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(54) **METHODS OF DIAGNOSING ALZHEIMER'S DISEASE**

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(57) **ABSTRACT**

Methods and compositions relating to Alzheimer's disease are provided. Specifically, proteins that are differentially expressed in the Alzheimer's disease state relative to their expression in the normal state are provided. Proteins associated with Alzheimer's disease are identified and described. Methods of diagnosis of Alzheimer's disease using the differentially expressed proteins are also provided, as are methods for the identification and therapeutic use of compounds for the prevention and treatment of Alzheimer's disease.

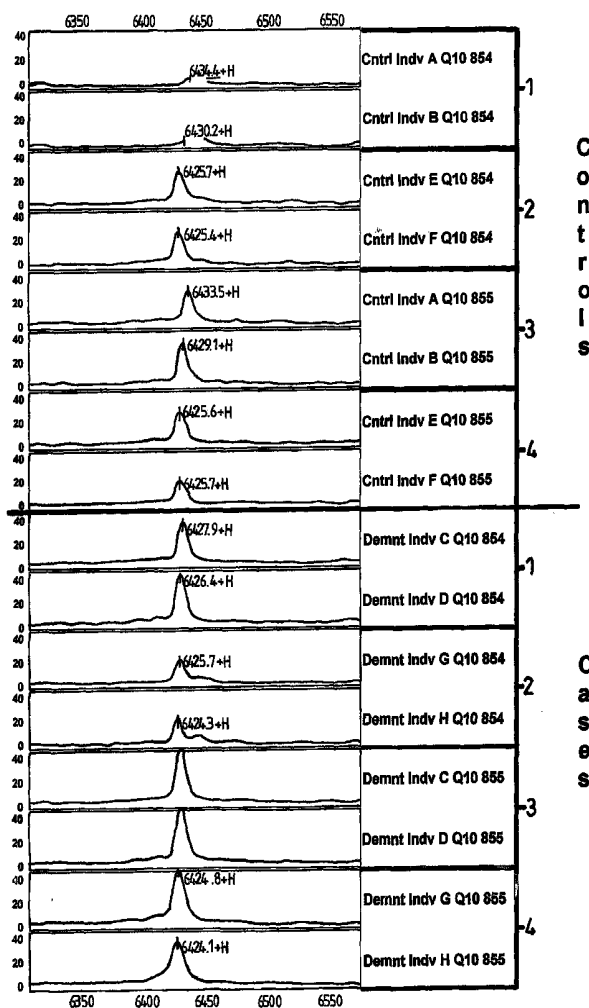


Figure 1

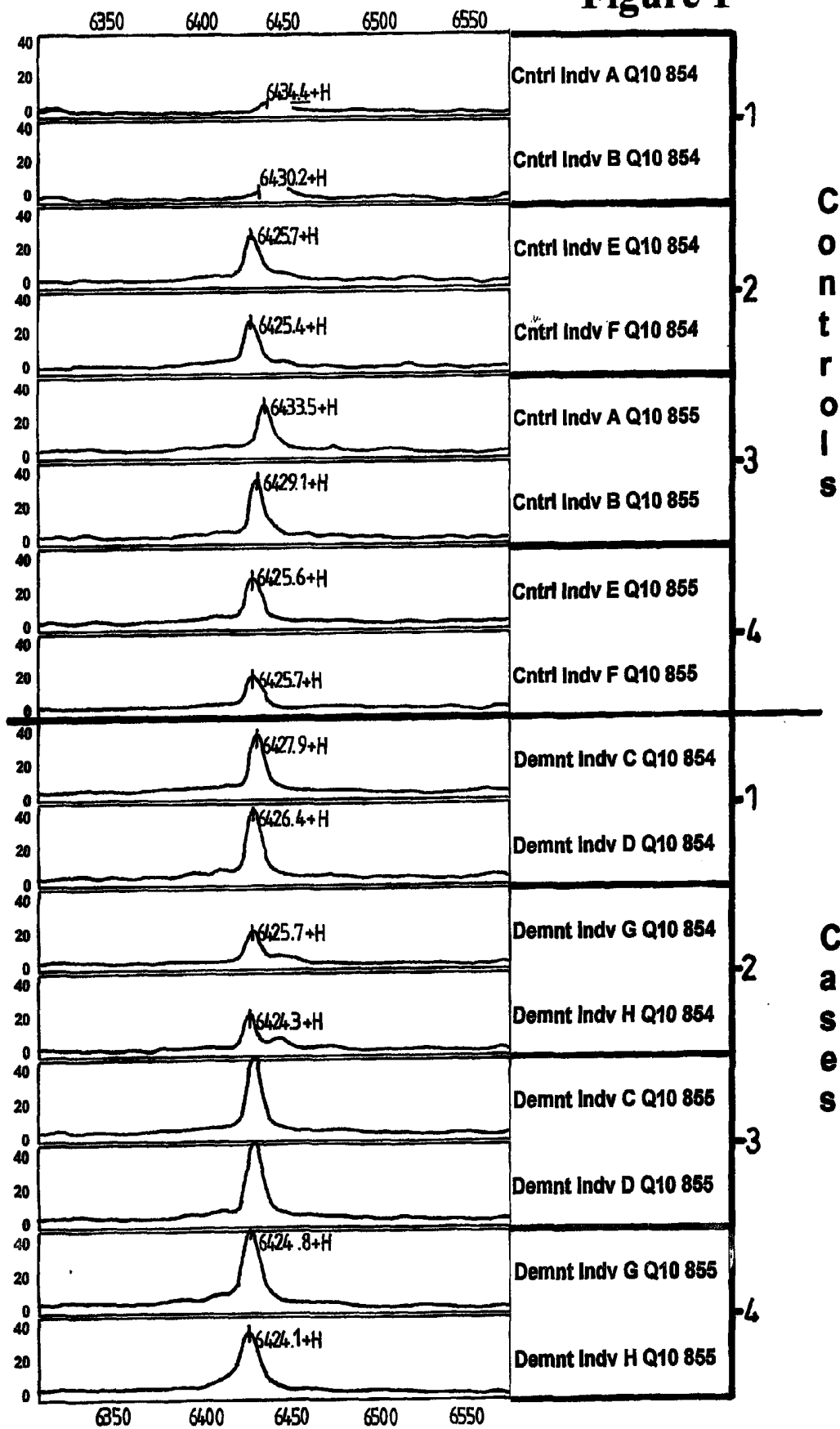
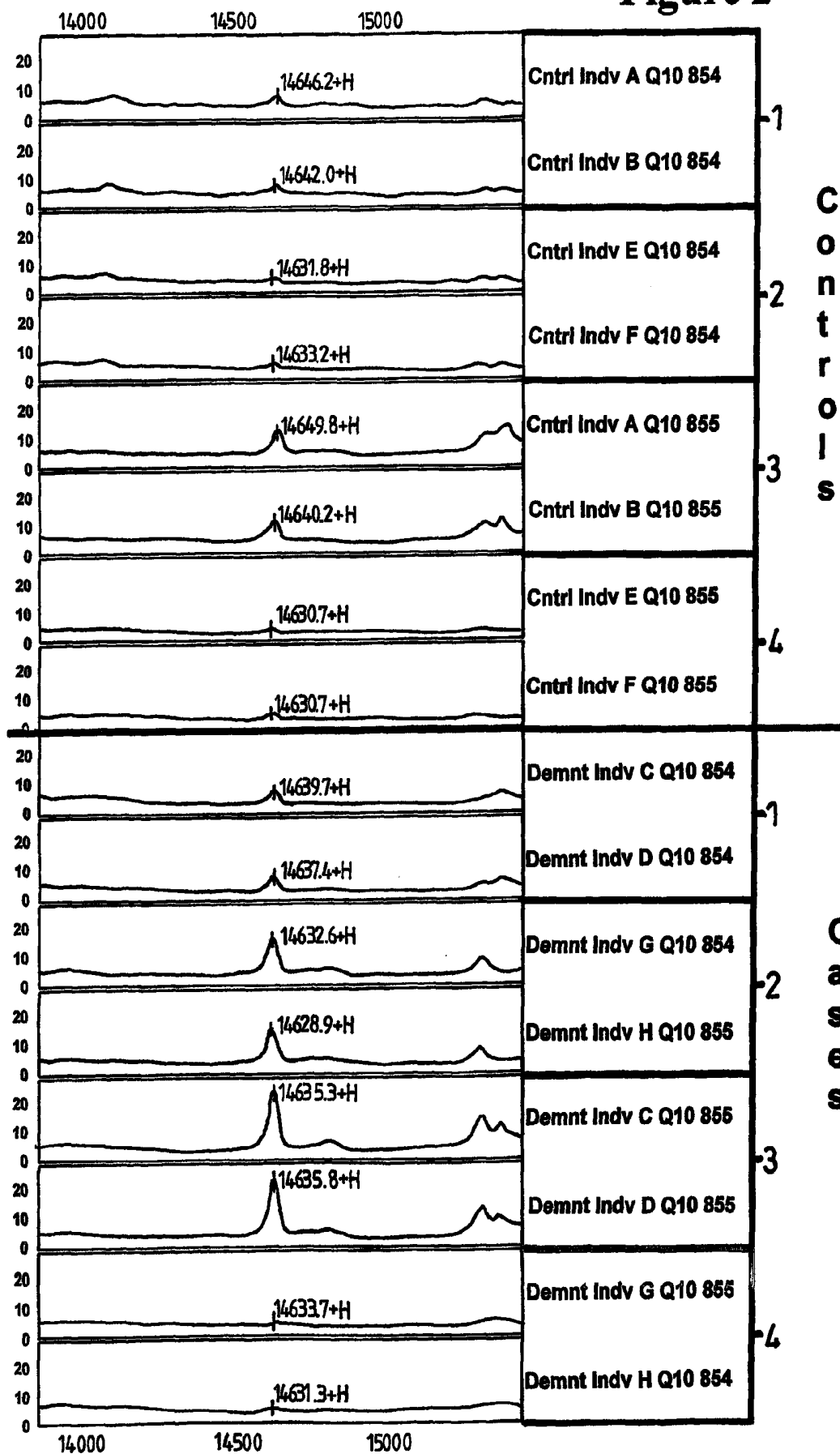


Figure 2



**Figure 3**

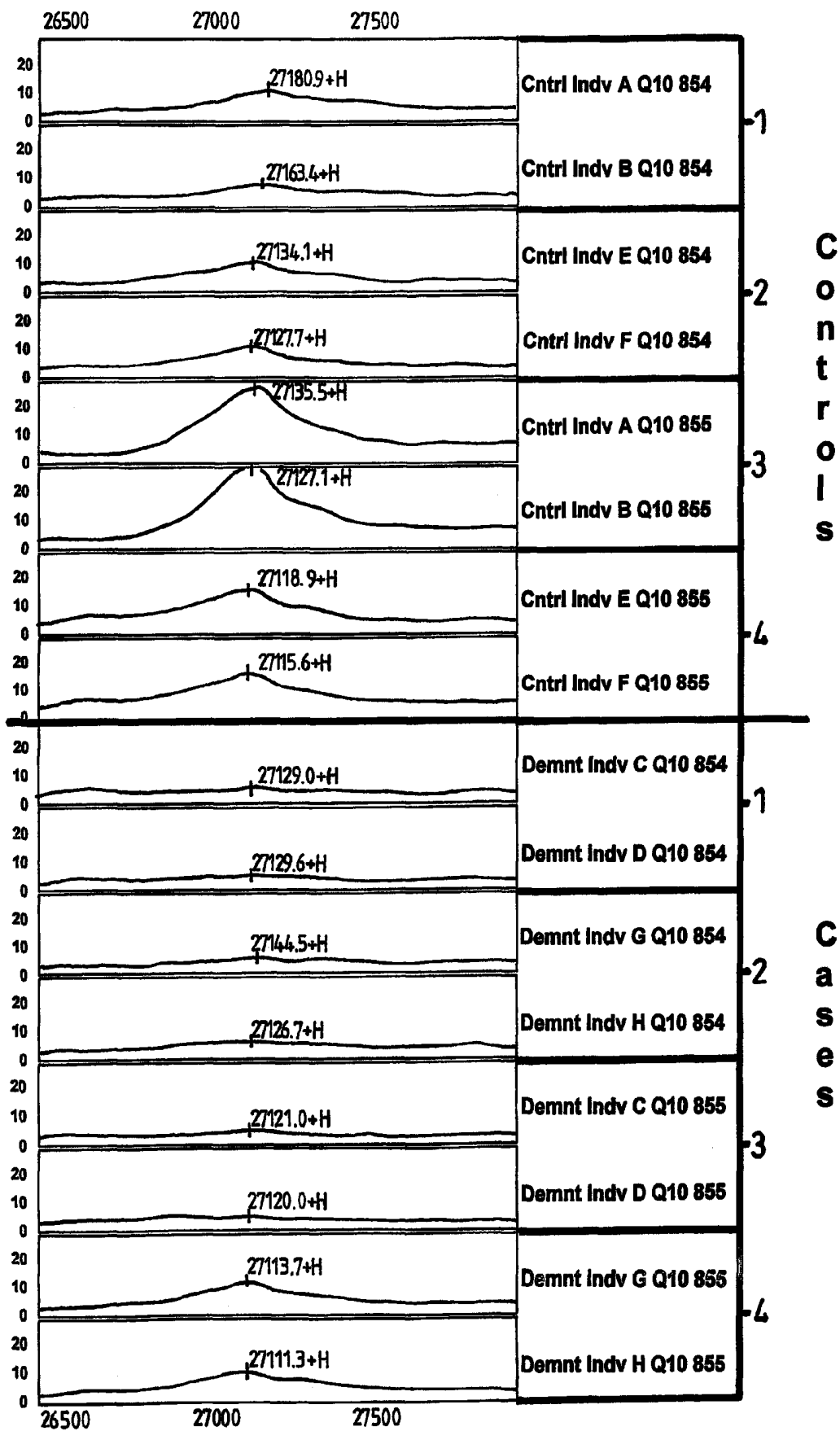
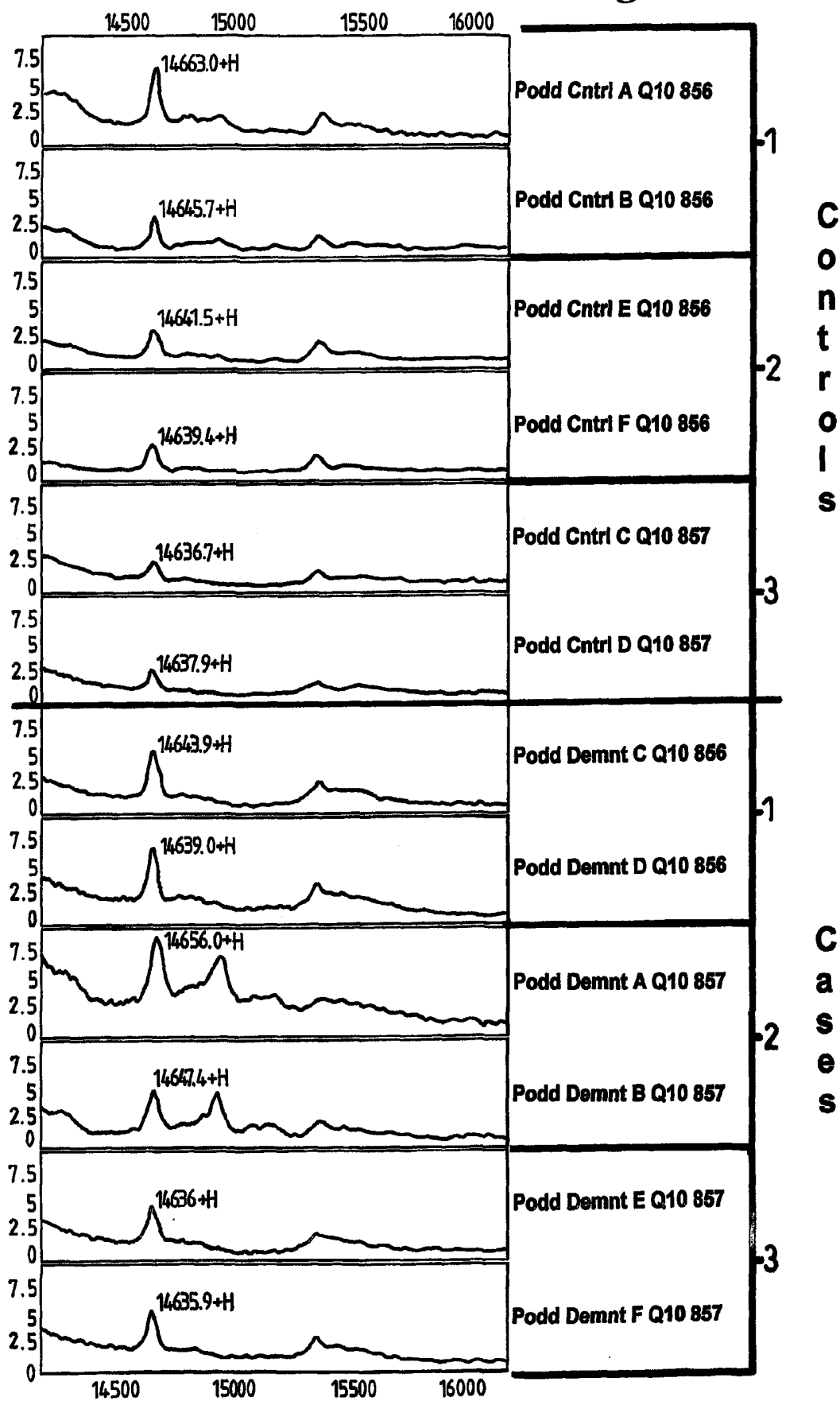


Figure 4



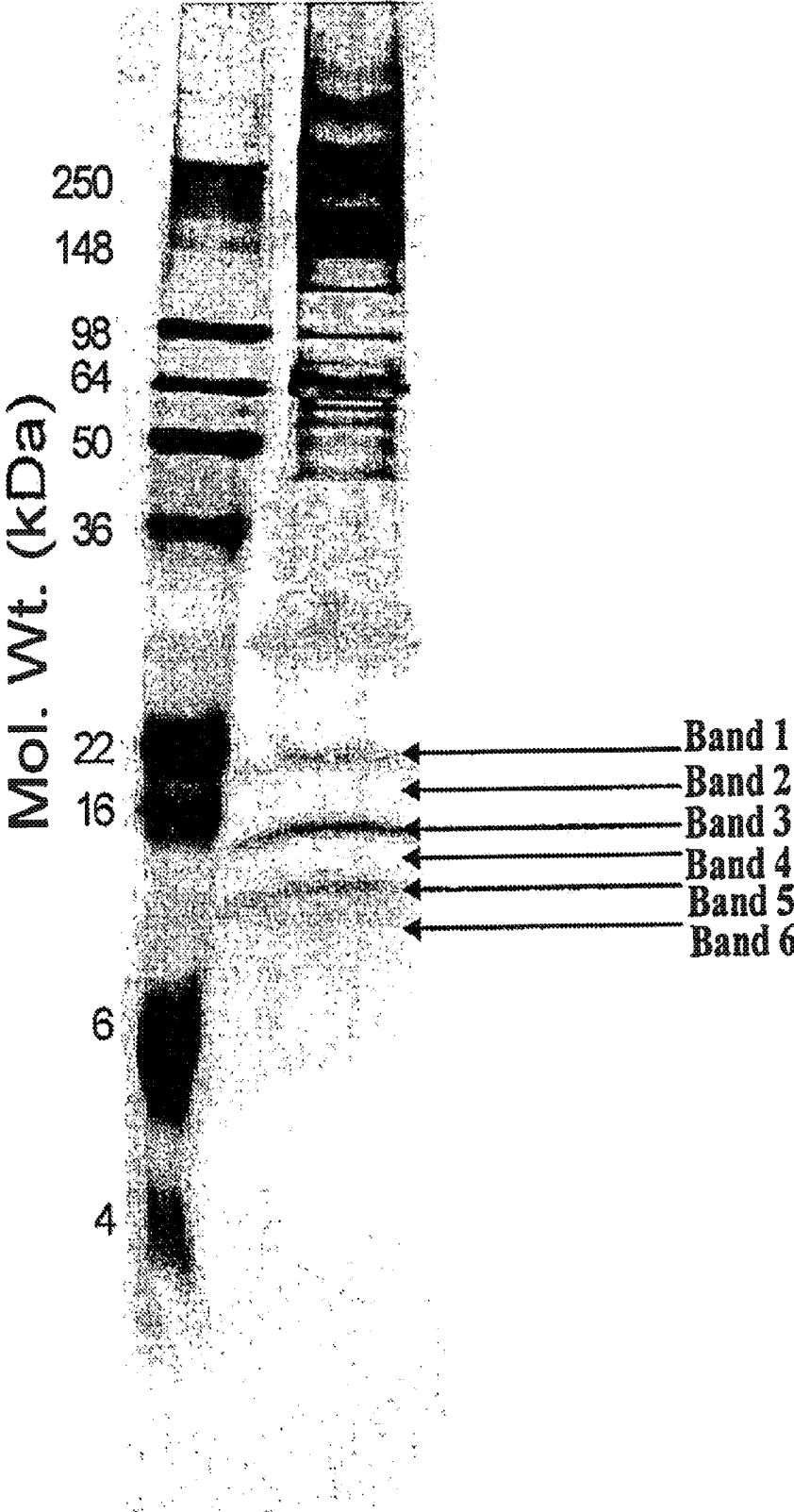


Figure 5

Spot No.	Rank	p Value	Fold Difference	State Change	Protein I.D.	Accession No.	Search Log No.
196	1	0.00030199	1.78	↑ AD	Desmoplakin (DP) (250/210 kDa paraneoplastic pemphigus antigen) Ig kappa chain C region	P15924 P01834 P01617 P02743	7495 7542 7542 7951
171	2	0.001255545	2.11	↑ AD	Ig kappa chain C region Serum albumin precursor	P01834 P02768 P47929	5623 7954 5623
2 (old)	3	0.001447694	13.75	↑ AD	Galectin-7 (Gal-7) (HKL-14) (PI7) (p53-induced protein 1) Complement factor H precursor (H factor 1) Serum albumin precursor	P08603 P02768 P01023 P00450	6672 6672 6672 6672
184	4	0.005360087	2.43	↑ AD	Alpha-2-macroglobulin precursor (Alpha-2-M) Ceruloplasmin precursor (EC 1.16.3.1) (Ferroxidase) Ig lambda chain C regions	P01842 P80748 P02768 P36980	7818 7818 7818 7818
177 (old)	5	0.005382883	1.92	↑ AD	Complement factor H-related protein 2 precursor (FHR-2) Ig lambda chain C regions Serum albumin precursor	P01842 P02768 P80748 P01834 P01023	5627 7955 7955 7955 7827
4	6	0.005985336	8.83	↑ AD	Alpha-2-macroglobulin precursor (Alpha-2-M)		
170	7	0.01167553	1.98	↑ AD			
13	8	0.015500401	4.23	↓ AD	Inter-alpha-trypsin inhibitor heavy chain H4 precursor (ITI heavy chain H4)	Q14624 P00450	7829 7829
165 (old)	9	0.018305158	1.58	↓ AD	Ceruloplasmin precursor (EC 1.16.3.1) (Ferroxidase) Serum albumin precursor	P02768	5625
164	10	0.020647469	2.03	↓ AD	Complement C4 precursor [Contains: C4a anaphylatoxin; C4b]	P01028	7821

Figure 6

14 (old)	11	0.025004429	10.82	↓ AD	Ig gamma-1 chain C region Serum albumin precursor Histone H2B.a/g/h/k/l (H2B.1 A) (H2B/a) (H2B/g) (H2B/h) (H2B/k) (H2B/l)	P01857 P02768 P62807	7821 6227 6227
126	12	0.028979402	1.6	↓ AD	CD5 antigen-like precursor (SP-alpha) (CT-2) (IgM-associated peptide) Serum albumin precursor	O43866 P02768 P01871	7493 7952 7952
176	13	0.029106689	1.75	↑ AD	Ig mu chain C region Ig lambda chain C regions Serum albumin precursor Ig lambda chain V-III region LOI	P01842 P02768 P80748	7816 7816 7816
123	14	0.031441346	1.36	↑ AD	Serum albumin precursor	P02768	7462
1	15	0.034723104	3.32	↑ AD	Alpha-2-macroglobulin precursor (Alpha-2-M) Ig alpha-1 chain C region	P01023 P01876	7823 7823

Figure 6 (continued)

Band No.	Protein I.D.	Species Accession		MW (Da)	pI	No. Peptides Matched	Percentages Coverage	Error (ppm)	Search Log	
		No.	Gen						No.	Peptide Matched
SPI_1C	Haptoglobin precursor	Human: P00738	19300	45177	6.13	5	9%	16	5288	
SPI_2C	Transferrin	Human: g13398S	18200	12835	5.33	8	87%	223	5275	
	Serum albumin precursor	Human: P02768	18200	69248	5.82	3	6%	236	5274	
	Complement C4 precursor	Human: P01028	18200	192650	6.65	1	0%	236	5274	GEEELQPSLQSK
	Fibrinogen alpha1alpha-1 chain precursor	Human: P02671	18200	94914	5.7	1	3%	224	5274	EVPFSEDSGDCPEAMDGLTSLGSLDGR
SPI_3C	Chain A, Transferrin	Human: g14432S	14900	13753	5.35	10	92%	102	5251	
	Apolipoprotein A-IV precursor (Apo-AIV)	Human: P06727	14900	45343	5.28	2	10%	100	5247	
	Serum albumin precursor	Human: P02768	14900	69321	5.92	1	1%	109	5247	
SPI_4C	Transferrin precursor	Human: P02766	14200	15877	5.52	5	60%	170	5260	
	Hemoglobin beta chain	Human: P02023	14200	15857	6.81	4	28%	12	5260	
	Serum albumin precursor	Human: P02768	14200	69321	5.92	4	7%	7	5260	
SPI_5C	Haptoglobin-related protein precursor	Human: P00739	12600	38983	6.42	5	10%	102	5294	
	Transferrin precursor	Human: P02766	12600	15877	5.52	2	16%	129	5294	
	Serum albumin precursor	Human: P02768	12600	69321	5.92	4	10%	120	5294	
	Apolipoprotein C-III precursor (Apo-CIII)	Human: P02656	12600	10845	5.23	1	16%	122	5294	DALSSVQESFAQQAR
	Hemoglobin alpha	Human: P01922	12600	15227	9.84	2	17%	118	5294	
	Hemoglobin beta chain	Human: P02023	12600	15857	6.81	1	15%	131	5294	SAVTALWGVWVDEYGGALGR
SPI_6C	Serum albumin precursor	Human: P02768	11600	69321	5.92	6	11%	244	5280	
	Apolipoprotein C-III precursor (Apo-CIII)	Human: P02656	11600	10845	5.23	3	37%	244	5280	
	Haptoglobin precursor	Human: P00738	11600	45177	6.13	2	6%	233	5280	
	Vitronectin precursor (Serum spreading factor) (S-protein)	Human: P04004	11600	54271	5.55	1	3%	249	5280	SLAQYWLGPAPGEL

