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(54) BITTER TASTE RECEPTORS

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(57) ABSTRACT

The present invention relates to bitter-taste receptors and their role in bitter taste transduction. The invention also relates to assays for screening molecules that modulate, e.g. suppress or block, bitter taste transduction, or enhance bitter taste response.

BITTER TASTE RECEPTORS

CROSS-REFERENCE TO RELATED APPLICATION

[0001] This application is a divisional application of copending application Ser. No. 10/528,630, filed Mar. 22, 2005; which is a National Stage Application of international Application Number PCT/EP2003/010691, filed Sep. 25, 2003; which claims the benefit of U.S. Provisional Application Ser. No. 60/413,298, filed Sep. 25, 2002, all of which are incorporated herein in their entirety.

BACKGROUND OF THE INVENTION

[0002] Investigators have recently turned their attention to understanding the biological mechanisms of taste, and in particular bitter taste. For a review of the literature see, for example, *Science* 291, 1557-1560. (2001); *Cell* 100, 607-610 (2000); *Neuron* 25, 507-510 (2000); *Nature* 413, 219-225. (2001); and *J Biol. Chem.* 277, 1-4 (2001).

[0003] Bitter taste is aversive, and as such provides humans with a mechanism of protection against poisonous substances, which are generally bitter-tasting compounds. More subtly, bitter-tastants also affect the palatability of food, beverages, thereby influencing human nutritional habits as is more fully discussed by Drewnowski in "The Science and Complexity of Bitter Taste", *Nutr. Rev.* 59, 163-169 (2001). They also affect the palatability of other ingestibles such as orally administered pharmaceuticals and nutraceuticals. Understanding the mechanism of bitter taste transduction has implications for the food and pharmaceutical industries. If the bitter taste transduction pathway can be manipulated, it may be possible to suppress or eliminate bitter taste to render foods more palatable and increase patient compliance in oral pharmaceutics.

[0004] Taste transduction involves the interaction of molecules, i.e., tastants with taste receptor-expressing cells which reside in the taste buds located in the papillae of the tongue. Taste buds relay information to the brain on the nutrient content of food and the presence of poisons. Recent advances in biochemical and physiological studies have enabled researchers to conclude that bitter taste transduction is mediated by so-called G-protein coupled receptors (GPCRs). GPCRs are 7 transmembrane domain cell surface proteins that amplify signals generated at a cell surface when the receptor interacts with a ligand (a tastant) whereupon they activate heterotrimeric G-proteins. The G-proteins are protein complexes that are composed of alpha and beta-gamma subunits. They are usually referred to by their alpha subunits and classified generally into 4 groups: $G_{alpha\ s,\ i,\ q\ and\ 12}.$ The G_{alpha q} type couple with GPCRs to activate phospholipase C which leads to the increase in cellular Ca²⁺. There are many G_a-type G-proteins that are promiscuous and can couple to GPCRs, including taste receptors, and these so-called "promiscuous" G-proteins are well known to the man skilled in the art. These G-proteins dissociate into alpha and betagamma subunits upon activation, resulting in a complex cascade of cellular events that results in the cell producing cell messengers, such as calcium ions, that enable the cells to send a signal to the brain indicating a bitter response.

[0005] There is also anatomical evidence that GPCRs mediate bitter taste transduction: clusters of these receptors are found in mammalian taste cells containing gustducin. Gustducin is a G-protein subunit that is implicated in the

perception of bitter taste in mammals, see for example Chandrashekar, J. et al., *Cell* 100, 703-711 (2000); Matsunami H. et al., *Nature* 404, 601-604 (2000); or Adler E. et al., *Cell* 100, 693-702 (2000). cDNAs encoding such GPCRs have been identified, isolated, and used as templates to compare with DNA libraries using in-silico data-mining techniques to identify other related receptors. In this manner it has been possible to identify a family of related receptors, the so-called T2R family of receptors, that have been putatively assigned as bitter receptors.

[0006] To-date, however, it is not clear as to whether all the bitter taste receptors have been discovered. Further, of those that have been discovered, many have not been matched, or paired, with ligands, and applicant is aware of very few published studies wherein rigorous matching has been undertaken. Chandrashekar, J. et al. in Cell 100, 703-711 (2000), has expressed a human T2R receptor, the so-called hT2R4 receptor, in heterologous systems and looked at the in vitro response of this receptor. They found that it provided a response to the bitter compounds denatonium and 6-n-propyl-2-thiouracil. However, the concentrations of bitter tastants needed to activate the hT2R4 receptor were two orders of magnitude higher than the thresholds reported in human taste studies, and so it is not clear that the protein encoded by the hT2R4 gene is a functional bitter receptor. The authors of the Chandrashekar et al. article also looked at a number of mouse T2R receptors with a range of stock bittertasting chemicals of disparate chemical structure. However, no study has looked at receptor responses to bitter ligands that are problematic in the food and pharmaceutical industries, and means of suppressing the bitter response to these ligands. [0007] The universe of compounds that provoke a bitter

[0007] The universe of compounds that provoke a bitter response in humans is structurally very diverse. Therefore, if research into bitter receptors is to be of any practical significance to the food and pharmaceutical industries, all bitter receptors will need to be identified, and once identified, there has to be a rigorous understanding of how specific receptors are matched to particular structural classes of bitter compounds. Unfortunately, although much basic research has been conducted in the area of bitter taste receptors, there are potentially many more bitter receptors to be discovered, and little is still known as to whether the known members of the human T2R family of bitter receptors actually respond to bitter tastants, and if so what, if any, specificity they show to ligand substructures.

BRIEF SUMMARY OF THE INVENTION

[0008] The present invention relates to bitter-taste receptors and their role in bitter taste transduction. The invention also relates to assays for screening molecules that modulate, e.g. suppress or block bitter taste transduction, or enhance bitter taste response.

DETAILED DESCRIPTION OF THE INVENTION

[0009] Unless otherwise defined, all technical and scientific terms used herein have the same meaning as commonly understood by one of ordinary skill in the art to which this invention pertains. In case of conflict, the present document, including definitions, will control. Preferred methods and materials are described below, although methods and materials similar or equivalent to those described herein can be used

in the practice or testing of the present invention. All publications, patent applications, patents and other references mentioned herein are incorporated by reference in their entirety. The materials, methods, and examples disclosed herein are illustrative only and not intended to be limiting.

[0010] Other features and advantages of the invention, e.g., screening for compounds that inhibit bitter taste, will be apparent from the following description, from the drawings and from the claims.

[0011] Surprisingly, applicant has now found a new group of putative bitter taste receptors, and in respect of certain known bitter receptors, applicant has found that they respond with specificity towards classes of bitter compounds that are important in the food and pharmaceutical industries.

[0012] In a first aspect of the invention there is provided a new group of putative bitter receptors. The genetics of bitter tasting has been extensively studied in mice and rats. Therefore, applicant compared the nucleotide sequences encoding polypeptides previously proposed to be bitter receptors with publicly available human nucleic acid sequences in the NCBI database using the BLAST® search methodology (Parameters: Expect=0.01, Filter=default). Surprisingly, the search identified 24 DNA sequences (from human chromosomes 5, 7, and 12) that, because of their homology to a mouse nucleic acid sequence that encodes a polypeptide (T2R5) previously designated as a bitter receptor, we designated as bitter receptor-encoding. Bitter taste receptors were originally assigned identifiers starting with the three characters "T2R" (identifying the receptor family) followed by a number (e.g., 1, 2, 3, etc.) that identifies a particular receptor, e.g., T2R5. More recently a different system has been used in which the identifiers start with five characters "TAS2R" (identifying the receptor family) followed, as previously, by a number (e.g., 1, 2, 3, etc.) that identifies a particular receptor, e.g., TAS2R5. A lower case letter in front of the identifier indicates the species of the receptor (e.g., "h" for human, "r" for rat, and "m" for mouse). Thus, for example, mTAS2R5 is a mouse bitter receptor and hTAS2R2 is a human bitter receptor. For consistency the new TAS2R identifier system is used throughout the rest of this application.

[0013] Of the 24 coding sequences identified by the search, 12 are believed to be novel; the polypeptides encoded by these novel sequences are designated hTAS2R38-41, and 43-50. The DNA sequences encoding the polypeptides are assigned SEQ ID NOs: 2 (hTAS2R38), 4 (hTAS2R39), 6 (hTAS2R40), 8 (hTAS2R41), 10 (hTAS2R43), 12 (hTAS2R44), 14(hTAS2R45), 16 (hTAS2R46), 18 (hTAS2R47), 20 (hTAS2R48), 22 (hTAS2R49), and 24 (hTAS2R50), respectively, and the amino acid sequences of the polypeptides are assigned SEQ ID NOs: 1, 3, 5, 7, 9, 11, 13, 15, 17, 19, 21, and 23, respectively.

[0014] The 12 additional (possibly novel) sequences identified by the search encode polypeptides, which are designated hTAS2R1, 4, 5, 7-10, 13, 14, 16, 3, 42 and 60. The DNA sequences encoding the polypeptides are assigned SEQ ID NOs: 26 (hTAS2R1), 28 (hTAS2R4), 30 (hTAS2R5), 32 (hTAS2R7), 34 (hTAS2R8), 36 (hTAS2R9), 38 (hTAS2R10), 40 (hTAS2R13), 42 (hTAS2R14), 44 (hTAS2R16), 46 (hTAS2R3), 48 (hTAS2R42), and 50 (hTAS2R60), respectively, and the amino acid sequences of the polypeptides are assigned SEQ ID NOs: 25, 27, 29, 31, 33, 35, 37, 39, 41, 43, 45, 47 and 49, respectively.

[0015] Thus, one aspect of the present invention is a polynucleotide selected from the group consisting of

[0016] (a) polynucleotides encoding at least the mature form of the polypeptide having the deduced amino acid sequence as shown in SEQ ID NOs 1, 3, 5, 7, 9, 11, 13, 15, 17, 19, 21, and 23;

[0017] (b) polynucleotides having the coding sequence, as shown in SEQ ID NOs: 2, 4, 6, 8, 10, 12, 14, 16, 18, 20, 22, and 24 encoding at least the mature form of the polypeptide:

[0018] (c) polynucleotides encoding a fragment or derivative of a polypeptide encoded by a polynucleotide of any one of (a) to (b), wherein in said derivative one or more amino acid residues are conservatively substituted compared to said polypeptide, and said fragment or derivative has bitter substance binding activity;

[0019] (d) polynucleotides which are at least 50% identical to a polynucleotide as defined in any one of (a) to (c) and which code for a polypeptide having bitter substance binding activity; and

[0020] (e) polynucleotides the complementary strand of which hybridizes, preferably under stringent conditions to a polynucleotide as defined in any one of (a) to (d) and which code for a polypeptide having bitter substance binding activity;

or the complementary strand of such a polynucleotide.

[0021] A polypeptide that exhibits bitter substance binding activity is a polypeptide that has at least 20% (e.g., at least: 20%; 30%; 40%; 50%; 60%; 70%; 80%; 90%; 95%; 98%; 99%; 99.5%; or 100% or even more) of the ability of the respective full-length TAS2R to bind to a given bitter substance. Binding assays and bitter substances are described herein below.

[0022] In a preferred embodiment the polynucleotide of the present invention encodes a polypeptide that still exhibits essentially the same activity as the respective mature bitter taste receptor, i.e. has "bitter taste receptor activity". Preferably the polypeptide has at least 20% (e.g., at least: 20%; 30%; 40%; 50%; 60%; 70%; 80%; 90%; 95%; 98%; 99%; 99.5%; or 100% or even more) of the ability of the respective full-length TAS2R to release intracellular calcium in a heterologous cell expression system like, for example, HEK293/ 15-cells, which stably express the alpha-subunit of promiscuous G-proteins, e.g. the mouse G₁₅ subunit, in response to bitter tastants, which is dependent on the expression of polypeptides encoded by the polynucleotides of the present invention. The amount of intracellular calcium release can be monitored by, for example, the in vitro FLIPR assay described herein below.

[0023] The TAS2R nucleic acid molecules of the invention can be DNA, cDNA, genomic DNA, synthetic DNA, or, RNA, and can be double-stranded or single-stranded, the sense and/or an antisense strand. Segments of these molecules are also considered within the scope of the invention, and can be produced by, for example, the polymerase chain reaction (PCR) or generated by treatment with one or more restriction endonucleases. A ribonucleic acid (RNA) molecule can be produced by in vitro transcription.

[0024] The polynucleotide molecules of the invention can contain naturally occurring sequences, or sequences that differ from those that occur naturally, but, due to the degeneracy of the genetic code, encode the same polypeptide (for example, the polypeptides with SEQ ID NOs: 1, 3, 5, 7, 9, 11, 13, 15, 17, 19, 21, and 23). In addition, these nucleic acid

molecules are not limited to coding sequences, e.g., they can include some or all of the non-coding sequences that lie upstream or downstream from a coding sequence.

[0025] The polynucleotide molecules of the invention can be synthesized in vitro (for example, by phosphoramidite-based synthesis) or obtained from a cell, such as the cell of a bacteria mammal. The nucleic acids can be those of a human but also derived from a non-human primate, mouse, rat, guinea pig, cow, sheep, horse, pig, rabbit, dog, or cat as long as they fulfill the criteria set out above. Combinations or modifications of the nucleotides within these types of nucleic acids are also encompassed.

[0026] In addition, the isolated nucleic acid molecules of the invention encompass segments that are not found as such in the natural state. Thus, the invention encompasses recombinant nucleic acid molecules incorporated into a vector (for example, a plasmid or viral vector) or into the genome of a heterologous cell (or the genome of a homologous cell, at a position other than the natural chromosomal location). Recombinant nucleic acid molecules and uses therefore are discussed further below.

[0027] A polynucleotide belonging to a family of any of the TAS2R disclosed herein or a protein can be identified based on its similarity to the relevant TAS2R gene or protein, respectively. For example, the identification can be based on sequence identity. In certain preferred embodiments the invention features isolated nucleic acid molecules which are at least 50% (or 55%, 65%, 75%, 85°/a, 95%, or 98%) identical to: (a) a nucleic acid molecule that encodes the polypeptide of SEQ ID NOs:1, 3, 5, 7, 9, 11, 13, 15, 17, 19, 21, or 23; (b) the nucleotide sequence of SEQ ID NOs: 2, 4, 6, 8, 10, 12, 14, 16, 18, 20, 22, or 24; and (c) a nucleic acid molecule which includes a segment of at least 30 (e.g., at least 30, 40, 50, 60, 80, 100, 125, 150, 175, 200, 250, 300, 400, 500, 600, 700, 800, 850, 900, 950, 1000, or 1010) nucleotides of SEQ ID NOs: 2, 4, 6, 8, 10, 12, 14, 16, 18, 20, 22, or 24.

[0028] The determination of percent identity between two sequences is accomplished using the mathematical algorithm of Karlin and Altschul, Proc. Natl. Acad. Sci. USA 90, 5873-5877, 1993. Such an algorithm is incorporated into the BLASTN and BLASTP programs of Altschul et al. (1990) J. Mol. Biol. 215, 403-410. BLAST nucleotide searches are performed with the BLASTN program, score=100, wordlength=12, to obtain nucleotide sequences homologous to HIN-1-encoding nucleic acids. BLAST protein searches are performed with the BLASTP program, score=50, wordlength=3, to obtain amino acid sequences homologous to the TAS2R polypeptide. To obtain gapped alignments for comparative purposes, Gapped BLAST is utilized as described in Altschul et al. (1997) Nucleic Acids Res. 25, 3389-3402. When utilizing BLAST and Gapped BLAST programs, the default parameters of the respective programs.

[0029] Hybridization can also be used as a measure of homology between two nucleic acid sequences. A nucleic acid sequence encoding any of the TAS2R disclosed herein, or a portion thereof, can be used as a hybridization probe according to standard hybridization techniques. The hybridization of a TAS2R probe to DNA or RNA from a test source (e.g., a mammalian cell) is an indication of the presence of the relevant TAS2R DNA or RNA in the test source. Hybridization conditions are known to those skilled in the art and can be found in Current Protocols in Molecular Biology, John Wiley & Sons, N.Y., 6.3.1-6.3.6, 1991. Moderate hybridization conditions are defined as equivalent to hybridization in 2×

sodium chloride/sodium citrate (SSC) at 30° C., followed by a wash in 1 X SSC, 0.1% SDS at 50° C. Highly stringent conditions are defined as equivalent to hybridization in 6× sodium chloride/sodium citrate (SSC) at 45° C., followed by a wash in 0.2×SSC, 0.1% SDS at 65° C.

[0030] An "isolated DNA" is either (1) a DNA that contains sequence not identical to that of any naturally occurring sequence, or (2), in the context of a DNA with a naturallyoccurring sequence (e.g., a cDNA or genomic DNA), a DNA free of at least one of the genes that flank the gene containing the DNA of interest in the genome of the organism in which the gene containing the DNA of interest naturally occurs. The term therefore includes a recombinant DNA incorporated into a vector, into an autonomously replicating plasmid or virus, or into the genomic DNA of a prokaryote or eukaryote. The term also includes a separate molecule such as a cDNA where the corresponding genomic DNA has introns and therefore a different sequence; a genomic fragment that lacks at least one of the flanking genes; a fragment of cDNA or genomic DNA produced by polymerase chain reaction (PCR) and that lacks at least one of the flanking genes; a restriction fragment that lacks at least one of the flanking genes; a DNA encoding a non-naturally occurring protein such as a fusion protein, mutein, or fragment of a given protein; and a nucleic acid which is a degenerate variant of a cDNA or a naturally occurring nucleic acid. In addition, it includes a recombinant nucleotide sequence that is part of a hybrid gene, i.e., a gene encoding a non-naturally occurring fusion protein. It will be apparent from the foregoing that isolated DNA does not mean a DNA present among hundreds to millions of other DNA molecules within, for example, cDNA or genomic DNA libraries or genomic DNA restriction digests in, for example, a restriction digest reaction mixture or an electrophoretic gel

[0031] A further aspect of the present invention is a vector containing the polynucleotide(s) of the present invention or a protein encoded by a polynucleotide of the present invention. The term "vector" refers to a protein or a polynucleotide or a mixture thereof which is capable of being introduced or of introducing the proteins and/or nucleic acid comprised into a cell. It is preferred that the proteins encoded by the introduced polynucleotide are expressed within the cell upon introduction of the vector.

[0032] In a preferred embodiment the vector of the present invention comprises plasmids, phagemids, phages, cosmids, artificial mammalian chromosomes, knock-out or knock-in constructs, viruses, in particular adenoviruses, vaccinia viruses, attenuated vaccinia viruses, canary pox viruses, lentivirus (Chang, L. J. and Gay, E. E. (20001) Curr. Gene Therap. 1:237-251), herpes viruses, in particular Herpes simplex virus (HSV-1, Carlezon, W. A. et al. (2000) Crit. Rev. Neurobiol.), baculovirus, retrovirus, adeno-associated-virus (AAV, Carter, P. J. and Samulski, R. J. (2000) J. Mol. Med. 6:17-27), rhinovirus, human immune deficiency virus (HIV), filovirus and engineered versions thereof (see, for example, Cobinger G. P. et al (2001) Nat. Biotechnol. 19:225-30), virosomes, "naked" DNA liposomes, and nucleic acid coated particles, in particular gold spheres. Particularly preferred are viral vectors like adenoviral vectors or retroviral vectors (Lindemann et al. (1997) Mol. Med. 3:466-76 and Springer et al. (1998) Mol. Cell. 2:549-58). Liposomes are usually small unilamellar or multilamellar vesicles made of cationic, neutral and/or anionic lipids, for example, by ultrasound treatment of liposomal suspensions. The DNA can, for example,

be ionically bound to the surface of the liposomes or internally enclosed in the liposome. Suitable lipid mixtures are known in the art and comprise, for example, DOTMA (1,2-Dioleyloxpropyl-3-trimethylammoniumbromid) and DPOE (Dioleoylphosphatidyl-ethanolamin) which both have been used on a variety of cell lines.

[0033] Nucleic acid coated particles are another means for the introduction of nucleic acids into cells using so called "gene guns", which allow the mechanical introduction of particles into the cells. Preferably the particles itself are inert, and therefore, are in a preferred embodiment made out of gold spheres.

[0034] In a further aspect the polynucleotide of the present invention is operatively linked to expression control sequences allowing expression in prokaryotic and/or eukaryotic host cells. The transcriptional/translational regulatory elements referred to above include but are not limited to inducible and non-inducible, constitutive, cell cycle regulated, metabolically regulated promoters, enhancers, operators, silencers, repressors and other elements that are known to those skilled in the art and that drive or otherwise regulate gene expression. Such regulatory elements include but are not limited to regulatory elements directing constitutive expression like, for example, promoters transcribed by RNA polymerase III like, e.g., promoters for the snRNA U6 or scRNA 7SK gene, the cytomegalovirus hCMV immediate early gene, the early or late promoters of SV40 adenovirus, viral promoter and activator sequences derived from, e.g., NBV, HCV, HSV, HPV, EBV, HTLV, MMTV or HIV; which allow inducible expression like, for example, CUP-1 promoter, the tetrepressor as employed, for example, in the tet-on or tet-off systems, the lac system, the trp system; regulatory elements directing tissue specific expression, preferably taste bud specific expression, e.g., PLCβ2 promoter or gust-ducin promoter, regulatory elements directing cell cycle specific expression like, for example, cdc2, cdc25C or cyclin A; or the TAC system, the TRC system, the major operator and promoter regions of phage A, the control regions of fd coat protein, the promoter for 3-phosphoglycerate kinase, the promoters of acid phosphatase, and the promoters of the yeast aor a-mating factors.

[0035] As used herein, "operatively linked" means incorporated into a genetic construct so that expression control sequences effectively control expression of a coding sequence of interest.

[0036] Similarly, the polynucleotides of the present invention can form part of a hybrid gene encoding additional polypeptide sequences, for example, a sequence that functions as a marker or reporter. Examples of marker and reporter genes include β-lactamase, chloramphenicol acetyltransferase (CAT), adenosine deaminase (ADA), aminoglycoside phosphotransferase (neo^r, G418^r), dihydrofolate reductase (DHFR), hygromycin-B-phosphotransferase (HPH), thymidine kinase (TK), lacZ (encoding β-galactosidase), and xanthine guanine phosphoribo-syltransferase (XGPRT). As with many of the standard procedures associated with the practice of the invention, skilled artisans will be aware of additional useful reagents, for example, additional sequences that can serve the function of a marker or reporter. Generally, the hybrid polypeptide will include a first portion and a second portion; the first portion being a TAS2R polypeptide and the second portion being, for example, the reporter described above or an Ig constant region or part of an Ig constant region, e.g., the CH2 and CH3 domains of IgG2a heavy chain. Other hybrids could include an antigenic tag or His tag to facilitate purification and/or detection. Recombinant nucleic acid molecules can also contain a polynucleotide sequence encoding a TAS2R polypeptide operatively linked to a heterologous signal sequence. Such signal sequences can direct the protein to different compartments within the cell and are well known to someone of skill in the art. A preferred signal sequence is a sequence that facilitates secretion of the resulting protein.

