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(57) Abrégé/Abstract:
The present invention relates to transgenic animals, as well as compositions and methods relating to the characterization of gene function. Specifically, the present invention provides transgenic mice comprising disruptions in PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO102, PRO788, PRO792, PRO840, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 genes. Such in vivo studies and characterizations may provide valuable identification and discovery of therapeutics and/or treatments useful in the prevention, amelioration or correction of diseases or dysfunctions associated with gene disruptions such as neurological disorders; cardiovascular, endothelial or angiogenic disorders, eye abnormalities; immunological disorders; oncological disorders; bone metabolic abnormalities or disorders; lipid metabolic disorders; or developmental abnormalities.
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(54) Title: NOVEL GENE DISRUPTIONS, COMPOSITIONS AND METHODS RELATING THERETO

(55) Abstract: The present invention relates to transgenic animals, as well as compositions and methods relating to the characterization of gene function. Specifically, the present invention provides transgenic mice comprising disruptions in PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO672, PRO929, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO247, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1855, PRO1879, PRO3446, PRO3543, PRO4329, PRO5733, PRO859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99048, PRO2894, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO30332, PRO38465 or PRO3464 genes. Such in vivo studies and characterizations may provide valuable identification and discovery of therapeutics and/or treatments useful in the prevention, amelioration or correction of diseases or dysfunctions associated with gene disruptions such as neurological disorders; cardiovascular, endothelial or angiogenic disorders; eye abnormalities; immunological disorders; oncological disorders; bone metabolic abnormalities or disorders; lipid metabolic disorders; or developmental abnormalities.
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LA PRÉSENTE PARTIE DE CETTE DEMANDE OU CE BREVET COMPRENDS PLUS D’UN TOME.

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NOVEL GENE DISRUPTIONS, COMPOSITIONS AND METHODS RELATING THERETO

FIELD OF THE INVENTION

The present invention relates to compositions, including transgenic and knockout animals and methods of using such compositions for the diagnosis and treatment of diseases or disorders.

BACKGROUND OF THE INVENTION

Extracellular proteins play important roles in, among other things, the formation, differentiation and maintenance of multicellular organisms. The fate of many individual cells, e.g., proliferation, migration, differentiation, or interaction with other cells, is typically governed by information received from other cells and/or the immediate environment. This information is often transmitted by secreted polypeptides (for instance, mitogenic factors, survival factors, cytotoxic factors, differentiation factors, neuropeptides, and hormones) which are, in turn, received and interpreted by diverse cell receptors or membrane-bound proteins. These secreted polypeptides or signaling molecules normally pass through the cellular secretory pathway to reach their site of action in the extracellular environment.

Secreted proteins have various industrial applications, including as pharmaceuticals, diagnostics, biosensors and bioreactors. Most protein drugs available at present, such as thrombolytic agents, interferons, interleukins, erythropoietins, colony stimulating factors, and various other cytokines, are secretory proteins. Their receptors, which are membrane proteins, also have potential as therapeutic or diagnostic agents. Efforts are being undertaken by both industry and academia to identify new, native secreted proteins. Many efforts are focused on the screening of mammalian recombinant DNA libraries to identify the coding sequences for novel secreted proteins. Examples of screening methods and techniques are described in the literature [see, for example, Klein et al., Proc. Natl. Acad. Sci. 93:7108-7113 (1996); U.S. Patent No. 5,536,637].

Membrane-bound proteins and receptors can play important roles in, among other things, the formation, differentiation and maintenance of multicellular organisms. The fate of many individual cells, e.g., proliferation, migration, differentiation, or interaction with other cells, is typically governed by information received from other cells and/or the immediate environment. This information is often transmitted by secreted polypeptides (for instance, mitogenic factors, survival factors, cytotoxic factors, differentiation factors, neuropeptides, and hormones) which are, in turn, received and interpreted by diverse cell receptors or membrane-bound proteins. Such membrane-bound proteins and cell receptors include, but are not limited to, cytokine receptors, receptor kinases, receptor phosphatases, receptors involved in cell-cell interactions, and cellular adhesion molecules like selectins and integrins. For instance, transduction of signals that regulate cell growth and differentiation is regulated in part by phosphorylation of various cellular proteins. Protein tyrosine kinases, enzymes that catalyze that process, can also act as growth factor receptors. Examples include fibroblast growth factor receptor and nerve growth factor receptor.

Membrane-bound proteins and receptor molecules have various industrial applications, including as pharmaceutical and diagnostic agents. Receptor immuno-adhesions, for instance, can be employed as therapeutic
agents to block receptor-ligand interactions. The membrane-bound proteins can also be employed for screening
of potential peptide or small molecule inhibitors of the relevant receptor/ligand interaction.

Efforts are being undertaken by both industry and academia to identify new, native receptor or membrane-bound proteins. Many efforts are focused on the screening of mammalian recombinant DNA libraries to identify the coding sequences for novel receptor or membrane-bound proteins.

Given the importance of secreted and membrane-bound proteins in biological and disease processes, in vivo studies and characterizations may provide valuable identification and discovery of therapeutics and/or treatments useful in the prevention, amelioration or correction of diseases or dysfunctions. In this regard, genetically engineered mice have proven to be invaluable tools for the functional dissection of biological processes relevant to human disease, including immunology, cancer, neuro-biology, cardiovascular biology, obesity and many others. Gene knockouts can be viewed as modeling the biological mechanism of drug action by presaging the activity of highly specific antagonists in vivo. Knockout mice have been shown to model drug activity; phenotypes of mice deficient for specific pharmaceutical target proteins can resemble the human clinical phenotype caused by the corresponding antagonist drug. Gene knockouts enable the discovery of the mechanism of action of the target, the predominant physiological role of the target, and mechanism-based side-effects that might result from inhibition of the target in mammals. Examples of this type include mice deficient in the angiotensin converting enzyme (ACE) [Esther, C.R. et al., Lab. Invest., 74:953-965 (1996)] and cyclooxygenase-1 (COX1) genes [Langenbach, R. et al., Cell, 83:483-492 (1995)]. Conversely, knocking the gene out in the mouse can have an opposite phenotypic effect to that observed in humans after administration of an agonist drug to the corresponding target. Examples include the erythropoietin knockout [Wu, C.S. et al., Cell, 83:59-67 (1996)], in which a consequence of the mutation is deficient red blood cell production, and the GABA(A)-R-B3 knockout [DeLorey, T.M., J. Neurosci., 18:8505-8514 (1998)], in which the mutant mice show hyperactivity and hyper-responsiveness. Both these phenotypes are opposite to the effects of erythropoietin and benzodiazepine administration in humans. A striking example of a target validated using mouse genetics is the ACC2 gene. Although the human ACC2 gene had been identified several years ago, interest in ACC2 as a target for drug development was stimulated only recently after analysis of ACC2 function using a knockout mouse. ACC2 mutant mice eat more than their wild-type littermates, yet burn more fat and store less fat in their adipocytes, making this enzyme a probable target for chemical antagonism in the treatment of obesity [Abu-Elheiga, L. et al., Science, 291:2613-2616 (2001)].

In the instant application, mutated gene disruptions have resulted in phenotypic observations related to various disease conditions or dysfunctions including: CNS/neurological disturbances or disorders such as anxiety; eye abnormalities and associated diseases; cardiovascular, endothelial or angiogenic disorders including atherosclerosis; abnormal metabolic disorders including diabetes and dyslipidemias associated with elevated serum triglycerides and cholesterol levels; immunological and inflammatory disorders; oncological disorders; bone metabolic abnormalities or disorders such as arthritis, osteoporosis and osteopetrosis; or a developmental disease such as embryonic lethality.

SUMMARY OF THE INVENTION
A. **Embodiments**

The invention provides an isolated nucleic acid molecule comprising a nucleotide sequence that encodes a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide.

In one aspect, the isolated nucleic acid molecule comprises a nucleotide sequence having at least about 80% nucleic acid sequence identity, alternatively at least about 81% nucleic acid sequence identity, alternatively at least about 82% nucleic acid sequence identity, alternatively at least about 83% nucleic acid sequence identity, alternatively at least about 84% nucleic acid sequence identity, alternatively at least about 85% nucleic acid sequence identity, alternatively at least about 86% nucleic acid sequence identity, alternatively at least about 87% nucleic acid sequence identity, alternatively at least about 88% nucleic acid sequence identity, alternatively at least about 89% nucleic acid sequence identity, alternatively at least about 90% nucleic acid sequence identity, alternatively at least about 91% nucleic acid sequence identity, alternatively at least about 92% nucleic acid sequence identity, alternatively at least about 93% nucleic acid sequence identity, alternatively at least about 94% nucleic acid sequence identity, alternatively at least about 95% nucleic acid sequence identity, alternatively at least about 96% nucleic acid sequence identity, alternatively at least about 97% nucleic acid sequence identity, alternatively at least about 98% nucleic acid sequence identity and alternatively at least about 99% nucleic acid sequence identity to (a) a DNA molecule encoding a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide having a full-length amino acid sequence as disclosed herein, an amino acid sequence lacking the signal peptide as disclosed herein, an extracellular domain of a transmembrane protein, with or without the signal peptide, as disclosed herein or any other specifically defined fragment of the full-length amino acid sequence as disclosed herein, or (b) the complement of the DNA molecule of (a).

In other aspects, the isolated nucleic acid molecule comprises a nucleotide sequence having at least about 80% nucleic acid sequence identity, alternatively at least about 81% nucleic acid sequence identity, alternatively at least about 82% nucleic acid sequence identity, alternatively at least about 83% nucleic acid sequence identity, alternatively at least about 84% nucleic acid sequence identity, alternatively at least about 85% nucleic acid sequence identity, alternatively at least about 86% nucleic acid sequence identity, alternatively at least about 87% nucleic acid sequence identity, alternatively at least about 88% nucleic acid sequence identity, alternatively at least about 89% nucleic acid sequence identity, alternatively at least about 90% nucleic acid sequence identity, alternatively at least about 91% nucleic acid sequence identity, alternatively at least about 92% nucleic acid sequence identity, alternatively at least about 93% nucleic acid sequence identity, alternatively at least about 94% nucleic acid sequence identity, alternatively at least about 95% nucleic acid sequence identity, alternatively at least about 96% nucleic acid sequence identity, alternatively at least about 97% nucleic acid sequence identity.
alternatively at least about 98% nucleic acid sequence identity and alternatively at least about 99% nucleic acid sequence identity to (a) a DNA molecule comprising the coding sequence of a full-length PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide cDNA as disclosed herein, the coding sequence of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide lacking the signal peptide as disclosed herein, the coding sequence of an extracellular domain of a transmembrane PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, with or without the signal peptide, as disclosed herein or the coding sequence of any other specifically defined fragment of the full-length amino acid sequence as disclosed herein, or (b) the complement of the DNA molecule of (a).

In a further aspect, the invention concerns an isolated nucleic acid molecule comprising a nucleotide sequence having at least about 80% nucleic acid sequence identity, alternatively at least about 81% nucleic acid sequence identity, alternatively at least about 82% nucleic acid sequence identity, alternatively at least about 83% nucleic acid sequence identity, alternatively at least about 84% nucleic acid sequence identity, alternatively at least about 85% nucleic acid sequence identity, alternatively at least about 86% nucleic acid sequence identity, alternatively at least about 87% nucleic acid sequence identity, alternatively at least about 88% nucleic acid sequence identity, alternatively at least about 89% nucleic acid sequence identity, alternatively at least about 90% nucleic acid sequence identity, alternatively at least about 91% nucleic acid sequence identity, alternatively at least about 92% nucleic acid sequence identity, alternatively at least about 93% nucleic acid sequence identity, alternatively at least about 94% nucleic acid sequence identity, alternatively at least about 95% nucleic acid sequence identity, alternatively at least about 96% nucleic acid sequence identity, alternatively at least about 97% nucleic acid sequence identity, alternatively at least about 98% nucleic acid sequence identity and alternatively at least about 99% nucleic acid sequence identity to (a) a DNA molecule that encodes the same mature polypeptide encoded by any of the human protein cDNAs deposited with the ATCC as disclosed herein, or (b) the complement of the DNA molecule of (a).

Another aspect of the invention provides an isolated nucleic acid molecule comprising a nucleotide sequence encoding a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543,
PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide which is either transmembrane domain-deleted or transmembrane domain-inactivated, or is complementary to such encoding nucleotide sequence, wherein the transmembrane domain(s) of such polypeptide are disclosed herein. Therefore, soluble extracellular domains of the herein described PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides are contemplated.

The invention also provides fragments of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide coding sequence, or the complement thereof, that may find use as, for example, hybridization probes, for encoding fragments of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide that may optionally encode a polypeptide comprising a binding site for an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody or as antisense oligonucleotide probes. Such nucleic acid fragments usually are or are at least about 10 nucleotides in length, alternatively are or are at least about 15 nucleotides in length, alternatively are or are at least about 20 nucleotides in length, alternatively are or are at least about 30 nucleotides in length, alternatively are or are at least about 40 nucleotides in length, alternatively are or are at least about 50 nucleotides in length, alternatively are or are at least about 60 nucleotides in length, alternatively are or are at least about 70 nucleotides in length, alternatively are or are at least about 80 nucleotides in length, alternatively are or are at least about 90 nucleotides in length, alternatively are or are at least about 100 nucleotides in length, alternatively are or are at least about 110 nucleotides in length, alternatively are or are at least about 120 nucleotides in length, alternatively are or are at least about 130 nucleotides in length, alternatively are or are at least about 140 nucleotides in length, alternatively are or are at least about 150 nucleotides in length, alternatively are or are at least about 160 nucleotides in length, alternatively are or are at least about 170 nucleotides in length.
about 170 nucleotides in length, alternatively are or are at least about 180 nucleotides in length, alternatively are or are at least about 190 nucleotides in length, alternatively are or are at least about 200 nucleotides in length, alternatively are or are at least about 250 nucleotides in length, alternatively are or are at least about 300 nucleotides in length, alternatively are or are at least about 350 nucleotides in length, alternatively are or are at least about 400 nucleotides in length, alternatively are or are at least about 450 nucleotides in length, alternatively are or are at least about 500 nucleotides in length, alternatively are or are at least about 600 nucleotides in length, alternatively are or are at least about 700 nucleotides in length, alternatively are or are at least about 800 nucleotides in length, alternatively are or are at least about 900 nucleotides in length and alternatively are or are at least about 1000 nucleotides in length, wherein in this context the term “about” means the referenced nucleotide sequence length plus or minus 10% of that referenced length. It is noted that novel fragments of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO9904, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide-encoding nucleotide sequence may be determined in a routine manner by aligning the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO9904, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide-encoding nucleotide sequence with other known nucleotide sequences using any of a number of well known sequence alignment programs and determining which PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO9904, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide-encoding nucleotide sequence fragment(s) are novel. All of such PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO9904, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide-encoding nucleotide sequences are contemplated herein. Also contemplated are the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO9904, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide fragments encoded by these nucleotide molecule fragments, preferably those PRO218, PRO228, PRO271, PRO273, PRO295,
PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide fragments that comprise a binding site for an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

The invention provides isolated PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, comprising an amino acid sequence having at least about 80% amino acid sequence identity, alternatively at least about 81% amino acid sequence identity, alternatively at least about 82% amino acid sequence identity, alternatively at least about 83% amino acid sequence identity, alternatively at least about 84% amino acid sequence identity, alternatively at least about 85% amino acid sequence identity, alternatively at least about 86% amino acid sequence identity, alternatively at least about 87% amino acid sequence identity, alternatively at least about 88% amino acid sequence identity, alternatively at least about 89% amino acid sequence identity, alternatively at least about 90% amino acid sequence identity, alternatively at least about 91% amino acid sequence identity, alternatively at least about 92% amino acid sequence identity, alternatively at least about 93% amino acid sequence identity, alternatively at least about 94% amino acid sequence identity, alternatively at least about 95% amino acid sequence identity, alternatively at least about 96% amino acid sequence identity, alternatively at least about 97% amino acid sequence identity, alternatively at least about 98% amino acid sequence identity and alternatively at least about 99% amino acid sequence identity to a PRO218, PRO228, PRO271, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089,
PRO19563, PRO19675, PRO20084, PRO21434, PROS0332, PRO38465 or PRO346 polypeptide having a full-length amino acid sequence as disclosed herein, an amino acid sequence lacking the signal peptide as disclosed herein, an extracellular domain of a transmembrane protein, with or without the signal peptide, as disclosed herein or any other specifically defined fragment of the full-length amino acid sequence as disclosed herein.

In a further aspect, the invention concerns an isolated PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO10948, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PROS0332, PRO38465 or PRO346 polypeptide comprising an amino acid sequence having at least about 80% amino acid sequence identity, alternatively at least about 81% amino acid sequence identity, alternatively at least about 82% amino acid sequence identity, alternatively at least about 83% amino acid sequence identity, alternatively at least about 84% amino acid sequence identity, alternatively at least about 85% amino acid sequence identity, alternatively at least about 86% amino acid sequence identity, alternatively at least about 87% amino acid sequence identity, alternatively at least about 88% amino acid sequence identity, alternatively at least about 89% amino acid sequence identity, alternatively at least about 90% amino acid sequence identity, alternatively at least about 91% amino acid sequence identity, alternatively at least about 92% amino acid sequence identity, alternatively at least about 93% amino acid sequence identity, alternatively at least about 94% amino acid sequence identity, alternatively at least about 95% amino acid sequence identity, alternatively at least about 96% amino acid sequence identity, alternatively at least about 97% amino acid sequence identity, alternatively at least about 98% amino acid sequence identity and alternatively at least about 99% amino acid sequence identity to an amino acid sequence encoded by any of the human protein cDNAs deposited with the ATCC as disclosed herein.

In one aspect, the invention concerns PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO10948, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PROS0332, PRO38465 or PRO346 variant polypeptides which are or are at least about 10 amino acids in length, alternatively are or are at least about 20, 30, 40, 50, 60, 70, 80, 90, 100, 110, 120, 130, 140, 150, 160, 170, 180, 190, 200, 210, 220, 230, 240, 250, 260, 270, 280, 290, 300, 310, 320, 330, 340, 350, 360, 370, 380, 390, 400, 410, 420, 430, 440, 450, 460, 470, 480, 490, 500, 510, 520, 530, 540, 550, 560, 570, 580, 590, 600 amino acids in length, or more. Optionally, PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99098, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PROS0332, PRO38465 or PRO346 variant polypeptides will have or have no more than one conservative amino acid substitution as compared to the native PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113,
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PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide sequence, alternatively will have or will have no more than 2, 3, 4, 5, 6, 7, 8, 9, or 10 conservative amino acid substitution as compared to the native PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide sequence.

In a specific aspect, the invention provides an isolated PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide without the N-terminal signal sequence and/or the initiating methionine and is encoded by a nucleotide sequence that encodes such an amino acid sequence as hereinbefore described. Processes for producing the same are also herein described, wherein those processes comprise culturing a host cell comprising a vector which comprises the appropriate encoding nucleic acid molecule under conditions suitable for expression of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide and recovering the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide from the cell culture.

Another aspect the invention provides an isolated PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide which is either transmembrane domain-deleted or transmembrane domain-inactivated. Processes for producing the same are also herein described, wherein those processes comprise culturing a host cell comprising a vector which comprises the appropriate encoding nucleic acid molecule under conditions suitable for expression of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016,
PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99094, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide and recovering the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO92, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99094, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide from the cell culture.

The invention provides agonists and antagonists of a native PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO92, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99094, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide as defined herein. In particular, the agonist or antagonist is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO92, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO99094, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody or a small molecule.

The invention provides a method of identifying agonists or antagonists to a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO92, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99094, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide which comprise contacting the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO92, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99094, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide with a candidate molecule and monitoring a biological activity mediated by said PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO92, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99094, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434,
PRO50332, PRO38465 or PRO346 polypeptide. Preferably, the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide is a native PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide.

The invention provides a composition of matter comprising a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, or an agonist or antagonist of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, anti-PRO38465 or anti-PRO346 antibody, in combination with a carrier. Optionally, the carrier is a pharmaceutically acceptable carrier.

The invention provides the use of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, or an agonist or antagonist thereof as hereinbefore described, or an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody, in combination with a carrier.
PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-
PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-
PRO9909, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434,
anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody, for the preparation of a medicament useful in the
treatment of a condition which is responsive to the anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273,
anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO306, anti-PRO655, anti-PRO162, anti-PRO788, anti-
PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238,
anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-
PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO5733, anti-PRO9859, anti-
PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-
PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346
antibody.

The invention provides vectors comprising DNA encoding any of the herein described polypeptides. Host
cell comprising any such vector are also provided. By way of example, the host cells may be CHO cells, E. coli,
or yeast. A process for producing any of the herein described polypeptides is further provided and comprises
culturing host cells under conditions suitable for expression of the desired polypeptide and recovering the desired
polypeptide from the cell culture.

The invention provides chimeric molecules comprising any of the herein described polypeptides fused to
a heterologous polypeptide or amino acid sequence. Example of such chimeric molecules comprise any of the
herein described polypeptides fused to an epitope tag sequence or a Fc region of an immunoglobulin.

The invention provides an antibody which binds, preferably specifically, to any of the above or below
described polypeptides. Optionally, the antibody is a monoclonal antibody, humanized antibody, antibody
fragment or single-chain antibody.

The invention provides oligonucleotide probes which may be useful for isolating genomic and cDNA
nucleotide sequences, measuring or detecting expression of an associated gene or as antisense probes, wherein
those probes may be derived from any of the above or below described nucleotide sequences. Preferred probe
lengths are described above.

The invention also provides a method of identifying a phenotype associated with a disruption of a gene
which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386,
PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238,
PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543,
PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694,
PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the
method comprising:

(a) providing a non-human transgenic animal whose genome comprises a disruption of the gene which
encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655,
PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069,
PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329,
PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089,
PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide;
(b) measuring a physiological characteristic of the non-human transgenic animal; and
(c) comparing the measured physiological characteristic with that of a gender matched wild-type animal,
wherein the physiological characteristic of the non-human transgenic animal that differs from the physiological
characteristic of the wild-type animal is identified as a phenotype resulting from the gene disruption in the non-
human transgenic animal. In one aspect, the non-human transgenic animal is a mammal. In another aspect, the
mammal is a rodent. In still another aspect, the mammal is a rat or a mouse. In one aspect, the non-human
transgenic animal is heterozygous for the disruption of a gene which encodes for a PRO218, PRO228, PRO271,
PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940,
PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130,
PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859,
PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675,
PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide. In another aspect, the phenotype
exhibited by the non-human transgenic animal as compared with gender matched wild-type littermates is at least
one of the following: a neurological disorder; a cardiovascular, endothelial or angiogenic disorder; an eye
abnormality; an immunological disorder; an oncological disorder; a bone metabolic abnormality or disorder; a lipid
metabolic disorder; or a developmental abnormality.

In yet another aspect, the neurological disorder is an increased anxiety-like response during open field
activity testing. In yet another aspect, the neurological disorder is a decreased anxiety-like response during open
field activity testing. In yet another aspect, the neurological disorder is an abnormal circadian rhythm during home-
cage activity testing. In yet another aspect, the neurological disorder is an enhanced motor coordination during
inverted screen testing. In yet another aspect, the neurological disorder is impaired motor coordination during
inverted screen testing. In yet another aspect, the neurological disorder includes depression, generalized anxiety
disorders, attention deficit disorder, sleep disorder, hyperactivity disorder, obsessive compulsive disorder,
schizophrenia, cognitive disorders, hyperalgesia and sensory disorders. Such neurological disorders include the
category defined as “anxiety disorders” which include but are not limited to: mild to moderate anxiety, anxiety
disorder due to a general medical condition, anxiety disorder not otherwise specified, generalized anxiety disorder,
panic attack, panic disorder with agoraphobia, panic disorder without agoraphobia, posttraumatic stress disorder,
social phobia, social anxiety, autism, specific phobia, substance-induced anxiety disorder, acute alcohol
withdrawal, obsessive compulsive disorder, agoraphobia, monopolar disorders, bipolar disorder I or II, bipolar
disorder not otherwise specified, cyclothymic disorder, depressive disorder, major depressive disorder, mood
disorder, substance-induced mood disorder, enhancement of cognitive function, loss of cognitive function
associated with but not limited to Alzheimer’s disease, stroke, or traumatic injury to the brain, seizures resulting
from disease or injury including but not limited to epilepsy, learning disorders/disabilities, cerebral palsy. In
addition, anxiety disorders may apply to personality disorders including but not limited to the following types:
paranoid, antisocial, avoidant behavior, borderline personality disorders, dependent, histrionic, narcissistic,
obssessive-compulsive, schizoid, and schizotypal.

In another aspect, the eye abnormality is a retinal abnormality. In still another aspect, the eye abnormality
is consistent with vision problems or blindness. In yet another aspect, the retinal abnormality is consistent with
retinitis pigmentosa or is characterized by retinal degeneration or retinal dysplasia.

In still another aspect, the retinal abnormalities are consistent with retinal dysplasia, various retinopathies, including retinopathy of prematurity, retrolental fibrosa, neovascular glaucoma, age-related macular degeneration, diabetic macular edema, corneal neovascularization, corneal graft neovascularization, corneal graft rejection, retinal/choroidal neovascularization, neovascularization of the angle (rubecosis), ocular neovascular disease, vascular restenosis, arteriovenous malformations (AVM), menigioma, hemangioma, angiofibroma, thyroid hyperplasias (including Grave's disease), corneal and other tissue transplantation, retinal artery obstruction or occlusion; retinal degeneration causing secondary atrophy of the retinal vasculature, retinitis pigmentosa, macular dystrophies, Stargardt's disease, congenital stationary night blindness, choroideremia, gyrate atrophy, Leber's congenital amaurosis, retinoschisis disorders, Wagner's syndrome, Usher syndromes, Zellweger syndrome, Saldino-Mainzer syndrome, Senior-Loken syndrome, Bardet-Biedl syndrome, Alport's syndrome, Alstrom's syndrome, Cockayne's syndrome, dysplasia spondyloepiphysaria congenita, Flynn-Aird syndrome, Friedrich ataxia, Hallgren syndrome, Marshall syndrome, Albers-Schonberg disease, Refsum's disease, Kearn's-Sayre syndrome, Waardenburg's syndrome, Alagille syndrome, myotonic dystrophy, olivopontocerebellar atrophy, Pierre-Marie dundrude, Stickler syndrome, carotidocemia, cystinosis, Wolfman syndrome, Bassen-Kornzweig syndrome, abetalipoproteinemia, incontinentia pigmenti, Batten's disease, mucopolysaccharidoses, homocystinuria, or mannosidosis.

In still another aspect, the eye abnormality is a cataract. In still yet another aspect, the cataract is a systemic disease such as human Down's syndrome, Hallman-Streiff syndrome, Lowe syndrome, galactosemia, Marfan syndrome, Trisomy 13-15, Alport syndrome, myotonic dystrophy, Fabry disease, hypoparathyroidism or Conradi syndrome.

In still another aspect, the developmental abnormality comprises embryonic lethality or reduced viability.

In still yet another aspect, the cardiovascular, endothelial or angiogenic disorders are arterial diseases, such as diabetes mellitus; papilledema; optic atrophy; atherosclerosis; angina; myocardial infarctions such as acute myocardial infarctions, cardiac hypertrophy, and heart failure such as congestive heart failure; hypertension; inflammatory vasculitides; Reynaud's disease and Reynaud's phenomenon; aneurysms and arterial restenosis; venous and lymphatic disorders such as thrombophlebitis, lymphangitis, and lymphedema; peripheral vascular disease; cancer such as vascular tumors, e.g., hemangioma (capillary and cavernous), glomus tumors, telangiectasia, bacillary angiomatosis, hemangioendothelioma, angiosarcoma, haemangioepicytoma, Kaposi's sarcoma, lymphangioma, and lymphangiosarcoma; tumor angiogenesis; trauma such as wounds, burns, and other injured tissue, implant fixation, scarring; ischemia reperfusion injury; rheumatoid arthritis; cerebrovascular disease; renal diseases such as acute renal failure, or osteoporosis.

In still another aspect, the immunological disorders are consistent with systemic lupus erythematosus; rheumatoid arthritis; juvenile chronic arthritis; spondyloarthropathies; systemic sclerosis (scleroderma); idiopathic inflammatory myopathies (dermatomyositis, polymyositis); SJögren's syndrome; systemic vasculitis; sarcoidosis; autoimmune hemolytic anemia (immune pancytopenia, paroxysmal nocturnal hemoglobinuria); autoimmune thrombocytopenia (idiopathic thrombocytopenic purpura, immune-mediated thrombocytopenia); thyroiditis (Grave's disease, Hashimoto's thyroiditis, juvenile lymphocytic thyroiditis, atrophic thyroiditis); diabetes mellitus; immune-mediated renal disease (glomerulonephritis, tubulointerstitial nephritis); demyelinating diseases of the
central and peripheral nervous systems such as multiple sclerosis, idiopathic demyelinating polyneuropathy or Guillain-Barré syndrome, and chronic inflammatory demyelinating polyneuropathy; hepatobiliary diseases such as infectious hepatitis (hepatitis A, B, C, D, E and other non-hepatotropic viruses), autoimmune chronic active hepatitis, primary biliary cirrhosis, granulomatous hepatitis, and sclerosing cholangitis; inflammatory bowel disease (ulcerative colitis: Crohn's disease); gluten-sensitive enteropathy, and Whipple's disease; autoimmune or immune-mediated skin diseases including bullous skin diseases, erythema multiforme and contact dermatitis, psoriasis; allergic diseases such as asthma, allergic rhinitis, atopic dermatitis, food hypersensitivity and urticaria; immunologic diseases of the lung such as eosinophilic pneumonia, idiopathic pulmonary fibrosis and hypersensitivity pneumonitis; or transplantation associated diseases including graft rejection and graft-versus-host disease.

In still another aspect, the bone metabolic abnormality or disorder is arthritis, osteoporosis, osteopenia or osteopetrosis.

In another aspect, the non-human transgenic animal exhibits at least one of the following physiological characteristics compared with gender matched wild-type littermates: increased anxiety-like response during open field testing; hyperactivity during open field testing; decreased anxiety during open field testing; decreased locomotor activity during open field testing; abnormal circadian rhythm during home-cage activity testing (low activity during the light phase; altered sleep/wake cycle); abnormal circadian rhythm during home-cage activity testing including increased ambulatory counts; hypoactivity with no circadian rhythm; abnormal circadian rhythm during home-cage activity testing including increased ambulatory counts; decreased rearing; increased sensitivity to stress induced hyperthermia (increased anxiety); impaired motor coordination during inverted screen testing; head tilt and retropulsion; increased prepulse inhibition response indicating enhanced sensorimotor gating/attention; decreased startle response during prepulse inhibition testing; no startle response indicating deafness or impaired hearing; decreased prepulse inhibition with impaired sensorimotor gating/attention; increased latency to respond in hot plate testing; decreased latency to respond in hot plate testing; ophthalmological abnormalities; impaired vision; white deposits of optic disc region; ocular infection and neutrophilia; bilateral optic disc lesion; decreased tear production; decreased heart rate; increased mean systolic blood pressure; decreased mean systolic blood pressure; increased mean fasting serum glucose levels; decreased mean serum glucose levels; increased mean serum cholesterol levels; decreased mean serum cholesterol levels; increased mean serum triglyceride levels; decreased mean serum triglyceride levels; impaired glucose tolerance; increased mean serum albumin, alanine amino transferase and phosphorus levels; increased mean serum alkaline phosphatase levels; urinary nitrites present; increased total white blood cell (WBC) count; decreased total white blood cell (WBC) count and absolute neutrophil count; increased mean absolute neutrophil count; increased mean absolute lymphocyte count; increased mean platelet count; increased mean red cell distribution width; decreased mean platelet count; reduced percentage of CD4 spleen thymocytes; decreased percentages of CD4 cells in the periphery resulting in increased percentages of B cells in lymph organs; CD4 cells exhibit a more activated/memory phenotype (CD62Llow, CD44hi); developmental defect in CD4+ cells; decreased percentages of CD4 cells and increased percentages of B cells in blood; decreased percentages of CD4 cells and increased percentages of B cells in tissues; increase in percentages of B cells in Peyer’s patches (; decreased germinal center, isotype-switched B cells in Peyer’s patches (CD38low; IgM negative); decreased CD23 intensity in spleen; increased mean percentages of B220 Med/CD23-
cells and B220+/CD11b-Low/CD23- cells in peritoneal lavage; increased mean percentages of B cells in peripheral blood; decreased CD4 and CD8 T cells and increased B cells; increase in peritoneal B cells; reduction in CD11b-Hi cells in peritoneal cavity; decreased mean CD4 to CD8 ratio in spleen; decreased CD8 cells; decreased mean percentages of B220+/CD23+ cells and B220+/CD11bLow/CD23- cells in peritoneal lavage; increased mean serum IgG1 response to ovalbumin challenge; increased mean serum IgG2a response to ovalbumin challenge; increased mean serum IL-6 response to LPS challenge; increased mean serum TNF alpha response to LPS challenge; increased mean serum MCP-1 response to LPS challenge; increased mean serum IgM level; increased mean serum IgA; increase mean serum IgG1; increased mean serum IgG2a; increased mean serum IgG2b; decreased mean serum IgG1 response to ovalbumin challenge; decreased mean serum IgG2a response to ovalbumin challenge; failure in ovalbumin response; decreased mean serum IgA level; decreased mean serum IgG2a level; decreased skin fibroblast proliferation rate; increased mean percent of total body fat and total fat mass; increased mean body weight; increased mean body length; increased total tissue mass (TTM); increased bone mineral density (BMD); increase in bone mineral content (BMC); increased mean femoral midshaft cortical thickness; decreased mean percent of total body fat and total fat mass; decreased mean body weight; decreased mean body length; decreased mean body weight and length in heterozygotes; decreased total tissue mass (TTM); decreased lean body mass (LBM); decreased femoral bone mineral density (BMD); decreased vertebral bone mineral density (BMD); decreased bone mineral density (BMD) in total body; decreased bone mineral content (BMC); decreased bone mineral density index; decreased volumetric bone mineral density (vBMD); decreased mean femoral midshaft cortical thickness; decreased mean femoral midshaft cross-sectional area; decreased mean vertebral trabecular bone volume, number and connectivity density; osteoporosis; osteopenia; moderate kidney hydronephrosis; hydrocephalus; enlarged liver; induced in activated T cells; induced in activated NK cells and dendritic cells; myeloid B cell expression; hyperplasia of sebaceous glands and multifocal hyperplasia of the epidermis (acanthosis and hyperkeratosis); moderate dermatitis; increased extramedullary hematopoiesis in liver and spleen; myeloid hyperplasia of the bone marrow; encephalitis due to Group B streptococcus; meningitis due to E. Coli infection; lymphocytic infiltrates in salivary glands, pancreas and lungs; poor breeders requiring foster mothers; decreased litter size; homozygous mice were small and dehydrated; vacuolar degeneration of testes resulting in decreased sperm production and infertility; defective spermatogenesis in the testes; hypospermia and defective spermatogenesis in the epididymus; male infertility; decreased testes weight; growth retardation; small mice and failure to thrive; reduced viability; reduced viability with situs inversus; and homozygous embryonic lethality.

The invention also provides an isolated cell derived from a non-human transgenic animal whose genome comprises a disruption of the gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide. In one aspect, the isolated cell is a murine cell. In yet another aspect, the murine cell is an embryonic stem cell. In still another aspect, the isolated cell is derived from a non-human transgenic animal which exhibits at least one of the following phenotypes compared with gender matched wild-type littermates: a neurological disorder; a cardiovascular, endothelial or angiogenic disorder; an eye abnormality; an
immunological disorder; an oncological disorder; a bone metabolic abnormality or disorder; a lipid metabolic disorder; or a developmental abnormality. The invention also provides a method of identifying an agent that modulates a phenotype associated with a disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising:

(a) providing a non-human transgenic animal whose genome comprises a disruption of the gene which encodes for the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide;

(b) measuring a physiological characteristic of the non-human transgenic animal of (a);

(c) comparing the measured physiological characteristic of (b) with that of a gender matched wild-type animal, wherein the physiological characteristic of the non-human transgenic animal that differs from the physiological characteristic of the wild-type animal is identified as a phenotype resulting from the gene disruption in the non-human transgenic animal;

(d) administering a test agent to the non-human transgenic animal of (a); and

(e) determining whether the test agent modulates the identified phenotype associated with gene disruption in the non-human transgenic animal.

In one aspect, the phenotype associated with the gene disruption comprises a neurological disorder; a cardiovascular, endothelial or angiogenic disorder; an eye abnormality; an immunological disorder; an oncological disorder; a bone metabolic abnormality or disorder; a lipid metabolic disorder; or a developmental abnormality.

In yet another aspect, the neurological disorder is an increased anxiety-like response during open field activity testing. In yet another aspect, the neurological disorder is a decreased anxiety-like response during open field activity testing. In yet another aspect, the neurological disorder is an abnormal circadian rhythm during home-cage activity testing. In yet another aspect, the neurological disorder is an enhanced motor coordination during inverted screen testing. In yet another aspect, the neurological disorder is impaired motor coordination during inverted screen testing. In yet another aspect, the neurological disorder includes depression, generalized anxiety disorders, attention deficit disorder, sleep disorder, hyperactivity disorder, obsessive compulsive disorder, schizophrenia, cognitive disorders, hyperalgesia and sensory disorders. Such neurological disorders include the category defined as “anxiety disorders” which include but are not limited to: mild to moderate anxiety, anxiety disorder due to a general medical condition, anxiety disorder not otherwise specified, generalized anxiety disorder, panic attack, panic disorder with agoraphobia, panic disorder without agoraphobia, posttraumatic stress disorder, social phobia, social anxiety, autism, specific phobia, substance-induced anxiety disorder, acute alcohol withdrawal, obsessive compulsive disorder, agoraphobia, monopolar disorders, bipolar disorder I or II, bipolar disorder not otherwise specified, cyclothymic disorder, depressive disorder, major depressive disorder, mood
disorder, substance-induced mood disorder, enhancement of cognitive function, loss of cognitive function associated with but not limited to Alzheimer's disease, stroke, or traumatic injury to the brain, seizures resulting from disease or injury including but not limited to epilepsy, learning disorders/disabilities, cerebral palsy. In addition, anxiety disorders may apply to personality disorders including but not limited to the following types: paranoid, antisocial, avoidant behavior, borderline personality disorders, dependent, histrionic, narcissistic, obsessive-compulsive, schizoid, and schizotypal.

In yet another aspect, the eye abnormality is a retinal abnormality. In still another aspect, the eye abnormality is consistent with vision problems or blindness. In yet another aspect, the retinal abnormality is consistent with retinitis pigmentosa or is characterized by retinal degeneration or retinal dysplasia.

In still another aspect, the retinal abnormalities are consistent with retinal dysplasia, various retinopathies, including retinopathy of prematurity, retrolental fibroplasia, neovascular glaucoma, age-related macular degeneration, diabetic macular edema, corneal neovascularization, corneal graft neovascularization, corneal graft rejection, retinal/choroidal neovascularization, neovascularization of the angle (rubeosis), ocular neovascular disease, vascular restenosis, arteriovenous malformations (AVM), meningioma, hemangioma, angiofibroma, thyroid hyperplasias (including Grave's disease), corneal and other tissue transplantation, retinal artery obstruction or occlusion; retinal degeneration causing secondary atrophy of the retinal vasculature, retinitis pigmentosa, macular dystrophies, Stargardt's disease, congenital stationary night blindness, chorioideremia, gyrate atrophy, Leber's congenital amaurosis, retinoschisis disorders, Wagner's syndrome, Usher syndromes, Zellweger syndrome, Saldino-Mainzer syndrome, Senior-Loken syndrome, Bardet-Biedl syndrome, Alport's syndrome, Alström's syndrome, Cockayne's syndrome, dysplasia spondyloepiphysaria congenita, Flynn-Aird syndrome, Friedrich ataxia, Hallgren syndrome, Marshall syndrome, Albers-Schönberg disease, Refsum's disease, Kearns-Sayre syndrome, Waardenburg's syndrome, Alagille syndrome, myotonic dystrophy, olivopontocerebellar atrophy, Pierre-Marie dunsdrome, Stickler syndrome, carotinemia, cystinosis, Wolfram syndrome, Bassen-Kornzweig syndrome, abetalipoproteinemia, incontinentia pigmenti, Batten's disease, mucopolysaccharidoses, homocystinuria, or mannosidosis.

In still another aspect, the eye abnormality is a cataract. In still yet another aspect, the cataract is a systemic disease such as human Down's syndrome, Hallerman-Streiff syndrome, Lowe syndrome, galactosemia, Marfan syndrome, Trisomy 13-15, Alport syndrome, myotonic dystrophy, Fabry disease, hypoparathyroidism, or Conradi syndrome.

In still another aspect, the developmental abnormality comprises embryonic lethality or reduced viability.

In still another aspect, the cardiovascular, endothelial or angiogenic disorders are arterial diseases, such as diabetes mellitus; papilledema; optic atrophy; atherosclerosis; angina; myocardial infarctions such as acute myocardial infarctions, cardiac hypertrophy, and heart failure such as congestive heart failure; hypertension; inflammatory vasculitides; Reynaud's disease and Reynaud's phenomenon; aneurysms and arterial restenosis; venous and lymphatic disorders such as thrombophlebitis, lymphangitis, and lymphedema; peripheral vascular disease; cancer such as vascular tumors, e.g., hemangioma (capillary and cavernous), glomus tumors, telangiectasia, bacillary angiomatosis, hemangioendothelioma, angiosarcoma, haemangiopericytoma, Kaposi's sarcoma, lymphangioma, and lymphangiosarcoma; tumor angiogenesis; trauma such as wounds, burns, and other injured tissue, implant fixation, scarring; ischemia reperfusion injury; rheumatoid arthritis; cerebrovascular disease;
renal diseases such as acute renal failure, or osteoporosis.

In still another aspect, the immunological disorders are consistent with systemic lupus erythematosus; rheumatoid arthritis; juvenile chronic arthritis; spondyloarthopathies; systemic sclerosis (scleroderma); idiopathic inflammatory myopathies (dermatomyositis, polymyositis); Sjögren's syndrome; systemic vasculitis; sarcoidosis; autoimmune hemolytic anemia (immune pancytopenia, paroxysmal nocturnal hemoglobinuria); autoimmune thrombocytopenia (idiopathic thrombocytopenic purpura, immune-mediated thrombocytopenia); thyroiditis (Grave's disease, Hashimoto's thyroiditis, juvenile lymphocytic thyroiditis, atrophic thyroiditis); diabetes mellitus; immune-mediated renal disease (glomerulonephritis, tubulointerstitial nephritis); demyelinating diseases of the central and peripheral nervous systems such as multiple sclerosis, idiopathic demyelinating polyneuropathy or Guillain-Barré syndrome, and chronic inflammatory demyelinating polyneuropathy; hepatobiliary diseases such as infectious hepatitis (hepatitis A, B, C, D, E and other non-hepatotropic viruses), autoimmune chronic active hepatitis, primary biliary cirrhosis, granulomatous hepatitis, and sclerosing cholangitis; inflammatory bowel disease (ulcerative colitis; Crohn's disease); gluten-sensitive enteropathy, and Whipple's disease; autoimmune or immune-mediated skin diseases including bullous skin diseases, erythema multiforme and contact dermatitis, psoriasis; allergic diseases such as asthma, allergic rhinitis, atopic dermatitis, food hypersensitivity and urticaria; immunologic diseases of the lung such as eosinophilic pneumonia, idiopathic pulmonary fibrosis and hypersensitivity pneumonitis; or transplantation associated diseases including graft rejection and graft-versus-host disease.

In yet another aspect, the bone metabolic abnormality or disorder is arthritis, osteoporosis, osteopenia or osteopetrosis.

In another aspect, the non-human transgenic animal exhibits at least one of the following physiological characteristics compared with gender matched wild-type littermates: increased anxiety-like response during open field testing; hyperactivity during open field testing; decreased anxiety during open field testing; decreased locomotor activity during open field testing; abnormal circadian rhythm during home-cage activity testing (low activity during the light phase; altered sleep/wake cycle); abnormal circadian rhythm during home-cage activity testing including decreased ambulatory counts; hypoactivity with no circadian rhythm; abnormal circadian rhythm during home-cage activity testing including increased ambulatory counts; decreased rearing; increased sensitivity to stress induced hyperthermia (increased anxiety); impaired motor coordination during inverted screen testing; head tilt and retropulsion; increased prepulse inhibition response indicating enhanced sensorimotor gating/attention; decreased startle response during prepulse inhibition testing; no startle response indicating deafness or impaired hearing; decreased prepulse inhibition with impaired sensorimotor gating/attention; increased latency to respond in hot plate testing; decreased latency to respond in hot plate testing; ophthalmological abnormalities; impaired vision; white deposits of optic disc region; ocular infection and neutrophilia; bilateral optic disc lesion; decreased tear production; decreased heart rate; increased mean systolic blood pressure; decreased mean systolic blood pressure; increased mean fasting serum glucose levels; decreased mean serum glucose levels; increased mean serum cholesterol levels; decreased mean serum cholesterol levels; increased mean serum triglyceride levels; decreased mean serum triglyceride levels; impaired glucose tolerance; increased mean serum albumin, alanine amino transferase and phosphorous levels; increased mean serum alkaline phosphatase levels; urinary nitrites present; increased total white blood cell (WBC) count; decreased total white blood cell (WBC) count and absolute
neutrophil count; increased mean absolute neutrophil count; increased mean absolute lymphocyte count; increased mean platelet count; increased mean red cell distribution width; decreased mean platelet count; reduced percentage of CD4 spleen thymocytes; decreased percentages of CD4 cells in the periphery resulting in increased percentages of B cells in lymph organs; CD4 cells exhibit a more activated/memory phenotype (CD62Llow, CD44hi); developmental defect in CD4+ cells; decreased percentages of CD4 cells and increased percentages of B cells in blood; decreased percentages of CD4 cells and increased percentages of B cells in tissues; increase in percentages of B cells in Peyer’s patches; decreased germinal center, isotype-switched B cells in Peyer’s patches (CD38low, IgM negative); decreased CD23 intensity in spleen; increased mean percentages of B220 Med/CD23-cells and B220+/CD11b-Low/CD23- cells in peritoneal lavage; increased mean percentages of B cells in peripheral blood; decreased C4D and CD8 T cells and increased B cells; increase in peritoneal B cells; reduction in CD11b-Hi cells in peritoneal cavity; decreased mean CD4 to CD8 ratio in spleen; decreased CD8 cells; decreased mean percentages of B220+/CD23+ cells and B220+/CD11bLow/CD23- cells in peritoneal lavage; increased mean serum IgG1 response to ovalbumin challenge; increased mean serum IgG2a response to ovalbumin challenge; increased mean serum IL-6 response to LPS challenge; increased mean serum TNF alpha response to LPS challenge; increased mean serum MCP-1 response to LPS challenge; increased mean serum IgM level; increased mean serum IgA; increased mean serum IgG1; increased mean serum IgG2a; increased mean serum IgG2b; decreased mean serum IgG1 response to ovalbumin challenge; decreased mean serum IgG2a response to ovalbumin challenge; failure in ovalbumin response; decreased mean serum IgA level; increased mean serum IgG2a level; skin fibroblast proliferation rate; increased mean percent of total body fat and total fat mass; increased mean body weight; increased mean body length; increased total tissue mass (TTM); increased bone mineral density (BMD); increase in bone mineral content (BMC); increased mean femoral midshaft cortical thickness; decreased mean percent of total body fat and total fat mass; decreased mean body weight; decreased mean body length; decreased mean body weight and length in heterozygotes; decreased total tissue mass (TTM); decreased lean body mass (LBM); decreased femoral bone mineral density (BMD); decreased vertebral bone mineral density (BMD); decreased bone mineral density (BMD) in total body; decreased bone mineral content (BMC); decreased bone mineral density index; decreased volumetric bone mineral density (vBMD); decreased mean femoral midshaft cortical thickness; decreased mean femoral midshaft cross-sectional area; decreased mean vertebral trabecular bone volume, number and connectivity density; osteopenia; osteoporosis; moderate kidney hydropnephrosis; hydrocephalus; enlarged liver; induced in activated T cells; induced in activated NK cells and dendritic cells; myeloid B cell expression; hyperplasia of sebaceous glands and multifocal hyperplasia of the epidermis (acanthosis and hyperkeratosis); moderate dermatitis; increased extramedullary hematopoiesis in liver and spleen; myeloid hyperplasia of the bone marrow; encephalitis due to Group B streptococcius; meningitis due to E. Coli infection; lymphocytic infiltrates in salivary glands, pancreas and lungs; poor breeders requiring foster mothers; decreased litter size; homozygous mice were small and dehydrated; vascular degeneration of testes resulting in decreased sperm production and infertility; defective spermatogenesis in the testes; hypospermia and defective spermatozoa in the epididymus; male infertility; decreased testes weight; growth retardation; small mice and failure to thrive; reduced viability; reduced viability with situs inversus; and homozygous embryonic lethality.

The invention also provides an agent which modulates the phenotype associated with gene disruption. In one aspect, the agent is an agonist or antagonist of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302,
PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide. In yet another aspect, the agonist agent is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody. In still another aspect, the antagonist agent is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

The invention also provides a method of identifying an agent that modulates a physiological characteristic associated with a disruption of the gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising:

(a) providing a non-human transgenic animal whose genome comprises a disruption of the gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide;

(b) measuring a physiological characteristic exhibited by the non-human transgenic animal of (a);

(c) comparing the measured physiological characteristic of (b) with that of a gender matched wild-type animal, wherein the physiological characteristic exhibited by the non-human transgenic animal that differs from the physiological characteristic exhibited by the wild-type animal is identified as a physiological characteristic associated with gene disruption;

(d) administering a test agent to the non-human transgenic animal of (a); and

(e) determining whether the physiological characteristic associated with gene disruption is modulated.
In one aspect, the non-human transgenic animal exhibits at least one of the following physiological characteristics compared with gender matched wild-type littermates:

In another aspect, the non-human transgenic animal exhibits at least one of the following physiological characteristics compared with gender matched wild-type littermates: increased anxiety-like response during open field testing; hyperactivity during open field testing; decreased anxiety during open field testing; decreased locomotor activity during open field testing; abnormal circadian rhythm during home-cage activity testing (low activity during the light phase; altered sleep/wake cycle); abnormal circadian rhythm during home-cage activity testing including decreased ambulatory counts; hypoactivity with no circadian rhythm; abnormal circadian rhythm during home-cage activity testing including increased ambulatory counts; decreased rearing; increased sensitivity to stress induced hyperthermia (increased anxiety); impaired motor coordination during inverted screen testing; head tilt and retropulsion; increased prepulse inhibition response indicating enhanced sensorimotor gating/attention; decreased startle response during prepulse inhibition testing; no startle response indicating deafness or impaired hearing; decreased prepulse inhibition with impaired sensorimotor gating/attention; increased latency to respond in hot plate testing; decreased latency to respond in hot plate testing; ophthalmological abnormalities; impaired vision; white deposits of optic disc region; ocular infection and neutrophilia; bilateral optic disc lesion; decreased tear production; decreased heart rate; increased mean systolic blood pressure; decreased mean systolic blood pressure; increased mean fasting serum glucose levels; decreased mean serum glucose levels; increased mean serum cholesterol levels; decreased serum cholesterol levels; increased mean serum triglyceride levels; decreased mean serum triglyceride levels; impaired glucose tolerance; increased mean serum albumin, alanine amino transferase and phosphorus levels; increased mean serum alkaline phosphatase levels; urinary nitrites present; increased total white blood cell (WBC) count; decreased total white blood cell (WBC) count and absolute neutrophil count; increased mean absolute neutrophil count; increased mean absolute lymphocyte count; increased mean platelet count; increased mean red cell distribution width; decreased mean platelet count; reduced percentage of CD4 spleen thymocytes; decreased percentages of CD4 cells in the periphery resulting in increased percentages of B cells in lymph organs; CD4 cells exhibit a more activated/memory phenotype (CD62Llow, CD44hi); developmental defect in CD4+ cells; decreased percentages of CD4 cells and increased percentages of B cells in blood; decreased percentages of CD4 cells and increased percentages of B cells in tissues; increase in percentages of B cells in Peyer's patches; decreased germinal center, isotype-switched B cells in Peyer's patches (CD38low; IgM negative); decreased CD23 intensity in spleen; increased mean percentages of B220 Med/CD23-cells and B220+/CD11b-Low/CD23- cells in peritoneal lavage; increased mean percentages of B cells in peripheral blood; decreased CD4 and CD8 T cells and increased B cells; increase in peritoneal B cells; reduction in CD11b-Hi cells in peritoneal cavity; decreased mean CD4 to CD8 ratio in spleen; decreased CD8 cells; decreased mean percentages of B220+/CD23+ cells and B220+/CD11bLow/CD23- cells in peritoneal lavage; increased mean serum IgG1 response to ovalbumin challenge; increased mean serum IgG2a response to ovalbumin challenge; increased mean serum IL-6 response to LPS challenge; increased mean serum TNF alpha response to LPS challenge; increased mean serum MCP-1 response to LPS challenge; increased mean serum IgM level; increased mean serum IgA; increase mean serum IgG1; increased mean serum IgG2a; increased mean serum IgG2b; decreased mean serum IgG1 response to ovalbumin challenge; decreased mean serum IgG2a response to ovalbumin challenge; failure in ovalbumin response; decreased mean serum IgA level; decreased mean serum IgG2a level; decreased skin
fibroblast proliferation rate; increased mean percent of total body fat and total fat mass; increased mean body weight; increased mean body length; increased total tissue mass (TTM); increased bone mineral density (BMD); increase in bone mineral content (BMC); increased mean femoral midshaft cortical thickness; decreased mean percent of total body fat and total fat mass; decreased mean body weight; decreased mean body length; decreased mean body weight and length in heterozygotes; decreased total tissue mass (TTM); decreased lean body mass (LBM); decreased femoral bone mineral density (BMD); decreased vertebral bone mineral density (BMD); decreased bone mineral density (BMD) in total body; decreased bone mineral content (BMC); decreased bone mineral density index; decreased volumetric bone mineral density (vBMD); decreased mean femoral midshaft cortical thickness; decreased mean femoral midshaft cross-sectional area; decreased mean vertebral trabecular bone volume, number and connectivity density; osteopetrosis; osteoporosis; moderate kidney hydronephrosis; hydrocephalus; enlarged liver; induced in activated T cells; induced in activated NK cells and dendritic cells; myeloid B cell expression; hyperplasia of sebaceous glands and multifocal hyperplasia of the epidermis (acanthosis and hyperkeratosis); moderate dermatitis; increased extramedullary hematopoiesis in liver and spleen; myeloid hyperplasia of the bone marrow; encephalitis due to Group B streptococcus; meningitis due to E. Coli infection; lymphocytic infiltrates in salivary glands, pancreas and lungs; poor breeders requiring foster mothers; decreased litter size; homozygous mice were small and dehydrated; vacuolar degeneration of testes resulting in decreased sperm production and infertility; defective spermatogenesis in the testes; hyposperma and defective spermatozoa in the epididymus; male infertility; decreased testes weight; growth retardation; small mice and failure to thrive; reduced viability; reduced viability with situs inversus; and homozygous embryonic lethality.

The invention also provides an agent that modulates a physiological characteristic which is associated with gene disruption. In one aspect, the agent is an agonist or antagonist of the phenotype associated with a disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO10023, PRO1546, PRO1966, PRO21434, PRO25332, PRO38465 or PRO346 polypeptide. In yet another aspect, the agent is an agonist or antagonist of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO10023, PRO1546, PRO1966, PRO21434, PRO25332, PRO38465 or PRO346 polypeptide. In yet another aspect, the agonist agent is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO99048, anti-PRO2694, anti-PRO16089, anti-PRO19663, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO5332, anti-PRO38465 or anti-PRO346 polypeptide. In yet another aspect, the antagonist agent is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO99048, anti-PRO2694, anti-PRO16089, anti-PRO19663, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO5332, anti-PRO38465 or anti-PRO346 polypeptide. In still another aspect, the antagonist
agent is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1131, anti-PRO7195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO99048, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

The invention also provides a method of identifying an agent which modulates a behavior associated with a disruption of the gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1131, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99048, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising:

(a) providing a non-human transgenic animal whose genome comprises a disruption of the gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1131, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99048, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide;

(b) observing the behavior exhibited by the non-human transgenic animal of (a);

(c) comparing the observed behavior of (b) with that of a gender matched wild-type animal, wherein the observed behavior exhibited by the non-human transgenic animal that differs from the observed behavior exhibited by the wild-type animal is identified as a behavior associated with gene disruption;

(d) administering a test agent to the non-human transgenic animal of (a); and

(e) determining whether the agent modulates the behavior associated with gene disruption.

In one aspect, the observed behavior is an increased anxiety-like response during open field activity testing. In yet another aspect, the observed behavior is a decreased anxiety-like response during open field activity testing. In yet another aspect, the observed behavior is an abnormal circadian rhythm during home-cage activity testing. In yet another aspect, the observed behavior is an enhanced motor coordination during inverted screen testing. In yet another aspect, the observed behavior is impaired motor coordination during inverted screen testing. In yet another aspect, the observed behavior includes depression, generalized anxiety disorders, attention deficit disorder, sleep disorder, hyperactivity disorder, obsessive compulsive disorder, schizophrenia, cognitive disorders, hyperalgesia and sensory disorders. Such disorders include the category defined as “anxiety disorders” which include but are not limited to: mild to moderate anxiety, anxiety disorder due to a general medical condition, anxiety disorder not otherwise specified, generalized anxiety disorder, panic attack, panic disorder with agoraphobia, panic disorder without agoraphobia, posttraumatic stress disorder, social phobia, social anxiety, autism, specific phobia, substance-induced anxiety disorder, acute alcohol withdrawal, obsessive compulsive disorder, agoraphobia, monopolar disorders, bipolar disorder I or II, bipolar disorder not otherwise specified,
cyclothymic disorder, depressive disorder, major depressive disorder, mood disorder, substance-induced mood disorder, enhancement of cognitive function, loss of cognitive function associated with but not limited to Alzheimer’s disease, stroke, or traumatic injury to the brain, seizures resulting from disease or injury including but not limited to epilepsy, learning disorders/disabilities, cerebral palsy. In addition, anxiety disorders may apply to personality disorders including but not limited to the following types: paranoid, antisocial, avoidant behavior, borderline personality disorders, dependent, histrionic, narcissistic, obsessive-compulsive, schizoid, and schizotypal.

The invention also provides an agent that modulates a behavior which is associated with gene disruption. In one aspect, the agent is an agonist or antagonist of the phenotype associated with a disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide. In yet another aspect, the agent is an agonist or antagonist of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide.

In yet another aspect, the agent is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody. In still another aspect, the antagonist agent is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

The invention also provides a method of identifying an agent that ameliorates or modulates a neurological disorder; a cardiovascular, endothelial or angiogenic disorder; an eye abnormality; an immunological disorder; an oncological disorder; a bone metabolic abnormality or disorder; a lipid metabolic disorder; or a developmental abnormality associated with a disruption in the gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195,
PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising:

(a) providing a non-human transgenic animal whose genome comprises a disruption of the gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide;

(b) administering a test agent to said non-human transgenic animal; and

(c) determining whether the test agent ameliorates or modulates the neurological disorder; cardiovascular, endothelial or angiogenic disorder; eye abnormality; immunological disorder; oncological disorder; bone metabolic abnormality or disorder; lipid metabolic disorder; or developmental abnormality associated with the gene disruption in the non-human transgenic animal.

In yet another aspect, the neurological disorder is an increased anxiety-like response during open field activity testing. In yet another aspect, the neurological disorder is a decreased anxiety-like response during open field activity testing. In yet another aspect, the neurological disorder is an abnormal circadian rhythm during home-cage activity testing. In yet another aspect, the neurological disorder is an enhanced motor coordination during inverted screen testing. In yet another aspect, the neurological disorder is impaired motor coordination during inverted screen testing. In yet another aspect, the neurological disorder includes depression, generalized anxiety disorders, attention deficit disorder, sleep disorder, hyperactivity disorder, obsessive compulsive disorder, schizophrenia, cognitive disorders, hyperalgesia and sensory disorders. Such neurological disorders include the category defined as “anxiety disorders” which include but are not limited to: mild to moderate anxiety, anxiety disorder due to a general medical condition, anxiety disorder not otherwise specified, generalized anxiety disorder, panic attack, panic disorder with agoraphobia, panic disorder without agoraphobia, posttraumatic stress disorder, social phobia, social anxiety, autism, specific phobia, substance-induced anxiety disorder, acute alcohol withdrawal, obsessive compulsive disorder, agoraphobia, monopolar disorders, bipolar disorder I or II, bipolar disorder not otherwise specified, cyclothymic disorder, depressive disorder, major depressive disorder, mood disorder, substance-induced mood disorder, enhancement of cognitive function, loss of cognitive function associated with but not limited to Alzheimer’s disease, stroke, or traumatic injury to the brain, seizures resulting from disease or injury including but not limited to epilepsy, learning disorders/disabilities, cerebral palsy. In addition, anxiety disorders may apply to personality disorders including but not limited to the following types: paranoid, antisocial, avoidant behavior, borderline personality disorders, dependent, histrionic, narcissistic, obsessive-compulsive, schizoid, and schizotypal.

In another aspect, the eye abnormality is a retinal abnormality. In still another aspect, the eye abnormality is consistent with vision problems or blindness. In yet another aspect, the retinal abnormality is consistent with retinitis pigmentosa or is characterized by retinal degeneration or retinal dysplasia.

In still another aspect, the retinal abnormalities the retinal abnormalities are consistent with retinal dysplasia, various retinopathies, including retinopathy of prematurity, retrolental fibroplasia, neovascular glaucoma,

In still another aspect, the eye abnormality is a cataract. In still yet another aspect, the cataract is a systemic disease such as human Down's syndrome, Hallerman-Streiff syndrome, Lowe syndrome, galactosemia, Marfan syndrome, Trisomy 13-15, Alport syndrome, myotonic dystrophy, Fabry disease, hypoparathyroidism, or Conradi syndrome.

In still another aspect, the developmental abnormality comprises embryonic lethality or reduced viability.

In yet another aspect, the cardiovascular, endothelial or angiogenic disorders are arterial diseases, such as diabetes mellitus; papilledema; optic atrophy; atherosclerosis; angina; myocardial infarctions such as acute myocardial infarctions, cardiac hypertrophy, and heart failure such as congestive heart failure; hypertension; inflammatory vasculitides; Reynaud's disease and Reynaud's phenomenon; aneurysms and arterial restenosis; venous and lymphatic disorders such as thrombophlebitis, lymphangitis, and lymphedema; peripheral vascular disease; cancer such as vascular tumors, e.g., hemangioma (capillary and cavernous), glomus tumors, telangiectasia, bacillary angiomatosis, hemangioendothelioma, angiosarcoma, haemangiopericytoma, Kaposi's sarcoma, lymphangioma, and lymphangiosarcoma; tumor angiogenesis; trauma such as wounds, burns, and other injured tissue, implant fixation, scarring; ischemia reperfusion injury; rheumatoid arthritis; cerebrovascular disease; renal diseases such as acute renal failure, or osteoporosis.

In still yet another aspect, the immunological disorders are consistent with systemic lupus erythematosus; rheumatoid arthritis; juvenile chronic arthritis; spondyloarthropathies; systemic sclerosis (scleroderma); idiopathic inflammatory myopathies (dermatomyositis, polymyositis); Sjögren's syndrome; systemic vasculitis; sarcoidosis; autoimmune hemolytic anemia (immune pancytopenia, paroxysmal nocturnal hemoglobinuria); autoimmune thrombocytopenia (idiopathic thrombocytopenic purpura, immune-mediated thrombocytopenia); thyroiditis (Grave's disease, Hashimoto's thyroiditis, juvenile lymphocytic thyroiditis, atrophic thyroiditis); diabetes mellitus; immune-mediated renal disease (glomerulonephritis, tubulo-interstitial nephritis); demyelinating diseases of the central and peripheral nervous systems such as multiple sclerosis, idiopathic demyelinating polyneuropathy or Guillain-Barré syndrome, and chronic inflammatory demyelinating polyneuropathy; hepatobiliary diseases such as infectious hepatitis (hepatitis A, B, C, D, E and other non-hepatotropic viruses), autoimmune chronic active
hepatitis, primary biliary cirrhosis, granulomatous hepatitis, and sclerosing cholangitis; inflammatory bowel disease (ulcerative colitis; Crohn's disease); gluten-sensitive enteropathy, and Whipple's disease; autoimmune or immune-mediated skin diseases including bullous skin diseases, erythema multiforme and contact dermatitis, psoriasis; allergic diseases such as asthma, allergic rhinitis, atopic dermatitis, food hypersensitivity and urticaria; immunologic diseases of the lung such as eosinophilic pneumonia, idiopathic pulmonary fibrosis and hypersensitivity pneumonitis; or transplantation associated diseases including graft rejection and graft-versus-host disease.

In yet another aspect, the bone metabolic abnormality or disorder is arthritis, osteoporosis, osteopenia or osteopetrosis.

In another aspect, the non-human transgenic animal exhibits at least one of the following physiological characteristics compared with gender matched wild-type littermates: increased anxiety-like response during open field testing; hyperactivity during open field testing; decreased anxiety during open field testing; decreased locomotor activity during open field testing; abnormal circadian rhythm during home-cage activity testing (low activity during the light phase; altered sleep/wake cycle); abnormal circadian rhythm during home-cage activity testing including decreased ambulatory counts; hypoactivity with no circadian rhythm; abnormal circadian rhythm during home-cage activity testing including increased ambulatory counts; decreased rearing; increased sensitivity to stress induced hyperthermia (increased anxiety); impaired motor coordination during inverted screen testing; head tilt and retropulsion; increased prepulse inhibition response indicating enhanced sensorimotor gating/attention; decreased startle response during prepulse inhibition testing; no startle response indicating deafness or impaired hearing; decreased prepulse inhibition with impaired sensorimotor gating/attention; increased latency to respond in hot plate testing; decreased latency to respond in hot plate testing; ophthalmological abnormalities; impaired vision; white deposits of optic disc region; ocular infection and neutrophilia; bilateral optic disc lesion; decreased tear production; decreased heart rate; increased mean systolic blood pressure; decreased mean systolic blood pressure; increased mean fasting serum glucose levels; decreased mean serum glucose levels; increased mean serum cholesterol levels; decreased mean serum cholesterol levels; increased mean serum triglyceride levels; decreased mean serum triglyceride levels; impaired glucose tolerance; increased mean serum albumin, alanine amino transferase and phosphorus levels; increased mean serum alkaline phosphatase levels; urinary nitrites present; increased total white blood cell (WBC) count; decreased total white blood cell (WBC) count and absolute neutrophil count; increased mean absolute neutrophil count; increased mean absolute lymphocyte count; increased mean platelet count; increased mean red cell distribution width; decreased mean platelet count; reduced percentage of CD4 spleen thymocytes; decreased percentages of CD4 cells in the periphery resulting in increased percentages of B cells in lymph organs; CD4 cells exhibit a more activated/memory phenotype (CD62L low, CD44 hi); developmental defect in CD4+ cells; decreased percentages of CD4 cells and increased percentages of B cells in blood; decreased percentages of CD4 cells and increased percentages of B cells in tissues; increase in percentages of B cells in Peyer’s patches; decreased germinal center, isotype-switched B cells in Peyer’s patches (CD38 low; IgM negative); decreased CD23 intensity in spleen; increased mean percentages of B220 Med/CD23-cells and B220+/CD11b-Low/CD23- cells in peritoneal lavage; increased mean percentages of B cells in peripheral blood; decreased CD4 and CD8 T cells and increased B cells; increase in peritoneal B cells; reduction in CD11b-Hi cells in peritoneal cavity; decreased mean CD4 to CD8 ratio in spleen; decreased CD8 cells; decreased mean
percentages of B220+/CD23+ cells and B220+/CD1bLow/CD23- cells in peritoneal lavage; increased mean serum IgG1 response to ovalbumin challenge; increased mean serum IgG2a response to ovalbumin challenge; increased mean serum IL-6 response to LPS challenge; increased mean serum TNF alpha response to LPS challenge; increased mean serum MCP-1 response to LPS challenge; increased mean serum IgM level; increased mean serum IgA; increase mean serum IgG1; increased mean serum IgG2a; increased mean serum IgG2b; decreased mean serum IgG1 response to ovalbumin challenge; decreased mean serum IgG2a response to ovalbumin challenge; failure in ovalbumin response; decreased mean serum IgA level; decreased mean serum IgG2a level; decreased skin fibroblast proliferation rate; increased mean percent of total body fat and total fat mass; increased mean body weight; increased mean body length; increased total tissue mass (TTM); increased bone mineral density (BMD); increase in bone mineral content (BMC); increased mean femoral midshaft cortical thickness; decreased mean percent of total body fat and total fat mass; decreased mean body weight; decreased mean body length; decreased mean body weight and length in heterozygotes; decreased total tissue mass (TTM); decreased lean body mass (LBM); decreased femoral bone mineral density (BMD); decreased vertebral bone mineral density (BMD); decreased bone mineral density (BMD) in total body; decreased bone mineral content (BMC); decreased bone mineral density index; decreased volumetric bone mineral density (vBMD); decreased mean femoral midshaft cortical thickness; decreased mean femoral midshaft cross-sectional area; decreased mean vertebral trabecular bone volume, number and connectivity density; osteopenosis; osteoporosis; moderate kidney hydronephrosis; hydrocephalus; enlarged liver; induced in activated T cells; induced in activated NK cells and dendritic cells; myeloid B cell expression; hyperplasia of sebaceous glands and multifocal hyperplasia of the epidermis (acanthosis and hyperkeratosis); moderate dermatitis; increased extramedullary hematopoiesis in liver and spleen; myeloid hyperplasia of the bone marrow; encephalitis due to Group B streptococcus; meningitis due to E. Coli infection; lymphocytic infiltrates in salivary glands, pancreas and lungs; poor breeders requiring foster mothers; decreased litter size; homozygous mice were small and dehydrated; vacuolar degeneration of testes resulting in decreased sperm production and infertility; defective spermatogenesis in the testes; hypospermia and defective spermatozoa in the epididymus; male infertility; decreased testes weight; growth retardation; small mice and failure to thrive; reduced viability; reduced viability with situs inversus; and homozygous embryonic lethality.

The invention also provides an agent that ameliorates or modulates a neurological disorder; a cardiovascular, endothelial or angiogenic disorder; an eye abnormality; an immunological disorder; an oncological disorder; a bone metabolic abnormality or disorder; a lipid metabolic disorder; or a developmental abnormality which is associated with gene disruption. In one aspect, the agent is an agonist or antagonist of the phenotype associated with a disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO528, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO50332, PRO38465 or PRO346 polypeptide. In yet another aspect, the agent is an agonist or antagonist of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO528, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352,
PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide. In yet another aspect, the agonist agent is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1195, anti-PRO1271, anti-PRO1879, anti-PRO1889, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody. In still another aspect, the antagonist agent is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1195, anti-PRO1271, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

The invention also provides a therapeutic agent for the treatment of a neurological disorder; a cardiovascular, endothelial or angiogenic disorder; an eye abnormality; an immunological disorder; an oncological disorder; a bone metabolic abnormality or disorder; a lipid metabolic disorder; or a developmental abnormality.

The invention also provides a method of identifying an agent that modulates the expression of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1195, PRO1271, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising:

(a) contacting a test agent with a host cell expressing a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1195, PRO1271, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide; and

(b) determining whether the test agent modulates the expression of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1195, PRO1271, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide by the host cell.

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The invention also provides an agent that modulates the expression of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide. In one aspect, the agent is an agonist or antagonist of the phenotype associated with a disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide. In yet another aspect, the agent is an agonist or antagonist of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide. In yet another aspect, the agonist agent is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody. In still another aspect, the antagonist agent is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

The invention also provides a method of evaluating a therapeutic agent capable of affecting a condition associated with a disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising:
(a) providing a non-human transgenic animal whose genome comprises a disruption of the gene which encodes for the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide;
(b) measuring a physiological characteristic of the non-human transgenic animal of (a);
(c) comparing the measured physiological characteristic of (b) with that of a gender matched wild-type animal, wherein the physiological characteristic of the non-human transgenic animal that differs from the physiological characteristic of the wild-type animal is identified as a condition resulting from the gene disruption in the non-human transgenic animal;
(d) administering a test agent to the non-human transgenic animal of (a); and
(e) evaluating the effects of the test agent on the identified condition associated with gene disruption in the non-human transgenic animal.

In one aspect, the condition is a neurological disorder; a cardiovascular, endothelial or angiogenic disorder; an eye abnormality; an immunological disorder; an oncological disorder; a bone metabolic abnormality or disorder; a lipid metabolic disorder; or a developmental abnormality.

The invention also provides a therapeutic agent which is capable of affecting a condition associated with gene disruption. In one aspect, the agent is an agonist or antagonist of the phenotype associated with a disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide. In yet another aspect, the agent is an agonist or antagonist of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide. In yet another aspect, the agonist agent is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody. In still another aspect, the antagonist agent is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941,
anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

The invention also provides a pharmaceutical composition comprising a therapeutic agent capable of affecting the condition associated with gene disruption.

The invention also provides a method of treating or preventing or ameliorating a neurological disorder; cardiovascular, endothelial or angiogenic disorder; immunological disorder; oncological disorder; bone metabolic abnormality or disorder, or embryonic lethality associated with the disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising administering to a subject in need of such treatment whom may already have the disorder, or may be prone to have the disorder or may be in whom the disorder is to be prevented, a therapeutically effective amount of a therapeutic agent, or agonists or antagonists thereof, thereby effectively treating or preventing or ameliorating said disorder or disease.

In yet another aspect, the neurological disorder is an increased anxiety-like response during open field activity testing. In yet another aspect, the neurological disorder is a decreased anxiety-like response during open field activity testing. In yet another aspect, the neurological disorder is an abnormal circadian rhythm during home-cage activity testing. In yet another aspect, the neurological disorder is an enhanced motor coordination during inverted screen testing. In yet another aspect, the neurological disorder is impaired motor coordination during inverted screen testing. In yet another aspect, the neurological disorder includes depression, generalized anxiety disorders, attention deficit disorder, sleep disorder, hyperactivity disorder, obsessive compulsive disorder, schizophrenia, cognitive disorders, hyperalgesia and sensory disorders. Such neurological disorders include the category defined as “anxiety disorders” which include but are not limited to: mild to moderate anxiety, anxiety disorder due to a general medical condition, anxiety disorder not otherwise specified, generalized anxiety disorder, panic attack, panic disorder with agoraphobia, panic disorder without agoraphobia, posttraumatic stress disorder, social phobia, social anxiety, autism, specific phobia, substance-induced anxiety disorder, acute alcohol withdrawal, obsessive compulsive disorder, agoraphobia, monopolar disorders, bipolar disorder I or II, bipolar disorder not otherwise specified, cyclothymic disorder, depressive disorder, major depressive disorder, mood disorder, substance-induced mood disorder, enhancement of cognitive function, loss of cognitive function associated with but not limited to Alzheimer’s disease, stroke, or traumatic injury to the brain, seizures resulting from disease or injury including but not limited to epilepsy, learning disorders/disabilities, cerebral palsy. In addition, anxiety disorders may apply to personality disorders including but not limited to the following types: paranoid, antisocial, avoidant behavior, borderline personality disorders, dependent, histrionic, narcissistic, obsessive-compulsive, schizoid, and schizotypal.
In another aspect, the eye abnormality is a retinal abnormality. In still another aspect, the eye abnormality is consistent with vision problems or blindness. In yet another aspect, the retinal abnormality is consistent with retinitis pigmentosa or is characterized by retinal degeneration or retinal dysplasia.