Figure 7

No	IPI Accession no	SWISS-PROT Accession no	Name	No of matched peptides	regulation (control / disease)	CV (%)
1	IPI00166866	P01876	MGC27165 PROTEIN	2	0,38	7
2	IPI00336074	P01876	IG ALPHA-1 CHAIN C REGION	2	0,35	4
3	IPI00423461	P01842	HYPOTHETICAL PROTEIN DKFZP686C0222 0 (FRAGMENT)	2	0,35	24
4	IPI00431645	P00738	HAPTOGLOBIN PRECURSOR	1	0,33	-
5	IPI00478493	P00738	HAPTOGLOBIN PRECURSOR	1	0,34	-

Figure 8

1 MFLKAVVLT ALVAVAGARA EVSADQVATV MWDYFSQLSN NAKEAVEHLQ  
51 KSELTQQLNA LFQDKLGEVN TYAGDLQKKL VPFATELHER LAKDSEKLKE  
101 EIGKELEELR ARLLPHANEV SQKIGDNLRE LQQRLEPYAD QLRTQVNTQA  
151 EQLRRQLTPY AQRMERVLRE NADSLQASLR PHADELKAKI DQNVEELKGR  
201 LTPYADEFKV KIDQTVEELR RSLAPYAQDT QEKLNHQLEG LTFQMKNAB  
251 ELKARISASA EELRQRLAPL AEDVRGNLKG NTEGLQKSLA ELGGHLDQQV  
301 EEFRRRVEPY GENFNKALVQ QMEQLRQKLG PHAGDVEGHL SFLEKDLRDK  
351 VNSFFSTFKE KESQDKTSL PELEQQQEQ QEQQQEQVQM LAPLES

Figure 9

1 MRLWGLIWA SSFFTLISLQK PRLLLFSPSV VHLGVPLSVG VQLQDVPRGQ  
 51 VVKGSVFLRN PSRNNVPCSP KVDFTLSSEK DFALLSLQVP LKDAKSCGLH  
 101 QLLRGPEVQL VAHSPWLKDS LSRTTNIQGI NLLFS SRRGH LFLQTDQPIY  
 151 NPGQVRVRYV FALDQKMRPS TDTITVMVEN SHGLRVRKKE VYMPSSIFQD  
 201 DFVIPDISEP GTWKISARFS DGLESNSSTQ FEVKKYVLPN FEVKITPGKP  
 251 YILTVPGHLD EMQLDIQARY IYGKPVQGVA YVRFGLLDED GKKTFFRGLE  
 301 SQTCLVNGQS HISLSKAEFQ DALEKLNMG I TDLQGLRLYV AAAIESPGG  
 351 EMEEAEELTSW YFVSSPFLD LSKTKRHLVP GAPFLQLQALV REMSGSPASG  
 401 IPVKVSATVS SPGSVPEVQD IQQNTDGSQ VSIPI IIPQT ISELQLSVSA  
 451 GSPHPAIARL TVAAPPSSGGP GFLSIERPDS RPPRVGDTLN LNLRAVSGGA  
 501 TFSHYYYMIL SRGQIVFMNR EPKRTLTSVS VFVDHHLAPS FYFVAFYYHG  
 551 DHPVANSLRV DVQAGACEGK LELSVDGAKQ YRNGESVKLH LETDSLALVA  
 601 LGALDTALYA AGSKSHKPLN MGKVFEMNS YDLGCGPGGG DSALQVFQAA  
 651 GLAFSDGDQW TLRKRLSCP KEKTRKRN VNFQKAIN EK LGQYASPTAK  
 701 RCCQDGVTRL PMMRSCEQRA ARVQQPDCRE PFLSCCQFAE SLRKKSRDKG  
 751 QAGLQRALEI LQEDLIDED DIPVRSFFPE NWLWRVETVD RFQILTLWLP  
 801 DSLTTWEIHG LSLSKTKGLC VATPVQLRVF REFHLHLRLP MSVRRFEQLE  
 851 LRPVLYNYLD KNLTVSVHVS PVEGLCLAGG GGLAQQVLVP AGSARPVAFS  
 901 VVPTAAAVS LKVVARGSFE FVVGDAVSKV LQIEKEGAIH REELVYELNP  
 951 LDHRGRTLEI PGNSDPNMIP DGDFNSYVRV TASDPLDTLG SEGALSPGGV  
 1001 ASLLRLPRGC GEQTMIIY LAP TLAASRYLDK TEQWSTLPPE TKDHAVDLIQ  
 1051 KGYMRIQQFR KADGSYAOWL SRDSSTWLTA FVLKVLSLAQ EQVGGSPPEKL  
 1101 QETS NWLLSQ QQADGSFQDP CPVLD RSMQG GLVGNDETVA LTAFVTIALH  
 1151 HGLAVFQDEG AEPLKORVEA SISKANSFLG EKASAGLLGA HAAAITAYAL  
 1201 SLTKAPVDLL GVAHNNLMAM AQETGDNLYW GSVTGSQSNA VSPTPAPRNP  
 1251 SDPMPQAPAL WIETTAYALL HLLHEGKAE MADQASAWLT RQGSFOGGFR  
 1301 STQDTVIALD ALSAYWIASH TTEERGLNVT LSSTGERNGFK SHALQLNNRQ  
 1351 IRGLEEELQF SLGSKINVKV GGNSKGT LKV LRTYNVLD MK NTTCQDLQIE  
 1401 VTVKGHVEYT MEANEDYEDY EYDELPAKDD PDAPLQPVTP LQLFEGRRNR  
 1451 RRREAPKVVE EQESRVHYTV CIWRNGKVGL SGMALADVT L LSGFHALRAD  
 1501 LEKLTSLSDR YVSHFETEGP HVLLYFDSVP TSRECVGF EA VQEVVGLVQ  
 1551 PASATLYDYY NPERRCSV FY GAPSKSRLLA TLCSAEVCQC AEGKCPRQRR  
 1601 ALERGLQDED GYRMKFACY PRVEYGFQVK VLREDSRAAF RLFETKITQV  
 1651 LHFTKDVKAA ANQMRNFLVR ASCRLRLEPG KEYLIMGLDG ATYDLEGHPQ  
 1701 YLLDSNSWIE EMPSERLCRS TRQRAACAQL NDFLQIEYGTQ GCQV

Figure 10

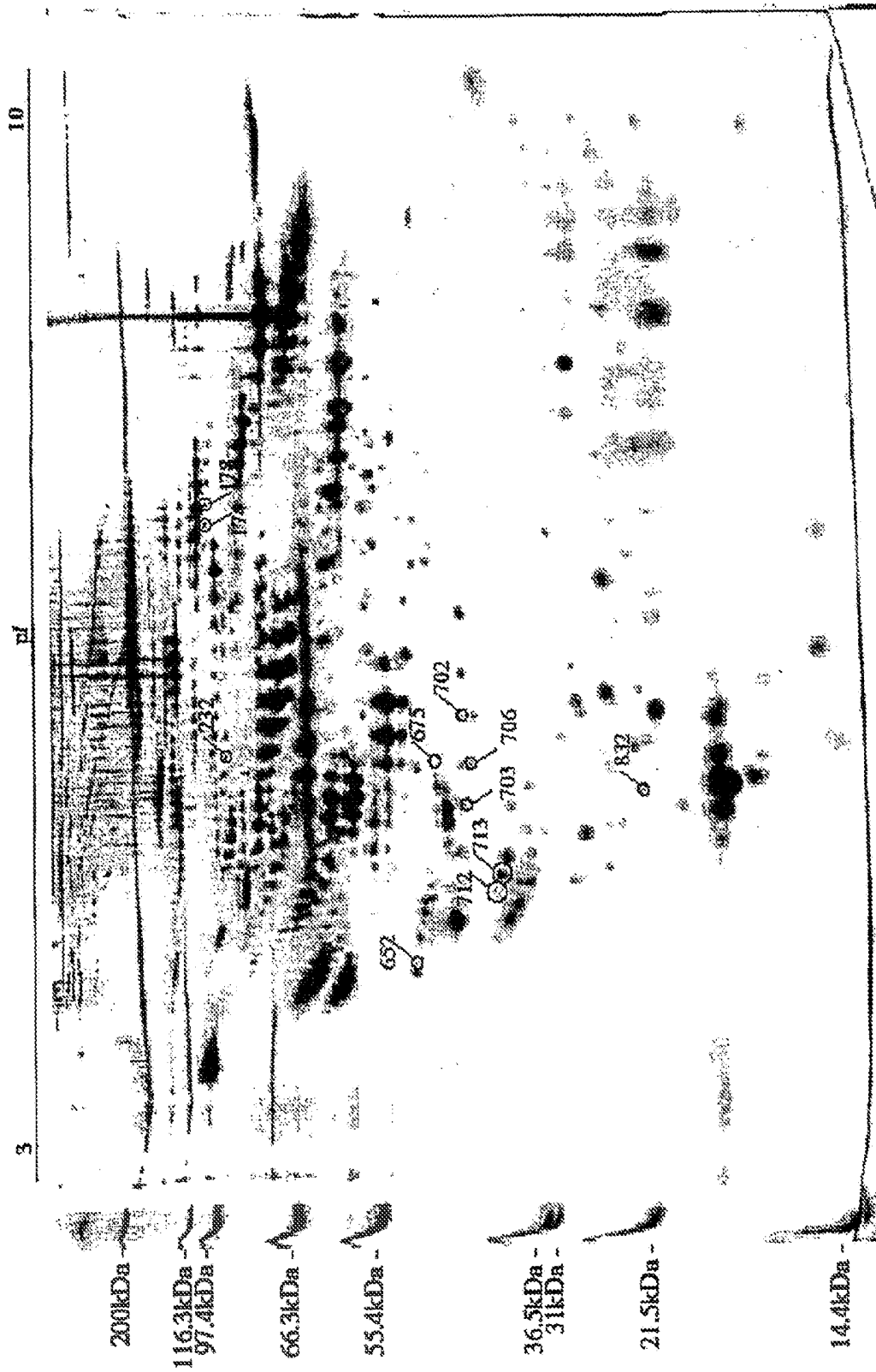


Figure 11

Spot no.	Protein Name	Acc. No.	Norm. Vol	CV (%)	Norm. Vol	CV (%)	Expressio n ratio	T-test (p)	Detection ratio	Theoretical Mr	pI	Cover age (%)
			Control	Disease								
174	alpha-2-macroglobulin precursor	P01023	0,36604	52	0,14510	57	0,40	2,50852E-06	28/21	160796 Da	5,95	13,8
178	alpha-2-macroglobulin precursor	P01023	0,28348	53	0,12475	57	0,43	1,25521E-05	27/20	160796 Da	5,95	11,7
232	Inter-alpha-trypsin inhibitor heavy chain H4 precursor	Q14624	0,32468	81	0,14481	64	0,45	0,001793752	28/27	103358 Da	6,51	25,6
712	Complement C3 precursor	P01024	0,81225	74	0,36429	62	0,45	0,001371446	28/24	184967 Da	6,00	14,9
712	Clusterin precursor	P10909	0,81225	74	0,36446	62	0,45	0,001371446	28/24	50062105	5,89	22,9
719	Complement C3 precursor	P01024	3,45893	67	1,44590	63	0,42	9,67927E-05	29/29	184967 Da	6,00	16,9
652	Complement C4 precursor	P01028	0,18067	122	0,39844	69	2,21	0,003595593	24/25	192771 Da	6,60	5,9
675	Actin cytoplasmic 2 (Gamma/beta actin)	P63261	0,21268	89	0,45443	74	2,14	0,002107165	25/28	41793 Da	5,31	49,1
702	Haptoglobin precursor	P00738	0,09210	86	0,40084	115	4,35	0,002920373	20/25	43349 Da	6,13	21,2
703	Haptoglobin precursor	P00738	1,54479	95	4,64500	76	3,01	0,000171685	24/28	43349 Da	6,13	23,9
706	Haptoglobin precursor	P00738	1,00814	112	3,26743	89	3,24	0,000607583	21/28	43349 Da	6,13	23,6
832	Complement C4 precursor	P01028	0,24743	126	0,61914	88	2,50	0,003005256	28/28	192771 Da	6,60	4,5

Figure 12

1 MGKNKLLHPS LVLLLLVLLP TDASVSGKPQ YMVLVPSLLH TETTEKGCVL LSYLNETVTV  
61 SASLESVRGN RSLFTDLEAE NDVLHCVAFA VPKSSSNEEV MFLTVQVKGP TQEFKKRTTV  
121 MVKNEDSLVF VQTDKSIYKP GQTVKFRVVS MDENFHPLNE LIPLVYIQDP KGNRIAQWQS  
181 FQLEGGLKQF SFPLSSEPFQ GSYKVVVQKK **SGGRTEHPFT** **VEEFVLPKFE** VQVTVPKIIT  
241 ILEEE MNVSV CGLYTYGKPV PGHVTVSICR KYSDASDCHG EDSQAFCEKF **SGQLNSHGCF**  
301 **YQQVKT**KVFQ LKRKEYEMKL **HTEAQIQEEG** **TVVELTGRQS** SEITRTITKL **SFVKVDSHFR**  
361 **QGIPFFGQVR** **LVDGKGVPIP** NKVIFIRGNE ANYYSNATTD EHGLVQFSIN TTNVMGTSLT  
421 VRVNYKDRSP CYGYQWVSEE HEEAHHTAYL VFSPSKSFVH LEPMSELPC GHTQTVQAHY  
481 IILNGGTLGL KKL SFYYLIM AKGGIVRTGT HGLLVKQEDM **KGHFSISIPV** **KSDIAPVARL**  
541 LIYAVLPTGD VIGDSAKYDV ENCLANKVDL **SFSPSQSLPA** **SHAHLRVTAA** **PQSVCALRAV**  
601 DQSVLLMKPD AELSASSVYN LLPEKDLTGF PGPLNDQDDE DCINRHNVIYI NGITYTPVSS  
661 TNEKDMYSFL EDMGLKAFTN SKIRKPKMCP **QLQQYEMHGP** **EGLRVGFYES** DVMGRGHARL  
721 VHVEEPHTET VRKYFPETWI WDLVVVNSAG VAEVGVTPD TITEWKAGAF CLSEDAGLGI  
781 SSTASLRAFQ PFFVELTMPY SVIRGEAFTL KATVLNLYPK CIRVSVQLEA SPAFLAVPVE  
841 KEQAPHCICA NGRQTVSWAV TPKSLGNVNF TVSAEALESQ ELCGTEVPSV PEHGRKDTVI  
901 KPLLVEPEGL EKETT FNSSL CPSGGEVSEE LSLKLPNNV EESARASVSV LGDILGSAMQ  
961 NTQNLLQMPY GCGEQNMVLF APNIYVLDYL NETQQLTPEV KSKAIGYLNT GYQRQLNYKH  
1021 YDGSYSTFGE RYGRNQGNTW LTAFVLKTFQ QARAYIFIDE AHITQALIWL SQRQKDNCGF  
1081 RSSGSLNNA IKGGVEDEV T LSAYITIALL EIPLTVTHPV VRNALFCLES ANKTAQEGDH  
1141 GSHVYTKALL AYAFALAGNQ DKRKEVLKSL NEEAVKKNNS VHWERPQKPK APVGHFYEPQ  
1201 APSAEVENTS YVLLAYLTAQ PAPTSEDLTN ATNIVKWITK QQNAQGGFSS TQDTVVALHA  
1261 LSKYGAATFT RTGKAAQVTI QSSGTFSSKF QVDNRRLLL QQVSLPELPG EYSMKVTGEG  
1321 CVYLQTSKY NILPEKEEFP FALGVQTLPO TCDEPKAHTS FQISLSVSYT GSRASANMAI  
1381 VDVKMVSGFI PLKPTVKMLE RSNHVSRTVE SSNHVLIYLD KVSNOTLSLF FTVLQDVPVR  
1441 DLKPAIVKVY DYYETDEFI AEYNAPCSKD LGNA

Figure 13

1 MKPPRPVRTC SKVLVLLSLL AIHQTTTAEK NGIDIYSLTV DSRVSSREAH TVVTSRVVNR  
61 ANTVQEATFQ MELPKKAFIT NFSMNIDGMT YPGIIEKAE AQAQYSAAVA KGKSAGLVKA  
121 TGRNMEQFQV SVSVAPNAKI TFELVYEELL KRRLGVYELL LKVRPQQLVK HLQMDIHIFE  
181 POGISFLETE STFMTNQLVD ALTTWQNKTK AHIRFKPTLS QOQKSPEQQE TVLDGNLIIR  
241 YDVDRAISGG SIQIENGYFV HYFAPEGLTT MPKNVVFVID KSGSMSGRKI QOTREALIKI  
301 LDDLSPRDQF NLIVFSTEAT QWRPSLVPAS AENVNKARSE AAGIQALGGT NINDAMLMAV  
361 QLLDSSNQEE RLPEGSVSLI ILLTDGDPTV GETNPSIQN NVREAVSGRY SLFCLGFGFD  
421 VSYAFLEKLA LDNGGLARRI HEDSDSALQL QDFYQEVANP LLTAVTFEYP SNAVEEVTQN  
481 NERLLEFKGSE MVVAGKLQDR GPDVLTATVS GKLPTQNITF QTESSVAEQE AEFQSPKYIF  
541 HNEMERLWAY LTIQQLLEQT VSASDADQQA LRNQALNLSL AYSEVTPPTS MVVTKPDDQE  
601 QSQVAEKPME GESRNRNVHS GSTFFKYYLQ GAKIPKPEAS FSPRRGWNRO AGAAGSRMNF  
661 RPGVLSSRQL GLPGPPDVED HAAYHPFRRL AILPASAPPA TSNPDPAVSR VMNMKIEETT  
721 MTTQTPAPIQ APSAILPLPG QSVRLCVDP RHRQGPVNLL SDPEQGVVET GQYEREKAGF  
781 SWIEVTFKNP LVVWHASPEH VVTRNRRSS AYKWKETLFS VMPGLKMTMD KTGLLLLSDP  
841 DKVTIGLLFW DGRGEGRLRL LRDTDRFSSH VGGTLGQFYQ EVLWGSPAAS DDGRRTLVRQ  
901 GNDHSATRER RLDYQEGPPG VEISCWSVEL

Figure 14

1 MGPTSGPSSL LLLLTHLPLA LGSPMYSIIT PNILRESEE TMVLEAHAQ GDVPTVTVH  
 61 DFPGKKLVLS SEKTVLTPAT NHMGNVTFTI PANREFKSEK GRNKFVTVQA TFGTQVVEKV  
 121 VLVSLQSGYL FIQTDKTIYT PGSTVLYRIF TVNHKLLPVG RTVMVNIENP EGIPVKQDSL  
 181 SSONQLGVLP LSWDIPELVN MGQWKIRAYY ENSPQQVFST EFEVKEYVLP SFEVIVEPTE  
 241 KFYIYNEKG LEVTITARFL YGKKVEGTAF VIFGIQDGEQ RISLPESLKR IPIEDGSGEV  
 301 VLSRKVLLDG VQNLRAEDLV GKSLYVSATV ILHSGSDMVQ AERSGIPIVT SPYQIHFTKT  
 361 PKYFKPGMPF DLMVFVTNPD GSPAYRVPVA VQGEDTVQSL TQGDGVAKLS INTGPSQKPL  
 421 SITVRTKKQE LSEAEQATRT MQALPYSTVG NSNNYLHLSV LRTELRPGET LNVNELLRMD  
 481 RAHEAKIRYY TYLIMNKGR LKAGRQVREP GQDLVVLPLS ITTDFIPSFR LVAYYTLIGA  
 541 SGQREVVADS VWVDVKDSCV GSLVVKSGQS EDRQPVPGQQ MTLKIEGDHG ARVVLVAVDK  
 601 GVFLNKKKNK LTQSKIWDVV EKADIGCTPG SGKDYAGVFS DAGLFTFTSSS GQQAQRAEL  
 661 QCPQPAARRR RSVQLTEKRM DKVGKYPKEL RKCCEDGMRE NPMRFSCQRR **TRFISLGEAC**  
 721 KKVFLDCCNY ITELRRQHAR ASHLGLARSN LDEDIAEEN IVSRSEFPES WLWNVEDLKE  
 781 PPKNGISTKL MNIFLKDSIT TWEILAVSMS DKKGICVADP FEVTVMQDFE IDLRLPYSVV  
 841 RNEQVEIRAV LYNRQNOEL KVRVELLHNP AFCSLATTKR RHQQTVTIPP KSSLSVPYVI  
 901 VPLKTGLEQEV EVKAAVYHHF ISDGVKSLK VVPEGIRMNK TVAVRTLDPD RLGREGVQKE  
 961 **DIPPADLSDQ** **VPDTESETRI** **LLQGTQVQOM** **TEDAUDAERL** **KHLIVTPSGC** **GEQNMIGMTP**  
 1021 TVIAVHYLDE TEQWEKFGLE KRQGALELIK KGYTQQLAFR **QPSSAFAAFV** **KRAPSTWLTA**  
 1081 YVVKVFSLAV **NLIAIDSQVL** **CGAVKWLILE** **KQKPDGVFQE** **DAPVIHQEMI** **GGLRNNNEKD**  
 1141 MALTAFLVLIS **LQEAKDICEE** **QVNSLPGSIT** **KAGDFLEANY** **MNLQRSYTV** **IAGYALQMG**  
 1201 RLKGPLLNKE LTTAKDKNRW EDPGKQLYNV **EATSYALLAL** **LQLKDFDFVP** **PVVRWLNEQR**  
 1261 **YGGGYGSTQ** **ATFMVQALA** **QYQKDAPDHQ** **ELNLDVSLQL** **PSRSSKITHR** **IHWESASLLR**  
 1321 SEETKENEGB **TVTAEKGQCG** **TLSVVTMYHA** **KAKDQLTCNK** **FDLKVTIKPA** **PETEKRPQDA**  
 1381 KNTMILEICT RYRGDQDATM SILDISMGTG FAPDTDDLKQ LANGVDRYIS KYELDKAFSD  
 1441 RNTLIIYLDK VSHSEDDCLA FKVHQYFNVE LIQPGAVKVY AYYNLEESCT RFYHPEKEDG  
 1501 KLNKLCRDEL CRCAEENCFI QKSDDKVTLE ERLDKACEPG VDYVYKTRLV KVQLSNDFDE  
 1561 YIMAIEQTIK SGSDEVQVGQ QRTFISPIKC REALKLEEK HYLWGLSSD FWGKPNLSY  
 1621 IIGKDTWVEH WPEEDECQDE ENQKQCQDLG AFTESMVVFG CPN

Figure 15

1 MMKTLLLFVG LLLTWESGQV LGDOTVSDNE LQEMSNQGSK YVNKEIQNAV NGVKQIKTLI  
61 EKTNEERKTL LSNLEEAKKK KEDALNETRE SETKLKELPG **VCNETMMALW** **EECKPCLKQT**  
121 CMKFYARVCR SGSGLVGRQL EEFLNQSSPF YFWMNGDRID SLENDRQQT HMLDVMQDHF  
181 SRASSIIDEL **FQDRFFTREP** **QDTYHYLPFS** **LPHRRPHFFF** PKSRIVRSLM PFSPYEPLNF  
241 HAMFQPFLEM IHEAQQAMDI HEHSPAFQHP PTEFIREGDD DRTVCREIRH NSTGCLRMKD  
301 QCDKCREILS VDCSTNNPSQ AKLRRELD**ES** **LQVAERLTRK** YNELLKSYQW KMLNTSSLLE  
361 QLNEQFNWVS RLANLTQGED QYYLRVTTVA SHTSDSDVPS GVTEVVVKLE **DSDPITVTVP**  
421 **VEVSRKNPKF** METVAEKALQ EYRKKHREE

**Figure 16**

1 MRLWGLIWA SSEFTLSLQK PRLLLFSPSV VHLGVPLSVG VQLQDVPRGQ VVKGSVFLRN  
61 PSRNNVPCSP KVDFTLSSER DFALLSLQVP LKDAKSCGLH QLLRGPEVQL VAHSPWLKDS  
121 LSRTTNIQGI NLLFSSRRGH LFLQTDQPIY NPGQVRVRYV FALDQKMRPS TDTITVMVEN  
181 SHGLRVRKKE VYMPSSIFQD DFVIPDISEP GTWKISARFS DGLESNSSTQ FEVKKYVLPN  
241 FEVKITPGKP YILTVPGHLD EMQLDIQARY IYGKPVQGVA YVRFGLLDED GKKTFFRGLE  
301 SQTKLNVGQS HISLSKAEFQ DALEKLNMG I TDLOGLRLYV AAATIESPGG EMEEAELTSW  
361 YFVSSPFLSD LSKTKRHLVP GAPFLQALV REMSGSPASG IPVKVSATVS SPGSVPEVQD  
421 IQQNTDGSQG VSIPIIIPQT ISELQLSVSA GSPHPAIARL TVAAPPSGGP GFLSIERPDS  
481 RPRVGDTLN LNLRAVGSGA TFSHYYYMIL SRQIVFMNR EPKRTLTSVS VFVDHHLAPS  
541 FVFVAFYYHG DHPVANSLRV DVQAGACEGK LELSVDGAKQ YRNGESVKLH LETDSLALVA  
601 LGALDTALYA AGSKSHKPLN MGKVFEAMNS YDLGCGPGGG DSALQVFQAA GLAFSDGDQW  
661 TLSRKRLSCP **KEKTTRKRN** VNFQKAIN EK **LGQYASPTAK** RCCQDGVTRL PMMRSCEQRA  
721 ARVQQPDCRE PFLSCCFAE SLRKKSRDKG QAGLQRALEI LQEEDLDED DIPVRSFFPE  
781 NWLWRVETVD RFQILTLWLP DSLTTWEIHG LSLSKTKGLC **VATPVQLRVF** **REFHLHLRLP**  
841 **MSVRRFEQLE** LRPVLYNYLD KNLTVSVHVS PVEGLCLAGG GGLAQQVLVP AGSARPVAFS  
901 VVPTAAAAYS LKVVARGSFE **FPVGDVSKV** **LQIEKEGAIH** **REELVYELNP** LDHRGRITL E I  
961 **PGNSDPNMIP** **DGFNSYVRV** **TASDPLDTLG** **SEGALSPGGV** **ASLLRLPRGC** **GEQTMYYLAP**  
1021 **TLAASRYLDK** TEQWSTLPPE TKDHAVDLIQ KGYMRIQQFR KADGSYAAWL SRDSSTWLTA  
1081 FVLKVLSLAQ EQVGGSPPEL QETSNWLLSQ QQADGSFQDP CPVLDRSMQG GLVGNDETVA  
1141 LTAFVTIALH HGLAVFQDEG AEPLKQVEA SISKANSFLG EKASAGLLGA HAAAITAYAL  
1201 SLTKAPVDLL GVAHNNLMAM AQETGDNLYW GSVTGSQSNA VSPTPAPRNP SDPMPQAPAL  
1261 WIETTAYALL HLLHEGKAE MADQASAWLT **RQGSFQGGFR** STQDTVIALD ALSAYWIASH  
1321 TTEERGLNVT LSSTGRNGFK SHALQLNNRQ IRGLEEBELQF SLGSKINVKV GGNSKGTLLK  
1381 LRTYNVLDK NNTCODLQIE VTVKGHVEYT MEANEDYEDY EYDELPAKDD PDAPLQPVTP  
1441 LQLFEGRNR RRREAPKVE EQESRVHYTV CIWRNGKVGL SGMADVTL LSGFHALRAD  
1501 LEKLTSLSDR YVSHFETEGP HVLLYFDSVP TSRECVGFEA VQEVVGLVQ PASATLYDYY  
1561 NPERRCSVYF GAPSKSRLLA TLCSAEVCQC AEGKCPQRR ALERGLQDED GYRMKFACY  
1621 PRVEYGFQVK VLREDSRAAF RLFETKITQV LHFTKDVKAA ANQMRNFLVR ASCRLRLEPG  
1681 KEYLIMGLDG ATYDLEGHPQ YLLDSNSWIE EMPSERLCRS TRQRAACAQL NDFLQEYGTQ  
1741 GCQV

Figure 17

1 MEEEIAALVI DNGSGMCKAG FAGDDAPRAY **FPSIVGRPRH** QGVMVGMGQK **DSYVGDEAQS**  
61 **KRGILTLKYP** IEHGIVTNWD DMEKIWHHTF YNELRVAPEE **HFVLLTEAPL** **NPKANREKMT**  
121 QIMFETFNTP AMYVAIQAVL SLYASGR**TTG** **IVMDSGDGVT** **HTVPIYEGYA** **LPHAILRLDL**  
181 **AGRDLTDYLM** KILTERGYSE TTTAEREIVR DIKEKLCYVA LD**FEQEMATA** **ASSSSLEKSY**  
241 **ELPDGQVITI** GNERFRCPEA LFQPSFLGME SCGIHETTFN **SIMKCDVDIR** **KDLYANTVLS**  
301 **GGTMYPGIA** DRMQKEITAL APSTMKIKII **APPERKYSVW** **IGGSILASLS** **TFQOMWISKQ**  
361 **EYDESGPSIV** **HRKCF**

**Figure 18**

1 MSALGAVIAL LLWGQLFAVD SGNDVTDIAD DGCPKPPEIA HGYVEHSVRY QCKNYYKLRT  
61 EGDGVYTLND KKQWINKAVG DKLPECEADD GCPKPPEIAH GYVEHSVRYQ CKNYYKL RTE  
121 GDGVYTLNNE KQWINKAVGD KLPECEAVCG KPKNPANPVQ RILGGHLD AK GSFPWQAKMV  
181 SHHNLTTGAT LINEQWLLTT AKNLFLNHSE NATAKDIAPT LTLYVGKKQL VEIEKVV LHP  
241 NYSQVDIGLI KLKQKSVNE RVMPICLPSK DYAEVGRVGY VSGWGRNANF KFTDHLKYVM  
301 LPVADQDQCI RHYEGSTVPE KKT PKSPVGV QPILNEHTFC AGMSKYQEDT CYGDAGSAFA  
361 VHDLEEDTWY ATGILSEDKS CAVAEYGVYV KVTSIQDWVQ KTIAEN

**Figure 19**

## METHODS OF DIAGNOSING ALZHEIMER'S DISEASE

### FIELD OF THE INVENTION

**[0001]** The present invention relates to methods and compositions relating to Alzheimer's disease. Specifically, the present invention identifies and describes proteins that are differentially expressed in the Alzheimer's disease state relative to their expression in the normal state and, in particular, identifies and describes proteins associated with Alzheimer's disease. Further, the present invention provides methods of diagnosis of Alzheimer's disease using the differentially expressed proteins. Still further, the present invention provides methods for the identification and therapeutic use of compounds for the prevention and treatment of Alzheimer's disease.