[0037] Another aspect of the present invention is a host cell genetically engineered with the polynucleotide or the vector as outlined above. The host cells that may be used for purposes of the invention include but are not limited to prokaryotic cells such as bacteria (for example, E. coli and B. subtilis), which can be transformed with, for example, recombinant bacteriophage DNA, plasmid DNA, or cosmid DNA expression vectors containing the polynucleotide molecules of the invention; simple eukaryotic cells like yeast (for example, Saccharomyces and Pichia), which can be transformed with, for example, recombinant yeast expression vectors containing the polynucleotide molecule of the invention; insect cell systems like, for example, Sf9 of Hi5 cells, which can be infected with, for example, recombinant virus expression vectors (for example, baculovirus) containing the polynucleotide molecules of the invention; Xenopus oocytes, which can be injected with, for example, plasmids; plant cell systems, which can be infected with, for example, recombinant virus expression vectors (for example, cauliflower mosaic virus (CaMV) or tobacco mosaic virus (TMV)) or transformed with recombinant plasmid expression vectors (for example, Ti plasmid) containing a TAS2R nucleotide sequence; or mammalian cell systems (for example, COS, CHO, BHK, HEK293, VERO, HeLa, MDCK, Wi38, and NIH 3T3 cells), which can be transformed with recombinant expression constructs containing, for example, promoters derived, for example, from the genome of mammalian cells (for example, the metallothionein promoter) from mammalian viruses (for example, the adenovirus late promoter and the vaccinia virus 7.5K promoter) or from bacterial cells (for example, the tet-repressor binding its employed in the tet-on and tetoff systems). Also useful as host cells are primary or secondary cells obtained directly from a mammal and transfected with a plasmid vector or infected with a viral vector. Depending on the host cell and the respective vector used to introduce the polynucleotide of the invention the polynucleotide can integrate, for example, into the chromosome or the mitochondrial DNA or can be maintained extrachromosomally like, for example, episomally or can be only transiently comprised in the cells.

[0038] In a preferred embodiment, the TAS2R encoded by the polynucleotides of the present invention and which are expressed by such cells are functional, i.e., upon binding to one or more bitter molecules they trigger an activation pathway in the cell. The cells are preferably mammalian (e.g., human, non-human primate, horse, bovine, sheep, goat, pig, dog, cat, goat, rabbit mouse, rat, guinea pig, hamster, or gerbil) cells, insect cells, bacterial cells, or fungal (including yeast) cells.

[0039] A further aspect of the present invention is a transgenic non-human animal containing a polynucleotide, a vector and/or a host cell as described above. The animal can be a mosaic animal, which means that only part of the cells making up the body comprise polynucleotides, vectors, and/or cells of the present invention or the animal can be a transgenic animal which means that all cells of the animal comprise the

polynucleotides and/or vectors of the present invention or are derived from a cell of the present invention. Mosaic or transgenic animals can be either homo- or heterozygous with respect to the polynucleotides of the pre-sent invention contained in the cell. In a preferred embodiment the transgenic animals are either homo- or heterozygous knock-out or knock-in animals with respect to the genes which code for the proteins of the present invention. The animals can in principal be any animal, preferably, however, it is a mammal, selected from the group of non-human pimate horse, bovine, sheep, goat, pig, dog, cat, goat, rabbit, mouse, rat, guinea pig, hamster, or gerbil. Another aspect of the present invention is a process for producing a polypeptide encoded by a polynucleotide of the present invention comprising: culturing the host cell described above and recovering the polypeptide encoded by said polynucleotide. Preferred combinations of host cells and vectors are outlined above and further combination will be readily apparent to someone of skill in the art. Depending on the intended later use of the recovered peptide a suitable cell type can be chosen. Eukaryotic cells are preferably chosen, if it is desired that the proteins produced by the cells exhibit an essentially natural pattern of glycosylation and prokaryotic cells are chosen, if, for example, glycosylation or other modifications, which are normally introduced into proteins only in eukaryotic cells, are not desired or not needed.

[0040] A further aspect of the invention is a process for producing cells capable of expressing at least one of the bitter taste receptor polypeptides comprising genetically engineering cells in vitro with at least one of the vectors described above, wherein said bitter taste receptor polypeptide(s) is(are) encoded by a polynucleotide of the present invention. Another aspect of the invention is a polypeptide having the amino acid sequence encoded by a polynucleotide of the invention or obtainable by the process mentioned above. The polypeptides of the invention include all those disclosed herein and functional fragments of these polypeptides. "Polypeptide" and "protein" are used interchangeably and mean any peptidelinked chain of amino acids, regardless of length or posttranslational modification. As used herein, a functional fragment of a TAS2R is a fragment of the TAS2R that is shorter than the full-length TAS2R but that has at least 20% (e.g., at least: 20%; 30%; 40%; 50%; 60%; 70%; 80%; 90%; 95%; 98%; 99%; 99.5%; or 100% or even more) of the ability of the full-length TAS2R to bind to a bitter substance to which the full-length TAS2R binds. Binding assays and bitter substances are described herein. Further bitter substances can be identified by the binding assays and bitter taste receptor activity assays described herein. The polypeptides embraced by the invention also include fusion proteins that contain either a full-length TAS2R polypeptide or a functional fragment of it fused to an unrelated amino acid sequence. The unrelated sequences can be additional functional domains or signal peptides. Signal peptides are described in greater detail and exemplified below.

[0041] The polypeptides can be any of those described above but with not more than 50 (e.g., not more than: 50, 45, 40, 35, 30, 25, 20, 15, 14, 13, 12, 11, 10, nine, eight, seven, six, five, four, three, two, or one) conservative substitutions. Conservative substitutions typically include substitutions within the following groups: glycine and alanine; valine, isoleucine, and leucine; aspartic acid and glutainic acid; asparagine, glutamine, serine and threonine; lysine, histidine and arginine; and phenylalanine and tyrosine. All that is required of a polypeptide having one or more conservative substitutions is

that it has at least 20% (e.g., at least: 20%; 30%; 40%; 50%; 60%; 70%; 80%; 90%; 95%; 98%; 99%; 99.5%; or 100% or even more) of the ability of the wild-type, full-length TAS2R to bind to a bitter substance, preferably the ability to release intracellular calcium, when expressed in a cellular system.

[0042] The polypeptides can be purified from natural sources (e.g., blood, serum, plasma, tissues or cells such as normal tongue cells or any cell that naturally produces the relevant TAS2R polypeptides). Smaller peptides (less than 50 amino acids long) can also be conveniently synthesized by standard chemical means. In addition, both polypeptides and peptides can be produced by standard in vitro recombinant DNA techniques and in vivo transgenesis, using nucleotide sequences encoding the appropriate polypeptides or peptides. Methods well-known to those skilled in the art can be used to construct expression vectors containing relevant coding sequences and appropriate transcriptional/translational control signals. See, for example, the techniques described in Sambrook et al., Molecular Cloning: A Laboratory Manual (2nd Ed.) [Cold Spring Harbor Laboratory, N.Y., 1989], and Ausubel et al., Current Protocols in Molecular Biology [Green Publishing Associates and Wiley Interscience, N.Y., 1989].

[0043] Polypeptides and fragments of the invention also include those described above, but modified for in vivo use by the addition, at the amino- and/or carboxyl-terminal ends, of blocking agents to facilitate survival of the relevant polypeptide in vivo. This can be useful in those situations in which the peptide termini tend to be degraded by proteases prior to cellular uptake. Such blocking agents can include, without limitation, additional related or unrelated peptide sequences that can be attached to the amino and/or carboxyl terminal residues of the peptide to be administered. This can be done either chemically during the synthesis of the peptide or by recombinant DNA technology by methods familiar to artisans of average skill.

[0044] Alternatively, blocking agents such as pyroglutamic acid or other molecules known in the art can be attached to the amino and/or carboxyl terminal residues, or the amino group at the amino terminus or carboxyl group at the carboxyl terminus can be replaced with a different moiety. Likewise, the peptides can be covalently or noncovalently coupled to pharmaceutically acceptable "carrier" proteins prior to administration.

[0045] Also of interest are peptidomimetic compounds that are designed based upon the amino acid sequences of the functional peptides or peptide fragments. Peptidomimetic compounds are synthetic compounds having a three-dimensional conformation (i.e., a "peptide motif") that is substantially the same as the three-dimensional conformation of a selected peptide. The peptide motif provides the peptidomimetic compound with the ability to bind to a bitter compound in a manner qualitatively identical to that of the TAS2R functional fragment from which the peptidomimetic was derived. Peptidomimetic compounds can have additional characteristics that enhance their therapeutic utility, such as increased cell permeability and prolonged biological half-life.

[0046] The peptidomimetics typically have a backbone that is partially or completely nonpeptide, but with side groups that are identical to the side groups of the amino acid residues that occur in the peptide on which the peptidomimetic is based. Several types of chemical bonds, e.g., ester, thioester, thioamide, retroamide, reduced carbonyl, dimethylene and ketomethylene bonds, are known in the art to be generally

useful substitutes for peptide bonds in the construction of protease-resistant peptidomimetics.

[0047] The term "isolated" polypeptide or peptide fragment as used herein refers to a polypeptide or a peptide fragment which either has no naturally-occurring counterpart or has been separated or purified from components which naturally accompany it, e.g., in tissues such as tongue, pancreas, liver, spleen, ovary, testis, muscle, joint tissue, neural tissue, gastrointestinal tissue or tumor tissue, or body fluids such as blood, serum, or urine. Typically, the polypeptide or peptide fragment is considered "isolated" when it is at least 70%, by dry weight, free from the proteins and other naturally-occurring organic molecules with which it is naturally associated. Preferably, a preparation of a polypeptide (or peptide fragment thereof) of the invention is at least 80%, more preferably at least 90%, and most preferably at least 99%, by dry weight, the polypeptide (or the peptide fragment thereof), respectively, of the invention. Thus, for example, a preparation of polypeptide x is at least 80%, more preferably at least 90%, and most preferably at least 99%, by dry weight, polypeptide x. Since a polypeptide that is chemically synthesized is, by its nature, separated from the components that naturally accompany it, the synthetic polypeptide is "isolated.'

[0048] An isolated polypeptide (or peptide fragment) of the invention can be obtained, for example, by extraction from a natural source (e.g., from tissues or bodily fluids); by expression of a recombinant nucleic acid encoding the polypeptide; or by chemical synthesis. A polypeptide that is produced in a cellular system different from the source from which it naturally originates is "isolated," because it will necessarily be free of components which naturally accompany it. The degree of isolation or purity can be measured by any appropriate method, e.g., column chromatography, polyacrylamide gel electrophoresis, or HPLC analysis.

[0049] A further aspect of the invention is an antibody, which specifically binds to the polypeptide encoded by a polynucleotide of the invention or obtainable by the process mentioned above. The term "antibody" comprises monoclonal and polyclonal antibodies and binding fragments thereof, in particular Fe-fragments as well as so called "single-chain-antibodies" (Bird R. E. et al (1988) Science 242:423-6), chimeric, humanized, in particular CDR-grafted antibodies, and dia or tetrabodies (Holliger P. et al (1993) Proc. Natl. Acad. Sci. U.S.A. 90:6444-8). Also comprised are immunoglobulin like proteins that are selected through techniques including, for example, phage display to specifically bind to the polypeptides of the present invention. Preferred antibodies bind to the extracellular domain of bitter receptors and in particular to those domains responsible for binding to bitter tastants.

[0050] In yet another embodiment there is provided a molecule, or collections of molecules containing a molecule, that act to antagonise aforementioned receptors in particular the bitter taste response, and methods for screening for such molecules.

[0051] Therefore, a further aspect of the invention is a nucleic acid molecule which specifically hybridizes to a polynucleotide of the present invention. In particular this nucleic acid molecule is an inhibiting RNA. Preferred inhibiting RNAs are antisense constructs hybridizing to a polynucleotide of the present invention, RNAi, siRNA or a ribozyme. The design of such inhibiting RNAs would be readily apparent to someone of skill in the art.

[0052] Another type of antagonist/inhibitor against the polypeptides of the present invention is an antibody, which is preferably directed against the extracellular domain of the

respective bitter taste receptor and even more preferably binds to the site(s) of the receptor that interact(s) with the bitter substance(s) essentially without triggering the release of intracellular calcium. Further antagonists to the bitter taste response of a receptor are fragments of the receptor which have the capability to bind to the bitter substances as defined above. Such fragments can bind to the bitter substance and, thus, competitively antagonize the activity of the respective TAS2R. If such antagonists are, for example, employed within foodstuff to suppress the bitter taste of a specific bitter substance they might be exposed to a proteolytic environment and in this case the modifications of the polypeptides outlined above could be used to stabilize the competitive bitter receptor antagonist. However, various additional modifications, which stabilize such fragments will be readily apparent to the skilled person.

[0053] Antagonists and agonists of the bitter taste receptors described herein are of great importance for specific stimulation of a given bitter taste receptor or to antagonize it. The bitter taste response of the receptor is elicited by the specific binding of the respective bitter substance. Therefore, the present invention is also directed at a process for isolating a compound that binds to a polypeptide encoded by a polynucleotide selected from the group consisting of:

[0054] (a) polynucleotides encoding at least the mature form of the polypeptide having the deduced amino acid sequence as shown in SEQ ID NOs1, 3, 5, 7, 9, 11, 13, 15, 17, 19, 21, 23, 25, 27, 29, 31, 33, 35, 37, 39, 41, 43, 45, 47 and 49;

[0055] (b) polynucleotides having the coding sequence, as shown in SEQ ID NOs: 2, 4, 6, 8, 10, 12, 14, 16, 18, 20, 22, 24, 26, 28, 30, 32, 34, 36, 38, 40, 42, 44, 46, 48 and 50 encoding at least the mature form of the polypeptide;

[0056] (c) polynucleotides encoding a fragment or derivative of a polypeptide encoded by a polynucleotide of any one of (a) to (b), wherein in said derivative one or more amino acid residues are conservatively substituted compared to said polypeptide, and said fragment or derivative has bitter substance binding activity;

[0057] (d) polynucleotides which are at least 50% identical to a polynucleotide as defined in any one of (a) to (c) and which code for a polypeptide having bitter substance binding activity; and

[0058] (e) polynucleotides the complementary strand of which hybridizes, preferably under stringent conditions to a polynucleotide as defined in any one of (a) to (d) and which code for a polypeptide having bitter substance binding activity;

comprising:

[0059] (1) contacting said polypeptide or a host cell genetically engineered with said polynucleotide or with a vector containing said polynucleotide with a compound;

[0060] (2) detecting the presence of the compound which binds to said polypeptide; and

[0061] (3) determining whether the compound binds said polypeptide.

[0062] A polynucleotide employed in this process is in preferred embodiments of the invention at least 50% (or 55%, 65%, 75%, 85°/a, 95%, or 98%) identical to: (a) a nucleic acid molecule that encodes the polypeptide of SEQ ID NOs: 1, 3, 5, 7, 9, 11, 13, 15, 17, 19, 21, 23, 25, 27, 29, 31, 33, 35, 37, 39, 41, 43, 45, 47 or 49; (b) the nucleotide sequence of SEQ ID NOs: 2, 4, 6, 8, 10, 12, 14, 16, 18, 20, 22, 24, 26, 28, 30, 32, 34, 36, 38, 40, 42, 44, 46, 48 or 50; and has a length of at least

30 (e.g., at least 30, 40, 50, 60, 80, 100, 125, 150, 175, 200, 250, 300, 400, 500, 600, 700, 800, 850, 900, 950, 1000, or 1010) of the nucleotides of SEQ ID NOs: 2, 4, 6, 8, 10, 12, 14, 16, 18, 20, 22, 24, 26, 28, 30, 32, 34, 36, 38, 40, 42, 44, 46, or 50.

[0063] Furthermore for all of the above described hTAS2Rs, which can be employed in a process for isolating

binding compounds, with the exception of hTAS2R40, single nucleotide polymorphisms are known. 79 of these are listed in Table I below. 61 of those result in an amino acid change. Polynucleotides or polypeptides that differ from the respectively in SEQ ID 1-50 indicated sequences by the nucleotide and amino acid change as indicted in Table I can similarly be employed for the process of the present invention.

TABLE I

Gen +		Su	bstitution	Pos	sition	_
Acession			Amino	Base	Amino	Allelic
No.	Name of SNP	Base	acid	pair	acid	frequency
hTAS2R1	rs2234231	C/T	P/L	128	43	unknown
NM019599	rs41469	G/A	R/H	332	111	A 0.46/G 0.54
	rs223432	G/A	C/Y	422	141	unknown
	rs2234233	C/T	R/W S/S	616	206 225	C 0.87/T 0.13
	rs2234234 rs2234235	C/T T/C	5/5 L/L	675 850	284	unknown unknown
hTAS2R3	rs227009	C/T	G/G	807	369	unknown
NM016943	13227009	C/I	0/0	007	307	unknown
hTAS2R4	ss3181498	G/A	R/Q	8	3	unknown
NM016944	rs2233996	G/C	R/R	9	3	unknown
	rs2233997	A/C	Y/C	17	6	unknown
	rs2233998	T/C	F/S	20	7	unknown
	rs2233999 rs2234000	T/A C/T	F/L T/M	186 221	62 74	unknown C 0.94/T 0.56
	rs2234000	G/C	V/L	286	96	C 0.78/G 0.22
	rs2234002	G/A	S/N	512	171	A 0.78/G 0.22
	rs2234003	A/G	I/V	571	191	unknown
hTAS2R5	rs2234013	G/A	G/S	58	20	unknown
NM018980	rs2227264	G/T	S/I	77	26	unknown
	rs2234014	C/T	P/L	338	113	unknown
	rs2234015	G/A	R/Q	638	213	unknown
1	rs2234016	G/T	R/L	294	881	unknown
hTAS2R7 NM023919	rs3759251	A/T	T/S	787	263 276	A 0.97/T 0.03 unknown
NM023919	rs3759252 rs619381	C/A G/A	I/I M/I	828	304	
hTAS2R8	ss2391467	G/A	L/L	912 549	183	unknown unknown
NM023918	rs2537817	A/G	M/V	922	308	unknown
hTAS2R9	rs3741845	T/C	V/A	560	187	C 073/T 027
NM23917	rs3944035	C/T	L/F	910	304	unknown
INIVI23917	rs2159903	C/T	P/L	926	309	unknown
hTAS2R10	rs597468	C/T	T/M	467	156	unknown
NM23921	18377400	C/I	1/141	707	150	unknown
hTAS2R13	ss1478988	A/G	N/S	776	259	C 0.73/T 0.27
NM23920						
hTAS2R14	rs3741843	G/A	R/R	375	125	A 0.97/G 0.03
NM23922						
hTAS2R16	rs2233988	C/T	T/T	300	100	unknown
NM016945	rs2692396	G/C	V/V	303	101	unknown
	rs2233989	T/C	L/L	460	154	unknown
	rs846664	T/G	N/K	516	172	A 0.71/C 0.29
	rs860170	G/A	R/H	665	222	A 0.55/G 0.45
hTAS2R38	PTC Paper	G/A	V/I	886	296	G 0.38/A 0.62
AF494321	rs1726866	T/C	V/A	785	262	G 0.38/T 0.62
	rs713598	C/T	A/P	49	145	C 0.36/G 0.64
	hTAS2R38 SNP1	A/T	N/I	557	186	C 0.60/G 0.40
hTAS2R39	hTAS2R39 SNP1	A/AA	frameshift	967	323	unknown
AF494230	1404625	. /G	m /m	1.00		1
hTAS2R41	rs1404635	A/G	T/T	189	64	unknown
AF494232	hTAS2R41 SNP1	T/C	L/P	380	127	unknown
LTA COD 40	hTAS2R41 SNP2	A/G	S/S	885	295	unknown
hTAS2R42 AX097739	rs1650017 rs1669411	G/C	A/P	931	311	unknown
AA091139		T/C	N/N	930	310	unknown
	rs1669412 rs1451772	G/A A/G	R/Q Y/C	875 794	292 265	unknown unknown
		G/T				unknown
	rs1669413 rs1650019	A/G	G/W L/L	763 561	255 187	unknown
hTAS2R43	rs3759246	G/C	L/L R/T	893	298	unknown
AF494237	hTAS2R43 SNP1	C/G	S/W	104	35	unknown
14 12 Taul	hTAS2R43 SNP2	G/A	R/H	635	212	unknown
	hTAS2R43 SNP3	G/C	T/T	663	221	unknown
				303		

TABLE I-continued

Gen +		Sı	ubstitution	Pos	ition	_
Acession No.	Name of SNP	Base	Amino acid	Base pair	Amino acid	Allelic frequency
hTAS2R44	rs3759247	G/A	W/stop	900	300	unknown
AF494228	rs3759246	G/C	R/T	893	298	unknown
	hTAS2R44 SNP1	A/T	M/L	162	484	unknown
	hTAS2R44 SNP2	T/A	F/Y	869	290	unknown
	hTAS2R44 SNP3	G/A	V/M	899	297	unknown
hTAS2R45	rs3759247	A/G	G/stop	900	300	unknown
AF494226	rs3759246	G/C	R/T	893	298	unknown
	rs3759245	C/T	R/C	712	238	unknown
	rs3759244	T/C	F/L	703	235	unknown
hTAS2R46	rs2708381	G/A	W/stop	749	250	unknown
AF494227	rs2708380	T/A	L/M	682	228	unknown
	rs2598002	T/G	F/V	106	36	unknown
	hTAS2R46 SNP1	A/T	Q/H	888	296	unknown
	hTAS2R46 SNP2	A/G	M/V	889	297	unknown
	hTAS2R46 SNP3	T/C	F/F	108	36	unknown
hTAS2R47	rs2597924	G/A	R/H	920	307	unknown
AF494233	rs1669405	T/G	L/W	842	281	unknown
	rs2599404	T/G	F/L	756	252	unknown
	rs2600355	T/G	V/V	54	18	unknown
hTAS2R48	rs1868769	T/C	L/L	418	140	unknown
AF494234						
hTAS2R49	hTAS2R49	A/G	K/R	164	55	unknown
AF494236	SNP1					
hTAS2R50	rs1376521	A/G	Y/C	608	203	G 0.66/A0.34
AF494235	hTAS2R50 SNP1	A/G	P/P	777	259	unknown

[0064] A polypeptide that exhibits bitter substance binding activity is a polypeptide that has at least 20% (e.g., at least: 20%; 30%; 40%; 50%; 60%; 70%; 80%; 90%; 95%; 98%; 99%; 99.5%; or 100% or even more) of the ability of the respective full-length TAS2R to bind to a given bitter substance. Binding assays and bitter substances are described herein

[0065] The term "contacting" in the context of the present invention means any interaction between the compound with the polypeptide of the invention, whereby any of the at least two components can be independently of each other in a liquid phase, for example in solution, or in suspension or can be bound to a solid phase, for example, in the form of an essentially planar surface or in the form of particles, pearls or the like. In a preferred embodiment a multitude of different compounds are immobilized on a solid surface like, for example, on a compound library chip and the protein of the present invention is subsequently contacted with such a chip. In another preferred embodiment the cells genetically engineered with the polynucleotide of the invention or with a vector containing such a polynucleotide express the bitter taste receptor at the cell surface and are contacted separately in small containers, e.g., microtitre plates, with various com-

[0066] Detecting the presence and the binding of the compound to the polypeptide can be carried out, for example, by measuring a marker that can be attached either to the protein or to the compound. Suitable markers are known in the art and comprise, for example, fluorescence, enzymatic or radioactive markers. The binding of the two components can, however, also be measured by the change of an electrochemical parameter of the binding compound or of the protein, e.g. a change of the redox properties of either the protein or the binding compound, upon binding. Suitable methods of detecting such changes comprise, for example, potentiomet-

ric methods. Further methods for detecting and/or measuring the binding of the two components to each other are known in the art and can without limitation also be used to measure the binding of the compound to the polypeptide. The effect of the binding of the compound on the activity of the polypeptide can also be measured by assessing changes in the cells that express the polypeptides, for example, by assaying the intracellular release of calcium upon binding of the compound.

[0067] As a further step after measuring the binding of a compound and after having measured the binding strength of at least two different compounds at least one compound can be selected, for example, on grounds of a higher binding strength or on grounds of the detected intracellular release of calcium.

