 3a and 3b are isometric views of a forming ice
crystal and ice crystal with AFPS bound to steps thereon,
respectively, which together show how AFPS interact with
the forming ice crystal step according to the adsorption
inhibition hypothesis;

FIG. 4 is a base pair sequence of a synthesized gene w8r
and a synthesized amino acid sequence for an AFP polypep-
tide WFSR wherein the gene w8r codes for the AFP WFSR;

FIG. 5 is a 33 base pair sequence of a DNA insert
fragment which can be inserted in the w8r gene of FIG. 4
at the PST1 restriction site which extends the length of
WFSR AFP by one repeating unit to produce W9R;

FIG. 6 is a 66 base pair sequence of a dimer of the DNA
insert fragment shown in FIG. 5;

FIG. 7 is a 33 base pair sequence of a DNA insert
fragment similar to that shown in FIG. 5, but in which a
threonine codon has been replaced with a cysteine codon to
enable cross-linking of two AFP chains through disulfide
bridges;

FIGS. 8a–8g are seven synthetic restriction fragments
which were used in the process of producing the gene shown
in FIG. 4;

FIGS. 9a–9c are flow diagrams for constructing the gene
shown in FIG. 4 from the six synthetic restriction fragments
shown in FIGS. 8a–8f; and

FIG. 10 is a synthetic restriction fragment which will be
used to modify the gene shown in FIG. 4 to make the
N-terminal amino acids of the WFSR protein after digestion
with Clostripain identical to the N-terminal amino acids of
the native flounder AFP.

DETAILED DESCRIPTION OF A PREFERRED
EMBODIMENT OF THE INVENTION

Referring now to the drawings, and more particularly to
FIG. 4, there is shown a synthetic AFP peptide (SEQ ID
NO:2) and a synthetic gene (comprised of DNA SEQ ID
NO:3 and DNA SEQ ID NO:4) coding for the AFP peptide
having SEQ ID NO:2. The inventors have successfully
synthesized the gene (SEQ ID NO:3 and 4) having 336
nucleotides and expressed the gene in E. coli and have
produced the AFP peptide (SEQ ID NO:2) having 98 amino
acids according to the methods discussed below in the
experimental section. The synthetic gene was too large to be
produced as a single piece and was constructed from several
subfractions, therefore, the restriction endonuclease sites (i.e., ECO RI, Bgl II, Cla I, Sst II, Pst I, and Bam HI) where the subfractions were ligated together to form the synthetic gene are indicated above SEQ ID NO:3. Stars denote termination codons in the synthetic gene.

The AFP peptide in FIG. 4 includes five repeats of the following eleven amino acid sequence: thr-ala-thr-thr-ala-ala-thr-ala-ala-thr-ala-thr-ala (SEQ ID NO:5). The repeating SEQ ID NO:5 amino acid sequences are indicated by numbered blocked regions 2–6. In addition, the AFP peptide includes the following three amino acid sequences: thr-ala-ser-asp-ala-ala-ala-ala-ala-ala (SEQ ID NO:6) indicated as blocked region 1, thr-ala-thr-ala-ala-thr-ala-thr-ala (SEQ ID NO:7) indicated as blocked region 7, and thr-ala-ala-ala-ala-thr-ala-ala-thr-ala-ala (SEQ ID NO:8) indicated as blocked region 8.

A key feature of the synthetic and the native AFP is that there are several repeating eleven amino acid sequences, each of which typically includes two polar amino acids spaced apart by two nonpolar amino acids. The applicants are herein defining polar amino acids as those with side chains capable of forming hydrogen bonds and nonpolar amino acids as those with side chains not capable of forming hydrogen bonds. In the synthetic and the native AFPs, the nonpolar amino acids are usually alanine (ala) molecules; however, other nonpolar amino acids such as valine (val), isoleucine (ile), leucine (leu), glycine (gly), etc., could easily be substituted for theala molecules in the amino acid sequence. In the synthetic and the native AFPs, the polar amino acids are usually threonine (thr) or aspartic acid (asp); however, other polar amino acids such as asparagine (asn), glutamine (gln), glutamic acid (glu), serine (ser), etc., could easily be substituted for the thr and asp molecules in the amino acid sequence. The nonpolar amino acids serve as spacers to position the polar amino acids 4.5 Å apart in free space. As discussed above, the positions of the polar amino acids allows the AFP polypeptide to form hydrogen bonds with the forming ice crystal in competition with free water molecules and, thereby, block the steps for growth of the ice crystal. This process according to the “adsorption inhibition hypothesis”, in essence lowers the freezing point of water.

Seven of the eight repeating eleven amino acid sequences in FIG. 4, indicated by blocks 2–8 (SEQ ID NOs:5,7,8), fit the description of having a first polar amino acid, a second and third nonpolar amino acid, a fourth polar amino acid, and fifth through eleventh nonpolar amino acids. The eleven amino acid sequence indicated by block 1 (SEQ ID NO:6) differs from the sequences in blocks 2–8 only in that the third amino acid is a serine molecule. Serines, of course, are polar molecules which include a hydroxy group. The serine polypeptide may have some effect on the hydrogen bonding capability of the AFP and on its correct orientation relative to the face of the ice crystal prior to binding with it.

A major distinction between the synthetic AFP shown in FIG. 4 and native AFP found in winter flounder is the number of repeats of the eleven amino acid sequences containing the nine nonpolar amino acids and two polar amino acids. In the native AFP, the eleven amino acid sequence is repeated three to five times, but is not known to repeat more than five times in any known organism. In the synthetic AFP, the eleven amino acid sequence is repeated seven or eight times depending on how the function of the serine residue in the first block is understood.

