US Patent & Trademark Office Patent Public Search | Text View

United States Patent

Kind Code

B2

Date of Patent

Inventor(s)

12384855

B2

August 12, 2025

Bradley; Allan et al.

PCSK9 antagonists

Abstract

The invention relates to Proprotein Convertase Subtilisin Kexin type 9 (PCSK9) antagonists, such as antibodies and fragments, as well as methods, uses and combinations.

Inventors: Bradley; Allan (Cambridge, GB), Liang; Qi (Cambridge, GB), Lee; E-Chiang

(Cambridge, GB), Liou; Li-Ying (Xinyi District Taipei, TW), Huang; Yu-Hui (Xinyi District Taipei, TW), Chen; Yen-Ju (Xinyi District Taipei, TW), Chen; Li-Tzu (Xinyi

District Taipei, TW)

Applicant: KYMAB LIMITED (Cambridge, GB)

Family ID: 65147244

Assignee: KYMAB LIMITED (Cambridge, GB)

Appl. No.: 17/299147

Filed (or PCT Filed): December 18, 2019

PCT No.: PCT/GB2019/053607

PCT Pub. No.: WO2020/128467

PCT Pub. Date: June 25, 2020

Prior Publication Data

Document IdentifierUS 20220049017 A1

Publication Date
Feb. 17, 2022

Foreign Application Priority Data

GB 1820687 Dec. 19, 2018

Publication Classification

Int. Cl.: C07K16/40 (20060101); A61K39/00 (20060101); A61K45/06 (20060101); A61P3/06 (20060101)

U.S. Cl.:

CPC **C07K16/40** (20130101); **A61K45/06** (20130101); **A61P3/06** (20180101); A61K2039/505 (20130101); C07K2317/56 (20130101); C07K2317/565 (20130101)

Field of Classification Search

CPC: C07K (16/40); C07K (16/28); C07K (2317/92); A61K (45/06); A61K (2039/505); A61K (2039/545); A61P (3/06)

References Cited

U.S. PATENT DOCUMENTS

Patent No.	Issued Date	Patentee Name	U.S. Cl.	CPC
5859205	12/1998	Adair	530/387.3	C07K 16/465
7879984	12/2010	Martin	530/808	A61P 35/02
8883157	12/2013	Clube	N/A	C07K 16/40
8999341	12/2014	Clube	424/146.1	C12Q 1/6883
9017678	12/2014	Clube	N/A	N/A
9034332	12/2014	Clube	424/146.1	C07K 16/40
9045548	12/2014	Clube	N/A	C07K 16/40
9051378	12/2014	Clube	N/A	C07K 16/40
9255154	12/2015	Feldhaus et al.	N/A	N/A
9266961	12/2015	Wu	N/A	C07K 16/40
2012/0093818	12/2011	Jackson et al.	N/A	N/A
2013/0004501	12/2012	Towne	435/254.2	C07K 16/244

FOREIGN PATENT DOCUMENTS

Patent No.	Application Date	Country	CPC
106062004	12/2015	CN	N/A
2650016	12/2012	EP	N/A
2016-528193	12/2015	JP	N/A
WO 1998/050431	12/1997	WO	N/A
WO 2006/106905	12/2005	WO	N/A
WO 2007/110205	12/2006	WO	N/A
WO 2008/003103	12/2007	WO	N/A
WO 2010/077854	12/2009	WO	N/A
WO 2011/004192	12/2010	WO	N/A
WO 2011/143545	12/2010	WO	N/A
WO 2013/157954	12/2012	WO	N/A
WO 2017/063593	12/2013	WO	N/A
WO 2014/209384	12/2013	WO	N/A
WO 2015/092393	12/2014	WO	N/A
WO 2015/103072	12/2014	WO	N/A

OTHER PUBLICATIONS

Bowie Ju, et al. Deciphering the message in protein sequences: tolerance to amino acid substitutions. Science. Mar. 16, 1990;247(4948): 1306-10. doi: 10.1126/science.2315699. PMID: 2315699. (Year: 1990). cited by examiner Portolano S, et al. Lack of promiscuity in autoantigen-specific H and L chain combinations as revealed by human H and L chain "roulette". J Immunol. Feb. 1, 1993;150(3):880-7. PMID: 8423344. (Year: 1993). cited by examiner Padlan EA. Anatomy of the antibody molecule. Mol Immunol. Feb. 1994;31(3):169-217. doi: 10.1016/0161-5890(94)90001-9. PMID: 8114766. (Year: 1994). cited by examiner

Bork P, et al. Go hunting in sequence databases but watch out for the traps. Trends Genet. Oct. 1996;12(10):425-7. doi: 10.1016/0168-9525(96)60040-7. PMID: 8909140. (Year: 1996). cited by examiner

Brown M, et al. Tolerance of single, but not multiple, amino acid replacements in antibody VH CDR 2: a means of minimizing B cell wastage from somatic hypermutation? J Immunol. May 1, 1996;156(9):3285-91. PMID: 8617951. (Year: 1996). cited by examiner

MacCallum RM, et al. Antibody-antigen interactions: contact analysis and binding site topography. J Mol Biol. Oct. 11, 1996;262(5): 732-45. doi: 10.1006/jmbi.1996.0548. PMID: 8876650. (Year: 1996). cited by examiner Metzler WJ, et al. Solution structure of human CTLA-4 and delineation of a CD80/CD86 binding site conserved in CD28. Nat Struct Biol. Jul. 1997;4(7):527-31. doi: 10.1038/nsb0797-527. PMID: 9228944. (Year: 1997). cited by examiner

Smith TF, et al. The challenges of genome sequence annotation or "the devil is in the details". Nat Biotechnol. Nov.

1997; 15(12): 1222-3. doi: 10.1038/nbt1197-1222. PMID: 9359093. (Year: 1997). cited by examiner

Doerks T, et al. Protein annotation: detective work for function prediction. Trends Genet. Jun. 1998;14(6):248-50.

doi: 10.1016/s0168-9525(98)01486-3. PMID: 9635409. (Year: 1998). cited by examiner

Brenner SE. Errors in genome annotation. Trends Genet. Apr. 1999;15(4): 132-3. doi: 10.1016/s0168-

9525(99)01706-0. PMID: 10203816. (Year: 1999). cited by examiner

Attwood TK. Genomics. The Babel of bioinformatics. Science. Oct. 20, 2000;290(5491):471-3. doi:

10.1126/science.290.5491.471. PMID: 11183771. (Year: 2000). cited by examiner

Bork P. Powers and pitfalls in sequence analysis: the 70% hurdle. Genome Res. Apr. 2000;10(4):398-400. doi:

10.1101/gr.10.4.398. PMID: 10779480. (Year: 2000). cited by examiner

Skolnick J, et al. From genes to protein structure and function: novel applications of computational approaches in the genomic era. Trends Biotechnol. Jan. 2000;18(1):34-9. doi: 10.1016/s0167-7799(99)01398-0. PMID: 10631780. (Year: 2000). cited by examiner

Vajdos FF, et al. Comprehensive functional maps of the antigen-binding site of an anti-ErbB2 antibody obtained with shotgun scanning mutagenesis. J Mol Biol. Jul. 5, 2002;320(2):415-28. doi: 10.1016/S0022-2836(02)00264-4. PMID: 12079396. (Year: 2002). cited by examiner

Casset F, et al. A peptide mimetic of an anti-CD4 monoclonal antibody by rational design. Biochem Biophys Res Commun. Jul. 18, 2003;307(1):198-205. doi: 10.1016/s0006-291x(03)01131-8. PMID: 12850000. (Year: 2003). cited by examiner

International Human Genome Sequencing Consortium. Finishing the euchromatic sequence of the human genome. Nature. Oct. 21, 2004;431(7011):931-45. doi: 10.1038/nature03001. PMID: 15496913. (Year: 2004). cited by examiner

Zody MC et al. Analysis of the DNA sequence and duplication history of human chromosome 15. Nature. Mar. 30, 2006;440(7084): 671-5. doi: 10.1038/nature04601. PMID: 16572171. (Year: 2006). cited by examiner Boyd SD, et al. High-Throughput DNA Sequencing Analysis of Antibody Repertoires. Microbiol Spectr. Oct. 2014;2(5). doi: 10.1128/microbiolspec.AID-0017-2014. PMID: 26104353. (Year: 2014). cited by examiner Rabia LA, et al. Understanding and overcoming trade-offs between antibody affinity, specificity, stability and solubility. Biochem Eng J. Sep. 15, 2018;137:365-374. doi: 10.1016/j.bej.2018.06.003. Epub Jun. 5, 2018. PMID: 30666176; PMCID: PMC6338232. (Year: 2018). cited by examiner

Chan et al., "A Proprotein convertase subtilisin/kexin type 9 neutralizing antibody reduces serum cholesterol in mice and nonhuman primates", Proc. Natl. Acad. Sci. USA, Jun. 16, 2009, 106(24): 9820-9825. cited by applicant Search Report for Colombian Patent Application No. NC2021/0007653, dated Dec. 28, 2023. cited by applicant International Search Report and Written Opinion for PCT International Patent Application No.

PCT/GB2019/053607, mailed Jun. 23, 2020. cited by applicant

Anderson et al., 1997. Activation of the furin endoprotease is a multiple-step process: requirements for acidification and internal propeptide cleavage. EMBO J., 16, pp. 1508-1518. cited by applicant

Ason B. et al., 2011, "Improved efficacy for ezetimibe and rosuvastatin by attenuating the induction of PCSK9", J Lipid Res, 52(4): 679-687. cited by applicant

Ason, B. et al., 2014, "PCSK9 inhibition fails to alter hepatic LDLR, circulating cholesterol, and atherosclerosis in the absence of ApoE", J Lipid Res, 55(11):2370-9. cited by applicant

Benjannet et al., 2004. NARC-1/PCSK9 and its natural mutants: zymogen cleavage and effects on the low density lipoprotein (LDL) receptor and LDL cholesterol, J. Biol. Chem., 279 (2004), pp. 48865-48875. cited by applicant Chothia, C. & Lesk, A. M., 1987, "Canonical structures for the hypervariable regions of immunoglobulins", J. Mal. Biol., 196, 901-917. cited by applicant

De Knijff, P. et al., 1991, "Familial dysbetalipoproteinemia associated with apolipoprotein E3-Leiden in an extended multigeneration pedigree", J. Clin. Invest. 88: 643-655. cited by applicant

DiGiammarino et al., "Design and generation of DVD-Ig™ molecules for dual-specific targeting", Meth. Mo. Biol., 2012, 889, 145-156. cited by applicant

Fu et al., 2000. Folding pathway mediated by an intramolecular chaperone. The inhibitory and chaperone functions of the subtilisin propeptide are not obligatorily linked. J. Biol. Chem., 275, pp. 16871-16878. cited by applicant Henrich et al., 2003. The crystal structure of the proprotein processing proteinase furin explains its stringent specificity. Nat. Struct. Biol., 10, pp. 520-526. cited by applicant

Henrich et al., 2005. Proprotein convertase models based on the crystal structures of furin and kexin: explanation of their specificity. J. Mol. Biol., 345, pp. 211-227. cited by applicant

Horton et al., 2007. Molecular biology of PCSK9: its role in LDL metabolism, Trends Biochem. Sci., 32, pp. 71-77. cited by applicant

Kabat et al., (1971) Ann. NY Acad. Sci., 190:382-391. cited by applicant

Kabat, E. A. et al., 1991, "Sequences of Proteins of Immunological Interest", 5th edit., NIH Publication No. 91-3242, U.S. Department of Health and Human Services. cited by applicant

Kotowski et al., 2006. A spectrum of PCSK9 alleles contributes to plasma levels of low-density lipoprotein cholesterol. Am. J. Hum. Genet., 78, pp. 410-422. cited by applicant

Kuhnast, Set al, 2014, "Alirocumab inhibits atherosclerosis, improves the plaque morphology, and enhances the effects of a statin", J Lipid Res. Oct. 2014, 55(10): 2103-2112. cited by applicant

Lagace et al., 2006. Secreted PCSK9 decreases the number of LDL receptors in hepatocytes and in livers of parabiotic mice. J. Clin. Invest., 116, pp. 2995-3005. cited by applicant

Langer (1990) Science 249:1527-1533. cited by applicant

Lee et al, Nat Biotechnol. 2014, 32(4):356-63. doi:10.1038/nbt.2825, "Complete humanization of the mouse immunoglobulin loci enables efficient therapeutic antibody discovery". cited by applicant

Lee, Jeong Hyun, et al. "Antibodies to a conformational epitope on gp41 neutralize HIV-1 by destabilizing the Env spike.", Nature communications, 6, (2015). cited by applicant

Lefranc, M. P., 1997, "Unique database numbering system for immunogenetic analysis", Immunol. Today, 18, 50. cited by applicant

Ikemura et al., 1987. Requirement of pro-sequence for the production of active subtilisin E in *Escherichia coli*. J. Biol. Chem., 262, pp. 7859-7864. cited by applicant

Mathis (1995) Clinical Chemistry 41(9), 1391-1397. cited by applicant

Maxwell et al., 2005. Overexpression of PCSK9 accelerates the degradation of the LDLR in a post-endoplasmic reticulum compartment. Proc. Natl. Acad. Sci. USA, 102, pp. 2069-2074. cited by applicant

Naureckiene et al., 2003. Functional characterization of Narc 1, a novel proteinase related to proteinase K. Arch. Biochem. Biophys., 420, pp. 55-67. cited by applicant

Niederfellner, Gerhard, et al. "Epitope characterization and crystal structure of GA101 provide insights into the molecular basis for type 1/11 distinction of CD20 antibodies.", Blood, 118.2, (2011), 358-367. cited by applicant Park et al., 2004. Post-transcriptional regulation of low density lipoprotein receptor protein by proprotein convertase subtilisin/kexin type 9a in mouse liver. J. Biol. Chem., 279, pp. 50630-50638. cited by applicant

Powell et al. "Compendium of excipients for parenteral formulations" PDA (1998) J Pharm Sci Technol 52:238-311. cited by applicant

Rashid et al., 2005. Decreased plasma cholesterol and hypersensitivity to statins in mice lacking Pcsk9. Proc. Natl. Acad. Sci. USA, 102, pp. 5374-5379. cited by applicant

Rawlings et al., 2006. MEROPS: the peptidase database. Nucleic Acids Res., 34, pp. D270-D272. cited by applicant Sakai et al., 1998. Molecular identification of the sterol-regulated luminal protease that cleaves SREBPs and controls lipid composition of animal cells. Mol. Cell, 2, pp. 505-514. cited by applicant

Sefton (1987) CRC Crit. Ref. Biomed. Eng. 14:201. cited by applicant

Seidah and Prat, 2007. The proprotein convertases are potential targets in the treatment of dyslipidemia, J. Mol. Med., 10.1007/s00109-007-0172-7. cited by applicant

