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(54) **BIOMARKER FOR COGNITIVE
DYSFUNCTION DISEASES, AND METHOD
FOR DETECTION OF COGNITIVE
DYSFUNCTION DISEASES USING THE
BIOMARKER**

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2333/974 (2013.01); **G01N 2800/60** (2013.01);
G01N 2800/2814 (2013.01)

(57) **ABSTRACT**

The present invention aims to provide methods to detect cognitive impairment including mild cognitive impairment and Alzheimer disease by using a protein or its partial peptide that differs in presence or absence, or in quantity between non-cognitive impairment and patients with cognitive impairment and further aims to present biomarkers comprising said protein and said partial peptide to be used to detect cognitive impairment including Alzheimer disease or mild cognitive impairment. Specifically, a biomarker for diagnosis of psychiatry disease or cognitive impairment comprising protein fragment or peptide of not less than 5 amino acid residues arising from at least one protein or peptide selected from the group of proteins consisting of amino acid sequence expressed by SEQ ID NOS: 1, 3, 5, 7, 9, 11, 13, 15, 17, 19, 21, 23, and 25 and selected from the group of partial peptide in these proteins consisting of amino acid sequence expressed by SEQ ID NOS: 2, 4, 6, 8, 10, 12, 14, 16, 18, 20, 22, 24, 26, and 27. And further aims to provide diagnostic method using these biomarker.