### BACKGROUND OF THE INVENTION

**[0002]** Dementia is one of the major public health problems of the elderly, and in our ageing populations the increasing numbers of patients with dementia is imposing a major financial burden on health systems around the world. More than half of the patients with dementia have Alzheimer's disease (AD). The prevalence and incidence of AD have been shown to increase exponentially. The prevalence for AD in Europe is 0.3% for ages 60-69 years, 3.2% for ages 70-79 years, and 10.8% for ages 80-89 years (Rocca, Hofman et al. 1991). The survival time after the onset of AD is approximately from 5 to 12 years (Friedland 1993).

**[0003]** Alzheimer's disease (AD), the most common cause of dementia in older individuals, is a debilitating neurodegenerative disease for which there is currently no cure. It destroys neurons in parts of the brain, chiefly the hippocampus, which is a region involved in coding memories. Alzheimer's disease gives rise to an irreversible progressive loss of cognitive functions and of functional autonomy. The earliest signs of AD may be mistaken for simple forgetfulness, but in those who are eventually diagnosed with the disease, these initial signs inexorably progress to more severe symptoms of mental deterioration. While the time it takes for AD to develop will vary from person to person, advanced signs include severe memory impairment, confusion, language disturbances, personality and behaviour changes, and impaired judgement. Persons with AD may become non-communicative and hostile. As the disease ends its course in profound dementia, patients are unable to care for themselves and often require institutionalisation or professional care in the home setting. While some patients may live for years after being diagnosed with AD, the average life expectancy after diagnosis is eight years.

**[0004]** In the past, AD could only be definitively diagnosed by brain biopsy or upon autopsy after a patient died. These methods, which demonstrate the presence of the characteristic plaque and tangle lesions in the brain, are still considered the gold standard for the pathological diagnoses of AD. However, in the clinical setting brain biopsy is rarely performed and diagnosis depends on a battery of neurological, psychometric and biochemical tests, including the measurement of biochemical markers such as the ApoE and tau proteins or the beta-amyloid peptide in cerebrospinal fluid and blood.

**[0005]** Biomarkers may possibly possess the key in the next step for diagnosing AD and other dementias. A biological marker that fulfils the requirements for the diagnostic test for

AD would have several advantages. An ideal biological marker would be one that identifies AD cases at a very early stage of the disease, before there is degeneration observed in the brain imaging and neuropathological tests. A biomarker could be the first indicator for starting treatment as early as possible, and also very valuable in screening the effectiveness of new therapies, particularly those that are focussed on preventing the development of neuropathological changes. A biological marker would also be useful in the follow-up of the development of the disease.

**[0006]** Markers related to pathological characteristics of AD; plaques and tangles (A $\beta$  and tau) have been the most extensively studied. The most promising has been from studies of CSF concentration of A $\beta$ (1-40), A $\beta$ (1-42) and tau or the combination of both proteins in AD. Many studies have reported a decrease in A $\beta$ (1-42) in CSF, while the total A $\beta$  protein or A $\beta$ (1-40) concentration remain unchanged (Ida, Hartmann et al. 1996; Kanai, Matsubara et al. 1998; Andreassen, Hesse et al. 1999).

### SUMMARY OF THE INVENTION

**[0007]** Broadly, the present invention relates to methods and compositions for the diagnosis of Alzheimer's disease. More specifically, the present invention identifies and describes proteins that are differentially expressed in the Alzheimer's disease state relative to their expression in the normal state.

**[0008]** In a first aspect, the invention provides a method of diagnosing Alzheimer's disease in a subject, the method comprising detecting one or more of a differentially expressed protein identified by the methods described herein in a tissue sample or body fluid sample from said subject. Preferably, the method is an in vitro method.

**[0009]** In all aspects, the methods of the present invention may also be used in relation to pre-Alzheimer's stages such as mild cognitive impairment (MCI) as well as advanced Alzheimer's disease.

**[0010]** In another aspect, the present invention provides a method of determining the nature or degree of Alzheimer's disease in a human or animal subject, the method comprising detecting one or more of a differentially expressed protein identified by the methods described herein in a tissue sample or body fluid sample from said subject. Thus, the methods of the present invention encompass methods of monitoring the progress of Alzheimer's disease or of disease progression from MCI to Alzheimer's disease. Also encompassed are prognostic methods, for example prognosis of likely progression from MCI to Alzheimer's disease, or prognosis of likely duration or severity of Alzheimer's disease.

**[0011]** In a preferred embodiment the method comprises:

**[0012]** (a) establishing a paradigm in which at least one protein is differentially expressed in relevant tissue or body fluid sample from, or representative of, subjects having differential levels of Alzheimer's disease;

**[0013]** (b) obtaining a sample of the tissue or body fluid sample from the subject;

**[0014]** (c) determining the presence, absence or degree of expression of the differentially expressed protein or proteins in the sample; and

**[0015]** (d) relating the determination to the nature or degree of the Alzheimer's disease by reference to a previous correlation between such a determination and clinical information.

**[0016]** In one embodiment, the progression of the disorder may be tracked by using the methods of the invention to determine the severity of the disorder, e.g. global dementia severity). In another embodiment, the duration of the disorder up to the point of assessment may be determined using the methods of the invention. For example, expression of an Ig lambda chain C region (see spot 177, FIG. 6) may correlate with global dementia severity. Expression of a serum albumin precursor (see spot 165, FIG. 6) may show a negative correlation with the duration of the disease.

**[0017]** This method allows the type of Alzheimer's disease of a patient to be correlated to different types of prophylactic or therapeutic treatment available in the art, thereby enhancing the likely response of the patient to the therapy.

**[0018]** In some embodiments, more than one protein is differentially expressed, providing a multi-protein fingerprint of the nature or degree of the Alzheimer's disease. Preferably, at least four proteins are differentially expressed.

**[0019]** Conveniently, the patient sample used in the methods of the invention can be a tissue sample or body fluid sample such as a blood, plasma, serum or urine sample. Use of body fluids such as those listed is preferred because they can be more readily obtained from a subject. This has clear advantages in terms of cost, ease, speed and subject wellbeing. Blood, blood products such as plasma, and urine are particularly preferred.

**[0020]** The step of detecting the differentially expressed protein may be preceded by a depletion step to remove the most abundant proteins from the sample, as described below.

**[0021]** Preferably, at least one of the differentially expressed proteins is a protein shown in FIG. 6, FIG. 7 or FIG. 12. In preferred embodiments, the differentially expressed protein is apolipoprotein A-IV precursor, apolipoprotein C-III precursor, transthyretin, galectin 7, complement C4 precursor, alpha-2-macroglobulin precursor, Ig alpha-1 chain C, histone 2B, Ig lambda chain C region, fibrinogen gamma chain precursor, complement factor H, inter-alpha-trypsin heavy chain H4 precursor, complement C3 precursor, clusterin precursor, gamma or beta actin, haptoglobin precursor or the serum albumin precursor isoform found in spot ID no 2, 14, 15, 123, 165, 176 or 184 of FIG. 6 or fragments thereof. Preferred fragments are a C-terminal fragment of Apo-AIV or a C4 alpha region of complement C4 precursor Lacking the anaphylatoxin domain. For example, the fragment may comprise amino acid residues 270-309 of apolipoprotein A-IV; residues 1446-1744 of complement C4, or may be an N-terminal fragment of apolipoprotein A-IV which migrates as a polypeptide of 10-16 kD or a polypeptide of 28 kD in SDS-PAGE, or a fragment of any of the proteins in FIG. 7 with a molecular weight of 6430, 14640, 27147 or 14646 Da. Other preferred fragments comprise the areas indicated in bold in FIGS. 9, 10, and 13 to 19.

**[0022]** Preferred fragments are less than 50, less than 100, less than 150 less than 200, less than 250, less than 300, less than 350, less than 400, less than 500, less than 600, less than 700, less than 800, less than 900, less than 1000, less than 1100, less than 1200, less than 1300, less than 1400, less than 1500, less than 1600, less than 1700, less than 1800, less than 1900 or less than 2000 amino acids in length.

**[0023]** The expression of certain differentially expressed proteins may be increased in subjects with Alzheimer's disease as compared to control subjects. The expression of other differentially expressed proteins may be decreased in subjects with Alzheimer's disease as compared to control sub-

jects. FIGS. 6, 8 and 12 indicate whether the expression of the proteins disclosed therein is increased or decreased in Alzheimer's versus control subjects. It is thus clear from the figures whether an increase or decrease in expression is indicative of the disease state for all the proteins listed therein. Including the preferred proteins listed above.

**[0024]** Preferably, a differentially expressed protein shows a fold difference in expression of at least 1.5, at least 1.6, at least 1.7, at least 1.8, at least 1.9, at least 2.0, at least 2.5, at least 3, at least 3.5, at least 4, at least 5, at least 10 or more between the level found in patients with Alzheimer's versus control subjects.

**[0025]** The differentially expressed protein may be detected using an antibody specific to that protein, for example in an ELISA assay or Western blotting. Alternatively, the differentially expressed protein may be detected by, amongst others, 2D gel electrophoresis or mass spectrometry techniques including LS/MS/MS, MALDI-TOF or SELDI-TOF. The sample may be immobilised on a solid support for analysis.

**[0026]** In one embodiment, a diagnosis may be made solely on the basis of the pattern of spots on a 2D gel prepared from a subject sample. The pattern of spots obtained from Alzheimer's disease or MCI subjects may be compared directly with the pattern obtained from control subject samples, without the need for identifying individual proteins.

**[0027]** In one embodiment, an antibody sandwich technique where antibodies specific for one or more of the biomarkers is added and the immobilised antibodies capture the biomarker protein. The captured proteins are then detected using a second antibody that may be directly labelled with a signal generating agent (enzyme, fluorescent tag, radiolabel etc.) or may be detected using further amplification (labelled secondary antibody, streptavidin/biotin systems with enzyme, fluorophore, radiolabel etc.). Other immunological methods may include one-dimensional or two-dimensional gel electrophoresis of patient samples followed by transfer to a solid surface using techniques such as Western blotting and subsequent detection using antibodies specific for the AD biomarkers.

**[0028]** In an alternative embodiment, autoantibodies to the biomarkers may be detected by using the Western blotting approach described above using either samples from a patient or representative of AD and then detecting the presence of antibodies specific for the biomarker that are present in the blood of AD patients but not in controls.

**[0029]** The method may further comprise determining an effective therapy for treating the Alzheimer's disease.

**[0030]** In a further aspect, the present invention provides a method of treatment by the use of an agent that will restore the expression of one or more differentially expressed proteins in the Alzheimer's disease state towards that found in the normal state in order to prevent the development or progression of Alzheimer's disease. Preferably, the expression of the protein is restored to that of the normal state.

**[0031]** In a further aspect, the present invention provides a method whereby the pattern of differentially expressed proteins in a tissue sample or body fluid sample or urine of an individual with Alzheimer's disease is used to predict the most appropriate and effective therapy to alleviate the Alzheimer's disease.

**[0032]** Also provided is a method of screening an agent to determine its usefulness in treating a Alzheimer's disease, the method comprising:

- [0033] (a) obtaining a sample of relevant tissue taken from, or representative of, a subject having Alzheimer's disease symptoms, who or which has been treated with the agent being screened;
- [0034] (b) determining the presence, absence or degree of expression of the differentially expressed protein or proteins in the tissue from, or representative of, the treated subject; and,
- [0035] (c) selecting or rejecting the agent according to the extent to which it changes the expression, activity or amount of the differentially expressed protein or proteins in the treated subject having Alzheimer's disease symptoms.
- [0036] Optionally, the method may further comprise, prior to step (a), the step of establishing a paradigm in which at least one protein is differentially expressed in relevant tissue from, or representative of, subjects having Alzheimer's disease symptoms and normal subjects.
- [0037] Preferably, the agent is selected if it converts the expression of the differentially expressed protein towards that of a normal subject. More preferably, the agent is selected if it converts the expression of the protein or proteins to that of the normal subject.
- [0038] Also provided is a method of screening an agent to determine its usefulness in treating Alzheimer's disease, the method comprising:
- [0039] (a) obtaining over time samples of relevant tissue or body fluid taken from, or representative of, a subject having Alzheimer's disease symptoms, who or which has been treated with the agent being screened;
- [0040] (b) determining the presence, absence or degree of expression of a differentially expressed protein or proteins in said samples; and,
- [0041] (c) determining whether the agent affects the change over time in the expression of the differentially expressed protein in the treated subject having Alzheimer's disease symptoms.
- [0042] Optionally, the method may further comprise, prior to step (a), the step of establishing a paradigm in which at least one protein is differentially expressed in relevant tissue or body fluid from, or representative of, subjects having Alzheimer's disease symptoms and normal subjects; and/or
- [0043] establishing that expression of said differentially expressed protein diverges over time in subjects having Alzheimer's disease symptoms and normal subjects.
- [0044] Samples taken over time may be taken at intervals of weeks, months or years. For example, samples may be taken at monthly, two-monthly, three-monthly, four-monthly, six-monthly, eight-monthly or twelve-monthly intervals.
- [0045] A change in expression over time may be an increase or decrease in expression, compared to the initial level of expression in samples from the subject and/or compared to the level of expression in samples from normal subjects. The agent is selected if it slows or stops the change of expression over time.
- [0046] In the screening methods described above, subjects having differential levels of protein expression comprise:
- [0047] (a) normal subjects and subjects having Alzheimer's disease symptoms; and,
- [0048] (b) subjects having Alzheimer's disease symptoms which have not been treated with the agent and subjects Alzheimer's disease which have been treated with the agent.
- [0049] In alternative embodiments, the subjects having differential levels of protein expression comprise:
- [0050] (a) normal subjects who have and have not been treated with the agent; and one or both of
- [0051] (b) subjects having mild cognitive impairment who have and have not been treated with the agent; and
- [0052] (c) subjects having Alzheimer's disease symptoms who have and have not been treated with the agent.
- [0053] Preferably, the differential levels of protein expression are not observed in normal subjects who have and have not been treated with the agent.
- [0054] The subjects having Alzheimer's disease symptoms are preferably human subjects with Alzheimer's disease.
- [0055] Alternatively, the subjects having Alzheimer's disease symptoms may be an animal model such as mutant amyloid precursor protein (APP) transgenic mice, presenilin-1 (PS-1) transgenic mice, and/or double transgenic APP/PS-1 transgenic mice.
- [0056] The tissue or body fluid samples may be, for example, brain tissue, blood, plasma, serum, saliva or cerebro-spinal fluid samples.
- [0057] In one embodiment, the paradigm is established using two-dimensional (2D) gel electrophoresis carried out on the relevant tissue or a protein-containing extract thereof.
- [0058] In another embodiment, the paradigm is established using SELDI analysis of the relevant tissue or a protein-containing extract thereof. Preferably, the tissue or extract is immobilised on a solid support, for example a chip.
- [0059] Conveniently, a depletion step may be performed prior to 2D gel electrophoresis or SELDI analysis, to remove the most abundant proteins from the samples and reduce background.
- [0060] The method may further comprise the step of isolating a differentially expressed protein identified in the method, and optionally the step of characterising the isolated protein.
- [0061] Preferably, at least one of the differentially expressed proteins is a protein shown in FIG. 6, FIG. 7, FIG. 8 or FIG. 12 or a rodent equivalent thereof. In preferred embodiments, the differentially expressed protein is apolipoprotein A-IV precursor, apolipoprotein C-III precursor, transthyretin, galectin 7, complement C4 precursor, complement factor H, S100 calcium binding protein or ceruloplasmin, inter-alpha-trypsin heavy chain H4 precursor, complement C3 precursor, clusterin precursor, gamma or beta actin, haptoglobin precursor or fragments thereof. Preferred fragments are a C-terminal fragment of Apo-AIV or a C4 alpha region of complement C4 precursor lacking the anaphylatoxin domain. For example, the fragment may comprise amino acid residues 270-309 of apolipoprotein A-IV; residues 1446-1744 of complement C4.
- [0062] Preferred fragments will comprise one or more of the sequences highlighted in FIGS. 9, 10 and 13-19.
- [0063] In a further aspect, the invention provides a method of making a pharmaceutical composition which comprises having identified an agent using the method described above, the further step of manufacturing the agent and formulating it with an acceptable carrier to provide the pharmaceutical composition.
- [0064] In a further aspect, the invention provides a method of identifying a protein which is differentially expressed in relevant tissue or body fluid sample from subjects with mild cognitive impairment and/or subjects with Alzheimer's disease and normal subjects, comprising:

**[0065]** i) immobilising a tissue sample or body fluid sample or protein-containing extract thereof on a solid support

**[0066]** ii) analysing the immobilised proteins by surface enhanced laser desorption time of flight mass spectroscopy

**[0067]** iii) comparing the spectra obtained to detect differences in protein expression between Alzheimer's subjects and normal subjects.

**[0068]** Also provided is protein which is differentially expressed in relevant tissue from, or representative of subjects having differential levels of Alzheimer's disease symptoms and which is as obtainable by the methods described herein or by two-dimensional gel electrophoresis carried out on said tissue or a protein-containing extract thereof, the method comprising:

**[0069]** (a) providing non-linear immobilized pH gradient (ILG) strips of acrylamide polymer 3 mm×180 mm;

**[0070]** (b) rehydrating the IPG strips in a cassette containing 25 ml. of an aqueous solution of urea (8M), 3-[(cholamidopropyl)dimethylammonio]-1-propanesulphonate (CHAPS, 2% w/v), 0.5% IPG Pharmalyte and a trace of Bromophenol Blue;

**[0071]** (c) emptying the cassette of liquid, transferring the strips to an electrophoretic tray fitted with humid electrode wicks, electrodes and sample cups, covering the strips and cups with low viscosity paraffin oil;

**[0072]** (d) applying 200 micrograms of an aqueous solution of dried, powdered material of the relevant body tissue in urea (8M), CHAPS (4% w/v), Tris (40 mM), 0.5% IPG Pharmalyte and a trace of Bromophenol Blue to the sample cups, at the cathodic end of the IPG strips;

**[0073]** (e) carrying out isoelectric focusing on the gel at S1 500V step-n-hold (s/h) for 1 h; S2 500V s/h for 2 h; S3 1000V gradient (G) for 1 h; S4 1000V s/h for 2 h; S5 8000V G for 2 h and S6 8000V s/h for a time effective to enable the proteins to migrate in the strips to their pI-dependent final positions;

**[0074]** (f) equilibrating the strips within the tray with 100 ml of an aqueous solution containing Tris-HCl (50 mM) pH 6.8, urea (6M), glycerol (30% v/v), SDS (2% w/v) and DTT (10 mg/ml);

**[0075]** (g) replacing this solution by 100 ml. of an aqueous solution containing Tris-HCl (50 mM) pH 8.8, urea (6M), glycerol (30% v/v), SDS (2% w/v), iodoacetamide (25 mg/ml) and a trace of Bromophenol Blue and incubating for 20 minutes;

**[0076]** (h) providing a vertical gradient slab gel 160×200×1.5 mm of acrylamide/piperazine-diacrylyl cross-linker (9-16% T/2.6% C), polymerised in the presence of TEMED (0.5% w/v), ammonium persulphate (0.1% w/v) and sodium thiosulphate (5 mM), in Tris-HCl (0.375M) pH 8.8 as leading buffer;

**[0077]** (i) over-layering the gel with sec-butanol for about 2 hours, removing the overlay and replacing it with water;

**[0078]** (j) cutting the IPG gel strips to a size suitable for the second dimensional electrophoresis, removing 6 mm from the anode end and 14 mm from the cathode end;

**[0079]** (k) over-layering the slab gel with an aqueous solution of agarose (0.5% w/v) and Tris-glycine-SDS (25 mM-198 mM-0.1% w/v) as leading buffer, heated to 70° C. and loading the IPG gel strips onto the slab gel through this over-layered solution;

**[0080]** (l) running the second dimensional electrophoresis at a constant current of 40 mA at 8-12° C. for 5 hours; and (m) washing the gel.

**[0081]** This invention is based, in part, on systematic search strategies involving sensitive detection of proteins by 2D-electrophoresis. To aid the identification of differentially expressed protein a standard marker set of proteins such as those available from Genomic Solutions may be run on an extra lane to 2D electrophoresis.

**[0082]** The examples presented below demonstrate the successful use of the experimental paradigms of the invention to identify target proteins associated with Alzheimer's disease.

#### DEFINITIONS

**[0083]** "Differential expression", as used herein, refers to at least one recognisable difference in tissue or body fluid protein expression. It may be a quantitatively measurable, semi-quantitatively estimatable or qualitatively detectable difference in tissue protein expression. Thus, a differentially expressed protein (herein DEP) may be strongly expressed in tissue in the normal state and less strongly expressed or not expressed at all in tissue in the Alzheimer's disease state. Conversely, it may be strongly expressed in tissue in the Alzheimer's disease state and less strongly expressed or not expressed at all in the normal state. Further, expression may be regarded as differential if the protein undergoes any recognisable change between the two states under comparison.

**[0084]** The term "paradigm" means a prototype example, test model or standard.

**[0085]** Wherever a differentially expressible protein is used in the screening procedure, it follows that there must have been at some time in the past a preliminary step of establishing a paradigm by which the differential expressibility of the protein was pre-determined. Once the paradigm has been established, it need not be re-established on every occasion that a screening procedure is carried out. The term "establishing a paradigm" is to be construed accordingly.

**[0086]** "Relevant tissue" means any tissue involved in brain function, in particular tissue involved in Alzheimer's disease.

**[0087]** "Tissue/Body fluid . . . representative of . . . subjects" means any tissue or body fluid in which the above-mentioned biological change can be simulated for laboratory purposes and includes, for example, a primary cell culture or cell line derived ultimately from relevant tissue.

**[0088]** The term "subjects" includes human and animal subjects.

**[0089]** The treatments referred to above can comprise the administration of one or more drugs or foodstuffs, and/or other factors such as diet or exercise.

**[0090]** The differentially expressed proteins (DEPs) include "fingerprint proteins", "target proteins" or "pathway proteins".

**[0091]** The term "fingerprint protein", as used herein, means a DEP, the expression of which can be used, alone or together with other DEPs, to monitor or assess the condition of a patient suspected of suffering from Alzheimer's disease. Since these proteins will normally be used in combination, especially a combination of four or more, they are conveniently termed "fingerprint proteins", without prejudice to the possibility that on occasions they may be used singly or along with only one or two other proteins for this purpose. Such a fingerprint protein or proteins can be used, for example, to diagnose a particular type of Alzheimer's disease and thence to suggest a specific treatment for it.

**[0092]** The term “diagnosis”, as used herein, includes the provision of any information concerning the existence, non-existence or probability of the disorder in a patient. It further includes the provision of information concerning the type or classification of the disorder or of symptoms which are or may be experienced in connection with it. This may include, for example, diagnosis of the severity of the disorder. It encompasses prognosis of the medical course of the disorder, for example its duration, severity and the course of progression from MCI to Alzheimer’s disease.

**[0093]** Currently disease status is assessed by duration of disease from inception to present (longer duration equals more severe disease) and clinical assessment measures. These assessment measures include clinical tests for memory and other cognitions, clinical tests for function (abilities of daily living) and clinical assessments of global severity. Trials of potential therapies in AD are currently evaluated against these measures. The FDA and other medicines approval bodies require as part of these assessments measures of both cognition and global function. The Global Dementia Scale is one such measure of global function. It is assessed by rater assessment of severity including cognition and function against a standardised set of severity criteria.

**[0094]** The term “target protein”, as used herein, means a DEP, the level or activity of which can be modulated by treatment to alleviate Alzheimer’s disease. Modulation of the level or activity of the target protein in a patient may be achieved, for example, by administering the target protein, another protein or gene which interacts with it or an agent which counteracts or reduces it, for example an antibody to the protein, competitive inhibitor of the protein or an agent which acts in the process of transcription or translation of the corresponding gene.

**[0095]** The term “alleviate”, as used herein, in relation to Alzheimer’s disease means any form of reducing one or more undesired symptoms or effects thereof. Any amelioration of Alzheimer’s disease of the patient falls within the term “alleviation”. Amelioration may also include slowing down the progression of the disease.