As discussed above, there appears to be a relationship between AFP size and freezing point depression activity. If too small a region is blocked by an AFP on the forming ice crystal, the increase in the free energy barrier would be insignificant. In fact, in Chakrabartty et al., J. Bio. Chem. 264:11313–11316 (1989), it was reported that chemically synthesized AFP analogs with a minimum of three repeats of the basic sequence are required for ice suppression activity. Hence, larger molecules with a greater number of repeating sequences which can form a larger amount of hydrogen bonds should be more effective in preventing ice crystal growth. This phenomena is observed in natural systems where lower molecular weight antifreezes are less effective than high molecular weight antifreezes (see, Burcham et al., Biopolymers 23:1379–1395 (1984) and Schrag et al., Biochemica et Biophysica Acta). There is also evidence that on a molar basis, larger molecules are more effective in ice suppression than smaller ones. Kao et al., in Can. J. Zool. 64:578–582 (1986), pointed out that when the thermal hysteresis curves for peptide antifreezes are compared on a molar basis, the activity increases with increasing molecular weight. Scott et al., in Eur. J. Biochem. 168:629–633 (1987), analyzed various protein antifreezes from two different flounder species and have speculated that the reduced activity from the yellowtail (Lima nda ferrugineus) might be attributed to its size and smaller number of active sites compared to the AFP found in P. americanus.

The applicants’ synthesis of the AFP polypeptide shown in FIG. 4 in a bacterial host using the synthetic gene of FIG. 4 represents a significant advance over the previous methods of producing AFP polypeptides. Chakrabartty et al., J. Bio. Chem. 264:11313–11316 (1989), have only contemplated making extended chains of AFPs by direct chemical synthesis which is a more cumbersome process than biosynthetic techniques. Peters et al., in Protein Engineering 3:145–151 (1989), merely contemplated semisynthetic AFP where the native genes encoding for an AFP were inserted into a bacterial host for production of the AFP by biosynthetic techniques. However, the Peters et al. biosynthetic technique can only result in the bacteria producing AFPs having only 3–5 repeats of the eleven amino acid sequence responsible for blocking the steps in ice formation, since no known native system has more than 5 repeats of the sequence. It was the applicants’ foresight to design a completely new AFP polypeptide with a greater number of repeating sequences than the AFPs found in nature. The larger AFP polypeptide molecule has a greater number of hydrogen bonding sites and should result in greater depression of the freezing point of water than natural AFPs can achieve. Hence, the synthetic AFP should be more suitable to use in colder environments such as boat decks, airport runways, and agriculture.

Another important feature of the applicants invention is that the DNA sequence shown in FIG. 4 can readily be altered to create synthetic AFPS of even greater lengths than the AFP shown in FIG. 4. The applicants have synthesized the double stranded DNA insert fragment shown in FIG. 5.
wherein the two strands are represented by SEQ ID NO:9 and SEQ ID NO:10 and each strand has 33 nucleotides. The insert shown in FIG. 5 has sticky ends which are readily compatible with the PST1 restriction endonuclease site PST1 in the synthetic gene shown in FIG. 4. As can be seen by analyzing FIGS. 4 and 5 simultaneously, the sequence of 11 amino acids (SEQ ID NO:11 in FIG. 5), where the first and last nucleotides code for only part of the threonine amino acid, insertion of this 33 base pair fragment into the synthetic gene at the PST1 site according to well known biosynthetic techniques will regenerate a PST1 site at each end of the fragment in the newly created gene wfr. Furthermore, insertion will add a single repeat of the eleven amino acid sequence resulting in a 9 repeat AFP “WF9R”.

The insert fragment of FIG. 5 has been designed such that the spacing of the threonine residues is maintained and no frame shifts occur.

An added advantage of the insert fragment shown in FIG. 5 is that it is capable of “self polymerization”. That is, monomeric insert oligonucleotides like those shown in FIG. 5 can be ligated end to end to produce dimeric, trimeric and polymeric forms without losing the PST1 compatible ends present in the monomer. FIG. 6 shows an example of a dimer of the FIG. 5 sequence comprised of SEQ ID NO:12 and SEQ ID NO:13 where the ends of the dimeric form are identical to those of the monomer. Polymerized forms can easily be separated by size on gels and inserted in the PST1 site of the original WFSR sequence, creating “WF12R”, “WF2SR”, etc., forms of the AFP.

It should not be lost on the reader that while the insert fragments shown in FIGS. 5 and 6 add additional repeats of the eleven amino acid sequence responsible for hydrogen bonding to steps on an ice crystal, the AFP produced could merely be lengthened without adding additional repeating eleven amino acid sequences simply by inserting any 33 base pair nucleotide sequence having ends compatible with the PST1 site. If this is done, the spacing of the threonine residues is maintained and no frame shifts occur. Hence, the AFP produced will have the same first six repeating eleven amino acid sequences which can hydrogen bond to an ice crystal as in the WFSR of FIG. 4, an elongated polypeptide region of any particular quality desired, and the same eighth repeating eleven amino acid sequence as in the WFSR of FIG. 4.

Further in this line of reasoning, the applicants have determined that it would be quite easy to incorporate one or more cysteine residues into the sequence of an artificial AFP. If this were done, and if each such cysteine containing chain maintained a normal (i.e., active) tertiary structure, then more cysteine residues into the sequence of an artificial AFP could merely be lengthened without adding additional repeating eleven amino acid sequences merely by inserting any 33 nucleotide fragment into the synthetic gene at the PST1 site. If desired, the process of polymerization described in conjunction with FIG. 5 could be carried out with the fragment of FIG. 7 so that AFPs with more than one disulfide link could be created. The elongated single chain AFPs containing the cysteine residues will be expressed in bacteria, and later linked through the formation of interchain disulfide bridges after purification.