Seidah et al., 1999. Mammalian subtilisin/kexin isozyme SKI-1: a widely expressed proprotein convertase with a unique cleavage specificity and cellular localization. Proc. Natl. Acad. Sci. USA, 96, pp. 1321-1326. cited by applicant

Seidah, et al., 2003. The secretory proprotein convertase neural apoptosis-regulated convertase 1 (NARC-1): liver regeneration and neuronal differentiation. Proc. Natl. Acad. Sci. USA, 100, pp. 928-933. cited by applicant Shield et al. (2002) JBC 277:26733. cited by applicant

Spiess, C., et al., Mal. Immunol. (2015). cited by applicant

Suckau, Detlev, et al. "Molecular epitope identification by limited proteolysis of an immobilized antigen-antibody complex and mass spectrometric peptide mapping.", Proceedings of the National Academy of Sciences, 87.24, (1990), 9848-9852. cited by applicant

Tangrea et al., 2002. Solution structure of the pro-hormone convertase 1 pro-domain from Mus musculus. J. Mol. Biol., 320, pp. 801-812. cited by applicant

Tsuyoshi Nozue: "Lipid Lowering Therapy and Circulating PCSK9 Concentration" Journal Of Atherosclerosis And Thrombosis, vol. 24, No. 9, Jan. 1, 2017 (Jan. 1, 2017), pp. 895-907. cited by applicant

Van den Hoek AM. et al., 2014, "APOE*3Leiden.CETP transgenic mice as model for pharmaceutical treatment of the metabolic syndrome", Diabetes Obes Metab. Jun. 2014;16(6):537-44. cited by applicant

Westerterp, M. et al., 2006, "Cholesteryl ester transfer protein decreases high-density lipoprotein and severely aggravates atherosclerosis in APOE*3-Leidenmice", Arterioscler. Thromb. Vase. Biol. 26:2552- 2559. cited by applicant

Wu et al. (1987) J. Biol. Chem. 262:4429-4432. cited by applicant

Zhao et al., 2006. Molecular characterization of loss-of-function mutations in PCSK9 and identification of a compound heterozygote. Am. J. Hum. Genet., 79, pp. 514-523. cited by applicant

Primary Examiner: Yu; Misook

Assistant Examiner: Peters; Alec Jon

Attorney, Agent or Firm: MH2 Technology Law Group, LLP

Background/Summary

FIELD OF THE INVENTION

(1) The invention relates to Proprotein Convertase Subtilisin Kexin type 9 (PCSK9) antagonists, such as antibodies and fragments, as well as methods, uses and combinations.

BACKGROUND

- (2) Proprotein convertase subtilisin kexin type 9 (PCSK9) is a serine protease involved in regulating the levels of the low density lipoprotein receptor (LDLR) protein (Horton et al., 2007; Seidah and Prat, 2007). In vitro experiments have shown that adding PCSK9 to HepG2 cells lowers the levels of cell surface LDLR (Benjannet et al., 2004; Lagace et al., 2006; Maxwell et al., 2005; Park et al., 2004).
- (3) Experiments with mice have shown that increasing PCSK9 protein levels decreases levels of LDLR protein in the liver (Benjannet et al., 2004; Lagace et al., 2006; Maxwell et al., 2005; Park et al., 2004), while PCSK9 knockout mice have increased levels of LDLR in the liver (Rashid et al., 2005).
- (4) Additionally, various human PCSK9 mutations that result in either increased or decreased levels of plasma LDL have been identified (Kotowski et al., 2006; Zhao et al., 2006). PCSK9 has been shown to directly interact with the LDLR protein, be endocytosed along with the LDLR, and co-immunofluoresce with the LDLR throughout the endosomal pathway (Lagace et al., 2006).
- (5) PCSK9 is a prohormone-proprotein convertase in the subtilisin (S8) family of serine proteases (Seidah et al., 2003). Humans have nine prohormone-proprotein convertases that can be divided between the S8A and S8B subfamilies (Rawlings et al., 2006). Furin, PC1/PC3, PC2, PACE4, PC4, PC5/PC6 and PC7/PC8/LPC/SPC7 are classified in subfamily S8B. Crystal and NMR structures of different domains from mouse furin and PC1 reveal subtilisin-like pro- and catalytic domains, and a P domain directly C-terminal to the catalytic domain (Henrich et al., 2003; Tangrea et al., 2002). Based on the amino acid sequence similarity within this subfamily, all seven members are predicted to have similar structures (Henrich et al., 2005). SKI-1/SIP and PCSK9 are classified in subfamily S8A. Sequence comparisons with these proteins also suggest the presence of subtilisin-like pro- and catalytic domains (Sakai et al., 1998; Seidah et al., 2003; Seidah et al., 1999). In these proteins the amino acid sequence C-terminal to the catalytic domain is more variable and does not suggest the presence of a P domain.
- (6) Prohormone-proprotein convertases are expressed as zymogens and they mature through a multi step process. The function of the pro-domain in this process is two-fold. The pro-domain first acts as a chaperone and is required for proper folding of the catalytic domain (Ikemura et al., 1987). Once the catalytic domain is folded, autocatalysis occurs between the pro-domain and catalytic domain. Following this initial cleavage reaction, the pro-domain remains bound to the catalytic domain where it then acts as an inhibitor of catalytic activity (Fu et al., 2000). When conditions are correct, maturation proceeds with a second autocatalytic event at a site within the pro-domain (Anderson et al., 1997). After this second cleavage event occurs the pro-domain and catalytic domain dissociate, giving rise to an active protease.
- (7) Autocatalysis of the PCSK9 zymogen occurs between Gln152 and Ser153 (VFAQ|SIP (SEQ ID NO: 67)) (Naureckiene et al., 2003), and has been shown to be required for its secretion from cells (Seidah et al., 2003). A second autocatalytic event at a site within PCSK9's pro-domain has not been observed. Purified PCSK9 is made up of two species that can be separated by non-reducing SDS-PAGE; the pro-domain at 17 Kd, and the catalytic plus C-terminal domains at 65 Kd. PCSK9 has not been isolated without its inhibitory pro-domain, and measurements of PCSK9's catalytic activity have been variable (Naureckiene et al., 2003; Seidah et al., 2003).
- (8) In certain embodiments, a PCSK9 polypeptide includes terminal residues, such as, but not limited to, leader sequence residues, targeting residues, amino terminal methionine residues, lysine residues, tag residues and/or fusion protein residues. "PCSK9" has also been referred to as FH3, NARC1, HCHOLA3, proprotein convertase subtilisin/kexin type 9, and neural apoptosis regulated convertase 1. The PCSK9 gene encodes a proprotein convertase protein that belongs to the proteinase K subfamily of the secretory subtilase family. The term "PCSK9" denotes both the proprotein and the product generated following autocatalysis of the proprotein. When only the autocatalyzed product is being referred to (such as for an antigen binding protein or ligand that binds to the cleaved PCSK9), the protein can be referred to as the "mature," "cleaved", "processed" or "active" PCSK9. When only the inactive form is being referred to, the protein can be referred to as the "inactive", "pro-form", or "unprocessed" form of PCSK9. The term PCSK9 also encompasses PCSK9 molecules incorporating post-translational modifications of

the PCSK9 amino acid sequence, such as PCSK9 sequences that have been glycosylated, PCSK9 sequences from which its signal sequence has been cleaved, PCSK9 sequence from which its pro domain has been cleaved from the catalytic domain but not separated from the catalytic domain (see, e.g., FIGS. 1A and 1B of US20120093818A1). PCSK9 controls expression of the low-density lipoprotein (LDL) receptor in the liver by promoting lysosomal degradation of the receptor. Inhibition of PCSK9 leads to increased hepatocyte LDL receptor expression and results in decreased plasma cholesterol levels through increased clearance of LDL particles by the liver. Treatment with monoclonal antibodies against PCSK9 is a clinically successful therapeutic intervention in patients diagnosed with hyperlipidemia or hypercholesterolemia that is unresponsive to statin treatment. Anti-PCSK9 antibody alirocumab is marketed by Regeneron Pharmaceuticals, Inc under the name PraluentTM. Anti-PCSK9 antibody evolocumab is marketed by Amgen, Inc under the name RepathaTM.

STATEMENT OF INVENTION

- (9) In a first configuration the invention provides:
- (10) An antibody or fragment comprising a binding site which specifically binds to Proprotein Convertase Subtilisin Kexin type 9 (PCSK9), wherein the binding site comprises a VH domain that is encoded by a nucleotide sequence that is derived from the recombination of a human VH gene segment, DH gene segment and JH gene segment, wherein the VH gene segment is selected from IGHV4-31, IGHV4-59, IGHV4-4 and IGHV3-9.
- (11) In a second configuration the invention provides:
- (12) An antibody or fragment which specifically binds to PCSK9 and comprises the CDRH3 sequence of an anti-PCSK9 antibody according to the invention, or said CDRH3 sequence comprising 3, 2 or 1 amino acid substitution(s).
- (13) In a third configuration the invention provides:
- (14) An antibody or fragment (optionally according to any preceding claim) which specifically binds to PCSK9 and comprises a VH domain which comprises a CDRH3 sequence of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or said sequence comprising 3, 2 or 1 amino acid substitution(s).
- (15) In a fourth configuration the invention provides:
- (16) An antibody or fragment comprising a binding site which specifically binds to PCSK9, wherein the binding site comprises a VH domain that comprises the amino acid sequence of a VH domain of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or an amino acid that is at least 70% identical thereto.
- (17) In a fifth configuration the invention provides:
- (18) An antibody or fragment (optionally according to any preceding claim) comprising a binding site which specifically binds to PCSK9, wherein the binding site comprises a VL domain that is encoded by a nucleotide sequence that is derived from the recombination of a human VL gene segment and JL gene segment, wherein the VL gene segment is selected from IGKV3-11, IGKV2-28 and IGKV2-29.
- (19) In a sixth configuration the invention provides:
- (20) An antibody or fragment which specifically binds to PCSK9 and comprises the CDRL3 sequence of an anti-PCSK9 antibody of the invention, said CDRL3 sequence comprising 3, 2 or 1 amino acid substitution(s).
- (21) In a seventh configuration the invention provides:
- (22) An antibody or fragment which specifically binds to PCSK9 and comprises a VL domain which comprises a CDRL3 (and optionally a CDRH3) sequence of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or said sequence(s) each comprising 3, 2 or 1 amino acid substitution(s).
- (23) In a eighth configuration the invention provides:
- (24) An antibody or fragment comprising a binding site which specifically binds to PCSK9, wherein the binding site comprises a VL domain that comprises the amino acid sequence of a VL domain of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or an amino acid that is at least 70% identical thereto.
- (25) In a ninth configuration the invention provides:
- (26) An antibody or fragment which specifically binds to PCSK9 and comprises the heavy chain amino acid sequence of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or an amino acid that is at least 70% identical thereto.
- (27) In a tenth configuration the invention provides:
- (28) An antibody or fragment which specifically binds to PCSK9 and comprises the light chain amino acid sequence of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or an amino acid that is at least 70% identical thereto.
- (29) In a eleventh configuration the invention provides:
- (30) An antibody or fragment which specifically binds to a human PCSK9 epitope that is identical to an epitope to

- which the antibody of the invention (eg, CL-274711 or CL-148219QLT) binds.
- (31) In a twelfth configuration the invention provides:
- (32) An antibody or fragment which competes for binding to human PCSK9 with the antibody of the invention.
- (33) In a thirteenth configuration the invention provides:
- (34) An anti-PCSK9 antibody or fragment of the invention for treating or preventing a PCSK9-mediated disease or condition (optionally hyperlipidaemia or hypercholesterolaemia) in a subject.
- (35) In a fourteenth configuration the invention provides:
- (36) A combination of an amount of an anti-PCSK9 antibody or fragment and an amount of a statin (optionally comprising multiple doses of said antibody and/or statin), wherein the antibody or fragment is according to the invention.
- (37) A combination of an amount of an anti-PCSK9 antibody or fragment and an amount of an ANGPTL3 inhibitor (eg, an anti-ANGPTL3 antibody, eg, evinacumab) (optionally comprising multiple doses of said antibody and/or statin), wherein the antibody or fragment is according to the invention. Optionally, the combination also comprises a statin.
- (38) In a fifteenth configuration the invention provides:
- (39) Use of the antibody, fragment or combination of the invention in the manufacture of a medicament for administration to a subject for treating or preventing a PCSK9-mediated disease or condition, optionally hyperlipidaemia or hypercholesterolaemia.
- (40) In a sixteenth configuration the invention provides:
- (41) A method of treating or preventing a PCSK9-mediated disease or condition in a subject (optionally hyperlipidaemia or hypercholesterolaemia), the method comprising administering to said subject a therapeutically effective amount of an antibody, fragment or combination of the invention, wherein the PCSK9-mediated disease or condition is thereby treated or prevented.
- (42) In a seventeenth configuration the invention provides:
- (43) A pharmaceutical composition comprising an antibody, fragment or combination of the invention and a pharmaceutically acceptable excipient, diluent or carrier.
- (44) In a eighteenth configuration the invention provides:
- (45) A nucleic acid that encodes a VH domain and/or a VL domain of an antibody or fragment of the invention
- (46) In a nineteenth configuration the invention provides:
- (47) A nucleic acid that encodes a VH domain comprising the amino acid sequence of a VH domain of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or an amino acid that is at least 70% identical thereto.
- (48) In a twentieth configuration the invention provides:
- (49) A nucleic acid that encodes a heavy chain and/or a light chain of an antibody or fragment of the invention.
- (50) In a twenty-first configuration the invention provides:
- (51) A vector comprising the nucleic acid(s); optionally wherein the vector is a CHO or HEK293 vector.
- (52) In a twenty-second configuration the invention provides:
- (53) A host cell comprising the nucleic acid(s) or the vector.
- (54) In a twenty-third configuration the invention provides:
- (55) An antibody, fragment, combination, vector, host cell, use or method as herein described.