In still another aspect, the retinal abnormalities are consistent with retinal dysplasia, various retinopathies, including retinopathy of prematurity, retrolental fibroplasia, neovascular glaucoma, age-related macular degeneration, diabetic macular edema, corneal neovascularization, corneal graft neovascularization, corneal graft rejection, retinal/choroidal neovascularization, neovascularization of the angle (rubeosis), ocular neovascular disease, vascular restenosis, arteriovenous malformations (AVM), menigioma, hemangioma, angiofibroma, thyroid hyperplasias (including Grave's disease), corneal and other tissue transplantation, retinal artery obstruction or occlusion; retinal degeneration causing secondary atrophy of the retinal vasculature, retinitis pigmentosa, macular dystrophies, Stargardt's disease, congenital stationary night blindness, choroideremia, gyrate atrophy, Leber's congenital amaurosis, retinoschisis disorders, Wagner's syndrome, Usher syndromes, Zellweger syndrome, Saldino-Mainzer syndrome, Senior-Loken syndrome, Bardet-Biedl syndrome, Alport's syndrome, Alstrom's syndrome, Cockayne's syndrome, dysplasia spondyloepiphysaria congenita, Flynn-Aird syndrome, Fredreich ataxia, Hallgren syndrome, Marshall syndrome, Albers-Schonberg disease, Refsum's disease, Kearns-Sayre syndrome, Waardenburg's syndrome, Alagille syndrome, myotonic dystrophy, olivopontocerebellar atrophy, Pierre-Marie dunsdrome, Stickler syndrome, carotinemia, cystinosis, Wolfram syndrome, Bassen-Kornzweig syndrome, abetalipoproteinemia, incontinentia pigmenti, Batten's disease, mucopolysaccharidoses, homocystinuria, or mannosidosis.

In still another aspect, the eye abnormality is a cataract. In still yet another aspect, the cataract is a systemic disease such as human Down's syndrome, Hallerman-Streiff syndrome, Lowe syndrome, galactosemia, Marfan syndrome, Trisomy 13-15, Alport syndrome, myotonic dystrophy, Fabry disease, hypoparathyroidism or Conradi syndrome.

In still another aspect, the developmental abnormality comprises embryonic lethality or reduced viability.

In yet another aspect, the cardiovascular, endothelial or angiogenic disorders are arterial diseases, such as diabetes mellitus; papilledema; optic atrophy; atherosclerosis; angina; myocardial infarctions such as acute myocardial infarctions, cardiac hypertrophy, and heart failure such as congestive heart failure; hypertension; inflammatory vasculitides; Reynaud's disease and Reynaud's phenomenon; aneurysms and arterial restenosis; venous and lymphatic disorders such as thrombophlebitis, lymphangitis, and lymphedema; peripheral vascular disease; cancer such as vascular tumors, e.g., hemangioma (capillary and cavernous), glomus tumors, telangiectasia, basillary angiomatosis, hemangioendothelioma, angiosarcoma, haemangioencrityoma, Kaposis's sarcoma, lymphangiomia, and lymphangiosarcoma; tumor angiogenesis; trauma such as wounds, burns, and other injured tissue, implant fixation, scarring; ischemia reperfusion injury; rheumatoid arthritis; cerebrovascular disease; renal diseases such as acute renal failure, or ostecoporosis.

In still yet another aspect, the immunological disorders are consistent with systemic lupus erythematosus; rheumatoid arthritis; juvenile chronic arthritis; spondyloarthopathies; systemic sclerosis (scleroderma); idiopathic inflammatory myopathies (dermatomyositis, polymyositis); Sjögren's syndrome; systemic vasculitis; sarcoidosis; autoimmune hemolytic anemia (immune pancytopenia, paroxysmal nocturnal hemoglobinuria); autoimmune thrombocytopenia (idiopathic thrombocytopenic purpura, immune-mediated thrombocytopenia); thyroiditis.
(Grave's disease, Hashimoto's thyroiditis, juvenile lymphocytic thyroiditis, atrophic thyroiditis); diabetes mellitus; immune-mediated renal disease (glomerulonephritis, tubulointerstitial nephritis); demyelinating diseases of the central and peripheral nervous systems such as multiple sclerosis, idiopathic demyelinating polyneuropathy or Guillain-Barré syndrome, and chronic inflammatory demyelinating polyneuropathy; hepatobiliary diseases such as infectious hepatitis (hepatitis A, B, C, D, E and other non-hepatotropic viruses), autoimmune chronic active hepatitis, primary biliary cirrhosis, granulomatous hepatitis, and sclerosing cholangitis; inflammatory bowel disease (ulcerative colitis; Crohn's disease); gluten-sensitive enteropathy, and Whipple's disease; autoimmune or immune-mediated skin diseases including bullous skin diseases, erythema multiforme and contact dermatitis, psoriasis; allergic diseases such as asthma, allergic rhinitis, atopic dermatitis, food hypersensitivity and urticaria; immunologic diseases of the lung such as eosinophilic pneumonia, idiopathic pulmonary fibrosis and hypersensitivity pneumonitis; or transplantation associated diseases including graft rejection and graft-versus-host disease.

In yet another aspect, the bone metabolic abnormality or disorder is arthritis, osteoporosis, osteopenia or osteopetrosis.

In another aspect the therapeutic agent is an agonist or antagonist of the phenotype associated with a disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO10014, PRO28694, PRO16089, PRO19563, PRO19675, PRO200084, PRO21434, PRO30332, PRO38465 or PRO346 polypeptide. In yet another aspect, the agent is an agonist or antagonist of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO200084, PRO21434, PRO30332, PRO38465 or PRO346 polypeptide. In yet another aspect, the agonist agent is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO200084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody. In still another aspect, the antagonist agent is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-
The invention also provides a method of identifying an agent that ameliorates or modulates a neurological disorder; a cardiovascular, endothelial or angiogenic disorder; an eye abnormality; an immunological disorder; an oncological disorder; a bone metabolic abnormality or disorder; a lipid metabolic disorder; or a developmental abnormality associated with a disruption in the gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising:

(a) providing a non-human transgenic animal cell culture, each cell of said culture comprising a disruption of the gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide;

(b) administering a test agent to said cell culture; and

(c) determining whether the test agent ameliorates or modulates the neurological disorder; cardiovascular, endothelial or angiogenic disorder; eye abnormality; immunological disorder; oncological disorder; bone metabolic abnormality or disorder; lipid metabolic disorder; or developmental abnormality in said culture. In yet another aspect, the neurological disorder is an increased anxiety-like response during open field activity testing. In yet another aspect, the neurological disorder is a decreased anxiety-like response during open field activity testing. In yet another aspect, the neurological disorder is an abnormal circadian rhythm during home-cage activity testing.

In yet another aspect, the neurological disorder is an enhanced motor coordination during inverted screen testing. In yet another aspect, the neurological disorder is impaired motor coordination during inverted screen testing. In yet another aspect, the neurological disorder includes depression, generalized anxiety disorders, attention deficit disorder, sleep disorder, hyperactivity disorder, obsessive compulsive disorder, schizophrenia, cognitive disorders, hyperalgesia and sensory disorders. Such neurological disorders include the category defined as “anxiety disorders” which include but are not limited to: mild to moderate anxiety, anxiety disorder due to a general medical condition, anxiety disorder not otherwise specified, generalized anxiety disorder, panic attack, panic disorder with agoraphobia, panic disorder without agoraphobia, posttraumatic stress disorder, social phobia, social anxiety, autism, specific phobia, substance-induced anxiety disorder, acute alcohol withdrawal, obsessive compulsive disorder, agoraphobia, monopolar disorders, bipolar disorder I or II, bipolar disorder not otherwise specified, cyclothymic disorder, depressive disorder, major depressive disorder, mood disorder, substance-induced mood disorder, enhancement of cognitive function, loss of cognitive function associated with but not limited to Alzheimer’s disease, stroke, or traumatic injury to the brain, seizures resulting from disease or injury including but not limited to epilepsy, learning disorders/disabilities, cerebral palsy. In addition, anxiety disorders may apply to
personality disorders including but not limited to the following types: paranoid, antisocial, avoidant behavior, borderline personality disorders, dependent, histrionic, narcissistic, obsessive-compulsive, schizoid, and schizotypal.

In another aspect, the eye abnormality is a retinal abnormality. In still another aspect, the eye abnormality is consistent with vision problems or blindness. In yet another aspect, the retinal abnormality is consistent with retinitis pigmentosa or is characterized by retinal degeneration or retinal dysplasia.

In still another aspect, the retinal abnormalities are consistent with retinal dysplasia, various retinopathies, including retinopathy of prematurity, retrolental fibroplasia, neovascular glaucoma, age-related macular degeneration, diabetic macular edema, corneal neovascularization, corneal graft neovascularization, corneal graft rejection, retinal/choroidal neovascularization, neovascularization of the angle (rubeosis), ocular neovascular disease, vascular restenosis, arteriovenous malformations (AVM), meningioma, hemangioma, angiofibroma, thyroid hyperplasias (including Grave's disease), corneal and other tissue transplantation, retinal artery obstruction or occlusion; retinal degeneration causing secondary atrophy of the retinal vasculature, retinitis pigmentosa, macular dystrophies, Stargardt's disease, congenital stationary night blindness, choroideremia, gyrate atrophy, Leber's congenital amaurosis, retinoschisis disorders, Wagner's syndrome, Usher syndromes, Zellweger syndrome, Saldino-Mainzer syndrome, Senior-Loken syndrome, Bardet-Biedl syndrome, Alport's syndrome, Alstrom's syndrome, Cockayne's syndrome, dysplasia spondyloepiphysaria congenita, Flynn-Aird syndrome, Friedreich ataxia, Hallgren syndrome, Marshall syndrome, Albers-Schonberg disease, Refsum's disease, Kearns-Sayre syndrome, Waardenburg's syndrome, Alagille syndrome, myotonic dystrophy, olivopontocerebellar atrophy, Pierre-Marie dunsdrome, Stickler syndrome, carotinemia, cystinosis, Wolfram syndrome, Bassen-Kornzweig syndrome, abetalipoproteinemia, incontinentia pigmenti, Batten's disease, mucopolysaccharidoses, homocystinuria, or mannosidosis.

In still another aspect, the eye abnormality is a cataract. In still yet another aspect, the cataract is a systemic disease such as Down's syndrome, Hallerman-Streiff syndrome, Lowe syndrome, galactosemia, Marfan syndrome, Trisomy 13-15, Alport syndrome, myotonic dystrophy, Fabry disease, hypoparathyroidism or Conradi syndrome.

In still another aspect, the developmental abnormality comprises embryonic lethality or reduced viability.

In yet another aspect, the cardiovascular, endothelial or angiogenic disorders are arterial diseases, such as diabetes mellitus; papilledema; optic atrophy; atherosclerosis; angina; myocardial infarctions such as acute myocardial infarctions, cardiac hypertrophy, and heart failure such as congestive heart failure; hypertension; inflammatory vasculitides; Reynaud's disease and Reynaud's phenomenon; aneurysms and arterial restenosis; venous and lymphatic disorders such as thrombophlebitis, lymphangitis, and lymphedema; peripheral vascular disease; cancer such as vascular tumors, e.g., hemangioma (capillary and cavernous), glomus tumors, telangiectasia, bacillary angiomatosis, hemangioendothelioma, angiosarcoma, haemangiopericytoma, Kaposi's sarcoma, lymphangiomata, and lymphangiosarcoma; tumor angiogenesis; trauma such as wounds, burns, and other injured tissue, implant fixation, scarring; ischemia reperfusion injury; rheumatoid arthritis; cerebrovascular disease; renal diseases such as acute renal failure, or osteoporosis.

In still yet another aspect, the immunological disorders are consistent with systemic lupus erythematosis; rheumatoid arthritis; juvenile chronic arthritis; spondyloarthropathies; systemic sclerosis (scleroderma); idiopathic inflammatory myopathies (dermatomyositis, polymyositis); Sjögren's syndrome; systemic vasculitis; sarcoidosis;
autoimmune hemolytic anemia (immune pancytopenia, paroxysmal nocturnal hemoglobinuria); autoimmune thrombocytopenia (idiopathic thrombocytopenic purpura, immune-mediated thrombocytopenia); thyroiditis (Grave's disease, Hashimoto's thyroiditis, juvenile lymphocytic thyroiditis, atrophic thyroiditis); diabetes mellitus; immune-mediated renal disease (glomerulonephritis, tubulointerstitial nephritis); demyelinating diseases of the central and peripheral nervous systems such as multiple sclerosis, idiopathic demyelinating polyneuropathy or Guillain-Barré syndrome, and chronic inflammatory demyelinating polyneuropathy; hepatobiliary diseases such as infectious hepatitis (hepatitis A, B, C, D, E and other non-hepatotropic viruses), autoimmune chronic active hepatitis, primary biliary cirrhosis, granulomatous hepatitis, and sclerosing cholangitis; inflammatory bowel disease (ulcerative colitis; Crohn's disease); gluten-sensitive enteropathy, and Whipple's disease; autoimmune or immune-mediated skin diseases including bullous skin diseases, erythema multiforme and contact dermatitis, psoriasis; allergic diseases such as asthma, allergic rhinitis, atopic dermatitis, food hypersensitivity and urticaria; immunologic diseases of the lung such as eosinophilic pneumonia, idiopathic pulmonary fibrosis and hypersensitivity pneumonitis; or transplantation associated diseases including graft rejection and graft-versus-host disease.

In yet another aspect, the bone metabolic abnormality or disorder is arthritis, osteoporosis, osteopenia or osteopetrosis.

The invention also provides an agent that ameliorates or modulates a neurological disorder; a cardiovascular, endothelial or angiogenic disorder; an eye abnormality; an immunological disorder; an oncological disorder; a bone metabolic abnormality or disorder; a lipid metabolic disorder; or a developmental abnormality which is associated with gene disruption in said culture. In one aspect, the agent is an agonist or antagonist of the phenotype associated with a disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide. In yet another aspect, the agent is an agonist or antagonist of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide. In yet another aspect, the agonist agent is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody. In still another aspect, the antagonist agent is an anti-PRO218, anti-PRO228, anti-PRO271, anti-
PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO3429, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO99048, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

The invention also provides a method of modulating a phenotype associated with a disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99048, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising administering to a subject whom may already have the phenotype, or may be prone to have the phenotype or may be in whom the phenotype is to be prevented, an effective amount of an agent identified as modulating said phenotype, or agonists or antagonists thereof, thereby effectively modulating the phenotype.

The invention also provides a method of modulating a physiological characteristic associated with a disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99048, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising administering to a subject whom may already exhibit the physiological characteristic, or may be prone to exhibit the physiological characteristic or may be in whom the physiological characteristic is to be prevented, an effective amount of an agent identified as modulating said physiological characteristic, or agonists or antagonists thereof, thereby effectively modulating the physiological characteristic.

The invention also provides a method of modulating a behavior associated with a disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99048, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising administering to a subject whom may already exhibit the behavior, or may be prone to exhibit the behavior or may be in whom the exhibited behavior is to be prevented, an effective amount of an agent identified as modulating said behavior, or agonists or antagonists thereof, thereby effectively modulating the behavior.

The invention also provides a method of modulating the expression of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130,
PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising administering to a host cell expressing said PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, an effective amount of an agent identified as modulating said expression, or agonists or antagonists thereof, thereby effectively modulating the expression of said polypeptide.

The invention also provides a method of modulating a condition associated with a disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising administering to a subject whom may have the condition, or may be prone to have the condition or may be in whom the condition is to be prevented, a therapeutically effective amount of a therapeutic agent identified as modulating said condition, or agonists or antagonists thereof, thereby effectively modulating the condition.

The invention also provides a method of treating or preventing or ameliorating a neurological disorder, cardiovascular, endothelial or angiogenic disorder; immunological disorder; oncological disorder; bone metabolic abnormality or disorder; or embryonic lethality associated with the disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising administering to a non-human transgenic animal cell culture, each cell of said culture comprising a disruption of the gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, an effective amount of an agent identified as treating or preventing or ameliorating said disorder, or agonists or antagonists thereof, thereby effectively treating or preventing or ameliorating said disorder.

B. Further Embodiments

In yet further embodiments, the invention is directed to the following set of potential claims for this
application:

1. A method of identifying a phenotype associated with a disruption of a gene which encodes for a 
   PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, 
   PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, 
   PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, 
   PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, 
   PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising:
   
   (a) providing a non-human transgenic animal whose genome comprises a disruption of the gene which 
   encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, 
   PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, 
   PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, 
   PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, 
   PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide; 
   
   (b) measuring a physiological characteristic of the non-human transgenic animal; and 
   
   (c) comparing the measured physiological characteristic with that of a gender matched wild-type animal, 
   wherein the physiological characteristic of the non-human transgenic animal that differs from the physiological 
   characteristic of the wild-type animal is identified as a phenotype resulting from the gene disruption in the non-
   human transgenic animal.

2. The method of Claim 1, wherein the non-human transgenic animal is heterozygous for the disruption 
   of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, 
   PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, 
   PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, 
   PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, 
   PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 
   polypeptide.

3. The method of Claim 1, wherein the phenotype exhibited by the non-human transgenic animal as 
   compared with gender matched wild-type littersmates is at least one of the following: a neurological disorder; a 
   cardiovascular, endothelial or angiogenic disorder; an eye abnormality; an immunological disorder; an oncological 
   disorder; a bone metabolic abnormality or disorder; a lipid metabolic disorder; or a developmental abnormality.

4. The method of Claim 3, wherein the neurological disorder is an increased anxiety-like response during 
   open field activity testing.

5. The method of Claim 3, wherein the neurological disorder is a decreased anxiety-like response during 
   open field activity testing.

6. The method of Claim 3, wherein the neurological disorder is an abnormal circadian rhythm during home-
   cage activity testing.

7. The method of Claim 3, wherein the neurological disorder is an enhanced motor coordination during 
   inverted screen testing.

8. The method of Claim 3, wherein the neurological disorder is an impaired motor coordination during
inverted screen testing.

9. The method of Claim 3, wherein the neurological disorder is depression, generalized anxiety disorders, attention deficit disorder, sleep disorder, hyperactivity disorder, obsessive compulsive disorder, schizophrenia, cognitive disorders, hyperalgesia or sensory disorders.

10. The method of Claim 3, wherein the eye abnormality is a retinal abnormality.

11. The method of Claim 3, wherein the eye abnormality is consistent with vision problems or blindness.

12. The method of Claim 10, wherein the retinal abnormality is consistent with retinitis pigmentosa.

13. The method of Claim 10, wherein the retinal abnormality is characterized by retinal degeneration or retinal dysplasia.

14. The method of Claim 10, wherein the retinal abnormality is consistent with retinal dysplasia, various retinopathies, including retinopathy of prematurity, retrolental fibroplasia, neovascular glaucoma, age-related macular degeneration, diabetic macular edema, corneal neovascularization, corneal graft neovascularization, corneal graft rejection, retinal/choroidal neovascularization, neovascularization of the angle (rubecosis), ocular neovascular disease, vascular restenosis, arteriovenous malformations (AVM), meningioma, hemangioma, angiofibroma, thyroid hyperplasias (including Grave's disease), corneal and other tissue transplantation, retinal artery obstruction or occlusion; retinal degeneration causing secondary atrophy of the retinal vasculature, retinitis pigmentosa, macular dystrophies, Stargardt's disease, congenital stationary night blindness, choroideremia, gyrate atrophy, Leber's congenital amaurosis, retinoschisis disorders, Wagner's syndrome, Usher syndromes, Zellweger syndrome, Saldino-Mainzer syndrome, Senior-Loken syndrome, Bardet-Biedl syndrome, Alport's syndrome, Alstrom's syndrome, Cockayne's syndrome, dysplasia spondyloepiphysaria congenita, Flynn-Aird syndrome, Friedreich ataxia, Hallgren syndrome, Marshall syndrome, Albers-Schonberg disease, Reesum's disease, Kerns-Sayre syndrome, Waardenburg's syndrome, Alagille syndrome, myotonic dystrophy, olivopontocerebellar atrophy, Pierre-Marie dunderdrome, Stickler syndrome, carotinemia, cystinosis, Wolfman syndrome, Bassen-Kornzweig syndrome, abetalipoproteinemia, incontinentia pigmenti, Batten's disease, mucopolysaccharidoses, homocystinuria, or mannosidosis.

15. The method of Claim 3, wherein the eye abnormality is a cataract.

16. The method of Claim 15, wherein the cataract is consistent with systemic diseases such as human Down's syndrome, Hallerman-Streiff syndrome, Lowe syndrome, galactosemia, Marfan syndrome, Trisomy 13-15, Alport syndrome, myotonic dystrophy, Fabry disease, hypoparathyroidism or Conradi syndrome.

17. The method of Claim 3, wherein the developmental abnormality comprises embryonic lethality or reduced viability.

18. The method of Claim 3, wherein the cardiovascular, endothelial or angiogenic disorders are arterial diseases, such as diabetes mellitus; papilledema; optic atrophy; atherosclerosis; angina; myocardial infarctions such as acute myocardial infarctions, cardiac hypertrophy, and heart failure such as congestive heart failure; hypertension; inflammatory vasculitides; Reyonud's disease and Reynud's phenomenon; aneurysms and arterial restenosis; venous and lymphatic disorders such as thrombophlebitis, lymphangitis, and lymphedema; peripheral vascular disease; cancer such as vascular tumors, e.g., hemangioma (capillary and cavernous), glomus tumors, telangiectasia, bacillary angiomatosis, hemangioendothelioma, angiosarcoma, haemangiopericytoma, Kaposi's sarcoma, lymphangioma, and lymphangiosarcoma; tumor angiogenesis; trauma such as wounds, burns, and other
injured tissue, implant fixation, scarring; ischemia reperfusion injury; rheumatoid arthritis; cerebrovascular disease; renal diseases such as acute renal failure, or osteoporosis.

19. The method of Claim 3, wherein the immunological disorders are systemic lupus erythematosus; rheumatoid arthritis; juvenile chronic arthritis; spondyloarthopathies; systemic sclerosis (scleroderma); idiopathic inflammatory myopathies (dermatomyositis, polymyositis); Sjögren's syndrome; systemic vasculitis; sarcoidosis; autoimmune hemolytic anemia (immune pancytopenia, paroxysmal nocturnal hemoglobinuria); autoimmune thrombocytopenia (idiopathic thrombocytopenic purpura, immune-mediated thrombocytopenia); thyroiditis (Grave's disease, Hashimoto's thyroiditis, juvenile lymphocytic thyroiditis, atrophic thyroiditis); diabetes mellitus; immune-mediated renal disease (glomerulonephritis, tubulointerstitial nephritis); demyelinating diseases of the central and peripheral nervous systems such as multiple sclerosis, idiopathic demyelinating polyneuropathy or Guillain-Barré syndrome, and chronic inflammatory demyelinating polyneuropathy; hepatobiliary diseases such as infectious hepatitis (hepatitis A, B, C, D, E and other non-hepatotropic viruses), autoimmune chronic active hepatitis, primary biliary cirrhosis, granulomatous hepatitis, and sclerosing cholangitis; inflammatory bowel disease (ulcerative colitis; Crohn's disease); gluten-sensitive enteropathy, and Whipple's disease; autoimmune or immune-mediated skin diseases including bullous skin diseases, erythema multiforme and contact dermatitis, psoriasis; allergic diseases such as asthma, allergic rhinitis, atopic dermatitis, food hypersensitivity and urticaria; immunologic diseases of the lung such as eosinophilic pneumonia, idiopathic pulmonary fibrosis and hypersensitivity pneumonitis; or transplantation associated diseases including graft rejection and graft-versus-host disease.

20. The method of Claim 3, wherein the bone metabolic abnormality or disorder is arthritis, osteoporosis or osteopetrosis.

21. The method of Claim 1, wherein the non-human transgenic animal exhibits at least one of the following physiological characteristics compared with gender matched wild-type littermates: increased anxiety-like response during open field testing; hyperactivity during open field testing; decreased anxiety during open field testing; decreased locomotor activity during open field testing; abnormal circadian rhythm during home-cage activity testing (low activity during the light phase; altered sleep/wake cycle); abnormal circadian rhythm during home-cage activity testing including decreased ambulatory counts; hypoactivity with no circadian rhythm; abnormal circadian rhythm during home-cage activity testing including increased ambulatory counts; decreased rearing; increased sensitivity to stress induced hyperthermia (increased anxiety); impaired motor coordination during inverted screen testing; head tilt and retropulsion; increased prepulse inhibition response indicating enhanced sensorimotor gating/attention; decreased startle response during prepulse inhibition testing; no startle response indicating deafness or impaired hearing; decreased prepulse inhibition with impaired sensorimotor gating/attention; increased latency to respond in hot plate testing; decreased latency to respond in hot plate testing; ophthalmological abnormalities; impaired vision; white deposits of optic disc region; ocular infection and neutrophilia; bilateral optic disc lesion; decreased tear production; decreased heart rate; increased mean systolic blood pressure; decreased mean systolic blood pressure; increased mean fasting serum glucose levels; decreased mean serum glucose levels; increased mean serum cholesterol levels; decreased mean serum cholesterol levels; increased mean serum triglyceride levels; decreased mean serum triglyceride levels; impaired glucose tolerance; increased mean serum albumin, alanine amino transferase and phosphorus levels; increased mean serum alkaline phosphatase levels;
urinary nitrites present; increased total white blood cell (WBC) count; decreased total white blood cell (WBC) count and absolute neutrophil count; increased mean absolute neutrophil count; increased mean absolute lymphocyte count; increased mean platelet count; increased mean red cell distribution width; decreased mean platelet count; reduced percentage of CD4 spleen thymocytes; decreased percentages of CD4 cells in the periphery resulting in increased percentages of B cells in lymph organs; CD4 cells exhibit a more activated/memory phenotype (CD62Llow, CD44hi); developmental defect in CD4+ cells; decreased percentages of CD4 cells and increased percentages of B cells in blood; decreased percentages of CD4 cells and increased percentages of B cells in tissues; increased in percentages of B cells in Peyer’s patches; decreased germinal center, isotype-switched B cells in Peyer’s patches (CD38low;IgM negative); decreased CD23 intensity in spleen; increased mean percentages of B220 Med/CD23- cells and B220+/CD11b-Low/CD23- cells in peritoneal lavage; increased mean percentages of B cells in peripheral blood; decreased CD4 and CD8 T cells and increased B cells; increase in peritoneal B cells; reduction in CD11b-Hi cells in peritoneal cavity; decreased mean CD4 to CD8 ratio in spleen; decreased CD8 cells; decreased mean percentages of B220+/CD23+ cells and B220+/CD11b-Low/CD23- cells in peritoneal lavage; increased mean serum IgG1 response to ovalbumin challenge; increased mean serum IgG2a response to ovalbumin challenge; increased mean serum IL-6 response to LPS challenge; increased mean serum TNF alpha response to LPS challenge; increased mean serum MCP-1 response to LPS challenge; increased mean serum IgM level; increased mean serum IgA; increased mean serum IgG1; increased mean serum IgG2a; increased mean serum IgG2b; decreased mean serum IgG1 response to ovalbumin challenge; decreased mean serum IgG2a response to ovalbumin challenge; failure in ovalbumin response; decreased mean serum IgA level; decreased mean serum IgG2a level; decreased skin fibroblast proliferation rate; increased mean percent of total body fat and total fat mass; increased mean body weight; increased mean body length; increased total tissue mass (TTM); increased bone mineral density (BMD); increased in bone mineral content (BMC); increased mean femoral midshaft cortical thickness; decreased mean percent of total body fat and total fat mass; decreased mean body weight; decreased mean body length; decreased mean body weight and length in heterozygotes; decreased total tissue mass (TTM); decreased lean body mass (LBM); decreased femoral bone mineral density (BMD); decreased vertebral bone mineral density (BMD); decreased bone mineral density (BMD) in total body; decreased bone mineral content (BMC); decreased bone mineral density index; decreased volumetric bone mineral density (vBMD); decreased mean femoral midshaft cortical thickness; decreased mean femoral midshaft cross-sectional area; decreased mean vertebral trabecular bone volume, number and connectivity density; osteopetrosis; osteoporosis; moderate kidney hydropsphrosis; hydrocephalus; enlarged liver; induced in activated T cells; induced in activated NK cells and dendritic cells; myeloid B cell expression; hyperplasia of sebaceous glands and multifocal hyperplasia of the epidermis (acanthosis and hyperkeratosis); moderate dermatitis; increased extramedullary hematopoiesis in liver and spleen; myeloid hyperplasia of the bone marrow; encephalitis due to Group B streptococcus; meningitis due to E. Coli infection; lymphocytic infiltrates in salivary glands, pancreas and lungs; poor breeders requiring foster mothers; decreased litter size; homozygous mice were small and dehydrated; vacuolar degeneration of testes resulting in decreased sperm production and infertility; defective spermatogenesis in the testes; hypospermia and defective spermatozoa in the epididymus; male infertility; decreased testes weight; growth retardation; small mice and failure to thrive; reduced viability; reduced viability with situs inveritus; and homozygous embryonic lethality.
disruption of the gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99098, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide.

23. The isolated cell of Claim 22 which is a murine cell.
24. The isolated cell of Claim 23, wherein the murine cell is an embryonic stem cell.
25. The isolated cell of Claim 22, wherein the non-human transgenic animal exhibits at least one of the following phenotypes compared with gender matched wild-type littermates; a neurological disorder; a cardiovascular, endothelial or angiogenic disorder; an eye abnormality; an immunological disorder; an oncological disorder; a bone metabolic abnormality or disorder; a lipid metabolic disorder; or a developmental abnormality.
26. A method of identifying an agent that modulates a phenotype associated with a disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99098, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising:

   (a) providing a non-human transgenic animal whose genome comprises a disruption of the gene which encodes for the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99098, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide;

   (b) measuring a physiological characteristic of the non-human transgenic animal of (a);

   (c) comparing the measured physiological characteristic of (b) with that of a gender matched wild-type animal, wherein the physiological characteristic of the non-human transgenic animal that differs from the physiological characteristic of the wild-type animal is identified as a phenotype resulting from the gene disruption in the non-human transgenic animal;

   (d) administering a test agent to the non-human transgenic animal of (a); and

   (e) determining whether the test agent modulates the identified phenotype associated with gene disruption in the non-human transgenic animal.

27. The method of Claim 26, wherein the phenotype associated with the gene disruption comprises a neurological disorder; a cardiovascular, endothelial or angiogenic disorder; an eye abnormality; an immunological disorder; an oncological disorder; a bone metabolic abnormality or disorder; a lipid metabolic disorder; or a developmental abnormality.
28. The method of Claim 27, wherein the neurological disorder is an increased anxiety-like response during open field activity testing.
29. The method of Claim 27, wherein the neurological disorder is a decreased anxiety-like response during open field activity testing.
30. The method of Claim 27, wherein the neurological disorder is an abnormal circadian rhythm during home-cage activity testing.
31. The method of Claim 27, wherein the neurological disorder is an enhanced motor coordination during inverted screen testing.
32. The method of Claim 27, wherein the neurological disorder is an impaired motor coordination during inverted screen testing.
33. The method of Claim 27, wherein the neurological disorder is depression, generalized anxiety disorders, attention deficit disorder, sleep disorder, hyperactivity disorder, obsessive compulsive disorder, schizophrenia, cognitive disorders, hyperalgesia or sensory disorders.
34. The method of Claim 27, wherein the eye abnormality is a retinal abnormality.
35. The method of Claim 27, wherein the eye abnormality is consistent with vision problems or blindness.
36. The method of Claim 34, wherein the retinal abnormality is consistent with retinitis pigmentosa.
37. The method of Claim 34, wherein the retinal abnormality is characterized by retinal degeneration or retinal dysplasia.
38. The method of Claim 34, wherein the retinal abnormality is consistent with retinal dysplasia, various retinopathies, including retinopathy of prematurity, retrolental fibroplasia, neovascular glaucoma, age-related macular degeneration, diabetic macular edema, corneal neovascularization, corneal graft neovascularization, corneal graft rejection, retinal/choroidal neovascularization, neovascularization of the angle (rubecosis), ocular neovascular disease, vascular restenosis, arteriovenous malformations (AVM), meningioma, hemangioma, angiofibroma, thyroid hyperplasias (including Grave's disease), corneal and other tissue transplantation, retinal artery obstruction or occlusion; retinal degeneration causing secondary atrophy of the retinal vasculature, retinitis pigmentosa, macular dystrophies, Stargardt's disease, congenital stationary night blindness, choroideremia, gyrate atrophy, Leber's congenital amaurosis, retinoschisis disorders, Wagner's syndrome, Usher syndromes, Zellweger syndrome, Saldino-Mainzer syndrome, Senior-Loken syndrome, Bardet-Biedl syndrome, Alport's syndrome, Alstrom's syndrome, Cockayne's syndrome, dysplasia spondyloepiphysaria congenita, Flynn-Aird syndrome, Friedreich ataxia, Hallgren syndrome, Marshall syndrome, Albers-Schonberg disease, Reisum's disease, Kearns-Sayre syndrome, Waardenburg's syndrome, Alagille syndrome, myotonic dystrophy, olivopontocerebellar atrophy, Pierre-Marie-Drash syndrome, Stickler syndrome, carotinemia, cystinosis, Wollfram syndrome, Bassen-Kornzweig syndrome, abetalipoproteinemia, incontinentia pigmenti, Batten's disease, mucopolysaccharidoses, homocystinuria, or mannosidosis.
39. The method of Claim 27, wherein the eye abnormality is a cataract.
40. The method of Claim 39, wherein the cataract is consistent with systemic diseases such as human Down's syndrome, Hallerman-Streiff syndrome, Lowe syndrome, galactosemia, Marfan syndrome, Trisomy 13-15, Alport syndrome, myotonic dystrophy, Fabry disease, hypoparathyroidism or Conradi syndrome.
41. The method of Claim 39, wherein the developmental abnormality comprises embryonic lethality or reduced viability.
42. The method of Claim 27, wherein the cardiovascular, endothelial or angiogenic disorders are arterial
diseases, such as diabetes mellitus; papilledema; optic atrophy; atherosclerosis; angina; myocardial infarctions such as acute myocardial infarctions, cardiac hypertrophy, and heart failure such as congestive heart failure; hypertension; inflammatory vasculitides; Reynaud's disease and Reynaud's phenomenon; aneurysms and arterial restenosis; venous and lymphatic disorders such as thrombophlebitis, lymphangitis, and lymphedema; peripheral vascular disease; cancer such as vascular tumors, *e.g.*, hemangioma (capillary and cavernous), glomus tumors, telangietasia, bacillary angiomatosis, hemangioendothelioma, angiosarcoma, haemangiopericytoma, Kaposi's sarcoma, lymphangioma, and lymphangiosarcoma; tumor angiogenesis; trauma such as wounds, burns, and other injured tissue, implant fixation, scarring; ischemia reperfusion injury; rheumatoid arthritis; cerebrovascular disease; renal diseases such as acute renal failure, or osteoporosis.

43. The method of Claim 27, wherein the immunological disorders are systemic lupus erythematosus; rheumatoid arthritis; juvenile chronic arthritis; spondyloarthropathies; systemic sclerosis (scleroderma); idiopathic inflammatory myopathies (dermatomyositis, polymyositis); Sjögren's syndrome; systemic vasculitis; sarcoidosis; autoimmune hemolytic anemia (immune pancytopenia, paroxysmal nocturnal hemoglobinuria); autoimmune thrombocytopenia (idiopathic thrombocytopenic purpura, immune-mediated thrombocytopenia); thyroiditis (Grave's disease, Hashimoto's thyroiditis, juvenile lymphocytic thyroiditis, atrophic thyroiditis); diabetes mellitus; immune-mediated renal disease (glomerulonephritis, tubulointerstitial nephritis); demyelinating diseases of the central and peripheral nervous systems such as multiple sclerosis, idiopathic demyelinating polyneuropathy or Guillain-Barré syndrome, and chronic inflammatory demyelinating polyneuropathy; hepatobiliary diseases such as infectious hepatitis (hepatitis A, B, C, D, E and other non-hepatotropic viruses), autoimmune chronic active hepatitis, primary biliary cirrhosis, granulomatous hepatitis, and sclerosing cholangitis; inflammatory bowel disease (ulcerative colitis; Crohn's disease); gluten-sensitive enteropathy, and Whipple's disease; autoimmune or immune-mediated skin diseases including bullous skin diseases, erythema multiforme and contact dermatitis, psoriasis; allergic diseases such as asthma, allergic rhinitis, atopic dermatitis, food hypersensitivity and urticaria; immunologic diseases of the lung such as eosinophilic pneumonia, idiopathic pulmonary fibrosis and hypersensitivity pneumonitis; or transplantation-associated diseases including graft rejection and graft-versus-host disease.

44. The method of Claim 27, wherein said bone metabolic abnormality or disorder is arthritis, osteoporosis or osteopetrosis.

45. The method of Claim 26, wherein the non-human transgenic animal exhibits at least one of the following physiological characteristics compared with gender matched wild-type littermates: increased anxiety-like response during open field testing; hyperactivity during open field testing; decreased anxiety during open field testing; decreased locomotor activity during open field testing; abnormal circadian rhythm during home-cage activity testing (low activity during the light phase; altered sleep/wake cycle); abnormal circadian rhythm during home-cage activity testing including decreased ambulatory counts; hypoactivity with no circadian rhythm; abnormal circadian rhythm during home-cage activity testing including increased ambulatory counts; decreased rearing; increased sensitivity to stress induced hyperthermia (increased anxiety); impaired motor coordination during inverted screen testing; head tilt and retropulsion; increased prepulse inhibition response indicating enhanced sensorimotor gating/attention; decreased startle response during prepulse inhibition testing; no startle response indicating deafness or impaired hearing; decreased prepulse inhibition with impaired sensorimotor gating/attention; increased
latency to respond in hot plate testing; decreased latency to respond in hot plate testing; ophthalmological abnormalities; impaired vision; white deposits of optic disc region; ocular infection and neutrophilia; bilateral optic disc lesion; decreased tear production; decreased heart rate; increased mean systolic blood pressure; decreased mean systolic blood pressure; increased mean fasting serum glucose levels; decreased mean serum glucose levels; increased mean serum cholesterol levels; decreased mean serum cholesterol levels; increased mean serum triglyceride levels; decreased mean serum triglyceride levels; impaired glucose tolerance; increased mean serum albumin, alanine amino transferase and phosphorus levels; increased mean serum alkaline phosphatase levels; urinary nitrites present; increased total white blood cell (WBC) count; decreased total white blood cell (WBC) count and absolute neutrophil count; increased mean absolute neutrophil count; increased mean absolute lymphocyte count; increased mean platelet count; increased mean red cell distribution width; decreased mean platelet count; reduced percentage of CD4 spleen thymocytes; decreased percentages of CD4 cells in the periphery resulting in increased percentages of B cells in lymph organs; CD4 cells exhibit a more activated/memory phenotype (CD62Llow, CD44hi); developmental defect in CD4+ cells; decreased percentages of CD4 cells and increased percentages of B cells in blood; decreased percentages of CD4 cells and increased percentages of B cells in tissues; increase in percentages of B cells in Peyer’s patches; decreased germinal center, isotype-switched B cells in Peyer’s patches (CD38low, IgM negative); decreased CD23 intensity in spleen; increased mean percentages of B220 Med/CD23- cells and B220+/CD11b-Low/CD23- cells in peritoneal lavage; increased mean percentages of B cells in peripheral blood; decreased CD4 and CD8 T cells and increased B cells; increase in peritoneal B cells; reduction in CD11b-Hi cells in peritoneal cavity; decreased mean CD4 to CD8 ratio in spleen; decreased CD8 cells; decreased mean percentages of B220+/CD23+ cells and B220+/CD11bLow/CD23- cells in peritoneal lavage; increased mean serum IgG1 response to ovalbumin challenge; increased mean serum IgG2a response to ovalbumin challenge; increased mean serum IL-6 response to LPS challenge; increased mean serum TNF alpha response to LPS challenge; increased mean serum MCP-1 response to LPS challenge; increased mean serum IgM level; increased mean serum IgA; increased mean serum IgG1; increased mean serum IgG2a; increased mean serum IgG2b; decreased mean serum IgG1 response to ovalbumin challenge; decreased mean serum IgG2a response to ovalbumin challenge; failure in ovalbumin response; decreased mean serum IgA level; decreased mean serum IgG2a level; decreased skin fibroblast proliferation rate; increased mean percent of total body fat and total fat mass; increased mean body weight; increased mean body length; increased total tissue mass (TTM); increased bone mineral density (BMD); increase in bone mineral content (BMC); increased mean femoral midshaft cortical thickness; decreased mean percent of total body fat and total fat mass; decreased mean body weight; decreased mean body length; decreased mean body weight and length in heterozygotes; decreased total tissue mass (TTM); decreased lean body mass (LBM); decreased femoral bone mineral density (BMD); decreased vertebral bone mineral density (BMD); decreased bone mineral density (BMD) in total body; decreased bone mineral content (BMC); decreased bone mineral density index; decreased volumetric bone mineral density (vBMD); decreased mean femoral midshaft cortical thickness; decreased mean femoral midshaft cross-sectional area; decreased mean vertebral trabecular bone volume, number and connectivity density; osteopetrosis; osteoporosis; moderate kidney hydronephrosis; hydrocephalus; enlarged liver; induced in activated T cells; induced in activated NK cells and dendritic cells; myeloid B cell expression; hyperplasia of sebaceous glands and multifocal hyperplasia of the epidermis (acanthosis and hyperkeratosis); moderate dermatitis; increased extramedullary hematopoiesis in liver
and spleen; myeloid hyperplasia of the bone marrow; encephalitis due to Group B streptococcus; meningitis due to E. Coli infection; lymphocytic infiltrates in salivary glands, pancreas and lungs; poor breeders requiring foster mothers; decreased litter size; homozygous mice were small and dehydrated; vacuolar degeneration of testes resulting in decreased sperm production and infertility; defective spermatogenesis in the testes; hyposperma and defective spermatozoa in the epididymus; male infertility; decreased testes weight; growth retardation; small mice and failure to thrive; reduced viability; reduced viability with situs inveritus; and homozygous embryonic lethality.

46. An agent identified by the method of Claim 26.

47. The agent of Claim 46 which is an agonist or antagonist of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide.

48. The agent of Claim 47, wherein the agonist is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

49. The agent of Claim 47, wherein the antagonist is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

50. A method of identifying an agent that modulates a physiological characteristic associated with a disruption of the gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising:

(a) providing a non-human transgenic animal whose genome comprises a disruption of the gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069,
PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1879, PRO3446, PRO3543, PRO4329. 
PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, 
PRO19563, PRO19675, PRO20084, PRO21434, PRO5032, PRO38465 or PRO346 polypeptide; 
(b) measuring a physiological characteristic exhibited by the non-human transgenic animal of (a); 
(c) comparing the measured physiological characteristic of (b) with that of a gender matched wild-type 
animal, wherein the physiological characteristic exhibited by the non-human transgenic animal that differs from 
the physiological characteristic exhibited by the wild-type animal is identified as a physiological characteristic 
associated with gene disruption; 
(d) administering a test agent to the non-human transgenic animal of (a); and 
(e) determining whether the physiological characteristic associated with gene disruption is modulated.

51. The method of Claim 50, wherein the non-human transgenic animal exhibits at least one of the following 
physiological characteristics compared with gender matched wild-type littermates: increased anxiety-like response 
during open field testing; hyperactivity during open field testing; decreased anxiety during open field testing; 
decreased locomotor activity during open field testing; abnormal circadian rhythm during home-cage activity 
testing (low activity during the light phase; altered sleep/wake cycle); abnormal circadian rhythm during home-cage 
activity testing including decreased ambulatory counts; hypoactivity with no circadian rhythm; abnormal circadian 
rhythm during home-cage activity testing including increased ambulatory counts; decreased rearing; increased 
sensitivity to stress induced hyperthermia (increased anxiety); impaired motor coordination during inverted screen 
testing; head tilt and retropulsion; increased prepulse inhibition response indicating enhanced sensorimotor 
gating/attention; decreased startle response during prepulse inhibition testing; no startle response indicating 
deafness or impaired hearing; decreased prepulse inhibition with impaired sensorimotor gating/attention; increased 
lateness to respond in hot plate testing; decreased latency to respond in hot plate testing; ophthalmological 
abnormalities; impaired vision; white deposits of optic disc region; ocular infection and neutrophilia; bilateral optic 
disc lesion; decreased tear production; decreased heart rate; increased mean systolic blood pressure; decreased 
mean systolic blood pressure; increased mean fasting serum glucose levels; decreased mean serum glucose levels; 
increased mean serum cholesterol levels; decreased mean serum cholesterol levels; increased mean serum 
triglyceride levels; decreased mean serum triglyceride levels; impaired glucose tolerance; increased mean serum 
albumin, alanine amino transferase and phosphorus levels; increased mean serum alkaline phosphatase levels; 
urinary nitrites present; increased total white blood cell (WBC) count; decreased total white blood cell (WBC) 
count and absolute neutrophil count; increased mean absolute neutrophil count; increased mean absolute 
lymphocyte count; increased mean platelet count; increased mean red cell distribution width; decreased mean 
platelet count; reduced percentage of CD4 spleen thymocytes; decreased percentages of CD4 cells in the periphery 
resulting in increased percentages of B cells in lymph organs; CD4 cells exhibit a more activated/memory 
phenotype (CD62Llow, CD44hi); developmental defect in CD4+ cells; decreased percentages of CD4 cells and 
increased percentages of B cells in blood; decreased percentages of CD4 cells and increased percentages of B cells 
in tissues; increase in percentages of B cells in Peyer’s patches; decreased germinal center, isotype-switched B 
cells in Peyer’s patches (CD38low; IgM negative); decreased CD23 intensity in spleen; increased mean percentages 
of B220 Med/CD23- cells and B220+/CD11b-Low/CD23- cells in peritoneal lavage; increased mean percentages 
of B cells in peripheral blood; decreased CD4 and CD8 T cells and increased B cells; increase in peritoneal B cells; 

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reduction in CD11b-Hi cells in peritoneal cavity; decreased mean CD4 to CD8 ratio in spleen; decreased CD8 cells; decreased mean percentages of B220+/CD23+ cells and B220+/CD11bLow/CD23- cells in peritoneal lavage; increased mean serum IgG1 response to ovalbumin challenge; increased mean serum IgG2a response to ovalbumin challenge; increased mean serum IL-6 response to LPS challenge; increased mean serum TNF alpha response to LPS challenge; increased mean serum MCP-1 response to LPS challenge; increased mean serum IgM level; increased mean serum IgA; increase mean serum IgG1; increased mean serum IgG2a; increased mean serum IgG2b; decreased mean serum IgG1 response to ovalbumin challenge; decreased mean serum IgG2a response to ovalbumin challenge; failure in ovalbumin response; decreased mean serum IgA level; decreased mean serum IgG2a level; decreased skin fibroblast proliferation rate; increased mean percent of total body fat and total fat mass; increased mean body weight; increased mean body length; increased total tissue mass (TTM); increased bone mineral density (BMD); increased in bone mineral content (BMC); increased femoral midshaft cortical thickness; decreased mean percent of total body fat and total fat mass; decreased mean body weight; decreased mean body length; decreased mean body weight and length in heterozygotes; decreased total tissue mass (TTM); decreased lean body mass (LBM); decreased femoral bone mineral density (BMD); decreased vertebral bone mineral density (BMD); decreased bone mineral density (BMD) in total body; decreased bone mineral content (BMC); decreased bone mineral density index; decreased volumetric bone mineral density (vBMD); decreased mean femoral midshaft cortical thickness; decreased mean femoral midshaft cross-sectional area; decreased mean vertebral trabecular bone volume, number and connectivity density; osteopetrosis; osteoporosis; moderate kidney hydronephrosis; hydrocephalus; enlarged liver; induced in activated T cells; induced in activated NK cells and dendritic cells; myeloid B cell expression; hyperplasia of sebaceous glands and multifocal hyperplasia of the epidermis (acanthosis and hyperkeratosis); moderate dermatitis; increased extramedullary hematopoiesis in liver and spleen; myeloid hyperplasia of the bone marrow; encephalitis due to Group B streptococcus; meningitis due to E. Coli infection; lymphocytic infiltrates in salivary glands, pancreas and lungs; poor breeders requiring foster mothers; decreased litter size; homozygous mice were small and dehydrated; vacuolar degeneration of testes resulting in decreased sperm production and infertility; defective spermatogenesis in the testes; hypospermatemia and defective spermatogenesis in the epididymis; male infertility; decreased testes weight; growth retardation; small mice and failure to thrive; reduced viability; reduced viability with situs inverus; and homozygous embryonic lethality.

52. An agent identified by the method of Claim 50.

53. The agent of Claim 52 which is an agonist or antagonist of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide.

54. The agent of Claim 53, wherein the agonist is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-
PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

55. The agent of Claim 53, wherein the antagonist is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

56. A method of identifying an agent which modulates a behavior associated with a disruption of the gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising:

(a) providing a non-human transgenic animal whose genome comprises a disruption of the gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide;
(b) observing the behavior exhibited by the non-human transgenic animal of (a);
(c) comparing the observed behavior of (b) with that of a gender matched wild-type animal, wherein the observed behavior exhibited by the non-human transgenic animal that differs from the observed behavior exhibited by the wild-type animal is identified as a behavior associated with gene disruption;
(d) administering a test agent to the non-human transgenic animal of (a); and
(e) determining whether the agent modulates the behavior associated with gene disruption.

57. The method of Claim 56, wherein the behavior is an increased anxiety-like response during open field activity testing.

58. The method of Claim 56, wherein the behavior is a decreased anxiety-like response during open field activity testing.

59. The method of Claim 56, wherein the behavior is an abnormal circadian rhythm during home-cage activity testing.

60. The method of Claim 56, wherein the behavior is an enhanced motor coordination during inverted screen
61. The method of Claim 56, wherein the behavior is an impaired motor coordination during inverted screen testing.

62. The method of Claim 56, wherein the behavior is depression, generalized anxiety disorders, attention deficit disorder, sleep disorder, hyperactivity disorder, obsessive compulsive disorder, schizophrenia, cognitive disorders, hyperalgesia or sensory disorders.

63. An agent identified by the method of Claim 56.

64. The agent of Claim 63 which is an agonist or antagonist of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide.

65. The agent of Claim 64, wherein the agonist is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

66. The agent of Claim 64, wherein the antagonist is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

67. A method of identifying an agent that ameliorates or modulates a neurological disorder; a cardiovascular, endothelial or angiogenic disorder; an eye abnormality; an immunological disorder; an oncological disorder; a bone metabolic abnormality or disorder; a lipid metabolic disorder; or a developmental abnormality associated with a disruption in the gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising:
(a) providing a non-human transgenic animal whose genome comprises a disruption of the gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide;

(b) administering a test agent to said non-human transgenic animal; and

(c) determining whether said test agent ameliorates or modulates the neurological disorder; cardiovascular, endothelial or angiogenic disorder; eye abnormality; immunological disorder; oncological disorder; bone metabolic abnormality or disorder; lipid metabolic disorder; or developmental abnormality in the non-human transgenic animal.

68. The method of Claim 67, wherein the neurological disorder is an increased anxiety-like response during open field activity testing.

69. The method of Claim 67, wherein the neurological disorder is a decreased anxiety-like response during open field activity testing.

70. The method of Claim 67, wherein the neurological disorder is an abnormal circadian rhythm during home-cage activity testing.

71. The method of Claim 67, wherein the neurological disorder is an enhanced motor coordination during inverted screen testing.

72. The method of Claim 67, wherein the neurological disorder is an impaired motor coordination during inverted screen testing.

73. The method of Claim 73, wherein the neurological disorder is depression, generalized anxiety disorders, attention deficit disorder, sleep disorder, hyperactivity disorder, obsessive compulsive disorder, schizophrenia, cognitive disorders, hyperalgesia or sensory disorders.

74. The method of Claim 67, wherein the eye abnormality is a retinal abnormality.

75. The method of Claim 67, wherein the eye abnormality is consistent with vision problems or blindness.

76. The method of Claim 74, wherein the retinal abnormality is consistent with retinitis pigmentosa.

77. The method of Claim 74, wherein the retinal abnormality is characterized by retinal degeneration or retinal dysplasia.

78. The method of Claim 74, wherein the retinal abnormality is consistent with retinal dysplasia, various retinopathies, including retinopathy of prematurity, retrolental fibroplasia, neovascular glaucoma, age-related macular degeneration, diabetic macular edema, corneal neovascularization, corneal graft neovascularization, corneal graft rejection, retinal/choroidal neovascularization, neovascularization of the angle (rubeosis), ocular neovascular disease, vascular restenosis, arteriovenous malformations (AVM), menigioma, hemangioma, angiofibroma, thyroid hyperplasias (including Grave's disease), corneal and other tissue transplantation, retinal artery obstruction or occlusion; retinal degeneration causing secondary atrophy of the retinal vasculature, retinitis pigmentosa, macular dystrophies, Stargardt's disease, congenital stationary night blindness, choroideremia, gyrate atrophy, Leber's congenital amaurosis, retinoschisis disorders, Wagner's syndrome, Usher syndromes, Zellweger syndrome, Saldino-Mainzer syndrome, Senior-Loken syndrome, Bardet-Biedl syndrome, Alport's syndrome,

79. The method of Claim 67, wherein the eye abnormality is a cataract.

80. The method of Claim 79, wherein the cataract is a systemic disease such as human Down's syndrome, Hallerman-Streiff syndrome, Lowe syndrome, galactosemia, Marfan syndrome, Trisomy 15-15, Alport syndrome, myotonic dystrophy, Fabry disease, hypoparathyroidism or Conradi syndrome.

81. The method of Claim 67, wherein the developmental abnormality comprises embryonic lethality or reduced viability.

82. The method of Claim 67, wherein the cardiovascular, endothelial or angiogenic disorders are arterial diseases, such as diabetes mellitus; papilledema; optic atrophy; atherosclerosis; angina; myocardial infarctions such as acute myocardial infarctions, cardiac hypertrophy, and heart failure such as congestive heart failure; hypertension; inflammatory vasculitides; Reynaud's disease and Reynaud's phenomenon; aneurysms and arterial restenosis; venous and lymphatic disorders such as thrombophlebitis, lymphangitis, and lymphedema; peripheral vascular disease; cancer such as vascular tumors, e.g., hemangioma (capillary and cavernous), glomus tumors, telangiectasia, basillary angiomatosis, hemangioendothelioma, angiosarcoma, haemangiopericytoma, Kaposi's sarcoma, lymphangiomia, and lymphangiosarcroma; tumor angiogenesis; trauma such as wounds, burns, and other injured tissue, implant fixation, scarring; ischemia reperfusion injury; rheumatoid arthritis; cerebrovascular disease; renal diseases such as acute renal failure, or osteoporosis.

83. The method of Claim 67, wherein the immunological disorders are systemic lupus erythematosus; rheumatoid arthritis; juvenile chronic arthritis; spondyloarthopathies; systemic sclerosis (scleroderma); idiopathic inflammatory myopathies (dermatomyositis, polymyositis); Sjögren's syndrome; systemic vasculitis; sarcoidosis; autoimmune hemolytic anemia (immune pancytopenia, paroxysmal nocturnal hemoglobinuria); autoimmune thrombocytopenia (idiopathic thrombocytopenic purpura, immune-mediated thrombocytopenia); thyroiditis (Grave's disease, Hashimoto's thyroiditis, juvenile lymphocytic thyroiditis, atrophic thyroiditis); diabetes mellitus; immune-mediated renal disease (glomerulonephritis, tubulointerstitial nephritis); demyelinating diseases of the central and peripheral nervous systems such as multiple sclerosis, idiopathic demyelinating polynuropathy or Guillain-Barré syndrome, and chronic inflammatory demyelinating polynuropathy; hepatobiliary diseases such as infectious hepatitis (hepatitis A, B, C, D, E and other non-hepatotropic viruses), autoimmune chronic active hepatitis, primary biliary cirrhosis, granulomatous hepatitis, and sclerosing cholangitis; inflammatory bowel disease (ulcerative colitis; Crohn's disease); gluten-sensitive enteropathy, and Whipple's disease; autoimmune or immune-mediated skin diseases including bullous skin diseases, erythema multiforme and contact dermatitis, psoriasis; allergic diseases such as asthma, allergic rhinitis, atopic dermatitis, food hypersensitivity and urticaria; immunologic diseases of the lung such as eosinophilic pneumonia, idiopathic pulmonary fibrosis and hypersensitivity pneumonitis; or transplantation associated diseases including graft rejection and graft-versus-host disease.
84. The method of Claim 67, wherein said bone metabolic abnormality or disorder is arthritis, osteoporosis or osteopenosis.

85. The method of Claim 67, wherein the non-human transgenic animal exhibits at least one of the following physiological characteristics compared with gender matched wild-type littermates: increased anxiety-like response during open field testing; hyperactivity during open field testing; decreased anxiety during open field testing; decreased locomotor activity during open field testing; abnormal circadian rhythm during home-cage activity testing (low activity during the light phase; altered sleep/wake cycle); abnormal circadian rhythm during home-cage activity testing including decreased ambulatory counts; hypoactivity with no circadian rhythm; abnormal circadian rhythm during home-cage activity testing including increased ambulatory counts; decreased rearing; increased sensitivity to stress induced hyperthermia (increased anxiety); impaired motor coordination during inverted screen testing; head tilt and retropulsion; increased prepulse inhibition response indicating enhanced sensorimotor gating/attention; decreased startle response during prepulse inhibition testing; no startle response indicating deafness or impaired hearing; decreased prepulse inhibition with impaired sensorimotor gating/attention; increased latency to respond in hot plate testing; decreased latency to respond in hot plate testing; ophthalmological abnormalities; impaired vision; white deposits of optic disc region; ocular infection and neutrophilia; bilateral optic disc lesion; decreased tear production; decreased heart rate; increased mean systolic blood pressure; decreased mean systolic blood pressure; increased mean fasting serum glucose levels; decreased mean serum glucose levels; increased mean serum cholesterol levels; decreased mean serum cholesterol levels; increased mean serum triglyceride levels; decreased mean serum triglyceride levels; impaired glucose tolerance; increased mean serum albumin, alanine amino transferase and phosphorus levels; increased mean serum alkaline phosphatase levels; urinary nitrites present; increased total white blood cell (WBC) count; decreased total white blood cell (WBC) count and absolute neutrophil count; increased mean absolute neutrophil count; increased mean absolute lymphocyte count; increased mean platelet count; increased mean red cell distribution width; decreased mean platelet count; reduced percentage of CD4 spleen thymocytes; decreased percentages of CD4 cells in the periphery resulting in increased percentages of B cells in lymph organs; CD4 cells exhibit a more activated/memory phenotype (CD62Llow, CD44hi); developmental defect in CD4+ cells; decreased percentages of CD4 cells and increased percentages of B cells in blood; decreased percentages of CD4 cells and increased percentages of B cells in tissues; increase in percentages of B cells in Peyer’s patches; decreased germinal center, isotype-switched B cells in Peyer’s patches (CD38low; IgM negative); decreased CD23 intensity in spleen; increased mean percentages of B220 Med/CD23- cells and B220+/CD11b-Low/CD23- cells in peritoneal lavage; increased mean percentages of B cells in peripheral blood; decreased CD4 and CD8 T cells and increased B cells; increase in peritoneal B cells; reduction in CD11b-Hi cells in peritoneal cavity; decreased mean CD4 to CD8 ratio in spleen; decreased CD8 cells; decreased mean percentages of B220+/CD23+ cells and B220+/CD11bLow/CD23- cells in peritoneal lavage; increased mean serum IgG1 response to ovalbumin challenge; increased mean serum IgG2a response to ovalbumin challenge; increased mean serum IL-6 response to LPS challenge; increased mean serum TNF alpha response to LPS challenge; increased mean serum MCP-1 response to LPS challenge; increased mean serum IgM level; increased mean serum IgA; increase mean serum IgG1; increased mean serum IgG2a; increased mean serum IgG2b; decreased mean serum IgG1 response to ovalbumin challenge; decreased mean serum IgG2a response to ovalbumin challenge; failure in ovalbumin response; decreased mean serum IgA level; decreased mean serum
IgG2a level; decreased skin fibroblast proliferation rate; increased mean percent of total body fat and total fat mass; increased mean body weight; increased mean body length; increased total tissue mass (TTM); increased bone mineral density (BMD); increase in bone mineral content (BMC); increased mean femoral midshaft cortical thickness; decreased mean percent of total body fat and total fat mass; decreased mean body weight; decreased mean body length; decreased mean body weight and length in heterozygotes; decreased total tissue mass (TTM); decreased lean body mass (LBM); decreased femoral bone mineral density (BMD); decreased vertebral bone mineral density (BMD); decreased bone mineral density (BMD) in total body; decreased bone mineral content (BMC); decreased bone mineral density index; decreased volumetric bone mineral density (vBMD); decreased mean femoral midshaft cortical thickness; decreased mean femoral midshaft cross-sectional area; decreased mean vertebral trabecular bone volume, number and connectivity density; osteopetrosis; osteoporosis; moderate kidney hydronephrosis; hydrocephalus; enlarged liver; induced in activated T cells; induced in activated NK cells and dendritic cells; myeloid B cell expression; hyperplasia of sebaceous glands and multifocal hyperplasia of the epidermis (acanthosis and hyperkeratosis); moderate dermatitis; increased extramedullary hematopoiesis in liver and spleen; myeloid hyperplasia of the bone marrow; encephalitis due to Group B streptococcus; meningitis due to *E. Coli* infection; lymphocytic infiltrates in salivary glands, pancreas and lungs; poor breeders requiring foster mothers; decreased litter size; homozygous mice were small and dehydrated; vacuolar degeneration of testes resulting in decreased sperm production and infertility; defective spermatogenesis in the testes; hypospermatia and defective spermatogenesis in the epididymis; male infertility; decreased testes weight; growth retardation; small mice and failure to thrive; reduced viability; reduced viability with sitsus inversus; and homozygous embryonic lethality.

86. An agent identified by the method of Claim 67.

87. The agent of Claim 86 which is an agonist or antagonist of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide.

88. The agent of Claim 87 wherein the agonist is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

89. The agent of Claim 87, wherein the antagonist is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.
PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

90. A therapeutic agent identified by the method of Claim 67.

91. A method of identifying an agent that modulates the expression of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1115, PRO1117, PRO1119, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising:

(a) contacting a test agent with a host cell expressing a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1115, PRO1117, PRO1119, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide; and

(b) determining whether the test agent modulates the expression of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1115, PRO1117, PRO1119, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide by the host cell.

92. An agent identified by the method of Claim 91.

93. The agent of Claim 92 which is an agonist or antagonist of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1115, PRO1117, PRO1119, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide.

94. The agent of Claim 93, wherein the agonist is an anti-PRO218, anti-PRO228, anti-PRO271, anti-

95. The agent of Claim 93, wherein the antagonist is an anti-PRO218, anti-PRO228, anti-PRO271, anti-

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PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-
PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474,
anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-
PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-
PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO99048, anti-PRO28694, anti-
PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465
or anti-PRO346 antibody.

96. A method of evaluating a therapeutic agent capable of affecting a condition associated with a
disruption of a gene which encodes for a PROW218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305,
PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016,
PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879,
PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013,
PRO99048, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or
PRO346 polypeptide, the method comprising:

(a) providing a non-human transgenic animal whose genome comprises a disruption of the gene which
encodes for the PROW218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655,
PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069,
PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329,
PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99048, PRO28694, PRO16089,
PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide;

(b) measuring a physiological characteristic of the non-human transgenic animal of (a);

(c) comparing the measured physiological characteristic of (b) with that of a gender matched wild-type
animal, wherein the physiological characteristic of the non-human transgenic animal that differs from the
physiological characteristic of the wild-type animal is identified as a condition resulting from the gene disruption
in the non-human transgenic animal;

(d) administering a test agent to the non-human transgenic animal of (a); and

(e) evaluating the effects of the test agent on the identified condition associated with gene disruption
in the non-human transgenic animal.

97. The method of Claim 96, wherein the condition is a neurological disorder; a cardiovascular, endothelial
or angiogenic disorder; an eye abnormality; an immunological disorder; an oncological disorder; a bone metabolic
abnormality or disorder; a lipid metabolic disorder; or a developmental abnormality.

98. A therapeutic agent identified by the method of Claim 96.

99. The therapeutic agent of Claim 98 which is an agonist or antagonist of a PRO218, PRO228, PRO271,
PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940,
PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130,
PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859,
PRO9864, PRO9904, PRO9907, PRO10013, PRO99048, PRO28694, PRO16089, PRO19563, PRO19675,
PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide.

100. The therapeutic agent of Claim 99, wherein the agonist is an anti-PRO218, anti-PRO228, anti-PRO271,
anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

101. The therapeutic agent of Claim 99, wherein the antagonist is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

102. A pharmaceutical composition comprising the therapeutic agent of Claim 98.

103. A method of treating or preventing or ameliorating a neurological disorder; cardiovascular, endothelial or angiogenic disorder; immunological disorder; oncological disorder; bone metabolic abnormality or disorder, or embryonic lethality associated with the disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising administering to a subject in need of such treatment whom may already have the disorder, or may be prone to have the disorder or may be in whom the disorder is to be prevented, a therapeutically effective amount of the therapeutic agent of Claim 94, or agonists or antagonists thereof, thereby effectively treating or preventing or ameliorating said disorder.

104. The method of Claim 103, wherein the neurological disorder is an increased anxiety-like response during open field activity testing.

105. The method of Claim 103, wherein the neurological disorder is a decreased anxiety-like response during open field activity testing.

106. The method of Claim 103, wherein the neurological disorder is an abnormal circadian rhythm during home-cage activity testing.

107. The method of Claim 103, wherein the neurological disorder is an enhanced motor coordination during inverted screen testing.

108. The method of Claim 103, wherein the neurological disorder is an impaired motor coordination during inverted screen testing.
109. The method of Claim 103, wherein the neurological disorder is depression, generalized anxiety disorders, attention deficit disorder, sleep disorder, hyperactivity disorder, obsessive compulsive disorder, schizophrenia, cognitive disorders, hyperalgesia or sensory disorders.

110. The method of Claim 103, wherein the eye abnormality is a retinal abnormality.

111. The method of Claim 103, wherein the eye abnormality is consistent with vision problems or blindness.

112. The method of Claim 110, wherein the retinal abnormality is consistent with retinitis pigmentosa.

113. The method of Claim 110, wherein the retinal abnormality is characterized by retinal degeneration or retinal dysplasia.

114. The method of Claim 110, wherein the retinal abnormality is consistent with retinal dysplasia, various retinopathies, including retinopathy of prematurity, retrolental fibroplasia, neovascular glaucoma, age-related macular degeneration, diabetic macular edema, corneal neovascularization, corneal graft neovascularization, corneal graft rejection, retinal/choroidal neovascularization, neovascularization of the angle (rubeosis), ocular neovascular disease, vascular restenosis, arteriovenous malformations (AVM), meningioma, hemangioma, angiofibroma, thyroid hyperplasias (including Grave's disease), corneal and other tissue transplantation, retinal artery obstruction or occlusion; retinal degeneration causing secondary atrophy of the retinal vasculature, retinitis pigmentosa, macular dystrophies, Stargardt's disease, congenital stationary night blindness, choroideremia, gyrate atrophy, Leber's congenital amaurosis, retinoschisis disorders, Wagner's syndrome, Usher syndromes, Zellweger syndrome, Saldivo-Mainzer syndrome, Senior-Loken syndrome, Bardet-Biedl syndrome, Alport's syndrome, Alstrom's syndrome, Cockayne's syndrome, dysplasia spondyloepiphysaria congenita, Flynn-Aird syndrome, Friedreich ataxia, Hallgren syndrome, Marshall syndrome, Albers-Schonberg disease, Refsum's disease, Kearns-Sayre syndrome, Waardenburg's syndrome, Alagille syndrome, myotonic dystrophy, olivopontocerebellar atrophy, Pierre-Marie dunsdrome, Stickler syndrome, carotinemia, cystinosis, Wolfram syndrome, Bassen-Kornzweig syndrome, abetalipoproteinemia, incontinentia pigmenti, Batten's disease, mucopolysaccharidoses, homocystinuria, or marnosidosis.

115. The method of Claim 103, wherein the eye abnormality is a cataract.

116. The method of Claim 115, wherein the cataract is a systemic disease such as human Down's syndrome, Hallerman-Streiff syndrome, Lowe syndrome, galactosemia, Marfan syndrome, Trismoy 13-15, Alport syndrome, myotonic dystrophy, Fabry disease, hypoparathyroidism or Conradi syndrome.

117. The method of Claim 103, wherein the developmental abnormality comprises embryonic lethality or reduced viability.

118. The method of Claim 103, wherein the cardiovascular, endothelial or angiogenic disorders are arterial diseases, such as diabetes mellitus; papilledema; optic atrophy; atherosclerosis; angina; myocardial infarctions such as acute myocardial infarctions, cardiac hypertrophy, and heart failure such as congestive heart failure; hypertension; inflammatory vasculitides; Reynaud's disease and Reynaud's phenomenon; aneuysms and arterial restenosis; venous and lymphatic disorders such as thrombophlebitis, lymphangitis, and lymphedema; peripheral vascular disease; cancer such as vascular tumors, e.g., hemangioma (capillary and cavernous), glomus tumors, telangiectasia, bacillary angiomatosis, hemangioendothelioma, angiosarcoma, haemangiopericytoma, Kaposi's sarcoma, lymphangioma, and lymphangiosarcoma; tumor angiogenesis; trauma such as wounds, burns, and other injured tissue, implant fixation, scarring; ischemia reperfusion injury; rheumatoid arthritis; cerebrovascular disease;
renal diseases such as acute renal failure, or osteoporosis.

119. The method of Claim 103, wherein the immunological disorders are systemic lupus erythematosis; rheumatoid arthritis; juvenile chronic arthritis; spondyloarthropathies; systemic sclerosis (scleroderma); idiopathic inflammatory myopathies (dermatomyositis, polymyositis); Sjögren's syndrome; systemic vasculitis; sarcoidosis; autoimmune hemolytic anemia (immune pancytopenia, paroxysmal nocturnal hemoglobinuria); autoimmune thrombocytopenia (idiopathic thrombocytopenic purpura, immune-mediated thrombocytopenia); thyroiditis (Grave's disease, Hashimoto's thyroiditis, juvenile lymphocytic thyroiditis, atrophic thyroiditis); diabetes mellitus; immune-mediated renal disease (glomerulonephritis, tubulointerstitial nephritis); demyelinating diseases of the central and peripheral nervous systems such as multiple sclerosis, idiopathic demyelinating polyneuropathy or Guillain-Barré syndrome, and chronic inflammatory demyelinating polyneuropathy; hepatobiliary diseases such as infectious hepatitis (hepatitis A, B, C, D, E and other non-hepatotropic viruses), autoimmune chronic active hepatitis, primary biliary cirrhosis, granulomatous hepatitis, and sclerosing cholangitis; inflammatory bowel disease (ulcerative colitis; Crohn's disease); gluten-sensitive enteropathy, and Whipple's disease; autoimmune or immune-mediated skin diseases including bullous skin diseases, erythema multiforme and contact dermatitis, psoriasis; allergic diseases such as asthma, allergic rhinitis, atopic dermatitis, food hypersensitivity and urticaria; immunologic diseases of the lung such as eosinophilic pneumonia, idiopathic pulmonary fibrosis and hypersensitivity pneumonitis; or transplantation associated diseases including graft rejection and graft-versus-host disease.

120. The method of Claim 103, wherein said bone metabolic abnormality or disorder is arthritis, osteoporosis or osteopetrosis.

121. A method of identifying an agent that ameliorates or modulates a neurological disorder; a cardiovascular, endothelial or angiogenic disorder; an eye abnormality; an immunological disorder; an oncological disorder; a bone metabolic abnormality or disorder; a lipid metabolic disorder; or a developmental abnormality associated with a disruption in the gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1169, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO990948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising:

(a) providing a non-human transgenic animal cell culture, each cell of said culture comprising a disruption of the gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO990948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide;

(b) administering a test agent to said cell culture; and

(c) determining whether said test agent ameliorates or modulates the neurological disorder; cardiovascular, endothelial or angiogenic disorder; eye abnormality; immunological disorder; oncological disorder;
bone metabolic abnormality or disorder; lipid metabolic disorder; or developmental abnormality in said cell
culture.

122. The method of Claim 121, wherein the neurological disorder is an increased anxiety-like response during
open field activity testing.

123. The method of Claim 121, wherein the neurological disorder is a decreased anxiety-like response during
open field activity testing.

124. The method of Claim 121, wherein the neurological disorder is an abnormal circadian rhythm during
home-cage activity testing.

125. The method of Claim 121, wherein the neurological disorder is an enhanced motor coordination during
inverted screen testing.

126. The method of Claim 121, wherein the neurological disorder is an impaired motor coordination during
inverted screen testing.

127. The method of Claim 121, wherein the neurological disorder is depression, generalized anxiety disorders,
attention deficit disorder, sleep disorder, hyperactivity disorder, obsessive compulsive disorder, schizophrenia,
cognitive disorders, hyperalgiesia or sensory disorders.

128. The method of Claim 121, wherein the eye abnormality is a retinal abnormality.

129. The method of Claim 121, wherein the eye abnormality is consistent with vision problems or blindness.

130. The method of Claim 128, wherein the retinal abnormality is consistent with retinitis pigmentosa.

131. The method of Claim 128, wherein the retinal abnormality is characterized by retinal degeneration or
retinal dysplasia.

132. The method of Claim 128, wherein the retinal abnormality is consistent with retinal dysplasia, various
retinopathies, including retinopathy of prematurity, retrolental fibroplasia, neovascular glaucoma, age-related
macular degeneration, diabetic macular edema, corneal neovascularization, corneal graft neovascularization,
corneal graft rejection, retinal/choroidal neovascularization, neovascularization of the angle (rubeosis), ocular
neovascular disease, vascular restenosis, arteriovenous malformations (AVM), meningioma, hemangioma,
angiofibroma, thyroid hyperplasias (including Grave's disease), corneal and other tissue transplantation, retinal
artery obstruction or occlusion; retinal degeneration causing secondary atrophy of the retinal vasculature, retinitis
pigmentosa, macular dystrophies, Stargardt's disease, congenital stationary night blindness, choroideremia, gyrate
atrophy, Leber’s congenital amaurosis, retinoschisis disorders, Wagner’s syndrome, Usher syndromes, Zellweger
syndrome, Saldino-Mainzer syndrome, Senior-Loken syndrome, Bardet-Biedl syndrome, Alport's syndrome,
Alstrom's syndrome, Cockayne's syndrome, dysplaisa spondyloepiphysaria congenita, Flynn-Aird syndrome,
Friedreich ataxia, Hallgren syndrome, Marshall syndrome, Albers-Schoenberg disease, Refsum's disease,
Kearns-Sayre syndrome, Waardenburg's syndrome, Alagile syndrome, myotonic dystrophy, olivopontocerebellar
atrophy, Pierre-Marie dunsdrome, Stickler syndrome, carotinemia, cystinosis, Wolfram syndrome,
Bassen-Kornzweig syndrome, abetalipoproteinemia, incontinentia pigmenti, Batten's disease,
mucopolysaccharidoses, homocystinuria, or mnoonosisis.

133. The method of Claim 121, wherein the eye abnormality is a cataract.

134. The method of Claim 133, wherein the cataract is a systemic disease such as human Down’s syndrome,
Hallerman-Streiff syndrome, Lowe syndrome, galactosemia, Marfan syndrome, Trismoy 13-15, Alport syndrome,
myotonic dystrophy, Fabry disease, hypoparathyroidism or Conrad syndrome.