**EXAMPLE**

The following defines the synthesis of the wfr gene shown in FIG. 4 and its expression in *Escherichia coli.*

**Assembly of the wfr Coding Sequence**

The structural part of the wfr gene was assembled from six synthetic double stranded DNA restriction fragments: A, B1, B2, B3, B4, C and CA1 shown respectively in FIGS. 8a-8g. In each of the fragments A, B1, B2, B3, B4, C and CA1 the internal restriction enzyme recognition sites are indicated by lines above the sequences and the digestion pattern for each of these enzymes is indicated by small vertical arrows. Each of the fragments was commercially synthesized as two single-stranded oligonucleotides by the Promega Co. of Madison, Wis. FIG. 8a shows that fragment A is constructed from SEQ ID NO:16 and SEQ ID NO:17 and partially codes for SEQ ID NO:18. The stars denote termination codons. FIG. 8b shows that fragment B1 is constructed from SEQ ID NO:19 and SEQ ID NO:20 and codes for SEQ ID NO:21. FIG. 8c shows that fragment B2 is constructed from SEQ ID NO:22 and SEQ ID NO:23 and codes for SEQ ID NO:24. FIG. 8d shows that fragment B3 is constructed from SEQ ID NO:25 and SEQ ID NO:26 and codes for SEQ ID NO:27. FIG. 8e shows that fragment B4 is constructed from SEQ ID NO:28 and SEQ ID NO:29 and codes for SEQ ID NO:30. FIG. 8f shows that fragment C is constructed from SEQ ID NO:31 and SEQ ID NO:32 and codes for SEQ ID NO:33. FIG. 8g shows that fragment CA1, which is very similar except for the ending to fragment C in FIG. 8f, is constructed from SEQ ID NO:34 and SEQ ID NO:35 and codes for SEQ ID NO:36. The complimentary oligonucleotides were hybridized to each other and inserted into vectors according to the following process steps which are best illustrated in FIGS. 9a-9c.

The fragment A was digested with EcoRI and KpnI and ligated into EcoRI and KpnI cleaved pGem-3Z+(−). The resulting plasmid was named pGA. The identity of the insert in pGA (as well as in all the subsequent intermediate plasmids created in the course of this work) was confirmed by sequencing.

**Fragments B1 and B2** were ligated together and this ligation mixture was, together with fragment B3, added to the SstII and SmaI digested and dephosphorylated plasmid pGA and incubated with the T4 DNA ligase in a buffer. This reaction mixture was transformed into an *E. coli* host and the resulting clones were screened for presence of the plasmid pGAB3.

**Fragments B1 and C** were ligated together and the resulting fragment B1-C was excised from an agarose gel after electrophoresis of the ligation mixture. This fragment was then ligated with the fragment B3. This ligation mixture was again electrophoresed on an agarose gel and the B3-B1-C fragment was excised and ligated into BamHI and XmaIII digested pGAB3 to produce the plasmid pGAC.

**Cloning of the wfr Coding Sequence Under Control of the T7 Derived Regulatory Elements**

The applicants chose as their expression vector the plasmid pGEMEX-2 which is available from the Promega Co. of
Madison, Wis. Present in this plasmid is a fragment of the T7 bacteriophage gene 10 (T7g10), including its promoter, ribosome binding site, and 260 codons specifying the N-terminal amino acids of the gene 10 product (P10). As shown in FIG. 9b, downstream from this fragment is a multiple cloning site followed by a T7 transcription terminator sequence. When the pGEMEX-2 is transformed into the E.coli JM109(DE3), which harbors a chromosomal copy of the T7 RNA polymerase gene under control of the lacZ regulatory sequences, the P10 protein sequences will be transcribed by the T7 RNA polymerase upon induction by the isopropylthio-β-galactoside (IPTG) which is an inducer of the lacZ promoter. The T7 polymerase is very specific for the T7 promoters and will not transcribe any native E.coli genes. The protein 10 of the T7 phage is one of the most abundant proteins in the T7 infected cells due to high strength of its promoter and high efficiency of its ribosome binding site (see, Studier et al., J. Mol. Biol. 189:113–130 (1986) and Olin et al., Gene 73:227–235 (1988)). These features of the pGEMEX-2/JM109(DE3) vector/host combination have made it a promising system for high-efficiency expression of the wfr gene. The applicants’ objective, therefore, was to insert the wfr structural sequences in frame downstream from the T7 gene 10 and express it as a P10:WFR fusion protein.

FIG. 9b shows the EcoRI-BamHI fragment of pGAC, containing the wfr structural sequences, was subcloned into pGEMEX-2 to create the plasmid pGGAC. The in-frame fusion to the T7 gene 10 was created by cleaving off the EcoRI-ClaI fragment of wfr, filling the protruding ends and religating to produce the plasmid pX28.

To reduce the foreign (non-WFR) amino acid content of the N-terminal end of the fusion protein, the pX28 was digested with KpnI and NheI. Both termini were blunted by T4 DNA polymerase and the DNA was religated to produce the plasmid pX28. This procedure removed 257 codons from the P10 part of the fusion protein. Left in the resulting fusion protein were three N-terminal amino acids of the P10 followed by 24 amino acids coded by the multiple cloning site upstream from wfr sequences.

Expression of the Fusion Proteins Coded by the pX284 and pX28 Plasmids

The pX28 and pX284 were transformed into JM109 (DE3). The proteins produced in either the JM109(DE3) [pX284] or in the JM109(DE3)[pX284] after induction by IPTG in mid- or late-logarithmic growth phase, have been analyzed by sodium dodecyl sulfate (SDS)-polyacrylamide gel electrophoresis (SDS-PAGE). No fusion protein (expected MW=10.5 kd) was detected from the strain bearing pX28 while a low-level expression of the fusion protein (MW=37.3 kd) was observed in the pX284 bearing strain. Studies of cell viability in these strains have shown that both strains are adversely affected with respect to growth and stability of the plasmid (these effects were more pronounced in the pX28 harboring strain). The applicants concluded that the WFR must be toxic to the cells and that the cells were reacting to this toxicity by loss or mutation of the plasmid and by degradation of the proteins. To avert this problem the applicants decided to protect the fusion protein from intracellular degradation, as well as to protect the cells from the toxicity of WFR, by adding to the fusion a C-terminal end derived from E.coli β-galactosidase.