FIG.1

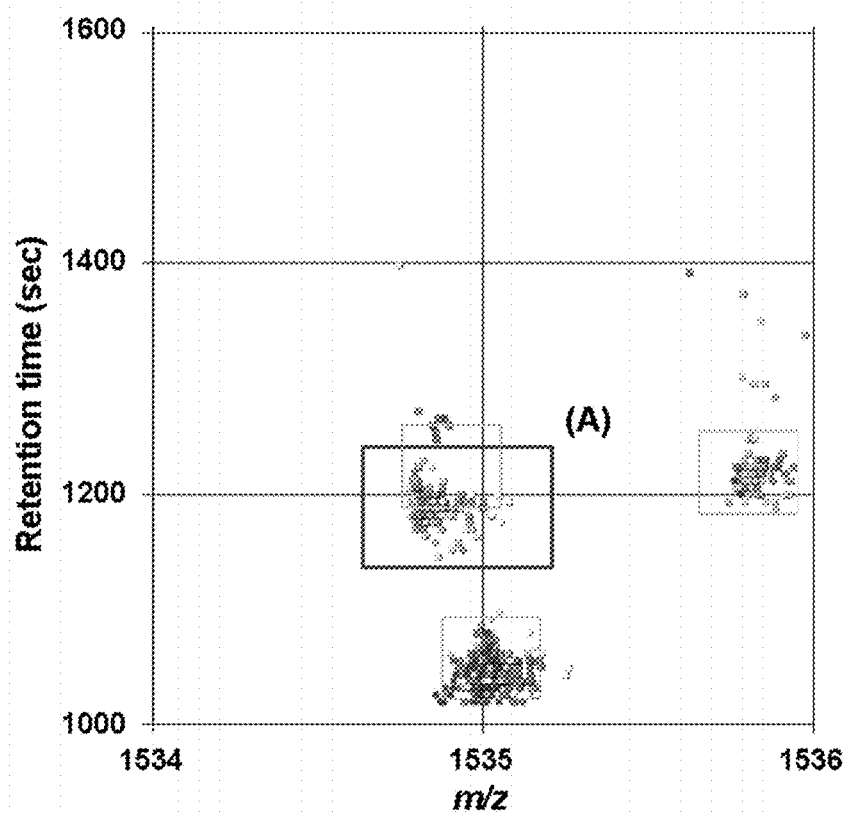
Marker A = CO₃

FIG.2

Marker A = CO3

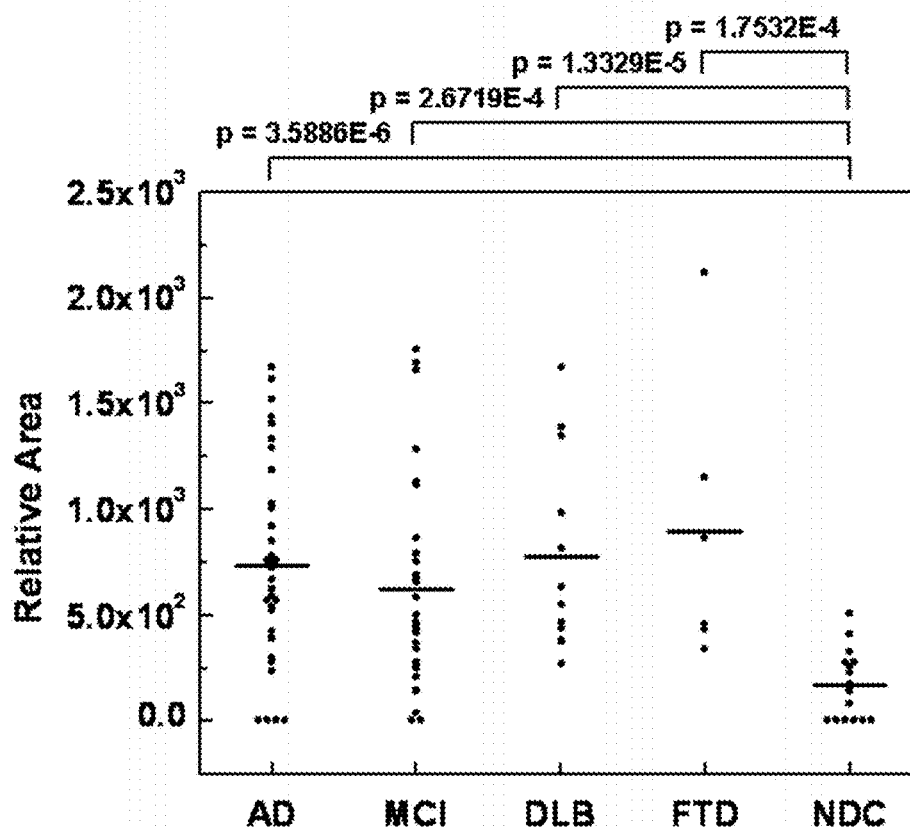


FIG.3

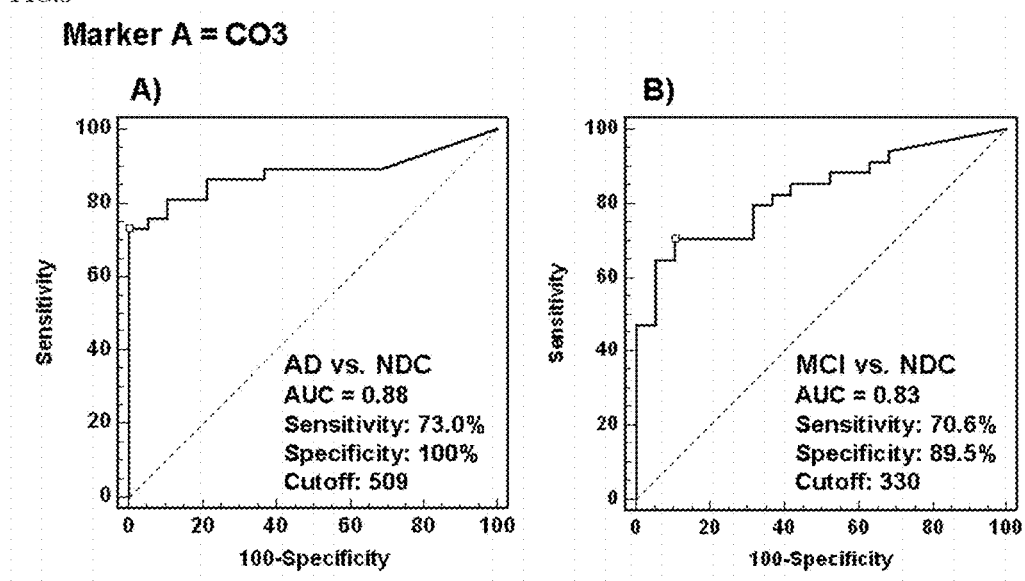


FIG. 4

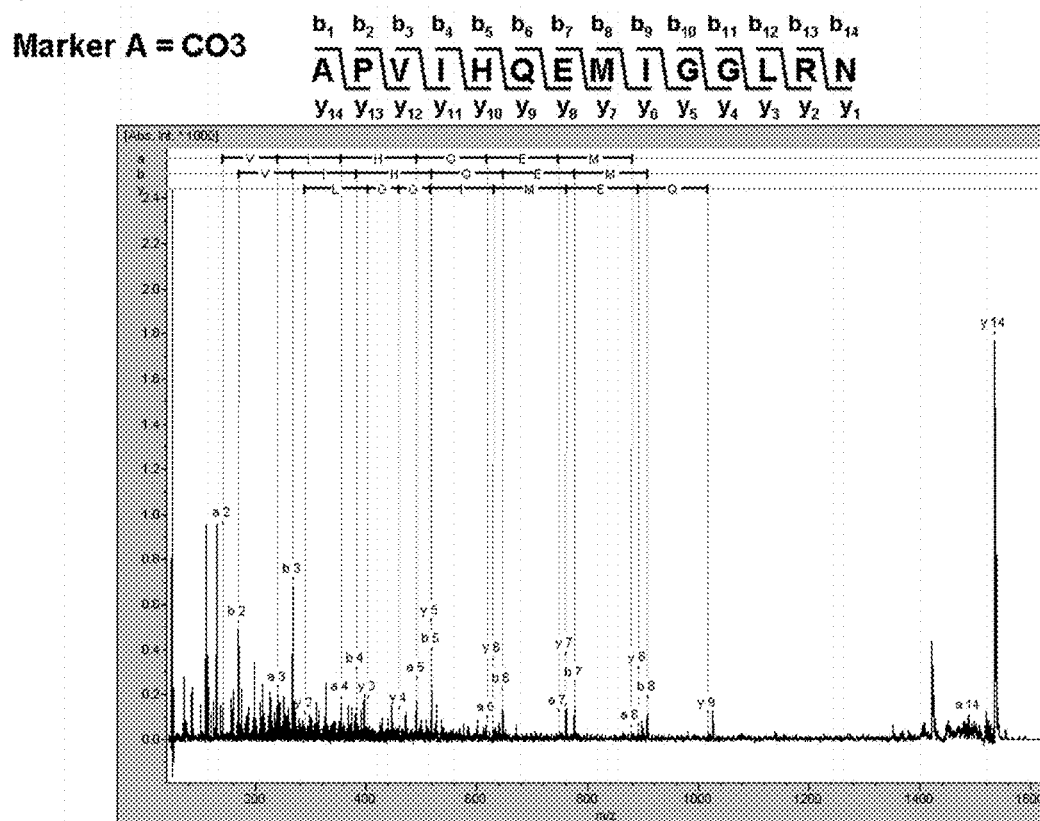


FIG.5

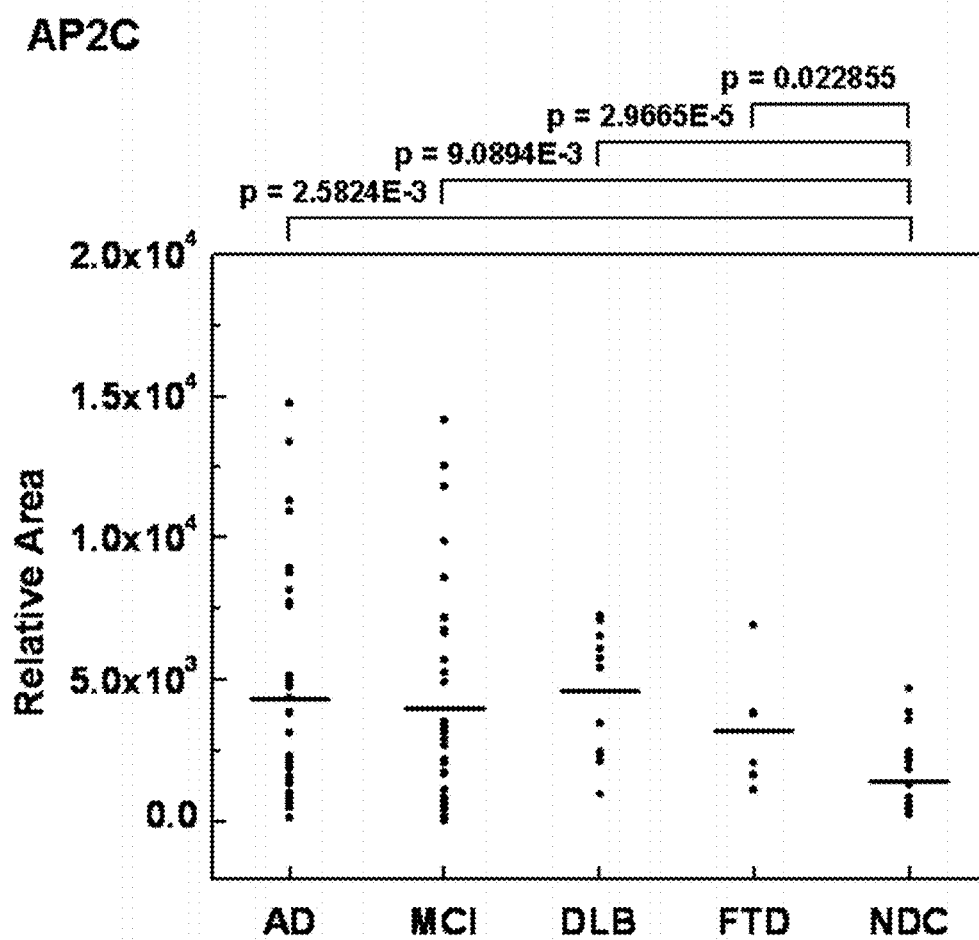


FIG.6

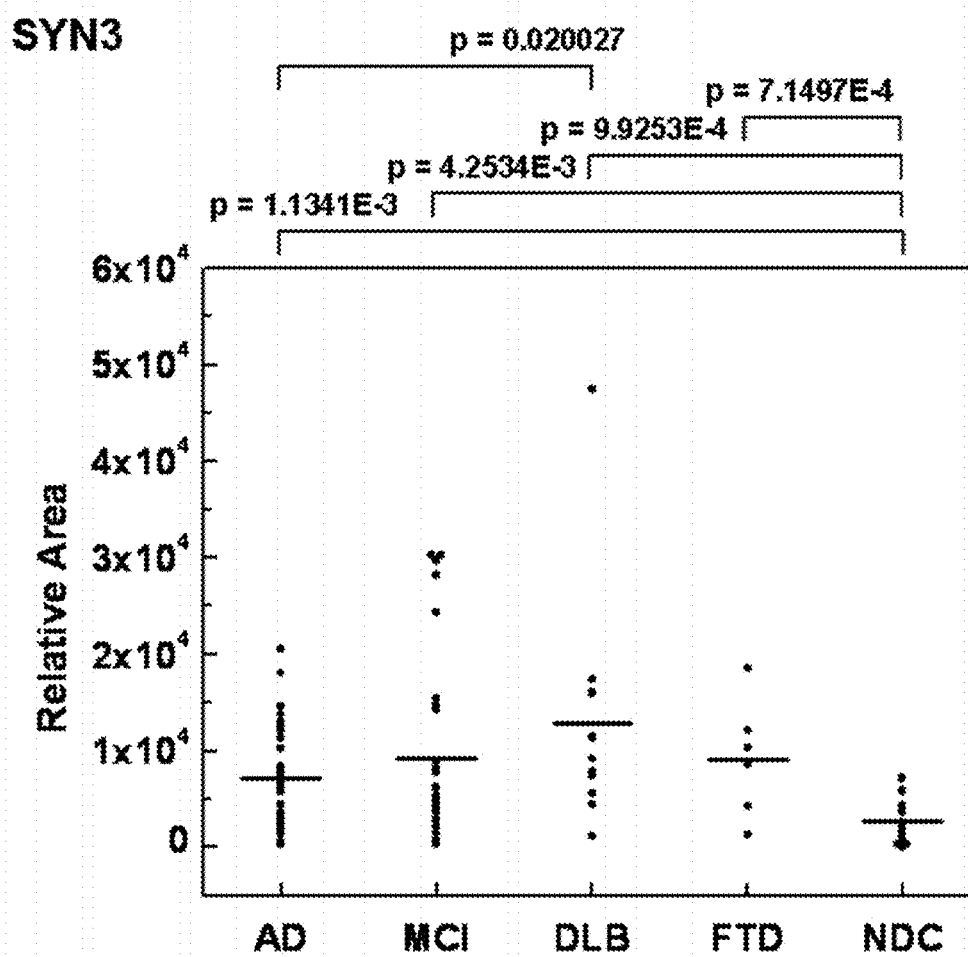


FIG. 7

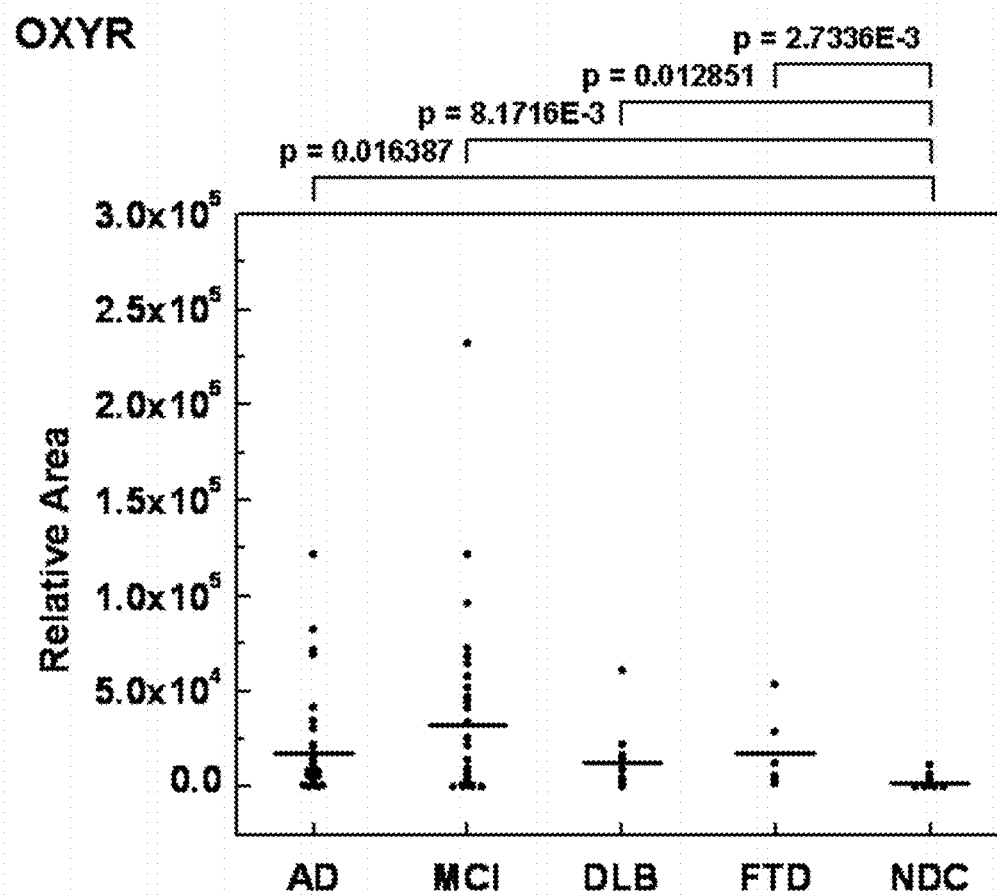


FIG.8

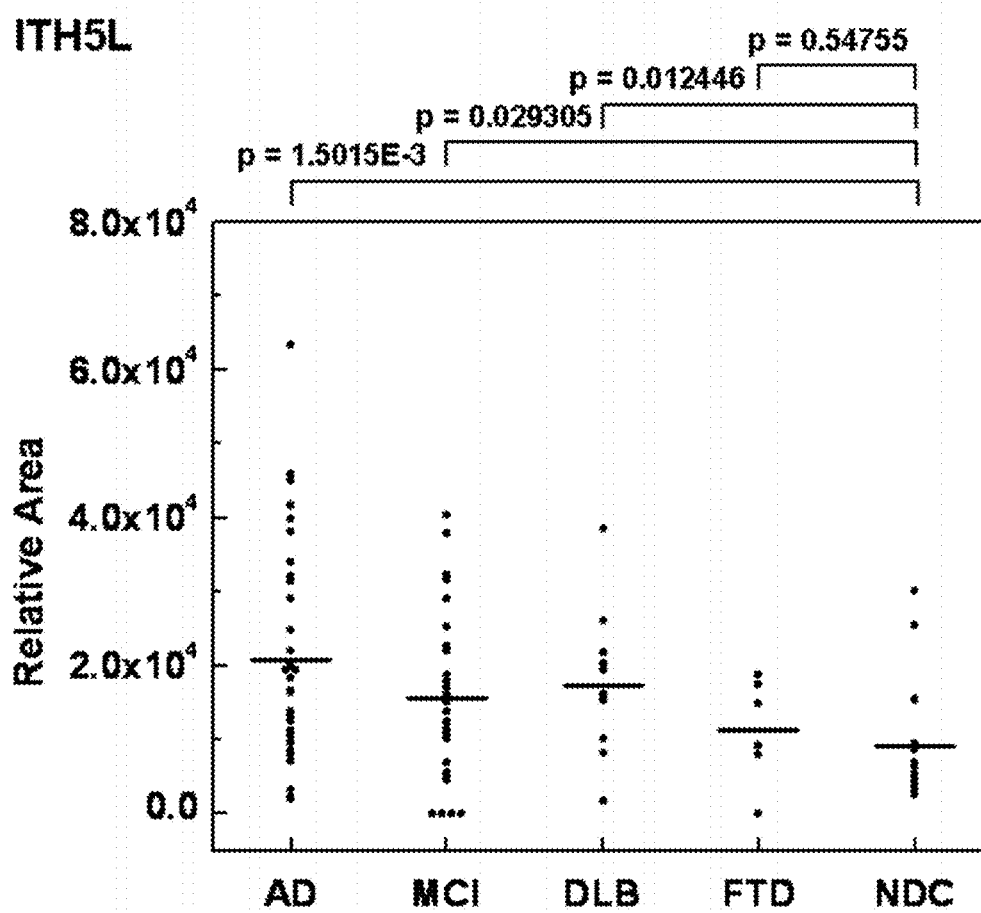


FIG.9

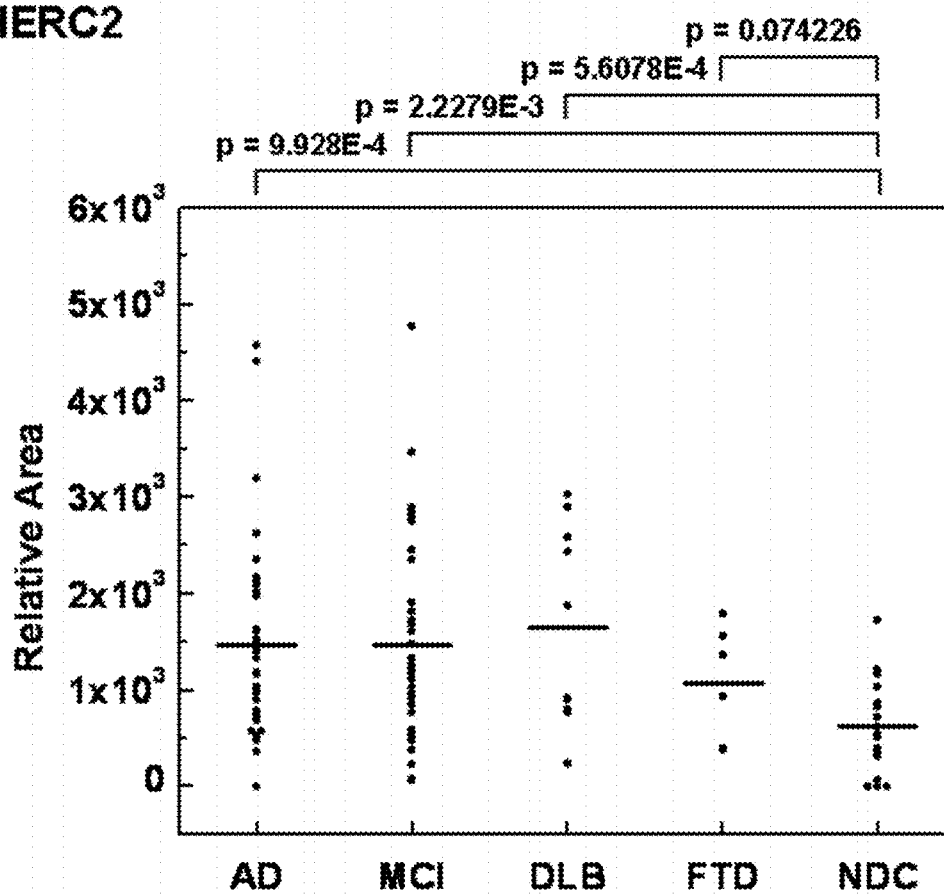
HERC2

FIG.10

THRB

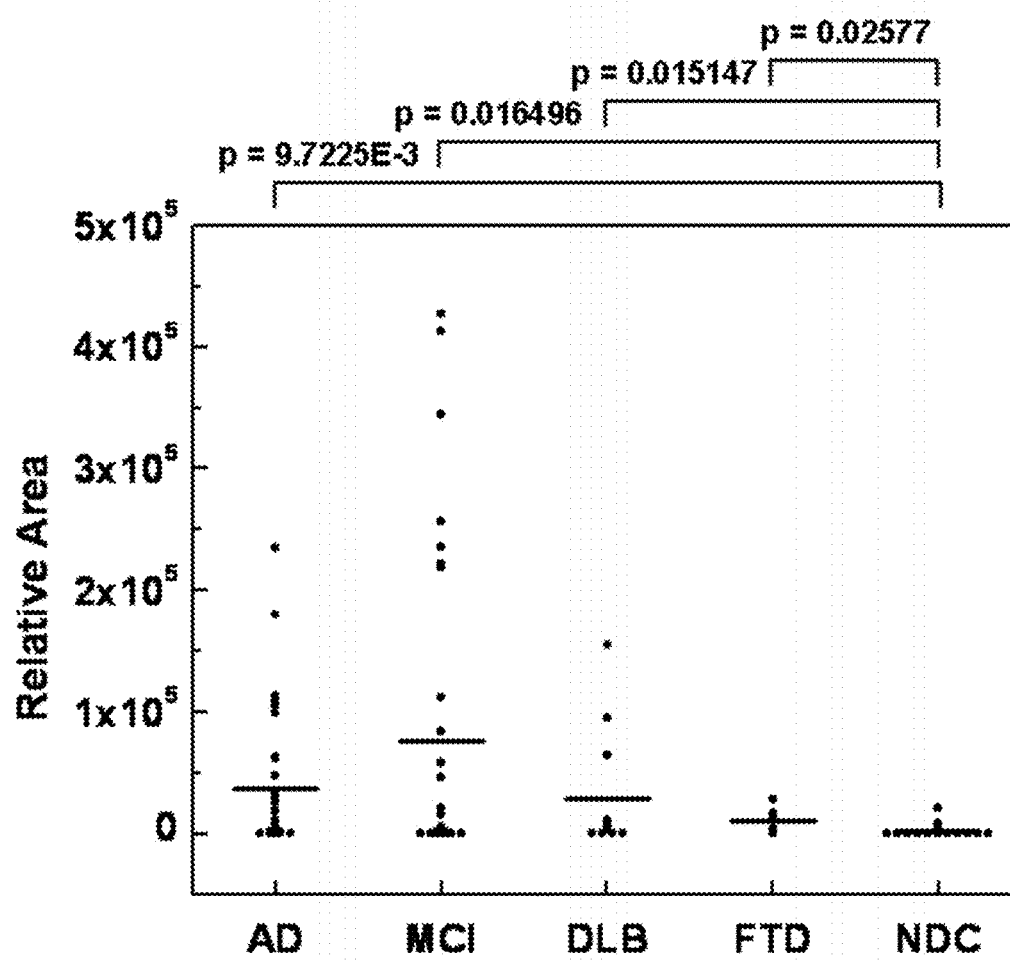


FIG.11

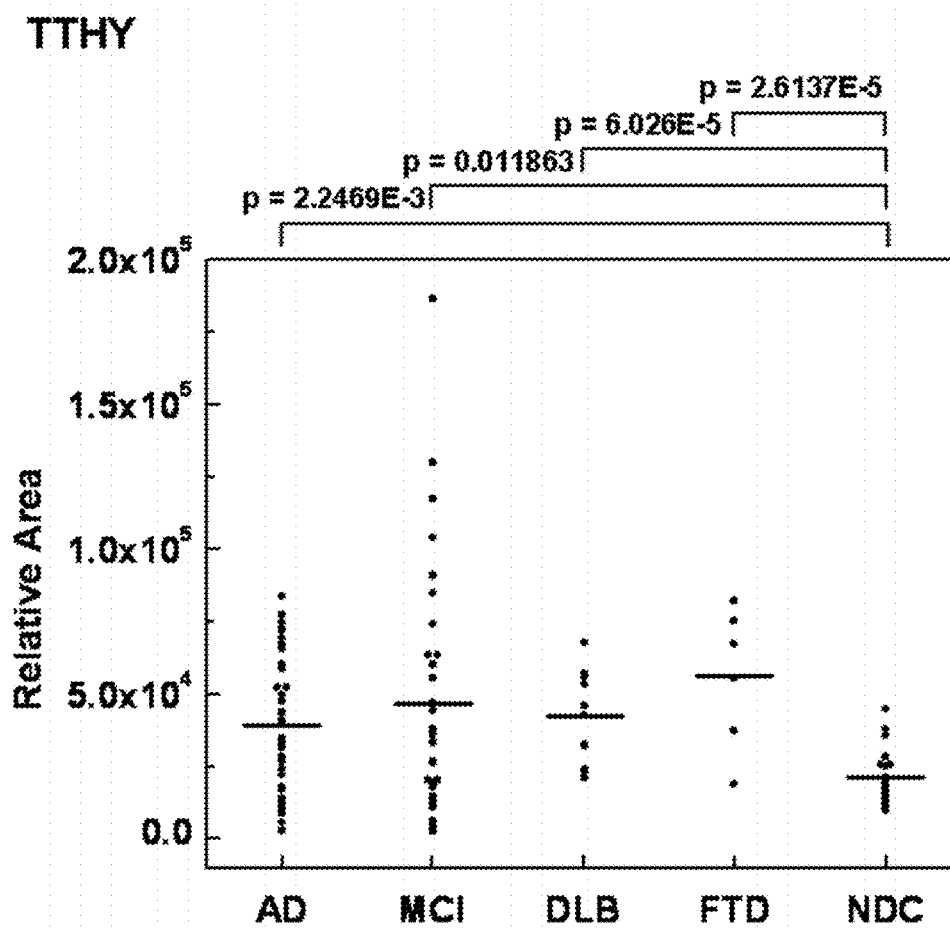


FIG.12

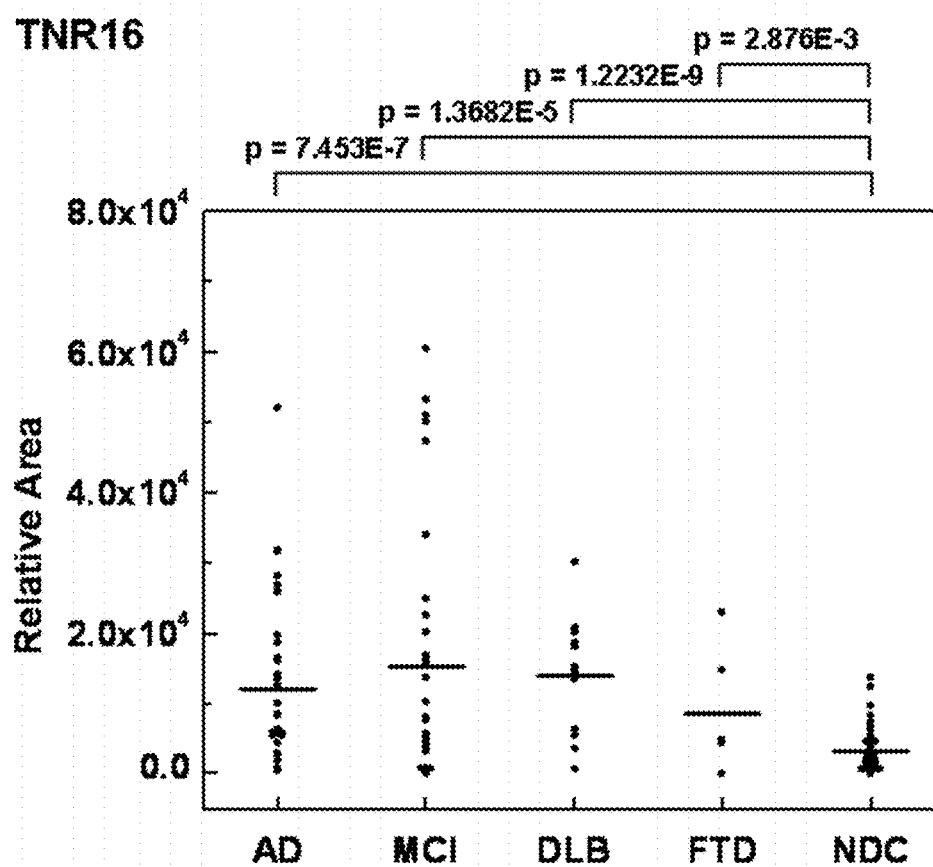


FIG.13

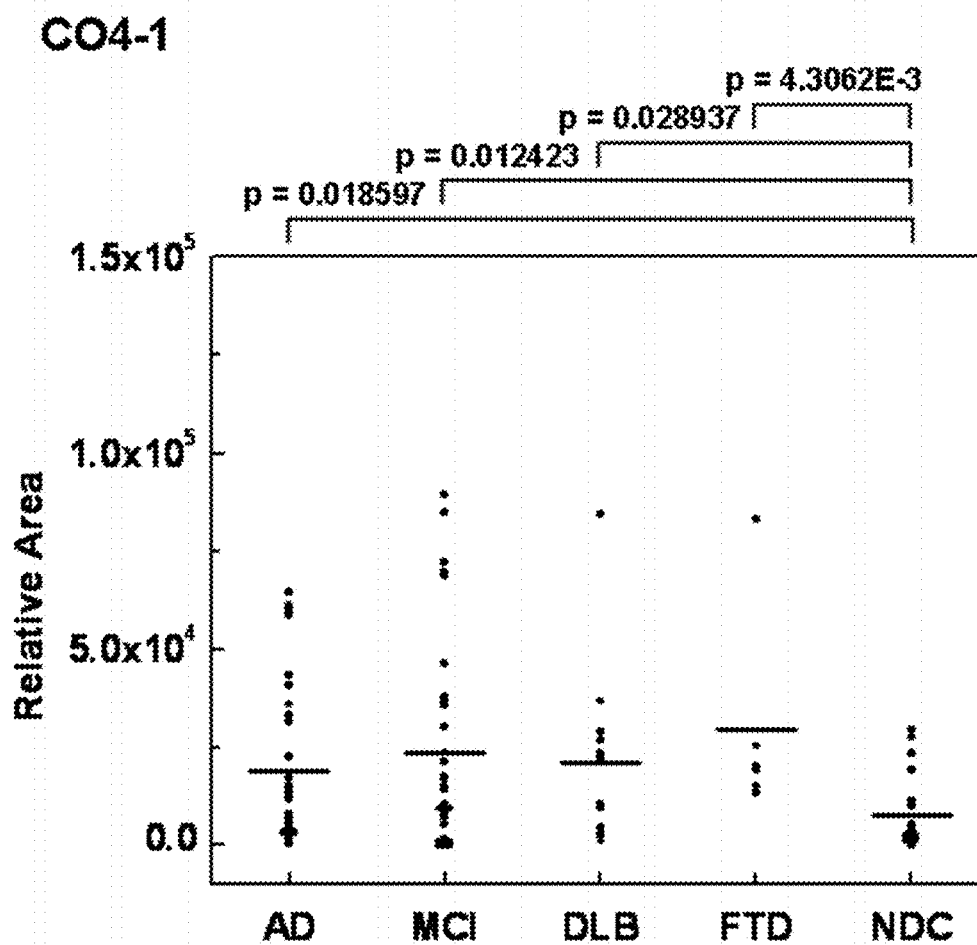


FIG.14

CO4-2

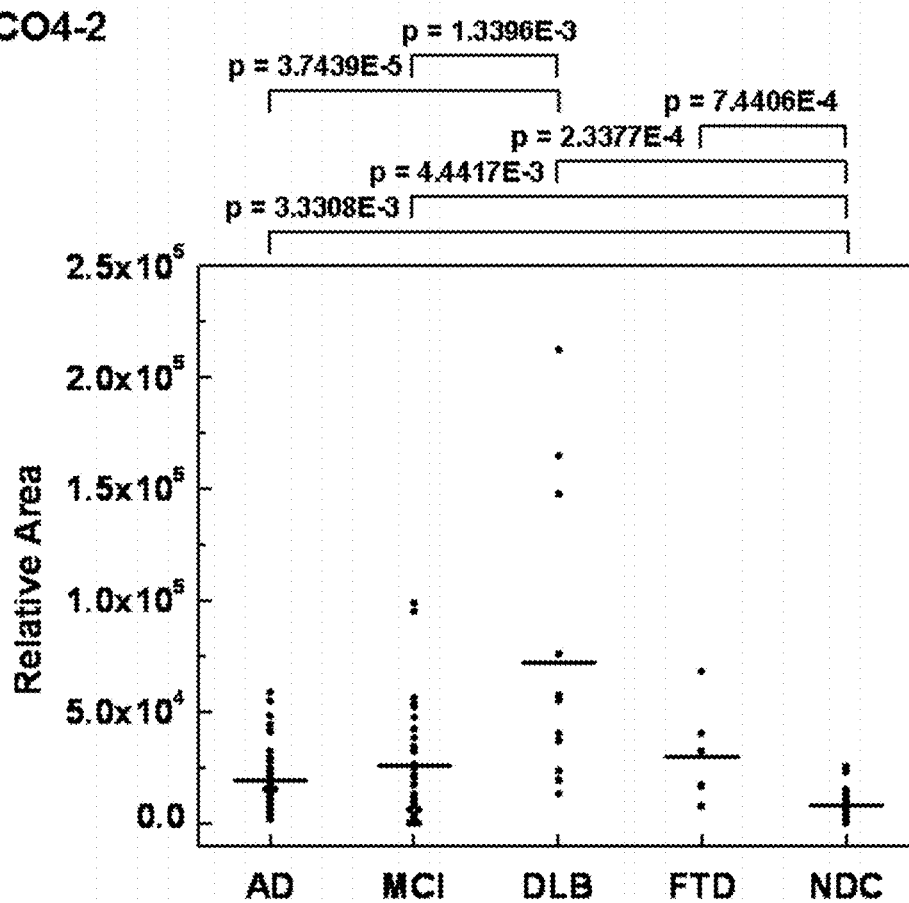


FIG.15

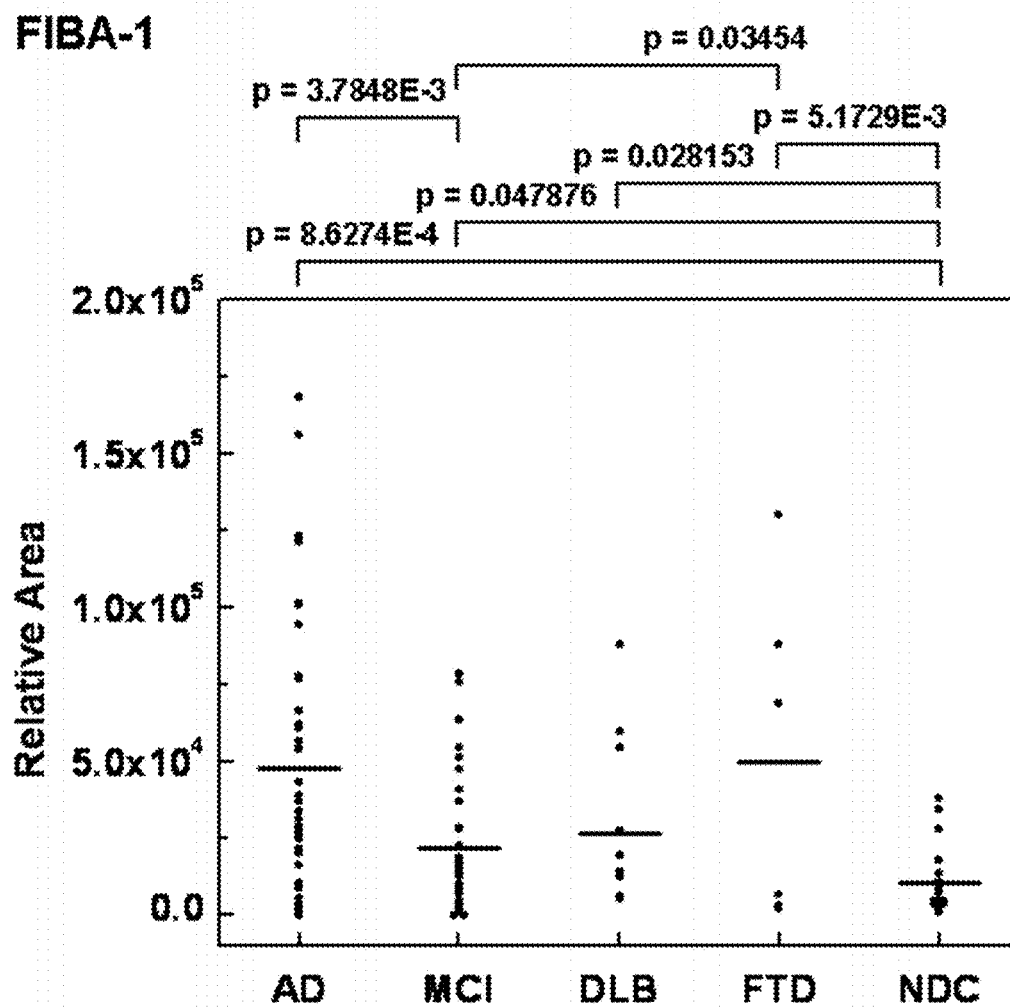


FIG.16

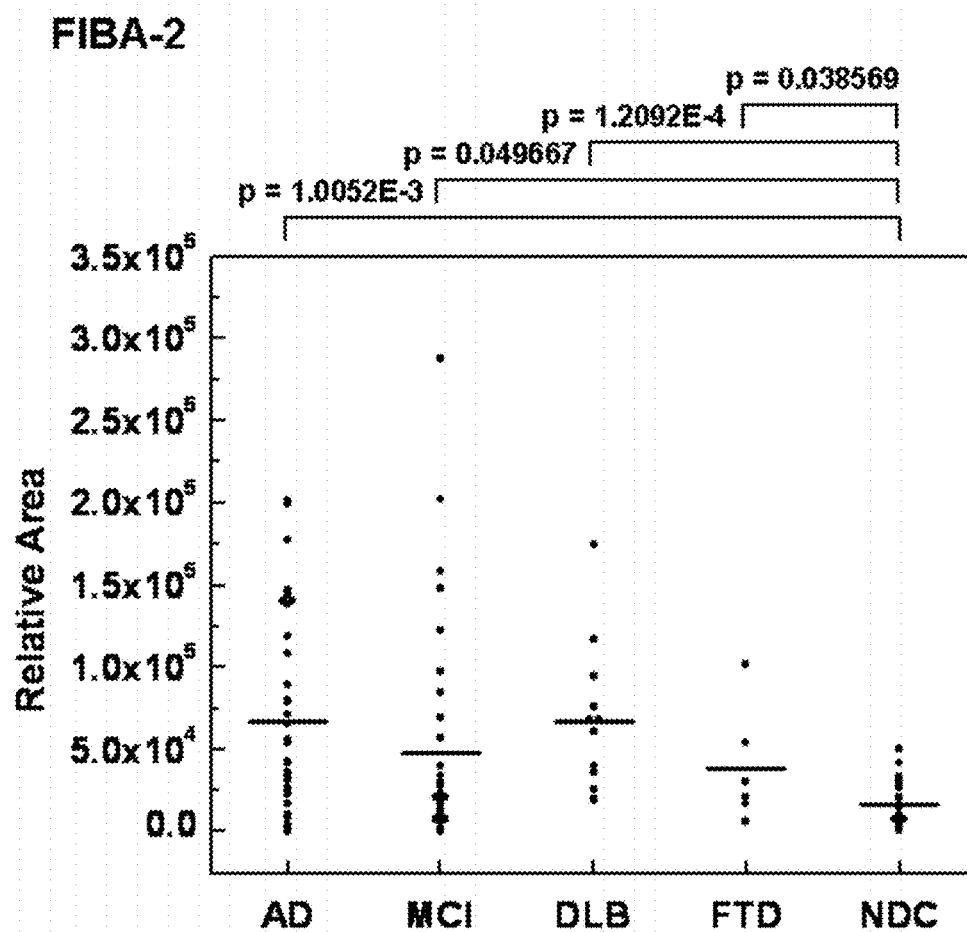
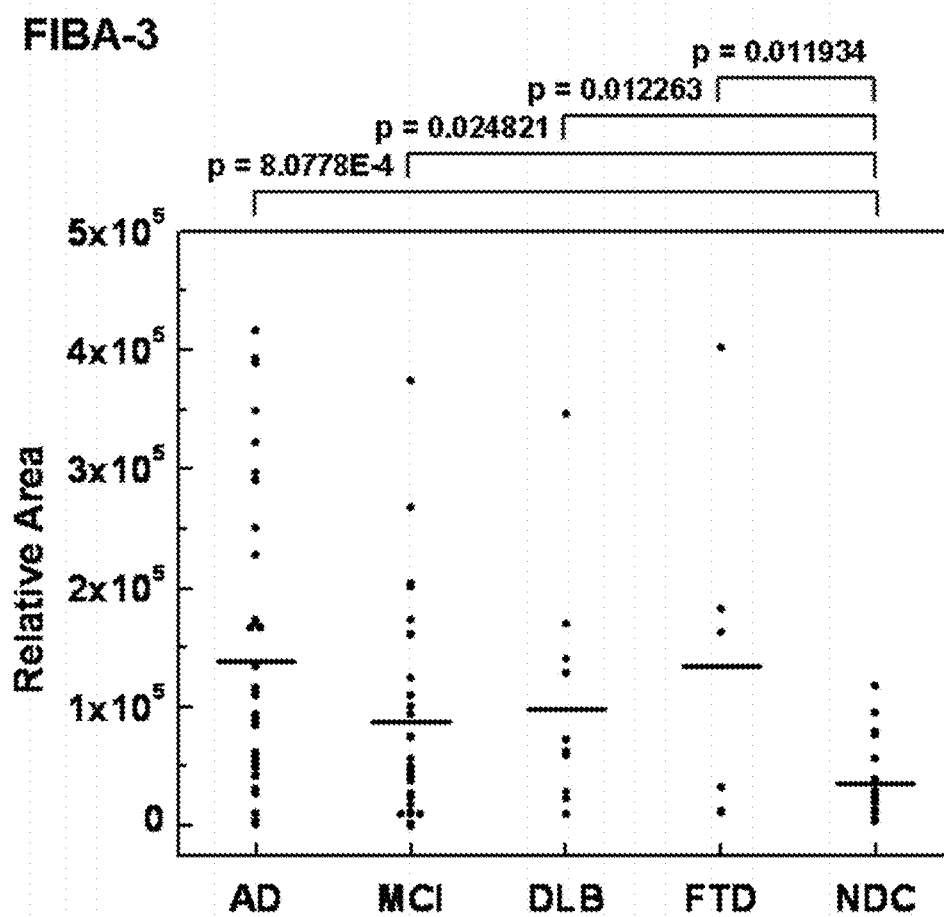


FIG.17



BIOMARKER FOR COGNITIVE DYSFUNCTION DISEASES, AND METHOD FOR DETECTION OF COGNITIVE DYSFUNCTION DISEASES USING THE BIOMARKER

CROSS-REFERENCE TO RELATED APPLICATIONS

[0001] This application is a Continuation of copending application Ser. No. 15/467,646, filed on Mar. 23, 2017, which is a Continuation of Ser. No. 14/582,778, filed on Dec. 24, 2014, which is a Continuation of application Ser. No. 13/995,682, filed on Sep. 3, 2013 (now abandoned), which was filed as PCT International Application No. PCT/JP2011/007150 on Dec. 21, 2011, which claims the benefit under 35 U.S.C. §119(a) to Patent Application No. 2010-285726, filed in JAPAN on Dec. 22, 2010, all of which are hereby expressly incorporated by reference into the present application.

FIELD OF THE INVENTION

[0002] The present invention relates to novel biomarkers for mild cognitive impairment or cognitive impairment including Alzheimer disease, and methods for detecting cognitive impairment using such biomarkers.

BACKGROUND OF THE INVENTION

[0003] The commonly used means to differentiate between normal and non-normal states of a human subject using his or her biological materials are mainly those which have been used in the field of diagnostics. Most frequently used are those methods which target biomarkers in blood. It has been practiced in this field to comparatively measure the amount of a specific protein or a peptide that is less than 10,000 in molecular weight or, in the case of enzyme protein, enzyme activities in samples from normal (healthy) subjects and those from diseased individuals to help diagnosis. Here, prior to testing real samples, measurements are done on a fixed number each of samples from healthy controls and patients with certain disease with respect to the amount (s) or activity (activities) of single or multiple specific proteins or peptides and the ranges of abnormal and normal values are respectively determined. The sample to be evaluated is then analyzed by the same method and the resultant value is judged with respect to whether it is in normal or abnormal range.

[0004] In the actual measurements, the amount(s) of specified protein(s) or peptide(s) in test samples, as such or after dilution, are determined by the use of enzyme-linked immunosorbent assay (ELISA) which uses a primary, or secondary, antibody labeled with an enzyme reacting with a substrate that yields a color upon reaction, chemiluminescent immunoassay (CLIA), radioimmunoassay (RIA) which uses a primary, or secondary, antibody labeled with a radioisotope, and, if the protein is an enzyme, the measurement of the activity of the enzyme by adding its substrate and determining the intensity of produced color, etc. These antibody-based methods are called as enzyme-, fluorescence- or radioisotope-labeled methods, respectively. In addition, there is a method where an enzyme reaction product derived from the corresponding substrate is determined by high performance liquid chromatography (HPLC). In further addition, there is a method where HPLC is

combined with mass spectrometer, called LC-MS/MS, and there is a method called selected reaction monitoring (SRM)/multiple reaction monitoring (MRM) that utilizes LC-MS/MS. In another method to determine the concentration in a sample, it is appropriately pretreated, and separation of proteins or peptides is attained by 2-dimensional polyacrylamide gel electrophoresis (2D-PAGE), and target protein or peptide is determined by silver staining, Coomassie blue staining or immunological staining (Western blotting) that uses an antibody to target protein or peptide. In still further addition, there is a method which utilizes mass spectrometry to determine the amount of target protein or peptide in samples fractionated by column chromatography. Instead of column chromatography, protein chips and magnetic beads may also be utilized for purpose of pretreatment.

[0005] Furthermore, these inventors have developed an immunoMS method, where target protein or peptide is captured by beads (including magnetic ones) with linked antibody to the protein or peptide, eluted from the beads, and determined by mass spectrometry. Further, intact proteins have been reported to be analyzed by mass spectrometry using above-mentioned methods after digestion with trypsin etc. (Patent Document 1). Here, intact target proteins are selected either by fractionation or by adsorption to an adsorbant specific to them and then determined by mass spectrometry.

[0006] Number of patients suffered from cognitive impairment like Alzheimer disease is increasing rapidly along with increasing of old-age population in Japan. It is estimated that number of patients is 1.3 million in 1995 and it will be 1.9 million in 2005 and will reach to about 3.0 million in 2020. It is reported that 60-90% of cognitive impairment is Alzheimer disease. As manifestation of Alzheimer disease is not only loss of memory but several disturbance in daily life, increase of patients of this disease is becoming an important social issue to be solved. In Japan, Donepezil-hydrochloride, anti-acetylcholine esterase inhibitor has been available for medical treatment for Alzheimer disease since 1999, and it let progress of cognitive impairment in these patients be 'slow-down' efficiently, if the patient is diagnosed at early stage. Thus, in medication of Alzheimer disease, most important issue is 'early diagnosis' to treat the patients effectively by drug available at present and new coming drug.

[0007] Followings are major criteria for diagnosis of Alzheimer disease described in DSM-IV, which is published by American Psychiatric Association.

[0008] A. The development of multiple cognitive deficits manifested by both

[0009] (1) memory impairment (impaired ability to learn new information or to recall previously learned information)

[0010] (2) one (or more) of the following cognitive disturbances:

[0011] a) aphasia (language disturbance)

[0012] b) apraxia (impaired ability to carry out motor activities despite intact motor function)

[0013] c) agnosia (failure to recognize or identify objects despite intact sensory function)

[0014] d) disturbance in executive functioning (i.e., planning, organizing, sequencing, abstracting)

[0015] B. The cognitive deficits in Criteria A1 and A2 each cause significant impairment in social or occupa-

tional functioning and represent a significant decline from a previous level of functioning. (Non-patent reference 1)

[0016] There are several types of neurological disorders related to Alzheimer disease (AD). As cognitive dysfunction appears gradually in dementia including AD, there is a disease status of pre-stage of dementia. This stage is called as mild cognitive impairment (MCI). In United States, 10% MCI develops to AD within 1 year, and 50% of MCI develops to AD within 4 years. MCI is defined as a condition characterized by newly acquired cognitive decline to an extent that is beyond that expected for age or educational background, yet not causing significant functional impairment, and not showing disturbance in daily life. Frontotemporal dementia (frontotemporal lobar degeneration) (FTD) shows loss of personal awareness, loss of social awareness, hyperorality, and stereotyped, perseverative behavior. These clinical characteristics are different from AD. FTD includes Pick's disease, which is characterized by microscopically Pick bodies usually found in limbic, paralimbic, and ventral temporal lobe cortex. Dementia with Lewy bodies (DLB) is characterized by progressive disease and psychiatric symptoms include anxiety, depression, hallucinations (usually visual) and delusions (false beliefs). DLB is thought to be the second most common subtype and 10-30% of dementia is DLB. The symptoms of DLB are caused by the build-up of Lewy bodies. FTD and DLB belong to demented neurological disease as they also lose of memory, their ability to solve problems and maintain emotional control. (Non-patent reference 1)

[0017] In description in present patent, cognitive impairment includes AD, MCI and the demented neurological disease.

[0018] The screening tests for dementia widely used are the Hasegawa Dementia Scale-revised (HDS-R) and Mini-Mental State Examination (MMSE). In these screening tests, inspector asks several questions and evaluates level of cognitive impairment of each subject by scores. HDS-R is revised version of HDS published in 1991. In HDS-R, test consists of 9 questions to analyses orientation, remembrance, calculation, retain and recall ability, and common sense. Full score is 30 and a person whose score is less than 23 is suspected as dementia. MMSE has been developed in United States to screen and diagnose dementia, and analyses global cognitive function, with items assessing orientation, word recall, attention and calculation, language abilities, and visuospatial (drawing) ability. This test consists of 11 questions, and full score is 30 and a person who has score less than 23 is suspected as dementia. The results of HDS-R and MMSE coincide with each other. Both are used for screening, not for diagnosis and not for staging of disease progression. (Non-patent reference 1)

[0019] Neuroimaging test for dementia are Computed tomography (CT) and Magnetic resonance imaging (MRI) which evaluate morphological changes like brain atrophy and ventricular dilation and single-photon emission computed tomography (SPECT) which analyses regional cerebral blood flow and PET which shows brain metabolism by measurement of consumption of oxygen and sugar. SPECT and PET, nuclear imaging technologies, can identify neuronal dysfunction at preclinical stage. However, these neuroimaging cannot be widely used in hospitals because they need special facilities to perform nuclear imaging, and

neuroimaging may not be objective test as imaging diagnosis is completely depend on the skill of physician who analyses the mages.

[0020] Thus, methods for screening and diagnosis of dementia including AD that are available at present is dependent on tests lacking objectivity and is dependent on expensive instruments, and so it is very difficult to use these tests for screening of early stage-cognitive impairment. If we get blood (serum/plasma) biomarker for cognitive impairment, which enables us objective test using specimens we can easily obtain, we can identify cognitive impairment at early stage (preclinical stage) by blood test using such biomarker. Present patent provides novel biomarkers and a novel and potent diagnostic method for cognitive impairment by using such biomarkers and biomarkers described here.

CITATION LIST

Patent Document

[0021] Patent Document 1, JP-A-2004-333274

[0022] Patent Document 2, JP-A-2006-308533

Non-Patent Document

[0023] Non-Patent Document 1, "The better understanding of Alzheimer's disease.," edited by Imaharu Nakano and Hldehiro Mizusawa., Nagai Shoten Co., Ltd., 2004 (in Japanese)

[0024] Non-Patent Document 2, Benkirane, N. et al., J. Biol. Chem. Vol. 268, 26279-26285, 1993

SUMMARY OF THE INVENTION

Technical Problem

[0025] The present invention aims to present methods to detect mild cognitive impairment or cognitive impairment including Alzheimer disease by using a protein or its partial peptide that differs in presence or absence, or in quantity between non-cognitive impairment subjects (including healthy people, the human subjects that may be affected with any disease and unaffected with psychiatry disease including cognitive impairment. These human subjects are allowed to match the age and gender of patient with cognitive impairment. And, these human subjects are called non-demented control, hereinafter abbreviated to NDC.) and patients with cognitive impairment and further aims to present biomarkers comprising said proteins and said partial peptides to be used to detect mild cognitive impairment or cognitive impairment including Alzheimer disease.

Solution to Problem

[0026] These inventors investigated to find out means to detect cognitive impairment and found a peptide capable of detecting mild cognitive impairment or cognitive impairment including Alzheimer disease in the serum. Said peptides found in the present invention are those with significance as a biomarker to detecting in the case of serum not only other biological materials such as blood, plasma, cerebrospinal fluid, and urine. Simultaneously, protein or peptide is the origin of these peptides (hereinafter referred to as intact proteins or peptides) also has significance as biomarkers.

[0027] Specifically, these inventors found that a biomarker comprising at least one protein or peptide selected from the group consisting of Complement C3 consisting of amino acid sequence expressed by SEQ ID NO: 1, Transcription factor AP-2 gamma consisting of amino acid sequence expressed by SEQ ID NO: 3, Synapsin-3 consisting of amino acid sequence expressed by SEQ ID NO: 5, Oxytocin receptor consisting of amino acid sequence expressed by SEQ ID NO: 7, Inter-alpha-trypsin inhibitor heavy chain H5-like protein consisting of amino acid sequence expressed by SEQ ID NO: 9, E3 ubiquitin-protein ligase HERC2 consisting of amino acid sequence expressed by SEQ ID NO: 11, Prothrombin consisting of amino acid sequence expressed by SEQ ID NO: 13, Transthyretin consisting of amino acid sequence expressed by SEQ ID NO: 15, Tumor necrosis factor receptor superfamily member 16 consisting of amino acid sequence expressed by SEQ ID NO: 17, Complement C4-A consisting of amino acid sequence expressed by SEQ ID NO: 19, Complement C4-B consisting of amino acid sequence expressed by SEQ ID NO: 21, Fibrinogen alpha chain (isoform 1) consisting of amino acid sequence expressed by SEQ ID NO: 23, and Fibrinogen alpha chain (isoform 2) consisting of amino acid sequence expressed by SEQ ID NO: 25; or a biomarker comprising protein fragment or peptide of not less than 5 amino acid residues arising from at least one protein or peptide selected from the group consisting of them, could be used as biomarkers to detect cognitive impairment.

[0028] Furthermore, these inventors found that a biomarker comprising from the group consisting of Complement C3-derived peptide CO3 consisting of amino acid sequence expressed by SEQ ID NO: 2, Transcription factor AP-2 gamma-derived peptide AP2C consisting of amino acid sequence expressed by SEQ ID NO: 4, Synapsin-3-derived peptide SYN3 consisting of amino acid sequence expressed by SEQ ID NO: 6, Oxytocin receptor-derived peptide OXYR consisting of amino acid sequence expressed by SEQ ID NO: 8, Inter-alpha-trypsin inhibitor heavy chain H5-like protein-derived peptide ITH5L consisting of amino acid sequence expressed by SEQ ID NO: 10, E3 ubiquitin-protein ligase HERC2-derived peptide HERC2 consisting of amino acid sequence expressed by SEQ ID NO: 12, Prothrombin-derived peptide THRB consisting of amino acid sequence expressed by SEQ ID NO: 14, Transthyretin-derived peptide TTHY consisting of amino acid sequence expressed by SEQ ID NO: 16, Tumor necrosis factor receptor superfamily member 16-derived peptide TNFR16 consisting of amino acid sequence expressed by SEQ ID NO: 18, Complement C4-derived peptide C04-1 consisting of amino acid sequence expressed by SEQ ID NO: 20, Complement C4-derived peptide C04-2 consisting of amino acid sequence expressed by SEQ ID NO: 22, Fibrinogen alpha chain-derived peptide FIBA-1 consisting of amino acid sequence expressed by SEQ ID NO: 24, Fibrinogen alpha chain-derived peptide FIBA-2 consisting of amino acid sequence expressed by SEQ ID NO: 26, and Fibrinogen alpha chain-derived peptide FIBA-3 consisting of amino acid sequence expressed by SEQ ID NO: 27 could be used as biomarkers to detect cognitive impairment.

[0029] These inventors brought the present invention to perfection by further succeeding in determining simultaneously these many proteins and its partial peptides by using

two-dimensional high performance liquid chromatography-MALDI TOF-MS method (mass spectrometry) and immunoMS method.

[0030] The features of the present invention are shown below.

[0031] [1] A biomarker for detection of cognitive impairment comprising protein fragment or peptide of not less than 5 amino acid residues arising from at least one protein or peptide selected from the group consisting of Complement C3 consisting of amino acid sequence expressed by SEQ ID NO: 1, Transcription factor AP-2 gamma consisting of amino acid sequence expressed by SEQ ID NO: 3, Synapsin-3 consisting of amino acid sequence expressed by SEQ ID NO: 5, Oxytocin receptor consisting of amino acid sequence expressed by SEQ ID NO: 7, Inter-alpha-trypsin inhibitor heavy chain H5-like protein consisting of amino acid sequence expressed by SEQ ID NO: 9, E3 ubiquitin-protein ligase HERC2 consisting of amino acid sequence expressed by SEQ ID NO: 11, Prothrombin consisting of amino acid sequence expressed by SEQ ID NO: 13, Transthyretin consisting of amino acid sequence expressed by SEQ ID NO: 15, Tumor necrosis factor receptor superfamily member 16 consisting of amino acid sequence expressed by SEQ ID NO: 17, Complement C4-A consisting of amino acid sequence expressed by SEQ ID NO: 19, Complement C4-B consisting of amino acid sequence expressed by SEQ ID NO: 21, Fibrinogen alpha chain (isoform 1) consisting of amino acid sequence expressed by SEQ ID NO: 23, and Fibrinogen alpha chain (isoform 2) consisting of amino acid sequence expressed by SEQ ID NO: 25, or a biomarker for detection of cognitive impairment comprising at least one protein or peptide selected from the group consisting of them.

[0032] [2] A biomarker for detection of cognitive impairment comprising the peptide selected from the group consisting of Complement C3-derived peptide CO3 consisting of amino acid sequence expressed by SEQ ID NO: 2, Transcription factor AP-2 gamma-derived peptide AP2C consisting of amino acid sequence expressed by SEQ ID NO: 4, Synapsin-3-derived peptide SYN3 consisting of amino acid sequence expressed by SEQ ID NO: 6, Oxytocin receptor-derived peptide OXYR consisting of amino acid sequence expressed by SEQ ID NO: 8, Inter-alpha-trypsin inhibitor heavy chain H5-like protein-derived peptide ITH5L consisting of amino acid sequence expressed by SEQ ID NO: 10, E3 ubiquitin-protein ligase HERC2-derived peptide I-HERC2 consisting of amino acid sequence expressed by SEQ ID NO: 12, Prothrombin-derived peptide THRB consisting of amino acid sequence expressed by SEQ ID NO: 14, Transthyretin-derived peptide TTHY consisting of amino acid sequence expressed by SEQ ID NO: 16, Tumor necrosis factor receptor superfamily member 16-derived peptide TNFR16 consisting of amino acid sequence expressed by SEQ ID NO: 18, Complement C4-derived peptide C04-1 consisting of amino acid sequence expressed by SEQ ID NO: 20, Complement C4-derived peptide C04-2 consisting of amino acid sequence expressed by SEQ ID NO: 22, Fibrinogen alpha chain-derived peptide FIBA-1 consisting of amino acid sequence expressed by SEQ ID NO: 24, Fibrinogen alpha chain-derived peptide FIBA-2 consisting of amino acid sequence expressed by SEQ ID NO: 26, and Fibrinogen alpha chain-derived peptide FIBA-3 consisting of amino acid sequence expressed by SEQ ID NO: 27, or a biomarker for detection of cognitive impairment.

ment comprising at least one protein or peptide selected from the group consisting of them.

[0033] [3] A biomarker of cognitive impairment comprising the peptides selected from the group consisting of amino acid sequence expressed by SEQ ID NOS: 2, 4, 6, 8, 10, 12, 14, 16, 18, 20, 22, 24, 26, and 27 that is appeared or increased in biological material of patients of cognitive impairment as compared to biological material of subjects not suffering from psychiatry disease.

[0034] [4] A biomarker of Alzheimer disease comprising the peptides selected from the group consisting of amino acid sequence expressed by SEQ ID NOS: 2, 4, 6, 8, 10, 12, 14, 16, 18, 20, 22, 24, 26, and 27 that is appeared or increased in biological material of patients of Alzheimer disease as compared to biological material of subjects not suffering from psychiatry disease.

[0035] [5] A biomarker of mild cognitive impairment comprising the peptides selected from the group consisting of amino acid sequence expressed by SEQ ID NOS: 2, 4, 6, 8, 10, 12, 14, 16, 18, 20, 22, 24, 26, and 27 that is appeared or increased in biological material of patients of mild cognitive impairment as compared to biological material of subjects not suffering from psychiatry disease.

[0036] [6] Method for detection of cognitive impairment involving determination in biological material of at least one biomarker for cognitive impairment described in any of [1] to [5].

[0037] [7] Method for detection of psychiatry disease described in [6] wherein detection is made either by immunoblot procedure, Western blotting, enzyme-, fluorescence-, or radioisotope-labeled antibody method, mass spectrometry, immunoMS method or surface plasmon resonance method.

[0038] [8] A kit for detection of cognitive impairment to determine at least one biomarker described in any of [1] to [5].

[0039] [9] A kit for detection of psychiatry disease containing antibody or aptamer to at least one biomarker described in any of [1] to [5].

Advantageous Effect of the Invention

[0040] According to the present invention, it is possible to diagnose the subject such as suffering from mild cognitive impairment or cognitive impairment including Alzheimer's disease, when to increase or appear compared to the biological sample of subjects not suffering from psychiatry disease by determining amount of at least one biomarker comprising protein fragment or peptide of not less than 5 amino acid residues arising from at least one protein or peptide selected from the group consisting of Complement C3 consisting of amino acid sequence expressed by SEQ ID NO: 1, Transcription factor AP-2 gamma consisting of amino acid sequence expressed by SEQ ID NO: 3, Synapsin-3 consisting of amino acid sequence expressed by SEQ ID NO: 5, Oxytocin receptor consisting of amino acid sequence expressed by SEQ ID NO: 7, Inter-alpha-trypsin inhibitor heavy chain H5-like protein consisting of amino acid sequence expressed by SEQ ID NO: 9, E3 ubiquitin-protein ligase HERC2 consisting of amino acid sequence expressed by SEQ ID NO: 11, Prothrombin consisting of amino acid sequence expressed by SEQ ID NO: 13, Transthyretin consisting of amino acid sequence expressed by SEQ ID NO: 15, Tumor necrosis factor receptor superfamily member 16 consisting of amino acid sequence expressed by

SEQ ID NO: 17, Complement C4-A consisting of amino acid sequence expressed by SEQ ID NO: 19, Complement C4-B consisting of amino acid sequence expressed by SEQ ID NO: 21, Fibrinogen alpha chain (isoform 1) consisting of amino acid sequence expressed by SEQ ID NO: 23, and Fibrinogen alpha chain (isoform 2) consisting of amino acid sequence expressed by SEQ ID NO: 25.

[0041] In addition, according to the present invention, it is possible to diagnose the subject such as suffering from mild cognitive impairment or cognitive impairment including Alzheimer's disease, when to increase or appear compared to the biological sample of subjects not suffering from psychiatry disease by determining kind or amount at least one peptide selected from the group consisting of Complement C3-derived peptide CO3 consisting of amino acid sequence expressed by SEQ ID NO: 2, Transcription factor AP-2 gamma-derived peptide AP2C consisting of amino acid sequence expressed by SEQ ID NO: 4, Synapsin-3-derived peptide SYN3 consisting of amino acid sequence expressed by SEQ ID NO: 6, Oxytocin receptor-derived peptide OXYR consisting of amino acid sequence expressed by SEQ ID NO: 8, Inter-alpha-trypsin inhibitor heavy chain H5-like protein-derived peptide ITH5L consisting of amino acid sequence expressed by SEQ ID NO: 10, E3 ubiquitin-protein ligase HERC2-derived peptide HERC2 consisting of amino acid sequence expressed by SEQ ID NO: 12, Prothrombin-derived peptide THRB consisting of amino acid sequence expressed by SEQ ID NO: 14, Transthyretin-derived peptide TTHY consisting of amino acid sequence expressed by SEQ ID NO: 16, Tumor necrosis factor receptor superfamily member 16-derived peptide TNR16 consisting of amino acid sequence expressed by SEQ ID NO: 18, Complement C4-derived peptide C04-1 consisting of amino acid sequence expressed by SEQ ID NO: 20, Complement C4-derived peptide C04-2 consisting of amino acid sequence expressed by SEQ ID NO: 22, Fibrinogen alpha chain-derived peptide FIBA-1 consisting of amino acid sequence expressed by SEQ ID NO: 24, Fibrinogen alpha chain-derived peptide FIBA-2 consisting of amino acid sequence expressed by SEQ ID NO: 26, and Fibrinogen alpha chain-derived peptide FIBA-3 consisting of amino acid sequence expressed by SEQ ID NO: 27.

[0042] The present invention provides a diagnostic system that is high in both accuracy and specificity. The present invention enables highly accurate diagnosis of cognitive impairment in which there have been no specific test methods for such biological materials as blood. Furthermore, the biomarkers disclosed in the present invention are highly useful in judgment of drug efficacy.

BRIEF DESCRIPTION OF DRAWINGS

[0043] FIG. 1 illustrates the cluster map of Marker A. The dots within the rectangle indicated by (A) are m/z and retention time of the mass peak of Marker A detected from the serum of the individual subject using reverse phase chromatography. The dots in a cluster can be regarded as the same retention time and the same m/z in the error range, and the dots in a cluster are defined to be derived from the same peptide.

[0044] FIG. 2 illustrates the results of differential analysis in the case of Marker A. As shown in the amino acid sequences resulting of MS/MS analysis in FIG. 4, Marker A is Complement C3-derived peptides CO3. FIG. 2 shows the

comparison between NDC and cognitive impairment (AD, MCI, DLB and FTD) related to CO3.

[0045] FIG. 3 illustrates the ROC curves of CO3 expressed by SEQ ID NO: 2. Definition of the ROC curve, see the section on the results of Example. FIG. 3A) shows the ROC curve of the comparison of AD vs. NDC. FIG. 3B) shows the ROC curve of the comparison of MCI vs. NDC.

[0046] FIG. 4 illustrates the MS/MS spectrum of CO3 by TOF/TOF mass spectrometer. In FIG. 4 top, it was shown the amino acid sequence of CO3, and it was shown y-ions and b-ions that appear in the MS/MS spectrum.

[0047] FIG. 5 illustrates the results of differential analysis of AP2C expressed by SEQ ID NO: 4. This figure shows a comparison between cognitive impairment patients (AD, MCI, DLB, FTD) and subjects not suffering from psychiatry disease (NDC).

[0048] FIG. 6 illustrates the results of differential analysis of SYN3 expressed by SEQ ID NO: 6. This figure shows a comparison between cognitive impairment patients (AD, MCI, DLB, FTD) and subjects not suffering from psychiatry disease (NDC).

[0049] FIG. 7 illustrates the results of differential analysis of OXYR expressed by SEQ ID NO: 8. This figure shows a comparison between cognitive impairment patients (AD, MCI, DLB, FTD) and subjects not suffering from psychiatry disease (NDC).

[0050] FIG. 8 illustrates the results of differential analysis of ITH5L expressed by SEQ ID NO: 10. This figure shows a comparison between cognitive impairment patients (AD, MCI, DLB, FTD) and subjects not suffering from psychiatry disease (NDC).

[0051] FIG. 9 illustrates the results of differential analysis of HERC2 expressed by SEQ ID NO: 12. This figure shows a comparison between cognitive impairment patients (AD, MCI, DLB, FTD) and subjects not suffering from psychiatry disease (NDC).

[0052] FIG. 10 illustrates the results of differential analysis of THRB expressed by SEQ ID NO: 14. This figure shows a comparison between cognitive impairment patients (AD, MCI, DLB, FTD) and subjects not suffering from psychiatry disease (NDC).

[0053] FIG. 11 illustrates the results of differential analysis of TTHY expressed by SEQ ID NO: 16. This figure shows a comparison between cognitive impairment patients (AD, MCI, DLB, FTD) and subjects not suffering from psychiatry disease (NDC).

[0054] FIG. 12 illustrates the results of differential analysis of TNFR16 expressed by SEQ ID NO: 18. This figure shows a comparison between cognitive impairment patients (AD, MCI, DLB, FTD) and subjects not suffering from psychiatry disease (NDC).

[0055] FIG. 13 illustrates the results of differential analysis of C04-1 expressed by SEQ ID NO: 20. This figure shows a comparison between cognitive impairment patients (AD, MCI, DLB, FTD) and subjects not suffering from psychiatry disease (NDC).

[0056] FIG. 14 illustrates the results of differential analysis of C04-2 expressed by SEQ ID NO: 22. This figure shows a comparison between cognitive impairment patients (AD, MCI, DLB, FTD) and subjects not suffering from psychiatry disease (NDC).

[0057] FIG. 15 illustrates the results of differential analysis of FIBA-1 expressed by SEQ ID NO: 24. This figure

shows a comparison between cognitive impairment patients (AD, MCI, DLB, FTD) and subjects not suffering from psychiatry disease (NDC).

[0058] FIG. 16 illustrates the results of differential analysis of FIBA-2 expressed by SEQ ID NO: 26. This figure shows a comparison between cognitive impairment patients (AD, MCI, DLB, FTD) and subjects not suffering from psychiatry disease (NDC).

[0059] FIG. 17 illustrates the results of differential analysis of FIBA-3 expressed by SEQ ID NO: 27. This figure shows a comparison between cognitive impairment patients (AD, MCI, DLB, FTD) and subjects not suffering from psychiatry disease (NDC).

DESCRIPTION OF EMBODIMENTS

[0060] The present invention is a method for determining the kind and the amount of intact protein and/or its partial peptide when test subject is suffering from cognitive impairment as well as for diagnosing whether test subject is suffering from cognitive impairment. A peptide is generally said to be a chemical entity, made by polymerizing a number of amino acids, of less than 10,000 in molecular weight or by polymerizing several to less than about 50 amino acid residues. While in the present invention a partial peptide of an intact protein can be used as a biomarker for detection of cognitive impairment, such partial peptide is defined as a peptide of less than 10,000 in molecular weight consisting of a part of the amino acid sequence of the intact protein. Such peptide may arise as a partial peptide during the expression by transcription followed by synthesis by translation before maturing into an intact protein or as a peptide produced by enzyme digestion in the body after the intact protein has been synthesized. It is possible that, when the body is in abnormal state suffering from such disease as cognitive impairment, the mechanism for protein synthesis and regulation is de-regulated. In other words, the present invention is also a method for determining if test subject is in normal state or is suffering from cognitive impairment by using the degree of protein synthesis and/or protein digestion as an indicator. The detection of cognitive impairment in the present invention means evaluation and differentiation, i.e., diagnosis of test subject as to whether the subject is suffering from cognitive impairment. The present invention can also include the evaluation of patient's risk of suffering from more serious cognitive impairment.

[0061] Specifically, in the method of the present invention, the examples of intact protein that can be used as a cognitive impairment include Complement C3 consisting of amino acid sequence expressed by SEQ ID NO: 1, Transcription factor AP-2 gamma consisting of amino acid sequence expressed by SEQ ID NO: 3, Synapsin-3 consisting of amino acid sequence expressed by SEQ ID NO: 5, Oxytocin receptor consisting of amino acid sequence expressed by SEQ ID NO: 7, Inter-alpha-trypsin inhibitor heavy chain H5-like protein consisting of amino acid sequence expressed by SEQ ID NO: 9, E3 ubiquitin-protein ligase HERC2 consisting of amino acid sequence expressed by SEQ ID NO: 11, Prothrombin consisting of amino acid sequence expressed by SEQ ID NO: 13, Transthyretin consisting of amino acid sequence expressed by SEQ ID NO: 15, Tumor necrosis factor receptor superfamily member 16 consisting of amino acid sequence expressed by SEQ ID NO: 17, Complement C4-A consisting of amino acid sequence expressed by SEQ ID NO: 19, Complement C4-B consisting

of amino acid sequence expressed by SEQ ID NO: 21, Fibrinogen alpha chain (isoform 1) consisting of amino acid sequence expressed by SEQ ID NO: 23, and Fibrinogen alpha chain (isoform 2) consisting of amino acid sequence expressed by SEQ ID NO: 25, and further, the peptide fragments that comprise of partial peptides of not less than 5 amino acid residues of these intact proteins can be used as same purpose.