135. The method of Claim 121, wherein the developmental abnormality comprises embryonic lethality or reduced viability.

136. The method of Claim 121, wherein the cardiovascular, endothelial or angiogenic disorders are arterial diseases, such as diabetes mellitus; papilledema; optic atrophy; atherosclerosis; angina; myocardial infarctions such as acute myocardial infarctions, cardiac hypertrophy, and heart failure such as congestive heart failure; hypertension; inflammatory vasculitides; Reynaud's disease and Reynaud's phenomenon; aneurysms and arterial restenosis; venous and lymphatic disorders such as thrombophlebitis, lymphangitis, and lymphedema; peripheral vascular disease; cancer such as vascular tumors, e.g., hemangioma (capillary and cavernous), glomus tumors, telangiectasia, bacillary angiomatosis, hemangioendothelioma, angiosarcoma, haemangioepicytoma, Kaposi's sarcoma, lymphangioma, and lymphangiosarcoma; tumor angiogenesis; trauma such as wounds, burns, and other injured tissue, implant fixation, scarring; ischemia reperfusion injury; rheumatoid arthritis; cerebrovascular disease; renal diseases such as acute renal failure, or osteoporosis.

137. The method of Claim 121, wherein the immunological disorders are systemic lupus erythematosus; rheumatoid arthritis; juvenile chronic arthritis; spondyloarthopathies; systemic sclerosis (scleroderma); idiopathic inflammatory myopathies (dermatomyositis, polymyositis); Sjögren's syndrome; systemic vasculitis; sarcoidosis; autoimmune hemolytic anemia (immune pancytopenia, paroxysmal nocturnal hemoglobinuria); autoimmune thrombocytopenia (idiopathic thrombocytopenic purpura, immune-mediated thrombocytopenia); thyroiditis (Grave's disease, Hashimoto's thyroiditis, juvenile lymphocytic thyroiditis, atrophic thyroiditis); diabetes mellitus; immune-mediated renal disease (glomerulonephritis, tubulointerstitial nephritis); demyelinating diseases of the central and peripheral nervous systems such as multiple sclerosis, idiopathic demyelinating polyneuropathy or Guillain-Barré syndrome, and chronic inflammatory demyelinating polyneuropathy; hepatobiliary diseases such as infectious hepatitis (hepatitis A, B, C, D, E and other non-hepatotropic viruses), autoimmune chronic active hepatitis, primary biliary cirrhosis, granulomatous hepatitis, and sclerosing cholangitis; inflammatory bowel disease (ulcerative colitis; Crohn's disease); gluten-sensitive enteropathy, and Whipple's disease; autoimmune or immune-mediated skin diseases including bullous skin diseases, erythema multiforme and contact dermatitis, psoriasis; allergic diseases such as asthma, allergic rhinitis, atopic dermatitis, food hypersensitivity and urticaria; immunologic diseases of the lung such as eosinophilic pneumonia, idiopathic pulmonary fibrosis and hypersensitivity pneumonitis; or transplantation associated diseases including graft rejection and graft-versus-host disease.

138. The method of Claim 121, wherein said bone metabolic abnormality or disorder is arthritis, osteoporosis or osteopetrosis.

139. An agent identified by the method of Claim 121.

140. The agent of Claim 139 which is an agonist or antagonist of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide.
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The agent of Claim 140, wherein the agonist is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

The agent of Claim 140, wherein the antagonist is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

A therapeutic agent identified by the method of Claim 121.

A method of modulating a phenotype associated with a disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising administering to a subject whom may already have the phenotype, or may be prone to have the phenotype or may be in whom the phenotype is to be prevented, an effective amount of the agent of Claim 46, or agonists or antagonists thereof, thereby effectively modulating the phenotype.

A method of modulating a physiological characteristic associated with a disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising administering to a subject whom may already exhibit the physiological characteristic, or may be prone to exhibit the physiological characteristic or may be in whom the physiological characteristic is to be prevented, an effective amount of the agent of Claim 52, or agonists or antagonists thereof, thereby effectively modulating the physiological characteristic.

A method of modulating a behavior associated with a disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising administering to a subject whom may already exhibit the physiological characteristic, or may be prone to exhibit the physiological characteristic or may be in whom the physiological characteristic is to be prevented, an effective amount of the agent of Claim 52, or agonists or antagonists thereof, thereby effectively modulating the physiological characteristic.
PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO2084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising administering to a subject whom may already exhibit the behavior, or may be prone to exhibit the behavior or may be in whom the exhibited behavior is to be prevented, an effective amount of the agent of Claim 63, or agonists or antagonists thereof, thereby effectively modulating the behavior.

147. A method of modulating the expression of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO19563, PRO19675, PRO2084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising administering to a host cell expressing said PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO2084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, an effective amount of the agent of Claim 92, or agonists or antagonists thereof, thereby effectively modulating the expression of said polypeptide.

148. A method of modulating a condition associated with a disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO2084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising administering to a subject whom may have the condition, or may be prone to have the condition or may be in whom the condition is to be prevented, a therapeutically effective amount of the therapeutic agent of Claim 98, or agonists or antagonists thereof, thereby effectively modulating the condition.

149. A method of treating or preventing or ameliorating a neurological disorder; cardiovascular, endothelial or angiogenic disorder; immunological disorder; oncological disorder; bone metabolic abnormality or disorder, or embryonic lethality associated with the disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO2084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising administering to a non-human transgenic animal cell culture, each cell of said culture comprising a disruption of the gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO2084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide.
BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 shows a nucleotide sequence (SEQ ID NO:1) of a native sequence PRO218 cDNA, wherein SEQ ID NO:1 is a clone designated herein as “DNA30867-1335” (UNQ192).

Figure 2 shows the amino acid sequence (SEQ ID NO:2) derived from the coding sequence of SEQ ID NO:1 shown in Figure 1.

Figure 3 shows a nucleotide sequence (SEQ ID NO:3) of a native sequence PRO228 cDNA, wherein SEQ ID NO:3 is a clone designated herein as “DNA33092-1202” (UNQ202).

Figure 4 shows the amino acid sequence (SEQ ID NO:4) derived from the coding sequence of SEQ ID NO:3 shown in Figure 3.

Figure 5 shows a nucleotide sequence (SEQ ID NO:5) of a native sequence PRO271 cDNA, wherein SEQ ID NO:5 is a clone designated herein as “DNA39423-1182” (UNQ238).

Figure 6 shows the amino acid sequence (SEQ ID NO:6) derived from the coding sequence of SEQ ID NO:5 shown in Figure 5.

Figure 7 shows a nucleotide sequence (SEQ ID NO:7) of a native sequence PRO273 cDNA, wherein SEQ ID NO:7 is a clone designated herein as “DNA39523-1192” (UNQ240).

Figure 8 shows the amino acid sequence (SEQ ID NO:8) derived from the coding sequence of SEQ ID NO:7 shown in Figure 7.

Figure 9 shows a nucleotide sequence (SEQ ID NO:9) of a native sequence PRO295 cDNA, wherein SEQ ID NO:9 is a clone designated herein as “DNA38268-1188” (UNQ258).

Figure 10 shows the amino acid sequence (SEQ ID NO:10) derived from the coding sequence of SEQ ID NO:9 shown in Figure 9.

Figure 11 shows a nucleotide sequence (SEQ ID NO:11) of a native sequence PRO302 cDNA, wherein SEQ ID NO:11 is a clone designated herein as “DNA40370-1217” (UNQ265).

Figure 12 shows the amino acid sequence (SEQ ID NO:12) derived from the coding sequence of SEQ ID NO:11 shown in Figure 11.

Figure 13 shows a nucleotide sequence (SEQ ID NO:13) of a native sequence PRO305 cDNA, wherein SEQ ID NO:13 is a clone designated herein as “DNA40619-1220” (UNQ268).

Figure 14 shows the amino acid sequence (SEQ ID NO:14) derived from the coding sequence of SEQ ID NO:13 shown in Figure 13.

Figure 15 shows a nucleotide sequence (SEQ ID NO:15) of a native sequence PRO326 cDNA, wherein SEQ ID NO:15 is a clone designated herein as “DNA37140-1234” (UNQ287).

Figure 16 shows the amino acid sequence (SEQ ID NO:16) derived from the coding sequence of SEQ ID NO:15 shown in Figure 15.
Figure 17 shows a nucleotide sequence (SEQ ID NO:17) of a native sequence PRO386 cDNA, wherein SEQ ID NO:17 is a clone designated herein as "DNA45415-1318" (UNQ326).

Figure 18 shows the amino acid sequence (SEQ ID NO:18) derived from the coding sequence of SEQ ID NO:17 shown in Figure 17.

Figure 19 shows a nucleotide sequence (SEQ ID NO:19) of a native sequence PRO655 cDNA, wherein SEQ ID NO:19 is a clone designated herein as "DNA50960-1224" (UNQ360).

Figure 20 shows the amino acid sequence (SEQ ID NO:20) derived from the coding sequence of SEQ ID NO:19 shown in Figure 19.

Figure 21 shows a nucleotide sequence (SEQ ID NO:21) of a native sequence PRO162 cDNA, wherein SEQ ID NO:21 is a clone designated herein as "DNA56965-1356" (UNQ429).

Figure 22 shows the amino acid sequence (SEQ ID NO:22) derived from the coding sequence of SEQ ID NO:21 shown in Figure 21.

Figure 23 shows a nucleotide sequence (SEQ ID NO:23) of a native sequence PRO788 cDNA, wherein SEQ ID NO:23 is a clone designated herein as "DNA56405-1357" (UNQ430).

Figure 24 shows the amino acid sequence (SEQ ID NO:24) derived from the coding sequence of SEQ ID NO:23 shown in Figure 23.

Figure 25 shows a nucleotide sequence (SEQ ID NO:25) of a native sequence PRO792 cDNA, wherein SEQ ID NO:25 is a clone designated herein as "DNA6352-1358" (UNQ431).

Figure 26 shows the amino acid sequence (SEQ ID NO:26) derived from the coding sequence of SEQ ID NO:25 shown in Figure 25.

Figure 27 shows a nucleotide sequence (SEQ ID NO:27) of a native sequence PRO940 cDNA, wherein SEQ ID NO:27 is a clone designated herein as "DNA54002-1367" (UNQ477).

Figure 28 shows the amino acid sequence (SEQ ID NO:28) derived from the coding sequence of SEQ ID NO:27 shown in Figure 27.

Figure 29 shows a nucleotide sequence (SEQ ID NO:29) of a native sequence PRO941 cDNA, wherein SEQ ID NO:29 is a clone designated herein as "DNA53906-1368" (UNQ478).

Figure 30 shows the amino acid sequence (SEQ ID NO:30) derived from the coding sequence of SEQ ID NO:29 shown in Figure 29.

Figure 31 shows a nucleotide sequence (SEQ ID NO:31) of a native sequence PRO1004 cDNA, wherein SEQ ID NO:31 is a clone designated herein as "DNA57844-1410" (UNQ488).

Figure 32 shows the amino acid sequence (SEQ ID NO:32) derived from the coding sequence of SEQ ID NO:31 shown in Figure 31.

Figure 33 shows a nucleotide sequence (SEQ ID NO:33) of a native sequence PRO1012 cDNA, wherein SEQ ID NO:33 is a clone designated herein as "DNA56439-1376" (UNQ495).

Figure 34 shows the amino acid sequence (SEQ ID NO:34) derived from the coding sequence of SEQ ID NO:33 shown in Figure 33.

Figure 35 shows a nucleotide sequence (SEQ ID NO:35) of a native sequence PRO1016 cDNA, wherein SEQ ID NO:35 is a clone designated herein as "DNA56113-1378" (UNQ499).

Figure 36 shows the amino acid sequence (SEQ ID NO:36) derived from the coding sequence of SEQ ID NO:35 shown in Figure 35.
ID NO:35 shown in Figure 35.

Figure 37 shows a nucleotide sequence (SEQ ID NO:37) of a native sequence PRO474 cDNA, wherein
SEQ ID NO:37 is a clone designated herein as “DNA56045-1380” (UNQ502).

Figure 38 shows the amino acid sequence (SEQ ID NO:38) derived from the coding sequence of SEQ
ID NO:37 shown in Figure 37.

Figure 39 shows a nucleotide sequence (SEQ ID NO:39) of a native sequence PROS238 cDNA, wherein
SEQ ID NO:39 is a clone designated herein as “DNA257845” (UNQ503).

Figure 40 shows the amino acid sequence (SEQ ID NO:40) derived from the coding sequence of SEQ
ID NO:39 shown in Figure 39.

Figure 41 shows a nucleotide sequence (SEQ ID NO:41) of a native sequence PRO1069 cDNA, wherein
SEQ ID NO:41 is a clone designated herein as “DNA59211-1450” (UNQ526).

Figure 42 shows the amino acid sequence (SEQ ID NO:42) derived from the coding sequence of SEQ
ID NO:41 shown in Figure 41.

Figure 43 shows a nucleotide sequence (SEQ ID NO:43) of a native sequence PRO1111 cDNA, wherein
SEQ ID NO:43 is a clone designated herein as “DNA58721-1475” (UNQ554).

Figure 44 shows the amino acid sequence (SEQ ID NO:44) derived from the coding sequence of SEQ
ID NO:43 shown in Figure 43.

Figure 45 shows a nucleotide sequence (SEQ ID NO:45) of a native sequence PRO1113 cDNA, wherein
SEQ ID NO:45 is a clone designated herein as “DNA57254-1477” (UNQ556).

Figure 46 shows the amino acid sequence (SEQ ID NO:46) derived from the coding sequence of SEQ
ID NO:45 shown in Figure 45.

Figure 47 shows a nucleotide sequence (SEQ ID NO:47) of a native sequence PRO1130 cDNA, wherein
SEQ ID NO:47 is a clone designated herein as “DNA59814-1486” (UNQ567).

Figure 48 shows the amino acid sequence (SEQ ID NO:48) derived from the coding sequence of SEQ
ID NO:47 shown in Figure 47.

Figure 49 shows a nucleotide sequence (SEQ ID NO:49) of a native sequence PRO1195 cDNA, wherein
SEQ ID NO:49 is a clone designated herein as “DNA65412-1523” (UNQ608).

Figure 50 shows the amino acid sequence (SEQ ID NO:50) derived from the coding sequence of SEQ
ID NO:49 shown in Figure 49.

Figure 51 shows a nucleotide sequence (SEQ ID NO:51) of a native sequence PRO1271 cDNA, wherein
SEQ ID NO:51 is a clone designated herein as “DNA66309-1538” (UNQ641).

Figure 52 shows the amino acid sequence (SEQ ID NO:52) derived from the coding sequence of SEQ
ID NO:51 shown in Figure 51.

Figure 53 shows a nucleotide sequence (SEQ ID NO:53) of a native sequence PRO1865 cDNA, wherein
SEQ ID NO:53 is a clone designated herein as “DNA81757-2512” (UNQ856).

Figure 54 shows the amino acid sequence (SEQ ID NO:54) derived from the coding sequence of SEQ
ID NO:53 shown in Figure 53.

Figure 55 shows a nucleotide sequence (SEQ ID NO:55) of a native sequence PRO1879 cDNA, wherein
SEQ ID NO:55 is a clone designated herein as “DNA54009-2517” (UNQ863).
Figure 56 shows the amino acid sequence (SEQ ID NO:56) derived from the coding sequence of SEQ ID NO:55 shown in Figure 55.

Figure 57 shows a nucleotide sequence (SEQ ID NO:57) of a native sequence PRO3446 cDNA, wherein SEQ ID NO:57 is a clone designated herein as “DNA92219-2541” (UNQ1833).

Figure 58 shows the amino acid sequence (SEQ ID NO:58) derived from the coding sequence of SEQ ID NO:57 shown in Figure 57.

Figure 59 shows a nucleotide sequence (SEQ ID NO:59) of a native sequence PRO3543 cDNA, wherein SEQ ID NO:51 is a clone designated herein as “DNA86571-2551” (UNQ1835).

Figure 60 shows the amino acid sequence (SEQ ID NO:60) derived from the coding sequence of SEQ ID NO:59 shown in Figure 59.

Figure 61 shows a nucleotide sequence (SEQ ID NO:61) of a native sequence PRO4329 cDNA, wherein SEQ ID NO:61 is a clone designated herein as “DNA77629-2573” (UNQ1885).

Figure 62 shows the amino acid sequence (SEQ ID NO:62) derived from the coding sequence of SEQ ID NO:61 shown in Figure 61.

Figure 63 shows a nucleotide sequence (SEQ ID NO:63) of a native sequence PRO4352 cDNA, wherein SEQ ID NO:63 is a clone designated herein as “DNA87976-2593” (UNQ1906).

Figure 64 shows the amino acid sequence (SEQ ID NO:64) derived from the coding sequence of SEQ ID NO:63 shown in Figure 63.

Figure 65 shows a nucleotide sequence (SEQ ID NO:65) of a native sequence PRO5733 cDNA, wherein SEQ ID NO:65 is a clone designated herein as “DNA82343” (UNQ2453).

Figure 66 shows the amino acid sequence (SEQ ID NO:66) derived from the coding sequence of SEQ ID NO:65 shown in Figure 65.

Figure 67 shows a nucleotide sequence (SEQ ID NO:67) of a native sequence PRO9859 cDNA, wherein SEQ ID NO:67 is a clone designated herein as “DNA125170-2780” (UNQ3043).

Figure 68 shows the amino acid sequence (SEQ ID NO:68) derived from the coding sequence of SEQ ID NO:67 shown in Figure 67.

Figure 69 shows a nucleotide sequence (SEQ ID NO:69) of a native sequence PRO9864 cDNA, wherein SEQ ID NO:69 is a clone designated herein as “DNA125151-2784” (UNQ3048).

Figure 70 shows the amino acid sequence (SEQ ID NO:70) derived from the coding sequence of SEQ ID NO:69 shown in Figure 69.

Figure 71 shows a nucleotide sequence (SEQ ID NO:71) of a native sequence PRO9904 cDNA, wherein SEQ ID NO:71 is a clone designated herein as “DNA129549-2798” (UNQ3072).

Figure 72 shows the amino acid sequence (SEQ ID NO:72) derived from the coding sequence of SEQ ID NO:71 shown in Figure 71.

Figure 73 shows a nucleotide sequence (SEQ ID NO:73) of a native sequence PRO9907 cDNA, wherein SEQ ID NO:73 is a clone designated herein as “DNA142392-2800” (UNQ3075).

Figure 74 shows the amino acid sequence (SEQ ID NO:74) derived from the coding sequence of SEQ ID NO:73 shown in Figure 73.

Figure 75 shows a nucleotide sequence (SEQ ID NO:75) of a native sequence PRO10013 cDNA, wherein
SEQ ID NO: 75 is a clone designated herein as “DNA125181-2804” (UNQ5082).

Figure 76 shows the amino acid sequence (SEQ ID NO: 76) derived from the coding sequence of SEQ ID NO: 75 shown in Figure 75.

Figure 77 shows a nucleotide sequence (SEQ ID NO: 77) of a native sequence PRO90948 cDNA, wherein SEQ ID NO: 77 is a clone designated herein as “DNA336882” (UNQ5043).

Figure 78 shows the amino acid sequence (SEQ ID NO: 78) derived from the coding sequence of SEQ ID NO: 77 shown in Figure 77.

Figure 79 shows a nucleotide sequence (SEQ ID NO: 79) of a native sequence PRO28694 cDNA, wherein SEQ ID NO: 79 is a clone designated herein as “DNA184073” (UNQ5384).

Figure 80 shows the amino acid sequence (SEQ ID NO: 80) derived from the coding sequence of SEQ ID NO: 79 shown in Figure 79.

Figure 81 shows a nucleotide sequence (SEQ ID NO: 81) of a native sequence PRO16089 cDNA, wherein SEQ ID NO: 81 is a clone designated herein as “DNA150163-2842” (UNQ5782).

Figure 82 shows the amino acid sequence (SEQ ID NO: 82) derived from the coding sequence of SEQ ID NO: 81 shown in Figure 81.

Figure 83 shows a nucleotide sequence (SEQ ID NO: 83) of a native sequence PRO19563 cDNA, wherein SEQ ID NO: 83 is a clone designated herein as “DNA96861-2844” (UNQ5785).

Figure 84 shows the amino acid sequence (SEQ ID NO: 84) derived from the coding sequence of SEQ ID NO: 83 shown in Figure 83.

Figure 85 shows a nucleotide sequence (SEQ ID NO: 85) of a native sequence PRO19675 cDNA, wherein SEQ ID NO: 85 is a clone designated herein as “DNA131658-2875” (UNQ5835).

Figure 86 shows the amino acid sequence (SEQ ID NO: 86) derived from the coding sequence of SEQ ID NO: 85 shown in Figure 85.

Figure 87 shows a nucleotide sequence (SEQ ID NO: 87) of a native sequence PRO20084 cDNA, wherein SEQ ID NO: 87 is a clone designated herein as “DNA168061-2897” (UNQ6124).

Figure 88 shows the amino acid sequence (SEQ ID NO: 88) derived from the coding sequence of SEQ ID NO: 87 shown in Figure 87.

Figure 89 shows a nucleotide sequence (SEQ ID NO: 89) of a native sequence PRO21434 cDNA, wherein SEQ ID NO: 89 is a clone designated herein as “DNA147253-2983” (UNQ6509).

Figure 90 shows the amino acid sequence (SEQ ID NO: 90) derived from the coding sequence of SEQ ID NO: 89 shown in Figure 89.

Figure 91 shows a nucleotide sequence (SEQ ID NO: 91) of a native sequence PRO50332 cDNA, wherein SEQ ID NO: 91 is a clone designated herein as “DNA255255” (UNQ11645).

Figure 92 shows the amino acid sequence (SEQ ID NO: 92) derived from the coding sequence of SEQ ID NO: 91 shown in Figure 91.

Figure 93 shows a nucleotide sequence (SEQ ID NO: 93) of a native sequence PRO38465 cDNA, wherein SEQ ID NO: 93 is a clone designated herein as “DNA228002” (UNQ15965).

Figure 94 shows the amino acid sequence (SEQ ID NO: 94) derived from the coding sequence of SEQ ID NO: 93 shown in Figure 93.
Figure 95 shows a nucleotide sequence (SEQ ID NO:95) of a native sequence PRO346 cDNA, wherein SEQ ID NO:95 is a clone designated herein as “DNA44167-1243” (UNQ305).

Figure 96 shows the amino acid sequence (SEQ ID NO:96) derived from the coding sequence of SEQ ID NO:95 shown in Figure 95.

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

I. Definitions

The terms "PRO polypeptide" and “PRO” as used herein and when immediately followed by a numerical designation refer to various polypeptides, wherein the complete designation (i.e., PRO/number) refers to specific polypeptide sequences as described herein. The terms “PRO/number polypeptide” and “PRO/number” wherein the term "number" is provided as an actual numerical designation as used herein encompass native sequence polypeptides and polypeptide variants (which are further defined herein). The PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides described herein may be isolated from a variety of sources, such as from human tissue types or from another source, or prepared by recombinant or synthetic methods. The term "PRO polypeptide" refers to each individual PRO/number polypeptide disclosed herein. All disclosures in this specification which refer to the "PRO polypeptide" refer to each of the polypeptides individually as well as jointly. For example, descriptions of the preparation of, purification of, derivation of, formation of antibodies to or against, administration of, compositions containing, treatment of a disease with, etc., pertain to each polypeptide of the invention individually. The term “PRO polypeptide” also includes variants of the PRO/number polypeptides disclosed herein.

A "native sequence PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide" comprises a polypeptide having the same amino acid sequence as the corresponding PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide derived from nature. Such native sequence PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948,
PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides can be isolated from nature or can be produced by recombinant or synthetic means. The term "native sequence" PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide" specifically encompasses naturally-occurring truncated or secreted forms of the specific PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide (e.g., an extracellular domain sequence), naturally-occurring variant forms (e.g., alternatively spliced forms) and naturally-occurring allelic variants of the polypeptide. The invention provides native sequence PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide comprising the full-length amino acids sequences shown in the accompanying figures. Start and stop codons are shown in bold font and underlined in the figures. However, while the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide disclosed in the accompanying figures are shown to begin with methionine residues designated herein as amino acid position 1 in the figures, it is conceivable and possible that other methionine residues located either upstream or downstream from the amino acid position 1 in the figures may be employed as the starting amino acid residue for the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides.

The PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089,
PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide “extracellular domain” or “ECD” refers to a form of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide which is essentially free of the transmembrane and cytoplasmic domains. Ordinarily, a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide ECD will have less than 1% of such transmembrane and/or cytoplasmic domains and preferably, will have less than 0.5% of such domains. It will be understood that any transmembrane domains identified for the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides of the present invention are identified pursuant to criteria routinely employed in the art for identifying that type of hydrophobic domain. The exact boundaries of a transmembrane domain may vary but most likely by no more than about 5 amino acids at either end of the domain as initially identified herein. Optionally, therefore, an extracellular domain of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide may contain from about 5 or fewer amino acids on either side of the transmembrane domain/extracellular domain boundary as identified in the Examples or specification and such polypeptides, with or without the associated signal peptide, and nucleic acid encoding them, are contemplated by the present invention.

The approximate location of the “signal peptides” of the various PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides disclosed herein are shown in the present specification and/or the accompanying figures. It is noted, however, that the C-terminal boundary of a signal peptide may vary, but most likely by no more than about 5 amino acids on either side of the signal peptide C-terminal boundary as initially identified herein, wherein the C-terminal boundary of the signal peptide may be.
identified pursuant to criteria routinely employed in the art for identifying that type of amino acid sequence element (e.g., Nielsen et al., Prot. Eng. 10:1-6 (1997) and von Heinje et al., Nucl. Acids Res. 14:4683-4690 (1986)). Moreover, it is also recognized that, in some cases, cleavage of a signal sequence from a secreted polypeptide is not entirely uniform, resulting in more than one secreted species. These mature polypeptides, where the signal peptide is cleaved within no more than about 5 amino acids on either side of the C-terminal boundary of the signal peptide as identified herein, and the polymonucleotides encoding them, are contemplated by the present invention.

"PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide variant" means a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, preferably an active PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, as defined herein having at least about 80% amino acid sequence identity with a full-length native sequence PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide sequence as disclosed herein, a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide sequence lacking the signal peptide as disclosed herein, an extracellular domain of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, with or without the signal peptide, as disclosed herein.
or any other fragment of a full-length PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide sequence as disclosed herein (such as those encoded by a nucleic acid that represents only a portion of the complete coding sequence for a full-length PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide). Such PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides wherein one or more amino acid residues are added, or deleted, at the N- or C-terminus of the full-length native amino acid sequence. Ordinarily, a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide variant will have or will have at least about 80% amino acid sequence identity, alternatively will have or will have at least about 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, or 99% amino acid sequence identity, to a full-length native sequence PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide sequence as disclosed herein, a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907,
PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide sequence lacking the signal peptide as disclosed herein, an extracellular domain of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, with or without the signal peptide, as disclosed herein or any other specifically defined fragment of a full-length PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO655, PRO162, PRO788, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide sequence as disclosed herein. Ordinarily, PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 variant polypeptides are or are at least about 10 amino acids in length, alternatively are or are at least about 20, 30, 40, 50, 60, 70, 80, 90, 100, 110, 120, 130, 140, 150, 160, 170, 180, 190, 200, 210, 220, 230, 240, 250, 260, 270, 280, 290, 300, 310, 320, 330, 340, 350, 360, 370, 380, 390, 400, 410, 420, 430, 440, 450, 460, 470, 480, 490, 500, 510, 520, 530, 540, 550, 560, 570, 580, 590, 600 amino acids in length, or more. Optionally, PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 variant polypeptides will have no more than one conservative amino acid substitution as compared to the native PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide sequence, alternatively will have or will have no more than 2, 3, 4, 5, 6, 7, 8, 9, or 10 conservative amino acid substitution as compared to the native PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide sequence.
"Percent (%) amino acid sequence identity" with respect to the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9904, PRO9907, PRO9909, PRO9909, PRO10013, PRO10013, PRO10013, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide sequences identified herein is defined as the percentage of amino acid residues in a candidate sequence that are identical with the amino acid residues in the specific PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9904, PRO9907, PRO9909, PRO9909, PRO10013, PRO10013, PRO10013, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide sequence, after aligning the sequences and introducing gaps, if necessary, to achieve the maximum percent sequence identity, and not considering any conservative substitutions as part of the sequence identity. Alignment for purposes of determining percent amino acid sequence identity can be achieved in various ways that are within the skill in the art, for instance, using publicly available computer software such as BLAST, BLAST-2, ALIGN or Megalign (DNASTAR) software. Those skilled in the art can determine appropriate parameters for measuring alignment, including any algorithms needed to achieve maximal alignment over the full length of the sequences being compared. For purposes herein, however, % amino acid sequence identity values are generated using the sequence comparison computer program ALIGN-2, wherein the complete source code for the ALIGN-2 program is provided in Table 1 below. The ALIGN-2 sequence comparison computer program was authored by Genentech, Inc. and the source code shown in Table 1 below has been filed with user documentation in the U.S. Copyright Office, Washington D.C., 20559, where it is registered under U.S. Copyright Registration No. TXU510087. The ALIGN-2 program is publicly available through Genentech, Inc., South San Francisco, California or may be compiled from the source code provided in Table 1 below. The ALIGN-2 program should be compiled for use on a UNIX operating system, preferably digital UNIX V4.0D. All sequence comparison parameters are set by the ALIGN-2 program and do not vary.

In situations where ALIGN-2 is employed for amino acid sequence comparisons, the % amino acid sequence identity of a given amino acid sequence A to, with, or against a given amino acid sequence B (which can alternatively be phrased as a given amino acid sequence A that has or comprises a certain % amino acid sequence identity to, with, or against a given amino acid sequence B) is calculated as follows:

\[
\text{100 times the fraction } \frac{X}{Y}
\]

where X is the number of amino acid residues scored as identical matches by the sequence alignment program ALIGN-2 in that program's alignment of A and B, and where Y is the total number of amino acid residues in B.

It will be appreciated that where the length of amino acid sequence A is not equal to the length of amino acid sequence B, the % amino acid sequence identity of A to B will not equal the % amino acid sequence identity of B to A. As examples of % amino acid sequence identity calculations using this method, Tables 2 and 3
demonstrate how to calculate the % amino acid sequence identity of the amino acid sequence designated “Comparison Protein” to the amino acid sequence designated “PRO”, wherein "PRO" represents the amino acid sequence of a hypothetical PRO polypeptide of interest, "Comparison Protein" represents the amino acid sequence of a polypeptide against which the "PRO" polypeptide of interest is being compared, and "X", "Y" and "Z" each represent different hypothetical amino acid residues. Unless specifically stated otherwise, all % amino acid sequence identity values used herein are obtained as described in the immediately preceding paragraph using the ALIGN-2 computer program.

"PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO662, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 variant polynucleotide” or “PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO662, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 variant nucleic acid sequence” means a nucleic acid molecule which encodes a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO662, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, preferably an active PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO662, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, as defined herein and which has at least about 80% nucleic acid sequence identity with a nucleotide acid sequence encoding a full-length native sequence PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO662, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide sequence as disclosed herein, a full-length native sequence PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or
PRO346 polypeptide sequence lacking the signal peptide as disclosed herein, an extracellular domain of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, with or without the signal peptide, as disclosed herein or any other fragment of a full-length PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide sequence as disclosed herein (such as those encoded by a nucleic acid that represents only a portion of the complete coding sequence for a full-length PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 variant polynucleotide will have or will have at least about 80% nucleic acid sequence identity, alternatively will have or will have at least about 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, or 99% nucleic acid sequence identity with a nucleic acid sequence encoding a full-length native sequence PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide sequence as disclosed herein, a full-length native sequence PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide sequence lacking the signal peptide as disclosed herein, an extracellular domain of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111,
PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, with or without the signal sequence, as disclosed herein or any other fragment of a full-length PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide sequence as disclosed herein. Variants do not encompass the native nucleotide sequence.

Ordinarily, PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 variant polynucleotides are or are at least about 5 nucleotides in length, alternatively are or are at least about 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 35, 40, 45, 50, 55, 60, 65, 70, 75, 80, 85, 90, 95, 100, 105, 110, 115, 120, 125, 130, 135, 140, 145, 150, 155, 160, 165, 170, 175, 180, 185, 190, 195, 200, 210, 220, 230, 240, 250, 260, 270, 280, 290, 300, 310, 320, 330, 340, 350, 360, 370, 380, 390, 400, 410, 420, 430, 440, 450, 460, 470, 480, 490, 500, 510, 520, 530, 540, 550, 560, 570, 580, 590, 600, 610, 620, 630, 640, 650, 660, 670, 680, 690, 700, 710, 720, 730, 740, 750, 760, 770, 780, 790, 800, 810, 820, 830, 840, 850, 860, 870, 880, 890, 900, 910, 920, 930, 940, 950, 960, 970, 980, 990, or 1000 nucleotides in length, wherein in this context the term “about” means the referenced nucleotide sequence length plus or minus 10% of that referenced length.

"Percent (%) nucleic acid sequence identity" with respect to PRO218-, PRO228-, PRO271-, PRO273-, PRO295-, PRO302-, PRO305-, PRO326-, PRO386-, PRO655-, PRO162-, PRO788-, PRO792-, PRO940-, PRO941-, PRO1004-, PRO1012-, PRO1016-, PRO474-, PRO5238-, PRO1069-, PRO1111-, PRO1113-, PRO1130-, PRO1195-, PRO1271-, PRO1865-, PRO1879-, PRO3446-, PRO3543-, PRO4329-, PRO4352-, PRO5733-, PRO9859-, PRO9864-, PRO9904-, PRO9907-, PRO10013-, PRO90948-, PRO28694-, PRO16089-, PRO19563-, PRO19675-, PRO20084-, PRO21434-, PRO50332-, PRO38465- or PRO346-encoding nucleic acid sequences identified herein is defined as the percentage of nucleotides in a candidate sequence that are identical with the nucleotides in the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 nucleic acid sequence of interest, after aligning the sequences and introducing gaps, if necessary, to achieve the maximum percent sequence identity. Alignment for purposes of determining percent nucleic acid sequence identity can be achieved in various ways that are within the skill in the art, for instance, using publicly available computer software such as BLAST, BLAST-2, ALIGN or Megalign (DNASTAR) software. For purposes herein, however,
% nucleic acid sequence identity values are generated using the sequence comparison computer program ALIGN-2, wherein the complete source code for the ALIGN-2 program is provided in Table 1 below. The ALIGN-2 sequence comparison computer program was authored by Genentech, Inc. and the source code shown in Table 1 below has been filed with user documentation in the U.S. Copyright Office, Washington D.C., 20559, where it is registered under U.S. Copyright Registration No. TX-U10087. The ALIGN-2 program is publicly available through Genentech, Inc., South San Francisco, California or may be compiled from the source code provided in Table 1 below. The ALIGN-2 program should be compiled for use on a UNIX operating system, preferably digital UNIX V4.0D. All sequence comparison parameters are set by the ALIGN-2 program and do not vary.

In situations where ALIGN-2 is employed for nucleic acid sequence comparisons, the % nucleic acid sequence identity of a given nucleic acid sequence C to, with, or against a given nucleic acid sequence D (which can alternatively be phrased as a given nucleic acid sequence C that has or comprises a certain % nucleic acid sequence identity to, with, or against a given nucleic acid sequence D) is calculated as follows:

100 times the fraction W/Z

where W is the number of nucleotides scored as identical matches by the sequence alignment program ALIGN-2 in that program’s alignment of C and D, and where Z is the total number of nucleotides in D. It will be appreciated that where the length of nucleic acid sequence C is not equal to the length of nucleic acid sequence D, the % nucleic acid sequence identity of C to D will not equal the % nucleic acid sequence identity of D to C. As examples of % nucleic acid sequence identity calculations, Tables 4 and 5, demonstrate how to calculate the % nucleic acid sequence identity of the nucleic acid sequence designated “Comparison DNA” to the nucleic acid sequence designated “PRO-DNA”, wherein "PRO-DNA" represents a hypothetical PRO-encoding nucleic acid sequence of interest, "Comparison DNA" represents the nucleotide sequence of a nucleic acid molecule against which the "PRO-DNA" nucleic acid molecule of interest is being compared, and "N", "L" and "V" each represent different hypothetical nucleotides. Unless specifically stated otherwise, all % nucleic acid sequence identity values used herein are obtained as described in the immediately preceding paragraph using the ALIGN-2 computer program.

The invention also provides PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO92894, PRO10098, PRO10956, PRO119675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 variant polynucleotides which are nucleic acid molecules that encode a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO90948, PRO92894, PRO10098, PRO10956, PRO119675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide and which are capable of hybridizing, preferably under stringent hybridization and wash conditions, to nucleotide sequences encoding a full-length
PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide as disclosed herein. PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 variant polypeptides may be those that are encoded by a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 variant polynucleotid.

The term “full-length coding region” when used in reference to a nucleic acid encoding a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide refers to the sequence of nucleotides which encode the full-length PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide of the invention (which is often shown between start and stop codons, inclusive thereof, in the accompanying figures). The term “full-length coding region” when used in reference to an ATCC deposited nucleic acid refers to the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide-encoding portion of the cDNA that is inserted into the vector deposited with the ATCC (which is often shown between start and stop codons, inclusive thereof, in the accompanying figures).

"Isolated," when used to describe the various polypeptides disclosed herein, means polypeptide that has been identified and separated and/or recovered from a component of its natural environment. Contaminant components of its natural environment are materials that would typically interfere with diagnostic or therapeutic
uses for the polypeptide, and may include enzymes, hormones, and other proteinaceous or non-proteinaceous solutes. The invention provides that the polypeptide will be purified (1) to a degree sufficient to obtain at least 15 residues of N-terminal or internal amino acid sequence by use of a spinning cup sequenator, or (2) to homogeneity by SDS-PAGE under non-reducing or reducing conditions using Coomassie blue or, preferably, silver stain. Isolated polypeptide includes polypeptide in situ within recombinant cells, since at least one component of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide natural environment will not be present. Ordinarily, however, isolated polypeptide will be prepared by at least one purification step.

An "isolated" PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide-encoding nucleic acid or other polypeptide-encoding nucleic acid is a nucleic acid molecule that is identified and separated from at least one contaminant nucleic acid molecule with which it is ordinarily associated in the natural source of the polypeptide-encoding nucleic acid. An isolated polypeptide-encoding nucleic acid molecule is other than in the form or setting in which it is found in nature. Isolated polypeptide-encoding nucleic acid molecules therefore are distinguished from the specific polypeptide-encoding nucleic acid molecule as it exists in natural cells. However, an isolated polypeptide-encoding nucleic acid molecule includes polypeptide-encoding nucleic acid molecules contained in cells that ordinarily express the polypeptide where, for example, the nucleic acid molecule is in a chromosomal location different from that of natural cells.

The term "control sequences" refers to DNA sequences necessary for the expression of an operably linked coding sequence in a particular host organism. The control sequences that are suitable for prokaryotes, for example, include a promoter, optionally an operator sequence, and a ribosome binding site. Eukaryotic cells are known to utilize promoters, polyadenylation signals, and enhancers.

Nucleic acid is "operably linked" when it is placed into a functional relationship with another nucleic acid sequence. For example, DNA for a presequence or secretory leader is operably linked to DNA for a polypeptide if it is expressed as a preprotein that participates in the secretion of the polypeptide; a promoter or enhancer is operably linked to a coding sequence if it affects the transcription of the sequence; or a ribosome binding site is operably linked to a coding sequence if it is positioned so as to facilitate translation. Generally, "operably linked" means that the DNA sequences being linked are contiguous, and, in the case of a secretory leader, contiguous and in reading phase. However, enhancers do not have to be contiguous. Linking is accomplished by ligation at convenient restriction sites. If such sites do not exist, the synthetic oligonucleotide adaptors or linkers are used in accordance with conventional practice.

"Stringency" of hybridization reactions is readily determinable by one of ordinary skill in the art, and
generally is an empirical calculation dependent upon probe length, washing temperature, and salt concentration. In general, longer probes require higher temperatures for proper annealing, while shorter probes need lower temperatures. Hybridization generally depends on the ability of denatured DNA to reanneal when complementary strands are present in an environment below their melting temperature. The higher the degree of desired homology between the probe and hybridizable sequence, the higher the relative temperature which can be used. As a result, it follows that higher relative temperatures would tend to make the reaction conditions more stringent, while lower temperatures less so. For additional details and explanation of stringency of hybridization reactions, see Ausubel et al., *Current Protocols in Molecular Biology*, Wiley Interscience Publishers, (1995).

"Stringent conditions" or "high stringency conditions", as defined herein, may be identified by those that: (1) employ low ionic strength and high temperature for washing, for example 0.015 M sodium chloride/0.0015 M sodium citrate/0.1% sodium dodecyl sulfate at 50°C; (2) employ during hybridization a denaturing agent, such as formamide, for example, 50% (v/v) formamide with 0.1% bovine serum albumin/0.1% Ficoll/0.1% polyvinylpyrrolidone/50mM sodium phosphate buffer at pH 6.5 with 750 mM sodium chloride, 75 mM sodium citrate at 42°C; or (3) employ 50% formamide, 5 x SSC (0.75 M NaCl, 0.075 M sodium citrate), 50 mM sodium phosphate (pH 6.8), 0.1% sodium pyrophosphate, 5 x Denhardt’s solution, sonicated salmon sperm DNA (50 μg/ml), 0.1% SDS, and 10% dextran sulfate at 42°C, with washes at 42°C in 0.2 x SSC (sodium chloride/sodium citrate) and 50% formamide at 55°C, followed by a high-stringency wash consisting of 0.1 x SSC containing EDTA at 55°C.

"Moderately stringent conditions" may be identified as described by Sambrook et al., *Molecular Cloning: A Laboratory Manual*, New York: Cold Spring Harbor Press, 1989, and include the use of washing solution and hybridization conditions (e.g., temperature, ionic strength and %SDS) less stringent than those described above. An example of moderately stringent conditions is overnight incubation at 37°C in a solution comprising: 20% formamide, 5 x SSC (150 mM NaCl, 15 mM trisodium citrate), 50 mM sodium phosphate (pH 7.6), 5 x Denhardt’s solution, 10% dextran sulfate, and 20 mg/ml denatured sheared salmon sperm DNA, followed by washing the filters in 1 x SSC at about 37-50°C. The skilled artisan will recognize how to adjust the temperature, ionic strength, etc. as necessary to accommodate factors such as probe length and the like.

The term "epitope tagged" when used herein refers to a chimeric polypeptide comprising a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide fused to a "tag polypeptide". The tag polypeptide has enough residues to provide an epitope against which an antibody can be made, yet is short enough such that it does not interfere with activity of the polypeptide to which it is fused. The tag polypeptide preferably also is fairly unique so that the antibody does not substantially cross-react with other epitopes. Suitable tag polypeptides generally have at least six amino acid residues and usually between about 8 and 50 amino acid residues (preferably, between about 10 and 20 amino acid residues).

"Active" or "activity" for the purposes herein refers to form(s) of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941,
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PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide which retain a biological and/or an immunological activity of native or naturally-occurring PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, wherein “biological” activity refers to a biological function (either inhibitory or stimulatory) caused by a native or naturally-occurring PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide other than the ability to induce the production of an antibody against an antigenic epitope possessed by a native or naturally-occurring PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide and an “immunological” activity refers to the ability to induce the production of an antibody against an antigenic epitope possessed by a native or naturally-occurring PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide.

The term "antagonist" is used in the broadest sense [unless otherwise qualified], and includes any molecule that partially or fully blocks, inhibits, or neutralizes a biological activity of a native PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide disclosed herein. In a similar manner, the term "agonist" is used in the broadest sense [unless otherwise qualified] and includes any molecule that mimics a biological activity of a native PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879,
PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99048, PRO28694, PRO16089, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide disclosed herein. Suitable agonist or antagonist molecules specifically include agonist or antagonist antibodies or antibody fragments, fragments or amino acid sequence variants of native PRO218, PRO228, PRO2271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO655, PRO162, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99048, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides, peptides, antisense oligonucleotides, small organic molecules, etc. Methods for identifying agonists or antagonists of a PRO218, PRO228, PRO2271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO655, PRO162, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99048, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide may comprise contacting a PRO218, PRO228, PRO2271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO655, PRO162, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99048, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide with a candidate agonist or antagonist molecule and measuring a detectable change in one or more biological activities normally associated with the PRO218, PRO228, PRO2271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO655, PRO162, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99048, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide.

"Treating" or "treatment" or "alleviation" refers to both therapeutic treatment and prophylactic or preventative measures, wherein the object is to prevent or slow down (lessen) the targeted pathologic condition or disorder. A subject in need of treatment may already have the disorder, or may be prone to have the disorder or may be in whom the disorder is to be prevented.

"Chronic" administration refers to administration of the agent(s) in a continuous mode as opposed to an acute mode, so as to maintain the initial therapeutic effect (activity) for an extended period of time. "Intermittent" administration is treatment that is not consecutively done without interruption, but rather is cyclic in nature.

"Mammal" for purposes of treatment refers to any animal classified as a mammal, including humans, rodents such as rats or mice, domestic and farm animals, and zoo, sports, or pet animals, such as dogs, cats, cattle, horses, sheep, pigs, goats, rabbits, etc. Preferably, the mammal is human.

Administration "in combination with" one or more further therapeutic agents includes simultaneous (concurrent) and consecutive administration in any order.

"Carriers" as used herein include pharmaceutically acceptable carriers, excipients, or stabilizers which
are nontoxic to the cell or mammal being exposed thereto at the dosages and concentrations employed. Often the physiologically acceptable carrier is an aqueous pH buffered solution. Examples of physiologically acceptable carriers include buffers such as phosphate, citrate, and other organic acids; antioxidants including ascorbic acid; low molecular weight (less than about 10 residues) polypeptide; proteins, such as serum albumin, gelatin, or immunoglobulins; hydrophilic polymers such as polyvinylpyrrolidone; amino acids such as glycine, glutamine, asparagine, arginine or lysine; monosaccharides, disaccharides, and other carbohydrates including glucose, mannose, or dextrins; chelating agents such as EDTA; sugar alcohols such as mannitol or sorbitol; salt-forming counterions such as sodium; and/or nonionic surfactants such as TWEEN™, polyethylene glycol (PEG), and PLURONICS™.

By "solid phase" is meant a non-aqueous matrix to which the antibody of the present invention can adhere. Examples of solid phases encompassed herein include those formed partially or entirely of glass (e.g., controlled pore glass), polysaccharides (e.g., agarose), polyacrylamides, polystyrene, polyvinyl alcohol and silicones. Depending on the context, the solid phase can comprise the well of an assay plate; in others it is a purification column (e.g., an affinity chromatography column). This term also includes a discontinuous solid phase of discrete particles, such as those described in U.S. Patent No. 4,275,149.

A "liposome" is a small vesicle composed of various types of lipids, phospholipids and/or surfactant which is useful for delivery of a drug (such as a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1130, PRO1195, PRO121, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide or antibody thereto) to a mammal. The components of the liposome are commonly arranged in a bilayer formation, similar to the lipid arrangement of biological membranes.

A “small molecule” is defined herein to have a molecular weight below about 500 Daltons.

An “effective amount” of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO121, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO121, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody, a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO121, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352,
PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089,
PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 binding oligopeptide, a
PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162,
PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111,
PRO1113, PRO1119, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352,
PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089,
PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 binding organic molecule
or an agonist or antagonist thereof as disclosed herein is an amount sufficient to carry out a specifically stated
purpose. An “effective amount” may be determined empirically and in a routine manner, in relation to the stated
purpose.

The term “therapeutically effective amount” refers to an amount of an anti-PRO218, anti-PRO228, anti-
PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-
PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016,
anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1119, anti-PRO1271, anti-
PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-
PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-
PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332,
anti-PRO38465 or anti-PRO346 antibody, a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305,
PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016,
PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1119, PRO1271, PRO1865, PRO1879,
PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694;
PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 binding oligopeptide, a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655,
PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069,
PRO1111, PRO1113, PRO1119, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329,
PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089,
PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 binding organic molecule
or other drug effective to “treat” a disease or disorder in a subject or mammal. In the case of cancer, the
therapeutically effective amount of the drug may reduce the number of cancer cells; reduce the tumor size; inhibit
(i.e., slow to some extent and preferably stop) cancer cell infiltration into peripheral organs; inhibit (i.e., slow
to some extent and preferably stop) tumor metastasis; inhibit, to some extent, tumor growth; and/or relieve to some
extent one or more of the symptoms associated with the cancer. See the definition herein of “treating.” To the
extent the drug may prevent growth and/or kill existing cancer cells, it may be cytostatic and/or cytotoxic.

The phrases "cardiovascular, endothelial and angiogenic disorder", "cardiovascular, endothelial and
angiogenic dysfunction”, “cardiovascular, endothelial or angiogenic disorder” and “cardiovascular, endothelial or angiogenic dysfunction” are used interchangeably and refer in part to systemic disorders that affect vessels, such as diabetes mellitus, as well as diseases of the vessels themselves, such as of the arteries, capillaries, veins, and/or lymphatics. This would include indications that stimulate angiogenesis and/or cardiovascularization, and those that inhibit angiogenesis and/or cardiovascularization. Such disorders include, for example, arterial disease, such as atherosclerosis, hypertension, inflammatory vasculitides, Reynaud’s disease and Reynaud’s phenomenon, aneurysms, and arterial restenosis; venous and lymphatic disorders such as thrombophlebitis, lymphangitis, and lymphedema; and other vascular disorders such as peripheral vascular disease, cancer such as vascular tumors, e.g., hemangioma (capillary and cavernous), glomus tumors, telangiectasia, bacillary angiomatosis, hemangioendothelioma, angiosarcoma, haemangiopericytoma, Kaposi’s sarcoma, lymphangioma, and lymphangiosarcoma, tumor angiogenesis, trauma such as wounds, burns, and other injured tissue, implant fixation, scarring, ischemia reperfusion injury, rheumatoid arthritis, cerebrovascular disease, renal diseases such as acute renal failure, or osteoporosis. This would also include angina, myocardial infarctions such as acute myocardial infarctions, cardiac hypertrophy, and heart failure such as CHF.

“Hypertrophy”, as used herein, is defined as an increase in mass of an organ or structure independent of natural growth that does not involve tumor formation. Hypertrophy of an organ or tissue is due either to an increase in the mass of the individual cells (true hypertrophy), or to an increase in the number of cells making up the tissue (hyperplasia), or both. Certain organs, such as the heart, lose the ability to divide shortly after birth. Accordingly, “cardiac hypertrophy” is defined as an increase in mass of the heart, which, in adults, is characterized by an increase in myocyte cell size and contractile protein content without concomitant cell division. The character of the stress responsible for inciting the hypertrophy, (e.g., increased preload, increased afterload, loss of myocytes, as in myocardial infarction, or primary depression of contractility), appears to play a critical role in determining the nature of the response. The early stage of cardiac hypertrophy is usually characterized morphologically by increases in the size of myofibrils and mitochondria, as well as by enlargement of mitochondria and nuclei. At this stage, while muscle cells are larger than normal, cellular organization is largely preserved. At a more advanced stage of cardiac hypertrophy, there are preferential increases in the size or number of specific organelles, such as mitochondria, and new contractile elements are added in localized areas of the cells, in an irregular manner. Cells subjected to long-standing hypertrophy show more obvious disruptions in cellular organization, including markedly enlarged nuclei with highly lobulated membranes, which displace adjacent myofibrils and cause breakdown of normal Z-band registration. The phrase “cardiac hypertrophy” is used to include all stages of the progression of this condition, characterized by various degrees of structural damage of the heart muscle, regardless of the underlying cardiac disorder. Hence, the term also includes physiological conditions instrumental in the development of cardiac hypertrophy, such as elevated blood pressure, aortic stenosis, or myocardial infarction.

“Heart failure” refers to an abnormality of cardiac function where the heart does not pump blood at the rate needed for the requirements of metabolizing tissues. The heart failure can be caused by a number of factors, including ischemic, congenital, rheumatic, or idiopathic forms.

“Congestive heart failure” (CHF) is a progressive pathologic state where the heart is increasingly unable to supply adequate cardiac output (the volume of blood pumped by the heart over time) to deliver the oxygenated
blood to peripheral tissues. As CHF progresses, structural and hemodynamic damages occur. While these damages have a variety of manifestations, one characteristic symptom is ventricular hypertrophy. CHF is a common end result of a number of various cardiac disorders.

“Myocardial infarction” generally results from atherosclerosis of the coronary arteries, often with superimposed coronary thrombosis. It may be divided into two major types: transmural infarcts, in which myocardial necrosis involves the full thickness of the ventricular wall, and subendocardial (nontransmural) infarcts, in which the necrosis involves the subendocardium, the intramyocardial myocardium, or both, without extending all the way through the ventricular wall to the epicardium. Myocardial infarction is known to cause both a change in hemodynamic effects and an alteration in structure in the damaged and healthy zones of the heart. Thus, for example, myocardial infarction reduces the maximum cardiac output and the stroke volume of the heart. Also associated with myocardial infarction is a stimulation of the DNA synthesis occurring in the interstice as well as an increase in the formation of collagen in the areas of the heart not affected.

As a result of the increased stress or strain placed on the heart in prolonged hypertension due, for example, to the increased total peripheral resistance, cardiac hypertrophy has long been associated with “hypertension”. A characteristic of the ventricle that becomes hypertrophic as a result of chronic pressure overload is an impaired diastolic performance. Fouad et al., J. Am. Coll. Cardiol., 4: 1500-1506 (1984); Smith et al., J. Am. Coll. Cardiol., 5: 869-874 (1985). A prolonged left ventricular relaxation has been detected in early essential hypertension, in spite of normal or supranormal systolic function. Hartford et al., Hypertension, 6: 329-338 (1984). However, there is no close parallelism between blood pressure levels and cardiac hypertrophy. Although improvement in left ventricular function in response to antihypertensive therapy has been reported in humans, patients variously treated with a diuretic (hydrochlorothiazide), a β-blocker (propranolol), or a calcium channel blocker (diltiazem), have shown reversal of left ventricular hypertrophy, without improvement in diastolic function. Inouye et al., Am. J. Cardiol., 53: 1583-7 (1984).


Supravalvular “aortic stenosis” is an inherited vascular disorder characterized by narrowing of the ascending aorta, but other arteries, including the pulmonary arteries, may also be affected. Untreated aortic
stenosis may lead to increased intracardiac pressure resulting in myocardial hypertrophy and eventually heart failure and death. The pathogenesis of this disorder is not fully understood, but hypertrophy and possibly hyperplasia of medial smooth muscle are prominent features of this disorder. It has been reported that molecular variants of the elastin gene are involved in the development and pathogenesis of aortic stenosis. U.S. Patent No. 5,650,282 issued July 22, 1997.

"Valvular regurgitation" occurs as a result of heart diseases resulting in disorders of the cardiac valves. Various diseases, like rheumatic fever, can cause the shrinking or pulling apart of the valve orifice, while other diseases may result in endocarditis, an inflammation of the endocardium or lining membrane of the atrioventricular orifices and operation of the heart. Defects such as the narrowing of the valve stenosis or the defective closing of the valve result in an accumulation of blood in the heart cavity or regurgitation of blood past the valve. If uncorrected, prolonged valvular stenosis or insufficiency may result in cardiac hypertrophy and associated damage to the heart muscle, which may eventually necessitate valve replacement.

The term "immune related disease" means a disease in which a component of the immune system of a mammal causes, mediates or otherwise contributes to a morbidity in the mammal. Also included are diseases in which stimulation or intervention of the immune response has an ameliorative effect on progression of the disease. Excluded within this term are immune-mediated inflammatory diseases, non-immune-mediated inflammatory diseases, infectious diseases, immunodeficiency diseases, neoplasia, etc.

The term "T cell mediated disease" means a disease in which T cells directly or indirectly mediate or otherwise contribute to a morbidity in a mammal. The T cell mediated disease may be associated with cell mediated effects, lymphokine mediated effects, etc., and even effects associated with B cells if the B cells are stimulated, for example, by the lymphokines secreted by T cells.

Examples of immune-related and inflammatory diseases, some of which are immune or T cell mediated, include systemic lupus erythematosus, rheumatoid arthritis, juvenile chronic arthritis, spondyloarthopathies, systemic sclerosis (scleroderma), idiopathic inflammatory myopathies (dermatomyositis, polymyositis), Sjögren's syndrome, systemic vasculitis, sarcoidosis, autoimmune hemolytic anemia (immune pancytopenia, paroxysmal nocturnal hemoglobinuria), autoimmune thrombocytopenia (idiopathic thrombocytopenic purpura, immune-mediated thrombocytopenia), thyroiditis (Grave's disease, Hashimoto's thyroiditis, juvenile lymphocytic thyroiditis, atrophic thyroiditis), diabetes mellitus, immune-mediated renal disease (glomerulonephritis, tubulointerstitial nephritis), demyelinating diseases of the central and peripheral nervous systems such as multiple sclerosis, idiopathic demyelinating polyneuropathy or Guillain-Barré syndrome, and chronic inflammatory demyelinating polyneuropathy, hepatobiliary diseases such as infectious hepatitis (hepatitis A, B, C, D, E and other non-hepatotropic viruses), autoimmune chronic active hepatitis, primary biliary cirrhosis, granulomatous hepatitis, and sclerosing cholangitis, inflammatory bowel disease (ulcerative colitis: Crohn's disease), gluten-sensitive enteropathy, and Whipple's disease, autoimmune or immune-mediated skin diseases including bullous skin diseases, erythema multiforme and contact dermatitis, psoriasis, allergic diseases such as asthma, allergic rhinitis, atopic dermatitis, food hypersensitivity and urticaria, immunologic diseases of the lung such as eosinophilic pneumonia, idiopathic pulmonary fibrosis and hypersensitivity pneumonitis, or transplantation associated diseases including graft rejection and graft-versus-host-disease. Infectious diseases including viral diseases such as AIDS (HIV infection), hepatitis A, B, C, D, and E, herpes, etc., bacterial infections, fungal infections, protozoal
infections and parasitic infections.

An "autoimmune disease" herein is a disease or disorder arising from and directed against an individual's own tissues or organs or a co-segregate or manifestation thereof or resulting condition therefrom. In many of these autoimmune and inflammatory disorders, a number of clinical and laboratory markers may exist, including, but not limited to, hypergammaglobulinemia, high levels of autoantibodies, antigen-antibody complex deposits in tissues, benefit from corticosteroid or immunosuppressive treatments, and lymphoid cell aggregates in affected tissues. Without being limited to any one theory regarding B-cell mediated autoimmune disease, it is believed that B cells demonstrate a pathogenic effect in human autoimmune diseases through a multitude of mechanistic pathways, including autoantibody production, immune complex formation, dendritic and T-cell activation, cytokine synthesis, direct chemokine release, and providing a nidus for ectopic neo-lymphogenesis. Each of these pathways may participate to different degrees in the pathology of autoimmune diseases.

"Autoimmune disease" can be an organ-specific disease (i.e., the immune response is specifically directed against an organ system such as the endocrine system, the hematopoietic system, the skin, the cardiopulmonary system, the gastrointestinal and liver systems, the renal system, the thyroid, the ears, the neuromuscular system, the central nervous system, etc.) or a systemic disease which can affect multiple organ systems (for example, systemic lupus erythematosus (SLE), rheumatoid arthritis, polymyositis, etc.). Preferred such diseases include autoimmune rheumatologic disorders (such as, for example, rheumatoid arthritis, Sjögren's syndrome, scleroderma, lupus such as SLE and lupus nephritis, polymyositis/dermatomyositis, cryoglobulinemia, anti-phospholipid antibody syndrome, and psoriatic arthritis), autoimmune gastrointestinal and liver disorders (such as, for example, inflammatory bowel diseases (e.g., ulcerative colitis and Crohn's disease), autoimmune gastritis and pernicious anemia, autoimmune hepatitis, primary biliary cirrhosis, primary sclerosing cholangitis, and celiac disease), vasculitis (such as, for example, ANCA-associated vasculitis, including Churg-Strauss vasculitis, Wegener's granulomatosis, and polyarteritis), autoimmune neurological disorders (such as, for example, multiple sclerosis, opsoclonus myoclonus syndrome, myasthenia gravis, neuromyelitis optica, Parkinson's disease, Alzheimer's disease, and autoimmune polyneuropathies), renal disorders (such as, for example, glomerulonephritis, Goodpasture's syndrome, and Berger's disease), autoimmune dermatologic disorders (such as, for example, psoriasis, urticaria, hives, pemphigus vulgaris, bullous pemphigoid, and cutaneous lupus erythematosus), hematologic disorders (such as, for example, thrombocytopenic purpura, thrombotic thrombocytopenic purpura, post-transfusion purpura, and autoimmune hemolytic anemia), atherosclerosis, uveitis, autoimmune hearing diseases (such as, for example, inner ear disease and hearing loss), Behcet's disease, Raynaud's syndrome, organ transplant, and autoimmune endocrine disorders (such as, for example, diabetic-related autoimmune diseases such as insulin-dependent diabetes mellitus (IDDM), Addison's disease, and autoimmune thyroid disease (e.g., Graves' disease and thyroiditis)). More preferred such diseases include, for example, rheumatoid arthritis, ulcerative colitis, ANCA-associated vasculitis, lupus, multiple sclerosis, Sjögren's syndrome, Graves' disease, IDDM, pernicious anemia, thyroiditis, and glomerulonephritis.

Specific examples of other autoimmune diseases as defined herein, which in some cases encompass those listed above, include, but are not limited to, arthritis (acute and chronic, rheumatoid arthritis including juvenile-onset rheumatoid arthritis and stages such as rheumatoid synovitis, gout or gouty arthritis, acute immunological arthritis, chronic inflammatory arthritis, degenerative arthritis, type II collagen-induced arthritis, infectious arthritis, Lyme
arthritis, proliferative arthritis, psoriatic arthritis, Still's disease, vertebral arthritis, osteoarthritis, arthritis chronica progrediente, arthritis deformans, polyarthritis chronica primaria, reactive arthritis, menopausal arthritis, estrogen-depletion arthritis, and ankylosing spondylitis/rheumatoid spondylitis), autoimmune lymphoproliferative disease, inflammatory hyperproliferative skin diseases, psoriasis such as plaque psoriasis, guttate psoriasis, pustular psoriasis, and psoriasis of the nails, atopy including atopic diseases such as hay fever and Job's syndrome, dermatitis including contact dermatitis, chronic contact dermatitis, exfoliative dermatitis, allergic dermatitis, allergic contact dermatitis, hives, dermatitis herpetiformis, nummular dermatitis, seborrheic dermatitis, non-specific dermatitis, primary irritant contact dermatitis, and atop dermatitis, x-linked hyper IgM syndrome, allergic intraocular inflammatory diseases, urticaria such as chronic allergic urticaria and chronic idiopathic urticaria, including chronic autoimmune urticaria, myositis, polymyositis/dermatomyositis, juvenile dermatomyositis, toxic epidermal necrolysis, scleroderma (including systemic scleroderma), sclerosis such as systemic sclerosis, multiple sclerosis (MS) such as spino-optical MS, primary progressive MS (PPMS), and relapsing remitting MS (RRMS), progressive systemic sclerosis, atherosclerosis, arteriosclerosis, sclerosis disseminata, atactic sclerosis, neuromyelitis optica (NMO), inflammatory bowel disease (IBD) (for example, Crohn's disease, autoimmune-mediated gastrointestinal diseases, gastrointestinal inflammation, colitis such as ulcerative colitis, colitis ulcerosa, microscopic colitis, collagenous colitis, colitis polyposa, necrotizing enterocolitis, and transmural colitis, and autoimmune inflammatory bowel disease), bowel inflammation, pyoderma gangrenosum, erythema nodosum, primary sclerosing cholangitis, respiratory distress syndrome, including adult or acute respiratory distress syndrome (ARDS), meningitis, inflammation of all or part of the uvea, iritis, choroiditis, an autoimmune hematological disorder, graft-versus-host disease, angioedema such as hereditary angioedema, cranial nerve damage as in meningitis, herpes gestationis, pemphigoid gestationis, pruritis scrofi, autoimmune premature ovarian failure, sudden hearing loss due to an autoimmune condition, IgE-mediated diseases such as anaphylaxis and allergic and atopic rhinitis, encephalitis such as Rasmussen's encephalitis and limbic and/or brainstem encephalitis, uveitis, such as anterior uveitis, acute anterior uveitis, granulomatous uveitis, nongranulomatous uveitis, phacoantigenic uveitis, posterior uveitis, or autoimmune uveitis, glomerulonephritis (GN) with and without nephrotic syndrome such as chronic or acute glomerulonephritis such as primary GN, immune-mediated GN, membranous GN (membranous nephropathy), idiopathic membranous GN or idiopathic membranous nephropathy, membrano- or membranous proliferative GN (MPGN), including Type I and Type II, and rapidly progressive GN (RPGN), proliferative nephritis, autoimmune polyglandular endocrine failure, balanitis including balanitis circumspecta plasmacellularis, balanoposthitis, erythema annulare centrifugum, erythema dyschormicum perstans, erythema multiform, granuloma annulare, lichen nitidus, lichen sclerosus et atrophicus, lichen simplex chronicus, lichen spinulosus, lichen planus, lamellar ichthyosis, epidermolytic hyperkeratosis, pemphigant keratosis, pyoderma gangrenosum, allergic conditions and responses, food allergies, drug allergies, insect allergies, rare allergic disorders such as mastocytosis, allergic reaction, eczema including allergic or atopic eczema, astreatic eczema, dyshidrotic eczema, and vesicular palmoplantar eczema, asthma such as asthma bronchiale, bronchial asthma, and auto-immune asthma, conditions involving infiltration of T cells and chronic inflammatory responses, immune reactions against foreign antigens such as fetal A-B-O blood groups during pregnancy, chronic pulmonary inflammatory disease, autoimmune myocarditis, leukocyte adhesion deficiency, lupus, including lupus nephritis, lupus cerebritis, pediatric lupus, non-renal lupus, extra-renal lupus, discoid lupus and discoid lupus erythematosus,
alopecia lupus, SLE, such as cutaneous SLE or subacute cutaneous SLE, neonatal lupus syndrome (NLE), and lupus erythematosus disseminatus, juvenile onset (Type I) diabetes mellitus, including pediatric IDDM, adult onset diabetes mellitus (Type II diabetes), autoimmune diabetes, idiopathic diabetes insipidus, diabetic retinopathy, diabetic nephropathy, diabetic colitis, diabetic large-artery disorder, immune responses associated with acute and delayed hypersensitivity mediated by cytokines and T-lymphocytes, tuberculosis, sarcoidosis, granulomatosis including lymphomatoid granulomatosis, Wegener's granulomatosis, agranulocytosis, vasculitides, including vasculitis, large-vessel vasculitis (including polyarthritis rheumatica and giant-cell (Takayasu's) arteritis), medium-vessel vasculitis (including Kawasaki's disease and polyarteritis nodosa/periarteritis nodosa), microscopic polyarteritis, immunovasculitis, CNS vasculitis, cutaneous vasculitis, hypersensitivity vasculitis, necrotizing vasculitis such as systemic necrotizing vasculitis, and ANCA-associated vasculitis, such as Churg-Strauss vasculitis or syndrome (CSS) and ANCA-associated small-vessel vasculitis, temporal arteritis, aplastic anemia, autoimmune aplastic anemia, Coombs positive anemia, Diamond Blackfan anemia, hemolytic anemia or immune hemolytic anemia including autoimmune hemolytic anemia (AIHA), pernicious anemia (anemia perniciosa), Addison's disease, pure red cell anemia or aplasia (PRCA), Factor VIII deficiency, hemophilia A, autoimmune neutropenia(s), cytopenias such as pancytopenia, leukopenia, diseases involving leukocyte diapedesis, CNS inflammatory disorders, Alzheimer's disease, Parkinson's disease, multiple organ injury syndrome such as those secondary to septicemia, trauma or hemorrhage, antigen-antibody complex- mediated diseases, anti-glomerular basement membrane disease, anti-phospholipid antibody syndrome, motoneuritis, allergic neuritis, Behcet's disease/syndrome, Castleman's syndrome, Goodpasture's syndrome, Reynaud's syndrome, Sjögren's syndrome, Stevens-Johnson syndrome, pemphigoid such as pemphigoid bullous and skin pemphigoid, pemphigus (including pemphigus vulgaris, pemphigus foliaceus, pemphigus mucos-membrane pemphigoid, and pemphigus erythematosus), autoimmune polyendocrinopathies, Reiter's disease or syndrome, thermal injury due to an autoimmune condition, preeclampsia, an immune complex disorder such as immune complex nephritis, antibody-mediated nephritis, neuroinflammatory disorders, polynephropathies, chronic neuropathy such as IgM polynephropathies or IgM-mediated neuropathy, thrombocytopenia (as developed by myocardial infarction patients, for example), including thrombotic thrombocytopenic purpura (TTP), post-transfusion purpura (PTP), heparin-induced thrombocytopenia, and autoimmune or immune-mediated thrombocytopenia including, for example, idiopathic thrombocytopenic purpura (ITP) including chronic or acute ITP, scleritis such as idiopathic cerato-scleritis, episcleritis, autoimmune disease of the testis and ovary including autoimmune orchitis and oophoritis, primary hypothyroidism, hypoparathyroidism, autoimmune endocrine diseases including thyroiditis such as autoimmune thyroiditis, Hashimoto's disease, chronic thyroiditis (Hashimoto's thyroiditis), or subacute thyroiditis, autoimmune thyroid disease, idiopathic hypothyroidism, Grave's disease, polyglandular syndromes such as autoimmune polyglandular syndromes, for example, type I (or polyglandular endocrinopathy syndromes), paraneoplastic syndromes, including neurologic paraneoplastic syndromes such as Lambert-Eaton myasthenic syndrome or Eaton-Lambert syndrome, stiff-man or stiff-person syndrome, encephalomyelitis such as allergic encephalomyelitis or encephalomyelitis allergic and experimental allergic encephalomyelitis (EAE), myasthenia gravis such as thymoma-associated myasthenia gravis, cerebellar degeneration, neuromyotonia, opsoconitus or opsoconus myoclonus syndrome (OMS), and sensory neuropathy, multifocal motor neuropathy, Sheehan's syndrome, autoimmune hepatitis, chronic hepatitis, lupoid hepatitis, giant-cell hepatitis, chronic active hepatitis.
or autoimmune chronic active hepatitis, pneumonitis such as lymphoid interstitial pneumonitis (LIP), bronchiolitis obliterans (non-transplant) vs NSIP, Guillain-Barré syndrome, Berger's disease (IgA nephropathy), idiopathic IgA nephropathy, linear IgA dermatosis, acute febrile neutrophilic dermatosis, subcorneal pustular dermatosis, transient acantholytic dermatosis, cirrhosis such as primary biliary cirrhosis and pneumonocirrhosis, autoimmune enteropathy syndrome, Celiac or Coeliac disease, celiac sprue (gluten enteropathy), refractory sprue, idiopathic sprue, cryoglobulinemia such as mixed cryoglobulinemia, amyloidotic lateral sclerosis (ALS; Lou Gehrig's disease), coronary artery disease, autoimmune ear disease such as autoimmune inner ear disease (AIED), autoimmune hearing loss, osteochondritis such as refractory or relapsed or relapsing osteochondritis, pulmonary alveolar proteinosis, Cogan's syndrome/nonsyphilitic interstitial keratitis, Bell's palsy, Sweet's disease/syndrome, rosacea autoimmune, zoster-associated pain, amyloidosis, a non-cancerous lymphocytosis, a primary lymphocytosis, which includes monoclonal B cell lymphocytosis (e.g., benign monoclonal gammopathy and monoclonal gammopathy of undetermined significance, MGUS), peripheral neuropathy, paraneoplastic syndrome, channelopathies such as epilepsy, migraine, arrhythmia, muscular disorders, deafness, blindness, periodic paralysis, and channelopathies of the CNS, autism, inflammatory myopathy, focal or segmental or focal segmental glomerulosclerosis (FSGS), endocrine ophthalmopathy, uveoretinitis, chorioretinitis, autoimmune hepaticological disorder, fibromyalgia, multiple endocrine failure, Schmidt's syndrome, adenitis, gastric atrophy, presenile dementia, demyelinating diseases such as autoimmune demyelinating diseases and chronic inflammatory demyelinating polyneuropathy, Dressler's syndrome, alopecia areata, alopecia totalis, CREST syndrome (calcinosis, Raynaud's phenomenon, esophageal dysmotility, sclerodactyly, and telangiectasia), male and female autoimmune infertility, e.g., due to anti-spermatozoan antibodies, mixed connective tissue disease, Chagas' disease, rheumatic fever, recurrent abortion, farmer's lung, erythema multiforme, post-cardiotomy syndrome, Cushing's syndrome, bird-fancier's lung, allergic granulomatous angiitis, benign lymphocytic angiitis, Alport's syndrome, alveolitis such as allergic alveolitis and fibrosing alveolitis, interstitial lung disease, transfusion reaction, leprosy, malaria, parasitic diseases such as leishmaniasis, kypamosomiasis, schitosomiasis, ascariasis, aspergillosis, Sampeter's syndrome, Caplan's syndrome, dengue, endocarditis, endomyocardial fibrosis, diffuse interstitial pulmonary fibrosis, interstitial lung fibrosis, fibrosing mediastinitis, pulmonary fibrosis, idiopathic pulmonary fibrosis, cystic fibrosis, endophthalmitis, erythema elevatum et diutinum, erythroblastosis fetalis, eosinophilic faciitis, Shulman's syndrome, Felty's syndrome, flarisia, cycelitis such as chronic cyclitis, heterochronic cyclitis, iridocyclitis (acute or chronic), or Fuch's cyclitis, Henoch-Schönlein purpura, human immunodeficiency virus (HIV) infection, SCID, acquired immune deficiency syndrome (AIDS), echovirus infection, sepsis (systemic inflammatory response syndrome (SIRS)), endotoxemia, pancreatitis, thyroxicosis, parovirus infection, rubella virus infection, post-vaccination syndromes, congenital rubella infection, Epstein-Barr virus infection, mumps, Evan's syndrome, autoimmune gonadal failure, Sydenham's chorea, post-streptococcal nephritis, thromboangiitis obliterans, thyrotoxicosis, tabes dorsalis, chorioiditis, giant-cell polymyalgia, chronic hypersensitivity pneumonitis, conjunctivitis, such as vernal catarrh, keratoconjunctivitis sicca, and epidemic keratoconjunctivitis, idiopathic nephritic syndrome, minimal change nephropathy, benign familial and ischemia-reperfusion injury, transplant organ reperfusion, retinal autoimmunity, joint inflammation, bronchitis, chronic obstructive airway/pulmonary disease, silicosis, aphthae, aphthous stomatitis, arteriosclerotic disorders (cerebral vascular insufficiency) such as arteriosclerotic encephalopathy and arteriosclerotic retinopathy, aspermiogenese, autoimmune hemolysis, Boeck's
disease, cryoglobulinemia, Dupuytren's contracture, endophthalmitis phacoinfected, enteritis allergica, erythema nodosum leprosum, idiopathic facial paralysis, chronic fatigue syndrome, febris rheumatica, Hamman-Rich's disease, sensorineural hearing loss, haemoglobinuria paroxysmality, hypogonadism, ileitis regionalis, leucopenia, mononucleosis infectiosa, traverse myelitis, primary idiopathic myxedema, nephrosis, ophthalmia symptomatica, orchitis granulomatosa, pancreatitis, polycladulitits acute, pyoderma gangrenosum, Quervain's thyreoiditis, acquired spenic atrophy, non-malignant thymoma, lymphofollicular thymitis, vitiligo, toxic-shock syndrome, food poisoning, conditions involving infiltration of T cells, leukocyte-adhesion deficiency, immune responses associated with acute and delayed hypersensitivity mediated by cytokines and T-lymphocytes, diseases involving leukocyte diapedesis, multiple organ injury syndrome, antigen-antibody complex-mediated diseases, antilgmural basement membrane disease, autoimmune polyendocrinopathies, oophoritis, primary myxedema, autoimmune atrophic gastritis, sympathetic ophthalmia, rheumatic diseases, mixed connective tissue disease, nephrotic syndrome, insulinitis, polyendocrine failure, autoimmune polyglandular syndromes, including polyendocrinopathic syndrome type I, adult-onset idiopathic hypoparathyroidism (AIH), cardiomyopathy such as dilated cardiomyopathy, epidermolysis bullosa acquisita (EBA), hemochromatosis, myocarditis, nephrotic syndrome, primary sclerosing cholangitis, purulent or nonpurulent sinusitis, acute or chronic sinusitis, ethmoid, frontal, maxillary, or sphenoid sinusitis, allergic sinusitis, an eosinophil-related disorder such as eosinophilia, pulmonary infiltration eosinophilia, eosinophilia-nyalgia syndrome, Loffler's syndrome, chronic eosinophilic pneumonia, tropical pulmonary eosinophilia, bronchopneumonic aspergillosis, aspergilloma, or granulomas containing eosinophils, anaphylaxis, spondyloarthropathies, seronegative spondyloarthritides, polyendocrine autoimmune disease, sclerosing cholangitis, sclera, episclera, chronic mucocutaneous candidiasis, Bruton's syndrome, transient hypogammaglobulinemia of infancy, Wiskott-Aldrich syndrome, ataxia telangiectasia syndrome, angiectasis, autoimmune disorders associated with collagen disease, rheumatism such as chronic arthralgias, lymphadenitis, reduction in blood pressure response, vascular dysfunction, tissue injury, cardiovascular ischemia, hyperalgesia, renal ischemia, cerebral ischemia, and disease accompanying vascularization, allergic hypersensitivity disorders, glomerulonephritis, reperfusion injury, ischemic reperfusion disorder, reperfusion injury of myocardial or other tissues, lymphomatous tracheobronchitis, inflammatory dermatoses, dermatoses with acute inflammatory components, multiple organ failure, bullous diseases, renal cortical necrosis, acute purulent meningitis or other central nervous system inflammatory disorders, ocular and orbital inflammatory disorders, granulocyte transfusion-associated syndromes, cytokine-induced toxicity, narcolepsy, acute serious inflammation, chronic intractable inflammation, pyelitis, endarterial hyperplasia, peptic ulcer, valvulitis, and endometriosis.