Generation of the P10:WFR:β-galactosidase Fusion Gene

The plasmid pMC1403 contains a fragment of the lac operon truncated at its 5’-end for all the regulatory sequences and the first seven codons of the lacZ gene (see, Casadaban et al., J. Bacteriol. 143:971–980 (1980)). The 5th codon of lacZ in this plasmid is attached at the BamHI site to a short multiple cloning cassette consisting of EcoRI, SmaI, and BamHI recognition sequences. To produce a translational fusion between the construct pX28 and the lacZ fragment from pMC1403 the C part of the wfr had to be modified to remove the two tandem termination codons present at its 3’-end (FIG. 8). The applicants therefore substituted the C fragment of pX28 with the fragment CA1 (FIG. 8g). The CA1 has a point deletion at the end of the first termination codon. This mutation introduced a frame shift which added three new amino acid codons to the wfr sequence and provided a potential for an in-frame read-through to the lacZ sequences upon ligation at the BamHI site. The resulting plasmid, pX28CA1, was digested with SmaI and BamHI and the released fragment, containing the entire fusion gene, including the regulatory sequences, was inserted into the SmaI and BamHI digested pMC1403, generating the plasmid pMX28L. Colonies of a lac- E.coli strain transformed with pMX28L were colored when plated on agar substituted with chromogenic indicator of β-galactosidase activity. This result confirmed the in-frame nature of the fusion with lacZ, as well as the expression of the fusion protein. This result has also shown that the E.coli RNA polymerase recognizes the T7 promoter and can initiate a low-level transcription from this promoter (this result was later confirmed on an SDS-PAGE). To increase the level of expression of the P10:WFR:β-gal fusion protein (pMC1403 is a lower copy number plasmid than pGEMEX-2 and the expression of the fusion protein in PMX28L have proven to interfere with the expression of the amp gene) the applicants subcloned the SmaI-Sall fragment of the pMX28L into pGEMEX-2, creating the plasmid pGX28L (FIG. 9C).

Expression of the Fusion Protein (P10:WFR:β-gal) in the JM109(DE3)[pGX28L] Cells

One half hour after addition of IPTG to the growth media, the JM109(DE3)[pGX28L] cells produce a protein of a higher apparent molecular weight (124 kd) then the β-galactosidase (116 kd). The double fusion protein was identified in gels with antibodies to the β-galactosidase portion. In the applicants initial attempts of expression only small amounts of this fusion protein were produced after 3-hr post-induction in complex media at 37° C. The applicants improved the yield of the protein dramatically (by 57.5%) by renewal of media at the time of induction. Further increase in yield (up to 630% of the initial value) was achieved by growing the induced cells at 25° C. overnight. At 37° C the yield of the fusion protein did not increase significantly at growth times longer than 1-hr post induction. The improvement in yield was measured by comparing the protein bands on SDS-PAGE using as an internal control a band representing a prominent cellular protein of 70 kD apparent MW. The optimal yield of the fusion protein is 4.5–6.5 mg of the protein per gram of wet cell pellet, or 19–20 mg of fusion protein per liter of cell culture.

Purification of the Fusion Protein Produced by the JM109 (DE3)[pGX28L] Cells

The cells produce the fusion protein partially as insoluble aggregates and partially as soluble protein. Recently it was learned that the percentage of soluble protein that can be recovered from the cells depends both on the method of growth (42–45% soluble protein recovered from the cells grown at 25° C. vs. 7–9% of soluble protein recovered from the 37° C. grown cells) as well as on the method of cell lysis (the insoluble aggregates of fusion protein can be solubilized by a pressure treatment in the French Pressure Cell). The
The fusion protein is purified as follows. The harvested cell pellet is resuspended in Tris hydrochloride (pH=8) and ethylenediaminetetraacetate (EDTA) containing buffer (TE), the cells are lysed in a French Pressure Cell (15000 psi) and the content of insoluble fusion protein in the cell extract is collected by centrifugation. The pellet is resuspended again in the TE buffer and pressure treated in the French Pressure Cell again and the resulting solution is then centrifuged.

The resulting pellet contains the rest of the aggregated fusion protein, which is essentially pure of most of the cellular proteins, except two (MW 20—30 kd) that always co-purify with this fraction. The supernatant from the second centrifugation is combined with the supernatant of the first centrifugation and the resulting solution is then centrifuged. The resulting pellet contains the rest of the aggregated fusion protein, which is essentially pure of most of the cellular proteins, except two (MW 20—30 kd) that always co-purify with this fraction. The supernatant from the second centrifugation is then combined with the supernatant of the first centrifugation and the fusion protein content of this combined supernatant is precipitated with 10% ammonium sulfate. The precipitate, containing 80—90% of the total produced fusion protein constitutes the partially purified protein used for further manipulations.

Excision of the WF8R Protein from the P10:WF8R:β-gal Fusion Protein

The WF8R is excised from the fusion protein by the enzyme Clostripain available from the Promega Co. of Madison, Wis. This enzyme digests proteins at the carboxylic site of arginine residues. As seen in FIG. 8f, cleavage at the carboxylic site of arginine will leave the WF8R protein with a C-terminal arginine (end of first row of the C fragment). At the N-terminal end of the WF8R protein, the fusion with P10 has added an arginine residue three amino acid residues in front of the serine residue initiating the WF8R in the design of the war gene (underlined in fragment A of FIG. 8a). Thus, the WF8R protein excised from the fusion protein by the Clostripain enzyme will have the following N-terminal amino acids: N-Asn-Ser-Met-Ser-(SEQ ID NO:43) where the underlined serine residue is the one underlined in FIG. 8a of fragment A.