**[0096]** Alternatively or additionally, the DEPs can interact with at least one other protein or with a gene involved in the regulation of brain function. Such other proteins are termed herein “pathway proteins” (PPs). The term is applied to the protein with which the DEP interacts, not to the DEP itself, although a pathway protein can be another DEP.

**[0097]** By way of example, embodiments of the present invention will now be described in more detail with reference to the accompanying figures.

#### BRIEF DESCRIPTION OF THE FIGURES

**[0098]** FIG. 1 shows spectra for the 6430 Da peak identified by SELDI analysis in normal (top) and Alzheimer’s disease (bottom) subjects.

**[0099]** FIG. 2 shows spectra for the 14640 Da peak identified by SELDI analysis in normal (top) and Alzheimer’s disease (bottom) subjects.

**[0100]** FIG. 3 shows spectra for the 27147 Da peak identified by SELDI analysis in normal (top) and Alzheimer’s disease (bottom) subjects.

**[0101]** FIG. 4 shows spectra for the 14646 Da peak identified by SELDI analysis in pooled normal (top) and Alzheimer’s disease (bottom) subjects.

**[0102]** FIG. 5 shows a silver stained gel obtained from material extracted from the chips used for SELDI analysis. The bands (1-6) excised and analysed by LC/MS/MS are indicated by arrows.

**[0103]** FIG. 6 shows differentially expressed proteins identified by 2D gel analysis and mass spectroscopy.

**[0104]** FIG. 7 shows differentially expressed proteins identified by SELDI and LC/MS/MS.

**[0105]** FIG. 8 shows differentially expressed proteins identified by qPST.

**[0106]** FIG. 9 shows the sequence coverage (indicated in bold) obtained for apolipoprotein A-IV (P06727) in the 14.6 kDa band isolated on the Q10 SAX2 SELDI chip. C-terminal residues 270-396 are underlined.

**[0107]** FIG. 10 shows sequence coverage (indicated in bold) obtained for Complement C4 precursor (P01028) in 2DE spot 164.

**[0108]** FIG. 11 shows a 2D gel obtained from the pre-depletion analysis. The differentially expressed spots are circled.

**[0109]** FIG. 12 lists the differentially expressed spots identified by the pre-depletion analysis. Column 3 gives the accession number for the human protein, column 4 the mean normalised spot volume in the control samples, column 6 the mean normalised spot volume in the disease samples, column 8 the mean normalised spot volume in the disease sample divided by that in the control sample, column 9 the significance (p-value) of the difference in spot volumes compared by Student’s t-test, column 10 the number of gels in the control group in which the spot was detected. CV is coefficient of variation.

**[0110]** FIG. 13 shows sequence coverage (indicated in bold) obtained for alpha-2 macroglobulin (P01023) in the pre-depletion analysis. The signal sequence is underlined.

**[0111]** FIG. 14 shows sequence coverage (indicated in bold) obtained for inter-alpha trypsin inhibitor heavy chain H4 precursor (Q14624) in the pre-depletion analysis. The signal sequence is underlined.

**[0112]** FIG. 15 shows sequence coverage (indicated in bold) obtained for complement C3 precursor (P01024) in the pre-depletion analysis. The signal sequence is underlined.

**[0113]** FIG. 16 shows sequence coverage (indicated in bold) obtained for clusterin precursor (P10909) in the pre-depletion analysis. The signal sequence is underlined.

**[0114]** FIG. 17 shows sequence coverage (indicated in bold for spot 832 and bold italic for spot 652) obtained for complement C4 precursor (P01028) in the pre-depletion analysis. The signal sequence is underlined.

**[0115]** FIG. 18 shows sequence coverage (indicated in bold) obtained for gamma actin (P63261) in the pre-depletion analysis. The signal sequence is underlined.

**[0116]** FIG. 19 shows sequence coverage (indicated in bold) obtained for haptoglobin precursor (P00738) in the pre-depletion analysis. The signal sequence is underlined.

#### DETAILED DESCRIPTION

**[0117]** Methods and compositions for the treatment of Alzheimer’s disease. Proteins termed ‘target proteins’ and/or fingerprint proteins are described which are differentially expressed in Alzheimer’s disease states relative to their expression in normal states. Methods for the identification of such fingerprint and target proteins are also described.

**[0118]** ‘Differential expression’ as used herein indicates that a protein is present at different levels in samples from normal and diseased subjects.

**[0119]** Also described below are methods for prognostic and diagnostic evaluation of Alzheimer’s disease states and for the identification of subjects exhibiting a predisposition to Alzheimer’s disease.

#### 1. Identification of Differentially Expressed and Pathway Proteins

**[0120]** In one embodiment, the present invention concerns methods for the identification of proteins which are involved in Alzheimer’s disease. Such proteins may represent proteins which are differentially expressed in Alzheimer’s disease states relative to their expression in normal states. Such differentially expressed proteins may represent ‘target’ or ‘fingerprint’ proteins.

**[0121]** Methods for the identification of such proteins are described in Section 1. Methods for the further characterisation of such differentially expressed proteins and for their identification as target and/or fingerprint proteins are presented below in Section 1.1.

**[0122]** ‘Differential expression’, as used herein, refers to both qualitative as well as quantitative differences in protein expression. Thus a differentially expressed protein may qualitatively have its expression activated or completely inactivated in normal versus Alzheimer’s disease state. Such a qualitatively regulated protein will exhibit an expression pattern within a given tissue, cell type or body fluid sample which is detectable in either control or Alzheimer’s disease subject, but not detectable in both. Alternatively, such a qualitatively regulated protein will exhibit an expression pattern within a given tissue, cell type or body fluid sample, which is detectable in either control or Alzheimer’s disease subjects but not detectable in both. ‘Detectable’, as used herein, refers to a protein expression pattern, which are detectable using techniques described herein.

**[0123]** Alternatively, a differentially expressed protein may have its expression modulated, i.e. quantitatively increased or decreased, in normal versus Alzheimer’s disease states. The degree to which expression differs in normal versus Alzheimer’s disease states need only be large enough to be visualised via standard characterisation techniques, such as silver staining of 2D-electrophoretic gels. Other such standard characterisation techniques by which expression differences may be visualised are well known to those skilled in the art. These include successive chromatographic separations of fractions and comparisons of the peaks, capillary electrophoresis, separations using micro-channel networks, including on a micro-chip, SELDI analysis and qPST analysis.

**[0124]** Chromatographic separations can be carried out by high performance liquid chromatography as described in Pharmacia literature, the chromatogram being obtained in the form of a plot of absorbance of light at 280 nm against time of separation. The material giving incompletely resolved peaks is then re-chromatographed and so on.

**[0125]** Capillary electrophoresis is a technique described in many publications, for example in the literature “Total CE Solutions” supplied by Beckman with their P/ACE 5000 system. The technique depends on applying an electric potential across the sample contained in a small capillary tube. The tube has a charged surface, such as negatively charged silicate glass. Oppositely charged ions (in this instance, positive ions) are attracted to the surface and then migrate to the appropriate

electrode of the same polarity as the surface (in this instance, the cathode). In this electroosmotic flow (EOF) of the sample, the positive ions move fastest, followed by uncharged material and negatively charged ions. Thus, proteins are separated essentially according to charge on them.

**[0126]** Micro-channel networks function somewhat like capillaries and can be formed by photocablation of a polymeric material. In this technique, a UV laser is used to generate high energy light pulses that are fired in bursts onto polymers having suitable UV absorption characteristics, for example polyethylene terephthalate or polycarbonate. The incident photons break chemical bonds with a confined space, leading to a rise in internal pressure, mini-explosions and ejection of the ablated material, leaving behind voids which form micro-channels. The micro-channel material achieves a separation based on EOF, as for capillary electrophoresis. It is adaptable to micro-chip form, each chip having its own sample injector, separation column and electrochemical detector: see J. S. Rossier et al., 1999, *Electrophoresis* 20: pages 727-731.

**[0127]** Surface enhanced laser desorption ionisation time of flight mass spectrometry (SELDI-TOF-MS) combined with ProteinChip technology can also provide a rapid and sensitive means of profiling proteins and is used as an alternative to 2D gel electrophoresis in a complementary fashion. The ProteinChip system consists of aluminium chips to which protein samples can be selectively bound on the surface chemistry of the chip (eg. anionic, cationic, hydrophobic, hydrophilic etc). Bound proteins are then co-crystallised with a molar excess of small energy-absorbing molecules. The chip is then analysed by short intense pulses of N2 320 nm UV laser with protein separation and detection being by time of flight mass spectrometry. Spectral profiles of each group within an experiment are compared and any peaks of interest can be further analysed using techniques as described below to establish the identity of the protein.

**[0128]** Quantitative protein sequence tag (qPST) technology may also be used to detect differentially expressed proteins. Briefly, the proteins in the samples for comparison are labelled with a stable isotope tag. A different isotope is used for each sample. The proteins are enzymatically cleaved and the labelled peptides in each sample are quantified by mass spectrometry. In this way, expression of equivalent proteins in the different samples can be compared directly by comparing the intensities of their respective isotopic peaks.

**[0129]** Detection of differentially expressed proteins may be preceded by a depletion step to remove the most abundant proteins from the sample. The large majority of the protein composition of serum/plasma consists of just a few proteins. For example, albumin, which is present at a concentration of 35-50 mg/ml, represents approximately 54% of the total protein content with IgG adding other 16%. In contrast, proteins changing in response to disease, for example as a result of tissue leakage, may circulate at 10 ng/ml. This vast dynamic range of protein concentrations represents a major analytical challenge and to overcome the problem, a multiple affinity depletion column can be used to remove the most highly abundant proteins (eg the 5, 6, 7, 8, 9 or 10 most highly abundant proteins). This enables the detection of changes in lower abundance ranges because more starting material can be used and there is less interference from the highly abundant molecules. Such a depletion strategy can be applied before any detection method.

**[0130]** Differentially expressed proteins may be further described as target proteins and/or fingerprint proteins. ‘Fin-

gerprint proteins', as used herein, refer to a differentially expressed protein whose expression pattern may be utilised as part of a prognostic or diagnostic Alzheimer's disease evaluation or which, alternatively, may be used in methods for identifying compounds useful for the treatment of Alzheimer's disease. A fingerprint protein may also have characteristics of a target protein or a pathway protein.

**[0131]** 'Target protein', as used herein, refers to a differentially expressed protein involved in Alzheimer's disease such that modulation of the level or activity of the protein may act to prevent the development of Alzheimer's disease. A target protein may also have the characteristics of a fingerprint protein or a pathway protein.

### 1.1 Characterisation of Differentially Expressed Proteins

**[0132]** Differentially expressed proteins, such as those identified via the methods discussed above, may be further characterised by using, for example, methods such as those discussed herein. Such proteins will be referred to herein as 'identified proteins'.

**[0133]** Analyses such as those described herein, yield information regarding the biological function of the identified proteins. An assessment of the biological function of the differentially expressed proteins, in addition, will allow for their designation as target and/or fingerprint proteins.

**[0134]** Specifically, any of the differentially expressed proteins whose further characterisation indicates that a modulation of the proteins expressed or a modulation of the proteins activity may ameliorate Alzheimer's disease will be designated 'target proteins', as defined above, in Section 1.

**[0135]** Any of the differentially expressed proteins whose expression pattern contributes to a protein 'fingerprint' profile correlative of Alzheimer's disease, will be designated a 'fingerprint protein'. 'Fingerprint profiles' will be more fully discussed below. It should be noted that each of the target proteins may also function as fingerprint proteins.

**[0136]** A variety of techniques can be utilised to further characterise the identified proteins. First, the corresponding nucleotide sequence of the identified protein may be obtained by utilising standard techniques well known to those of skill in the art, may, for example, be used to reveal homologies to one or more known sequence motifs which may yield information regarding the biological function of the identified protein.

**[0137]** Secondly, the biological function of the identified proteins may be more directly assessed by utilising relevant *in vivo* and *in vitro* systems. *In vivo* systems may include, but are not limited to, animal systems which naturally exhibit Alzheimer's disease-like symptoms and/or pathology, or ones which have been engineered to exhibit such symptom and/or pathology. Further, such systems may include systems for the further characterisation of Alzheimer's disease, and may include, but are not limited to, naturally occurring and transgenic animal systems.

**[0138]** *In vitro* systems may include cell lines derived from such animals or Alzheimer's disease subjects. Animal models may be used to generate cell lines, containing one or more cell types involved in Alzheimer's disease, that can be used as cell culture models for this disorder. While primary cultures derived from the transgenic animals of the invention may be utilised, the generation of continuous cell lines is preferred. For examples of techniques which may be used to derive a continuous cell line from the transgenic animals, see Small, et al., 1985, *Mol. Cell. Biol.* 5: 642-648.

**[0139]** Preferred transgenic animal models of Alzheimer's disease include mice overexpressing glycogen synthase kinase (GSK) (see Lucas et al (2001) *EMBO J.* 20, p27-39), mice overexpressing mutant alleles of APP or PS1 and double (APP/PS1) transgenic mouse models overexpressing mutant alleles of both APP and PS1. Double transgenic mice resulting from a cross between a mutant APP line Tg2576 and a mutant PS1M146L transgenic line is reported in Holcomb et al., *Nat. Med.* 1998 January; 4(1):97-100).

**[0140]** In further characterising the biological function of the identified proteins, the expression of these proteins may be modulated within the *in vivo* and/or *in vitro* systems, i.e. either overexpressed or underexpressed in, for example, transgenic animals and/or cell lines, and its subsequent effect on the system then assayed. Alternatively, the activity of the identified protein may be modulated by either increasing or decreasing the level of activity in the *in vivo* and/or *in vitro* system of interest, and its subsequent effect then assayed.

**[0141]** The information obtained through such characterisations may suggest relevant methods for the treatment of Alzheimer's disease using the protein of interest. For example, treatment may include a modulation of protein expression and/or protein activity. Characterisation procedures such as those described herein may indicate where such modulation should involve an increase or a decrease in the expression or activity of the protein of interest. Such methods of treatment are discussed below in Section 4.

## 2. Differentially Expressed Proteins

**[0142]** Identified proteins, which include differentially expressed proteins such as those identified in Section 1 above, are described herein. Specifically, the amino acid sequences of such identified proteins are described. Further, antibodies directed against the identified protein, and cell- and animal-based models by which the identified proteins may be further characterised and utilised are also discussed in this Section.

### 2.1 Antibodies Specific for Differentially Expressed or Pathway Proteins

**[0143]** The present invention also relates to methods for the production of antibodies capable of specifically recognising one or more differentially expressed or pathway protein epitopes. Such antibodies may include, but are not limited to, polyclonal antibodies, monoclonal antibodies (mAbs), humanised or chimeric antibodies, single chain antibodies, Fab fragments, F(ab')<sub>2</sub> fragments, fragments produced by a Fab expression library, anti-idiotypic (anti-Id) antibodies, and epitope-binding fragments of any of the above. Such antibodies may be utilised as part of Alzheimer's disease treatment methods, and/or may be used as part of diagnostic techniques whereby patients may be tested for abnormal levels of fingerprint, target, or pathway gene proteins, or for the presence of abnormal forms of such proteins.

**[0144]** For the production of antibodies to a differentially expressed or pathway protein, various host animals may be immunised by injection with a differentially expressed or pathway protein, or a portion thereof. Such host animals may include, but are not limited to, rabbits, mice and rats, to name but a few. Various adjuvants may be used to increase the immunological response, depending on the host species, including active substances such as lysolecithin, Pluronic polyols, polyanions, peptides, oil emulsions, keyhole limpet

hemocyanin, dinitrophenol, and potentially useful human adjuvant such as BCG bacille Calmette-Fuerin) and *Corynebacterium parvum*.

**[0145]** Polyclonal antibodies are heterogeneous populations of antibody molecules derived from the sera of animals immunised with an antigen, such as target proteins, or an antigenic functional derivative thereof. For the production of polyclonal antibodies, host animals such as those described above, may be immunised by injection with differentially expressed or pathway protein supplemented with adjuvants as also described above.

**[0146]** Monoclonal antibodies, which are homogeneous populations of antibodies to a particular antigen, may be obtained by any technique, which provides for the production of antibody molecules by continuous cell lines in culture. These include, but are not limited to, the hybridoma technique of Kohler and Milstein (1975, *Nature* 256: 495-497; and U.S. Pat. No. 4,376,110), the human  $\beta$ -cell hybridoma technique (Kosbor, et al., 1983, *Immunology Today* 4: 72; Cole, et al., 1983, *Proc. Natl. Acad. Sci. USA* 80: 2026-2030), and the EBV-hybridoma technique (Cole, et al., 1985, *Monoclonal Antibodies and Cancer Therapy*, Alan R. Liss Inc., pp. 77-96). Such antibodies may be of any immunoglobulin class including IgG, IgM, IgE, IgA, IgD and any subclass thereof. The hybridoma producing the mAb of this invention may be cultivated *in vitro* or *in vivo*. Production of high titers of mAbs *in vivo* makes this the presently preferred method of production.

**[0147]** In addition, techniques developed for the production of 'chimeric antibodies' (Morrison, et al., 1984, *Proc. Natl. Acad. Sci.* 81: 6851-6855; Neuberger, et al., 1984, *Nature* 312: 604-608; Takeda, et al., 1985, *Nature* 314: 452-454) by splicing the genes from a mouse antibody molecule of appropriate antigen specificity together with genes from a human antibody molecule of appropriate biological activity can be used. A chimeric antibody is a molecule in which different portions are derived from different animal species, such as those having a variable region derived from a murine mAb and a human immunoglobulin constant region.

**[0148]** Alternatively, techniques described for the production of single chain antibodies (U.S. Pat. No. 4,946,778; Bird, 1988, *Science* 242: 423-426; Huston, et al., 1988, *Proc. Natl. Acad. Sci. USA* 85: 5879-5883; and Ward, et al., 1989, *Nature* 334: 544-546) can be adapted to produce differentially expressed or pathway protein-single chain antibodies. Single chain antibodies are formed by linking the heavy and light chain fragments of the Fv region via an amino acid bridge, resulting in a single chain polypeptide.

**[0149]** Antibody fragments, which recognise specific epitopes, may be generated by known techniques. For example, such fragments include, but are not limited to, the F(ab')<sub>2</sub> fragments which can be produced by pepsin digestion of the antibody molecule and the Fab fragments which can be generated by reducing the disulfide bridges of the F(ab')<sub>2</sub> fragments. Alternative, Fab expression libraries may be constructed (Huse, et al., 1989, *Science* 246: 1275-1281) to allow rapid and easy identification of monoclonal Fab fragments with the desired specificity.

### 3 Assays for Amelioration of Alzheimer's Disease Symptoms

**[0150]** The differentially expressed proteins described herein may be used to test compounds for the ability to prevent or ameliorate Alzheimer's disease.

**[0151]** Such compounds may be tested in human subjects in clinical trials. Any compound which restores the expression

of a differentially expressed protein or proteins towards the normal level may be of potential use in treating Alzheimer's disease, i.e. reducing Alzheimer's disease symptoms or slowing the progression of Alzheimer's disease.

**[0152]** With regard to intervention, any treatments that restore or partially restore marker protein expression to normal levels should be considered as candidates for Alzheimer's disease therapeutic intervention. Dosages of test agents may be determined by deriving dose-response curves, as discussed in Section 6 below.

**[0153]** Similarly, any treatments that can prevent the development of Alzheimer's disease or prevent progression to levels of more advanced Alzheimer's disease should be considered as candidates for the Alzheimer's disease therapeutic intervention.

**[0154]** In addition, animal models of Alzheimer's disease, such as those described above, may be used to identify compounds capable of treating Alzheimer's disease symptoms. Such animal models may be used as test substrates for the identification of drugs, pharmaceuticals, therapies and interventions which may be effective in treating such disorders. The response of the animals to the exposure may be monitored by assessing the expression of the marker proteins and comparing it to that of wild-type mice.

**[0155]** Protein expression patterns may be utilised in conjunction with animal model systems to assess the ability of a compound to ameliorate Alzheimer's disease symptoms, or prevent the progression of Alzheimer's disease. For example, the expression pattern of one or more fingerprint proteins may form part of a fingerprint profile, which may then be used in such an assessment. Fingerprint profiles may be characterised for Alzheimer's disease states within the animal model systems. Subsequently, these known fingerprint profiles may be compared to ascertain the effect a test compound has to modify such fingerprint profiles, and to cause the profile to more closely resemble that of a more desirable fingerprint. For example, administration of a compound may cause the fingerprint profile of an Alzheimer's disease model system to more closely resemble the control system, or may prevent further changes in fingerprint profile. Administration of a compound may, alternatively, cause the fingerprint profile of a control system to begin to mimic an Alzheimer's disease state, which may, for example, be used in further characterising the compound of interest, or may be used in the generation of additional animal models.

### 4. Compounds and Methods for Treatment of Alzheimer's Disease

**[0156]** Described below are methods and compositions whereby Alzheimer's disease symptoms may be ameliorated or the progression of Alzheimer's disease slowed or halted. It is possible that Alzheimer's disease symptoms may be brought about, at least in part, by an abnormal level of target protein, or by the presence of a target protein exhibiting an abnormal activity. As such, the reduction in the level and/or activity of such target protein would bring about the amelioration Alzheimer's disease symptoms. Techniques for the reduction of target protein gene expression levels or target protein activity levels are discussed in Section 4.1.

**[0157]** Alternatively, it is possible that Alzheimer's disease symptoms may be brought about, at least in part, by the absence or reduction of the level of target protein expression, or a reduction in the level of a target protein's activity. As such, an increase in the level of target protein gene expression

and/or the activity of such proteins would bring about the amelioration Alzheimer's disease symptoms. Techniques for increasing target protein gene expression levels or target protein activity levels are discussed in Section 4.2.

#### 4.1 Compounds that Inhibit Expression, Synthesis or Activity of Target Proteins

**[0158]** As discussed above, target proteins involved in Alzheimer's disease may cause such disorders via an increased level of target protein activity. A variety of techniques may be utilised to inhibit the expression, synthesis, or activity of such target genes and/or proteins.

**[0159]** For example, compounds which exhibit inhibitory activity, may be used in accordance with the invention to prevent mild cognitive impairment or Alzheimer's disease symptoms. Such molecules may include, but are not limited to, peptides (such as, for example, peptides representing soluble extracellular portions of target protein transmembrane receptors), phosphopeptides, small organic or inorganic molecules, or antibodies (including, for example, polyclonal, monoclonal, humanised, anti-idiotypic, chimeric or single chain antibodies, and FAb, F(ab')<sub>2</sub> and FAb expression library fragments, and epitope-binding fragments thereof). Techniques for determination of effective doses and administration of such compounds are described below, in Section 6.1. Inhibitory antibody techniques are further described below, in Section 4.1.2.

**[0160]** Further, antisense, siRNA and ribozyme molecules, which inhibit expression of the target protein gene, may also be used in accordance with the invention to inhibit the aberrant target protein gene activity. Such techniques are described below, in Section 4.1.1; triple helix molecules may be utilised in inhibiting the aberrant target protein gene activity.

##### 4.1.1 Inhibitory Antisense, Ribozyme and Triple Helix Approaches

**[0161]** Antisense, ribozyme and triple helix molecules may be designed to reduce or inhibit either wild type, or if appropriate, mutant target protein gene activity. Techniques for the production and use of such molecules are well known to those of skill in the art.

**[0162]** Antisense RNA and DNA molecules act to directly block the translation of mRNA by hybridising to targeted mRNA and preventing protein translation. With respect to antisense DNA, oligodeoxy-ribonucleotides derived from the translation initiation site, e.g. between the -10 and +10 regions of the target gene nucleotide sequence of interest, are preferred.

**[0163]** Ribozymes are enzymatic RNA molecules capable of catalysing the specific cleavage of RNA. (For a review, see Rossi, J., 1994, *Current Biology* 4: 469-471). The mechanism of ribozyme action involves sequence specific hybridisation of the ribozyme molecule to complementary target RNA, followed by an endonucleolytic cleavage. The composition of ribozyme molecules must include one or more sequences complementary to the target protein mRNA, and must include the well known catalytic sequence responsible for mRNA cleavage. For this sequence, see U.S. Pat. No. 5,093,246. As such, within the scope of the invention are engineered hammerhead motif ribozyme molecules that specifically and efficiently catalyse endonucleolytic cleavage of RNA sequences encoding target proteins.

**[0164]** Specific ribozyme cleavage sites within any potential RNA target are initially identified by scanning the mol-

ecule of interest for ribozyme cleavage sites which include the following sequences, GUA, GUU and GUC. Once identified, short TNA sequences of between 15 and 20 ribonucleotides corresponding to the region of the target protein gene, containing the cleavage site may be evaluated for predicted structural features, such as secondary structure, that may render the oligonucleotide sequence unsuitable. The suitability of candidate sequences may also be evaluated by testing their accessibility to hybridise with complementary oligonucleotides, using ribonuclease protection assays.

**[0165]** RNA interference (RNAi) is a process of sequence-specific, post-transcriptional gene silencing in animals and plants, initiated by double-stranded RNA (dsRNA) that is homologous in sequence to the silenced gene. RNAi is mediated by short double-stranded RNA molecules (small interfering RNAs or siRNAs). siRNAs may be introduced into a cell as short RNA oligonucleotides of 10-15 bp, or as longer dsRNAs which are subsequently cleaved to produce siRNAs. The RNA may be introduced into the cell as RNA, or may be transcribed from a DNA or RNA vector.

**[0166]** siRNA molecules may be synthesized using standard solid or solution phase synthesis techniques which are known in the art. Alternatively, siRNA molecules or longer dsRNA molecules may be made recombinantly by transcription of a nucleic acid sequence, preferably contained within a vector as described below.

**[0167]** Another alternative is the expression of a short hairpin RNA molecule (shRNA) in the cell. shRNAs are more stable than synthetic siRNAs. A shRNA consists of short inverted repeats separated by a small loop sequence. One inverted repeat is complementary to the gene target. The shRNA is then processed into an siRNA which degrades the target gene mRNA and suppresses expression. shRNAs can be produced within a cell by transfecting the cell with a DNA construct encoding the shRNA sequence under control of a RNA polymerase III promoter, such as the human Hi or 7SK promoter. Alternatively, the shRNA may be synthesised exogenously and introduced directly into the cell. Preferably, the shRNA sequence is between 40 and 100 bases in length, more preferably between 40 and 70 bases in length. The stem of the hairpin is preferably between 19 and 30 base pairs in length. The stem may contain G-U pairings to stabilise the hairpin structure.

**[0168]** Nucleic acid molecules to be used in triple helix formation for the inhibition of transcription should be single stranded and composed of deoxynucleotides. The base composition of these oligonucleotides must be designed to promote triple helix formation via Hoogsteen base pairing rules, which generally require sizeable stretches of either purines or pyrimidines to be present on one strand of a duplex. Nucleotide sequences may be pyrimidine-based, which will result in TAT and CGC<sup>+</sup> triplets across the three associated strands of the resulting triple helix. The pyrimidine-rich molecules provide base complementary to a purine-rich region of a single strand of the duplex in a parallel orientation to that strand. In addition, nucleic acid molecules may be chosen that are purine-rich, for example, containing a stretch of G residues. These molecules will form a triple helix with a DNA duplex that is rich in GC pairs, in which the majority of the purine residues are located on a single strand of the targeted duplex, resulting in GGC triplets across the three strands in the triple helix.