Description

BRIEF DESCRIPTION OF THE DRAWINGS

- (1) FIG. 1: CL-148219 hPCSK9 neutralisation versus benchmark antibodies;
- (2) FIG. 2: CL-148219 restoration of LDL uptake versus benchmark antibodies;
- (3) FIG. 3: CL-148219 affinity versus benchmark antibodies;
- (4) FIG. 4: CL-148219 plasma non-HDL reduction versus benchmark antibodies;
- (5) FIG. 5: CL-148489 hPCSK9 neutralisation versus benchmark antibodies;
- (6) FIG. **6**: CL-148489 restoration of LDL uptake versus benchmark antibodies;
- (7) FIG. 7: CL-148489 affinity versus benchmark antibodies;
- (8) FIG. 8: CL-148489 plasma non-HDL reduction versus benchmark antibodies;
- (9) FIG. **9**: Study design;
- (10) FIG. **10**: Total cholesterol;
- (11) FIG. 11: Total Non-HDL Cholesterol;
- (12) FIG. **12**: HDL cholesterol;
- (13) FIG. **13**: Triglycerides;
- (14) FIGS. **14***a***-14***b*: FIG. **14***a* shows the phospholipid profiles for lipoprotein fractions after treatment with different

- anti-PCSK9 antibodies; FIG. **14***b* shows the cholesterol profiles for lipoprotein fractions after treatment with different anti-PCSK9 antibodies;
- (15) FIG. **15**: Shows the effects of a single 1 mg/kg subcutaneous dose of CL-148219 QLT, Benchmark Antibody F (a commercially-marketed antibody) or a single dose of 10 mg/kg of Isotype Control subcutaneously on total cholesterol levels (±SEM) in APOE*3Leiden.CETP transgenic mice feed a Western-type diet containing 0.15% cholesterol and 15% saturated fat;
- (16) FIG. **16**: Shows the effects of a single 3 mg/kg subcutaneous dose of CL-148219 QLT, Benchmark Antibody F or a single dose of 10 mg/kg of Isotype Control subcutaneously on total cholesterol levels (±SEM) in APOE*3Leiden.CETP transgenic mice feed a Western-type diet containing 0.15% cholesterol and 15% saturated fat; (17) FIG. **17**: Shows the effects of a single 10 mg/kg subcutaneous dose of CL-148219 QLT, Benchmark Antibody F or Isotype Control on total cholesterol levels (±SEM) in APOE*3Leiden.CETP transgenic mice feed a Western-type diet containing 0.15% cholesterol and 15% saturated fat;
- (18) FIG. **18**: Comparison of the effects of a single 1 mg/kg subcutaneous dose of CL-148219 QLT with a 1 and 3 mg/kg dose of Benchmark Antibody F on total cholesterol levels(±SEM) in APOE*3Leiden.CETP transgenic mice feed a Western-type diet containing 0.15% cholesterol and 15% saturated fat;
- (19) FIG. **19**: Comparison of the effects of a single 3 mg/kg subcutaneous dose of CL-148219 QLT with a 3 and 10 mg/kg dose of Benchmark Antibody F on total cholesterol levels (±SEM) in APOE*3Leiden.CETP transgenic mice feed a Western-type diet containing 0.15% cholesterol and 15% saturated fat;
- (20) FIG. **20**: Shows the effects of a single 1 (A), 3 (B) or 10 mg/kg (C) subcutaneous dose of CL-148219 QLT/Benchmark Antibody F, 1 and 3 mg/kg of CL-274711 (A,B) or 10 mg/kg of Isotype Control on non-HDL cholesterol levels (±SEM) in APOE*3Leiden.CETP transgenic mice feed a Western-type diet containing 0.15% cholesterol and 15% saturated fat:
- (21) FIG. **21**: Comparison of the effects of a single 1 (A) and 3 mg/kg (B) subcutaneous dose of CL-148219 QLT with 1 and 3 (A) or 3 and 10 mg/kg (B) doses of Benchmark Antibody F on non-HDL cholesterol levels(±SEM) in APOE*3Leiden.CETP transgenic mice feed a Western-type diet containing 0.15% cholesterol and 15% saturated fat; (22) FIG. **22**: Shows the lipoprotein profiles (A—VLDL Cholesterol, B—VLDL Phospholipids, C—LDL Cholesterol, D—LDL Phospholipids) following a single subcutaneous 1, 3 or 10 mg/kg dose of Benchmark Antibody F/CL-148219 QLT, 3 or 10 mg/kg of CL-274711 or 10 mg/kg dose of isotype control in APOE*3Leiden.CETP transgenic mice feed a Western-type diet containing 0.15% cholesterol and 15% saturated fat; and
- (23) FIG. **23**: Shows the antibody concentrations (±SEM) over time following a single 3 mg/kg subcutaneous dose of Benchmark Antibody F, CL-148219 QLT or CL-274711 in male, C57/B16 mice. FIG. **1**. CL-148219 neuralised human PCSK9 internalisation more than benchmarks in a cell-based assay in vitro. HepG2 cells were treated with AF647-labelled human PCSK9 gain-of-function mutant in the presence of isotype control, benchmarks, or CL-148219, followed by treatment with BODIPY LDL. Cells were then collected and the fluorescent signals of AF647 was detected by CytoFlex™ flow cytometer.
- (24) FIG. **24**. CL-148219 neuralised human PCSK9 internalisation more than benchmarks in a cell-based assay in vitro. HepG2 cells were treated with AF647-labelled human PCSK9 gain-of-function mutant in the presence of isotype control, benchmarks, or CL-148219, followed by treatment with BODIPY LDL. Cells were then collected and the fluorescent signals of AF647 was detected by CytoFlex[™] flow cytometer.
- (25) FIG. **25**: CL-148219 increased LDL uptake better than benchmarks in a cell-based assay in vitro. HepG2 cells were treated with AF647-labelled human PCSK9 gain-of-function mutant in the presence of isotype control, benchmarks, or CL-148219, followed by treatment with BODIPY LDL. Cells were then collected and the fluorescent signals of BODIPY was detected by CytoFlex flow cytometer.
- (26) FIG. **26**: Affinity of CL-148219. The affinity was determined by KD. The KD of CL-148219 was measured against human and mouse PCSK9 at pH7.6 by Proteon™ SPR analysis system.
- (27) FIG. 27: CL-148219 showed superior function to reduce non-HDL cholesterol compared to benchmarks in a hyperlipidemia mouse model (E3L.CETP mice). E3L.CETP mice were treated with western diet for 4 weeks followed by antibody injection. At different time after Ab injection, blood was collected and the amount of plasma HDL and total cholesterol were measured. The amount of plasma non-HDL cholesterol was then calculated by subtracting HDL from total cholesterol.
- (28) FIG. **28**: CL-148219 QLT neuralised human PCSK9 internalisation more than benchmarks in a cell-based assay in vitro. HepG2 cells were treated with AF647-labelled human PCSK9 gain-of-function mutant in the presence of isotype control, benchmarks, or CL-148219 QLT, followed by treatment with BODIPY LDL. Cells were then collected and the fluorescent signals of AF647 was detected by CytoFlex flow cytometer.
- (29) FIG. **29**: CL-148219 QLT increased LDL uptake better than benchmarks in a cell-based assay in vitro. HepG2 cells were treated with AF647-labelled human PCSK9 gain-of-function mutant in the presence of isotype control, benchmarks, or CL-148219 QLT, followed by treatment with BODIPY LDL. Cells were then collected and the

- fluorescent signals of BODIPY was detected by CytoFlex flow cytometer.
- (30) FIG. **30**: Affinity of CL-148219 QLT. The affinity was determined by KD. The KD of CL-148219 QLT was measured against human and mouse PCSK9 at pH7.6 by Proteon SPR analysis system.
- (31) FIG. **31**: CL-148219 QLT showed superior function to reduce non-HDL cholesterol compared to benchmarks in a hyperlipidemia mouse model (E3L.CETP mice). E3L.CETP mice were treated with western diet for 4 weeks followed by antibody injection. At different time after Ab injection, blood was collected and the amount of plasma HDL and total cholesterol were measured. The amount of plasma non-HDL cholesterol was then calculated by subtracting HDL from total cholesterol.
- (32) FIG. **32**: CL-148489 moderately neuralised human PCSK9 internalisation in a cell-based assay in vitro. HepG2 cells were treated with AF647-labelled human PCSK9 gain-of-function mutant in the presence of isotype control, benchmarks, or CL-148489, followed by treatment with BODIPY LDL. Cells were then collected and the fluorescent signals of AF647 was detected by CytoFlex flow cytometer.
- (33) FIG. **33**: CL-148489 increased LDL uptake better than benchmarks in a cell-based assay in vitro. HepG2 cells were treated with AF647-labelled human PCSK9 gain-of-function mutant in the presence of isotype control, benchmarks, or CL-148489, followed by treatment with BODIPY LDL. Cells were then collected and the fluorescent signals of BODIPY was detected by CytoFlex flow cytometer.
- (34) FIG. **34**: Affinity of CL-148489. The affinity was determined by KD. The KD of CL-148489 was measured against human and mouse PCSK9 at pH7.6 by Proteon SPR analysis system.
- (35) FIG. **35**: CL-148489 showed superior function to reduce non-HDL cholesterol compared to benchmarks in a hyperlipidemia mouse model (E3L.CETP mice). E3L.CETP mice were treated with western diet for 4 weeks followed by antibody injection. At different time after Ab injection, blood was collected and the amount of plasma HDL and total cholesterol were measured. The amount of plasma non-HDL cholesterol was then calculated by subtracting HDL from total cholesterol.
- (36) FIG. **36**: CL-274698 neuralised human PCSK9 internalisation more than benchmarks in a cell-based assay in vitro. HepG2 cells were treated with AF647-labelled human PCSK9 gain-of-function mutant in the presence of isotype control, benchmarks, or CL-274698, followed by treatment with BODIPY LDL. Cells were then collected and the fluorescent signals of AF647 was detected by CytoFlex flow cytometer.
- (37) FIG. **37**: CL-274698 increased LDL uptake better than benchmarks in a cell-based assay in vitro. HepG2 cells were treated with AF647-labelled human PCSK9 gain-of-function mutant in the presence of isotype control, benchmarks, or CL-274698, followed by treatment with BODIPY LDL. Cells were then collected and the fluorescent signals of BODIPY was detected by CytoFlex flow cytometer.
- (38) FIG. **38**: Affinity of CL-274698. The affinity was determined by KD. The KD of CL-274698 was measured against human and mouse PCSK9 at pH7.6 by Proteon SPR analysis system.
- (39) FIG. **39**: CL-274698 showed superior function to reduce non-HDL cholesterol compared to benchmarks in a hyperlipidemia mouse model (E3L.CETP mice). E3L.CETP mice were treated with western diet for 4 weeks followed by antibody injection. At different time after Ab injection, blood was collected and the amount of plasma HDL and total cholesterol were measured. The amount of plasma non-HDL cholesterol was then calculated by subtracting HDL from total cholesterol.
- (40) FIG. **40**: CL-274711 neuralised human PCSK9 internalisation more than benchmarks in a cell-based assay in vitro. HepG2 cells were treated with AF647-labelled human PCSK9 gain-of-function mutant in the presence of isotype control, benchmarks, or CL-274711, followed by treatment with BODIPY LDL. Cells were then collected and the fluorescent signals of AF647 was detected by CytoFlex flow cytometer.
- (41) FIG. **41**: CL-274711 increased LDL uptake better than benchmarks in a cell-based assay in vitro. HepG2 cells were treated with AF647-labelled human PCSK9 gain-of-function mutant in the presence of isotype control, benchmarks, or CL-274711, followed by treatment with BODIPY LDL. Cells were then collected and the fluorescent signals of BODIPY was detected by CytoFlex flow cytometer.
- (42) FIG. **42**: Affinity of CL-274711. The affinity was determined by KD. The KD of CL-274711 was measured against human and mouse PCSK9 at pH7.6 by Proteon SPR analysis system.
- (43) FIG. **43**: CL-274711 showed superior function to reduce non-HDL cholesterol compared to benchmarks in a hyperlipidemia mouse model (E3L.CETP mice). E3L.CETP mice were treated with western diet for 4 weeks followed by antibody injection. At different time after Ab injection, blood was collected and the amount of plasma HDL and total cholesterol were measured. The amount of plasma non-HDL cholesterol was then calculated by subtracting HDL from total cholesterol.

DETAILED DESCRIPTION

Definitions

(44) Unless otherwise defined herein, scientific and technical terms shall have the meanings that are commonly understood by those of ordinary skill in the art. Further, unless otherwise required by context, singular terms shall include pluralities and plural terms shall include the singular. The singular terms "a," "an," and "the" include plural

referents unless context clearly indicates otherwise. Similarly, the word "or" is intended to include "and" unless the context clearly indicates otherwise. Although methods and materials similar or equivalent to those described herein can be used in the practice or testing of this disclosure, suitable methods and materials are described below. The abbreviation, "e.g." is derived from the Latin exempli gratia and is used herein to indicate a non-limiting example. Thus, the abbreviation "e.g." is synonymous with the term "for example." In the specification and claims, the term "about" is used to modify, for example, the quantity of an ingredient in a composition, concentration, volume, process temperature, process time, yield, flow rate, pressure, and like values, and ranges thereof, employed in describing the embodiments of the disclosure. The term "about" refers to variation in the numerical quantity that can occur, for example, through typical measuring and handling procedures used for making compounds, compositions, concentrates or use formulations; through inadvertent error in these procedures; through differences in the manufacture, source, or purity of starting materials or ingredients used to carry out the methods, and like proximate considerations. The term "about" also encompasses amounts that differ due to aging of a formulation with a particular initial concentration or mixture, and amounts that differ due to mixing or processing a formulation with a particular initial concentration or mixture. Where modified by the term "about" the claims appended hereto include equivalents to these quantities.