The phrase “anxiety related disorders” refers to disorders of anxiety, mood, and substance abuse, including but not limited to: depression, generalized anxiety disorders, attention deficit disorder, sleep disorder, hyperactivity disorder, obsessive compulsive disorder, schizophrenia, cognitive disorders, hyperalgesia and sensory disorders. Such disorders include the mild to moderate anxiety, anxiety disorder due to a general medical condition, anxiety disorder not otherwise specified, generalized anxiety disorder, panic attack, panic disorder with agoraphobia, panic disorder without agoraphobia, posttraumatic stress disorder, social phobia, social anxiety, autism, specific phobia, substance-induced anxiety disorder, acute alcohol withdrawal, obsessive compulsive disorder, agoraphobia, monopolar disorders, bipolar disorder I or II, bipolar disorder not otherwise specified,
cyclothymic disorder, depressive disorder, major depressive disorder, mood disorder, substance-induced mood disorder, enhancement of cognitive function, loss of cognitive function associated with but not limited to Alzheimer’s disease, stroke, or traumatic injury to the brain, seizures resulting from disease or injury including but not limited to epilepsy, learning disorders/disabilities, cerebral palsy. In addition, anxiety disorders may apply to personality disorders including but not limited to the following types: paranoid, antisocial, avoidant behavior, borderline personality disorders, dependent, histrionic, narcissistic, obsessive-compulsive, schizoid, and schizotypal.

The term “lipid metabolic disorder” refers to abnormal clinical chemistry levels of cholesterol and triglycerides, wherein elevated levels of these lipids is an indication for atherosclerosis. Additionally, abnormal serum lipid levels may be an indication of various cardiovascular diseases including hypertension, stroke, coronary artery diseases, diabetes and/or obesity.

The phrase “eye abnormality” refers to such potential disorders of the eye as they may be related to atherosclerosis or various ophthalmological abnormalities. Such disorders include but are not limited to the following: retinal dysplasia, various retinopathies, restenosis, retinal artery obstruction or occlusion; retinal degeneration causing secondary atrophy of the retinal vasculature, retinitis pigmentosa, macular dystrophies, Stargardt’s disease, congenital stationary night blindness, choroideremia, gyrate atrophy, Leber’s congenital amaurosis, retinoschisis disorders, Wagner’s syndrome, Usher syndromes, Zellweger syndrome, Saldino-Mainzer syndrome, Senior-Loken syndrome, Bardet-Biedl syndrome, Alport’s syndrome, Alstrom’s syndrome, Cockayne’s syndrome, dysplasia spondyloepiphysaria congenita, Flynn-Aird syndrome, Friedreich ataxia, Hallgren syndrome, Marshall syndrome, Albers-Schonberg disease, Refsum’s disease, Kearns-Sayre syndrome, Waardenburg’s syndrome, Alagile syndrome, myotonic dystrophy, olovopontocerebellar atrophy, Pierre-Marie dursdrome, Sticker syndrome, carotinemia, cystinosis, Wolfram syndrome, Bassen-Kornzweig syndrome, abetalipoproteinemia, incontinentia pigmenti, Batten’s disease, mucopolysaccharidoses, homocystinuria, or memnonidosis. Cataracts are also considered an eye abnormality and are associated with such systemic diseases as: Human Down’s syndrome, Hallerman-Streiff syndrome, Lowe syndrome, galactosemia, Marfan syndrome, Trisomy 15-15 condition, Alport syndrome, myotonic dystrophy, Fabry disease, hypothyroidism, or Conradi syndrome. Other ocular developmental anomalies include: Aniridia, anterior segment and dysgenesis syndrome. Cataracts may also occur as a result of an intraocular infection or inflammation (uveitis).

A “growth inhibitory amount” of an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1271, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO1013, anti-PRO09948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody, PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089,
PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 binding oligopeptide or PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 binding organic molecule is an amount capable of inhibiting the growth of a cell, especially tumor, e.g., cancer cell, either in vitro or in vivo. A “growth inhibitory amount” of an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 binding organic molecule for purposes of inhibiting neoplastic cell growth may be determined empirically and in a routine manner.

A “cytotoxic amount” of an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 binding oligopeptide or PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 binding organic molecule
PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO919563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody, PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 binding oligopeptide or PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 binding organic molecule is an amount capable of causing the destruction of a cell, especially tumor, e.g., cancer cell, either in vitro or in vivo.

A "cytotoxic amount" of an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody, PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 binding oligopeptide or PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111,
PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 binding organic molecule for purposes of inhibiting neoplastic cell growth may be determined empirically and in a routine manner.

The term "antibody" is used in the broadest sense and specifically covers, for example, single anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody monoclonal antibodies (including agonist, antagonist, and neutralizing antibodies), anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody compositions with polyepitopic specificity, polyclonal antibodies, single chain anti-PRO218, anti-

PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibodies, and fragments of anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibodies (see below) as long as they exhibit the desired biological or immunological activity. The term "immunoglobulin" (Ig) is used interchangeably with antibody herein.

An "isolated antibody" is one which has been identified and separated and/or recovered from a component of its natural environment. Contaminant components of its natural environment are materials which would interfere with diagnostic or therapeutic uses for the antibody, and may include enzymes, hormones, and other proteinaceous or nonproteinaceous solutes. The invention provides that the antibody will be purified (1) to greater
than 95% by weight of antibody as determined by the Lowry method, and most preferably more than 99% by weight, (2) to a degree sufficient to obtain at least 15 residues of N-terminal or internal amino acid sequence by use of a spinning cup sequenator, or (3) to homogeneity by SDS-PAGE under reducing or nonreducing conditions using Coomassie blue or, preferably, silver stain. Isolated antibody includes the antibody in situ within recombinant cells since at least one component of the antibody's natural environment will not be present. Ordinarily, however, isolated antibody will be prepared by at least one purification step.

The basic 4-chain antibody unit is a heterotetrameric glycoprotein composed of two identical light (L) chains and two identical heavy (H) chains (an IgM antibody consists of 5 of the basic heterotetramer unit along with an additional polypeptide called J chain, and therefore contain 10 antigen binding sites, while secreted IgA antibodies can polymerize to form polyvalent assemblages comprising 2-5 of the basic 4-chain units along with J chain). In the case of IgGs, the 4-chain unit is generally about 150,000 daltons. Each L chain is linked to a H chain by one covalent disulfide bond, while the two H chains are linked to each other by one or more disulfide bonds depending on the H chain isotype. Each H and L chain also has regularly spaced intrachain disulfide bridges. Each H chain has at the N-terminus, a variable domain (V_\text{H}) followed by three constant domains (C_\text{H}) for each of the \( \alpha \) and \( \gamma \) chains and four C_\text{H} domains for \( \mu \) and \( \epsilon \) isotypes. Each L chain has at the N-terminus, a variable domain (V_L) followed by a constant domain (C_L) at its other end. The V_L is aligned with the V_\text{H} and the C_L is aligned with the first constant domain of the heavy chain (C_\text{H}1). Particular amino acid residues are believed to form an interface between the light chain and heavy chain variable domains. The pairing of a V_\text{H} and V_L together forms a single antigen-binding site. For the structure and properties of the different classes of antibodies, see, e.g., Basic and Clinical Immunology, 8th edition, Daniel P. Stites, Abba I. Terr and Tristram G. Parslow (eds.), Appleton & Lange, Norwalk, CT, 1994, page 71 and Chapter 6.

The L chain from any vertebrate species can be assigned to one of two clearly distinct types, called kappa and lambda, based on the amino acid sequences of their constant domains. Depending on the amino acid sequence of the constant domain of their heavy chains (C_\text{H}), immunoglobulins can be assigned to different classes or isotypes. There are five classes of immunoglobulins: IgA, IgD, IgE, IgG, and IgM, having heavy chains designated \( \alpha \), \( \delta \), \( \epsilon \), \( \gamma \), and \( \mu \), respectively. The \( \gamma \) and \( \alpha \) classes are further divided into subclasses on the basis of relatively minor differences in C_\text{H} sequence and function, e.g., humans express the following subclasses: IgG1, IgG2, IgG3, IgG4, IgA1, and IgA2.

The term "variable" refers to the fact that certain segments of the variable domains differ extensively in sequence among antibodies. The V domain mediates antigen binding and define specificity of a particular antibody for its particular antigen. However, the variability is not evenly distributed across the 110-amino acid span of the variable domains. Instead, the V regions consist of relatively invariant stretches called framework regions (FRs) of 15-30 amino acids separated by shorter regions of extreme variability called "hypervariable regions" that are each 9-12 amino acids long. The variable domains of native heavy and light chains each comprise four FRs, largely adopting a \( \beta \)-sheet configuration, connected by three hypervariable regions, which form loops connecting, and in some cases forming part of, the \( \beta \)-sheet structure. The hypervariable regions in each chain are held together in close proximity by the FRs and, with the hypervariable regions from the other chain, contribute to the formation of the antigen-binding site of antibodies (see Kabat et al., Sequences of Proteins of Immunological Interest, 5th Ed. Public Health Service, National Institutes of Health, Bethesda, MD. (1991)). The constant
domains are not involved directly in binding an antibody to an antigen, but exhibit various effector functions, such as participation of the antibody in antibody dependent cellular cytotoxicity (ADCC).

The term "hypervariable region" when used herein refers to the amino acid residues of an antibody which are responsible for antigen-binding. The hypervariable region generally comprises amino acid residues from a "complementarity determining region" or "CDR" (e.g. around residues 24-34 (L1), 50-56 (L2) and 89-97 (L3) in the V_{L}, and around 1-35 (H1), 50-65 (H2) and 95-102 (H3) in the V_{H} in Kabat et al., *Sequences of Proteins of Immunological Interest*, 5th Ed. Public Health Service, National Institutes of Health, Bethesda, MD. (1991)) and/or those residues from a "hypervariable loop" (e.g. residues 26-32 (L1), 50-52 (L2) and 91-96 (L3) in the V_{L}, and 26-32 (H1), 52-55 (H2) and 96-101 (H3) in the V_{H} in Chothia and Lesk *J. Mol. Biol.* 196:901-917 (1987)).

The term "monoclonal antibody" as used herein refers to an antibody obtained from a population of substantially homogeneous antibodies, i.e., the individual antibodies comprising the population are identical except for possible naturally occurring mutations that may be present in minor amounts. Monoclonal antibodies are highly specific, being directed against a single antigenic site. Furthermore, in contrast to polyclonal antibody preparations which include different antibodies directed against different determinants (epitopes), each monoclonal antibody is directed against a single determinant on the antigen. In addition to their specificity, the monoclonal antibodies are advantageous in that they may be synthesized uncontaminated by other antibodies. The modifier "monoclonal" is not to be construed as requiring production of the antibody by any particular method. For example, the monoclonal antibodies useful in the present invention may be prepared by the hybridoma methodology first described by Kohler et al., *Nature*, 256:495 (1975), or may be made using recombinant DNA methods in bacterial, eukaryotic animal or plant cells (see, e.g., U.S. Patent No. 4,816,567). The "monoclonal antibodies" may also be isolated from phage antibody libraries using the techniques described in Clackson et al., *Nature*, 352:624-628 (1991) and Marks et al., *J Mol Biol.*, 222:581-597 (1991), for example.

The monoclonal antibodies herein include "chimeric" antibodies in which a portion of the heavy and/or light chain is identical with or homologous to corresponding sequences in antibodies derived from a particular species or belonging to a particular antibody class or subclass, while the remainder of the chain(s) is identical with or homologous to corresponding sequences in antibodies derived from another species or belonging to another antibody class or subclass, as well as fragments of such antibodies, so long as they exhibit the desired biological activity (see U.S. Patent No. 4,816,567; and Morrison et al., *Proc. Natl. Acad. Sci. USA*, 81:6851-6855 (1984)). Chimeric antibodies of interest herein include "humanized" antibodies comprising variable domain antigen-binding sequences derived from a non-human primate (e.g. Old World Monkey, Ape etc.), and human constant region sequences.

An "intact" antibody is one which comprises an antigen-binding site as well as a C_{L} and at least heavy chain constant domains, C_{H} 1, C_{H} 2 and C_{H} 3. The constant domains may be native sequence constant domains (e.g. human native sequence constant domains) or amino acid sequence variant thereof. Preferably, the intact antibody has one or more effector functions.

"Antibody fragments" comprise a portion of an intact antibody, preferably the antigen binding or variable region of the intact antibody. Examples of antibody fragments include Fab, Fab', F(ab')_{2}, and Fv fragments; diabodies; linear antibodies (see U.S. Patent No. 5,641,870, Example 2; Zapata et al., *Protein Eng.*, 8(10):
1057-1062 [1995]); single-chain antibody molecules; and multispecific antibodies formed from antibody fragments.

Papain digestion of antibodies produces two identical antigen-binding fragments, called "Fab" fragments, and a residual "Fc" fragment, a designation reflecting the ability to crystallize readily. The Fab fragment consists of an entire L chain along with the variable region domain of the H chain (VH), and the first constant domain of one heavy chain (C\text{H}1). Each Fab fragment is monovalent with respect to antigen binding, i.e., it has a single antigen-binding site. Pepsin treatment of an antibody yields a single large F(ab')2 fragment which roughly corresponds to two disulfide linked Fab fragments having divalent antigen-binding activity and is still capable of cross-linking antigen. Fab' fragments differ from Fab fragments by having additional few residues at the carboxy terminus of the C\text{H}1 domain including one or more cysteines from the antibody hinge region. Fab'-SH is the designation herein for Fab' in which the cysteine residue(s) of the constant domains bear a free thiol group. F(ab')2 antibody fragments originally were produced as pairs of Fab' fragments which have hinge cysteines between them. Other chemical couplings of antibody fragments are also known.

The Fc fragment comprises the carboxy-terminal portions of both H chains held together by disulfides. The effector functions of antibodies are determined by sequences in the Fc region, which region is also the part recognized by Fc receptors (FcR) found on certain types of cells.

"Fv" is the minimum antibody fragment which contains a complete antigen-recognition and-binding site. This fragment consists of a dimer of one heavy- and one light-chain variable region domain in tight, non-covalent association. From the folding of these two domains emanate six hypervariable loops (3 loops each from the H and L chain) that contribute the amino acid residues for antigen binding and confer antigen binding specificity to the antibody. However, even a single variable domain (or half of an Fv comprising only three CDRs specific for an antigen) has the ability to recognize and bind antigen, although at a lower affinity than the entire binding site.

"Single-chain Fv" also abbreviated as "sFv" or "scFv" are antibody fragments that comprise the VH and VL antibody domains connected into a single polypeptide chain. Preferably, the sFv polypeptide further comprises a polypeptide linker between the VH and VL domains which enables the sFv to form the desired structure for antigen binding. For a review of sFv, see Pluckthun in The Pharmacology of Monoclonal Antibodies, vol. 113, Rosenberg and Moore eds., Springer-Verlag, New York, pp. 269-315 (1994); Borrebaeck 1995, infra.

The term "diabodies" refers to small antibody fragments prepared by constructing sFv fragments (see preceding paragraph) with short linkers (about 5-10 residues) between the VH and VL domains such that inter-chain but not intra-chain pairing of the V domains is achieved, resulting in a bivalent fragment, i.e., fragment having two antigen-binding sites. Bispecific diabodies are heterodimers of two "crossover" sFv fragments in which the VH and VL domains of the two antibodies are present on different polypeptide chains. Diabodies are described more fully in, for example, EP 404,097; WO 93/11161; and Hollinger et al., Proc. Natl. Acad. Sci. USA, 90:6444-6448 (1993).

"Humanized" forms of non-human (e.g., rodent) antibodies are chimeric antibodies that contain minimal sequence derived from the non-human antibody. For the most part, humanized antibodies are human immunoglobulins (recipient antibody) in which residues from a hypervariable region of the recipient are replaced by residues from a hypervariable region of a non-human species (donor antibody) such as mouse, rat, rabbit or non-human primate having the desired antibody specificity, affinity, and capability. In some instances, framework
region (FR) residues of the human immunoglobulin are replaced by corresponding non-human residues. Furthermore, humanized antibodies may comprise residues that are not found in the recipient antibody or in the donor antibody. These modifications are made to further refine antibody performance. In general, the humanized antibody will comprise substantially all of at least one, and typically two, variable domains, in which all or substantially all of the hypervariable loops correspond to those of a non-human immunoglobulin and all or substantially all of the FRs are those of a human immunoglobulin sequence. The humanized antibody optionally also will comprise at least a portion of an immunoglobulin constant region (Fc), typically that of a human immunoglobulin. For further details, see Jones et al., Nature 321:522-525 (1986); Riechmann et al., Nature 332:323-329 (1988); and Presta, Curr. Opin. Struct. Biol. 2:593-596 (1992).

A "species-dependent antibody," e.g., a mammalian anti-human IgG antibody, is an antibody which has a stronger binding affinity for an antigen from a first mammalian species than it has for a homologue of that antigen from a second mammalian species. Normally, the species-dependent antibody "binds specifically" to a human antigen (i.e., has a binding affinity (Kd) value of no more than about 1 x 10^-7 M, preferably no more than about 1 x 10^-8 and most preferably no more than about 1 x 10^-9 M) but has a binding affinity for a homologue of the antigen from a second non-human mammalian species which is at least about 50 fold, or at least about 500 fold, or at least about 1000 fold, weaker than its binding affinity for the human antigen. The species-dependent antibody can be of any of the various types of antibodies as defined above, but preferably is a humanized or human antibody.

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PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 binding oligopeptides usually are or are at least about 5 amino acids in length, alternatively are or are at least about 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, or 100 amino acids in length or more, wherein such oligopeptides that are capable of binding, preferably specifically, to a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO656, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide as described herein. PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 binding oligopeptides may be identified without undue experimentation using well known techniques. In this regard, it is noted that techniques for screening oligopeptide libraries for oligopeptides that are capable of specifically binding to a polypeptide target are well known in the art (see, e.g., U.S. Patent Nos. 5,556,762, 5,750,733, 4,708,871, 4,833,092, 5,223,409, 5,403,484, 5,571,689, 5,663,143; PCT Publication Nos. WO 84/03506 and WO84/03564; Geysen et al., Proc. Natl. Acad. Sci. U.S.A., 81:3998-4002 (1984); Geysen et al., Proc. Natl. Acad. Sci. U.S.A., 82:178-182 (1985); Geysen et al., in Synthetic Peptides as Antigens, 130-149 (1986); Geysen et al., J. Immunol. Meth., 102:259-274 (1987); Schoofs et al., J. Immunol., 140:611-616 (1988), Cwirla, S. E. et al. (1990) Proc. Natl. Acad. Sci. USA, 87:6378; Lowman, H. B. et al. (1991) Biochemistry, 30:10832; Clackson, T. et al. (1991) Nature, 352: 624; Marks, J. D. et al. (1991), J. Mol. Biol., 222:581; Kang, A.S. et al. (1991) Proc. Natl. Acad. Sci. USA, 88:8363, and Smith, G. P. (1991) Current Opin. Biotechnol., 2:668).

A "PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO656, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 binding organic molecule" is an organic molecule other than an oligopeptide or antibody as defined herein that binds, preferably specifically, to a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide as described herein. PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 binding organic molecule" is an organic molecule other than an oligopeptide or antibody as defined herein that binds, preferably specifically,
PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 binding organic molecules may be identified and chemically synthesized using known methodology (see, e.g., PCT Publication Nos. WO00/00823 and WO00/39585). PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 binding organic molecules are usually less than about 2000 daltons in size, alternatively less than about 1500, 750, 500, 250 or 200 daltons in size, wherein such organic molecules that are capable of binding, preferably specifically, to a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide as described herein may be identified without undue experimentation using well known techniques. In this regard, it is noted that techniques for screening organic molecule libraries for molecules that are capable of binding to a polypeptide target are well known in the art (see, e.g., PCT Publication Nos. WO00/00823 and WO00/39585).

An antibody, oligopeptide or other organic molecule "which binds" an antigen of interest, e.g. a tumor-associated polypeptide antigen target, is one that binds the antigen with sufficient affinity such that the antibody, oligopeptide or other organic molecule is preferably useful as a diagnostic and/or therapeutic agent in targeting a cell or tissue expressing the antigen, and does not significantly cross-react with other proteins. The extent of binding of the antibody, oligopeptide or other organic molecule to a "non-target" protein will be less than about 10% of the binding of the antibody, oligopeptide or other organic molecule to its particular target protein as determined by fluorescence activated cell sorting (FACS) analysis or radioimmunoprecipitation (RIA). With regard to the binding of an antibody, oligopeptide or other organic molecule to a target molecule, the term "specific binding" or "specifically binds to" or is "specific for" a particular polypeptide or an epitope on a particular polypeptide target means binding that is measurably different from a non-specific interaction. Specific binding can be measured, for example, by determining binding of a molecule compared to binding of a control molecule, which generally is a molecule of similar structure that does not have binding activity. For example, specific binding can be determined by competition with a control molecule that is similar to the target, for example, an excess of non-labeled target. In this case, specific binding is indicated if the binding of the labeled target to a probe is competitively inhibited by excess unlabeled target. The term "specific binding" or "specifically binds to" or is "specific for" a particular polypeptide or an epitope on a particular polypeptide target as used herein can be exhibited, for example, by a molecule having a Kd for the target of at least about 10^-4 M, alternatively at least about 10^-5 M, alternatively at least about 10^-6 M, alternatively at least about 10^-7 M, alternatively at least about 10^-8 M, alternatively at least about 10^-9 M, alternatively at least about 10^-10 M, alternatively at least about 10^-11 M, alternatively at least about 10^-12 M, or greater. The term "specific binding" refers to binding where a molecule
binds to a particular polypeptide or epitope on a particular polypeptide without substantially binding to any other polypeptide or polypeptide epitope.

An antibody, oligopeptide or other organic molecule that "inhibits the growth of tumor cells expressing a “PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO1017, PRO474, PRO5238, PRO5169, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346" or a "growth inhibitory" antibody, oligopeptide or other organic molecule is one which results in measurable growth inhibition of cancer cells expressing or overexpressing the appropriate PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide. The PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide may be a transmembrane polypeptide expressed on the surface of a cancer cell or may be a polypeptide that is produced and secreted by a cancer cell. Preferred growth inhibitory anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibodies, oligopeptides or organic molecules inhibit growth of PRO218-, PRO228-, PRO271-, PRO273-, PRO295-, PRO302-, PRO305-, PRO326-, PRO386-, PRO655-, PRO162-, PRO788-, PRO792-, PRO940-, PRO941-, PRO1004-, PRO1012-, PRO1016-, PRO474-, PRO5238-, PRO1069-, PRO1111-, PRO1113-, PRO1130-, PRO1195-, PRO1271-, PRO1865-, PRO1879-, PRO3446-, PRO3543-, PRO4329-, PRO4352-, PRO5733-, PRO9859-, PRO9864-, PRO9904-, PRO9907-, PRO10013-, PRO90948-, PRO28694-, PRO16089-, PRO19563-, PRO19675-, PRO20084-, PRO21434-, PRO50332-, PRO38465- or PRO346-expressing tumor cells by or by greater than 20%, preferably from about 20% to about 50%, and even more preferably, by or by greater than 50% (e.g., from about 50% to about 100%) as compared to the appropriate control, the control typically being tumor cells not treated with the antibody, oligopeptide or other organic molecule being tested. Growth inhibition can be measured at an antibody concentration of about 0.1 to 30 μg/ml or about 0.5 nM to 200 nM in cell culture, where the growth inhibition is determined 1-10 days after exposure of the tumor cells to the antibody. Growth inhibition of tumor cells in vivo can be determined in various
The antibody is growth inhibitory \textit{in vivo} if administration of the anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO28694, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody at about 1 \( \mu \text{g/kg} \) to about 100 \( \mu \text{g/kg} \) body weight results in reduction in tumor size or tumor cell proliferation within about 5 days to 3 months from the first administration of the antibody, preferably within about 5 to 30 days.

An antibody, oligopeptide or other organic molecule which "induces apoptosis" is one which induces programmed cell death as determined by binding of annexin V, fragmentation of DNA, cell shrinkage, dilation of endoplasmic reticulum, cell fragmentation, and/or formation of membrane vesicles (called apoptotic bodies). The cell is usually one which overexpresses a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99094, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide. Preferably the cell is a tumor cell, e.g., a prostate, breast, ovarian, stomach, endometrial, lung, kidney, colon, bladder cell. Various methods are available for evaluating the cellular events associated with apoptosis. For example, phosphatidyl serine (PS) translocation can be measured by annexin binding; DNA fragmentation can be evaluated through DNA laddering; and nuclear/chromatin condensation along with DNA fragmentation can be evaluated by any increase in hypodiploid cells. Preferably, the antibody, oligopeptide or other organic molecule which induces apoptosis is one which results in or in about to 50 fold, preferably in or in about 5 to 50 fold, and most preferably in or in about 10 to 50 fold, induction of annexin binding relative to untreated cell in an annexin binding assay.

Antibody "effector functions" refer to those biological activities attributable to the Fc region (a native sequence Fc region or amino acid sequence variant Fc region) of an antibody, and vary with the antibody isotype. Examples of antibody effector functions include: C1q binding and complement dependent cytotoxicity; Fc receptor binding; antibody-dependent cell-mediated cytotoxicity (ADCC); phagocytosis; down regulation of cell surface receptors (e.g., B cell receptor); and B cell activation.

"Antibody-dependent cell-mediated cytotoxicity" or "ADCC" refers to a form of cytotoxicity in which secreted Ig bound onto Fc receptors (FcRs) present on certain cytotoxic cells (e.g., Natural Killer (NK) cells, neutrophils, and macrophages) enable these cytotoxic effector cells to bind specifically to an antigen-bearing target cell and subsequently kill the target cell with cytotoxins. The antibodies "arm" the cytotoxic cells and are absolutely required for such killing. The primary cells for mediating ADCC, NK cells, express Fc\( \gamma \)RII only, whereas monocytes express Fc\( \gamma \)RI, Fc\( \gamma \)RII and Fc\( \gamma \)RIII. FcR expression on hematopoietic cells is summarized in Table 3 on page 464 of Ravetch and Kinet, \textit{Annu. Rev. Immunol.} 9:457-92 (1991). To assess ADCC activity of a molecule of interest, an \textit{in vitro} ADCC assay, such as that described in US Patent No. 5,500,362 or 5,821,337.
may be performed. Useful effector cells for such assays include peripheral blood mononuclear cells (PBMC) and Natural Killer (NK) cells. Alternatively, or additionally, ADCC activity of the molecule of interest may be assessed in vivo, e.g., in an animal model such as that disclosed in Clynes et al. Proc. Natl. Acad. Sci. U.S.A. 95:652-656 (1998).

"Fc receptor" or "FcR" describes a receptor that binds to the Fc region of an antibody. The preferred FcR is a native sequence human FcR. Moreover, a preferred FcR is one which binds an IgG antibody (a gamma receptor) and includes receptors of the FcγRI, FcγRII and FcγRIII subclasses, including allelic variants and alternatively spliced forms of these receptors. FcγRII receptors include FcγRIIA (an "activating receptor") and FcγRIIB (an "inhibiting receptor"), which have similar amino acid sequences that differ primarily in the cytoplasmic domains thereof. Activating receptor FcγRIIA contains an immunoreceptor tyrosine-based activation motif (ITAM) in its cytoplasmic domain. Inhibiting receptor FcγRIIB contains an immunoreceptor tyrosine-based inhibition motif (ITIM) in its cytoplasmic domain. (see review M. in Daecon. Annu. Rev. Immunol. 15:203-234 (1997)). FcRs are reviewed in Ravetch and Kinet, Annu. Rev. Immunol. 9:457-492 (1991); Capel et al., Immunomethods 4:25-34 (1994); and de Haas et al., J. Lab. Clin. Med. 126:330-41 (1995). Other FcRs, including those to be identified in the future, are encompassed by the term "FcR" herein. The term also includes the neonatal receptor, FcRn, which is responsible for the transfer of maternal IgGs to the fetus (Guyer et al., J. Immunol. 117:587 (1976) and Kim et al., J. Immunol. 24:249 (1994)).

"Human effector cells" are leukocytes which express one or more FcRs and perform effector functions. Preferably, the cells express at least FcγRII and perform ADCC effector function. Examples of human leukocytes which mediate ADCC include peripheral blood mononuclear cells (PBMC), natural killer (NK) cells, monocytes, cytotoxic T cells and neutrophils; with PBMCs and NK cells being preferred. The effector cells may be isolated from a native source, e.g., from blood.

"Complement dependent cytotoxicity" or "CDC" refers to the lysis of a target cell in the presence of complement. Activation of the classical complement pathway is initiated by the binding of the first component of the complement system (C1q) to antibodies (of the appropriate subclass) which are bound to their cognate antigen. To assess complement activation, a CDC assay, e.g., as described in Gazzano-Santoro et al., J. Immunol. Methods 202:163 (1996), may be performed.

The terms "cancer" and "cancerous" refer to or describe the physiological condition in mammals that is typically characterized by unregulated cell growth. Examples of cancer include but are not limited to, carcinoma, lymphoma, blastoma, sarcoma, and leukemia. More particular examples of such cancers include squamous cell cancer, lung cancer (including small-cell lung cancer, non-small cell lung cancer, adenocarcinoma of the lung, and squamous carcinoma of the lung), cancer of the peritoneum, hepatocellular cancer, gastric or stomach cancer (including gastrointestinal cancer), pancreatic cancer, glioblastoma, cervical cancer, ovarian cancer, liver cancer, bladder cancer, hepatoma, breast cancer, colon cancer, colorectal cancer, endometrial or uterine carcinoma, salivary gland carcinoma, kidney or renal cancer, liver cancer, prostate cancer, vulval cancer, thyroid cancer, hepatic carcinoma and various types of head and neck cancer, as well as B-cell lymphoma (including low grade/follicular non-Hodgkin's lymphoma (NHL); small lymphocytic (SL) NHL; intermediate grade/follicular NHL; intermediate grade diffuse NHL; high grade immunoblastic NHL; high grade lymphoblastic NHL; high grade small non-cleaved cell NHL; bulky disease NHL; mantle cell lymphoma; AIDS-related lymphoma; and
Waldenstrom's Macroglobulinemia); chronic lymphocytic leukemia (CLL); acute lymphoblastic leukemia (ALL); hairy cell leukemia; chronic myeloblastic leukemia; and post-transplant lymphoproliferative disorder (PTLD). Preferably, the cancer comprises a tumor that expresses an IGF receptor, more preferably breast cancer, lung cancer, colorectal cancer, or prostate cancer, and most preferably breast or prostate cancer.

A "chemotherapeutic agent" is a chemical compound useful in the treatment of cancer. Examples of chemotherapeutic agents include alkylating agents such as thiotepa and CYTOXAN® cyclophosphamide; alkyl sulfonates such as busulfan, imposulfan and pipsosulfan; aziridines such as benzdopa, carboquone, mertredopa, and uredopa; ethylenimines and methylamelines including altretamine, triethylenemelamine, trietylenephosphoramide, triethylene thiophosphoramide and trimethylolmelamine; acetogenins (especially bullatacin and bullatacinone); a camptothecin (including the synthetic analogue topotecan); bryostatin; calyssatin; CC-1065 (including its adozelesin, carzelesin and bizelesin synthetic analogues); cryptophycins (particularly cryptophycin 1 and cryptophycin 8); dolastatin; duocarmycin (including the synthetic analogues, KW-2189 and CB1-TM1); eleutherobin; pancratistatin; a sarcodictyin; spogwortin; nitrogen mustard such as chlorambucil, clomaphazine, clorophosphamide, estramustine, ifosfamide, mechlorethamine, mechlorethamine oxide hydrochloride, melfalan, novembichin, phenesterine, prednimustine, trofosfamide, uracil mustard; nitrosourea such as carmustine, chlorozotocin, fotemustine, lomustine, nimustine, and ranimustine; antibiotics such as the enediyne antibiotics (e.g., calicheamicin, especially calicheamicin gamma I1 and calicheamicin omega I1 (see, e.g., Agnew, Chem Int. Ed. Engl., 33: 183-186 (1994)); dynemicin, including dynemicin A; bisphosphonates, such as clodronate; an esperamicin; as well as neocarzinostatin chromophore and related chromoprotein enediyne antibiotic chromophores), aclacinomycins, actinomycin, authramycin, azaserine, bleomycins, caetnonymycin, carabacin, carminomycin, carzinophilin, chromomycin, dactinomycin, daunorubicin, detorubicin, 6-diazo-5-oxo-L-norleucine, ADRIAMYCIN® doxorubicin (including morpholino-doxorubicin, cyanomorpholino-doxorubicin, 2-pyrrolino-doxorubicin and deoxydoxorubicin), epirubicin, esorubicin, idarubicin, marcellomycin, mitomycins such as mitomycin C, mycophenolic acid, norgalamycin, olivomycins, peplomycin, potifromycin, puromycin, quelamycin, rorubicin, streptonigrin, streptozocin, tabercidin, ubenimex, zinostatin, zorubicin; anti metabolites such as methotrexate and 5-fluorouracil (5-FU); folic acid analogues such as denopterin, methotrexate, pteropterin, trimetrexate; purine analogs such as fludarabine, 6-mercaptopurine, thioguanine, pyrimidine analogs such as ancitabine, azacitidine, 6-azauridine, carnofur, cytarabine, deoxouridine, doxifluridine, enocitabine, floxuridine; androgens such as calusterone, dromostanolone propionate, epithiotanol, meptiotostane, testolactone; anti- andrinals such as aminoglutethimide, mitotane, trilostane; folic acid replensher such as frolinic acid; acetalone; aldophosphamide glycoside; aminolevulinic acid; eniluracil; ansamycin; bestrabucil; bisantrene; edatrazate; defofamine; demecocoline; diaziqone; elfornithine; elliptinium acetate; an epithelone; etogolucid; gallium nitrate; hydroxyurea; lentinam; lonidamine; maytansinoids such as maytansine and ansamitocins; mitoguazone; mitoxantrone; mopidamol; nitraerine; pentostatin; phenemet; pirarubicin; losoxantrone; podophyllinic acid; 2-ethylhydrazide; procabazine; PSK® polysaccharide complex (JHS Natural Products, Eugene, OR); razonexane; rhizoxin; sizofiran; spirogermanium; tenuazonic acid; triaziqone; 2,2',2'-trichlorotriethylamine; trichotheccenes (especially T-2 toxin, verrucarin A, rosidin A and anguidine); urethan; vindesine; dacarbazine; mannomustine; mitobronitol; mitolactol; pipobroman; gacytosine; arabinoside ("Ara-C"); cyclophosphamide; thiotope; taxoids, e.g., TAXOL® paclitaxel (Bristol-Myers Squibb Oncology, Princeton, N.J.),
ABRAXANE™, cremophor-free, albumin-engineered nanoparticle formulation of paclitaxel (American Pharmaceutical Partners, Schaumberg, Illinois), and TAXOTERE® doxetaxel (Rhône-Poulenc Rorer, Antony, France); chlorambucil; GEMZAR® gemcitabine; 6-thioguanine; mercaptopurine; methotrexate; platinum analogs such as cisplatin and carboplatin; vinblastine; platinum; etoposide (VP-16); ifosfamide; mitoxantrone; vincristine; NAVELBINE® vinorelbine; novantrone; teniposide; etadretaxel; daunomycin; aminopterin; xeloda; i bandronate; CPT-11; topoisomerase inhibitor RFS 2000; difluoromethylornithine (DMFO); retinoids such as retinoic acid; capecitabine; and pharmaceutically acceptable salts, acids or derivatives of any of the above.

Also included in this definition are anti-hormonal agents that act to regulate or inhibit hormone action on tumors such as anti-estrogens and selective estrogen receptor modulators (SERMs), including, for example, tamoxifen (including NOLVADEX® tamoxifen), raloxifene, droloxifene, 4-hydroxytamoxifen, trioxifene, keoxifene, LY117018, onapristone, and FARESTON® toremifene; aromatase inhibitors that inhibit the enzyme aromatase, which regulates estrogen production in the adrenal glands, such as, for example, 4(5)-imidazoles, aminogluthethimide, MEGASE® megestrol acetate, AROMASIN® exemestane, formestane, fadrozole, RIVISOR® vorozole, FEMARA® letrozole, and ARIMIDEX® anastrozole; and anti-androgens such as flutamide, nilutamide, bicalutamide, leuprolide, and goserelin; as well as troxacetimide (a 1,3-dioxolane nucleoside cytosine analog); antisense oligonucleotides, particularly those which inhibit expression of genes in signaling pathways implicated in abherant cell proliferation, such as, for example, PKC-alpha, Ralfl and H-Ras; ribozymes such as a VEGF expression inhibitor (e.g., ANGIOZYME® ribozyme) and a HER2 expression inhibitor; vaccines such as gene therapy vaccines, for example, ALLOVECTIN® vaccine, LFUVECTIN® vaccine, and VAXID® vaccine; PROLEUKIN® rIL-2; LURTOTECAN® topoisomerase 1 inhibitor; ABARELIX® rmRH; and pharmaceutically acceptable salts, acids or derivatives of any of the above.

The terms "cell proliferative disorder" and "proliferative disorder" refer to disorders that are associated with some degree of abnormal cell proliferation. In one aspect of the invention, the cell proliferative disorder is cancer.

"Tumor", as used herein, refers to all neoplastic cell growth and proliferation, whether malignant or benign, and all pre-cancerous and cancerous cells and tissues.

An antibody, oligopeptide or other organic molecule which "induces cell death" is one which causes a viable cell to become nonviable. The cell is one which expresses a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO1047, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, preferably a cell that overexpresses a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide as compared to a normal cell of the same tissue type. The PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305,
PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO36 polypeptide may be a transmembrane polypeptide expressed on the surface of a cancer cell or may be a polypeptide that is produced and secreted by a cancer cell. Preferably, the cell is a cancer cell, e.g., a breast, ovarian, stomach, endometrial, salivary gland, lung, kidney, colon, thyroid, pancreatic or bladder cell. Cell death in vitro may be determined in the absence of complement and immune effector cells to distinguish cell death induced by antibody-dependent cell-mediated cytotoxicity (ADCC) or complement dependent cytotoxicity (CDC). Thus, the assay for cell death may be performed using heat inactivated serum (i.e., in the absence of complement) and in the absence of immune effector cells. To determine whether the antibody, oligopeptide or other organic molecule is able to induce cell death, loss of membrane integrity as evaluated by uptake of propidium iodide (PI), trypan blue (see Moore et al., Cytotherapy 17:1-11 (1995)) or 7AAD can be assessed relative to untreated cells. Preferred cell death-inducing antibodies, oligopeptides or other organic molecules are those which induce PI uptake in the PI uptake assay in BT474 cells.

As used herein, the term "immunoadhesion" designates antibody-like molecules which combine the binding specificity of a heterologous protein (an "adhesion") with the effector functions of immunoglobulin constant domains. Structurally, the immunoadhesions comprise a fusion of an amino acid sequence with the desired binding specificity which is other than the antigen recognition and binding site of an antibody (i.e., is "heterologous"), and an immunoglobulin constant domain sequence. The adhesion part of an immunoadhesion molecule typically is a contiguous amino acid sequence comprising at least the binding site of a receptor or a ligand. The immunoglobulin constant domain sequence in the immunoadhesion may be obtained from any immunoglobulin, such as IgG-1, IgG-2, IgG-3, or IgG-4 subtypes, IgA (including IgA-1 and IgA-2), IgE, IgD or IgM.

The word "label" when used herein refers to a detectable compound or composition which is conjugated directly or indirectly to the antibody so as to generate a "labeled" antibody. The label may be detectable by itself (e.g. radioisotope labels or fluorescent labels) or, in the case of an enzymatic label, may catalyze chemical alteration of a substrate compound or composition which is detectable.

"Replication-preventing agent" is an agent wherein replication, function, and/or growth of the cells is inhibited or prevented, or cells are destroyed, no matter what the mechanism, such as by apoptosis, angiostasis, cytosis, tumoricide, mytosis inhibition, blocking cell cycle progression, arresting cell growth, binding to tumors, acting as cellular mediators, etc. Such agents include a chemotherapeutic agent, cytotoxic agent, cytokine, growth-inhibitory agent, or anti-hormonal agent, e.g., an anti-estrogen compound such as tamoxifen, an anti-progesterone such as onapristone (see, EP 616 812); or an anti-androgen such as flutamide, as well as aromidase inhibitors, or a hormonal agent such as an androgen.

The term "cytotoxic agent" as used herein refers to a substance that inhibits or prevents the function of cells and/or causes destruction of cells. The term is intended to include radioactive isotopes (e.g., At211, I131, T125, Y90, Re186, Re188, Sm153, Bi212, P32 and radioactive isotopes of Lu), chemotherapeutic agents e.g. methotrexate, adriamicin, vinca alkaloids (vincreistine, vinblastine, etoposide), doxorubicin, melphalan, mitomycin C,
chlorambucil, daunorubicin or other intercalating agents, enzymes and fragments thereof such as nucleolytic enzymes, antibiotics, and toxins such as small molecule toxins or enzymatically active toxins of bacterial, fungal, plant or animal origin, including fragments and/or variants thereof, and the various antitumor or anticancer agents disclosed below. Other cytotoxic agents are described below. A tumoricidal agent causes destruction of tumor cells.

Preferred cytotoxic agents herein for the specific tumor types to use in combination with the antagonists herein are as follows:
1. Prostate cancer: androgens, docetaxel, paclitaxel, estramustine, doxorubicin, mitoxantrone, antibodies to ErbB2 domain(s) such as 2C4 (WO 01/00245; hybridoma ATCC HB-12697), which binds to a region in the extracellular domain of ErbB2 (e.g., any one or more residues in the region from about residue 22 to about residue 584 of ErbB2, inclusive), AVASTIN™ anti-vascular endothelial growth factor (VEGF), TARCEVA™ OSI-774 (erlotinib) (Genentech and OSI Pharmaceuticals), or other epidermal growth factor receptor tyrosine kinase inhibitors (EGFR TKIs).

2. Stomach cancer: 5-fluorouracil (5FU), XELODA™ capecitabine, methotrexate, etoposide, cisplatin/carboplatin, paclitaxel, docetaxel, gemcitabine, doxorubicin, and CPT-11 (camptothecin-11; irinotecan, USA Brand Name: CAMPTOSAR®).

3. Pancreatic cancer: gemcitabine, 5FU, XELODA™ capecitabine, CPT-11, docetaxel, paclitaxel, cisplatin, carboplatin, TARCEVA™ erlotinib, and other EGFR TKIs.

4. Colorectal cancer: 5FU, XELODA™ capecitabine, CPT-11, oxaliplatin, AVASTIN™ anti-VEGF, TARCEVA™ erlotinib and other EGFR TKIs, and ERBITUX™ (formerly known as IMC-C225) human-murine-chimerized monoclonal antibody that binds to EGFR and blocks the ability of EGF to initiate receptor activation and signaling to the tumor.

5. Renal cancer: IL-2, interferon alpha, AVASTIN™ anti-VEGF, MEGACE™ (Megestrol acetate) progesterin, vinblastine, TARCEVA™ erlotinib, and other EGFR TKIs.

(vincristine and vinblastine), taxanes, and topoisomerase II inhibitors such as doxorubicin, epirubicin, daunorubicin, etoposide, and bleomycin. Those agents that arrest G1 also spill over into S-phase arrest, for example, DNA alkylating agents such as tamoxifen, prednisone, dacarbazine, mechloroethamine, cisplatin, methotrexate, 5-fluorouracil, and ara-C. Further information can be found in The Molecular Basis of Cancer, Mendelsohn and Israel, eds., Chapter 1, entitled "Cell cycle regulation, oncogenes, and antineoplastic drugs" by Murakami et al. (WB Saunders: Philadelphia, 1995), especially p. 13. The taxanes (paclitaxel and docetaxel) are anticancer drugs both derived from the yew tree. Docetaxel (TAXOTERE®, Rhone-Poulenc Rorer), derived from the European yew, is a semisynthetic analogue of paclitaxel (TAXOL®, Bristol-Myers Squibb). Paclitaxel and docetaxel promote the assembly of microtubules from tubulin dimers and stabilize microtubules by preventing depolymerization, which results in the inhibition of mitosis in cells.

"Doxorubicin" is an anthracycline antibiotic. The full chemical name of doxorubicin is (8S-cis)-10-[[3-amino-2,3,6-trideoxy-α-L-lyxo-hexopyranosyl(oxy)]-7,8,9,10-tetrahydro-6,8,11-trihydroxy-8-(hydroxyacetyl)-1-methoxy-5,12-naphthacenedione.

The term "cytokine" is a generic term for proteins released by one cell population which act on another cell as intercellular mediators. Examples of such cytokines are lymphokines, monokines, and traditional polypeptide hormones. Included among the cytokines are growth hormone such as human growth hormone, N-methionyl human growth hormone, and bovine growth hormone; parathyroid hormone; thyroxine; insulin; proinsulin; relaxin; prorelaxin; glycoprotein hormones such as follicle stimulating hormone (FSH), thyroid stimulating hormone (TSH), and luteinizing hormone (LH); hepatic growth factor; fibroblast growth factor; prolactin; placental lactogen; tumor necrosis factor-α and -β; Mullerian-inhibiting substance; mouse gonadotropin-associated peptide; inhibin; activin; vascular endothelial growth factor; integrin; thrombopoietin (TPO); nerve growth factors such as NGF-β; platelet-growth factor; transforming growth factors (TGFs) such as TGF-α and TGF-β; insulin-like growth factor-I and -II; erythropoietin (EPO); osteoinductive factors; interferons such as interferon-α, -β, and -γ; colony stimulating factors (CSFs) such as macrophage-CSF (M-CSF); granulocyte-macrophage-CSF (GM-CSF); and granulocyte-CSF (G-CSF); interleukins (ILs) such as IL-1, IL-1a, IL-2, IL-3, IL-4, IL-5, IL-6, IL-7, IL-8, IL-9, IL-11, IL-12; a tumor necrosis factor such as TNF-α or TNF-β; and other polypeptide factors including LIF and kit ligand (KL). As used herein, the term cytokine includes proteins from natural sources or from recombinant cell culture and biologically active equivalents of the native sequence cytokines.

The term “package insert” is used to refer to instructions customarily included in commercial packages of therapeutic products, that contain information about the indications, usage, dosage, administration, contraindications and/or warnings concerning the use of such therapeutic products.

The term “gene” refers to (a) a gene containing at least one of the DNA sequences disclosed herein; (b) any DNA sequence that encodes the amino acid sequence encoded by the DNA sequences disclosed herein and/or; © any DNA sequence that hybridizes to the complement of the coding sequences disclosed herein. Preferably, the term includes coding as well as noncoding regions, and preferably includes all sequences necessary for normal gene expression.

The term “gene targeting” refers to a type of homologous recombination that occurs when a fragment of genomic DNA is introduced into a mammalian cell and that fragment locates and recombines with endogenous
homologous sequences. Gene targeting by homologous recombination employs recombinant DNA technologies to replace specific genomic sequences with exogenous DNA of particular design.

The term “homologous recombination” refers to the exchange of DNA fragments between two DNA molecules or chromatids at the site of homologous nucleotide sequences.

The term “target gene” (alternatively referred to as “target gene sequence” or “target DNA sequence”) refers to any nucleic acid molecule, polynucleotide, or gene to be modified by homologous recombination. The target sequence includes an intact gene, an exon or intron, a regulatory sequence or any region between genes. The target gene may comprise a portion of a particular gene or genetic locus in the individual’s genomic DNA.

“Disruption” of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 gene occurs when a fragment of genomic DNA locates and recombines with an endogenous homologous sequence wherein the disruption is a deletion of the native gene or a portion thereof, or a mutation in the native gene or wherein the disruption is the functional inactivation of the native gene. Alternatively, sequence disruptions may be generated by nonspecific insertional inactivation using a gene trap vector (i.e. non-human transgenic animals containing and expressing a randomly inserted transgene; see for example U.S. Pat. No. 6,436,707 issued August 20, 2002). These sequence disruptions or modifications may include insertions, missense, frameshift, deletion, or substitutions, or replacements of DNA sequence, or any combination thereof. Insertions include the insertion of entire genes, which may be of animal, plant, fungal, insect, prokaryotic, or viral origin. Disruption, for example, can alter the normal gene product by inhibiting its production partially or completely or by enhancing the normal gene product’s activity. Preferably, the disruption is a null disruption, wherein there is no significant expression of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 gene.

The term “native expression” refers to the expression of the full-length polypeptide encoded by the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 gene, at expression levels present in the wild-type mouse. Thus, a disruption in which there is “no native expression” of the endogenous
PRO19563, PRO19675, PRO20084, PRO21434, PROS0332, PRO38465 or PRO346 gene refers to a partial or complete reduction of the expression of at least a portion of a polypeptide encoded by an endogenous PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9864, PRO9865, PRO9904, PRO9902, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PROS0332, PRO38465 or PRO346 gene of a single cell, selected cells, or all of the cells of a mammal.

The term “knockout” refers to the disruption of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9865, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO20084, PRO21434, PROS0332, PRO38465 or PRO346 gene wherein the disruption results in: the functional inactivation of the native gene; the deletion of the native gene or a portion thereof; or a mutation in the native gene.


The term “construct” or “targeting construct” refers to an artificially assembled DNA segment to be transferred into a target tissue, cell line or animal. Typically, the targeting construct will include a gene or a nucleic acid sequence of particular interest, a marker gene and appropriate control sequences. As provided herein, the targeting construct comprises a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9865, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PROS0332, PRO38465 or PRO346 targeting construct. A “PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9865, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PROS0332, PRO38465 or PRO346 targeting construct” includes a DNA sequence homologous to at least one portion of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130,
PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 gene and is capable of producing a disruption in a PRO218, PRO228, PRO271, PRO273, PRO295, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO10013, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 gene in a host cell.

The term “transgenic cell” refers to a cell containing within its genome a PRO218, PRO228, PRO271, PRO273, PRO295, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO10013, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 gene that has been disrupted, modified, altered, or replaced completely or partially by the method of gene targeting.

The term “transgenic animal” refers to an animal that contains within its genome a specific gene that has been disrupted or otherwise modified or mutated by the methods described herein or methods otherwise well known in the art. Preferably the non-human transgenic animal is a mammal. More preferably, the mammal is a rodent such as a rat or mouse. In addition, a “transgenic animal” may be a heterozygous animal (i.e., one defective allele and one wild-type allele) or a homozygous animal (i.e., two defective alleles). An embryo is considered to fall within the definition of an animal. The provision of an animal includes the provision of an embryo or foetus in utero, whether by mating or otherwise, and whether or not the embryo goes to term.

As used herein, the terms “selective marker” and position selection marker” refer to a gene encoding a product that enables only the cells that carry the gene to survive and/or grow under certain conditions. For example, plant and animal cells that express the introduced neomycin resistance (Neo') gene are resistant to the compound G418. Cells that do not carry the Neo' gene marker are killed by G418. Other positive selection markers are known to, or are within the purview of, those of ordinary skill in the art.

The term “modulates” or “modulation” as used herein refers to the decrease, inhibition, reduction, amelioration, increase or enhancement of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO10013, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 gene function, expression, activity, or alternatively a phenotype associated with PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO10013, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 gene.
The term “ameliorates” or “amelioration” as used herein refers to a decrease, reduction or elimination of a condition, disease, disorder, or phenotype, including an abnormality or symptom.

The term “abnormality” refers to any disease, disorder, condition, or phenotype in which PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 is implicated, including pathological conditions and behavioral observations.
Table 1

/*
 * C-C increased from 12 to 15
 * Z is average of EQ
 * B is average of ND
 * match with stop is _M; stop-stop = 0; J (joker) match = 0
 */
#define _M 8 /* value of a match with a stop */

int _day[26][26] = {
    /* A */ 2, 0, 2, 0, 0, 4, 1, -1, 0, 1, 0, 2, 1, 1, 0, 0, 6, 0, 3, 0,
    /* B */ 0, 3, -4, 3, 2, -5, 0, 1, 2, 0, 0, -3, 2, 2, _M, 1, 1, 0, 0, 0, 0, -2, 5, 0, -3, -1,
    /* C */ -2, -4, 15, 5, 5, 4, -3, 3, -2, 0, 5, -5, 5, -4, 0, 2, 0, -2, 8, -8, 0, 5,
    /* D */ 0, 3, 5, 4, 3, -6, 1, 1, 2, 0, 0, 4, 3, -2, _M, -1, 2, 1, 0, 0, 0, 2, -7, 0, -4, -2,
    /* E */ 0, 2, 5, 3, 4, -5, 0, 1, 2, 0, 0, 3, -2, 1, _M, 1, 2, -1, 0, 0, 0, 2, 7, 0, 4, 3,
    /* F */ -4, 5, -4, 6, -5, 9, -5, 2, 1, 0, 5, 2, 0, -4, _M, -5, 5, -4, 3, 3, 0, 1, -1, 0, 7, 5,
    /* G */ 1, 0, -3, 1, 0, 5, 5, -2, -3, 0, 2, -4, 3, 0, _M, -1, 1, -3, 1, 0, 0, 1, 7, 5, 0, -5,
    /* H */ -1, -1, -3, 1, 1, 2, -2, 6, 2, 0, 0, -2, 2, _M, 0, 3, 2, -1, -1, 0, -2, 3, 0, 2, 0,
    /* I */ -1, -2, 2, 2, 2, 1, -3, 2, 5, 0, 2, 2, 2, 2, _M, 1, 2, -2, 1, 0, 0, 4, 5, 0, -1, -2,
    /* J */ 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, _M, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0,
    /* K */ -1, -1, 0, 5, 0, 0, 5, 2, -2, -2, 0, 5, 3, 0, 1, _M, -1, 3, 0, 0, 0, 0, -2, 3, 0, -4, 0,
    /* L */ -2, 3, -4, 4, 3, 2, -4, 3, -2, 0, 3, -6, 4, -3, _M, 3, 2, -3, 3, -1, 0, 2, -2, 0, 1, 2,
    /* M */ -1, -2, 5, -3, 3, 2, 0, 3, -2, 2, 0, 0, 4, 6, 2, _M, 2, -1, 0, 2, -1, 0, 2, 4, 0, -2, 1,
    /* N */ 0, 2, -4, 3, 1, 2, 4, -1, 0, 2, 2, 0, 1, -3, 2, _M, -1, 1, 0, 1, 0, 0, -2, 4, -3, 0, 2,
    /* P */ 1, -1, -3, 1, -1, 5, -1, 0, 2, 0, 1, -3, -2, -1, _M, 6, 0, 0, 1, 0, 0, -1, 6, 0, -5, 0,
    /* Q */ 0, 1, 5, 2, 2, -5, 1, -3, 2, 0, 1, -2, -1, _M, 0, 4, 1, 1, 1, 0, 2, -2, 5, 0, -4, 3,
    /* R */ -2, 0, 4, 1, -1, 4, -3, 2, 2, 0, 3, -3, 0, _M, 0, 1, 6, 0, -1, 0, 2, 2, 0, -4, 0,
    /* S */ 1, 0, 0, 0, 0, -3, 1, -1, -1, 0, 0, -3, -2, _M, 1, 1, 0, 2, 1, 0, -1, -2, 0, -3, 0,
    /* T */ 1, 0, -2, 0, 0, 3, 0, -1, 0, 0, 0, 1, 0, _M, 0, -1, -1, 1, 3, 0, 0, 5, 0, -3, 0,
    /* U */ 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, _M, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0,
    /* V */ 0, 2, -2, 2, -2, -1, -1, -2, 4, 0, -2, 2, 2, _M, 1, -2, -2, 1, 0, 0, 4, -6, -2, 12,
    /* W */ -6, 5, 8, -7, 7, 0, 7, -3, -5, 0, 3, -2, -4, 4, _M, -6, -5, 2, -2, 5, 0, 0, 17, 0, -12,
    /* X */ 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, _M, 0, 0, 0, 0, 0, 0, 0, 0, 0,
    /* Y */ -3, -3, 0, 4, -4, 7, -5, 0, 1, 0, 4, -1, 2, -2, _M, -5, -4, 3, -3, 0, -2, 0, 0, -10, 4,
    /* Z */ 0, 1, -5, 2, 3, -5, 0, 2, -2, 0, 0, -2, -1, _M, 0, 3, 0, 0, 0, 0, -2, -6, 0, -4, 4
};
/*
#include <stdio.h>
#include <ctype.h>

#define MAXJMP 16    /* max jumps in a diag */
#define MAXGAP 24    /* don't continue to penalize gaps larger than this */
#define JMPS 1024    /* max jmps in an path */
#define MX 4         /* save if there's at least MX-1 bases since last jmp */

#define DMAT 3       /* value of matching bases */
#define DMIS 0       /* penalty for mismatched bases */
#define DINS0 8      /* penalty for a gap */
#define DINS1 1      /* penalty per base */
#define PINS0 8      /* penalty for a gap */
#define PINS1 4      /* penalty per residue */

struct jmp {
    short     n[MAXJMP]; /* size of jmp (neg for dely) */
    unsigned short x[MAXJMP]; /* base no. of jmp in seq x */
}; /* limits seq to 2^16 -1 */

struct diag {
    int score;  /* score at last jmp */
    long offset; /* offset of prev block */
    short jmp;  /* current jmp index */
    struct jmp *jp; /* list of jmps */
};

struct path {
    int spec;  /* number of leading spaces */
    int n[JMP]; /* size of jmp (gap) */
    int x[JMP]; /* loc of jmp (last elem before gap) */
};

char *ofile; /* output file name */
char *namex[2]; /* seq names: getseq() */
char *prog; /* prog name for err msgs */
char *seqx[2]; /* seqs: getseq() */
int dmax; /* best diag: nw() */
int dmax0; /* final diag */
int dna; /* set if dna: main() */
int endgaps; /* set if penalizing end gaps */
int gapx, gapy; /* total gaps in seqs */
int len0, len1; /* seq lens */
int ngapx, ngapy; /* total size of gaps */
int smax; /* max score: nw() */
int *xbm; /* bitmap for matching */
long offset; /* current offset in jmp file */
struct diag *dx; /* holds diagnals */
struct path pp[2]; /* holds path for seqs */
char *calloc(), *malloc(), *index(), *strcpy();
char *getseq(), *g_calloc();
/* Needleman-Wunsch alignment program */

/* usage: progs file1 file2 
where file1 and file2 are two dna or two protein sequences. 
The sequences can be in upper- or lower-case as may contain ambiguity 
Any lines beginning with '!', '>', or '<' are ignored 
Max file length is 65535 (limited by unsigned short x in the jmp struct) 
A sequence with 1/3 or more of its elements ACGTU is assumed to be DNA 
Output is in the file "align.out" */

/* The program may create a tmp file in /tmp to hold info about traceback. 
Original version developed under BSD 4.3 on a vax 8650 */

#include "nw.h"
#include "day.h"

static _dbval[26] = {
  1,14,2,13,0,0,4,11,0,0,12,0,3,15,0,0,0,5,6,8,8,7,9,0,10,0
};

static _pbval[26] = {
  1, 2[1<<('D'-A'))][1<<('N'-A')], 4, 8, 16, 32, 64,
  128, 256, 0xFFFFFFF, 1<<10, 1<<11, 1<<12, 1<<13, 1<<14,
  1<<15, 1<<16, 1<<17, 1<<18, 1<<19, 1<<20, 1<<21, 1<<22,
  1<<23, 1<<24, 1<<25}(1<<('E'-A'))(1<<('Q'-A'))
};

main(ac, av)

int    ac;
char  *av[];

{ 
    prog = av[0];
    if (ac != 3) {
        fprintf(stderr, "usage: %s file1 file2
", prog);
        fprintf(stderr, "where file1 and file2 are two dna or two protein sequences
");
        fprintf(stderr, "The sequences can be in upper- or lower-case
");
        fprintf(stderr, "Any lines beginning with ", " or ", " are ignored
");
        fprintf(stderr, "Output is in the file "align.out"");
        exit(1);
    }
    namex[0] = av[1];
    namex[1] = av[2];
    seqx[0] = getseq(namex[0], &len0);
    seqx[1] = getseq(namex[1], &len1);
    xbm = (dna) _dbval : _pbval;

    endgaps = 0; /* 1 to penalize endgaps */
    ofile = "align.out"; /* output file */

    nw(); /* fill in the matrix, get the possible jmp */
    readjumps(); /* get the actual jmp */
    print(); /* print stats, alignment */
    cleanup(); /* unlink any tmp files */
}
Table 1 (cont')
/* do the alignment, return best score: main() */
/* dna values in Fitch and Smith, PNAS, 80, 1382-1386, 1983 */
/* pro: PAM 250 values */
/* When scores are equal, we prefer mismatches to any gap, prefer */
/* a new gap to extending an ongoing gap, and prefer a gap in seqx */
/* to a gap in seqy. */
*/
/* nwl() */

{ char *px, *py; /* seqs and ptrs */
  int *ndely, *dely; /* keep track of dely */
  int ndels, delx; /* keep track of delx */
  int *tmp; /* for swapping row0, row1 */
  int mis; /* score for each type */
  int ins0, ins1; /* insertion penalties */
  register id; /* diagonal index */
  register ii; /* jmp index */
  register *col0, *col1; /* score for curr, last row */
  register xx, yy; /* index into seqs */

  int dx = (struct diag *)g_malloc("to get diags", len0=len1+1, sizeof(struct diag));
  int ndely = (int *)g_malloc("to get ndely", len1+1, sizeof(int));
  int dely = (int *)g_malloc("to get dely", len1+1, sizeof(int));
  int col0 = (int *)g_malloc("to get col0", len1+1, sizeof(int));
  int col1 = (int *)g_malloc("to get col1", len1+1, sizeof(int));
  int ins0 = (int)DINS0 : PINSO;  
  int ins1 = (int)DINS1 : PINSL; 
  int smax = -10000;

  if (ends) { /* Waterman Bull Math Biol 84 */
    for (col0[0] = dely[0] = -ins0, yy = 1; yy <= len1; yy++) {
      col0[yy] = dely[yy] = col0[yy-1] - ins1;
      ndely[yy] = yy;
    }
    col0[0] = 0;
  } else for (yy = 1; yy <= len1; yy++) {
    dely[yy] = -ins0;

    /* fill in match matrix */
    /* initialize first entry in col */
    if (ends) { /*/ 
      if (xx == 1) 
        col1[0] = delx = (ins0+ins1);
      else 
        col1[0] = delx = col0[0] - ins1;
      ndelx = xx;
    }
  }

  else { /*/ 
    col1[0] = 0;
    delx = -ins0;
    ndelx = 0;
  }

  /* */
Table 1 (cont')

for (py = seqx[1], yy = 1; yy <= len1; py++, yy++) {
    mis = col0[yy-1];
    if (dna)
        mis += (xbn[*px-'A']&xbn[*py-'A'])? DMAT : DMIS;
    else
        mis += _day[*px-'A']||*py-'A'];
/* update penalty for del in x seq;
   favor new del over ongoing del
   ignore MAXGAP if weighting endpoints */
    if (endgaps || ndely[yy] < MAXGAP) {
        if (col0[yy] - ins0 >= dely[yy]) {
            dely[yy] = col0[yy] - (ins0+ins1);
            ndely[yy] = 1;
        } else {
            dely[yy] -= ins1;
            ndely[yy]++;
        }
    } else {
        if (col0[yy] - (ins0+ins1) >= dely[yy]) {
            dely[yy] = col0[yy] - (ins0+ins1);
            ndely[yy] = 1;
        }
    } else
        ndely[yy]++;
/* update penalty for del in y seq;
   favor new del over ongoing del */
    if (endgaps || ndelx < MAXGAP) {
        if (col1[yy-1] - ins0 >= delx) {
            delx = col1[yy-1] - (ins0+ins1);
            ndelx = 1;
        } else {
            delx -= ins1;
            ndelx++;
        }
    } else {
        if (col1[yy-1] - (ins0+ins1) >= delx) {
            delx = col1[yy-1] - (ins0+ins1);
            ndelx = 1;
        } else
            ndelx++;
/* pick the maximum score; we're favoring
   mis over any del and delx over dely */

id = xx - yy + len1 - 1;
if ((mis >= delx && mis >= dely[yy])
    col1[yy] = mis;
...nw

id = xx - yy + len1 - 1;
if ((mis >= delx && mis >= dely[yy])
    col1[yy] = mis;
...nw
Table 1 (cont')

    else if (delx >= dely[yy]) {
        col1[yy] = delx;
        i = dx[i][i].jmp;
        if (dx[i].p.n[0] && (d & MAXJMP
            && xx > dx[i].p.x[i][i]+MX) || mis > dx[i].score+DINS0) {
            dx[i].jmp += 1;
            if (i >= MAXJMP) {
                wx = jmps(i);
                i = dx[i].jmp = 0;
                dx[i].offset = offset;
                offset = sizeof(struct jmp) + sizeof(offset);
            }
        dx[i].p.n[i] = -delx;
        dx[i].p.x[i][i] = xx;
        dx[i].score = delx;
    } else {
        col1[yy] = dely[yy];
        i = dx[i].jmp;
        if (dx[i].p.n[0] && (d & MAXJMP
            && xx > dx[i].p.x[i][i]+MX) || mis > dx[i].score+DINS0) {
            dx[i].jmp += 1;
            if (i >= MAXJMP) {
                wx = jmps(i);
                i = dx[i].jmp = 0;
                dx[i].offset = offset;
                offset = sizeof(struct jmp) + sizeof(offset);
            }
        dx[i].p.n[i] = -dely[yy];
        dx[i].p.x[i][i] = xx;
        dx[i].score = -dely[yy];
    }

    } if (xx <= len0 && yy < len1) {
        /* last col */

        if (ends) {
            col1[yy] -= ins0+ins1*(len1-yy);
        }

        if (col1[yy] > smax) {
            smax = col1[yy];
            dmax = id;
        }

    }

    if (ends && xx < len0) {
        col1[yy-1] -= ins0+ins1*(len0-xx);
        if (col1[yy-1] > smax) {
            smax = col1[yy-1];
            dmax = id;
        }
        tmp = col0; col0 = col1; col1 = tmp;
    }

(vid) free((char *)ndely);
(vid) free((char *)dcly);
(vid) free((char *)co0);
(vid) free((char *)co1);
/*
 * print() -- only routine visible outside this module
 *
 * static:
 * getmat() -- trace back best path, count matches: print()
 * pr_align() -- print alignment of described in array p[]: print()
 * dumpblock() -- dump a block of lines with numbers, stars: pr_align()
 * nums() -- put out a number line: dumpblock()
 * putline() -- put out a line (name, [num], seq, [num]): dumpblock()
 * stars() -- put a line of stars: dumpblock()
 * stripname() -- strip any path and prefix from a seqname
 */

#include "nw.h"

#define SPC 3
#define P_LINE 256 /* maximum output line */
#define P_SPC 3 /* space between name or num and seq */

extern _day[26][26];
int olen; /* set output line length */
FILE *fpx; /* output file */

print()
{
    int lx, ly, firstgap, lastgap; /* overlap */

    if ((fpx = fopen(ofile, "w")) == 0) {
        fprintf(stderr, "can't write %s in", ofile);
        cleanup(1);
    }

    fprintf(fpx, "<First sequence: %s (length = %d) in
            name[0], len0);" name[1], len1);

    if (dmax < len1 - 1) { /* leading gap in x */
        pp[0].spc = firstgap = len1 - dmax - 1;
        ly = pp[0].spc;
    }
    else if (dmax > len1 - 1) { /* leading gap in y */
        pp[1].spc = firstgap = dmax - (len1 - 1);
        lx = pp[1].spc;
    }

    if (dmax0 < len0 - 1) { /* trailing gap in x */
        lastgap = len0 - dmax0 -1;
        lx = lastgap;
    }
    else if (dmax0 > len0 - 1) { /* trailing gap in y */
        lastgap = dmax0 - (len0 - 1);
        ly = lastgap;
    }

    getmat(lx, ly, firstgap, lastgap);
    pr_align();
}
/* trace back the best path, count matches */

static

getmat(lx, ly, firstgap, lastgap)

int lx, ly; /* "core" (minus endgaps) */
int firstgap, lastgap; /* leading trailing overlap */
{
  int nm, i0, i1, siz0, siz1;
  char outx[32];
  double pctl;
  register n0, n1;
  register char *p0, *p1;
  /* get total matches, score */

  i0 = i1 = siz0 = siz1 = 0;
  p0 = seqs[0] + pp[0].spc;
  n0 = pp[1].spc + 1;
  n1 = pp[0].spc + 1;
  nm = 0;
  while (*p0 & *p1) {
    if (sz0) {
      p1++;
      n1++;
      siz0--;
    } else if (sz1) {
      p0++;
      n0++;
      siz1--;
    } else {
      if (xbm[*p0-1] & xbm[*p1-1])
        nm++;
      if (n0++ == pp[0].n[i0++])
        siz0 = pp[0].n[i0++];
      if (n1++ == pp[1].n[i1++])
        siz1 = pp[1].n[i1++];
    }
  p0++;
  p1++;
}

/* pct homology: */
/* if penalizing endgaps, base is the shorter seq */
/* else, knock off overhangs and take shorter core */
if (endgaps)
  lx = (len0 < len1) ? len0 : len1;
else
  lx = (lx < ly) ? lx : ly;
  pct = 100.0*(double)nm/(double)lx;
  fprintf(fp, "%n",
  fprintf(fp, "<%.2f match%%s in an overlap of %.2f percent similarity\n",
      nm, (nm == 1) ? "es", lx, pct);
Table 1 (cont')

fprint(fx, "<gaps in first sequence: %d", gapx);
if (gapx) {
    (void) sprintf(outx, "(%d %.3f\%s)",
        ngapx, (dna) ? "base", "residue", (ngapx == 1) ? "": "s");
    fprint(fx, "%s", outx);
    fprint(fx, ", gaps in second sequence: %d", gapy);
    if (gapy) {
        (void) sprintf(outx, "(%d %.3f\%s)",
            ngapy, (dna) ? "base", "residue", (ngapy == 1) ? "": "s");
        fprint(fx, "%s", outx);
    }
}
if (dna)
    fprint(fx,
        "m<score: %d (match = %d, mismatch = %d, gap penalty = %d + %d per base)\n", smax, DMAT, DMIS, DINS0, DINS1);
else
    fprint(fx,
        "m<score: %d (Dayhoff PAM 250 matrix, gap penalty = %d + %d per residue)\n", smax, PINS0, PINS1);
if (cendgaps)
    fprint(fx,
        "<cendgaps penalized. left endgap: %d %.3f\%s, right endgap: %d %.3f\%s\n",
        firstgap, (dna) ? "base", "residue", (firstgap == 1) ? "": "s",
        lastgap, (dna) ? "base", "residue", (lastgap == 1) ? "": "s");
else
    fprint(fx, "<cendgaps not penalized\n";)

static int nn; /* matches in core -- for checking */
static int lmax; /* lengths of stripped file names */
static int ij[2]; /* jmp index for a path */
static int nc[2]; /* number at start of current line */
static int ni[2]; /* current elem number -- for gapping */
static int siz[2];
static char *ps[2]; /* ptr to current element */
static char *po[2]; /* ptr to next output char slot */
static char out[2][P_LINE]; /* output line */
static char star[P_LINE]; /* set by stars() */
/*
 * print alignment of described in struct path pp[]
 */

static pr_align()
{
    int nn; /* char count */
    int more;
    register i;

    for (i = 0, lmax = 0; i < 2; i++) {
        nn = strlen(names[i]);
        if (nn > lmax)
            lmax = nn;
        nc[i] = 1;
        ni[i] = 1;
        siz[i] = ij[i] = 0;
        ps[i] = sequ[i];
        po[i] = out[i];
    }
```c
for (nn = nn = 0, more = 1; more; ) {
    for (l = more = 0; l < 2; l++) {
        /*
         * do we have more of this sequence?
         */
        if (*ps[i])
            continue;
        more++;  
        if (pp[i].spc) /* leading space */
            *po[i]++ = ' ';  
                        pp[i].spc--;  
        else if (siz[i]) /* in a gap */
            *po[i]++ = ' ';  
                        siz[i]--;  
        else /* we're putting a seq element */
            *po[i] = *ps[i];
        if (islower(*ps[i]))
            *ps[i] = toupper(*ps[i]);
        po[i]++;  
        ps[i]++;  
        /*
         * are we at next gap for this seq?
         */
        if (ni[i] == pp[i].x[i[j]] ) {
            /*
            * we need to merge all gaps
            * at this location
            */
            siz[i] = pp[i].a[i[j][i]++ ];
            while (ni[i] == pp[i].x[i[j][i] ] )
                siz[i] += pp[i].a[i[j][i]++ ];
            ni[i]++;  
        }
    }  
    if (++nn == olen || !more && nn) {
        dumpblock();
        for (l = 0; l < 2; l++)
            po[i] = out[i];
        nn = 0;  
    }
}  
/*
 * dump a block of lines, including numbers, stars: pr_align()
 */
static
    dumpblock()
{
    register l;
    for (l = 0; l < 2; l++)
        *po[i]-- = '0';
}
```

(void) putc(\'u\', fx);
for (i = 0; i < 2; i++) {
    if (*out[i] && (*out[i] != \'\' || *(po[i]) != \'\')) {
        if (i == 0)
            nums(i);
        if (i == 0 && *out[1])
            stars(i);
        putline(i);
        if (i == 0 && *out[1])
            fprintf(fx, star);
        if (i == 1)
            nums(i);
    }
}

/*
 * put out a number line: dumpblock()
 */

static

int ix;    /* index in out[] holding seq */
nums(ix)
{
    int inl[P_LINE];
    register i, j;
    register char *pn, *px, *py;
    for (pn = inl, i = 0; i < inmax + P_SPC; i++, pn++)
        *pn = '\';
    for (i = ic[ix], py = out[ix]; *py; py++, pn++)
        if (*py == '\n' || *py == '\')
            *pn = '\';
        else if (p%0 == 0 || (i == 1 && ic[ix] != 1)) {
            j = (i < 0) ? -1 : 1;
            for (px = pn; j /= 10, px--)
                *px = j%10 + '0';
            if (i < 0)
                *px = '\';
        } else
        *pn = '\';
    I++;
}
*pn = '\0';
nc[ix] = I;
for (pn = inl; *pn; pn++)
    (void) putc(*pn, fx);
(void) putc('u\', fx);

/*
 * put out a line (name, [num], seq, [num]): dumpblock()
 */

static

int ix;
putline(ix)
{
for (px = name[index], i = 0; *px && *px != '; px++, I++)
(void) putc(*px, fx);
for (I < max + P_SPC; I++)
(void) putc(' ', fx);
/* these count from 1:
 * nil[] is current element (from 1)
 * nec[] is number at start of current line
 */
for (px = out[index]; *px; px++)
(void) putc(*px & 0x7F, fx);
(void) putc('n', fx);
}

/*
 * put a line of stars (seqs always in out[0], out[1]): dumpblock()
 */
static
stars()
{
int I;
register char *p0, *p1, cx, *px;

if (!(out[0] == '=' && *(p0) == '=') ||
    return;

px = star;
for (I = max + P_SPC; I--; I--)
    *px++ = ' ';

for (p0 = out[0], p1 = out[1]; *p0 && *p1; p0++, p1++)
{
if (!isalpha(*p0) && isalpha(*p1))
{
    if (xbm[0]'p0'-A'][xbm[0]'p1'-A'])
        cx = 'n';
    nm++;
}
else if (!isalpha & isday[*p0-'A'][*p1-'A'] > 0)
    cx = '.';
else
cx = 'n';

else
cx = 'n';

*p0 = *cx;
}

*p0 = 'n';
*p0 = 'n';

}
/ * strip path or prefix from pn, return len: pr_align() */
static
5

char *pn;  /* file name (may be path) */
{
    register char *px, *py;

    py = 0;
    for (px = pn; *px; px++)
        if (*px == '/')
            py = px + 1;

    if (py)
        (void) strcpy(pn, py);
    return(strlen(pn));
}
Table 1 (cont')

/*
 * cleanup() -- cleanup any tmp file
 * getseq() -- read in seq, set dna, len, maxlen
 * g_calloc() -- calloc() with error checking
 * readjumps() -- get the good jmps, from tmp file if necessary
 * writejumps() -- write a filled array of jmps to a tmp file: nw()
 */
#include "nw.h"
#include <sys/file.h>

char *jname = "/tmp/homgXXXXXX";         /* tmp file for jmps */
FILE *jf;
int cleanup();                         /* cleanup tmp file */
long lseek();

/*
 * remove any tmp file if we blow
 */
cleanup()
{
    int I;
    if (fj)
    {
        (void) unlink(jname);
        exit(1);
    }
    /*
     * read, return ptr to seq, set dna, len, maxlen
     * skip lines starting with ';','<', or '>
     * seq in upper or lower case
     */

    char *
    getseq(file, len)
    {
        char *file;        /* file name */
        int *len;         /* seq len */
        char line[1024], *pseq;
        register char *px, *py;
        int natgc, tlen;
        FILE *fp;
        if ((fp = fopen(file,"r")) == 0) {
            fprintf(stderr,"%s: can't read %s\n", prog, file);
            exit(1);
        }
        tlen = natgc = 0;
        while (fgets(line, 1024, fp)) {
            if (*line == ';' || *line == '<' || *line == '>')
                continue;
            for (px = line; *px != 'n'; px++)
                if (isupper(*px) || islower(*px))
                    tlen++;
        }
        if ((pseq = malloc((unsigned)(tlen+6))) == 0) {
            fprintf(stderr,"%s: malloc() failed to get %d bytes for %s\n", prog, tlen+6, file);
            exit(1);
        }
    }
Table 1 (cont')

```c
...getseq

py = pseq + 4;
*len = tlen;
rewind(fp);
while ((fgets(line, 1024, fp)) {
    if(*line == ':' || *line == '<' || *line == '>')
        continue;
    for (px = line; *px != 'n'; px++) {
        if(isupper(*px))
            *py++ = *px;
        else if(islower(*px))
            *py++ = toupper(*px);
        if(index("ATGCU", *(py-1)))
            nmatches++;
    }
    *py++ = '0';
    *py = '0';
    (void) fclose(fp);
    dna = nmatches > (tlen/3);
    return(pseq+4);
}
char *
g_calloc(msg, nx, sz)

25  char *msg; /* program, calling routine */
int nx, sz; /* number and size of elements */
{ char *px, *calloc();
  if ((px = calloc(unsigned)nx, (unsigned)sz)) == 0) {
    if(*msg) {
        fprintf(stderr, "%s: g_calloc() failed %s (%d, %d)\n", prog, msg, nx, sz);
        exit(1);
    }
  }
  return(px);
}
/*
* get final jmps from dx[] or tmp file, set pp[], reset dmax: main()
*/

40  readjumps()
{ int fd = -1;
  int siz, i0, i1;
  register l, j, xx;
  if(fd) {
      (void) fclose(fd);
      if ((fd = open(jname, O_RDWR, 0)) < 0) {
        fprintf(stderr, "%s: can't open() %s\n", prog, jname);
        cleanup(1);
      }
  }
  for (I = 0) = i1 = 0, dmax0 = dmax, xx = len0; ; I++) {
    while (I) {
      for (j = dx[dmax].jmp; j >= 0; dx[dmax].jp.x[j] >= xx; j--) ;
    }
```
Table 1 (cont')

...readjumps

if (j < 0 && dx[dnax].offset && f) {
    (void) fseek(fd, dx[dnax].offset, 0);
    (void) read(fd, (char *)&dx[dnax].jp, sizeof(struct jmp));
    (void) read(fd, (char *)&dx[dnax].offset, sizeof(dx[dnax].offset));
    dx[dnax].jmp = MAXJMP - 1;
}
else
    break;
}

if (f >= JMPS) {
    fprintf(stderr, "%s: too many gaps in alignment\n", prog);
    cleanup(1);
}

if (j >= 0) {
    siz = dx[dnax].jp.n[j];
    xx = dx[dnax].jp.x[j];
    dmax += siz;
    if (siz < 0) {
        /* gap in second seq */
        pp[1].a[i1] = -siz;
        xx -= siz;
        /* id = xx - yy + len1 - 1 */
        pp[1].x[i1] = xx - dmax + len1 - 1;
        gapy++;
        ngapy += siz;
    } /* ignore MAXGAP when doing endgaps */
    siz = (-siz < MAXGAP || endgaps) ? -siz : MAXGAP;
    i1++;
    } else if (siz > 0) /* gap in first seq */
        pp[0].a[i0] = siz;
    pp[0].x[i0] = xx;
    gapx++;
    ngapx += siz;
    /* ignore MAXGAP when doing endgaps */
    siz = (-siz < MAXGAP || endgaps) ? siz : MAXGAP;
    i0++;
}
else
    break;
/* reverse the order of jmps */
for (j = 0, i0--; j < i0; j++, i0--) {
    1 = pp[0].a[i0];
    pp[0].a[i0] = pp[0].a[i0];
    pp[0].a[i0] = 1;
    1 = pp[0].x[i0];
    pp[0].x[i0] = pp[0].x[i0];
    pp[0].x[i0] = 1;
}
for (j = 0, i1--; j < i1; j++, i1--) {
    1 = pp[1].a[i1];
    pp[1].a[i1] = pp[1].a[i1];
    pp[1].a[i1] = 1;
    1 = pp[1].x[i1];
    pp[1].x[i1] = pp[1].x[i1];
    pp[1].x[i1] = 1;
}
/* close file */
if (fd >= 0)
    (void) close(fd);
if (f)
    (void) unlink(jname);
fj = 0;
    offset = 0;
}
/*
 * write a filled jmp struct offset of the prev one (if any): nw()
 */

5  writejumps(ix)
   int   ix;
{
   char  *mktemp();

10  if (!fj) {
      if (mktemp(jname) < 0) {
        fprintf(stderr, "%s: can't mktemp() %s\n", prog, jname);
        cleanup(1);
      }

15  if ((fj = fopen(jname, "w")) == 0) {
      fprintf(stderr, "%s: can't write %s\n", prog, jname);
      exit(1);
    }

20  (void) fwrite((char *)dx[ix].jp, sizeof(struct jmp), 1, fj);
    (void) fwrite((char *)dx[ix].offset, sizeof(dx[ix].offset), 1, fj);
}
**Table 2**

<table>
<thead>
<tr>
<th>PRO</th>
<th>XXXXXXXXXXXXXXXX (Length = 15 amino acids)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Comparison Protein</td>
<td>XXXXXXXYYYYYY (Length = 12 amino acids)</td>
</tr>
</tbody>
</table>

5 \%

amino acid sequence identity =

(the number of identically matching amino acid residues between the two polypeptide sequences as determined by ALIGN-2) divided by (the total number of amino acid residues of the PRO polypeptide) =

5 divided by 15 = 33.3%

**Table 3**

<table>
<thead>
<tr>
<th>PRO</th>
<th>XXXXXXXXXXXX (Length = 10 amino acids)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Comparison Protein</td>
<td>XXXXXXXYYYYYZYYZ (Length = 15 amino acids)</td>
</tr>
</tbody>
</table>

15 \%

amino acid sequence identity =

(the number of identically matching amino acid residues between the two polypeptide sequences as determined by ALIGN-2) divided by (the total number of amino acid residues of the PRO polypeptide) =

5 divided by 10 = 50%

**Table 4**

<table>
<thead>
<tr>
<th>PRO-DNA</th>
<th>NNNNNNNNNNNNNNN (Length = 14 nucleotides)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Comparison DNA</td>
<td>NNNNNNNLLLLLLLL (Length = 16 nucleotides)</td>
</tr>
</tbody>
</table>

% nucleic acid sequence identity =

(the number of identically matching nucleotides between the two nucleic acid sequences as determined by ALIGN-2) divided by (the total number of nucleotides of the PRO-DNA nucleic acid sequence) =

6 divided by 14 = 42.9%
Table 5

<table>
<thead>
<tr>
<th>PRO-DNA</th>
<th>NNNNNNNNNNNN</th>
<th>(Length = 12 nucleotides)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Comparison DNA</td>
<td>NNNNLLLVV</td>
<td>(Length = 9 nucleotides)</td>
</tr>
</tbody>
</table>

% nucleic acid sequence identity =

(the number of identically matching nucleotides between the two nucleic acid sequences as determined by ALIGN-2) divided by (the total number of nucleotides of the PRO-DNA nucleic acid sequence) =

4 divided by 12 = 33.3%

II. Compositions and Methods of the Invention

A. Full-Length PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO972, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 Polypeptides

The present invention provides newly identified and isolated nucleotide sequences encoding polypeptides referred to in the present application as PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO972, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides. In particular, cDNAs encoding various PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO972, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides have been identified and isolated, as disclosed in further detail in the Examples below. It is noted that proteins produced in separate expression rounds may be given different PRO numbers but the UNQ number is unique for any given DNA and the encoded protein, and will not be changed. However, for sake of simplicity, in the present specification the protein encoded by the full length native nucleic acid molecules disclosed herein as well as all further native homologues and variants included in the foregoing definition of PRO, will be referred to as “PRO/number”, regardless of their origin or mode of preparation.