**[0169]** Alternatively, the potential sequences that can be targeted for triple helix formation may be increased by cre-

ating a so-called "switchback" nucleic acid molecule. Switchback molecules are synthesised in an alternating 5'-3', 3'-5' manner, such that they base pair with first one strand of a duplex and then the other, eliminating the necessity for a sizeable stretch of either purines or pyrimidines to be present on one strand of a duplex.

[0170] Anti-sense RNA and DNA, siRNAs, ribozyme and triple helix molecules of the invention may be prepared by any method known in the art for the synthesis of DNA and RNA molecules. They include techniques for chemically synthesising oligodeoxyribonucleotides and oligo-ribonucleotides well known in the art such as, for example, solid phase phosphoramidite chemical synthesis. Alternatively, RNA molecules may be generated by in vitro and in vivo transcription of DNA sequences encoding the antisense RNA molecule. Such DNA sequences may be incorporated into a wide variety of vectors, which incorporate suitable RNA polymerase promoters such as the T7 or SP6 polymerase promoters. Alternatively, antisense cDNA constructs that synthesise antisense RNA constitutively inducibly, depending on the promoter used, can be introduced stably into cell lines.

#### 4.1.2 Antibodies for the Inhibition of Target Protein

[0171] Antibodies that are both specific for target protein and interfere with its activity may be used to inhibit target protein function. Where desirable, antibodies specific for mutant target protein, which interferes with the activity of such mutant target product, may also be used. Such antibodies may be generated, using standard techniques described in Section 2. above, against the proteins themselves or against peptides corresponding to portions of the proteins. The antibodies include, but are not limited to, polyclonal, monoclonal, Fab fragments, single chain antibodies, chimeric antibodies, etc.

[0172] In instances where the target gene protein is intracellular and whole antibodies are used, internalising antibodies may be preferred. However, lipofectin or liposomes may be used to deliver the antibody or a fragment of the Fab region, which binds to the target protein epitope into cells. Where fragments of the antibody are used, the smallest inhibitory fragment, which binds to the target protein's binding domain, is preferred. For example, peptides having an amino acid sequence corresponding to the domain of the variable region of the antibody that binds to the target protein may be used. Such peptides may be synthesised chemically or produced via recombinant DNA technology using methods well known in the art (e.g. see Creighton, 1983, supra; and Sambrook et al, 1989, supra).

[0173] Alternatively, single chain neutralising antibodies, which bind to intracellular target protein epitopes, may also be administered. Such single chain antibodies may be administered, for example, by expressing nucleotide sequences encoding single-chain antibodies within the target cell population by utilising, for example, techniques such as those described in Marasco et al (Marasco, W. et al, 1993, Proc. Natl. Acad. Sci. USA, 90: 7889-7893).

[0174] In instances where the target protein is extracellular, or is a transmembrane protein, any of the administration techniques described below, in Section 6, which are appropriate for peptide administration may be utilised to effectively administer inhibitory target protein antibodies to their site of action.

#### 4.2 Methods for Restoring Target Protein Activity

[0175] Target proteins that cause Alzheimer's disease may be underexpressed in Alzheimer's disease disorder situations.

Alternatively, the activity of target protein may be diminished, leading to the development of Alzheimer's disease symptoms. Described in this Section are methods whereby the level of target protein may be increased to levels wherein Alzheimer's disease symptoms are prevented or ameliorated. The level of target protein activity may be increased, for example, by either increasing the level of target protein present or by increasing the level of active target protein which is present.

[0176] For example, a target protein, at a level sufficient to ameliorate Alzheimer's disease symptoms may be administered to a patient exhibiting such symptoms. Any of the techniques discussed below, in Section 6, may be utilised for such administration. One of skill in the art will readily know how to determine the concentration of effective, non-toxic doses of the normal target protein, utilising techniques such as those described below.

[0177] Further, patients may be treated by gene replacement therapy. One or more copies of a normal target protein gene or a portion of the gene that directs the production of a normal target protein with target protein gene function, may be inserted into cells, using vectors which include, but are not limited to, adenovirus, adeno-associated virus, and retrovirus vectors, in addition to other particles that introduce DNA into cells, such as liposomes. Additionally, techniques such as those described above may be utilised for the introduction of normal target protein gene sequences into human cells.

[0178] Cells, preferably autologous cells, containing normal target protein gene sequences may then be introduced or reintroduced into the patient at positions which allow for the prevention or amelioration of Alzheimer's disease symptoms. Such cell replacement techniques may be preferred, for example, when the target protein is a secreted, extracellular protein.

[0179] Additionally, antibodies may be administered which specifically bind to a target protein and by binding, serve to, either directly or indirectly, activate the target protein function. Such antibodies can include, but are not limited to, polyclonal, monoclonal, Fab fragments, single chain antibodies, chimeric antibodies and the like. The antibodies may be generated using standard techniques such as those described above, in Section 2.3, and may be generated against the protein themselves or against proteins corresponding to portions of the proteins. The antibodies may be administered, for example, according to the techniques described above.

#### 5. Pharmaceutical Preparations and Methods of Administration

[0180] The identified compounds, nucleic acid molecules and cells that affect target protein expression, synthesis and/or activity can be administered to a patient at therapeutically effective doses to prevent or to treat or to ameliorate Alzheimer's disease. A therapeutically effective dose refers to that amount of the compound sufficient to result in amelioration of symptoms Alzheimer's disease, or alternatively, to that amount of a nucleic acid molecule sufficient to express a concentration of protein which results in the amelioration of such symptoms.

##### 5.1 Effective Dose

[0181] Toxicity and therapeutic efficacy of such compounds can be determined by standard pharmaceutical procedures in cell cultures or experimental animals, e.g. for

determining by  $ED_{50}$  (the dose therapeutically effective in 50% of the population) and by determining the  $ED_{50}$  of any side-effects (toxicity— $TD_{50}$ ). The dose ratio between toxic and therapeutic effects is the therapeutic index and it can be expressed as the ratio  $TD_{50}/ED_{50}$ . Compounds, which exhibit large therapeutic indices, are preferred. While compounds that exhibit toxic side effects may be used, care should be taken to design a delivery system that targets such compounds to the site of affected tissue in order to minimise potential damage to uninfected cells and, thereby, reduce side effects.

**[0182]** The data obtained from the animal studies can be used in formulating a range of dosage for use in humans. The dosage of such compounds lies preferably within a range of circulating concentrations that include the  $ED_{50}$  with little or no toxicity. The dosage may vary within this range depending upon the dosage form employed and the route of administration utilised.

## 5.2 Formulations and Use

**[0183]** Pharmaceutical compositions for use in accordance with the present invention may be formulated in conventional manner using one or more physiologically acceptable carriers or excipients.

**[0184]** Thus, the compounds and their physiologically acceptable salts and solvates may be formulated for administration by inhalation or insufflation (either through the mouth or the nose) or oral, buccal, parenteral and rectal administration.

**[0185]** For oral administration, the pharmaceutical compositions may take the form of, for example, tablets or capsules prepared by conventional means with pharmaceutically acceptable excipients such as binding agents (e.g. pre-gelatinised maize starch, polyvinylpyrrolidone or hydroxypropyl methyl-cellulose); fillers (e.g. lactose, microcrystalline cellulose or calcium hydrogen phosphate); lubricants (e.g. magnesium stearate, talc or silica); disintegrants (e.g. potato starch or sodium starch glycolate); or wetting agents (e.g. sodium lauryl sulphate). The tablets may be coated by methods well known in the art. Liquid preparations for oral administration may take the form of, for example, solutions, syrups or suspensions, or they may be presented as a dry product for constitution with water or other suitable vehicle before use. Such liquid preparations may be prepared by conventional means with pharmaceutically acceptable additives such as suspending agents (e.g. sorbitol syrup, cellulose derivatives or hydrogenated edible fats); emulsifying agents (e.g. lecithin or acacia); and preservatives (e.g. methyl or propyl-p-hydroxybenzoates or sorbic acid). The preparations may also contain buffer salts, flavouring, colouring and sweetening agents as appropriate.

**[0186]** Preparations for oral administration may be suitably formulated to give controlled release of the active compound.

**[0187]** For buccal administration the compositions may take the form of tablets or lozenges formulated in conventional manner.

**[0188]** For administration by inhalation, the compounds for use according to the present invention are conveniently delivered in the form of an aerosol spray presentation from pressurised packs or a nebuliser, with the use of a suitable propellant, e.g. dichlorodifluoromethane, trichlorofluoromethane, dichloro-tetrafluoroethane, carbon dioxide or other suitable gas. In the case of a pressurised aerosol the dosage unit may be determined by providing a valve to deliver a metered amount. Capsules and cartridges of, e.g. gelatin, for use in an

inhaler or insufflator may be formulated containing a powder mix of the compound and a suitable powder base such as lactose or starch.

**[0189]** The compounds may be formulated for parenteral administration by injection, e.g. by bolus injection or continuous infusion. Formulations for injection may be presented in unit dosage form, e.g. in ampoules or in multi-dose containers, with an added preservative. The compositions may take such forms as suspensions, solutions or emulsions in oily or aqueous vehicles, and may contain formulatory agents such as suspending, stabilising and/or dispersing agents. Alternatively, the active ingredient may be in powder form for constitution with a suitable vehicle, e.g. sterile pyrogen-free water, before use.

**[0190]** The compounds may also be formulated in rectal compositions such as suppositories or retention enemas, e.g. containing conventional suppository bases such as cocoa butter or other glycerides.

**[0191]** In addition to the formulations described previously, the compounds may also be formulated as a depot preparation. Such long acting formulations may be administered by implantation, for example, subcutaneously or intramuscularly) or by intramuscular injection. Thus, for example, the compounds may be formulated with suitable polymeric or hydrophobic materials (for example, as an emulsion in an acceptable oil) or ion exchange resins, or as sparingly soluble derivatives, for example, as a sparingly soluble salt.

**[0192]** The compositions may, if desired, be presented in a pack or dispenser device, which may contain one or more unit dosage forms containing the active ingredient. The pack may, for example, comprise metal or plastic foil, such as blister pack. The pack or dispenser device may be accompanied by instructions for administration.

## 6. Diagnosis of Alzheimer's Disease

**[0193]** A variety of methods may be employed for the diagnosis of Alzheimer's disease, monitoring progression of mild cognitive impairment and Alzheimer's disease, the predisposition to Alzheimer's disease, and for monitoring the efficacy of any Alzheimer's disease compounds during, for example, clinical trials and for monitoring patients undergoing clinical evaluation for the treatment of Alzheimer's disease. The differentially expressed and fingerprint proteins can also be used to define the nature or degree of Alzheimer's disease to aid in the identification and/or selection of treatments for the disorder.

**[0194]** Alzheimer's disease is characterised by a progressive, insidious onset, two or more deficits in cognitive function, and the absence of any other illnesses that could account for the dementia

**[0195]** In addition to memory loss, there may be disorientation, poor attention span, and language impairment. There is likely to be a decline in the activity of daily living, and possibly also impaired perception and personality changes. Behavioural symptoms include delusions, aggression, agitation, anger, wandering, hallucinations, and sleep disturbance.

**[0196]** A simple test assessing orientation, registration, calculations and attention, recall, language, and visual-spatial function may be used for an initial diagnosis.

**[0197]** Structural imaging by standard CT or MRI may also be used. Typically a non-contrast head CT scan suffices, but MRI is preferred for those who have hypertension or diabetes, who are at risk for cerebral vascular disease.

[0198] Alzheimer's disease may be confirmed histologically by autopsy or brain biopsy showing neurofibrillary tangles and senile plaques.

[0199] Identifying individuals at risk from Alzheimer's disease may involve diagnosis of mild cognitive impairment (MCI). (MCI) may be a transitional state between normal aging and dementia. There are different types of MCI. There may be cognitive impairment in multiple areas of cognitive function, in addition to memory. In some cases, memory is normal but some other domain of cognitive function is abnormal.

[0200] Amnesic MCI appears to be a risk state for the development of Alzheimer's disease. Amnesic impairment is defined by subjective memory complaints. These patients have poor memory performance for their age and education on formal testing when compared to age-matched peers. General cognitive functions and the ability to perform the activities of daily living should be entirely normal. The amnesic type of MCI is associated with hippocampal atrophy, neurofibrillary tangles in the medial temporal lobes, and elevated levels of Tau in the cerebrospinal fluid (CSF).

[0201] Methods for diagnosing Alzheimer's disease or predisposition to Alzheimer's disease may also, for example, utilise reagents such as the differentially expressed and fingerprint proteins described above, and antibodies directed against differentially expressed, as described above. Specifically, such reagents may be used for the detection of either an over- or an under-abundance of target protein relative to the normal state.

[0202] The methods described herein may be performed, for example, by utilising pre-packaged diagnostic kits comprising at least one specific differentially expressed/fingerprint protein or anti-differentially expressed/fingerprint protein antibody reagent described herein, which may be conveniently used, e.g. in clinical settings, to diagnose patients exhibiting Alzheimer's disease symptoms.

[0203] Any cell type, tissue or body fluid in which the fingerprint protein is expressed may be utilised in the diagnostics described herein. Examples of suitable samples types include cell samples, tissue samples, and fluid samples such as blood, urine, serum, saliva, cerebrospinal fluid or plasma.

[0204] Among the methods which can be utilised herein, are methods for monitoring the efficacy of compounds in clinical trials for the treatment of Alzheimer's disease. Such compounds can, for example, be compounds such as those described above, in Section 4. Such a method comprises detecting, in a patient sample, a protein, which is differentially expressed in the Alzheimer's disease state relative to its expression in a normal state.

[0205] During clinical trials, for example, the expression of a single differentially expressed protein, or alternatively, a fingerprint pattern of a cell involved in Alzheimer's disease can be determined in the presence or absence of the compound being tested. The efficacy of the compound can be followed by comparing the expression data obtained to the corresponding known expression patterns in a normal state. Compounds exhibiting efficacy are those which alter the protein expression and/or the fingerprint pattern to more closely resemble that of the normal state, or which stabilise protein expression and/or the fingerprint pattern i.e. prevent progression of the disease.

[0206] The detection of the protein differentially expressed in the Alzheimer's disease state relative to their expression in a normal state can also be used for monitoring the efficacy of

potential compounds for the treatment of Alzheimer's disease during clinical trials. During clinical trials, for example, the level and/or activity of the differentially expressed protein can be determined in relevant cells and/or tissues and/or body fluids in the presence or absence of the compound being tested. The efficacy of the compound can be followed by comparing the protein level and/or activity data obtained to the corresponding known levels/activities for the cells and/or tissues and/or body fluids in a normal state. Compounds exhibiting efficacy are those which alter the pattern of the cell and/or tissue sample and/or body fluid from an Alzheimer's disease subject to more closely resemble that of the normal state or which stabilise the pattern i.e. prevent progression of the disease.

## EXPERIMENTAL

### Subjects

[0207] The study population is derived from a large, longitudinally assessed, community based population of people with AD (NINCDS-ADRDA probable), other dementias and normal elderly persons. Samples are available on over 1000 subjects, all whom have detailed clinical assessment. Clinical research data includes systematic diagnostic, cognitive and behavioural assessments. Approximately 50 ml blood (4x10 ml in BD vacutainer K3E 15% tubes and 1x10 ml in extainer) is drawn from each subject. Subjects have had no food or fluid intake for more than 2 hours prior to collection. One BD vacutainer K3E (plasma) and extainer (serum) is used for proteomics study. The serum/plasma samples collected for proteomics are spun at 3000 rpm for 8 min within 2 h of collection.

### Analysis

[0208] Serum/plasma samples were lysed and rehydrated in a 2D lysis buffer consisting of 8M Urea, 2% w/v CHAPS, 0.5% IPG Pharmalyte (pH 3-10; Amersham Biotech, UK). The lysed samples were then subjected to isoelectrofocusing 18 cm 3-10 NL Immobiline pH gradient strips. IPG electrofocusing of the rehydrated strips was carried out for 16 h using the following protocol: S1 500V step-n-hold (s/h; i.e. the electric current applied to the strip is gradually increased in steps holding at particular settings for the times indicated) for 1 h; S2 500V s/h for 2 h; S3 1000V gradient (G) for 1 h; S4 1000V s/h for 2 h; S5 8000V G for 2 h and a final step S6 8000V s/h for 8 h with the IPGphor™.

[0209] Electrofocused IPG strips were then equilibrated in a SDS equilibration buffer (50 mM Tris-HCl pH8.8, 6M urea, 30% (v/v) glycerol, 2% SDS, and trace amount of bromophenol blue) with 10 mg/ml dithiothreitol (DTT) for 20 min, followed by 20 min step with 25 mg/ml Iodoacetamide. The equilibrated strips were then separated on a 10% acrylamide second dimension electrophoresis gel using the Ettan Dalt II system.

[0210] Following the electrophoresis the gels were placed in separate staining boxes and fixed using 40% ethanol/10% acetic acid for 1 h at room temperature and then stained according to Hochstrasse protocol (Table 1). Gel analysis was performed using the Melanie 3 software and Mann and Whitney rank sum test and False Discovery Rate statistical analysis was carried out to compare subject groups.

TABLE 1

Hochstrasse staining protocol	
Staining step	Time
Fix 40% ethanol/10% acetic acid	1 h
Soak in 5% ethanol/5% acetic acid	3 hr or overnight
Wash in water	5 min
Soak in 0.5M Sodium acetate, 1% gluteraldehyde	1.5 h
Wash	4 × 15 min
Soak in 0.05% Naphthalene sulphonic acid	2 × 30 min
Rinse in water	4 × 15 min
Silver stain (12 g silver, 20 ml ammonium hydroxide and 3 ml 10M sodium hydroxide)	25 min
Wash	4 × 4 min
Develop (0.005% citric acid and 0.1% formaldehyde)	As required
Stop solution (5% tris and 2% acetic acid)	1-2 h
Storage solution (35% ethanol and 5% glycerol)	

#### Sample Preparation

[0211] In-gel reduction, alkylation and digestion (with trypsin) were performed prior to subsequent analysis by mass spectrometry. Cysteine residues were reduced with DTT and derivatized by treatment with iodoacetamide to form stable carbamidomethyl (CAM) derivatives. Trypsin digestion was carried out overnight at room temperature after an initial 1 hr incubation at 37° C.

#### MALDI-TOF Mass Spectrometry

[0212] The digested sample (3 µl) was desalted and concentrated using ZipTipC18 microtips (Millipore). Peptides were eluted in 4 µl 50% acetonitrile/0.1% trifluoroacetic acid. 0.5 µl was then loaded onto a target plate with 0.5 µl matrix ( $\alpha$ -Cyano-4-hydroxy-cinnamic acid). Peptide mass fingerprints were acquired using a Voyager De-Pro, MALDI-TOF mass spectrometer (Applied Biosystems). The mass spectra were acquired in reflectron mode with delayed extraction. An autolytic tryptic peptide of mass 2163.0569 Da was then used to lock mass the acquired spectra, to achieve a mass accuracy of better than 30 ppm.

#### LC/MS/MS

[0213] Peptides were extracted from the gel pieces by a series of acetonitrile and aqueous washes. The extract was pooled with the initial supernatant and lyophilised. Each sample was then resuspended in 6 µL of 50 mM ammonium bicarbonate and analysed by LC/MS/MS. Chromatographic separations were performed using an Ultimate LC system (Dionex, UK). Peptides were resolved by reverse phase chromatography on a 75 µm C18 PepMap column. A gradient of acetonitrile in 0.05% formic acid was delivered to elute the peptides at a flow rate of 200 nl/min. Peptides were ionised by eLectrospray ionisation using a Z-spray source fitted to a QTOFmicro (Waters Corporation). The instrument was set to run in automated switching mode, selecting precursor ions based on their intensity, for sequencing by collision-induced fragmentation. The MS/MS analyses were conducted using

collision energy profiles that were chosen based on the  $m/z$  and the charge state of the peptide.

#### Results

[0214] Analysis of all control group (n=50) and case group [0215] (n=50) 2D gel images and subjecting them to statistical analysis. A total of 16 protein spots show a significant result ( $p < 0.05$ ) (see FIG. 6).

[0216] The results shown in FIG. 6 are unambiguous matches as they are based on exact matching of multiple MS/MS spectra. The sequence of selected proteins showing the peptide coverage obtained is given in FIGS. 8 to 10.

#### Class Prediction Using Peptide Fingerprinting

[0217] A class prediction analysis was performed in order to determine whether the pattern of peptide spots on 2DGE could predict caseness as determined clinically. Support Vector Machines (SVM), a supervised machine learning algorithm for prediction of class set in a group based upon a training set of data<sup>13</sup>, was used. SVM is most typically used in microarray analyses. However the statistical challenges are similar for proteomics and SVM has previously been used as a class prediction model for various proteomic studies<sup>14,15</sup>. Using GeneSpring (Silicon Genetics) the original 25 cases and 25 controls were designated as a training set and then the replication 25 cases and 25 controls designated as a test set. All identified proteins were used as possible identifiers and with the parameters Polynomial Dot Product Order 1 and Diagonal Scaling Factor 1; 34 of the 50 test-samples were correctly identified as being either cases or controls. Sensitivity was 56% and specificity 80% using SVM analysis of 2DGE data alone.

#### Identification of Peptides that Differentiate Between Cases and Controls

[0218] The normalised spot optical density in both the initial set of cases and controls and the replication set was compared. Mean differences between patients and controls at each spot were compared using the Wilcoxon rank-sum (Mann-Whitney) tests. The p-values for the nul hypothesis of no mean differences were saved, sorted by increasing value and ranked. A false discovery rate index (FDR) was computed as the ratio of the rank number and the theoretical probability (which is the rank number divided by the total number of spots). Fifteen spots were identified to have a FDR of less than 0.50 These were then identified using LC-MS/MS (FIG. 6).

#### Correlation of Peptide Spots with Clinical Parameters

[0219] Although the cases and controls were similarly aged it was possible that the observed peptide or spot differences were due to an association with age, gender or APOE genotype. A correlation analysis was thus performed for the 15 spots that differed between cases and controls in all 100 subjects with age, gender and APOE genotype. Data was first scaled to unit variance so as to standardise the scales upon which the variables were compared (i.e. each value was divided by the standard deviation of all the values for that particular variable). The Pearson correlation coefficient was then calculated. Cases with missing values were excluded pairwise. There were no strong correlations of any spot with age, gender or APOE. Two spots weakly correlated with age, two with gender and one with APOE genotype.

[0220] An ideal biomarker would not only be different between cases and controls but would be a marker of disease progression. The 15 spots showing case-control differences

in all 50 cases were thus correlated with duration of dementia and severity as measured by MMSE and GDS. Two spots correlated moderately and significantly with measures of disease progression and global dementia severity ( $r^2=0.29$  with spot 177) and duration of disease ( $r^2=-0.29$  with spot 166). Thus, one peptide—an Ig lambda chain C region (spot 177) correlates with global dementia severity. The other marker of disease progression examined, duration, shows a negative correlation with albumin (Spot 165).

#### Pre-Depletion Analysis

**[0221]** In these experiments, human plasma samples were depleted to remove the 6 most abundant proteins before the 2D gel electrophoresis step.

#### Methods

**[0222]** 60 human plasma samples (30 Controls and 30 disease subjects) were depleted using a removal column from Agilent. The samples were separated by 2D electrophoresis (pH 3-10 NL, 10% SDS-PAGE, 75  $\mu$ g protein load). Gels were silver-stained, scanned (8 bit, 200 dpi) and quantitatively analysed with Progenesis. To pick gel plugs from preparative gels, several control samples were mixed together and 3 gels run (2 gels with 205  $\mu$ g protein load and 1 gel with 350  $\mu$ g protein load). The same strategy was used with disease samples to run preparative gels. Protein spots were then destained, trypsinated and polypeptides were spotted onto MALDI target with Spot handling workstations (GE Amersham Biosciences). Peptide profiles generated were analysed with Ms-Fit programme in combination with the Swiss-Prot database.

#### Results

**[0223]** Gel images of proteins extracted from control and disease samples were analysed with Progenesis (v2005). Each group (Control and Disease) were based on 29 analytical gels. Spot detection, matching were performed with Progenesis, then the spot data were exported to Excel and a macro developed in-house was used to calculate coefficient of variation (CV %), T-Test and Regulation factor or change.

**[0224]** 11 spots were selected for analysis based on the following selection criteria: spots have to be found within at least 60% of gels, 2-fold up/down regulation and  $p$  value < 0.005. FIG. 11 displays the location of these 11 spots in the reference gel. This image corresponds to the 2D profile of proteins extracted from a control sample. The normalised volumes of the 11 spots detected in gels was analysed and is given in FIG. 12.

**[0225]** All protein spots were picked from 3 to 6 different preparative gels and submitted to MS analyses. All protein spots were successfully identified as shown in FIG. 12. In the down regulated spots, we found two spots of alpha-2-macroglobulin precursor (174; 178), one spot of inter-alpha-trypsin inhibitor heavy chain H4 precursor (232), one spot of a mix of complement C3 precursor with clusterin precursor (712) as indicated in grey in FIG. 12 and one spot of complement C3 precursor (713). In the up regulated spots, we found two spots of complement C4 precursor (652; 832), one spot of actin (675) and three spots of haptoglobin precursor (702; 703; 706).

**[0226]** To estimate the coverage of proteins identified and to discriminate the different chains or isoforms, for each spot, a common list of peptide masses was established.

**[0227]** This list regroups all peptide masses matched corresponding to the same spot picked in 3 to 6 preparative gels. The amino acids belonging to the peptides matched are underlined in FIGS. 13 to 19.

#### Discussion

**[0228]** The 11 spots analysed identified 7 regulated proteins between control- and disease samples, namely alpha-2-macroglobulin, inter-alpha-trypsin inhibitor heavy chain H4, complement C3, complement C4, actin cytoplasmic and haptoglobin.

**[0229]** Alpha-2-macroglobulin protein is able to inhibit all four classes of proteinases by a unique "trapping" mechanism. The observed molecular weight of the gel spots (~100 kDa, FIG. 11, spots 174; 178), matched by PMF, cover mainly the N-terminus of the protein (FIG. 14). The protein identified may thus correspond to a fragment of the full-length sequence of alpha-2-macroglobulin. As spots identified as alpha-2-macroglobulin belong to the same chain of spots (FIG. 11), it is possible that the difference between the two spots may be due to a post-translation modification.

**[0230]** There are two isoforms of inter-alpha-trypsin inhibitor heavy chain H4. Isoform 1 has 930 amino-acids and isoform 2 has 914 amino-acids. This protein is cleaved by plasma kallikrein to yield 100 kDa and a 35 kDa fragments. The resulting 100 kDa fragment is further converted to a 70 kDa fragment. The masses matched by PMF cover the sequence up to amino acid (aa) 688. This sequence corresponds to isoform 1 and may include the 70 kDa fragment and a short potentially active peptide. In this case, there is good agreement between the theoretical molecular weight and pI (74 kDa and 6.04 respectively) and the observed ones from the gel spot (see FIG. 11, spot 232).