[0068] The thus selected compound is than in a preferred embodiment modified in a further step. Modification can be effected by a variety of methods known in the art, which include without limitation the introduction of one or more, preferably two, three or four novel side chains or residues or the exchange of one or more functional groups like, for example, introduction or exchange of halogens, in particular F, Cl or Br; the introduction or exchange of lower alkyl residues, preferably having one to five carbon atoms like, for example, methyl, ethyl, n-propyl, isopropyl, n-butyl, isobutyl, tert-butyl, n-pentyl or iso-pentyl residues; lower alkenyl residues, preferably having two, three, four or five carbon atoms; lower alkinyl residues, preferably having two, three, four or five carbon atoms, which can in a preferred embodiment be further substituted with F, Cl, Br, NH_2 , NO_2 , OH, SH, NH, CN, aryl, heteroaryl, COH or COOH group; or the introduction of, for example, one or more residue(s) selected from the group consisting of NH₂, NO₂, OH, SH, NH, CN, aryl, alkylaryl, heteroaryl, alkylheteroaryl, COH or COOH group. [0069] The thus modified binding substances are than individually tested with the method of the pre-sent invention, i.e.

they are contacted with the polypeptide as such or with the polypeptide expressed in a cell, and subsequently binding of the modified compounds is measured. In this step both the binding per se can be measured and/or the effect of the function of the protein like, e.g. the intracellular calcium release. If needed the steps of selecting the compound, modifying the compound, contacting the compound with a polypeptide of the invention and measuring the binding of the modified compound to the polypeptide can be repeated a third or any given number of times as required. The above described method is also termed "directed evolution" of the compound since it involves a multitude of steps including modification and selection, whereby binding compounds are selected in an "evolutionary" process optimizing their capabilities with respect to a particular property, e.g. its binding activity, its ability to activate, inhibit or modulate the activity, in particular inhibit the intracellular release of calcium mediated by the polypeptides of the present invention.

[0070] Of particular interest are compounds that antagonize the bitter taste receptor activity of the TAS2Rs disclosed and described herein. The specification thereby enables the skilled person to design intelligent compound libraries to screen for antagonists to the bitter response of these receptors, which in turn enables the development of compounds and compositions to suppress or eliminate bitter tasting components of foods, in particular animal foods, nutrients and dietary supplements and pharmaceutical or homeopathic preparations containing such phyto-chemicals. Similarly, the invention also enables the skilled person to screen for additional bitter ligands, or even to screen for compounds that enhance a bitter response, such as might be useful in the food industry. Therefore, another aspect of the invention is a process for isolating an antagonist of the bitter taste receptor activity of the polypeptide encoded by a polynucleotide selected from the group consisting of:

[0071] (a) polynucleotides encoding at least the mature form of the polypeptide having the deduced amino acid sequence as shown in SEQ ID NONOs: 1, 3, 5, 7, 9, 11, 13, 15, 17, 19, 21, 23, 25, 27, 29, 31, 33, 35, 37, 39, 41, 43, 45, 47 and 49;

[0072] (b) polynucleotides having the coding sequence, as shown in SEQ ID NOs: 2, 4, 6, 8, 10, 12, 14, 16, 18, 20, 22, 24, 26, 28, 30, 32, 34, 36, 38, 40, 42, 44, 46, 48 and 50 encoding at least the mature form of the polypeptide;

[0073] (c) polynucleotides encoding a fragment or derivative of a polypeptide encoded by a polynucleotide of any one of (a) to (b), wherein in said derivative one or more amino acid residues are conservatively substituted compared to said polypeptide, and said fragment or derivative has bitter taste receptor activity;

[0074] (d) polynucleotides which are at least 50% identical to a polynucleotide as defined in any one of (a) to (c) and which code for a polypeptide having bitter taste receptor activity; and

[0075] (e) polynucleotides the complementary strand of which hybridizes, preferably under stringent conditions to a polynucleotide as defined in any one of (a) to (d) and which code for a polypeptide having bitter taste receptor activity;

comprising:

[0076] (1) contacting said polypeptide or a host cell genetically engineered with said polynucleotide or with a vector containing said polynucleotide with a potential antagonist;

[0077] (2) determining whether the potential antagonists antagonizes the bitter taste receptor activity of said polypeptide.

[0078] The polynucleotide employed in this process encodes a polypeptide that still exhibits essentially the same activity as the respective mature bitter taste receptor, i.e. has "bitter taste receptor activity". Preferably the polypeptide has at least 20% (e.g., at least: 20%; 30%; 40%; 50%; 60%; 70%; 80%; 90%; 95%; 98%; 99%; 99.5%; or 100% or even more) of the activity of the respective full-length TAS2R. One preferred way of measuring TAS2R activity is the ability to release intracellular calcium in a heterologous cell expression system like, for example, (HEK293/15) that stably expresses the alpha-subunit of promiscuous G-proteins, e.g. the mouse G₁₅ subunit or chimeric, in response to bitter tastants, which is dependent on the expression of polypeptides encoded by the polynucleotides of the present invention. The amount of intracellular calcium released can be monitored by, for example, the in vitro FLIPR assay described herein but also by the measurement of one of a variety of other parameters including, for example, IP3 or cAMP. Additional ways of measuring G-protein coupled receptor activity are known in the art and comprise without limitation electrophysiological methods, transcription assays, which measure, e.g. activation or repression of reporter genes which are coupled to regulatory sequences regulated via the respective G-protein coupled signaling pathway, such reporter proteins comprise, e.g., CAT or LUC; assays measuring internalization of the receptor; or assays in frog melanophore systems, in which pigment movement in melanophores is used as a read out for the activity of adenylate cyclase or phospholipase C (PLC), which in turn are coupled via G-proteins to exogenously expressed receptors (see, for example, McClintock T. S. et al. (1993) Anal. Biochem, 209: 298-305; McClintock T. S. and Lerner M. R. (1997) Brain Res. Brain, Res. Protoc. 2: 59-68, Potenza M N (1992) Pigment Cell Res. 5: 372-328, and Potenza M. N. (1992) Anal. Biochem. 206: 315-322)

[0079] As described above with the exception of hTAS2R40, single nucleotide polymorphisms are known for all of the above hTAS2R5, which can be employed in a process for isolating an antagonist of the bitter taste receptor activity. Polynucleotides or polypeptides that differ from the respectively in SEQ ID 1-50 indicated sequences by the nucleotide and amino acid change as indicted in Table I can similarly be employed for the process of the present invention.

[0080] The term "contacting" has the meaning as outlined above. A potential antagonist is a sub-stance which lowers the respective bitter taste receptor activity determined in the absence of the antagonist by at least 10% (e.g., at least: 1%, 15% 20%; 30%; 40%; 50%; 60%; 70%; 80%; 90%; 95%; 98%; 99%; 99.5%; or 100%) once contacted with the bitter taste receptor.

[0081] In a preferred embodiment the process further comprises the contacting of the polypeptide with an agonist of the respective bitter taste receptor activity. The contacting of the bitter taste receptor with the agonist can be carried out prior, concomitantly or after contacting the polypeptide with the potential antagonist.

[0082] It has been demonstrated by the inventors that the bitter receptors hTAS2R10, hTAS2R14, hTAS2R16, hTAS2R38, hTAS2R43, hTAS2R44, hTAS2R45, hTAS2R46 and hTAS2R48 respond with specificity to (a) defined classe (s) of ligand(s) that include a class of useful phyto-chemicals

in a functional expression assay. Therefore, in an even more preferred embodiment the polypeptides and agonist employed together in above process are selected from the group consisting of:

[0083] (a) the polypeptide encoded by the polynucleotide outlined above as determined by SEQ ID NO: 1 and SEQ ID NO: 2 and the agonist selected from the group consisting of acetyl-thiourea, N,N-dimethylthioformamide, N,N'-diphenylthiourea, N-ethylthiourea, 2-imidazolidinethione, 4(6)-methyl-2-thiouracil, N-methylthiourea, phenylthiocarbamid, 6-phenyl-2-thiouracil, 6-propyl-2-thiouracil, tetramethylthiourea, thioacetamide, thioacetanilide, 2-thiobarbituric acid, and 2-thiouracil and functional derivatives thereof;

[0084] (b) the polypeptide encoded by the polynucleotide of claim 1 or 2 as determined by SEQ ID NO: 9 and SEQ ID NO: 10 and the agonist selected from the group consisting of saccharin and functional derivatives thereof;

[0085] (c) the polypeptide encoded by the polynucleotide of claim 1 or 2 as determined by SEQ ID NO: 11 and SEQ ID NO: 12 and the agonist selected from the group consisting of saccharin and acesulfame K and functional derivatives thereof;

[0086] (d) the polypeptide encoded by the polynucleotide of claim 1 or 2 as determined by SEQ ID NO: 13 and SEQ ID NO: 14 and the agonist selected from the group consisting of absinthine and functional derivatives thereof;

[0087] (e) the polypeptide encoded by the polynucleotide of claim 1 or 2 as determined by SEQ ID NO: 15 and SEQ ID NO: 16 and the agonist selected from the group consisting of absinthine and functional derivatives thereof;

[0088] (f) the polypeptide encoded by the polynucleotide of claim 1 or 2 as determined by SEQ ID NO: 19 and SEQ ID NO: 20 and the agonist selected from the group consisting of absinthine and functional derivatives thereof;

[0089] (g) the polypeptide encoded by the polynucleotide of claim 1 or 2 as determined by SEQ ID NO: 37 and SEQ ID NO: 38 and the agonist selected from the group consisting of strychnine, brucine, denatonium benzoate, and absinthine and functional derivatives thereof;

[0090] (h) the polypeptide encoded by the polynucleotide of claim 1 or 2 as determined by SEQ ID NO: 41 and SEQ ID NO: 42 and the agonist selected from the group consisting of tyrosine, preferably L-tyrosine, and other bitter tasting amino acids including, e.g., leucine, histidine phenylalanine and tryptophan, and functional derivatives thereof; and

[0091] (i) the polypeptide encoded by the polynucleotide of claim 1 or 2 as determined by SEQ ID NO: 43 and SEQ ID NO: 44 and the agonist selected from the group consisting of naphtyl-β-D-glucoside, phenyl-β-D-glucoside, salicin, helicin, arbutin, 2-nitrophenyl-β-D-glucoside, 4-nitrophenyl-β-D-glucoside, methyl-β-D-glucoside, esculin, 4-nitrophenyl-β-D-thioglucoside, 4-nitrophenyl-β-D-mannoside, and amygdalin and functional derivatives thereof.

[0092] The term "functional derivatives thereof" refers to substances, which are derived from the respectively indicated bitter substance by chemical modification and which elicit at least 20% (e.g., at least: 20%; 30%; 40%; 50%; 60%; 70%; 80%; 90%; 95%; 98%; 99%; 99.5%; or 100% or even more) of the bitter taste receptor activity, if compared to the respective unmodified bitter substance. Chemical modification includes without limitation the introduction of one or more, preferably two, three or four novel side chains or residues or

the exchange of one or more functional groups like, for example, introduction or exchange of H; linear or branched alkyl, in particular lower alkyl (C1, C2, C3, C4, and C5, e.g. methyl, ethyl, n-propyl, isopropyl, n-butyl, isobutyl, tert-butyl, n-pentyl or iso-pentyl); substituted linear or branched alkyl, in particular lower substituted alkyl; linear or branched alkenyl, in particular lower alkenyl (C₂, C₃, C₄ and C₅, e.g. ethenyl, 1-propenyl, 2-propenyl, iso-propenyl, 1-butenyl, 2-butenyl, 3-butenyl; substituted linear or branched alkenyl, in particular lower substituted alkenyl; linear or branched alkinyl, in particular lower alkinyl (C2, C3, C4 and C5); substituted linear or branched alkinyl, in particular lower substituted alkinyl; linear or branched alkanol, in particular lower alkanol (C₁, C₂, C₃, C₄, and C₅); linear or branched alkanal, in particular lower alkanal (C1, C2, C3, C4, and C5, e.g. COH, CH2COH, CH2CH2COH; aryl, in particular phenyl; substituted aryl, in particular substituted aryl; heteroaryl; substituted heteroaryl; alkylaryl, in particular benzyl; substituted alkylaryl; in particular substituted benzyl; alkylheteroaryl; substituted alkylheteroaryl; aminoalkyl, C₁, C₂, C₃, C₄ and C_5 , e.g. —NHCH₃, —NHCH₂CH₃, —N(CH₃)₂; substituted aminoalkyl; aminoketone, in particular —NHCOCH3; substituted aminoketone; aminoaryl, in particular -NH-Ph; substituted aminoaryl, in particular substituted —NH-Ph; CN; NH₂; Halogen, in particular F, Cl, and Br; NO₂; OH; SH; NH; CN; or COOH group. If the residues mentioned above are substituted they are preferably mono, di, or tri substituted with a substituent selected from the group of halogen, in particular F, Cl, and Br, NH₂, NO₂, OH, SH, NH, CN, aryl, alkylaryl, heteroaryl, alkylheteroaryl, COH or COOH.

[0093] In particular the hTAS2R16 receptor has been shown to respond specifically to a narrow class of interesting phyto-chemicals selected from the group consisting of bitter beta-glucopyranosides and mannopyranosides.

[0094] The beta-glucopyranosides and beta-mannosepyranosides are a group of bitter compounds consisting of a hydrophobic residue attached to glucose and mannose, respectively, by a beta-glycosidic bond.

[0095] Preferred compounds that bind to the hTAS2R16 taste receptor are chosen from beta glucopyranosides and beta-mannopyranosides defined by the formula:

[0096] These compounds were studied in vitro (see Table I and IV below) and also by human panelists (see Table I below) as is described in greater detail below.

[0097] From these studies certain inferences can be drawn regarding the affinity of the compounds towards activation of the hTAS2R16 receptor. Thus, for the promotion of activation the steric position at C2 can be either alpha or beta and the beta-configuration of the glycosidic bond and the alpha steric position of the hydroxyl group at C4 of the pyranose ring are preferred. Whereas R can be hydrogen, it is preferred that R is a substituent selected from C_1 - C_8 alkyl which may be branched, linear or cyclic as appropriate; lower alkenyl resi-

dues, preferably having two, three, four or five carbon atoms; lower alkinyl residues, preferably having two, three, four or five carbon atoms, which can in a preferred embodiment be further substituted with F, Cl, Br, NH₂, NO₂, OH, SH, NH, CN, aryl, heteroaryl, COH or COOH group; heteroaryl, e.g. benzofuran and cumarin; aryl, e.g. phenyl, naphtyl; or the same of other sugar residue, e.g. glucopyranoside, which itself can carry a substituent R with the meaning as outlined above. Bulkier groups at Cl may increase the activation of the receptor. The aryl or heteroaryl may be further substituted with one or more substituents. Preferred substituents of the aryl or heteroaryl group are F, Br, Cl, NO₂, lower alkyl with one, two, three, four, five, six, seven or eight carbon atoms and CH₂OH. The phenyl group is preferably mono, di, or trisubstituted in ortho, para and/or meta position(s). The substituent at C6 is shown as an hydroxyl group above. However, the compounds activity as agonists are little effected by further or alternative substitution at this position, and there is design freedom at this part of the compound. Furthermore, without intending to be bound by theory, it is thought that the substituent "R" is not responsible for bitterness in these compounds. Rather, bitterness is thought to derive from a hydrogen acceptor and donor site provided by two hydroxyl groups on the ring. In another embodiment the β -glycosidic bond of the compounds outlined above can be a S-glycosidic bond, as exemplified by the bitter substance 4-nitrophenyl-β-D-thio-

[0098] Most preferred compounds are selected from the group consisting of naphtyl-β-D-glucoside, phenyl-β-D-glucoside, salicin, helicin, arbutin, 2-nitrophenyl-β-D-glucoside, 4-nitrophenyl-β-glucoside, methyl-β-D-glucoside, esculin, 4-nitrophenyl-β-D-thioglucoside, 4-nitrophenyl-β-D-mannoside, and amygdalin.

[0099] The beta-glucopyranosides are phytonutrients that represent an important class of compounds found in plantderived foods that may be useful as dietary supplements, or in functional foods or medicaments for the prevention of disease states. However, due to their bitter after-taste they are aversive to consumers and so they are routinely removed from foods during production and processing as is further described in Drewnowski, A. & Gomez-Carneros, C. Bitter taste, phytonutrients, and the consumer: a review. Am. J. Clin. Nutr. 72, 1424-1435 (2000). Removal is laborious and therefore expensive. The alternative is to mask the off-flavor using encapsulation technologies or organoleptic compounds as masking agents. However, encapsulation technology may not be appropriate in pharmaceutics as this may affect the absorption characteristics of the active compound, whereas the use of masking agents may impart their own characteristic flavor which may unbalance the flavor of food or beverages.

[0100] Without wishing to be bound by any particular theory as to their mechanism of action, applicant believes that the bitter receptors activate a G-protein and thereby initiate the aforementioned cellular activation cascade as a result of conformational changes in the receptor after binding by a ligand. Potential antagonists of the bitter response will contain functionality (i.e., will compete for binding at the receptor, and/or act at another binding site through an allosteric mechanism, and/or stabilize the receptor in the inactive conformation, and/or bind reversibly or irreversibly, and/or weaken receptor G protein interaction, and/or interfere with G protein activation).

[0101] Similarly, in another embodiment of the invention, it has been found that the so-called hTAS2R10 receptor is acti-

vated by strychnine, and strychnine analogues such as brucine as well as by denatonium benzoate, absinthine and other alkaloids with (a) ring system(s). Strychnine and its analogues are also useful phytochemicals that find use in medicines and homeopathic treatments.

[0102] In another embodiment of the invention, it has been found that the so-called hTAS2R14 receptor is activated by tyrosine, in particular L-tyrosine, and other bitter tasting amino acids including leucine, histidine, phenylalanine and tryptophan.

[0103] In another embodiment of the invention, it has been found that the so-called hTAS2R38 receptor is activated by acetylthiourea, N,N-dimethylthioformamide, N,N'-diphenylthiourea, N-ethylthiourea, 2-imidazolidinethione, 4(6)-methyl-2-thiouracil, N-methylthiourea, phenylthio-carbamid, 6-phenyl-2-thiouracil, 6-propyl-2-thiouracil, tetramethylthiourea, thioacetamide, thioacetanilide, 2-thiobarbituric acid, and 2-thiouracil.

[0104] From these studies certain inferences can be drawn regarding the affinity of the compounds, which activate the hTAS2R38 receptor. Thus, for the promotion of activation derivatives of 2-thiouracil according to following formula are preferred compounds.

[0105] Whereas R in this formula can be hydrogen, it is preferred that R is a substituent selected from C₁-C₁₀ alkyl, which may be branched, linear or cyclic as appropriate, particularly preferred alkyls are methyl, ethyl, n-propyl, isopropyl, n-butyl, isobutyl, tert-butyl, n-pentyl or iso-pentyl residues; lower alkenyl residues, preferably having two, three, four or five carbon atoms; lower alkynyl residues, preferably having two, three, four or five carbon atoms, which can in a preferred embodiment be further substituted with F, Cl, Br, NH₂, NO₂, OH, SH, NH, CN, aryl, heteroaryl, COH or COOH group; heteroaryl, e.g. benzofuran and cumarin; aryl, e.g. phenyl, naphtyl; F, Cl, Br, NH₂, NO₂, OH, SH, NH, CN, aryl, alkylaryl, heteroaryl, alkylheteroaryl, COH or COOH group. ID a further embodiment the carbon atom at the 4 position can substituted with $-O-R_1$, in which R_1 can have the same meaning as outlined above for R.

[0106] Another general structure of compounds having affinity for hTAS2R38 and which are thus suitable for activation of hTAS2R38 is depicted by the following formula:

$$R_4$$
— C — N

[0107] In this formula R_2 , R_3 , and R_4 can each independently of each other have the meaning H; alkyl, in particular lower alkyl (C_1 - C_5 , e.g. methyl, ethyl, n-propyl, isopropyl, n-butyl, isobutyl, tert-butyl, n-pentyl or iso-pentyl); substituted alkyl; alkenyl, in particular lower alkenyl (C_2 - C_5); sub-

stituted alkenyl; alkinyl, in particular lower alkinyl (C_2 - C_5); substituted alkinyl: alkanal, in particular lower alkanal (e.g. —COCH₃, —COCH₂CH₃, —COCH₂CH₂CH₃); aryl, in particular phenyl; substituted aryl; heteroaryl; substituted heteroaryl; alkylaryl, in particular benzyl; substituted alkylaryl; alkylheteroaryl; substituted alkylheteroaryl aminoalkyl, in particular —NHCH₃, —NHCH₂CH₃, —N(CH₃)₂; substituted aminoalkyl; aminoketone, in particular —NHCOCH3; substituted aminoketone; aminoaryl, in particular —NH-Ph; substituted aminoaryl; CN; NH₂; Halogen, in particular F, Cl, and Br; NO₂. In a preferred embodiment R₂ or R₃ and R₄ can form a ring, preferably a four, five, six, seven or eight membered hetero cycle, which in a preferred embodiment is an aromatic hetero cycle. The residue of R₂ or R₃, which is not involved in the formation of the ring structure can have any of the meanings as outlined above. In a further preferred embodiment at least one of R₂ or R₃ has the meaning alkanal, preferably lower alkanal as outlined above. In case that only one of R₂ or R₃ has the meaning alkanal, than the other substituent preferably has the meaning H.

[0109] In another embodiment of the invention, it has been found that the so-called hTAS2R43 receptor is activated by saccharin, derivatives thereof and other sulfonamides.

[0110] In another embodiment of the invention, it has been found that the so-called hTAS2R44 receptor is activated by saccharin and acesulfame K, derivatives thereof and other sulfonamides.

[0111] In another embodiment of the invention, it has been found that the so-called hTAS2R45, hTAS2R46 and hTAS2R48 receptor is activated by absinthine derivatives thereof and other sulfonamides.

[0112] The skilled person will appreciate that having regard to the structure-function information provided by the present invention, it is possible to compile libraries of molecules to find inhibitors of the bitter response of the disclosed hTAS2R in particular of the hTAS2R10, 14, 16, 38, 43, 44, 45, 46, and 48, which are triggered by the above outlined specific bitter substance(s). Such inhibitors, and libraries comprising same, form other aspects of the present invention. A still farther aspect of the invention relates to the use of such inhibitors in food or pharmaceutical compositions containing bitter tastants such as referred to herein above, for the elimination or suppression of bitter taste perception.

[0113] In practicing the various aspects and embodiments of the present invention in relation to cloning receptors, elucidating ligand-receptor pairs, and finding modulators of the bitter response of receptors, recourse is made to conventional techniques in molecular biology, microbiology and recombinant technology. Accordingly, the skilled person is fully apprised of such techniques and as such they are hereafter treated only summarily in order to more fully describe the context of the present invention.

[0114] In order to express cDNAs encoding the receptors, one typically subclones receptor cDNA into an expression vector that contains a strong promoter to direct transcription, a transcription/translation terminator, and a ribosome-binding site for translational initiation. Suitable bacterial promoters are well known in the art, e.g., *E. Coli, Bacillus* sp., and *Salmonella*, and kits for such expression systems are com-

mercially available. Similarly eukaryotic expression systems for mammalian cells, yeast, and insect cells are well known in the art and are also commercially available. The eukaryotic expression vector may be, for example an adenoviral vector, an adeno-associated vector, or a retroviral vector.

[0115] In addition to the promoter, the expression vector typically contains a transcription unit or expression cassette that contains all the additional elements required for the expression of the receptor-encoding nucleic acid in host cells. A typical expression cassette thus contains a promoter operatively linked to the nucleic acid sequence encoding the receptor and signals required for efficient polyadenylation of the transcript, ribosome binding sites, and translation termination. The nucleic acid sequence encoding the receptor may typically be linked to a membrane-targeting signal such as the N-terminal 45 amino acids of the rat Somatostatin-3 receptor sequence to promote efficient cell-surface expression of the recombinant receptor. Additional elements of the cassette may include, for example enhancers.

[0116] An expression cassette should also contain a transcription termination region downstream of the structural gene to provide for efficient termination. The termination region may be obtained from the same gene as the promoter sequence or may be obtained from different genes.