[0062] Still further, an example of biomarkers for cognitive impairment of the present invention includes the partial peptides consisting of amino acid sequence expressed by SEQ ID NO: 2 of Complement C3-derived peptide CO3, SEQ ID NO: 4 of Transcription factor AP-2 gamma-derived peptide AP2C, SEQ ID NO: 6 of Synapsin-3-derived peptide SYN3, SEQ ID NO: 8 of Oxytocin receptor-derived peptide OXYR, SEQ ID NO: 10 of Inter-alpha-trypsin inhibitor heavy chain H5-like protein-derived peptide ITH5L, SEQ ID NO: 12 of E3 ubiquitin-protein ligase HERC2-derived peptide HERC2, SEQ ID NO: 14 of Prothrombin-derived peptide THRB, SEQ ID NO: 16 of Transthyretin-derived peptide TTHY, SEQ ID NO: 18 of Tumor necrosis factor receptor superfamily member 16-derived peptide TNFR16, SEQ ID NO: 20 of Complement C4-derived peptide C04-1, SEQ ID NO: 22 of Complement C4-derived peptide C04-2, SEQ ID NO: 24 of Fibrinogen alpha chain-derived peptide FIBA-1, SEQ ID NO: 26 of Fibrinogen alpha chain-derived peptide FIBA-2, and SEQ ID NO: 27 of Fibrinogen alpha chain-derived peptide FIBA-3. In the present invention, proteins and peptides consisting of amino acid sequences derived from SEQ ID NOS: 1 through 27 by deletion, exchange, and/or addition of one or a few amino acids can be used as biomarkers and are included in the present invention. "One or a few" herein means "one or three," "one or two," or "one." Furthermore, the partial peptides that can be used as biomarkers in the present invention include those peptide fragments consisting of not less than 5 amino acid residues arising respectively from SEQ ID NOS: 1 through 27. The basis for the limitation of peptide fragments consisting of not less than 5 amino acid residues is in the description below in Non-patent Document 2. The document reported that an antibody obtained by using the peptide IRGERA as immunogen, which was the C-terminus (130-135) of histone H3, recognized the peptide IKGERA derived by exchange of K for R and the peptide CGGGERA which was derived by deletion of IR followed by addition of CGG. This demonstrates that the immunogenicity (antigenicity) is recognized by a peptide of not less than 4 amino acid residues. In order to expand this finding to other peptides than the C-terminus of histone H3, the number of amino acid residue is defined as not less than 5 instead of 4 in the present invention. To make such a low molecular weight peptide as the subject of the present invention is important when the method of detection and differentiation uses immunological means including immunoblot, ELISA and immunoMS.

[0063] It is to be noted that there are cases where a sugar chain or sugar chains have been added to an intact protein or its partial peptide to form glycosylated entities. Proteins and partial peptides in glycosylated form can also be used as biomarkers for detection of cognitive impairment.

[0064] It is also to be noted that, in the present invention, biomarker can be quantified or its presence or absence can be determined qualitatively.

[0065] Two-dimensional electrophoresis (2-DE) or 2-dimensional chromatography (2-DC) can be used in the pres-

ent invention to separate biomarkers in biological materials including serum. Known chromatographic methods can be selected from ion-exchange chromatography, reverse-phase chromatography and gel-filtration chromatography. It is also possible to make quantification with the SRM/MRM method in LC-MS/MS technology. Furthermore, the immunoMS method which these inventors have developed, where target protein or peptide is captured by beads (including magnetic ones) with antibody linked to the protein or peptide, eluted from the beads, and determined by mass spectrometry enables convenient determination of presence or absence or the amount of target protein, protein fragment or peptide without the use of 2-DE or chromatography.

[0066] It is possible with the use of the method disclosed in the present invention to evaluate at the stage of mild of cognitive dysfunction in test subject and therefore it can be useful in prophylactic medicine. Further, when psychotherapy and/or drug therapy is given to patients with cognitive impairment, it is reflected in the amount of proteins and partial peptides in biological materials such as serum if the progression of the disorder has been inhibited. Therefore, by measuring these proteins and partial peptides, it is possible to evaluate and determine therapeutic effect.

[0067] The kind and amount of a protein in biological materials can be determined by various methods. If target protein (including protein fragment and partial peptide) has been characterized and when an antibody (primary antibody) to it has already been obtained, the following methods can be used:

1. Immunoblot

[0068] This is one of the simplest methods. Test serum in a fixed amount (about 1 microliter) after stepwise dilution is dropped onto an appropriate membrane such as of nitrocellulose and dried in air. The membrane is treated with a blocking solution containing a protein such as BSA, washed, reacted with primary antibody, and washed. Thereafter, the membrane is reacted with labeled secondary antibody to detect the primary antibody. The membrane is washed and the label is visualized to measure its density.

2. Western Blotting

[0069] After separation with one-dimensional or two-dimensional electrophoresis involving isoelectric focusing or SDS-PAGE, proteins are transferred onto such an appropriate membrane as of PVDF and their amounts are determined, as in above-mentioned immunoblot, using primary antibody and labeled secondary antibody.

3. ELISA

[0070] Antibody to protein or its partial peptide is fixed to such a plate as a chemically modified microtiter plate. Appropriate amounts of samples after stepwise dilution are applied to the plate and incubated. Proteins and peptides not captured are removed by washing. Next, the plate is incubated with secondary antibody labeled with fluorescent or chemiluminescent substance or enzyme. After addition of respective substrate, fluorescence, chemiluminescence or visible light due to enzyme reaction is measured for evaluation and judgment.

[0071] Additional examples of methods are illustrated below (see Patent Document 2) but the invention is not limited by these examples.

4. Methods that Use Microarray (Microchip)

[0072] A microarray is a general term for devices where solidified materials with affinity for target substances are arrayed on solid support (plate). In the present invention, antibodies or aptamer to proteins and partial peptides are arrayed. A sample of biological material is placed on the microarray for fixation of target proteins or partial peptides and the microarray is then incubated with secondary antibody labeled with fluorescent or chemiluminescent substance or enzyme. After addition of respective substrate, fluorescence, chemiluminescence or visible light due to enzyme reaction is measured.

5. Mass Spectrometry

[0073] In mass spectrometry, for example, antibody to a specified protein or partial peptide is attached to chemically modified microbeads or plate (protein chip). The microbeads could be magnetic beads. There are no requirements for the material of the plate. The antibody to be used could be (1) an antibody which recognizes the full length form of the specified protein only, (2) an antibody which recognizes a partial peptide only, (3) all of antibodies which recognizes both the specified protein and its partial peptide, or a combination of (1) and (2), (1) and (3), or (2) and (3). Samples after stepwise dilution with original solvent or buffer are added to the microbeads or plate carrying antibody or antibodies and incubated. Those proteins and partial peptides not captured are removed by washing. The protein or partial peptide captured by microbeads or plate is eluted, and analyzed by mass spectrometry with MALDI-TOF-MS, SELDI-TOF-MS, etc. Measurements are made with respect to the mass and intensity of the peak due to the protein, protein fragment or partial peptide. Prior to the measurements a fixed amount of substance serving as the internal standard is added to the original biological material and the intensity of its peak is also measured. The concentration of the target in the original biological material can be calculated from the ratio of peak intensity of the target to the peak intensity of the internal standard. This is called immunoMS method. Further, it is possible to make quantification, after the sample is diluted with original solvent or buffer, or after part of proteins are removed, by separation with HPLC followed by mass spectrometry with electrospray ionization (ESI) method. Therein the SRM/MRM method can be utilized for absolute quantification with the use of an isotope-labeled internal standard peptide.

[0074] Furthermore, in addition to the above-mentioned methods, it is possible to analyze proteins and partial peptides by using 2-DE, surface plasmon resonance, etc.

[0075] The present invention includes the method to detect cognitive impairment from the presence or absence or amount of the above-mentioned biomarker after applying biological material obtained from test subject to 2-DE or surface plasmon resonance.

EXAMPLES

[0076] Discovery of a marker peptide for detection of cognitive impairment using two-dimensional liquid chromatography-mass spectrometry (2D-LC-MALDI TOF-MS).

(1) Serum Samples.

[0077] Followings, the characters before the parenthesis are an abbreviation.

[0078] A sera obtained from 40 AD (Alzheimer's disease), 35 MCI (mild cognitive impairment), 13 DLB (Dementia with Lewy bodies), 7 FTD (frontotemporal lobar degeneration), and 21 NDC (subjects not suffering from psychiatry disease) were used.

(2) Methods

[0079] After 475 μ l of 0.1% trifluoroacetic acid (TFA) were added in each of 25 μ l of sera, samples were boiled for 15 min at 100 degrees. Subsequently, in order to recover peptides of molecular weight of 10,000 or less, ultrafiltration were performed by using YM-10 filter unit (Millipore Corp.). Then the analysis using 2D-LC-MALDI TOF-MS were performed as follows. In other words, recovering samples were fractionated to 382 fractions per sample by using two-dimensional HPLC (SCX cation exchange column at one-dimension and C18 reverse-phase column at two-dimension). The samples were fractionated into two fractions by SCX cation exchange column, namely, SCX 1 fraction is through fraction, SCX 2 fraction is the fraction that eluted with 100% salt solution. Two fractions that were fractionated by SCX, respectively, were fractionated 191 fractions by C18 reverse phase column chromatography. It was eluted with 6 seconds in one fraction, and the retention times were calculated by multiplying the number of minus 1 from number of eluted fractions to 6 seconds. All fractionated samples were spotted on MALDI target plate (MTP AnchorChip™ 600/384 plate, BRUKER DALTONICS) for MALDI TOF/TOF mass spectrometer (ultraflex TOF/TOF, BRUKER DALTONICS) using a spotting robot (AccuSpot, SHIMADZU) that is connected online, and matrix solution (alpha-cyano-hydroxycinnamic acid, CHCA) were mixed and crystallized. After mounting MALDI target plate into ultraflex TOF/TOF, the mass and the peak area of the mass were measured automatically in reflectron mode by irradiating to crystallized sample by laser. Peak area was normalized with 250 mole of per each well of bradykinin 1-7 fragment that was added into matrix solution in advance. In other words, the area value was calculated dividing the peak area of specific mass in sample by the peak area obtained from bradykinin1-7 fragment. This area value is corresponding in 25 μ l of sample serum. Detection of difference in abundance of peptides in serum between groups (called differential analysis) was performed using multi-group statistical analysis software Parnassum™ (MCBI) developed by us. Peptide that was observed to difference in abundance was directly determined amino acid sequence in MS/MS analysis by ultraflex TOF/TOF, and intact proteins or peptides of their origin were identified.

(3) Results

[0080] The following shows the result of differential analysis by Parnassum software for data of serum individual subjects obtained using 2D-LC MALDI TOF-MS. FIG. 1 shows the result that was obtained from sample that was applied to 2D-LC-MALDI TOF-MS. Sample was fractionated into 2 fractions by SCX cation exchange column in the first dimension, then first fractions from SCX column (SCX 1) were fractionated into 191 fractions by C18 reverse-phase column. Mass spectra of 191 fractions were obtained by MALDI TOF-MS measuring. As the horizontal axis is the m/z and the vertical axis is the fractions of reverse-phase column chromatography, FIG. 1 was visualized by Parnassum.

sum software developed by present inventors. The dots in FIG. 1 shows respectively TOF-MS peak derived from the individual subject. The sections that dots are gathered can be regarded as the same retention time and the same m/z in the error range, and the dots in the sections are defined to be derived from the same peptide. These sections are referred to as clusters. Section (A) of FIG. 1 shows cluster of Marker A.

[0081] FIG. 2 shows the results of differential analysis in the case of Marker A. As shown in FIG. 4, Marker A is Complement C3-derived peptides CO3. FIG. 2 shows the comparison between subjects not suffering from psychiatry disease (NDC) and cognitive impairment (AD, MCI, DLB and FTD) related to CO3. In the results of t-test, area values of cognitive impairment (AD, MCI, DLB and FTD) were significantly higher than NDC ($p < 0.05$).

[0082] From the results of FIG. 2, in order to evaluate the extent to which the Marker A is useful as biomarker, the analysis by receiver operating characteristic (ROC) curve was performed. A) and B) in FIG. 3 shows respectively the ROC curve of the comparison of AD vs. NDC and MCI vs. NDC. If the area value (hereinafter referred to as the AUC value) of under the ROC curve is close to 1, the usefulness as biomarker of Marker A will be higher. In A) and B) of FIG. 3, the typical values of sensitivity and specificity are the values of the point (open square in the figure) of the coordinate on ROC curve that the distance is minimized when a straight line was drawn to ROC curve from the point of 100% on y-axis. The value of cut-off giving this point becomes a useful threshold to distinguish between the different groups, and the values of sensitivity and specificity at that time (i.e., above the typical values) becomes an indicator of the usefulness of biomarkers together with AUC values. In A) of FIG. 3, as typical values in AD vs. NDC, the sensitivity was 73.0%, the specificity was 100%, and the AUC value was 0.88. In B) of FIG. 3, as typical values in MCI vs. NDC, the sensitivity was 70.6%, the specificity was 89.5%, and the AUC value was 0.83.

[0083] Thus, it was revealed that Marker A was useful to distinguish AD and MCI with NDC. In particular, since MCI is the state of previous stage of AD, Marker A is considered to be an extremely useful marker to detect MCI for early diagnosis of potential subjects to migrate to AD.

[0084] FIG. 4, for Marker A, illustrates the results of MS/MS spectrum using ultraflex TOF/TOF. The signals that show y-ions and b-ions have enough appeared, and the amino acid sequence could be readily identified. Mascot search was performed on this result and the protein of origin or the peptide (hereinafter referred to as intact proteins or peptides) is Complement C3, and the detected peptide was found that the sequence is APVIHQEMIGGLRN (SEQ ID NO: 2). CO3 of entry name of Swiss-Prot against Complement C3 will use as an abbreviation of the peptide name. Followings, for peptides other than CO3, entry name will use as peptide name, similarly.

[0085] Including the Marker A, the peptides that have difference in abundance between the groups in serum were measured MS/MS spectra using ultraflex TOF/TOF, and in addition to determining the amino acid sequence, the results identified intact proteins or peptides were shown below. For peptides other than Marker A, the signals that show y-ions and b-ions has enough appeared, and the amino acid sequence could be readily identified. The following amino acid sequence that shows a set of two sequences, the first sequence shows the amino acid sequence of intact proteins, and the second sequence shows the amino acid sequence of peptide detected by 2D-LC MALDI TOF-MS. The peptide comprising of the underlined portion in the first sequence correspond to the sequence of peptide detected by 2D-LC MALDI TOF-MS. The amino acid sequence starting at 0001 in the sequence shows the sequence of the N-terminus side.

(1) Complement C3-Derived Peptide CO3

[0086] CO3 shown as SEQ ID NO: 2 had formed a cluster by clustering using Parnassus software.

[0087] As shown in FIG. 2, area values of cognitive impairment (AD, MCI, DLB and FTD) were significantly higher than NDC (t-test, $p < 0.05$). Thus, it was revealed that CO3 shown as SEQ ID NO: 2 was useful to distinguish patient of cognitive impairment (AD, MCI, DLB and FTD) with subjects not suffering from psychiatry disease (NDC). According to the analysis by receiver operating characteristic (ROC) curve, CO3 was clearly useful to distinguish AD and MCI with NDC. (See FIG. 3A), 3B) and Table 1).

Intact Protein/Peptide

[0088]

(SEQ ID NO: 1)

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0001 SPMYSIITPN ILRLESEETM VLEAHDAQGD VPVTVTVHDF PGKKLVLSSE
0051 KTVLTPATNH MGNVTFTIPA NREFKSEKGR NKFVTVQATF GTQVVEKVV
0101 VSLQSGYLF I QTDKTIYTPG STVLYRIFTV NHKLLPVGRT VMVNIENPEG
0151 IPVKQDSLSS QNQLGVLP L S WDIPELVNMG QWKIRAYYEN SPQQVFSTEF
0201 EVKEYVLPSF EVIVEPTEKF YYIYNEKGLE VTITARFLYG KKVEGTAFVI
0251 FGIQDGEQRI SLPESLKRIP IEDGSGEVVL SRKVLLDGVQ NPRAEDLVGK
0301 SLYVSATVIL HSGSDMVQAE RSGIPIVTSP YQIHFTKTPK YFKPGMPFDL
0351 MVFVTNPDGS PAYRVPVAVQ GEDTVQSLTQ GDGVAKLSIN THPSQKPLSI
0401 TVRTKKQELS EAEQATRMTQ ALPYSTVGNS NNYLHLSVLR TELRPGETLN
0451 VNFLLRMDRA HEAKIRYYTY LIMNKGRLK AGRQVREPGQ DLVVLP LSLIT
0501 TDFIPSRFLV AYYTLIGASG QREVVADSVW VDVKDCVGS LVVKSQGSSE

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0551 RQPVPGQQMT LKIEGDHGAR VVLVAVDKGV FVLNKKNKLT QSKIWDVVEK
 0601 ADIGCTPGSG KDYAGVFSDA GLTFTSSSGQ QTAQRAELQC PQPAARRRRS
 0651 VQLTEKRMCK VGKYPKELRK CCEDGMREN P MRFSCQRRTR FISLGEACKK
 0701 VFLDCCNYIT ELRRQHARAS HLGLARSNLD EDIIAENIV SRSEFPESWL
 0751 WNVEDLKEPP KNGISTKLMN IFLKDSITTW EILAVSMSDK KGICVADPFE
 0801 VTMQDFFID LRLPYSVVRN EQVEIRAVLY NYRQNQELKV RVELLHNPAP
 0851 CSLATTKRHR QQTVTIPPKS SLSVPYVIVP LKTGLQEVEV KAAVYHHFIS
 0901 DGVRKSLKVV PEGIRMNKT V AVRTLDPERL GREGVQKEDI PPADLSDQVP
 0951 DTESETRILL QGTPVAQMT E DAVDAERLKH LIVTPSGCGE QNMIGMTPTV
 1001 IAVHYLDETE QWEKFGLEKR QGALELIKKG YTQQLAFRQP SSAAFAFVKR
 1051 APSTWLTAYV VKVFSLAVNL IAIDSQVLCG AVKWLILEKQ KPDGVFQEDA
 1101 PVIHQEMIGG LRNNNEKDMA LTAFVLISLQ EAKDICEEQV NSLPGSITKA
 1151 GDFLEANYMN LQRSYTVAIA GYALAQMGRL KGPLLNKFLT TAKDKNRWD
 1201 PGKQLYNVEA TSYALLALLQ LKDFDFVPPV VRWLNEQRY Y GGGYGSTQAT
 1251 FMVFQALAQY QKDAPDHQEL NLDVSLQLPS RSSKITHRIH WESASLLRSE
 1301 ETKENEGFTV TAEGKGQGT L SVVTMYHAKA KDQLTCNKFD LKVTIKPAPE
 1351 TEKRPQDAKN TMILEICTRY RGDQDATMSI LDISMMTGFA PDTDDLKQLA
 1401 NGVDRIYSKY ELDKAFSDRN TLIIYLDKVS HSEDDCLAFK VHQYFNVELI
 1451 QPGAVKVYAY YNLEESCTRF YHPEKEDGKL NKLCRDELCR CAEENCFIQK
 1501 SDDKVTLEER LDKACEPGVD YVYKTRLVKV QLSNDFDEYI MAIEQTIKSG
 1551 SDEVQVGQQR TFISPIKRE ALKLEEKHY LMWGLSSDFW GEKPNLSYII
 1601 GKDTWVEHWP EEDECQDEEN QKQCQDLGAF TESMVVFGCP N

Complement C3-Derived Peptide CO3

[0089]

(SEQ ID NO: 2)
 APVIHQEMIGGLRN

(2) Transcription Factor AP-2 Gamma-Derived Peptide AP2C

[0090] For AP2C shown as SEQ ID NO: 4, area values of cognitive impairment (AD, MCI, DLB and FTD) were significantly higher than NDC. (t-test, $p < 0.05$) (see FIG. 5)

[0091] Thus, it was revealed that AP2C shown as SEQ ID NO: 4 was useful to distinguish patient of cognitive impairment (AD, MCI, DLB and FTD) with subjects not suffering from psychiatry disease (NDC). According to the analysis by receiver operating characteristic (ROC) curve, CO3 was clearly useful to distinguish AD and MCI with NDC. (See Table 1)

Intact Protein/Peptide

[0092]

(SEQ ID NO: 3)
 0001 MLWKITDNVK YEEDCEDRHD GSSNGNPRVP HLSSAGQHLY SPAPPLSHTG
 0051 VAEYQPPPYF PPPYQQLAYS QSADPYSHLG EAYAAAINPL HQPAPTGSQQ
 0101 QAWFGRQSQE GAGLPSHHGR PAGLLPHLSG LEAGAVSARR DAYRRSLLLL
 0151 PHAHALDAAG LAENLGLHDM PHQMDEVQNV DDQHLLLDHQ TVIRKGPISM
 0201 TKNPLNLPCQ KELVGAVMNP TEVFCVPGR LSLLSSTSKY KVTVAEVQRR

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0251 LSPPECLNAS LLGGVLRRAK SKNGGRSLRE KLDKIGLNLP AGRRKAAHVT
 0301 LLTSLVEGEA VHLARDFAYV CEAFFPSKPV AEYLTRPHLG GRNEMAARKN
 0351 MLLAAQQLCK EFTTELSQDR TPHGTSRLAP VLETNIQNCL SHFSLITHGF
 0401 GSQAICAAVS ALQNYIKEAL IVIDKSYMNP GDQSPADSNK TLEKMEKHRK

Transcription Factor AP-2 Gamma-Derived Peptide AP2C

Synapsin-3-Derived Peptide SYN3

[0093]

[0097]

(SEQ ID NO: 4)
 PGRQSQEGAGLP SHHG

(SEQ ID NO: 6)
 EMFGGLDICA VKAVHSHK

(3) Synapsin-3-Derived Peptide SYN3

[0094] For SYN3 shown as SEQ ID NO: 6, area values of cognitive impairment (AD, MCI, DLB and FTD) were significantly higher than NDC. (t-test, $p < 0.05$) (see FIG. 6)

[0095] Thus, it was revealed that SYN3 shown as SEQ ID NO: 6 was useful to distinguish patient of cognitive impairment (AD, MCI, DLB and FTD) with subjects not suffering from psychiatry disease (NDC). According to the analysis by receiver operating characteristic (ROC) curve, CO3 was clearly useful to distinguish AD and MCI with NDC. (See Table 1)

Intact Protein/Peptide

[0096]

(SEQ ID NO: 5)
 0001 MNFLRRRLSD SSFMANLPNG YMTDLQRPDS STSSPASPAM ERRHPQPLAA
 0051 SFSSPGSSLF SSLSSAMKQA PQATSGLMPEP PGPSTPIVQR PRILLVIDDA
 0101 HTDWSKYFHG KKVNGEIEIR VEQAEFSELN LAAYVTGGCM VDMQVVRNGT
 0151 KVSRSFSPKD FILVRQHAYS MALGEDYRSL VIGLQYGGLP AVNSLYSVYN
 0201 FCSKPWFVSQ LIKIFHSLGP EKFP LVEQTF FPNHKPMVTA PHFPVVVKLG
 0251 HAHAGMGKIK VENQLDFQDI TSVVAMAKTY ATTEAFIDSK YDIRIQKIGS
 0301 NYKAYMRTSI SGNWKANTGS AMLEQVAMTE RYRLWVDSCS EMFGGLDICA
 0351 VKAVHSHKDGR DYIIEVMDSS MPLIGEHVEE DRQLMADLVV SKMSQLPMPG
 0401 GTAPSPLRPW APQIKSAKSP GQAQLGPQLG QPQPRPPQ GPRQAQSPQP
 0451 QRSGPSQQR LSPQGQQPLS PQSGSPQQQR SPGSPQLSRA SSGSSPNQAS
 0501 KPGATLASQP RPPVQGRSTS QQGEESKKPA PPHPLNKSQ SLTNSLSTSD
 0551 TSQRGTPSED EAKAETIRNL RKSFAFLFSD

(4) Oxytocin Receptor-Derived Peptide OXYR

[0098] For OXYR shown as SEQ ID NO: 8, area values of cognitive impairment (AD, MCI, DLB and FTD) were significantly higher than NDC. (t-test, $p < 0.05$) (see FIG. 7) Thus, it was revealed that OXYR shown as SEQ ID NO: 8 was useful to distinguish patient of cognitive impairment (AD, MCI, DLB and FTD) with subjects not suffering from psychiatry disease (NDC). According to the analysis by receiver operating characteristic (ROC) curve, CO3 was clearly useful to distinguish AD and MCI with NDC. (See Table 1)

Intact Protein/Peptide

[0099]

(SEQ ID NO: 7)

0001 MEGALANWS AEAANASAP PGAEGNRTAG PPRNEALAR VEVAVLCIL

0051 LLALSGNACV LLALRTTRQK HSRLEFFMKH LSIADLVVAV FQVLPQLLWD

0101 ITFRFYGPD LCLRVKYLQV VGMFASTYLL LLMSLDRCLA ICQPLRSLRR

0151 RTDRLAVLAT WLGCLVASAP QVHIFSLREV ADGVFDCWAV FIQPWGPKAY

0201 ITWITLAVYI VPVIVLAACY GLISFKIWQN LRLKTAATAA AEAPEGAAAG

0251 DGGREVALARV SSVKLISKAK IRTVKMTFII VLAFLVCWTP FFFVQMWVW

0301 DANAPKEASA FIIVMLLASL NSCCNPWIYM LFTGHLFHEL VQRFLCCSAS

0351 YLKGRRRLGET SASKSNSSS FVLSHRSSSQ RSCSQPSTA

Oxytocin Receptor-Derived Peptide OXYR

[0100]

(SEQ ID NO: 8)

AAPPGAEGNRT

[0101] (5) Inter-Alpha-Trypsin Inhibitor Heavy Chain H5-like Protein-Derived Peptide ITH5L**[0102]** For ITH5L shown as SEQ ID NO: 10, area values of cognitive impairment (AD, MCI and DLB) were significantly higher than NDC. (t-test, $p < 0.05$) (see FIG. 8)

[0103] Thus, it was revealed that ITH5L shown as SEQ ID NO: 10 was useful to distinguish patient of cognitive impairment (AD, MCI and DLB) with subjects not suffering from psychiatry disease (NDC). According to the analysis by receiver operating characteristic (ROC) curve, CO3 was clearly useful to distinguish AD and MCI with NDC. (See Table 1)

Intact Protein/Peptide

[0104]

(SEQ ID NO: 9)

0001 GPPVPASSST KLLMTSYSMR STVVSRYAHT LVTSVLFNPH AEAHEAIFDL

0051 DLPHLAFISN FTMTINKVY IAEVKEKHQA KKIYEEAHQQ GKTAHVGIR

0101 DRESEKFRIS TSLAAGTEVT FSLAYEELLQ RHQGQYQLVV SLRPGQLVKR

0151 LSIEVTVSER TGISYVHIPP LRTGRLRTNA HASEVDSPPS TRIERGETCV

0201 RITYCPTLQD QSSISGSGIM ADPLVQYDVV MEDIIGDVQI YDDYFIHYFA

0251 PRGLPPMEKN VVFVIDVSS MFGTKMEQTK TAMNVILSDL QANDYFNIIIS

0301 FSDTVNVWKA GGSIQATIQN VHSADYLLHC MEADGWTDVN SALLAAASVL

0351 NHSNQEPGRG PSVGRIPLII FLTDGEPTAG VTPSVILSN VRQALGHRVS

0401 LFSLAFGDDA DFTLLRRLSL ENRGIAIRIY EDTDAALQLK GLYEIISMLP

0451 LADVRLNYLG GLVGASPWAV FPNYFGGSEL VVAGQVQPGK QELGIHLAAR

0501 GPKDQLLVAH HSEGATNNSQ KAFGCPGEPA PNVAFIRRL WAYVTIGELL

0551 DAHFQARDTT TRHLLAAKVL NLSLEYNFVT PLTSLVMVQP KQASEETRQ

0601 TSTSAGPDTI MPSSSRHGL GVSTAQPALV PKVISPKSRP VKPKFYLST

0651 TTASTKKMLS SKELEPLGES PHTLSMPTYP KAKIPAQQDS GTLAQPTLRT

0701 KPTILVPSNS GTLLPLKPGS LSHQNPDIPL TNSRTQVPPV KPGIPASPKA

0751 DTVKCVTPLH SKPGAPSHQ LGALTSQAPK GLPQSRPGVS TLQVPKYPLH

0801 TRPRVPAPKT RNNMPHLGPG ILLSKTPKIL LSLKPSAPPH QISTISLSK

0851 PETPNPHMPQ TPLPPRPDRP RPPLPESLST FPNTISSSTG PSSTTTTSLV

0901 GEPLPMFPTP TLPPGRFWHQ YDLLPGPQRT RQVLGSPSRPG VPTMSLLNSS

0951 RPTPEGSPPN LPILLPSSIL PEAISLLLLL EELLESSESM VESKFVESLN

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1001 PPAFYTFLTP DEDGSPNWDG NSEEILGGAG GSMESQGSSV GLAKGTLPSI
 1051 FTFSSSVDDG PHFVIQIPHS EEKICFTLNG HPGDLLQLIE DPKAGLHVSG
 1101 KLLGAPPRPG HKDQTRTYFQ IITVTTDKPR AYTITISRSS ISLRGEGTLR
 1151 LSWDQPALLK RPQLELYVAA AARLTLRLGP YLEFLVLRHR YRHPSTLQLP
 1201 HLGIFYVANGS GLSPSARGLI GQFQHADIRL VTGPMGPCLR RHGPDVVPVI
 1251 LGKRLLKDSP RLLPRWASCV LVKRSHVELL LGHPYLSYVL

Inter-Alpha-Trypsin Inhibitor Heavy Chain H5-like Protein-Derived Peptide ITH5L

RVSLFSLAFGDDAD (SEQ ID NO: 10)

(6) E3 Ubiquitin-Protein Ligase HERC2-Derived Peptide HERC2

[0105] For HERC2 shown as SEQ ID NO: 12, area values of cognitive impairment (AD, MCI and DLB) were significantly higher than NDC. (t-test, $p < 0.05$) (see FIG. 9)

[0106] Thus, it was revealed that HERC2 shown as SEQ ID NO: 12 was useful to distinguish patient of cognitive impairment (AD, MCI and DLB) with subjects not suffering from psychiatry disease (NDC). According to the analysis by receiver operating characteristic (ROC) curve, CO3 was clearly useful to distinguish AD and MCI with NDC. (See Table 1)

Intact Protein/Peptide

[0107]

(SEQ ID NO: 11)

0001 MPSESFCLAA QARLDSKWLK TDIQLAFTRD GLCGLWNEMV KDGEIVYTGT
 0051 ESTQNGELPP RKDDSVESPG TKKEDLNDKE KKDEETPAP IYRAKSILDS
 0101 WVGKQPDVN ELKECLSVLV KEQQALAVQS ATTTLSALRL KQRLVILERY
 0151 FIALNRTVFQ ENVVKWKSS GISLPPVDKK SSRPAGKGVE GLARVGSRAA
 0201 LSFAPAFRLR AWRSGEDADL CSELLQESLD ALRALPEASL FDESTVSSVW
 0251 LEVVERATRF LRSVVTGDVH GTPATKGPGS IPLQDQHLAL AILLELAVQR
 0301 GTLSQMLSAI LLLLQLWDSG AQETDNERSA QGTSAPLLPL LQRFQSIIICR
 0351 KDAPHSEGDM HLLSGPLSPN ESFLRYLTLP QDNELALDLR QTAVVVMABL
 0401 DRLATPCMP L CSSPTSHKG SLQEVIGWGL IGWKYANVI GPIQCEGLAN
 0451 LGVTQIACAE KRFLILSRNG RVYTQAYNSD TLAPQLVQGL ASRNIVKIAA
 0501 HSDGHHYAL AATGEVYSWG CGDGGR LGHG DTVPLEEPKV ISAFSGKQAG
 0551 KHVVHIACGS TYSAAITAEG ELYTWGRGNY GRLGHGSSSED EAIPMLVAGL
 0601 KGLKVIDVAC GSGDAQTLAV TENGQVWSWG DGDYGKLG RG GSDGCKTPKL
 0651 IEKLQDLVV KVRCSQFSI ALTKDGQVYS WGKGNQRLG HGTEEHVRY
 0701 KLEGLQGKK VIDVAAGSTH CLALTEDSEV HSWGSDNQCC HPDTRLRVTKP
 0751 EPAALPGLDT KHIVGIACGP AQSFWSWSSCS EWSIGLRVPF VVDICSMTFE
 0801 QLDLLLRQVS EGMDGSADWP PPQKECEVAV ATLNLLRLQL HAAISHQVDP
 0851 EFLGLGLGSI LLNSLKQTVV TCLASSAGVLS TVQSAAQAVL QSGWSVLLPT
 0901 AEERARALSA LLPCAVSGNE VNISPGRRFM IDLLVGSLMA DGGLESALHA
 0951 AITAEIQDIE AKKEAQKEKE IDEQEANAST FHRSTPLDK DLINTGICES
 1001 SGKQCLPLVQ LIQQLLRNIA SQTVARLKDV ARRISCLDF EQHSRERSAS
 1051 LDLLLRFRQL LISKLYPGES IGQTSDISSP ELMGVGSLLK KYTALLCTHI
 1101 GDILPVAASI ASTSWRHFAE VAYIVEGDFT GVLLPELVVS IVLLLSKNAG

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1151 LMQEAGAVPL LGGLLEHLDR FNHLAPGKER DDHEELAWPG IMESFFTQGN
1201 CRNNEEVTLI RKADLENHNK DGGFWTVIDG KVDIKDFQT QSLTGNSILA
1251 QFAGEDPVVA LEAALQFEDT RESMHAFVCG QYLEPDQEIV TIPDLGSLSS
1301 PLIDTERNLG LLLGLHASYL AMSTPLSPVE IECAKWLQSS IFSGGLQTSQ
1351 IHYSYNEEKD EDHCSSPGGT PASKSRLCSH RRALGDHSQA FLQAIADNNI
1401 QDHNVKDFLC QIERYCRQCH LTTPIMFPE HPVEEVGRLL LCCLLKHEDL
1451 GHVALSLVHA GALGIEQVKH RPLPKSVVDV CRVVYQAKCS LIKTHQEQGR
1501 SYKEVCAPVI ERLRFLFNL RPAVCNDLSI MSKFKLLSSL PRWRRIAQKI
1551 IRERRKKRVP KKPESTDDEE KIGNEESDLE EACILPHSPI NVDKRPIAIK
1601 SPKDKWQPLL STVTGVHKYK WLKQNVQGLY PQSPLLSTIA EFALKEEPVD
1651 VEKMRKCLKL QLERAERVLE GIDTILKLAS KNFLLPVQY AMFCGWQRLI
1701 PEGIDIGEPL TDCLKDVDLI PPFNRMLLEV TFGKLYAWAV QNIRNVLMDA
1751 SAKFKELGIQ PVPLQTITNE NPSGPSLGTI PQARFLLVML SMLTLQHGAN
1801 NLDLLLSNGM LALTQTALRL IGPSCDNVEE DMNASAQGAS ATVLEETRKE
1851 TAPVQLPVSG PELAAAMKIG TRVMRGVDWK WGDQDGPPEG LGRVIGELGE
1901 DGWIRVQWDT GSTNSYRMGK EGKYDLKLA LPAAQPSAE DSDTEDDSEA
1951 EQTERNIHPT AMMFTSTINL LQTLCLSAGV HAEIMQSEAT KTLGGLRML
2001 VESGTTDKTS SPNRLVYREQ HRSWCTLGFV RSIALTPQVC GALSSPQWIT
2051 LLMKVVEGHA PFTATSLQRQ ILAVHLLQAV LPSWDKTERA RDMKCLVEKL
2101 FDFLGSLLTT CSSDVPLLRE STLRRRRVRP QASLTATHSS TLAEEVVALL
2151 RTLHSLTQWN GLINKYINSQ LRSITHSVFG RPSEGAQLED YFPDSENPEV
2201 GGLMAVLAVI GGIDGRLRLG GQVMHDEFGE GTVTRITPKG KITVQFSDMR
2251 TCRVCPLNQL KLPFAVAFNV NNLPFTEPML SVWAQLVNLA GSKLEKHKIK
2301 KSTKQAFAGQ VDLDLLRCQQ LKLYILKAGR ALLSHQDKLR QILSQPAVQE
2351 TGTVHTDDGA VVSPDLGDMS PEGPQPPMIL LQQLLASATQ PSPVKAIFDK
2401 QELEAAALAV CQCLAVESTH PSSPGFEDCS SSEATTPVAV QHIRPARVKR
2451 RKQSPVPALP IVVQLMEMGF SRRNIEFALK SLTGASGNAS SLPGVEALVG
2501 WLLDHSDIQV TELSDADTVS DEYSDEEVVE DVDDAAYSMS TGAVVTESQT
2551 YKKRADFLSN DDYAVYVREN IQVGMVRCC RAYEEVCEGD VGKVIKLD RD
2601 GLHDLNVQCD WQKKGTYWV RYIHVELIGY PPPSSSHIK IGDKVRVKAS
2651 VTPPKYKWS VTHQSVGVVK AFSANGKDII VDFPQQSHWT GLLSEMELVP
2701 SIHPGVTCDG CQMPFINGSR FKCRNCDDFD FCETCFKTKK HNTRHTFGRI
2751 NEPGQSAVFC GRSGKQLKRC HSSQPGMLLD SWSRMVKSLN VSSSVNQASR
2801 LIDGSEPCWQ SSGSQGKHWI RLEIFPDVLV HRLKMIVDPA DSSYMPSLVV
2851 VSGGNSLNNL IELKTININP SDTTVPLND CTEYHRYIEI AIKQCRSSGI
2901 DCKIHGLILL GRIRAEEDL AAVPFLASDN EEEDEKGNs GSLIRKKAAG
2951 LESAAITRTK VFVWGLNDKD QLGGLKGSKI KVPSFSETLS ALNVVQVAGG
3001 SKSLFAVTVE GKVYACGEAT NGRLGLGISS GTVPIPRQIT ALSSYVVKV
3051 AVHSGGRHAT ALTVDGKVFS WGEEDDGKLG HFSRMNCDKP RLIEALKTKR