**[0231]** Complement C3 precursor plays a central role in the activation of the complement system. This protein contains two chains (alpha and beta). We identified peptide masses covering the sequence from aa 714 to aa 1360 (FIG. 15), which corresponds to the alpha chain of complement C3. The theoretical molecular weight and pI of the alpha chain (115 kDa and 5.55 respectively) are not in agreement with the observed ones from the gel spots (see FIG. 11, spots 712, 713). The alpha chain is processed into different fragments. It appears that a temporary peptide appearing during the activation of complement system. As spots identified as complement C3 belong to the same chain of spots (FIG. 11), it is possible that the difference is due to a post-translational modification.

**[0232]** Complement C4 plays a central role in the activation of the classical pathway of the complement system. This protein contains three chains (alpha, beta and gamma). We identified peptide masses covering the alpha and beta chains for spot 832 and only alpha chain for spot 652 (FIG. 17). The theoretical molecular weights and pIs of these chains differ from the observed ones from the gel spots (see FIG. 11, spots 652; 832). As for complement C3, clusterin precursor protein contains two chains (alpha and beta). We identified peptide masses covering the alpha and beta chains (FIG. 16). The theoretical molecular weight and pI of clusterin (50 kDa and 5.89 respectively) are in agreement with those from the gel (FIG. 11, spot 712). It appears the full-length protein was identified.

Surface Enhanced Laser Desorption Ionisation Time of Flight Mass Spectrometry [SELDI-TOF-MS].

**[0233]** SELDI-TOF-MS and ProteinChip technology were combined to identify protein peaks differing between Alzhe-

imer's and control subjects, followed by extraction of material from the chips to allow further characterisation of the material and identification of the components present.

#### METHOD (SELDI Analysis)

**[0234]** The SELDI analysis comprises of a comparison of AD cases and control samples and data has been obtained for both a set of individual samples as well as a pooled set of samples. In each case spectral profiles of sera from control and AD cases were compared.

#### A). Analysis of a Set of Individuals

**[0235]** Control and AD sera from individuals were run on Q10-SAX2 chips:

n=4 control

n=4 AD

**[0236]** Serum samples were prepared fresh by diluting 20  $\mu$ l serum with 30  $\mu$ l SELDI lysis buffer. Five microlitres of sample were spotted onto each spot as necessary.

**[0237]** The chips were processed using the following protocol:

#### Chip Preparation

**[0238]** A hydrophobic ring is drawn around each spot using a PAP pen and the PAP allowed to dry thoroughly by placing chip on the SELDI machine for up to 25 minutes.

#### Sample Preparation

**[0239]** Serum diluted in SELDI lysis buffer using a 40:60 ratio (40  $\mu$ l serum+60  $\mu$ l lysis buffer). Typically, this dilution will render the sample at a 20 mg/mL to 30 mg/mL concentration. Therefore, using a 5  $\mu$ l aliquot of the lysis buffer sample will enable between 100  $\mu$ g to 150  $\mu$ g protein to be loaded on each spot.

**[0240]** Samples are vortexed and incubated on ice until ready to use, then briefly centrifuged samples immediately before use (30 secs, 14,000 rpm).

#### Chip Equilibration

**[0241]** The chip is placed in a 15 mL Falcon tube and 10-15 mL 100 mM Tris buffer pH 9 at room temperature added, then mixed on a rotary mixer for 5 minutes. The procedure is repeated twice.

#### Sample Application

**[0242]** After the last equilibration step, the chip is removed and dried carefully with soft tissue. 5  $\mu$ l of sample is pipetted onto each spot, the chip is placed in a sealed humidity chamber and placed on a shaker for 30 minutes.

#### Chip Washing

**[0243]** After incubation, the sample is carefully removed from each spot and the chip replaced in the Falcon tube. 10-15 mL 100 mM Tris buffer pH 9 is added, and the Falcon tube mixed on a rotary mixer and for 5 minutes. This is repeat four more times, then the chip washed twice in double distilled water.

#### Chip Drying

**[0244]** After the last wash step, the chip is removed and dried carefully with soft tissue, then left to air-dry at room temperature for 25 minutes.

#### SPA Application

**[0245]** 2x0.6  $\mu$ l saturated SPA matrix (freshly made) is pipette onto each spot. The first application is allowed to dry

before applying the second 0.6  $\mu$ l aliquot. The SPA is then left to dry for up to 10 minutes on the SELDI machine.

**[0246]** The chips are then read on the SELDI machine.

**[0247]** The following criteria were applied for data analysis:

**[0248]** Clustering criteria: 5 s/n; 100% spectra; 0.3% mass; 2 s/n; add est. peaks.

**[0249]** Normalisation: Total ion count between 3,000 and 30,000 Da only.

#### Results

**[0250]** Spot to spot reproducibility between loadings of the same sample was very good. Good correlation was achieved.

**[0251]** Patient to patient variability in both control and dementia groups was observed. This may be due to differences in protein amount as well as idiosyncratic differences. Using very stringent clustering, 3 peaks were found to be statistically significant ( $p=0.05$ ) and these were visually verified to check validity. The three peaks of interest (see FIGS. 1-3) are as follows:

Mr 6,430 Da abundance in AD	1.62 fold increase in $p = 0.027$
Mr 14,640 Da abundance in AD	2.29 fold increase in $p = 0.036$
Mr 27,147 Da abundance in AD	2.82 fold increase in $p = 0.004574$

#### B) Analysis of Pooled Sets

**[0252]** A set of pooled samples were analysed using exactly the same methods and criteria as described above. Here, however, we analysed 3 pooled AD samples versus 3 pooled controls where each pool contains serum from at least 25 individuals. In this manner we have encompassed samples from over 75 individuals with AD and compared them against a control cohort representing 75 number of individuals. Pooled groups are described as: AD Pool 1, 2 and 3 comprising of 25, 25 and 25 unique individuals respectively. Similarly, the pooled controls are described as: Control Pool 1, 2 and 3 comprising of 25, 25 and 25 unique individuals respectively.

#### Results

**[0253]** Using very stringent clustering, 1 peak was found to be statistically significant ( $p=0.05$ ) and this was visually verified to check validity.

**[0254]** The peak of interest (see FIG. 4) is as follows:

Mr 14,646 Da abundance in AD	1.72 fold increase in $p = 0.037$
---------------------------------	--------------------------------------

#### SELDI Peak Identification Strategy

**[0255]** The differentially expressed proteins identified by SELDI analysis were further analysed by SDS-PAGE. Bands corresponding to the MW of differentially expressed proteins were excised for analysis by mass spectroscopy.

**[0256]** Material was extracted from chips Q10 854 & 855 ("individual" samples) by boiling for 10 minutes in Laemmli buffer and control and disease spots were pooled into separate Eppendorf tubes. Extracted material was separated using

SDS-PAGE (18%, tris-glycine, Novex) and the gel was initially stained with Colloidal Coomassie Blue (CCB) but no bands were visualised. Subsequently the same gel was restained using modified (MS-compatible) silver stain (FIG. 5).

**[0257]** Six bands, between 11 and 20 kDa, were visualised and these were excised from the 1st control lane for analysis by LC/MS/MS as described above.

**[0258]** Identified proteins are shown in FIG. 7.

#### Further Analysis of Identified Proteins

##### Apolipoprotein A-IV

**[0259]** Sequence coverage obtained for apolipoprotein A-IV (P06727) is shown in FIG. 9 for the 14.6 kDa band isolated on the Q10 SAX2 SELDI chip

**[0260]** The molecular weight of the biomarker of interest observed within the SELDI profiling experiments was determined to be 14640 $\pm$ 6 Da. The 14.6 kDa species is thought to be a fragment of ApoA-IV based on the facts that the intact protein should be observed at higher mass (45 kDa) and that the peptides observed in the LC/MS/MS analysis only represented the C-terminal region of the protein. The observed molecular weight is in good agreement with the average molecular weight of 14636 Da predicted for residues 270-396 of the sequence defined for apolipoprotein A-IV within the Swiss Prot database entry P06727.

**[0261]** Both authentic full length apolipoprotein A-IV and a C-terminal fragment of apolipoprotein A-IV comprising of residues 270-396 may thus represent serum biomarkers of Alzheimer's disease.

##### Complement C4 Precursor

**[0262]** Sequence coverage obtained for Complement C4 precursor (P01028) in 2DE spot 164 is shown in FIG. 10. Spot 164 was identified on the basis of several peptides indicated in underlined bold and this defines the protein in Spot 164 as a C-Terminal fragment extending from residues 1466-1744.

##### Quantitative Protein Sequence Tag (qPST) Analysis

**[0263]** 10 disease samples and 10 control samples were individually immunodepleted for the 6 most highly abundant proteins. 2 pools consisting of either the disease or the control samples were generated and applied to the qPST procedure (precleavage with CNBr, labelling with dimethylglycine, trypsination and fractionation by strong cation exchange). The obtained SCX fractions were analysed by LC-MS and LC-MSMS using the QTOF-II instrument following the standard approach (LC-MS and LC-MSMS by three different data acquisition methods)

#### Results

##### Identification of Proteins

**[0264]** As stated above, three different MSMS acquisition strategies were employed:

**[0265]** 1. Data Dependent Analysis to obtain as many as possible peptide ID's (1 mass window).

**[0266]** 2. Data acquisition by an 'include list' containing regulated pairs, ie peptides whose intensity varied between disease and control samples (regulation criteria:  $\geq 2/\geq 0.5$ )

**[0267]** 3. Data acquisition by an 'include list' containing non-paired MS-signals.

**[0268]** Taking all results from these three approaches into account and correcting them for redundancy, 88 protein IDs were obtained.

Directed Searching for Regulated Proteins by Include List (Pairs with a Regulation  $\geq 2.0/\geq 0.5$ ) MSMS Strategy and Crossmatching:

**[0269]** 8 peptides were identified which could be cross-matched to regulations. These 8 peptides represent five proteins (the peptide grouping to obtain protein ID's was achieved by the ProteinProphet algorithm).

**[0270]** The ID's of these five proteins are shown in FIG. 8.

**[0271]** The 2 peptides which represent protein 1 also occur in Ig alpha-1 chain C region, so that the protein ID's 1 and 2 in fact represent one ID (Ig alpha-1 chain C region).

**[0272]** The hypothetical protein DKFZP686C02220 is a unique one (in fact, one peptide is unique, the second one can occur in several proteins). This protein has typical signatures of immunoglobulins (regarding InterPro entries), and the second peptide also occurs in Ig alpha-2 chain C region.

**[0273]** The proteins 4 and 5 represent one protein ID (haptoglobin precursor) because both peptides occur also in haptoglobin precursor, but the corresponding peptides were grouped as individual proteins by the algorithm used.

**[0274]** Validation of APO-AIV Fragments Using Western Blotting

**[0275]** Western blotting has been undertaken to confirm that the 14.6 kDa species was a fragment of APO-AIV.

**[0276]** Plasma samples were diluted 1:10 with double distilled water and assayed using a Bradford dye-binding method (diluted samples permit handling of suitably sized aliquot volumes).

**[0277]** SDS-PAGE was carried out using 20  $\mu$ g sample per lane (2  $\mu$ g if sample is a denatured primary or secondary antibody) on 16% acrylamide gels, 1.5 mm thick, 10 wells (NOVEX) for 1 hr 80 V; 1% hrs 125 V. This was followed by Western Blotting onto nitrocellulose membrane at 50 V for 1 1/2 hrs. The blots were probed with the following antibodies:

**[0278]** Anti-ApoA-IV (N-terminal specific), Santa Cruz Biotechnology, Inc.

**[0279]** Anti-ApoA-IV (C-terminal specific), Santa Cruz Biotechnology, Inc.

**[0280]** Both antibodies are affinity purified goat polyclonals raised against a peptide mapping near the amino (N-terminal) or carboxy (C-terminal) terminus of ApoA-IV of human origin. These antibodies were chosen since probing for the N- and C-terminals should increase the chance of detection of the ApoA-IV protein and/or fragments.

**[0281]** Several bands were found that appear to be ApoA-IV specific and also discriminatory for AD. These bands do not appear in the secondary antibody-only control blot for control or AD samples.

**[0282]** Bands 3-6 which are observed in the 10-16 kDa region are discriminatory for AD bands 3-6, but also appear to align with bands in the denatured ApoA-IV antibody lanes. It has also been observed that bands 3-6 are much stronger on blots where the N-terminal specific anti-ApoA-IV antibody has been used.

**[0283]** Two other key bands are observed. Band 1 is observed at approximately 45 kDa and appears to correspond to the full length mature APO-AIV protein. Band 2 is observed at approximately 28 kDa and appears to be an N-terminal fragment of APO-AIV.

##### Complement Factor H Validation.

##### Methods

##### Sample Dilution

**[0284]** Plasma samples were diluted to 1 in 8 in Phosphate buffered saline (PBS). An equal volume of Laemmli 2 $\times$  sample buffer was added and then boiled for 10 min until use.

## Western Blot

[0285] SDS gel electrophoresis was performed using the Fisher Scientific 36 well, 1.5 mm gels (all solutions were purchased from National Diagnostics). Samples were separated on a 10% resolving gel with a 4% stacking gel (all solutions were purchased from National Diagnostics). Samples (20  $\mu$ l) were separated initially for 30 min at 110V and then for 60 min at 150V until the dye front just began to enter the running buffer.

[0286] The gel was transferred to PVDF (Amersham Biosciences) using a Semi-dry transblot (Bio-Rad) for 45 min at 15V. The membrane was then blocked in 5% milk made in PBS-Tween and probed with Complement factor H primary antibody (Abcam, UK) overnight at 4° C. The bands were detected with a chemiluminescence Western detection kit (ECL+, Amersham Biosciences) and the membranes were scanned using Storm fluorescence scanner (Amersham Biosciences).

[0287] An immunoreactive band was observed at 139 kDa (CFH) and the optical density was quantified using the Image Quant (Amersham Biosciences) software. Analysis was by non-parametric Mann-Whitney using the SPSS package.

## Results

[0288] Western blot data was acquired from plasma from 128 people with NINCDS-AD/DA probable AD and 78 normal healthy elderly controls. Cases with AD had a 32% increase in CFH (Mann-Whitney; table 2)

TABLE 2

Diagnosis	Number	Mean CFH	SD	SEM
Controls	128	65.6	65.5	5.8
Probable AD	78	96.0	96.8	11.0

[0289] There was a gender difference with a relatively higher CFH value in females overall relative to males (p=0.05). However CFH was higher in cases with AD relative to controls even when considering genders separately (p<0.01; table 3)

TABLE 3

Females only	Number	Mean CFH	SEM
Controls	78	73.0	8.9
Probable AD	64	102.7	13.0
Total	142	86.4	7.7

[0290] A receiver operator curve (ROC) analysis showed that CFH performs better than chance as a diagnostic test.

[0291] The references mentioned herein are all expressly incorporated by reference.

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Arg Asn Phe Leu Val Arg Ala Ser Cys Arg Leu Arg Leu Glu Pro Gly 1665	1670	1675 1680
Lys Glu Tyr Leu Ile Met Gly Leu Asp Gly Ala Thr Tyr Asp Leu Glu 1685	1690	1695
Gly His Pro Gln Tyr Leu Leu Asp Ser Asn Ser Trp Ile Glu Glu Met 1700	1705	1710
Pro Ser Glu Arg Leu Cys Arg Ser Thr Arg Gln Arg Ala Ala Cys Ala 1715	1720	1725
Gln Leu Asn Asp Phe Leu Gln Glu Tyr Gly Thr Gln Gly Cys Gln Val 1730	1735	1740
 <210> SEQ ID NO 3 <211> LENGTH: 1474 <212> TYPE: PRT <213> ORGANISM: Homo Sapiens  <400> SEQUENCE: 3		
Met Gly Lys Asn Lys Leu Leu His Pro Ser Leu Val Leu Leu Leu Leu 1	5	10 15
Val Leu Leu Pro Thr Asp Ala Ser Val Ser Gly Lys Pro Gln Tyr Met 20	25	30
Val Leu Val Pro Ser Leu Leu His Thr Glu Thr Thr Glu Lys Gly Cys 35	40	45
Val Leu Leu Ser Tyr Leu Asn Glu Thr Val Thr Val Ser Ala Ser Leu 50	55	60
Glu Ser Val Arg Gly Asn Arg Ser Leu Phe Thr Asp Leu Glu Ala Glu 65	70	75 80
Asn Asp Val Leu His Cys Val Ala Phe Ala Val Pro Lys Ser Ser Ser 85	90	95
Asn Glu Glu Val Met Phe Leu Thr Val Gln Val Lys Gly Pro Thr Gln 100	105	110
Glu Phe Lys Lys Arg Thr Thr Val Met Val Lys Asn Glu Asp Ser Leu 115	120	125
Val Phe Val Gln Thr Asp Lys Ser Ile Tyr Lys Pro Gly Gln Thr Val 130	135	140
Lys Phe Arg Val Val Ser Met Asp Glu Asn Phe His Pro Leu Asn Glu 145	150	155 160
Leu Ile Pro Leu Val Tyr Ile Gln Asp Pro Lys Gly Asn Arg Ile Ala 165	170	175

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Gln Trp Gln Ser Phe Gln Leu Glu Gly Gly Leu Lys Gln Phe Ser Phe  
 180 185 190

Pro Leu Ser Ser Glu Pro Phe Gln Gly Ser Tyr Lys Val Val Val Gln  
 195 200 205

Lys Lys Ser Gly Gly Arg Thr Glu His Pro Phe Thr Val Glu Glu Phe  
 210 215 220

Val Leu Pro Lys Phe Glu Val Gln Val Thr Val Pro Lys Ile Ile Thr  
 225 230 235 240

Ile Leu Glu Glu Glu Met Asn Val Ser Val Cys Gly Leu Tyr Thr Tyr  
 245 250 255

Gly Lys Pro Val Pro Gly His Val Thr Val Ser Ile Cys Arg Lys Tyr  
 260 265 270

Ser Asp Ala Ser Asp Cys His Gly Glu Asp Ser Gln Ala Phe Cys Glu  
 275 280 285

Lys Phe Ser Gly Gln Leu Asn Ser His Gly Cys Phe Tyr Gln Gln Val  
 290 295 300

Lys Thr Lys Val Phe Gln Leu Lys Arg Lys Glu Tyr Glu Met Lys Leu  
 305 310 315 320

His Thr Glu Ala Gln Ile Gln Glu Glu Gly Thr Val Val Glu Leu Thr  
 325 330 335

Gly Arg Gln Ser Ser Glu Ile Thr Arg Thr Ile Thr Lys Leu Ser Phe  
 340 345 350

Val Lys Val Asp Ser His Phe Arg Gln Gly Ile Pro Phe Phe Gly Gln  
 355 360 365

Val Arg Leu Val Asp Gly Lys Gly Val Pro Ile Pro Asn Lys Val Ile  
 370 375 380

Phe Ile Arg Gly Asn Glu Ala Asn Tyr Tyr Ser Asn Ala Thr Thr Asp  
 385 390 395 400

Glu His Gly Leu Val Gln Phe Ser Ile Asn Thr Thr Asn Val Met Gly  
 405 410 415

Thr Ser Leu Thr Val Arg Val Asn Tyr Lys Asp Arg Ser Pro Cys Tyr  
 420 425 430

Gly Tyr Gln Trp Val Ser Glu Glu His Glu Glu Ala His His Thr Ala  
 435 440 445

Tyr Leu Val Phe Ser Pro Ser Lys Ser Phe Val His Leu Glu Pro Met  
 450 455 460

Ser His Glu Leu Pro Cys Gly His Thr Gln Thr Val Gln Ala His Tyr  
 465 470 475 480

Ile Leu Asn Gly Gly Thr Leu Leu Gly Leu Lys Lys Leu Ser Phe Tyr  
 485 490 495

Tyr Leu Ile Met Ala Lys Gly Gly Ile Val Arg Thr Gly Thr His Gly  
 500 505 510

Leu Leu Val Lys Gln Glu Asp Met Lys Gly His Phe Ser Ile Ser Ile  
 515 520 525

Pro Val Lys Ser Asp Ile Ala Pro Val Ala Arg Leu Leu Ile Tyr Ala  
 530 535 540

Val Leu Pro Thr Gly Asp Val Ile Gly Asp Ser Ala Lys Tyr Asp Val  
 545 550 555 560

Glu Asn Cys Ile Ala Asn Lys Val Asp Leu Ser Phe Ser Pro Ser Gln  
 565 570 575

Ser Leu Pro Ala Ser His Ala His Leu Arg Val Thr Ala Ala Pro Gln

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580					585					590					
Ser	Val	Cys	Ala	Leu	Arg	Ala	Val	Asp	Gln	Ser	Val	Leu	Leu	Met	Lys
		595					600					605			
Pro	Asp	Ala	Glu	Leu	Ser	Ala	Ser	Ser	Val	Tyr	Asn	Leu	Leu	Pro	Glu
		610					615					620			
Lys	Asp	Leu	Thr	Gly	Phe	Pro	Gly	Pro	Leu	Asn	Asp	Gln	Asp	Asp	Glu
		625					630					635			640
Asp	Cys	Ile	Asn	Arg	His	Asn	Val	Tyr	Ile	Asn	Gly	Ile	Thr	Tyr	Thr
				645					650					655	
Pro	Val	Ser	Ser	Thr	Asn	Glu	Lys	Asp	Met	Tyr	Ser	Phe	Leu	Glu	Asp
				660					665					670	
Met	Gly	Leu	Lys	Ala	Phe	Thr	Asn	Ser	Lys	Ile	Arg	Lys	Pro	Lys	Met
				675					680					685	
Cys	Pro	Gln	Leu	Gln	Gln	Tyr	Glu	Met	His	Gly	Pro	Glu	Gly	Leu	Arg
				690					695					700	
Val	Gly	Phe	Tyr	Glu	Ser	Asp	Val	Met	Gly	Arg	Gly	His	Ala	Arg	Leu
				705					710					715	720
Val	His	Val	Glu	Glu	Pro	His	Thr	Glu	Thr	Val	Arg	Lys	Tyr	Phe	Pro
				725					730					735	
Glu	Thr	Trp	Ile	Trp	Asp	Leu	Val	Val	Val	Asn	Ser	Ala	Gly	Val	Ala
				740					745					750	
Glu	Val	Gly	Val	Thr	Val	Pro	Asp	Thr	Ile	Thr	Glu	Trp	Lys	Ala	Gly
				755					760					765	
Ala	Phe	Cys	Leu	Ser	Glu	Asp	Ala	Gly	Leu	Gly	Ile	Ser	Ser	Thr	Ala
				770					775					780	
Ser	Leu	Arg	Ala	Phe	Gln	Pro	Phe	Phe	Val	Glu	Leu	Thr	Met	Pro	Tyr
				785					790					795	800
Ser	Val	Ile	Arg	Gly	Glu	Ala	Phe	Thr	Leu	Lys	Ala	Thr	Val	Leu	Asn
				805					810					815	
Tyr	Leu	Pro	Lys	Cys	Ile	Arg	Val	Ser	Val	Gln	Leu	Glu	Ala	Ser	Pro
				820					825					830	
Ala	Phe	Leu	Ala	Val	Pro	Val	Glu	Lys	Glu	Gln	Ala	Pro	His	Cys	Ile
				835					840					845	
Cys	Ala	Asn	Gly	Arg	Gln	Thr	Val	Ser	Trp	Ala	Val	Thr	Pro	Lys	Ser
				850					855					860	
Leu	Gly	Asn	Val	Asn	Phe	Thr	Val	Ser	Ala	Glu	Ala	Leu	Glu	Ser	Gln
				865					870					875	880
Glu	Leu	Cys	Gly	Thr	Glu	Val	Pro	Ser	Val	Pro	Glu	His	Gly	Arg	Lys
				885					890					895	
Asp	Thr	Val	Ile	Lys	Pro	Leu	Leu	Val	Glu	Pro	Glu	Gly	Leu	Glu	Lys
				900					905					910	
Glu	Thr	Thr	Phe	Asn	Ser	Leu	Leu	Cys	Pro	Ser	Gly	Gly	Glu	Val	Ser
				915					920					925	
Glu	Glu	Leu	Ser	Leu	Lys	Leu	Pro	Pro	Asn	Val	Val	Glu	Glu	Ser	Ala
				930					935					940	
Arg	Ala	Ser	Val	Ser	Val	Leu	Gly	Asp	Ile	Leu	Gly	Ser	Ala	Met	Gln
				945					950					955	960
Asn	Thr	Gln	Asn	Leu	Leu	Gln	Met	Pro	Tyr	Gly	Cys	Gly	Glu	Gln	Asn
				965					970					975	
Met	Val	Leu	Phe	Ala	Pro	Asn	Ile	Tyr	Val	Leu	Asp	Tyr	Leu	Asn	Glu
				980					985					990	

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Thr Gln Gln Leu Thr Pro Glu Val Lys Ser Lys Ala Ile Gly Tyr Leu  
 995 1000 1005

Asn Thr Gly Tyr Gln Arg Gln Leu Asn Tyr Lys His Tyr Asp Gly Ser  
 1010 1015 1020

Tyr Ser Thr Phe Gly Glu Arg Tyr Gly Arg Asn Gln Gly Asn Thr Trp  
 1025 1030 1035 1040

Leu Thr Ala Phe Val Leu Lys Thr Phe Ala Gln Ala Arg Ala Tyr Ile  
 1045 1050 1055

Phe Ile Asp Glu Ala His Ile Thr Gln Ala Leu Ile Trp Leu Ser Gln  
 1060 1065 1070

Arg Gln Lys Asp Asn Gly Cys Phe Arg Ser Ser Gly Ser Leu Leu Asn  
 1075 1080 1085

Asn Ala Ile Lys Gly Gly Val Glu Asp Glu Val Thr Leu Ser Ala Tyr  
 1090 1095 1100

Ile Thr Ile Ala Leu Leu Glu Ile Pro Leu Thr Val Thr His Pro Val  
 1105 1110 1115 1120

Val Arg Asn Ala Leu Phe Cys Leu Glu Ser Ala Trp Lys Thr Ala Gln  
 1125 1130 1135

Glu Gly Asp His Gly Ser His Val Tyr Thr Lys Ala Leu Leu Ala Tyr  
 1140 1145 1150

Ala Phe Ala Leu Ala Gly Asn Gln Asp Lys Arg Lys Glu Val Leu Lys  
 1155 1160 1165

Ser Leu Asn Glu Glu Ala Val Lys Lys Asp Asn Ser Val His Trp Glu  
 1170 1175 1180

Arg Pro Gln Lys Pro Lys Ala Pro Val Gly His Phe Tyr Glu Pro Gln  
 1185 1190 1195 1200

Ala Pro Ser Ala Glu Val Glu Met Thr Ser Tyr Val Leu Leu Ala Tyr  
 1205 1210 1215

Leu Thr Ala Gln Pro Ala Pro Thr Ser Glu Asp Leu Thr Ser Ala Thr  
 1220 1225 1230

Asn Ile Val Lys Trp Ile Thr Lys Gln Gln Asn Ala Gln Gly Gly Phe  
 1235 1240 1245

Ser Ser Thr Gln Asp Thr Val Val Ala Leu His Ala Leu Ser Lys Tyr  
 1250 1255 1260

Gly Ala Ala Thr Phe Thr Arg Thr Gly Lys Ala Ala Gln Val Thr Ile  
 1265 1270 1275 1280