As disclosed in the Examples below, various cDNA clones have been deposited with the ATCC. The actual nucleotide sequences of those clones can readily be determined by the skilled artisan by sequencing of the
deposited clone using routine methods in the art. The predicted amino acid sequence can be determined from the nucleotide sequence using routine skill. For the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO656, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides and encoding nucleic acids described herein. Applicants have identified what is believed to be the reading frame best identifiable with the sequence information available at the time.

B. PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 Polypeptide Variants

In addition to the full-length native sequence PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 variants can be prepared. PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 variants can be prepared by introducing appropriate nucleotide changes into the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 DNA, and/or by synthesis of the desired PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865,

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PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide. Those skilled in the art will appreciate that amino acid changes may alter post-translational processes of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1271, PRO1865, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, such as changing the number or position of glycosylation sites or altering the membrane anchoring characteristics.

Variations in the native full-length sequence PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1271, PRO1865, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide or in various domains of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1271, PRO1865, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide described herein, can be made, for example, using any of the techniques and guidelines for conservative and non-conservative mutations set forth, for instance, in U.S. Patent No. 5,364,934. Variations may be a substitution, deletion or insertion of one or more codons encoding the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1271, PRO1865, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide that results in a change in the amino acid sequence of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1271, PRO1865, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide as compared with the native sequence PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1271, PRO1865, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide. Optionally the variation is by substitution of at least one amino
acid with any other amino acid in one or more of the domains of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO19904, PRO19907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide. Guidance in determining which amino acid residue may be inserted, substituted or deleted without adversely affecting the desired activity may be found by comparing the sequence of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO19904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide with that of homologous known protein molecules and minimizing the number of amino acid sequence changes made in regions of high homology. Amino acid substitutions can be the result of replacing one amino acid with another amino acid having similar structural and/or chemical properties, such as the replacement of a leucine with a serine, i.e., conservative amino acid replacements. Insertions or deletions may optionally be in the range of about 1 to 5 amino acids. The variation allowed may be determined by systematically making insertions, deletions or substitutions of amino acids in the sequence and testing the resulting variants for activity exhibited by the full-length or mature native sequence.

PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO19904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide fragments are provided herein. Such fragments may be truncated at the N-terminus or C-terminus, or may lack internal residues, for example, when compared with a full length native protein. Certain fragments lack amino acid residues that are not essential for a desired biological activity of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO19904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide.

PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO19904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 fragments may be prepared by any of a number of conventional techniques. Desired peptide fragments may be chemically synthesized. An alternative approach involves generating PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305,
PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 fragments by enzymatic digestion, e.g., by treating the protein with an enzyme known to cleave proteins at sites defined by particular amino acid residues, or by digesting the DNA with suitable restriction enzymes and isolating the desired fragment. Yet another suitable technique involves isolating and amplifying a DNA fragment encoding a desired polypeptide fragment, by polymerase chain reaction (PCR). Oligonucleotides that define the desired termini of the DNA fragment are employed at the 5’ and 3’ primers in the PCR. Preferably, PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide fragments share at least one biological and/or immunological activity with the native PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide disclosed herein.

Conservative substitutions of interest are shown in Table 6 under the heading of preferred substitutions. If such substitutions result in a change in biological activity, then more substantial changes, denominated exemplary substitutions in Table 6, or as further described below in reference to amino acid classes, are preferably introduced and the products screened.

<table>
<thead>
<tr>
<th>Original Residue</th>
<th>Exemplary Substitutions</th>
<th>Preferred Substitutions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ala (A)</td>
<td>Val, Leu, Ile</td>
<td>Val</td>
</tr>
<tr>
<td>Arg (E)</td>
<td>Lys, Gln, Asn</td>
<td>Lys</td>
</tr>
<tr>
<td>Asn (N)</td>
<td>Gln, His, Asp, Lys, Arg</td>
<td>Gln</td>
</tr>
<tr>
<td>Asp (D)</td>
<td>Glu, Asn</td>
<td>Glu</td>
</tr>
<tr>
<td>Cys (C)</td>
<td>Ser, Ala</td>
<td>Ser</td>
</tr>
<tr>
<td>Gln (Q)</td>
<td>Asn, Glu</td>
<td>Asn</td>
</tr>
<tr>
<td>Glu (E)</td>
<td>Asp, Gln</td>
<td>Asp</td>
</tr>
<tr>
<td>Gly (G)</td>
<td>Ala</td>
<td>Ala</td>
</tr>
<tr>
<td>His (H)</td>
<td>Asn, Gln, Lys, Arg</td>
<td>Arg</td>
</tr>
<tr>
<td>Ile (I)</td>
<td>Leu, Val, Met, Ala, Phe, Norleucine</td>
<td>Leu</td>
</tr>
<tr>
<td>Leu (L)</td>
<td>Norleucine; Ile; Val; Met; Ala; Phe</td>
<td>Ile</td>
</tr>
<tr>
<td>--------</td>
<td>----------------------------------</td>
<td>-----</td>
</tr>
<tr>
<td>Lys (K)</td>
<td>Arg; Gln; Asn</td>
<td>Arg</td>
</tr>
<tr>
<td>Met (M)</td>
<td>Leu; Phe; Ile</td>
<td>Leu</td>
</tr>
<tr>
<td>Phe (F)</td>
<td>Trp; Leu; Val; Ile; Ala; Tyr</td>
<td>Tyr</td>
</tr>
<tr>
<td>Pro (P)</td>
<td>Ala</td>
<td>Ala</td>
</tr>
<tr>
<td>Ser (S)</td>
<td>Thr</td>
<td>Thr</td>
</tr>
<tr>
<td>Thr (T)</td>
<td>Val; Ser</td>
<td>Ser</td>
</tr>
<tr>
<td>Trp (W)</td>
<td>Tyr; Phe</td>
<td>Tyr</td>
</tr>
<tr>
<td>Tyr (Y)</td>
<td>Trp; Phe; Thr; Ser</td>
<td>Phe</td>
</tr>
<tr>
<td>Val (V)</td>
<td>Ile; Leu; Met; Phe; Ala; Norleucine</td>
<td>Leu</td>
</tr>
</tbody>
</table>

Substantial modifications in function or immunological identity of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide are accomplished by selecting substitutions that differ significantly in their effect on maintaining (a) the structure of the polypeptide backbone in the area of the substitution, for example, as a sheet or helical conformation, (b) the charge or hydrophobicity of the molecule at the target site, or (c) the bulk of the side chain. Naturally occurring residues are divided into groups based on common side-chain properties:

Amino acids may be grouped according to similarities in the properties of their side chains (in A. L. Lehninger, in Biochemistry, second ed., pp. 73-75, Worth Publishers, New York (1975)):

1. non-polar: Ala (A), Val (V), Leu (L), Ile (I), Pro (P), Phe (F), Trp (W), Met (M)
2. uncharged polar: Gly (G), Ser (S), Thr (T), Cys (C), Tyr (Y), Asn (N), Gln (Q)

(3) acidic: Asp (D), Glu (E)
(4) basic: Lys (K), Arg (R), His(H)

Alternatively, naturally occurring residues may be divided into groups based on common side-chain properties:

1. hydrophobic: Norleucine, Met, Ala, Val, Leu, Ile;
2. neutral hydrophilic: Cys, Ser, Thr, Asn, Gln;
3. acidic: Asp, Glu;
4. basic: His, Lys, Arg;
5. residues that influence chain orientation: Gly, Pro;
6. aromatic: Trp, Tyr, Phe.

Non-conservative substitutions will entail exchanging a member of one of these classes for another class. Such substituted residues also may be introduced into the conservative substitution sites or, more preferably, into the remaining (non-conserved) sites.
The variations can be made using methods known in the art such as oligonucleotide-mediated (site-directed) mutagenesis, alanine scanning, and PCR mutagenesis. Site-directed mutagenesis [Carter et al., Nucl. Acids Res., 13:4331 (1986); Zoller et al., Nucl. Acids Res., 10:6487 (1987)], cassette mutagenesis [Wells et al., Gene, 34:315 (1985)], restriction selection mutagenesis [Wells et al., Philos. Trans. R. Soc. London Ser. A, 317:415 (1986)] or other known techniques can be performed on the cloned DNA to produce the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO1047, PRO5238, PRO5239, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 variant DNA.

Scanning amino acid analysis can also be employed to identify one or more amino acids along a contiguous sequence. Among the preferred scanning amino acids are relatively small, neutral amino acids. Such amino acids include alanine, glycine, serine, and cysteine. Alanine is typically a preferred scanning amino acid among this group because it eliminates the side-chain beyond the beta-carbon and is less likely to alter the main-chain conformation of the variant [Cunningham and Wells, Science, 244: 1081-1085 (1989)]. Alanine is also typically preferred because it is the most common amino acid. Further, it is frequently found in both buried and exposed positions [Creighton, The Proteins, (W.H. Freeman & Co., N.Y.); Chothia, J. Mol. Biol., 150:1 (1976)]. If alanine substitution does not yield adequate amounts of variant, an isoteric amino acid can be used.

C. Modifications of PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 Polypeptides

Covalent modifications of PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides are included within the scope of this invention. One type of covalent modification includes reacting targeted amino acid residues of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide with an organic derivatizing agent that is capable of reacting with selected side chains or the N- or C-terminal residues of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238,
PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9864, PRO9890, PRO9907, PRO10004, PRO10012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO10016, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10004, PRO10012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10004, PRO10013, PRO10016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10004, PRO10013, PRO9904, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides to a water-insoluble support matrix or surface for use in the method for purifying anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO655, anti-PRO655, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO9904, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibodies, and vice versa. Commonly used crosslinking agents include, e.g., 1,1-bis(diazoacetyl)-2-phenylethane, glutaraldehyde, N-hydroxy succinimide esters, for example, esters with 4-azidosalicylic acid, homobifunctional imidoesters, including disuccinimidyld esters such as 3,3'-dithiobis(succinimidyl)propionate), bifunctional maleimides such as bis-N-maleimido-1,8-octane and agents such as methyl-3-[[p-azidophenyl]dithio]propioimidate.

Other modifications include deamidation of glutaminyl and asparaginyl residues to the corresponding glutamyl and aspartyl residues, respectively, hydroxylation of proline and lysine, phosphorylation of hydroxyl groups of seryl or threonyl residues, methylation of the α-amino groups of lysine, arginine, and histidine side chains [T.E. Creighton, Proteins: Structure and Molecular Properties, W.H. Freeman & Co., San Francisco, pp. 79-86 (1983)], acetylation of the N-terminal amine, and amidation of any C-terminal carboxyl group.

Another type of covalent modification of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO9904, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide included within the scope of this invention comprises altering the native glycosylation pattern of the polypeptide. "Altering the native glycosylation pattern" is intended for purposes herein to mean deleting one or more carbohydrate moieties found in native sequence PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO9904, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides (either by removing the underlying
glycosylation site or by deleting the glycosylation by chemical and/or enzymatic means), and/or adding one or more glycosylation sites that are not present in the native sequence PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide. In addition, the phrase includes qualitative changes in the glycosylation of the native proteins, involving a change in the nature and proportions of the various carbohydrate moieties present.

Addition of glycosylation sites to the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide may be accomplished by altering the amino acid sequence. The alteration may be made, for example, by the addition of, or substitution by, one or more serine or threonine residues to the native sequence PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 (for O-linked glycosylation sites). The PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 amino acid sequence may optionally be altered through changes at the DNA level, particularly by mutating the DNA encoding the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide at preselected bases such that codons are generated that will translate into the desired amino acids.

Another means of increasing the number of carbohydrate moieties on the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide is by chemical or enzymatic coupling of

Removal of carbohydrate moieties present on the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO202084, PRO50332, PRO38465 or PRO346 polypeptide may be accomplished chemically or enzymatically or by mutational substitution of codons encoding for amino acid residues that serve as targets for glycosylation. Chemical deglycosylation techniques are known in the art and described, for instance, by Hakimuddin, et al., Arch. Biochem. Biophys., 259:52 (1987) and by Edge et al., Anal. Biochem., 118:131 (1981). Enzymatic cleavage of carbohydrate moieties on polypeptides can be achieved by the use of a variety of endo- and exo-glycosidases as described by Thotakura et al., Meth. Enzymol., 138:350 (1987).

Another type of covalent modification of PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide comprises linking the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide to one of a variety of nonproteinaceous polymers, e.g., polyethylene glycol (PEG), polypropylene glycol, or polyoxyalkylenes, in the manner set forth in U.S. Patent Nos. 4,640,835; 4,496,689; 4,301,444; 4,670,417; 4,791,192 or 4,179,337.

The PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides of the present invention may also be modified in a way to form a chimeric molecule comprising the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide fused to another, heterologous polypeptide or amino acid sequence.

Such a chimeric molecule comprises a fusion of the PRO218, PRO228, PRO271, PRO273, PRO295,
PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide with a tag polypeptide which provides an epitope to which an anti-tag antibody can selectively bind. The epitope tag is generally placed at the amino- or carboxyl- terminus of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide. The presence of such epitope-tagged forms of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide can be detected using an antibody against the tag polypeptide. Also, provision of the epitope tag enables the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide to be readily purified by affinity purification using an anti-tag antibody or another type of affinity matrix that binds to the epitope tag. Various tag polypeptides and their respective antibodies are well known in the art. Examples include poly-histidine (poly-his) or poly-histidine-glycine (poly-his-gly) tags; the flu HA tag polypeptide and its antibody 12CAS [Field et al., Mol. Cell. Biol., 8:2159-2165 (1988)]; the c-myec tag and the 8F9, 3C7, 6E10, G4, B7 and 9E10 antibodies thereto [Evan et al., Molecular and Cellular Biology, 5:3610-3616 (1985)]; and the Herpes Simplex virus glycoprotein D (gD) tag and its antibody [Paborsky et al., Protein Engineering, 3(6):547-553 (1990)]. Other tag polypeptides include the Flag-peptide [Hopp et al., BioTechnology, 6:1204-1210 (1988)]; the KT3 epitope peptide [Martin et al., Science, 255:192-194 (1992)]; an α-tubulin epitope peptide [Skinner et al., J. Biol. Chem., 266:15163-15166 (1991)]; and the T7 gene 10 protein peptide tag [Lutz-Freyerimuth et al., Proc. Natl. Acad. Sci. USA, 87:6393-6397 (1990)].

The chimeric molecule may comprise a fusion of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide with an immunoglobulin or a particular region of an immunoglobulin. For a bivalent form of the chimeric molecule (also referred to as an “immunoadhesin”), such
a fusion could be to the Fc region of an IgG molecule. The Ig fusions preferably include the substitution of a soluble (transmembrane domain deleted or inactivated) form of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO34329, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide in place of at least one variable region within an Ig molecule. In a particularly preferred aspect of the invention, the immunoglobulin fusion includes the hinge, CH2 and CH3, or the hinge, CH1, CH2 and CH3 regions of an IgG1 molecule. For the production of immunoglobulin fusions see also US Patent No. 5,428,130 issued June 27, 1995.

D. Preparation of PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 Polypeptides

The description below relates primarily to production of PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides by culturing cells transformed or transfected with a vector containing PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 nucleic acid. It is, of course, contemplated that alternative methods, which are well known in the art, may be employed to prepare PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides. For instance, the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 sequence, or portions thereof,
may be produced by direct peptide synthesis using solid-phase techniques [see, e.g., Stewart et al., *Solid-Phase Peptide Synthesis*, W.H. Freeman Co., San Francisco, CA (1969); Merrifield, *J. Am. Chem. Soc.*, 85:2149-2154 (1963)]. *In vitro* protein synthesis may be performed using manual techniques or by automation. Automated synthesis may be accomplished, for instance, using an Applied Biosystems Peptide Synthesizer (Foster City, CA) using manufacturer’s instructions. Various portions of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99048, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide may be chemically synthesized separately and combined using chemical or enzymatic methods to produce the full-length PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99048, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide.

1. **Isolation of DNA Encoding** PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99048, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 Polypeptides

DNA encoding PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99048, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides may be obtained from a cDNA library prepared from tissue believed to possess the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99048, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 mRNA and to express it at a detectable level. Accordingly, human PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99048, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084,
PRO21434-, PRO50332-, PRO38465- or PRO346-DNA can be conveniently obtained from a cDNA library prepared from human tissue, such as described in the Examples. The PRO218-, PRO228-, PRO271-, PRO273-, PRO295-, PRO302-, PRO305-, PRO326-, PRO386-, PRO655-, PRO162-, PRO788-, PRO792-, PRO940-, PRO941-, PRO1004-, PRO1012-, PRO1016-, PRO474-, PRO5238-, PRO1069-, PRO1111-, PRO1113-, PRO1130-, PRO1195-, PRO1271-, PRO1865-, PRO1879-, PRO3446-, PRO3543-, PRO4329-, PRO4352-, PRO5733-, PRO9859-, PRO9864-, PRO9904-, PRO9907-, PRO10013-, PRO90948-, PRO28694-, PRO16089-, PRO19563-, PRO19675-, PRO20084-, PRO21434-, PRO50332-, PRO38465- or PRO346-encoding gene may also be obtained from a genomic library or by known synthetic procedures (e.g., automated nucleic acid synthesis).

Libraries can be screened with probes (such as antibodies to the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide or oligonucleotides of at least about 20-80 bases) designed to identify the gene of interest or the protein encoded by it. Screening the cDNA or genomic library with the selected probe may be conducted using standard procedures, such as described in Sambrook et al., Molecular Cloning: A Laboratory Manual (New York: Cold Spring Harbor Laboratory Press, 1989). An alternative means to isolate the gene encoding PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 is to use PCR methodology [Sambrook et al., supra; Dieffenbach et al., PCR Primer: A Laboratory Manual (Cold Spring Harbor Laboratory Press, 1995)].

The Examples below describe techniques for screening a cDNA library. The oligonucleotide sequences selected as probes should be of sufficient length and sufficiently unambiguous that false positives are minimized. The oligonucleotide is preferably labeled such that it can be detected upon hybridization to DNA in the library being screened. Methods of labeling are well known in the art, and include the use of radiolabels like 32P-labeled ATP, biotinylation or enzyme labeling. Hybridization conditions, including moderate stringency and high stringency, are provided in Sambrook et al., supra.

Sequences identified in such library screening methods can be compared and aligned to other known sequences deposited and available in public databases such as GenBank or other private sequence databases. Sequence identity (at either the amino acid or nucleotide level) within defined regions of the molecule or across the full-length sequence can be determined using methods known in the art and as described herein.

Nucleic acid having protein coding sequence may be obtained by screening selected cDNA or genomic libraries using the deduced amino acid sequence disclosed herein for the first time, and, if necessary, using conventional primer extension procedures as described in Sambrook et al., supra, to detect precursors and processing intermediates of mRNA that may not have been reverse-transcribed into cDNA.
Selection and Transformation of Host Cells

Host cells are transfected or transformed with expression or cloning vectors described herein for PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide production and cultured in conventional nutrient media modified as appropriate for inducing promoters, selecting transformants, or amplifying the genes encoding the desired sequences. The culture conditions, such as media, temperature, pH and the like, can be selected by the skilled artisan without undue experimentation. In general, principles, protocols, and practical techniques for maximizing the productivity of cell cultures can be found in Mammalian Cell Biotechnology: a Practical Approach, M. Butler, ed. (IRL Press, 1991) and Sambrook et al., supra.

Methods of eukaryotic cell transfection and prokaryotic cell transformation are known to the ordinarily skilled artisan, for example, CaCl₂, CaPO₄, liposome-mediated and electroporation. Depending on the host cell used, transformation is performed using standard techniques appropriate to such cells. The calcium treatment employing calcium chloride, as described in Sambrook et al., supra, or electroporation is generally used for prokaryotes. Infection with Agrobacterium tumefaciens is used for transformation of certain plant cells, as described by Shaw et al., Gene, 23:315 (1983) and WO 89/05859 published 29 June 1989. For mammalian cells without such cell walls, the calcium phosphate precipitation method of Graham and van der Eb, Virology, 52:456-457 (1978) can be employed. General aspects of mammalian cell host system transfections have been described in U.S. Patent No. 4,399,216. Transformations into yeast are typically carried out according to the method of Van Solingen et al., J. Bact., 130:946 (1977) and Hsiao et al., Proc. Natl. Acad. Sci. (USA), 76:3829 (1979). However, other methods for introducing DNA into cells, such as by nuclear microinjection, electroporation, bacterial protoplast fusion with intact cells, or polyectations, e.g., polybren, polynithine, may also be used. For various techniques for transforming mammalian cells, see Kewton et al., Methods in Enzymology, 185:527-537 (1990) and Mansour et al., Nature, 336:348-352 (1988).

Suitable host cells for cloning or expressing the DNA in the vectors herein include prokaryote, yeast, or higher eukaryote cells. Suitable prokaryotes include but are not limited to eubacteria, such as Gram-negative or Gram-positive organisms, for example, Enterobacteriaceae such as E. coli. Various E. coli strains are publicly available, such as E. coli K12 strain MM294 (ATCC 31,446); E. coli X1776 (ATCC 31,537); E. coli strain W3110 (ATCC 27,325) and K5 772 (ATCC 53,635). Other suitable prokaryotic host cells include Enterobacteriaceae such as Escherichia, e.g., E. coli, Enterobacter, Erwinia, Klebsiella, Proteus, Salmonella, e.g., Salmonella typhimurium, Serratia, e.g., Serratia marcescans, and Shigella, as well as Bacilli such as B. subtilis and B. licheniformis (e.g., B. licheniformis 41P disclosed in DD 266,710 published 12 April 1989), Pseudomonas such as P. aeruginosa, and Streptomyces. These examples are illustrative rather than limiting. Strain W3110 is one particularly preferred host or parent host because it is a common host strain for recombinant DNA product fermentations. Preferably, the host cell secretes minimal amounts of proteolytic enzymes. For example, strain W3110 may be modified to effect a genetic mutation in the genes encoding proteins endogenous to the host, with examples of such hosts including E. coli W3110 strain 1A2, which has the complete genotype tonA; E. coli

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W3110 strain 9E4, which has the complete genotype \( \text{tonA} \) \( \text{ptr3} \); \( \text{E. coli} \) W3110 strain 27C7 (ATCC 55,244), which has the complete genotype \( \text{tonA} \) \( \text{ptr3} \) \( \text{phaA} \) E15 (argF-lac)\( ^{16} \) degP ompT kan'; \( \text{E. coli} \) W3110 strain 37D6, which has the complete genotype \( \text{tonA} \) \( \text{ptr3} \) \( \text{phaE} \) E15 (argF-lac)\( ^{16} \) degP ompT rbs7 ivyG kan'; \( \text{E. coli} \) W3110 strain 40B4, which is strain 37D6 with a non-kanamycin resistant degP deletion mutation; and an \( \text{E. coli} \) strain having mutant periplasmic protease disclosed in U.S. Patent No. 4,946,783 issued 7 August 1990. Alternatively, \textit{in vitro} methods of cloning, e.g., PCR or other nucleic acid polymerase reactions, are suitable.


Suitable host cells for the expression of glycosylated PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO656, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides are derived from multicellular organisms. Examples of invertebrate cells include insect cells such as \textit{Drosophila} S2 and \textit{Spodoptera} S19, as well as plant cells. Examples of useful mammalian host cell lines include Chinese hamster ovary (CHO) and COS cells. More specific examples include monkey kidney CV1 line transformed by SV40 (COS-7, ATCC CRL 1651); human embryonic kidney line (293 or 293 cells subcloned for growth in suspension culture, Graham et al., \textit{J. Gen Virol.}, 36:59 [1977]); Chinese
hamster ovary cells/-DHFR (CHO, Urlaub and Chasin, Proc. Natl. Acad. Sci. USA, 77:4216 (1980)); mouse sertoli cells (TM4, Mather, Biol. Reprod., 23:243-251 (1980)); human lung cells (W138, ATCC CCL 75); human liver cells (Hep G2, HB 8065); and mouse mammary tumor (MMT 060562, ATCC CCL51). The selection of the appropriate host cell is deemed to be within the skill in the art.

3. Selection and Use of a Replicable Vector

The nucleic acid (e.g., cDNA or genomic DNA) encoding PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO9908, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides may be inserted into a replicable vector for cloning (amplification of the DNA) or for expression. Various vectors are publicly available. The vector may, for example, be in the form of a plasmid, cosmid, viral particle, or phage. The appropriate nucleic acid sequence may be inserted into the vector by a variety of procedures. In general, DNA is inserted into an appropriate restriction endonuclease site(s) using techniques known in the art. Vector components generally include, but are not limited to, one or more of a signal sequence, an origin of replication, one or more marker genes, an enhancer element, a promoter, and a transcription termination sequence. Construction of suitable vectors containing one or more of these components employs standard ligation techniques which are known to the skilled artisan.

The PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide may be produced recombinantly not only directly, but also as a fusion polypeptide with a heterologous polypeptide, which may be a signal sequence or other polypeptide having a specific cleavage site at the N-terminus of the mature protein or polypeptide. In general, the signal sequence may be a component of the vector, or it may be a part of the PRO218-, PRO228-, PRO271-, PRO273-, PRO295-, PRO302-, PRO305-, PRO326-, PRO386-, PRO655-, PRO162-, PRO788-, PRO792-, PRO940-, PRO941-, PRO1004-, PRO1012-, PRO1016-, PRO474-, PRO5238-, PRO1069-, PRO1111-, PRO1130-, PRO1195-, PRO1271-, PRO1865-, PRO1879-, PRO3446-, PRO5733-, PRO9859-, PRO9864-, PRO9904-, PRO9907-, PRO10013-, PRO90948-, PRO28694-, PRO16089-, PRO19563-, PRO19675-, PRO20084-, PRO21434-, PRO50332-, PRO38465- or PRO346-encoding DNA that is inserted into the vector. The signal sequence may be a prokaryotic signal sequence selected, for example, from the group of the alkaline phosphatase, penicillinase, lpp, or heat-stable enterotoxin II leaders. For yeast secretion the signal sequence may be, e.g., the yeast invertase leader, alpha factor leader (including Saccharomyces and Kluyveromyces α-factor leaders, the latter described in U.S. Patent No. 5,010,182), or acid phosphatase leader, the C. albicans glucoamylase leader (EP 362,179 published 4 April 1990), or the signal described in WO 90/13646 published 15 November 1990. In mammalian cell expression, mammalian signal sequences may be used to direct secretion of the protein, such as signal sequences from secreted polypeptides of
the same or related species, as well as viral secretory leaders.

Both expression and cloning vectors contain a nucleic acid sequence that enables the vector to replicate in one or more selected host cells. Such sequences are well known for a variety of bacteria, yeast, and viruses. The origin of replication from the plasmid pBR322 is suitable for most Gram-negative bacteria, the 2μ plasmid origin is suitable for yeast, and various viral origins (SV40, polyoma, adenovirus, VSV or BPV) are useful for cloning vectors in mammalian cells.

Expression and cloning vectors will typically contain a selection gene, also termed a selectable marker. Typical selection genes encode proteins that (a) confer resistance to antibiotics or other toxins, e.g., ampicillin, neomycin, methotrexate, or tetracycline, (b) complement auxotrophic deficiencies, or (c) supply critical nutrients not available from complex media, e.g., the gene encoding D-alanine racemase for Bacilli.

An example of suitable selectable markers for mammalian cells are those that enable the identification of cells competent to take up the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346-encoding nucleic acid, such as DHFR or thymidine kinase. An appropriate host cell when wild-type DHFR is employed is the CHO cell line deficient in DHFR activity, prepared and propagated as described by Urlaub et al., Proc. Natl. Acad. Sci. USA, 77:4216 (1980). A suitable selection gene for use in yeast is the trp1 gene present in the yeast plasmid YRp7 [Stinchcomb et al., Nature, 282:39 (1979); Kingsman et al., Gene, 7:141 (1979); Tschemper et al., Gene, 10:157 (1980)]. The trp1 gene provides a selection marker for a mutant strain of yeast lacking the ability to grow in tryptophan, for example, ATCC No. 44076 or PEP4-1 [Jones, Genetics, 85:12 (1977)].

Expression and cloning vectors usually contain a promoter operably linked to the PRO218-, PRO228-, PRO271-, PRO273-, PRO295-, PRO302-, PRO305-, PRO326-, PRO386-, PRO655-, PRO162-, PRO788-, PRO792-, PRO940-, PRO941-, PRO1004-, PRO1012-, PRO1016-, PRO474-, PRO5238-, PRO1069-, PRO1111-, PRO1130-, PRO1195-, PRO1271-, PRO1865-, PRO1879-, PRO3446-, PRO3543-, PRO4329-, PRO4352-, PRO5733-, PRO9859-, PRO9864-, PRO9904-, PRO9907-, PRO10013-, PRO90948-, PRO28694-, PRO16089-, PRO19563-, PRO19675-, PRO20084-, PRO21434-, PRO50332-, PRO38465- or PRO346-encoding nucleic acid sequence to direct mRNA synthesis. Promoters recognized by a variety of potential host cells are well known. Promoters suitable for use with prokaryotic hosts include the β-lactamase and lactose promoter systems [Chang et al., Nature, 275:615 (1978); Goeddel et al., Nature, 281:544 (1979)], alkaline phosphatase, a tryptophan (trp) promoter system [Goeddel, Nucleic Acids Res., 8:4057 (1980); EP 36,776], and hybrid promoters such as the tac promoter [deBoer et al., Proc. Natl. Acad. Sci. USA, 80:21-25 (1983)]. Promoters for use in bacterial systems also will contain a Shine-Dalgarno (S.D.) sequence operably linked to the DNA encoding PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465- or PRO346-encoding nucleic acid sequence.
Examples of suitable promoting sequences for use with yeast hosts include the promoters for 3-
phosphoglycerate kinase [Hitzeman et al., J. Biol. Chem., 255:2073 (1980)] or other glycolytic enzymes [Hess et
al., J. Adv. Enzyme Reg., 7:149 (1968); Holland, Biochemistry, 17:4900 (1978)], such as enolase, glyceraldehyde-
3-phosphate dehydrogenase, hexokinase, pyruvate decarboxylase, phosphofructokinase, glucose-6-phosphate
isomerase, 3-phosphoglycerate mutase, pyruvate kinase, triosephosphate isomerase, phosphoglucone isomerase,
and glucokinase.

Other yeast promoters, which are inducible promoters having the additional advantage of transcription
controlled by growth conditions, are the promoter regions for alcohol dehydrogenase 2, isocitratechrome C, acid
phosphatase, degradative enzymes associated with nitrogen metabolism, metallothionein, glyceraldehyde-3-
phosphate dehydrogenase, and enzymes responsible for maltose and galactose utilization. Suitable vectors and
promoters for use in yeast expression are further described in EP 73,657.

PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655,
PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO7069,
PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329,
PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089,
PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 transcription from vectors
in mammalian host cells is controlled, for example, by promoters obtained from the genomes of viruses such as
polyoma virus, fowlpox virus (UK 2,211,504 published 5 July 1989), adenovirus (such as Adenovirus 2), bovine
papilloma virus, avian sarcoma virus, cytomegalovirus, a retrovirus, hepatitis-B virus and Simian Virus 40 (SV40),
from heterologous mammalian promoters, e.g., the actin promoter or an immunoglobulin promoter, and from heat-
shock promoters, provided such promoters are compatible with the host cell systems.

Transcription of a DNA encoding the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305,
PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016,
PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879,
PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013,
PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or
PRO346 polypeptide by higher eukaryotes may be increased by inserting an enhancer sequence into the vector.
Enhancers are cis-acting elements of DNA, usually about from 10 to 300 bp, that act on a promoter to increase its
transcription. Many enhancer sequences are now known from mammalian genes (globin, elastase, albumin, e-
fetoprotein, and insulin). Typically, however, one will use an enhancer from a eukaryotic cell virus. Examples
include the SV40 enhancer on the late side of the replication origin (bp 100-270), the cytomegalovirus early
promoter enhancer, the polyoma enhancer on the late side of the replication origin, and adenovirus enhancers. The
enhancer may be spliced into the vector at a position 5' or 3' to the PRO218, PRO228, PRO271, PRO273, PRO295,
PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004,
PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271,
PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904,
PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434,
PRO50332, PRO38465 or PRO346 coding sequence, but is preferably located at a site 5' from the promoter.

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Expression vectors used in eukaryotic host cells (yeast, fungi, insect, plant, animal, human, or nucleated cells from other multicellular organisms) will also contain sequences necessary for the termination of transcription and for stabilizing the mRNA. Such sequences are commonly available from the 5' and, occasionally 3', untranslated regions of eukaryotic or viral DNAs or cDNAs. These regions contain nucleotide segments transcribed as polyadenylated fragments in the untranslated portion of the mRNA encoding PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides.

Still other methods, vectors, and host cells suitable for adaptation to the synthesis of PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides in recombinant vertebrate cell culture are described in Gething et al., Nature, 293:620-625 (1981); Mantei et al., Nature, 281:40-46 (1979); EP 117,060; and EP 117,058.

4. **Detecting Gene Amplification/Expression**

Gene amplification and/or expression may be measured in a sample directly, for example, by conventional Southern blotting, Northern blotting to quantitate the transcription of mRNA [Thomas, Proc. Natl. Acad. Sci. USA, 77:5201-5205 (1980)], dot blotting (DNA analysis), or in situ hybridization, using an appropriately labeled probe, based on the sequences provided herein. Alternatively, antibodies may be employed that can recognize specific duplexes, including DNA duplexes, RNA duplexes, and DNA-RNA hybrid duplexes or DNA-protein duplexes. The antibodies in turn may be labeled and the assay may be carried out where the duplex is bound to a surface, so that upon the formation of duplex on the surface, the presence of antibody bound to the duplex can be detected.

Gene expression, alternatively, may be measured by immunological methods, such as immunohistochemical staining of cells or tissue sections and assay of cell culture or body fluids, to quantitate directly the expression of gene product. Antibodies useful for immunohistochemical staining and/or assay of sample fluids may be either monoclonal or polyclonal, and may be prepared in any mammal. Conveniently, the antibodies may be prepared against a native sequence PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide or against a synthetic peptide based on the DNA sequences provided herein or against exogenous sequence fused to PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016,
5. **Purification of Polypeptide**

Forms of PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99094, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 DNA and encoding a specific antibody epitope.

It may be desired to purify PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99094, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides from recombinant cell proteins or polypeptides. The following procedures are exemplary of suitable purification procedures: by fractionation on an ion-exchange column; ethanol precipitation; reverse phase HPLC; chromatography on silica or on a cation-exchange resin such as DEAE; chromatofocusing; SDS-PAGE; ammonium sulfate precipitation; gel filtration using, for example, Sephadex G-75; protein A Sepharose columns to remove contaminants such as IgG; and metal chelating columns to bind epitope-tagged forms of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99094, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide. Various methods of protein purification may be employed and such methods are known in the art and described for example in Deutscher, *Methods in Enzymology*, 182 (1990); Scopes, *Protein Purification: Principles and Practice*, Springer-Verlag, New York (1982). The purification step(s) selected will depend, for example, on the nature of the production process used and the particular PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305,
PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide produced.

5 E. Uses for PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 Polypeptides

Nucleotide sequences (or their complement) encoding PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides have various applications in the art of molecular biology, including uses as hybridization probes, in chromosome and gene mapping and in the generation of anti-sense RNA and DNA. PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 nucleic acid will also be useful for the preparation of PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides by the recombinant techniques described herein.

The full-length native sequence PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 gene, or portions thereof, may be used as hybridization probes for a cDNA library to isolate the full-length PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089,
PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 cDNA or to isolate still other cDNAs (for instance, those encoding naturally-occurring variants of PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides or PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides from other species) which have a desired sequence identity to the native PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 sequence disclosed herein.

Optionally, the length of the probes will be about 20 to about 50 bases. The hybridization probes may be derived from at least partially novel regions of the full length native nucleotide sequence wherein those regions may be determined without undue experimentation or from genomic sequences including promoters, enhancer elements and introns of native sequence PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346.

By way of example, a screening method will comprise isolating the coding region of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 gene using the known DNA sequence to synthesize a selected probe of about 40 bases. Hybridization probes may be labeled by a variety of labels, including radionucleotides such as $^{32}$P or $^{35}$S, or enzymatic labels such as alkaline phosphatase coupled to the probe via avidin/biotin coupling systems. Labeled probes having a sequence complementary to that of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 gene of the present invention.
can be used to screen libraries of human cDNA, genomic DNA or mRNA to determine which members of such libraries the probe hybridizes to. Hybridization techniques are described in further detail in the Examples below. Any EST sequences disclosed in the present application may similarly be employed as probes, using the methods disclosed herein.

Other useful fragments of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 nucleic acids include antisense or sense oligonucleotides comprising a single-stranded nucleic acid sequence (either RNA or DNA) capable of binding to target PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 mRNA (sense) or PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 DNA (antisense) sequences. Antisense or sense oligonucleotides, according to the present invention, comprise a fragment of the coding region of PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 DNA. Such a fragment generally comprises at least about 14 nucleotides, preferably from about 14 to 30 nucleotides. The ability to derive an antisense or a sense oligonucleotide, based upon a cDNA sequence encoding a given protein is described in, for example, Stein and Cohen (Cancer Res. 48:2659, 1988) and van der Krol et al. (BioTechniques 6:958, 1988).

Binding of antisense or sense oligonucleotides to target nucleic acid sequences results in the formation of duplexes that block transcription or translation of the target sequence by one of several means, including enhanced degradation of the duplexes, premature termination of transcription or translation, or by other means. The antisense oligonucleotides thus may be used to block expression of PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346. Antisense or sense oligonucleotides further comprise
oligonucleotides having modified sugar-phosphodiester backbones (or other sugar linkages, such as those described in WO 91/06629) and wherein such sugar linkages are resistant to endogenous nucleases. Such oligonucleotides with resistant sugar linkages are stable in vivo (i.e., capable of resisting enzymatic degradation) but retain sequence specificity to be able to bind to target nucleotide sequences.

Other examples of sense or antisense oligonucleotides include those oligonucleotides which are covalently linked to organic moieties, such as those described in WO 90/10048, and other moieties that increases affinity of the oligonucleotide for a target nucleic acid sequence, such as poly-L-lysine). Further still, intercalating agents, such as ellipticine, and alkylating agents or metal complexes may be attached to sense or antisense oligonucleotides to modify binding specificities of the antisense or sense oligonucleotide for the target nucleotide sequence.

Antisense or sense oligonucleotides may be introduced into a cell containing the target nucleic acid sequence by any gene transfer method, including, for example, CaPO₄-mediated DNA transfection, electroporation, or by using gene transfer vectors such as Epstein-Barr virus. In a preferred procedure, an antisense or sense oligonucleotide is inserted into a suitable retroviral vector. A cell containing the target nucleic acid sequence is contacted with the recombinant retroviral vector, either in vivo or ex vivo. Suitable retroviral vectors include, but are not limited to, those derived from the murine retrovirus M-MuLV, N2 (a retrovirus derived from M-MuLV), or the double copy vectors designated DCT5A, DCT5B and DCT5C (see WO 90/13641).

Sense or antisense oligonucleotides also may be introduced into a cell containing the target nucleotide sequence by formation of a conjugate with a ligand binding molecule, as described in WO 91/04753. Suitable ligand binding molecules include, but are not limited to, cell surface receptors, growth factors, other cytokines, or other ligands that bind to cell surface receptors. Preferably, conjugation of the ligand binding molecule does not substantially interfere with the ability of the ligand binding molecule to bind to its corresponding molecule or receptor, or block entry of the sense or antisense oligonucleotide or its conjugated version into the cell.

Alternatively, a sense or an antisense oligonucleotide may be introduced into a cell containing the target nucleic acid sequence by formation of an oligonucleotide-lipid complex, as described in WO 90/10448. The sense or antisense oligonucleotide-lipid complex is preferably dissociated within the cell by an endogenous lipase.

Antisense or sense RNA or DNA molecules are generally at least about 5 bases in length, about 10 bases in length, about 15 bases in length, about 20 bases in length, about 25 bases in length, about 30 bases in length, about 35 bases in length, about 40 bases in length, about 45 bases in length, about 50 bases in length, about 55 bases in length, about 60 bases in length, about 65 bases in length, about 70 bases in length, about 75 bases in length, about 80 bases in length, about 85 bases in length, about 90 bases in length, about 95 bases in length, about 100 bases in length, or more.

The probes may also be employed in PCR techniques to generate a pool of sequences for identification of closely related PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 coding sequences.
Nucleotide sequences encoding a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99048, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide can also be used to construct hybridization probes for mapping the gene which encodes that PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99048, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide and for the genetic analysis of individuals with genetic disorders. The nucleotide sequences provided herein may be mapped to a chromosome and specific regions of a chromosome using known techniques, such as in situ hybridization, linkage analysis against known chromosomal markers, and hybridization screening with libraries.

When the coding sequences for PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99048, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 encode a protein which binds to another protein (for example, where the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99048, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 is a receptor), the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99048, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide can be used in assays to identify the other proteins or molecules involved in the binding interaction. By such methods, inhibitors of the receptor/ligand binding interaction can be identified. Proteins involved in such binding interactions can also be used to screen for peptide or small molecule inhibitors or agonists of the binding interaction. Also, the receptor PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99048, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 can be used to isolate correlative ligand(s). Screening assays can be designed to find lead compounds that mimic the biological activity of a native
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PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide or a receptor for

PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides. Such screening

assays will include assays amenable to high-throughput screening of chemical libraries, making them particularly suitable for identifying small molecule drug candidates. Small molecules contemplated include synthetic organic or inorganic compounds. The assays can be performed in a variety of formats, including protein-protein binding assays, biochemical screening assays, immunoassays and cell based assays, which are well characterized in the art.

Nucleic acids which encode PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or

PRO346 polypeptides or its modified forms can also be used to generate either transgenic animals or "knock out" animals which, in turn, are useful in the development and screening of therapeutically useful reagents. A transgenic animal (e.g., a mouse or rat) is an animal having cells that contain a transgene, which transgene was introduced into the animal or an ancestor of the animal at a prenatal, e.g., an embryonic stage. A transgene is a DNA which is integrated into the genome of a cell from which a transgenic animal develops. The invention provides cDNA encoding a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or

PRO346 polypeptide which can be used to clone genomic DNA encoding a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide in accordance with established techniques and the genomic

sequences used to generate transgenic animals that contain cells which express DNA encoding PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113,
PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides. Any technique known in the art may be used to introduce a target gene transgene into animals to produce the founder lines of transgenic animals. Such techniques include, but are not limited to pronuclear microinjection (U.S. Pat. Nos. 4,873,191, 4,736,866 and 4,870,009); retrovirus mediated gene transfer into germ lines (Van der Putten, et al., Proc. Natl. Acad. Sci. USA, 82:6148-6152 (1985)); gene targeting in embryonic stem cells (Thompson, et al., Cell, 56:313-321 (1989)); nonspecific insertional inactivation using a gene trap vector (U.S. Pat. No. 6,436,707); electroporation of embryos (Lo, Mol. Cell. Biol., 3:1803-1814 (1983)); and sperm-mediated gene transfer (Lavitano, et al., Cell, 52:717-723 (1989)); etc. Typically, particular cells would be targeted for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 transgene incorporation with tissue-specific enhancers.

Transgenic animals that include a copy of a transgene encoding a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide introduced into the germ line of the animal at an embryonic stage can be used to examine the effect of increased expression of DNA encoding PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides. Such animals can be used as tester animals for reagents thought to confer protection from, for example, pathological conditions associated with its overexpression. In accordance with this facet of the invention, an animal is treated with the reagent and a reduced incidence of the pathological condition, compared to untreated animals bearing the transgene, would indicate a potential therapeutic intervention for the pathological condition. Alternatively, non-human homologues of PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO306, PRO326, PRO386, PRO655, PRO162, PRO788, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides can be used to construct a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO306, PRO326, PRO386, PRO655, PRO162, PRO788, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides.
PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 "knock out" animal which has a defective or altered gene encoding PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1113, PRO1119, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides and altered genomic DNA encoding PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides introduced into an embryonic stem cell of the animal. Preferably the knock out animal is a mammal. More preferably, the mammal is a rodent such as a rat or mouse. For example, cDNA encoding PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides can be used to clone genomic DNA encoding PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides in accordance with established techniques. A portion of the genomic DNA encoding the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide can be deleted or replaced with another gene, such as a gene encoding a selectable marker which can be used to monitor integration. Typically, several kilobases of unaltered flanking DNA (both at the 5' and 3' ends) are included in the vector [see e.g., Thomas and Capecchi, Cell, 51:503 (1987) for a description of homologous recombination vectors]. The vector is introduced
into an embryonic stem cell line (e.g., by electroporation) and cells in which the introduced DNA has homologously recombined with the endogenous DNA are selected [see e.g., Li et al., *Cell*, 69:915 (1992)]. The selected cells are then injected into a blastocyst of an animal (e.g., a mouse or rat) to form aggregation chimeras [see e.g., Bradley, in *Teratocarcinomas and Embryonic Stem Cells: A Practical Approach*, E. J. Robertson, ed. (IRL, Oxford, 1987), pp. 113-152]. A chimeric embryo can then be implanted into a suitable pseudopregnant female foster animal and the embryo brought to term to create a "knock out" animal. Progeny harboring the homologously recombined DNA in their germ cells can be identified by standard techniques and used to breed animals in which all cells of the animal contain the homologously recombined DNA. Knockout animals can be characterized for instance, for their ability to defend against certain pathological conditions and for their development of pathological conditions due to absence of the gene encoding the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO3545, PRO3552, PRO3733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide.

In addition, knockout mice can be highly informative in the discovery of gene function and pharmaceutical utility for a drug target, as well as in the determination of the potential on-target side effects associated with a given target. Gene function and physiology are so well conserved between mice and humans, since they are both mammals and contain similar numbers of genes, which are highly conserved between the species. It has recently been well documented, for example, that 98% of genes on mouse chromosome 16 have a human ortholog (Mural et al., *Science* 296:1661-71 (2002)).

Although gene targeting in embryonic stem (ES) cells has enabled the construction of mice with null mutations in many genes associated with human disease, not all genetic diseases are attributable to null mutations. One can design valuable mouse models of human diseases by establishing a method for gene replacement (knock-in) which will disrupt the mouse locus and introduce a human counterpart with mutation. Subsequently one can conduct *in vivo* drug studies targeting the human protein (Kitamoto et. Al., *Biochemical and Biophysical Res. Commun.* 222:742-47 (1996)).

Nucleic acid encoding the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides may also be used in gene therapy. In gene therapy applications, genes are introduced into cells in order to achieve *in vivo* synthesis of a therapeutically effective genetic product, for example for replacement of a defective gene. "Gene therapy" includes both conventional gene therapy where a lasting effect is achieved by a single treatment, and the administration of gene therapeutic agents, which involves the one time or repeated administration of a therapeutically effective DNA or mRNA. Antisense RNAs and DNAs can be used as therapeutic agents for blocking the expression of certain genes *in vivo*. It has already been shown that short antisense oligonucleotides can be imported into cells where they act as inhibitors, despite their low intracellular
concentrations caused by their restricted uptake by the cell membrane. (Zamecnik et al., Proc. Natl. Acad. Sci. USA 83:4143-4146 [1986]). The oligonucleotides can be modified to enhance their uptake, e.g. by substituting their negatively charged phosphodiester groups by uncharged groups.

There are a variety of techniques available for introducing nucleic acids into viable cells. The techniques vary depending upon whether the nucleic acid is transferred into cultured cells in vitro, or in vivo in the cells of the intended host. Techniques suitable for the transfer of nucleic acid into mammalian cells in vitro include the use of liposomes, electroporation, microinjection, cell fusion, DEAE-dextran, the calcium phosphate precipitation method, etc. The currently preferred in vivo gene transfer techniques include transfection with viral (typically retroviral) vectors and viral coat protein-liposome mediated transfection (Dzau et al., Trends in Biotechnology 11, 205-210 [1993]). In some situations it is desirable to provide the nucleic acid source with an agent that targets the target cells, such as an antibody specific for a cell surface membrane protein or the target cell, a ligand for a receptor on the target cell, etc. Where liposomes are employed, proteins which bind to a cell surface membrane protein associated with endocytosis may be used for targeting and/or to facilitate uptake, e.g. capsid proteins or fragments thereof tropic for a particular cell type, antibodies for proteins which undergo internalization in cycling, proteins that target intracellular localization and enhance intracellular half-life. The technique of receptor-mediated endocytosis is described, for example, by Wu et al., J. Biol. Chem. 262, 4429-4432 (1987); and Wagner et al., Proc. Natl. Acad. Sci. USA 87, 3410-3414 (1990). For review of gene marking and gene therapy protocols see Anderson et al., Science 256, 808-813 (1992).

The PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides described herein may also be employed as molecular weight markers for protein electrophoresis purposes and the isolated nucleic acid sequences may be used for recombinantly expressing those markers.

The nucleic acid molecules encoding the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides or fragments thereof described herein are useful for chromosome identification. In this regard, there exists an ongoing need to identify new chromosome markers, since relatively few chromosome marking reagents, based upon actual sequence data are presently available. Each PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 nucleic acid molecule of the present invention can be used as a chromosome marker.
The PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides and nucleic acid molecules of the present invention may also be used diagnostically for tissue typing, wherein the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides of the present invention may be differentially expressed in one tissue as compared to another, preferably in a diseased tissue as compared to a normal tissue of the same tissue type. PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides described herein may also be employed as therapeutic agents. The PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides of the present invention can be formulated according to known methods to prepare pharmaceutically useful compositions, whereby the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 product hereof is combined in admixture with a pharmaceutically acceptable carrier vehicle. Therapeutic formulations are prepared for storage by mixing the active ingredient having the desired degree of purity with optional physiologically acceptable carriers, excipients or stabilizers (Remington's Pharmaceutical Sciences, 16th edition, Osol, A. Ed. (1980)), in the form of lyophilized formulations or aqueous
solutions. Acceptable carriers, excipients or stabilizers are nontoxic to recipients at the dosages and concentrations employed, and include buffers such as phosphate, citrate and other organic acids; antioxidants including ascorbic acid; low molecular weight (less than about 10 residues) polypeptides; proteins, such as serum albumin, gelatin or immunoglobulins; hydrophilic polymers such as polyvinylpyrrolidone, amino acids such as glycine, glutamine, asparagine, arginine or lysine; monosaccharides, disaccharides and other carbohydrates including glucose, mannose, or dextrins; chelating agents such as EDTA; sugar alcohols such as mannitol or sorbitol; salt-forming counterions such as sodium; and/or nonionic surfactants such as TWEEN™, PLURONICS™ or PEG.

The formulations to be used for in vivo administration must be sterile. This is readily accomplished by filtration through sterile filtration membranes, prior to or following lyophilization and reconstitution.

Therapeutic compositions herein generally are placed into a container having a sterile access port, for example, an intravenous solution bag or vial having a stopper pierceable by a hypodermic injection needle.

The route of administration is in accord with known methods, e.g. injection or infusion by intravenous, intraperitoneal, intracerebral, intramuscular, intraocular, intraarterial or intracapsular routes, topical administration, or by sustained release systems.

Dosages and desired drug concentrations of pharmaceutical compositions of the present invention may vary depending on the particular use envisioned. The determination of the appropriate dosage or route of administration is well within the skill of an ordinary physician. Animal experiments provide reliable guidance for the determination of effective doses for human therapy. Interspecies scaling of effective doses can be performed following the principles laid down by Mordenti, J. and Chappell, W. "The use of interspecies scaling in toxicokinetics" In Toxicokinetics and New Drug Development, Yacobi et al., Eds., Pergamon Press, New York 1989, pp. 42-96.

When in vivo administration of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide or agonist or antagonist thereof is employed, normal dosage amounts may vary from about 10 μg/kg to up to 100 μg/kg of mammal body weight or more per day, preferably about 1 μg/kg/day to 10 μg/kg/day, depending upon the route of administration. Guidance as to particular dosages and methods of delivery is provided in the literature; see, for example, U.S. Pat. Nos. 4,657,760; 5,206,344; or 5,225,212. It is anticipated that different formulations will be effective for different treatment compounds and different disorders, that administration targeting one organ or tissue, for example, may necessitate delivery in a manner different from that to another organ or tissue.

Where sustained-release administration of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide is desired in a formulation with release characteristics suitable for the treatment
of any disease or disorder requiring administration of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1115, PRO1115, PRO1119, PRO1217, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, microencapsulation of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1115, PRO1119, PRO1217, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide is contemplated. Microencapsulation of recombinant proteins for sustained release has been successfully performed with human growth hormone (rhGH), interferon- (rIFN- ), interleukin-2, and MN rgp120. Johnson et al., Nat. Med., 2:795-799 (1996); Yasuda, Biomed. Ther., 27:1221-1223 (1993); Hora et al., Bio/Technology, 8:755-758 (1990); Cleland, "Design and Production of Single Immunization Vaccines Using Polylactide Polyglycolide Microsphere Systems," in Vaccine Design: The Subunit and Adjuvant Approach, Powell and Newman, eds, (Plenum Press: New York, 1995), pp. 439-462; WO 97/03692, WO 96/0072, WO 96/07339; and U.S. Pat. No. 5,654,010.

The sustained-release formulations of these proteins were developed using poly-lactic-coglycolic acid (PLGA) polymer due to its biocompatibility and wide range of biodegradable properties. The degradation products of PLGA, lactic and glycolic acids, can be cleared quickly within the human body. Moreover, the degradability of this polymer can be adjusted from months to years depending on its molecular weight and composition. Lewis, "Controlled release of bioactive agents from lactide/glycolide polymer," in: M. Chasin and R. Langer (Eds.), Biodegradable Polymers as Drug Delivery Systems (Marcel Dekker: New York, 1990), pp. 1-41.

This invention encompasses methods of screening compounds to identify those that mimic the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1115, PRO1119, PRO1217, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide (agonists) or prevent the effect of the PRO218, PRO228, PRO271, PRO273, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1115, PRO1119, PRO1217, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide (antagonists). Agonists that mimic a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1115, PRO1119, PRO1217, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or
PRO346 polypeptide would be especially valuable therapeutically in those instances where a negative phenotype is observed based on findings with the non-human transgenic animal whose genome comprises a disruption of the gene which encodes for the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide. Antagonists that prevent the effects of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1016, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide would be especially valuable therapeutically in those instances where a positive phenotype is observed based upon observations with the non-human transgenic knockout animal. Screening assays for antagonist drug candidates are designed to identify compounds that bind or complex with the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1016, PRO1012, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide encoded by the genes identified herein, or otherwise interfere with the interaction of the encoded polypeptide with other cellular proteins. Such screening assays will include assays amenable to high-throughput screening of chemical libraries, making them particularly suitable for identifying small molecule drug candidates.

The assays can be performed in a variety of formats, including protein-protein binding assays, biochemical screening assays, immunoassays, and cell-based assays, which are well characterized in the art.

All assays for antagonists are common in that they call for contacting the drug candidate with a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide encoded by a nucleic acid identified herein under conditions and for a time sufficient to allow these two components to interact.

In binding assays, the interaction is binding and the complex formed can be isolated or detected in the reaction mixture. The PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide encoded by the gene identified herein or the drug candidate is immobilized on a solid phase, e.g., on a microtiter
plate, by covalent or non-covalent attachments. Non-covalent attachment generally is accomplished by coating the solid surface with a solution of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO10948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide and drying. Alternatively, an immobilized antibody, e.g., a monoclonal antibody, specific for the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO9048, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide to be immobilized can be used to anchor it to a solid surface. The assay is performed by adding the non-immobilized component, which may be labeled by a detectable label, to the immobilized component, e.g., the coated surface containing the anchored component. When the reaction is complete, the non-reacted components are removed, e.g., by washing, and complexes anchored on the solid surface are detected. When the originally non-immobilized component carries a detectable label, the detection of label immobilized on the surface indicates that complexing occurred. Where the originally non-immobilized component does not carry a label, complexing can be detected, for example, by using a labeled antibody specifically binding the immobilized complex.

If the candidate compound interacts with but does not bind to a particular PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO9048, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide encoded by a gene identified herein, its interaction with that polypeptide can be assayed by methods well known for detecting protein-protein interactions. Such assays include traditional approaches, such as, e.g., cross-linking, co-immunoprecipitation, and co-purification through gradients or chromatographic columns. In addition, protein-protein interactions can be monitored by using a yeast-based genetic system described by Fields and co-workers (Fields and Song, Nature (London), 340:245-246 (1989); Chien et al., Proc. Natl. Acad. Sci. USA, 88:9578-9582 (1991)) as disclosed by Chevray and Nathans, Proc. Natl. Acad. Sci. USA, 89: 5789-5793 (1991). Many transcriptional activators, such as yeast GAL4, consist of two physically discrete modular domains, one acting as the DNA-binding domain, the other one functioning as the transcription-activation domain. The yeast expression system described in the foregoing publications (generally referred to as the "two-hybrid system") takes advantage of this property, and employs two hybrid proteins, one in which the target protein is fused to the DNA-binding domain of GAL4, and another, in which candidate activating proteins are fused to the activation domain. The expression of a GAL1-lacZ reporter gene under control of a GAL4-activated promoter depends on reconstitution of GAL4 activity via protein-protein interaction. Colonies containing interacting polypeptides are detected with a chromogenic substrate for β-galactosidase. A complete kit (MATCHMAKER™) for identifying protein-protein interactions between two
specific proteins using the two-hybrid technique is commercially available from Clontech. This system can also be extended to map protein domains involved in specific protein interactions as well as to pinpoint amino acid residues that are crucial for these interactions.

Compounds that interfere with the interaction of a gene encoding a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide identified herein and other intra- or extracellular components can be tested as follows: usually a reaction mixture is prepared containing the product of the gene and the intra- or extracellular component under conditions and for a time allowing for the interaction and binding of the two products. To test the ability of a candidate compound to inhibit binding, the reaction is run in the absence and in the presence of the test compound. In addition, a placebo may be added to a third reaction mixture, to serve as positive control. The binding (complex formation) between the test compound and the intra- or extracellular component present in the mixture is monitored as described hereinafore. The formation of a complex in the control reaction(s) but not in the reaction mixture containing the test compound indicates that the test compound interferes with the interaction of the test compound and its reaction partner.

To assay for antagonists, the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide may be added to a cell along with the compound to be screened for a particular activity and the ability of the compound to inhibit the activity of interest in the presence of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide indicates that the compound is an antagonist to the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or
PRO346 polypeptide and a potential antagonist with membrane-bound PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide receptors or recombinant receptors under appropriate conditions for a competitive inhibition assay. The PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide can be labeled, such as by radioactivity, such that the number of PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide molecules bound to the receptor can be used to determine the effectiveness of the potential antagonist. The gene encoding the receptor can be identified by numerous methods known to those of skill in the art, for example, ligand binding and FACS sorting. Coligan et al., Current Protocols in Immun., 1(2): Chapter 5 (1991). Preferably, expression cloning is employed wherein polyadenylated RNA is prepared from a cell responsive to the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide and a cDNA library created from this RNA is divided into pools and used to transfect COS cells or other cells that are not responsive to the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide. Transfected cells that are grown on glass slides are exposed to labeled PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide. The PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012,
PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO909948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide can be labeled by a variety of means including iodination or inclusion of a recognition site for a site-specific protein kinase. Following fixation and incubation, the slides are subjected to autoradiographic analysis. Positive pools are identified and sub-pools are prepared and re-transfected using an interactive sub-pooling and re-screening process, eventually yielding a single clone that encodes the putative receptor.

As an alternative approach for receptor identification, the labeled PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO99097, PRO10013, PRO909948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide can be photoaffinity-linked with cell membrane or extract preparations that express the receptor molecule. Cross-linked material is resolved by PAGE and exposed to X-ray film. The labeled complex containing the receptor can be excised, resolved into peptide fragments, and subjected to protein micro-sequencing. The amino acid sequence obtained from micro-sequencing would be used to design a set of degenerate oligonucleotide probes to screen a cDNA library to identify the gene encoding the putative receptor.

Another approach in assessing the effect of an antagonist to a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO99097, PRO10013, PRO909948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, would be administering a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO99097, PRO10013, PRO909948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 antagonist to a wild-type mouse in order to mimic a known knockout phenotype. Thus, one would initially knockout the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO99097, PRO10013, PRO909948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 gene of interest and observe the resultant phenotype as a consequence of knocking out or disrupting the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865,
Likewise, one could assess the effect of an agonist to a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide by administering an antagonist to the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide to a wild-type mouse. An effective antagonist would be expected to mimic the phenotypic effect that was initially observed in the knockout animal.

PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 gene. Subsequently, one could then assess the effectiveness of an antagonist to the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 gene.
PRO792, PRO940, PRO944, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide by administering an agonist to the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO944, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide to a non-human transgenic mouse. An effective agonist would be expected to ameliorate the negative phenotypic effect that was initially observed in the knockout animal.

In another assay for antagonists, mammalian cells or a membrane preparation expressing the receptor would be incubated with a labeled PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO944, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide in the presence of the candidate compound. The ability of the compound to enhance or block this interaction could then be measured.

More specific examples of potential antagonists include an oligonucleotide that binds to the fusions of immunoglobulin with the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO944, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, and, in particular, antibodies including, without limitation, poly- and monoclonal antibodies and antibody fragments, single-chain antibodies, anti-idiotypic antibodies, and chimeric or humanized versions of such antibodies or fragments, as well as human antibodies and antibody fragments. Alternatively, a potential antagonist may be a closely related protein, for example, a mutated form of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide that recognizes the receptor but imparts no effect, thereby competitively inhibiting the action of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or
PRO346 polypeptide.

Another potential PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO652, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9864, PRO9865, PRO9904, PRO9907, PRO9913, PRO9948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide antagonist is an antisense RNA or DNA construct prepared using antisense technology, where, e.g., an antisense RNA or DNA molecule acts to block directly the translation of mRNA by hybridizing to targeted mRNA and preventing protein translation. Antisense technology can be used to control gene expression through triple-helix formation or antisense DNA or RNA, both of which methods are based on binding of a polynucleotide to DNA or RNA. For example, the 3' coding portion of the polynucleotide sequence, which encodes the mature PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO652, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9864, PRO9904, PRO9907, PRO10013, PRO9948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides herein, is used to design an antisense RNA oligonucleotide of from about 10 to 40 base pairs in length. A DNA oligonucleotide is designed to be complementary to a region of the gene involved in transcription (triple helix - see Lee et al., Nucl. Acids Res., 6:3073 (1979); Cooney et al., Science, 241:456 (1988); Dervan et al., Science, 251:1360 (1991)), thereby preventing transcription and the production of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO652, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9864, PRO9904, PRO9907, PRO10013, PRO9948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide. The antisense RNA oligonucleotide hybridizes to the mRNA in vivo and blocks translation of the mRNA molecule into the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO652, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9864, PRO9904, PRO9907, PRO10013, PRO9948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide (antisense - Okano, Neurochem., 56:560 (1991); Oligodeoxynucleotides as Antisense Inhibitors of Gene Expression (CRC Press: Boca Raton, FL., 1988)). The oligonucleotides described above can also be delivered to cells such that the antisense RNA or DNA may be expressed in vivo to inhibit production of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO652, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9864, PRO9904, PRO9907, PRO10013, PRO9948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide. When antisense DNA is used, oligodeoxyribonucleotides derived from the translation-
initiation site, e.g., between about -10 and +10 positions of the target gene nucleotide sequence, are preferred.

Potential antagonists include small molecules that bind to the active site, the receptor binding site, or growth factor or other relevant binding site of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO10111, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, thereby blocking the normal biological activity of the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide. Examples of small molecules include, but are not limited to, small peptides or peptide-like molecules, preferably soluble peptides, and synthetic non-peptidyl organic or inorganic compounds.

Ribozymes are enzymatic RNA molecules capable of catalyzing the specific cleavage of RNA. Ribozymes act by sequence-specific hybridization to the complementary target RNA, followed by endonucleolytic cleavage. Specific ribozyme cleavage sites within a potential RNA target can be identified by known techniques. For further details see, e.g., Rossi, Current Biology, 4:469-471 (1994), and PCT publication No. WO 97/33551 (published September 18, 1997).

Nucleic acid molecules in triple-helix formation used to inhibit transcription should be single-stranded and composed of deoxynucleotides. The base composition of these oligonucleotides is designed such that it promotes triple-helix formation via Hoogsteen base-pairing rules, which generally require sizeable stretches of purines or pyrimidines on one strand of a duplex. For further details see, e.g., PCT publication No. WO 97/33551, supra.

These small molecules can be identified by any one or more of the screening assays discussed hereinabove and/or by any other screening techniques well known for those skilled in the art.

Diagnostic and therapeutic uses of the herein disclosed molecules may also be based upon the positive functional assay hits disclosed and described below.

PRO346 Antibodies

The present invention provides anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1150, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibodies which may find use herein as therapeutic and/or diagnostic agents. Exemplary antibodies include polyclonal, monoclonal, humanized, bispecific, and heteroconjugate antibodies.