The applicants are currently in the process of making a minor modification in the C fragment of thewf8r gene present in the pGS28L plasmid. This modification will make the N-terminal codons of the WF8R protein after digestion with Clostripain identical to the N-terminal codons of the native flounder AFP. The applicants are substituting the current EcoRI-SstII fragment with the DNA sequence (SEQ ID NO:39 and SEQ ID NO:40) and amino acid codon sequence (SEQ ID NO:41) shown in FIG. 10. The four codons which are different from the current version of the EcoRI-SstII fragment in the gene are underlined. In the new version of the gene the methionine codon will be substituted with an arginine codon and the following serine codon, which will thus encode the first amino acid of the WF8R protein released from the fusion with Clostripain cleavage, will become asparagine as in the native fish protein.

Purification of the WF8R Protein

The WF8R protein excised by the Clostripain digestion from the fusion protein should be purified by reverse phase high-performance liquid chromatography (HPLC) and assayed for its antifreeze activity.

The plasmid is a Pgs28L and is expressed in the bacterium JM109(DE3). This bacterium is not suitable for long term maintenance of cultures. Therefore, a deposit has been made with the American Type Culture Collection (ATCC) of Rockville, Md., of the plasmid PgX28L in the E. coli strain DH5α and has ATCC number.

While the invention has been described in terms of its preferred embodiments where a synthetic AFP longer than that which occurs in nature was biosynthetically produced in an E. coli host which had been modified with a synthetic gene having a particular coding sequence, those skilled in the art will recognize that the invention can be practiced with modification within the spirit and scope of the appended claims.

**SEQUENCE LISTING**

(1) GENERAL INFORMATION:

(iii) NUMBER OF SEQUENCES: 43

(2) INFORMATION FOR SEQ ID NO:1:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 38 amino acids
(B) TYPE: amino acid
(C) STRANDEDNESS: linear
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

Asp Thr Ala Ser Asp Ala Ala Ala Ala Ala Ala Leu Thr Ala Ala Asp
1 5 10 15
Asp Ala Ala Ala Ala Ala Leu Thr Ala Asp Ala Ala Ala Ala Ala
20 25 30
Ala Ala Thr Ala Ala Xaa
35

(2) INFORMATION FOR SEQ ID NO:2:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 98 amino acids
(B) TYPE: amino acid

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

Ala Ala Ala Ala Ala Leu Thr Ala Ala Asp Ala Ala Ala Ala Ala
1 5 10 15
Asp Ala Ala Ala Ala Ala Leu Thr Ala Asp Ala Ala Ala Ala Ala
20 25 30
Ala Ala Thr Ala Ala Xaa
35
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:

Met Ser Thr Ala Ser Asp Ala Ala Ala Ala Ala Ala Thr Ala Ala
1 5 10 15
Thr Ala Ala Ala Ala Ala Thr Ala Ala Ala Ala Ala Ala Ala Ala
20 25 30
Ala Ala Ala Thr Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala
35 40 45
Ala Thr Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala
50 55 60
Ala Ala Ala Thr Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala
65 70 75 80
Ala Ala Asn Ala Ala Ala Ala Ala Ala Thr Ala Ala Ala Ala Ala
85 90 95
Arg Gly

(2) INFORMATION FOR SEQ ID NO: 3:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 336 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: double
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(ix) FEATURE:
(A) NAME/KEY: CDS
(B) LOCATION: 31..324

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:

GAATTGCAG ATCTTAAAG AGTTTTATCG AGT TCC ACC GCT TCC GAC GCC

54

GCA GCA GCA GCT GCT ACT GCC GCG ACC GCA GCA GCC GCT TCC GAC GCC GCA GCT
Ala Ala Ala Ala Thr Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala
1 5 10 15 20

ACT GCC ACC GCA GCA GCC GCT GCG GCA GCC GCT GCC GCC GCC GCC ACC GCA GCA GCC GCT TCC GAC GCC GCA GCT
Thr Ala Thr Ala Thr Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Ala Al

(ii) MOLECULE TYPE: DNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

GGATCCTTAT CAACCACGAG GCOSGAGGCAG CGCCTGCGTT 60
TGCAGCTTCT GCAGGCGCAG CTTTAGCTGC AGCTAGCTGC CAGCCGCTGC 120
TGCAGCTCCT GCAGGCGCAG TGCTAGCTGTA GCAGCGGTAG GTCCCGCCAG 180
GCCTGCGCT GCAGGCGCAG TCCTGCGGTGC CGCCTGCGGC CAGCCGCTGC 240
GCCAGCTGCT GCTGGCGCTG CGCAGCTGCT GCACTGCTGC CGCAGCAGC 300
GGATCCTTAT AAACCTGCTT ACAGATCTAC GAAATC 336

(2) INFORMATION FOR SEQ ID NO:5:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 11 amino acids
(B) TYPE: amino acid
(C) STRANDEDNESS:
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

Thr Ala Ala Thr Ala Ala Ala Ala Ala Ala Ala 1 5 10

(2) INFORMATION FOR SEQ ID NO:6:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 11 amino acids
(B) TYPE: amino acid
(C) STRANDEDNESS:
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

Thr Ala Ala Ser Asp Ala Ala Ala Ala Ala Ala Ala Ala 1 5 10

(2) INFORMATION FOR SEQ ID NO:7:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 11 amino acids
(B) TYPE: amino acid
(C) STRANDEDNESS:
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:

Thr Ala Ala Thr Ala Ala Ala Ala Ala Leu Ala Ala Ala Ala 1 5 10

(2) INFORMATION FOR SEQ ID NO:8:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 11 amino acids
(B) TYPE: amino acid
(C) STRANDEDNESS:
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:

Thr Ala Ala Asn Ala Ala Ala Ala Ala Ala Ala Ala Ala 1 5 10
(2) INFORMATION FOR SEQ ID NO:9:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 33 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: double
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:
GCTGCGGCCG CGGCAGCCAC CGCAGCAACT GCA

(2) INFORMATION FOR SEQ ID NO:10:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 33 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: double
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:
GTAGCTGCGG TGGCTGCCGC GGCCGCAGCT GCA

(2) INFORMATION FOR SEQ ID NO:11:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 11 amino acids
(B) TYPE: amino acid
(C) STRANDEDNESS:
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:
Ala Ala Ala Ala Ala Ala Ala Thr Ala Ala Thr