- (45) As used herein, "administer" or "administration" refers to the act of injecting or otherwise physically delivering a substance as it exists outside the body (e.g., an anti-hPCSK9 antibody provided herein) into a patient, such as by mucosal, intradermal, intravenous, intramuscular delivery and/or any other method of physical delivery described herein or known in the art. When a disease, or a symptom thereof, is being treated, administration of the substance typically occurs after the onset of the disease or symptoms thereof. When a disease, or symptoms thereof, are being prevented, administration of the substance typically occurs before the onset of the disease or symptoms thereof. (46) The term "antibody", "immunoglobulin" or "Ig" may be used interchangeably herein and means an immunoglobulin molecule that recognizes and specifically binds to a target, such as a protein, polypeptide, peptide, carbohydrate, polynucleotide, lipid, or combinations of the foregoing through at least one antigen recognition site within the variable region of the immunoglobulin molecule. As used herein, the term "antibody" encompasses intact polyclonal antibodies, intact monoclonal antibodies, antibody fragments (such as Fab, Fab', F(ab'), sub. 2, and Fy fragments), single chain Fy (scFy) mutants, multispecific antibodies such as bispecific antibodies (including dual binding antibodies), chimeric antibodies, humanized antibodies, human antibodies, fusion proteins comprising an antigen determination portion of an antibody, and any other modified immunoglobulin molecule comprising an antigen recognition site so long as the antibodies exhibit the desired biological activity. The term "antibody" can also refer to a Y-shaped glycoprotein with a molecular weight of approximately 150 kDa that is made up of four polypeptide chains: two light (L) chains and two heavy (H) chains. There are five types of mammalian Ig heavy chain isotypes denoted by the Greek letters alpha (a), delta (δ), epsilon (ϵ), gamma (γ), and mu (μ). The type of heavy chain defines the class of antibody, i.e., IgA, IgD, IgE, IgG, and IgM, respectively. The γ and α classes are further divided into subclasses on the basis of differences in the constant domain sequence and function, e.g., IgG1, hIgG2, mIgG2A, mIgG2B, IgG3, IgG4, IgA1 and IgA2. In mammals, there are two types of immunoglobulin light chains, A and K. The "variable region" or "variable domain" of an antibody refers to the amino-terminal domains of the heavy or light chain of the antibody. The variable domains of the heavy chain and light chain may be referred to as "V.sub.H" and "V.sub.L", respectively. These domains are generally the most variable parts of the antibody (relative to other antibodies of the same class) and contain the antigen binding sites. An example of antibodies are heavy chain-only (ie, H2) antibodies that comprise a dimer of a heavy chain (5'-VH-(optional Hinge)-CH2-CH3-3') and are devoid of a light chain.
- (47) The antibodies described herein may be oligoclonal, polyclonal, monoclonal (including full-length monoclonal antibodies), camelised, chimeric, CDR-grafted, multi-specific, bi-specific (including dual-binding antibodies), catalytic, chimeric, humanized, fully human, anti-idiotypic, including antibodies that can be labelled in soluble or bound form as well as fragments, variants or derivatives thereof, either alone or in combination with other amino acid sequences provided by known techniques. An antibody may be from any species. Antibodies described herein can be naked or conjugated to other molecules such as toxins, radioisotopes, etc.
- (48) The term "antigen binding site," "antigen binding domain," "antigen binding region," "antigen binding fragment," and similar terms refer to that portion of an antibody which comprises the amino acid residues that interact with an antigen and confer on the binding agent its specificity and affinity for the antigen (e.g. the complementarity determining regions (CDRs)). The antigen binding region can be derived from any animal species, such as rodents (e.g. rabbit, rat or hamster) and humans. Preferably, the antigen binding region will be of human origin.
- (49) Antigen binding fragments described herein can include single-chain Fvs (scFv), single-chain antibodies, single domain antibodies, domain antibodies, Fv fragments, Fab fragments, F(ab') fragments, F(ab').sub.2 fragments, antibody fragments that exhibit the desired biological activity, disulfide-stabilised variable region (dsFv), dimeric variable region (diabody), anti-idiotypic (anti-Id) antibodies (including, e.g. anti-Id antibodies to antibodies),

described herein can include immunoglobulin molecules and immunologically active fragments of immunoglobulin molecules, i.e., molecules that contain an antigen-binding site. Digestion of antibodies with the enzyme, papain, results in two identical antigen-binding fragments, known also as "Fab" fragments, and a "Fc" fragment, having no antigen-binding activity but having the ability to crystallize. "Fab" when used herein refers to a fragment of an antibody that includes one constant and one variable domain of each of the heavy and light chains. The term "Fc region" herein is used to define a C-terminal region of an immunoglobulin heavy chain, including native-sequence Fc regions and variant Fc regions. The "Fc fragment" refers to the carboxy-terminal portions of both H chains held together by disulfides. The effector functions of antibodies are determined by sequences in the Fc region, the region which is also recognized by Fc receptors (FcR) found on certain types of cells. Digestion of antibodies with the enzyme, pepsin, results in a F(ab').sub.2 fragment in which the two arms of the antibody molecule remain linked and comprise two-antigen binding sites. The F(ab').sub.2 fragment has the ability to crosslink antigen. (50) The term "derived from the recombination of" in relation to gene segments will be readily apparent to the skilled person, who will understand that B-cells recombine their variable region gene segments to produce coding sequence for variable domains. For example "derived from the recombination of a human VH gene segment, DH gene segment and JH gene segment" relates to the recombination of one human VH gene segment, with one DH gene segment and one JH gene segment together to form a rearranged VDJ sequence encoding a heavy chain antibody variable domain. Junctional and somatic hypermutation may also be features of the process, whereby the resulting recombined VDJ sequence includes one or more nucleotide additions, substitutions or deletions (eg, padditions and/or n-additions) that are not comprised by the germline V, D and J sequences. The equivalent will be said of V.sub.K and J.sub.K gene segments for a kappa light chain variable domain, and of Vλ and Jλ for a lambda light chain variable domain. It is intended that any post-translational modifications may additionally encompassed in variable domains.

intrabodies, linear antibodies, single-chain antibody molecules and multispecific antibodies formed from antibody fragments and epitope-binding fragments of any of the above. In particular, antibodies and antibody fragments

- (51) "Fv" when used herein refers to the minimum fragment of an antibody that retains both antigen-recognition and antigen-binding sites. This region consists of a dimer of one heavy and one light chain variable domain in tight, noncovalent or covalent association. It is in this configuration that the three CDRs of each variable domain interact to define an antigen-binding site on the surface of the V.sub.H-V.sub.L dimer. Collectively, the six CDRs confer antigen-binding specificity to the antibody. However, even a single variable domain (or half of an F.sub.V comprising only three CDRs specific for an antigen) has the ability to recognize and bind antigen, although at a lower affinity than the entire binding site.
- (52) The term "monoclonal antibody" as used herein refers to an antibody obtained from a population of substantially homogeneous antibodies, i.e. the individual antibodies comprising the population are identical except for possible naturally occurring mutations and/or post-translation modifications (e.g. isomerizations, amidations) that may be present in minor amounts. Monoclonal antibodies are highly specific and are directed against a single antigentic determinant or epitope. In contrast, polyclonal antibody preparations typically include different antibodies directed against different antigenic determinants (or epitopes). The term "monoclonal antibody" as used herein encompasses both intact and full-length monoclonal antibodies as well as antibody fragments (such as Fab, Fab', F(ab').sub.2, Fv), single chain (scFv) mutants, fusion proteins comprising an antibody portion, and any other modified immunoglobulin molecule comprising an antigen recognition site. Furthermore, "monoclonal antibody" refers to such antibodies made in any number of ways including, but not limited to, hybridoma, phage selection, recombinant expression, and transgenic animals. The monoclonal antibodies herein can include "chimeric" antibodies (immunoglobulins) in which a portion of the heavy and/or light chain is identical with or homologous to corresponding sequences in antibodies derived from a particular species or belonging to a particular antibody class or subclass, while the remainder of the chain(s) is(are) identical with or homologous to corresponding sequences in antibodies derived from another species or belonging to another antibody class or subclass, as well as fragments of such antibodies that exhibit the desired biological activity.
- (53) The term "humanised antibody" refers to a subset of chimeric antibodies in which a "hypervariable region" from a non-human immunoglobulin (the donor antibody) replaces residues from a hypervariable region in a human immunoglobulin (recipient antibody). In general, a humanized antibody will include substantially all of at least one, and typically two, variable domains, in which all or substantially all of the hypervariable loops correspond to those of a non-human immunoglobulin sequence, and all or substantially all of the framework regions are those of a human immunoglobulin sequence, although the framework regions may include one or more substitutions that improve antibody performance, such as binding affinity, isomerization, immunogenicity, etc.
- (54) The term "bispecific antibody" means an antibody which comprises specificity for two target molecules, and includes, but is not limited to, formats such as DVD-Ig (see DiGiammarino et al., "Design and generation of DVD-Ig™ molecules for dual-specific targeting", Meth. Mo. Biol., 2012, 889, 145-156), mAb.sup.2 (see WO2008/003103, the description of the mAb.sup.2 format is incorporated herein by reference), FIT-Ig (see

WO2015/103072, the description of the FIT-Ig scaffold is incorporated herein by reference), mAb-dAb, dock and lock, Fab-arm exchange, SEEDbody, Triomab, LUZ-Y, Fcab, Kλ-body, orthogonal Fab, scDiabody-Fc, diabody-Fc, tandem scFv-Fc, Fab-scFv-Fc, Fab-scFv, intrabody, BiTE, diabody, DART, TandAb, scDiabody, scDiabody-CH3, Diabody-CH3, Triple body, Miniantibody, minibody, TriBi minibody, scFv-CH3 KIH, scFv-CH-CL-scFv, F(ab')2scFv, scFv-KIH, Fab-scFv-Fc, tetravalent HCab, ImmTAC, knobs-in-holes, knobs-in-holes with common light chain, knobs-in-holes with common light chain and charge pairs, charge pairs, charge pairs with common light chain, DT-IgG, DutaMab, IgG(H)-scFv, scFv-(H)IgG, IgG(L)-scFv, scFv-(L)IgG, IgG(L,H)-Fv, IgG(H)-V, V(H)— IgG, IgG(L)-V, V(L)-IgG, KIH IgG-scFab, 2scFv-IgG, IgG-2scFv, scFv4-Ig and zybody. For a review of bispecific formats, see Spiess, C., et al., Mol. Immunol. (2015). In another embodiment, the bispecific molecule comprises an antibody which is fused to another non-Ig format, for example a T-cell receptor binding domain; an immunoglobulin superfamily domain; an agnathan variable lymphocyte receptor; a fibronectin domain (e.g. an Adnectin™); an antibody constant domain (e.g. a CH.sub.3 domain, e.g., a CH.sub.2 and/or CH.sub.3 of an Fcab™) wherein the constant domain is not a functional CH.sub.1 domain; an scFv; an (scFv).sub.2; an sc-diabody; an scFab; a centyrin and an epitope binding domain derived from a scaffold selected from CTLA-4 (Evibody™); a lipocalin domain; Protein A such as Z-domain of Protein A (e.g. an AffibodyTM or SpA); an A-domain (e.g. an AvimerTM or Maxibody™); a heat shock protein (such as and epitope binding domain derived from GroEl and GroES); a transferrin domain (e.g. a trans-body); ankyrin repeat protein (e.g. a DARPinTM); peptide aptamer; C-type lectin domain (e.g. TetranectinTM); human γ-crystallin or human ubiquitin (an affilin); a PDZ domain; scorpion toxin; and a kunitz type domain of a human protease inhibitor. See, eg, U.S. Pat. No. 5,731,568 and WO98/50431 (both incorporated herein by reference) for non-limiting examples of knob-in-hole technology. (55) The principle is to engineer paired CH3 domains of heterodimeric heavy chains so that one CH3 domain contains a "knob" and the other CH3 domains contains a "hole" at a sterically opposite position. Knobs are created by replacing small amino acid side chain at the interface between the CH3 domains, while holes are created by replacing large side chains with smaller ones. The knob is designed to insert into the hole, to favour heterodimerisation of the different CH3 domains while destabilising homodimer formation. In in a mixture of antibody heavy and light chains that assemble to form a bispecific antibody, the proportion of IgG molecules having paired heterodimeric heavy chains is thus increased, raising yield and recovery of the active molecule (56) Mutations Y349C and/or T366W may be included to form "knobs" in an IgG CH3 domain. Mutations E356C, T366S, L368A and/or Y407V may be included to form "holes" in an IgG CH3 domain. Knobs and holes may be introduced into any human IgG CH3 domain, e.g., an IgG1, IgG2, IgG3 or IgG4 CH3 domain. A preferred example is IgG4. As noted, the IgG4 may include further modifications such as the "P" and/or "E" mutations. The IgG4 type a ("ra") sequence contains substitutions Y349C and T366W ("knobs"), and the IgG4 type b ("yb") sequence contains substitutions E356C, T366S, L368A, and Y407V ("holes"). Both ra and vb also contain the "P" substitution at position 228 in the hinge (S228P), to stabilise the hinge region of the heavy chain. Both ra and yb also contain the "E" substitution in the CH2 region at position 235 (L235S), to abolish binding to FcyR. Thus the relevant sequence of the IgG4-PE heavy chain is ppcpPcpapefEggps.

(57) A further advance in bispecific IgG engineering was the idea of using a common light chain, as described in WO98/50431. Bispecific antibodies comprising two heavy-light chain pairs were described, in which the variable light chains of both heavy-light chain pairs had a common sequence. WO98/50431 described combining the common light chain approach with specific complementary interactions in the heavy chain heterodimerisation interface (such as knobs-into-holes) to promote heterodimer formation and hinder homodimer formation. In combination, these approaches enhance formation of the desired heterodimer relative to undesired heterodimers and homodimers. While knobs-into-holes technology involves engineering amino acid side chains to create complementary molecular shapes at the interface of the paired CH3 domains in the bispecific heterodimer, another way to promote heterodimer formation and hinder homodimer formation is to engineer the amino acid side chains to have opposite charges. Association of CH3 domains in the heavy chain heterodimers is favoured by the pairing of oppositely charged residues, while paired positive charges or paired negative charges would make homodimer formation less energetically favourable. WO2006/106905 described a method for producing a heteromultimer composed of more than one type of polypeptide (such as a heterodimer of two different antibody heavy chains) comprising a substitution in an amino acid residue forming an interface between said polypeptides such that heteromultimer association will be regulated, the method comprising: (a) modifying a nucleic acid encoding an amino acid residue forming the interface between polypeptides from the original nucleic acid, such that the association between polypeptides forming one or more multimers will be inhibited in a heteromultimer that may form two or more types of multimers; (b) culturing host cells such that a nucleic acid sequence modified by step (a) is expressed; and (c) recovering said heteromultimer from the host cell culture. wherein the modification of step (a) is modifying the original nucleic acid so that one or more amino acid residues

are substituted at the interface such that two or more amino acid residues, including the mutated residue(s), forming

the interface will carry the same type of positive or negative charge.