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3101 IRDIACGSSH SAALTSSGEL YTWGLGEYGR LGHGDNTTQL KPKMVKVLLG
3151 HRVIQVACGS RDAQTLALTD EGLVFSWGDG DFGKLGRGGS EGCNIPQNIE
3201 RLNGQGVCQI ECGAQFSLAL TKSGVVWTWG KGDYFRLGHG SDVHVRKPQV
3251 VEGLRGKKIV HVAVGALHCL AVTDSGQVYA WGDNDHGQQG NGTTTVNRKP
3301 TLVQGLEGQK ITRVACGSSH SWAWTTVDVA TPSVHEPVLF QTARDPLGAS
3351 YLGVPDADS SAASNKISGA SNSKPNRPSL AKILLSLDGN LAKQQALSHI
3401 LTALQIMYAR DAVVGALMPA AMIAPVECPS FSSAAPSDAS AMASPMNGEE
3451 CMLAVDIEDR LSPNPWQEK R EIVSSEDAVT PSAVTPSAPS ASARPPFIPVT
3501 DDLGAASIIA ETMTKTKEDV ESQNKAGPE PQALDEFTSL LIADDTRVVV
3551 DLLKLSVCSR AGDRGRDVL AVLSGMTAY PQVADMLLEL CVTELEDVAT
3601 DSQSGRLSSQ PVVVESSHYP TDDTSTSGTV KIPGAEGLRV EFDRQCSTER
3651 RHDPLTVMDG VNRIVSVRSG REWSDWSSEL RIPGDELKWK FIDSGSVNGW
3701 GWRFTVYPIM PAAGPKELLS DRCVLSCPSP DLVTCLLDFR LNLASNRSIV
3751 PRLAASLAAC AQLSALAASH RMWALQRLRK LLTTEFGQSI NINRLLEGND
3801 GETRALSFTG SALAALVKGL PEALQRQFEY EDPIVRGGKQ LLHSPFFKVL
3851 VALACDLELD TLPCCAETHK WAWFRRYCMA SRVAVALDKR TPLPRFLDE
3901 VAKKIRELMA DSENMDVLHE SHDIFKREQD EQLVQWMNRR PDDWTLSAGG
3951 SGTIYGWGHN HRGQLGGIEG AKVKVPTPCE ALATLRPVQL IGGEQTLFAV
4001 TADGKLYATG YGAGGRLGIG GTESVSTPTL LESIQHVFIK KVAVNSGGKH
4051 CLALSSEGEV YSWGEAEDGK LGHGNRSPCD RPRVIESLRG IEVVDVAAGG
4101 AHSACVTAAG DLYTWGKGRY GRLGHSDSED QLKPKLVEAL QGHRVVDIAC
4151 GSGDAQTLCL TDDDTVWSWG DGDYGKLGRG GSDGCKVPMK IDSLTGLGVV
4201 KVECGSQFSV ALTKSGAVYT WGKGDYHRLG HGSDDHVRRP RQVQGLQGKK
4251 VIAIATGSLH CVCCTEDGEV YTWGDNDGQ LGDGTTNAIQ RPRLVAALQG
4301 KKNRVACGS AHTLAWSTSK PASAGKLPAQ VPMEYNHLQE IPIIALRNRL
4351 LLLHHLSELF CPCIPMFDLE GSLDETGLGP SVGFDTLRGI LISQGKEAAF
4401 RKVVQATMVR DRQHGPVVEL NRIQVKRSRS KGGLAGPDGT KSVFGQMCAC
4451 MSSFGPDSLL LPHRVWKVKF VGESVDDCGG GYSESIAEIC EELQNGLTPL
4501 LIVTPNGRDE SGANRDCYLL SPAARAPVHS SMFRFLGVLL GIAIRTSPL
4551 SLNLAEPVWK QLAGMSLTIA DLSEVDKDFI PGLMYIRDNE ATSEEFEMS
4601 LPFTVPSASG QDIQLSSKHT HITLDNRAEY VRLAINYRLH EFDEQVAAR
4651 EGMARVVPVP LLSLFTGYEL ETMVCSPDI PLHLLKSVAT YKGIEPSASL
4701 IQWFWEVMES FSNTERSLFL RFVWGRTRLP RTIADFRGRD FVIQVLDKYN
4751 PPDHFLPESY TCFLLKLPR YSCKQVLEEK LKYAIHFCKS IDTDDYARIA
4801 LTGEPAADDS SDDSDNEDVD SFASDSTQDY LTGH

E3 Ubiquitin-Protein Ligase HERC2-Derived Peptide HERC2

[0108]

(SEQ ID NO: 12)
KLAELPAAQPSAEDSD

(7) Prothrombin-Derived Peptide THRB

[0109] For THRB shown as SEQ ID NO: 14, area values of cognitive impairment (AD, MCI, DLB and FTD) were significantly higher than NDC. (t-test, $p < 0.05$) (see FIG. 10)

(8) Transthyretin-Derived Peptide TTHY

[0113] For TTHY shown as SEQ ID NO: 16, area values of cognitive impairment (AD, MCI, DLB and FTD) were significantly higher than NDC. (t-test, $p < 0.05$) (see FIG. 11)

[0114] Thus, it was revealed that TTHY shown as SEQ ID NO: 16 was useful to distinguish patient of cognitive impairment (AD, MCI, DLB and FTD) with subjects not suffering from psychiatry disease (NDC). According to the analysis by receiver operating characteristic (ROC) curve, CO3 was clearly useful to distinguish AD and MCI with NDC. (See Table 1)

Intact Protein/Peptide

[0115]

(SEQ ID NO: 15)
0001 GPTGTGESKC PLMVKVLDAV RGSPAINVAV HVFRKAADDT WEPPASGKTS
0051 ESGELHGLTT EEEFVEGIYK VEIDTKSYWK ALGISPFHEH AEVVFANDS
0101 GPRRYTIAAL LSPYSYSTTA VVTNPKE

[0110] Thus, it was revealed that THRB shown as SEQ ID NO: 14 was useful to distinguish patient of cognitive impairment (AD, MCI, DLB and FTD) with subjects not suffering from psychiatry disease (NDC). According to the analysis by receiver operating characteristic (ROC) curve, CO3 was clearly useful to distinguish AD and MCI with NDC. (See Table 1)

Intact Protein/Peptide

[0111]

(SEQ ID NO: 13)
0001 ANTFLFEEVRK GNLERECVEE TCSYEEAFEA LESSTATDVF WAKYTACETA
0051 RTPRDKLAAC LEGNCAEGLG TNYRGHVNIT RSGIECQLWR SRYPHKPEIN
0101 STTHPGADLQ ENFCRNPDS TTGPWCYTDD PTVRRQECSE PVCQDQVTV
0151 AMTPRSEGS VNLSPPLEQC VPDRGQQYQG RLAVTTHGLP CLAWASQAQAK
0201 ALSKHQDFNS AVQLVENFCR NPDGDEEGVW CYVAGKPGDF GYCDLNYCEE
0251 AVEEETGDGL DEDSDRAIEG RTATSEYQTF FNPRTFGSGE ADCGLRPLFE
0301 KKSLEDKTER ELLESYIDGR IVEGSDAEIG MSPWQVLMFR KSPQELLCGA
0351 SLISDRWVLT AAHCLLYPPW DKNFTENDLL VRIGKHSRTR YERNIEKISM
0401 LEKIYIHPRY NWRENLDRI ALMKLKKPVA FSDYIHPVCL PDRETAASLL
0451 QAGYKGRVTG WGNLKETWTA NVGKGQPSVL QVVNLPIVER PVCKDSTRIR
0501 ITDNMFACGY KPDEGKRGDA CEGDSGGPFV MKSPFNNRWY QMGIVSWGEG
0551 CDRDGKYGFI THVFRLLKKWI QKVIDQFGE

Prothrombin-Derived Peptide THRB

[0112]

(SEQ ID NO: 14)
TATSEYQTFNPRTFGSGEAD

Transthyretin-Derived Peptide TTHY

[0116]

(SEQ ID NO: 16)
AVRGSPAINVAVHVFRKAAD

(9) Tumor Necrosis Factor Receptor Superfamily Member 16-Derived Peptide TNR16

[0117] For TNR16 shown as SEQ ID NO: 18, area values of cognitive impairment (AD, MCI, DLB and FTD) were significantly higher than NDC. (t-test, $p < 0.05$) (see FIG. 12)

[0118] Thus, it was revealed that TNR16 shown as SEQ ID NO: 18 was useful to distinguish patient of cognitive impairment (AD, MCI, DLB and FTD) with subjects not suffering from psychiatry disease (NDC). According to the analysis by receiver operating characteristic (ROC) curve, CO3 was clearly useful to distinguish AD and MCI with NDC. (See Table 1)

Intact Protein/Peptide
[0119]

(SEQ ID NO: 17)

0001 KEACPTGLYT HSGECKKACN LGEGVAQPCG ANQTVCEPCL DSVTFSDVVS

0051 ATEPCKPCTE CVGLQSMSAP CVEADDAVCR CAYGYYQDET TGRCEACRVC

0101 EAGSGLVFSC QDKQNTVCEE CPDGTYSDEA NHVDPCLPCT VCDTERQLR

0151 ECTRWADAEC EEIPGRWITR STPPEGSDST APSTQEPEAP PEQDLIASTV

0201 AGVVTTVMGS SQPVVTRGTT DNLIPVYCSI LAAVVVGLVA YIAFKRWNSC

0251 KQNKQGANSR PVNQTPPPEG EKLHSDSGIS VDSQSLHDQQ PHTQTASGQA

0301 LKGDGGGLYS LPPAKREEVE KLLNGSAGDT WRHLAGELGY QPEHIDSFTH

0351 EACPVRALLA SWATQDSATL DALLAALRRI QRADLVESLC SESTATSPV

Tumor Necrosis Factor Receptor Superfamily Member
16-Derived Peptide TNR16
[0120]

(SEQ ID NO: 18)

QTASGQALKGDGGGLYS

(10) Complement C4-A-Derived Peptide CO4-1

[0121] For CO4-1 shown as SEQ ID NO: 20, area values of cognitive impairment (AD, MCI, DLB and FTD) were significantly higher than NDC. (t-test, p<0.05) (see FIG. 13)

[0122] Thus, it was revealed that CO4-1 shown as SEQ ID NO: 20 was useful to distinguish patient of cognitive impair-

ment (AD, MCI, DLB and FTD) with subjects not suffering from psychiatry disease (NDC). According to the analysis by receiver operating characteristic (ROC) curve, CO3 was clearly useful to distinguish AD and MCI with NDC. (See Table 1)

[0123] After Biosynthesis, Complement C4-A protein is divided into C4 beta chain, Complement C4-A alpha chain and Complement C4 gamma chain by processing.

[0124] SEQ ID NO: 19 is amino acid sequence of intact Complement C4-A protein containing all of these processed peptides.

Intact Protein/Peptide
[0125]

(SEQ ID NO: 19)

0001 KPRLLLFSPS VVHLGVPLSV GVQLQDVPRG QVVKGSVFLR NPSRNNVPCS

0051 PKVDFTLSSE RDFALLSLQV PLKDAKSCGL HQLLRGPEVQ LVAHSPWLKD

0101 SLSRTTNIQG INLLFSSRRG HLFQLTDQPI YNPGQVRVYR VFALDQKMRP

0151 STDITITVMVE NSHGLRVRKK EVYMPSSIFQ DDFVIPDISE PGTWKISARF

0201 SDGLESNSST QFEVKKYVLP NFEVKITPGK PYILTPVPHL DEMQLDIQAR

0251 YIYGKPVQGV AYVRFGLLDE DGKKTFFRGL ESQTKLVNGQ SHISLSKAEF

0301 QDALEKLNMG ITDLQGLRLY VAAAIIESPG GEMEEAELTS WYFVSSPFSL

0351 DLSKTKRHLV PGAPFLLQAL VREMSGSPAS GIPVKVSATV SSPGSVPEVQ

0401 DIQONTDGSQ QVSIPIIIIPQ TISELQLSVS AGSPHPAIAR LTVAAPPSGG

0451 PGFLSIERPD SRPPRVGDTL NLNLRVAVSG ATFSHYYYMI LSRGQIVFMN

0501 REPKRTLTSV SVFVDHHLAP SFYFVAFYYH GDHPVANSR VDVQAGACEG

0551 KLELSVDGAK QYRNAGESVKL HLETDSLALV ALGALDTALY AAGSKSHKPL

0601 NMGKVFEAMN SYDLGCGPGG GDSALQVFQA AGLAFSDGDQ WTLRSRKLSC

0651 PKEKTTRKKR NVNFQKAIN KLGQYASPTA KRCCQDGVTR LPMMSCEQR

0701 AARVQQPDCR EPFLSCCQFA ESLRKKSRDK GQAGLQRALE ILQEEDLIDE

0751 DDIPVRSFFP ENWLRVETV DRFQILTLWL PDSLTTWEIH GLSLSKTKGL

0801 CVATPVQLRV FREFHLHLRL PMSVRRFEQL ELRPVLYNYL DKNLTVSVHV

0851 SPVEGLCLAG GGGLAQQVLV PAGSARPAF SVVPTAAAV SLKVVARGSF

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0901 EFPVGDAVSK VLQIEKEGAI HREELVYELN PLDHRGRTLE IPGNSDPNMI
 0951 PDGDFNSYVR VTASDPLDTL GSEGALSPGG VASLLRLPRG CGEQTMIIYLA
 1001 PTLAASRYLD KTEQWSTLPP ETKDHAVDLI QKGYMRIQQF RKADGSYAAW
 1051 LSRDSSSTWLT AFVLKVLSLA QEQVGGSPEK LQETSNWLLS QQQADGSFQD
 1101 PCPVLDRSMQ GGLVGNDQTV ALTAFTVIAL HHGLAVFQDE GAEPLKQORVE
 1151 ASISKANSFL GEKASAGLLG AHAAAITAYA LSLTKAPVDL LGVAHNNLMA
 1201 MAQETGDNLY WGSVTGSQSN AVSPTPAPRN PSDPMPQAPA LWIETTAYAL
 1251 LHLLLHEGKA EMADQASAWL TRQGSFQGGF RSTQDTVIAL DALSAWIAS
 1301 HTTEERGLNV TLSSTGRNGF KSHALQLNNR QIRGLEEELQ FSLGSKINVK
 1351 VGGNSKGTLL VLRTYNVLDL KNTTCQDLQI EVTVKGHVEY TMEANEDYED
 1401 YEYDELPAKD DPDAPLQPVV PLQLFEGRRN RRRREAPKVV EEQESRVHYT
 1451 VCIWRNGKVG LSGMAIADVT LLSGFHALRA DLEKLTSLSD RYVSHFETEG
 1501 PHVLLYFDSV PTSRECVGFE AVQEVVGLV QPASATLYDY YNPERRCSVF
 1551 YGAPSKSRLI ATLCSAEVCQ CAEGKCPQRQ RALERGLQDE DGYRMKFACY
 1601 YPRVEYGFQV KVLREDSRAA FRLFETKITQ VLHFTKDVKA AANQMRNFLV
 1651 RASCLRLLEP GKEYLIMGLD GATYDLEGHP QYLLDSNSWI EEMPSERLCR
 1701 STRQRAACAQ LNDFLQEYGT QGCQV

Complement C4-Derived Peptide C04-1

[0126]

(SEQ ID NO: 20)
 NGFKSHALQLNNRQIR

(11) Complement C4-B-Derived Peptide C04-1

[0127] From the results of MS/MS analysis and MASCOT database search, A sequence of C04-1 peptide as shown SEQ ID NO: 20 is an amino acid sequence present in the part

of topological region that is common to Complement C4-A protein (SEQ ID NO: 19) and Complement C4-B protein. After Biosynthesis, Complement C4-B protein is divided into C4 beta chain, Complement C4-B alpha chain and Complement C4 gamma chain by processing.

[0128] SEQ ID NO: 21 is amino acid sequence of intact Complement C4-B protein containing all of these processed peptides.

Intact Protein/Peptide

[0129]

(SEQ ID NO: 21)
 0001 KPRLLLFSPS VVHLGVPLSV GVQLQDVPRG QVVKGSVFLR NPSRNNVPSC
 0051 PKVDFTLSSE RDFALLSLQV PLKDAKSCGL HQLLRGPEVQ LVAHSPWLKD
 0101 SLRSTTNIQG INLLFSSRRG HFLQTDQPI YNPGQVRVYR VFALDQKMRP
 0151 STDTITVMVE NSHGLRVRKK EVYMPSSIFQ DDFVIPDISE PGTWKISARF
 0201 SDGLESNSST QFEVKYVLP NFEVKITPGK PYILTPVPHL DEMQLDIQAR
 0251 YIYGKPVQGV AYVRFGLLDE DGKKTFFRGL ESQTKLVNGQ SHISLSKAEF
 0301 QDALEKLNMG ITDLQGLRLY VAAAIIESPG GEMEEAELTS WYFVSSPFSL
 0351 DLSKTKRHLV PGAPFLQAL VREMSGSPAS GIPVKVSATV SSPGSVPEVQ
 0401 DIQNTDGSQ QVSIPIIIPQ TISELQLSVS AGSPHPAIAR LTVAAPPSGG
 0451 PGFLSIERP SRPPRVGDTL NLNLRAVGSG ATFSHYYYMI LSRGQIVFMN
 0501 REPKRTLTSV SVFVDHHLAP SFYFVAFYYH GDHPVANSR VDVQAGACEG
 0551 KLELSVDGAK QYRNGESVKL HLETDSLALV ALGALDTALY AAGSKSHKPL

-continued

0601 NMGKVFAMN SYDLGCGPGG GDSALQVFQA AGLAFSDGDQ WTLSRKRLSC
 0651 PKEKTRKKR NVNFQKAIN KLGQYASPTA KRCCQDGVTR LPMMSCEQR
 0701 AARVQQPDCR EPFLSCCQFA ESLRKSRDK GQAGLQRALE ILQEEDLIDE
 0751 DDIPVRSFFP ENWLWRVETV DRFQILTTLWL PDSLTWEIH GLSLSKTKGL
 0801 CVATPVQLRV FREFHLHLRL PMSVRRFEQL ELRPVLYNYL DKNLTVSVHV
 0851 SPVEGLCLAG GGGLAQQVLV PAGSARPVAF SVVPTAAAV SLKVVARGSF
 0901 EFPVGDAVSK VLQIEKEGAI HREELVYELN PLDHRGRTLE IPGNSDPNMI
 0951 PDGDFNSYVR VTASDPLDTL GSEGALSPGG VASLLRLPRG CGEQMTIYLA
 1001 PTLAASRYLD KTEQWSTLPP ETKDHAVDLI QKGYMRIQQF RKADGSYAAW
 1051 LSRDSSTWLT AFVLKVLSLA QEQVGGSPK LQETSNWLLS QQQADGSFQD
 1101 LSPVIHRSMQ GGLVGNDTV ALTAFTVIAL HHGLAVFQDE GAEPLKQVE
 1151 ASISKANSFL GEKASAGLLG AHAAAITAYA LSLTKAPVDL LGVAHNLMMA
 1201 MAQETGDNLY WGSVTGSQSN AVSPTAPARN PSDPMPQAPA LWIETTAYAL
 1251 LHLLEHGA EMADQASAWL TRQGSFQGGF RSTQDTVIAL DALSAWIAS
 1301 HTTEERGLNV TLSTGRNGF KSHALQLNNR QIRGLEEELQ FSLGSKINVK
 1351 VGGNSKGTLK VLRTYNVLDL KNTTCQDLQI ETVKGVHVEY TMEANEDYED
 1401 YEYDELPAK DPDAPLQPV PLQLFEGRRN RRRREAPKV EEQESRVHYT
 1451 VCIWRNGKVG LSGMAIADVT LLSGFHALRA DLEKLTSLS RYVSHFETEG
 1501 PHVLLYFDSV PTSRECVGFE AVQEVVGLV QPASATLYDY YNPERRCVSF
 1551 YGAPSKSRL ATLCSAEVCQ CAEGKCPQR RALERGLQDE DGYRMKFACY
 1601 YPRVEYGFQV KVLREDSRAA FRLFETKITQ VLHFTKDVKA AANQMRNFLV
 1651 RASCRLRLEP GKEYLIMGLD GATYDLEGHP QYLLDSNSWI EEMPSERLCR
 1701 STRQRAACAQ LNDFLQEYGT QGCQV

Just in case, following shows the sequence of CO4-1.

ComplementC4-Derived Peptide CO4-1

[0130]

(SEQ ID NO: 20)
 NGFKSHALQLNNRQIR

(12) Complement C4-A-Derived Peptide CO4-2

[0131] For CO4-2 shown as SEQ ID NO: 22, area values of cognitive impairment (AD, MCI, DLB and FTD) were significantly higher than NDC. (t-test, $p < 0.05$) (see FIG. 14)

[0132] Thus, it was revealed that CO4-2 shown as SEQ ID NO: 22 was useful to distinguish patient of cognitive impairment (AD, MCI, DLB and FTD) with subjects not suffering from psychiatry disease (NDC). According to the analysis by receiver operating characteristic (ROC) curve, CO3 was clearly useful to distinguish AD and MCI with NDC. (See Table 1) After Biosynthesis, Complement C4-A protein is divided into C4 beta chain, Complement C4-A alpha chain and Complement C4 gamma chain by processing.

[0133] SEQ ID NO: 19 is amino acid sequence of intact Complement C4-A protein containing all of these processed peptides.

Intact Protein/Peptide

[0134]

(SEQ ID NO: 19)
 0001 KPRLLLFSPS VVHLGVPLSV GVQLQDVPRG QVVKGSVFLR NPSRNNVPCS
 0051 PKVDFTLSSE RDFALLSLQV PLKDAKSCGL HQLLRGPEVQ LVAHSPWLKD
 0101 SLRSTTNIQG INLLFSSRRG HLFLQTDQPI YNPGQVRVYR VFALDQKMRP
 0151 STDITVMVE NSHGLRVKKE EVYMPSSIFQ DDFVIPDISE PGTWKISARF
 0201 SDGLESNSST QFEVKYVLP NFEVKITPGK PYILTVPGHL DEMQLDIQAR

-continued

0251 YIYGKPVQGV AYVRFGLLDE DGKKTFFRGL ESQTKLVNGQ SHISLSKAEF
 0301 QDALEKLNMG ITDLQGLRLY VAAAIIESPG GEMEEAELTS WYFVSSPFSL
 0351 DLSKTKRHLV PGAPFLLQAL VREMSGSPAS GIPVKVSATV SSPGSVPEVQ
 0401 DIQQNTDGSQ QVSIPIIIPQ TISELQLSVS AGSPHPAIAR LTVAAPPSGG
 0451 PGFLSIERPD SRPPRVGDTL NLNLRVAVSG ATFSHYYYMI LSRGQIVFMN
 0501 REPKRTLTSV SVFVDHHLAP SFYFVAFYYH GDHPVANSLR VDVQAGACEG
 0551 KLELSVDGAK QYRNGESVKL HLETDSLALV ALGALDTALY AAGSKSHKPL
 0601 NMGKVFAMN SYDLGCGPGG GDSALQVFQA AGLAFSDGDQ WTLNRKRLSC
 0651 PKEKTTKRKR NVNFQKAIN KLGQYASPTA KRCCQDGVTR LPMRSCEQR
 0701 AARVQQPDCR EPFLSCCQFA ESLRKSRDK GQAGLQRALE ILQEEDLIDE
 0751 DDIPVRSFFP ENWLWRVETV DRFQILTTLWL PDSLTTWEIH GLSLSKTKGL
 0801 CVATPVQLRV FREFHLHLRL PMSVRRFEQL ELRPVLYNYL DKNLTVSVHV
 0851 SPVEGLCLAG GGGLAQQLV PAGSARPAF SVVPTAAAV SLKVVARGSF
 0901 EFPVGDAVSK VLQIEKGAH HREELVYELN PLDHRGRTLE IPGNSDPNMI
 0951 PDGDFNSYVR VTASDPLDTL GSEGALSPGG VASLLRLPRG CGEQTMIIYA
 1001 PTLAASRYLD KTEQWSTLPP ETKDHAVDLI QKGYMRIQQF RKADGSYAAW
 1051 LSRDSSTWLT AFVLKVLSLA QEQVGSPEK LQETSNWLLS QQQADGSFQD
 1101 PCPVLDRSMQ GGLVGNDETALV ALTAFTVIAL HHGLAVFQDE GAEPLKQVE
 1151 ASISKANSFL GEKASAGLLG AHAHAITAYA LSLTKAPVDL LGVAHNNLMA
 1201 MAQETGDNLY WGSVTGSQSN AVSPTAPARN PSDPMPQAPA LWIETTAYAL
 1251 LHLLEHGA EMADQASAWL TRQGSFQGGF RSTQDTVIAL DALSAYWIAS
 1301 HTTEERGLNV TLSSTGRNGF KSHALQLNNR QIRGLEELQ FSLGSKINVK
 1351 VGGNSKGTLLK VLRTYNVLDL KNTTCQDLQI EVTVKGHVEY TMEANEDYED
 1401 YEYDELPAKD DPDAPLQPV PLQLFEGRRN RRRREAPKVV EEQESRVHYT
 1451 VCIWRNGKVG LSGMAIADVT LLSGFHALRA DLEKLTSLSD RYVSHFETEG
 1501 PHVLLYPDSV PTSRECVGFE AVQEVVGLV QPASATLYDY YNPERRCSVF
 1551 YGAPSKSRLD ATLCSEAECQ CAEGKCPQR RALERGLQDE DGYRMKFACY
 1601 YPRVEYGFQV KVLREDSRAA FRLFETKITQ VLHFTKDVKA AANQMRNFLV
 1651 RASCLRLRLEP GKEYLIMGLD GATYDLEGHP QYLLDSNSWI EEMPSERLCR
 1701 STRQRAACAQ LNDFLQEYGT QGCQV

Complement C4-Derived Peptide CO4-2

[0135]

(SEQ ID NO: 22)

APLQPVTPQLFEGRRN

(13) Complement C4-B-Derived Peptide CO4-2

[0136] From the results of MS/MS analysis and MASCOT database search, A sequence of CO4-2 peptide as shown SEQ ID NO: 22 is an amino acid sequence present in the part of topological region that is common to Complement C4-A protein (SEQ ID NO: 19) and Complement C4-B protein. After Biosynthesis, Complement C4-B protein is divided into C4 beta chain, Complement C4-B alpha chain and Complement C4 gamma chain by processing. SEQ ID NO: 21 is amino acid sequence of intact Complement C4-B protein containing all of these processed peptides.

Intact Protein/Peptide

[0137]

(SEQ ID NO: 21)

0001 KPRLLLFSPS VVHLGVPLSV GVQLQDVPRG QVVKGSVFLR NPSRNNVPCS
0051 PKVDFTLSSE RDFALLSLQV PLKDAKSCGL HQLLRGPEVQ LVAHSPWLKD
0101 SLSRTTNIQG INLLFSSRRG HLFLQTDQPI YNPGQVRVYR VFALDQKMRP
0151 STDITITVMVE NSHGLRVRKK EYMPSSIFQ DDFVIPDISE PGTWKISARF
0201 SDGLESNSST QFEVKKYVLP NFEVKITPGK PYILTVPGHL DEMQLDIQAR
0251 YIYGKPVQGV AYVRFGLLDE DGKKTFFRGL ESQTKLVNGQ SHISLSKAEF
0301 QDALEKLNMG ITDLQGLRLY VAAAIIESPG GEMEEAELTS WYFVSSPFFSL
0351 DLSKTKRHLV PGAPFLLQAL VREMSGSPAS GIPVKVSATV SSPGSVPEVQ
0401 DIQQNTDGGG QVSIPIIIPQ TISELQLSVS AGSPHPAIAR LTVAAPPSGG
0451 PGFLSIERPD SRPPRVGDTL NLNLRAVGSG ATFSHYYMI LSRGQIVFMN
0501 REPKRTLTSV SVFVDHHLAP SFYFVAFYYH GDHPVANSLR VDVQAGACEG
0551 KLELSVDGAK QYRNGESVKL HLETDSLALV ALGALDTALY AAGSKSHKPL
0601 NMGKVFAMN SYDLGCGPGG GDSALQVFQA AGLAFSDGDQ WTLRKRLSC
0651 PKEKTTTRKKR NVNFQKAIN KLGQYASPTA KRCCQDGVTR LPMMRSCQQR
0701 AARVQQPDCR EPFLSCCQFA ESLRKSRDK GQAGLQRALE ILQEEDLIDE
0751 DDIPVRSEFP ENWLWRVETV DRFQILTLWL PDSLTTWEIH GLSLSKTKGL
0801 CVATPVQLRV FREFHLHLRL PMSVRRFEQL ELRPVLYNYL DKNLTVSVHV
0851 SPVEGLCLAG GGGLAQQVLV PAGSARPAF SVVPTAAAV SLKVVARGSF
0901 EFPVGDAVSK VLQIEKEGAI HREELVYELN PLDHRGRTLE IPGNSDPNMI
0951 PDGDFNSYVR VTASDPLDTL GSEGALSPGG VASLLRLPRG CGEQTMIIYA
1001 PTLAASRYLD KTEQWSTLPP ETKDHAVDLI KGYMRIQQF RKADGSYAAW
1051 LSRDSSTWLT AFVLKVLSLA QEQVGSPEK LQETSNWLLS QQQADGSFQD
1101 LSPVIHRSMQ GGLVGNDTV ALTAFTIAL HHGLAVFQDE GAEPLKQRE
1151 ASISKANSFL GEKASAGLLG AHAATAIYAY LSLTKAPVDL LGVAHNLMMA
1201 MAQETGDNLY WGSVTGSQSN AVSPTPAPRN PSDPMPQAPA LWIETTAYAL
1251 LHLLEHEGKA EMADQASAWL TRQGSFQGGF RSTQDTVIAL DALSAWIAS
1301 HTTEERGLNV TLSSTGRNGF KSHALQLNRR QIRGLEELQ FSLGSKINVK
1351 VGGNSKGLK VLRTYNVLDL KNTTCQDLQI EVTVKGHVEY TMEANEDYED
1401 YEYDELPAKD DPDAPLQPV PLQLFEGRRN RRRREAPKV EEQESRVHYT
1451 VCIWRNGKVG LSGMAIADVT LLSGFHALRA DLEKLTSLSL RYVSHFETEG
1501 PHVLLYFDSV PTSRECVGFE AVQEVVGLV QPASATLYDY YNPERRCSVF
1551 YGAPSKSRL ATLCSAEVCQ CAEGKCPQR RALERGLQDE DGYRMKFACY
1601 YPRVEYGFQV KVLREDSRAA FRLFETKITQ VLHFTKDVKA AANQMRNFLV

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1651 RASCRLRLREP GKEYLIMGLD GATYDLEGHP QYLSDNSWI EEMPSERLCR

1701 STRQRAACAQ LNDFLQEQYGT QGCQV

[0138] Just in case, following shows the sequence of Fibrinogen Alpha Chain-Derived Peptide FIBA-1 CO4-2.

Complement C4-Derived Peptide CO4-2

[0139]**[0143]**

(SEQ ID NO: 22)
APLQPVTPPLQLFEGRRN

(SEQ ID NO: 24)
SSSYSKQFTSSTSYNRGDSTFES

(14) Fibrinogen Alpha Chain (Isoform 1)-Derived Peptide FIBA-1

[0140] For FIBA-1 shown as SEQ ID NO: 24, area values of cognitive impairment (AD, MCI, DLB and FTD) were significantly higher than NDC. (t-test, $p < 0.05$) (see FIG. 15)
[0141] Thus, it was revealed that FIBA-1 shown as SEQ ID NO: 24 was useful to distinguish patient of cognitive impairment (AD, MCI, DLB and FTD) with subjects not suffering from psychiatry disease (NDC). According to the analysis by receiver operating characteristic (ROC) curve, CO3 was clearly useful to distinguish AD and MCI with NDC.

(See Table 1)

Intact Protein/Peptide

[0142]

(15) Fibrinogen Alpha Chain (Isoform 2)-Derived Peptide FIBA-1

[0144] From the results of MS/MS analysis and MASCOT database search, A sequence of FIBA-1 peptide as shown SEQ ID NO: 24 is an amino acid sequence present in the part of topological region that is common to Fibrinogen alpha chain (isoform 1) (SEQ ID NO: 23) and Fibrinogen alpha chain (isoform 2). Followings, as SEQ ID NO: 25, an amino acid sequence of intact protein of Fibrinogen alpha chain (isoform 2) were shown.

(SEQ ID NO: 23)

0001 GPRVVERHQ S ACKDSWPFC SDEDWNYKCP SGC RMKGLID EVNQDFTNRI

0051 NKLKNSLF EY QKNNKDSHSL TTNIMEILRG DFSSANNRDN TYNRVSEDLR

0101 SRIEVLKRK V IEKVQHIQLL QKNVRAQLVD MKRLEVDIDI KIRSCRGSCS

0151 RALAREVDL K DYEDQKQLE QVIAKDLLPS RDRQHLPLIK MKPVPDLVPG

0201 NFKSQLQKVP PEWKALTDMP QMRMELERPG GNEITRGGST SYGTGSETES

0251 PRNPSSAGS W NSGSSGPGST GNRNPGSSGT GGTATWKPGS SGPGSTGSWN

0301 SGSSGTGSTG NQNP GS PRPG STGTWNP GSS ERGSAGH WTS ESSVSGSTGQ

0351 WHSESGSFRP DSPGSGNARP NNPDWGTFEE VSGNVSPGTR REYHTEKLV T

0401 SKGDKELRTG KEKVTSGSTT TTRRSCSKTV TKTVIGPDGH KEVTKEVVT S

0451 EDGSDCPEAM DLGTL SGIGT LDGFRHRHPD EAAFFDTAST GKTFP GFFSP

0501 MLGEFVSETE SRGSESGIFT NTKESSSHHP GIAEFPSRGK SSSYSKQFTS

0551 STSYNRGDST FESKSYKMAD EAGSEADHEG THSTKRGHAK SRPVRDCDDV

0601 LQTHPSGTQS GIFNIKLP GS SKIFSVYCDQ ETSLGWLLI QQRM DGSLNF

0651 NRTWQDYKRG FGSLNDEGEG EFWLGNDYLH LLTQRGSVLR VELEDWAGNE

0701 AYA EYHFRVG SEAEGYALQV SSYEGTAGDA LIEGSVEEGA EYTSHNNMQF

0751 STFDRDADQW EENCAEVYGG GWWYNNCQAA NLNGIYYPGG SYDPRNNSPY

0801 EIENG VVWVS FRGADYSLRA VRMKIRPLVT Q

Intact Protein/Peptide
[0145]

(SEQ ID NO: 25)

0001 GPRVVERHQS ACKDSWPFPC SDEDWNYKCP SGCRMKGLID EVNQDFTNRI

0051 NKLKNSLFY QKNNKDSHSL TTNIMEILRG DFSSANNRDN TYNRVSEDLR

0101 SRIEVLKRKV IEKVQHIQLL QKNVRAQLVD MKRLEVDIDI KIRSCRGSCS

0151 RALAREVDLK DYEDQQKQLE QVIAKDLLPS RDRQHLPLIK MKPVPDLVPG

0201 NFKSQQLKVP PEWKALTDMP QMRMELERPG GNEITRGGST SYGTGSETES

0251 PRNPSSAGSW NSGSSGPGST GNRNPGSSGT GGTATWKPGS SGPGSTGSWN

0301 SGSSGTGSTG NQNPSPRP G STGTWNP GSS ERGSAGH WTS ESSVSGSTGQ

0351 WHSESGSFRP DSPGSGNARP NNPDWGTFEE VSGNVSPGTR REYHTEKLV

0401 SKGDKELRTG KEKVTSGST TTRRSCSKTV TKTVIGPDGH KEVTKEVTS

0451 EDGSDCPEAM DLGTLGIGT LDGFRHRHPD EAAFFDTAST GKTFFGFFSP

0501 MLGEFVSETE SRGSESGIFT NTKESSHHP GIAEFPSRGK SSSYSKQFTS

0551 STSYNRGDST FESKSYKMAD EAGSEADHEG THSTKRGHAK SRPVRGIHTS

0601 PLGKPSLSP

[0146] Just in case, following shows the sequence of FIBA-1.

Fibrinogen Alpha Chain-Derived Peptide FIBA-1

[0147]

(SEQ ID NO: 24)

SSSYSKQFTSSTSYNRGDSTFES

(16) Fibrinogen Alpha Chain (Isoform 1)-Derived Peptide FIBA-2

[0148] For FIBA-2 shown as SEQ ID NO: 26, area values of cognitive impairment (AD, MCI, DLB and FTD) were significantly higher than NDC. (t-test, p<0.05) (see FIG. 16)

[0149] Thus, it was revealed that FIBA-2 shown as SEQ ID NO: 26 was useful to distinguish patient of cognitive impairment (AD, MCI, DLB and FTD) with subjects not suffering from psychiatry disease (NDC). According to the analysis by receiver operating characteristic (ROC) curve, CO3 was clearly useful to distinguish AD and MCI with NDC.

(See Table 1)

Intact Protein/Peptide

[0150]

(SEQ ID NO: 23)

0001 GPRVVERHQS ACKDSWPFPC SDEDWNYKCP SGCRMKGLID EVNQDFTNRI

0051 NKLKNSLFY QKNNKDSHSL TTNIMEILRG DFSSANNRDN TYNRVSEDLR

0101 SRIEVLKRKV IEKVQHIQLL QKNVRAQLVD MKRLEVDIDI KIRSCRGSCS

0151 RALAREVDLK DYEDQQKQLE QVIAKDLLPS RDRQHLPLIK MKPVPDLVPG

0201 NFKSQQLKVP PEWKALTDMP QMRMELERPG GNEITRGGST SYGTGSETES

0251 PRNPSSAGSW NSGSSGPGST GNRNPGSSGT GGTATWKPGS SGPGSTGSWN

0301 SGSSGTGSTG NQNPSPRP G STGTWNP GSS ERGSAGH WTS ESSVSGSTGQ

0351 WHSESGSFRP DSPGSGNARP NNPDWGTFEE VSGNVSPGTR REYHTEKLV

0401 SKGDKELRTG KEKVTSGST TTRRSCSKTV TKTVIGPDGH KEVTKEVTS

0451 EDGSDCPEAM DLGTLGIGT LDGFRHRHPD EAAFFDTAST GKTFFGFFSP

0501 MLGEFVSETE SRGSESGIFT NTKESSHHP GIAEFPSRGK SSSYSKQFTS

0551 STSYNRGDST FESKSYKMAD EAGSEADHEG THSTKRGHAK SRPVRCDDV

0601 LQTHPSGTQS GIFNIKLP GS SKIFSVYCDQ ETSLGWLLI QQRMDSGLNF

-continued

0651 NRTWQDYKRG FGS LNDEGEG EFWLGNDYLH LLTQRGSVLR VELEDWAGNE

0701 AYAETHFRVG SEAEGYALQV SSYEGTAGDA LIEGSVEEGA EYTSHNNMQF

0751 STFD RDADQW EENCAEVYGG GWYNNCQAA NLNGIYYPGG SYDPRNNSPY

0801 EIENGVVWVS FRGADYSLRA VRMKIRPLVT Q

Fibrinogen Alpha Chain-Derived Peptide FIBA-2

[0151]

(SEQ ID NO: 26)
SSSYSKQFTSSTS YNRGDSTFESKS

[0154] Just in case, following shows the sequence of FIBA-2.

Fibrinogen Alpha Chain-Derived Peptide FIBA-2

[0155]

(SEQ ID NO: 26)
SSSYSKQFTSSTS YNRGDSTFESKS

(17) Fibrinogen Alpha Chain (Isoform 2)-Derived Peptide FIBA-2

[0152] From the results of MS/MS analysis and MASCOT database search, A sequence of FIBA-2 peptide as shown SEQ ID NO: 26 is an amino acid sequence present in the part of topological region that is common to Fibrinogen alpha chain (isoform 1) (SEQ ID NO: 23) and Fibrinogen alpha chain (isoform 2). Followings, as SEQ ID NO: 25, an amino acid sequence of intact protein of Fibrinogen alpha chain (isoform 2) were shown.