Gln Ser Ser Gly Thr Phe Ser Ser Lys Phe Gln Val Asp Asn Asn Asn  
 1285 1290 1295

Arg Leu Leu Leu Gln Gln Val Ser Leu Pro Glu Leu Pro Gly Glu Tyr  
 1300 1305 1310

Ser Met Lys Val Thr Gly Glu Gly Cys Val Tyr Leu Gln Thr Ser Leu  
 1315 1320 1325

Lys Tyr Asn Ile Leu Pro Glu Lys Glu Glu Phe Pro Phe Ala Leu Gly  
 1330 1335 1340

Val Gln Thr Leu Pro Gln Thr Cys Asp Glu Pro Lys Ala His Thr Ser  
 1345 1350 1355 1360

Phe Gln Ile Ser Leu Ser Val Ser Tyr Thr Gly Ser Arg Ser Ala Ser  
 1365 1370 1375

Asn Met Ala Ile Val Asp Val Lys Met Val Ser Gly Phe Ile Pro Leu  
 1380 1385 1390

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Lys Pro Thr Val Lys Met Leu Glu Arg Ser Asn His Val Ser Arg Thr  
 1395 1400 1405

Glu Val Ser Ser Asn His Val Leu Ile Tyr Leu Asp Lys Val Ser Asn  
 1410 1415 1420

Gln Thr Leu Ser Leu Phe Phe Thr Val Leu Gln Asp Val Pro Val Arg  
 1425 1430 1435 1440

Asp Leu Lys Pro Ala Ile Val Lys Val Tyr Asp Tyr Tyr Glu Thr Asp  
 1445 1450 1455

Glu Phe Ala Ile Ala Glu Tyr Asn Ala Pro Cys Ser Lys Asp Leu Gly  
 1460 1465 1470

Asn Ala

<210> SEQ ID NO 4  
 <211> LENGTH: 930  
 <212> TYPE: PRT  
 <213> ORGANISM: Homo Sapiens

<400> SEQUENCE: 4

Met Lys Pro Pro Arg Pro Val Arg Thr Cys Ser Lys Val Leu Val Leu  
 1 5 10 15

Leu Ser Leu Leu Ala Ile His Gln Thr Thr Thr Ala Glu Lys Asn Gly  
 20 25 30

Ile Asp Ile Tyr Ser Leu Thr Val Asp Ser Arg Val Ser Ser Arg Phe  
 35 40 45

Ala His Thr Val Val Thr Ser Arg Val Val Asn Arg Ala Asn Thr Val  
 50 55 60

Gln Glu Ala Thr Phe Gln Met Glu Leu Pro Lys Lys Ala Phe Ile Thr  
 65 70 75 80

Asn Phe Ser Met Asn Ile Asp Gly Met Thr Tyr Pro Gly Ile Ile Lys  
 85 90 95

Glu Lys Ala Glu Ala Gln Ala Gln Tyr Ser Ala Ala Val Ala Lys Gly  
 100 105 110

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

Gln Val Ser Val Ser Val Ala Pro Asn Ala Lys Ile Thr Phe Glu Leu  
 130 135 140

Val Tyr Glu Glu Leu Leu Lys Arg Arg Leu Gly Val Tyr Glu Leu Leu  
 145 150 155 160

Leu Lys Val Arg Pro Gln Gln Leu Val Lys His Leu Gln Met Asp Ile  
 165 170 175

His Ile Phe Glu Pro Gln Gly Ile Ser Phe Leu Glu Thr Glu Ser Thr  
 180 185 190

Phe Met Thr Asn Gln Leu Val Asp Ala Leu Thr Thr Trp Gln Asn Lys  
 195 200 205

Thr Lys Ala His Ile Arg Phe Lys Pro Thr Leu Ser Gln Gln Gln Lys  
 210 215 220

Ser Pro Glu Gln Gln Glu Thr Val Leu Asp Gly Asn Leu Ile Ile Arg  
 225 230 235 240

Tyr Asp Val Asp Arg Ala Ile Ser Gly Gly Ser Ile Gln Ile Glu Asn  
 245 250 255

Gly Tyr Phe Val His Tyr Phe Ala Pro Glu Gly Leu Thr Thr Met Pro  
 260 265 270

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Lys Asn Val Val Phe Val Ile Asp Lys Ser Gly Ser Met Ser Gly Arg  
 275 280 285

Lys Ile Gln Gln Thr Arg Glu Ala Leu Ile Lys Ile Leu Asp Asp Leu  
 290 295 300

Ser Pro Arg Asp Gln Phe Asn Leu Ile Val Phe Ser Thr Glu Ala Thr  
 305 310 315 320

Gln Trp Arg Pro Ser Leu Val Pro Ala Ser Ala Glu Asn Val Asn Lys  
 325 330 335

Ala Arg Ser Phe Ala Ala Gly Ile Gln Ala Leu Gly Gly Thr Asn Ile  
 340 345 350

Asn Asp Ala Met Leu Met Ala Val Gln Leu Leu Asp Ser Ser Asn Gln  
 355 360 365

Glu Glu Arg Leu Pro Glu Gly Ser Val Ser Leu Ile Ile Leu Leu Thr  
 370 375 380

Asp Gly Asp Pro Thr Val Gly Glu Thr Asn Pro Arg Ser Ile Gln Asn  
 385 390 395 400

Asn Val Arg Glu Ala Val Ser Gly Arg Tyr Ser Leu Phe Cys Leu Gly  
 405 410 415

Phe Gly Phe Asp Val Ser Tyr Ala Phe Leu Glu Lys Leu Ala Leu Asp  
 420 425 430

Asn Gly Gly Leu Ala Arg Arg Ile His Glu Asp Ser Asp Ser Ala Leu  
 435 440 445

Gln Leu Gln Asp Phe Tyr Gln Glu Val Ala Asn Pro Leu Leu Thr Ala  
 450 455 460

Val Thr Phe Glu Tyr Pro Ser Asn Ala Val Glu Glu Val Thr Gln Asn  
 465 470 475 480

Asn Phe Arg Leu Leu Phe Lys Gly Ser Glu Met Val Val Ala Gly Lys  
 485 490 495

Leu Gln Asp Arg Gly Pro Asp Val Leu Thr Ala Thr Val Ser Gly Lys  
 500 505 510

Leu Pro Thr Gln Asn Ile Thr Phe Gln Thr Glu Ser Ser Val Ala Glu  
 515 520 525

Gln Glu Ala Glu Phe Gln Ser Pro Lys Tyr Ile Phe His Asn Phe Met  
 530 535 540

Glu Arg Leu Trp Ala Tyr Leu Thr Ile Gln Gln Leu Leu Glu Gln Thr  
 545 550 555 560

Val Ser Ala Ser Asp Ala Asp Gln Gln Ala Leu Arg Asn Gln Ala Leu  
 565 570 575

Asn Leu Ser Leu Ala Tyr Ser Phe Val Thr Pro Leu Thr Ser Met Val  
 580 585 590

Val Thr Lys Pro Asp Asp Gln Glu Gln Ser Gln Val Ala Glu Lys Pro  
 595 600 605

Met Glu Gly Glu Ser Arg Asn Arg Asn Val His Ser Gly Ser Thr Phe  
 610 615 620

Phe Lys Tyr Tyr Leu Gln Gly Ala Lys Ile Pro Lys Pro Glu Ala Ser  
 625 630 635 640

Phe Ser Pro Arg Arg Gly Trp Asn Arg Gln Ala Gly Ala Ala Gly Ser  
 645 650 655

Arg Met Asn Phe Arg Pro Gly Val Leu Ser Ser Arg Gln Leu Gly Leu  
 660 665 670

Pro Gly Pro Pro Asp Val Pro Asp His Ala Ala Tyr His Pro Phe Arg

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675					680					685					
Arg	Leu	Ala	Ile	Leu	Pro	Ala	Ser	Ala	Pro	Pro	Ala	Thr	Ser	Asn	Pro
690					695					700					
Asp	Pro	Ala	Val	Ser	Arg	Val	Met	Asn	Met	Lys	Ile	Glu	Glu	Thr	Thr
705					710					715					720
Met	Thr	Thr	Gln	Thr	Pro	Ala	Pro	Ile	Gln	Ala	Pro	Ser	Ala	Ile	Leu
				725					730					735	
Pro	Leu	Pro	Gly	Gln	Ser	Val	Glu	Arg	Leu	Cys	Val	Asp	Pro	Arg	His
			740					745					750		
Arg	Gln	Gly	Pro	Val	Asn	Leu	Leu	Ser	Asp	Pro	Glu	Gln	Gly	Val	Glu
		755					760						765		
Val	Thr	Gly	Gln	Tyr	Glu	Arg	Glu	Lys	Ala	Gly	Phe	Ser	Trp	Ile	Glu
		770				775							780		
Val	Thr	Phe	Lys	Asn	Pro	Leu	Val	Trp	Val	His	Ala	Ser	Pro	Glu	His
				790							795				800
Val	Val	Val	Thr	Arg	Asn	Arg	Arg	Ser	Ser	Ala	Tyr	Lys	Trp	Lys	Glu
				805					810					815	
Thr	Leu	Phe	Ser	Val	Met	Pro	Gly	Leu	Lys	Met	Thr	Met	Asp	Lys	Thr
				820					825					830	
Gly	Leu	Leu	Leu	Leu	Ser	Asp	Pro	Asp	Lys	Val	Thr	Ile	Gly	Leu	Leu
			835				840						845		
Phe	Trp	Asp	Gly	Arg	Gly	Glu	Gly	Leu	Arg	Leu	Leu	Leu	Arg	Asp	Thr
		850					855						860		
Asp	Arg	Phe	Ser	Ser	His	Val	Gly	Gly	Thr	Leu	Gly	Gln	Phe	Tyr	Gln
				870							875				880
Glu	Val	Leu	Trp	Gly	Ser	Pro	Ala	Ala	Ser	Asp	Asp	Gly	Arg	Arg	Thr
				885					890					895	
Leu	Arg	Val	Gln	Gly	Asn	Asp	His	Ser	Ala	Thr	Arg	Glu	Arg	Arg	Leu
			900						905					910	
Asp	Tyr	Gln	Glu	Gly	Pro	Pro	Gly	Val	Glu	Ile	Ser	Cys	Trp	Ser	Val
		915					920						925		
Glu	Leu														
				930											

&lt;210&gt; SEQ ID NO 5

&lt;211&gt; LENGTH: 1663

&lt;212&gt; TYPE: PRT

&lt;213&gt; ORGANISM: Homo Sapiens

&lt;400&gt; SEQUENCE: 5

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

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Lys Ser Glu Lys Gly Arg Asn Lys Phe Val Thr Val Gln Ala Thr Phe  
 100 105 110

Gly Thr Gln Val Val Glu Lys Val Val Leu Val Ser Leu Gln Ser Gly  
 115 120 125

Tyr Leu Phe Ile Gln Thr Asp Lys Thr Ile Tyr Thr Pro Gly Ser Thr  
 130 135 140

Val Leu Tyr Arg Ile Phe Thr Val Asn His Lys Leu Leu Pro Val Gly  
 145 150 155 160

Arg Thr Val Met Val Asn Ile Glu Asn Pro Glu Gly Ile Pro Val Lys  
 165 170 175

Gln Asp Ser Leu Ser Ser Gln Asn Gln Leu Gly Val Leu Pro Leu Ser  
 180 185 190

Trp Asp Ile Pro Glu Leu Val Asn Met Gly Gln Trp Lys Ile Arg Ala  
 195 200 205

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

Lys Glu Tyr Val Leu Pro Ser Phe Glu Val Ile Val Glu Pro Thr Glu  
 225 230 235 240

Lys Phe Tyr Tyr Ile Tyr Asn Glu Lys Gly Leu Glu Val Thr Ile Thr  
 245 250 255

Ala Arg Phe Leu Tyr Gly Lys Lys Val Glu Gly Thr Ala Phe Val Ile  
 260 265 270

Phe Gly Ile Gln Asp Gly Glu Gln Arg Ile Ser Leu Pro Glu Ser Leu  
 275 280 285

Lys Arg Ile Pro Ile Glu Asp Gly Ser Gly Glu Val Val Leu Ser Arg  
 290 295 300

Lys Val Leu Leu Asp Gly Val Gln Asn Leu Arg Ala Glu Asp Leu Val  
 305 310 315 320

Gly Lys Ser Leu Tyr Val Ser Ala Thr Val Ile Leu His Ser Gly Ser  
 325 330 335

Asp Met Val Gln Ala Glu Arg Ser Gly Ile Pro Ile Val Thr Ser Pro  
 340 345 350

Tyr Gln Ile His Phe Thr Lys Thr Pro Lys Tyr Phe Lys Pro Gly Met  
 355 360 365

Pro Phe Asp Leu Met Val Phe Val Thr Asn Pro Asp Gly Ser Pro Ala  
 370 375 380

Tyr Arg Val Pro Val Ala Val Gln Gly Glu Asp Thr Val Gln Ser Leu  
 385 390 395 400

Thr Gln Gly Asp Gly Val Ala Lys Leu Ser Ile Asn Thr His Pro Ser  
 405 410 415

Gln Lys Pro Leu Ser Ile Thr Val Arg Thr Lys Lys Gln Glu Leu Ser  
 420 425 430

Glu Ala Glu Gln Ala Thr Arg Thr Met Gln Ala Leu Pro Tyr Ser Thr  
 435 440 445

Val Gly Asn Ser Asn Asn Tyr Leu His Leu Ser Val Leu Arg Thr Glu  
 450 455 460

Leu Arg Pro Gly Glu Thr Leu Asn Val Asn Phe Leu Leu Arg Met Asp  
 465 470 475 480

Arg Ala His Glu Ala Lys Ile Arg Tyr Tyr Thr Tyr Leu Ile Met Asn  
 485 490 495

Lys Gly Arg Leu Leu Lys Ala Gly Arg Gln Val Arg Glu Pro Gly Gln



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Lys Ala Ala Val Tyr His His Phe Ile Ser Asp Gly Val Arg Lys Ser  
 915 920 925  
 Leu Lys Val Val Pro Glu Gly Ile Arg Met Asn Lys Thr Val Ala Val  
 930 935 940  
 Arg Thr Leu Asp Pro Glu Arg Leu Gly Arg Glu Gly Val Gln Lys Glu  
 945 950 955 960  
 Asp Ile Pro Pro Ala Asp Leu Ser Asp Gln Val Pro Asp Thr Glu Ser  
 965 970 975  
 Glu Thr Arg Ile Leu Leu Gln Gly Thr Pro Val Ala Gln Met Thr Glu  
 980 985 990  
 Asp Ala Val Asp Ala Glu Arg Leu Lys His Leu Ile Val Thr Pro Ser  
 995 1000 1005  
 Gly Cys Gly Glu Gln Asn Met Ile Gly Met Thr Pro Thr Val Ile Ala  
 1010 1015 1020  
 Val His Tyr Leu Asp Glu Thr Glu Gln Trp Glu Lys Phe Gly Leu Glu  
 1025 1030 1035 1040  
 Lys Arg Gln Gly Ala Leu Glu Leu Ile Lys Lys Gly Tyr Thr Gln Gln  
 1045 1050 1055  
 Leu Ala Phe Arg Gln Pro Ser Ser Ala Phe Ala Ala Glu Val Lys Arg  
 1060 1065 1070  
 Ala Pro Ser Thr Trp Leu Thr Ala Tyr Val Val Lys Val Phe Ser Leu  
 1075 1080 1085  
 Ala Val Asn Leu Ile Ala Ile Asp Ser Gln Val Leu Cys Gly Ala Val  
 1090 1095 1100  
 Lys Trp Leu Ile Leu Glu Lys Gln Lys Pro Asp Gly Val Phe Gln Glu  
 1105 1110 1115 1120  
 Asp Ala Pro Val Ile His Gln Glu Met Ile Gly Gly Leu Arg Asn Asn  
 1125 1130 1135  
 Asn Glu Lys Asp Met Ala Leu Thr Ala Phe Val Leu Ile Ser Leu Gln  
 1140 1145 1150  
 Glu Ala Lys Asp Ile Cys Glu Glu Gln Val Asn Ser Leu Pro Gly Ser  
 1155 1160 1165  
 Ile Thr Lys Ala Gly Asp Phe Leu Glu Ala Asn Tyr Met Asn Leu Gln  
 1170 1175 1180  
 Arg Ser Tyr Thr Val Ala Ile Ala Gly Tyr Ala Leu Ala Gln Met Gly  
 1185 1190 1195 1200  
 Arg Leu Lys Gly Pro Leu Leu Asn Lys Phe Leu Thr Thr Ala Lys Asp  
 1205 1210 1215  
 Lys Asn Arg Trp Glu Asp Pro Gly Lys Gln Leu Tyr Asn Val Glu Ala  
 1220 1225 1230  
 Thr Ser Tyr Ala Leu Leu Ala Leu Leu Gln Leu Lys Asp Phe Asp Phe  
 1235 1240 1245  
 Val Pro Pro Val Val Arg Trp Leu Asn Glu Gln Arg Tyr Tyr Gly Gly  
 1250 1255 1260  
 Gly Tyr Gly Ser Thr Gln Ala Thr Phe Met Val Phe Gln Ala Leu Ala  
 1265 1270 1275 1280  
 Gln Tyr Gln Lys Asp Ala Pro Asp His Gln Glu Leu Asn Leu Asp Val  
 1285 1290 1295  
 Ser Leu Gln Leu Pro Ser Arg Ser Ser Lys Ile Thr His Arg Ile His  
 1300 1305 1310

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Trp Glu Ser Ala Ser Leu Leu Arg Ser Glu Glu Thr Lys Glu Asn Glu  
 1315 1320 1325

Gly Phe Thr Val Thr Ala Glu Gly Lys Gly Gln Gly Thr Leu Ser Val  
 1330 1335 1340

Val Thr Met Tyr His Ala Lys Ala Lys Asp Gln Leu Thr Cys Asn Lys  
 1345 1350 1355 1360

Phe Asp Leu Lys Val Thr Ile Lys Pro Ala Pro Glu Thr Glu Lys Arg  
 1365 1370 1375

Pro Gln Asp Ala Lys Asn Thr Met Ile Leu Glu Ile Cys Thr Arg Tyr  
 1380 1385 1390

Arg Gly Asp Gln Asp Ala Thr Met Ser Ile Leu Asp Ile Ser Met Met  
 1395 1400 1405

Thr Gly Phe Ala Pro Asp Thr Asp Asp Leu Lys Gln Leu Ala Asn Gly  
 1410 1415 1420

Val Asp Arg Tyr Ile Ser Lys Tyr Glu Leu Asp Lys Ala Phe Ser Asp  
 1425 1430 1435 1440

Arg Asn Thr Leu Ile Ile Tyr Leu Asp Lys Val Ser His Ser Glu Asp  
 1445 1450 1455

Asp Cys Leu Ala Phe Lys Val His Gln Tyr Phe Asn Val Glu Leu Ile  
 1460 1465 1470

Gln Pro Gly Ala Val Lys Val Tyr Ala Tyr Tyr Asn Leu Glu Glu Ser  
 1475 1480 1485

Cys Thr Arg Phe Tyr His Pro Glu Lys Glu Asp Gly Lys Leu Asn Lys  
 1490 1495 1500

Leu Cys Arg Asp Glu Leu Cys Arg Cys Ala Glu Glu Asn Cys Phe Ile  
 1505 1510 1515 1520

Gln Lys Ser Asp Asp Lys Val Thr Leu Glu Glu Arg Leu Asp Lys Ala  
 1525 1530 1535

Cys Glu Pro Gly Val Asp Tyr Val Tyr Lys Thr Arg Leu Val Lys Val  
 1540 1545 1550

Gln Leu Ser Asn Asp Phe Asp Glu Tyr Ile Met Ala Ile Glu Gln Thr  
 1555 1560 1565

Ile Lys Ser Gly Ser Asp Glu Val Gln Val Gly Gln Gln Arg Thr Phe  
 1570 1575 1580

Ile Ser Pro Ile Lys Cys Arg Glu Ala Leu Lys Leu Glu Glu Lys Lys  
 1585 1590 1595 1600

His Tyr Leu Met Trp Gly Leu Ser Ser Asp Phe Trp Gly Glu Lys Pro  
 1605 1610 1615

Asn Leu Ser Tyr Ile Ile Gly Lys Asp Thr Trp Val Glu His Trp Pro  
 1620 1625 1630

Glu Glu Asp Glu Cys Gln Asp Glu Glu Asn Gln Lys Gln Cys Gln Asp  
 1635 1640 1645

Leu Gly Ala Phe Thr Glu Ser Met Val Val Phe Gly Cys Pro Asn  
 1650 1655 1660

<210> SEQ ID NO 6  
 <211> LENGTH: 449  
 <212> TYPE: PRT  
 <213> ORGANISM: Homo Sapiens  
 <400> SEQUENCE: 6

Met Met Lys Thr Leu Leu Leu Phe Val Gly Leu Leu Leu Thr Trp Glu  
 1 5 10 15

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Ser Gly Gln Val Leu Gly Asp Gln Thr Val Ser Asp Asn Glu Leu Gln  
 20 25 30  
 Glu Met Ser Asn Gln Gly Ser Lys Tyr Val Asn Lys Glu Ile Gln Asn  
 35 40 45  
 Ala Val Asn Gly Val Lys Gln Ile Lys Thr Leu Ile Glu Lys Thr Asn  
 50 55 60  
 Glu Glu Arg Lys Thr Leu Leu Ser Asn Leu Glu Glu Ala Lys Lys Lys  
 65 70 75 80  
 Lys Glu Asp Ala Leu Asn Glu Thr Arg Glu Ser Glu Thr Lys Leu Lys  
 85 90 95  
 Glu Leu Pro Gly Val Cys Asn Glu Thr Met Met Ala Leu Trp Glu Glu  
 100 105 110  
 Cys Lys Pro Cys Leu Lys Gln Thr Cys Met Lys Phe Tyr Ala Arg Val  
 115 120 125  
 Cys Arg Ser Gly Ser Gly Leu Val Gly Arg Gln Leu Glu Glu Phe Leu  
 130 135 140  
 Asn Gln Ser Ser Pro Phe Tyr Phe Trp Met Asn Gly Asp Arg Ile Asp  
 145 150 155 160  
 Ser Leu Leu Glu Asn Asp Arg Gln Gln Thr His Met Leu Asp Val Met  
 165 170 175  
 Gln Asp His Phe Ser Arg Ala Ser Ser Ile Ile Asp Glu Leu Phe Gln  
 180 185 190  
 Asp Arg Phe Phe Thr Arg Glu Pro Gln Asp Thr Tyr His Tyr Leu Pro  
 195 200 205  
 Phe Ser Leu Pro His Arg Arg Pro His Phe Phe Phe Pro Lys Ser Arg  
 210 215 220  
 Ile Val Arg Ser Leu Met Pro Phe Ser Pro Tyr Glu Pro Leu Asn Phe  
 225 230 235 240  
 His Ala Met Phe Gln Pro Phe Leu Glu Met Ile His Glu Ala Gln Gln  
 245 250 255  
 Ala Met Asp Ile His Phe His Ser Pro Ala Phe Gln His Pro Pro Thr  
 260 265 270  
 Glu Phe Ile Arg Glu Gly Asp Asp Asp Arg Thr Val Cys Arg Glu Ile  
 275 280 285  
 Arg His Asn Ser Thr Gly Cys Leu Arg Met Lys Asp Gln Cys Asp Lys  
 290 295 300  
 Cys Arg Glu Ile Leu Ser Val Asp Cys Ser Thr Asn Asn Pro Ser Gln  
 305 310 315 320  
 Ala Lys Leu Arg Arg Glu Leu Asp Glu Ser Leu Gln Val Ala Glu Arg  
 325 330 335  
 Leu Thr Arg Lys Tyr Asn Glu Leu Leu Lys Ser Tyr Gln Trp Lys Met  
 340 345 350  
 Leu Asn Thr Ser Ser Leu Leu Glu Gln Leu Asn Glu Gln Phe Asn Trp  
 355 360 365  
 Val Ser Arg Leu Ala Asn Leu Thr Gln Gly Glu Asp Gln Tyr Tyr Leu  
 370 375 380  
 Arg Val Thr Thr Val Ala Ser His Thr Ser Asp Ser Asp Val Pro Ser  
 385 390 395 400  
 Gly Val Thr Glu Val Val Val Lys Leu Phe Asp Ser Asp Pro Ile Thr  
 405 410 415

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Val Thr Val Pro Val Glu Val Ser Arg Lys Asn Pro Lys Phe Met Glu
      420                      425                      430

Thr Val Ala Glu Lys Ala Leu Gln Glu Tyr Arg Lys Lys His Arg Glu
      435                      440                      445

Glu

<210> SEQ ID NO 7
<211> LENGTH: 1744
<212> TYPE: PRT
<213> ORGANISM: Homo Sapiens

<400> SEQUENCE: 7

Met Arg Leu Leu Trp Gly Leu Ile Trp Ala Ser Ser Phe Phe Thr Leu
 1          5          10          15

Ser Leu Gln Lys Pro Arg Leu Leu Leu Phe Ser Pro Ser Val Val His
      20          25          30

Leu Gly Val Pro Leu Ser Val Gly Val Gln Leu Gln Asp Val Pro Arg
      35          40          45

Gly Gln Val Val Lys Gly Ser Val Phe Leu Arg Asn Pro Ser Arg Asn
      50          55          60

Asn Val Pro Cys Ser Pro Lys Val Asp Phe Thr Leu Ser Ser Glu Arg
 65          70          75          80

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

Cys Gly Leu His Gln Leu Leu Arg Gly Pro Glu Val Gln Leu Val Ala
      100         105         110

His Ser Pro Trp Leu Lys Asp Ser Leu Ser Arg Thr Thr Asn Ile Gln
      115         120         125

Gly Ile Asn Leu Leu Phe Ser Ser Arg Arg Gly His Leu Phe Leu Gln
      130         135         140

Thr Asp Gln Pro Ile Tyr Asn Pro Gly Gln Arg Val Arg Tyr Arg Val
 145         150         155         160

Phe Ala Leu Asp Gln Lys Met Arg Pro Ser Thr Asp Thr Ile Thr Val
      165         170         175

Met Val Glu Asn Ser His Gly Leu Arg Val Arg Lys Lys Glu Val Tyr
      180         185         190

Met Pro Ser Ser Ile Phe Gln Asp Asp Phe Val Ile Pro Asp Ile Ser
      195         200         205

Glu Pro Gly Thr Trp Lys Ile Ser Ala Arg Phe Ser Asp Gly Leu Glu
      210         215         220

Ser Asn Ser Ser Thr Gln Phe Glu Val Lys Lys Tyr Val Leu Pro Asn
 225         230         235         240

Phe Glu Val Lys Ile Thr Pro Gly Lys Pro Tyr Ile Leu Thr Val Pro
      245         250         255

Gly His Leu Asp Glu Met Gln Leu Asp Ile Gln Ala Arg Tyr Ile Tyr
      260         265         270

Gly Lys Pro Val Gln Gly Val Ala Tyr Val Arg Phe Gly Leu Leu Asp
      275         280         285

Glu Asp Gly Lys Lys Thr Phe Phe Arg Gly Leu Glu Ser Gln Thr Lys
      290         295         300