[0117] The particular expression vector used to transport the genetic information into the cell is not particularly critical. Any of the conventional vectors used for expression in eukaryotic or prokaryotic cells may be used. Standard bacterial expression vectors include plasmids such as pBR322 based plasmids, pSKF, pET23D, and fusion expression systems such as GST and LacZ, but there are many more known in the art to the skilled person that can be usefully employed.

[0118] Expression vectors containing regulatory elements from eukaryotic viruses are typically used in eukaryotic expression vectors, e.g., SV40 vectors, papilloma virus vectors, and vectors derived from Epstein-Barr virus. Other exemplary eukaryotic vectors include pMSG, pAV009/A. sup.+, pMTO10/A.sup.+, pMAMneo-5, baculovirus pDSVE, pcDNA3.1, pIRES and any other vector allowing expression of proteins under the direction of the SV40 early promoter, SV40 late promoter, metallothionein promoter, murine mammary tumor virus promoter, Rous sarcoma virus promoter, polyhedrin promoter, or other promoters shown effective for expression in eukaryotic cells.

[0119] Some expression systems have markers that provide gene amplification such as thymidine kinase, hygromycin B phosphotransferase, and dihydrofolate reductase. Alternatively, high yield expression systems not involving gene amplification are also suitable.

[0120] The elements that are typically included in expression vectors also include a replicon that functions in *E. Coli*, a gene encoding drug resistance to permit selection of bacteria that harbor recombinant plasmids, and unique restriction sites in nonessential regions of the plasmid to allow insertion of eukaryotic sequences. The particular drug resistance gene chosen is not critical, any of the many drug resistance genes known in the art are suitable. The prokaryotic sequences are optionally chosen such that they do not interfere with the replication of the DNA in eukaryotic cells, if necessary.

[0121] Standard transfection methods can be used to produce bacterial, mammalian, yeast or insect cell lines that express large quantities of the receptor, which are then purified using standard techniques.

[0122] Any of the well known procedures for introducing foreign nucleotide sequences into host cells may be used. These include the use of calcium phosphate transfection, polybrene, protoplast fusion, electroporation, liposomes, microinjection, plasma vectors, viral vectors and any of the other well known methods for introducing cloned genomic DNA, cDNA, synthetic DNA or other foreign genetic material into a host cell. It is only necessary that the particular genetic engineering procedure used be capable of successfully introducing at least one gene into the host cell capable of expressing the receptor.

[0123] After the expression vector is introduced into the cells, the transfected cells may be cultured under conditions favoring expression of the receptor, which is recovered from the culture using standard techniques. For example the cells may be burst open either mechanically or by osmotic shock before being subject to precipitation and chromatography steps, the nature and sequence of which will depend on the particular recombinant material to be recovered. Alternatively, the recombinant protein may be recovered from the culture medium in which the recombinant cells had been cultured.

[0124] The activity of any of the receptors described herein can be assessed using a variety of in vitro and in vivo assays to determine functional, chemical, and physical effects, e.g., measuring ligand binding, secondary messengers (e.g., cAMP, cGMP, IP₃, DAG, or Ca²⁺) ion flux, phosphorylation levels, transcription levels, neurotransmitter levels, and the like. Furthermore, such assays can be used to test for inhibitors of the receptors as is well known in the art.

[0125] Samples or assays that are treated with a potential receptor inhibitor may be compared to control samples without the test compound, to examine the extent of modulation. Control samples (untreated with inhibitors) are assigned a relative receptor activity value of 100. Inhibition of receptor activity is achieved when the receptor activity value relative to the control is lower, and conversely receptor activity is enhanced when activity relative to the control is higher.

[0126] The effects of the test compounds upon the function of the receptors can be measured by examining any of the parameters described above. Any suitable physiological change that affects receptor activity can be used to assess the influence of a test compound on the receptors of this invention. When the functional consequences are determined using intact cells or animals, one can measure a variety of effects such as changes in intracellular secondary messengers such as Ca²⁺, IP₃ or cAMP.

[0127] Preferred assays for G-protein coupled receptors include cells that are loaded with ion sensitive dyes to report receptor activity. In assays for identifying modulatory compounds, changes in the level of ions in the cytoplasm or membrane voltage will be monitored using an ion sensitive or membrane voltage fluorescent indicator, respectively. For G-protein coupled receptors, promiscuous G-proteins such as G.alpha. 15 and G.alpha. 16 and chimeric G-proteins can be used in the assay of choice (see, for example, Wilkie et al., Proc. Nat. Acad. Sci. USA 88, 10049-10053 (1991)). Such promiscuous G-proteins allow coupling of a wide range of receptors.

[0128] Receptor activation typically initiates subsequent intracellular events, e.g., increases in second messengers such as IP₃, which releases intracellular stores of calcium ions. Activation of some G-protein coupled receptors stimulates the formation of inositol triphosphate (1133) through phos-

pholipase C-mediated hydrolysis of phosphatidylinositol (Berridge & Irvine, Nature 312, 315-21 (1984)). IP3 in turn stimulates the release of intracellular calcium ion stores. Thus, a change in cytoplasmic calcium ion levels, or a change in second messenger levels such as $\rm IP_3$ can be used to assess G-protein coupled receptor function. Cells expressing such G-protein coupled receptors may exhibit increased cytoplasmic calcium levels as a result of contribution from both intracellular stores and via activation of ion channels, in which case it may be desirable, although not necessary, to conduct such assays in calcium-free buffer, optionally supplemented with a chelating agent such as EGTA, to distinguish fluorescence response resulting from calcium release from internal stores.

[0129] In a preferred embodiment, receptor activity is measured by expressing the receptor in a heterologous cell with a promiscuous G-protein, such as G.alpha. 15, 16, or a chimeric G-protein that links the receptor to a phospholipase C signal transduction pathway. Optionally the cell line is HEK-293, although other mammalian cells are also preferred such as CHO and COS cells. Modulation of taste transduction is assayed by measuring changes in intracellular Ca²⁺ levels, which change in response to modulation of the receptor signal transduction pathway via administration of a molecule that associates with the receptor. Changes in Ca²⁺ levels are optionally measured using fluorescent Ca⁺ indicator dyes and fluorometric imaging.

[0130] The type of assay described above with respect to G-protein coupled bitter taste receptors can, however, also be employed for the identification of binding compounds, in particular agonists or antagonists of any G-protein coupled signalling molecule, in particular G-protein coupled receptor. Therefore, another aspect of the present invention relates to a process for the identification of agonists or antagonists of G-protein coupled signalling molecules comprising the steps of:

[0131] (1) contacting a cell comprising a promiscuous G-protein like, for example, G.alpha.15, 16, or a chimeric G-protein, and a G-protein coupled signalling molecule, in particular receptor, with a the potential agonist or antagonists of the signalling molecule;

[0132] (2) determining whether the potential agonist or antagonists agonizes or antagonizes the activity of the signalling molecule.

[0133] The activity of the signalling molecule and the increase or decrease of that activity in response to the potential agonist or antagonist can be determined as outlined above with respect to the identification of bitter receptor taste activity. The respectively indicated percent increases or decreases of the activity, which are required to qualify as antagonist or agonist do apply mutatis mutandis. Additionally the term "contacting" has the meaning as outlined above. Preferably the signalling molecule and/or the promiscuous G-protein has been introduced into the cell. The type of cell, which are preferred are those indicated above.

[0134] In yet another embodiment, the ligand-binding domains of the receptors can be employed in vitro in soluble or solid-state reactions to assay for ligand binding. Ligand binding in a receptor, or a domain of a receptor, can be tested in solution, in a bilayer membrane attached to a solid phase in a lipid monolayer or vesicles. Thereby, the binding of a modulator to the receptor, or domain, can be observed using changes in spectroscopic characteristics, e.g. fluorescence,

absorbance or refractive index; or hydrodynamic (e.g. shape), chromatographic, or solubility properties, as is generally known in the art.

[0135] The compounds tested as modulators of the receptors can be any small chemical compound, or a biological entity, such as a protein, sugar, nucleic acid or lipid. Typically, test compounds will be small chemical molecules. Essentially any chemical compound can be used as a potential modulator or ligand in the assays of the invention, although knowledge of the ligand specificity of an individual receptor would enable the skilled person to make an intelligent selection of interesting compounds. The assays may be designed to screen large chemical libraries by automating the assay steps and providing compounds from any convenient source to assays, which are typically run in parallel (e.g., in microtiter formats on microtiter plates in robotic assays). The skilled person will understand that there are many suppliers of libraries of chemical compounds.

[0136] Assays may be run in high throughput screening methods that involve providing a combinatorial chemical or peptide library containing a large number of potential therapeutic, or tastant compounds (that are potential ligand compounds). Such libraries are then screened in one or more assays, as described herein, to identify those library members (particular chemical species or subclasses) that display a desired characteristic activity. The compounds thus identified can serve as lead compounds to further develop modulators for final products, or can themselves be used as actual modulators.

[0137] A combinatorial chemical library is a collection of diverse chemical compounds generated by either chemical synthesis or biological synthesis, by combining a number of chemical "building blocks" such as reagents. For example, a linear combinatorial chemical library such as a polypeptide library is formed by combining a set of chemical building blocks (amino acids) in every possible way for a given compound length (i.e., the number of amino acids in a polypeptide compound). Millions of chemical compounds can be synthesized through such combinatorial mixing of chemical building blocks.

[0138] Preparation and screening of combinatorial chemical libraries is well known to those of skill in the art and no more needs to be stated here.

[0139] In the high throughput assays of the invention, it is possible to screen up to several thousand different modulators or ligands in a single day. In particular, each well of a microtiter plate can be used to run a separate assay against a selected potential modulator, or, if concentration or incubation time effects are to be observed, every 5-10 wells can test a single modulator. Thus, a single standard microtiter plate can assay about 100 (e.g., 96) modulators. If 1536 well plates are used, then a single plate can easily assay from about 100 to about 1500 different compounds. It is possible to assay several different plates per day; assay screens for up to about 6,000-20,000 different compounds is possible using the integrated systems of the invention.

[0140] Lead compounds found by assay technology herein above described, or development compounds formed from such leads can be administered directly to a human subject to modulate bitter taste. Alternatively, such compounds can be formulated with other ingredients of preparations to be taken orally, for example, foods, including animal food, and beverages, pharmaceutical or nutraceutical or homeopathic preparations.

[0141] Therefore, another aspect of the invention is a process for the production of foodstuffs or any precursor material or additive employed in the production of foodstuffs comprising the steps of the above described processes for the identification of a compound binding to hTAS2R or an antagonist of hTAS2R and the subsequent step of admixing the identified compound or antagonist with foodstuffs or any precursor material or additive employed in the production of foodstuffs. [0142] Bitter taste is a particular problem when orally administering pharmaceuticals, which often have an unpleasant bitter taste. In particular in elderly persons, children and chronically ill patients this taste can lead to a lack of compliance with a treatment regimen. In addition in veterinary applications the oral administration of bitter tasting pharmaceuticals can be problematic. Therefore, a further aspect of the invention is a process for the production of a nutraceutical or pharmaceutical composition comprising the steps of the processes of a compound binding to hTAS2R or an antagonist of hTAS2R and the subsequent step of formulating the compound or antagonist with an active agent in a pharmaceutically acceptable form.

[0143] Consequently, a further aspect of the invention is a foodstuff, in particular animal food, or any precursor material or additive employed in the production of foodstuffs comprising an antagonist/inhibitor described above, preferably an antibody directed against one of the hTAS2Rs described herein, the extracellular domain of one of the hTAS2Rs described herein or an inhibiting RNA.

[0144] Also comprised is a nutraceutical or pharmaceutical composition comprising an antagonist/inhibitor as described above, preferably an antibody directed against one of the hTAS2Rs described herein, the extracellular domain of one of the hTAS2Rs described herein or an inhibiting RNA and an active agent, which preferably inhibits a bitter taste, and optionally a pharmaceutically acceptable carrier.

[0145] The amount of compound to be taken orally must be sufficient to effect a beneficial response in the human subject, and will be determined by the efficacy of the particular taste modulators and the existence, nature, and extent of any adverse side-effects that accompany the administration of a particular compound. There now follows a series of examples that serve to illustrate the invention, not to limit.

[0146] A further aspect of the present invention is the use of a polynucleotide as described above, a vector as described above, an antibody as described above or an antagonist/inhibitor of as described above, preferably an antibody directed against one of the hTAS2Rs described herein, the extracellular domain of one of the hTAS2Rs described herein or an inhibiting RNA for the manufacture of a medicament for the treatment of an abnormally increased or decreased sensitivity towards a bitter substance.

[0147] Techniques associated with detection or regulation of genes are well known to skilled artisans. Such techniques can be used, for example, for basic research on bitter receptors and to diagnose and/or treat disorders associated with aberrant bitter receptor expression.

[0148] The following examples are merely illustrative of the present invention and should not be construed to limit the scope of the invention as indicated by the appended claims in any way. The contents of the U.S. provisional application Ser. No. 60/413,298 the priority of which is claimed is hereby incorporated by reference in its entirety.

EXAMPLE 1

Cloning of the hTAS2R Genes

[0149] Human genomic DNA was isolated from HEK293 cells using the E.Z.N.A. Blood DNA Kit II (Peqlab) and the

various hTAS2Rs were amplified by PCR using gene-specific primers that span the complete coding region of the individual hTAS2R genes. Reaction parameters were: 4 cycles; 1 min, 94° C.; 1 min, 64° C.; 1.5 min 68° C. using Advantage 2 polymerase (Clontech). 5% of the reaction served then as template for further amplification with Pfu DNA polymerase (Promega): 30 cycles; 1 min, 94 C; 1 min, 64° C.; 3 min, 72° C. The hTAS2R amplicons were then sub-cloned into a cassette based on pcDNA5-FRT (Invitrogen). The cloning cassette contains the first 45 amino acids of the rat somatostatin type 3 receptor (as is further described by Meyerhof et al., Proc. Nat. Acad. Sci. USA, 89, 10267-10271 (1992)) as a cell surface-targeting signal at the N-terminus. The C-terminus contained the herpes simplex virus (HSV) glycoprotein D epitope which does not interfere with signaling of heptahelical receptors and can be used for immunocytochemistry using an antibody that binds specifically to the HSV glycoprotein D epitope (see Roosterman et al, J. Neuroendocrinol, 9, 741-751 (1997)). Comparison of the DNA sequences of at least four clones identified mutations generated during PCR and this avoided picking mutated clones. We compared the amino acid sequences using the AlignX program of the Vector NTITM Suite (InforMax).

[0150] Using the above-described method, DNA sequences encoding all 24 bitter receptors identified by applicant were cloned. As indicated above, they were derived by a PCR-based method using genomic DNA as the template. Since all of the 24 genomic sequences lack introns, the DNA clones obtained had the same sequences as corresponding cDNA clones derived by reverse transcription-PCR (RT-PCR) of mRNA from cells expressing the relevant polypeptides would have.

EXAMPLE 2

Immunocytochemistry

[0151] Batches of HE 93 cells were separately transiently transfected with expression vectors (pCDN5/FRT; Invitrogen) containing each of the 24 above described coding sequences using lipofectamine 2000 (Invitrogen) and aliquots of the resulting cell populations were separately seeded on polylysine-coated coverslips. At 24 h post transfection they were washed with phosphate buffered saline (PBS), cooled on ice and added 20 microgram/ml biotin-labeled concanavalin A (Sigma) for 1 h, which binds to cell surface glycoproteins. Thereafter, the cells were fixed for 5 min in methanol/acetone (1:1) and then permeabilized for 4 min with 0.25% Triton X-100. In order to reduce nonspecific binding the coverslips were incubated in 2% goat serum. Thereafter, anti-HSV glycoprotein D antiserum (Novagen, 1:10,000) was added to detect the chimeric receptors that, as described above, would have a HSV glycoprotein epitope fused to their C-temini, and Texas Red-Avidin D (Vector, 1:200) has added to stain the cell surface and incubation continued overnight at 4° C. Such C-termini are intracellular and for this reason it is necessary to permeabilize the cells to permit entry of the HSV glycoprotein D epitope-specific antibody molecules into them. After washing (5x in PBS, RT) Alexa488-conjugated goat anti-mouse antiserum (Molecular Probes, 1:1000) was added and incubation continued at room temperature for 1 h. Finally, the cells were embedded in Fluorescent Mounting Medium (Dako) and analyzed using a Leica TCS SP2 Laser Scan Inverted microscope. The preparations were scanned sequentially with an argon/krypton laser (488 nm) to excite the Alexa488 dye and with a greenhelium-neon laser (543 nm) to excite the Texas Red dye. The spectral detector recorded light emission at 510-560 nm and 580-660 nm, respectively. Images of 1024×1024 pixels were processed with Corel PHOTO-PAINT 10.0 (Corel Corporation) and printed on a Tektronix color laser printer. The immunocytochemical data permitted calculation of the proportion of cells expressing recombinant receptors (green fluorescent cells divided by total cell number in a microscopic field) and the proportion of cells that display expression of TAS2Rs at the plasma membrane level (number of cells with colocalization of green and red fluorescence divided by the number of green fluorecent cells). Of the 24 transfectant lines tested, all were found to express the encoded polypeptides. The proportion of receptor-expressing cells in the various transfectant lines ranged from about 10% to about 35%.

EXAMPLE 3

Heterologous Expression of hTAS2R Receptors

[0152] A fluorescence imaging plate reader (FLIPR, Molecular Devices) was used to functionally screen cell populations transiently transfected with expression vectors encoding the above-described 24 bitter receptors and to establish concentration-response curves for hTAS2R16 and hTASR10. The single-cell calcium imaging technique was also employed to demonstrate receptor selectivity and crossdesensitization. For the FLIPR experiments the HEK293/15 cells were grown to 50% confluence. The cells were then seeded at a density of 3×10³ cells per well into 96-well blackwall, clear-bottom microtiter plates (Greiner). After 48 h the cells in each well were transfected using Lipofectamine 2000 and 24-30 h later were loaded with Fluo4AM (Molecular Probes). Thereafter they were stimulated with bitter compounds (SigmaAldrich, further purified by reversed-phase HPLC to >99% purity). Calcium signals were recorded simultaneously from each well at 1 Hz at 510 nm after excitation at 488 nm and the recordings were corrected for cell density. The responses of five wells containing cells expressing the same receptor and that received the same stimulus (i.e., the same compound at the same concentration) were averaged. Calcium traces were subtracted that were determined in triplicate of mock-transfected cells stimulated with the same concentration of tastant. The calculations rest on at least four independent transfection experiments. Plots of the amplitudes versus concentrations fitted by nonlinear regression to the function $f(x)=100/(1+(EC_{50}/x)_{p}H)$, with x=agonist concentration and nH=Hill coefficient permitted calculation of EC_{50} values and threshold values of activation. [0153] EC₅₀ and threshold values obtained with hTAS2R16-expressing transfectants are shown in Table 1 below and the results are described in Example 4.

[0154] In separate experiments, hTAS2R10-expressing transfectants were found to have a threshold of activation of approximately 0.1 μM and a EC $_{50}$ of 5-20 μM using strychnine as the test compound. Similar results were obtained with brucine.

[0155] Single-cell Ca²⁺ imaging was performed with the hTAS2R16-transfected HEK293/15 cells as described in *Cell* 95, 917-926 (1998), but with the following modifications: The Till Photonics imaging system (Munich, Germany) was used in which a monochromator is connected by a quartz fiber lightguide and an epifluorescence condenser to an inverted Olympus IX50 microscope equipped with a UApo/340 40×1.

35 oil-immersion lens. 30 h post-transfection, FURA-2AM-loaded cells were sequentially illuminated in 5 s intervals for 3-10 ms, first at 340 nm, then at 380 nm, online ratioed light emissions at 510 nm (340/380) and monitored the images via an intensified, cooled CCD camera. The 5 s interval camera pictures of all cells in the microscope field of vision permanently were stored and analyzed offline. 10-15% of all cells in the camera field responded to agonists in transient transfection experiments. The proportion of responders was about half of that found by immunocytochemistry, probably reflecting a sub-optimal signal transduction. Responses were not observed in mocktransfected cells. Isoproterenol (10 micro-Molar) was used at the end of all experiments to stimulate endogenous betaadrenergic receptors, proving a functional G_{totals} dependent signal transduction cascade.

[0156] For RT-PCR and in-situ hybridization work, human RNA (Clontech) was purchased or it was isolated firm surgical tongue specimens with peqGOLD RNAPure (Peqlab) and the preparations digested with DNase I (Invitrogen). Following cDNA synthesis (Smart cDNA synthesis Kit, Clontech) hTAS2R16 cDNA was PCR-amplified (39 cycles, 1 min 94° C., 1 min 64° C., 1 min 72° C.) using specific forward and reverse primers with overhangs containing EcoRI or NotI sites SEQ ID Nos 51 and 52 and the amplicons analyzed on agarose gels. Subcloning and sequencing demonstrated the identity of the amplified bands. Approximately 15 micrometer cryo-sections of human tongue specimens containing vallate papilla at 65° C. were processed and hybridized with a hTAS2R16 riboprobe spanning the complete coding region and generated from hTAS2R16 cDNA. The in-situ hybridization method used was essentially the same as that described in Nature, 413, 631-635 (2001) except that the riboprobe was conjugated with biotin and an alkaline phosphatase-avidin conjugate was used for detection. This experiment indicated that TAS2R16 mRNA is expressed in vallate papilla which are known to perceive bitter taste.

EXAMPLE 4

Human Taste Experiments

[0157] 15 experienced panelists in a sensory panel room at 22-25° C. determined bitter thresholds on three different sessions using a triangle test with tap water as solvent, according to methodology set out in *J. Agric. Food Chem.*, 49, 231-238 (2001), or Mailgaard M et al, "Sensory Evaluation Techniques" (CRC Press LLC, New York 1999). For dose-response relations, bitter tastant concentration series were presented to 10 trained panelists in random order. The panelists ranked the samples in increasing order of intensity and, for each concentration, evaluated bitterness intensity on a scale from 0 to 5 (ref. 24). The dose-response curves of three different sessions were averaged. The intensity values between individuals and separate sessions differed by not more than 0.5 units.

[0158] To investigate adaptation, the 8 panelists first maintained aqueous solutions (5 ml) of phenyl- β -D-glucopyranoside (8 mM), phenyl-alpha-D-glucopyranoside (180 mM), salicin (8 mM), or helicin (8 mM) for 15 s in their oral cavities and evaluated the bitter intensity as described above. After 30 min, they kept a denatonium benzoate solution (5 ml, 0.0003 mM) for 15 s in their mouth and evaluated its bitterness. The panelists spat off the denatonium benzoate solution, took up the phenyl- β -D-glucopyranoside or the phenyl-alpha-D-glucopyranoside solutions orally for 120 s or 180 s and judged

their bitterness intensity after 15, 30, 60, 120 and 180s. Thereafter, the panelists spat off these solutions and then sequentially took up salicin, helicin (5 ml, 8 mM) and denatonium benzoate (5 ml, 0.0003 mM) and evaluated bitterness intensities of these solutions after 15 s. After an additional 30 min, the first experiment was repeated. The data of three different sessions for each panelist were averaged. Intensity values between individuals and separate sessions differed by not more than ±0.5 units.

[0159] Results of in vitro assays (FLIPR) and human taste experiments are shown in Table 1 below.