Intact Protein/Peptide

[0153]

(SEQ ID NO: 25)

0001 GPRVVERHQS ACKDSWPFC SDEDWNYKCP SGC RMKGLID EVNQDFTNRI

0051 NKLNLSLFEY QKNNKDSHSL YFNIMEILRG DFSSANNRDN TYNRVSEDLR

0101 SRIEVLKRKV IEKVQHIQLL QKNVRAQLVD MKRLEVDIDI KIRSCRGSCS

0151 RALAREVDLK DYEDQKQLE QVIAKDLLPS RDRQHLPLIK MKPVPDLVPG

0201 NFKSQQLQKVP PEWKALTDMP QMRMELERPG GNEITRGGST SYGTGSETES

0251 PRNPSSAGSW NSGSSGPGST GNRNPGSSGT GGTATWKPGS SGPSTGSWN

0301 SGSSGTGSTG NQNPSPRPG STGTWNPSS ERGSAGHWS ESSVSGSTGQ

0351 WHSESGSFRP DSPGSGNARP NNPDWGTFEE VSGNVSPGTR REYHTEKLVT

0401 SKGDKELRTG KEKVTSGSTT TTRRSCSKTV TKTVIGPDGH KEVTKEVVTs

0451 EDGSDCPEAM DLGTLGGIGT LDGFRHRHPD EAAFFDTAST GKTFFGFFSP

0501 MLGEFVSETE SRGSESGIFT NTKESSHHP GIAEFPSRGK SSSYSKQFTS

0551 STSYNRGDST FESKSYKMAD EAGSEADHEG THSTKRGHAK SRPVRGIHTS

0601 PLGKPSLSP

(18) Fibrinogen Alpha Chain (Isoform 1)-Derived Peptide FIBA-3

[0156] For FIBA-3 shown as SEQ ID NO: 27, area values of cognitive impairment (AD, MCI, DLB and FTD) were significantly higher than NDC. (t-test, $p < 0.05$) (see FIG. 17)

[0157] Thus, it was revealed that FIBA-3 shown as SEQ ID NO: 27 was useful to distinguish patient of cognitive impairment (AD, MCI, DLB and FTD) with subjects not suffering from psychiatry disease (NDC). According to the analysis by receiver operating characteristic (ROC) curve, CO3 was clearly useful to distinguish AD and MCI with NDC. (See Table 1)

Intact Protein/Peptide
[0158]

(SEQ ID NO: 23)

0001 GPRVVERHQS ACKDSWPFCSDEDWNYKCP SGCRMKGLID EVNQDFTNRI

0051 NKLNLSLFEY QKNKDSHSL TTNIMEILRG DFSSANNRDN TYNRVSEDLR

0101 SRIEVLKRKV IEKVQHIQLL QKNVRAQLVD MKRLEVDIDI KIRSCRGSCS

0151 RALAREVDLK DYEDQQKQLE QVIAKDLLPS RDRQHLPLIK MKPVPDLVPG

0201 NFKSQQLQKVP PEWKALTDMP QMRMELERPG GNEITRGGST SYGTGSETES

0251 PRNPSSAGSW NSGSSGPGST GNRNPGSSGT GGTATWKPGS SGPGSTGSWN

0301 SGSSGTGSTG NQNPSPRPSTGTWNPSS ERGSAGHWTS ESSVSGSTGQ

0351 WHSESGSFRP DSPSGNARP NNPDWGTFEE VSGNVSPGTR REYHTEKLV

0401 SKGDKELRTG KEKVTSGSTT TTRRSCSKTV TKTVIGPDGH KEVTKEVTS

0451 EDGSDCPEAM DLGTLSGIGT LDGFRHRHPD EAAFFDTAST GKTFFGFFSP

0501 MLGEFVSETE SRGSESGIFT NTKESSHHP GIAEFPSRGK SSSYSKQFTS

0551 STSYNRGDST FESKSYKMAD EAGSEADHEG THSTKRGHAK SRPVRDCDDV

0601 LQTHPSGTQS GIFNIKLPSSKIFSVYCDQ ETSLGGWLLI QQRMDSGLNF

0651 NRTWQDYKRG FGSLNDEGEG EFWLGNDYLH LLTQRGSVLR VELEDWAGNE

0701 AYAETHFRVG SEAEYALQV SSYEGTAGDA LIEGSVEEGA EYTSNNMQF

0751 STFDRDADQW EENCAEVYGG GWYNNCQAA NLNGIYYPGG SYDPRNNSPY

0801 EIENGVVWVS FRGADYSLRA VRMKIRPLVT Q

Fibrinogen Alpha Chain-Derived Peptide FIBA-3
[0159]

(SEQ ID NO: 27)

SSSYSKQFTSSTSYNRGDSTFESKSY

SEQ ID NO: 27 is an amino acid sequence present in the part of topological region that is common to Fibrinogen alpha chain (isoform 1) (SEQ ID NO: 23) and Fibrinogen alpha chain (isoform 2). Followings, as SEQ ID NO: 25, an amino acid sequence of intact protein of Fibrinogen alpha chain (isoform 2) were shown.

(19) Fibrinogen Alpha Chain (Isoform 2)-Derived Peptide FIBA-3

[0160] From the results of MS/MS analysis and MASCOT database search, A sequence of FIBA-3 peptide as shown

Intact Protein/Peptide
[0161]

(SEQ ID NO: 25)

0001 GPRVVERHQS ACKDSWPFCSDEDWNYKCP SGCRMKGLID EVNQDFTNRI

0051 NKLNLSLFEY QKNKDSHSL TTNIMEILRG DFSSANNRDN TYNRVSEDLR

0101 SRIEVLKRKV IEKVQHIQLL QKNVRAQLVD MKRLEVDIDI KIRSCRGSCS

0151 RALAREVDLK DYEDQQKQLE QVIAKDLLPS RDRQHLPLIK MKPVPDLVPG

0201 NFKSQQLQKVP PEWKALTDMP QMRMELERPG GNEITRGGST SYGTGSETES

0251 PRNPSSAGSW NSGSSGPGST GNRNPGSSGT GGTATWKPGS SGPGSTGSWN

0301 SGSSGTGSTG NQNPSPRPSTGTWNPSS ERGSAGHWTS ESSVSGSTGQ

0351 WHSESGSFRP DSPSGNARP NNPDWGTFEE VSGNVSPGTR REYHTEKLV

0401 SKGDKELRTG KEKVTSGSTT TTRRSCSKTV TKTVIGPDGH KEVTKEVTS

0451 EDGSDCPEAM DLGTLSGIGT LDGFRHRHPD EAAFFDTAST GKTFFGFFSP

0501 MLGEFVSETE SRGSESGIFT NTKESSHHP GIAEFPSRGK SSSYSKQFTS

-continued

0551 STSYNRGDST FESKSYKMAD EAGSEADHEG THSTKRGHAK SRPVRIHTS

0601 PLGKPSLSP

[0162] Just in case, following shows the sequence of FIBA-3.

Fibrinogen Alpha Chain-Derived Peptide FIBA-3

[0163]

(SEQ ID NO: 27)

SSSYSKQFTSSTSYNRGDSTFESKSY

TABLE 1

Marker Peptide			
Sequence No.	Sequence name	AD vs. NDC AUC value	MCI vs. NDC AUC value
2	CO3	0.88	0.83
4	AP2C	0.78	0.70
6	SYN3	0.77	0.77
8	OXYR	0.81	0.77
10	ITH5L	0.79	0.70
12	HERC2	0.76	0.73
14	THRB	0.85	0.79
16	TTHY	0.73	0.69
18	TNR16	0.75	0.74
20	CO4-1	0.73	0.67
22	CO4-2	0.76	0.74
24	FIBA-1	0.77	0.64

TABLE 1-continued

Marker Peptide			
Sequence No.	Sequence name	AD vs. NDC AUC value	MCI vs. NDC AUC value
26	FIBA-2	0.74	0.61
27	FIBA-3	0.80	0.64

[0164] Table 1 shows AUC values obtained by the analysis by receiver operating characteristic (ROC) curve in the detection of cognitive impairment of each marker peptides.

[0165] Using these marker peptides in singly or in combination, using or without using liquid chromatography and/or any other suitable separation methods, directly measuring the abundance in serum using other methods such as mass spectrometry or immunological methods or enzymatic methods, on the diagnosis, it is possible to distinguish between non-psychiatry disease subjects including normal healthy subjects and subjects of cognitive impairment like AD, MCI, DLB and FTD.

INDUSTRIAL APPLICABILITY

[0166] By using the biomarkers disclosed in the present invention, mild cognitive impairment and cognitive impairment including Alzheimer disease can be detected. This invention is applicable to the field of medical diagnostics including diagnostic reagent.

SEQUENCE LIST

[0167] 10P01009 Sequence.txt

SEQUENCE LISTING

<160> NUMBER OF SEQ ID NOS: 27

<210> SEQ ID NO 1

<211> LENGTH: 1641

<212> TYPE: PRT

<213> ORGANISM: Homo sapiens

<400> SEQUENCE: 1

Ser Pro Met Tyr Ser Ile Ile Thr Pro Asn Ile Leu Arg Leu Glu Ser
1 5 10 15

Glu Glu Thr Met Val Leu Glu Ala His Asp Ala Gln Gly Asp Val Pro
20 25 30

Val Thr Val Thr Val His Asp Phe Pro Gly Lys Lys Leu Val Leu Ser
35 40 45

Ser Glu Lys Thr Val Leu Thr Pro Ala Thr Asn His Met Gly Asn Val
50 55 60

Thr Phe Thr Ile Pro Ala Asn Arg Glu Phe Lys Ser Glu Lys Gly Arg
65 70 75 80

Asn Lys Phe Val Thr Val Gln Ala Thr Phe Gly Thr Gln Val Val Glu
85 90 95

Lys Val Val Leu Val Ser Leu Gln Ser Gly Tyr Leu Phe Ile Gln Thr
100 105 110

Asp Lys Thr Ile Tyr Thr Pro Gly Ser Thr Val Leu Tyr Arg Ile Phe
115 120 125

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Thr	Val	Asn	His	Lys	Leu	Leu	Pro	Val	Gly	Arg	Thr	Val	Met	Val	Asn
130						135					140				
Ile	Glu	Asn	Pro	Glu	Gly	Ile	Pro	Val	Lys	Gln	Asp	Ser	Leu	Ser	Ser
145					150					155					160
Gln	Asn	Gln	Leu	Gly	Val	Leu	Pro	Leu	Ser	Trp	Asp	Ile	Pro	Glu	Leu
				165						170				175	
Val	Asn	Met	Gly	Gln	Trp	Lys	Ile	Arg	Ala	Tyr	Tyr	Glu	Asn	Ser	Pro
			180					185					190		
Gln	Gln	Val	Phe	Ser	Thr	Glu	Phe	Glu	Val	Lys	Glu	Tyr	Val	Leu	Pro
			195					200				205			
Ser	Phe	Glu	Val	Ile	Val	Glu	Pro	Thr	Glu	Lys	Phe	Tyr	Tyr	Ile	Tyr
	210					215					220				
Asn	Glu	Lys	Gly	Leu	Glu	Val	Thr	Ile	Thr	Ala	Arg	Phe	Leu	Tyr	Gly
225					230					235					240
Lys	Lys	Val	Glu	Gly	Thr	Ala	Phe	Val	Ile	Phe	Gly	Ile	Gln	Asp	Gly
				245						250				255	
Glu	Gln	Arg	Ile	Ser	Leu	Pro	Glu	Ser	Leu	Lys	Arg	Ile	Pro	Ile	Glu
			260					265					270		
Asp	Gly	Ser	Gly	Glu	Val	Val	Leu	Ser	Arg	Lys	Val	Leu	Leu	Asp	Gly
			275					280				285			
Val	Gln	Asn	Pro	Arg	Ala	Glu	Asp	Leu	Val	Gly	Lys	Ser	Leu	Tyr	Val
			290			295					300				
Ser	Ala	Thr	Val	Ile	Leu	His	Ser	Gly	Ser	Asp	Met	Val	Gln	Ala	Glu
305					310					315					320
Arg	Ser	Gly	Ile	Pro	Ile	Val	Thr	Ser	Pro	Tyr	Gln	Ile	His	Phe	Thr
				325					330					335	
Lys	Thr	Pro	Lys	Tyr	Phe	Lys	Pro	Gly	Met	Pro	Phe	Asp	Leu	Met	Val
			340					345					350		
Phe	Val	Thr	Asn	Pro	Asp	Gly	Ser	Pro	Ala	Tyr	Arg	Val	Pro	Val	Ala
			355				360					365			
Val	Gln	Gly	Glu	Asp	Thr	Val	Gln	Ser	Leu	Thr	Gln	Gly	Asp	Gly	Val
	370					375					380				
Ala	Lys	Leu	Ser	Ile	Asn	Thr	His	Pro	Ser	Gln	Lys	Pro	Leu	Ser	Ile
385					390					395					400
Thr	Val	Arg	Thr	Lys	Lys	Gln	Glu	Leu	Ser	Glu	Ala	Glu	Gln	Ala	Thr
				405					410					415	
Arg	Thr	Met	Gln	Ala	Leu	Pro	Tyr	Ser	Thr	Val	Gly	Asn	Ser	Asn	Asn
			420					425					430		
Tyr	Leu	His	Leu	Ser	Val	Leu	Arg	Thr	Glu	Leu	Arg	Pro	Gly	Glu	Thr
	435						440					445			
Leu	Asn	Val	Asn	Phe	Leu	Leu	Arg	Met	Asp	Arg	Ala	His	Glu	Ala	Lys
	450					455					460				
Ile	Arg	Tyr	Tyr	Thr	Tyr	Leu	Ile	Met	Asn	Lys	Gly	Arg	Leu	Leu	Lys
465					470					475					480
Ala	Gly	Arg	Gln	Val	Arg	Glu	Pro	Gly	Gln	Asp	Leu	Val	Val	Leu	Pro
				485					490					495	
Leu	Ser	Ile	Thr	Thr	Asp	Phe	Ile	Pro	Ser	Phe	Arg	Leu	Val	Ala	Tyr
			500					505					510		
Tyr	Thr	Leu	Ile	Gly	Ala	Ser	Gly	Gln	Arg	Glu	Val	Val	Ala	Asp	Ser
	515						520						525		

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Val	Trp	Val	Asp	Val	Lys	Asp	Ser	Cys	Val	Gly	Ser	Leu	Val	Val	Lys
530						535					540				
Ser	Gly	Gln	Ser	Glu	Asp	Arg	Gln	Pro	Val	Pro	Gly	Gln	Gln	Met	Thr
545					550					555					560
Leu	Lys	Ile	Glu	Gly	Asp	His	Gly	Ala	Arg	Val	Val	Leu	Val	Ala	Val
				565					570					575	
Asp	Lys	Gly	Val	Phe	Val	Leu	Asn	Lys	Lys	Asn	Lys	Leu	Thr	Gln	Ser
			580					585					590		
Lys	Ile	Trp	Asp	Val	Val	Glu	Lys	Ala	Asp	Ile	Gly	Cys	Thr	Pro	Gly
		595					600					605			
Ser	Gly	Lys	Asp	Tyr	Ala	Gly	Val	Phe	Ser	Asp	Ala	Gly	Leu	Thr	Phe
610						615					620				
Thr	Ser	Ser	Ser	Gly	Gln	Gln	Thr	Ala	Gln	Arg	Ala	Glu	Leu	Gln	Cys
625					630					635					640
Pro	Gln	Pro	Ala	Ala	Arg	Arg	Arg	Arg	Ser	Val	Gln	Leu	Thr	Glu	Lys
				645					650					655	
Arg	Met	Asp	Lys	Val	Gly	Lys	Tyr	Pro	Lys	Glu	Leu	Arg	Lys	Cys	Cys
			660					665					670		
Glu	Asp	Gly	Met	Arg	Glu	Asn	Pro	Met	Arg	Phe	Ser	Cys	Gln	Arg	Arg
		675					680					685			
Thr	Arg	Phe	Ile	Ser	Leu	Gly	Glu	Ala	Cys	Lys	Lys	Val	Phe	Leu	Asp
690						695					700				
Cys	Cys	Asn	Tyr	Ile	Thr	Glu	Leu	Arg	Arg	Gln	His	Ala	Arg	Ala	Ser
705					710					715					720
His	Leu	Gly	Leu	Ala	Arg	Ser	Asn	Leu	Asp	Glu	Asp	Ile	Ile	Ala	Glu
				725					730					735	
Glu	Asn	Ile	Val	Ser	Arg	Ser	Glu	Phe	Pro	Glu	Ser	Trp	Leu	Trp	Asn
			740					745					750		
Val	Glu	Asp	Leu	Lys	Glu	Pro	Pro	Lys	Asn	Gly	Ile	Ser	Thr	Lys	Leu
		755					760					765			
Met	Asn	Ile	Phe	Leu	Lys	Asp	Ser	Ile	Thr	Thr	Trp	Glu	Ile	Leu	Ala
770						775					780				
Val	Ser	Met	Ser	Asp	Lys	Lys	Gly	Ile	Cys	Val	Ala	Asp	Pro	Phe	Glu
785					790					795					800
Val	Thr	Val	Met	Gln	Asp	Phe	Phe	Ile	Asp	Leu	Arg	Leu	Pro	Tyr	Ser
				805					810					815	
Val	Val	Arg	Asn	Glu	Gln	Val	Glu	Ile	Arg	Ala	Val	Leu	Tyr	Asn	Tyr
			820					825					830		
Arg	Gln	Asn	Gln	Glu	Leu	Lys	Val	Arg	Val	Glu	Leu	Leu	His	Asn	Pro
		835					840					845			
Ala	Phe	Cys	Ser	Leu	Ala	Thr	Thr	Lys	Arg	Arg	His	Gln	Gln	Thr	Val
850						855					860				
Thr	Ile	Pro	Pro	Lys	Ser	Ser	Leu	Ser	Val	Pro	Tyr	Val	Ile	Val	Pro
865					870					875					880
Leu	Lys	Thr	Gly	Leu	Gln	Glu	Val	Glu	Val	Lys	Ala	Ala	Val	Tyr	His
				885					890					895	
His	Phe	Ile	Ser	Asp	Gly	Val	Arg	Lys	Ser	Leu	Lys	Val	Val	Pro	Glu
			900					905					910		
Gly	Ile	Arg	Met	Asn	Lys	Thr	Val	Ala	Val	Arg	Thr	Leu	Asp	Pro	Glu
		915					920					925			
Arg	Leu	Gly	Arg	Glu	Gly	Val	Gln	Lys	Glu	Asp	Ile	Pro	Pro	Ala	Asp

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930	935	940
Leu Ser Asp Gln Val	Pro Asp Thr Glu Ser Glu Thr Arg Ile Leu Leu	
945	950	955 960
Gln Gly Thr Pro Val Ala Gln Met Thr Glu Asp Ala Val Asp Ala Glu		
	965	970 975
Arg Leu Lys His Leu Ile Val Thr Pro Ser Gly Cys Gly Glu Gln Asn		
	980	985 990
Met Ile Gly Met Thr Pro Thr Val Ile Ala Val His Tyr Leu Asp Glu		
	995	1000 1005
Thr Glu Gln Trp Glu Lys Phe Gly Leu Glu Lys Arg Gln Gly Ala		
1010	1015	1020
Leu Glu Leu Ile Lys Lys Gly Tyr Thr Gln Gln Leu Ala Phe Arg		
1025	1030	1035
Gln Pro Ser Ser Ala Phe Ala Ala Phe Val Lys Arg Ala Pro Ser		
1040	1045	1050
Thr Trp Leu Thr Ala Tyr Val Val Lys Val Phe Ser Leu Ala Val		
1055	1060	1065
Asn Leu Ile Ala Ile Asp Ser Gln Val Leu Cys Gly Ala Val Lys		
1070	1075	1080
Trp Leu Ile Leu Glu Lys Gln Lys Pro Asp Gly Val Phe Gln Glu		
1085	1090	1095
Asp Ala Pro Val Ile His Gln Glu Met Ile Gly Gly Leu Arg Asn		
1100	1105	1110
Asn Asn Glu Lys Asp Met Ala Leu Thr Ala Phe Val Leu Ile Ser		
1115	1120	1125
Leu Gln Glu Ala Lys Asp Ile Cys Glu Glu Gln Val Asn Ser Leu		
1130	1135	1140
Pro Gly Ser Ile Thr Lys Ala Gly Asp Phe Leu Glu Ala Asn Tyr		
1145	1150	1155
Met Asn Leu Gln Arg Ser Tyr Thr Val Ala Ile Ala Gly Tyr Ala		
1160	1165	1170
Leu Ala Gln Met Gly Arg Leu Lys Gly Pro Leu Leu Asn Lys Phe		
1175	1180	1185
Leu Thr Thr Ala Lys Asp Lys Asn Arg Trp Glu Asp Pro Gly Lys		
1190	1195	1200
Gln Leu Tyr Asn Val Glu Ala Thr Ser Tyr Ala Leu Leu Ala Leu		
1205	1210	1215
Leu Gln Leu Lys Asp Phe Asp Phe Val Pro Pro Val Val Arg Trp		
1220	1225	1230
Leu Asn Glu Gln Arg Tyr Tyr Gly Gly Gly Tyr Gly Ser Thr Gln		
1235	1240	1245
Ala Thr Phe Met Val Phe Gln Ala Leu Ala Gln Tyr Gln Lys Asp		
1250	1255	1260
Ala Pro Asp His Gln Glu Leu Asn Leu Asp Val Ser Leu Gln Leu		
1265	1270	1275
Pro Ser Arg Ser Ser Lys Ile Thr His Arg Ile His Trp Glu Ser		
1280	1285	1290
Ala Ser Leu Leu Arg Ser Glu Glu Thr Lys Glu Asn Glu Gly Phe		
1295	1300	1305
Thr Val Thr Ala Glu Gly Lys Gly Gln Gly Thr Leu Ser Val Val		
1310	1315	1320

-continued

Thr Met	Tyr His Ala Lys Ala	Lys Asp Gln Leu Thr	Cys Asn Lys
1325	1330	1335	
Phe Asp	Leu Lys Val Thr Ile	Lys Pro Ala Pro Glu	Thr Glu Lys
1340	1345	1350	
Arg Pro	Gln Asp Ala Lys Asn	Thr Met Ile Leu Glu	Ile Cys Thr
1355	1360	1365	
Arg Tyr	Arg Gly Asp Gln Asp	Ala Thr Met Ser Ile	Leu Asp Ile
1370	1375	1380	
Ser Met	Met Thr Gly Phe Ala	Pro Asp Thr Asp Asp	Leu Lys Gln
1385	1390	1395	
Leu Ala	Asn Gly Val Asp Arg	Tyr Ile Ser Lys Tyr	Glu Leu Asp
1400	1405	1410	
Lys Ala	Phe Ser Asp Arg Asn	Thr Leu Ile Ile Tyr	Leu Asp Lys
1415	1420	1425	
Val Ser	His Ser Glu Asp Asp	Cys Leu Ala Phe Lys	Val His Gln
1430	1435	1440	
Tyr Phe	Asn Val Glu Leu Ile	Gln Pro Gly Ala Val	Lys Val Tyr
1445	1450	1455	
Ala Tyr	Tyr Asn Leu Glu Glu	Ser Cys Thr Arg Phe	Tyr His Pro
1460	1465	1470	
Glu Lys	Glu Asp Gly Lys Leu	Asn Lys Leu Cys Arg	Asp Glu Leu
1475	1480	1485	
Cys Arg	Cys Ala Glu Glu Asn	Cys Phe Ile Gln Lys	Ser Asp Asp
1490	1495	1500	
Lys Val	Thr Leu Glu Glu Arg	Leu Asp Lys Ala Cys	Glu Pro Gly
1505	1510	1515	
Val Asp	Tyr Val Tyr Lys Thr	Arg Leu Val Lys Val	Gln Leu Ser
1520	1525	1530	
Asn Asp	Phe Asp Glu Tyr Ile	Met Ala Ile Glu Gln	Thr Ile Lys
1535	1540	1545	
Ser Gly	Ser Asp Glu Val Gln	Val Gly Gln Gln Arg	Thr Phe Ile
1550	1555	1560	
Ser Pro	Ile Lys Cys Arg Glu	Ala Leu Lys Leu Glu	Glu Lys Lys
1565	1570	1575	
His Tyr	Leu Met Trp Gly Leu	Ser Ser Asp Phe Trp	Gly Glu Lys
1580	1585	1590	
Pro Asn	Leu Ser Tyr Ile Ile	Gly Lys Asp Thr Trp	Val Glu His
1595	1600	1605	
Trp Pro	Glu Glu Asp Glu Cys	Gln Asp Glu Glu Asn	Gln Lys Gln
1610	1615	1620	
Cys Gln	Asp Leu Gly Ala Phe	Thr Glu Ser Met Val	Val Phe Gly
1625	1630	1635	
Cys Pro	Asn		
1640			

<210> SEQ ID NO 2

<211> LENGTH: 14

<212> TYPE: PRT

<213> ORGANISM: Homo sapiens

<400> SEQUENCE: 2

Ala Pro Val Ile His Gln Glu Met Ile Gly Gly Leu Arg Asn

1	5								10							
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<211> LENGTH: 450																
<212> TYPE: PRT																
<213> ORGANISM: Homo sapiens																
<400> SEQUENCE: 3																
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Asp	Arg	His	Asp 20	Gly	Ser	Ser	Asn	Gly 25	Asn	Pro	Arg	Val	Pro 30	His	Leu	
Ser	Ser	Ala 35	Gly	Gln	His	Leu	Tyr 40	Ser	Pro	Ala	Pro 45	Leu	Ser	His		
Thr	Gly 50	Val	Ala	Glu	Tyr 55	Gln	Pro	Pro	Pro	Tyr	Phe 60	Pro	Pro	Pro	Tyr	
Gln 65	Gln	Leu	Ala	Tyr 70	Ser	Gln	Ser	Ala	Asp 75	Pro	Tyr	Ser	His	Leu	Gly 80	
Glu	Ala	Tyr	Ala 85	Ala	Ile	Asn	Pro	Leu 90	His	Gln	Pro	Ala	Pro 95	Thr		
Gly	Ser	Gln	Gln 100	Gln	Ala	Trp	Pro	Gly 105	Arg	Gln	Ser	Gln 110	Glu	Gly	Ala	
Gly	Leu 115	Pro	Ser	His	His	Gly	Arg 120	Pro	Ala	Gly	Leu 125	Pro	His	Leu		
Ser	Gly 130	Leu	Glu	Ala	Gly	Ala 135	Val	Ser	Ala	Arg	Arg 140	Asp	Ala	Tyr	Arg	
Arg 145	Ser	Asp	Leu	Leu 150	Leu	Pro	His	Ala	His 155	Ala	Leu	Asp	Ala	Ala	Gly 160	
Leu	Ala	Glu	Asn 165	Leu	Gly	Leu	His	Asp	Met 170	Pro	His	Gln	Met	Asp 175	Glu	
Val	Gln	Asn 180	Val	Asp	Asp	Gln	His	Leu 185	Leu	Leu	His	Asp 190	Gln	Thr	Val	
Ile	Arg	Lys 195	Gly	Pro	Ile	Ser	Met 200	Thr	Lys	Asn	Pro 205	Leu	Asn	Leu	Pro	
Cys 210	Gln	Lys	Glu	Leu	Val	Gly 215	Ala	Val	Met	Asn	Pro 220	Thr	Glu	Val	Phe	
Cys 225	Ser	Val	Pro	Gly	Arg 230	Leu	Ser	Leu	Leu	Ser 235	Ser	Thr	Ser	Lys	Tyr 240	
Lys	Val	Thr	Val	Ala 245	Glu	Val	Gln	Arg	Arg 250	Leu	Ser	Pro	Pro	Glu 255	Cys	
Leu	Asn	Ala 260	Ser	Leu	Leu	Gly	Gly	Val 265	Leu	Arg	Arg	Ala	Lys 270	Ser	Lys	
Asn	Gly	Gly 275	Arg	Ser	Leu	Arg	Glu 280	Lys	Leu	Asp	Lys	Ile 285	Gly	Leu	Asn	
Leu 290	Pro	Ala	Gly	Arg	Arg	Lys 295	Ala	Ala	His	Val	Thr 300	Leu	Leu	Thr	Ser	
Leu 305	Val	Glu	Gly	Glu	Ala 310	Val	His	Leu	Ala	Arg 315	Asp	Phe	Ala	Tyr	Val 320	
Cys	Glu	Ala	Glu 325	Phe	Pro	Ser	Lys	Pro	Val 330	Ala	Glu	Tyr	Leu	Thr	Arg	
Pro	His	Leu 340	Gly	Gly	Arg	Asn	Glu	Met 345	Ala	Ala	Arg	Lys	Asn 350	Met	Leu	

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Leu Ala Ala Gln Gln Leu Cys Lys Glu Phe Thr Glu Leu Leu Ser Gln
   355                               360               365

Asp Arg Thr Pro His Gly Thr Ser Arg Leu Ala Pro Val Leu Glu Thr
   370                               375               380

Asn Ile Gln Asn Cys Leu Ser His Phe Ser Leu Ile Thr His Gly Phe
   385                               390               395               400

Gly Ser Gln Ala Ile Cys Ala Ala Val Ser Ala Leu Gln Asn Tyr Ile
   405                               410               415

Lys Glu Ala Leu Ile Val Ile Asp Lys Ser Tyr Met Asn Pro Gly Asp
   420                               425               430

Gln Ser Pro Ala Asp Ser Asn Lys Thr Leu Glu Lys Met Glu Lys His
   435                               440               445

Arg Lys
   450

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<210> SEQ ID NO 4
<211> LENGTH: 16
<212> TYPE: PRT
<213> ORGANISM: Homo sapiens

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<400> SEQUENCE: 4

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Pro Gly Arg Gln Ser Gln Glu Gly Ala Gly Leu Pro Ser His His Gly
1           5           10           15

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<210> SEQ ID NO 5
<211> LENGTH: 580
<212> TYPE: PRT
<213> ORGANISM: Homo sapiens

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<400> SEQUENCE: 5

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Met Asn Phe Leu Arg Arg Arg Leu Ser Asp Ser Ser Phe Met Ala Asn
1           5           10           15

Leu Pro Asn Gly Tyr Met Thr Asp Leu Gln Arg Pro Asp Ser Ser Thr
   20           25           30

Ser Ser Pro Ala Ser Pro Ala Met Glu Arg Arg His Pro Gln Pro Leu
   35           40           45

Ala Ala Ser Phe Ser Ser Pro Gly Ser Ser Leu Phe Ser Ser Leu Ser
   50           55           60

Ser Ala Met Lys Gln Ala Pro Gln Ala Thr Ser Gly Leu Met Glu Pro
   65           70           75           80

Pro Gly Pro Ser Thr Pro Ile Val Gln Arg Pro Arg Ile Leu Leu Val
   85           90           95

Ile Asp Asp Ala His Thr Asp Trp Ser Lys Tyr Phe His Gly Lys Lys
  100          105          110

Val Asn Gly Glu Ile Glu Ile Arg Val Glu Gln Ala Glu Phe Ser Glu
  115          120          125

Leu Asn Leu Ala Ala Tyr Val Thr Gly Gly Cys Met Val Asp Met Gln
  130          135          140

Val Val Arg Asn Gly Thr Lys Val Val Ser Arg Ser Phe Lys Pro Asp
  145          150          155          160

Phe Ile Leu Val Arg Gln His Ala Tyr Ser Met Ala Leu Gly Glu Asp
  165          170          175