- (58) An example of this is to suppress association between heavy chains by introducing electrostatic repulsion at the interface of the heavy chain homodimers, for example by modifying amino acid residues that contact each other at the interface of the CH3 domains, including: positions 356 and 439 positions 357 and 370 positions 399 and 409, the residue numbering being according to the EU numbering system.
- (59) By modifying one or more of these pairs of residues to have like charges (both positive or both negative) in the CH3 domain of a first heavy chain, the pairing of heavy chain homodimers is inhibited by electrostatic repulsion. By engineering the same pairs or pairs of residues in the CH3 domain of a second (different) heavy chain to have an opposite charge compared with the corresponding residues in the first heavy chain, the heterodimeric pairing of the first and second heavy chains is promoted by electrostatic attraction.
- (60) Amino acids at the heavy chain constant region CH3 interface were modified to introduce charge pairs, the mutations being listed in Table 1 of WO2006/106905. It was reported that modifying the amino acids at heavy chain positions 356, 357, 370, 399, 409 and 439 to introduce charge-induced molecular repulsion at the CH3 interface had the effect of increasing efficiency of formation of the intended bispecific antibody. For example, one heavy chain constant region may be an IgG4 constant region containing mutation K439E (positively charged Lys replaced by negatively charged Glu) and the other heavy chain constant region may be an IgG4 constant region containing mutation E356K (negatively charged Glu replaced by positively charged Lys), using EU numbering. "Charge pairing" results from spatial proximity of residues 439 and 356 in an Fc region assembled from heterodimerisation of these two constant regions.
- (61) Where two different heavy chain constant regions are used, these may be connected to the two different VH domains of the antibody in either orientation.
- (62) WO2006/106905 also exemplified bispecific IgG antibodies in which the CH3 domains of IgG4 were engineered with knobs-into-holes mutations. Type a Type a (IgG4 γ a) was an IgG4 substituted at Y349C and T366W, and type b (IgG4 γ b) was an IgG4 substituted at E356C, T366S, L368A, and Y407V. In another example, introduction of charge pairs in the antibody VH and VL domains was used to inhibit the formation of "incorrect" VH-VL pairs (pairing of VH from one antibody with VL of the other antibody). In one example, Q residues in the VH and VL were changed to K or R (positive), or to E or D (negative), to inhibit hydrogen bonding between the Q side chains and to introduce electrostatic repulsion.
- (63) Further examples of charge pairs were disclosed in WO2013/157954, which described a method for producing a heterodimeric CH3 domain-comprising molecule from a single cell, the molecule comprising two CH3 domains capable of forming an interface. The method comprised providing in the cell (a) a first nucleic acid molecule encoding a first CH3 domain-comprising polypeptide chain, this chain comprising a K residue at position 366 according to the EU numbering system and (b) a second nucleic acid molecule encoding a second CH3 domain-comprising polypeptide chain, this chain comprising a D residue at position 351 according to the EU numbering system,
- the method further comprising the step of culturing the host cell, allowing expression of the two nucleic acid molecules and harvesting the heterodimeric CH3 domain-comprising molecule from the culture.
- (64) Further methods of engineering electrostatic interactions in polypeptide chains to promote heterodimer formation over homodimer formation were described in WO2011/143545.
- (65) Another example of engineering at the CH3-CH3 interface is strand-exchange engineered domain (SEED) CH3 heterodimers. The CH3 domains are composed of alternating segments of human IgA and IgG CH3 sequences, which form pairs of complementary SEED heterodimers referred to as "SEED-bodies" [WO2007/110205].
- (66) Bispecifics have also been produced with heterodimerised heavy chains that are differentially modified in the CH3 domain to alter their affinity for binding to a purification reagent such as Protein A. WO2010/151792 described a heterodimeric bispecific antigen-binding protein comprising a first polypeptide comprising, from N-terminal to C-terminal, a first epitope-binding region that selectively binds a first epitope, an immunoglobulin constant region that comprises a first CH3 region of a human IgG selected from IgG1, IgG2, and IgG4; and a second polypeptide comprising, from N-terminal to C-terminal, a second epitope-binding region that selectively binds a second epitope, an immunoglobulin constant region that comprises a second CH3 region of a human IgG selected from IgG1, IgG2, and IgG4, wherein the second CH3 region comprises a modification that reduces or eliminates binding of the second CH3 domain to Protein A.
- (67) The Fc region may thus comprise one or more mutations to promote differential purification of the active heterodimer from homodimer species. The CH3 of one heavy chain constant region may comprise the mutation His435Arg and/or Tyr436Phe (EU numbering) [] while the CH3 of the other heavy chain constant region lacks said mutations. Emicizumab, for example, comprises an Fc region in which one CH3 comprises His435 and the other CH3 comprises His435Arg.
- (68) The bispecifics of the present invention may employ any of these bispecifics techniques and molecular formats as desired.
- (69) In one embodiment, the bispecific antibody is a mAb.sup.2. A mAb.sup.2 comprises a V.sub.H and V.sub.L

domain from an intact antibody, fused to a modified constant region, which has been engineered to form an antigenbinding site, known as an "Fcab". The technology behind the Fcab/mAb.sup.2 format is described in more detail in WO2008/003103, and the description of the mAb.sup.2 format is incorporated herein by reference.

- (70) In another embodiment, the bispecific antibody is a "dual binding antibody". As used herein, the term "dual binding antibody" is a bispecific antibody wherein both antigen-binding domains are formed by a V.sub.H/V.sub.L pair, and includes FIT-Ig (see WO2015/103072, incorporated herein by reference), mAb-dAb, dock and lock, Fab-arm exchange, SEEDbody, Triomab, LUZ-Y, Fcab, K λ -body, orthogonal Fab, scDiabody-Fc, diabody-Fc, tandem scFv-Fc, Fab-scFv, intrabody, BiTE, diabody, DART, TandAb, scDiabody, scDiabody-CH3, Diabody-CH3, Triple body, Miniantibody, minibody, scFv-CH.sub.3 KIH, scFv-CH-CL-scFv, F(ab').sub.2-scFv, scFv-KIH, Fab-scFv-Fc, tetravalent HCab, ImmTAC, knobs-in-holes, knobs-in-holes with common light chain, knobs-in-holes with common light chain and charge pairs, charge pairs with common light chain, DT-IgG, DutaMab, IgG(H)-scFv, scFv-(H)IgG, IgG(L)-scFv, scFv-(L)IgG, IgG(L,H)-Fv, IgG(H)-V, V(H)—IgG, IgG(L)-V, V(L)-IgG, KIH IgG-scFab, 2scFv-IgG, IgG-2scFv and scFv4-Ig.
- (71) The term "hypervariable region", "CDR region" or "CDR" refers to the regions of an antibody variable domain which are hypervariable in sequence and/or form structurally defined loops. Generally, antigen binding sites of an antibody include six hypervariable regions: three in the V.sub.H (CDRH1, CDRH2, CDRH3), and three in the V.sub.L (CDRL1, CDRL2, CDRL3). These regions of the heavy and light chains of an antibody confer antigen-binding specificity to the antibody. CDRs may be defined according to the Kabat system (see Kabat, E. A. et al., 1991, "Sequences of Proteins of Immunological Interest", 5.sup.th edit., NIH Publication no. 91-3242, U.S. Department of Health and Human Services).
- (72) Other systems may be used to define CDRs, which as the system devised by Chothia et al (see Chothia, C. & Lesk, A. M., 1987, "Canonical structures for the hypervariable regions of immunoglobulins", J. Mol. Biol., 196, 901-917) and the IMGT system (see Lefranc, M. P., 1997, "Unique database numbering system for immunogenetic analysis", Immunol. Today, 18, 50). An antibody typically contains 3 heavy chain CDRs and 3 light chain CDRs. The term CDR or CDRs is used here to indicate one or several of these regions. A person skilled in the art is able to readily compare the different systems of nomenclature and determine whether a particular sequence may be defined as a CDR.
- (73) A "human antibody" is an antibody that possesses an amino-acid sequence corresponding to that of an antibody produced by a human and/or has been made using any of the techniques for making human antibodies and specifically excludes a humanized antibody comprising non-human antigen-binding residues. The term "specifically binds to" refers to measurable and reproducible interactions such as binding between a target and an antibody, which is determinative of the presence of the target in the presence of a heterogeneous population of molecules including biological molecules. For example, an antibody that specifically binds to a target (which can be an epitope) is an antibody that binds this target with greater affinity, avidity, more readily, and/or with greater duration than it binds to other targets. In one embodiment, the extent of binding of an antibody to an unrelated target is less than about 10% of the binding of the antibody to the target as measured, e.g. by a radioimmunoassay (RIA).
- (74) An antibody or a fragment thereof that specifically binds to a hPCSK9 antigen may be cross-reactive with related antigens. Preferably, an antibody or a fragment thereof that specifically binds to a hPCSK9 antigen does not cross-react with other antigens (but may optionally cross-react with PCSK9 of a different species, e.g. rhesus, or murine). An antibody or a fragment thereof that specifically binds to a hPCSK9 antigen can be identified, for example, by immunoassays, BIAcore™, or other techniques known to those of skill in the art. An antibody or a fragment thereof binds specifically to a PCSK9 antigen when it binds to a hPCSK9 antigen with higher affinity than to any cross-reactive antigen as determined using experimental techniques, such as radioimmunoassays (RIA) and enzyme-linked immunosorbent assays (ELISAs). Typically, a specific or selective reaction will be at least twice background signal or noise and more typically more than 10 times (such as more than 15 times, more than 20 times, more than 50 times or more than 100 times) background. See, e.g. Paul, ed., 1989, Fundamental Immunology Second Edition, Raven Press, New York at pages 332-336 for a discussion regarding antibody specificity.
- (75) The term "aliphatic amino acid" means that the amino acid R groups are nonpolar and hydrophobic. Hydrophobicity increases with increasing number of C atoms in the hydrocarbon chain. Glycine, Alanine, Valine, Leucine and Isoleucine are aliphatic amino acids.
- (76) The term "aromatic amino acid" means that the amino acid R groups contain an aromatic ring system. Phenylalanine, Tyrosine and Tryptophan are aromatic amino acids.
- (77) The term "hydroxyl-containing amino acid" means that the amino acid R groups contain a hydroxyl group and are hydrophilic. Serine, Cysteine, Threonine and Methionine are hydroxyl-containing amino acids.
- (78) The term "basic amino acid" means that the amino acid R groups are nitrogen containing and are basic at neutral pH. Histidine, Lysine and Arginine are basic amino acids.
- (79) The term "cyclic amino acid" means that the amino acid R groups have an aliphatic cyclic structure. Proline is the only cyclic aliphatic amino acid.

- (80) The term "acidic amino acid" means that the amino acid R groups are polar and are negatively charged at physiological pH. Aspartate and Glutamate are acidic amino acids.
- (81) The term "amide amino acid" means that the amino acid R groups contain an amide group. Asparagine and Glutamine are amide amino acids.
- (82) As used herein, "authorization number" or "marketing authorization number" refers to a number issued by a regulatory agency upon that agency determining that a particular medical product and/or composition may be marketed and/or offered for sale in the area under the agency's jurisdiction. As used herein "regulatory agency" refers to one of the agencies responsible for evaluating, e.g. the safety and efficacy of a medical product and/or composition and controlling the sales/marketing of such products and/or compositions in a given area. The Food and Drug Administration (FDA) in the US and the European Medicines Agency (EPA) in Europe are but two examples of such regulatory agencies. Other non-limiting examples can include SDA, MPA, MHPRA, IMA, ANMAT, Hong Kong Department of Health-Drug Office, CDSCO, Medsafe, and KFDA.
- (83) As used herein, a "buffer" refers to a chemical agent that is able to absorb a certain quantity of acid or base without undergoing a strong variation in pH.
- (84) As used herein, the term "carrier" refers to a diluent, adjuvant (e.g., Freund's adjuvant (complete and incomplete)), excipient, or vehicle with which the therapeutic is administered. Such pharmaceutical carriers can be sterile liquids, such as water and oils, including those of petroleum, animal, vegetable or synthetic origin, such as peanut oil, soybean oil, mineral oil, sesame oil and the like. Water is a preferred carrier when the pharmaceutical composition is administered intravenously. Saline solutions and aqueous dextrose and glycerol solutions can also be employed as liquid carriers, particularly for injectable solutions.
- (85) As used herein, the term "composition" is intended to encompass a product containing the specified ingredients (e.g. an antibody of the invention) in, optionally, the specified amounts, as well as any product which results, directly or indirectly, from combination of the specified ingredients in, optionally, the specified amounts.
- (86) As used herein the term "comprising" or "comprises" is used with reference to antibodies, fragments, uses, compositions, methods, and respective component(s) thereof, that are essential to the method or composition, yet open to the inclusion of unspecified elements, whether essential or not.
- (87) The term "consisting of" refers to antibodies, fragments, uses, compositions, methods, and respective components thereof as described herein, which are exclusive of any element not recited in that description of the embodiment.
- (88) As used herein the term "consisting essentially of" refers to those elements required for a given embodiment. The term permits the presence of elements that do not materially affect the basic and novel or functional characteristic(s) of that embodiment.
- (89) In the context of a polypeptide, the term "derivative" as used herein includes a polypeptide that comprises an amino acid sequence of a hPCSK9 polypeptide, a fragment of a hPCSK9 polypeptide, or an antibody or fragment that specifically binds to a hPCSK9 polypeptide which has been altered by the introduction of amino acid residue substitutions, deletions or additions. The term "derivative" as used herein also includes a hPCSK9 polypeptide, a fragment of a hPCSK9 polypeptide, or an antibody that specifically binds to a hPCSK9 polypeptide which has been chemically modified, e.g. by the covalent attachment of any type of molecule to the polypeptide. For example, but not by way of limitation, a hPCSK9 polypeptide, a fragment of a hPCSK9 polypeptide, or a hPCSK9 antibody may be chemically modified, e.g. by glycosylation, acetylation, pegylation, phosphorylation, amidation, derivatization by known protecting/blocking groups, proteolytic cleavage, linkage to a cellular ligand or other protein, etc. The derivatives are modified in a manner that is different from naturally occurring or starting peptide or polypeptides, either in the type or location of the molecules attached. Derivatives further include deletion of one or more chemical groups which are naturally present on the peptide or polypeptide. A derivative of a hPCSK9 polypeptide, a fragment of a hPCSK9 polypeptide, or a hPCSK9 antibody may be chemically modified by chemical modifications using techniques known to those of skill in the art, including, but not limited to specific chemical cleavage, acetylation, formulation, metabolic synthesis of tunicamycin, etc. Further, a derivative of a hPCSK9 polypeptide, a fragment of a hPCSK9 polypeptide, or a hPCSK9 antibody may contain one or more non-classical amino acids. A polypeptide derivative possesses a similar or identical function as a hPCSK9 polypeptide, a fragment of a hPCSK9 polypeptide, or a hPCSK9 antibody described herein.
- (90) The term "effector function" (or "effector-enabled") as used herein refers to one or more of antibody dependent cell mediated cytotoxic activity (ADCC), complement-dependent cytotoxic activity (CDC) mediated responses, Fcmediated phagocytosis or antibody dependent cellular phagocytosis (ADCP) and antibody recycling via the FcRn receptor.
- (91) An "effective amount" refers to an amount effective, at dosages and for periods of time necessary, to achieve the desired effect, including a therapeutic or prophylactic result. A "therapeutically effective amount" refers to the minimum concentration required to effect a measurable improvement or prevention of a particular disorder. A therapeutically effective amount herein may vary according to factors such as the disease state, age, sex, and weight