1. Polyclonal Antibodies

Polyclonal antibodies are preferably raised in animals by multiple subcutaneous (sc) or intraperitoneal (ip) injections of the relevant antigen and an adjuvant. It may be useful to conjugate the relevant antigen (especially when synthetic peptides are used) to a protein that is immunogenic in the species to be immunized. For example, the antigen can be conjugated to keyhole limpet hemocyanin (KLH), serum albumin, bovine thyroglobulin, or soybean trypsin inhibitor, using a bifunctional or derivatizing agent, e.g., maleimidobenzoyl sulfosuccinimide ester (conjugation through cysteine residues), N-hydroxysuccinimide (through lysine residues), glutaraldehyde, succinic anhydride, SOCl₂, or RN=C=NR, where R and R¹ are different alkyl groups.

Animals are immunized against the antigen, immunogenic conjugates, or derivatives by combining, e.g., 100 μg or 5 μg of the protein or conjugate (for rabbits or mice, respectively) with 3 volumes of Freund's complete adjuvant and injecting the solution intradermally at multiple sites. One month later, the animals are boosted with 1/5 to 1/10 the original amount of peptide or conjugate in Freund's complete adjuvant by subcutaneous injection at multiple sites. Seven to 14 days later, the animals are bled and the serum is assayed for antibody titer. Animals are boosted until the titer plateaus. Conjugates also can be made in recombinant cell culture as protein fusions. Also, aggregating agents such as alum are suitably used to enhance the immune response.

2. Monoclonal Antibodies

Monoclonal antibodies may be made using the hybridoma method first described by Kohler et al., Nature, 256:495 (1975), or may be made by recombinant DNA methods (U.S. Patent No. 4,816,567).

In the hybridoma method, a mouse or other appropriate host animal, such as a hamster, is immunized as described above to elicit lymphocytes that produce or are capable of producing antibodies that will specifically bind to the protein used for immunization. Alternatively, lymphocytes may be immunized in vitro. After immunization, lymphocytes are isolated and then fused with a myeloma cell line using a suitable fusing agent, such as polyethylene glycol, to form a hybridoma cell (Goding, Monoclonal Antibodies: Principles and Practice, pp.59-103 (Academic Press, 1986)).

The hybridoma cells thus prepared are seeded and grown in a suitable culture medium which medium preferably contains one or more substances that inhibit the growth or survival of the unfused, parental myeloma cells (also referred to as fusion partner). For example, if the parental myeloma cells lack the enzyme hypoxanthine

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guanine phosphoribosyl transferase (HGPRT or HPRT), the selective culture medium for the hybridomas typically will include hypoxanthine, aminopterin, and thymidine (HAT medium), which substances prevent the growth of HGPRT-deficient cells.

Preferred fusion partner myeloma cells are those that fuse efficiently, support stable high-level production of antibody by the selected antibody-producing cells, and are sensitive to a selective medium that selects against the unfused parental cells. Preferred myeloma cell lines are murine myeloma lines, such as those derived from MOPC-21 and MPC-11 mouse tumors available from the Salk Institute Cell Distribution Center, San Diego, California USA, and SP-2 and derivatives e.g., X63-Ag8-653 cells available from the American Type Culture Collection, Manassas, Virginia, USA. Human myeloma and mouse-human heteromyeloma cell lines also have been described for the production of human monoclonal antibodies (Kozbor, J. Immunol., 133:3001 (1984); and Brodeur et al., Monoclonal Antibody Production Techniques and Applications, pp. 51-63 (Marcel Dekker, Inc., New York, 1987)).

Culture medium in which hybridoma cells are growing is assayed for production of monoclonal antibodies directed against the antigen. Preferably, the binding specificity of monoclonal antibodies produced by hybridoma cells is determined by immunoprecipitation or by an in vitro binding assay, such as radioimmunoassay (RIA) or enzyme-linked immunosorbent assay (ELISA).

The binding affinity of the monoclonal antibody can, for example, be determined by the Scatchard analysis described in Munson et al., Anal. Biochem., 107:220 (1980).

Once hybridoma cells that produce antibodies of the desired specificity, affinity, and/or activity are identified, the clones may be subcloned by limiting dilution procedures and grown by standard methods (Goding, Monoclonal Antibodies: Principles and Practice, pp.59-103 (Academic Press, 1986)). Suitable culture media for this purpose include, for example, D-MEM or RPMI-1640 medium. In addition, the hybridoma cells may be grown in vivo as ascites tumors in an animal e.g., by i.p. injection of the cells into mice.

The monoclonal antibodies secreted by the subclones are suitably separated from the culture medium, ascites fluid, or serum by conventional antibody purification procedures such as, for example, affinity chromatography (e.g., using protein A or protein G-Sepharose) or ion-exchange chromatography, hydroxylapatite chromatography, gel electrophoresis, dialysis, etc.

DNA encoding the monoclonal antibodies is readily isolated and sequenced using conventional procedures (e.g., by using oligonucleotide probes that are capable of binding specifically to genes encoding the heavy and light chains of murine antibodies). The hybridoma cells serve as a preferred source of such DNA. Once isolated, the DNA may be placed into expression vectors, which are then transfected into host cells such as E. coli cells, simian COS cells, Chinese Hamster Ovary (CHO) cells, or myeloma cells that do not otherwise produce antibody protein, to obtain the synthesis of monoclonal antibodies in the recombinant host cells. Review articles on recombinant expression in bacteria of DNA encoding the antibody include Skerra et al., Curr. Opinion in Immunol., 5:256-262 (1993) and Plückthun, Immunol. Revs, 130:151-188 (1992).

human antibodies by chain shuffling (Marks et al., *Bio/Technology*, 10:779-783 (1992)), as well as combinatorial infection and *in vivo* recombination as a strategy for constructing very large phage libraries (Waterhouse et al., *Nuc. Acids. Res.*, 21:2265-2266 (1993)). Thus, these techniques are viable alternatives to traditional monoclonal antibody hybridoma techniques for isolation of monoclonal antibodies.

The DNA that encodes the antibody may be modified to produce chimeric or fusion antibody polypeptides, for example, by substituting human heavy chain and light chain constant domain (C\text{H} and C\text{L}) sequences for the homologous murine sequences (U.S. Patent No. 4,816,567; and Morrison, et al., *Proc. Natl Acad. Sci. USA*, 81:6851 (1984)), or by fusing the immunoglobulin coding sequence with all or part of the coding sequence for a non-immunoglobulin polypeptide (heterologous polypeptide). The non-immunoglobulin polypeptide sequences can substitute for the constant domains of an antibody, or they are substituted for the variable domains of one antigen-combining site of an antibody to create a chimeric bivalent antibody comprising one antigen-combining site having specificity for an antigen and another antigen-combining site having specificity for a different antigen.

3. Human and Humanized Antibodies

The anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO665, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO50465 or anti-PRO346 antibodies of the invention may further comprise humanized antibodies or human antibodies. Humanized forms of non-human (e.g., murine) antibodies are chimeric immunoglobulins, immunoglobulin chains or fragments thereof (such as Fv, Fab, Fab', F(ab')\text{2}, or other antigen-binding subsequences of antibodies) which contain minimal sequence derived from non-human immunoglobulin. Humanized antibodies include human immunoglobulins (recipient antibody) in which residues from a complementary determining region (CDR) of the recipient are replaced by residues from a CDR of a non-human species (donor antibody) such as mouse, rat or rabbit having the desired specificity, affinity and capacity. In some instances, Fv framework residues of the human immunoglobulin are replaced by corresponding non-human residues. Humanized antibodies may also comprise residues which are found neither in the recipient antibody nor in the imported CDR or framework sequences. In general, the humanized antibody will comprise substantially all of at least one, and typically two, variable domains, in which all or substantially all of the CDR regions correspond to those of a non-human immunoglobulin and all or substantially all of the FR regions are those of a human immunoglobulin consensus sequence. The humanized antibody optimally also will comprise at least a portion of an immunoglobulin constant region (Fc), typically that of a human immunoglobulin [Jones et al., *Nature*, 321:522-525 (1986); Riechmann et al., *Nature*, 332:323-329 (1988); and Presta, *Curr. Op. Struct. Biol.*, 2:593-596 (1992)].

Methods for humanizing non-human antibodies are well known in the art. Generally, a humanized antibody has one or more amino acid residues introduced into it from a source which is non-human. These non-
human amino acid residues are often referred to as "import" residues, which are typically taken from an "import" variable domain. Humanization can be essentially performed following the method of Winter and co-workers [Jones et al., Nature, 321:522-525 (1986); Riechmann et al., Nature, 332:323-327 (1988); Verhoeyen et al., Science, 239:1534-1536 (1988)], by substituting rodent CDRs or CDR sequences for the corresponding sequences of a human antibody. Accordingly, such "humanized" antibodies are chimeric antibodies (U.S. Patent No. 4,816,567), wherein substantially less than an intact human variable domain has been substituted by the corresponding sequence from a non-human species. In practice, humanized antibodies are typically human antibodies in which some CDR residues and possibly some FR residues are substituted by residues from analogous sites in rodent antibodies.

The choice of human variable domains, both light and heavy, to be used in making the humanized antibodies is very important to reduce antigenicity and HAMA response (human anti-mouse antibody) when the antibody is intended for human therapeutic use. According to the so-called "best-fit" method, the sequence of the variable domain of a rodent antibody is screened against the entire library of known human variable domain sequences. The human V domain sequence which is closest to that of the rodent is identified and the human framework region (FR) within it accepted for the humanized antibody (Sims et al., J. Immunol., 151:2296 (1993); Chothia et al., J. Mol. Biol., 196:901 (1987)). Another method uses a particular framework region derived from the consensus sequence of all human antibodies of a particular subgroup of light or heavy chains. The same framework may be used for several different humanized antibodies (Carter et al., Proc. Natl. Acad. Sci. USA, 89:4285 (1992); Presta et al., J. Immunol., 151:2623 (1993)).

It is further important that antibodies be humanized with retention of high binding affinity for the antigen and other favorable biological properties. To achieve this goal, according to a preferred method, humanized antibodies are prepared by a process of analysis of the parental sequences and various conceptual humanized products using three-dimensional models of the parental and humanized sequences. Three-dimensional immunoglobulin models are commonly available and are familiar to those skilled in the art. Computer programs are available which illustrate and display probable three-dimensional conformational structures of selected candidate immunoglobulin sequences. Inspection of these displays permits analysis of the likely role of the residues in the functioning of the candidate immunoglobulin sequence, i.e., the analysis of residues that influence the ability of the candidate immunoglobulin to bind its antigen. In this way, FR residues can be selected and combined from the recipient and import sequences so that the desired antibody characteristic, such as increased affinity for the target antigen(s), is achieved. In general, the hypervariable region residues are directly and most substantially involved in influencing antigen binding.

Various forms of a humanized anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody are contemplated. For example, the humanized antibody may be an antibody fragment, such as a Fab,
which is optionally conjugated with one or more cytotoxic agent(s) in order to generate an immunoconjugate. Alternatively, the humanized antibody may be an intact antibody, such as an intact IgG1 antibody.

As an alternative to humanization, human antibodies can be generated. For example, it is now possible to produce transgenic animals (e.g., mice) that are capable, upon immunization, of producing a full repertoire of human antibodies in the absence of endogenous immunoglobulin production. For example, it has been described that the homozygous deletion of the antibody heavy-chain joining region (JH) gene in chimeric and germ-line mutant mice results in complete inhibition of endogenous antibody production. Transfer of the human germ-line immunoglobulin gene array into such germ-line mutant mice will result in the production of human antibodies upon antigen challenge. See, e.g., Jakobovits et al., Proc. Natl. Acad. Sci. USA, 90:2551 (1993); Jakobovits et al., Nature, 362:255-258 (1993); Bruggemann et al., Year in Immunol., 7:33 (1993); U.S. Patent Nos. 5,545,806, 5,569,825, 5,591,669 (all of GenPharm); 5,545,807; and WO 97/17852.

Alternatively, phage display technology (McCafferty et al., Nature 348:525-533 [1990]) can be used to produce human antibodies and antibody fragments in vitro, from immunoglobulin variable (V) domain gene repertoires from unimmunized donors. According to this technique, antibody V domain genes are cloned in-frame into either a major or minor coat protein gene of a filamentous bacteriophage, such as M13 or fd, and displayed as functional antibody fragments on the surface of the phage particle. Because the filamentous particle contains a single-stranded DNA copy of the phage genome, selections based on the functional properties of the antibody also result in selection of the gene encoding the antibody exhibiting those properties. Thus, the phage mimics some of the properties of the B-cell. Phage display can be performed in a variety of formats, reviewed in, e.g., Johnson, Kevin S. and Chiswell, David J., Current Opinion in Structural Biology, 3:564-571 (1993). Several sources of V-gene segments can be used for phage display. Clackson et al., Nature, 352:624-628 (1991) isolated a diverse array of anti-oxazolone antibodies from a small random combinatorial library of V genes derived from the spleens of immunized mice. A repertoire of V genes from unimmunized human donors can be constructed and antibodies to a diverse array of antigens (including self-antigens) can be isolated essentially following the techniques described by Marks et al., J. Mol. Biol., 222:581-597 (1991), or Griffith et al., EMBO J., 12:725-734 (1993). See, also, U.S. Patent Nos. 5,565,332 and 5,573,905.

As discussed above, human antibodies may also be generated by in vitro activated B cells (see U.S. Patents 5,567,610 and 5,229,275).

4. Antibody fragments

In certain circumstances there are advantages of using antibody fragments, rather than whole antibodies. The smaller size of the fragments allows for rapid clearance, and may lead to improved access to solid tumors.

Various techniques have been developed for the production of antibody fragments. Traditionally, these fragments were derived via proteolytic digestion of intact antibodies (see, e.g., Morimoto et al., Journal of Biochemical and Biophysical Methods 24:107-117 (1992); and Brennan et al., Science, 229:81 (1985)). However, these fragments can now be produced directly by recombinant host cells. Fab, Fv and ScFv antibody fragments can all be expressed in and secreted from E. coli, thus allowing the facile production of large amounts of these fragments. Antibody fragments can be isolated from the antibody phage libraries discussed above. Alternatively, Fab'-SH fragments can be directly recovered from E. coli and chemically coupled to form F(ab')2 fragments (Carter
et al., *Bio/Technology* 10:163-167 (1992)). According to another approach, F(ab')2, fragments can be isolated directly from recombinant host cell culture. Fab and F(ab')2, fragment with increased in vivo half-life comprising a salvage receptor binding epitope residues are described in U.S. Patent No. 5,869,046. Other techniques for the production of antibody fragments will be apparent to the skilled practitioner. The antibody of choice is a single chain Fv fragment (scFv). See WO 93/16185; U.S. Patent No. 5,571,894; and U.S. Patent No. 5,587,458. Fv and sFv are the only species with intact combining sites that are devoid of constant regions; thus, they are suitable for reduced nonspecific binding during in vivo use. sFv fusion proteins may be constructed to yield fusion of an effector protein at either the amino or the carboxy terminus of an sFv. See *Antibody Engineering*, ed. Borrebaeck, supra. The antibody fragment may also be a “linear antibody”, e.g., as described in U.S. Patent 5,641,870 for example. Such linear antibody fragments may be monospecific or bispecific.

5. **Bispecific Antibodies**

Bispecific antibodies are antibodies that have binding specificities for at least two different epitopes. Exemplary bispecific antibodies may bind to two different epitopes of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19675, PRO20084, PRO2864, PRO38465 or PRO346 protein as described herein. Other such antibodies may combine a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 binding site with a binding site for another protein. Alternatively, an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 arm may be combined with an arm which binds to a triggering molecule on a leukocyte such as a T-cell receptor molecule (e.g. CD3), or Fc receptors for IgG (FcγR), such as FcγRI (CD64), FcγRII (CD32) and FcγRIII (CD16), so as to focus and localize cellular defense mechanisms to the PRO218-, PRO228-, PRO271-, PRO273-, PRO295-, PRO302-, PRO305-, PRO326-, PRO386-, PRO655-, PRO162-, PRO788-, PRO792-, PRO940-, PRO941-, PRO1004-, PRO1012-, PRO1016-, PRO474-, PRO5238-, PRO1069-, PRO1111-, PRO1113-, PRO1130-, PRO1195-, PRO1271-, PRO1865-, PRO1879-, PRO3446-, PRO3543-, PRO4329-, PRO4352-, PRO5733-, PRO9859-, PRO9864-, PRO9904-, PRO9907-, PRO10013-, PRO90948-, PRO28694-, PRO16089-, PRO19675-, PRO20084-, PRO21434-, PRO50332-, PRO38465- or PRO346-expressing cell. Bispecific antibodies
may also be used to localize cytotoxic agents to cells which express a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9903, PRO9907, PRO9908, PRO28694, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide. These antibodies possess a PRO218-, PRO228-, PRO271-, PRO273-, PRO295-, PRO302-, PRO305-, PRO326-, PRO386-, PRO655-, PRO162-, PRO788-, PRO792-, PRO940-, PRO941-, PRO1004-, PRO1012-, PRO1016-, PRO474-, PRO238-, PRO1069-, PRO1111-, PRO1130-, PRO1195-, PRO1271-, PRO1865-, PRO1879-, PRO3446-, PRO3543-, PRO4329-, PRO4352-, PRO5733-, PRO9859-, PRO9864-, PRO9903-, PRO9907-, PRO9908-, PRO10013-, PRO90948-, PRO16089-, PRO19563-, PRO919675-, PRO20084-, PRO21434-, PRO50332-, PRO38465- or PRO346-binding arm and an arm which binds the cytotoxic agent (e.g., saporin, anti-interferon-α, vinca alkaloid, ricin A chain, methotrexate or radioactive isotope hapten). Bispecific antibodies can be prepared as full length antibodies or antibody fragments (e.g., F(ab')2 bispecific antibodies).


Methods for making bispecific antibodies are known in the art. Traditional production of full length bispecific antibodies is based on the co-expression of two immunoglobulin heavy chain-light chain pairs, where the two chains have different specificities (Millstein et al., Nature 305:537-539 (1983)). Because of the random assortment of immunoglobulin heavy and light chains, these hybridomas (quadromas) produce a potential mixture of 10 different antibody molecules, of which only one has the correct bispecific structure. Purification of the correct molecule, which is usually done by affinity chromatography steps, is rather cumbersome, and the product yields are low. Similar procedures are disclosed in WO 93/08829, and in Traunecker et al., EMBO J. 10:3655-3659 (1991).

According to a different approach, antibody variable domains with the desired binding specificity (antibody-antigen combining sites) are fused to immunoglobulin constant domain sequences. Preferably, the fusion is with an Ig heavy chain constant domain, comprising at least part of the hinge, C_{H}2, and C_{H}3 regions. It is preferred to have the first heavy-chain constant region (C_{H}1) containing the site necessary for light chain bonding, present in at least one of the fusions. DNAs encoding the immunoglobulin heavy chain fusions and, if desired, the immunoglobulin light chain, are inserted into separate expression vectors, and are co-transfected into a suitable host cell. This provides for greater flexibility in adjusting the mutual proportions of the three polypeptide fragments when unequal ratios of the three polypeptide chains used in the construction provide the optimum yield of the desired bispecific antibody. It is, however, possible to insert the coding sequences for two or all three polypeptide chains into a single expression vector when the expression of at least two polypeptide chains in equal ratios results in high yields or when the ratios have no significant affect on the yield of the desired chain combination.

The invention provides bispecific antibodies which are composed of a hybrid immunoglobulin heavy chain with a first binding specificity in one arm, and a hybrid immunoglobulin heavy chain-light chain pair
(providing a second binding specificity) in the other arm. It was found that this asymmetric structure facilitates the separation of the desired bispecific compound from unwanted immunoglobulin chain combinations, as the presence of an immunoglobulin light chain in only one half of the bispecific molecule provides for a facile way of separation. This approach is disclosed in WO 94/04690. For further details of generating bispecific antibodies see, for example, Suresh et al., Methods in Enzymology 121:210 (1986).

According to another approach described in U.S. Patent No. 5,731,168, the interface between a pair of antibody molecules can be engineered to maximize the percentage of heterodimers which are recovered from recombinant cell culture. The preferred interface comprises at least a part of the C_{10}3 domain. In this method, one or more small amino acid side chains from the interface of the first antibody molecule are replaced with larger side chains (e.g., tyrosine or tryptophan). Compensatory "cavities" of identical or similar size to the large side chain(s) are created on the interface of the second antibody molecule by replacing large amino acid side chains with smaller ones (e.g., alanine or threonine). This provides mechanism for increasing the yield of the heterodimer over other unwanted end-products such as homodimers.

Bispecific antibodies include cross-linked or "heteroconjugate" antibodies. For example, one of the antibodies in the heteroconjugate can be coupled to avidin, the other to biotin. Such antibodies have, for example, been proposed to target immune system cells to unwanted cells (U.S. Patent No. 4,676,980), and for treatment of HIV infection (WO 91/00360, WO 92/200373, and EP 03089). Heteroconjugate antibodies may be made using any convenient cross-linking methods. Suitable cross-linking agents are well known in the art, and are disclosed in U.S. Patent No. 4,676,980, along with a number of cross-linking techniques.

Techniques for generating bispecific antibodies from antibody fragments have also been described in the literature. For example, bispecific antibodies can be prepared using chemical linkage. Brennan et al., Science 229:81 (1985) describe a procedure wherein intact antibodies are proteolytically cleaved to generate F(ab')_{2} fragments. These fragments are reduced in the presence of the dithiol complexing agent, sodium arsenite, to stabilize vicinal dithiols and prevent intermolecular disulfide formation. The Fab' fragments generated are then converted to thionitrobenzoate (TNB) derivatives. One of the Fab'-TNB derivatives is then reconverted to the Fab'-thiol by reduction with mercaptoethanolamine and is mixed with an equimolar amount of the other Fab'-TNB derivative to form the bispecific antibody. The bispecific antibodies produced can be used as agents for the selective immobilization of enzymes.

Recent progress has facilitated the direct recovery of Fab'-SH fragments from E. coli, which can be chemically coupled to form bispecific antibodies. Shalaby et al., J. Exp. Med. 175; 217-225 (1992) describe the production of a fully humanized bispecific antibody F(ab')_{2} molecule. Each Fab' fragment was separately secreted from E. coli and subjected to directed chemical coupling in vitro to form the bispecific antibody. The bispecific antibody thus formed was able to bind to cells overexpressing the ErbB2 receptor and normal human T cells, as well as trigger the lytic activity of human cytotoxic lymphocytes against human breast tumor targets. Various techniques for making and isolating bispecific antibody fragments directly from recombinant cell culture have also been described. For example, bispecific antibodies have been produced using leucine zippers. Kostelnky et al., J. Immunol. 148(5):1547-1553 (1992). The leucine zipper peptides from the Fos and Jun proteins were linked to the Fab' portions of two different antibodies by gene fusion. The antibody homodimers were reduced at the hinge region to form monomers and then re-oxidized to form the antibody heterodimers. This method can also be
utilized for the production of antibody homodimers. The "diabody" technology described by Hollinger et al., Proc. Natl. Acad. Sci. USA 90:6444-6448 (1993) has provided an alternative mechanism for making bispecific antibody fragments. The fragments comprise a $V_{H}$ connected to a $V_{L}$ by a linker which is too short to allow pairing between the two domains on the same chain. Accordingly, the $V_{H}$ and $V_{L}$ domains of one fragment are forced to pair with the complementary $V_{H}$ and $V_{L}$ domains of another fragment, thereby forming two antigen-binding sites. Another strategy for making bispecific antibody fragments by the use of single-chain Fv (sFv) dimers has also been reported. See Gruber et al., J. Immunol., 152:5368 (1994).

Antibodies with more than two valencies are contemplated. For example, trispecific antibodies can be prepared. Tutt et al., J. Immunol., 147:60 (1991).

6. Heteroconjugate Antibodies

Heteroconjugate antibodies are also within the scope of the present invention. Heteroconjugate antibodies are composed of two covalently joined antibodies. Such antibodies have, for example, been proposed to target immune system cells to unwanted cells [U.S. Patent No. 4,676,980], and for treatment of HIV infection [WO 91/00360; WO 92/200373; EP 030897]. It is contemplated that the antibodies may be prepared in vitro using known methods in synthetic protein chemistry, including those involving crosslinking agents. For example, immunotoxins may be constructed using a disulfide exchange reaction or by forming a thioether bond. Examples of suitable reagents for this purpose include iminoothiolate and methyl-4-mercaptobutyrimidate and those disclosed, for example, in U.S. Patent No. 4,676,980.

7. Multivalent Antibodies

A multivalent antibody may be internalized (and/or catabolized) faster than a bivalent antibody by a cell expressing an antigen to which the antibodies bind. The antibodies of the present invention can be multivalent antibodies (which are other than of the IgM class) with three or more antigen binding sites (e.g. tetravalent antibodies), which can be readily produced by recombinant expression of nucleic acid encoding the polypeptide chains of the antibody. The multivalent antibody can comprise a dimerization domain and three or more antigen binding sites. The preferred dimerization domain comprises (or consists of) an Fc region or a hinge region. In this scenario, the antibody will comprise an Fc region and three or more antigen binding sites amino-terminal to the Fc region. The preferred multivalent antibody herein comprises (or consists of) three to about eight, but preferably four, antigen binding sites. The multivalent antibody comprises at least one polypeptide chain (and preferably two polypeptide chains), wherein the polypeptide chain(s) comprise two or more variable domains. For instance, the polypeptide chain(s) may comprise $V_{D}1-(X1)_{n}-V_{D}2-(X2)_{n}-F_{c}$, wherein $V_{D}1$ is a first variable domain, $V_{D}2$ is a second variable domain, $F_{c}$ is one polypeptide chain of an Fc region, $X1$ and $X2$ represent an amino acid or polypeptide, and $n$ is 0 or 1. For instance, the polypeptide chain(s) may comprise: VH-CH1-flexible linker-VH-CH1-Fc region chain, or VH-CH1-VH-CH1-Fc region chain. The multivalent antibody herein preferably further comprises at least two (and preferably four) light chain variable domain polypeptides. The multivalent antibody herein may, for instance, comprise from about two to about eight light chain variable domain polypeptides. The light chain variable domain polypeptides contemplated here comprise a light chain variable domain and, optionally, further comprise a CL domain.
8. **Effector Function Engineering**

It may be desirable to modify the antibody of the invention with respect to effector function, e.g., so as to enhance antigen-dependent cell-mediated cytotoxicity (ADCC) and/or complement dependent cytotoxicity (CDC) of the antibody. This may be achieved by introducing one or more amino acid substitutions in an Fc region of the antibody. Alternatively or additionally, cysteine residue(s) may be introduced in the Fc region, thereby allowing interchain disulfide bond formation in this region. The homodimeric antibody thus generated may have improved internalization capability and/or increased complement-mediated cell killing and antibody-dependent cellular cytotoxicity (ADCC). See Caron et al., *J Exp Med.*, 176:1191-1195 (1992) and Shopes, B. J. *Immunol.* 148:2918-2922 (1992). Homodimeric antibodies with enhanced anti-tumor activity may also be prepared using heterobifunctional cross-linkers as described in Wolff et al., *Cancer Research* 53:2560-2565 (1993). Alternatively, an antibody can be engineered which has dual Fc regions and may thereby have enhanced complement lysis and ADCC capabilities. See Stevenson et al., *Anti-Cancer Drug Design* 3:219-230 (1989). To increase the serum half-life of the antibody, one may incorporate a salvage receptor binding epitope into the antibody (especially an antibody fragment) as described in U.S. Patent 5,739,277, for example. As used herein, the term "salvage receptor binding epitope" refers to an epitope of the Fc region of an IgG molecule (e.g., IgG1, IgG2, IgG3, or IgG4) that is responsible for increasing the in vivo serum half-life of the IgG molecule.

9. **Immunocoujugates**

The invention also pertains to immunocoujugates comprising an antibody conjugated to a cytotoxic agent such as a chemotherapeutic agent, a growth inhibitory agent, a toxin (e.g., an enzymatically active toxin of bacterial, fungal, plant, or animal origin, or fragments thereof), or a radioactive isotope (i.e., a radioconjugate).

Chemotherapeutic agents useful in the generation of such immunocoujugates have been described above. Enzymatically active toxins and fragments thereof that can be used include diphtheria A chain, non-binding active fragments of diphtheria toxin, exotoxin A chain (from *Pseudomonas aeruginosa*), ricin A chain, abrin A chain, modeccin A chain, alpha-sarcin, *Aleurites fordii* proteins, dianthin proteins, *Phyllolaca americana* proteins (PAPI, PAPII, and PAP-S), momordica charantia inhibitor, curcin, crotin, saponaaria officinalis inhibitor, gelonin, mitogellin, restrictocin, phenomycin, enomycin, and the tricothecenes. A variety of radionuclides are available for the production of radioconjugated antibodies. Examples include $^{212}$Bi, $^{131}$I, $^{133}$In, $^{89}$Y, and $^{186}$Re. Conjugates of the antibody and cytotoxic agent are made using a variety of bifunctional protein-coupling agents such as N-succinimidyl-3-(2-pyridyldithio) propionate (SPDP), iminolithiane (IT), bifunctional derivatives of imidoesters (such as dimethyl adipimide HCL), active esters (such as disuccinimidyl suberate), aldehydes (such as glutaraldehyde), bis-azido compounds (such as bis (p-azidobenzoyl) hexanediamine), bis-diazonium derivatives (such as bis-(p-diazoniumbenzoyl)-ethylenediamine), diisocyanates (such as tolylene 2,6-diisocyanate), and bis-active fluorine compounds (such as 1,5-difluoro-2,4-dinitrobenzene). For example, a ricin immunotoxin can be prepared as described in Vitetta et al., *Science*, 238: 1098 (1987). Carbon-14-labeled 1-isothiocyanatobenzyl-3-methylidithyene triminepentanetric acid (MX-DTPA) is an exemplary chelating agent for conjugation of radionucleotide to the antibody. See WO94/11026.

Conjugates of an antibody and one or more small molecule toxins, such as a calicheamicin, maytansinoids, a trichothene, and CC1065, and the derivatives of these toxins that have toxin activity, are also
contemplated herein.

**Maytansine and maytansinoids**

The invention provides an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody (full length or fragments) which is conjugated to one or more maytansinoid molecules.

Maytansinoids are mitotic inhibitors which act by inhibiting tubulin polymerization. Maytansine was first isolated from the east African shrub *Maytenus serrata* (U.S. Patent No. 3,896,111). Subsequently, it was discovered that certain microbes also produce maytansinoids, such as maytansinol and C-3 maytansinol esters (U.S. Patent No. 4,151,042). Synthetic maytansinol and derivatives and analogues thereof are disclosed, for example, in U.S. Patent Nos. 4,137,230; 4,248,870; 4,256,746; 4,260,608; 4,265,814; 4,294,757; 4,307,016; 4,308,268; 4,308,269; 4,309,428; 4,313,946; 4,315,929; 4,317,821; 4,322,348; 4,331,598; 4,361,650; 4,364,866; 4,424,219; 4,450,254; 4,362,663; and 4,371,533, the disclosures of which are hereby expressly incorporated by reference.

**Maytansinoid-antibody conjugates**

In an attempt to improve their therapeutic index, maytansine and maytansinoids have been conjugated to antibodies specifically binding to tumor cell antigens. Immunoconjugates containing maytansinoids and their therapeutic use are disclosed, for example, in U.S. Patent Nos. 5,208,020, 5,416,064 and European Patent EP 0 425 235 B1, the disclosures of which are hereby expressly incorporated by reference. Liu et al., *Proc. Natl. Acad. Sci. USA* 93:8618-8623 (1996) described immunoconjugates comprising a maytansinoid designated DM1 linked to the monoclonal antibody C242 directed against human colorectal cancer. The conjugate was found to be highly cytotoxic towards cultured colon cancer cells, and showed antitumor activity in an *in vivo* tumor growth assay. Chari et al., *Cancer Research* 52:127-131 (1992) describe immunoconjugates in which a maytansinoid was conjugated via a disulfide linker to the murine antibody A7 binding to an antigen on human colon cancer cell lines, or to another murine monoclonal antibody TA.1 that binds the HER-2/neu oncogene. The cytotoxicity of the TA.1-maytansinoid conjugate was tested *in vitro* on the human breast cancer cell line SK-BR-3, which expresses 3 x 10^4 HER-2 surface antigens per cell. The drug conjugate achieved a degree of cytotoxicity similar to the free maytansinoid drug, which could be increased by increasing the number of maytansinoid molecules per antibody molecule. The A7-maytansinoid conjugate showed low systemic cytotoxicity in mice.

5 Anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody-maytansinoid conjugates are prepared by chemically linking an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody to a maytansinoid molecule without significantly diminishing the biological activity of either the antibody or the maytansinoid molecule. An average of 3-4 maytansinoid molecules conjugated per antibody molecule has shown efficacy in enhancing cytotoxicity of target cells without negatively affecting the function or solubility of the antibody, although even one molecule of toxin/antibody would be expected to enhance cytotoxicity over the use of naked antibody. Maytansinoids are well known in the art and can be synthesized by known techniques or isolated from natural sources. Suitable maytansinoids are disclosed, for example, in U.S. Patent No. 5,208,020 and in the other patents and nonpatent publications referred to hereinafore. Preferred maytansinoids are maytansinol and maytansinol analogues modified in the aromatic ring or at other positions of the maytansinol molecule, such as various maytansinol esters.

There are many linking groups known in the art for making antibody-maytansinoid conjugates, including, for example, those disclosed in U.S. Patent No. 5,208,020 or EP Patent 0 425 235 B1, and Chari et al., Cancer Research 52:127-131 (1992). The linking groups include disulfide groups, thioether groups, acid labile groups, photolabile groups, peptidase labile groups, or esterase labile groups, as disclosed in the above-identified patents, disulfide and thioether groups being preferred.

Conjugates of the antibody and maytansinoid may be made using a variety of bifunctional protein coupling agents such as N-succinimidyl-3-(2-pyridyl-dithio) propionate (SPDP), succinimidyl-4-(N-maleimidomethyl)cyclohexane-1-carboxylate, iminothiolane (IT), bifunctional derivatives of imidoesters (such as dimethyl adipimidate HCl), active esters (such as disuccinimidyl suberate), aldehydes (such as glutareddehyde), bis-azido compounds (such as bis (p-azidobenzoyl) hexane-diamine), bis-diazoimid derivatives (such as bis-(p-diazoniumbenzoyl)-ethylenediamine), diisocyanates (such as toluene 2,6-diisocyanate), and bis-active fluorine
compounds (such as 1,5-difluoro-2,4-dinitrobenzene). Particularly preferred coupling agents include N-succinimidyl-3-(2-pyridyldithio) propionate (SPDP) (Carlsson et al., Biochem. J. 173:723-737 [1978]) and N-succinimidyl-4-(2-pyridylthio)pentanoate (SPP) to provide for a disulfide linkage.

The linker may be attached to the maytansinoid molecule at various positions, depending on the type of the link. For example, an ester linkage may be formed by reaction with a hydroxyl group using conventional coupling techniques. The reaction may occur at the C-3 position having a hydroxyl group, the C-14 position modified with hydroxymethyl, the C-15 position modified with a hydroxyl group, and the C-20 position having a hydroxyl group. The linkage is formed at the C-3 position of maytansinol or a maytansinol analogue.

Calicheamicin

Another immunoconjugate of interest comprises an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO100013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody conjugated to one or more calicheamicin molecules. The calicheamicin family of antibiotics is capable of producing double-stranded DNA breaks at sub-picomolar concentrations. For the preparation of conjugates of the calicheamicin family, see U.S. patents 5,712,374, 5,714,586, 5,739,116, 5,767,285, 5,770,701, 5,770,710, 5,773,001, 5,877,296 (all to American Cyanamid Company). Structural analogues of calicheamicin which may be used include, but are not limited to, γ\textsubscript{1}, α\textsubscript{1}, α\textsubscript{1}, N-acetyl-γ\textsubscript{1}, PSAG and θ\textsubscript{1} (Hinman et al., Cancer Research 53:3336-3342 (1993), Lode et al., Cancer Research 58:2925-2928 (1998) and the aforementioned U.S. patents to American Cyanamid). Another anti-tumor drug that the antibody can be conjugated is QFA which is an antifolate. Both calicheamicin and QFA have intracellular sites of action and do not readily cross the plasma membrane. Therefore, cellular uptake of these agents through antibody mediated internalization greatly enhances their cytotoxic effects.

Other cytotoxic agents

Other antitumor agents that can be conjugated to the anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO100013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibodies of the invention include BCNU, streptozocin, vincristine and 5-fluorouracil, the family of agents known collectively LL-E33288 complex described in U.S. patents 5,053,394, 5,770,710, as well as esperamicin (U.S. patent 5,877,296).
Enzymatically active toxins and fragments thereof which can be used include diphtheria A chain, nonbinding active fragments of diphtheria toxin, exotoxin A chain (from *Pseudomonas aeruginosa*), ricin A chain, abrin A chain, modecin A chain, alpha-sarcin, *Aleurites fordii* proteins, dianthin proteins, *Phytolaca americana* proteins (PAPI, PAPII, and PAP-S), momordica charantia inhibitor, curcin, eritin, saponaria officinalis inhibitor, gelonin, mitogellin, restrictocin, phenomycin, enomycin and the tricothecenes. See, for example, WO 93/21232 published October 28, 1993.

The present invention further contemplates an immunoconjugate formed between an antibody and a compound with nucleolytic activity (e.g., a ribonuclease or a DNA endonuclease such as a deoxyribonuclease; DNase).

For selective destruction of the tumor, the antibody may comprise a highly radioactive atom. A variety of radioactive isotopes are available for the production of radioconjugated anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibodies. Examples include At$^{31}$, I$^{131}$, I$^{125}$, Y$^{90}$, Re$^{186}$, Re$^{188}$, Sm$^{153}$, Bi$^{212}$, P$^{32}$, Pb$^{212}$ and radioactive isotopes of Lu. When the conjugate is used for diagnosis, it may comprise a radioactive atom for scintigraphic studies, for example tc$^{99m}$ or I$^{123}$, or a spin label for nuclear magnetic resonance (NMR) imaging (also known as magnetic resonance imaging, mri), such as iodine-123 again, iodine-131, indium-111, fluorine-19, carbon-13, nitrogen-15, oxygen-17, gadolinium, manganese or iron.

The radio- or other labels may be incorporated in the conjugate in known ways. For example, the peptide may be biosynthesized or may be synthesized by chemical amino acid synthesis using suitable amino acid precursors involving, for example, fluorine-19 in place of hydrogen. Labels such as tc$^{99m}$ or I$^{123}$, Re$^{186}$, Re$^{188}$ and I$^{121}$ can be attached via a cysteine residue in the peptide. Yttrium-90 can be attached via a lysine residue. The IODOGEN method (Fraker et al 1978) Biochem. Biophys. Res. Commun. 80: 49-57 can be used to incorporate iodine-123. "Monoclonal Antibodies in Immunoscintigraphy" (Chatal, CRC Press 1989) describes other methods in detail.

Conjugates of the antibody and cytotoxic agent may be made using a variety of bifunctional protein coupling agents such as N-succinimidyl-3-(2-pyridyl)dithio) propionate (SPDP), succinimidyl-4-(N-maleimidomethyl) cyclohexane-1-carboxylate, iminothiolane (IT), bifunctional derivatives of imidoesters (such as dimethyl adipimidate HCL), active esters (such as disuccinimidyl suberate), aldehydes (such as glutareddehyde), bis-azido compounds (such as bis (p-azidobenzoyl) hexanedianime), bis-diazonium derivatives (such as bis-(p-diazoniumbenzoyl)-ethylenediamine), diisocyantes (such as tolyene 2,6-dioscyante), and bis-active fluorine compounds (such as 1,5-difluoro-2,4-dinitrobenzene). For example, the ricin immunotoxin can be prepared as described in Vitetta et al., *Science* 238: 1098 (1987). Carbon-14-labeled 1-isothiocyanaotbenzyl-3-methyleneylene triaminepentaacetic acid (MX-DTPA) is an exemplary chelating agent for conjugation of radionucleotide to the antibody. See WO94/11026. The linker may be a "cleavable linker" facilitating release of the cytotoxic drug in
the cell. For example, an acid-labile linker, peptidase-sensitive linker, photolabile linker, dimethyl linker or disulfide-containing linker (Chari et al., Cancer Research 52:127-131 (1992); U.S. Patent No. 5,208,020) may be used.

Alternatively, a fusion protein comprising the anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody and cytotoxic agent may be made, e.g., by recombinant techniques or peptide synthesis. The length of DNA may comprise respective regions encoding the two portions of the conjugate either adjacent one another or separated by a region encoding a linker peptide which does not destroy the desired properties of the conjugate.

The invention provides that the antibody may be conjugated to a "receptor" (such streptavidin) for utilization in tumor pre-targeting wherein the antibody-receptor conjugate is administered to the patient, followed by removal of unbound conjugate from the circulation using a clearing agent and then administration of a "ligand" (e.g., avidin) which is conjugated to a cytotoxic agent (e.g., a radionucleotide).

10. **Immunoliposomes**

The anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibodies disclosed herein may also be formulated as immunoliposomes. A "liposome" is a small vesicle composed of various types of lipids, phospholipids and/or surfactant which is useful for delivery of a drug to a mammal. The components of the liposome are commonly arranged in a bilayer formation, similar to the lipid arrangement of biological membranes.

Liposomes containing the antibody are prepared by methods known in the art, such as described in Epstein et al., Proc. Natl. Acad. Sci. USA 82:3688 (1985); Hwang et al., Proc. Natl. Acad. Sci. USA 77:4030 (1980); U.S. Pat. Nos. 4,485,045 and 4,544,545; and WO97/38731 published October 23, 1997. Liposomes with enhanced circulation time are disclosed in U.S. Patent No. 5,013,556.

Particularly useful liposomes can be generated by the reverse phase evaporation method with a lipid composition comprising phosphatidylcholine, cholesterol and PEG-derivatized phosphatidylethanolamine (PEG-PE). Liposomes are extruded through filters of defined pore size to yield liposomes with the desired diameter. Fab' fragments of the antibody of the present invention can be conjugated to the liposomes as described in Martin et al., J. Biol. Chem. 257:286-288 (1982) via a disulfide interchange reaction. A chemotherapeutic agent is

11. **Pharmaceutical Compositions of Antibodies**

Antibodies specifically binding a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99048, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide identified herein, as well as other molecules identified by the screening assays disclosed hereinbefore, can be administered for the treatment of various disorders in the form of pharmaceutical compositions.

If the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO99048, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide is intracellular and whole antibodies are used as inhibitors, internalizing antibodies are preferred. However, lipofections or liposomes can also be used to deliver the antibody, or an antibody fragment, into cells. Where antibody fragments are used, the smallest inhibitory fragment that specifically binds to the binding domain of the target protein is preferred. For example, based upon the variable-region sequences of an antibody, peptide molecules can be designed that retain the ability to bind the target protein sequence. Such peptides can be synthesized chemically and/or produced by recombinant DNA technology. See, e.g., Marasco et al., Proc. Natl. Acad. Sci. USA, 90: 7889-7893 (1993). The formulation herein may also contain more than one active compound as necessary for the particular indication being treated, preferably those with complementary activities that do not adversely affect each other. Alternatively, or in addition, the composition may comprise an agent that enhances its function, such as, for example, a cytotoxic agent, cytokine, chemotherapeutic agent, or growth-inhibitory agent. Such molecules are suitably present in combination in amounts that are effective for the purpose intended.

The active ingredients may also be entrapped in microcapsules prepared, for example, by coacervation techniques or by interfacial polymerization, for example, hydroxyethylcellulose or gelatin-microcapsules and poly-(methylmethacrylate) microcapsules, respectively, in colloidal drug delivery systems (for example, liposomes, albumin microspheres, microemulsions, nano-particles, and nanocapsules) or in macroemulsions. Such techniques are disclosed in Remington's *Pharmaceutical Sciences*, *supra*.

The formulations to be used for *in vivo* administration must be sterile. This is readily accomplished by filtration through sterile filtration membranes.

Sustained-release preparations may be prepared. Suitable examples of sustained-release preparations include semipermeable matrices of solid hydrophobic polymers containing the antibody, which matrices are in the form of shaped articles, e.g., films, or microcapsules. Examples of sustained-release matrices include polyesters, hydrogels (for example, poly(2-hydroxyethyl-methacrylate), or poly(vinylalcohol)), polylactides (U.S. Pat. No. 3,773,919), copolymers of L-glutamic acid and γ ethyl-L-glutamate, non-degradable ethylene-vinyl acetate,
degradable lactic acid-glycolic acid copolymers such as the LUPRON DEPOT™ (injectable microspheres composed of lactic acid-glycolic acid copolymer and leuprolide acetate), and poly-D-(−)-3-hydroxybutyric acid. While polymers such as ethylene-vinyl acetate and lactic acid-glycolic acid enable release of molecules for over 100 days, certain hydrogels release proteins for shorter time periods. When encapsulated antibodies remain in the body for a long time, they may denature or aggregate as a result of exposure to moisture at 37°C, resulting in a loss of biological activity and possible changes in immunogenicity. Rational strategies can be devised for stabilization depending on the mechanism involved. For example, if the aggregation mechanism is discovered to be intermolecular S-S bond formation through thio-disulfide interchange, stabilization may be achieved by modifying sulfhydryl residues, lyophilizing from acidic solutions, controlling moisture content, using appropriate additives, and developing specific polymer matrix compositions.


The anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibodies of the invention have various therapeutic and/or diagnostic utilities for a neurological disorder; a cardiovascular, endothelial or angiogenic disorder; an immunological disorder; an oncological disorder; an embryonic developmental disorder or lethality, or a metabolic abnormality. For example, anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibodies may be used in diagnostic assays for PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904,
PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346, e.g., detecting its expression (and in some cases, differential expression) in specific cells, tissues, or serum. Various diagnostic assay techniques known in the art may be used, such as competitive binding assays, direct or indirect sandwich assays and immunoprecipitation assays conducted in either heterogeneous or homogeneous phases [Zola, Monoclonal Antibodies: A Manual of Techniques, CRC Press, Inc. (1987) pp. 147-158]. The antibodies used in the diagnostic assays can be labeled with a detectable moiety. The detectable moiety should be capable of producing, either directly or indirectly, a detectable signal. For example, the detectable moiety may be a radioisotope, such as $^3$H, $^{14}$C, $^{32}$P, $^{35}$S, or $^{125}$I, a fluorescent or chemiluminescent compound, such as fluorescein isothiocyanate, rhodamine, or luciferin, or an enzyme, such as alkaline phosphatase, beta-galactosidase or horseradish peroxidase. Any method known in the art for conjugating the antibody to the detectable moiety may be employed, including those methods described by Hunter et al., *Nature*, 144:945 (1962); David et al., *Biochemistry*, 12:1014 (1974); Pain et al., *J. Immunol. Meth.*, 40:219 (1981); and Nygren, *J. Histochem. and Cytochem.*, 30:407 (1982).

Anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO99048, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibodies also are useful for the affinity purification of PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO99048, PRO10013, PRO9904, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides from recombinant cell culture or natural sources. In this process, the antibodies against PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO99048, PRO10013, PRO9904, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides are immobilized on a suitable support, such as a Sephadex resin or filter paper, using methods well known in the art. The immobilized antibody then is contacted with a sample containing the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO99048, PRO10013, PRO9904, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide to be purified, and thereafter the support is washed with a suitable solvent that will remove substantially all the material in the sample except the PRO218, PRO228, PRO271, PRO273, PRO295,
PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, which is bound to the immobilized antibody. Finally, the support is washed with another suitable solvent that will release the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide from the antibody.

The following examples are offered for illustrative purposes only, and are not intended to limit the scope of the present invention in any way.

All patent and literature references cited in the present specification are hereby incorporated by reference in their entirety.

EXAMPLES

Commercially available reagents referred to in the examples were used according to manufacturer's instructions unless otherwise indicated. The source of those cells identified in the following examples, and throughout the specification, by ATCC accession numbers is the American Type Culture Collection, Manassas, VA.

EXAMPLE 1: Extracellular Domain Homology Screening to Identify Novel Polypeptides and cDNA Encoding Therefor

The extracellular domain (ECD) sequences (including the secretion signal sequence, if any) from about 950 known secreted proteins from the Swiss-Prot public database were used to search EST databases. The EST databases included public databases (e.g., Dayhoff, GenBank), and proprietary databases (e.g. LIFSEQ™, Incyte Pharmaceuticals, Palo Alto, CA). The search was performed using the computer program BLAST or BLAST-2 (Altschul et al., Methods in Enzymology, 266:460-480 (1996)) as a comparison of the ECD protein sequences to a 6 frame translation of the EST sequences. Those comparisons with a BLAST score of 70 (or in some cases 90) or greater that did not encode known proteins were clustered and assembled into consensus DNA sequences with the program "phrap" (Phil Green, University of Washington, Seattle, WA).

Using this extracellular domain homology screen, consensus DNA sequences were assembled relative to the other identified EST sequences using phrap. In addition, the consensus DNA sequences obtained were often (but not always) extended using repeated cycles of BLAST or BLAST-2 and phrap to extend the consensus sequence as far as possible using the sources of EST sequences discussed above.

Based upon the consensus sequences obtained as described above, oligonucleotides were then synthesized and used to identify by PCR a cDNA library that contained the sequence of interest and for use as probes to isolate a clone of the full-length coding sequence for a PRO polypeptide. Forward and reverse PCR primers generally
range from 20 to 30 nucleotides and are often designed to give a PCR product of about 100-1000 bp in length. The probe sequences are typically 40-55 bp in length. In some cases, additional oligonucleotides are synthesized when the consensus sequence is greater than about 1-1.5kbp. In order to screen several libraries for a full-length clone, DNA from the libraries was screened by PCR amplification, as per Ausubel et al., Current Protocols in Molecular Biology, with the PCR primer pair. A positive library was then used to isolate clones encoding the gene of interest using the probe oligonucleotide and one of the primer pairs.

The cDNA libraries used to isolate the cDNA clones were constructed by standard methods using commercially available reagents such as those from Invitrogen, San Diego, CA. The cDNA was primed with oligo dT containing a NotI site, linked with blunt to SalI hemin kinase adaptors, cleaved with NotI, sized appropriately by gel electrophoresis, and cloned in a defined orientation into a suitable cloning vector (such as pBK or pKD; pRK5B is a precursor of pRK5D that does not contain the SfiI site; see, Holmes et al., Science, 253:1278-1280 (1991)) in the unique XhoI and NotI sites.

EXAMPLE 2: Isolation of cDNA clones by Amylase Screening

1. Preparation of oligo dT primed cDNA library

mRNA was isolated from a human tissue of interest using reagents and protocols from Invitrogen, San Diego, CA (Fast Track 2). This RNA was used to generate an oligo dT primed cDNA library in the vector pRK5D using reagents and protocols from Life Technologies, Gaithersburg, MD (Super Script Plasmid System). In this procedure, the double stranded cDNA was sized to greater than 1000 bp and the SalI/NotI linker cleaved cDNA was cloned into XhoI/NotI cleaved vector. pRK5D is a cloning vector that has an sp6 transcription initiation site followed by an SfiI restriction enzyme site preceding the XhoI/NotI cDNA cloning sites.

2. Preparation of random primed cDNA library

A secondary cDNA library was generated in order to preferentially represent the 5' ends of the primary cDNA clones. Sp6 RNA was generated from the primary library (described above), and this RNA was used to generate a random primed cDNA library in the vector pSST-AMY.0 using reagents and protocols from Life Technologies (Super Script Plasmid System, referenced above). In this procedure, the double stranded cDNA was sized to 500-1000 bp, linker cleaved blunt to NotI adaptors, cleaved with SfiI, and cloned into SfiI/NotI cleaved vector. pSST-AMY.0 is a cloning vector that has a yeast alcohol dehydrogenase promoter preceding the cDNA cloning sites and the mouse amylase sequence (the mature sequence without the secretion signal) followed by the yeast alcohol dehydrogenase terminator, after the cloning sites. Thus, cDNAs cloned into this vector that are fused in frame with amylase sequence will lead to the secretion of amylase from appropriately transfected yeast colonies.

3. Transformation and Detection

DNA from the library described in paragraph 2 above was chilled on ice to which was added electrocompetent DH10B bacteria (Life Technologies, 20 ml). The bacteria and vector mixture was then electroporated as recommended by the manufacturer. Subsequently, SOC media (Life Technologies, 1 ml) was added and the mixture was incubated at 37°C for 30 minutes. The transformants were then plated onto 20 standard 150 mm LB plates containing ampicillin and incubated for 16 hours (37°C). Positive colonies were scraped off
the plates and the DNA was isolated from the bacterial pellet using standard protocols, e.g. CsCl-gradient. The purified DNA was then carried on to the yeast protocols below.

The yeast methods were divided into three categories: (1) Transformation of yeast with the plasmid/cDNA combined vector; (2) Detection and isolation of yeast clones secreting amylase; and (3) PCR amplification of the insert directly from the yeast colony and purification of the DNA for sequencing and further analysis.

The yeast strain used was HD56-5A (ATCC-90785). This strain has the following genotype: MAT alpha, ura3-52, leu2-3, leu2-112, his3-11, his3-15, MAL1, SUC2, GAL1. Preferably, yeast mutants can be employed that have deficient post-translational pathways. Such mutants may have translocation deficient alleles in sec71, sec72, sec62, with truncated sec71 being most preferred. Alternatively, antagonists (including antisense nucleotides and/or ligands) which interfere with the normal operation of these genes, other proteins implicated in this post translation pathway (e.g., SEC61p, SEC72p, SEC62p, SEC63p, TDJ1p or SSA1p-4p) or the complex formation of these proteins may also be preferably employed in combination with the amylase-expressing yeast.

Transformation was performed based on the protocol outlined by Gietz et al., Nucl. Acid. Res., 20:1425 (1992). Transformed cells were then inoculated from agar into YEPD complex media broth (100 ml) and grown overnight at 30°C. The YEPD broth was prepared as described in Kaiser et al., Methods in Yeast Genetics, Cold Spring Harbor Press, Cold Spring Harbor, NY, p. 207 (1994). The overnight culture was then diluted to about 2 x 10^6 cells/ml (approx. OD_600=0.1) into fresh YEPD broth (500 ml) and regrown to 1 x 10^7 cells/ml (approx. OD_600=0.4-0.5).

The cells were then harvested and prepared for transformation by transfer into GS3 rotor bottles in a Sorval GS3 rotor at 5,000 rpm for 5 minutes, the supernatant discarded, and then resuspended into sterile water, and centrifuged again in 50 ml falcon tubes at 3,500 rpm in a Beckman GS-6KR centrifuge. The supernatant was discarded and the cells were subsequently washed with LiAc/TE (10 ml, 10 mM Tris-HCl, 1 mM EDTA pH 7.5, 100 mM Li_2OOCCH_3), and resuspended into LiAc/TE (2.5 ml).

Transformation took place by mixing the prepared cells (100 μl) with freshly denatured single stranded salmon testes DNA (Lofstrand Labs, Gaithersburg, MD) and transforming DNA (1 μg, vol. < 10 μl) in microfuge tubes. The mixture was mixed briefly by vortexing, then 40% PEG/TE (600 μl, 40% polyethylene glycol-4000, 10 mM Tris-HCl, 1 mM EDTA, 100 mM Li_2OOCCH_3, pH 7.5) was added. This mixture was gently mixed and incubated at 30°C while agitating for 30 minutes. The cells were then heat shocked at 42°C for 15 minutes, and the reaction vessel centrifuged in a microfuge at 12,000 rpm for 5-10 seconds, decanted and resuspended into TE (500 μl, 10 mM Tris-HCl, 1 mM EDTA pH 7.5) followed by recentrifugation. The cells were then diluted into TE (1 ml) and aliquots (200 μl) were spread onto the selective media previously prepared in 150 mm growth plates (VWR).

Alternatively, instead of multiple small reactions, the transformation was performed using a single, large scale reaction, wherein reagent amounts were scaled up accordingly.

The selective media used was a synthetic complete dextrose agar lacking uracil (SCD-Ura) prepared as described in Kaiser et al., Methods in Yeast Genetics, Cold Spring Harbor Press, Cold Spring Harbor, NY, p. 208-210 (1994). Transformants were grown at 30°C for 2-3 days.

The detection of colonies secreting amylase was performed by including red starch in the selective growth media. Starch was coupled to the red dye (Reactive Red-120, Sigma) as per the procedure described by Biely et
al., Anal. Biochem., 172:176-179 (1988). The coupled starch was incorporated into the SCD-Ura agar plates at a final concentration of 0.15% (w/v), and was buffered with potassium phosphate to a pH of 7.0 (50-100 mM final concentration).

The positive colonies were picked and streaked across fresh selective media (onto 150 mm plates) in order to obtain well isolated and identifiable single colonies. Well isolated single colonies positive for amylase secretion were detected by direct incorporation of red starch into buffered SCD-Ura agar. Positive colonies were determined by their ability to break down starch resulting in a clear halo around the positive colony visualized directly.

4. Isolation of DNA by PCR Amplification

When a positive colony was isolated, a portion of it was picked by a toothpick and diluted into sterile water (30 μl) in a 96 well plate. At this time, the positive colonies were either frozen and stored for subsequent analysis or immediately amplified. An aliquot of cells (5 μl) was used as a template for the PCR reaction in a 25 μl volume containing: 0.5 μl KlenTaq (Clontech, Palo Alto, CA); 4.0 μl 10 mM dNTP’s (Perkin Elmer-Cetus); 2.5 μl KlenTaq buffer (Clontech); 0.25 μl forward oligo 1; 0.25 μl reverse oligo 2; 12.5 μl distilled water. The sequence of the forward oligonucleotide 1 was:

5’-TGTTAACACGAGCCAGTTAAATAGACCTGCAATTTATTACT-3’ (SEQ ID NO:97)

The sequence of reverse oligonucleotide 2 was:

5’-CAGGAAACAGCTATGACCACCTGACACCTGCAAATCCATT-3’ (SEQ ID NO:98)

PCR was then performed as follows:

<table>
<thead>
<tr>
<th>a.</th>
<th>Denature</th>
<th>92°C, 5 minutes</th>
</tr>
</thead>
<tbody>
<tr>
<td>b.</td>
<td>3 cycles of:</td>
<td>Denature: 92°C, 30 seconds</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Anneal: 59°C, 30 seconds</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Extend: 72°C, 60 seconds</td>
</tr>
<tr>
<td>c.</td>
<td>3 cycles of:</td>
<td>Denature: 92°C, 30 seconds</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Anneal: 57°C, 30 seconds</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Extend: 72°C, 60 seconds</td>
</tr>
<tr>
<td>d.</td>
<td>25 cycles of:</td>
<td>Denature: 92°C, 30 seconds</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Anneal: 55°C, 30 seconds</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Extend: 72°C, 60 seconds</td>
</tr>
<tr>
<td>e.</td>
<td>Hold</td>
<td>4°C</td>
</tr>
</tbody>
</table>

The underlined regions of the oligonucleotides annealed to the ADH promoter region and the amylase region, respectively, and amplified a 307 bp region from vector pSSS-AMY.0 when no insert was present. Typically, the first 18 nucleotides of the 5’ end of these oligonucleotides contained annealing sites for the sequencing primers. Thus, the total product of the PCR reaction from an empty vector was 343 bp. However, signal sequence-fused cDNA resulted in considerably longer nucleotide sequences.

Following the PCR, an aliquot of the reaction (5 μl) was examined by agarose gel electrophoresis in a 1% agarose gel using a Tris-Borate-EDTA (TBE) buffering system as described by Sambrook et al., supra. Clones resulting in a single strong PCR product larger than 400 bp were further analyzed by DNA sequencing after purification with a 96 Qiaquick PCR clean-up column (Qiagen Inc., Chatsworth, CA).
EXAMPLE 3: Isolation of cDNA Clones Using Signal Algorithm Analysis

Various polypeptide-encoding nucleic acid sequences were identified by applying a proprietary signal sequence finding algorithm developed by Genentech, Inc. (South San Francisco, CA) upon ESTs as well as clustered and assembled EST fragments from public (e.g., GenBank) and/or private (LIFESEQ®, Incyte Pharmaceuticals, Inc., Palo Alto, CA) databases. The signal sequence algorithm computes a secretion signal score based on the character of the DNA nucleotides surrounding the first and optionally the second methionine codon(s) (ATG) at the 5'-end of the sequence or sequence fragment under consideration. The nucleotides following the first ATG must code for at least 35 unambiguous amino acids without any stop codons. If the first ATG has the required amino acids, the second is not examined. If neither meets the requirement, the candidate sequence is not scored. In order to determine whether the EST sequence contains an authentic signal sequence, the DNA and corresponding amino acid sequences surrounding the ATG codon are scored using a set of seven sensors (evaluation parameters) known to be associated with secretion signals. Use of this algorithm resulted in the identification of numerous polypeptide-encoding nucleic acid sequences.

Using the techniques described in Examples 1 to 3 above, numerous full-length cDNA clones were identified as encoding PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptides as disclosed herein. These cDNAs were then deposited under the terms of the Budapest Treaty with the American Type Culture Collection, 10801 University Blvd., Manassas, VA 20110-2209, USA (ATCC) as shown in Table 7 below. In addition, the sequence of DNA257845 encoding PRO5238 polypeptides was identified from GenBank accession no.: AF369794; the sequence of DNA82343 encoding PRO5733 polypeptides was identified from GenBank accession no.: BC017089; the sequence of DNA336882 encoding PRO90948 polypeptides was identified from GenBank accession no.: AK045869; the sequence of DNA184073 encoding PRO28694 polypeptides was identified from GenBank accession no.: AX281784; the sequence of DNA255255 encoding PRO50332 polypeptides was identified from GenBank accession no.: AB040120; and the sequence of DNA228002 encoding PRO38465 polypeptides was identified from GenBank accession no.: AF142409.

Table 7

<table>
<thead>
<tr>
<th>Material</th>
<th>ATCC Dep. No.</th>
<th>Deposit Date</th>
</tr>
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<tbody>
<tr>
<td>DNA30867-1335</td>
<td>209807</td>
<td>April 28, 1998</td>
</tr>
<tr>
<td>DNA33092-1202</td>
<td>209420</td>
<td>October 18, 1998</td>
</tr>
<tr>
<td>DNA39423-1182</td>
<td>209387</td>
<td>October 17, 1997</td>
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<td>DNA39523-1192</td>
<td>209424</td>
<td>October 31, 1997</td>
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<td>DNA38268-1188</td>
<td>209421</td>
<td>October 28, 1997</td>
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</tr>
<tr>
<td>DNA37140-1234</td>
<td>209489</td>
<td>November 12, 1997</td>
</tr>
</tbody>
</table>
These deposits were made under the provisions of the Budapest Treaty on the International Recognition of the Deposit of Microorganisms for the Purpose of Patent Procedure and the Regulations thereunder (Budapest Treaty). This assures maintenance of a viable culture of the deposit for 30 years from the date of deposit. The deposits will be made available by ATCC under the terms of the Budapest Treaty, and subject to an agreement
between Genentech, Inc. and ATCC, which assures permanent and unrestricted availability of the progeny of the culture of the deposit to the public upon issuance of the pertinent U.S. patent or upon laying open to the public of any U.S. or foreign patent application, whichever comes first, and assures availability of the progeny to one determined by the U.S. Commissioner of Patents and Trademarks to be entitled thereto according to 35 USC § 122 and the Commissioner's rules pursuant thereto (including 37 CFR § 1.14 with particular reference to 886 OG 638).

The assignee of the present application has agreed that if a culture of the materials on deposit should die or be lost or destroyed when cultivated under suitable conditions, the materials will be promptly replaced on notification with another of the same. Availability of the deposited material is not to be construed as a license to practice the invention in contravention of the rights granted under the authority of any government in accordance with its patent laws.

EXAMPLE 4: Isolation of cDNA clones Encoding Human PRO218 Polypeptides [UNQ192]

A consensus sequence was obtained relative to a variety of EST sequences as described in Example 1 above, wherein the consensus sequence obtained is herein designated DNA17411. Two proprietary Genentech EST sequences were employed in the consensus assembly. Based on the DNA17411 consensus sequence, oligonucleotides were synthesized: 1) to identify by PCR a cDNA library that contained the sequence of interest, and 2) for use as probes to isolate a clone of the full-length coding sequence for PRO218.

A pair of PCR primers (forward and reverse) were synthesized:

forward PCR primer 5’-AAGTGGAGCCGGAGCGTCTCC-3’ (SEQ ID NO:99);
reverse PCR primer 5’-TCGTTGTATGATAGTCGG-3’ (SEQ ID NO:100).

Additionally, a synthetic oligonucleotide hybridization probe was constructed from the consensus DNA17411 sequence which had the following nucleotide sequence:

hybridization probe
5’-ATGTGTTAAGACTATGATAGTCAGTGATTGTTACAGG-3’ (SEQ ID NO:101).

In order to screen several libraries for a source of a full-length clone, DNA from the libraries was screened by PCR amplification with the PCR primer pair identified above. A positive library was then used to isolate clones encoding the PRO218 gene using the probe oligonucleotide and one of the PCR primers. RNA for construction of the cDNA libraries was isolated from human fetal kidney tissue (LIB28).

DNA sequencing of the clones isolated as described above gave the full-length DNA sequence for PRO218 [herein designated as UNQ192 (DNA30867-1335)] (SEQ ID NO:1) and the derived protein sequence for PRO218.

The entire nucleotide sequence of UNQ192 (DNA30867-1335) is shown in Figure 1 (SEQ ID NO:1). Clone UNQ192 (DNA30867-1335) contains a single open reading frame with an apparent translational initiation site at nucleotide positions 150-152 and ending at the stop codon at nucleotide positions 1515-1517 (Figure 1). The predicted polypeptide precursor is 455 amino acids long (Figure 2; SEQ ID NO:2). The full-length PRO218 protein shown in Figure 4 has an estimated molecular weight of about 52,917 daltons and a pI of about 9.5. Clone UNQ192 (DNA30867-1335) has been deposited with the ATCC on April 28, 1998 with ATCC deposit number 209807. Regarding the sequence, it is understood that the deposited clone contains the correct sequence, and the sequences provided herein are based on known sequencing techniques.
Analysis of the amino acid sequence of the full-length PRO218 polypeptide suggests that PRO218 may be a novel transmembrane protein.

Still analyzing the amino acid sequence of SEQ ID NO:2, the putative signal peptide is at about amino acids 1 through 23 of SEQ ID NO:2. Transmembrane domains are potentially at about amino acids 37-55, 81-102, 150-168, 288-311, 338-356, 375-398, and 425-444 of SEQ ID NO:2. N-glycosylation sites are at about amino acids 67, 180, and 243 of SEQ ID NO:2. Eukaryotic cobalamin-binding protein is at about amino acids 151-160 of SEQ ID NO:2. The corresponding nucleotides can be routinely determined given the sequences provided herein.

EXAMPLE 5: Isolation of cDNA clones Encoding Human PRO228 Polypeptides [UNQ202]

A consensus DNA sequence was assembled relative to other EST sequences using phrap as described in Example 1 above. This consensus sequence is herein designated DNA28758. An EST proprietary to Genentech was employed in the consensus assembly. This EST is herein designated as DNA21951.

Based on the DNA28758 consensus sequence, oligonucleotides were synthesized: 1) to identify by PCR a cDNA library that contained the sequence of interest, and 2) for use as probes to isolate a clone of the full-length coding sequence for PRO228.

PCR primers (forward and reverse) were synthesized:

<table>
<thead>
<tr>
<th>Primer Type</th>
<th>Primer Sequence (5' to 3')</th>
<th>SEQ ID NO</th>
</tr>
</thead>
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<tr>
<td>forward PCR</td>
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<td>NO:102</td>
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<tr>
<td>forward PCR</td>
<td>GGAGTGAGAAACGCATCGG-3'</td>
<td>NO:103</td>
</tr>
<tr>
<td>forward PCR</td>
<td>CACCTGATACCATAGATGCG-3'</td>
<td>NO:104</td>
</tr>
<tr>
<td>reverse PCR</td>
<td>CGAGCTGAAATTCAATCG-3'</td>
<td>NO:105</td>
</tr>
<tr>
<td>reverse PCR</td>
<td>GGATCTCTCTGAGCTCAGG-3'</td>
<td>NO:106</td>
</tr>
<tr>
<td>reverse PCR</td>
<td>CCTAGTTGGATGATCCTGTA-3'</td>
<td>NO:107</td>
</tr>
</tbody>
</table>

Additionally, a synthetic oligonucleotide hybridization probe was constructed from the consensus DNA28758 sequence which had the following nucleotide sequence

hybridization probe

5'-ATGAGACCCACACCTCATGCGCTGTAATCCTGACACATTGCAATT-3' (SEQ ID NO:108)

In order to screen several libraries for a source of a full-length clone, DNA from the libraries was screened by PCR amplification with the PCR primer pairs identified above. A positive library was then used to isolate clones encoding the PRO228 gene using the probe oligonucleotide and one of the PCR primers.

RNA for construction of the cDNA libraries was isolated from human fetal kidney tissue.

DNA sequencing of the clones isolated as described above gave the full-length DNA sequence for PRO228 [herein designated as DNA33092-1202] (SEQ ID NO:3) and the derived protein sequence for PRO228.

The entire nucleotide sequence of DNA33092-1202 is shown in Figure 3 (SEQ ID NO:3). Clone DNA33092-1202 contains a single open reading frame with an apparent translational initiation site at nucleotide positions 24-26 of SEQ ID NO:3 and ending at the stop codon after nucleotide position 2093 of SEQ ID NO:3. The predicted polypeptide precursor is 690 amino acids long (Figure 4; SEQ ID NO:4). Clone DNA33092-1202 has been deposited with ATCC on October 18, 1997 and is assigned ATCC deposit no. ATCC 209420.

Analysis of the amino acid sequence of the full-length PRO228 polypeptide suggests that portions of it possess significant homology to the secretin-related proteins CD97 and EMR1 as well as the secretin member,
latrophilin, thereby indicating that PRO228 may be a new member of the secretin related proteins.

EXAMPLE 6: Isolation of cDNA clones Encoding Human PRO271 Polypeptides [UNQ238]

A consensus DNA sequence was assembled relative to other EST sequences using phrap as described in Example 1 above. This consensus sequence is herein designated DNA35737. Based on the DNA35737 consensus sequence, oligonucleotides were synthesized: 1) to identify by PCR a cDNA library that contained the sequence of interest, and 2) for use as probes to isolate a clone of the full-length coding sequence for PRO271.

Forward and reverse PCR primers were synthesized:

forward PCR primer 1 5'-TGCTTCGCTACTGCCCTC-3'  (SEQ ID NO:109)
forward PCR primer 2 5'-TTCCCTTGTGGGTGGAG-3'  (SEQ ID NO:110)
forward PCR primer 3 5'-AGGGCTGGAAAGCCAGTTT-3'  (SEQ ID NO:111)
reverse PCR primer 1 5'-AGCAGTGAGGAAATGCG-3'  (SEQ ID NO:112)
reverse PCR primer 2 5'-TGCACAAAGTACACACACCTGAGG-3'  (SEQ ID NO:113)

Additionally, a synthetic oligonucleotide hybridization probe was constructed from the consensus DNA35737 sequence which had the following nucleotide sequence

hybridization probe
5'-GATGCCACCGATCGCCAAGGTTGGAGACGCTGCTTCGCGCTGAAAG-3'  (SEQ ID NO:114)

In order to screen several libraries for a source of a full-length clone, DNA from the libraries was screened by PCR amplification with the PCR primer pair identified above. A positive library was then used to isolate clones encoding the PRO271 gene using the probe oligonucleotide and one of the PCR primers.

RNA for construction of the cDNA libraries was isolated from human fetal brain tissue.

DNA sequencing of the clones isolated as described above gave the full-length DNA sequence for PRO271 [herein designated as DNA39423-1182] (SEQ ID NO:5) and the derived protein sequence for PRO271.

The entire nucleotide sequence of DNA39423-1182 is shown in Figure 5 (SEQ ID NO:5). Clone DNA39423-1182 contains a single open reading frame with an apparent translational initiation site at nucleotide positions 101-103 and ending at the stop codon at nucleotide positions 1181-1183 (Figure 5). The predicted polypeptide precursor is 360 amino acids long (Figure 6; SEQ ID NO:6). Clone DNA39423-1182 has been deposited with ATCC and on October 17, 1997 and is assigned ATCC deposit no. ATCC 209387.

Analysis of the amino acid sequence of the full-length PRO271 polypeptide suggests that it possess significant homology to the proteoglycan link protein, thereby indicating that PRO271 may be a link protein homolog.

EXAMPLE 7: Isolation of cDNA clones Encoding Human PRO273 Polypeptides [UNQ240]

A consensus sequence was obtained relative to a variety of EST sequences as described in Example 1 above, wherein the consensus sequence obtained is herein designated DNA36465. Based on the DNA36465 consensus sequence, oligonucleotides were synthesized: 1) to identify by PCR a cDNA library that contained the sequence of interest, and 2) for use as probes to isolate a clone of the full-length coding sequence for PRO273.

A pair of PCR primers (forward and reverse) were synthesized:

forward PCR primer 5'-CAGCGCCCTCCCATGTCCCTG-3'  (SEQ ID NO:115)
reverse PCR primer 5'-TCCCAACTGTTGGAGTTCCCTCCC-3'  (SEQ ID NO:116)
Additionally, a synthetic oligonucleotide hybridization probe was constructed from the consensus DNA36465 sequence which had the following nucleotide sequence
hybridization probe
5'-CTCCGGTCAGCATGAGGCTTGGCGGCGCTGCTCTGCTGCTG-3' (SEQ ID NO:117)

In order to screen several libraries for a source of a full-length clone, DNA from the libraries was screened by PCR amplification with the PCR primer pair identified above. A positive library was then used to isolate clones encoding the PRO273 gene using the probe oligonucleotide and one of the PCR primers. RNA for construction of the cDNA libraries was isolated from human fetal kidney tissue.

DNA sequencing of the clones isolated as described above gave the full-length DNA sequence for PRO273 [herein designated as UNQ240 (DNA39523-1192)] (SEQ ID NO:7) and the derived protein sequence for PRO273.

The entire nucleotide sequence of UNQ240 (DNA39523-1192) is shown in Figure 7 (SEQ ID NO:7). Clone UNQ240 (DNA39523-1192) contains a single open reading frame with an apparent translational initiation site at nucleotide positions 167-169 and ending at the stop codon at nucleotide positions 500-502 (Figure 7). The predicted polypeptide precursor is 111 amino acids long (Figure 8; SEQ ID NO:8). Clone UNQ240 (DNA39523-1192) has been deposited with the ATCC on October 31, 1997 and is assigned ATCC number 209424. It is understood that the deposited clone contains the actual sequence and that the sequences provided herein are merely representative based on current sequencing techniques. Moreover, given the sequences provided herein and knowledge of the universal genetic code, the corresponding nucleotides for any given amino acid can be routinely identified and vice versa.

Analysis of the amino acid sequence of the full-length PRO273 polypeptide suggests that portions of it possess sequence identity with human macrophage inflammatory protein-2, cytokine-induced neutrophil chemoattractant 2, and neutrophil chemotactic factor 2-beta, thereby indicating that PRO273 is a novel chemokine.

As discussed further below, the cDNA was subeloned into a baculovirus vector and expressed in insect cells as a C-terminally tagged IgG fusion protein. N-terminal sequencing of the resultant protein identified the signal sequence cleavage site, yielding a mature polypeptide of 77 amino acids. The mature sequence, showing 31-40% identity to other human CXC chemokines, includes the four canonical cysteine residues but lacks the ELR motif. Northern analysis demonstrates expression at least in the small intestine, colon, spleen, lymph node and kidney. By in situ hybridization, also described in detail below, mRNA is localized to the lamina propria of intestinal villi and to renal tubules.

EXAMPLE 8: Isolation of cDNA clones Encoding Human PRO295 Polypeptides [UNQ258]

A consensus DNA sequence was assembled relative to other EST sequences using phrap as described in Example 1 above. This consensus sequence is herein designated DNA35814. Based on the DNA35814 consensus sequence, oligonucleotides were synthesized: 1) to identify by PCR a cDNA library that contained the sequence of interest, and 2) for use as probes to isolate a clone of the full-length coding sequence for PRO295.