TABLE II

	Threshold Value (mM)		EC.	50 (mM)
Compound	FLIPR	Human	FLIPR	Human
1	0.07 +/- 0.02	0.1 +/- 0.05	1.1 +/- 0.1	0.7 +/- 0.2
2	0.07 +/- 0.02	0.2 + / - 0.1	1.4 +/- 0.2	1.1 + / - 0.3
3	0.3 + / - 0.1	0.4 + / - 0.1	2.3 +/- 0.4	2.2 + / - 0.7
4	0.5 + / - 0.2	0.9 + / - 0.3	5.8 +/- 0.9	5.4 + / - 1.8
5	1.5 + / - 0.5	n.d.	n.d.	n.d.
6	0.4 + / - 0.1	0.2 + / - 0.1	1.0 + / - 0.1	1.4 + / - 0.4
7	15 +/- 6	32 +/- 11	n.d.	320 +/- 108
8	2.3 + / - 0.9	n.d.	20 +/- 3.4	n.d.
9	4 +/- 2	4 +/- 1	n.d.	n.d.
10	n.r.	40 +/- 13	n.r.	n.d.
11	n.r.	9 +/- 3	n.r.	50 +/- 17

1 = phenyl-beta-D-glucopyranoside; 2 = salicin; 3 = helicin; 4 = arbutin; 5 = 2-nitro-phenyl-beta-D-glucopyranoside; 6 = naphthyl-beta-D-pyranoside; 7 = methyl-beta-D-lucopyranoside; 8 = amygdalin; 9 = esculin; 10 = phenyl-beta-D-galactopyranoside; 11 = phenyl-alpha-D-glucopyranoside. n.d. = not data due to solubility problems or toxicity or artifacts in vitro. n.r. = No response up to 100 mM.

[0160] The FLIPR results provide the threshold concentration of the compounds (nM) at which point the receptor detects the compounds. The EC_{50} results express the concentration of the compound wherein the receptor signal is at 50%, and is a representation of the affinity of a receptor for a compound.

[0161] The results show that the in vitro FLIPR measurements for salicin closely resemble the human taste study results. This bitter-tasting compound has known anti-pyretic and analgesic action, and the results suggest that in vitro assays using hTAS2R16 may represent a useful tool to find compounds that suppress or eliminate the bitter response to this compound. Also, for all the other tested beta-glucopyranosides, the close correspondence of Threshold Concentration and EC $_{50}$ results suggest that hTAS2R16 is a cognate human receptor for these class of bitter compounds. In contrast, the related structures (see compounds 10 and 11) show 90- to 400-fold higher Threshold Concentrations, which indicates that this receptor is rather selective, and that these bitter compounds activate different receptors.

[0162] Adaptation frequently occurs in sensory systems and means that stimuli elicit reduced responses upon prolonged or repeated stimulus presentations. Repeated stimulation of hTAS2R16-expressing cells with phenyl-beta-D-glucopyranoside resulted in largely diminished responses to salicin as well. This cross-desensitization occurred among the other tested beta-pyranosides and was fully reversible. It resembles homologous desensitization of agonist-occupied heptahelical receptors mediated by GRKs, i.e. specific kinases, and arresting. We also observed adaptation in the human test panel that initially scored phenyl-beta-D-glucopyranoside, salicin and helicin as equally intensely bitter.

The bitterness of phenyl-beta-D glucopyranoside declined during prolonged stimulation and the test panel perceived salicin and helicin also as less bitter, but not the unrelated bitter substance denatorium benzoate, which cannot activate TAS2R16. Adaptation was fully reversible. On the opposite, the phenyl-alpha-D-glucopyranoside failed to cross-adapt with all tested beta-D-glucopyranosides, although its own bitter response desensitized strongly. This indicates that betaglucopyranosides signal through a common mechanism most likely involving hTAS2R16 as a bitter taste receptor while the alpha-isomer activates a separate receptor. A recent human psychophysical study also revealed cross-adaptation amongst two bitter amino acids but not between the two bitter amino acids and urea, suggesting the existence of distinct receptors for the bitter amino acids and urea. Although most, if not all, bitter receptors are present in the same subset of taste receptor cells, adaptation to specific bitter stimuli can be explained if bitter receptors were subject to homologous desensitization.

EXAMPLE 5

Heterologous Expression of hTAS2R

[0163] Transient transfection of TAS2Rs into HEK-293T- $G\alpha16$ gustducin44 cells. We cloned the DNAs of all human putative bitter responsive receptors into pcDNA5/FRT (Invitrogen) by PCR-methods and transiently transfected the plasmids with lipofectamine 2000 (Invitrogen) into HEK-293T- $G\alpha16$ gustducin44 cells grown to 50% confluence. These cells stably express a chimeric G protein constructed from human $G\alpha16$ and rat gustducin. Finally, we seeded the transfected cells at a density of 3×10^3 cells per well into 96-well black-wall, clear-bottom microtiter plates (Greiner).

[0164] Co-transfection of TAS2Rs with gustducin and phospholipase-C β 2 into HEK-293 cells. Alternatively, we transfected simultaneously plasmid DNAs encoding one of the TAS2Rs, phospholipase-C β 2 and α -gustducin into HEK-293 cells using the lipofectamine method. Additional cotrans-

fection of G-protein β and γ -subunits may improve the bitter tastant-induced responses. Thereafter, the transfected cells were seeded at a density of 3×10^3 cells per well into 96-well black-wall, clear-bottom microtiter plates (Greiner).

[0165] Fluorometric Imaging Plate Reader (FLIPR) assay 24-30 h later, the cells were loaded with 4 μ M FLUO-4/AM (Molecular Probes) and 0.04% Pluronic F-127 (Molecular Probes) in Hepes-buffered saline (HBS), 140 mM NaCl, 5 mM KCl, 2.5 mM CaCl₂, 10 mM Hepes, 10 mM glucose and 2.5 mM probenicide, pH 7.4, for 1 hour at 37° C. Thereafter, cells were gently washed in HBS by an automated plate washer (Denley Cellwash, Labsystenms) and transferred to the FLIPR (Molecular Devices). The FLIPR integrates an argon laser excitation source, a 96-well pipettor, and a detection system utilizing a Charged Coupled Device imaging camera. Fluorescence emissions from the 96 wells were monitored at an emission wavelength of 510 nm, after excitation with 488 nm (F488). Fluorescence data were collected 1 min before and 10 min after stimulation. Data were collected every 6 s before and every 1 s after agonist stimulation. 50 μl of 3× concentrated agonists were delivered within 2 by the integrated 96-well pipettor to the wells containing 100 μl HBS. Agonist responses were quantified using the amplitudes of the fluorescence peaks. We averaged the responses of five wells containing cells expressing the same receptor and that received the same stimulus. Calcium traces were determined in triplicate of mock-transfected cells stimulated with the same concentration of tastant. EC₅₀ values and plots of the amplitudes versus concentrations were derived from fitting the data by nonlinear regression to the function f(x)=100/[1+ $(EC_{50}/x)^{nH}H$], where x is the agonist concentration and nH is the Hill coefficient. The results for hTAS2R10 (Table II), hTAS2R14 (Table III), hTAS2R16 (Table IV), hTAS2R38 (Table V), hTAS2R43 (Table VI), hTAS2R44 (Table VII), hTAS2R45 (Table VIII), hTAS2R46 (Table IX) and hTAS2R (Table X) are shown below.

TABLE III

	Identified agonists of hTAS2R10		
Substance	Structure	Approx. threshold [mM]	EC ₅₀ [mM]
Strychnine*	H H H H H H H H H H H H H H H H H H H	0.003	0.04
Brucine	CH ₃ O N H H O	0.01	0.06

TABLE III-continued

	Identified agonists of hTAS2R10		
Substance	Structure	Approx. threshold [mM]	EC ₅₀ [mM]
Denatonium benzoate	· C-0-	0.003	0.07
	CH ₃ O CH ₂ CH ₃		
Absinthine	HO, M, H, M,	0.01	

TABLE IV

Substance	Structure	Reacts at
L-Tyrosine	$\begin{array}{c c} H_2N & H & O \\ \hline \\ CH_2-C-C-C-OH \end{array}$	1 mM

TABLE V

	Identified agonists of hTAS2R16		
Substance	Structure	Threshold [mM]	EC ₅₀ [mM]
Naphtyl-β-D-Glucoside	HOIM OH	0.4 ± 0.1	1.0 ± 0.1
Phenyl-β-D-Glucoside	HOUTH OH OH	0.07 ± 0.02	1.1 ± 0.1

TABLE V-continued

	Identified agonists of hTAS2R16		
Substance	Structure	Threshold [mM]	EC ₅₀ [mM]
Salicin	HOCH ₂ OH OH	0.07 ± 0.02	1.4 ± 0.2
Helicin	HOILING HCO	0.3 ± 0.1	2.3 ± 0.4
Arbutin	HO OH HOCH ₂ O HO OH	0.5 ± 0.2	5.8 ± 0.9
2-Nitrophenyl-β-D-Glucoside	OHIM OH NO2	0.3 – 1	Not determined
4-Nitrophenyl-β-D-Glucoside	OH IIIII OH NO2	1 – 3	Not determined
Methyl-β-D-Glucoside*	HOCH ₂ OCH ₃ HOOOH	15 ± 6	32 ± 11
Esculin	HOCH ₂ OH HO	4 ± 2	Not determined

TABLE V-continued

	Identified agonists of hTAS2R16		
Substance	Structure	Threshold [mM]	EC ₅₀ [mM]
4-Nitrophenyl-β-D-Thioglucoside	OH IIII NO2	1 - 5	Not determined
4-Nitrophenyl-β-D-Mannoside	OH IIII OH NO2	1 – 3	Not determined
Amygdalin	HOID OH HO OH	2.3 ± 0.9	20 ± 3.4

TABLE VI

	Identified agonists of hTAS2R38		
Substance	Structure	Approx. Threshold $[\mu M]$	EC ₅₀ [μΜ]
Acetylthiourea	$\begin{array}{c} O & S \\ \parallel & \parallel \\ CH_3 - C - NH - C - NH_2 \end{array}$	2	15
N,N-Dimethyl- thioformamide	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	10	55
N,N'-Diphenylthiourea		0.3	2.3
N-Ethylthiourea	$\begin{array}{c} S \\ \parallel \\ \text{CH}_3\text{CH}_2\text{NH} \longrightarrow \text{C} \longrightarrow \text{NH}_2 \end{array}$	30	260
$ \begin{array}{ll} \hbox{2-Imidazolidinethione} \\ (=\!\!-\!\!\!-\!\!\!-\!\!\!-\!\!\!-\!\!\!-\!\!\!\!-\!\!\!\!-\!\!\!\!$	NH	10	not determined

TABLE VI-continued

	TABLE VI-continued Identified agonists of hTAS2R38		
g.l.,		Approx. Threshold	
Substance 4(6)-Methyl-2-thiouracil	Structure	[μM] 20	[μM] 180
N-Methylthiourea	$\begin{array}{c} \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\$	100	estimated 600-800
Phenylthiocarbamid (PTC)	S NH	0.3	2
6-Phenyl-2-thiouracil		0.15	0.5
	N N N N S		
6-Propyl-2-thiouracil (PROP)	O N H	0.3	2
Tetramethylthiourea	$ \begin{array}{c} \text{CH}_{3} \\ \text{N} \\ \text{CH}_{3} \end{array} $ $ \begin{array}{c} \text{CH}_{3} \\ \text{CH}_{3} \end{array} $	10-30	100
Thioacetamide	CH_3 — C — NH_2	100	not determined
Thioacetanilide	CH_3 — C — NH — C	3	18
2-Thiobarbituric acid	HO NH S	reacts a	at 10 mM

TABLE VI-continued

	Identified agonists of hTA	AS2R38_	
Substance	Structure	Approx. Threshold $[\mu M]$	EC ₅₀ [μΜ]
2-Thiouracil	O H S	300	estimated 2000

TABLE VII

TABLE VIII

	Identified agonists of hTA	AS2R43	_	_	Identified agonists of hTA	AS2R44	
Substance	Structure	Approx. Thresh- old [mM]	EC ₅₀ [mM]	Substance	Structure	Approx. Threshold [mM]	EC ₅₀ [mM]
Saccharin	Na ⁺	0.2	1.1	Saccharin	Na ⁺	0.2	estimated 2-5
Acesulfame K	O H	No response	up to 10 mM	Acesulfame K	O N H	0.5	3

TABLE IX

	Identified agonists of hTAS2R	.45	
Substance	Structure	Approx. Threshold [mM]	EC ₅₀ [mM]
Absinthine	H. M. OH	0.003	Not determined

TABLE X

	Identified agonists of	hTAS2R46	
Substance	Structure	Approx. Threshold [mM]	EC ₅₀ [mM]
Absinthine HO,,,	H.M. OH	0.001	Not determined

TABLE XI

Substance Structure Approx. Threshold [mM] EC₅₀ [mM] Absinthine OH 0.03 Not determined

SEQUENCE LISTING

											-	con	tin	ued							
				85					90					95							
Met	Leu	Trp	Met 100	Ile	Ala	Asn	Gln	Ala 105	Asn	Leu	Trp	Leu	Ala 110	Ala	Cys						
Leu	Ser	Leu 115	Leu	Tyr	Cys	Ser	Lys 120	Leu	Ile	Arg	Phe	Ser 125	His	Thr	Phe						
Leu	Ile 130	CÀa	Leu	Ala	Ser	Trp 135	Val	Ser	Arg	Lys	Ile 140	Ser	Gln	Met	Leu						
Leu 145	Gly	Ile	Ile	Leu	Сув 150	Ser	CÀa	Ile	Cys	Thr 155	Val	Leu	Сув	Val	Trp 160						
CAa	Phe	Phe	Ser	Arg 165	Pro	His	Phe	Thr	Val 170	Thr	Thr	Val	Leu	Phe 175	Met						
Asn	Asn	Asn	Thr 180	Arg	Leu	Asn	Trp	Gln 185	Asn	Lys	Asp	Leu	Asn 190	Leu	Phe						
Tyr	Ser	Phe 195	Leu	Phe	CAa	Tyr	Leu 200	Trp	Ser	Val	Pro	Pro 205	Phe	Leu	Leu						
Phe	Leu 210	Val	Ser	Ser	Gly	Met 215	Leu	Thr	Val	Ser	Leu 220	Gly	Arg	His	Met						
Arg 225	Thr	Met	Lys	Val	Tyr 230	Thr	Arg	Asn	Ser	Arg 235	Asp	Pro	Ser	Leu	Glu 240						
Ala	His	Ile	Lys	Ala 245	Leu	ГÀа	Ser	Leu	Val 250	Ser	Phe	Phe	Cya	Phe 255	Phe						
Val	Ile	Ser	Ser 260	CÀa	Val	Ala	Phe	Ile 265	Ser	Val	Pro	Leu	Leu 270	Ile	Leu						
Trp	Arg	Asp 275	Lys	Ile	Gly	Val	Met 280	Val	Сла	Val	Gly	Ile 285	Met	Ala	Ala						
CAa	Pro 290	Ser	Gly	His	Ala	Ala 295	Ile	Leu	Ile	Ser	Gly 300	Asn	Ala	Lys	Leu						
Arg 305	_	Ala	Val	Met	Thr 310	Ile	Leu	Leu	Trp	Ala 315	Gln	Ser	Ser	Leu	Lys 320						
Val	Arg	Ala	Asp	His 325	ГЛа	Ala	Asp	Ser	Arg 330	Thr	Leu	СЛа									
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	O> SE																				
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480

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tcccagatgc tcctgggtat tattctttgc tcctgcatct gcactgtcct ctgtgtttgg

tgctttttta gcagacctca cttcacagtc acaactgtgc tattcatgaa taacaataca

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Ile Ile Ala Asn Gly Phe Ile Met Ala Ile His Ala Ala Glu Trp Val 50 55 60										
Gln Asn Lys Ala Val Ser Thr Ser Gly Arg Ile Leu Val Phe Leu Ser 65 70 75 80										
Val Ser Arg Ile Ala Leu Gln Ser Leu Met Met Leu Glu Ile Thr Ile 85 90 95										
Ser Ser Thr Ser Leu Ser Phe Tyr Ser Glu Asp Ala Val Tyr Tyr Ala										
Phe Lys Ile Ser Phe Ile Phe Leu Asn Phe Cys Ser Leu Trp Phe Ala 115 120 125										
Ala Trp Leu Ser Phe Phe Tyr Phe Val Lys Ile Ala Asn Phe Ser Tyr 130 135 140										
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Leu Leu Trp Leu Ser Val Phe Ile Ser Phe Ser His Ser Met Phe Cys 165 170 175										
Ile Asn Ile Cys Thr Val Tyr Cys Asn Asn Ser Phe Pro Ile His Ser 180 185 190										
Ser Asn Ser Thr Lys Lys Thr Tyr Leu Ser Glu Ile Asn Val Val Gly 195 200 205										
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Ile Leu Thr Ala Thr Leu Leu Ile Leu Ser Leu Lys Arg His Thr Leu 225 230 235 240										
His Met Gly Ser Asn Ala Thr Gly Ser Asn Asp Pro Ser Met Glu Ala 245 250 255										
His Met Gly Ala Ile Lys Ala Ile Ser Tyr Phe Leu Ile Leu Tyr Ile 260 265 270										
Phe Asn Ala Val Ala Leu Phe Ile Tyr Leu Ser Asn Met Phe Asp Ile 275 280 285										
Acr Con Lou Two Acr Acr Lou Cuc Cla Tlo Tlo Mot Ale Ale The Dec										

Asn Ser Leu Trp Asn Asn Leu Cys Gln Ile Ile Met Ala Ala Tyr Pro

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Ala Ser His Ser Ile Leu Leu Ile Gln Asp Asn Pro Gly Leu Arg Arg 305 310 315 320	
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Arg Gly Lys Thr Leu Pro Thr Gly Asp Arg Ile Met Leu Met Leu Ser 50 55 60	
Phe Ser Arg Leu Leu Gln Ile Trp Met Met Leu Glu Asn Ile Phe 65 70 75 80	

Ser Leu Leu Phe Arg Ile Val Tyr Asn Gln Asn Ser Val Tyr Ile Leu

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85 90 95
Phe Lys Val Ile Thr Val Phe Leu Asn His Ser Asn Leu Trp Phe Ala 100 105 110
Ala Trp Leu Lys Val Phe Tyr Cys Leu Arg Ile Ala Asn Phe Asn His 115 120 125
Pro Leu Phe Phe Leu Met Lys Arg Lys Ile Ile Val Leu Met Pro Trp 130 135 140
Leu Leu Arg Leu Ser Val Leu Val Ser Leu Ser Phe Ser Phe Pro Leu 145 150 155 160
Ser Arg Asp Val Phe Asn Val Tyr Val Asn Ser Ser Ile Pro Ile Pro 165 170 175
Ser Ser Asn Ser Thr Glu Lys Lys Tyr Phe Ser Glu Thr Asn Met Val 180 185 190
Asn Leu Val Phe Phe Tyr Asn Met Gly Ile Phe Val Pro Leu Ile Met 195 200 205
Phe Ile Leu Ala Ala Thr Leu Leu Ile Leu Ser Leu Lys Arg His Thr 210 215 220
Leu His Met Gly Ser Asn Ala Thr Gly Ser Arg Asp Pro Ser Met Lys 225 230 235 240
Ala His Ile Gly Ala Ile Lys Ala Thr Ser Tyr Phe Leu Ile Leu Tyr 245 250 255
Ile Phe Asn Ala Ile Ala Leu Phe Leu Ser Thr Ser Asn Ile Phe Asp 260 265 270
Thr Tyr Ser Ser Trp Asn Ile Leu Cys Lys Ile Ile Met Ala Ala Tyr 275 280 285
Pro Ala Gly His Ser Val Gln Leu Ile Leu Gly Asn Pro Gly Leu Arg 290 295 300
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Gln Thr Leu
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Arg Glu Trp Leu Arg Tyr Gly Arg Leu Leu Pro Leu Asp Met Ile Leu 35 40 45	
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His Asn Phe Tyr Tyr Ser Ala Gln Lys Val Glu Tyr Ser Gly Gly Leu 65 70 75 80	
Gly Arg Gln Phe Phe His Leu His Trp His Phe Leu Asn Ser Ala Thr 85 90 95	
Phe Trp Phe Cys Ser Trp Leu Ser Val Leu Phe Cys Val Lys Ile Ala	
Asn Ile Thr His Ser Thr Phe Leu Trp Leu Lys Trp Arg Phe Leu Gly 115 120 125	
Trp Val Pro Trp Leu Leu Gly Ser Val Leu Ile Ser Phe Ile Ile 130 135 140	
Thr Leu Leu Phe Phe Trp Val Asn Tyr Pro Val Tyr Gln Glu Phe Leu 145 150 155 160	
Ile Arg Lys Phe Ser Gly Asn Met Thr Tyr Lys Trp Asn Thr Arg Ile 165 170 175	
Glu Thr Tyr Tyr Phe Pro Ser Leu Lys Leu Val Ile Trp Ser Ile Pro 180 185 190	
Phe Ser Val Phe Leu Val Ser Ile Met Leu Leu Ile Asn Ser Leu Arg 195 200 205	
Arg His Thr Gln Arg Met Gln His Asn Gly His Ser Leu Gln Asp Pro 210 215 220	
Ser Thr Gln Ala His Thr Arg Ala Leu Lys Ser Leu Ile Ser Phe Leu 225 230 235 240	
Ile Leu Tyr Ala Leu Ser Phe Leu Ser Leu Ile Ile Asp Ala Ala Lys 245 250 255	
Phe Ile Ser Met Gln Asn Asp Phe Tyr Trp Pro Trp Gln Ile Ala Val 260 265 270	
Tyr Leu Cys Ile Ser Val His Pro Phe Ile Leu Ile Phe Ser Asn Leu 275 280 285	
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Asn	. Met	Thr	Trp	Lys 165	Ile	Lys	Leu	Lys	Ser	Ala	Met	Tyr	Phe	Ser 175	Asn		
Met	Thr	Val	Thr		Val	Ala	Asn	Leu 185		Pro	Phe	Thr	Leu 190		Leu		
Leu	Ser			Leu	Leu	Ile	_		Leu	Cys	Lys			Lys	Tàa		
Met	Gln	195 Leu	Arg	Gly	Lys	Gly	200 Ser	Gln	Asp	Pro	Ser	205 Thr	Lys	Val	His		
	210 Lys		_	-	-	215					220		-				
225	-				230					235					240		
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ys :05	Thr	Ser	Ser	Pro													
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:40	0> SI	EQUE	ICE :	10													
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Thr Glu Trp Val Lys Arg Gln Lys Ile Ser Phe Ala Asp Gln Ile Val \$35\$

Asn Trp Tyr Ser Thr Val Leu Asn Pro Ala Phe Cys Ser Val Glu Leu 65 70 75 80

Arg Thr Thr Ala Tyr Asn Ile Trp Ala Val Thr Gly His Phe Ser Asn $85 \hspace{1.5cm} 90 \hspace{1.5cm} 95$

Trp Pro Ala Thr Ser Leu Ser Ile Phe Tyr Leu Leu Lys Ile Ala Asn 100 \$100\$

Phe Ser Asn Leu Ile Phe Leu Arg Leu Lys Arg Arg Val Lys Ser Val 115 \$120\$

Ile Leu Val Val Leu Leu Gly Pro Leu Leu Phe Leu Ala Cys His Leu 130 135 140

Phe Val Val Asn Met Asn Gln Ile Val Trp Thr Lys Glu Tyr Glu Gly 145 $$ 150 $$ 155 $$ 160

Asn Met Thr Trp Lys Ile Lys Leu Arg Arg Ala Met Tyr Leu Ser Asp \$165\$ \$170\$ \$175\$