of the patient, and the ability of the antibody to elicit a desired response in the individual. A therapeutically effective amount is also one in which toxic or detrimental effects of the antibody are outweighed by the therapeutically beneficial effects. A "prophylactically effective amount" refers to an amount effective, at the dosages and for periods of time necessary, to achieve the desired prophylactic result. In some embodiments, the effective amount of an antibody of the invention is from about 0.1 mg/kg (mg of antibody per kg weight of the subject) to about 100 mg/kg. In certain embodiments, an effective amount of an antibody provided therein is about 0.1 mg/kg, about 0.5 mg/kg, about 1 mg/kg, 3 mg/kg, 5 mg/kg, about 10 mg/kg, about 15 mg/kg, about 20 mg/kg, about 25 mg/kg, about 30 mg/kg, about 35 mg/kg, about 40 mg/kg, about 45 mg/kg, about 50 mg/kg, about 60 mg/kg, about 70 mg/kg, about 80 mg/kg about 90 mg/kg or about 100 mg/kg (or a range therein). In some embodiments, "effective amount" as used herein also refers to the amount of an antibody of the invention to achieve a specified result (e.g. inhibition of a hPCSK9 biological activity of a cell). The term "epitope" as used herein refers to a localized region on the surface of an antigen, such as hPCSK9 polypeptide or hPCSK9 polypeptide fragment, that is capable of being bound to one or more antigen binding regions of an antibody, and that has antigenic or immunogenic activity in an animal, preferably a mammal, and most preferably in a human, that is capable of eliciting an immune response. An epitope having immunogenic activity is a portion of a polypeptide that elicits an antibody response in an animal. An epitope having antigenic activity is a portion of a polypeptide to which an antibody specifically binds as determined by any method well known in the art, for example, by the immunoassays described herein. Antigenic epitopes need not necessarily be immunogenic. Epitopes usually consist of chemically active surface groupings of molecules such as amino acids or sugar side chains and have specific three-dimensional structural characteristics as well as specific charge characteristics. A region of a polypeptide contributing to an epitope may be contiguous amino acids of the polypeptide or the epitope may come together from two or more non-contiguous regions of the polypeptide. The epitope may or may not be a three-dimensional surface feature of the antigen. In certain embodiments, a hPCSK9 epitope is a three-dimensional surface feature of a hPCSK9 polypeptide (e.g. in a trimeric form of a hPCSK9 polypeptide). In other embodiments, a hPCSK9 epitope is linear feature of a hPCSK9 polypeptide (e.g. in a trimeric form or monomeric form of the hPCSK9 polypeptide). Antibodies provided herein may specifically bind to an epitope of the monomeric (denatured) form of hPCSK9, an epitope of the trimeric (native) form of hPCSK9, or both the monomeric (denatured) form and the trimeric (native) form of hPCSK9. In specific embodiments, the antibodies provided herein specifically bind to an epitope of the trimeric form of hPCSK9 but do not specifically bind the monomeric form of hPCSK9.

- (92) The term "excipients" as used herein refers to inert substances which are commonly used as a diluent, vehicle, preservatives, binders, or stabilizing agent for drugs and includes, but not limited to, proteins (e.g. serum albumin, etc.), amino acids (e.g. aspartic acid, glutamic acid, lysine, arginine, glycine, histidine, etc.), fatty acids and phospholipids (e.g. alkyl sulfonates, caprylate, etc.), surfactants (e.g. SDS, polysorbate, nonionic surfactant, etc.), saccharides (e.g. sucrose, maltose, trehalose, etc.) and polyols (e.g. mannitol, sorbitol, etc.). See, also, Remington's Pharmaceutical Sciences (1990) Mack Publishing Co., Easton, Pa., which is hereby incorporated by reference in its entirety.
- (93) As used herein, "authorization number" or "marketing authorization number" refers to a number issued by a regulatory agency upon that agency determining that a particular medical product and/or composition may be marketed and/or offered for sale in the area under the agency's jurisdiction. As used herein "regulatory agency" refers to one of the agencies responsible for evaluating, e.g, the safety and efficacy of a medical product and/or composition and controlling the sales/marketing of such products and/or compositions in a given area. The Food and Drug Administration (FDA) in the US and the European Medicines Agency (EPA) in Europe are but two examples of such regulatory agencies. Other non-limiting examples can include SDA, MPA, MHPRA, IMA, ANMAT, Hong Kong Department of Health-Drug Office, CDSCO, Medsafe, and KFDA.
- (94) As used herein, "injection device" refers to a device that is designed for carrying out injections, an injection including the steps of temporarily fluidically coupling the injection device to a person's tissue, typically the subcutaneous tissue. An injection further includes administering an amount of liquid drug into the tissue and decoupling or removing the injection device from the tissue. In some embodiments, an injection device can be an intravenous device or IV device, which is a type of injection device used when the target tissue is the blood within the circulatory system, e.g., the blood in a vein. A common, but non-limiting example of an injection device is a needle and syringe.
- (95) As used herein, a "buffer" refers to a chemical agent that is able to absorb a certain quantity of acid or base without undergoing a strong variation in pH.
- (96) As used herein, "packaging" refers to how the components are organized and/or restrained into a unit fit for distribution and/or use. Packaging can include, e.g., boxes, bags, syringes, ampoules, vials, tubes, clamshell packaging, barriers and/or containers to maintain sterility, labeling, etc.
- (97) As used herein, "instructions" refers to a display of written, printed or graphic matter on the immediate container of an article, for example the written material displayed on a vial containing a pharmaceutically active

agent, or details on the composition and use of a product of interest included in a kit containing a composition of interest. Instructions set forth the method of the treatment as contemplated to be administered or performed. (98) In the context of a peptide or polypeptide, the term "fragment" as used herein refers to a peptide or polypeptide that comprises less than the full length amino acid sequence. Such a fragment may arise, for example, from a truncation at the amino terminus, a truncation at the carboxy terminus, and/or an internal deletion of a residue(s) from the amino acid sequence. Fragments may, for example, result from alternative RNA splicing or from in vivo protease activity. In certain embodiments, PCSK9 fragments include polypeptides comprising an amino acid sequence of at least 5 contiguous amino acid residues, at least 10 contiguous amino acid residues, at least 15 contiguous amino acid residues, at least 20 contiguous amino acid residues, at least 25 contiguous amino acid residues, at least 40 contiguous amino acid residues, at least 50 contiguous amino acid residues, at least 60 contiguous amino residues, at least 70 contiguous amino acid residues, at least 80 contiguous amino acid residues, at least 90 contiguous amino acid residues, at least contiguous 100 amino acid residues, at least 125 contiguous amino acid residues, at least 150 contiguous amino acid residues, at least 175 contiguous amino acid residues, at least 200 contiguous amino acid residues, or at least 250 contiguous amino acid residues of the amino acid sequence of a hPCSK9 polypeptide or an antibody that specifically binds to a hPCSK9 polypeptide. In a specific embodiment, a fragment of a hPCSK9 polypeptide or an antibody that specifically binds to a hPCSK9 antigen retains at least 1, at least 2, or at least 3 functions of the polypeptide or antibody.

(99) The term "free" can refer to a polypeptide, for example, PCSK9 or fragments and variants thereof, that is combined with a buffer, wherein the polypeptide is not associated with a cell surface or cell membrane. As such, the term "free" can refer to a polypeptide that is capable of surface expression (i.e. includes one or more transmembrane domains or membrane association domains), but that is not, in its present state, expressed on the surface of a cell or bound to a protein that is expressed on the surface of a cell. A free polypeptide can also refer to a free recombinant or native or unbound polypeptide. In the context of phage display, a free antigen can be selected in solution (referred to herein as a "soluble selection") or adsorbed to a surface, for example, adsorbed to the surface of a 96-well plate (referred to herein as "biopanning selection").

(100) The term "fusion protein" as used herein refers to a polypeptide that comprises an amino acid sequence of an antibody and an amino acid sequence of a heterologous polypeptide or protein (i.e. a polypeptide or protein not normally a part of the antibody (e.g. a non-anti-PCSK9 antigen antibody)). The term "fusion" when used in relation to PCSK9 or to an anti-PCSK9 antibody refers to the joining of a peptide or polypeptide, or fragment, variant and/or derivative thereof, with a heterologous peptide or polypeptide. Preferably, the fusion protein retains the biological activity of the PCSK9 or anti-PCSK9 antibody. In certain embodiments, the fusion protein comprises a PCSK9 antibody VH domain, VL domain, VH CDR (one, two or three VH CDRs), and/or VL CDR (one, two or three VL CDRs), wherein the fusion protein specifically binds to a PCSK9 epitope.

(101) The term "heavy chain" when used with reference to an antibody refers to five distinct types, called alpha (α) , delta (δ), epsilon (ϵ), gamma (γ) and mu (μ), based on the amino acid sequence of the heavy chain constant domain. These distinct types of heavy chains are well known and give rise to five classes of antibodies, IgA, IgD, IgE, IgG and IgM, respectively, including four subclasses of IgG, namely IgG1, IgG2, IgG3 and IgG4. Preferably the heavy chain is a human heavy chain. In the human population, multiple heavy chain constant region alleles, of each immunoglobulin or immunoglobulin subclass, exist. The nucleotide and amino acid sequences of these allelic variants are accessible on publicly available databases such as IMGT, ENSEMBL Swiss-Prot and Uniprot. Allelic variants may also be identified in various genome sequencing projects. In one embodiment, the antibodies and antibody fragments disclosed herein comprise a heavy chain encoded by a IgG1 constant region allele, which includes, but is not limited to, human IGHG1*01 (Seq ID Nos:340, 341 & 537), IGHG1*02 (Seq ID Nos:340, 341 &537), IGHG1*03 (Seq ID Nos:523 & 524), IGHG1*04 (Seq ID Nos:525 & 526) and IGHG1*05 (Seq ID Nos:340, 341 & 537). In one embodiment, the antibodies and antibody fragments disclosed herein comprise a protein encoded by a IgG2 constant region allele, which includes, but is not limited to, human IGHG2*01 (Seq ID Nos:527 & 528), IGHG2*02 (Seq ID Nos:529 & 530), IGHG2*03 (Seq ID Nos:527 & 528), IGHG2*04 (Seq ID Nos:531 & 532), IGHG2*05 (Seg ID Nos:527 & 528) and IGHG2*06 (Seg ID Nos:533 & 534). In one embodiment, the antibodies or antibody fragments disclosed herein comprise a protein encoded by a IgG3 constant region allele, which includes but is not limited to human IGHG3*01, IGHG3*02, IGHG3*03, IGHG3*04, IGHG3*05, IGHG3*06, IGHG3*07, IGHG3*08. IGHG3*09. IGHG3*10. IGHG3*11. IGHG3*12. IGHG3*13. IGHG3*14. IGHG3*15. IGHG3*16. IGHG3*17, IGHG3*18 and IGHG3*19. In one embodiment, the antibodies or antibody fragments disclosed herein comprise a protein encoded by a IgG4 constant region allele, which includes but is not limited to human IGHG4*01 (see, eg, the sequence table herein), IGHG4*02 (see, eg, the sequence table herein), IGHG4*03 (see, eg, the sequence table herein) and IGHG4*04 (see, eg, the sequence table herein). In another example, the heavy chain is a disabled IgG isotype, e.g. a disabled IgG4. In certain embodiments, the antibodies of the invention comprise a human gamma 4 constant region. In another embodiment, the heavy chain constant region does not bind Fc-y receptors, and e.g. comprises a Leu235Glu mutation. In another embodiment, the heavy chain constant region

comprises a Ser228Pro mutation to increase stability.

(102) In another embodiment, the heavy chain constant region is IgG4-PE (see, eg, the sequence table herein). In another embodiment, the antibodies and antibody fragments disclosed herein comprise a heavy chain constant region encoded by a murine IgG1 constant region allele, which includes but is not limited to mouse IGHG1*01 or IGHG1*02. In one embodiment, the antibodies and antibody fragments disclosed herein comprise a heavy chain constant region encoded by a murine IgG2 constant region allele, which includes, but is not limited to, mouse IGHG2A*01, IGHG2A*02, IGHG2B*01, IGHG2B*02, IGHG2C*01, IGHG2C*02 or IGHG2C*03. In one embodiment, the antibodies or antibody fragments disclosed herein comprise a protein encoded by a murine IgG3 constant region allele, which includes but is not limited to mouse IGHG3*01.

(103) The term "host" as used herein refers to an animal, preferably a mammal, and most preferably a human. (104) The term "host cell" as used herein refers to the particular subject cell transfected with a nucleic acid molecule and the progeny or potential progeny of such a cell. Progeny of such a cell may not be identical to the parent cell transfected with the nucleic acid molecule due to mutations or environmental influences that may occur in succeeding generations or integration of the nucleic acid molecule into the host cell genome.

(105) The term "in combination" in the context of the administration of other therapies refers to the use of more than one therapy. The use of the term "in combination" does not restrict the order in which therapies are administered to a subject with a disease. A first therapy can be administered before (e.g. 1 minute, 45 minutes, 30 minutes, 45 minutes, 1 hour, 2 hours, 4 hours, 6 hours, 12 hours, 24 hours, 48 hours, 72 hours, 96 hours, 1 week, 2 weeks, 3 weeks, 4 weeks, 5 weeks, 6 weeks, 8 weeks, or 12 weeks), concurrently, or after (e.g. 1 minute, 45 minutes, 30 minutes, 45 minutes, 1 hour, 2 hours, 4 hours, 6 hours, 12 hours, 24 hours, 48 hours, 72 hours, 96 hours, 1 week, 2 weeks, 3 weeks, 4 weeks, 5 weeks, 6 weeks, 8 weeks, or 12 weeks) the administration of a second therapy to a subject which had, has, or is susceptible to a PCSK9-mediated disease. Any additional therapy can be administered in any order with the other additional therapies. In certain embodiments, the antibodies of the invention can be administered in combination with one or more therapies (e.g. therapies that are not the antibodies of the invention that are currently administered to prevent, treat, manage, and/or ameliorate a PCSK9-mediated disease. Non-limiting examples of therapies that can be administered in combination with an antibody of the invention include analgesic agents, anaesthetic agents, antibiotics, or immunomodulatory agents or any other agent listed in the U.S. Pharmacopoeia and/or Physician's Desk Reference.