Forward and reverse PCR primers were synthesized:
forward PCR primer (f1) 5'-GCAGAGCGGAGATGCAGGGCTTG-3' (SEQ ID NO:118)
forward PCR primer (f2) 5'-CCCAGCATGACTGCGCCAG-3' (SEQ ID NO:119)
forward PCR primer (.f3) 5′-TTGGCACGCTTACATGGAGG-3′ (SEQ ID NO:120)
forward PCR primer (.f4) 5′-CCTGGGAATAATGCAAC-3′ (SEQ ID NO:121)
reverse PCR primer (.r1) 5′-CTGCCAGCTTGGGCCACCTCCC-3′ (SEQ ID NO:122)
Additionally, a synthetic oligonucleotide hybridization probe was constructed from the consensus DNA35814 sequence which had the following nucleotide sequence
hybridization probe
5′-GGCTCCTAGTACCGGCAGGAGGCGCCACCCCTCAATGAGATG-3′ (SEQ ID NO:123)

In order to screen several libraries for a source of a full-length clone, DNA from the libraries was screened by PCR amplification with the PCR primer pairs identified above. A positive library was then used to isolate clones encoding the PRO295 gene using the probe oligonucleotide and one of the PCR primers.

RNA for construction of the cDNA libraries was isolated from human fetal lung tissue.

DNA sequencing of the clones isolated as described above gave the full-length DNA sequence for PRO295 [herein designated as DNA38268-1188] (SEQ ID NO:9) and the derived protein sequence for PRO295.

The entire nucleotide sequence of DNA38268-1188 is shown in Figure 9 (SEQ ID NO:9). Clone DNA38268-1188 contains a single open reading frame with an apparent translational initiation site at nucleotide positions 153-155 and ending at the stop codon at nucleotide positions 1202-1204 (Figure 9). The predicted polypeptide precursor is 350 amino acids long (Figure 10; SEQ ID NO:10). Clone DNA38268-1188 has been deposited with ATCC on October 28, 1997 and is assigned ATCC deposit no. 209421.

Analysis of the amino acid sequence of the full-length PRO295 polypeptide suggests that portions of it possess significant homology to the integrin proteins, thereby indicating that PRO295 may be a novel integrin.

EXAMPLE 9: Isolation of cDNA clones Encoding Human PRO302 Polypeptides [UNQ265]

Consensus DNA sequences were assembled relative to other EST sequences using phrap as described in Example 1 above. These consensus sequences are herein designated DNA35953. Based on the DNA35953 consensus sequence, oligonucleotides were synthesized: 1) to identify by PCR a cDNA library that contained the sequence of interest, and 2) for use as probes to isolate a clone of the full-length coding sequence for PRO302.

PCR primers (forward and reverse) were synthesized:
forward PCR primer 1 5′-GTCCGCAAGGATGCTACATGTTC-3′ (SEQ ID NO:124)
forward PCR primer 2 5′-GCAGAGGTGTCTAAGGTTG-3′ (SEQ ID NO:125)
reverse PCR primer 5′-AGCTCTAGACCAATGCAGCTTCC-3′ (SEQ ID NO:126)
Also, a synthetic oligonucleotide hybridization probe was constructed from the consensus DNA35953 sequence which had the following nucleotide sequence
hybridization probe
5′-GCCACCAACTCTGCAAGAACTTCTCAGAACTGCCCCCTTGGTCATG-3′ (SEQ ID NO:127)

In order to screen several libraries for a source of a full-length clone, DNA from the libraries was screened by PCR amplification with the PCR primer pairs identified above. A positive library was then used to isolate clones encoding the PRO302 gene using the probe oligonucleotide and one of the PCR primers.

RNA for construction of the cDNA libraries was isolated from human fetal kidney tissue (LIB228).

DNA sequencing of the clones isolated as described above gave the full-length DNA sequence for
PRO302 [herein designated as DNA40370-1217] (SEQ ID NO: 11) and the derived protein sequence for PRO302.

The entire nucleotide sequence of DNA40370-1217 is shown in Figure 11 (SEQ ID NO:11). Clone DNA40370-1217 contains a single open reading frame with an apparent translational initiation site at nucleotide positions 34-36 and ending at the stop codon at nucleotide positions 1390-1392 (Figure 11). The predicted polypeptide precursor is 452 amino acids long (Figure 12; SEQ ID NO:12). Various unique aspects of the PRO302 protein are shown in Figure 12. Clone DNA40370-1217 has been deposited with the ATCC on November 21, 1997 and is assigned ATCC deposit no. ATCC 209485.

EXAMPLE 10: Isolation of cDNA clones Encoding Human PRO305 Polypeptides [UNO268]

The extracellular domain (ECD) sequences (including the secretion signal, if any) of from about 950 known secreted proteins from the Swiss-Prot public protein database were used to search expressed sequence tag (EST) databases. The EST databases included public EST databases (e.g., GenBank) and a proprietary EST DNA database (LIFESEQTM, Incyte Pharmaceuticals, Palo Alto, CA). The search was performed using the computer program BLAST or BLAST2 (Altschul et al., Methods in Enzymology 266:460-480 (1996)) as a comparison of the ECD protein sequences to a 6 frame translation of the EST sequence. Those comparisons resulting in a BLAST score of 70 (or in some cases 90) or greater that did not encode known proteins were clustered and assembled into consensus DNA sequences with the program "phrap" (Phil Green, University of Washington, Seattle, Washington).

A consensus DNA sequence was assembled relative to other EST sequences using phrap. This consensus sequence is herein designated DNA36440-from dna. In some cases, the consensus DNA sequence was extended using repeated cycles of BLAST and phrap to extend the consensus sequence as far as possible using the sources of EST sequences discussed above (the initial sequence used is designated DNA36440-init).

Based on the DNA36440-from dna consensus sequence, oligonucleotides were synthesized: 1) to identify by PCR a cDNA library that contained the sequence of interest, and 2) for use as probes to isolate a clone of the full-length coding sequence for PRO305. Forward and reverse PCR primers generally range from 20 to 30 nucleotides and are often designed to give a PCR product of about 100-1000 bp in length. The probe sequences are typically 40-55 bp in length. In some cases, additional oligonucleotides are synthesized when the consensus sequence is greater than about 1.5kbp. In order to screen several libraries for a full-length clone, DNA from the libraries was screened by PCR amplification, as per Ausubel et al., Current Protocols in Molecular Biology, with the PCR primer pair. A positive library was then used to isolate clones encoding the gene of interest by the in vivo cloning procedure using the probe oligonucleotide and one of the primer pairs.

A pair of PCR primers (forward and reverse) were synthesized:
forward PCR primer 5'-TGCGACGGCGTGTGGTTTTGAAAC-3' (SEQ ID NO:128)
reverse PCR primer 5'-AAAGCATTCTGGCCATTGTGAAG-3' (SEQ ID NO:129)

Additionally, a synthetic oligonucleotide hybridization probe was constructed from the consensus DNA36440-from dna sequence which had the following nucleotide sequence

hybridization probe
5'-CGCTCGTCTGGCCTTTTGTCTGGGAATAGCCTCCGCTGTTC-3' (SEQ ID NO:130)

In order to screen several libraries for a source of a full-length clone, DNA from the libraries was
screened by PCR amplification with the PCR primer pair identified above. A positive library was then used to isolate clones encoding the PRO305 gene using the probe oligonucleotide and one of the PCR primers.

RNA for construction of the cDNA libraries was isolated from human fetal lung tissue. The cDNA libraries used to isolate the cDNA clones were constructed by standard methods using commercially available reagents such as those from Invitrogen, San Diego, CA. The cDNA was primed with oligo dT containing a NotI site, linked with blunt to SalI hemikinased adaptors, cleaved with NotI, sized appropriately by gel electrophoresis, and cloned in a defined orientation into a suitable cloning vector (such as pRKIB or pRKD; pRK5B is a precursor of pRK5D that does not contain the SfiI site; see, Holmes et al., Science, 253:1278-1280 (1991)) in the unique XhoI and NotI sites.

DNA sequencing of the clones isolated as described above gave the full-length DNA sequence for PRO305 [herein designated as UNQ268 (DNA40619-seqmin)] (SEQ ID NO:13) and the derived protein sequence for PRO305.

The entire nucleotide sequence of UNQ268 (DNA40619-seqmin) is shown in Figure 13 (SEQ ID NO:13). Clone UNQ268 (DNA40619-seqmin) contains a single open reading frame with an apparent translational initiation site at nucleotide positions 251-253 and ending at the stop codon at nucleotide positions 1253-1255 (Figure 13). The predicted polypeptide precursor is 334 amino acids long (Figure 14; SEQ ID NO:14). Clone UNQ268 (DNA40619-seqmin) has been deposited with ATCC on December 10, 1997 and is assigned ATCC deposit no. 209525.

Analysis of the amino acid sequence of the full-length PRO305 polypeptide suggests that portions of it possess significant homology to the human procathepsin L protein thereby indicating that PRO305 is a novel member of the cathepsin family.

Analysis of the amino acid sequence of Figure 14 (SEQ ID NO:14) shows the following characteristics. The signal peptide is from amino acids 1 through 17. The start of the mature peptide begins with amino acid 18. The cysteine proteases cysteine active site is from amino acids 132 through 143. The cysteine proteases histidine active site is from amino acids 275 through 285. Potential N-glycosylation sites are at amino acids 221 and 292. The active site by homology to "CATL-PIG" is from amino acids 301 through 334.

EXAMPLE 11: Isolation of cDNA clones Encoding Human PRO326 Polypeptides [UNQ287]

A consensus DNA sequence was assembled relative to other EST sequences using phrap as described in Example 1 above. This consensus sequence is herein designated DNA36685. Based on the DNA36685 consensus sequence, and Incyte EST sequence no. 2228990, oligonucleotides were synthesized: 1) to identify by PCR a cDNA library that contained the sequence of interest, and 2) for use as probes to isolate a clone of the full-length coding sequence for PRO326.

Forward and reverse PCR primers were synthesized for the determination of PRO326:

<table>
<thead>
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<th>Primer Type</th>
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<th>SEQ ID No.</th>
</tr>
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<tbody>
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<td>Forward</td>
<td>5'-ACTCCAAGGAAATCCGGATCCGTTC-3'</td>
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</tr>
<tr>
<td>Reverse</td>
<td>5'-TTAGCAGCTGAGGATGGGCCAACA-3'</td>
<td>132</td>
</tr>
</tbody>
</table>

Additionally, a synthetic oligonucleotide hybridization probe was constructed for the determination of PRO331 which had the following nucleotide sequence hybridization probe
In order to screen several libraries for a source of a full-length clone, DNA from the libraries was screened by PCR amplification with the PCR primer pairs identified above. A positive library was then used to isolate clones encoding the PRO326 gene using the probe oligonucleotide and one of the PCR primers.

DNA sequencing of the cDNA libraries was isolated from human fetal kidney tissue.

The entire nucleotide sequences is shown in Figure 19, deposited with the ATCC on November 21, 1997 and is assigned ATCC deposit number 209489.

Analysis of the amino acid sequence of the full-length PRO326 polypeptide suggests that portions of it possess significant homology to the LIG-1 protein, thereby indicating that PRO326 may be a novel LIG-1-related protein.

**Example 12: Isolation of cDNA clones Encoding Human PRO386 Polypeptides [UNQ326]**

A consensus sequence was obtained relative to a variety of EST sequences as described in Example 1 above, wherein the consensus sequence obtained is herein designated DNA40674. Two proprietary Genentech EST sequences were employed in the consensus sequence assembly, wherein those EST sequences are herein designated DNA23350 and DNA23536. Based on the DNA40674 consensus sequence, oligonucleotides were synthesized: 1) to identify by PCR a cDNA library that contained the sequence of interest, and 2) for use as probes to isolate a clone of the full-length coding sequence for PRO386.

A pair of PCR primers (forward and reverse) were synthesized:

Forward PCR primer 5'-ACGGAGCATGGAGGTCACAGTC-3' (SEQ ID NO:134)

Reverse PCR primer 5'-GCACGTTTCTCACGATCCGACAGACGACG-3' (SEQ ID NO:135)

Additionally, a synthetic oligonucleotide hybridization probe was constructed from the consensus DNA40674 sequence which had the following nucleotide sequence:

5'-CGCCCTGACCTGACCTTCAACTCCTGCTACAGTAACCACAAACAGTT-3' (SEQ ID NO:136)

In order to screen several libraries for a source of a full-length clone, DNA from the libraries was screened by PCR amplification with the PCR primer pair identified above. A positive library was then used to isolate clones encoding the PRO386 gene using the probe oligonucleotide and one of the PCR primers. RNA for construction of the cDNA libraries was isolated from human fetal brain tissue (LIB153).

DNA sequencing of the clones isolated as described above gave the full-length DNA sequence for PRO386 [herein designated as UNQ326 (DNA45415-1318)] (SEQ ID NO:17) and the derived protein sequence for PRO386.

The entire nucleotide sequence of UNQ326 (DNA45415-1318) is shown in Figure 17 (SEQ ID NO:17). Clone UNQ326 (DNA45415-1318) contains a single open reading frame with an apparent translational initiation site at nucleotide positions 146-148 and ending at the stop codon at nucleotide positions 791-793 (Figure 17).
predicted polypeptide precursor is 215 amino acids long (Figure 18; SEQ ID NO:18). The full-length PRO386 protein shown in Figure 18 has an estimated molecular weight of about 24,326 daltons and a pI of about 6.32. Analysis of the full-length PRO386 sequence shown in Figure 18 (SEQ ID NO:18) evidences the presence of the following: a signal peptide from about amino acid 1 to about amino acid 20, a transmembrane domain from about amino acid 161 to about amino acid 179, an immunoglobulin-like fold from about amino acid 83 to about amino acid 127 and potential N-glycosylation sites from about amino acid 42 to about amino acid 45, from about amino acid 66 to about amino acid 69 and from about amino acid 74 to about amino acid 77. Clone UNQ326 (DNA45415-1318) has been deposited with ATCC on April 28, 1998 and is assigned ATCC deposit no. 209810.

Analysis of the amino acid sequence of the full-length PRO386 polypeptide suggests that it possesses significant sequence similarity to the sodium channel beta-2 subunit, thereby indicating that PRO386 is a novel homolog thereof. More specifically, an analysis of the Dayhoff database (version 35.45 SwissProt 35) evidenced significant homology between the PRO386 amino acid sequence and the following Dayhoff sequences, A57843, MYP0_HUMAN, GEN14531, JC4024, HS46KDA_1, HSU90716_1, D86996_2, MUSIGLVD_1, DMU42768_1 and S19247.

EXAMPLE 13: Isolation of cDNA clones Encoding Human PRO655 Polypeptides [UNQ360]

An expressed sequence tag (EST) DNA database (LIFESEQTM, Incyte Pharmaceuticals, Palo Alto, CA) was searched and an EST was identified which showed homology to interferon-ε. Possible homology was noted between Incyte EST 3728969 (subsequently renamed as DNA49668) and mammalian alpha interferons, in particular IFN-14. The homology was confirmed by inspection.

The following PCR primers and oligonucleotide probe were synthesized:

49668.r1:
TCTCTGCTTCCAGTCCCATGAGTGC (SEQ ID NO:137)

49668.r2:
GCTTCCAGTCCCATGAGTGCCTTCTAGG (SEQ ID NO:138)

49668.p1:
G GCCATTCTCCATGAGTGCCTCAGCA ATCTTCAGCCTTCAGGGCAA (SEQ ID NO:139)

In order to screen several libraries for a source of a full-length clone, DNA from the libraries was screened using the r1 and r2 probes identified above. A positive library was then used to isolate clones encoding the IFN-ε-encoding gene using the probe oligonucleotide.

Three million clones from a size selected (500-4000 bp) oligo dT primed cDNA library from human small intestine (LIB 99) constructed in a pRK5-based vector screened by hybridization. The cDNA libraries used to isolate the cDNA clones were constructed by standard methods using commercially available reagents such as those from Invitrogen, San Diego, CA. The cDNA was primed with oligo dT containing a NotI site, linked with blunt to SalI hemikinased adaptors, cleaved with NotI, sized appropriately by gel electrophoresis, and cloned in a defined orientation into a suitable cloning vector (such as pRKB or pRKD; pRK518 is a precursor of pRK5D that does not contain the SflI site; see, Holmes et al., Science, 253:1278-1280 (1991)) in the unique XhoI and NotI sites. Only one positive clone was found out of 3.6 x 106 cfu. The clone was sequenced in both directions and was found to cover the entire reading frame (ORF). A BAC clone (F480) was identified by screening a BAC array
DEMANDE OU BREVET VOLUMINEUX

LA PRÉSENTE PARTIE DE CETTE DEMANDE OU CE BREVET COMPREND PLUS D’UN TOME.

CECI EST LE TOME 1 DE 2
CONTENANT LES PAGES 1 À 213

NOTE : Pour les tomes additionnels, veuillez contacter le Bureau canadien des brevets

JUMBO APPLICATIONS/PATENTS

THIS SECTION OF THE APPLICATION/PATENT CONTAINS MORE THAN ONE VOLUME

THIS IS VOLUME 1 OF 2
CONTAINING PAGES 1 TO 213

NOTE: For additional volumes, please contact the Canadian Patent Office

NOM DU FICHIER / FILE NAME :

NOTE POUR LE TOME / VOLUME NOTE:
WHAT IS CLAIMED IS:

1. A method of identifying a phenotype associated with a disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO1017, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO1885, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising:
   
   (a) providing a non-human transgenic animal whose genome comprises a disruption of a gene which is an ortholog of a human gene that encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO1017, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide;

   (b) measuring a physiological characteristic of the non-human transgenic animal; and

   (c) comparing the measured physiological characteristic with that of a gender matched wild-type animal, wherein the physiological characteristic of the non-human transgenic animal that differs from the physiological characteristic of the wild-type animal is identified as a phenotype resulting from the gene disruption in the non-human transgenic animal.

2. The method of Claim 1, wherein the non-human transgenic animal is heterozygous for the disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO1017, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide.

3. The method of Claim 1, wherein the phenotype exhibited by the non-human transgenic animal as compared with gender matched wild-type littermates is at least one of the following: a neurological disorder; a cardiovascular, endothelial or angiogenic disorder; an eye abnormality; an immunological disorder; an oncological disorder; a bone metabolic abnormality or disorder; a lipid metabolic disorder; or a developmental abnormality.

4. The method of Claim 3, wherein the neurological disorder is an increased anxiety-like response during open field activity testing.
5. The method of Claim 3, wherein the neurological disorder is a decreased anxiety-like response during open field activity testing.

6. The method of Claim 3, wherein the neurological disorder is an abnormal circadian rhythm during home-cage activity testing.

7. The method of Claim 3, wherein the neurological disorder is an enhanced motor coordination during inverted screen testing.

8. The method of Claim 3, wherein the neurological disorder is an impaired motor coordination during inverted screen testing.

9. The method of Claim 3, wherein the neurological disorder is depression, generalized anxiety disorders, attention deficit disorder, sleep disorder, hyperactivity disorder, obsessive compulsive disorder, schizophrenia, cognitive disorders, hyperalgesia or sensory disorders.

10. The method of Claim 3, wherein the eye abnormality is a retinal abnormality.

11. The method of Claim 3, wherein the eye abnormality is consistent with vision problems or blindness.

12. The method of Claim 10, wherein the retinal abnormality is consistent with retinitis pigmentosa.

13. The method of Claim 10, wherein the retinal abnormality is characterized by retinal degeneration or retinal dysplasia.

14. The method of Claim 10, wherein the retinal abnormality is consistent with retinal dysplasia, various retinopathies, including retinopathy of prematurity, retrolental fibroplasia, neovascular glaucoma, age-related macular degeneration, diabetic macular edema, corneal neovascularization, corneal graft neovascularization, corneal graft rejection, retinal/choroidal neovascularization, neovascularization of the angle (rubeosis), ocular neovascular disease, vascular restenosis, arteriovenous malformations (AVM), meningioma, hemangioma, angiofibroma, thyroid hyperplasias (including Grave's disease), corneal and other tissue transplantation, retinal artery obstruction or occlusion; retinal degeneration causing secondary atrophy of the retinal vasculature, retinitis pigmentosa, macular dystrophies, Stargardt’s disease, congenital stationary night blindness, choroideremia, gyrate atrophy, Leber’s congenital amaurosis, retinoschisis disorders, Wagner’s syndrome, Usher syndromes, Zellweger syndrome, Saldino-Mainzer syndrome, Senior-Loken syndrome, Bardet-Biedl syndrome, Alport's syndrome, Alstrom's syndrome, Cockayne's syndrome, dysplasia spondyloepiphysaria congenita, Flynn-Aird syndrome, Friedreich ataxia, Hallgren syndrome, Marshall syndrome, Albers-Schönberg disease, Refsum's disease, Kearns-Sayre syndrome, Waardenburg's syndrome, Alagille syndrome, myotonic dystrophy, olivopontocerebellar atrophy, Pierre-Marie dunsdrome, Stickler syndrome, carotinemia, cystinosis, Wolfram syndrome,
Bassen-Kornzweig syndrome, abetalipoproteinemia, incontinentia pigmenti, Batten's disease, mucopolysaccharidoses, homocystinuria, or mannosidosis.

15. The method of Claim 3, wherein the eye abnormality is a cataract.

16. The method of Claim 15, wherein the cataract is consistent with systemic diseases such as human Down's syndrome, Hallerman-Streiff syndrome, Lowe syndrome, galactosemia, Marfan syndrome, Trisomy 13-15, Alport syndrome, myotonic dystrophy, Fabry disease, hypoparathyroidism or Conradi syndrome.

17. The method of Claim 3, wherein the developmental abnormality comprises embryonic lethality or reduced viability.

18. The method of Claim 3, wherein the cardiovascular, endothelial or angiogenic disorders are arterial diseases, such as diabetes mellitus; papilledema; optic atrophy; atherosclerosis; angina; myocardial infarctions such as acute myocardial infarctions, cardiac hypertrophy, and heart failure such as congestive heart failure; hypertension; inflammatory vasculitides; Reynaud's disease and Reynaud's phenomenon; aneurysms and arterial restenosis; venous and lymphatic disorders such as thrombophlebitis, lymphangitis, and lymphedema; peripheral vascular disease; cancer such as vascular tumors, e.g., hemangioma (capillary and cavernous), glomus tumors, telangiectasia, bacillary angiomatosis, hemangioendothelioma, angiosarcoma, haemangiopericytoma, Kaposi's sarcoma, lymphangioma, and lymphangiosarcoma; tumor angiogenesis; trauma such as wounds, burns, and other injured tissue, implant fixation, scarring; ischemia reperfusion injury; rheumatoid arthritis; cerebrovascular disease; renal diseases such as acute renal failure, or osteoporosis.

19. The method of Claim 3, wherein the immunological disorders are systemic lupus erythematosus; rheumatoid arthritis; juvenile chronic arthritis; spondyloarthropathies; systemic sclerosis (scleroderma); idiopathic inflammatory myopathies (dermatomyositis, polymyositis); Sjögren's syndrome; systemic vasculitis; sarcoidosis; autoimmune hemolytic anemia (immune pancytopenia, paroxysmal nocturnal hemoglobinuria); autoimmune thrombocytophenia (idiopathic thrombocytopenic purpura, immune-mediated thrombocytopenia); thyroiditis (Grave's disease, Hashimoto's thyroiditis, juvenile lymphocytic thyroiditis, atrophic thyroiditis); diabetes mellitus; immune-mediated renal disease (glomerulonephritis, tubulointerstitial nephritis); demyelinating diseases of the central and peripheral nervous systems such as multiple sclerosis, idiopathic demyelinating polyneuropathy or Guillain-Barré syndrome, and chronic inflammatory demyelinating polyneuropathy; hepatobiliary diseases such as infectious hepatitis (hepatitis A, B, C, D, E and other non-hepatotropic viruses), autoimmune chronic active hepatitis, primary biliary cirrhosis, granulomatous hepatitis, and sclerosing cholangitis; inflammatory bowel disease (ulcerative colitis; Crohn's disease); gluten-sensitive enteropathy, and Whipple's disease; autoimmune or immune-mediated skin diseases including bullous skin diseases, erythema multiforme and contact dermatitis, psoriasis; allergic diseases such as asthma, allergic rhinitis, atopic dermatitis, food hypersensitivity and urticaria;
immunologic diseases of the lung such as eosinophilic pneumonias, idiopathic pulmonary fibrosis and hypersensitivity pneumonitis; or transplantation associated diseases including graft rejection and graft-versus-host disease.

20. The method of Claim 3, wherein the bone metabolic abnormality or disorder is arthritis, osteoporosis or osteopetrosis.

21. The method of Claim 1, wherein the non-human transgenic animal exhibits at least one of the following physiological characteristics compared with gender matched wild-type littermates: increased anxiety-like response during open field testing; hyperactivity during open field testing; decreased anxiety during open field testing; decreased locomotor activity during open field testing; abnormal circadian rhythm during home-cage activity testing (low activity during the light phase; altered sleep/wake cycle); abnormal circadian rhythm during home-cage activity testing including decreased ambulatory counts; hypoactivity with no circadian rhythm; abnormal circadian rhythm during home-cage activity testing including increased ambulatory counts; decreased rearing; increased sensitivity to stress induced hyperthermia (increased anxiety); impaired motor coordination during inverted screen testing; head tilt and retropulsion; increased prepulse inhibition response indicating enhanced sensorimotor gating/attention; decreased startle response during prepulse inhibition testing; no startle response indicating deafness or impaired hearing; decreased prepulse inhibition with impaired sensorimotor gating/attention; increased latency to respond in hot plate testing; decreased latency to respond in hot plate testing; ophthalmological abnormalities; impaired vision; white deposits of optic disc region; ocular infection and neutrophilia; bilateral optic disc lesion; decreased tear production; decreased heart rate; increased mean systolic blood pressure; decreased mean systolic blood pressure; increased mean fasting serum glucose levels; decreased mean serum glucose levels; increased mean serum cholesterol levels; decreased mean serum cholesterol levels; increased mean serum triglyceride levels; decreased mean serum triglyceride levels; impaired glucose tolerance; increased mean serum albumin, alanine amino transferase and phosphorus levels; increased mean serum alkaline phosphatase levels; urinary nitrites present; increased total white blood cell (WBC) count; decreased total white blood cell (WBC) count and absolute neutrophil count; increased mean absolute neutrophil count; increased mean absolute lymphocyte count; increased mean platelet count; increased mean red cell distribution width; decreased mean platelet count; reduced percentage of CD4 spleen thymocytes; decreased percentages of CD4 cells in the periphery resulting in increased percentages of B cells in lymph organs; CD4 cells exhibit a more activated/memory phenotype (CD62Llow, CD44hi); developmental defect in CD4+ cells; decreased percentages of CD4 cells and increased percentages of B cells in blood; decreased percentages of CD4 cells and increased percentages of B cells in tissues; increase in percentages of B cells in Peyer’s patches; decreased germinal center, isotype-switched B cells in Peyer’s patches (CD38low:IgM negative); decreased CD23 intensity in spleen; increased mean percentages of B220 Med/CD23- cells and B220+/CD11b-Low/CD23- cells in peritoneal lavage; increased mean percentages of B cells in peripheral blood; decreased CD4 and CD8 T cells and increased B cells; increase in peritoneal B cells; reduction in CD11b-Hi cells in peritoneal cavity; decreased mean CD4 to CD8 ratio in spleen; decreased CD8 cells; decreased mean percentages of B220+/CD23+ cells and B220+/CD11bLow/CD23- cells in peritoneal lavage; increased mean serum IgG1 response to ovalbumin challenge; increased mean serum IgG2a response to ovalbumin.
challenge; increased mean serum IL-6 response to LPS challenge; increased mean serum TNF alpha response to LPS challenge; increased mean serum MCP-1 response to LPS challenge; increased mean serum IgM level; increased mean serum IgA; increase mean serum IgG1; increased mean serum IgG2a; increased mean serum IgG2b; decreased mean serum IgG1 response to ovalbumin challenge; decreased mean serum IgG2a response to ovalbumin challenge; failure in ovalbumin response; decreased mean serum IgA level; decreased mean serum IgG2a level; decreased skin fibroblast proliferation rate; increased mean percent of total body fat and total fat mass; increased mean body weight; increased mean body length; increased total tissue mass (TTM); increased bone mineral density (BMD); increase in bone mineral content (BMC); increased mean femoral midshaft cortical thickness; decreased mean percent of total body fat and total fat mass; decreased mean body weight; decreased mean body length; decreased mean body weight and length in heterozygotes; decreased total tissue mass (TTM); decreased lean body mass (LBM); decreased femoral bone mineral density (BMD); decreased vertebral bone mineral density (BMD); decreased bone mineral density (BMD) in total body; decreased bone mineral content (BMC); decreased bone mineral density index; decreased volumetric bone mineral density (vBMD); decreased mean femoral midshaft cortical thickness; decreased mean femoral midshaft cross-sectional area; decreased mean vertebral trabecular bone volume, number and connectivity density; osteopenosis; osteoporosis; moderate kidney hydroureter; hydrourephalus; enlarged liver; induced in activated T cells; induced in activated NK cells and dendritic cells; myeloid B cell expression; hyperplasia of sebaceous glands and multifocal hyperplasia of the epidermis (acanthosis and hyperkeratosis); moderate dermatitis; increased extramedullary hematopoiesis in liver and spleen; myeloid hyperplasia of the bone marrow; encephalitis due to Group B streptococcus; meningitis due to E. Coli infection; lymphocytic infiltrates in salivary glands, pancreas and lungs; poor breeders requiring foster mothers; decreased litter size; homozygous mice were small and dehydrated; vacuolar degeneration of testes resulting in decreased sperm production and infertility; defective spermatogenesis in the testes; hypospermia and defective spermatozoa in the epididymus; male infertility; decreased testes weight; growth retardation; small mice and failure to thrive; reduced viability; reduced viability with situs inversus; and homozygous embryonic lethality.

22. An isolated cell derived from a non-human transgenic animal whose genome comprises a disruption of a gene which is an ortholog of a human gene that encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide.

23. The isolated cell of Claim 22 which is a murine cell.

24. The isolated cell of Claim 23, wherein the murine cell is an embryonic stem cell.

25. The isolated cell of Claim 22, wherein the non-human transgenic animal exhibits at least one of the following phenotypes compared with gender matched wild-type littermates: a neurological disorder; a
cardiovascular, endothelial or angiogenic disorder; an eye abnormality; an immunological disorder; an oncological disorder; a bone metabolic abnormality or disorder; a lipid metabolic disorder; or a developmental abnormality.

26. A method of identifying an agent that modulates a phenotype associated with a disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising:

(a) providing a non-human transgenic animal whose genome comprises a disruption of a gene which is an ortholog of a human gene that encodes for the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide;

(b) measuring a physiological characteristic of the non-human transgenic animal of (a);

(c) comparing the measured physiological characteristic of (b) with that of a gender matched wild-type animal, wherein the physiological characteristic of the non-human transgenic animal that differs from the physiological characteristic of the wild-type animal is identified as a phenotype resulting from the gene disruption in the non-human transgenic animal;

(d) administering a test agent to the non-human transgenic animal of (a); and

(e) determining whether the test agent modulates the identified phenotype associated with gene disruption in the non-human transgenic animal.

27. The method of Claim 26, wherein the phenotype associated with the gene disruption comprises a neurological disorder; a cardiovascular, endothelial or angiogenic disorder; an eye abnormality; an immunological disorder; an oncological disorder; a bone metabolic abnormality or disorder; a lipid metabolic disorder; or a developmental abnormality.

28. The method of Claim 27, wherein the neurological disorder is an increased anxiety-like response during open field activity testing.

29. The method of Claim 27, wherein the neurological disorder is a decreased anxiety-like response during open field activity testing.

30. The method of Claim 27, wherein the neurological disorder is an abnormal circadian rhythm during home-cage activity testing.
31. The method of Claim 27, wherein the neurological disorder is an enhanced motor coordination during inverted screen testing.

32. The method of Claim 27, wherein the neurological disorder is an impaired motor coordination during inverted screen testing.

33. The method of Claim 27, wherein the neurological disorder is depression, generalized anxiety disorders, attention deficit disorder, sleep disorder, hyperactivity disorder, obsessive compulsive disorder, schizophrenia, cognitive disorders, hyperalgiesia or sensory disorders.

34. The method of Claim 27, wherein the eye abnormality is a retinal abnormality.

35. The method of Claim 27, wherein the eye abnormality is consistent with vision problems or blindness.

36. The method of Claim 34, wherein the retinal abnormality is consistent with retinitis pigmentosa.

37. The method of Claim 34, wherein the retinal abnormality is characterized by retinal degeneration or retinal dysplasia.

38. The method of Claim 34, wherein the retinal abnormality is consistent with retinal dysplasia, various retinopathies, including retinopathy of prematurity, retrolental fibroplasia, neovascular glaucoma, age-related macular degeneration, diabetic macular edema, corneal neovascularization, corneal graft neovascularization, corneal graft rejection, retinal/choroidal neovascularization, neovascularization of the angle (rubeosis), ocular neovascular disease, vascular restenosis, arteriovenous malformations (AVM), meningioma, hemangioma, angiofibroma, thyroid hyperplasias (including Grave's disease), corneal and other tissue transplantation, retinal artery obstruction or occlusion; retinal degeneration causing secondary atrophy of the retinal vasculature, retinitis pigmentosa, macular dystrophies, Stargardt’s disease, congenital stationary night blindness, choroideremia, gyrate atrophy, Leber’s congenital amaurosis, retinoschisis disorders, Wagner’s syndrome, Usher syndromes, Zellweger syndrome, Saldino-Mainzer syndrome, Senior-Loken syndrome, Bardet-Biedl syndrome, Alport's syndrome, Alstrom's syndrome, Cockayne's syndrome, dysplasia spondyloepiphysaria congenita, Flynn-Aird syndrome, Friedreich ataxia, Hallgren syndrome, Marshall syndrome, Albers-Schonberg disease, Refsum's disease, Kearns-Sayre syndrome, Waardenburg's syndrome, Alagile syndrome, myotonic dystrophy, olivopontocerebellar atrophy, Pierre-Marie dunsdrome, Stickler syndrome, carotinemia, cystinosis, Wolfman syndrome, Bassen-Kornzweig syndrome, abetalipoproteinemia, incontinentia pigmenti, Batten's disease, mucopolysaccharidosis, homocystinuria, or mannosidosis.

39. The method of Claim 27, wherein the eye abnormality is a cataract.

40. The method of Claim 39, wherein the cataract is consistent with systemic diseases such as human
41. The method of Claim 27, wherein the developmental abnormality comprises embryonic lethality or reduced viability.

42. The method of Claim 27, wherein the cardiovascular, endothelial or angiogenic disorders are arterial diseases, such as diabetes mellitus; papilledema; optic atrophy; atherosclerosis; angina; myocardial infarctions such as acute myocardial infarctions, cardiac hypertrophy, and heart failure such as congestive heart failure; hypertension; inflammatory vasculitides; Reynaud's disease and Reynaud's phenomenon; aneurysms and arterial restenosis; venous and lymphatic disorders such as thrombophlebitis, lymphangitis, and lymphedema; peripheral vascular disease; cancer such as vascular tumors, e.g., hemangioma (capillary and cavernous), glomus tumors, telangiectasia, bacillary angiomatosis, hemangiendothelioma, angiosarcoma, haemangiopericytoma, Kaposi's sarcoma, lymphangioma, and lymphangiosarcoma; tumor angiogenesis; trauma such as wounds, burns, and other injured tissue, implant fixation, scarring; ischemia reperfusion injury; rheumatoid arthritis; cerebrovascular disease; renal diseases such as acute renal failure, or osteoporosis.

43. The method of Claim 27, wherein the immunological disorders are systemic lupus erythematosus; rheumatoid arthritis; juvenile chronic arthritis; spondyloarthopathies; systemic sclerosis (scleroderma); idiopathic inflammatory myopathies (dermatomyositis, polymyositis); Sjögren's syndrome; systemic vasculitis; sarcoidosis; autoimmune hemolytic anemia (immune pancytopenia, paroxysmal nocturnal hemoglobinuria); autoimmune thrombocytopenia (idiopathic thrombocytopenic purpura, immune-mediated thrombocytopenia); thyroiditis (Grave's disease, Hashimoto's thyroiditis, juvenile lymphocytic thyroiditis, atrophic thyroiditis); diabetes mellitus; immune-mediated renal disease (glomerulonephritis, tubulointerstitial nephritis); demyelinating diseases of the central and peripheral nervous systems such as multiple sclerosis, idiopathic demyelinating polyneuropathy or Guillain-Barré syndrome, and chronic inflammatory demyelinating polyneuropathy; hepatobiliary diseases such as infectious hepatitis (hepatitis A, B, C, D, E and other non-hepatotropic viruses), autoimmune chronic active hepatitis, primary biliary cirrhosis, granulomatous hepatitis, and sclerosing cholangitis; inflammatory bowel disease (ulcerative colitis; Crohn's disease); gluten-sensitive enteropathy, and Whipple's disease; autoimmune or immune-mediated skin diseases including bullous skin diseases, erythema multiforme and contact dermatitis, psoriasis; allergic diseases such as asthma, allergic rhinitis, atopic dermatitis, food hypersensitivity and urticaria; immunologic diseases of the lung such as eosinophilic pneumonia, idiopathic pulmonary fibrosis and hypersensitivity pneumonitis; or transplantation-associated diseases including graft rejection and graft-versus-host disease.

44. The method of Claim 27, wherein said bone metabolic abnormality or disorder is arthritis, osteoporosis or osteopetrosis.
45. The method of Claim 26, wherein the non-human transgenic animal exhibits at least one of the following physiological characteristics compared with gender matched wild-type littermates: increased anxiety-like response during open field testing; hyperactivity during open field testing; decreased anxiety during open field testing; decreased locomotor activity during open field testing; abnormal circadian rhythm during home-cage activity testing (low activity during the light phase; altered sleep/wake cycle); abnormal circadian rhythm during home-cage activity testing including decreased ambulatory counts; hypoactivity with no circadian rhythm; abnormal circadian rhythm during home-cage activity testing including increased ambulatory counts; decreased rearing; increased sensitivity to stress induced hyperthermia (increased anxiety); impaired motor coordination during inverted screen testing; head tilt and retropulsion; increased prepulse inhibition response indicating enhanced sensorimotor gating/attention; decreased startle response during prepulse inhibition testing; no startle response indicating deafness or impaired hearing; decreased prepulse inhibition with impaired sensorimotor gating/attention; increased latency to respond in hot plate testing; decreased latency to respond in hot plate testing; ophthalmological abnormalities; impaired vision; white deposits of optic disc region; ocular infection and neutrophilia; bilateral optic disc lesion; decreased tear production; decreased heart rate; increased mean systolic blood pressure; decreased mean systolic blood pressure; increased mean fasting serum glucose levels; decreased mean serum glucose levels; increased mean serum cholesterol levels; decreased mean serum cholesterol levels; increased mean serum triglyceride levels; decreased mean serum triglyceride levels; increased serum albumin, alanine amino transferase and phosphorus levels; increased mean serum alkaline phosphatase levels; urinary nitrates present; increased total white blood cell (WBC) count; decreased total white blood cell (WBC) count and absolute neutrophil count; increased mean absolute neutrophil count; increased mean absolute lymphocyte count; increased mean platelet count; increased mean red cell distribution width; decreased mean platelet count; reduced percentage of CD4 spleen thymocytes; decreased percentages of CD4 cells in the periphery resulting in increased percentages of B cells in lymph organs; CD4 cells exhibit a more activated/memory phenotype (CD62Llow, CD44hi); developmental defect in CD4+ cells; decreased percentages of CD4 cells and increased percentages of B cells in blood; decreased percentages of CD4 cells and increased percentages of B cells in tissues; increase in percentages of B cells in Peyer’s patches; decreased germinal center, isotype-switched B cells in Peyer’s patches (CD38low; IgM negative); decreased CD23 intensity in spleen; increased mean percentages of B220 Med/CD23- cells and B220+/CD11b-Low/CD23- cells in peritoneal lavage; increased mean percentages of B cells in peripheral blood; decreased CD4 and CD8 T cells and increased B cells; increase in peritoneal B cells; reduction in CD11b-Hi cells in peritoneal cavity; decreased mean CD4 to CD8 ratio in spleen; decreased CD8 cells; decreased mean percentages of B220+/CD23+ cells and B220+/CD11bLow/CD23- cells in peritoneal lavage; increased mean serum IgG1 response to ovalbumin challenge; increased mean serum IgG2a response to ovalbumin challenge; increased mean serum IL-6 response to LPS challenge; increased mean serum TNF alpha response to LPS challenge; increased mean serum MCP-1 response to LPS challenge; increased mean serum IgM level; increased mean serum IgA; increased mean serum IgG1; increased mean serum IgG2a; increased mean serum IgG2b; decreased mean serum IgG1 response to ovalbumin challenge; decreased mean serum IgG2a response to ovalbumin challenge; failure in ovalbumin response; decreased mean serum IgA level; decreased mean serum IgG2a level; decreased skin fibroblast proliferation rate; increased mean percent of total body fat and total fat mass; increased mean body weight; increased mean body length; increased total tissue mass (TTM); increased
bone mineral density (BMD); increase in bone mineral content (BMC); increased mean femoral midshaft cortical thickness; decreased mean percent of total body fat and total fat mass; decreased mean body weight; decreased mean body length; decreased mean body weight and length in heterozygotes; decreased total tissue mass (TTM); decreased lean body mass (LBM); decreased femoral bone mineral density (BMD); decreased vertebral bone mineral density (BMD); decreased bone mineral density (BMD) in total body; decreased bone mineral content (BMC); decreased bone mineral density index; decreased volumetric bone mineral density (vBMD); decreased mean femoral midshaft cortical thickness; decreased mean femoral midshaft cross-sectional area; decreased mean vertebral trabecular bone volume, number and connectivity density; osteopetrosis; osteoporosis; moderate kidney hydronephrosis; hydrocephalus; enlarged liver; induced in activated T cells; induced in activated NK cells and dendritic cells; myeloid B cell expression; hyperplasia of sebaceous glands and multifocal hyperplasia of the epidermis (acanthosis and hyperkeratosis); moderate dermatitis; increased extramedullary hematopoiesis in liver and spleen; myeloid hyperplasia of the bone marrow; encephalitis due to Group B streptococcus; meningitis due to E. Coli infection; lymphocytic infiltrates in salivary glands, pancreas and lungs; poor breeders requiring foster mothers; decreased litter size; homozygous mice were small and dehydrated; vacuolar degeneration of testes resulting in decreased sperm production and infertility; defective spermatogenesis in the testes; hypospermatia and defective spermatozoa in the epididymus; male infertility; decreased testes weight; growth retardation; small mice and failure to thrive; reduced viability; reduced viability with situs inversus; and homozygous embryonic lethality.

46. An agent identified by the method of Claim 26.

47. The agent of Claim 46 which is an agonist or antagonist of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide.

48. The agent of Claim 47, wherein the agonist is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

49. The agent of Claim 47, wherein the antagonist is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-
PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9907, anti-PRO9907, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO10084, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

50. A method of identifying an agent that modulates a physiological characteristic associated with a disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO9907, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising:

(a) providing a non-human transgenic animal whose genome comprises a disruption of a gene which is an ortholog of a human gene that encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide;

(b) measuring a physiological characteristic exhibited by the non-human transgenic animal of (a);

(c) comparing the measured physiological characteristic of (b) with that of a gender matched wild-type animal, wherein the physiological characteristic exhibited by the non-human transgenic animal that differs from the physiological characteristic exhibited by the wild-type animal is identified as a physiological characteristic associated with gene disruption;

(d) administering a test agent to the non-human transgenic animal of (a); and

(e) determining whether the physiological characteristic associated with gene disruption is modulated.

51. The method of Claim 50, wherein the non-human transgenic animal exhibits at least one of the following physiological characteristics compared with gender matched wild-type littermates: increased anxiety-like response during open field testing; hyperactivity during open field testing; decreased anxiety during open field testing; decreased locomotor activity during open field testing; abnormal circadian rhythm during home-cage activity testing (low activity during the light phase; altered sleep/wake cycle); abnormal circadian rhythm during home-cage activity testing including decreased ambulatory counts; hypoactivity with no circadian rhythm; abnormal circadian rhythm during home-cage activity testing including increased ambulatory counts; decreased rearing; increased sensitivity to stress induced hyperthermia (increased anxiety); impaired motor coordination during inverted screen testing; head tilt and retropulsion; increased prepulse inhibition response indicating enhanced sensorimotor gating/attention; decreased startle response during prepulse inhibition testing; no startle response indicating
deafness or impaired hearing; decreased prepulse inhibition with impaired sensorimotor gating/attention; increased latency to respond in hot plate testing; decreased latency to respond in hot plate testing; opthalmological abnormalities; impaired vision; white deposits of optic disc region; ocular infection and neutrophilia; bilateral optic disc lesion; decreased tear production; decreased heart rate; increased mean systolic blood pressure; decreased mean systolic blood pressure; increased mean fasting serum glucose levels; decreased mean serum glucose levels; increased mean serum cholesterol levels; decreased mean serum cholesterol levels; increased mean serum triglyceride levels; decreased mean serum triglyceride levels; impaired glucose tolerance; increased mean serum albumin, alanine amino transferase and phosphorus levels; increased mean serum alkaline phosphatase levels; urinary nitrites present; increased total white blood cell (WBC) count; decreased total white blood cell (WBC) count and absolute neutrophil count; increased mean absolute neutrophil count; increased mean absolute lymphocyte count; increased mean platelet count; increased mean red cell distribution width; decreased mean platelet count; reduced percentage of CD4 spleen thymocytes; decreased percentages of CD4 cells in the periphery resulting in increased percentages of B cells in lymph organs; CD4 cells exhibit a more activated/memory phenotype (CD62Llow, CD44hi); developmental defect in CD4+ cells; decreased percentages of CD4 cells and increased percentages of B cells in blood; decreased percentages of CD4 cells and increased percentages of B cells in tissues; increase in percentages of B cells in Peyer’s patches;; decreased germinal center, isotype-switched B cells in Peyer’s patches (CD38low; IgM negative); decreased CD23 intensity in spleen; increased mean percentages of B220 Med/CD23- cells and B220+/CD11b-Low/CD23- cells in peritoneal lavage; increased mean percentages of B cells in peripheral blood; decreased CD4 and CD8 T cells and increased B cells; increase in peritoneal B cells; reduction in CD11b-Hi cells in peritoneal cavity; decreased mean CD4 to CD8 ratio in spleen; decreased CD8 cells; decreased mean percentages of B220+/CD23+ cells and B220+/CD11b-Low/CD23- cells in peritoneal lavage; increased mean serum IgG1 response to ovalbumin challenge; increased mean serum IgG2a response to ovalbumin challenge; increased mean serum IL-6 response to LPS challenge; increased mean serum TNF alpha response to LPS challenge; increased mean serum MCP-1 response to LPS challenge; increased mean serum IgM level; increased mean serum IgA; increase mean serum IgG1; increased mean serum IgG2a; increased mean serum IgG2b; decreased mean serum IgG1 response to ovalbumin challenge; decreased mean serum IgG2a response to ovalbumin challenge; failure in ovalbumin response; decreased mean serum IgA level; decreased mean serum IgG2a level; decreased skin fibroblast proliferation rate; increased mean percent of total body fat and total fat mass; increased mean body weight; increased mean body length; increased total tissue mass (TTM); increased bone mineral density (BMD); increase in bone mineral content (BMC); increased mean femoral midshaft cortical thickness; decreased mean percent of total body fat and total fat mass; decreased mean body weight; decreased mean body length; decreased mean body weight and length in heterozygotes; decreased total tissue mass (TTM); decreased lean body mass (LBM); decreased femoral bone mineral density (BMD); decreased vertebral bone mineral density (BMD); decreased bone mineral density (BMD) in total body; decreased bone mineral content (BMC); decreased bone mineral density index; decreased volumetric bone mineral density (vBMD); decreased mean femoral midshaft cortical thickness; decreased mean femoral midshaft cross-sectional area; decreased mean vertebral trabecular bone volume, number and connectivity density; osteopetrosis; osteoporosis; moderate kidney hydropnephrosis; hydrocephalus; enlarged liver; induced in activated T cells; induced in activated NK cells and dendritic cells; myeloid B cell expression; hyperplasia of sebaceous glands and multifocal hyperplasia of the
epidermis (acanthosis and hyperkeratosis); moderate dermatitis; increased extramedullary hematopoiesis in liver and spleen; myeloid hyperplasia of the bone marrow; encephalitis due to Group B streptococcus; meningitis due to *E. Coli* infection; lymphocytic infiltrates in salivary glands, pancreas and lungs; poor breeders requiring foster mothers; decreased litter size; homozygous mice were small and dehydrated; vacuolar degeneration of testes resulting in decreased sperm production and infertility; defective spermatogenesis in the testes; hypospermaia and defective spermatozoa in the epididymus; male infertility; decreased testes weight; growth retardation; small mice and failure to thrive; reduced viability; reduced viability with situs inversus; and homozygous embryonic lethality.

52. An agent identified by the method of Claim 50.

53. The agent of Claim 52 which is an agonist or antagonist of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide.

54. The agent of Claim 53, wherein the agonist is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

55. The agent of Claim 53, wherein the antagonist is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

56. A method of identifying an agent which modulates a behavior associated with a disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238,
PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising:

(a) providing a non-human transgenic animal whose genome comprises a disruption of a gene which is an ortholog of a human gene that encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide;

(b) observing the behavior exhibited by the non-human transgenic animal of (a);

(c) comparing the observed behavior of (b) with that of a gender matched wild-type animal, wherein the observed behavior exhibited by the non-human transgenic animal that differs from the observed behavior exhibited by the wild-type animal is identified as a behavior associated with gene disruption;

(d) administering a test agent to the non-human transgenic animal of (a); and

(e) determining whether the agent modulates the behavior associated with gene disruption.

57. The method of Claim 56, wherein the behavior is an increased anxiety-like response during open field activity testing.

58. The method of Claim 56, wherein the behavior is a decreased anxiety-like response during open field activity testing.

59. The method of Claim 56, wherein the behavior is an abnormal circadian rhythm during home-cage activity testing.

60. The method of Claim 56, wherein the behavior is an enhanced motor coordination during inverted screen testing.

61. The method of Claim 56, wherein the behavior is an impaired motor coordination during inverted screen testing.

62. The method of Claim 56, wherein the behavior is depression, generalized anxiety disorders, attention deficit disorder, sleep disorder, hyperactivity disorder, obsessive compulsive disorder, schizophrenia, cognitive disorders, hyperalgesia or sensory disorders.

63. An agent identified by the method of Claim 56.
64. The agent of Claim 63 which is an agonist or antagonist of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide.

65. The agent of Claim 64, wherein the agonist is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

66. The agent of Claim 64, wherein the antagonist is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

67. A method of identifying an agent that ameliorates or modulates a neurological disorder; a cardiovascular, endothelial or angiogenic disorder; an eye abnormality; an immunological disorder; an oncological disorder; a bone metabolic abnormality or disorder; a lipid metabolic disorder; or a developmental abnormality associated with a disruption in a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising:

(a) providing a non-human transgenic animal whose genome comprises a disruption of a gene which is an ortholog of a human gene that encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865,
PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide;

(b) administering a test agent to said non-human transgenic animal; and

(c) determining whether said test agent ameliorates or modulates the neurological disorder; cardiovascular, endothelial or angiogenic disorder; eye abnormality; immunological disorder; oncological disorder; bone metabolic abnormality or disorder; lipid metabolic disorder; or developmental abnormality in the non-human transgenic animal.

68. The method of Claim 67, wherein the neurological disorder is an increased anxiety-like response during open field activity testing.

69. The method of Claim 67, wherein the neurological disorder is a decreased anxiety-like response during open field activity testing.

70. The method of Claim 67, wherein the neurological disorder is an abnormal circadian rhythm during home-cage activity testing.

71. The method of Claim 67, wherein the neurological disorder is an enhanced motor coordination during inverted screen testing.

72. The method of Claim 67, wherein the neurological disorder is an impaired motor coordination during inverted screen testing.

73. The method of Claim 67, wherein the neurological disorder is depression, generalized anxiety disorders, attention deficit disorder, sleep disorder, hyperactivity disorder, obsessive compulsive disorder, schizophrenia, cognitive disorders, hyperalgesia or sensory disorders.

74. The method of Claim 67, wherein the eye abnormality is a retinal abnormality.

75. The method of Claim 67, wherein the eye abnormality is consistent with vision problems or blindness.

76. The method of Claim 74, wherein the retinal abnormality is consistent with retinitis pigmentosa.

77. The method of Claim 74, wherein the retinal abnormality is characterized by retinal degeneration or retinal dysplasia.

78. The method of Claim 74, wherein the retinal abnormality is consistent with retinal dysplasia, various retinopathies, including retinopathy of prematurity, retrolental fibroplasia, neovascular glaucoma, age-related

79. The method of Claim 67, wherein the eye abnormality is a cataract.

80. The method of Claim 79, wherein the cataract is a systemic disease such as human Down's syndrome, Hallerman-Streiff syndrome, Lowe syndrome, galactosemia, Marfan syndrome, Trisomy 13-15, Alport syndrome, myotonic dystrophy, Fabry disease, hypoparathyroidism or Conradi syndrome.

81. The method of Claim 67, wherein the developmental abnormality comprises embryonic lethality or reduced viability.

82. The method of Claim 67, wherein the cardiovascular, endothelial or angiogenic disorders are arterial diseases, such as diabetes mellitus; papilledema; optic atrophy; atherosclerosis; angina; myocardial infarctions such as acute myocardial infarctions, cardiac hypertrophy, and heart failure such as congestive heart failure; hypertension; inflammatory vasculitides; Reynaud's disease and Reynaud's phenomenon; aneurysms and arterial restenosis; venous and lymphatic disorders such as thrombophlebitis, lymphangitis, and lymphedema; peripheral vascular disease; cancer such as vascular tumors, e.g., hemangioma (capillary and cavernous), glomus tumors, telangiectasia, bacillary angiomatosis, hemangiodendothelioma, angiosarcoma, haemangiopericytoma, Kaposi's sarcoma, lymphangioma, and lymphangiosarcoma; tumor angiogenesis; trauma such as wounds, burns, and other injured tissue, implant fixation, scarring; ischemia reperfusion injury; rheumatoid arthritis; cerebrovascular disease; renal diseases such as acute renal failure, or osteoporosis.

83. The method of Claim 67, wherein the immunological disorders are systemic lupus erythematosus; rheumatoid arthritis; juvenile chronic arthritis; spondyloarthropathies; systemic sclerosis (scleroderma); idiopathic inflammatory myopathies (dermatomyositis, polymyositis); Sjögren's syndrome; systemic vasculitis; sarcoidosis; autoimmune hemolytic anemia (immune pancytopenia, paroxysmal nocturnal hemoglobinuria); autoimmune
thrombocytopenia (idiopathic thrombocytopenic purpura, immune-mediated thrombocytopenia); thyroiditis (Grave's disease, Hashimoto's thyroiditis, juvenile lymphocytic thyroiditis, atrophic thyroiditis); diabetes mellitus; immune-mediated renal disease (glomerulonephritis, tubulointerstitial nephritis); demyelinating diseases of the central and peripheral nervous systems such as multiple sclerosis, idiopathic demyelinating polyneuropathy or Guillain-Barré syndrome, and chronic inflammatory demyelinating polyneuropathy; hepatobiliary diseases such as infectious hepatitis (hepatitis A, B, C, D, E and other non-hepatotropic viruses), autoimmune chronic active hepatitis, primary biliary cirrhosis, granulomatous hepatitis, and sclerosing cholangitis; inflammatory bowel disease (ulcerative colitis; Crohn's disease); gluten-sensitive enteropathy, and Whipple's disease; autoimmune or immune-mediated skin diseases including bullous skin diseases, erythema multiforme and contact dermatitis, psoriasis; allergic diseases such as asthma, allergic rhinitis, atopic dermatitis, food hypersensitivity and urticaria; immunologic diseases of the lung such as eosinophilic pneumonia, idiopathic pulmonary fibrosis and hypersensitivity pneumonitis; or transplantation associated diseases including graft rejection and graft-versus-host disease.

84. The method of Claim 67, wherein said bone metabolic abnormality or disorder is arthritis, osteoporosis or osteopetrosis.

85. The method of Claim 67, wherein the non-human transgenic animal exhibits at least one of the following physiological characteristics compared with gender matched wild-type littermates: increased anxiety-like response during open field testing; hyperactivity during open field testing; decreased anxiety during open field testing; decreased locomotor activity during open field testing; abnormal circadian rhythm during home-cage activity testing (low activity during the light phase; altered sleep/wake cycle); abnormal circadian rhythm during home-cage activity testing including decreased ambulatory counts; hypoactivity with no circadian rhythm; abnormal circadian rhythm during home-cage activity testing including increased ambulatory counts; decreased rearing; increased sensitivity to stress induced hyperthermia (increased anxiety); impaired motor coordination during inverted screen testing; head tilt and retropulsion; increased prepulse inhibition response indicating enhanced sensorimotor gating/attention; decreased startle response during prepulse inhibition testing; no startle response indicating deafness or impaired hearing; decreased prepulse inhibition with impaired sensorimotor gating/attention; increased latency to respond in hot plate testing; decreased latency to respond in hot plate testing; ophthalmological abnormalities; impaired vision; white deposits of optic disc region; ocular infection and neutrophilia; bilateral optic disc lesion; decreased tear production; decreased heart rate; increased mean systolic blood pressure; decreased mean systolic blood pressure; increased mean fasting serum glucose levels; decreased mean serum glucose levels; increased mean serum cholesterol levels; decreased mean serum cholesterol levels; increased mean serum triglyceride levels; decreased mean serum triglyceride levels; impaired glucose tolerance; increased mean serum albumin, alanine amino transferase and phosphorous levels; increased mean serum alkaline phosphatase levels; urinary nitrates present; increased total white blood cell (WBC) count; decreased total white blood cell (WBC) count and absolute neutrophil count; increased mean absolute neutrophil count; increased mean absolute lymphocyte count; increased mean platelet count; increased mean red cell distribution width; decreased mean platelet count; reduced percentage of CD4 spleen thymocytes; decreased percentages of CD4 cells in the periphery...
resulting in increased percentages of B cells in lymph organs; CD4 cells exhibit a more activated/memory phenotype (CD62Llow, CD44hi); developmental defect in CD4+ cells; decreased percentages of CD4 cells and increased percentages of B cells in blood; decreased percentages of CD4 cells and increased percentages of B cells in tissues; increase in percentages of B cells in Peyer’s patches; decreased germinal center, isotype-switched B cells in Peyer’s patches (CD38low; IgM negative); decreased CD23 intensity in spleen; increased mean percentages of B220Med/CD23- cells and B220+/CD11b-Low/CD23- cells in peritoneal lavage; increased mean percentages of B cells in peripheral blood; decreased CD4 and CD8 T cells and increased B cells; increase in peritoneal B cells; reduction in CD11b-Hi cells in peritoneal cavity; decreased mean CD4 to CD8 ratio in spleen; decreased CD8 cells; decreased mean percentages of B220+/CD23+ cells and B220+/CD11bLow/CD23- cells in peritoneal lavage; increased mean serum IgG1 response to ovalbumin challenge; increased mean serum IgG2a response to ovalbumin challenge; increased mean serum IL-6 response to LPS challenge; increased mean serum TNF alpha response to LPS challenge; increased mean serum MCP-1 response to LPS challenge; increased mean serum IgM level; increased mean serum IgA; increase mean serum IgG1; increased mean serum IgG2a; increased mean serum IgG2b; decreased mean serum IgG1 response to ovalbumin challenge; decreased mean serum IgG2a response to ovalbumin challenge; failure in ovalbumin response; decreased mean serum IgA level; decreased mean serum IgG2a level; decreased skin fibroblast proliferation rate; increased mean percent of total body fat and total fat mass; increased mean body weight; increased mean body length; increased total tissue mass (TTM); increased bone mineral density (BMD); increase in bone mineral content (BMC); increased mean femoral midshaft cortical thickness; decreased mean percent of total body fat and total fat mass; decreased mean body weight; decreased mean body length; decreased mean body weight and length in heterozygotes; decreased total tissue mass (TTM); decreased lean body mass (LBM); decreased femoral bone mineral density (BMD); decreased vertebral bone mineral density (BMD); decreased bone mineral density (BMD) in total body; decreased bone mineral content (BMC); decreased bone mineral density index; decreased volumetric bone mineral density (vBMD); decreased mean femoral midshaft cortical thickness; decreased mean femoral midshaft cross-sectional area; decreased mean vertebral trabecular bone volume, number and connectivity density; osteopetrosis; osteoporosis; moderate kidney hydronephrosis; hydrocephalus; enlarged liver; induced in activated T cells; induced in activated NK cells and dendritic cells; myeloid B cell expression; hyperplasia of sebaceous glands and multifocal hyperplasia of the epidermis (acanthosis and hyperkeratosis); moderate dermatitis; increased extramedullary hematopoiesis in liver and spleen; myeloid hyperplasia of the bone marrow; encephalitis due to Group B streptococcus; meningitis due to *E. Coli* infection; lymphocytic infiltrates in salivary glands, pancreas and lungs; poor breeders requiring foster mothers; decreased litter size; homozygous mice were small and dehydrated; vacuolar degeneration of testes resulting in decreased sperm production and infertility; defective spermatogenesis in the testes; hypospermidia and defective spermatozoa in the epididymus; male infertility; decreased testes weight; growth retardation; small mice and failure to thrive; reduced viability; reduced viability with situs inversus; and homozygous embryonic lethality.

86. An agent identified by the method of Claim 67.

87. The agent of Claim 86 which is an agonist or antagonist of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941,
PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide.

88. The agent of Claim 87, wherein the agonist is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 polypeptide.

89. The agent of Claim 87, wherein the antagonist is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 polypeptide.

90. A therapeutic agent identified by the method of Claim 67.

91. A method of identifying an agent that modulates the expression of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1008, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising:

(a) contacting a test agent with a host cell expressing a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide; and

(b) determining whether the test agent modulates the expression of the PRO218, PRO228, PRO271,
PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide by the host cell.

92. An agent identified by the method of Claim 91.

93. The agent of Claim 92 which is an agonist or antagonist of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide.

94. The agent of Claim 93, wherein the agonist is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

95. The agent of Claim 93, wherein the antagonist is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

96. A method of evaluating a therapeutic agent capable of affecting a condition associated with a disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013,
PRO09048, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising:

(a) providing a non-human transgenic animal whose genome comprises a disruption of a gene which is an ortholog of a human gene that encodes for the PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide;

(b) measuring a physiological characteristic of the non-human transgenic animal of (a);

(c) comparing the measured physiological characteristic of (b) with that of a gender matched wild-type animal, wherein the physiological characteristic of the non-human transgenic animal that differs from the physiological characteristic of the wild-type animal is identified as a condition resulting from the gene disruption in the non-human transgenic animal;

(d) administering a test agent to the non-human transgenic animal of (a); and

(e) evaluating the effects of the test agent on the identified condition associated with gene disruption in the non-human transgenic animal.

97. The method of Claim 96, wherein the condition is a neurological disorder; a cardiovascular, endothelial or angiogenic disorder; an eye abnormality; an immunological disorder; an oncological disorder; a bone metabolic abnormality or disorder; a lipid metabolic disorder; or a developmental abnormality.

98. A therapeutic agent identified by the method of Claim 96.

99. The therapeutic agent of Claim 98 which is an agonist or antagonist of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide.

100. The therapeutic agent of Claim 99, wherein the agonist is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.
101. The therapeutic agent of Claim 99, wherein the antagonist is an anti-PRO218, anti-PRO228, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1135, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

102. A pharmaceutical composition comprising the therapeutic agent of Claim 98.

103. A method of treating or preventing or ameliorating a neurological disorder; cardiovascular, endothelial or angiogenic disorder; immunological disorder; oncological disorder; bone metabolic abnormality or disorder, or embryonic lethality associated with the disruption of a gene which encodes for a PRO218, PRO228, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1135, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising administering to a subject in need of such treatment whom may already have the disorder, or may be prone to have the disorder or may be in whom the disorder is to be prevented, a therapeutically effective amount of the therapeutic agent of Claim 90, or agonists or antagonists thereof, thereby effectively treating or preventing or ameliorating said disorder.

104. The method of Claim 103, wherein the neurological disorder is an increased anxiety-like response during open field activity testing.

105. The method of Claim 103, wherein the neurological disorder is a decreased anxiety-like response during open field activity testing.

106. The method of Claim 103, wherein the neurological disorder is an abnormal circadian rhythm during home-cage activity testing.

107. The method of Claim 103, wherein the neurological disorder is an enhanced motor coordination during inverted screen testing.

108. The method of Claim 103, wherein the neurological disorder is an impaired motor coordination during inverted screen testing.
109. The method of Claim 103, wherein the neurological disorder is depression, generalized anxiety disorders, attention deficit disorder, sleep disorder, hyperactivity disorder, obsessive compulsive disorder, schizophrenia, cognitive disorders, hyperalgesia or sensory disorders.

110. The method of Claim 103, wherein the eye abnormality is a retinal abnormality.

111. The method of Claim 103, wherein the eye abnormality is consistent with vision problems or blindness.

112. The method of Claim 110, wherein the retinal abnormality is consistent with retinitis pigmentosa.

113. The method of Claim 110, wherein the retinal abnormality is characterized by retinal degeneration or retinal dysplasia.

114. The method of Claim 110, wherein the retinal abnormality is consistent with retinal dysplasia, various retinopathies, including retinopathy of prematurity, retrolental fibroplasia, neovascular glaucoma, age-related macular degeneration, diabetic macular edema, corneal neovascularization, corneal graft neovascularization, corneal graft rejection, retinal/choroidal neovascularization, neovascularization of the angle (rubeosis), ocular neovascular disease, vascular restenosis, arteriovenous malformations (AVM), meningioma, hemangioma, angiofibroma, thyroid hyperplasias (including Grave's disease), corneal and other tissue transplantation, retinal artery obstruction or occlusion; retinal degeneration causing secondary atrophy of the retinal vasculature, retinitis pigmentosa, macular dystrophies, Stargardt's disease, congenital stationary night blindness, choroideremia, gyrate atrophy, Leber's congenital amaurosis, retinoschisis disorders, Wagner's syndrome, Usher syndromes, Zellweger syndrome, Saldino-Mainzer syndrome, Senior-Loken syndrome, Bardet-Biedl syndrome, Alport's syndrome, Alstrom's syndrome, Cockayne's syndrome, dysplasia spondyloepiphysaria congenita, Flynn-Aird syndrome, Friedreich ataxia, Hallgren syndrome, Marshall syndrome, Albers-Schönberg disease, Reisn'm's disease, Kearns-Sayre syndrome, Waardenburg's syndrome, Alagille syndrome, myotonic dystrophy, olivopontocerebellar atrophy, Pierre-Marie-Dundrino, Stickler syndrome, carotinemia, cystinosis, Wolfram syndrome, Bassen-Kornzweig syndrome, abetalipoproteinemia, incontinentia pigmenti, Batten's disease, mucopolysaccharidoses, homocystinuria, or mamonidosis.

115. The method of Claim 103, wherein the eye abnormality is a cataract.

116. The method of Claim 115, wherein the cataract is a systemic disease such as human Down's syndrome, Hallerman-Streiff syndrome, Lowe syndrome, galactosemia, Marfan syndrome, Trisomy 13-15, Alport syndrome, myotonic dystrophy, Fabry disease, hypoparathyroidism or Conradi syndrome.

117. The method of Claim 103, wherein the developmental abnormality comprises embryonic lethality or reduced viability.
118. The method of Claim 103, wherein the cardiovascular, endothelial or angiogenic disorders are arterial diseases, such as diabetes mellitus; papilledema; optic atrophy; atherosclerosis; angina; myocardial infarctions such as acute myocardial infarctions, cardiac hypertrophy, and heart failure such as congestive heart failure; hypertension; inflammatory vasculitides; Reynaud's disease and Reynaud's phenomenon; aneurysms and arterial restenosis; venous and lymphatic disorders such as thrombophlebitis, lymphangitis, and lymphedema; peripheral vascular disease; cancer such as vascular tumors, e.g., hemangiomata (capillary and cavernous), glomus tumors, telangiectasia, bacillary angiomatosis, hemangioendothelioma, angiosarcoma, haemangiopericytoma, Kaposi's sarcoma, lymphangioma, and lymphangiosarcoma; tumor angiogenesis; trauma such as wounds, burns, and other injured tissue, implant fixation, scarring; ischemia reperfusion injury; rheumatoid arthritis; cerebrovascular disease; renal diseases such as acute renal failure, or osteoporosis.

119. The method of Claim 103, wherein the immunological disorders are systemic lupus erythematosus; rheumatoid arthritis; juvenile chronic arthritis; spondyloarthropathies; systemic sclerosis (scleroderma); idiopathic inflammatory myopathies (dermatomyositis, polymyositis); Sjögren's syndrome; systemic vasculitis; sarcoidosis; autoimmune hemolytic anemia (immune pancytopenia, paroxysmal nocturnal hemoglobinuria); autoimmune thrombocytopenia (idiopathic thrombocytopenic purpura, immune-mediated thrombocytopenia); thyroiditis (Grave's disease, Hashimoto's thyroiditis, juvenile lymphocytic thyroiditis, atrophic thyroiditis); diabetes mellitus; immune-mediated renal disease (glomerulonephritis, tubulointerstitial nephritis); demyelinating diseases of the central and peripheral nervous systems such as multiple sclerosis, idiopathic demyelinating polyneuropathy or Guillain-Barré syndrome, and chronic inflammatory demyelinating polyneuropathy; hepatobiliary diseases such as infectious hepatitis (hepatitis A, B, C, D, E and other non-hepatotropic viruses), autoimmune chronic active hepatitis, primary biliary cirrhosis, granulomatous hepatitis, and sclerosing cholangitis; inflammatory bowel disease (ulcerative colitis; Crohn's disease); gluten-sensitive enteropathy, and Whipple's disease; autoimmune or immune-mediated skin diseases including bullous skin diseases, erythema multiforme and contact dermatitis, psoriasis; allergic diseases such as asthma, allergic rhinitis, atopic dermatitis, food hypersensitivity and urticaria; immunologic diseases of the lung such as eosinophilic pneumonia, idiopathic pulmonary fibrosis and hypersensitivity pneumonitis; or transplantation associated diseases including graft rejection and graft-versus-host disease.

120. The method of Claim 103, wherein said bone metabolic abnormality or disorder is arthritis, osteoporosis or osteopetrosis.

121. A method of modulating a phenotype associated with a disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3445, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising administering to a subject whom may already have the phenotype, or may be prone to have the
phenotype or may be in whom the phenotype is to be prevented, an effective amount of the agent of Claim 46, or agonists or antagonists thereof, thereby effectively modulating the phenotype.

122. A method of modulating a physiological characteristic associated with a disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising administering to a subject whom may already exhibit the physiological characteristic, or may be prone to exhibit the physiological characteristic or may be in whom the physiological characteristic is to be prevented, an effective amount of the agent of Claim 52, or agonists or antagonists thereof, thereby effectively modulating the physiological characteristic.

123. A method of modulating a behavior associated with a disruption of a gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising administering to a subject whom may already exhibit the behavior, or may be prone to exhibit the behavior or may be in whom the exhibited behavior is to be prevented, an effective amount of the agent of Claim 63, or agonists or antagonists thereof, thereby effectively modulating the behavior.

124. A method of modulating the expression of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising administering to a host cell expressing said PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, an effective amount of the agent of Claim 92, or agonists or antagonists thereof, thereby effectively modulating the expression of said polypeptide.

125. A method of modulating a condition associated with a disruption of a gene which encodes for a PRO218,
PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising administering to a subject whom may have the condition, or may be prone to have the condition or may be in whom the condition is to be prevented, a therapeutically effective amount of the therapeutic agent of Claim 98, or agonists or antagonists thereof, thereby effectively modulating the condition.

126. A method of identifying an agent that mimics a condition or phenotype associated with a disruption in a gene which encodes a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising:

(a) providing a non-human transgenic animal whose genome comprises a disruption of a gene which
is an ortholog of a human gene that encodes a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide;

(b) measuring a physiological characteristic of the non-human transgenic animal of (a);

(c) comparing the measured physiological characteristic of (b) with that of a gender matched wild-type animal, wherein the physiological characteristic of the non-human transgenic animal that differs from the physiological characteristic of the gender matched wild-type animal is identified as a condition or phenotype resulting from the gene disruption in the non-human transgenic animal;

(d) administering a test agent to said gender matched wild-type animal; and

(e) determining whether said test agent mimics the condition or phenotype initially observed in the non-human transgenic animal.

127. The method of Claim 126, wherein the condition or phenotype associated with the disruption of the gene which is an ortholog of a human gene that encodes a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide is enhanced glucose tolerance.
128. The method of Claim 126, wherein the condition or phenotype associated with the disruption of the gene which is an ortholog of a human gene that encodes a PRO218, PRO228, PRO227, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide is increased insulin sensitivity.

129. An agent identified by the method of Claim 126.

130. The agent of Claim 129 which is an antagonist of a PRO218, PRO228, PRO2271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide.

131. The agent of Claim 130, wherein the antagonist is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

132. A method of mimicking a condition or phenotype associated with a disruption of a gene which encodes a PRO218, PRO228, PRO227, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising administering to a subject in whom the condition or phenotype is to be mimicked, an effective amount of the agent of Claim 129 or an antagonist of a PRO218, PRO228, PRO2271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, thereby effectively mimicking the condition or phenotype.
133. The method of Claim 132, wherein the condition or phenotype associated with the disruption of the gene which is an ortholog of a human gene that encodes a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide is enhanced glucose tolerance.

134. The method of Claim 132, wherein the condition or phenotype associated with the disruption of the gene which is an ortholog of a human gene that encodes a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide is increased insulin sensitivity.

135. A method of evaluating a therapeutic agent capable of mimicking a condition or phenotype associated with a disruption of a gene which encodes a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising:

(a) providing a non-human transgenic animal whose genome comprises a disruption of a gene which is an ortholog of a human gene that encodes a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide;

(b) measuring a physiological characteristic of the non-human transgenic animal of (a);

(c) comparing the measured physiological characteristic of (b) with that of a gender matched wild-type animal, wherein the physiological characteristic of the non-human transgenic animal that differs from the physiological characteristic of the gender matched wild-type animal is identified as a condition or phenotype resulting from the gene disruption in the non-human transgenic animal;

(d) administering a test agent to said gender matched wild-type animal of (c); and

(e) evaluating the ability of the test agent to mimic the condition or phenotype associated with gene disruption in the non-human transgenic animal.
136. A therapeutic agent identified by the method of Claim 135.

137. The therapeutic agent of Claim 136 which is an antagonist of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide.

138. The therapeutic agent of Claim 137, wherein the antagonist is an anti-PRO218, anti-PRO228, anti-PRO271, anti-PRO273, anti-PRO295, anti-PRO302, anti-PRO305, anti-PRO326, anti-PRO386, anti-PRO655, anti-PRO162, anti-PRO788, anti-PRO792, anti-PRO940, anti-PRO941, anti-PRO1004, anti-PRO1012, anti-PRO1016, anti-PRO474, anti-PRO5238, anti-PRO1069, anti-PRO1111, anti-PRO1113, anti-PRO1130, anti-PRO1195, anti-PRO1271, anti-PRO1865, anti-PRO1879, anti-PRO3446, anti-PRO3543, anti-PRO4329, anti-PRO4352, anti-PRO5733, anti-PRO9859, anti-PRO9864, anti-PRO9904, anti-PRO9907, anti-PRO10013, anti-PRO90948, anti-PRO28694, anti-PRO16089, anti-PRO19563, anti-PRO19675, anti-PRO20084, anti-PRO21434, anti-PRO50332, anti-PRO38465 or anti-PRO346 antibody.