Tyr Arg Ser Leu Val Ile Gly Leu Gln Tyr Gly Gly Leu Pro Ala Val
  180          185          190

```

Asn	Ser	Leu	Tyr	Ser	Val	Tyr	Asn	Phe	Cys	Ser	Lys	Pro	Trp	Val	Phe
		195				200						205			
Ser	Gln	Leu	Ile	Lys	Ile	Phe	His	Ser	Leu	Gly	Pro	Glu	Lys	Phe	Pro
210					215						220				
Leu	Val	Glu	Gln	Thr	Phe	Phe	Pro	Asn	His	Lys	Pro	Met	Val	Thr	Ala
225					230				235		240				
Pro	His	Phe	Pro	Val	Val	Val	Lys	Leu	Gly	His	Ala	His	Ala	Gly	Met
				245					250				255		
Gly	Lys	Ile	Lys	Val	Glu	Asn	Gln	Leu	Asp	Phe	Gln	Asp	Ile	Thr	Ser
		260						265				270			
Val	Val	Ala	Met	Ala	Lys	Thr	Tyr	Ala	Thr	Thr	Glu	Ala	Phe	Ile	Asp
		275				280						285			
Ser	Lys	Tyr	Asp	Ile	Arg	Ile	Gln	Lys	Ile	Gly	Ser	Asn	Tyr	Lys	Ala
290					295						300				
Tyr	Met	Arg	Thr	Ser	Ile	Ser	Gly	Asn	Trp	Lys	Ala	Asn	Thr	Gly	Ser
305					310				315		320				
Ala	Met	Leu	Glu	Gln	Val	Ala	Met	Thr	Glu	Arg	Tyr	Arg	Leu	Trp	Val
				325					330				335		
Asp	Ser	Cys	Ser	Glu	Met	Phe	Gly	Gly	Leu	Asp	Ile	Cys	Ala	Val	Lys
		340						345				350			
Ala	Val	His	Ser	Lys	Asp	Gly	Arg	Asp	Tyr	Ile	Ile	Glu	Val	Met	Asp
		355				360						365			
Ser	Ser	Met	Pro	Leu	Ile	Gly	Glu	His	Val	Glu	Glu	Asp	Arg	Gln	Leu
370					375						380				
Met	Ala	Asp	Leu	Val	Val	Ser	Lys	Met	Ser	Gln	Leu	Pro	Met	Pro	Gly
385					390				395		400				
Gly	Thr	Ala	Pro	Ser	Pro	Leu	Arg	Pro	Trp	Ala	Pro	Gln	Ile	Lys	Ser
				405					410				415		
Ala	Lys	Ser	Pro	Gly	Gln	Ala	Gln	Leu	Gly	Pro	Gln	Leu	Gly	Gln	Pro
		420						425				430			
Gln	Pro	Arg	Pro	Pro	Pro	Gln	Gly	Gly	Pro	Arg	Gln	Ala	Gln	Ser	Pro
		435				440						445			
Gln	Pro	Gln	Arg	Ser	Gly	Ser	Pro	Ser	Gln	Gln	Arg	Leu	Ser	Pro	Gln
450					455						460				
Gly	Gln	Gln	Pro	Leu	Ser	Pro	Gln	Ser	Gly	Ser	Pro	Gln	Gln	Gln	Arg
465					470				475		480				
Ser	Pro	Gly	Ser	Pro	Gln	Leu	Ser	Arg	Ala	Ser	Ser	Gly	Ser	Ser	Pro
		485						490				495			
Asn	Gln	Ala	Ser	Lys	Pro	Gly	Ala	Thr	Leu	Ala	Ser	Gln	Pro	Arg	Pro
		500						505				510			
Pro	Val	Gln	Gly	Arg	Ser	Thr	Ser	Gln	Gln	Gly	Glu	Glu	Ser	Lys	Lys
		515				520						525			
Pro	Ala	Pro	Pro	His	Pro	His	Leu	Asn	Lys	Ser	Gln	Ser	Leu	Thr	Asn
530					535						540				
Ser	Leu	Ser	Thr	Ser	Asp	Thr	Ser	Gln	Arg	Gly	Thr	Pro	Ser	Glu	Asp
545					550				555		560				
Glu	Ala	Lys	Ala	Glu	Thr	Ile	Arg	Asn	Leu	Arg	Lys	Ser	Phe	Ala	Ser

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<210> SEQ ID NO 6
<211> LENGTH: 17
<212> TYPE: PRT
<213> ORGANISM: Homo sapiens

<400> SEQUENCE: 6

Glu Met Phe Gly Gly Leu Asp Ile Cys Ala Val Lys Ala Val His Ser
1 5 10 15

Lys

<210> SEQ ID NO 7
<211> LENGTH: 389
<212> TYPE: PRT
<213> ORGANISM: Homo sapiens

<400> SEQUENCE: 7

Met Glu Gly Ala Leu Ala Ala Asn Trp Ser Ala Glu Ala Ala Asn Ala
1 5 10 15

Ser Ala Ala Pro Pro Gly Ala Glu Gly Asn Arg Thr Ala Gly Pro Pro
20 25 30

Arg Arg Asn Glu Ala Leu Ala Arg Val Glu Val Ala Val Leu Cys Leu
35 40 45

Ile Leu Leu Leu Ala Leu Ser Gly Asn Ala Cys Val Leu Leu Ala Leu
50 55 60

Arg Thr Thr Arg Gln Lys His Ser Arg Leu Phe Phe Phe Met Lys His
65 70 75 80

Leu Ser Ile Ala Asp Leu Val Val Ala Val Phe Gln Val Leu Pro Gln
85 90 95

Leu Leu Trp Asp Ile Thr Phe Arg Phe Tyr Gly Pro Asp Leu Leu Cys
100 105 110

Arg Leu Val Lys Tyr Leu Gln Val Val Gly Met Phe Ala Ser Thr Tyr
115 120 125

Leu Leu Leu Leu Met Ser Leu Asp Arg Cys Leu Ala Ile Cys Gln Pro
130 135 140

Leu Arg Ser Leu Arg Arg Arg Thr Asp Arg Leu Ala Val Leu Ala Thr
145 150 155 160

Trp Leu Gly Cys Leu Val Ala Ser Ala Pro Gln Val His Ile Phe Ser
165 170 175

Leu Arg Glu Val Ala Asp Gly Val Phe Asp Cys Trp Ala Val Phe Ile
180 185 190

Gln Pro Trp Gly Pro Lys Ala Tyr Ile Thr Trp Ile Thr Leu Ala Val
195 200 205

Tyr Ile Val Pro Val Ile Val Leu Ala Ala Cys Tyr Gly Leu Ile Ser
210 215 220

Phe Lys Ile Trp Gln Asn Leu Arg Leu Lys Thr Ala Ala Ala Ala Ala
225 230 235 240

Ala Glu Ala Pro Glu Gly Ala Ala Ala Gly Asp Gly Gly Arg Val Ala
245 250 255

Leu Ala Arg Val Ser Ser Val Lys Leu Ile Ser Lys Ala Lys Ile Arg
260 265 270

Thr Val Lys Met Thr Phe Ile Ile Val Leu Ala Phe Ile Val Cys Trp
275 280 285

Thr Pro Phe Phe Phe Val Gln Met Trp Ser Val Trp Asp Ala Asn Ala
290 295 300

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Pro Lys Glu Ala Ser Ala Phe Ile Ile Val Met Leu Leu Ala Ser Leu
 305 310 315 320

Asn Ser Cys Cys Asn Pro Trp Ile Tyr Met Leu Phe Thr Gly His Leu
 325 330 335

Phe His Glu Leu Val Gln Arg Phe Leu Cys Cys Ser Ala Ser Tyr Leu
 340 345 350

Lys Gly Arg Arg Leu Gly Glu Thr Ser Ala Ser Lys Lys Ser Asn Ser
 355 360 365

Ser Ser Phe Val Leu Ser His Arg Ser Ser Ser Gln Arg Ser Cys Ser
 370 375 380

Gln Pro Ser Thr Ala
 385

<210> SEQ ID NO 8
 <211> LENGTH: 11
 <212> TYPE: PRT
 <213> ORGANISM: Homo sapiens

<400> SEQUENCE: 8

Ala Ala Pro Pro Gly Ala Glu Gly Asn Arg Thr
 1 5 10

<210> SEQ ID NO 9
 <211> LENGTH: 1290
 <212> TYPE: PRT
 <213> ORGANISM: Homo sapiens

<400> SEQUENCE: 9

Gly Pro Pro Val Pro Ala Ser Ser Ser Thr Lys Leu Leu Met Thr Ser
 1 5 10 15

Tyr Ser Met Arg Ser Thr Val Val Ser Arg Tyr Ala His Thr Leu Val
 20 25 30

Thr Ser Val Leu Phe Asn Pro His Ala Glu Ala His Glu Ala Ile Phe
 35 40 45

Asp Leu Asp Leu Pro His Leu Ala Phe Ile Ser Asn Phe Thr Met Thr
 50 55 60

Ile Asn Asn Lys Val Tyr Ile Ala Glu Val Lys Glu Lys His Gln Ala
 65 70 75 80

Lys Lys Ile Tyr Glu Glu Ala His Gln Gln Gly Lys Thr Ala Ala His
 85 90 95

Val Gly Ile Arg Asp Arg Glu Ser Glu Lys Phe Arg Ile Ser Thr Ser
 100 105 110

Leu Ala Ala Gly Thr Glu Val Thr Phe Ser Leu Ala Tyr Glu Glu Leu
 115 120 125

Leu Gln Arg His Gln Gly Gln Tyr Gln Leu Val Val Ser Leu Arg Pro
 130 135 140

Gly Gln Leu Val Lys Arg Leu Ser Ile Glu Val Thr Val Ser Glu Arg
 145 150 155 160

Thr Gly Ile Ser Tyr Val His Ile Pro Pro Leu Arg Thr Gly Arg Leu
 165 170 175

Arg Thr Asn Ala His Ala Ser Glu Val Asp Ser Pro Pro Ser Thr Arg
 180 185 190

Ile Glu Arg Gly Glu Thr Cys Val Arg Ile Thr Tyr Cys Pro Thr Leu
 195 200 205

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Gln	Asp	Gln	Ser	Ser	Ile	Ser	Gly	Ser	Gly	Ile	Met	Ala	Asp	Phe	Leu
210						215					220				
Val	Gln	Tyr	Asp	Val	Val	Met	Glu	Asp	Ile	Ile	Gly	Asp	Val	Gln	Ile
225					230					235					240
Tyr	Asp	Asp	Tyr	Phe	Ile	His	Tyr	Phe	Ala	Pro	Arg	Gly	Leu	Pro	Pro
				245					250					255	
Met	Glu	Lys	Asn	Val	Val	Phe	Val	Ile	Asp	Val	Ser	Ser	Ser	Met	Phe
			260					265					270		
Gly	Thr	Lys	Met	Glu	Gln	Thr	Lys	Thr	Ala	Met	Asn	Val	Ile	Leu	Ser
		275					280					285			
Asp	Leu	Gln	Ala	Asn	Asp	Tyr	Phe	Asn	Ile	Ile	Ser	Phe	Ser	Asp	Thr
290						295					300				
Val	Asn	Val	Trp	Lys	Ala	Gly	Gly	Ser	Ile	Gln	Ala	Thr	Ile	Gln	Asn
305					310					315					320
Val	His	Ser	Ala	Lys	Asp	Tyr	Leu	His	Cys	Met	Glu	Ala	Asp	Gly	Trp
				325					330					335	
Thr	Asp	Val	Asn	Ser	Ala	Leu	Leu	Ala	Ala	Ala	Ser	Val	Leu	Asn	His
			340					345					350		
Ser	Asn	Gln	Glu	Pro	Gly	Arg	Gly	Pro	Ser	Val	Gly	Arg	Ile	Pro	Leu
		355					360					365			
Ile	Ile	Phe	Leu	Thr	Asp	Gly	Glu	Pro	Thr	Ala	Gly	Val	Thr	Thr	Pro
370					375						380				
Ser	Val	Ile	Leu	Ser	Asn	Val	Arg	Gln	Ala	Leu	Gly	His	Arg	Val	Ser
385					390					395					400
Leu	Phe	Ser	Leu	Ala	Phe	Gly	Asp	Asp	Ala	Asp	Phe	Thr	Leu	Leu	Arg
				405					410					415	
Arg	Leu	Ser	Leu	Glu	Asn	Arg	Gly	Ile	Ala	Arg	Arg	Ile	Tyr	Glu	Asp
			420					425					430		
Thr	Asp	Ala	Ala	Leu	Gln	Leu	Lys	Gly	Leu	Tyr	Glu	Glu	Ile	Ser	Met
		435					440					445			
Pro	Leu	Leu	Ala	Asp	Val	Arg	Leu	Asn	Tyr	Leu	Gly	Gly	Leu	Val	Gly
450						455					460				
Ala	Ser	Pro	Trp	Ala	Val	Phe	Pro	Asn	Tyr	Phe	Gly	Gly	Ser	Glu	Leu
465					470					475					480
Val	Val	Ala	Gly	Gln	Val	Gln	Pro	Gly	Lys	Gln	Glu	Leu	Gly	Ile	His
				485					490					495	
Leu	Ala	Ala	Arg	Gly	Pro	Lys	Asp	Gln	Leu	Leu	Val	Ala	His	His	Ser
			500					505					510		
Glu	Gly	Ala	Thr	Asn	Asn	Ser	Gln	Lys	Ala	Phe	Gly	Cys	Pro	Gly	Glu
		515					520					525			
Pro	Ala	Pro	Asn	Val	Ala	His	Phe	Ile	Arg	Arg	Leu	Trp	Ala	Tyr	Val
530						535					540				
Thr	Ile	Gly	Glu	Leu	Leu	Asp	Ala	His	Phe	Gln	Ala	Arg	Asp	Thr	Thr
545					550					555					560
Thr	Arg	His	Leu	Leu	Ala	Ala	Lys	Val	Leu	Asn	Leu	Ser	Leu	Glu	Tyr
				565					570					575	
Asn	Phe	Val	Thr	Pro	Leu	Thr	Ser	Leu	Val	Met	Val	Gln	Pro	Lys	Gln
			580					585					590		
Ala	Ser	Glu	Glu	Thr	Arg	Arg	Gln	Thr	Ser	Thr	Ser	Ala	Gly	Pro	Asp
595							600						605		

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Thr	Ile	Met	Pro	Ser	Ser	Ser	Ser	Arg	His	Gly	Leu	Gly	Val	Ser	Thr	610	615	620
Ala	Gln	Pro	Ala	Leu	Val	Pro	Lys	Val	Ile	Ser	Pro	Lys	Ser	Arg	Pro	625	630	635
Val	Lys	Pro	Lys	Phe	Tyr	Leu	Ser	Ser	Thr	Thr	Thr	Ala	Ser	Thr	Lys	645	650	655
Lys	Met	Leu	Ser	Ser	Lys	Glu	Leu	Glu	Pro	Leu	Gly	Glu	Ser	Pro	His	660	665	670
Thr	Leu	Ser	Met	Pro	Thr	Tyr	Pro	Lys	Ala	Lys	Ile	Pro	Ala	Gln	Gln	675	680	685
Asp	Ser	Gly	Thr	Leu	Ala	Gln	Pro	Thr	Leu	Arg	Thr	Lys	Pro	Thr	Ile	690	695	700
Leu	Val	Pro	Ser	Asn	Ser	Gly	Thr	Leu	Leu	Pro	Leu	Lys	Pro	Gly	Ser	705	710	715
Leu	Ser	His	Gln	Asn	Pro	Asp	Ile	Leu	Pro	Thr	Asn	Ser	Arg	Thr	Gln	725	730	735
Val	Pro	Pro	Val	Lys	Pro	Gly	Ile	Pro	Ala	Ser	Pro	Lys	Ala	Asp	Thr	740	745	750
Val	Lys	Cys	Val	Thr	Pro	Leu	His	Ser	Lys	Pro	Gly	Ala	Pro	Ser	His	755	760	765
Pro	Gln	Leu	Gly	Ala	Leu	Thr	Ser	Gln	Ala	Pro	Lys	Gly	Leu	Pro	Gln	770	775	780
Ser	Arg	Pro	Gly	Val	Ser	Thr	Leu	Gln	Val	Pro	Lys	Tyr	Pro	Leu	His	785	790	795
Thr	Arg	Pro	Arg	Val	Pro	Ala	Pro	Lys	Thr	Arg	Asn	Asn	Met	Pro	His	805	810	815
Leu	Gly	Pro	Gly	Ile	Leu	Leu	Ser	Lys	Thr	Pro	Lys	Ile	Leu	Leu	Ser	820	825	830
Leu	Lys	Pro	Ser	Ala	Pro	Pro	His	Gln	Ile	Ser	Thr	Ser	Ile	Ser	Leu	835	840	845
Ser	Lys	Pro	Glu	Thr	Pro	Asn	Pro	His	Met	Pro	Gln	Thr	Pro	Leu	Pro	850	855	860
Pro	Arg	Pro	Asp	Arg	Pro	Arg	Pro	Pro	Leu	Pro	Glu	Ser	Leu	Ser	Thr	865	870	875
Phe	Pro	Asn	Thr	Ile	Ser	Ser	Ser	Thr	Gly	Pro	Ser	Ser	Thr	Thr	Thr	885	890	895
Thr	Ser	Val	Leu	Gly	Glu	Pro	Leu	Pro	Met	Pro	Phe	Thr	Pro	Thr	Leu	900	905	910
Pro	Pro	Gly	Arg	Phe	Trp	His	Gln	Tyr	Asp	Leu	Leu	Pro	Gly	Pro	Gln	915	920	925
Arg	Thr	Arg	Gln	Val	Leu	Gly	Pro	Ser	Arg	Pro	Gly	Val	Pro	Thr	Met	930	935	940
Ser	Leu	Leu	Asn	Ser	Ser	Arg	Pro	Thr	Pro	Glu	Gly	Ser	Pro	Pro	Asn	945	950	955
Leu	Pro	Ile	Leu	Leu	Pro	Ser	Ser	Ile	Leu	Pro	Glu	Ala	Ile	Ser	Leu	965	970	975
Leu	Leu	Leu	Pro	Glu	Glu	Leu	Glu	Leu	Ser	Glu	Ser	Met	Val	Glu		980	985	990
Ser	Lys	Phe	Val	Glu	Ser	Leu	Asn	Pro	Pro	Ala	Phe	Tyr	Thr	Phe	Leu	995	1000	1005
Thr	Pro	Asp	Glu	Asp	Gly	Ser	Pro	Asn	Trp	Asp	Gly	Asn	Ser	Glu				

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1010	1015	1020
Glu Ile Leu Gly Gly Ala Gly Gly Ser Met Glu Ser Gln Gly Ser		
1025	1030	1035
Ser Val Gly Leu Ala Lys Gly Thr Leu Pro Ser Ile Phe Thr Phe		
1040	1045	1050
Ser Ser Ser Val Asp Gly Asp Pro His Phe Val Ile Gln Ile Pro		
1055	1060	1065
His Ser Glu Glu Lys Ile Cys Phe Thr Leu Asn Gly His Pro Gly		
1070	1075	1080
Asp Leu Leu Gln Leu Ile Glu Asp Pro Lys Ala Gly Leu His Val		
1085	1090	1095
Ser Gly Lys Leu Leu Gly Ala Pro Pro Arg Pro Gly His Lys Asp		
1100	1105	1110
Gln Thr Arg Thr Tyr Phe Gln Ile Ile Thr Val Thr Thr Asp Lys		
1115	1120	1125
Pro Arg Ala Tyr Thr Ile Thr Ile Ser Arg Ser Ser Ile Ser Leu		
1130	1135	1140
Arg Gly Glu Gly Thr Leu Arg Leu Ser Trp Asp Gln Pro Ala Leu		
1145	1150	1155
Leu Lys Arg Pro Gln Leu Glu Leu Tyr Val Ala Ala Ala Ala Arg		
1160	1165	1170
Leu Thr Leu Arg Leu Gly Pro Tyr Leu Glu Phe Leu Val Leu Arg		
1175	1180	1185
His Arg Tyr Arg His Pro Ser Thr Leu Gln Leu Pro His Leu Gly		
1190	1195	1200
Phe Tyr Val Ala Asn Gly Ser Gly Leu Ser Pro Ser Ala Arg Gly		
1205	1210	1215
Leu Ile Gly Gln Phe Gln His Ala Asp Ile Arg Leu Val Thr Gly		
1220	1225	1230
Pro Met Gly Pro Cys Leu Arg Arg His His Gly Pro Asp Val Pro		
1235	1240	1245
Val Ile Leu Gly Lys Arg Leu Leu Lys Asp Ser Pro Arg Leu Leu		
1250	1255	1260
Pro Arg Trp Ala Ser Cys Trp Leu Val Lys Arg Ser His Val Glu		
1265	1270	1275
Leu Leu Leu Gly His Pro Tyr Leu Ser Tyr Val Leu		
1280	1285	1290

<210> SEQ ID NO 10
 <211> LENGTH: 14
 <212> TYPE: PRT
 <213> ORGANISM: Homo sapiens

<400> SEQUENCE: 10

Arg Val Ser Leu Phe Ser Leu Ala Phe Gly Asp Asp Ala Asp
 1 5 10

<210> SEQ ID NO 11
 <211> LENGTH: 4834
 <212> TYPE: PRT
 <213> ORGANISM: Homo sapiens

<400> SEQUENCE: 11

Met Pro Ser Glu Ser Phe Cys Leu Ala Ala Gln Ala Arg Leu Asp Ser

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1	5	10	15
Lys Trp Leu Lys Thr Asp Ile Gln Leu Ala Phe Thr Arg Asp Gly Leu	20	25	30
Cys Gly Leu Trp Asn Glu Met Val Lys Asp Gly Glu Ile Val Tyr Thr	35	40	45
Gly Thr Glu Ser Thr Gln Asn Gly Glu Leu Pro Pro Arg Lys Asp Asp	50	55	60
Ser Val Glu Pro Ser Gly Thr Lys Lys Glu Asp Leu Asn Asp Lys Glu	65	70	75
Lys Lys Asp Glu Glu Glu Thr Pro Ala Pro Ile Tyr Arg Ala Lys Ser	85	90	95
Ile Leu Asp Ser Trp Val Trp Gly Lys Gln Pro Asp Val Asn Glu Leu	100	105	110
Lys Glu Cys Leu Ser Val Leu Val Lys Glu Gln Gln Ala Leu Ala Val	115	120	125
Gln Ser Ala Thr Thr Thr Leu Ser Ala Leu Arg Leu Lys Gln Arg Leu	130	135	140
Val Ile Leu Glu Arg Tyr Phe Ile Ala Leu Asn Arg Thr Val Phe Gln	145	150	155
Glu Asn Val Lys Val Lys Trp Lys Ser Ser Gly Ile Ser Leu Pro Pro	165	170	175
Val Asp Lys Lys Ser Ser Arg Pro Ala Gly Lys Gly Val Glu Gly Leu	180	185	190
Ala Arg Val Gly Ser Arg Ala Ala Leu Ser Phe Ala Phe Ala Phe Leu	195	200	205
Arg Arg Ala Trp Arg Ser Gly Glu Asp Ala Asp Leu Cys Ser Glu Leu	210	215	220
Leu Gln Glu Ser Leu Asp Ala Leu Arg Ala Leu Pro Glu Ala Ser Leu	225	230	235
Phe Asp Glu Ser Thr Val Ser Ser Val Trp Leu Glu Val Val Glu Arg	245	250	255
Ala Thr Arg Phe Leu Arg Ser Val Val Thr Gly Asp Val His Gly Thr	260	265	270
Pro Ala Thr Lys Gly Pro Gly Ser Ile Pro Leu Gln Asp Gln His Leu	275	280	285
Ala Leu Ala Ile Leu Leu Glu Leu Ala Val Gln Arg Gly Thr Leu Ser	290	295	300
Gln Met Leu Ser Ala Ile Leu Leu Leu Leu Leu Trp Asp Ser Gly	305	310	315
Ala Gln Glu Thr Asp Asn Glu Arg Ser Ala Gln Gly Thr Ser Ala Pro	325	330	335
Leu Leu Pro Leu Leu Gln Arg Phe Gln Ser Ile Ile Cys Arg Lys Asp	340	345	350
Ala Pro His Ser Glu Gly Asp Met His Leu Leu Ser Gly Pro Leu Ser	355	360	365
Pro Asn Glu Ser Phe Leu Arg Tyr Leu Thr Leu Pro Gln Asp Asn Glu	370	375	380
Leu Ala Ile Asp Leu Arg Gln Thr Ala Val Val Val Met Ala His Leu	385	390	395
Asp Arg Leu Ala Thr Pro Cys Met Pro Pro Leu Cys Ser Ser Pro Thr	405	410	415

Ser	His	Lys	Gly	Ser	Leu	Gln	Glu	Val	Ile	Gly	Trp	Gly	Leu	Ile	Gly
			420				425								
Trp	Lys	Tyr	Tyr	Ala	Asn	Val	Ile	Gly	Pro	Ile	Gln	Cys	Glu	Gly	Leu
			435				440								
Ala	Asn	Leu	Gly	Val	Thr	Gln	Ile	Ala	Cys	Ala	Glu	Lys	Arg	Phe	Leu
			450				455								
Ile	Leu	Ser	Arg	Asn	Gly	Arg	Val	Tyr	Thr	Gln	Ala	Tyr	Asn	Ser	Asp
			465							475			480		
Thr	Leu	Ala	Pro	Gln	Leu	Val	Gln	Gly	Leu	Ala	Ser	Arg	Asn	Ile	Val
			485						490			495			
Lys	Ile	Ala	Ala	His	Ser	Asp	Gly	His	His	Tyr	Leu	Ala	Leu	Ala	Ala
			500						505			510			
Thr	Gly	Glu	Val	Tyr	Ser	Trp	Gly	Cys	Gly	Asp	Gly	Gly	Arg	Leu	Gly
			515						520			525			
His	Gly	Asp	Thr	Val	Pro	Leu	Glu	Glu	Pro	Lys	Val	Ile	Ser	Ala	Phe
			530						535			540			
Ser	Gly	Lys	Gln	Ala	Gly	Lys	His	Val	Val	His	Ile	Ala	Cys	Gly	Ser
			545						555			560			
Thr	Tyr	Ser	Ala	Ala	Ile	Thr	Ala	Glu	Gly	Glu	Leu	Tyr	Thr	Trp	Gly
			565						570			575			
Arg	Gly	Asn	Tyr	Gly	Arg	Leu	Gly	His	Gly	Ser	Ser	Glu	Asp	Glu	Ala
			580						585			590			
Ile	Pro	Met	Leu	Val	Ala	Gly	Leu	Lys	Gly	Leu	Lys	Val	Ile	Asp	Val
			595						600			605			
Ala	Cys	Gly	Ser	Gly	Asp	Ala	Gln	Thr	Leu	Ala	Val	Thr	Glu	Asn	Gly
			610						620						
Gln	Val	Trp	Ser	Trp	Gly	Asp	Gly	Asp	Tyr	Gly	Lys	Leu	Gly	Arg	Gly
			625						630			635			
Gly	Ser	Asp	Gly	Cys	Lys	Thr	Pro	Lys	Leu	Ile	Glu	Lys	Leu	Gln	Asp
			645						650			655			
Leu	Asp	Val	Val	Lys	Val	Arg	Cys	Gly	Ser	Gln	Phe	Ser	Ile	Ala	Leu
			660						665			670			
Thr	Lys	Asp	Gly	Gln	Val	Tyr	Ser	Trp	Gly	Lys	Gly	Asp	Asn	Gln	Arg
			675						680			685			
Leu	Gly	His	Gly	Thr	Glu	Glu	His	Val	Arg	Tyr	Pro	Lys	Leu	Leu	Glu
			690						695			700			
Gly	Leu	Gln	Gly	Lys	Lys	Val	Ile	Asp	Val	Ala	Ala	Gly	Ser	Thr	His
			705						710			715			
Cys	Leu	Ala	Leu	Thr	Glu	Asp	Ser	Glu	Val	His	Ser	Trp	Gly	Ser	Asn
			725						730			735			
Asp	Gln	Cys	Gln	His	Phe	Asp	Thr	Leu	Arg	Val	Thr	Lys	Pro	Glu	Pro
			740						745			750			
Ala	Ala	Leu	Pro	Gly	Leu	Asp	Thr	Lys	His	Ile	Val	Gly	Ile	Ala	Cys
			755						760			765			
Gly	Pro	Ala	Gln	Ser	Phe	Ala	Trp	Ser	Ser	Cys	Ser	Glu	Trp	Ser	Ile
			770						775			780			
Gly	Leu	Arg	Val	Pro	Phe	Val	Val	Asp	Ile	Cys	Ser	Met	Thr	Phe	Glu
			785						790			795			
Gln	Leu	Asp	Leu	Leu	Leu	Arg	Gln	Val	Ser	Glu	Gly	Met	Asp	Gly	Ser
			805						810			815			

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Ala	Asp	Trp	Pro	Pro	Pro	Gln	Glu	Lys	Glu	Cys	Val	Ala	Val	Ala	Thr	820	825	830
Leu	Asn	Leu	Leu	Arg	Leu	Gln	Leu	His	Ala	Ala	Ile	Ser	His	Gln	Val	835	840	845
Asp	Pro	Glu	Phe	Leu	Gly	Leu	Gly	Leu	Gly	Ser	Ile	Leu	Leu	Asn	Ser	850	855	860
Leu	Lys	Gln	Thr	Val	Val	Thr	Leu	Ala	Ser	Ser	Ala	Gly	Val	Leu	Ser	865	870	880
Thr	Val	Gln	Ser	Ala	Ala	Gln	Ala	Val	Leu	Gln	Ser	Gly	Trp	Ser	Val	885	890	895
Leu	Leu	Pro	Thr	Ala	Glu	Glu	Arg	Ala	Arg	Ala	Leu	Ser	Ala	Leu	Leu	900	905	910
Pro	Cys	Ala	Val	Ser	Gly	Asn	Glu	Val	Asn	Ile	Ser	Pro	Gly	Arg	Arg	915	920	925
Phe	Met	Ile	Asp	Leu	Leu	Val	Gly	Ser	Leu	Met	Ala	Asp	Gly	Gly	Leu	930	935	940
Glu	Ser	Ala	Leu	His	Ala	Ala	Ile	Thr	Ala	Glu	Ile	Gln	Asp	Ile	Glu	945	950	955
Ala	Lys	Lys	Glu	Ala	Gln	Lys	Glu	Lys	Glu	Ile	Asp	Glu	Gln	Glu	Ala	965	970	975
Asn	Ala	Ser	Thr	Phe	His	Arg	Ser	Arg	Thr	Pro	Leu	Asp	Lys	Asp	Leu	980	985	990
Ile	Asn	Thr	Gly	Ile	Cys	Glu	Ser	Ser	Gly	Lys	Gln	Cys	Leu	Pro	Leu	995	1000	1005
Val	Gln	Leu	Ile	Gln	Gln	Leu	Leu	Arg	Asn	Ile	Ala	Ser	Gln	Thr		1010	1015	1020
Val	Ala	Arg	Leu	Lys	Asp	Val	Ala	Arg	Arg	Ile	Ser	Ser	Cys	Leu		1025	1030	1035
Asp	Phe	Glu	Gln	His	Ser	Arg	Glu	Arg	Ser	Ala	Ser	Leu	Asp	Leu		1040	1045	1050
Leu	Leu	Arg	Phe	Gln	Arg	Leu	Leu	Ile	Ser	Lys	Leu	Tyr	Pro	Gly		1055	1060	1065
Glu	Ser	Ile	Gly	Gln	Thr	Ser	Asp	Ile	Ser	Ser	Pro	Glu	Leu	Met		1070	1075	1080
Gly	Val	Gly	Ser	Leu	Leu	Lys	Lys	Tyr	Thr	Ala	Leu	Leu	Cys	Thr		1085	1090	1095
His	Ile	Gly	Asp	Ile	Leu	Pro	Val	Ala	Ala	Ser	Ile	Ala	Ser	Thr		1100	1105	1110
Ser	Trp	Arg	His	Phe	Ala	Glu	Val	Ala	Tyr	Ile	Val	Glu	Gly	Asp		1115	1120	1125
Phe	Thr	Gly	Val	Leu	Leu	Pro	Glu	Leu	Val	Val	Ser	Ile	Val	Leu		1130	1135	1140
Leu	Leu	Ser	Lys	Asn	Ala	Gly	Leu	Met	Gln	Glu	Ala	Gly	Ala	Val		1145	1150	1155
Pro	Leu	Leu	Gly	Gly	Leu	Leu	Glu	His	Leu	Asp	Arg	Phe	Asn	His		1160	1165	1170
Leu	Ala	Pro	Gly	Lys	Glu	Arg	Asp	Asp	His	Glu	Glu	Leu	Ala	Trp		1175	1180	1185
Pro	Gly	Ile	Met	Glu	Ser	Phe	Phe	Thr	Gly	Gln	Asn	Cys	Arg	Asn		1190	1195	1200
Asn	Glu	Glu	Val	Thr	Leu	Ile	Arg	Lys	Ala	Asp	Leu	Glu	Asn	His				

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1205	1210	1215
Asn Lys Asp Gly Gly Phe Trp Thr Val Ile Asp Gly Lys Val Tyr		
1220	1225	1230
Asp Ile Lys Asp Phe Gln Thr Gln Ser Leu Thr Gly Asn Ser Ile		
1235	1240	1245
Leu Ala Gln Phe Ala Gly Glu Asp Pro Val Val Ala Leu Glu Ala		
1250	1255	1260
Ala Leu Gln Phe Glu Asp Thr Arg Glu Ser Met His Ala Phe Cys		
1265	1270	1275
Val Gly Gln Tyr Leu Glu Pro Asp Gln Glu Ile Val Thr Ile Pro		
1280	1285	1290
Asp Leu Gly Ser Leu Ser Ser Pro Leu Ile Asp Thr Glu Arg Asn		
1295	1300	1305
Leu Gly Leu Leu Leu Gly Leu His Ala Ser Tyr Leu Ala Met Ser		
1310	1315	1320
Thr Pro Leu Ser Pro Val Glu Ile Glu Cys Ala Lys Trp Leu Gln		
1325	1330	1335
Ser Ser Ile Phe Ser Gly Gly Leu Gln Thr Ser Gln Ile His Tyr		
1340	1345	1350
Ser Tyr Asn Glu Glu Lys Asp Glu Asp His Cys Ser Ser Pro Gly		
1355	1360	1365
Gly Thr Pro Ala Ser Lys Ser Arg Leu Cys Ser His Arg Arg Ala		
1370	1375	1380
Leu Gly Asp His Ser Gln Ala Phe Leu Gln Ala Ile Ala Asp Asn		
1385	1390	1395
Asn Ile Gln Asp His Asn Val Lys Asp Phe Leu Cys Gln Ile Glu		
1400	1405	1410
Arg Tyr Cys Arg Gln Cys His Leu Thr Thr Pro Ile Met Phe Pro		
1415	1420	1425
Pro Glu His Pro Val Glu Glu Val Gly Arg Leu Leu Leu Cys Cys		
1430	1435	1440
Leu Leu Lys His Glu Asp Leu Gly His Val Ala Leu Ser Leu Val		
1445	1450	1455
His Ala Gly Ala Leu Gly Ile Glu Gln Val Lys His Arg Thr Leu		
1460	1465	1470
Pro Lys Ser Val Val Asp Val Cys Arg Val Val Tyr Gln Ala Lys		
1475	1480	1485
Cys Ser Leu Ile Lys Thr His Gln Glu Gln Gly Arg Ser Tyr Lys		
1490	1495	1500
Glu Val Cys Ala Pro Val Ile Glu Arg Leu Arg Phe Leu Phe Asn		
1505	1510	1515
Glu Leu Arg Pro Ala Val Cys Asn Asp Leu Ser Ile Met Ser Lys		
1520	1525	1530
Phe Lys Leu Leu Ser Ser Leu Pro Arg Trp Arg Arg Ile Ala Gln		
1535	1540	1545
Lys Ile Ile Arg Glu Arg Arg Lys Lys Arg Val Pro Lys Lys Pro		
1550	1555	1560
Glu Ser Thr Asp Asp Glu Glu Lys Ile Gly Asn Glu Glu Ser Asp		
1565	1570	1575
Leu Glu Glu Ala Cys Ile Leu Pro His Ser Pro Ile Asn Val Asp		
1580	1585	1590

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Lys Arg	Pro Ile Ala Ile	Lys Ser	Pro Lys Asp	Lys Trp Gln Pro
1595		1600		1605
Leu Leu	Ser Thr Val Thr	Gly Val His Lys Tyr	Lys Trp Leu Lys	
1610		1615	1620	
Gln Asn	Val Gln Gly Leu Tyr	Pro Gln Ser Pro	Leu Leu Ser Thr	
1625		1630	1635	
Ile Ala	Glu Phe Ala Leu Lys	Glu Glu Pro Val	Asp Val Glu Lys	
1640		1645	1650	
Met Arg	Lys Cys Leu Leu Lys	Gln Leu Glu Arg	Ala Glu Val Arg	
1655		1660	1665	
Leu Glu	Gly Ile Asp Thr	Ile Leu Lys Leu Ala	Ser Lys Asn Phe	
1670		1675	1680	
Leu Leu	Pro Ser Val Gln Tyr	Ala Met Phe Cys Gly	Trp Gln Arg	
1685		1690	1695	
Leu Ile	Pro Glu Gly Ile Asp	Ile Gly Glu Pro	Leu Thr Asp Cys	
1700		1705	1710	
Leu Lys	Asp Val Asp Leu Ile	Pro Pro Phe Asn Arg	Met Leu Leu	
1715		1720	1725	
Glu Val	Thr Phe Gly Lys Leu	Tyr Ala Trp Ala Val	Gln Asn Ile	
1730		1735	1740	
Arg Asn	Val Leu Met Asp Ala	Ser Ala Lys Phe Lys	Glu Leu Gly	
1745		1750	1755	
Ile Gln	Pro Val Pro Leu Gln	Thr Ile Thr Asn Glu	Asn Pro Ser	
1760		1765	1770	
Gly Pro	Ser Leu Gly Thr Ile	Pro Gln Ala Arg Phe	Leu Leu Val	
1775		1780	1785	
Met Leu	Ser Met Leu Thr Leu	Gln His Gly Ala Asn	Asn Leu Asp	
1790		1795	1800	
Leu Leu	Leu Asn Ser Gly Met	Leu Ala Leu Thr Gln	Thr Ala Leu	
1805		1810	1815	
Arg Leu	Ile Gly Pro Ser Cys	Asp Asn Val Glu Glu	Asp Met Asn	
1820		1825	1830	
Ala Ser	Ala Gln Gly Ala Ser	Ala Thr Val Leu Glu	Glu Thr Arg	
1835		1840	1845	
Lys Glu	Thr Ala Pro Val Gln	Leu Pro Val Ser Gly	Pro Glu Leu	
1850		1855	1860	
Ala Ala	Met Met Lys Ile Gly	Thr Arg Val Met Arg	Gly Val Asp	
1865		1870	1875	
Trp Lys	Trp Gly Asp Gln Asp	Gly Pro Pro Pro Gly	Leu Gly Arg	
1880		1885	1890	
Val Ile	Gly Glu Leu Gly Glu	Asp Gly Trp Ile Arg	Val Gln Trp	
1895		1900	1905	
Asp Thr	Gly Ser Thr Asn Ser	Tyr Arg Met Gly Lys	Glu Gly Lys	
1910		1915	1920	
Tyr Asp	Leu Lys Leu Ala Glu	Leu Pro Ala Ala Ala	Gln Pro Ser	
1925		1930	1935	
Ala Glu	Asp Ser Asp Thr Glu	Asp Asp Ser Glu Ala	Glu Gln Thr	
1940		1945	1950	
Glu Arg	Asn Ile His Pro Thr	Ala Met Met Phe Thr	Ser Thr Ile	
1955		1960	1965	

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Asn	Leu	Leu	Gln	Thr	Leu	Cys	Leu	Ser	Ala	Gly	Val	His	Ala	Glu
1970						1975					1980			
Ile	Met	Gln	Ser	Glu	Ala	Thr	Lys	Thr	Leu	Cys	Gly	Leu	Leu	Arg
1985						1990					1995			
Met	Leu	Val	Glu	Ser	Gly	Thr	Thr	Asp	Lys	Thr	Ser	Ser	Pro	Asn
2000						2005					2010			
Arg	Leu	Val	Tyr	Arg	Glu	Gln	His	Arg	Ser	Trp	Cys	Thr	Leu	Gly
2015						2020					2025			
Phe	Val	Arg	Ser	Ile	Ala	Leu	Thr	Pro	Gln	Val	Cys	Gly	Ala	Leu
2030						2035					2040			
Ser	Ser	Pro	Gln	Trp	Ile	Thr	Leu	Leu	Met	Lys	Val	Val	Glu	Gly
2045						2050					2055			
His	Ala	Pro	Phe	Thr	Ala	Thr	Ser	Leu	Gln	Arg	Gln	Ile	Leu	Ala
2060						2065					2070			
Val	His	Leu	Leu	Gln	Ala	Val	Leu	Pro	Ser	Trp	Asp	Lys	Thr	Glu
2075						2080					2085			
Arg	Ala	Arg	Asp	Met	Lys	Cys	Leu	Val	Glu	Lys	Leu	Phe	Asp	Phe
2090						2095					2100			
Leu	Gly	Ser	Leu	Leu	Thr	Thr	Cys	Ser	Ser	Asp	Val	Pro	Leu	Leu
2105						2110					2115			
Arg	Glu	Ser	Thr	Leu	Arg	Arg	Arg	Arg	Val	Arg	Pro	Gln	Ala	Ser
2120						2125					2130			
Leu	Thr	Ala	Thr	His	Ser	Ser	Thr	Leu	Ala	Glu	Glu	Val	Val	Ala
2135						2140					2145			
Leu	Leu	Arg	Thr	Leu	His	Ser	Leu	Thr	Gln	Trp	Asn	Gly	Leu	Ile
2150						2155					2160			
Asn	Lys	Tyr	Ile	Asn	Ser	Gln	Leu	Arg	Ser	Ile	Thr	His	Ser	Phe
2165						2170					2175			
Val	Gly	Arg	Pro	Ser	Glu	Gly	Ala	Gln	Leu	Glu	Asp	Tyr	Phe	Pro
2180						2185					2190			
Asp	Ser	Glu	Asn	Pro	Glu	Val	Gly	Gly	Leu	Met	Ala	Val	Leu	Ala
2195						2200					2205			
Val	Ile	Gly	Gly	Ile	Asp	Gly	Arg	Leu	Arg	Leu	Gly	Gly	Gln	Val
2210						2215					2220			
Met	His	Asp	Glu	Phe	Gly	Glu	Gly	Thr	Val	Thr	Arg	Ile	Thr	Pro
2225						2230					2235			
Lys	Gly	Lys	Ile	Thr	Val	Gln	Phe	Ser	Asp	Met	Arg	Thr	Cys	Arg
2240						2245					2250			
Val	Cys	Pro	Leu	Asn	Gln	Leu	Lys	Pro	Leu	Pro	Ala	Val	Ala	Phe
2255						2260					2265			
Asn	Val	Asn	Asn	Leu	Pro	Phe	Thr	Glu	Pro	Met	Leu	Ser	Val	Trp
2270						2275					2280			
Ala	Gln	Leu	Val	Asn	Leu	Ala	Gly	Ser	Lys	Leu	Glu	Lys	His	Lys
2285						2290					2295			
Ile	Lys	Lys	Ser	Thr	Lys	Gln	Ala	Phe	Ala	Gly	Gln	Val	Asp	Leu
2300						2305					2310			
Asp	Leu	Leu	Arg	Cys	Gln	Gln	Leu	Lys	Leu	Tyr	Ile	Leu	Lys	Ala
2315						2320					2325			
Gly	Arg	Ala	Leu	Leu	Ser	His	Gln	Asp	Lys	Leu	Arg	Gln	Ile	Leu
2330						2335					2340			
Ser	Gln	Pro	Ala	Val	Gln	Glu	Thr	Gly	Thr	Val	His	Thr	Asp	Asp

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2345	2350	2355
Gly Ala Val Val Ser Pro Asp Leu Gly Asp Met Ser Pro Glu Gly		
2360	2365	2370
Pro Gln Pro Pro Met Ile Leu Leu Gln Gln Leu Leu Ala Ser Ala		
2375	2380	2385
Thr Gln Pro Ser Pro Val Lys Ala Ile Phe Asp Lys Gln Glu Leu		
2390	2395	2400
Glu Ala Ala Ala Leu Ala Val Cys Gln Cys Leu Ala Val Glu Ser		
2405	2410	2415
Thr His Pro Ser Ser Pro Gly Phe Glu Asp Cys Ser Ser Ser Glu		
2420	2425	2430
Ala Thr Thr Pro Val Ala Val Gln His Ile Arg Pro Ala Arg Val		
2435	2440	2445
Lys Arg Arg Lys Gln Ser Pro Val Pro Ala Leu Pro Ile Val Val		
2450	2455	2460
Gln Leu Met Glu Met Gly Phe Ser Arg Arg Asn Ile Glu Phe Ala		
2465	2470	2475
Leu Lys Ser Leu Thr Gly Ala Ser Gly Asn Ala Ser Ser Leu Pro		
2480	2485	2490
Gly Val Glu Ala Leu Val Gly Trp Leu Leu Asp His Ser Asp Ile		
2495	2500	2505
Gln Val Thr Glu Leu Ser Asp Ala Asp Thr Val Ser Asp Glu Tyr		
2510	2515	2520
Ser Asp Glu Glu Val Val Glu Asp Val Asp Asp Ala Ala Tyr Ser		
2525	2530	2535
Met Ser Thr Gly Ala Val Val Thr Glu Ser Gln Thr Tyr Lys Lys		
2540	2545	2550
Arg Ala Asp Phe Leu Ser Asn Asp Asp Tyr Ala Val Tyr Val Arg		
2555	2560	2565
Glu Asn Ile Gln Val Gly Met Met Val Arg Cys Cys Arg Ala Tyr		
2570	2575	2580
Glu Glu Val Cys Glu Gly Asp Val Gly Lys Val Ile Lys Leu Asp		
2585	2590	2595
Arg Asp Gly Leu His Asp Leu Asn Val Gln Cys Asp Trp Gln Gln		
2600	2605	2610
Lys Gly Gly Thr Tyr Trp Val Arg Tyr Ile His Val Glu Leu Ile		
2615	2620	2625
Gly Tyr Pro Pro Pro Ser Ser Ser Ser His Ile Lys Ile Gly Asp		
2630	2635	2640
Lys Val Arg Val Lys Ala Ser Val Thr Thr Pro Lys Tyr Lys Trp		
2645	2650	2655
Gly Ser Val Thr His Gln Ser Val Gly Val Val Lys Ala Phe Ser		
2660	2665	2670
Ala Asn Gly Lys Asp Ile Ile Val Asp Phe Pro Gln Gln Ser His		
2675	2680	2685
Trp Thr Gly Leu Leu Ser Glu Met Glu Leu Val Pro Ser Ile His		
2690	2695	2700
Pro Gly Val Thr Cys Asp Gly Cys Gln Met Phe Pro Ile Asn Gly		
2705	2710	2715
Ser Arg Phe Lys Cys Arg Asn Cys Asp Asp Phe Asp Phe Cys Glu		
2720	2725	2730

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Thr Cys	Phe Lys Thr Lys Lys	His Asn Thr Arg	His Thr Phe Gly
2735	2740	2745	
Arg Ile	Asn Glu Pro Gly Gln	Ser Ala Val Phe Cys	Gly Arg Ser
2750	2755	2760	
Gly Lys	Gln Leu Lys Arg Cys	His Ser Ser Gln Pro	Gly Met Leu
2765	2770	2775	
Leu Asp	Ser Trp Ser Arg Met	Val Lys Ser Leu Asn	Val Ser Ser
2780	2785	2790	
Ser Val	Asn Gln Ala Ser Arg	Leu Ile Asp Gly Ser	Glu Pro Cys
2795	2800	2805	
Trp Gln	Ser Ser Gly Ser Gln	Gly Lys His Trp Ile	Arg Leu Glu
2810	2815	2820	
Ile Phe	Pro Asp Val Leu Val	His Arg Leu Lys Met	Ile Val Asp
2825	2830	2835	
Pro Ala	Asp Ser Ser Tyr Met	Pro Ser Leu Val Val	Val Ser Gly
2840	2845	2850	
Gly Asn	Ser Leu Asn Asn Leu	Ile Glu Leu Lys Thr	Ile Asn Ile
2855	2860	2865	
Asn Pro	Ser Asp Thr Thr Val	Pro Leu Leu Asn Asp	Cys Thr Glu
2870	2875	2880	
Tyr His	Arg Tyr Ile Glu Ile	Ala Ile Lys Gln Cys	Arg Ser Ser
2885	2890	2895	
Gly Ile	Asp Cys Lys Ile His	Gly Leu Ile Leu Leu	Gly Arg Ile
2900	2905	2910	
Arg Ala	Glu Glu Glu Asp Leu	Ala Ala Val Pro Phe	Leu Ala Ser
2915	2920	2925	
Asp Asn	Glu Glu Glu Glu Asp	Glu Lys Gly Asn Ser	Gly Ser Leu
2930	2935	2940	
Ile Arg	Lys Lys Ala Ala Gly	Leu Glu Ser Ala Ala	Thr Ile Arg
2945	2950	2955	
Thr Lys	Val Phe Val Trp Gly	Leu Asn Asp Lys Asp	Gln Leu Gly
2960	2965	2970	
Gly Leu	Lys Gly Ser Lys Ile	Lys Val Pro Ser Phe	Ser Glu Thr
2975	2980	2985	
Leu Ser	Ala Leu Asn Val Val	Gln Val Ala Gly Gly	Ser Lys Ser
2990	2995	3000	
Leu Phe	Ala Val Thr Val Glu	Gly Lys Val Tyr Ala	Cys Gly Glu
3005	3010	3015	
Ala Thr	Asn Gly Arg Leu Gly	Leu Gly Ile Ser Ser	Gly Thr Val
3020	3025	3030	
Pro Ile	Pro Arg Gln Ile Thr	Ala Leu Ser Ser Tyr	Val Val Lys
3035	3040	3045	
Lys Val	Ala Val His Ser Gly	Gly Arg His Ala Thr	Ala Leu Thr
3050	3055	3060	
Val Asp	Gly Lys Val Phe Ser	Trp Gly Glu Gly Asp	Asp Gly Lys
3065	3070	3075	
Leu Gly	His Phe Ser Arg Met	Asn Cys Asp Lys Pro	Arg Leu Ile
3080	3085	3090	
Glu Ala	Leu Lys Thr Lys Arg	Ile Arg Asp Ile Ala	Cys Gly Ser
3095	3100	3105	

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Ser 3110	His	Ser	Ala	Ala	Leu	Thr 3115	Ser	Ser	Gly	Glu	Leu 3120	Tyr	Thr	Trp
Gly 3125	Leu	Gly	Glu	Tyr	Gly	Arg 3130	Leu	Gly	His	Gly	Asp 3135	Asn	Thr	Thr
Gln 3140	Leu	Lys	Pro	Lys	Met	Val 3145	Lys	Val	Leu	Leu	Gly 3150	His	Arg	Val
Ile 3155	Gln	Val	Ala	Cys	Gly	Ser 3160	Arg	Asp	Ala	Gln	Thr 3165	Leu	Ala	Leu
Thr 3170	Asp	Glu	Gly	Leu	Val	Phe 3175	Ser	Trp	Gly	Asp	Gly 3180	Asp	Phe	Gly
Lys 3185	Leu	Gly	Arg	Gly	Gly	Ser 3190	Glu	Gly	Cys	Asn	Ile 3195	Pro	Gln	Asn
Ile 3200	Glu	Arg	Leu	Asn	Gly	Gln 3205	Gly	Val	Cys	Gln	Ile 3210	Glu	Cys	Gly
Ala 3215	Gln	Phe	Ser	Leu	Ala	Leu 3220	Thr	Lys	Ser	Gly	Val 3225	Val	Trp	Thr
Trp 3230	Gly	Lys	Gly	Asp	Tyr	Phe 3235	Arg	Leu	Gly	His	Gly 3240	Ser	Asp	Val
His 3245	Val	Arg	Lys	Pro	Gln	Val 3250	Val	Glu	Gly	Leu	Arg 3255	Gly	Lys	Lys
Ile 3260	Val	His	Val	Ala	Val	Gly 3265	Ala	Leu	His	Cys	Leu 3270	Ala	Val	Thr
Asp 3275	Ser	Gly	Gln	Val	Tyr	Ala 3280	Trp	Gly	Asp	Asn	Asp 3285	His	Gly	Gln
Gln 3290	Gly	Asn	Gly	Thr	Thr	Thr 3295	Val	Asn	Arg	Lys	Pro 3300	Thr	Leu	Val
Gln 3305	Gly	Leu	Glu	Gly	Gln	Lys 3310	Ile	Thr	Arg	Val	Ala 3315	Cys	Gly	Ser
Ser 3320	His	Ser	Val	Ala	Trp	Thr 3325	Thr	Val	Asp	Val	Ala 3330	Thr	Pro	Ser
Val 3335	His	Glu	Pro	Val	Leu	Phe 3340	Gln	Thr	Ala	Arg	Asp 3345	Pro	Leu	Gly
Ala 3350	Ser	Tyr	Leu	Gly	Val	Pro 3355	Ser	Asp	Ala	Asp	Ser 3360	Ser	Ala	Ala
Ser 3365	Asn	Lys	Ile	Ser	Gly	Ala 3370	Ser	Asn	Ser	Lys	Pro 3375	Asn	Arg	Pro
Ser 3380	Leu	Ala	Lys	Ile	Leu	Leu 3385	Ser	Leu	Asp	Gly	Asn 3390	Leu	Ala	Lys
Gln 3395	Gln	Ala	Leu	Ser	His	Ile 3400	Leu	Thr	Ala	Leu	Gln 3405	Ile	Met	Tyr
Ala 3410	Arg	Asp	Ala	Val	Val	Gly 3415	Ala	Leu	Met	Pro	Ala 3420	Ala	Met	Ile
Ala 3425	Pro	Val	Glu	Cys	Pro	Ser 3430	Phe	Ser	Ser	Ala	Ala 3435	Pro	Ser	Asp
Ala 3440	Ser	Ala	Met	Ala	Ser	Pro 3445	Met	Asn	Gly	Glu	Glu 3450	Cys	Met	Leu
Ala 3455	Val	Asp	Ile	Glu	Asp	Arg 3460	Leu	Ser	Pro	Asn	Pro 3465	Trp	Gln	Glu