Thr Thr Val Thr Met Leu Ala Asn Leu Val Pro Phe Thr Val Thr Leu
180 185 190

Ile Ser Phe Leu Leu Leu Val Cys Ser Leu Cys Lys His Leu Lys Lys Met Gln Leu His Gly Lys Gly Ser Gln Asp Pro Ser Thr Lys Val His 215 Ile Lys Val Leu Gln Thr Val Ile Ser Phe Phe Leu Leu Arg Ala Ile Tyr Phe Val Ser Val Ile Ile Ser Val Trp Ser Phe Lys Asn Leu Glu Asn Lys Pro Val Phe Met Phe Cys Gln Ala Ile Gly Phe Ser Cys Ser Ser Ala His Pro Phe Ile Leu Ile Trp Gly Asn Lys Lys Leu Lys Gln 280 Thr Tyr Leu Ser Val Leu Trp Gln Met Arg Tyr 290 295 <210> SEO ID NO 14 <211> LENGTH: 897 <212> TYPE: DNA <213> ORGANISM: Homo sapiens <400> SEOUENCE: 14 atgataactt ttctgcccat catattttcc attctagtag tggttacatt tgttattgga 60 aattttgcta atggcttcat agcgttggta aattccaccg agtgggtgaa gagacaaaag 120 atctcctttg ctgaccaaat tgtcactgct ctggcggtct ccagagttgg tttgctctgg 180 gtgttattat taaattggta ttcaactgtg ttgaatccag ctttttgtag tgtagaatta 240 agaactactg cttataatat ctgggcagta accggccatt tcagcaactg gcctgctact agecteagea tattttattt geteaagatt geeaatttet eeaacettat ttttettege ttaaagagga gagttaagag tgtcattctg gtggtgctgt tggggccttt gctatttttg gettgteate tttttgtggt aaacatgaat cagattgtat ggacaaaaga atatgaagga aacatgactt ggaagatcaa attgaggcgt gcaatgtacc tttcagatac gactgtaacc 540 atgctagcaa acttagtacc ctttactgta accctgatat cttttctgct gttagtctgt 600 tctctgtgta aacatctcaa gaagatgcag ctccatggca aaggatctca agatcccagt 660 accaaggtcc acataaaagt tttgcaaact gtgatctcct tcttcttgtt acgtgccatt 720 tactttgtgt ctgtaataat atcagtttgg agttttaaga atctggaaaa caaacctgtc 780 ttcatgttct gccaagctat tggattcagc tgttcttcag cccacccgtt catcctgatt 840 897 tggggaaaca agaagctaaa gcagacttat ctttcagttt tgtggcaaat gaggtac <210> SEO ID NO 15 <211> LENGTH: 299 <212> TYPE: PRT <213> ORGANISM: Homo sapiens <400> SEQUENCE: 15 Met Ile Thr Phe Leu Pro Ile Ile Phe Ser Ile Leu Ile Val Val Thr Phe Val Ile Gly Asn Phe Ala Asn Gly Phe Ile Ala Leu Val Asn Ser Ile Glu Trp Phe Lys Arg Gln Lys Ile Ser Phe Ala Asp Gln Ile Leu

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Asn Trp Tyr Ala Thr Glu Leu Asn Pro Ala Phe Asn Ser Ile Glu Val 65 70 75 80	
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Trp Leu Ala Thr Ser Leu Ser Ile Phe Tyr Leu Leu Lys Ile Ala Asn 100 105 110	
Phe Ser Asn Leu Ile Phe Leu His Leu Lys Arg Arg Val Lys Ser Val 115 120 125	
Val Leu Val Ile Leu Leu Gly Pro Leu Leu Phe Leu Val Cys His Leu 130 135 140	
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Asn Met Thr Trp Lys Ile Lys Leu Arg Ser Ala Met Tyr Leu Ser Asn 165 170 175	
Thr Thr Val Thr Ile Leu Ala Asn Leu Val Pro Phe Thr Leu Thr Leu 180 185 190	
Ile Ser Phe Leu Leu Leu Ile Cys Ser Leu Cys Lys His Leu Lys Lys 195 200 205	
Met Gln Leu His Gly Lys Gly Ser Gln Asp Pro Ser Met Lys Val His	
Ile Lys Ala Leu Gln Thr Val Thr Ser Phe Leu Leu Cys Ala Ile 225 230 235 240	
Tyr Phe Leu Ser Ile Ile Met Ser Val Trp Ser Phe Glu Ser Leu Glu 245 250 255	
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Ser Leu Arg Leu

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		Val	Ile	Gly		Phe	Ala	Asn	Gly		Ile	Ala	Leu	Val	Asn	Ser		
		~ 1	_	20	_		~7		25		D.			30	-1			
	IIe	GIU	Trp 35	val	гув	Arg	GIn	ьуs 40	IIe	ser	Pne	val	Asp 45	GIN	Ile	ьeu		
	Thr	Ala 50	Leu	Ala	Val	Ser	Arg 55	Val	Gly	Leu	Leu	Trp	Val	Leu	Leu	Leu		
	His	Trp	Tyr	Ala	Thr	Gln	Leu	Asn	Pro	Ala	Phe	Tyr	Ser	Val	Glu	Val		
	65					70					75					80		
	Arg	Ile	Thr	Ala	Tyr 85	Asn	Val	Trp	Ala	Val 90	Thr	Asn	His	Phe	Ser 95	Ser		
	Trp	Leu	Ala	Thr	Ser	Leu	Ser	Met	Phe 105	Tyr	Leu	Leu	Arg	Ile 110	Ala	Asn		
	Phe	Ser	Asn 115	Leu	Ile	Phe	Leu	Arg 120	Ile	Lys	Arg	Arg	Val 125	ГЛа	Ser	Val		
	Val	Leu 130	Val	Ile	Leu	Leu	Gly 135	Pro	Leu	Leu	Phe	Leu 140	Val	Сув	His	Leu		
	Phe		Ile	Asn	Met	Asp		Thr	Val	Trp	Thr		Glu	Tyr	Glu	Gly		
	145					150				•	155	•		•		160		
	Asn	Val	Thr	Trp	Lys 165	Ile	ГЛа	Leu	Arg	Ser 170	Ala	Met	Tyr	His	Ser 175	Asn		
	Met	Thr	Leu	Thr	Met	Leu	Ala	Asn	Phe	Val	Pro	Leu	Thr	Leu 190	Thr	Leu		
	Ile	Ser			Leu	Leu	Ile	-		Leu	Cys	Lys			Lys	Lys		
	Mer	G 3	195	772 -	ar-	T	63 -	200	a.	3	Decem	G	205		777	772 cc		
	мet	Gln 210	ьeu	HIS	GIY	гуз	Gly 215	ser	GIN	Asp	Pro	Ser 220	Thr	гув	Val	HIS		
	Ile 225	Lys	Ala	Leu	Gln	Thr 230	Val	Thr	Ser	Phe	Leu 235	Leu	Leu	Cys	Ala	Ile 240		
	Tyr	Phe	Leu	Ser			Ile	Ser	Val	_	Asn	Phe	Gly	Arg	Leu	Glu		
	Lare	Gl 2	Dro	ΓeV.	245 Phe		Dhe	ر <i>ب</i> ر ۲۰ د	درات	250 Δla	Tla	Tla	Dhe	Sor	255 Tyr	Pro		
	пув	GIII	FIO	260	- 116	net	L116	CYB	265	AId	116	116	- 116	270	тут	£10		
	Ser	Thr	His 275	Pro	Phe	Ile	Leu	Ile 280	Leu	Gly	Asn	Lys	Lys 285	Leu	ГÀа	Gln		
	Ile		Leu	Ser	Val	Leu		His	Val	Arg	Tyr		Val	Lys	Asp	Arg		
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Ala Val Ile Thr Met Asp Glu Arg Val Trp Thr Lys Glu Tyr Glu Gly Asn Val Thr Trp Lys Ile Lys Leu Arg Asn Ala Ile His Leu Ser Ser Leu Thr Val Thr Thr Leu Ala Asn Leu Ile Pro Phe Thr Leu Ser Leu 185 Ile Cys Phe Leu Leu Leu Ile Cys Ser Leu Cys Lys His Leu Lys Lys 200 Met Arg Leu His Ser Lys Gly Ser Gln Asp Pro Ser Thr Lys Val His 215 Ile Lys Ala Leu Gln Thr Val Thr Ser Phe Leu Met Leu Phe Ala Ile 230 235 Tyr Phe Leu Cys Ile Ile Thr Ser Thr Trp Asn Leu Arg Thr Gln Gln 245 250 Ser Lys Leu Val Leu Leu Cys Gln Thr Val Ala Ile Met Tyr Pro 260 265 Ser Phe His Ser Phe Ile Leu Ile Met Gly Ser Arg Lys Leu Lys Gln 280 Thr Phe Leu Ser Val Leu Trp Gln Met Thr 290

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<212> TYPE: PRT

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Ile	Ala	Trp 35	Val	Lys	Arg	Gln	Lys 40	Ile	Ser	Ser	Ala	Asp 45	Gln	Ile	Ile		
Ala	Ala 50	Leu	Ala	Val	Ser	Lys 55	Val	Gly	Leu	Leu	Trp 60	Val	Ile	Leu	Leu		
His 65	Trp	Tyr	Ser	Thr	Val 70	Leu	Asn	Pro	Thr	Ser 75	Ser	Asn	Leu	Lys	Val 80		
Ile	Ile	Phe	Ile	Ser 85	Asn	Ala	Trp	Ala	Val 90	Thr	Asn	His	Phe	Ser 95	Ile		
Trp	Leu	Ala	Thr 100	Ser	Leu	Ser	Ile	Phe 105	Tyr	Leu	Leu	Lys	Ile 110	Val	Asn		
Phe	Ser	Arg 115	Leu	Ile	Phe	His	His 120	Leu	Lys	Arg	Lys	Ala 125	Lys	Ser	Val		
Val	Leu 130	Val	Ile	Val	Leu	Gly 135	Ser	Leu	Phe	Phe	Leu 140	Val	Cys	His	Leu		
Val 145	Met	Lys	His	Thr	Tyr 150	Ile	Asn	Val	Trp	Thr 155	Glu	Glu	СЛа	Glu	Gly 160		
Asn	Val	Thr	Trp	Lys 165	Ile	ГЛа	Leu	Arg	Asn 170	Ala	Met	His	Leu	Ser 175	Asn		
Leu	Thr	Val	Ala 180	Met	Leu	Ala	Asn	Leu 185	Ile	Pro	Phe	Thr	Leu 190	Thr	Leu		
Ile	Ser	Phe 195	Leu	Leu	Leu	Ile	Tyr 200	Ser	Leu	СЛа	ГЛа	His 205	Leu	Lys	Lys		
Met	Gln 210	Leu	His	Gly	ГЛа	Gly 215	Ser	Gln	Asp	Pro	Ser 220	Thr	Lys	Ile	His		
Ile 225	Lys	Ala	Leu	Gln	Thr 230	Val	Thr	Ser	Phe	Leu 235	Ile	Leu	Leu	Ala	Ile 240		
Tyr	Phe	Leu	Сув	Leu 245	Ile	Ile	Ser	Phe	Trp 250	Asn	Phe	Lys	Met	Arg 255	Pro		
ГÀа	Glu	Ile	Val 260	Leu	Met	Leu	СЛа	Gln 265	Ala	Phe	Gly	Ile	Ile 270	Tyr	Pro		
Ser	Phe	His 275						Trp	_		-			Lys	Gln		
Thr	Phe 290	Leu	Ser	Val	Leu	Trp 295	Gln	Val	Thr	Сув	Trp 300	Ala	Lys	Gly	Gln		
Asn 305	Gln	Ser	Thr	Pro													
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atct	ccto	ag c	tgat	caaa	at ta	attgo	ctgct	ctg	gcag	gtct	ccaa	aagtt	gg t	tttg	ctctgg	1	.80

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ataa	attt	ta t	ttci	taat	gc ct	tggg	cagta	a aco	caato	catt	tcaç	gcat	ctg q	gette	getaet
agc	ctcaç	gca t	atti	ttati	tt go	ctcaa	agato	gto	caatt	tct	ccaç	gacti	at 1	tttt	catcac
ttaa	aaaa	gga a	aggct	taaga	ag to	gtagi	tete	g gtg	gataç	gtgt	tgg	ggtci	tt (gttct	ttttg
gtti	gtca	acc t	tgt	gatga	aa a	caca	cgtat	ata	aaatq	gtgt	gga	cagaa	aga a	atgto	gaagga
aac	gtaad	ctt (ggaaq	gate	aa a	ctgaç	ggaat	gca	aatgo	cacc	ttt	ccaa	ett (gacto	gtagcc
atgo	ctago	caa a	actt	gata	cc at	ttcad	ctctc	g ac	cctga	atat	ctti	tct	get (gttaa	atctac
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tact	ttct	gt q	gtcta	aatca	at at	tcgt1	ttgg	g aat	ttta	aaga	tgc	gacca	aaa a	agaaa	attgtc
ttaa	atgct	tt (gccaa	agcti	tt to	ggaat	cata	tat	ccat	cat	tcca	actca	att (catto	ctgatt
tgg	gggaa	aca a	agac	gctaa	aa go	caga	ccttt	cti	tcaç	gttt	tgt	ggca	ggt (gactt	gctgg
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	D> SE L> LE														
	2> TY 3> OF			Homo	sap	piens	3								
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Phe	Val	Leu	Gly 20	Asn	Phe	Ala	Asn	Gly 25	Phe	Ile	Ala	Leu	Val 30	Asn	Phe
Ile	Asp	Trp 35	Val	Lys	Arg	Lys	Lys 40	Ile	Ser	Ser	Ala	Asp 45	Gln	Ile	Leu
Thr	Ala 50	Leu	Ala	Val	Ser	Arg 55	Ile	Gly	Leu	Leu	Trp	Ala	Leu	Leu	Leu
Asn 65	Trp	Tyr	Leu	Thr	Val 70	Leu	Asn	Pro	Ala	Phe 75	Tyr	Ser	Val	Glu	Leu 80
Arg	Ile	Thr	Ser	Tyr 85	Asn	Ala	Trp	Val	Val 90	Thr	Asn	His	Phe	Ser 95	Met
Trp	Leu	Ala	Ala 100	Asn	Leu	Ser	Ile	Phe 105	Tyr	Leu	Leu	Lys	Ile 110	Ala	Asn
Phe	Ser	Asn 115	Leu	Leu	Phe	Leu	His 120	Leu	Lys	Arg	Arg	Val 125	Arg	Ser	Val
Ile	Leu 130	Val	Ile	Leu	Leu	Gly 135	Thr	Leu	Ile	Phe	Leu 140	Val	Сув	His	Leu
Leu 145	Val	Ala	Asn	Met	Asp 150	Glu	Ser	Met	Trp	Ala 155	Glu	Glu	Tyr	Glu	Gly 160
Asn	Met	Thr	Gly	Lys 165	Met	Lys	Leu	Arg	Asn 170	Thr	Val	His	Leu	Ser 175	Tyr
Leu	Thr	Val	Thr 180	Thr	Leu	Trp	Ser	Phe 185	Ile	Pro	Phe	Thr	Leu 190	Ser	Leu
Ile	Ser	Phe 195	Leu	Met	Leu	Ile	Cys	Ser	Leu	Tyr	ГÀа	His 205	Leu	Lys	Lys
Met	Gln	Leu	His	Gly	Glu	Gly	Ser	Gln	Asp	Leu	Ser	Thr	Lys	Val	His

Ile Lys Ala Leu Gln Thr Leu Ile Ser Phe Leu Leu Cys Ala Ile

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225 230	235 240
Phe Phe Leu Phe Leu Ile Val Ser Val 245	Trp Ser Pro Arg Arg Leu Arg 250 255
Asn Asp Pro Val Val Met Val Ser Lya 260 269	
Ala Phe Asp Ser Phe Ile Leu Ile Trj 275 280	Arg Thr Lys Lys Leu Lys His 285
Thr Phe Leu Leu Ile Leu Cys Gln Ile 290 295	e Arg Cys
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atotootoag otgaccaaat totoactgot of	ggeggtet ccagaattgg tttgetetgg 180
gcattattat taaattggta tttaactgtg t	gaatccag ctttttatag tgtagaatta 240
agaattactt cttataatgc ctgggttgta ac	caaccatt tcagcatgtg gcttgctgct 300
aacctcagca tattttattt gctcaagatt g	caatttct ccaaccttct ttttcttcat 360
ttaaagagga gagttaggag tgtcattctg g	gatactgt tggggacttt gatatttttg 420
gtttgtcatc ttcttgtggc aaacatggat g	gagtatgt gggcagaaga atatgaagga 480
aacatgactg ggaagatgaa attgaggaat a	agtacatc tttcatattt gactgtaact 540
accetatgga getteatace etttactetg to	ecetgatat ettttetgat getaatetgt 600
tototgtata aacatotoaa gaagatgoag o	ccatggag aaggatcgca agatctcagc 660
accaaggtcc acataaaagc tttgcaaact c	gateteet teetettgtt atgtgeeatt 720
ttetttetat teetaategt tteggtttgg ag	tcctagga ggctgcggaa tgacccagtt 780
gtcatggtta gcaaggctgt tggaaacata ta	stottgoat togactoatt catcotaatt 840
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Phe Leu Leu Gly Ile Phe Thr Asn Gly 20 25	v Ile Ile Val Val Asn Gly 30
Ile Asp Leu Ile Lys His Arg Lys Met	Ala Pro Leu Asp Leu Leu Leu 45
Ser Cys Leu Ala Val Ser Arg Ile Pho 50 55	Leu Gln Leu Phe Ile Phe Tyr 60
Val Asn Val Ile Val Ile Phe Phe Ile 65 70	e Glu Phe Ile Met Cys Ser Ala 75 80

Asn Cys Ala Ile Leu Leu Phe Ile Asn Glu Leu Glu Leu Trp Leu Ala

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85 90 95	
Thr Trp Leu Gly Val Phe Tyr Cys Ala Lys Val Ala Ser Val Arg His 100 105 110	
Pro Leu Phe Ile Trp Leu Lys Met Arg Ile Ser Lys Leu Val Pro Trp 115 120 125	
Met Ile Leu Gly Ser Leu Leu Tyr Val Ser Met Ile Cys Val Phe His 130 135 140	
Ser Lys Tyr Ala Gly Phe Met Val Pro Tyr Phe Leu Arg Lys Phe Phe 145 150 155 160	
Ser Gln Asn Ala Thr Ile Gln Lys Glu Asp Thr Leu Ala Ile Gln Ile 165 170 175	
Phe Ser Phe Val Ala Glu Phe Ser Val Pro Leu Leu Ile Phe Leu Phe 180 185 190	
Ala Val Leu Leu Ile Phe Ser Leu Gly Arg His Thr Arg Gln Met 195 200 205	
Arg Asn Thr Val Ala Gly Ser Arg Val Pro Gly Arg Gly Ala Pro Ile 210 215 220	
Ser Ala Leu Leu Ser Ile Leu Ser Phe Leu Ile Leu Tyr Phe Ser His 225 230 235 240	
Cys Met Ile Lys Val Phe Leu Ser Ser Leu Lys Phe His Ile Arg Arg 245 250 255	
Phe Ile Phe Leu Phe Phe Ile Leu Val Ile Gly Ile Tyr Pro Ser Gly 260 265 270	
His Ser Leu Ile Leu Ile Leu Gly Asn Pro Lys Leu Lys Gln Asn Ala 275 280 285	
Lys Lys Phe Leu Leu His Ser Lys Cys Cys Gln 290 295	
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atggctccgc tggatctcct tetttettgt etggcagttt etagaatttt tetgcagttg	180
ttcatcttct acgttaatgt gattgttatc ttcttcatag aattcatcat gtgttctgcg	240
aattgtgcaa ttctcttatt tataaatgaa ttggaacttt ggcttgccac atggctcggc	300

360

420

480

540

600

660

840

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tgtgttttcc atagcaaata tgcagggttt atggtcccat acttcctaag gaaatttttc

tcccaaaatg ccacaattca aaaagaagat acactggcta tacagatttt ctcttttgtt

gctgagttct cagtgccatt gcttatcttc ctttttgctg ttttgctctt gattttctct

ctggggaggc acacccggca aatgagaaac acagtggccg gcagcagggt tcctggcagg

ggtgcaccca tcagcgcgtt gctgtctatc ctgtccttcc tgatcctcta cttctcccac
tgcatgataa aagttttct ctcttctcta aagtttcaca tcagaaggtt catctttctg
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Asn	Phe	Val	Gly 20	Ile	Ile	Met	Asn	Leu 25	Phe	Ile	Thr	Val	Val 30	Asn	Cha	
FÀa	Thr	Trp 35	Val	Lys	Ser	His	Arg 40	Ile	Ser	Ser	Ser	Asp 45	Arg	Ile	Leu	
Phe	Ser 50	Leu	Gly	Ile	Thr	Arg 55	Phe	Leu	Met	Leu	Gly 60	Leu	Phe	Leu	Val	
Asn 65	Thr	Ile	Tyr	Phe	Val 70	Ser	Ser	Asn	Thr	Glu 75	Arg	Ser	Val	Tyr	Leu 80	
Ser	Ala	Phe	Phe	Val 85	Leu	CÀa	Phe	Met	Phe 90	Leu	Asp	Ser	Ser	Ser 95	Val	
Trp	Phe	Val	Thr 100	Leu	Leu	Asn	Ile	Leu 105	Tyr	Cya	Val	Lys	Ile 110	Thr	Asn	
Phe	Gln	His 115	Ser	Val	Phe	Leu	Leu 120	Leu	Lys	Arg	Asn	Ile 125	Ser	Pro	Lys	
Ile	Pro 130	Arg	Leu	Leu	Leu	Ala 135	Cys	Val	Leu	Ile	Ser 140	Ala	Phe	Thr	Thr	
Суя 145	Leu	Tyr	Ile	Thr	Leu 150	Ser	Gln	Ala	Ser	Pro 155	Phe	Pro	Glu	Leu	Val 160	
Thr	Thr	Arg	Asn	Asn 165	Thr	Ser	Phe	Asn	Ile 170	Ser	Glu	Gly	Ile	Leu 175	Ser	
Leu	Val	Val	Ser 180	Leu	Val	Leu	Ser	Ser 185	Ser	Leu	Gln	Phe	Ile 190	Ile	Asn	
Val	Thr	Ser 195	Ala	Ser	Leu	Leu	Ile 200	His	Ser	Leu	Arg	Arg 205	His	Ile	Gln	
Lys	Met 210	Gln	ГÀа	Asn	Ala	Thr 215	Gly	Phe	Trp	Asn	Pro 220	Gln	Thr	Glu	Ala	
His 225	Val	Gly	Ala	Met	Lys 230	Leu	Met	Val	Tyr	Phe 235	Leu	Ile	Leu	Tyr	Ile 240	
Pro	Tyr	Ser	Val	Ala 245	Thr	Leu	Val	Gln	Tyr 250	Leu	Pro	Phe	Tyr	Ala 255	Gly	
Met	Asp	Met	Gly 260	Thr	Lys	Ser	Ile	Сув 265	Leu	Ile	Phe	Ala	Thr 270	Leu	Tyr	
Ser	Pro	Gly 275	His	Ser	Val	Leu	Ile 280	Ile	Ile	Thr	His	Pro 285	Lys	Leu	Lys	
Thr	Thr 290	Ala	ГÀа	Lys	Ile	Leu 295	CÀa	Phe	Lys	Lys						
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				-COIICII	lueu	
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atcattatga	atctgtttat	tacagtggtc	aattgcaaaa	cttgggtcaa	aagccataga	120
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<213> ORGANISM: Homo sapiens

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Phe Leu Ile Gly Leu Ile Gly Asn Gly Ser Leu Val Val Trp Ser Phe