139. A pharmaceutical composition comprising the therapeutic agent of Claim 136.

140. A method of mimicking a condition or phenotype associated with a disruption of a gene which encodes a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising administering to a subject in whom the condition or phenotype disorder is to be mimicked, a therapeutically effective amount of the therapeutic agent of Claim 136, or an antagonist of a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305, PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, thereby effectively mimicking the condition or phenotype.

141. A method of identifying an agent that ameliorates or modulates a neurological disorder; a cardiovascular, endothelial or angiogenic disorder; an eye abnormality; an immunological disorder; an oncological disorder; a bone metabolic abnormality or disorder; a lipid metabolic disorder; or a developmental abnormality associated with a disruption in the gene which encodes for a PRO218, PRO228, PRO271, PRO273, PRO295, PRO302, PRO305,
PRO326, PRO386, PRO655, PRO162, PRO788, PRO792, PRO940, PRO941, PRO1004, PRO1012, PRO1016, PRO474, PRO5238, PRO1069, PRO1111, PRO1113, PRO1130, PRO1195, PRO1271, PRO1865, PRO1879, PRO3446, PRO3543, PRO4329, PRO4352, PRO5733, PRO9859, PRO9864, PRO9904, PRO9907, PRO10013, PRO90948, PRO28694, PRO16089, PRO19563, PRO19675, PRO20084, PRO21434, PRO50332, PRO38465 or PRO346 polypeptide, the method comprising administering to a subject whom may have the neurological disorder; cardiovascular, endothelial or angiogenic disorder; eye abnormality; immunological disorder; oncological disorder; bone metabolic abnormality or disorder; lipid metabolic disorder; or developmental abnormality, a therapeutic agent of Claim 90, or agonists or antagonists thereof, thereby ameliorating or modulating the disorder.
FIGURE 2

MSFLIDSLMITSQILFFGFLFMRQLFHYYLEFHVYQQVIIFDSVFSTCMFELIIFEILGFVLNSSSRFH
WKNLNVVLLLVFMVFFYIGYIIVSNIRLQHQRLLFSCLLVLTGMYFFWKLGDFFILSPKISLIEQILIS
RVGIVTLMALLSGFGAVNCPYTIMYFIRNVTDLALERRLQTMQISKKRMAMARRTMEQKGEVHN
KFSGFWGMKSVTTSSAGSENLTIQEQEVALEELSEQLFLETADLYATKERIEYSKTFKGKYFNFLQFYFFSIY
CVWKIFMATINIVFDVGKTDPVTGIEITVNLIGIQFDVKFWSQHISFILVGIIIVTISIRGLLHRLTTKFFYAI
SSKSSNVIVLLEAQIMGYFVSSVLLIRMSPLEYRTITIEVLGELQFNFYHRWFVIFLVSALSSILFYLA
HKQAPEKQMAP

Important features:
Signal peptide:
amino acids 1-23

Potential transmembrane domains:

N-glycosylation sites.
amino acids 67-70, 180-183 and 243-246

Eukaryotic cobalamin-binding proteins
amino acids 151-160
**FIGURE 4**

MKRLPLLVVFSTLLNCSYTQNCCTKTPCLPNAKCEIRNGIEACYNMGFSNGVTCEDDNECGNLTQSCGENANCNTNTREGYCMCPVGFRSSNQDRFITNDGTVCIEVNANCHLDNVIQAAANNKLTLKIRSIKEPVALLOEVRNVSVDLSPVDITYEIELAESSSSLGYKNNTIASKDTLSNSLTEFVKTVNFINQVFQDFTFVWKLDSVNHRRTHLTHLMTLHTEQATLRRISQSFFKTEFDTNSTDIALKVFFDSYNMKHIHPHNMGDYINIFPKRKAAYDSNSGAVAFLYYKSI

**Signal peptide:**
amino acids 1-19

**Transmembrane domain:**

**N-glycosylation site.**
amino acids 15-19, 21-25, 64-68, 74-78, 127-131, 177-181, 188-192, 249-253, 381-385, 395-399

**Glycosaminoglycan attachment site.**
amino acids 49-53

**cAMP- and cGMP-dependent protein kinase phosphorylation site.**
amino acids 360-364

**Casein kinase II phosphorylation site.**

**Tyrosine kinase phosphorylation site.**
amino acids 36-44, 669-677, 670-678

**N-myristoylation site.**
amino acids 38-44, 50-56, 52-58, 80-86, 382-388, 388-394, 434-440, 480-486, 521-527

**Aspartic acid and asparagine hydroxylation site.**
amino acids 75-87
FIGURE 5

GGACAGCTCCTCGGCCCCAGGAGCTCTAGGCGACGTGCTGGGGACGTCTGCTCTCCCTGCTGATTGGACTGAGCTGTGCTGGCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTC
FIGURE 6

MGLLLLVPLLPGSYPGLPYGNYGYYNSANDQNLGNGHKDNLNVKLKVETPEETLFYTQGASVILPCHRYPALVSPRRVVKWWKLSXNGAPEKDVLLVAIGLRHRSFDFDYQGRVHRLQDKEHDVSLQDRLQEDYRCEVIDGLEDESLVELELRGVVPPQYQSPNQRYQFNFHEGQQVAEQAAVVASFEQLFRAWEEGLQCNAGWLQDATVQYPIMLPRQPCGFPGLAPGVRSGYPARHRRLHRYDVFCFATALKGKVYLEYHPEKLTLEAREACQEDDATIAKVQFLFAWKFHGLDRCAGWADGTVRYPVHPPNCGPPEPGVRSFGFPDPQSRLGYVYCYQH

Signal sequence:
amino acids 1-17

Casein kinase II phosphorylation site.
amino acids 29-33, 53-57, 111-115, 278-282

Tyrosine kinase phosphorylation site.
amino acids 137-145

N-myristoylation site.

FIGURE 8

MSLLPRAPPVSMRLAAALLLLLLALAYTARVDGSKCKCSRKPKIRYSVDKKGKEMKPKYPHCEEKMVIITTKS
VSRYRQEHCLHPKQSTKRFIUKWNAWNEKRRYEE

Signal sequence:

amino acids 1-34
FIGURE 10

MRQRLGATLLCLLLAAAVPTAPAFAPTATSAPVKPGFALSYPQEEATLNEMFREVEELMEDT
QHKLRSAVEMEAEEAAKASSEVNLAPPSYHNETNDTKVGNNTIHVREIHKITNNQ
TGQMFSETVITSVDEEGRRSHECIDEDCGPSMYCQFASFQYTQPCRGQRMLCTRDS
CCGDQLCVRGHCTKMATRGSNGTICDNQRDCQPLCCAFQRGLFPVCTPLPVEGELCHDP
ASRLDLITWELPLEPDGALDRCPCASGLLCQPSSHSLVYVCKPTFGSVDQDGEILLPREVP
DEYEVGSFMEEVQRELEDLERSLTEEMALGEPAAAAALLGEEI

Signal sequence:
amino acids 1-19

N-glycosylation site.
amino acids 96-100, 106-110, 121-125, 204-208

Casein kinase II phosphorylation site.
amino acids 46-50, 67-71, 98-102, 135-139, 206-210, 312-316, 327-331

N-myristoylation site.
amino acids 202-208, 217-223

Amidation site.
amino acids 140-144
FIGURE 12

MELALRRSFVPRWLLLPPLLGLNAGAVIDNPEEGKEVDYVTVRKDAYMFWWWLYATNS
CKNFSELPLVMWLGPGGGSTGFNGFEEIGPLDSLKPRKTLWQAASSLFDNVPVGTGF
SYVNGSGAYAKDLAMVSDMVMVLLKTFSSCHKEFQTVPFYIFSSEYGGKMAAGIGLELYKA
IQRGTDKCNFAGVALGDSWISPVSALSWSGPYLYSMLLEDKGLAEVSKVAEQVLNAVNGK
LYREATELWGKAEMIEEIQNTDGVFNILKSTPTSTTMESSELFQSHLVCLCQRHVRHLQ
RDALSQMLNGPIRKKLIIPEQDGSWGGATNVFVNMEEDFMKPVISIVDELLEAGINTVY
NGQLDLIVDTMQEAWVRLKWPFLPFSLQKWKALYSDPKSLETSAFVKSYYKNLAFYWIL
KAGHMVPSQGDMLKMMRLTQOE

Signal sequence:
amino acids 1-25

N-glycosylation site.
amino acids 64-68, 126-130, 362-366

cAMP- and cGMP-dependent protein kinase phosphorylation site.
amino acids 101-105

Casein kinase II phosphorylation site.

N-myristoylation site.
amino acids 22-28, 76-82, 79-85, 80-86, 119-125, 169-175,
**FIGURE 14**

MNLSLVLAACLIGASAVPKFDQNLDTKWyQWKATHRRLYGANEEGWRRAVWEKNMKMIELHNGEYSQGKHGMNAMAFDMTNEEFQMRMCFCFRNQCFKFRGFMRFPFLFLPKSVGWRKKGYVTMPKQCGCWAFSATGAGEQMFRKTGKLVSQNLVDCSRPQGNQGCNGGFMAARAFQYVEKENGGLDSEESYPYVAVDEICKYRPENSVANDTGFVTVPVGKEKALMKAVATVGPISVAMDAGHSFQFYKSIYFEPDCSSKNLDHVGLVGVGFEGANSNNSKYWLKVNSWGP

**Important features:**

**Signal sequence**

amino acids 1-17

**N-glycosylation sites.**

amino acids 2-6, 221-225, 292-296

**N-myristoylation sites.**


**Eukaryotic thiol (cysteine) proteases cysteine active site.**

amino acids 132-144

**Eukaryotic thiol (cysteine) proteases histidine active site.**

amino acids 275-286
FIGURE 16

MSAPSRLARAAKLGLLCAVLGRAGRSDSGGRGELQPSGVAARFCPTTCCRCLGDLLDCS
RKRLARLPEPLSPWSVARLDDLHNRSLQIKASSMLQSLREVKLNNILELIPNLGPD
ITLSLAGRNVEILPEHLKQEFSLETLDLSSNNISELQTAPFAALQKLYLYLNSRVTSEM
PGYFNDNLANTLVLKLNRRSNISAFPKMFKLPLQQHELRNKRKNVNDGFLTGQGLKSL
KMQRNQGVTKLMDGAFWGSSLNMEILQDLHDNNLTEITKGWLYGGLMLQELHLSQAINIRISPD
AWEFCQKLSDDLDLTNFNSRLDDSSLGLSLNTNLHIGNRRVSYIADCAFREQLSSLKTLDDL
KNNEISWTEDMNGAFSGLDKRLRLILQGNRISRITKAKTFGLDAELHDLSDNAMLSQG
NAFSGMKKQLQPLLNTSSLCDCQLKWLPGWQVAENNQSFVNASCHAFQLLLLGRKSF1AVSF
DGFVCDDFKQPQITVPETQSAIJKSNLSFICSAASSSDSMFTFSAWKKDNLHDAEMENY
AHLRAGQGEYMEYTSTILRLREVFASEGMKYQCVISNHFGSSYSVKALTvNMLPSFTKTPM
DLTRAGAMARLECAAVGHAPQIAWQKDGTDFPAARERHHMPEDDVFIVDVKIEDI
GVYSCATAQNSARGISANATLTVLEPSFLRPLLDRTVTKETAVLQCIAGGSPPPKLNWTK
DDPLVVTHERHAAAGNQLIIIVSDSDVDAKYGTCMSNLT6TERGNVLSVIPPTCDSP
QMTAFSLDDDDGATVGVVIIAVCVTSVWWVVIYYHTRRRNEDCSITNNTDENTLPADI
PSYLLSQQGTALDRQDGYVSESSESSHQVFTSSGAGFFLPQHDSSTGCHDIDNSEDVAAAT
DLFLCPFLGSTGPMYLKGNVYSDGFETYHTGCSPDPRTVLMDHYEPSYIKKCEYPSCHP
SEECSERSFSNISWFHVRKLLNHTYSHNEGPQMKLNCNKSSLDFSANPEVASSNSNF
MTGFKGKALRRPHTDAYSFSFQPSDCQPRAFYLKAhSSDLGSEEEDGKERTDFQEEHIC
TFQKQLENRTFNQFSYXLDLT

Signal sequence:
amino acids 1-27

Transmembrane domain:
amino acids 808-828

N-glycosylation site:
amino acids 122-126, 156-160, 274-278, 442-446, 469-473,

Glycosaminoglycan attachment site:
amino acids 886-890

Casein kinase II phosphorylation site:
amino acids 99-103, 180-184, 263-267, 314-318, 324-328, 374-378,
383-387, 407-411, 524-528, 608-612, 692-696, 709-713, 731-735,
799-803, 843-847, 863-867, 907-911, 1003-1007, 1018-1022,
1073-1077, 1079-1083, 1081-1085

Tyrosine kinase phosphorylation site:
amino acids 667-675

N-myristoylation site:
amino acids 14-20, 36-42, 239-245, 257-263, 380-386, 427-433,
513-519, 588-594, 672-678, 683-687, 774-780, 933-939

Leucine zipper pattern:
amino acids 58-80, 65-87
FIGURE 18

</usr/seqdb2/sst/DNA/Dnaseqs.min/ss.DNA45415
<subunit 1 of 1, 215 aa, 1 stop
<MW: 24326, pI: 6.32, NX(S/T): 4
MHRDAWLPRPAFSLTGLSLFSLVPPGRSMEVTPATLNVNLGSDARLPCTFNCHYTVNHKQFSLWNTYQECNN
CSEEMFLQFRMKIINLKERFQDRVEFSGNNPSKYDVSVMLRVNPQDEGIYNCYIMNPPDRHRHGHGKIHLQVLM
EEPPERDSTVAVIGASVGGFLAVILVMVVKCVRKKEQKLVSTDLKTEEEGKTDGEGNPDDGAK

Important features:
Signal peptide:
amino acids 1-20

Transmembrane domain:
amino acids 161-179

Immunoglobulin-like fold:
amino acids 83-127

N-glycosylation sites.
amino acids 42-45, 66-69 and 74-77
Signal sequence: Amino acids 1-21
N-glycosylation sites: Amino acids 95-99; 104-108
Casein kinase II phosphorylation site: Amino acids 181-185
N-myristoylation site: Amino acids 133-139
Interferon alpha, beta and delta family signature: Amino acids 147-166

MIIKHFFGTVLVLLASTTIFSLDLKLIIFQQRQVEQLKLLNLQTLSIGQQCLPHRRNFL
LPQKSLSQPQYQKHTAILHEMLQQISFLFRANISLDGWEENHTEKFLQLHQQLEYLEA
LMGLEAEKLSGTLLGSDNLRLQVKMRYFRRIHDYLENQDYSTCWAIVQVEISRCCLFFVFSLT
EKLSKQGRPLNDMKQELTTEFSPR
FIGURE 22

MLFPMALPSVSWLLSCLIIILCQVQGETQKELEKPRISPCKGSKAYGSPCYALFLSPKSWMDADLACQRRPSG
KLVSVLSGAESGPSYSSLVRSISMSSYIWIGLHDPQGEFSGSDGWESSTNGMNYFAKEKPNSTILMPGHCGS
LSRSTGFLKWKDYCDAKLVPVCKPRD

Important features:
Signal peptide:
amino acids 1-26

C-type lectin domain signature.
amino acids 146-171
FIGURE 23

CCAGCTCTGTCCACCTCACCTTGGTGTCTGCTGCCTCCCACCAGGCAAGCCTGGGTTGAGAG
CACAGAGGAGTAGGGCCGGGACCAGTATGCGGGGACGCCTGGCGCTGCCCTGGCGTGGTGCTG
GCTGCCCTGCGGAGAGCTGGCGGCACGTGCCCTGCTGCTAGCTCTGTCCGGAGACCCACAGGAG
TGTCGGACTGTGTCACCATCGCACCTGACACCACAAACGAAACCAGATGTGCAAGAAGCAACACT
CTACTCCCCGGGAGATACTGTACCCTTTCCAGGGGACTCCACGCTGGACCAAGTGCTGTGCC
AGCAAGTGTAAGCCCTCGGATGTGGATGGCATCGGGGACACCTGCCCCTGGTGCTCTGCTGCA
ATACTGAGCTGTGCAATGTAGACGGGGCCGCCCCTCTGCAACAGCTCCACCTGCGGGGCTC
CACGCTCCTCCCACCTTTGAGCCTCGACTGTAAGTCGCCGCCACCCCATGCCCCTAT
GCGGCCAGCCCGATGCCTTGAAGAAGTGCCCTCTGCAACCAGAAAGAAAGAAAGAAAGAA
FIGURE 24

"/usr/seqdb2/sst/DNA/Dnaseqs.min/ss.DNA56405
<subunit 1 of 1, 125 aa, 1 stop
<MW: 13115, pI: 5.90, NX(S/T): 1
MRGTRLALLALVLAACGELAPALRCYVCPEPTGVSDCVTIATCTTMCKTTLYSREIVYPFPQGDSTVTKS
CA SKCKPSVDGIGQTLVSCCNTELCNVDAPALNSLHGCALTLPLLSSRL

Important features:
Signal peptide:
amino acids 1-17

N-glycosylation site.
amino acids 46-49
FIGURE 26

</usr/seqdb2/sst/DNA/Dnaseqs.min/ss.DNA56352
<subunit 1 of 1, 293 aa, 1 stop
<MW: 32562, pI: 6.53, NX(S/T): 2
MDTRYSKNGGSSEEVFPGPGWGRWGHWSRRPLFLALAVLVTTLWAVILSSILLSKASTERAALLDGHDLLRTNA
SKQTALGALKEEVGDCHECCSCGTQGQLTQTRAELGEAQAKLMEQESALRELREVRVQGLAEAGRGEREDVRTEL
FRALEAVRLQNNSCCEPCPTSWLSFEGSCYFFSVFKTTWAAADQDAADSAHLVTVGGLDEQFLTRGLTRNTRGRGYW
LGLRAVRHLGKVQGYQWVDGVLSSFSHWNQGEPENDAWGCRENMLHMLHTGLWNADPCDSEKDGWICEKRHNC

Important features:
Type II transmembrane domain:
amino acids 31-54

N-glycosylation sites.
amino acids 73-76 and 159-162

Leucine zipper pattern.
amino acids 102-123

N-myristoylation sites.
amino acids 18-23, 133-138 and 242-247

C-type lectin domain signature.
amino acids 264-287
FIGURE 28

<<subunit 1 of 1, 544 aa, 1 stop
<<MW: 60268, pI: 9.53, NX(S/T): 3
MLLPFLSLLGGSQAMDGRFWIRVQEVWMVPEGLCISVPCSFSYPRQWDTBAYGWFKAVTETTKGAPVA
THQSPREVEMSTRGFRPQTLGDPKGNCLVIRDAQMQDESQYFRVERGSVFTYNFMNGDPFLLKTVLSFTPRP
QDHNLTDLCTHVDPSRLGSVAQRTVRLRVAYAPRDLVISISRDNTPALEQFPGQGNPYLEAQKQGQFLRLCAADS
QPPATLSSVWLSQVRSLSSSHPWGPRLGELPGVKAGDSGRTCRAENRLGSQRALDSVQYPENLVRMVSLQA
NRTVLENLNGTSLPVLEGQLCLLCVTSSFCRSLSWTQGQVLSPQGSPDPVELELPVQOEHEGETCHAR
HPLGSQHVSLSSVHVKGLISTAFSGAFLGIGITALFLCLALIIKMKLIPRTQETTPRPFRSRHSTILDY
INVVPTAGPLAQKRNQKATPNSPRPQGAPSPESCKNQKQYQLPSFPEPKSSTQAESPQESQELHYATLN
FPGRPRPEARMKPQTDAYAEVKQF

Important features:
Signal peptide:
amino acids 1-15

Transmembrane domain:
amino acids 399-418

N-glycosylation site.
amino acids 100-103, 297-300 and 306-309

Immunoglobulins and major histocompatibility complex proteins signature.
amino acids 365-371
FIGURE 30

>subunit 1 of 1, 772 aa, 1 stop
>MW: 87002, pl: 4.64, NX(S/T): 8

**Important features:**

**Signal peptide:**
amino acids 1-21

**Transmembrane domain:**
amino acids 597-611

**N-glycosylation sites.**
amino acids 57-60, 74-77, 419-423, 437-440, 508-511, 515-518, 516-519 and 534-537

**Cadherins extracellular repeated domain signature.**
amino acids 136-146 and 244-254
FIGURE 31

GGGAAAGCCATTTGAAAACCCATCTATACAAACTATATATATATCTTTTATTGGCTGCTAGCT
GCCTTGCGCCTCAACATTTTCTATTCTGTCTGTCTGACTTTCAAGTTATATACCGTAGG
ATTTGATCCCAAACCATACATCGTGAAGGTATTTATATTGCTGTGACACCCTCACCCAAATT
CTGGTGAGTGTCTTCTTGGACAGGGATTTCCACCTTCATTTAATCATGAAACTCTGGCTGAC
AAAAAGGAGATTTGTGATTCTACTCTAAAAGTCAATATAGGACTTGCAAAAGAAGCTAGCAG
AAGACTCAACCTGGCACTCCATAAAACAGGACAGATTATTCAGGTGATGGCAAAAAATGGATT
CTCATCAACGGAGCTGCTAGAAGCCATGACAGATTTCACAAAAGAAAAACTCAAATGGA
GGCCACCCACAGAAACAGCATTTCTGGGCAAGGCTGTAAAATCAGAATTGTCGTCGTACATGC
TCACGACATTGCTTTTCTCCCAAAATTACACATTGTGGAGAGTGATGATACCTCTCCC
CTTCCTTTCTCTTCTCCATTTGCATTCAACGGATTTATATATTTTCAATGAAATTAAACCCCTGCA
GCAAGGGAACCTTATGCTGTATTCTGACTGTATGCTTTACCAAATGAGAAAAATGC
ATTCCCTGATCATCCTTTTCTAAACGTATTCCATTTGCAAAAAAAAAAAAAAAAAA
**FIGURE 34**

</usr/seqdb2/sst/DNA/Dnaseqs.min/ss.DNA56439

<subunit 1 of 1, 747 aa, 1 stop

<MW: 86127, pI: 7.46, NX(S/T): 2

MGWLNKDDYIRDLOIIICFLIVYNAILVGTQDFSYSSLLGVSKSTASSREIRQAFKLKLHFLDPKNFNPNAH
GDPLKINRAYVLDLDLRLPKYDKEKGLFDEQNGQYESWNYYRDPGIYDDDPEIITLEREFRDAAVNSEG
WFVNFYSPGCSHCHDLAPTWDFAKEVDGLRRIGAVNCGDDRMLCRMVNGSYPSLFIPRSGMAPVKBNGDSRK
ESLVSFAMQHVRSVTCTLWTGNFVSNQTAFAAAGWLIIFCSKGGDCLETQTRLRLSGMLFLNSLAKEYILE
VHNLPEDDELSANTEDLRAHHRWLLFFHGKNNENSNPELKKTLKLNDHIOQVRDFCSAIPSECSNLVVF
QPSSLAVFKQGTKEYEHHGKILYDLAFKAENVSHVTTLQPQNFPANDKPVNLVDFAPWCQCPARLLPEL
RRASNLLYQQLKFGLDCTVHEGLCMNQYQAPTTVVFQNSNIHEYEGHHSAEIQILEFIEDLMNPSVSILTPT
TFNELVTQKRKNEVWMDFYSFWCHPMCQVLMPEWKMRAMLTLGIVNGSIDCQYHSFCAQENVQRYPEIRFFF
PKSNKAYQYHSYNGWRRADSLRILWGLGLFPQVSTDTLQFSEKVLQGNHVIDFYAPWGCQCNFAPFEL
LARMIKGKVAKGVDCQYAQTQCAGIRAYPTVFKFYYERAKRFNQEEQINTRDAKALISKEKLETRNQG
KRNKDEL

**Important features:**

**Endoplasmic reticulum targeting sequence.**
amino acids 744-747

**Cytochrome c family heme-binding site signature.**
amino acids 158-163

**Nt-dnaJ domain signature.**
amino acids 77-96

**N-glycosylation site.**
amino acids 494-487
FIGURE 36

MDLAGLLKSQFLCHLVFCYVFASGLIINTIQLFTLLLPINKQLFRRKINCRSLSYCISSLQ
VLMLLEWWSGTECTIFTDPRAILKYGKENAIIVVLNHKFEIDFLCGWSLSEFGLLGSKVLA
KKELAYVPIIGWMWFTEMVFSRKWEQDRKTVATSLQHLRDYPEKYFFLIHCEGTRFTEK
KHEISMQVARAKGLPLKHHLLLRTKGFAITVRSLRNVSVAYVDCNLNFRNNENPTLLGVL
NGKKYHADLYVRRIPLEDIPEDDECSAWHLKLYQEDAFQEEYRTGTFTPETPMVPPRRP
WTLVNWLFASLVLYPPFQFLVSMIRSGSSLTLASFILVFFVASVGVRMIGVTEIDKGSA
YGNDSKQKLNĐ
FIGURE 38
</usr/seqdb2/sst/DNA/Dnaseqs.min/ss.DNA56045
<subunit 1 of 1, 270 aa, 1 stop
<MW: 28317, pI: 6.00, NX(S/T): 1
MATGFRYAGKVVTGGGRGIGAGIVRAFVNGARVVICDKDESGGRALEQELPGAVFILCDVTQEDDVKTLYS
ETIRRPGLDCVNNAGHPPPQRPSQTSAQGFRQLELNLGTYLTKLALPYLRKSRQGNYINISLVLGAIGQ
AQAVPYVATKGAVTAMTKALADESPYGVVRNCISPGNIWTPLWEELAALMPDPATIREGMLAQPLGRMGPAPA
EVGAANVLASEANFCTGIELLVTGGAEIYGCKASRSTPVDPDIPS

Important features:
N-glycosylation site.
amino acids 138-141

Short-chain alcohol dehydrogenase family protein
amino acids 10-22, 81-91, 134-171 and 176-185
FIGURE 39 Continued
FIGURE 40

>>Sequence Version 1, Fri Feb 7 12:06:40 2003 DNA257845 [min]
>>/usr/seqdb2/sst/DNA/Dnaseqs.min/ss.DNA257845
>>subunit 1 of 1, 977 aa, 1 stop
>>NW: 106477, pi: 7.04, NX(S/T): 9
MLLVILLVLAPYSQFARTPRPIILFLQPPWTVFQGERVTTLCCKGFRFYSQTKWYHR
YLGEILRTEDPNILEVQESGEYRCQAQGSPSLSPVHLDFFSASLILQAPLSYFEDGSSV
LRCRAKAEVLNITYNKNVLAFLNKRTEHPIACDKNAYRTGCYKESCCCPVSNST
VKIQVQEPFTRPVLRASSFQPISGNVPVTLCETQLLSRSDVPLRFRFRRDDQTLGLGWS
LSPNFQITAMVSDKSDFYWCKAATMPHSVISDPSRSQVIQIPASHPVLTLSPEKAILNE
GTKVTLHCETQEDSLRTLYRFYHEGVPRLHKSVRCEGASISFSLTTENSNGYCTADNG
LGAKPSKAVSLVTVPWSHPVNLSSPEQL海尔FQFHDAA
LERRANSAGGVAISFSLTAEHSGNYCTADNGFGPRQSRKAVSLSITVPVSHPLTLLSA
EALTFEGATVLHCQVRGSPQILYQFYHEDMLPSSTPSVGRSFSLTEHGSNGYN
CTADNGFPQROSEVSVLFTVTVSRFPILTRVPAQAVVGLLELHCEAPGRSPILYWF
YHEDTVLGSSAAPSASGLSLENLTAEHSGNYCANEAGLDVAQHDSTISLSVIVPVSRPI
LTFRAPPAQAVVGLLELHCEALRGSPILYMFYHEVDVLGKISAPSGGASFNLSTE
HSGYSECEADNGPAQRSEMTLKVAFVFSPFRVLTRAFTHAAVGLLELHCEALRGSF
LHYRFHEDVTLGNSRPSSGGLASNLTAEHSGNYCADERNLGAQROSETTVLYITGL
TANRSGPFATGVAGLLSIAAGALALLHCWLSRKRAGKFASDFPASPSDSQSEPTYH
NVPAMEELQPYNTNANPQEENGYYSEVRRQEEKKKHAVASDFPRLNKSPPYSEVKVA
STFVGSLFLASSAPHR
FIGURE 41

ATCGGTTCTCTGGCCTGCCGAGCCGCTCAGGTGACCTCGCAGGACAAGCTG
GTGAAGGACAGTGAGGAAACCTGCAGAGTCACACAGTTGCTGACCAATTGAGCTGTGAGCC
TGAGGACGATCCGTTGGCTGCGAGACCCCGCCCCAGTGCCCTCTCCCCGTGACGGCCTGGCC
CTCGAAGCTGTGACATGGAGAGATGACCGCTGGCCCTTCTCTACTGCGAGGCTGACTGCC
TTGGAAGCCAATGACCATTGGCAAATAAAGAGCATCCCTTTCTACTATGACTGGAAAAACC
TGCAGCTGAGCGGACTGATCTGCCGAGGCTCCTGCCCATTTGCTGGAATCGCGGCGAGTTCT
GAGTGGCAATGCAATACAAAGAGCAGCCAGAAGCAGCACAGTCCTGTGACCTGTAGAGGCC
ATCCCATCACTCACTCCAGCCCTGCGACACTTTGCTGACACAGGACTGCGCTCCAGGGA
TGCCCTGAGGCTAACACTGCGCCCGACGACCTCCTCCCCGTGGAGGCTTATCTCTCAAGG
AAGGACTTCTCTCAAGGGGAGGCTGTGTTAGGGCCCTTTCTGTGATCAGGAGGCTTTTCTATGA
ATTTAAGCTGCCCCACCACCCTCTCA
FIGURE 42
MERVTLALLLALGLTAELANDPFANKDDPYYDWNKLQLSGLICGGLLAIAGIAAVLSGKC
KYKSSQKQHSPVPEKAIPITFGSATTC
FIGURE 46
MEPPGRRRGRAGQPLLPLLALLALLGGGGGGGGGAAALPAGCKHDGPRGAGRAAGAEG
KVVCSSLEAQVPFTPVTLPNRTVTLILSNKISLKLNSFSGLSSLRELRDLRNNLISSIPD
FAFWGLSSLKRLDLTNRIGCLNADIFRGLTNVLNLSSGLNFSLSLQQGFTPFDYLASLRSLE
FQTEYLLCDCNLWMHMHWVKEKNITVRDTRCVYPKSLQAPQVTGVKQELLTCDDPELPSF
YMTPSHRQVFEGDSLQFCMAYSYIDQMDQYQLVYQDGRIVETDSESQIFVEKNMIHNCSLI
ASALTISNIQAGSTGNWVGCHVQTKRGNNTTRTVDIVVLESSAQYCPPERVVNNKGDPRWPR
TLAGITAYLQCTRTHSGGYPGNPQDERKAWRRCDRGGFWADDYSRCYANDVTIVERLYMF
NQMLNLTNAVATARQALLAYTVEAAANFSKMDVIFVAEMIEKFGRFTKEEKSKELGDVMVD
IASNIMLAERVLWLQREAKASRIVQCLQRIATYRLAGGHVYSTYSPNIALEAYVIKSTGFTGMCTVTQKVAASDRTGLSDYGRRDPEGNLKQLSFKCNVSNTFSSLALKVCYILQSF
FKTIYS

Signal peptide:
amino acids 1-33

Transmembrane domain:
amino acids 13-40 (type II)

N-glycosylation site.
amino acids 81-85, 98-102, 159-163, 206-210, 301-305, 332-336,
433-437, 453-457, 592-596

N-myristoylation site.
384-390, 403-409, 554-560
FIGURE 48

<usr/seqdb2/sst/DNA/Dnaseqs.min/ss.DNA59814
<subunit 1 of 1, 224 aa, 1 stop
<MW: 24963, pI: 9.64, NX(S/T): 1
MRVSGVLRLLALIFAIYTMSPFIRSYMSFSMKTLRPRWLAASPTKEIQVKYKCGLIPCPANYFAPKICSGA
ANVVGPTMCFDRMIMSPVK3NVQGLMLALVNGTGGAVLGQKADMYSGDVHLVVKFLKEIPGGALVVLVASYD
DPGTQDNDERSPXLFSDLGSSYAXQLGFDRDSWYFIGAKDLRGKSPFEQFLANSPDTNKYEGWPELLEMGCMPKPF

Important features:
Signal peptide:
amino acids 1-15

ATP/GTP-binding site motif A (P-loop).
amino acids 184-191

N-glycosylation site.
amino acids 107-110
FIGURE 49

CTGGGATCAGCCACTGCAGCTCCCTGAGACTCTCTACAGAGACGGACCCAGACATGA
GGAGGCTCTCCTGACCCAGCTGCTGCTGTCTGCTGGAGGGGAGGTGCGAGTCC
AGCACCAGAGGCTCCCTATCAAGAtGCAAGTCAAAACACTGCCCCTCAGAGCCAGGACCCAGAG
AAGGCCTGGGCGCCCTGTTGCTGGAGCTCCGAGAAGGAGGAGCAGGCTGCTGTGTGCCTG
TCCCTGTCGAAGCCGAAACTCTTTGACCACCCGGAGAGAGGACAGAGGGTCAGGGCAGGG
CCCATCTCCTCCAGGCACCCAGGCTGGAGACCAGGGGACCCCTTGGGCGGTGCTCTG
AGTCCCGAGCCCGACCCATGACAGCCCTGTACACCCTCGCCCTGAGGAGGACCAGGGCGAGG
AGAGGCCCGGTGTGAGGTGATGCAAAAATCAACCAGGTGCTCTGCTTGGAGCCGAGGAGACCA
AGACCCAGCATCTACCCACCCCATGAGGCTCCAGGGGCACTCCTGCCCCCGCCTGCTCCA
AGGCCCAGGCTGTGGGACGTGGACCCCTCCACCTGCCCCAGCTAGACAAATAAACC
AGCAGGCAAAAAAAAAAAAAAAAAA
FIGURE 50
MRRLLLVTSLVVLWEGAVPAPKVPQIKMQVKHPSEQDPEKAWGARVVEPEKDDQVFLFVQKPKLLEKPRGQGRFGPTGKAWMETEDTLGRVLSPEPDHSLYHPPEEOQG
EERPFLVMNHQVLGPEEDQDIYHFPQ
FIGURE 52
MGLGARGAWALLLLGTLQVLALLGAAHESAAMAASANIAENSGLPHNNSANSTETLQHVPSDHTNETSNSTVKTPTSVASDSSNTTVMKPTAASNTTTPGMVTSTNMSTTTLKSTPKTTSVQNTSQISTSTMVTNNSVTSAAASSVTITTTMHSEAACKGSKFDTGFSVGGIVLTLGVLSELIGCKMYYSRGIRYRTIDEHDAII
FIGURE 53

TTCTGAAGTAACGGAAGTCATCTGGTATAAGACCTCAACACTGCTGACCATGATCAGCGC
AGCTCTGGAGACATCTCTTCATGGGACTAAATAATTGGGCTGTCTCTCAGTACACCTC
TCAGTTATGGCTAAATCTGTCCACTCTGTGCTGCTGCTGATGCGGTTTCTATTTACTTGA
ATGATCGTCTCTTGAATTCTCTACATTCCAAACAGGAGAATACACGAGAAGGCTACACACTCT
TCAGAACACACAAATATAATAATGTGTTTGGATACCCTTATCAAGTTCTTTCTGGACAAAG
GAAAAGATTACAATAAATACCAAGATGTTTAGTAGAATTCTCTTCCAAACCCTCCAAGATTG
TAAAGAAGTCTTACAAATGAAGATGATCGTACCTTCTCTGCTGTCAGTAGATAGTAA
AATTCTCTATCTCGGTAATATATACATCTGGAACACATGGACACTTGTCTCTGAGTACTCGA
GAGGGAGACTCCTGAGACACGAGAACACTATATTCCAGACTGCTTCTCCTGCTCCGTAATACCTTA
GCACAAATCTGCTGTGGTCTCCAGACATTAGAAGAACACTTGCTGATAGTCAATGCAT
ATCCACTTATATCATCACCACATCTCTTCAGGCTCTACTGATTACCCAAACGGCCTGTGTTCTAGT
GGAAACCTGTGGACAAACTCATGTGATTTAGTGGAACAAGTAAAGTTTCTTCAACCTAGTATTTG
CAGAGCTGTCTCTGCTGGGAATCTTCGACTGCTGCACTGGCAAATACCTCCAGGACAA
CCTGAGGAAGCTTTATCTCAGAAATAACCATACACATCAGATCGGGTGCCTCCACCACAG
TATCTAAGCCAGCTCTATGCACTGGATATGTCCAAATAATACCTAATAGTATTTAACCTCAGG
GTATCTTTGATGTATTGGACATATATAAACAACTACACTGATTCTCTGCAACAAATCTTGGATTTG
CGGTGGGCAAGATGAATGGTAGACTGTGATTGTTACAAATCTACTGAGTCAAGCTCAGTG
CGTGGGAATGTGCCAGAGGCCCAGAAAGTTTGCTGGGATGCTATAGGATGCTCAATG
CAGAAGCTTGTGGATTTGAAGACAGTTGGGATTGTGTAAGCAACATTGAGATAAACACTGCAAT
ACCCCAACAGGGTATTCTCCGGAACAGACATGCGTGGACCACGGTGACAACAGCCAGAT
ATTAGGAAAACCAAGGTCATCATTTAAGGAGTCAACAAACCACAGGGAGTCTCCCTCAAGAAAAAC
TTCAAATTACTTGAGATCTCAGTCTCAGCTGATACATATATCTCTCTACGGAGAAGCTCTCT
ACCTATGCTGCTTTTGAGACTGCTGCTGGTAAATGCTGGCCTACGCGCATCGGACATG
ATACAGAAACAAATTTTGCAAGGGAACAGCAGTGATTGTGTCATGACAGCGCTGCTGC
ATTCACCCCTATAAAGATATGGCATGTTTCCCATGGGAAACACAGCTACTTACTATTGTAGTAA
AACCTCTGTTTGGATTGAGACTGAACCTGACCCCTCTGCAGATGTCACACCTGACAACCACC
CTCAATCGAGAGCAAGAGAGAAAGAGAAAACCTACTACAAATTTGGCTGCCATCA
TGGTGGGGCTGTGGCCCTGGTTACATCGGCTCCCCTTCTGTATTTATACTGTGTTGATATTTCA
TAGGAATGGAATCGCTCTTTCTCAAGGAAACTGTGCTATATGCAAAAGGGAGAGAAAGAGAT
GACATAGCAGAAGCTGGCAGCTAAGAGGACAACTCTAATCTGGAAATGCGGGAACCTCTCTT
TTCAAGATGTCTACAGAATACAGAAGCTTACCTCTGAGGATTTGTAATACACACCAT
ATTGCCTCTAAATGGAGATCTAGCTGATACAAACACATACAGATAGGAAACAGATAGTAACCGA
AGCTACAGAAGAAGATGGTAGATTCAGACTCAAGACTCAGACATCACAACACTCAAGGATG
ACAGAAGACTCTGTGGTTTGGGTTTTTTAACCCTAAGGGAGATGATGTT
FIGURE 54

>>/usr/seqdb2/sst/DNA/Dnaseqs.min/ss.DNA81757
<<subunit 1 of 1, 649 aa, 1 stop
<<MW: 72995, pI: 7.88, NX(S/T): 7
MISAAWSIFLIGTKIGLFLQVAPLSVMAKSCPSVCRCDAGFIYCNDRFLTSIPTGIPEDAT
TYLQLNNQINNAGIPSDLKNNLLKVERIYLYHNSLDEFPTNLKPYKVELHLQENNIRTITYD
SLISKIPYLEELHLDDNSVSASVIEEGAFRDSNYLRLFLSRLNHSLTIPWGLPRTIEELRLD
DNRISTISSPSLQGLTSKRLVLDGNLLNNHGLGDKVFFNLVNLTELSLVRNLSLTAAPVNL
PGTNLRKLQLQDNH1NIRVPFPNSYLRQLYRLDMSSNNLNSLPQGFDLDNITQLILRNN
PWYCCKMKKWDRWQLQGLPVKNVRGLMCQAPKVRGMAIKDLNAELFDCKDSGIVSTIQI
TTAIPNTVYPQQGQWAPVTKQPDLDNPKLTQDDQOTTGSPSRKTITITKSVSTSDTIHISW
KLALPMTALRLSWLKLHSPAFGSIETETIVGERSEYLVTALEPSYKVCVMVFMETSNL
LYLDPETPVCEITEETAPLRMYNTTLNREQKEPEYKNFNPLLAIIIGAVALVIALALLALVC
WYVHRNGSLFSRNCAYSKGRRKDDYAEATKKDNSILEIRETSFQMLPISNEPISEKEEFV
IHTIFFPPGMNLYKNNHSES3SNRSYRDGIPSDHSHS

Important features of the protein:

Signal peptide:
1-28 MISAAWSIFLIGTKIGLFLQVAPLSVMA

Transmembrane domain:
531-552 AIIGAVALVIALALLALVCWYV

N-glycosylation sites.
226 NLTE
282 NLSN
296 NITQ
555 NGSL
626 NHSE
633 NRS

Tyrosine kinase phosphorylation site.
515 REQKEPEY

N-myristoylation sites.
12 GTKIGL
172 GLPRTI
208 GNLLNN
359 GIVSTI
534 GGAVAL
556 GSLFSR
640 GIPSD

Amidation site.
567 KGR

Leucine zipper pattern.
159 LFLSRNHSLTIPWGLPRTIEEL

Phospholipase A2 aspartic acid active site.
34 VCRCDAGFIYC
FIGURE 56

MEKYGGDLVLAGPGGGGLGPFDVSARLTKYIVLLCFTKFLKAVGLFESYDLLKAVHIVVQFIFILKLGTAFFM
VLFQRFSSGKTITKIQWIKIFKHAVAGCIISSLWWFLGLTLCGLFRLLLFLFHEHSDIVVISLLSVLFTSSSGGPA
KTRGAAFFIIAVICLLLFDNDDLMARKMAEHFEGHHDSDALTMLYTA1AFGADVHDHGGVLLVIALCCKVGPHT
ASRLSVDVGGAGKLQALSHLVSLCLFNPWIVLVSLVSSTESKVESWFSLMPFATVFIPFNILDVYDAGICSVK
MVEVSKARYGSPFIPISALLFLGNGFWTHPITDQPAMMNKAHAQEESTEHVLSSGCVVSGAIPIFSANILSSSSPSKRGQ
KGTLLGVSPEGPTLYFMGDAQHSSQIQSPFPKEKSLQIKLEEDSRQIFYFCLNILLFTFVEFVGYVTNLSIG
LISDGPHMLFDSCSALVMGFAALSRSWSKATRFASYGRIEILSGFINGFLIVIAFPVFMESSVARLIDPPELD
THMFLPVSYYGLILVNLIGICAFSHAHSIIHAGASQSGSSCHSDDSHSIIIIMHDHGHGHSIHSAGGGMNAMRGV
FLHVLDATLGSIGVSTVLIIEQGFIADPLCSLSTAILLLFLSVVPLIKDADQVQLLRLPPPEYEKEHILAEK
IQRIEGLISYRDHFWRHSASIVAGTHIQVTSDVLEQRIIVQQVTGILKDAGVNNLTEIQVEKEAYFQHMSGLST
GFHDVLAMTQMESMKYCKDGTYM

Important features of the protein:
Signal peptide:
amino acids 1-46
Transmembrane domains:
amino acids 59-77, 101-119, 150-167, 205-223, 239-258, 267-284, 305-324,
343-360, 421-440, 452-469, 486-505, 522-539, 592-612, 621-641
N-glycosylation site.
amino acids 721-725
Glycosaminoglycan attachment site.
amino acids 143-147
cAMP- and cGMP-dependent protein kinase phosphorylation site.
amino acids 225-229
Tyrosine kinase phosphorylation sites.
amino acids 750-758, 756-764
N-myristoylation sites.
amino acids 14-20, 46-52, 102-108, 112-118, 144-150, 317-323, 347-353,
369-375, 372-378, 437-443, 462-468, 529-535, 549-555, 553-559, 579-585,
582-588, 583-589, 584-590, 605-611, 737-743
Multicopper oxidases protein:
amino acids 561-569
FIGURE 57

CACAGCTCCCTTCCAGGAGCTGAGAATCTGCTCTCTCACCATA
CAGCTGCTCTGTATCTGTTCCTCTGCTCTCCATCTTCTCTCCACGAAAGGGAAGG
CCTGCAAGGCGCTGTCGGACAGGAGAACCAGGCTCTGCTGGCAAGGAGTCCCTAGCCTAC
ACTCAACAACCTGAAAGGACATCATGTGAGGCTGCTGTAAACCTGTGCAAGGCTGAGCCAGA
GCCCGCTTTTGCTGGCTGGCCTCTGGGACACTCCACAGGTGTAGCACTCCAAAGCAAGACT
CCAGACAGCGGAGAACCTCATGCTGGCAGCTCAGGATACCGCAGCGCTCCTGTGCTCCCTT
TTGACGGCTCTGGAGCGTGGCAAGTGGTGGGAGGGCTTCATCTCGGGGCTGCAAGGACCC
TGGGAAAGTTCCAGAACCCCATCGCTTGTCTCAATTGTGACATCAAACCTTACAGACTTC
ATGAGCCAACCTACCCCAAGGCCTGACACATGTGGGCTCTCCAGTGCAACAC
CACCAGCATTTCCACCATGACCCTCGTCACAGCTACAAATCAGACGAGACATCCATCTGCTAG
AGTGCGAGGTGGCAAGCAAGCAGGGCTGCTGACCAAGACTGCAAGTGCTCTCCATCTTC
AGGTCCATTCAGGCTCTGGCATTTAATTACAGCATCCAGTGGTGGCAGGATCCCT
CCTAGCCTCTTGACATGATGTTGCTGGAAAGAGCATCCCAAGCAAGCTATAAAT
AAATAAAGCTCA
FIGURE 58

MRLVLSSLLeLCSIFSTEKRRPAKAWGRRRLCCHRVPSPNSTNLKHGTVLRCKPCLEPEPLMV
PGALPQV

Important features of the protein:
Signal peptide:
amino acids 1-21

N-glycosylation site.
amino acids 48-52

Amidation sites.
amino acids 23-27, 33-37
FIGURE 60

Important features of the protein:
Signal peptide:
amino acids 1-25

N-glycosylation sites.
amino acids 117-121, 139-143

N-myristoylation site.
amino acids 9-15
**FIGURE 62**

```plaintext
>>/usr/seqdb2/sst/DNA/Dnaseqs.min/ss.DNA77629
>>subunit 1 of l, 745 aa, 1 stop
>>MW: 78990, pI: 5.26, NX(S/T): 6
MFPLRALWLVALLGVAAGCSCEPCACVDKYAHTQFADCAKELKREVPEGLPANVTTLLSANIKTVLRGAFADV
TQVTSWLNKHSNRTVPETPGALVSLQLRLNDLSHNFQISSFWSDLRNLSALQLKHNRLGSLPRDLALGFDP
LRSLRLHNRLRRTAPGTDASLASHHQLHYHPFCGCGCLWQLQWAALAASLREPDSIACASPPALGQVPV
YRLPALCAPPVHLSAEPEPLAAPGTPILRAGLAVLHCADGHPTPLQWQLQIPFGTVEPVPSLQGEDDGVG
AEEEGEGEDGGGDLTTQOQTPAAPTAPFAPFAPTFRLALANSSLVFLLLSAKEAGVYTCRHLGANSTSRV
AVAAATGPKHAPGAGGEPDGQAPTQSETKSTAKGRGNSVLPSKPEGKIQQGLAKVSLGETEPEEDTSEGEE
AEDQ1LADFAEEQCRGNGDSPRSYVSNHANQSAELKPHVFELGVIALDVAEREAVQTLTPLAARMGPGPAGG
APRPRGRRPRLLLYPAGGGAAAVQWSRVEEGVNAYWFRLPRGNTNYSVCLLAGAEAHQVVVFSTKKEPSSLV
IVAVSVFLLVLATVPPLLGAACCHLLAKHPGCYRLLRPRQAPDPMEKRIAADFPRASYLESEKSYPAEGG
EEPEDVQGEGLDEDAEQGPDGSGLQRESLAACSLVESQSKANQEEFEAGSEYSRDLPLGAEAVNIAQEGWNY
RQTAG
```

**Important features of the protein:**

**Signal peptide:**
1-19 MFPLRALWLVALLGVAAGS

**Transmembrane domain:**
587-610 LPSLLIVAVSVFLLVLATVPPLL

**N-glycosylation site.**
52 NVT
tl
121 NLSA
337 NGSL
364 NSTS
474 NQSA
563 NYSV

**cAMP- and cGMP-dependent protein kinase phosphorylation site.**
397 RKST

**Casein kinase II phosphorylation site.**
19 SCPE
202 SLPE
289 SGED
346 SAKE
411 SKPE
431 TBTE
433 TEPE
440 SEGE
544 SREVE
583 TKKE
N-myristoylation site.
15  GVAGSC
48  GLPANV
165 GTFDAL
296 GAAEGE
351 GVYTCR
362 GANSTS
390 GQAPTS
419 GQGLAK
514 GGAGGA
536 GGGAAV
557 GLRPGT
561 GTNYSV
610 GAACCH
661 GGEAGG
716 GSEYSD

Amidation site.
522 PGRR

Prokaryotic membrane lipoprotein lipid attachment site.
10  VWALLGVAGSC
603 LATVPLLGAAC
AATGAAACAAAGCATAATTTTGAGCAATACTGAACATCAATAATACCCCTTTAGTTTTATATACCTATTATTTTATCTTTAAAGCATGCTACTTTTACTTTGCAATATTTTTCTTTATGTTAACCTTTTGCTGATGTATAAAACAGACATGCTTTATAATTGAAATAAAATTATAATCTGCTGAAATGAAATAAAAATAAAACATTTTGGAATGAAAAAA--------------------------
FIGURE 64

subunit 1 of 1, 800 aa, 1 stop
MW: 87621, pI: 4.77, NX(S/T): 7
MAVRELCFPRQVRQVFLFPLFWGVSAGSGFGRYSVTBEETKGSFVNLAKDLGLAEELAA
RGTRVSDDNKQLMLDSHTGNLNTNEKLDREKLCGPKCPMYLYFQLMDDPFIQYRAELR
VRDINDHAPVFQDKETVLKISENATAGTAFLARERQDPDGLNGIQNYTISPNSSFPHINIS
GGDEGMIYPELVDKALDDREEOQGELSLSTLTALDGGSPSRSGTSVTRIVVLVDVNDAPQFAQ
ALYETQAPENSPIGLIVKWEVDVSGNAVNEVSYFFDASENIRTFFQINPFSGEIFLRE
LLDYELVNSYKINIQAMDDGGLSARCRVLVEVLDTNDNPPELIVSSVSAENSPETPLA
VKINDRGDSNGKMKVCIQENLPFLKPSVENFYILITEGALDREIARAENITIVFTDLG
TPRLKTEHNITVLVSDVNDAPAFTQTSYTLFVRENNSPALHIGSVSATDRDSGTNAQVTY
SLPPQDHLPLASLVIINADNHFLALRSLDYEALQAFESPFRGVATDRGSPALSREALVRV
LVLDANDNPSFVLYPLQNGSAPOCETELVPRAEAPGYLVTKVVAVDGDGQNWLQSYQLLKAT
EPGLFGVWHANGEVRTARLLSERDAAKHRVLVVLKDNGEPPSSATATLHLLLVDGFSQYPL
PLPEAAAPQAQAQAEADLTLTVLVALASVSLFLLSVLFLFVAVRCRSLASVGRCSVPEG
PPFGLHVDFVRGAETLSQYQYEVCLTGGPGTSEFKFLKPVISDIQAAQPGRKGEENSTFRN
SFQNPNOQ

Important features of the protein:
Signal peptide:
Amino acids 1-26, Transmembrane domain: amino acids 687-711
N-glycosylation sites.
Amino acids 169-173, 181-185, 418-422, 436-440, 567-571, 788-792
Glycosaminoglycan attachment site.
Amino acids 28-32
Tyrosine kinase phosphorylation sites.
Amino acids 394-402, 578-585
N-myristoylation sites.
Amidation site.
Amino acids 781-785
Aminoacyl-transfer RNA synthetases class-II signature 1.
Amino acids 117-138
Carchers extracellular repeated domain signature.
Amino acids 121-132, 230-241, 335-346, 439-450, 549-560
FIGURE 66

MAFRMPFLLLLCLAKTGVLDIMFRSCAPGWYHKSNCYGFRLNKRNWSDELECSYS
YGNAHLASILSLKAEASTIAEYISGQRSPQPIWGLHPQKQWQLQWIDGMAMYRWSW
SMGGNKHCAEISSNNNFLTSSNECNKRQHFLCKYRP
FIGURE 69

TCCGCTGTGCCCCAGTCCCCGGGCGCTGAGGCAAGCAGGAACCTCC
CTCCCCCTCTCCCGGTGCTCCGGAGGCTCCCCTACGATCCGGCTGCAGTGGG
CACGTCGAGGACACGAGCTCTGGGAGTTAGGAGGGCCACGACGAGGCAGG
AGAATAGCCAGGAGCCAGGAGCAGCCGACAGAGGGAGGAGGCTGGCTG
CATCGAGAGATGTTGCTTCCATTTAATCCCTTACCTGTCTTTGCTG
CTTTGCCATCTCTTGCTGCTCTCTTTCTCTGTTGGTGTGCTGACAGC
tCTTGCTGCTGCCTCCGGAAAGAGCAGACAGCCAGCCACATCTGCAACAGCAG
GCAGCCTGGAACGAGCAGTCTACCTCTATGGAGATGAGCAGAGGCTCTG
GAACCTTCATAGCTCTGGAGATCCAACTGGCATGGCTTGGCCCTGCTCTTTTG
GAGTCGGACCTGGACCTCCCGCTCTCTCTGCTACAGCCTGTAACCCCGAGCCTCAC
CTCTACGATGAGCTCAAGTGCCAGCCAGGAGGAGAGGAGCAGCACTCTCCA
GAGACCCAGCTCTCTTGGGCTCGGCTAGAGAGACACTCACTCGG
GGCCCCAAATACTCAACTACCACCTTTGCTGCCCTGCTGCTCTCTCTCT
TGAGAGGAGGTAGGAGAA
CGGACAGAAGCTTGGAGAACTAATTGCTTTGGAGCGAGGCCCCAGCCACCCCAAGGG
CACCACTGCTGCTGCTGTACACCGGCCCTGGGCTGCTACCCAC
TAGGAGGCTGCTTCTACACGCTCTCTTGCCCCAGGCCCCCCAGGCTG
GAAGAGGCCCTTGCGCTGAGCCACTGGGAGGCTAGGAGGCCATCTGGG
TCTCTTTGGAGAGAATAATATTGTATTTATAGGTT


FIGURE 70

MERHCLLFILLTCLRMLCHDPQGSGARWPRVSPVEPCSRRLLAVLLLCLGVTAGC
VRFCCLRKQAAQPHLPARQFCIDVAVIFMDSSPVHSTVTSVSYSSVYPLLQMLRPLFUE
LDLDSTAPPAYSLTYTFEPFFPSYDEAVKNAKREEGFALSQKFSLLLGAQGLETTFVPQES
GPNTQLPPCSFGAP

55-61  N-myristoylation site
50-61  Prokaryotic membrane lipoprotein lipid attachment site
FIGURE 72

MALGVPISVYLLFNMATLTEEEAAVTVPITAAQQADNIEGPIALK
FSLHCLNLEDHNSYCIDGACAFHHELEKAIICRCFTGYTGGERCEHLTLT
SYAVDSYEKYIAIGIGVGLLSGFLVIFCYIRKRYEKDKI

Important features of the protein:

Signal peptide:

1-20 (weak)

Transmembrane domain:

103-117

Motif name: N-myristoylation site.

4-10
106-112
110-116

Motif name: EGF-like domain cysteine pattern signature.

75-87

Motif name: Integrins beta chain cysteine-rich domain proteins

66-88
Important features of the protein:
Signal peptide:

1-30

Transmembrane domain:

425-443

Motif name: N-glycosylation site.

58-62
126-130
291-295
501-505

Motif name: Tyrosine kinase phosphorylation site.

136-143

Motif name: N-myristoylation site.

29-35
61-67
247-253
267-273
271-277
331-337
502-508
512-518
562-568

Motif name: Glycosyl hydrolases family

310-319
Important features of the protein:

Signal peptide:

1-20

Transmembrane domain:

none

Motif name: cAMP- and cGMP-dependent protein kinase phosphorylation site.

242-246

Motif name: N-myristoylation site.

22-28
48-54
121-127
136-142
141-147
328-334
447-453

Motif name: Leucine zipper pattern.

295-317
FIGURE 77 Continued
FIGURE 78

>Sequence Version 1, Mon Mar 17 15:03:37 2003 DNA336882 [min]
><usr/seqdb2/sst/DNA/Dnaseqs.min/ss.DNA336882
><subunit 1 of 1, 318 aa, 1 stop
><MW: 35629, pI: 5.58, NX(S/T): 1
MTQLASAVVWLPTLLLLLLLFWLPGVCVLHGFSTMSGSVGESLSVSCRYEEKFKTSDKW
RVSILKILCKDIRKTSSSEEARSGRVTRIDHPDNLFTVTVYESLTLADTYMCAVDSL
DGSLGFKDYFKIELESVPSEDVSSPGPTLETDPVSTSLPTKGPALGSNTEGHEHDYQ
GLRPALLSVLALLLLFLVGLTSLLAWRMFKRKLVKADRPHELSONLQASEEQNECQVNL
QLHWSLREEFVLPSQVEVVEYSTLALPQELHYYSSVAPNSQRQDHSNGLHQPQDK
AETYEIQKPRKGLSDLYL
FIGURE 79
TTCCATAGACAGCATGCTCAAAAGGAAATCTCTTTAACCCTAGTTGCGCAGAGGCTA

FIGURE 79 Continued

GTCCCTAGCTGGAGCTTAGGAGGGCAGAGCTAGTCTGATTCGAGTCGAGCCACT
GATTGTCACACGCTTTCCCTGTCACTGCTCAAAAAATTGGCAATTCTCTTTGATTTTTAGT
TGTTGAAATTTGGCTGTTTCAAGCAATTGGTACATATTGAAAGTCTAAGGGTACGCAAGTCAG
TGGGAGGACTTTTTTCACTCCCTGGCATTAGCAAGCTCAGCCCTACTTTCCAGTGGCACCAAC
CTCCCTATTAAATAAGTAGCAGAAGAAAGTGATTGTACAGTGAGGAAACAGTGAGGGGAGG
CAGGGTTCTGCTCTCTTCTCACTTAAACCGGCAACACGCTTGGCCCTGTCTTTGCCC
CAAAGGTATTTTTGTCAGCTCTGCAAATTGGAGCTATTTCTCAGTGTCTTAACCCTTGG
GTAAAAAGAGAGCCTCCTCTGTTGGTCAGCTTAAGAGCTGATAGTAGTAAGTGCTCTCT
TCCAAAGAGATGGCAATAGCTGGGATCTACTTTAAAAAAGTTGGCTGAGTATTTTGCA
AGAGGTTAGGATTTTTATGGTTCTCTATTTCCCTTTACGTTCTGCAGTTCTCAGACGT
ATTTTTAAAATACCTCAGGTTATGAGAGAAGAATTAGAAAAGGAATTTATTATAGTGG
ACTTGAAATATTTATTTGTAGATCCTCTAAATAAAGCGTATATTTCTGTT
<Tue Mar 6 10:09:06 2001 DNA184073 [min]
</usr/seqdb2/sst/DNA/Dnaseqs.min/ss.DNA184073
<<subunit 1 of 1, 502 aa, 1 stop
<<MW: 55431, pI: 5.43, NX(S/T): 3
MHFFRVYSVTLKLKLHPAFQSCLLLTLGLWRTTPEAHASSLGAPAISAASFLQDLIHRYGEGDSLTLQQHLKALLNHLDVGVGRGNVTQHVGHRNLSCTFSSGDLFTAHNFSEQSRIGSSELQEFCLTIQLQDSRCTSENQNEENEQTEEGFRSAEVEWGVYGLCVCVIIISLCSLL
GASVVFPMKKTFFKRLLLFYIALAIGTLYSNALFQLIPEAFGPFDYVYSSAVVFPGEFYLFFTEKILKILLLQKNNEHHHHGHSHYASESLPSKKDQEEGVMKQLMQGLDHMIPOHCSELDGKAPMVDEKVIGSLVQDLQASQSACYWLGKVRYSDIGTLAWMIILSDGGLHNFIDGLAIGASFTSVFQGISTSVAILCEEFPHELDDVFVILNAGMSIQALFNFLSACCCYLGLAFGIAGHFSANWIFALAGGMFLYISLADMFPMNEVCQEDERKGSILIPFIINL
GLLTGFTIMVLMYSGIQIG
GGGCCAGTAGAGTGCTGCTGGCTAGCTGACTACTACATCAAGCTCCACGCTTTGAAA
ACACATGTGTCCGCCGCCAGATATTGAAACATTAATTGATATAATTAAAAGTAGC
GTTTCTCTACAATTGCTGAGAAGTGAACCTACGCCACACTCAATTTGCAGATCTGC
TGAGCGAAGAATAAACCGAGATGGAATAACCTAGAAACAGAAAGACAGCAGCTCCACAT
CTCCATTTTGCCTCAATGCTCCAGCTGTTGCTGTAACCTTTGCTGATTTGGTACTG
CTGAGGAGCTTTGGGATGAGGTTTTTGCACTATCTAACATTACAGATTTTGAGA
AAATTGGAACCTCCAGAAATCCATCCAAACAGCAGCAGGATAAATTTATCCCGCAACT
GGCGAACACTCCACATTTCTCTCTCATGGGAAGATTCTTACTGCAATAGATTCTCTC
ACTGAGAGGCGAGAACAAATAGGCCATCAAATCTGTGCCAGAGCTAATACATCTACCTC
AGACCAACAGATGTAATCCATGTCCTAAGTGTGGCAAATGTGCACAAAATATGGCTACTA
TTTACAAACAATAGGAGAAGAACAACCTTGCCCTAAACAGTGAAGAAAGAAGCTGAATAGACAGAA
CTCCACCTTAGTTAGAATAGACAGTGGTTGGAAGAAAAGGATTTTCTTATATGCAAGCATT
ACTCAGTTTTCTCTTCTTTGTGGCTGAGATTATCATGGGAACCTCTCGTGCAAGATTGGTT
CTGGGAAGATGCGCTGTCCTTCCTCTCCATTTGTACCTCTCAAATATTGAGGTTAAAAC
ACAAGCTTTTCCATGGAGATCTCGGAATATTGAATATATATGATTGTGAGAATTTATACAGA
CATAAAAAGGAGGTCAAACATAGTTCAAGAAAGAGCTCCATGAAACAAATATGGGAGA
ATTTAGTACTAAGAATCTGACCAGATCAATTGATCCAAAGGATGTGCTATTATTTCTAAAA
AGGAAATATTATATTCTCGCTAGTGCTGCTGAAATTATTTTGGATTGTGCGAGAAGACGC
TGCCCCAGTGAAGACTGAGGATTTTGAGATTTGATGCTCTCCTAAATTTCCCTGAAGACTA
AGAGACTCTTGAGTAAAGCTCCTATAGGAAAGAGGAAACTACGGTACCGAGCAAGGCAG
AATCTGCA
Important features of the protein:

Signal peptide:
None

Transmembrane domain:
42-62

Motif name: N-glycosylation site.
   91-95
   101-105
   176-180

Motif name: N-myristoylation site.
   17-23
   97-103
CTGGCTGCCCCGCAACAAGCTCGCCACCGTGGCAGTGGCGATCCACATCCGCAAAGCCAGCTGAGGGGCACA
GACAGAGGGATGAGGAAGAGAAGACGTCTCGACAGAGGAGGGCGTCCTGGAGAATCCATTTTCTCCTTCAGCA
CTTCTGTGCTGGTCCTGACCCGAGCCCAACCCGCACTAGGGAAGTGGAGCCCTCACGAGTAAGAAATT
TTATAGATCCTAATAGGCAGTCCGTAATTAGGCAGCTAAAGTTTTTCTCTAGGCATTTATCGAAGCGCTCCAG
GCCATTTTCAAAGATCACTGGAAGACTTCTATTTCTCTAACATGTAGGTCGAGTCGGCCGAGGAGTTGAGCCCTGAG
TGAGATCCTCTCCATCGGGGACTCTGCTGCTCCCCCTCTATCTGTTTTGGAAGATCAGCATCTTCCGCAGGGACTCA
GCCATGCTGACGCTCTGAGGCAAGTGGGAGCTGCTGCACTGGTCTCCTGACCTACCCCTCTGTGTTATAACT
CTACGCTATGAGCCAGAAGAGCGCTCAGGCTACTGCGAAGACGGGGAAAGCCAGATCAGGCTGGAGGAGATCGCTCTCTT
GTGACTTTTCTTTTGAAAATACAAATAAATCTGTTTATACCTTTGAAAAAAATAATTTAAAAATAGAATATATGATTAT
GCACTAGCTACGTCCCAACATATTGAGTTTTTCTCTTCTGTGATAGTTAATCTCAAAAAACAGCATTTTGAGATCAG
GTATCATTTGATTGGTTTACAGCTATGCATTCCACACAGAAATTTCCAGCCAAGGTGTTGTGCCCATAAGATC
ATATGTGGCTAGAAATTTCTCTGCTACCTAATTAGCAGCTTCTGTTGACCTTGTTATGATAGGCTAGCTAAATAT
GTCTTTTGTATCTTTAGCTTTTTAATAAGAATTTTTATTTAATTTTTAATTTTAAA
FIGURE 85

AGTCTAGCAGGAAGGAGAGGAGCTTTCCCCAGAAAGCCTCCTTGGACCAGCCCCAGGCTC
CTGTGCTGGTTGCACGCGCAGGGCCTGTACTGACCACCTCCACTCCAGTGCCACTGGGCTGTAA
GAGGAATGCGGCCGTTGGGCGACCTGCTTGGCCTGGGAGCCCTCTCTTCTGGGTAGGGGCCA
GAAACCGCTCTGACCCACAGCTGTGGCCTCTGAGGAAGCCCGGAGGCTCATTGGCCCCGCC
ACGGTGACACGCCTCTGCCCAGTGGGAGCCAGCCACCCGAGACCTTCCTCC
GCCGCTGACACACATCCTCCTGGCGAGCTGACAGCAGCCGCTCACTCACTCGTGCTGCA
CCGCCAGGCTTTATGCACGACTCTACCCCCGTGCTGTGGTGAAGCAGGATGGCTCCACCATC
CACATCCCGTCAGGAGCCACCGCGCATGCTGGCGATGGCCATAGATCTGGACACCCTGT
CTCCTGAGGAGCGCCGGCCAGGCCTGGCAAGGCTGAGGCTCAGCTCCAGTGAGGAG
GTACGAGCAGGAGCTCAGTGATGACTTGCTCATGGGAGCGCTACCGACAGTTCTGGACCAGG
ACCAAGAAGGGTCCGTGGCTTCCAGCACCCTCGGGGACATTGCTAAGATGGGAGGGCTGTTC
TAATCACTCGTTCTTGAAGCTGC
MAAVGSLLGLAASSWLLGQNASDHLWLLRRKPRGSSCPGTGHQLCRLRQSTVKATGPALR
RLHTSSWRADSRSALTRVHRQAYARLYPVLLVKQDSTIHRYREPRRMLAMPIDDLTL
SPEERRARLRRKREAQLQSRKEYEQLDSDLHVERYRFQFWTRTKK
FIGURE 88

> Wed Mar 29 15:44:14 2000 DNA168061 [min]
> /usr/seqdb2/sst/DNA/Dnaseqs.min/ss.DNA168061
> subunit 1 of 1, 207 aa, 1 stop
> MW: 25219, pI: 8.36, NX(S/T): 0
MSTKPDQIKCLWLEILMGIFIAGTLSLDCNLLNVHLRRTWQNLRLSSMSNSFPVECL
RENIAFELPOEFLQYTPMKRDIKAFYEMSLOQAFNIFSQHTFKYWKERHLKIQIGLDQ
QAEYLNQCLEEDENEDMKEMKENEMKPEARVPQLSSLRLSRLRYFRIDNFLKEKYSDF
CAEIVRVEIRRCLYYFYKFTALFRKR
FIGURE 89

CTGGGACTTGGCTTTCTCCGGATAAGCGGCGCCACCAGGCACGGCTCACGAGATGACCCGTGCAGAGA
CTCTGTTGCGCGCGGTGCTGGGTGGCCCTGGTTCACTCATTCTCACAACCTGAGGCGGTGGGCTGA
TCACCTCAACTCGGCTGCGAGACCTGCGAGGTGGCGAGCCAGCGAGCCTGGGCTGACGGCAGA
GAGTGCTGCTGTGGCTGGACAGGAGGCCACCCGGAGGGCGGAGAGCGCCACCTTGGGAGCCAGAGG
CAGGTGGACGCACATGACTGAGGCGGCTGGGCTGCTCCAGGAGCCAGCCGAGGCTTTCAAG
AGTCCCGAGGCAACCCGTCACAATGCACTGCTCCAGCATGTCGCGCCTGCAACCTGCTGCGC
GGCCGCTCACCCGACAGCCAGGCTCCACCTCTCCTGGGCTGGGGCTTGCGCGCTGCCCTGGTGA
TCACCCGACGCCCGTGGCTGGGAGGAGGACATGGCCGCTGCGATTCCCAACTGCGAGTGGTGT
TCCTGGGCTCATGGGCTTCGTGAATTTTTCAAGAATTTGGCCGATAACACCACCTGTCCTGGTC
CTGCTACCTGAAACATTTGGCGGCTGCGCTCCTGCGGACAGGCTGCGGCAAGCCATGCTACTCTG
AACATTCTCCACAAGAGGAGGAAGCTGACATGCAGGCCCGGGTGAGTCTGATCAGACGGCTGCC
TGACACCGCCTTTCCGCGTGGGCTGCAATGACTACGTGAGTCACCAGTGGA
CCTTCTCAACGCGTCCATCAACCGCACACCTGCTATGCTGGAAACAGCCCTAGAAACCAAGGAC
TCCACACAAGTCACCTCCCTGCTGTGGACAGGCAACGGGATGAGCTGGGTTGACCTCT
GGCCATGTGCGGCAGACATGTGCGTGGTTACTGTTATGTGGTGTCATATGCTGCTAGTG
CGTGGGCGAACCCTCGTCTGCGCTCAGC
FIGURE 90

important features of the protein:
Signal peptide:
Amino acids 1-25

Transmembrane domains:
Amino acids 105-125;139-157;169-188

N-glycosylation site:
Amino acids 164-168

cAMP- and cGMP-dependent protein kinase phosphorylation site:
Amino acids 39-43

Tyrosine kinase phosphorylation site:
Amino acids 214-222

N-myristoylation sites:
Amino acids 44-50;62-68;66-72;79-85

Amidation site:
Amino acids 37-41
FIGURE 92

>`</usr/seqdb2/sst/DNA/Dnaseqs.min/ss.DNA255255
>>subunit 1 of 1, 460 aa, 1 stop
>>MW: 49631, pI: 6.01, NX(S/T): 3
MAPGRAVACLILLAAAGLGVEAEQPGPLAPSEDVLSVPNGNLSSLQAQLQHLEQMGAA
RGVPEPGQLHFNQCTAAAEEFSLHGFSNATQITSSKFSCVCPAVLQQNLNFPCEDRPKHK
TPSHSEVWGYGFLSVTIINLASLLGILTPLIKSYFPKILTFFVGLAGTILFSNAIPQ
LIFEOAGFDPKVDYVEKAVAVFQGFFYLFFERMLKMLKTYQGCNTTHFGNDNFPGQE
KTHQPKALPAINGVTCYANPAVTEANHIGNFDDNVSVSLOQDKKPESSCTCLKGPKLSE
GTIAWMTLCALHNFIDGLAIGASCSTLSLLQGLSTSSAILCEEFPHELGDVILLNAGM
STRQALFNFSLACSVCYGLAFGILVGNNFAPNIFALAGGMFLYISLADMFPEDMNDMLR
EKVTGRKTDFTFMIONQAGMFTAILLLITYAGEIELE
FIGURE 93

AGAAACCGTTGATGGGACTGGAAGAAACCAGAATTAACCTTCTTGGAGCTTTCTGAGGACT
CAGCTGAAACCACCGGCAAGTGGCAACCCATTGAGATTTTCATGGTTAGCTGGCTGCTGATC
GGAGCATTCAGAATCTTGGTCTGGCCATGATGTCATCTGCTTCGCCAGAGGAGGGCAACCC
CCACCACCAGGGGACAGGATAGCTTGGAGGAAACATCTACAGGGAGAAACATCAAGTTT
CGGTAGCTATCAAGATCTTTGGTCTGGCCATGATGTCATCTGCTTCGCCAGAGGAGGGCAACCC
CTGTTGGTTTCCATATTCTTCGTCCTCAGGCAAACAGGCACTTTAAATCCTGCTTCCTACGAG
TGGTAGTGGGACAAAAATAATATAACACAAAGAAGTTATGTTTATTTATCTTATTATCATGAT
TCACCTTATACCCAGGAACTGCTATACAGCCAAAGCCAGCCTGCTGGAGTTCTCTCTCCTG
ATGCTGATTCTGCACTTCTGTCGAATTCTTGCTCTAGCTGTGCTCACTTGCTGTGCCTGGCTG
AAACAGCCATTACTCTGAGCTCTCCCTGGGTTGAGTGTGCCTGGCCCTAGCATTAGAGAAACCA
CCACCACATCCAGGTACACATGUAGGCGCCATTTGCTTAGGAAAGCAGAGGAGGGCAACCC
AAGCAGATGGAATTTGGATAGCTGATAAGCTGAAAAATTCTAGTTAGTAAATAGTAAA
TCATGAGAAATATCAACTGATTCAATCTTCATTTGTCAAGGGAATGAGACTGCG
GAAGTTAAATGACTGGCTTGCAATTATGCTATAGTTTGCTGTGCTTTGCTGAGACACTAG
ACCTGCTGTGCTCTCCCTTAGGGAAGAAACAAATTTCTGGCCCAACACACTAGTCCTCTTTA
AATGAGTTCTGGTAAAGGCAATTTACACGTAACCTGGAATCGGTGACAGCTTATG
CTCTGCAATTGCTGCTGGGACTTTAAAATTCGTTGCGCTTTAAGGCTATATTTAAAA
TGTATTTGGGAATCCAAAAAAAAAAAAAAAA
FIGURE 94

>>>subunit 1 of 1, 225 aa, 1 stop
>>>MW: 24317, pI: 8.07, NX(S/T): 3
MTSQPVPNETIVLPSUVINFSQAEKPEPTMQGQDSSLKHLHAIEIKVIGTIQILCGMV
SLGIILASASFSPNFTQVTSTLLNSAYPFIGPPFFIISGSLSIATEKRLTKLLVHSSLVG
SILSALSALVGFILSVKQATLNPASLQCELDKNNIPRSTYVSYPYHDSLYTDCYTAKA
SLAGSLSILMLICTLLEFCALVLTAVLRWKQAYSDFPGVSVLAGFT
FIGURE 96

Signal sequence:
amino acids 1-18

Transmembrane domain:
amino acids 341-359

N-glycosylation site.
amino acids 73-77, 92-96, 117-121, 153-157, 189-193, 204-208, 276-280, 308-312

Casein kinase II phosphorylation site.

Tyrosine kinase phosphorylation site.
amino acids 272-280

N-myristoylation site.

Prokaryotic membrane lipoprotein lipid attachment site.
amino acids 7-18