1  5  10

(2) INFORMATION FOR SEQ ID NO:12:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 66 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: double
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:
GCTGCGGCCG CGGCAGCCAC CGCAGCAACT GCAGCTGCGG CCGCGGCAGC CACCGCAGCA

(2) INFORMATION FOR SEQ ID NO:13:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 66 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: double
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:
GCTGCGGCCG CGGCAGCCAC CGCAGCAACT GCAGCTGCGG CCGCGGCAGC CACCGCAGCA

ACTGCA

60

(2) INFORMATION FOR SEQ ID NO:14:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 66 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: double
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:
GCTGCGGCCG CGGCAGCCAC CGCAGCAACT GCAGCTGCGG CCGCGGCAGC CACCGCAGCA

ACTGCA

66
(2) INFORMATION FOR SEQ ID NO:14:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 33 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: double
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:

GCTGCGGCCG CGGCAGCCTG CGCAGCAACT GCA

(2) INFORMATION FOR SEQ ID NO:15:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 33 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: double
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:

GTAGCTGCGC AGGCTGCCGC GGCCGCAGCT GCA

(2) INFORMATION FOR SEQ ID NO:16:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 67 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: double
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid
(A) DESCRIPTION: /desc = “synthetic DNA”

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:

GTGTGAATTC GTAGATCTGT AAGGAGGTTT ATCGATGTCC ACCGCTTCCG ACGCCGCGGT

ACCGAGG

(2) INFORMATION FOR SEQ ID NO:17:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 67 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: double
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid
(A) DESCRIPTION: /desc = “synthetic DNA”

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:17:

CCTCGGTACC GC GGCGTCGG AAGCGGTGGA CATCGATAAA CCTCCTTACA GATCTACGAA

TTCACAC

(2) INFORMATION FOR SEQ ID NO:1B:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 15 amino acids
(B) TYPE: amino acid
(C) STRANDEDNESS: linear

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1B:

Ala Gly Leu Ser Met Ser Thr Ala Ser Asp Ala Ala Ala Ala
(2) INFORMATION FOR SEQ ID NO:19:
(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 29 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: double
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid
(A) DESCRIPTION: /desc = "synthetic DNA"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:19:

GCCAGCAGCA GCTGCTACTG CTGCCACCG

(2) INFORMATION FOR SEQ ID NO:20:
(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 28 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: double
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid
(A) DESCRIPTION: /desc = "synthetic DNA"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:

TGGCAGCAGT AGCAGCTGCT GCTGCCGC

(2) INFORMATION FOR SEQ ID NO:21:
(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 10 amino acids
(B) TYPE: amino acid
(C) STRANDEDNESS: linear
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:21:

Ala Ala Ala Ala Ala Thr Ala Ala Thr Ala

1 5 10

(2) INFORMATION FOR SEQ ID NO:22:
(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 64 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: double
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid
(A) DESCRIPTION: /desc = "synthetic DNA"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:

CAGCGGCGGC TGCGGCAGCT ACTGCGGCGA CCGCAGCAGC GGCTGCGGCA GCTACTGCGG

CGAC

(2) INFORMATION FOR SEQ ID NO:23:
(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 72 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: double
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid
(A) DESCRIPTION: /desc = "synthetic DNA"
(Xi) SEQUENCE DESCRIPTION: SEQ ID NO:23:
CGGCCGTCGC CGCAGTAGCT GCCGCAGCCG CTGCTGCGGT CGCCGCAGTA GCTGCCGCAG
60
CGGCCTGC GG 72

(2) INFORMATION FOR SEQ ID NO:24:
(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 21 amino acids
(B) TYPE: amino acid
(C) STRANDEDNESS:
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: peptide

(Xi) SEQUENCE DESCRIPTION: SEQ ID NO:24:
Ala Ala Ala Ala Ala Ala Thr Ala Ala Thr Ala Ala Ala Ala Ala Ala Ala Ala Ala
1 5 10 15
Ala Thr Ala Ala Thr
20

(2) INFORMATION FOR SEQ ID NO:25:
(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 62 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: double
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid
(A) DESCRIPTION: /desc = "synthetic DNA"

(Xi) SEQUENCE DESCRIPTION: SEQ ID NO:25:
GGCCGCAGCG GCTGCGGCAG CTACCGCTGC TACCGCAGCA GCTGCTGCGG CAGCAACTGC
60
GG 62

(2) INFORMATION FOR SEQ ID NO:26:
(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 57 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: double
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid
(A) DESCRIPTION: /desc = "synthetic DNA"

(Xi) SEQUENCE DESCRIPTION: SEQ ID NO:26:
CCGCAGTTGC TGCCGCAGCA GCTGCTGCGG TAGCAGCGGT AGCTGCCGCA GCCGCTG
5

(2) INFORMATION FOR SEQ ID NO:27:
(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 20 amino acids
(B) TYPE: amino acid
(C) STRANDEDNESS:
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: peptide

(Xi) SEQUENCE DESCRIPTION: SEQ ID NO:27:
Ala Ala Ala Ala Ala Ala Ala Thr Ala Ala Thr Ala Ala Ala Ala Ala Ala Ala
1 5 10 15
Ala Ala Thr Ala
20

(2) INFORMATION FOR SEQ ID NO:28:
(i) SEQUENCE CHARACTERISTICS:
   (A) LENGTH: 41 base pairs
   (B) TYPE: nucleic acid
   (C) STRANDEDNESS: double
   (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid
   (A) DESCRIPTION: /desc = “synthetic DNA”

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:28:

CCACCGCAGC AGCGGCTGCG GCAGCTACCG CAGCTACTGC A

(2) INFORMATION FOR SEQ ID NO:29:

(i) SEQUENCE CHARACTERISTICS:
   (A) LENGTH: 37 base pairs
   (B) TYPE: nucleic acid
   (C) STRANDEDNESS: double
   (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid
   (A) DESCRIPTION: /desc = “synthetic DNA”