Arg Glu Trp Ile Arg Lys Phe Asn Trp Ser Ser Tyr Asn Leu Ile Ile

Leu Gly Leu Ala Gly Cys Arg Phe Leu Leu Gln Trp Leu Ile Ile Leu

Asp Leu Ser Leu Phe Pro Leu Phe Gln Ser Ser Arg Trp Leu Arg Tyr 70

Leu Ser Ile Phe Trp Val Leu Val Ser Gln Ala Ser Leu Trp Phe Ala 90

Thr Phe Leu Ser Val Phe Tyr Cys Lys Lys Ile Thr Thr Phe Asp Arg 100 $\,$

Pro Ala Tyr Leu Trp Leu Lys Gln Arg Ala Tyr Asn Leu Ser Leu Trp 115 \$120\$

Cys Leu Leu Gly Tyr Phe Ile Ile Asn Leu Leu Leu Thr Val Gln Ile

Gly Leu Thr Phe Tyr His Pro Pro Gln Gly Asn Ser Ser Ile Arg Tyr 150 155

Pro Phe Glu Ser Trp Gln Tyr Leu Tyr Ala Phe Gln Leu Asn Ser Gly 170

Ser Tyr Leu Pro Leu Val Val Phe Leu Val Ser Ser Gly Met Leu Ile 180 $$185\$

Val Ser Leu Tyr Thr His His Lys Lys Met Lys Val His Ser Ala Gly

Arg Arg Asp Val Arg Ala Lys Ala His Ile Thr Ala Leu Lys Ser Leu Gly Cys Phe Leu Leu Leu His Leu Val Tyr Ile Met Ala Ser Pro Phe Ser Ile Thr Ser Lys Thr Tyr Pro Pro Asp Leu Thr Ser Val Phe Ile Trp Glu Thr Leu Met Ala Ala Tyr Pro Ser Leu His Ser Leu Ile Leu 265 Ile Met Gly Ile Pro Arg Val Lys Gln Thr Cys Gln Lys Ile Leu Trp 280 Lys Thr Val Cys Ala Arg Arg Cys Trp Gly Pro 290 <210> SEO ID NO 30 <211> LENGTH: 897 <212> TYPE: DNA <213> ORGANISM: Homo sapiens <400> SEOUENCE: 30 atgctgagcg ctggcctagg actgctgatg ctggtggcag tggttgaatt tctcatcggt 60 120 ttaattqqaa atqqaaqcct qqtqqtctqq aqttttaqaq aatqqatcaq aaaattcaac 180 tggtcctcat ataacctcat tatcctgggc ctggctggct gccgatttct cctgcagtgg ctgatcattt tggacttaag cttgtttcca cttttccaga gcagccgttg gcttcgctat 240 cttagtatet tetgggteet ggtaageeag geeagettat ggtttgeeae etteeteagt gtcttctatt gcaagaagat cacgaccttc gatcgcccgg cctacttgtg gctgaagcag agggeetata acctgagtet etggtgeett etgggetaet ttataateaa tttgttaett acagtccaaa ttggcttaac attctatcat cctccccaag gaaacagcag cattcggtat ccctttgaaa gctggcagta cctgtatgca tttcagctca attcaggaag ttatttgcct ttagtggtgt ttcttgtttc ctctgggatg ctgattgtct ctttgtatac acaccacaag 600 aagatgaagg tocattcagc tggtaggagg gatgtccggg ccaaggctca catcactgcg 660 ctgaagtcct tgggctgctt cctcttactt cacctggttt atatcatggc cagcccttc 720 tocatcacct ccaagactta toctootgat otcaccagtg tottcatotg ggagacactc 780 atggcagcct atccttctct tcattctctc atattgatca tggggattcc tagggtgaag 840 cagacttgtc agaagatcct gtggaagacg gtgtgtgctc ggagatgctg gggccca 897 <210> SEO ID NO 31 <211> LENGTH: 318 <212> TYPE: PRT <213> ORGANISM: Homo sapiens <400> SEQUENCE: 31 Met Ala Asp Lys Val Gln Thr Thr Leu Leu Phe Leu Ala Val Gly Glu 10 Phe Ser Val Gly Ile Leu Gly Asn Ala Phe Ile Gly Leu Val Asn Cys Met Asp Trp Val Lys Lys Arg Lys Ile Ala Ser Ile Asp Leu Ile Leu $35 \hspace{1.5cm} 40 \hspace{1.5cm} 45$ Thr Ser Leu Ala Ile Ser Arg Ile Cys Leu Leu Cys Val Ile Leu Leu

Asp Cys 65	Phe I	le Leu	Val 70	Leu	Tyr	Pro	Asp	Val 75	Tyr	Ala	Thr	Gly	80		
Glu Met	Arg I	le Ile 85	Asp	Phe	Phe	Trp	Thr 90	Leu	Thr	Asn	His	Leu 95	Ser		
Ile Trp		la Thr	Сув	Leu	Ser	Ile 105	Tyr	Tyr	Phe	Phe	Lys 110	Ile	Gly		
Asn Phe	Phe H: 115	is Pro	Leu	Phe	Leu 120	Trp	Met	Lys	Trp	Arg 125	Ile	Asp	Arg		
Val Ile 130	Ser T	rp Ile	Leu	Leu 135	Gly	CÀa	Val	Val	Leu 140	Ser	Val	Phe	Ile		
Ser Leu 145	Pro A	la Thr	Glu 150	Asn	Leu	Asn	Ala	Asp 155	Phe	Arg	Phe	СЛа	Val 160		
Lys Ala	Lys Ai	rg Lys 165	Thr	Asn	Leu	Thr	Trp 170	Ser	Сла	Arg	Val	Asn 175	Tàa		
Thr Gln		la Ser 30	Thr	Lys	Leu	Phe 185	Leu	Asn	Leu	Ala	Thr 190	Leu	Leu		
Pro Phe	Cys Va 195	al Cys	Leu	Met	Ser 200	Phe	Phe	Leu	Leu	Ile 205	Leu	Ser	Leu		
Arg Arg 210	His I	le Arg	Arg	Met 215	Gln	Leu	Ser	Ala	Thr 220	Gly	СЛа	Arg	Asp		
Pro Ser 225	Thr G	lu Ala	His 230	Val	Arg	Ala	Leu	Lys 235	Ala	Val	Ile	Ser	Phe 240		
Leu Leu	Leu Pl	ne Ile 245	Ala	Tyr	Tyr	Leu	Ser 250	Phe	Leu	Ile	Ala	Thr 255	Ser		
Ser Tyr		et Pro	Glu	Thr	Glu	Leu 265	Ala	Val	Ile	Phe	Gly 270	Glu	Ser		
Ile Ala	Leu I: 275	le Tyr	Pro	Ser	Ser 280	His	Ser	Phe	Ile	Leu 285	Ile	Leu	Gly		
Asn Asn 290	Lys Le	eu Arg	His	Ala 295	Ser	Leu	Lys	Val	Ile 300	Trp	Lys	Val	Met		
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attgcctc														180	
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gaaatgag														300	
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tggatgaa	.gt gga	agaatt	ga ca	ıgggt	gatt	tco	etgga	attc	tact	ggg	gtg (cgtg	gttctc	420	
tctgtgtt	ta tta	agcctt	cc ag	gccad	ctgag	g aat	ttga	acg	ctga	attt	cag (gtttt	gtgtg	480	
aaggcaaa	.ga gga	aaaaca	aa ct	taad	cttgg	g agt	tgca	agag	taaa	ataa	aac 1	caad	catgct	540	
tctaccaa	gt tat	tttctc	aa co	tgg	caaco	g ctg	gctco	cct	tttg	gtgt	gtg (cctaa	atgtcc	600	

tttt	tcct	ct t	gato	ectet	c co	etgeg	ggaga	a cat	atca	ıggc	gaat	gcag	get o	cagto	jccaca
gggt	gcag	gag a	accc	cagca	ac aç	gaago	ccat	gtg	gagag	jece	tgaa	agct	gt d	attt	ccttc
ctto	ctcct	ct t	tatt	gcct	a ct	attt	gtco	ttt	ctca	ittg	ccac	cctcc	ag o	ctact	ttatg
ccaç	gagad	egg a	aatta	agcto	gt ga	attt	tggt	gag	jtcca	tag	ctct	aato	cta c	ccct	caagt
catt	catt	ta t	ccta	aatao	et ge	gggaa	acaat	aaa	ittaa	ıgac	atgo	catct	ct a	aagg	ıtgatt
tgga	aaagt	aa t	gtct	atto	ct aa	aaagg	gaaga	a aaa	ttcc	aac	aaca	ataaa	aca a	atc	
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Phe	Ile	Leu	Gly 20	Ile	Leu	Gly	Asn	Gly 25	Tyr	Ile	Ala	Leu	Val 30	Asn	Trp
Ile	Asp	Trp 35	Ile	Lys	Lys	Lys	Lys 40	Ile	Ser	Thr	Val	Asp 45	Tyr	Ile	Leu
Thr	Asn 50	Leu	Val	Ile	Ala	Arg 55	Ile	CÀa	Leu	Ile	Ser 60	Val	Met	Val	Val
Asn 65	Gly	Ile	Val	Ile	Val 70	Leu	Asn	Pro	Asp	Val 75	Tyr	Thr	ГЛа	Asn	80 Lys
Gln	Gln	Ile	Val	Ile 85	Phe	Thr	Phe	Trp	Thr 90	Phe	Ala	Asn	Tyr	Leu 95	Asn
Met	Trp	Ile	Thr 100	Thr	Сув	Leu	Asn	Val 105	Phe	Tyr	Phe	Leu	Lys 110	Ile	Ala
Ser	Ser	Ser 115	His	Pro	Leu	Phe	Leu 120	Trp	Leu	Lys	Trp	Lys 125	Ile	Asp	Met
Val	Val 130	His	Trp	Ile	Leu	Leu 135	Gly	СЛа	Phe	Ala	Ile 140	Ser	Leu	Leu	Val
Ser 145	Leu	Ile	Ala	Ala	Ile 150	Val	Leu	Ser	Сув	Asp 155	Tyr	Arg	Phe	His	Ala 160
Ile	Ala	Lys	His	Lys 165	Arg	Asn	Ile	Thr	Glu 170	Met	Phe	His	Val	Ser 175	Lys
Ile	Pro	Tyr	Phe 180	Glu	Pro	Leu	Thr	Leu 185	Phe	Asn	Leu	Phe	Ala 190	Ile	Val
Pro	Phe	Ile 195	Val	Ser	Leu	Ile	Ser 200	Phe	Phe	Leu	Leu	Val 205	Arg	Ser	Leu
Trp	Arg 210	His	Thr	Lys	Gln	Ile 215	Lys	Leu	Tyr	Ala	Thr 220	Gly	Ser	Arg	Asp
Pro 225	Ser	Thr	Glu	Val	His 230	Val	Arg	Ala	Ile	Lys 235	Thr	Met	Thr	Ser	Phe 240
Ile	Phe	Phe	Phe	Phe 245	Leu	Tyr	Tyr	Ile	Ser 250	Ser	Ile	Leu	Met	Thr 255	Phe
Ser	Tyr	Leu	Met 260	Thr	Lys	Tyr	Lys	Leu 265	Ala	Val	Glu	Phe	Gly 270	Glu	Ile
Ala	Ala	Ile 275	Leu	Tyr	Pro	Leu	Gly 280	His	Ser	Leu	Ile	Leu 285	Ile	Val	Leu

Asn Asn Lys Leu Arg Gln Thr Phe Val Arg Met Leu Thr Cys Arg Lys

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936

130	eu Ala	TIE LE	135	GIY	ser	riie	ьец	140	ser	цец	TIE	iie	
Ser Val F 145	Pro Lys	Asn Asj 15		Met	Trp	Tyr	His 155	Leu	Phe	Lys	Val	Ser 160	
His Glu G	Slu Asn	Ile Th	r Trp	Lys	Phe	Lys 170	Val	Ser	Lys	Ile	Pro 175	Gly	
Thr Phe I	ys Gln 180	Leu Th	r Leu	Asn	Leu 185	Gly	Val	Met	Val	Pro 190	Phe	Ile	
Leu Cys I	eu Ile 195	Ser Ph	e Phe	Leu 200	Leu	Leu	Phe	Ser	Leu 205	Val	Arg	His	
Thr Lys 0	3ln Ile	Arg Le	1 His 215	Ala	Thr	Gly	Phe	Arg 220	Asp	Pro	Ser	Thr	
Glu Ala F 225	His Met	Arg Ala 23		Lys	Ala	Val	Ile 235	Ile	Phe	Leu	Leu	Leu 240	
Leu Ile V	/al Tyr	Tyr Pro	o Val	Phe	Leu	Val 250	Met	Thr	Ser	Ser	Ala 255	Leu	
Ile Pro 0	In Gly 260	Lys Le	ı Val	Leu	Met 265	Ile	Gly	Asp	Ile	Val 270	Thr	Val	
Ile Phe E	Pro Ser 275	Ser Hi	s Ser	Phe 280	Ile	Leu	Ile	Met	Gly 285	Asn	Ser	Lys	
Leu Arg 0 290	3lu Ala	Phe Le	1 Lys 295	Met	Leu	Arg	Phe	Val 300	Lys	Cys	Phe	Leu	
Arg Arg A	Arg Lys	Pro Pho 31		Pro									
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attctaatt	a tggga	aaatag (caagtt	gagg	g gaa	agctt	ttc	tgaa	agato	gtt a	aagat	ttgtg	900

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Val Met Leu Ala Ile Leu Leu Gly Ser Phe Leu Ile Ser Leu Ile Ile

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<210> SEQ ID NO 39

<211> LENGTH: 303

<212> TYPE: PRT

<213> ORGANISM: Homo sapiens

<400> SEQUENCE: 39

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1 5 10 15

Phe Ile Ile Gly Asn Leu Ser Asn Gly Phe Ile Val Leu Ile Asn Cys $20 \\ 25 \\ 30$

Ile Asp Trp Val Ser Lys Arg Glu Leu Ser Ser Val Asp Lys Leu Leu 35 40 45

Ser Trp Phe Leu Ala Leu His Tyr Leu Ala Ile Phe Val Ser Gly Thr 65 70 75 80

Gly Leu Arg Ile Met Ile Phe Ser Trp Ile Val Ser Asn His Phe Asn 85 90 95

Leu Trp Leu Ala Thr Ile Phe Ser Ile Phe Tyr Leu Leu Lys Ile Ala $100 \hspace{1.5cm} 105 \hspace{1.5cm} 110$

Ser Phe Ser Ser Pro Ala Phe Leu Tyr Leu Lys Trp Arg Val Asn Lys

Val Ile Leu Met Ile Leu Leu Gly Thr Leu Val Phe Leu Phe Leu Asn 130 135

Leu Ile Gln Ile Asn Met His Ile Lys Asp Trp Leu Asp Arg Tyr Glu 145 \$150\$

Arg Asn Thr Thr Trp Asn Phe Ser Met Ser Asp Phe Glu Thr Phe Ser

Val Ser Val Lys Phe Thr Met Thr Met Phe Ser Leu Thr Pro Phe Thr 180 185 190

Val Ala Phe Ile Ser Phe Leu Leu Leu Ile Phe Ser Leu Gln Lys His Leu Gln Lys Met Gln Leu Asn Tyr Lys Gly His Arg Asp Pro Arg Thr Lys Val His Thr Asn Ala Leu Lys Ile Val Ile Ser Phe Leu Leu Phe 235 Tyr Ala Ser Phe Phe Leu Cys Val Leu Ile Ser Trp Ile Ser Glu Leu 250 Tyr Gln Ser Thr Val Ile Tyr Met Leu Cys Glu Thr Ile Gly Val Phe 260 265 Ser Pro Ser Ser His Ser Phe Leu Leu Ile Leu Gly Asn Ala Lys Leu 280 Arg Gln Ala Phe Leu Leu Val Ala Ala Lys Val Trp Ala Lys Arg 290 295 <210> SEQ ID NO 40 <211> LENGTH: 909 <212> TYPE: DNA <213> ORGANISM: Homo sapiens <400> SEQUENCE: 40 atqqaaaqtq ccctqccqaq tatcttcact cttqtaataa ttqcaqaatt cataattqqq aatttgagca atggatttat agtactgatc aactgcattg actgggtcag taaaagagag 120 ctgtcctcag tcgataaact cctcattatc ttggcaatct ccagaattgg gctgatctgg 180 gaaatattag taagttggtt tttagctctg cattatctag ccatatttgt gtctggaaca ggattaagaa ttatgatttt tagctggata gtttctaatc acttcaatct ctggcttgct acaatettea geatetttta tttgeteaaa atagegagtt tetetageee tgetttete tatttgaagt ggagagtaaa caaagtgatt ctgatgatac tgctaggaac cttggtcttc ttatttttaa atctgataca aataaacatg catataaaag actggctgga ccgatatgaa agaaacacaa cttggaattt cagtatgagt gactttgaaa cattttcagt gtcggtcaaa ttcactatga ctatgttcag tctaacacca tttactgtgg ccttcatctc ttttctcctg ttaattttct ccctgcagaa acatctccag aaaatgcaac tcaattacaa aggacacaga 660 gaccccagga ccaaggtcca tacaaatgcc ttgaaaattg tgatctcatt ccttttattc 720 tatgctagtt tctttctatg tgttctcata tcatggattt ctgagctgta tcagagcaca 780 gtgatctaca tgctttgtga gacgattgga gtcttctctc cttcaagcca ctcctttctt 840 ctgattctag gaaacgctaa gttaagacag gcctttcttt tggtggcagc taaggtatgg 900 qctaaacga 909 <210> SEQ ID NO 41 <211> LENGTH: 317 <212> TYPE: PRT <213> ORGANISM: Homo sapiens <400> SEQUENCE: 41 Met Gly Gly Val Ile Lys Ser Ile Phe Thr Phe Val Leu Ile Val Glu Phe Ile Ile Gly Asn Leu Gly Asn Ser Phe Ile Ala Leu Val Asn Cys

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Thr Ala 50	Leu	Ala	Ile	Ser	Arg 55	Ile	Ser	Leu	Val	Trp 60	Leu	Ile	Phe	Gly	
Ser Trp 65	Сув	Val	Ser	Val 70	Phe	Phe	Pro	Ala	Leu 75	Phe	Ala	Thr	Glu	80 Tàs	
Met Phe	Arg	Met	Leu 85	Thr	Asn	Ile	Trp	Thr 90	Val	Ile	Asn	His	Phe 95	Ser	
Val Trp	Leu	Ala 100	Thr	Gly	Leu	Gly	Thr 105	Phe	Tyr	Phe	Leu	Lys 110	Ile	Ala	
Asn Phe	Ser 115	Asn	Ser	Ile	Phe	Leu 120	Tyr	Leu	Lys	Trp	Arg 125	Val	ГÀа	TÀa	
Val Val 130	Leu	Val	Leu	Leu	Leu 135	Val	Thr	Ser	Val	Phe 140	Leu	Phe	Leu	Asn	
Ile Ala 145	Leu	Ile	Asn	Ile 150	His	Ile	Asn	Ala	Ser 155	Ile	Asn	Gly	Tyr	Arg 160	
Arg Asn	Lys	Thr	Сув 165	Ser	Ser	Asp	Ser	Ser 170	Asn	Phe	Thr	Arg	Phe 175	Ser	
Ser Leu	Ile	Val 180	Leu	Thr	Ser	Thr	Val 185	Phe	Ile	Phe	Ile	Pro 190	Phe	Thr	
Leu Ser	Leu 195	Ala	Met	Phe	Leu	Leu 200	Leu	Ile	Phe	Ser	Met 205	Trp	Lys	His	
Arg Lys 210	Lys	Met	Gln	His	Thr 215	Val	Lys	Ile	Ser	Gly 220	Asp	Ala	Ser	Thr	
Lys Ala 225	His	Arg	Gly	Val 230	Lys	Ser	Val	Ile	Thr 235	Phe	Phe	Leu	Leu	Tyr 240	
Ala Ile	Phe	Ser	Leu 245	Ser	Phe	Phe	Ile	Ser 250	Val	Trp	Thr	Ser	Glu 255	Arg	
Leu Glu	Glu	Asn 260	Leu	Ile	Ile	Leu	Ser 265	Gln	Val	Met	Gly	Met 270	Ala	Tyr	
Pro Ser	Сув 275	His	Ser	CAa	Val	Leu 280	Ile	Leu	Gly	Asn	Lys 285	Lys	Leu	Arg	
Gln Ala 290	Ser	Leu	Ser	Val	Leu 295	Leu	Trp	Leu	Arg	Tyr 300	Met	Phe	Lys	Asp	
Gly Glu 305	Pro	Ser	Gly	His 310	ГÀа	Glu	Phe	Arg	Glu 315	Ser	Ser				
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atctcttc ttaatatt															180 240
atgttcag															300
acaggeet															360
tacctaaa	ıgt (ggaga	agtta	aa aa	aaggt	ggtt	ttg	ggtg	ctgc	ttci	tgt	gac 1	teg	gtette	420

ttgt	tttt	aa a	atatt	gcad	et ga	ataaa	acato	cat	ataa	atg	ccaç	gtato	caa t	ggat	acaga
agaa	acaa	iga d	cttgo	cagtt	c to	gatto	caagt	aac	ttta	cac	gatt	ttco	ag t	ctta	ittgta
ttaa	eccaç	jca (ctgtg	gttca	at tt	tcat	acco	ttt	actt	tgt	ccct	ggca	aat g	gtttc	ttctc
ctca	atctt	ct o	ccato	gtgga	aa a	catco	gcaag	g aag	gatgo	agc	acac	etgto	caa a	atat	ccgga
gaco	gccag	jca (ccaaa	agcco	ca ca	agagg	gagtt	aaa	agto	jtga	tcac	tttc	ett o	ctac	tctat
gcca	tttt	ct o	ctctç	gtctt	t tt	tcat	atca	gtt	tgga	cct	ctga	aagg	gtt g	ggagg	jaaaat
ctaa	attat	tc t	ttc	ccago	gt ga	atggg	gaato	g gct	tato	ctt	cato	gtcac	etc a	atgto	ıttctg
atto	ettgg	jaa a	acaaq	gaago	et ga	agaca	aggco	tct	ctgt	cag	tgct	acto	gtg g	gctga	ıggtac
atgt	tcaa	ag a	atggg	ggago	ec et	cago	gtcac	aaa	ıgaat	tta	gaga	atca	atc t	;	
<211 <212 <213	.> LE ?> TY ß> OF	NGTH PE:	SM:	1 Homo	sap	oiens	ţ								
		~	ICE:			m1	77-7	Dl	Dl	M-+	T1 -	T 1 -	m	*** 7	T
Met 1	IIe	Pro	IIe	Gln 5	Leu	Thr	vai	Pne	Pne 10	Met	11e	IIe	Tyr	va1 15	Leu
Glu	Ser	Leu	Thr 20	Ile	Ile	Val	Gln	Ser 25	Ser	Leu	Ile	Val	Ala 30	Val	Leu
Gly	Arg	Glu 35	Trp	Leu	Gln	Val	Arg 40	Arg	Leu	Met	Pro	Val 45	Asp	Met	Ile
Leu	Ile 50	Ser	Leu	Gly	Ile	Ser 55	Arg	Phe	Cha	Leu	Gln 60	Trp	Ala	Ser	Met
Leu 65	Asn	Asn	Phe	CÀa	Ser 70	Tyr	Phe	Asn	Leu	Asn 75	Tyr	Val	Leu	Cys	Asn 80
Leu	Thr	Ile	Thr	Trp 85	Glu	Phe	Phe	Asn	Ile 90	Leu	Thr	Phe	Trp	Leu 95	Asn
Ser	Leu	Leu	Thr 100	Val	Phe	Tyr	Cys	Ile 105	ГЛа	Val	Ser	Ser	Phe 110	Thr	His
His	Ile	Phe 115	Leu	Trp	Leu	Arg	Trp 120	Arg	Ile	Leu	Arg	Leu 125	Phe	Pro	Trp
Ile	Leu 130	Leu	Gly	Ser	Leu	Met 135	Ile	Thr	Cys	Val	Thr 140	Ile	Ile	Pro	Ser
Ala 145	Ile	Gly	Asn	Tyr	Ile 150	Gln	Ile	Gln	Leu	Leu 155	Thr	Met	Glu	His	Leu 160
Pro	Arg	Asn	Ser	Thr 165	Val	Thr	Asp	Lys	Leu 170	Glu	Asn	Phe	His	Gln 175	Tyr
Gln	Phe	Gln	Ala 180	His	Thr	Val	Ala	Leu 185	Val	Ile	Pro	Phe	Ile 190	Leu	Phe
Leu	Ala	Ser 195	Thr	Ile	Phe	Leu	Met 200	Ala	Ser	Leu	Thr	Lys 205	Gln	Ile	Gln
His	His 210	Ser	Thr	Gly	His	Cys 215	Asn	Pro	Ser	Met	Lys 220	Ala	His	Phe	Thr
Ala 225	Leu	Arg	Ser	Leu	Ala 230	Val	Leu	Phe	Ile	Val 235	Phe	Thr	Ser	Tyr	Phe 240
Leu	Thr	Ile	Leu	Ile 245	Thr	Ile	Ile	Gly	Thr 250	Leu	Phe	Asp	Lys	Arg 255	Cys