Leu Val Asn Gly Gln Ser His Ile Ser Leu Ser Lys Ala Glu Phe Gln
 305         310         315         320

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Asp Ala Leu Glu Lys Leu Asn Met Gly Ile Thr Asp Leu Gln Gly Leu  
                   325                                  330                                  335

Arg Leu Tyr Val Ala Ala Ala Ile Ile Glu Ser Pro Gly Gly Glu Met  
                   340                                  345                                  350

Glu Glu Ala Glu Leu Thr Ser Trp Tyr Phe Val Ser Ser Pro Phe Ser  
                   355                                  360                                  365

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

Leu Leu Gln Ala Leu Val Arg Glu Met Ser Gly Ser Pro Ala Ser Gly  
 385                                  390                                  395                                  400

Ile Pro Val Lys Val Ser Ala Thr Val Ser Ser Pro Gly Ser Val Pro  
                   405                                  410                                  415

Glu Val Gln Asp Ile Gln Gln Asn Thr Asp Gly Ser Gly Gln Val Ser  
                   420                                  425                                  430

Ile Pro Ile Ile Ile Pro Gln Thr Ile Ser Glu Leu Gln Leu Ser Val  
                   435                                  440                                  445

Ser Ala Gly Ser Pro His Pro Ala Ile Ala Arg Leu Thr Val Ala Ala  
                   450                                  455                                  460

Pro Pro Ser Gly Gly Pro Gly Phe Leu Ser Ile Glu Arg Pro Asp Ser  
 465                                  470                                  475                                  480

Arg Pro Pro Arg Val Gly Asp Thr Leu Asn Leu Asn Leu Arg Ala Val  
                   485                                  490                                  495

Gly Ser Gly Ala Thr Phe Ser His Tyr Tyr Tyr Met Ile Leu Ser Arg  
                   500                                  505                                  510

Gly Gln Ile Val Phe Met Asn Arg Glu Pro Lys Arg Thr Leu Thr Ser  
                   515                                  520                                  525

Val Ser Val Phe Val Asp His His Leu Ala Pro Ser Phe Tyr Phe Val  
                   530                                  535                                  540

Ala Phe Tyr Tyr His Gly Asp His Pro Val Ala Asn Ser Leu Arg Val  
 545                                  550                                  555                                  560

Asp Val Gln Ala Gly Ala Cys Glu Gly Lys Leu Glu Leu Ser Val Asp  
                   565                                  570                                  575

Gly Ala Lys Gln Tyr Arg Asn Gly Glu Ser Val Lys Leu His Leu Glu  
                   580                                  585                                  590

Thr Asp Ser Leu Ala Leu Val Ala Leu Gly Ala Leu Asp Thr Ala Leu  
                   595                                  600                                  605

Tyr Ala Ala Gly Ser Lys Ser His Lys Pro Leu Asn Met Gly Lys Val  
                   610                                  615                                  620

Phe Glu Ala Met Asn Ser Tyr Asp Leu Gly Cys Gly Pro Gly Gly Gly  
 625                                  630                                  635                                  640

Asp Ser Ala Leu Gln Val Phe Gln Ala Ala Gly Leu Ala Phe Ser Asp  
                   645                                  650                                  655

Gly Asp Gln Trp Thr Leu Ser Arg Lys Arg Leu Ser Cys Pro Lys Glu  
                   660                                  665                                  670

Lys Thr Thr Arg Lys Lys Arg Asn Val Asn Phe Gln Lys Ala Ile Asn  
                   675                                  680                                  685

Glu Lys Leu Gly Gln Tyr Ala Ser Pro Thr Ala Lys Arg Cys Cys Gln  
                   690                                  695                                  700

Asp Gly Val Thr Arg Leu Pro Met Met Arg Ser Cys Glu Gln Arg Ala  
 705                                  710                                  715                                  720

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

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725					730					735					
Gln	Phe	Ala	Glu	Ser	Leu	Arg	Lys	Lys	Ser	Arg	Asp	Lys	Gly	Gln	Ala
			740					745					750		
Gly	Leu	Gln	Arg	Ala	Leu	Glu	Ile	Leu	Gln	Glu	Glu	Asp	Leu	Ile	Asp
		755					760					765			
Glu	Asp	Asp	Ile	Pro	Val	Arg	Ser	Phe	Phe	Pro	Glu	Asn	Trp	Leu	Trp
	770					775					780				
Arg	Val	Glu	Thr	Val	Asp	Arg	Phe	Gln	Ile	Leu	Thr	Leu	Trp	Leu	Pro
	785					790					795				800
Asp	Ser	Leu	Thr	Thr	Trp	Glu	Ile	His	Gly	Leu	Ser	Leu	Ser	Lys	Thr
			805						810					815	
Lys	Gly	Leu	Cys	Val	Ala	Thr	Pro	Val	Gln	Leu	Arg	Val	Phe	Arg	Glu
			820						825					830	
Phe	His	Leu	His	Leu	Arg	Leu	Pro	Met	Ser	Val	Arg	Arg	Phe	Glu	Gln
		835						840					845		
Leu	Glu	Leu	Arg	Pro	Val	Leu	Tyr	Asn	Tyr	Leu	Asp	Lys	Asn	Leu	Thr
	850						855					860			
Val	Ser	Val	His	Val	Ser	Pro	Val	Glu	Gly	Leu	Cys	Leu	Ala	Gly	Gly
	865					870					875				880
Gly	Gly	Leu	Ala	Gln	Gln	Val	Leu	Val	Pro	Ala	Gly	Ser	Ala	Arg	Pro
				885					890					895	
Val	Ala	Phe	Ser	Val	Val	Pro	Thr	Ala	Ala	Ala	Ala	Val	Ser	Leu	Lys
			900						905					910	
Val	Val	Ala	Arg	Gly	Ser	Phe	Glu	Phe	Pro	Val	Gly	Asp	Ala	Val	Ser
		915					920					925			
Lys	Val	Leu	Gln	Ile	Glu	Lys	Glu	Gly	Ala	Ile	His	Arg	Glu	Glu	Leu
	930					935						940			
Val	Tyr	Glu	Leu	Asn	Pro	Leu	Asp	His	Arg	Gly	Arg	Thr	Leu	Glu	Ile
	945					950					955				960
Pro	Gly	Asn	Ser	Asp	Pro	Asn	Met	Ile	Pro	Asp	Gly	Asp	Glu	Asn	Ser
			965						970					975	
Tyr	Val	Arg	Val	Thr	Ala	Ser	Asp	Pro	Leu	Asp	Thr	Leu	Gly	Ser	Glu
			980						985					990	
Gly	Ala	Leu	Ser	Pro	Gly	Gly	Val	Ala	Ser	Leu	Leu	Arg	Leu	Pro	Arg
		995					1000					1005			
Gly	Cys	Gly	Glu	Gln	Thr	Met	Ile	Tyr	Leu	Ala	Pro	Thr	Leu	Ala	Ala
	1010						1015					1020			
Ser	Arg	Tyr	Leu	Asp	Lys	Thr	Glu	Gln	Trp	Ser	Thr	Leu	Pro	Pro	Glu
	1025					1030						1035			1040
Thr	Lys	Asp	His	Ala	Val	Asp	Leu	Ile	Gln	Lys	Gly	Tyr	Met	Arg	Ile
			1045						1050					1055	
Gln	Gln	Phe	Arg	Lys	Ala	Asp	Gly	Ser	Tyr	Ala	Ala	Trp	Leu	Ser	Arg
			1060						1065					1070	
Asp	Ser	Ser	Thr	Trp	Leu	Thr	Ala	Phe	Val	Leu	Lys	Val	Leu	Ser	Leu
			1075				1080					1085			
Ala	Gln	Glu	Gln	Val	Gly	Gly	Ser	Pro	Glu	Lys	Leu	Gln	Glu	Thr	Ser
	1090						1095					1100			
Asn	Trp	Leu	Leu	Ser	Gln	Gln	Gln	Ala	Asp	Gly	Ser	Phe	Gln	Asp	Pro
	1105					1110						1115			1120
Cys	Pro	Val	Leu	Asp	Arg	Ser	Met	Gln	Gly	Gly	Leu	Val	Gly	Asn	Asp
			1125						1130					1135	

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Glu Thr Val Ala Leu Thr Ala Phe Val Thr Ile Ala Leu His His Gly  
 1140 1145 1150

Leu Ala Val Phe Gln Asp Glu Gly Ala Glu Pro Leu Lys Gln Arg Val  
 1155 1160 1165

Glu Ala Ser Ile Ser Lys Ala Asn Ser Phe Leu Gly Glu Lys Ala Ser  
 1170 1175 1180

Ala Gly Leu Leu Gly Ala His Ala Ala Ala Ile Thr Ala Tyr Ala Leu  
 1185 1190 1195 1200

Ser Leu Thr Lys Ala Pro Val Asp Leu Leu Gly Val Ala His Asn Asn  
 1205 1210 1215

Leu Met Ala Met Ala Gln Glu Thr Gly Asp Asn Leu Tyr Trp Gly Ser  
 1220 1225 1230

Val Thr Gly Ser Gln Ser Asn Ala Val Ser Pro Thr Pro Ala Pro Arg  
 1235 1240 1245

Asn Pro Ser Asp Pro Met Pro Gln Ala Pro Ala Leu Trp Ile Glu Thr  
 1250 1255 1260

Thr Ala Tyr Ala Leu Leu His Leu Leu Leu His Glu Gly Lys Ala Glu  
 1265 1270 1275 1280

Met Ala Asp Gln Ala Ser Ala Trp Leu Thr Arg Gln Gly Ser Phe Gln  
 1285 1290 1295

Gly Gly Glu Arg Ser Thr Gln Asp Thr Val Ile Ala Leu Asp Ala Leu  
 1300 1305 1310

Ser Ala Tyr Trp Ile Ala Ser His Thr Thr Glu Glu Arg Gly Leu Asn  
 1315 1320 1325

Val Thr Leu Ser Ser Thr Gly Arg Asn Gly Phe Lys Ser His Ala Leu  
 1330 1335 1340

Gln Leu Asn Asn Arg Gln Ile Arg Gly Leu Glu Glu Glu Leu Gln Phe  
 1345 1350 1355 1360

Ser Leu Gly Ser Lys Ile Asn Val Lys Val Gly Gly Asn Ser Lys Gly  
 1365 1370 1375

Thr Leu Lys Val Leu Arg Thr Tyr Asn Val Leu Asp Met Lys Asn Thr  
 1380 1385 1390

Thr Cys Gln Asp Leu Gln Ile Glu Val Thr Val Lys Gly His Val Glu  
 1395 1400 1405

Tyr Thr Met Glu Ala Asn Glu Asp Tyr Glu Asp Tyr Glu Tyr Asp Glu  
 1410 1415 1420

Leu Pro Ala Lys Asp Asp Pro Asp Ala Pro Leu Gln Pro Val Thr Pro  
 1425 1430 1435 1440

Leu Gln Leu Phe Glu Gly Arg Arg Asn Arg Arg Arg Arg Glu Ala Pro  
 1445 1450 1455

Lys Val Val Glu Glu Gln Glu Ser Arg Val His Tyr Thr Val Cys Ile  
 1460 1465 1470

Trp Arg Asn Gly Lys Val Gly Leu Ser Gly Met Ala Ile Ala Asp Val  
 1475 1480 1485

Thr Leu Leu Ser Gly Phe His Ala Leu Arg Ala Asp Leu Glu Lys Leu  
 1490 1495 1500

Thr Ser Leu Ser Asp Arg Tyr Val Ser His Phe Glu Thr Glu Gly Pro  
 1505 1510 1515 1520

His Val Leu Leu Tyr Phe Asp Ser Val Pro Thr Ser Arg Glu Cys Val  
 1525 1530 1535

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Gly Phe Glu Ala Val Gln Glu Val Pro Val Gly Leu Val Gln Pro Ala  
 1540 1545 1550

Ser Ala Thr Leu Tyr Asp Tyr Tyr Asn Pro Glu Arg Arg Cys Ser Val  
 1555 1560 1565

Phe Tyr Gly Ala Pro Ser Lys Ser Arg Leu Leu Ala Thr Leu Cys Ser  
 1570 1575 1580

Ala Glu Val Cys Gln Cys Ala Glu Gly Lys Cys Pro Arg Gln Arg Arg  
 1585 1590 1595 1600

Ala Leu Glu Arg Gly Leu Gln Asp Glu Asp Gly Tyr Arg Met Lys Phe  
 1605 1610 1615

Ala Cys Tyr Tyr Pro Arg Val Glu Tyr Gly Phe Gln Val Lys Val Leu  
 1620 1625 1630

Arg Glu Asp Ser Arg Ala Ala Phe Arg Leu Phe Glu Thr Lys Ile Thr  
 1635 1640 1645

Gln Val Leu His Phe Thr Lys Asp Val Lys Ala Ala Ala Asn Gln Met  
 1650 1655 1660

Arg Asn Phe Leu Val Arg Ala Ser Cys Arg Leu Arg Leu Glu Pro Gly  
 1665 1670 1675 1680

Lys Glu Tyr Leu Ile Met Gly Leu Asp Gly Ala Thr Tyr Asp Leu Glu  
 1685 1690 1695

Gly His Pro Gln Tyr Leu Leu Asp Ser Asn Ser Trp Ile Glu Glu Met  
 1700 1705 1710

Pro Ser Glu Arg Leu Cys Arg Ser Thr Arg Gln Arg Ala Ala Cys Ala  
 1715 1720 1725

Gln Leu Asn Asp Phe Leu Gln Glu Tyr Gly Thr Gln Gly Cys Gln Val  
 1730 1735 1740

<210> SEQ ID NO 8  
 <211> LENGTH: 375  
 <212> TYPE: PRT  
 <213> ORGANISM: Homo Sapiens

<400> SEQUENCE: 8

Met Glu Glu Glu Ile Ala Ala Leu Val Ile Asp Asn Gly Ser Gly Met  
 1 5 10 15

Cys Lys Ala Gly Phe Ala Gly Asp Asp Ala Pro Arg Ala Val Phe Pro  
 20 25 30

Ser Ile Val Gly Arg Pro Arg His Gln Gly Val Met Val Gly Met Gly  
 35 40 45

Gln Lys Asp Ser Tyr Val Gly Asp Glu Ala Gln Ser Lys Arg Gly Ile  
 50 55 60

Leu Thr Leu Lys Tyr Pro Ile Glu His Gly Ile Val Thr Asn Trp Asp  
 65 70 75 80

Asp Met Glu Lys Ile Trp His His Thr Phe Tyr Asn Glu Leu Arg Val  
 85 90 95

Ala Pro Glu Glu His Pro Val Leu Leu Thr Glu Ala Pro Leu Asn Pro  
 100 105 110

Lys Ala Asn Arg Glu Lys Met Thr Gln Ile Met Phe Glu Thr Phe Asn  
 115 120 125

Thr Pro Ala Met Tyr Val Ala Ile Gln Ala Val Leu Ser Leu Tyr Ala  
 130 135 140

Ser Gly Arg Thr Thr Gly Ile Val Met Asp Ser Gly Asp Gly Val Thr  
 145 150 155 160

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His Thr Val Pro Ile Tyr Glu Gly Tyr Ala Leu Pro His Ala Ile Leu
      165                               170                               175

Arg Leu Asp Leu Ala Gly Arg Asp Leu Thr Asp Tyr Leu Met Lys Ile
      180                               185                               190

Leu Thr Glu Arg Gly Tyr Ser Phe Thr Thr Thr Ala Glu Arg Glu Ile
      195                               200                               205

Val Arg Asp Ile Lys Glu Lys Leu Cys Tyr Val Ala Leu Asp Phe Glu
      210                               215                               220

Gln Glu Met Ala Thr Ala Ala Ser Ser Ser Ser Leu Glu Lys Ser Tyr
      225                               230                               235                               240

Glu Leu Pro Asp Gly Gln Val Ile Thr Ile Gly Asn Glu Arg Phe Arg
      245                               250                               255

Cys Pro Glu Ala Leu Phe Gln Pro Ser Phe Leu Gly Met Glu Ser Cys
      260                               265                               270

Gly Ile His Glu Thr Thr Phe Asn Ser Ile Met Lys Cys Asp Val Asp
      275                               280                               285

Ile Arg Lys Asp Leu Tyr Ala Asn Thr Val Leu Ser Gly Gly Thr Thr
      290                               295                               300

Met Tyr Pro Gly Ile Ala Asp Arg Met Gln Lys Glu Ile Thr Ala Leu
      305                               310                               315                               320

Ala Pro Ser Thr Met Lys Ile Lys Ile Ile Ala Pro Pro Glu Arg Lys
      325                               330                               335

Tyr Ser Val Trp Ile Gly Gly Ser Ile Leu Ala Ser Leu Ser Thr Phe
      340                               345                               350

Gln Gln Met Trp Ile Ser Lys Gln Glu Tyr Asp Glu Ser Gly Pro Ser
      355                               360                               365

Ile Val His Arg Lys Cys Phe
      370                               375

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<210> SEQ ID NO 9
<211> LENGTH: 406
<212> TYPE: PRT
<213> ORGANISM: Homo Sapiens

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<400> SEQUENCE: 9

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Met Ser Ala Leu Gly Ala Val Ile Ala Leu Leu Leu Trp Gly Gln Leu
 1      5      10      15

Phe Ala Val Asp Ser Gly Asn Asp Val Thr Asp Ile Ala Asp Asp Gly
 20      25      30

Cys Pro Lys Pro Pro Glu Ile Ala His Gly Tyr Val Glu His Ser Val
 35      40      45

Arg Tyr Gln Cys Lys Asn Tyr Tyr Lys Leu Arg Thr Glu Gly Asp Gly
 50      55      60

Val Tyr Thr Leu Asn Asp Lys Lys Gln Trp Ile Asn Lys Ala Val Gly
 65      70      75      80

Asp Lys Leu Pro Glu Cys Glu Ala Asp Asp Gly Cys Pro Lys Pro Pro
 85      90      95

Glu Ile Ala His Gly Tyr Val Glu His Ser Val Arg Tyr Gln Cys Lys
100      105      110

Asn Tyr Tyr Lys Leu Arg Thr Glu Gly Asp Gly Val Tyr Thr Leu Asn
115      120      125

Asn Glu Lys Gln Trp Ile Asn Lys Ala Val Gly Asp Lys Leu Pro Glu

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9. A method according to claim 4 wherein a decrease in the expression of said protein is detected compared to that of a control subject.

10. A method according to claim 1 wherein the differentially expressed protein is detected using an antibody specific to said protein, by detecting in the sample an autoantibody specific to said protein, or by mass spectrometry.

11. A method according to claim 1 wherein the differentially expressed protein is detected using 2D gel electrophoresis.

12. A method according to claim 10 wherein the sample is immobilised on a solid support.

13. A method according to claim 1 which comprises detecting more than one differentially expressed protein.

14. A method according to claim 13 which comprises detecting four or more differentially expressed proteins.

15. A method according to claim 13, whereby a pattern of said differentially expressed proteins in a tissue sample or body fluid sample of an individual with Alzheimer's disease is used to predict the most appropriate and effective therapy to alleviate the Alzheimer's disease and to monitor the success of that treatment.

16. A method according to claim 1 wherein said at least one differentially expressed protein is a protein shown in FIG. 6, FIG. 7, FIG. 8 or FIG. 12.

17. A method according to claim 5 wherein at least one of said differentially expressed proteins is Ig lambda chain C region with accession no P01834 found in Spot 177 as shown in FIG. 6.

18. A method according to claim 6 wherein at least one of said differentially expressed proteins is the serum albumin precursor isoform found in Spot 165 as shown in FIG. 6.

19. A method according to claim 16 wherein said at least one differentially expressed proteins is one of the following proteins shown in FIG. 6, FIG. 7, FIG. 8 or FIG. 12 or a fragment thereof: apolipoprotein A-IV precursor, apolipoprotein C-III precursor, transthyretin, galectin 7, complement C4 precursor, complement factor H, S100 calcium binding protein or ceruloplasmin, histone 2B, Ig lambda chain C region, fibrinogen gamma chain precursor, inter-alpha-trypsin heavy chain H4 precursor, complement C3 precursor, clusterin precursor, gamma or beta actin, haptoglobin precursor or the serum albumin precursor isoform found in spot ID no 2, 14, 15, 123, 165, 176 or 184 of FIG. 6.

20. A method according to claim 19, wherein said fragment comprises; residues 270-309 of apolipoprotein A-IV; residues 680-1446-1744 of complement C4; or wherein said fragment is an N-terminal fragment of apolipoprotein A-IV which migrates as a 28 kD fragment in SDS-PAGE.

21. A method according to claim 19 wherein said at least one differentially expressed proteins is one of the following proteins shown in FIG. 6, FIG. 7 or FIG. 12 or a fragment thereof: alpha-2-macroglobulin, Ig alpha-1 chain C, apolipoprotein A-IV, complement factor H or serum albumin precursor found in Spot 2 of FIG. 6

22. The method of claim 1 which further comprises determining an effective therapy for treating the Alzheimer's disease.

23. A method of treatment by the use of an agent that will restore the expression of one or more differentially expressed proteins in the Alzheimer's disease state to that found in the normal state in order to prevent the development or progression of Alzheimer's disease.

24. A method of screening an agent to determine its usefulness in treating Alzheimer's disease, the method comprising:

(a) obtaining a sample of relevant tissue or body fluid taken from, or representative of, a subject having Alzheimer's disease symptoms, who or which has been treated with the agent being screened;

(b) determining the presence, absence or degree of expression of a differentially expressed protein or proteins in the tissue from, or representative of, the treated subject; and,

(c) selecting or rejecting the agent according to the extent to which it changes the expression, activity or amount of the differentially expressed protein or proteins in the treated subject having Alzheimer's disease symptoms.

25. A method according to claim 24, which method further comprises, prior to step (a), the step of establishing a paradigm in which at least one protein is differentially expressed in relevant tissue or body fluid from, or representative of, subjects having Alzheimer's disease symptoms and normal subjects.

26. The method of claim 24, wherein the agent is selected if it converts the expression of the differentially expressed protein or proteins towards that of a normal subject.

27. The method of claim 24, wherein the agent is selected if it converts the expression of the protein or proteins to that of the normal subject.

28. A method of screening an agent to determine its usefulness in treating Alzheimer's disease, the method comprising:

(a) obtaining over time samples of relevant tissue or body fluid taken from, or representative of, a subject having Alzheimer's disease symptoms, who or which has been treated with the agent being screened;

(b) determining the presence, absence or degree of expression of a differentially expressed protein or proteins in said samples; and,

(c) determining whether the agent affects the change over time in the expression of the differentially expressed protein in the treated subject having Alzheimer's disease symptoms.

29. A method according to claim 28, which method further comprises, prior to step (a), the step of

establishing a paradigm in which at least one protein is differentially expressed in relevant tissue or body fluid from, or representative of, subjects having Alzheimer's disease symptoms and normal subjects; and establishing that expression of said differentially expressed protein diverges over time in subjects having Alzheimer's disease symptoms and normal subjects.

30. The method of claim 25, wherein the subjects having differential levels of protein expression comprise:

(a) normal subjects and subjects having Alzheimer's disease symptoms; and,

(b) subjects having Alzheimer's disease which have not been treated with the agent and subjects Alzheimer's disease symptoms which have been treated with the agent.

31. The method of claim 30, wherein the differential levels of protein expression are not observed in normal subjects who have and have not been treated with the agent.

32. The method of claim 25, wherein the subjects having Alzheimer's disease symptoms are human subjects with Alzheimer's disease.

**33.** The method of claim **25**, wherein the subjects having Alzheimer's disease symptoms are mutant amyloid precursor protein (APP) transgenic mice, presenilin-1 (PS-1) transgenic mice, double transgenic APP/PS-1 transgenic mice and/or glycogen synthase kinase transgenic mice, and the normal subjects are wild-type mice.

**34.** The method of claim **33**, wherein the tissue or body fluid samples are brain tissue samples.

**35.** The method of claim **33**, wherein the tissue or body fluid samples are urine, blood, plasma, serum, saliva or cerebro-spinal fluid samples.

**36.** The method of claim **4**, wherein the paradigm is established using two-dimensional gel electrophoresis carried out on the relevant tissue or a protein-containing extract thereof.

**37.** The method of claim **4**, wherein the paradigm is established using SELDI analysis of the relevant tissue or a protein-containing extract thereof.

**38.** The method of claim **24**, wherein the differentially expressed protein or proteins comprise at least one of the proteins shown in FIG. **6**, FIG. **7**, FIG. **8** and FIG. **12**, or a rodent equivalent thereof.

**39.** A method according to claim **38** wherein at least one of said differentially expressed proteins is one of the following proteins shown in FIG. **6**, FIG. **7**, FIG. **8** or FIG. **12** or a fragment thereof: apolipoprotein A-IV precursor, apolipoprotein C-III precursor, transthyretin, galectin 7, complement C4 precursor, histone 2B, Ig lambda chain C region, fibrinogen gamma chain precursor, complement factor H, inter-alpha-trypsin heavy chain H4 precursor, complement C3 precursor, clusterin precursor, gamma or beta actin, haptoglobin precursor or the serum albumin precursor isoform found in spot ID no 2, 14, 15, 123, 165, 176 or 184 of FIG. **6**, or a rodent equivalent thereof.

**40.** A method according to claim **40**, wherein said fragment comprises amino acid residues 270-309 of apolipoprotein A-IV; or residues 1446-1744 of complement C4; or a rodent equivalent thereof.

**41.** A method according to claim **39** wherein at least one of said differentially expressed proteins is one of the following proteins shown in FIG. **6**, FIG. **7** or FIG. **12**, or a fragment thereof: alpha-2-macroglobulin, Ig alpha-1 chain C apolipoprotein A-IV, complement factor H or serum albumin precursor found in Spot 2 of FIG. **6**; or a rodent equivalent thereof.

**42.** A method of making a pharmaceutical composition which comprises having identified an agent using the method of claim **24**, the further step of manufacturing the agent and formulating it with an acceptable carrier to provide the pharmaceutical composition.

**43.** A method of identifying a protein which is differentially expressed in relevant tissue or body fluid sample from subjects with Alzheimer's disease and normal subjects, comprising:

- i) immobilising a tissue sample or body fluid sample or protein-containing extract thereof on a solid support
- ii) analysing the immobilised proteins by surface enhanced laser desorption time of flight mass spectroscopy
- iii) comparing the spectra obtained to detect differences in protein expression between Alzheimer's subjects and normal subjects.

**44.** The method of claim **43**, wherein the tissue or body fluid samples are blood, serum or cerebro-spinal fluid samples.

**45.** The method of claim **43**, further comprising the step of isolating a differentially expressed protein identified in the method.

**46.** The method of claim **45**, further comprising the step of characterising the isolated protein.

**47.** The method of claim **28**, wherein the agent is selected if it prevents or slows the change over time in the expression of the differentially expressed protein.

\* \* \* \* \*

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摘要(译)

提供了与阿尔茨海默病有关的方法和组合物。具体地，提供了相对于它们在正常状态下的表达而在阿尔茨海默病状态中差异表达的蛋白质。鉴定并描述了与阿尔茨海默病相关的蛋白质。还提供了使用差异表达的蛋白质诊断阿尔茨海默病的方法，以及用于预防和治疗阿尔茨海默病的化合物的鉴定和治疗用途的方法。