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:29:

GTAGCTGCGG TAGCTGCCGC AGCCGCTGCT GCGGTGG

(2) INFORMATION FOR SEQ ID NO:30:

(i) SEQUENCE CHARACTERISTICS:
   (A) LENGTH: 14 amino acids
   (B) TYPE: amino acid
   (C) STRANDEDNESS: linear
   (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:

Ala Thr Ala Ala Ala Ala Ala Ala Ala Thr Ala Ala Thr A]

(2) INFORMATION FOR SEQ ID NO:31:

(i) SEQUENCE CHARACTERISTICS:
   (A) LENGTH: 82 base pairs
   (B) TYPE: nucleic acid
   (C) STRANDEDNESS: double
   (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid
   (A) DESCRIPTION: /desc = “synthetic DNA”

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:31:

GCTAAAGCTG CGCCTGCTAC TGCTGCAAAC GCAGCGGCTG CTGCGGCGGC GACTGCTGCG GCAGCGGCTC GTGGTTGATA AG

GCTAAAGCTG CGCCTGCTAC TGCTGCAAAC GCAGCGGCTG CTGCGGCGGC GACTGCTGCG GCAGCGGCTC GTGGTTGATA AG

(2) INFORMATION FOR SEQ ID NO:32:

(i) SEQUENCE CHARACTERISTICS:
   (A) LENGTH: 90 base pairs
   (B) TYPE: nucleic acid
   (C) STRANDEDNESS: double
   (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid
   (A) DESCRIPTION: /desc = “synthetic DNA”

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:32:

GATCCTTATCAACCACGGA CGCTGCGCCA GCAGCTGCCGC GCAGCCGAC GCCTGGCTT
(2) INFORMATION FOR SEQ ID NO:33:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 25 amino acids
(B) TYPE: amino acid
(C) STRANDEDNESS:
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:33:

Ala Lys Ala Ala Ala Leu Thr Ala Ala Asn Ala Ala Ala Ala Ala Ala
1 5 10 15
Ala Thr Ala Ala Ala Ala Ala Arg Gly
20 25

(2) INFORMATION FOR SEQ ID NO:34:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 81 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: double
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid

(a) DESCRIPTION: /desc = “synthetic DNA”

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:34:

GCTAAAGCTG CGGCTCTGAC TGCTGCAAAC GCAGCGGCTG CTGCGGCGGC GACTGCTGCG
GCAGCGGCTC GTGGTTGTAA G

(2) INFORMATION FOR SEQ ID NO:35:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 89 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: double
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid

(a) DESCRIPTION: /desc = “synthetic DNA”

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:35:

GATCCTTACA ACCACGAGCC GCTGCCGCAG CAGTCGCCGC CGCAGCAGCC GCTGCGTTTG
CAGCAGTCAG AGCCGCAGCT TTAGCTGCA

(2) INFORMATION FOR SEQ ID NO:36:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 28 amino acids
(B) TYPE: amino acid
(C) STRANDEDNESS:
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:36:

Ala Lys Ala Ala Ala Leu Thr Ala Ala Asn Ala Ala Ala Ala Ala Ala
1 5 10 15
Ala Thr Ala Ala Ala Ala Ala Arg Gly Cys Lys Asp
20 25

(2) INFORMATION FOR SEQ ID NO:37:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 35 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: double
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid
   (A) DESCRIPTION: /desc = “synthetic DNA”

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:37:

AATTCAGATC TGCGAGATAC CGCTTCCGAC GCCGC

(2) INFORMATION FOR SEQ ID NO:38:

(i) SEQUENCE CHARACTERISTICS:
   (A) LENGTH: 29 base pairs
   (B) TYPE: nucleic acid
   (C) STRANDEDNESS: double
   (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid
   (A) DESCRIPTION: /desc = “synthetic DNA”

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:38:

GGCGTCGGAA GCGGTATCTC GCAGATCTG

(2) INFORMATION FOR SEQ ID NO:39:

(i) SEQUENCE CHARACTERISTICS:
   (A) LENGTH: 12 amino acids
   (B) TYPE: amino acid
   (C) STRANDEDNESS:
   (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:39:

Asn Ser Asp Leu Arg Asp Thr Ala Ser Asp Ala Ala
1  5  10

(2) INFORMATION FOR SEQ ID NO:40:

(i) SEQUENCE CHARACTERISTICS:
   (A) LENGTH: 78 base pairs
   (B) TYPE: nucleic acid
   (C) STRANDEDNESS: single
   (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid
   (A) DESCRIPTION: /desc = “synthetic DNA”

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:40:

GGCCAAGTCG GCCGAGCTCG GAATTCGTCG ACCTCGAGGG ATCCGGGCCC TCTAGATGCG GCCGCATGCA TAAGCTTG

(2) INFORMATION FOR SEQ ID NO:41:

(i) SEQUENCE CHARACTERISTICS:
   (A) LENGTH: 17 base pairs
   (B) TYPE: nucleic acid
   (C) STRANDEDNESS: double
   (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid
   (A) DESCRIPTION: /desc = “synthetic DNA”

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:41:

GAATTCGCCG GGATCCC

(2) INFORMATION FOR SEQ ID NO:42:


Having thus described our invention, what we claim as new and desire to secure by Letters Patent is as follows:

1. A synthetic DNA sequence having a sequence of nucleotides coding for an antifreeze polypeptide modeled after the antifreeze polypeptide found in winter flounder having greater than five repeats of an eleven amino acid sequence where the first and fourth amino acids in said eleven amino acid sequence are selected from the group consisting of threonine, asparagine, glutamine, glutamic acid, serine and aspartic acid and the second, third, and fifth through eleventh amino acids in said eleven amino acid sequence are selected from the group consisting of alanine, glycine, lysine, isoleucine, valine, serine and leucine.