Trp Leu Trp Val Trp Glu Ala Phe Val Tyr Ala Phe Ile Leu Met His $260 \hspace{1cm} 265 \hspace{1cm} 265 \hspace{1cm} 270 \hspace{1cm}$

Ser Thr Ser Leu Met Leu Ser Ser Pro Thr Leu Lys Arg Ile Leu Lys Gly Lys Cys 290 <210> SEQ ID NO 44 <211> LENGTH: 873 <212> TYPE: DNA <213> ORGANISM: Homo sapiens <400> SEOUENCE: 44 atgataccca tccaactcac tgtcttcttc atgatcatct atgtgcttga gtccttgaca 60 attattgtgc agagcagcct aattgttgca gtgctgggca gagaatggct gcaagtcaga 120 aggetgatge etgtggaeat gatteteate ageetgggea tetetegett etgtetaeag 180 tgggcatcaa tgctgaacaa tttttgctcc tattttaatt tgaattatgt actttgcaac 240 ttaacaatca cctgggaatt ttttaatatc cttacattct ggttaaacag cttgcttacc 300 gtgttctact gcatcaaggt ctcttctttc acccatcaca tctttctctg gctgaggtgg 360 agaattttga ggttgtttcc ctggatatta ctgggttctc tgatgattac ttgtgtaaca 420 atcatccctt cagctattgg gaattacatt caaattcagt tactcaccat ggagcatcta 480 ccaagaaaca gcactgtaac tgacaaactt gaaaattttc atcagtatca gttccaggct 540 catacagttg cattggttat teettteate etgtteetgg cetecaceat ettteteatg 600 gcatcactga ccaagcagat acaacatcat agcactggtc actgcaatcc aagcatgaaa gcgcacttca ctgccctgag gtcccttgcc gtcttattta ttgtgtttac ctcttacttt ctaaccatac tcatcaccat tataggtact ctatttgata agagatgttg gttatgggtc tgggaagett ttgtctatge tttcatetta atgeatteea etteaetgat getgageage 840 cctacgttga aaaggattct aaagggaaag tgc 873 <210> SEQ ID NO 45 <211> LENGTH: 316 <212> TYPE: PRT <213> ORGANISM: Homo sapiens <400> SEOUENCE: 45 Met Met Gly Leu Thr Glu Gly Val Phe Leu Ile Leu Ser Gly Thr Gln 10 Phe Thr Leu Gly Ile Leu Val Asn Cys Phe Ile Glu Leu Val Asn Gly Ser Ser Trp Phe Lys Thr Lys Arg Met Ser Leu Ser Asp Phe Ile Ile 40 Thr Thr Leu Ala Leu Leu Arg Ile Ile Leu Leu Cys Ile Ile Leu Thr Asp Ser Phe Leu Ile Glu Phe Ser Pro Asn Thr His Asp Ser Gly Ile 75 Ile Met Gln Ile Ile Asp Val Ser Trp Thr Phe Thr Asn His Leu Ser Ile Trp Leu Ala Thr Cys Leu Gly Val Leu Tyr Cys Leu Lys Ile Ala Ser Phe Ser His Pro Thr Phe Leu Trp Leu Lys Trp Arg Val Ser Arg

Val	Met 130	Val	Trp	Met	Leu	Leu 135	Gly	Ala	Leu	Leu	Leu 140	Ser	Сув	Gly	Ser		
Thr 145	Ala	Ser	Leu	Ile	Asn 150	Glu	Phe	Lys	Leu	Tyr 155	Ser	Val	Phe	Arg	Gly 160		
Ile	Glu	Ala	Thr	Arg 165	Asn	Val	Thr	Glu	His 170	Phe	Arg	Lys	Lys	Arg 175	Ser		
Glu	Tyr	Tyr	Leu 180	Ile	His	Val	Leu	Gly 185	Thr	Leu	Trp	Tyr	Leu 190	Pro	Pro		
Leu	Ile	Val 195	Ser	Leu	Ala	Ser	Tyr 200	Ser	Leu	Leu	Ile	Phe 205	Ser	Leu	Gly		
Arg	His 210	Thr	Arg	Gln	Met	Leu 215	Gln	Asn	Gly	Thr	Ser 220	Ser	Arg	Asp	Pro		
Thr 225	Thr	Glu	Ala	His	Lys 230	Arg	Ala	Ile	Arg	Ile 235	Ile	Leu	Ser	Phe	Phe 240		
Phe	Leu	Phe	Leu	Leu 245	Tyr	Phe	Leu	Ala	Phe 250	Leu	Ile	Ala	Ser	Phe 255	Gly		
Asn	Phe	Leu	Pro 260	Lys	Thr	Lys	Met	Ala 265	Lys	Met	Ile	Gly	Glu 270	Val	Met		
Thr	Met	Phe 275	Tyr	Pro	Ala	Gly	His 280	Ser	Phe	Ile	Leu	Ile 285	Leu	Gly	Asn		
Ser	Lys 290	Leu	Lys	Gln	Thr	Phe 295	Val	Val	Met	Leu	Arg 300	Cys	Glu	Ser	Gly		
His 305	Leu	ГЛа	Pro	Gly	Ser 310	ГЛа	Gly	Pro	Ile	Phe 315	Ser						
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atto	tggt	ca a	attgt	ttca	at to	gagtt	ggto	c aat	ggta	agca	gct	ggtto	caa 🤅	gacca	agaga	12	0
atgt	cttt	gt o	ctgad	cttca	at ca	atcad	ccaco	c ctç	ggcad	ctct	tgag	ggato	cat 1	tatga	tgtgt	18	0
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ataa	itgca	aaa t	tatt	gato	gt ti	cat	ggaca	a ttt	acaa	aacc	atct	gago	cat 1	ttggd	ettgee	30	0
acct	gtct	tg g	gtgto	cctct	a ct	gcct	gaaa	a ato	egeca	agtt	tcto	ctcac	ccc (cacat	tecte	36	0
tggc	ctcaa	agt g	ggaga	agtt	ic ta	agggt	gato	g gta	atgga	atgc	tgtt	gggt	gc a	actgo	ctctta	42	0
tect	gtgg	gta g	gtaco	cgcat	c to	ctgat	caat	gag	gttta	aagc	tcta	attct	gt (cttta	agggga	48	0
attg	gaggo	cca c	ccago	gaat	gt ga	actga	aacac	tto	cagaa	aaga	agag	ggagt	ga 🤅	gtatt	atctg	54	0
atco	catgt	tc t	tggg	gacto	et gt	ggta	accto	g cct	ccct	taa	ttgt	gtc	ect (ggcct	cctac	60	0
tctt	tgct	ca t	ctto	ctcc	ct g	gggag	ggcac	c aca	acggo	caga	tgct	gcaa	aaa 1	tggga	acaagc	66	0
tcca	agaga	atc o	caaco	cacto	ga go	gecea	acaaç	g agg	ggcca	atca	gaat	cato	ect 1	ttcct	tette	72	0
tttc	ctctt	ct t	actt	tact	t to	cttgo	ctttc	tta	aatto	gcat	catt	tggt	caa 1	tttco	ctacca	78	0
aaaa	eccaa	aga t	ggct	caaga	at ga	attg	gcgaa	a gta	aatga	acaa	tgtt	ttat	ccc t	tgctg	ggccac	84	0
tcat	ttat	tc t	catt	ctg	gg ga	aacaç	gtaag	g cto	gaago	caga	catt	tgta	agt q	gatgo	ctccgg	90	0
tgtg	gagto	ctg g	gtcat	ctga	aa go	cctg	gatco	c aag	gggad	ccca	ttt	ctct	:			94	8

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gtcttctcag	ctgacttcat	cctcacctgc	ttggctatct	ccacaattgg	acaactgttg	180
gtgatactgt	ttgattcatt	tctagtggga	cttgcttcac	atttatatac	cacatataga	240
ctaggaaaaa	ctgttattat	gctttggcac	atgactaatc	acttgacaac	ctggcttgcc	300
acctgcctaa	gcattttcta	tttctttaag	atageceact	tcccccactc	ccttttcctc	360
tggctgaggt	ggaggatgaa	cggaatgatt	gttatgcttc	ttatattgtc	tttgttctta	420
ctgatttttg	acagtttagt	gctagaaata	tttattgata	tctcactcaa	tataatagat	480
aaaagtaatc	tgactttata	tttagatgaa	agtaaaactc	tctttgataa	actctctatt	540
ttaaaaactc	ttctcagctt	gaccagtttt	atcccctttt	ctctgtccct	gacctccttg	600
ctttttttat	ttctgtcctt	ggtgagacat	actagaaatt	tgaagctcag	ttccttgggc	660
tctagagact	ccagcacaga	ggcccatagg	agggccatga	aaatggtgat	gtctttcctt	720
ttcctcttca	tagttcattt	tttttcctta	caagtggcca	attggatatt	ttttatgttg	780
tggaacaaca	agtacataaa	gtttgtcatg	ttagccttaa	atgcctttcc	ctcgtgccac	840
tcatttattc	tcattctggg	aaacagcaag	ctgcgacaga	cagctgtgag	gctactgtgg	900
catcttagga	actatacaaa	aacaccaaat	gctttacctt	tg		942

<210> SEQ ID NO 49

<211> LENGTH: 318

<212> TYPE: PRT

<213> ORGANISM: Homo sapiens

<400> SEQUENCE: 49

Met Asn Gly Asp His Met Val Leu Gly Ser Ser Val Thr Asp Lys Lys 1 $$ 10 $$ 15

Ala Ile Ile Leu Val Thr Ile Leu Leu Leu Leu Arg Leu Val Ala Ile 20 \$25\$

Ala Gly Asn Gly Phe Ile Thr Ala Ala Leu Gly Val Glu Trp Val Leu $35 \hspace{1.5cm} 40 \hspace{1.5cm} 45 \hspace{1.5cm}$

Arg Arg Met Leu Leu Pro Cys Asp Lys Leu Leu Val Ser Leu Gly Ala 50 60

Ser Arg Phe Cys Leu Gln Ser Val Val Met Gly Lys Thr Ile Tyr Val 65 70 75 80

Phe Leu His Pro Met Ala Phe Pro Tyr Asn Pro Val Leu Gln Phe Leu 85 90 95

Ala Phe Gln Trp Asp Phe Leu Asn Ala Ala Thr Leu Trp Ser Ser Thr $100 \hspace{1.5cm} 105 \hspace{1.5cm} 110 \hspace{1.5cm}$

Trp Leu Ser Val Phe Tyr Cys Val Lys Ile Ala Thr Phe Thr His Pro 115 120 125

Val Phe Phe Trp Leu Lys His Lys Leu Ser Gly Trp Leu Pro Trp Met 130 135 140

Leu Phe Ser Ser Val Gly Leu Ser Ser Phe Thr Thr Ile Leu Phe Phe 145 150 150

Ile Gly Asn His Arg Met Tyr Gln Asn Tyr Leu Arg Asn His Leu Gln 165 170 175

Pro Trp Asn Val Thr Gly Asp Ser Ile Arg Ser Tyr Cys Glu Lys Phe \$180\$

Tyr Leu Phe Pro Leu Lys Met Ile Thr Trp Thr Met Pro Thr Ala Val Phe Phe Ile Cys Met Ile Leu Leu Ile Thr Ser Leu Gly Arg His Arg Lys Lys Ala Leu Leu Thr Thr Ser Gly Phe Arg Glu Pro Ser Val Gln Ala His Ile Lys Ala Leu Leu Ala Leu Leu Ser Phe Ala Met Leu Phe 250 Ile Ser Tyr Phe Leu Ser Leu Val Phe Ser Ala Ala Gly Ile Phe Pro 260 265 Pro Leu Asp Phe Lys Phe Trp Val Trp Glu Ser Val Ile Tyr Leu Cys 280 Ala Ala Val His Pro Ile Ile Leu Leu Phe Ser Asn Cys Arg Leu Arg 295 Ala Val Leu Lys Ser Arg Arg Ser Ser Arg Cys Gly Thr Pro 310 315 <210> SEO ID NO 50 <211> LENGTH: 957 <212> TYPE: DNA <213> ORGANISM: Homo sapiens <400> SEQUENCE: 50 atquatqqaq accacatqqt tctaqqatct tcqqtqactq acaaqaaqqc catcatcttq gttaccattt tactcctttt acgcctggta gcaatagcag gcaatggctt catcactgct gctctgggcg tggagtgggt gctacggaga atgttgttgc cttgtgataa gttattggtt agcctagggg cctctcgctt ctgtctgcag tcagtggtaa tgggtaagac catttatgtt ttcttgcatc cgatggcctt cccatacaac cctgtactgc agtttctagc tttccagtgg gactteetga atgetgeeac ettatggtee tetacetgge teagtgtett etattgtgtg aaaattgcta ccttcaccca ccctgtcttc ttctggctaa agcacaagtt gtctgggtgg 420 ctaccatgga tgctcttcag ctctgtaggg ctctccagct tcaccaccat tctatttttc ataggcaacc acagaatgta tcagaactat ttaaggaacc atctacaacc ttggaatgtc 540 actggcgata gcatacggag ctactgtgag aaattctatc tcttccctct aaaaatgatt 600 acttggacaa tgcccactgc tgtctttttc atttgcatga ttttgctcat cacatctctg 660 ggaagacaca ggaagaaggc tctccttaca acctcaggat tccgagagcc cagtgtgcag 720 gcacacataa aggetetget ggeteteete tettttgeca tgetetteat eteatattte 780 ctgtcactgg tgttcagtgc tgcaggtatt tttccacctc tggactttaa attctgggtg 840 tgggagtcag tgatttatct gtgtgcagca gttcacccca tcattctgct cttcagcaac 900 tgcaggctga gagctgtgct gaagagtcgt cgttcctcaa ggtgtgggac accttga 957 <210> SEQ ID NO 51 <211> LENGTH: 33 <212> TYPE: DNA <213> ORGANISM: Artificial <220> FEATURE: <223> OTHER INFORMATION: Primer comprising EcoRI restriction site

for PCR amplification of hTAS2R16

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<400> SEQUENCE: 52	
gaagcgcgct ttcatgctt	19

We claim:

- 1. A polynucleotide selected from the group consisting of:
- (a) polynucleotides encoding at least the mature form of the polypeptide having the deduced amino acid sequence as shown in SEQ ID NO: 12;
- (b) polynucleotides having the coding sequence, as shown in SEQ ID NO: 11 encoding at least the mature form of the polypeptide;
- (c) polynucleotides encoding a fragment or derivative of a polypeptide encoded by a polynucleotide of any one of (a) to (b), wherein in said derivative one or more amino acid residues are conservatively substituted compared to said polypeptide, and said fragment or derivative has bitter substance binding activity;
- (d) polynucleotides which are at least 50% identical to a polynucleotide as defined in any one of (a) to (c) and which code for a polypeptide having bitter substance binding activity; and
- (e) polynucleotides the complementary strand of which hybridizes under stringent conditions to a polynucleotide as defined in any one of (a) to (d) and which code for a polypeptide having bitter substance binding activity;
- or the complementary strand of such a polynucleotide.
- 2. The polynucleotide of claim 1 which is DNA, genomic DNA or RNA.
 - 3. A vector containing the polynucleotide of claim 1.
- **4.** The vector of claim **3** in which the polynucleotide is operatively linked to one or more expression control sequences allowing expression in prokaryotic and/or eukaryotic host cells.
- 5. A host cell genetically engineered to comprise a polynucleotide of claim 1.
- **6.** A transgenic non-human animal comprising a polynucleotide of claim **1**.
- 7. A process for producing a polypeptide encoded by a polynucleotide of claim 1, comprising: culturing a host cell genetically engineered to comprises a polynucleotide of claim 1 and recovering the polypeptide encoded by said polynucleotide.
- **8.** A process for producing cells capable of expressing at least one bitter taste receptor polypeptide comprising genetically engineering cells in vitro with a vector containing a polynucleotide of claim 1, wherein said bitter taste receptor polypeptide(s) is(are) encoded by a polynucleotide of claim 1.
- **9**. A polypeptide having the amino acid sequence encoded by a polynucleotide selected from the group consisting of:

- (a) polynucleotides encoding at least the mature form of the polypeptide having the deduced amino acid sequence as shown in SEQ ID NO: 12;
- (b) polynucleotides having the coding sequence, as shown in SEQ ID NO: 11 encoding at least the mature form of the polypeptide;
- (c) polynucleotides encoding a fragment or derivative of a polypeptide encoded by a polynucleotide of any one of (a) to (b), wherein in said derivative one or more amino acid residues are conservatively substituted compared to said polypeptide, and said fragment or derivative has bitter substance binding activity;
- (d) polynucleotides which are at least 50% identical to a polynucleotide as defined in any one of (a) to (c) and which code for a polypeptide having bitter substance binding activity; and
- (e) polynucleotides the complementary strand of which hybridizes under stringent conditions to a polynucleotide as defined in any one of (a) to (d) and which code for a polypeptide having bitter substance binding activity;
- or the complementary strand of such a polynucleotide; wherein said polypeptide is obtainable by culturing a host cell genetically engineered to comprise a polynucleotide as defined in any of (a) to (e), and recovering the polypeptide encoded by said polynucleotide.
- 10. An antibody that specifically binds to the polypeptide of claim 9.
- 11. A nucleic acid molecule which specifically hybridizes to a polynucleotide of claim 1.
- 12. An antagonist/inhibitor against the polypeptide of claim 9 which is an antibody, the extracellular domain of the polypeptide of claim 9 or a fragment thereof, or an inhibiting RNA.
- 13. The antagonist/inhibitor of claim 12, wherein said inhibiting RNA is an antisense construct, RNAi, siRNA or a ribozyme.
- **14.** A process for isolating a compound that binds to a polypeptide encoded by a polynucleotide selected from the group consisting of:
 - (a) polynucleotides encoding at least the mature form of the polypeptide having the deduced amino acid sequence as shown in SEQ ID NO: 12;
 - (b) polynucleotides having the coding sequence, as shown in SEQ ID NO: 11 encoding at least the mature form of the polypeptide;
 - (c) polynucleotides encoding a fragment or derivative of a polypeptide encoded by a polynucleotide of any one of (a) to (b), wherein in said derivative one or more amino acid residues are conservatively substituted compared to

- said polypeptide, and said fragment or derivative has bitter substance binding activity;
- (d) polynucleotides which are at least 50% identical to a polynucleotide as defined in any one of (a) to (c) and which code for a polypeptide having bitter substance binding activity; and
- (e) polynucleotides the complementary strand of which hybridizes under stringent conditions to a polynucleotide as defined in any one of (a) to (d) and which code for a polypeptide having bitter substance binding activity;

comprising:

- contacting said polypeptide or a host cell genetically engineered with said polynucleotide or with a vector containing said polynucleotide with a compound;
- (2) detecting the presence of the compound which binds to said polypeptide; and
- determining whether the compound binds said polypeptide.
- 15. A process for isolating an antagonist of the bitter taste receptor activity of the polypeptide encoded by a polynucle-otide selected from the group consisting of:
 - (a) polynucleotides encoding at least the mature form of the polypeptide having the deduced amino acid sequence as shown in SEQ ID NO: 12;
 - (b) polynucleotides having the coding sequence, as shown in SEQ ID NO: 11 encoding at least the mature form of the polypeptide;
 - (c) polynucleotides encoding a fragment or derivative of a polypeptide encoded by a polynucleotide of any one of (a) to (b), wherein in said derivative one or more amino acid residues are conservatively substituted compared to said polypeptide, and said fragment or derivative has bitter taste receptor activity;
 - (d) polynucleotides which are at least 50% identical to a
 polynucleotide as defined in any one of (a) to (c) and
 which code for a polypeptide having bitter taste receptor
 activity; and
 - (e) polynucleotides the complementary strand of which hybridizes under stringent conditions to a polynucleotide as defined in any one of (a) to (d) and which code for a polypeptide having bitter taste receptor activity; comprising:
 - (1) contacting said polypeptide or a host cell genetically engineered with said polynucleotide or with a vector containing said polynucleotide with a potential antagonist; and
 - (2) determining whether the potential antagonist antagonizes the bitter taste receptor activity of said polypeptide
- 16. The process of claim 15 further comprising the contacting of the polypeptide with an agonist of the respective bitter taste receptor activity.
- 17. The process of claim 16 in which said contacting with an agonist is carried out prior to, concomitantly with, or after step (1).
- **18**. The process of claim **16** in which said agonist is selected from the group consisting of saccharin and acesulfame K and functional derivatives thereof.
- 19. A process for the production of a food or any precursor material or additive employed in the production of foodstuffs comprising the steps of the process of claim 14 and the subsequent step of admixing the identified compound or antago-

- nist with foodstuffs or any precursor material or additive employed in the production of foodstuffs.
- 20. A process for the production of a nutraceutical or pharmaceutical composition comprising the steps of the process of claim 14 and the subsequent step of formulating the compound or antagonist with an active agent in a pharmaceutically acceptable form.
- 21. A food stuff or any precursor material or additive employed in the production of foodstuffs comprising an antagonist/inhibitor of claim 12.
- 22. A nutraceutical or pharmaceutical composition comprising an antagonist/inhibitor of claim 12 and an active agent and optionally a pharmaceutically acceptable carrier.
- 23. A method for treating an abnormally increased or decreased sensitivity towards a bitter substance wherein said method comprises administering, to a subject in need of such treatment, one or more agents selected from the group consisting of:
 - 1) A polynucleotide selected from the group consisting of:
 - (a) polynucleotides encoding at least the mature form of the polypeptide having the deduced amino acid sequence as shown in SEQ ID NO: 12;
 - (b) polynucleotides having the coding sequence, as shown in SEQ ID NO: 11 encoding at least the mature form of the polypeptide;
 - (c) polynucleotides encoding a fragment or derivative of a polypeptide encoded by a polynucleotide of any one of (a) to (b), wherein in said derivative one or more amino acid residues are conservatively substituted compared to said polypeptide, and said fragment or derivative has bitter substance binding activity;
 - (d) polynucleotides which are at least 50% identical to a polynucleotide as defined in any one of (a) to (c) and which code for a polypeptide having bitter substance binding activity; and
 - (e) polynucleotides the complementary strand of which hybridizes under stringent conditions to a polynucleotide as defined in any one of (a) to (d) and which code for a polypeptide having bitter substance binding activity;
- or the complementary strand of such a polynucleotide;
 - 2) a vector comprising a polynucleotide of part 1) above;
 - 3) an antibody that specifically binds to a polypeptide that is encoded by a polynucleotide of part 1) above and which is obtainable by culturing a host cell genetically engineered to comprise a polynucleotide of part 1) above, and recovering the polypeptide encoded by said polynucleotide; and
 - 4) an antagonist/inhibitor against the polypeptide of part 3) above, which is an antibody, the extracellular domain of the polypeptide of part 3) above, or a fragment thereof, or an inhibiting RNA.
- **24**. A process for the production of a food or any precursor material or additive employed in the production of foodstuffs comprising the steps of the process of claim **15** and the subsequent step of admixing the identified compound or antagonist with foodstuffs or any precursor material or additive employed in the production of foodstuffs.
- 25. A process for the production of a nutraceutical or pharmaceutical composition comprising the steps of the process of claim 15 and the subsequent step of formulating the compound or antagonist with an active agent in a pharmaceutically acceptable form.

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