(106) As used herein, "injection device" refers to a device that is designed for carrying out injections, an injection including the steps of temporarily fluidically coupling the injection device to a person's tissue, typically the subcutaneous tissue. An injection further includes administering an amount of liquid drug into the tissue and decoupling or removing the injection device from the tissue. In some embodiments, an injection device can be an intravenous device or IV device, which is a type of injection device used when the target tissue is the blood within the circulatory system, e.g. the blood in a vein. A common, but non-limiting example of an injection device is a needle and syringe. As used herein, "instructions" refers to a display of written, printed or graphic matter on the immediate container of an article, for example the written material displayed on a vial containing a pharmaceutically active agent, or details on the composition and use of a product of interest included in a kit containing a composition of interest. Instructions set forth the method of the treatment as contemplated to be administered or performed. (107) An "isolated" or "purified" antibody or protein is one that has been identified, separated and/or recovered from a component of its production environment (e.g. natural or recombinant). For example, the antibody or protein is substantially free of cellular material or other contaminating proteins from the cell or tissue source from which the antibody is derived, or substantially free of chemical precursors or other chemicals when chemically synthesized. The language "substantially free of cellular material" includes preparations of an antibody in which the antibody is separated from cellular components of the cells from which it is isolated or recombinantly produced. Thus, an antibody that is substantially free of cellular material includes preparations of antibody having less than about 30%, 20%, 10%, or 5% (by dry weight) of heterologous protein (also referred to herein as a "contaminating protein"). When the antibody is recombinantly produced, it is also preferably substantially free of culture medium, i.e. culture medium represents less than about 20%, 10%, or 5% of the volume of the protein preparation. When the antibody is produced by chemical synthesis, it is preferably substantially free of chemical precursors or other chemicals, i.e., it is separated from chemical precursors or other chemicals which are involved in the synthesis of the protein. Accordingly, such preparations of the antibody have less than about 30%, 20%, 10%, 5% (by dry weight) of chemical precursors or compounds other than the antibody of interest. In a preferred embodiment, antibodies of the invention are isolated or purified.

(108) The terms "Kabat numbering," and like terms are recognized in the art and refer to a system of numbering amino acid residues which are more variable (i.e. hypervariable) than other amino acid residues in the heavy chain variable regions of an antibody, or an antigen binding portion thereof (Kabat et al., (1971) Ann. NY Acad. Sci., 190:382-391 and, Kabat et al. (1991) Sequences of Proteins of Immunological Interest, Fifth Edition, U.S. Department of Health and Human Services, NIH Publication No. 91-3242). For the heavy chain variable region, the

hypervariable region typically ranges from amino acid positions 31 to 35 for CDR1, amino acid positions 50 to 65 for CDR2, and amino acid positions 95 to 102 for CDR3.

(109) "Label" or "labelled" as used herein refers to the addition of a detectable moiety to a polypeptide, for example, a radiolabel, fluorescent label, enzymatic label, chemiluminescent label or a biotinyl group or gold. Radioisotopes or radionuclides may include .sup.3H, .sup.14C, .sup.15N, .sup.355, .sup.90Y, .sup.99Tc, .sup.115In, .sup.125I, .sup.131I, fluorescent labels may include rhodamine, lanthanide phosphors or FITC and enzymatic labels may include horseradish peroxidase, 3-galactosidase, luciferase, alkaline phosphatase. Additional labels include, by way of illustration and not limitation: enzymes, such as glucose-6-phosphate dehydrogenase ("G6PDH"), alpha-Dgalactosidase, glucose oxydase, glucose amylase, carbonic anhydrase, acetylcholinesterase, lysozyme, malate dehydrogenase and peroxidase; dyes (e.g. cyanine dyes, e.g. Cy5TM, Cy5.5TM. or Cy7TM); additional fluorescent labels or fluorescers include, such as fluorescein and its derivatives, fluorochrome, GFP (GFP for "Green Fluorescent Protein"), other fluorescent proteins (e.g. mCherry, mTomato), dansyl, umbelliferone, phycoerythrin, phycocyanin, allophycocyanin, o-phthaldehyde, and fiuorescamine; fluorophores such as lanthanide cryptates and chelates e.g. Europium etc (Perkin Elmer and Cisbio Assays); chemoluminescent labels or chemiluminescers, such as isoluminol, luminol and the dioxetanes; sensitisers; coenzymes; enzyme substrates; particles, such as latex or carbon particles; metal sol; crystallite; liposomes; cells, etc., which may be further labelled with a dye, catalyst or other detectable group; molecules such as biotin, digoxygenin or 5-bromodeoxyuridine; toxin moieties, such as for example a toxin moiety selected from a group of *Pseudomonas* exotoxin (PE or a cytotoxic fragment or mutant thereof), Diptheria toxin or a cytotoxic fragment or mutant thereof, a botulinum toxin A, B, C, D, E or F, ricin or a cytotoxic fragment thereof e.g. ricin A, abrin or a cytotoxic fragment thereof, saporin or a cytotoxic fragment thereof, pokeweed antiviral toxin or a cytotoxic fragment thereof and bryodin 1 or a cytotoxic fragment thereof. (110) The term "light chain" when used in reference to an antibody refers to the immunoglobulin light chains, of which there are two types in mammals, lambda (λ) and kappa (κ). Preferably, the light chain is a human light chain. Preferably the light chain constant region is a human constant region. In the human population, multiple light chain constant region alleles exist. The nucleotide and amino acid sequences of these allelic variants are accessible on publicly available databases such as IMGT, ENSEMBL, Swiss-Prot and Uniprot, In one embodiment, the antibodies or antibody fragments disclosed herein comprise a protein encoded by a human K constant region allele, which includes, but is not limited to, IGKC*01 (see, eg, the sequence table herein), IGKC*02 (see, eg, the sequence table herein), IGKC*03 (see, eg, the sequence table herein), IGKC*04 (see, eg, the sequence table herein) and IGKC*05 (see, eg, the sequence table herein). In one embodiment, the antibodies or antibody fragments disclosed herein comprise a protein encoded by a human A constant region allele, which includes but is not limited to IGLC1*01 (see, eg, the sequence table herein), IGLC1*02 (see, eg, the sequence table herein), IGLC2*01 (see, eg, the sequence table herein), IGLC2*02 (see, eg, the sequence table herein), IGLC2*03 (see, eg, the sequence table herein), IGLC3*01 (see, eg, the sequence table herein), IGLC3*02 (see, eg, the sequence table herein), IGLC3*03 (see, eg, the sequence table herein), IGLC3*04 (see, eg, the sequence table herein), IGLC6*01 (see, eg, the sequence table herein), IGLC7*01 (see, eg, the sequence table herein), IGLC7*02 (see, eg, the sequence table herein), IGLC7*03 (see, eg, the sequence table herein). In another embodiment, the antibodies and antibody fragments disclosed herein comprise a light chain constant region encoded by a mouse K constant region allele, which includes, but is not limited to, IGKC*01, IGKC*03 or IGKC*03. In another embodiment, the antibodies and antibody fragments disclosed herein comprise a light chain constant region encoded by a mouse A constant region allele, which includes, but is not limited to, IGLC1*01, IGLC2*01 or IGLC3*01.

(111) "Percent (%) amino acid sequence identity" and "homology" with respect to a peptide, polypeptide or antibody sequence are defined as the percentage of amino acid residues in a candidate sequence that are identical with the amino acid residues in the specific peptide or polypeptide sequence, after aligning the sequences and introducing gaps, if necessary, to achieve the maximum percent sequence identity, and not considering any conservative substitutions as part of the sequence identity. Alignment for purposes of determining percent amino acid sequence identity can be achieved in various ways that are within the skill in the art, for instance, using publicly available computer software such as BLAST, BLAST-2, ALIGN or MEG ALIGN™ (DNASTAR) software. In one embodiment, the % homology is about 70%. In one embodiment, the % homology is about 75%. In one embodiment, the % homology is about 85%. In one embodiment, the % homology is about 92%. In one embodiment, the % homology is about 95%. In one embodiment, the % homology is about 97%. In one embodiment, the % homology is about 98%. In one embodiment, the % homology is about 99%. In one embodiment, the % homology is about 98%. In one embodiment, the % homology is about 99%. In one embodiment, the % homology is about 98%. In one embodiment, the % homology is about 99%. In one embodiment, the % homology is about 99%. In one embodiment, the % homology is about 99%. In one

(112) The term "naturally occurring" or "native" when used in connection with biological materials such as nucleic acid molecules, polypeptides, host cells, and the like, refers to those which are found in nature and not manipulated by a human being.

(113) As used herein, "packaging" refers to how the components are organized and/or restrained into a unit fit for

distribution and/or use. Packaging can include, e.g. boxes, bags, syringes, ampoules, vials, tubes, clamshell packaging, barriers and/or containers to maintain sterility, labelling, etc. The term "pharmaceutically acceptable" as used herein means being approved by a regulatory agency of the Federal or a state government, or listed in the U.S. Pharmacopeia, European Pharmacopeia or other generally recognized Pharmacopeia for use in animals, and more particularly in humans.

- (114) As used herein, the term "polynucleotide," "nucleotide," nucleic acid" "nucleic acid molecule" and other similar terms are used interchangeable and include DNA, RNA, mRNA and the like. As used herein, the terms "prevent", "preventing", and "prevention" refer to the total or partial inhibition of the development, recurrence, onset or spread of a hPCSK9-mediated disease and/or symptom related thereto, resulting from the administration of a therapy or combination of therapies provided herein (e.g. a combination of prophylactic or therapeutic agents, such as an antibody of the invention).
- (115) The term "soluble" refers to a polypeptide, such as PCSK9 and variants or fragments thereof, that is lacking one or more transmembrane or cytoplasmic domains found in the native or membrane-associated form. In one embodiment, the "soluble" form of PCSK9 lacks both the transmembrane domain and the cytoplasmic domain. (116) The term "subject" or "patient" refers to any animal, including, but not limited to, mammals. As used herein, the term "mammal" refers to any vertebrate animal that suckle their young and either give birth to living young (eutharian or placental mammals) or are egg-laying (metatharian or nonplacental mammals). Examples of mammalian species include, but are not limited to, humans and other primates, including non-human primates such as chimpanzees and other apes and monkey species; farm animals such as cattle, sheep, pigs, goats and horses; domestic mammals such as dogs and cats; laboratory animals including rodents such as mice, rats (including cotton rats) and guinea pigs; birds, including domestic, wild and game birds such as chickens, turkeys and other gallinaceous birds, ducks, geese, and the like.
- (117) As used herein "substantially all" refers to refers to at least about 60%, at least about 70%, at least about 75%, at least about 80%, at least about 85%, at least about 90%, at least about 95%, at least about 98%, at least about 99%, or about 100%.
- (118) As used herein, the term "therapeutic agent" refers to any agent that can be used in the treatment, management or amelioration of a PCSK9-mediated disease and/or a symptom related thereto. In certain embodiments, the term "therapeutic agent" refers to an antibody of the invention. In certain other embodiments, the term "therapeutic agent" refers to an agent other than an antibody of the invention. Preferably, a therapeutic agent is an agent which is known to be useful for, or has been or is currently being used for the treatment, management or amelioration of a PCSK9-mediated disease or one or more symptoms related thereto. In specific embodiments, the therapeutic agent is a fully human anti-PCSK9 antibody, such as a fully human anti-PCSK9 monoclonal antibody.
- (119) As used herein, the term "therapy" refers to any protocol, method and/or agent that can be used in the prevention, management, treatment and/or amelioration of a PCSK9-mediated disease (e.g. cancer). In certain embodiments, the terms "therapies" and "therapy" refer to a biological therapy, supportive therapy, and/or other therapies useful in the prevention, management, treatment and/or amelioration of a PCSK9-mediated disease known to one of skill in the art such as medical personnel.
- (120) The terms "treat", "treatment" and "treating" refer to the reduction or amelioration of the progression, severity, and/or duration of a hPCSK9-mediated disease (e.g. cancer) resulting from the administration of one or more therapies (including, but not limited to, the administration of one or more prophylactic or therapeutic agents, such as an antibody of the invention). In specific embodiments, such terms refer to the reduction or inhibition of the binding of hPCSK9 to a BMP receptor or HJV, and/or the inhibition or reduction of one or more symptoms associated with a PCSK9-mediated disease, such as hyperlipidaemia or hypercholesterolaemia.
- (121) The term "variable region" or "variable domain" refers to a portion of the light and heavy chains, typically about the amino-terminal 120 to 130 amino acids in the heavy chain and about 100 to 110 amino acids in the light chain, which differ extensively in sequence among antibodies and are used in the binding and specificity of each particular antibody for its particular antigen. The variability in sequence is concentrated in those regions called complimentarily determining regions (CDRs) while the more highly conserved regions in the variable domain are called framework regions (FR). The CDRs of the PCSK9 and heavy chains are primarily responsible for the interaction of the antibody with antigen. Numbering of amino acid positions used herein is according to the EU Index, as in Kabat et al. (1991) Sequences of proteins of immunological interest. (U.S. Department of Health and Human Services, Washington, D.C.) 5.sup.th ed. ("Kabat et al."). In preferred embodiments, the variable region is a human variable region.
- (122) Definitions of common terms in cell biology and molecular biology can be found in "The Merck Manual of Diagnosis and Therapy", 19.sup.th Edition, published by Merck Research Laboratories, 2006 (ISBN 0-911910-19-0); Robert S. Porter et al. (eds.), The Encyclopedia of Molecular Biology, published by Blackwell Science Ltd., 1994 (ISBN 0-632-02182-9); Benjamin Lewin, Genes X, published by Jones & Bartlett Publishing, 2009 (ISBN-10: 0763766321); Kendrew et al. (Eds.), Molecular Biology and Biotechnology: a Comprehensive Desk Reference,

published by VCH Publishers, Inc., 1995 (ISBN 1-56081-569-8) and Current Protocols in Protein Sciences 2009, Wiley Intersciences, Coligan et al., eds.

(123) Unless otherwise stated, the present invention was performed using standard procedures, as described, for example in Sambrook et al., Molecular Cloning: A Laboratory Manual (4 ed.), Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y., USA (2012); Davis et al., Basic Methods in Molecular Biology, Elsevier Science Publishing, Inc., New York, USA (1995); or Methods in Enzymology: Guide to Molecular Cloning Techniques Vol. 152, S. L. Berger and A. R. Kimmel Eds., Academic Press Inc., San Diego, USA (1987); Current Protocols in Protein Science (CPPS) (John E. Coligan, et al., ed., John Wiley and Sons, Inc.), Current Protocols in Cell Biology (CPCB) (Juan S. Bonifacino et al. ed., John Wiley and Sons, Inc.), and Culture of Animal Cells: A Manual of Basic Technique by R. Ian Freshney, Publisher: Wiley-Liss; 5th edition (2005), Animal Cell Culture Methods (Methods in Cell Biology, Vol. 57, Jennie P. Mather and David Barnes editors, Academic Press, 1st edition, 1998) which are all incorporated by reference herein in their entireties. Other terms are defined herein within the description of the various aspects of the invention.