Lys 3470	Arg	Glu	Ile	Val	Ser	Ser 3475	Glu	Asp	Ala	Val	Thr 3480	Pro	Ser	Ala
Val	Thr	Pro	Ser	Ala	Pro	Ser	Ala	Ser	Ala	Arg	Pro	Phe	Ile	Pro

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3485	3490	3495
Val Thr Asp Asp Leu Gly Ala	Ala Ser Ile Ile Ala	Glu Thr Met
3500	3505	3510
Thr Lys Thr Lys Glu Asp Val	Glu Ser Gln Asn Lys	Ala Ala Gly
3515	3520	3525
Pro Glu Pro Gln Ala Leu Asp	Glu Phe Thr Ser Leu	Leu Ile Ala
3530	3535	3540
Asp Asp Thr Arg Val Val Val	Asp Leu Leu Lys Leu	Ser Val Cys
3545	3550	3555
Ser Arg Ala Gly Asp Arg Gly	Arg Asp Val Leu Ser	Ala Val Leu
3560	3565	3570
Ser Gly Met Gly Thr Ala Tyr	Pro Gln Val Ala Asp	Met Leu Leu
3575	3580	3585
Glu Leu Cys Val Thr Glu Leu	Glu Asp Val Ala Thr	Asp Ser Gln
3590	3595	3600
Ser Gly Arg Leu Ser Ser Gln	Pro Val Val Val Glu	Ser Ser His
3605	3610	3615
Pro Tyr Thr Asp Asp Thr Ser	Thr Ser Gly Thr Val	Lys Ile Pro
3620	3625	3630
Gly Ala Glu Gly Leu Arg Val	Glu Phe Asp Arg Gln	Cys Ser Thr
3635	3640	3645
Glu Arg Arg His Asp Pro Leu	Thr Val Met Asp Gly	Val Asn Arg
3650	3655	3660
Ile Val Ser Val Arg Ser Gly	Arg Glu Trp Ser Asp	Trp Ser Ser
3665	3670	3675
Glu Leu Arg Ile Pro Gly Asp	Glu Leu Lys Trp Lys	Phe Ile Ser
3680	3685	3690
Asp Gly Ser Val Asn Gly Trp	Gly Trp Arg Phe Thr	Val Tyr Pro
3695	3700	3705
Ile Met Pro Ala Ala Gly Pro	Lys Glu Leu Leu Ser	Asp Arg Cys
3710	3715	3720
Val Leu Ser Cys Pro Ser Met	Asp Leu Val Thr Cys	Leu Leu Asp
3725	3730	3735
Phe Arg Leu Asn Leu Ala Ser	Asn Arg Ser Ile Val	Pro Arg Leu
3740	3745	3750
Ala Ala Ser Leu Ala Ala Cys	Ala Gln Leu Ser Ala	Leu Ala Ala
3755	3760	3765
Ser His Arg Met Trp Ala Leu	Gln Arg Leu Arg Lys	Leu Leu Thr
3770	3775	3780
Thr Glu Phe Gly Gln Ser Ile	Asn Ile Asn Arg Leu	Leu Gly Glu
3785	3790	3795
Asn Asp Gly Glu Thr Arg Ala	Leu Ser Phe Thr Gly	Ser Ala Leu
3800	3805	3810
Ala Ala Leu Val Lys Gly Leu	Pro Glu Ala Leu Gln	Arg Gln Phe
3815	3820	3825
Glu Tyr Glu Asp Pro Ile Val	Arg Gly Gly Lys Gln	Leu Leu His
3830	3835	3840
Ser Pro Phe Phe Lys Val Leu	Val Ala Leu Ala Cys	Asp Leu Glu
3845	3850	3855
Leu Asp Thr Leu Pro Cys Cys	Ala Glu Thr His Lys	Trp Ala Trp
3860	3865	3870

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Phe Arg 3875	Arg Tyr Cys Met 3880	Ala Ser Arg Val Ala Val 3885	Ala Leu Asp
Lys Arg 3890	Thr Pro Leu Pro Arg 3895	Leu Phe Leu Asp Glu 3900	Val Ala Lys
Lys Ile 3905	Arg Glu Leu Met Ala 3910	Asp Ser Glu Asn Met 3915	Asp Val Leu
His Glu 3920	Ser His Asp Ile Phe 3925	Lys Arg Glu Gln Asp 3930	Glu Gln Leu
Val Gln 3935	Trp Met Asn Arg Arg 3940	Pro Asp Asp Trp Thr 3945	Leu Ser Ala
Gly Gly 3950	Ser Gly Thr Ile Tyr 3955	Gly Trp Gly His Asn 3960	His Arg Gly
Gln Leu 3965	Gly Gly Ile Glu Gly 3970	Ala Lys Val Lys Val 3975	Pro Thr Pro
Cys Glu 3980	Ala Leu Ala Thr Leu 3985	Arg Pro Val Gln Leu 3990	Ile Gly Gly
Glu Gln 3995	Thr Leu Phe Ala Val 4000	Thr Ala Asp Gly Lys 4005	Leu Tyr Ala
Thr Gly 4010	Tyr Gly Ala Gly Gly 4015	Arg Leu Gly Ile Gly 4020	Gly Thr Glu
Ser Val 4025	Ser Thr Pro Thr Leu 4030	Leu Glu Ser Ile Gln 4035	His Val Phe
Ile Lys 4040	Lys Val Ala Val Asn 4045	Ser Gly Gly Lys His 4050	Cys Leu Ala
Leu Ser 4055	Ser Glu Gly Glu Val 4060	Tyr Ser Trp Gly Glu 4065	Ala Glu Asp
Gly Lys 4070	Leu Gly His Gly Asn 4075	Arg Ser Pro Cys Asp 4080	Arg Pro Arg
Val Ile 4085	Glu Ser Leu Arg Gly 4090	Ile Glu Val Val Asp 4095	Val Ala Ala
Gly Gly 4100	Ala His Ser Ala Cys 4105	Val Thr Ala Ala Gly 4110	Asp Leu Tyr
Thr Trp 4115	Gly Lys Gly Arg Tyr 4120	Gly Arg Leu Gly His 4125	Ser Asp Ser
Glu Asp 4130	Gln Leu Lys Pro Lys 4135	Leu Val Glu Ala Leu 4140	Gln Gly His
Arg Val 4145	Val Asp Ile Ala Cys 4150	Gly Ser Gly Asp Ala 4155	Gln Thr Leu
Cys Leu 4160	Thr Asp Asp Asp Thr 4165	Val Trp Ser Trp Gly 4170	Asp Gly Asp
Tyr Gly 4175	Lys Leu Gly Arg Gly 4180	Gly Ser Asp Gly Cys 4185	Lys Val Pro
Met Lys 4190	Ile Asp Ser Leu Thr 4195	Gly Leu Gly Val Val 4200	Lys Val Glu
Cys Gly 4205	Ser Gln Phe Ser Val 4210	Ala Leu Thr Lys Ser 4215	Gly Ala Val
Tyr Thr 4220	Trp Gly Lys Gly Asp 4225	Tyr His Arg Leu Gly 4230	His Gly Ser
Asp Asp 4235	His Val Arg Arg Pro 4240	Arg Gln Val Gln Gly 4245	Leu Gln Gly

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Lys	Lys	Val	Ile	Ala	Ile	Ala	Thr	Gly	Ser	Leu	His	Cys	Val	Cys
4250						4255					4260			
Cys	Thr	Glu	Asp	Gly	Glu	Val	Tyr	Thr	Trp	Gly	Asp	Asn	Asp	Glu
4265						4270					4275			
Gly	Gln	Leu	Gly	Asp	Gly	Thr	Thr	Asn	Ala	Ile	Gln	Arg	Pro	Arg
4280						4285					4290			
Leu	Val	Ala	Ala	Leu	Gln	Gly	Lys	Lys	Val	Asn	Arg	Val	Ala	Cys
4295						4300					4305			
Gly	Ser	Ala	His	Thr	Leu	Ala	Trp	Ser	Thr	Ser	Lys	Pro	Ala	Ser
4310						4315					4320			
Ala	Gly	Lys	Leu	Pro	Ala	Gln	Val	Pro	Met	Glu	Tyr	Asn	His	Leu
4325						4330					4335			
Gln	Glu	Ile	Pro	Ile	Ile	Ala	Leu	Arg	Asn	Arg	Leu	Leu	Leu	Leu
4340						4345					4350			
His	His	Leu	Ser	Glu	Leu	Phe	Cys	Pro	Cys	Ile	Pro	Met	Phe	Asp
4355						4360					4365			
Leu	Glu	Gly	Ser	Leu	Asp	Glu	Thr	Gly	Leu	Gly	Pro	Ser	Val	Gly
4370						4375					4380			
Phe	Asp	Thr	Leu	Arg	Gly	Ile	Leu	Ile	Ser	Gln	Gly	Lys	Glu	Ala
4385						4390					4395			
Ala	Phe	Arg	Lys	Val	Val	Gln	Ala	Thr	Met	Val	Arg	Asp	Arg	Gln
4400						4405					4410			
His	Gly	Pro	Val	Val	Glu	Leu	Asn	Arg	Ile	Gln	Val	Lys	Arg	Ser
4415						4420					4425			
Arg	Ser	Lys	Gly	Gly	Leu	Ala	Gly	Pro	Asp	Gly	Thr	Lys	Ser	Val
4430						4435					4440			
Phe	Gly	Gln	Met	Cys	Ala	Lys	Met	Ser	Ser	Phe	Gly	Pro	Asp	Ser
4445						4450					4455			
Leu	Leu	Leu	Pro	His	Arg	Val	Trp	Lys	Val	Lys	Phe	Val	Gly	Glu
4460						4465					4470			
Ser	Val	Asp	Asp	Cys	Gly	Gly	Gly	Tyr	Ser	Glu	Ser	Ile	Ala	Glu
4475						4480					4485			
Ile	Cys	Glu	Glu	Leu	Gln	Asn	Gly	Leu	Thr	Pro	Leu	Leu	Ile	Val
4490						4495					4500			
Thr	Pro	Asn	Gly	Arg	Asp	Glu	Ser	Gly	Ala	Asn	Arg	Asp	Cys	Tyr
4505						4510					4515			
Leu	Leu	Ser	Pro	Ala	Ala	Arg	Ala	Pro	Val	His	Ser	Ser	Met	Phe
4520						4525					4530			
Arg	Phe	Leu	Gly	Val	Leu	Leu	Gly	Ile	Ala	Ile	Arg	Thr	Gly	Ser
4535						4540					4545			
Pro	Leu	Ser	Leu	Asn	Leu	Ala	Glu	Pro	Val	Trp	Lys	Gln	Leu	Ala
4550						4555					4560			
Gly	Met	Ser	Leu	Thr	Ile	Ala	Asp	Leu	Ser	Glu	Val	Asp	Lys	Asp
4565						4570					4575			
Phe	Ile	Pro	Gly	Leu	Met	Tyr	Ile	Arg	Asp	Asn	Glu	Ala	Thr	Ser
4580						4585					4590			
Glu	Glu	Phe	Glu	Ala	Met	Ser	Leu	Pro	Phe	Thr	Val	Pro	Ser	Ala
4595						4600					4605			
Ser	Gly	Gln	Asp	Ile	Gln	Leu	Ser	Ser	Lys	His	Thr	His	Ile	Thr
4610						4615					4620			
Leu	Asp	Asn	Arg	Ala	Glu	Tyr	Val	Arg	Leu	Ala	Ile	Asn	Tyr	Arg

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4625	4630	4635
Leu His Glu Phe Asp Glu Gln Val Ala Ala Val Arg Glu Gly Met		
4640	4645	4650
Ala Arg Val Val Pro Val Pro Leu Leu Ser Leu Phe Thr Gly Tyr		
4655	4660	4665
Glu Leu Glu Thr Met Val Cys Gly Ser Pro Asp Ile Pro Leu His		
4670	4675	4680
Leu Leu Lys Ser Val Ala Thr Tyr Lys Gly Ile Glu Pro Ser Ala		
4685	4690	4695
Ser Leu Ile Gln Trp Phe Trp Glu Val Met Glu Ser Phe Ser Asn		
4700	4705	4710
Thr Glu Arg Ser Leu Phe Leu Arg Phe Val Trp Gly Arg Thr Arg		
4715	4720	4725
Leu Pro Arg Thr Ile Ala Asp Phe Arg Gly Arg Asp Phe Val Ile		
4730	4735	4740
Gln Val Leu Asp Lys Tyr Asn Pro Pro Asp His Phe Leu Pro Glu		
4745	4750	4755
Ser Tyr Thr Cys Phe Phe Leu Leu Lys Leu Pro Arg Tyr Ser Cys		
4760	4765	4770
Lys Gln Val Leu Glu Glu Lys Leu Lys Tyr Ala Ile His Phe Cys		
4775	4780	4785
Lys Ser Ile Asp Thr Asp Asp Tyr Ala Arg Ile Ala Leu Thr Gly		
4790	4795	4800
Glu Pro Ala Ala Asp Asp Ser Ser Asp Asp Ser Asp Asn Glu Asp		
4805	4810	4815
Val Asp Ser Phe Ala Ser Asp Ser Thr Gln Asp Tyr Leu Thr Gly		
4820	4825	4830

His

<210> SEQ ID NO 12
 <211> LENGTH: 17
 <212> TYPE: PRT
 <213> ORGANISM: Homo sapiens

<400> SEQUENCE: 12

Lys Leu Ala Glu Leu Pro Ala Ala Ala Gln Pro Ser Ala Glu Asp Ser
 1 5 10 15

Asp

<210> SEQ ID NO 13
 <211> LENGTH: 579
 <212> TYPE: PRT
 <213> ORGANISM: Homo sapiens

<400> SEQUENCE: 13

Ala Asn Thr Phe Leu Glu Glu Val Arg Lys Gly Asn Leu Glu Arg Glu
 1 5 10 15

Cys Val Glu Glu Thr Cys Ser Tyr Glu Glu Ala Phe Glu Ala Leu Glu
 20 25 30

Ser Ser Thr Ala Thr Asp Val Phe Trp Ala Lys Tyr Thr Ala Cys Glu
 35 40 45

Thr Ala Arg Thr Pro Arg Asp Lys Leu Ala Ala Cys Leu Glu Gly Asn
 50 55 60

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Cys	Ala	Glu	Gly	Leu	Gly	Thr	Asn	Tyr	Arg	Gly	His	Val	Asn	Ile	Thr	65	70	75	80
Arg	Ser	Gly	Ile	Glu	Cys	Gln	Leu	Trp	Arg	Ser	Arg	Tyr	Pro	His	Lys	85	90	95	
Pro	Glu	Ile	Asn	Ser	Thr	Thr	His	Pro	Gly	Ala	Asp	Leu	Gln	Glu	Asn	100	105	110	
Phe	Cys	Arg	Asn	Pro	Asp	Ser	Ser	Thr	Thr	Gly	Pro	Trp	Cys	Tyr	Thr	115	120	125	
Thr	Asp	Pro	Thr	Val	Arg	Arg	Gln	Glu	Cys	Ser	Ile	Pro	Val	Cys	Gly	130	135	140	
Gln	Asp	Gln	Val	Thr	Val	Ala	Met	Thr	Pro	Arg	Ser	Glu	Gly	Ser	Ser	145	150	155	160
Val	Asn	Leu	Ser	Pro	Pro	Leu	Glu	Gln	Cys	Val	Pro	Asp	Arg	Gly	Gln	165	170	175	
Gln	Tyr	Gln	Gly	Arg	Leu	Ala	Val	Thr	Thr	His	Gly	Leu	Pro	Cys	Leu	180	185	190	
Ala	Trp	Ala	Ser	Ala	Gln	Ala	Lys	Ala	Leu	Ser	Lys	His	Gln	Asp	Phe	195	200	205	
Asn	Ser	Ala	Val	Gln	Leu	Val	Glu	Asn	Phe	Cys	Arg	Asn	Pro	Asp	Gly	210	215	220	
Asp	Glu	Glu	Gly	Val	Trp	Cys	Tyr	Val	Ala	Gly	Lys	Pro	Gly	Asp	Phe	225	230	235	240
Gly	Tyr	Cys	Asp	Leu	Asn	Tyr	Cys	Glu	Glu	Ala	Val	Glu	Glu	Glu	Thr	245	250	255	
Gly	Asp	Gly	Leu	Asp	Glu	Asp	Ser	Asp	Arg	Ala	Ile	Glu	Gly	Arg	Thr	260	265	270	
Ala	Thr	Ser	Glu	Tyr	Gln	Thr	Phe	Phe	Asn	Pro	Arg	Thr	Phe	Gly	Ser	275	280	285	
Gly	Glu	Ala	Asp	Cys	Gly	Leu	Arg	Pro	Leu	Phe	Glu	Lys	Lys	Ser	Leu	290	295	300	
Glu	Asp	Lys	Thr	Glu	Arg	Glu	Leu	Leu	Glu	Ser	Tyr	Ile	Asp	Gly	Arg	305	310	315	320
Ile	Val	Glu	Gly	Ser	Asp	Ala	Glu	Ile	Gly	Met	Ser	Pro	Trp	Gln	Val	325	330	335	
Met	Leu	Phe	Arg	Lys	Ser	Pro	Gln	Glu	Leu	Leu	Cys	Gly	Ala	Ser	Leu	340	345	350	
Ile	Ser	Asp	Arg	Trp	Val	Leu	Thr	Ala	Ala	His	Cys	Leu	Leu	Tyr	Pro	355	360	365	
Pro	Trp	Asp	Lys	Asn	Phe	Thr	Glu	Asn	Asp	Leu	Leu	Val	Arg	Ile	Gly	370	375	380	
Lys	His	Ser	Arg	Thr	Arg	Tyr	Glu	Arg	Asn	Ile	Glu	Lys	Ile	Ser	Met	385	390	395	400
Leu	Glu	Lys	Ile	Tyr	Ile	His	Pro	Arg	Tyr	Asn	Trp	Arg	Glu	Asn	Leu	405	410	415	
Asp	Arg	Asp	Ile	Ala	Leu	Met	Lys	Leu	Lys	Lys	Pro	Val	Ala	Phe	Ser	420	425	430	
Asp	Tyr	Ile	His	Pro	Val	Cys	Leu	Pro	Asp	Arg	Glu	Thr	Ala	Ala	Ser	435	440	445	
Leu	Leu	Gln	Ala	Gly	Tyr	Lys	Gly	Arg	Val	Thr	Gly	Trp	Gly	Asn	Leu	450	455	460	
Lys	Glu	Thr	Trp	Thr	Ala	Asn	Val	Gly	Lys	Gly	Gln	Pro	Ser	Val	Leu				

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465	470	475	480
Gln Val Val Asn Leu Pro Ile Val Glu Arg Pro Val Cys Lys Asp Ser			
	485	490	495
Thr Arg Ile Arg Ile Thr Asp Asn Met Phe Cys Ala Gly Tyr Lys Pro			
	500	505	510
Asp Glu Gly Lys Arg Gly Asp Ala Cys Glu Gly Asp Ser Gly Gly Pro			
	515	520	525
Phe Val Met Lys Ser Pro Phe Asn Asn Arg Trp Tyr Gln Met Gly Ile			
	530	535	540
Val Ser Trp Gly Glu Gly Cys Asp Arg Asp Gly Lys Tyr Gly Phe Tyr			
	545	550	555
Thr His Val Phe Arg Leu Lys Lys Trp Ile Gln Lys Val Ile Asp Gln			
	565	570	575
Phe Gly Glu			

<210> SEQ ID NO 14
 <211> LENGTH: 21
 <212> TYPE: PRT
 <213> ORGANISM: Homo sapiens

<400> SEQUENCE: 14

Thr Ala Thr Ser Glu Tyr Gln Thr Phe Phe Asn Pro Arg Thr Phe Gly
1 5 10 15
Ser Gly Glu Ala Asp
20

<210> SEQ ID NO 15
 <211> LENGTH: 127
 <212> TYPE: PRT
 <213> ORGANISM: Homo sapiens

<400> SEQUENCE: 15

Gly Pro Thr Gly Thr Gly Glu Ser Lys Cys Pro Leu Met Val Lys Val
1 5 10 15
Leu Asp Ala Val Arg Gly Ser Pro Ala Ile Asn Val Ala Val His Val
20 25 30
Phe Arg Lys Ala Ala Asp Asp Thr Trp Glu Pro Phe Ala Ser Gly Lys
35 40 45
Thr Ser Glu Ser Gly Glu Leu His Gly Leu Thr Thr Glu Glu Glu Phe
50 55 60
Val Glu Gly Ile Tyr Lys Val Glu Ile Asp Thr Lys Ser Tyr Trp Lys
65 70 75 80
Ala Leu Gly Ile Ser Pro Phe His Glu His Ala Glu Val Val Phe Thr
85 90 95
Ala Asn Asp Ser Gly Pro Arg Arg Tyr Thr Ile Ala Ala Leu Leu Ser
100 105 110
Pro Tyr Ser Tyr Ser Thr Thr Ala Val Val Thr Asn Pro Lys Glu
115 120 125

<210> SEQ ID NO 16
 <211> LENGTH: 20
 <212> TYPE: PRT
 <213> ORGANISM: Homo sapiens

<400> SEQUENCE: 16

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Ala Val Arg Gly Ser Pro Ala Ile Asn Val Ala Val His Val Phe Arg
 1 5 10 15

Lys Ala Ala Asp
 20

<210> SEQ ID NO 17
 <211> LENGTH: 399
 <212> TYPE: PRT
 <213> ORGANISM: Homo sapiens

<400> SEQUENCE: 17

Lys Glu Ala Cys Pro Thr Gly Leu Tyr Thr His Ser Gly Glu Cys Cys
 1 5 10 15

Lys Ala Cys Asn Leu Gly Glu Gly Val Ala Gln Pro Cys Gly Ala Asn
 20 25 30

Gln Thr Val Cys Glu Pro Cys Leu Asp Ser Val Thr Phe Ser Asp Val
 35 40 45

Val Ser Ala Thr Glu Pro Cys Lys Pro Cys Thr Glu Cys Val Gly Leu
 50 55 60

Gln Ser Met Ser Ala Pro Cys Val Glu Ala Asp Asp Ala Val Cys Arg
 65 70 75 80

Cys Ala Tyr Gly Tyr Tyr Gln Asp Glu Thr Thr Gly Arg Cys Glu Ala
 85 90 95

Cys Arg Val Cys Glu Ala Gly Ser Gly Leu Val Phe Ser Cys Gln Asp
 100 105 110

Lys Gln Asn Thr Val Cys Glu Glu Cys Pro Asp Gly Thr Tyr Ser Asp
 115 120 125

Glu Ala Asn His Val Asp Pro Cys Leu Pro Cys Thr Val Cys Glu Asp
 130 135 140

Thr Glu Arg Gln Leu Arg Glu Cys Thr Arg Trp Ala Asp Ala Glu Cys
 145 150 155 160

Glu Glu Ile Pro Gly Arg Trp Ile Thr Arg Ser Thr Pro Pro Glu Gly
 165 170 175

Ser Asp Ser Thr Ala Pro Ser Thr Gln Glu Pro Glu Ala Pro Pro Glu
 180 185 190

Gln Asp Leu Ile Ala Ser Thr Val Ala Gly Val Val Thr Thr Val Met
 195 200 205

Gly Ser Ser Gln Pro Val Val Thr Arg Gly Thr Thr Asp Asn Leu Ile
 210 215 220

Pro Val Tyr Cys Ser Ile Leu Ala Ala Val Val Val Gly Leu Val Ala
 225 230 235 240

Tyr Ile Ala Phe Lys Arg Trp Asn Ser Cys Lys Gln Asn Lys Gln Gly
 245 250 255

Ala Asn Ser Arg Pro Val Asn Gln Thr Pro Pro Pro Glu Gly Glu Lys
 260 265 270

Leu His Ser Asp Ser Gly Ile Ser Val Asp Ser Gln Ser Leu His Asp
 275 280 285

Gln Gln Pro His Thr Gln Thr Ala Ser Gly Gln Ala Leu Lys Gly Asp
 290 295 300

Gly Gly Leu Tyr Ser Ser Leu Pro Pro Ala Lys Arg Glu Glu Val Glu
 305 310 315 320

Lys Leu Leu Asn Gly Ser Ala Gly Asp Thr Trp Arg His Leu Ala Gly
 325 330 335

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Glu Leu Gly Tyr Gln Pro Glu His Ile Asp Ser Phe Thr His Glu Ala
 340 345 350

Cys Pro Val Arg Ala Leu Leu Ala Ser Trp Ala Thr Gln Asp Ser Ala
 355 360 365

Thr Leu Asp Ala Leu Leu Ala Ala Leu Arg Arg Ile Gln Arg Ala Asp
 370 375 380

Leu Val Glu Ser Leu Cys Ser Glu Ser Thr Ala Thr Ser Pro Val
 385 390 395

<210> SEQ ID NO 18
 <211> LENGTH: 16
 <212> TYPE: PRT
 <213> ORGANISM: Homo sapiens

<400> SEQUENCE: 18

Gln Thr Ala Ser Gly Gln Ala Leu Lys Gly Asp Gly Gly Leu Tyr Ser
 1 5 10 15

<210> SEQ ID NO 19
 <211> LENGTH: 1725
 <212> TYPE: PRT
 <213> ORGANISM: Homo sapiens

<400> SEQUENCE: 19

Lys Pro Arg Leu Leu Leu Phe Ser Pro Ser Val Val His Leu Gly Val
 1 5 10 15

Pro Leu Ser Val Gly Val Gln Leu Gln Asp Val Pro Arg Gly Gln Val
 20 25 30

Val Lys Gly Ser Val Phe Leu Arg Asn Pro Ser Arg Asn Asn Val Pro
 35 40 45

Cys Ser Pro Lys Val Asp Phe Thr Leu Ser Ser Glu Arg Asp Phe Ala
 50 55 60

Leu Leu Ser Leu Gln Val Pro Leu Lys Asp Ala Lys Ser Cys Gly Leu
 65 70 75 80

His Gln Leu Leu Arg Gly Pro Glu Val Gln Leu Val Ala His Ser Pro
 85 90 95

Trp Leu Lys Asp Ser Leu Ser Arg Thr Thr Asn Ile Gln Gly Ile Asn
 100 105 110

Leu Leu Phe Ser Ser Arg Arg Gly His Leu Phe Leu Gln Thr Asp Gln
 115 120 125

Pro Ile Tyr Asn Pro Gly Gln Arg Val Arg Tyr Arg Val Phe Ala Leu
 130 135 140

Asp Gln Lys Met Arg Pro Ser Thr Asp Thr Ile Thr Val Met Val Glu
 145 150 155 160

Asn Ser His Gly Leu Arg Val Arg Lys Lys Glu Val Tyr Met Pro Ser
 165 170 175

Ser Ile Phe Gln Asp Asp Phe Val Ile Pro Asp Ile Ser Glu Pro Gly
 180 185 190

Thr Trp Lys Ile Ser Ala Arg Phe Ser Asp Gly Leu Glu Ser Asn Ser
 195 200 205

Ser Thr Gln Phe Glu Val Lys Lys Tyr Val Leu Pro Asn Phe Glu Val
 210 215 220

Lys Ile Thr Pro Gly Lys Pro Tyr Ile Leu Thr Val Pro Gly His Leu
 225 230 235 240

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Asp	Glu	Met	Gln	Leu	Asp	Ile	Gln	Ala	Arg	Tyr	Ile	Tyr	Gly	Lys	Pro	245	250	255
Val	Gln	Gly	Val	Ala	Tyr	Val	Arg	Phe	Gly	Leu	Leu	Asp	Glu	Asp	Gly	260	265	270
Lys	Lys	Thr	Phe	Phe	Arg	Gly	Leu	Glu	Ser	Gln	Thr	Lys	Leu	Val	Asn	275	280	285
Gly	Gln	Ser	His	Ile	Ser	Leu	Ser	Lys	Ala	Glu	Phe	Gln	Asp	Ala	Leu	290	295	300
Glu	Lys	Leu	Asn	Met	Gly	Ile	Thr	Asp	Leu	Gln	Gly	Leu	Arg	Leu	Tyr	305	310	315
Val	Ala	Ala	Ala	Ile	Ile	Glu	Ser	Pro	Gly	Gly	Glu	Met	Glu	Glu	Ala	325	330	335
Glu	Leu	Thr	Ser	Trp	Tyr	Phe	Val	Ser	Ser	Pro	Phe	Ser	Leu	Asp	Leu	340	345	350
Ser	Lys	Thr	Lys	Arg	His	Leu	Val	Pro	Gly	Ala	Pro	Phe	Leu	Leu	Gln	355	360	365
Ala	Leu	Val	Arg	Glu	Met	Ser	Gly	Ser	Pro	Ala	Ser	Gly	Ile	Pro	Val	370	375	380
Lys	Val	Ser	Ala	Thr	Val	Ser	Ser	Pro	Gly	Ser	Val	Pro	Glu	Val	Gln	385	390	395
Asp	Ile	Gln	Gln	Asn	Thr	Asp	Gly	Ser	Gly	Gln	Val	Ser	Ile	Pro	Ile	405	410	415
Ile	Ile	Pro	Gln	Thr	Ile	Ser	Glu	Leu	Gln	Leu	Ser	Val	Ser	Ala	Gly	420	425	430
Ser	Pro	His	Pro	Ala	Ile	Ala	Arg	Leu	Thr	Val	Ala	Ala	Pro	Pro	Ser	435	440	445
Gly	Gly	Pro	Gly	Phe	Leu	Ser	Ile	Glu	Arg	Pro	Asp	Ser	Arg	Pro	Pro	450	455	460
Arg	Val	Gly	Asp	Thr	Leu	Asn	Leu	Asn	Leu	Arg	Ala	Val	Gly	Ser	Gly	465	470	475
Ala	Thr	Phe	Ser	His	Tyr	Tyr	Tyr	Met	Ile	Leu	Ser	Arg	Gly	Gln	Ile	485	490	495
Val	Phe	Met	Asn	Arg	Glu	Pro	Lys	Arg	Thr	Leu	Thr	Ser	Val	Ser	Val	500	505	510
Phe	Val	Asp	His	His	Leu	Ala	Pro	Ser	Phe	Tyr	Phe	Val	Ala	Phe	Tyr	515	520	525
Tyr	His	Gly	Asp	His	Pro	Val	Ala	Asn	Ser	Leu	Arg	Val	Asp	Val	Gln	530	535	540
Ala	Gly	Ala	Cys	Glu	Gly	Lys	Leu	Glu	Leu	Ser	Val	Asp	Gly	Ala	Lys	545	550	555
Gln	Tyr	Arg	Asn	Gly	Glu	Ser	Val	Lys	Leu	His	Leu	Glu	Thr	Asp	Ser	565	570	575
Leu	Ala	Leu	Val	Ala	Leu	Gly	Ala	Leu	Asp	Thr	Ala	Leu	Tyr	Ala	Ala	580	585	590
Gly	Ser	Lys	Ser	His	Lys	Pro	Leu	Asn	Met	Gly	Lys	Val	Phe	Glu	Ala	595	600	605
Met	Asn	Ser	Tyr	Asp	Leu	Gly	Cys	Gly	Pro	Gly	Gly	Gly	Asp	Ser	Ala	610	615	620
Leu	Gln	Val	Phe	Gln	Ala	Ala	Gly	Leu	Ala	Phe	Ser	Asp	Gly	Asp	Gln	625	630	635
																		640

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Trp	Thr	Leu	Ser	Arg	Lys	Arg	Leu	Ser	Cys	Pro	Lys	Glu	Lys	Thr	Thr	645	650	655
Arg	Lys	Lys	Arg	Asn	Val	Asn	Phe	Gln	Lys	Ala	Ile	Asn	Glu	Lys	Leu	660	665	670
Gly	Gln	Tyr	Ala	Ser	Pro	Thr	Ala	Lys	Arg	Cys	Cys	Gln	Asp	Gly	Val	675	680	685
Thr	Arg	Leu	Pro	Met	Met	Arg	Ser	Cys	Glu	Gln	Arg	Ala	Ala	Arg	Val	690	695	700
Gln	Gln	Pro	Asp	Cys	Arg	Glu	Pro	Phe	Leu	Ser	Cys	Cys	Gln	Phe	Ala	705	710	715
Glu	Ser	Leu	Arg	Lys	Lys	Ser	Arg	Asp	Lys	Gly	Gln	Ala	Gly	Leu	Gln	725	730	735
Arg	Ala	Leu	Glu	Ile	Leu	Gln	Glu	Asp	Leu	Ile	Asp	Glu	Asp	Asp		740	745	750
Ile	Pro	Val	Arg	Ser	Phe	Phe	Pro	Glu	Asn	Trp	Leu	Trp	Arg	Val	Glu	755	760	765
Thr	Val	Asp	Arg	Phe	Gln	Ile	Leu	Thr	Leu	Trp	Leu	Pro	Asp	Ser	Leu	770	775	780
Thr	Thr	Trp	Glu	Ile	His	Gly	Leu	Ser	Leu	Ser	Lys	Thr	Lys	Gly	Leu	785	790	795
Cys	Val	Ala	Thr	Pro	Val	Gln	Leu	Arg	Val	Phe	Arg	Glu	Phe	His	Leu	805	810	815
His	Leu	Arg	Leu	Pro	Met	Ser	Val	Arg	Arg	Phe	Glu	Gln	Leu	Glu	Leu	820	825	830
Arg	Pro	Val	Leu	Tyr	Asn	Tyr	Leu	Asp	Lys	Asn	Leu	Thr	Val	Ser	Val	835	840	845
His	Val	Ser	Pro	Val	Glu	Gly	Leu	Cys	Leu	Ala	Gly	Gly	Gly	Gly	Leu	850	855	860
Ala	Gln	Gln	Val	Leu	Val	Pro	Ala	Gly	Ser	Ala	Arg	Pro	Val	Ala	Phe	865	870	875
Ser	Val	Val	Pro	Thr	Ala	Ala	Ala	Ala	Val	Ser	Leu	Lys	Val	Val	Ala	885	890	895
Arg	Gly	Ser	Phe	Glu	Phe	Pro	Val	Gly	Asp	Ala	Val	Ser	Lys	Val	Leu	900	905	910
Gln	Ile	Glu	Lys	Glu	Gly	Ala	Ile	His	Arg	Glu	Glu	Leu	Val	Tyr	Glu	915	920	925
Leu	Asn	Pro	Leu	Asp	His	Arg	Gly	Arg	Thr	Leu	Glu	Ile	Pro	Gly	Asn	930	935	940
Ser	Asp	Pro	Asn	Met	Ile	Pro	Asp	Gly	Asp	Phe	Asn	Ser	Tyr	Val	Arg	945	950	955
Val	Thr	Ala	Ser	Asp	Pro	Leu	Asp	Thr	Leu	Gly	Ser	Glu	Gly	Ala	Leu	965	970	975
Ser	Pro	Gly	Gly	Val	Ala	Ser	Leu	Leu	Arg	Leu	Pro	Arg	Gly	Cys	Gly	980	985	990
Glu	Gln	Thr	Met	Ile	Tyr	Leu	Ala	Pro	Thr	Leu	Ala	Ala	Ser	Arg	Tyr	995	1000	1005
Leu	Asp	Lys	Thr	Glu	Gln	Trp	Ser	Thr	Leu	Pro	Pro	Glu	Thr	Lys		1010	1015	1020
Asp	His	Ala	Val	Asp	Leu	Ile	Gln	Lys	Gly	Tyr	Met	Arg	Ile	Gln		1025	1030	1035
Gln	Phe	Arg	Lys	Ala	Asp	Gly	Ser	Tyr	Ala	Ala	Trp	Leu	Ser	Arg				

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1040	1045	1050
Asp Ser Ser Thr Trp Leu Thr	Ala Phe Val Leu Lys	Val Leu Ser
1055	1060	1065
Leu Ala Gln Glu Gln Val Gly	Gly Ser Pro Glu Lys	Leu Gln Glu
1070	1075	1080
Thr Ser Asn Trp Leu Leu Ser	Gln Gln Gln Ala Asp	Gly Ser Phe
1085	1090	1095
Gln Asp Pro Cys Pro Val Leu	Asp Arg Ser Met Gln	Gly Gly Leu
1100	1105	1110
Val Gly Asn Asp Glu Thr Val	Ala Leu Thr Ala Phe	Val Thr Ile
1115	1120	1125
Ala Leu His His Gly Leu Ala	Val Phe Gln Asp Glu	Gly Ala Glu
1130	1135	1140
Pro Leu Lys Gln Arg Val Glu	Ala Ser Ile Ser Lys	Ala Asn Ser
1145	1150	1155
Phe Leu Gly Glu Lys Ala Ser	Ala Gly Leu Leu Gly	Ala His Ala
1160	1165	1170
Ala Ala Ile Thr Ala Tyr Ala	Leu Ser Leu Thr Lys	Ala Pro Val
1175	1180	1185
Asp Leu Leu Gly Val Ala His	Asn Asn Leu Met Ala	Met Ala Gln
1190	1195	1200
Glu Thr Gly Asp Asn Leu Tyr	Trp Gly Ser Val Thr	Gly Ser Gln
1205	1210	1215
Ser Asn Ala Val Ser Pro Thr	Pro Ala Pro Arg Asn	Pro Ser Asp
1220	1225	1230
Pro Met Pro Gln Ala Pro Ala	Leu Trp Ile Glu Thr	Thr Ala Tyr
1235	1240	1245
Ala Leu Leu His Leu Leu Leu	His Glu Gly Lys Ala	Glu Met Ala
1250	1255	1260
Asp Gln Ala Ser Ala Trp Leu	Thr Arg Gln Gly Ser	Phe Gln Gly
1265	1270	1275
Gly Phe Arg Ser Thr Gln Asp	Thr Val Ile Ala Leu	Asp Ala Leu
1280	1285	1290
Ser Ala Tyr Trp Ile Ala Ser	His Thr Thr Glu Glu	Arg Gly Leu
1295	1300	1305
Asn Val Thr Leu Ser Ser Thr	Gly Arg Asn Gly Phe	Lys Ser His
1310	1315	1320
Ala Leu Gln Leu Asn Asn Arg	Gln Ile Arg Gly Leu	Glu Glu Glu
1325	1330	1335
Leu Gln Phe Ser Leu Gly Ser	Lys Ile Asn Val Lys	Val Gly Gly
1340	1345	1350
Asn Ser Lys Gly Thr Leu Lys	Val Leu Arg Thr Tyr	Asn Val Leu
1355	1360	1365
Asp Met Lys Asn Thr Thr Cys	Gln Asp Leu Gln Ile	Glu Val Thr
1370	1375	1380
Val Lys Gly His Val Glu Tyr	Thr Met Glu Ala Asn	Glu Asp Tyr
1385	1390	1395
Glu Asp Tyr Glu Tyr Asp Glu	Leu Pro Ala Lys Asp	Asp Pro Asp
1400	1405	1410
Ala Pro Leu Gln Pro Val Thr	Pro Leu Gln Leu Phe	Glu Gly Arg
1415	1420	1425

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<210> SEQ ID NO 20
<211> LENGTH: 16
<212> TYPE: PRT
<213> ORGANISM: Homo sapiens

<400> SEQUENCE: 20
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<210> SEQ ID NO 21
<211> LENGTH: 1725
<212> TYPE: PRT
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<213> ORGANISM: Homo sapiens

<400> SEQUENCE: 21

Lys Pro Arg Leu Leu Leu Phe Ser Pro Ser Val Val His Leu Gly Val
1 5 10 15
Pro Leu Ser Val Gly Val Gln Leu Gln Asp Val Pro Arg Gly Gln Val
20 25 30
Val Lys Gly Ser Val Phe Leu Arg Asn Pro Ser Arg Asn Asn Val Pro
35 40 45
Cys Ser Pro Lys Val Asp Phe Thr Leu Ser Ser Glu Arg Asp Phe Ala
50 55 60
Leu Leu Ser Leu Gln Val Pro Leu Lys Asp Ala Lys Ser Cys Gly Leu
65 70 75 80
His Gln Leu Leu Arg Gly Pro Glu Val Gln Leu Val Ala His Ser Pro
85 90 95
Trp Leu Lys Asp Ser Leu Ser Arg Thr Thr Asn Ile Gln Gly Ile Asn
100 105 110
Leu Leu Phe Ser Ser Arg Arg Gly His Leu Phe Leu Gln Thr Asp Gln
115 120 125
Pro Ile Tyr Asn Pro Gly Gln Arg Val Arg Tyr Arg Val Phe Ala Leu
130 135 140
Asp Gln Lys Met Arg Pro Ser Thr Asp Thr Ile Thr Val Met Val Glu
145 150 155 160
Asn Ser His Gly Leu Arg Val Arg Lys Lys Glu Val Tyr Met Pro Ser
165 170 175
Ser Ile Phe Gln Asp Asp Phe Val Ile Pro Asp Ile Ser Glu Pro Gly
180 185 190
Thr Trp Lys Ile Ser Ala Arg Phe Ser Asp Gly Leu Glu Ser Asn Ser
195 200 205
Ser Thr Gln Phe Glu Val Lys Lys Tyr Val Leu Pro Asn Phe Glu Val
210 215 220
Lys Ile Thr Pro Gly Lys Pro Tyr Ile Leu Thr Val Pro Gly His Leu
225 230 235 240
Asp Glu Met Gln Leu Asp Ile Gln Ala Arg Tyr Ile Tyr Gly Lys Pro
245 250 255
Val Gln Gly Val Ala Tyr Val Arg Phe Gly Leu Leu Asp Glu Asp Gly
260 265 270
Lys Lys Thr Phe Phe Arg Gly Leu Glu Ser Gln Thr Lys Leu Val Asn
275 280 285
Gly Gln Ser His Ile Ser Leu Ser Lys Ala Glu Phe Gln Asp Ala Leu
290 295 300
Glu Lys Leu Asn Met Gly Ile Thr Asp Leu Gln Gly Leu Arg Leu Tyr
305 310 315 320
Val Ala Ala Ala Ile Ile Glu Ser Pro Gly Gly Glu Met Glu Glu Ala
325 330 335
Glu Leu Thr Ser Trp Tyr Phe Val Ser Ser Pro Phe Ser Leu Asp Leu
340 345 350
Ser Lys Thr Lys Arg His Leu Val Pro Gly Ala Pro Phe Leu Leu Gln
355 360 365
Ala Leu Val Arg Glu Met Ser Gly Ser Pro Ala Ser Gly Ile Pro Val
370 375 380

-continued

Lys	Val	Ser	Ala	Thr	Val	Ser	Ser	Pro	Gly	Ser	Val	Pro	Glu	Val	Gln
385					390					395					400
Asp	Ile	Gln	Gln	Asn	Thr	Asp	Gly	Ser	Gly	Gln	Val	Ser	Ile	Pro	Ile
				405					410					415	
Ile	Ile	Pro	Gln	Thr	Ile	Ser	Glu	Leu	Gln	Leu	Ser	Val	Ser	Ala	Gly
				420				425						430	
Ser	Pro	His	Pro	Ala	Ile	Ala	Arg	Leu	Thr	Val	Ala	Ala	Pro	Pro	Ser
		435					440					445			
Gly	Gly	Pro	Gly	Phe	Leu	Ser	Ile	Glu	Arg	Pro	Asp	Ser	Arg	Pro	Pro
	450					455					460				
Arg	Val	Gly	Asp	Thr	Leu	Asn	Leu	Asn	Leu	Arg	Ala	Val	Gly	Ser	Gly
465					470					475					480
Ala	Thr	Phe	Ser	His	Tyr	Tyr	Tyr	Met	Ile	Leu	Ser	Arg	Gly	Gln	Ile
				485					490					495	
Val	Phe	Met	Asn	Arg	Glu	Pro	Lys	Arg	Thr	Leu	Thr	Ser	Val	Ser	Val
			500					505					510		
Phe	Val	Asp	His	His	Leu	Ala	Pro	Ser	Phe	Tyr	Phe	Val	Ala	Phe	Tyr
		515					520					525			
Tyr	His	Gly	Asp	His	Pro	Val	Ala	Asn	Ser	Leu	Arg	Val	Asp	Val	Gln
	530					535					540				
Ala	Gly	Ala	Cys	Glu	Gly	Lys	Leu	Glu	Leu	Ser	Val	Asp	Gly	Ala	Lys
545					550					555					560
Gln	Tyr	Arg	Asn	Gly	Glu	Ser	Val	Lys	Leu	His	Leu	Glu	Thr	Asp	Ser
				565					570					575	
Leu	Ala	Leu	Val	Ala	Leu	Gly	Ala	Leu	Asp	Thr	Ala	Leu	Tyr	Ala	Ala
			580					585					590		
Gly	Ser	Lys	Ser	His	Lys	Pro	Leu	Asn	Met	Gly	Lys	Val	Phe	Glu	Ala
		595					600					605			
Met	Asn	Ser	Tyr	Asp	Leu	Gly	Cys	Gly	Pro	Gly	Gly	Gly	Asp	Ser	Ala
	610					615					620				
Leu	Gln	Val	Phe	Gln	Ala	Ala	Gly	Leu	Ala	Phe	Ser	Asp	Gly	Asp	Gln
625					630					635					640
Trp	Thr	Leu	Ser	Arg	Lys	Arg	Leu	Ser	Cys	Pro	Lys	Glu	Lys	Thr	Thr
				645					650					655	
Arg	Lys	Lys	Arg	Asn	Val	Asn	Phe	Gln	Lys	Ala	Ile	Asn	Glu	Lys	Leu
			660					665					670		
Gly	Gln	Tyr	Ala	Ser	Pro	Thr	Ala	Lys	Arg	Cys	Cys	Gln	Asp	Gly	Val
		675					680					685			
Thr	Arg	Leu	Pro	Met	Met	Arg	Ser	Cys	Glu	Gln	Arg	Ala	Ala	Arg	Val
	690					695					700				
Gln	Gln	Pro	Asp	Cys	Arg	Glu	Pro	Phe	Leu	Ser	Cys	Cys	Gln	Phe	Ala
705					710					715					720
Glu	Ser	Leu	Arg	Lys	Lys	Ser	Arg	Asp	Lys	Gly	Gln	Ala	Gly	Leu	Gln
				725					730					735	
Arg	Ala	Leu	Glu	Ile	Leu	Gln	Glu	Asp	Leu	Ile	Asp	Glu	Asp	Asp	
			740					745				750			
Ile	Pro	Val	Arg	Ser	Phe	Phe	Pro	Glu	Asn	Trp	Leu	Trp	Arg	Val	Glu
		755					760					765			
Thr	Val	Asp	Arg	Phe	Gln	Ile	Leu	Thr	Leu	Trp	Leu	Pro	Asp	Ser	Leu
	770					775					780				
Thr	Thr	Trp	Glu	Ile	His	Gly	Leu	Ser	Leu	Ser	Lys	Thr	Lys	Gly	Leu

-continued

785	790	795	800
Cys Val Ala Thr Pro Val Gln Leu Arg Val Phe Arg Glu Phe His Leu	805	810	815
His Leu Arg Leu Pro Met Ser Val Arg Arg Phe Glu Gln Leu Glu Leu	820	825	830
Arg Pro Val Leu Tyr Asn Tyr Leu Asp Lys Asn Leu Thr Val Ser Val	835	840	845
His Val Ser Pro Val Glu Gly Leu Cys Leu Ala Gly Gly Gly Gly Leu	850	855	860
Ala Gln Gln Val Leu Val Pro Ala Gly Ser Ala Arg Pro Val Ala Phe	865	870	875
Ser Val Val Pro Thr Ala Ala Ala Ala Val Ser Leu Lys Val Val Ala	885	890	895
Arg Gly Ser Phe Glu Phe Pro Val Gly Asp Ala Val Ser Lys Val Leu	900	905	910
Gln Ile Glu Lys Glu Gly Ala Ile His Arg Glu Glu Leu Val Tyr Glu	915	920	925
Leu Asn Pro Leu Asp His Arg Gly Arg Thr Leu Glu Ile Pro Gly Asn	930	935	940
Ser Asp Pro Asn Met Ile Pro Asp Gly Asp Phe Asn Ser Tyr Val Arg	945	950	955
Val Thr Ala Ser Asp Pro Leu Asp Thr Leu Gly Ser Glu Gly Ala Leu	965	970	975
Ser Pro Gly Gly Val Ala Ser Leu Leu Arg Leu Pro Arg Gly Cys Gly	980	985	990
Glu Gln Thr Met Ile Tyr Leu Ala Pro Thr Leu Ala Ala Ser Arg Tyr	995	1000	1005
Leu Asp Lys Thr Glu Gln Trp Ser Thr Leu Pro Pro Glu Thr Lys	1010	1015	1020
Asp His Ala Val Asp Leu Ile Gln Lys Gly Tyr Met Arg Ile Gln	1025	1030	1035
Gln Phe Arg Lys Ala Asp Gly Ser Tyr Ala Ala Trp Leu Ser Arg	1040	1045	1050
Asp Ser Ser Thr Trp Leu Thr Ala Phe Val Leu Lys Val Leu Ser	1055	1060	1065
Leu Ala Gln Glu Gln Val Gly Gly Ser Pro Glu Lys Leu Gln Glu	1070	1075	1080
Thr Ser Asn Trp Leu Leu Ser Gln Gln Gln Ala Asp Gly Ser Phe	1085	1090	1095
Gln Asp Leu Ser Pro Val Ile His Arg Ser Met Gln Gly Gly Leu	1100	1105	1110
Val Gly Asn Asp Glu Thr Val Ala Leu Thr Ala Phe Val Thr Ile	1115	1120	1125
Ala Leu His His Gly Leu Ala Val Phe Gln Asp Glu Gly Ala Glu	1130	1135	1140
Pro Leu Lys Gln Arg Val Glu Ala Ser Ile Ser Lys Ala Asn Ser	1145	1150	1155
Phe Leu Gly Glu Lys Ala Ser Ala Gly Leu Leu Gly Ala His Ala	1160	1165	1170
Ala Ala Ile Thr Ala Tyr Ala Leu Ser Leu Thr Lys Ala Pro Val	1175	1180	1185

-continued

Asp	Leu	Leu	Gly	Val	Ala	His	Asn	Asn	Leu	Met	Ala	Met	Ala	Gln
1190						1195					1200			
Glu	Thr	Gly	Asp	Asn	Leu	Tyr	Trp	Gly	Ser	Val	Thr	Gly	Ser	Gln
1205						1210					1215			
Ser	Asn	Ala	Val	Ser	Pro	Thr	Pro	Ala	Pro	Arg	Asn	Pro	Ser	Asp
1220						1225					1230			
Pro	Met	Pro	Gln	Ala	Pro	Ala	Leu	Trp	Ile	Glu	Thr	Thr	Ala	Tyr
1235						1240					1245			
Ala	Leu	Leu	His	Leu	Leu	Leu	His	Glu	Gly	Lys	Ala	Glu	Met	Ala
1250						1255					1260			
Asp	Gln	Ala	Ser	Ala	Trp	Leu	Thr	Arg	Gln	Gly	Ser	Phe	Gln	Gly
1265						1270					1275			
Gly	Phe	Arg	Ser	Thr	Gln	Asp	Thr	Val	Ile	Ala	Leu	Asp	Ala	Leu
1280						1285					1290			
Ser	Ala	Tyr	Trp	Ile	Ala	Ser	His	Thr	Thr	Glu	Glu	Arg	Gly	Leu
1295						1300					1305			
Asn	Val	Thr	Leu	Ser	Ser	Thr	Gly	Arg	Asn	Gly	Phe	Lys	Ser	His
1310						1315					1320			
Ala	Leu	Gln	Leu	Asn	Asn	Arg	Gln	Ile	Arg	Gly	Leu	Glu	Glu	Glu
1325						1330					1335			
Leu	Gln	Phe	Ser	Leu	Gly	Ser	Lys	Ile	Asn	Val	Lys	Val	Gly	Gly
1340						1345					1350			
Asn	Ser	Lys	Gly	Thr	Leu	Lys	Val	Leu	Arg	Thr	Tyr	Asn	Val	Leu
1355						1360					1365			
Asp	Met	Lys	Asn	Thr	Thr	Cys	Gln	Asp	Leu	Gln	Ile	Glu	Val	Thr
1370						1375					1380			
Val	Lys	Gly	His	Val	Glu	Tyr	Thr	Met	Glu	Ala	Asn	Glu	Asp	Tyr
1385						1390					1395			
Glu	Asp	Tyr	Glu	Tyr	Asp	Glu	Leu	Pro	Ala	Lys	Asp	Asp	Pro	Asp
1400						1405					1410			
Ala	Pro	Leu	Gln	Pro	Val	Thr	Pro	Leu	Gln	Leu	Phe	Glu	Gly	Arg
1415						1420					1425			
Arg	Asn	Arg	Arg	Arg	Arg	Glu	Ala	Pro	Lys	Val	Val	Glu	Glu	Gln
1430						1435					1440			
Glu	Ser	Arg	Val	His	Tyr	Thr	Val	Cys	Ile	Trp	Arg	Asn	Gly	Lys
1445						1450					1455			
Val	Gly	Leu	Ser	Gly	Met	Ala	Ile	Ala	Asp	Val	Thr	Leu	Leu	Ser
1460						1465					1470			
Gly	Phe	His	Ala	Leu	Arg	Ala	Asp	Leu	Glu	Lys	Leu	Thr	Ser	Leu
1475						1480					1485			
Ser	Asp	Arg	Tyr	Val	Ser	His	Phe	Glu	Thr	Glu	Gly	Pro	His	Val
1490						1495					1500			
Leu	Leu	Tyr	Phe	Asp	Ser	Val	Pro	Thr	Ser	Arg	Glu	Cys	Val	Gly
1505						1510					1515			
Phe	Glu	Ala	Val	Gln	Glu	Val	Pro	Val	Gly	Leu	Val	Gln	Pro	Ala
1520						1525					1530			
Ser	Ala	Thr	Leu	Tyr	Asp	Tyr	Tyr	Asn	Pro	Glu	Arg	Arg	Cys	Ser
1535						1540					1545			
Val	Phe	Tyr	Gly	Ala	Pro	Ser	Lys	Ser	Arg	Leu	Leu	Ala	Thr	Leu
1550						1555					1560			


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<210> SEQ ID NO 22
<211> LENGTH: 17
<212> TYPE: PRT
<213> ORGANISM: Homo sapiens
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<400> SEQUENCE: 22

Ala Pro Leu Gln Pro Val Thr Pro Leu Gln Leu Phe Glu Gly Arg Arg
1 5 10 15

Asn