2. A synthetic DNA sequence as recited in claim 1 wherein a group of adjacent codons within an active transcription region of said sequence coding for said antifreeze polypeptide provides a restriction endonuclease site.

3. A synthetic DNA sequence as recited in claim 2 wherein said restriction endonuclease site is PST1.

4. A synthetic DNA sequence as recited in claim 1 wherein said first and fourth amino acids in said eleven amino acid sequence coded for by said sequence of nucleotides are selected from the group consisting of threonine and aspartic acid.

5. A synthetic DNA sequence having a sequence of nucleotides coding for an antifreeze polypeptide modeled after the antifreeze polypeptide found in winter flounder comprising a sequence of nucleotides coding for a protein which has an amino acid sequence defined by blocks 1–6 of FIG. 4 (SEQ ID NO:2).

6. A synthetic DNA sequence having a sequence of nucleotides coding for an antifreeze polypeptide modeled after the antifreeze polypeptide found in winter flounder comprising a sequence of nucleotides coding for a protein which has an amino acid sequence defined by blocks 2–7 of FIG. 4 (SEQ ID NO:2).

7. A synthetic DNA sequence having a sequence of nucleotides coding for an antifreeze polypeptide modeled after the antifreeze polypeptide found in winter flounder comprising a sequence of nucleotides coding for a protein which has an amino acid sequence defined by blocks 3–8 of FIG. 4.

8. A synthetic DNA sequence having a sequence of nucleotides coding for an antifreeze polypeptide modeled after the antifreeze polypeptide found in winter flounder comprising a sequence of nucleotides coding for a protein which has an amino acid sequence defined by blocks 1–8 of FIG. 4 (SEQ ID NO:2).

9. A synthetic DNA sequence having a sequence of nucleotides coding for an antifreeze polypeptide modeled after the antifreeze polypeptide found in winter flounder comprising a sequence of nucleotides coding for a protein which has an amino acid sequence with greater than five of the eight blocks shown in FIG. 4 (SEQ ID NO:2).

10. A gene having a deoxyribonucleic acid sequence as shown in FIG. 4, wherein said sequence is selected from the group consisting of SEQ ID NOS: 3 and 4.

11. A gene having a deoxyribonucleic acid sequence coding for a protein which has an amino acid sequence as shown in FIG. 4 (SEQ ID NO:2).

12. A bacterial strain including a synthetic DNA sequence having a sequence of nucleotides coding for an antifreeze polypeptide modeled after the antifreeze polypeptide found in winter flounder wherein said antifreeze polypeptide has of greater than five repeats of an eleven amino acid sequence where the first and fourth amino acids in said eleven amino acid sequence are selected from the group consisting of threonine, asparagine, glutamine, glutamic acid, serine and aspartic acid and the second, third, and fifth through eleventh amino acids in said eleven amino acid sequence are selected from the group consisting of alanine, glycine, lysine, isoleucine, valine, serine, and leucine, said synthetic DNA sequence being expressed in said bacterial strain.

13. A bacterial strain as recited in claim 12 wherein said first and fourth amino acids in said eleven amino acid sequence are selected from the group consisting of threonine and aspartic acid.

14. A bacterial host transformed by a gene having a deoxyribonucleic acid sequence as shown in FIG. 4, wherein said sequence is selected from the group consisting of SEQ ID NOS: 3 and 4, which can express an amino acid sequence as shown in FIG. 4 (SEQ ID NO: 2).

15. The transformed bacterial host recited in claim 14 wherein said bacterial host is Eschericia coli.
16. A bacterial host having all the identifying characteristics of ATCC deposit No. 68425.

17. A method of enlarging the size of a biosynthetically produced antifreeze polypeptide modeled after the antifreeze polypeptide found in winter flounder, comprising the steps of:

- providing a DNA sequence coding for an antifreeze polypeptide modeled after the antifreeze polypeptide found in winter flounder that has at least three adjacent eleven amino acid sequences where the first and fourth amino acids in said eleven amino acid sequences are selected from the group consisting of threonine, aspartic acid, glutamine, glutamic acid, serine, and asparagine, and where the second, third, and fifth through eleventh amino acids in said eleven amino acid sequence are selected from the group consisting of alanine, lysine, leucine, isoleucine, glycine, serine, and valine,

- providing a restriction endonuclease site in said DNA sequence positioned within a region coding for said three adjacent eleven amino acid sequences;

- providing a DNA insert fragment which is a multiple of thirty three nucleic acids in length, said DNA insert fragment having first and second sticky ends which are compatible with said restriction endonuclease site, said DNA insert fragment being insertable within said DNA sequence without a frame shift; and

- ligating said DNA insert fragment within said DNA sequence such that said DNA insert fragment extends said DNA sequence coding for said antifreeze polypeptide that has at least three adjacent eleven amino acid sequences by at least one additional eleven amino acid sequence.

18. A method as recited in claim 17 wherein said one additional eleven amino acid sequence includes a cysteine residue.

19. A method as recited in claim 17 wherein said DNA insert fragment has the nucleotide sequence shown in FIG. 7, wherein said sequence is selected from the group consisting of SEQ ID NOS: 14 and 15.

20. A method as recited in claim 17 wherein said one additional eleven amino acid sequence includes first and fourth amino acids selected from the group consisting of threonine, aspartic acid, glutamine, glutamic acid, serine and asparagine, and second, third, and fifth through eleventh amino acids selected from the group consisting of alanine, lysine, leucine, isoleucine, glycine, serine and valine.

21. A method as recited in claim 17 wherein said DNA insert fragment has the nucleotide sequence shown in FIG. 5, wherein said sequence is selected from the group consisting of SEQ ID NOS: 9 and 10.

22. A method as recited in claim 17 wherein said DNA insert fragment has the nucleotide sequence shown in FIG. 6, wherein said sequence is selected from the group consisting of SEQ ID NOS: 12 and 13.

23. A method as recited in claim 17 wherein said first and second sticky ends of said DNA insert fragment are compatible with each other.