```
<210> SEQ ID NO 23
<211> LENGTH: 831
<212> TYPE: PRT
<213> ORGANISM: Homo sapiens
```

<400> SEQUENCE: 23

Gly Pro Arg Val Val Glu Arg His Gln Ser Ala Cys Lys Asp Ser Asp
1 5 10 15

Trp Pro Phe Cys Ser Asp Glu Asp Trp Asn Tyr Lys Cys Pro Ser Gly
20 25 30

Cys Arg Met Lys Gly Leu Ile Asp Glu Val Asn Gln Asp Phe Thr Asn
35 40 45

Arg Ile Asn Lys Leu Lys Asn Ser Leu Phe Glu Tyr Gln Lys Asn Asn
50 55 60

Lys Asp Ser His Ser Leu Thr Thr Asn Ile Met Glu Ile Leu Arg Gly
65 70 75 80

Asp Phe Ser Ser Ala Asn Asn Arg Asp Asn Thr Tyr Asn Arg Val Ser
85 90 95

Glu Asp Leu Arg Ser Arg Ile Glu Val Leu Lys Arg Lys Val Ile Glu
100 105 110

Lys Val Gln His Ile Gln Leu Leu Gln Lys Asn Val Arg Ala Gln Leu

-continued

115						120						125					
Val	Asp	Met	Lys	Arg	Leu	Glu	Val	Asp	Ile	Asp	Ile	Lys	Ile	Arg	Ser		
130						135					140						
Cys	Arg	Gly	Ser	Cys	Ser	Arg	Ala	Leu	Ala	Arg	Glu	Val	Asp	Leu	Lys		
145					150					155					160		
Asp	Tyr	Glu	Asp	Gln	Gln	Lys	Gln	Leu	Glu	Gln	Val	Ile	Ala	Lys	Asp		
				165					170					175			
Leu	Leu	Pro	Ser	Arg	Asp	Arg	Gln	His	Leu	Pro	Leu	Ile	Lys	Met	Lys		
			180					185					190				
Pro	Val	Pro	Asp	Leu	Val	Pro	Gly	Asn	Phe	Lys	Ser	Gln	Leu	Gln	Lys		
		195					200					205					
Val	Pro	Pro	Glu	Trp	Lys	Ala	Leu	Thr	Asp	Met	Pro	Gln	Met	Arg	Met		
	210					215					220						
Glu	Leu	Glu	Arg	Pro	Gly	Gly	Asn	Glu	Ile	Thr	Arg	Gly	Gly	Ser	Thr		
225					230					235					240		
Ser	Tyr	Gly	Thr	Gly	Ser	Glu	Thr	Glu	Ser	Pro	Arg	Asn	Pro	Ser	Ser		
				245					250					255			
Ala	Gly	Ser	Trp	Asn	Ser	Gly	Ser	Ser	Gly	Pro	Gly	Ser	Thr	Gly	Asn		
			260					265					270				
Arg	Asn	Pro	Gly	Ser	Ser	Gly	Thr	Gly	Gly	Thr	Ala	Thr	Trp	Lys	Pro		
		275					280					285					
Gly	Ser	Ser	Gly	Pro	Gly	Ser	Thr	Gly	Ser	Trp	Asn	Ser	Gly	Ser	Ser		
	290					295					300						
Gly	Thr	Gly	Ser	Thr	Gly	Asn	Gln	Asn	Pro	Gly	Ser	Pro	Arg	Pro	Gly		
305					310					315					320		
Ser	Thr	Gly	Thr	Trp	Asn	Pro	Gly	Ser	Ser	Glu	Arg	Gly	Ser	Ala	Gly		
				325					330					335			
His	Trp	Thr	Ser	Glu	Ser	Ser	Val	Ser	Gly	Ser	Thr	Gly	Gln	Trp	His		
			340					345					350				
Ser	Glu	Ser	Gly	Ser	Phe	Arg	Pro	Asp	Ser	Pro	Gly	Ser	Gly	Asn	Ala		
		355					360					365					
Arg	Pro	Asn	Asn	Pro	Asp	Trp	Gly	Thr	Phe	Glu	Glu	Val	Ser	Gly	Asn		
	370					375					380						
Val	Ser	Pro	Gly	Thr	Arg	Arg	Glu	Tyr	His	Thr	Glu	Lys	Leu	Val	Thr		
385					390					395					400		
Ser	Lys	Gly	Asp	Lys	Glu	Leu	Arg	Thr	Gly	Lys	Glu	Lys	Val	Thr	Ser		
				405					410					415			
Gly	Ser	Thr	Thr	Thr	Thr	Arg	Arg	Ser	Cys	Ser	Lys	Thr	Val	Thr	Lys		
				420				425					430				
Thr	Val	Ile	Gly	Pro	Asp	Gly	His	Lys	Glu	Val	Thr	Lys	Glu	Val	Val		
		435				440						445					
Thr	Ser	Glu	Asp	Gly	Ser	Asp	Cys	Pro	Glu	Ala	Met	Asp	Leu	Gly	Thr		
	450					455					460						
Leu	Ser	Gly	Ile	Gly	Thr	Leu	Asp	Gly	Phe	Arg	His	Arg	His	Pro	Asp		
465					470					475					480		
Glu	Ala	Ala	Phe	Phe	Asp	Thr	Ala	Ser	Thr	Gly	Lys	Thr	Phe	Pro	Gly		
				485					490					495			
Phe	Phe	Ser	Pro	Met	Leu	Gly	Glu	Phe	Val	Ser	Glu	Thr	Glu	Ser	Arg		
			500					505					510				
Gly	Ser	Glu	Ser	Gly	Ile	Phe	Thr	Asn	Thr	Lys	Glu	Ser	Ser	Ser	His		
	515						520					525					

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His	Pro	Gly	Ile	Ala	Glu	Phe	Pro	Ser	Arg	Gly	Lys	Ser	Ser	Ser	Tyr
530						535					540				
Ser	Lys	Gln	Phe	Thr	Ser	Ser	Thr	Ser	Tyr	Asn	Arg	Gly	Asp	Ser	Thr
545					550					555					560
Phe	Glu	Ser	Lys	Ser	Tyr	Lys	Met	Ala	Asp	Glu	Ala	Gly	Ser	Glu	Ala
				565					570					575	
Asp	His	Glu	Gly	Thr	His	Ser	Thr	Lys	Arg	Gly	His	Ala	Lys	Ser	Arg
		580						585					590		
Pro	Val	Arg	Asp	Cys	Asp	Asp	Val	Leu	Gln	Thr	His	Pro	Ser	Gly	Thr
		595					600					605			
Gln	Ser	Gly	Ile	Phe	Asn	Ile	Lys	Leu	Pro	Gly	Ser	Ser	Lys	Ile	Phe
610					615					620					
Ser	Val	Tyr	Cys	Asp	Gln	Glu	Thr	Ser	Leu	Gly	Gly	Trp	Leu	Leu	Ile
625					630					635					640
Gln	Gln	Arg	Met	Asp	Gly	Ser	Leu	Asn	Phe	Asn	Arg	Thr	Trp	Gln	Asp
			645						650					655	
Tyr	Lys	Arg	Gly	Phe	Gly	Ser	Leu	Asn	Asp	Glu	Gly	Glu	Gly	Glu	Phe
			660					665					670		
Trp	Leu	Gly	Asn	Asp	Tyr	Leu	His	Leu	Leu	Thr	Gln	Arg	Gly	Ser	Val
		675					680				685				
Leu	Arg	Val	Glu	Leu	Glu	Asp	Trp	Ala	Gly	Asn	Glu	Ala	Tyr	Ala	Glu
		690				695				700					
Tyr	His	Phe	Arg	Val	Gly	Ser	Glu	Ala	Glu	Gly	Tyr	Ala	Leu	Gln	Val
705					710					715					720
Ser	Ser	Tyr	Glu	Gly	Thr	Ala	Gly	Asp	Ala	Leu	Ile	Glu	Gly	Ser	Val
			725					730						735	
Glu	Glu	Gly	Ala	Glu	Tyr	Thr	Ser	His	Asn	Asn	Met	Gln	Phe	Ser	Thr
			740					745					750		
Phe	Asp	Arg	Asp	Ala	Asp	Gln	Trp	Glu	Glu	Asn	Cys	Ala	Glu	Val	Tyr
		755					760				765				
Gly	Gly	Gly	Trp	Trp	Tyr	Asn	Asn	Cys	Gln	Ala	Ala	Asn	Leu	Asn	Gly
		770				775				780					
Ile	Tyr	Tyr	Pro	Gly	Gly	Ser	Tyr	Asp	Pro	Arg	Asn	Asn	Ser	Pro	Tyr
785					790					795					800
Glu	Ile	Glu	Asn	Gly	Val	Val	Trp	Val	Ser	Phe	Arg	Gly	Ala	Asp	Tyr
			805					810					815		
Ser	Leu	Arg	Ala	Val	Arg	Met	Lys	Ile	Arg	Pro	Leu	Val	Thr	Gln	
		820					825						830		

<210> SEQ ID NO 24

<211> LENGTH: 23

<212> TYPE: PRT

<213> ORGANISM: Homo sapiens

<400> SEQUENCE: 24

Ser	Ser	Ser	Tyr	Ser	Lys	Gln	Phe	Thr	Ser	Ser	Thr	Ser	Tyr	Asn	Arg
1				5					10					15	

Gly	Asp	Ser	Thr	Phe	Glu	Ser
						20

<210> SEQ ID NO 25

<211> LENGTH: 609

<212> TYPE: PRT

-continued

<213> ORGANISM: Homo sapiens

<400> SEQUENCE: 25

Gly Pro Arg Val Val Glu Arg His Gln Ser Ala Cys Lys Asp Ser Asp
1 5 10 15
Trp Pro Phe Cys Ser Asp Glu Asp Trp Asn Tyr Lys Cys Pro Ser Gly
20 25 30
Cys Arg Met Lys Gly Leu Ile Asp Glu Val Asn Gln Asp Phe Thr Asn
35 40 45
Arg Ile Asn Lys Leu Lys Asn Ser Leu Phe Glu Tyr Gln Lys Asn Asn
50 55 60
Lys Asp Ser His Ser Leu Thr Thr Asn Ile Met Glu Ile Leu Arg Gly
65 70 75 80
Asp Phe Ser Ser Ala Asn Asn Arg Asp Asn Thr Tyr Asn Arg Val Ser
85 90 95
Glu Asp Leu Arg Ser Arg Ile Glu Val Leu Lys Arg Lys Val Ile Glu
100 105 110
Lys Val Gln His Ile Gln Leu Leu Gln Lys Asn Val Arg Ala Gln Leu
115 120 125
Val Asp Met Lys Arg Leu Glu Val Asp Ile Asp Ile Lys Ile Arg Ser
130 135 140
Cys Arg Gly Ser Cys Ser Arg Ala Leu Ala Arg Glu Val Asp Leu Lys
145 150 155 160
Asp Tyr Glu Asp Gln Gln Lys Gln Leu Glu Gln Val Ile Ala Lys Asp
165 170 175
Leu Leu Pro Ser Arg Asp Arg Gln His Leu Pro Leu Ile Lys Met Lys
180 185 190
Pro Val Pro Asp Leu Val Pro Gly Asn Phe Lys Ser Gln Leu Gln Lys
195 200 205
Val Pro Pro Glu Trp Lys Ala Leu Thr Asp Met Pro Gln Met Arg Met
210 215 220
Glu Leu Glu Arg Pro Gly Gly Asn Glu Ile Thr Arg Gly Gly Ser Thr
225 230 235 240
Ser Tyr Gly Thr Gly Ser Glu Thr Glu Ser Pro Arg Asn Pro Ser Ser
245 250 255
Ala Gly Ser Trp Asn Ser Gly Ser Ser Gly Pro Gly Ser Thr Gly Asn
260 265 270
Arg Asn Pro Gly Ser Ser Gly Thr Gly Gly Thr Ala Thr Trp Lys Pro
275 280 285
Gly Ser Ser Gly Pro Gly Ser Thr Gly Ser Trp Asn Ser Gly Ser Ser
290 295 300
Gly Thr Gly Ser Thr Gly Asn Gln Asn Pro Gly Ser Pro Arg Pro Gly
305 310 315 320
Ser Thr Gly Thr Trp Asn Pro Gly Ser Ser Glu Arg Gly Ser Ala Gly
325 330 335
His Trp Thr Ser Glu Ser Ser Val Ser Gly Ser Thr Gly Gln Trp His
340 345 350
Ser Glu Ser Gly Ser Phe Arg Pro Asp Ser Pro Gly Ser Gly Asn Ala
355 360 365
Arg Pro Asn Asn Pro Asp Trp Gly Thr Phe Glu Glu Val Ser Gly Asn
370 375 380

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Val	Ser	Pro	Gly	Thr	Arg	Arg	Glu	Tyr	His	Thr	Glu	Lys	Leu	Val	Thr
385					390					395					400
Ser	Lys	Gly	Asp	Lys	Glu	Leu	Arg	Thr	Gly	Lys	Glu	Lys	Val	Thr	Ser
			405						410					415	
Gly	Ser	Thr	Thr	Thr	Thr	Arg	Arg	Ser	Cys	Ser	Lys	Thr	Val	Thr	Lys
			420					425					430		
Thr	Val	Ile	Gly	Pro	Asp	Gly	His	Lys	Glu	Val	Thr	Lys	Glu	Val	Val
		435					440					445			
Thr	Ser	Glu	Asp	Gly	Ser	Asp	Cys	Pro	Glu	Ala	Met	Asp	Leu	Gly	Thr
		450				455					460				
Leu	Ser	Gly	Ile	Gly	Thr	Leu	Asp	Gly	Phe	Arg	His	Arg	His	Pro	Asp
465					470					475					480
Glu	Ala	Ala	Phe	Phe	Asp	Thr	Ala	Ser	Thr	Gly	Lys	Thr	Phe	Pro	Gly
				485					490					495	
Phe	Phe	Ser	Pro	Met	Leu	Gly	Glu	Phe	Val	Ser	Glu	Thr	Glu	Ser	Arg
			500					505					510		
Gly	Ser	Glu	Ser	Gly	Ile	Phe	Thr	Asn	Thr	Lys	Glu	Ser	Ser	Ser	His
		515				520						525			
His	Pro	Gly	Ile	Ala	Glu	Phe	Pro	Ser	Arg	Gly	Lys	Ser	Ser	Ser	Tyr
		530				535					540				
Ser	Lys	Gln	Phe	Thr	Ser	Ser	Thr	Ser	Tyr	Asn	Arg	Gly	Asp	Ser	Thr
545					550					555					560
Phe	Glu	Ser	Lys	Ser	Tyr	Lys	Met	Ala	Asp	Glu	Ala	Gly	Ser	Glu	Ala
				565					570					575	
Asp	His	Glu	Gly	Thr	His	Ser	Thr	Lys	Arg	Gly	His	Ala	Lys	Ser	Arg
			580					585					590		
Pro	Val	Arg	Gly	Ile	His	Thr	Ser	Pro	Leu	Gly	Lys	Pro	Ser	Leu	Ser
		595					600					605			

Pro

<210> SEQ ID NO 26
 <211> LENGTH: 25
 <212> TYPE: PRT
 <213> ORGANISM: Homo sapiens

<400> SEQUENCE: 26

Ser	Ser	Ser	Tyr	Ser	Lys	Gln	Phe	Thr	Ser	Ser	Thr	Ser	Tyr	Asn	Arg
1				5					10					15	
Gly	Asp	Ser	Thr	Phe	Glu	Ser	Lys	Ser							
			20					25							

<210> SEQ ID NO 27
 <211> LENGTH: 26
 <212> TYPE: PRT
 <213> ORGANISM: Homo sapiens

<400> SEQUENCE: 27

Ser	Ser	Ser	Tyr	Ser	Lys	Gln	Phe	Thr	Ser	Ser	Thr	Ser	Tyr	Asn	Arg
1				5					10					15	
Gly	Asp	Ser	Thr	Phe	Glu	Ser	Lys	Ser	Tyr						
			20					25							

1. A method of detecting Complement C3-derived peptide CO3 in a patient, said method comprising:

- a. obtaining a biological material from a human patient; and
- b. detecting presence or an amount of the Complement C3-derived peptide CO3 consisting of amino acid sequence of SEQ ID NO: 2 in the biological material by contacting the biological material with an antibody or aptamer that specifically binds to the peptide CO3 and detecting binding between the Complement C3-derived peptide CO3 and the antibody or aptamer.

2. The method of claim 1, wherein the biological material is serum, blood, plasma, cerebrospinal fluid, or urine.

3. A method of diagnosing and treating cognitive impairment in a patient, said method comprising:

- a. obtaining a biological material from a human patient;
- b. detecting an amount of Complement C3-derived peptide CO3 consisting of amino acid sequence of SEQ ID NO: 2 in the biological material by contacting the biological material with an antibody or aptamer that

specifically binds to the peptide CO3 and detecting binding between the peptide CO3 and the antibody or aptamer;

- c. diagnosing the patient with cognitive impairment when a higher amount of the Complement C3-derived peptide CO3 in the biological material is detected by comparing the amount of the Complement C3-derived peptide CO3 in the patient with an amount of the Complement C3-derived peptide CO3 in a biological material obtained from a non-psychiatry disease subject; and

- d. administering an effective amount of an anti-acetylcholine esterase inhibitor to the diagnosed patient.

4. The method of claim 3, wherein the biological material is serum, blood, plasma, cerebrospinal fluid, or urine.

5. The method of claim 3, wherein the cognitive impairment includes Alzheimer's dementia, mild cognitive impairment, Dementia with Lewy bodies, and frontotemporal dementia.

6. The method of claim 3, wherein the anti-acetylcholine esterase inhibitor is Donepezil-hydrochloride.

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