(124) Anti-PCSK9 Antibodies & Fragments

- (125) The invention provides various anti-PCSK9 antibodies and fragments (such as Fab or scFv fragments), uses, methods and combinations (eg, with statin). Examples are set out in the following numbered Clauses. 1. An antibody or fragment comprising a binding site which specifically binds to Proprotein Convertase Subtilisin Kexin type 9 (PCSK9), wherein the binding site comprises a V.sub.H domain that is encoded by a nucleotide sequence that is derived from the recombination of a human V.sub.H gene segment, DH gene segment and JH gene segment, wherein the V.sub.H gene segment is selected from IGHV4-59 and IGHV3-9.
- (126) For example, the VH gene segment is IGHV4-59*01 and the DH gene segment and JH gene segments are human gene segments. For example, the V.sub.H gene segment is IGHV3-9*01 and the DH gene segment and JH gene segments are human gene segments.
- (127) In an example, specific binding is with a KD, K.sub.off and/or K.sub.on as described further below. In an example, specific binding is with a KD from 1 μ M to 5 nM.
- (128) The skilled person is familiar with databases and other sources for human and other species of antibody gene segments. For example, the IMGT database (www IMGT.org) is a suitable source, eg, the version as at 1 Sep. 2018. (129) Reference is made to the Examples, showing antibodies that are based on IGHV4-59 and IGHV3-9. Surprisingly, this human VH gene segment produces anti-PCSK9 antibodies which have desirable anti-PCSK9 properties, such as those described in, eg, in the Examples.
- (130) For example, the antibody or fragment of the invention is for administration to a subject for decreasing plasma total cholesterol and plasma non-HDL-cholesterol in a dose-dependent fashion in the subject.
- (131) For example, the antibody or fragment of the invention is for administration to a subject for decreasing plasma total cholesterol in the subject.
- (132) For example, the antibody or fragment of the invention is for administration to a subject for decreasing plasma non-HDL-cholesterol (optionally in a dose-dependent fashion) in the subject.
- (133) For example, the antibody or fragment of the invention is for administration to a subject for administration to a subject for decreasing plasma triglyceride levels) in the subject.
- (134) In an embodiment of any of these examples, HDL-cholesterol is not reduced or is not significantly reduced in the subject.
- (135) The examples show beneficial dosing of the antibodies of the invention compared to alirocumab benchmark. For example, the antibody or fragment of the invention is for administration to a subject for treating or preventing a PCSK9-mediated disease or condition (eg, heterozygous familial hypercholesterolaemia (HeFH) or homozygous familial hypercholesterolaemia (HoFH)), wherein the antibody or fragment is administered at a dose that is less than 75 mg every 2 weeks or 300 mg every 4 weeks. For example, the antibody or fragment of the invention is for administration to a subject for treating or preventing a PCSK9-mediated disease or condition (eg. heterozygous familial hypercholesterolaemia (HeFH) or homozygous familial hypercholesterolaemia (HoFH)), wherein the antibody or fragment is administered at a dose that is less than 140 mg every 2 weeks or less than 420 mg every 4 weeks. For example, the antibody or fragment of the invention is for administration to a subject for treating or preventing a PCSK9-mediated disease or condition (eg, heterozygous familial hypercholesterolaemia (HeFH) or homozygous familial hypercholesterolaemia (HoFH)), wherein the antibody or fragment is administered at a dose that is less than 2 weekly dose of Praluent or Repatha. For example, the antibody or fragment of the invention is for administration to a subject for treating or preventing a PCSK9-mediated disease or condition (eg., heterozygous familial hypercholesterolaemia (HeFH) or homozygous familial hypercholesterolaemia (HoFH)), wherein the antibody or fragment is administered at a dose that is less than 4 weekly dose of Praluent or Repatha. For example, the antibody or fragment of the invention is for administration to a subject for treating or preventing a PCSK9mediated disease or condition (eg, heterozygous familial hypercholesterolaemia (HeFH) or homozygous familial hypercholesterolaemia (HoFH)), wherein the antibody or fragment is administered at a 2 weekly or 4 weekly dose

that is less than 70, 65, 60, 50, 40, 30 or 25 mg. (136) The recommended starting dose of Praluent[™] is 75 mg once every 2 weeks administered subcutaneously, since the majority of patients achieve sufficient LDL-C reduction with this dosage or 2×150 mg every 4 weeks. (137) The recommended subcutaneous dosage of RepathaTM in adults with established cardiovascular disease or in adults with primary hyperlipidemia (including heterozygous familial hypercholesterolemia [HeFH]) is either 140 mg every 2 weeks or 420 mg once monthly, based on patient preference for dosing frequency and injection volume. The recommended subcutaneous dosage of RepathaTM in patients with HoFH is 420 mg once monthly. (138) For example, the antibody or fragment comprises a CDRH3 sequence of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT) and optionally a CDRL3 of said selected antibody. For example, the antibody or fragment comprises a CDRH1 and CDRH3 sequence of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT) and optionally a CDRL2 of said selected antibody. For example, the antibody or fragment comprises a CDRH1 and CDRH2 sequence of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT). For example, the antibody or fragment comprises a CDRH2 and CDRH3 sequence of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT). For example, the antibody or fragment comprises an anti-PCSK9 binding site, wherein the binding site comprises a VH domain comprising the CDRH3 sequence of CL-58838 paired with a VL domain of CL-58838. (139) For example, the antibody or fragment comprises an anti-PCSK9 binding site, wherein the binding site comprises a VH domain comprising SEQ ID NO: 1 optionally paired with a VL domain comprising respectively SEQ ID NO: 33. For example, the antibody or fragment comprises an anti-PCSK9 binding site, wherein the binding site comprises a VH domain comprising SEQ ID NO: 65 paired with a VL domain comprising SEQ ID NO: 95. (140) For example, the antibody or fragment comprises an anti-PCSK9 binding site, wherein the binding site comprises a VH domain of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT), optionally paired with a VL domain of the selected antibody. For example, the antibody or fragment comprises an anti-PCSK9 binding site, wherein the binding site comprises a VH domain of CL-58838 paired with a VL domain of CL-58838. 2. The antibody or fragment according to Clause 1, wherein (i) the VH gene segment is IGHV4-59 (eg, IGHV4-59*01) and the DH gene segment is human gene segment IGHD3-10 (eg, IGHD3-10*01); or (ii) the VH gene segment is IGHV3-9 (eg, IGHV3-9*01) and the DH gene segment is human gene segment IGHD3-9 (eg, IGHD3-9*01). 3. The antibody or fragment according to Clause 1 or 2, wherein the JH gene segment is a human gene segment is JH6 (eg, JH6*02). 4. An antibody or fragment which specifically binds to Proprotein Convertase Subtilisin Kexin type 9 (PCSK9) and comprises the CDRH3 sequence of an anti-PCSK9 antibody according to any preceding Clause. 5. An antibody or fragment which specifically binds to Proprotein Convertase Subtilisin Kexin type 9 (PCSK9) and comprises a VH domain which comprises the CDRH3 sequence of any anti-PCSK9 antibody disclosed herein (eg, CL-148219 or CL-148489), or said CDRH3 sequence comprising 3, 2 or 1 amino acid substitution(s). 6. An antibody or fragment (optionally according to any preceding Clause) which specifically binds to Proprotein Convertase Subtilisin Kexin type 9 (PCSK9) and comprises a VH domain which comprises a CDRH3 sequence of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or said sequence comprising 3, 2 or 1 amino acid substitution(s). (141) Optionally, the VH domain comprises a CDRH3 sequence selected from the CDRH3 sequences disclosed herein, or said selected sequence comprising 3, 2 or 1 amino acid substitution(s). 7. The antibody or fragment according to Clause 6, wherein the VH domain comprises (i) a CDRH3 sequence of an antibody selected from CL-274711, CL-148219OLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219OLT); or said

- herein, or said selected sequence comprising 3, 2 or 1 amino acid substitution(s). 7. The antibody or fragment according to Clause 6, wherein the VH domain comprises (i) a CDRH3 sequence of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or said CDRH3 sequence comprising 3, 2 or 1 amino acid substitution(s); and (ii) a CDRH1 sequence of said selected antibody; or said CDRH1 sequence comprising 3, 2 or 1 amino acid substitution(s). 8. The antibody or fragment according to Clause 6 or 7, wherein the VH domain comprises (iii) a CDRH3 sequence of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or said CDRH3 sequence comprising 3, 2 or 1 amino acid substitution(s); and (iv) a CDRH2 sequence of said selected antibody; or said CDRH2 sequence comprising 3, 2 or 1 amino acid substitution(s). 9. An antibody or fragment (optionally according to any preceding Clause) comprising a binding site which specifically binds to Proprotein Convertase Subtilisin Kexin type 9 (PCSK9), wherein the binding site comprises a VH domain that comprises the amino acid sequence of a VH domain of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or an amino acid that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto.
- (142) For example, the identity is at least 85%. For example, the identity is at least 90%. For example, the identity is at least 95%.
- (143) Optionally the VH domain of the antibody or fragment of comprises a VH amino acid sequence of disclosed

herein, or a heavy chain variable domain amino acid sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto. For example, the identity is at least 85%. For example, the identity is at least 90%. For example, the identity is at least 95%, 10. The antibody or fragment according to any preceding Clause comprising first and second copies of said VH domain.

(144) In an example, the antibody or fragment comprises a binding site comprising a VH domain of the invention paired with a VL domain of the invention, wherein the binding site is capable of specifically binding to PCSK9 (eg. mature PCSK9, eg human and/or cynomolgus monkey PCSK9). For example, the antibody or fragment comprise two of such binding sites. 11. An antibody or fragment (optionally according to any preceding Clause) comprising a binding site which specifically binds to Proprotein Convertase Subtilisin Kexin type 9 (PCSK9), wherein the binding site comprises a VL domain that is encoded by a nucleotide sequence that is derived from the recombination of a human VL gene segment and JL gene segment, wherein the VL gene segment is selected from IGKV2-28 (eg, IGKV2-28*01) and IGKV2-29 (eg, IGKV2-29*01). 12. The antibody or fragment according to Clause 11, wherein the VL is a V.sub.K and the JL gene segment is a human gene segment selected from IGKJ3 and IGKJ4. (145) Optionally, the JL gene segment is selected from IGKJ3*01 and IGKJ4*01. 13. An antibody or fragment which specifically binds to Proprotein Convertase Subtilisin Kexin type 9 (PCSK9) and comprises the CDRL3 sequence of an anti-PCSK9 antibody according to Clause 11 or 12. 14. An antibody or fragment (optionally according to any preceding Clause) which specifically binds to Proprotein Convertase Subtilisin Kexin type 9 (PCSK9) and comprises a VL domain which comprises the CDRL3 sequence of any anti-PCSK9 antibody disclosed herein (eg, CL-148219 or CL-148489) or said selected CDRL3 sequence comprising 3, 2 or 1 amino acid substitution(s). 15. The antibody or fragment of Clause 14, comprising a VH domain which comprises the CDRH3 sequence of said selected antibody. 16. An antibody or fragment (optionally according to any preceding Clause) which specifically binds to Proprotein Convertase Subtilisin Kexin type 9 (PCSK9) and comprises a VL domain which comprises a CDRL3 sequence selected from a CDRL3 sequence disclosed herein, or said selected CDRL3 sequence comprising 3, 2 or 1 amino acid substitution(s). 17. An antibody or fragment (optionally according to any preceding Clause) which specifically binds to Proprotein Convertase Subtilisin Kexin type 9 (PCSK9) and comprises a VL domain which comprises a CDRL3 (and optionally a CDRH3) sequence of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or said sequence(s) each comprising 3, 2 or 1 amino acid substitution(s). 18. The antibody or fragment according to Clause 17, wherein the VL domain comprises (i) a CDRL3 sequence (and optionally a CDRH3) of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or said CDR3 sequence(s) each comprising 3, 2 or 1 amino acid substitution(s); and (ii) a CDRL1 (and optionally a CDRH1) sequence of said selected antibody; or said CDR1 sequence(s) each comprising 3, 2 or 1 amino acid substitution(s). 19. The antibody or fragment according to Clause 17 or 18, wherein the VL domain comprises (iii) a CDRL3 (and optionally a CDRH3) sequence of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or said CDR3 sequence(s) each comprising 3, 2 or 1 amino acid substitution(s); and (iv) a CDRL2 (and optionally a CDRH2) sequence of said selected antibody; or said CDR2 sequence(s) each comprising 3, 2 or 1 amino acid substitution(s). 20. An antibody or fragment (optionally according to any preceding Clause) comprising a binding site which specifically binds to Proprotein Convertase Subtilisin Kexin type 9 (PCSK9), wherein the binding site comprises a VL domain that comprises the amino acid sequence of a V.sub.L domain of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or an amino acid that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto.

(146) For example, the identity is at least 85%. For example, the identity is at least 90%. For example, the identity is at least 95%. 21. The antibody or fragment according to any preceding Clause comprising first and second copies of said VL domain.

(147) In an example, the antibody or fragment comprises a binding site comprising a VL domain of the invention paired with a VH domain, wherein the binding site is capable of specifically binding to PCSK9 (eg, mature PCSK9, eg human and/or cynomolgus monkey PCSK9). For example, the antibody or fragment comprise two of such binding sites. 22. An antibody or fragment (optionally according to any preceding Clause) which specifically binds to Proprotein Convertase Subtilisin Kexin type 9 (PCSK9) and comprises the heavy chain amino acid sequence of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or an amino acid that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto. (148) In an example, the heavy chain sequence comprises the VH amino acid sequence of CL-148219 or CL-148489 fused to an antibody heavy chain constant region disclosed herein, eg, an IgG4-PE constant region, eg, SEQ ID NO: 3. Additionally or alternatively, in an example, the light chain sequence comprises the V.sub.L amino acid sequence of CL-148489 fused to an antibody light chain constant region disclosed herein, eg, a kapa constant

(149) For example, the identity is at least 85%. For example, the identity is at least 90%. For example, the identity is

region, eg, SEQ ID NO: 156.

at least 95%. 23. An antibody or fragment (optionally according to any preceding Clause) which specifically binds to Proprotein Convertase Subtilisin Kexin type 9 (PCSK9) and comprises the light chain amino acid sequence of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or an amino acid that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto. (150) For example, the identity is at least 85%. For example, the identity is at least 90%. For example, the identity is at least 95%. 24. The antibody or fragment of Clause 23, comprising the light chain amino acid sequence of said selected antibody; or an amino acid that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto. (151) For example, the identity is at least 85%. For example, the identity is at least 90%. For example, the identity is at least 95%. 25. An antibody or fragment (optionally according to any preceding Clause) which specifically binds to a human PCSK9 epitope that is identical to an epitope to which the antibody of any preceding Clause binds. 26. The antibody or fragment according to Clause 25, wherein the epitope is identified by unrelated amino acid scanning, or by X-ray crystallography.

- (152) Contact amino acid residues involved in the interaction of antibody and antigen may be determined by various known methods to those skilled in the art.
- (153) In one embodiment, sequential replacement of the amino acids of the antigen sequence (using standard molecular biology techniques to mutate the DNA of the coding sequence of the antigen), in this case PCSK9 with Alanine (a.k.a Alanine scan), or another unrelated amino acid, may provide residues whose mutation would reduce or ablate the ability of the antibody to recognise the antigen in question. Binding may be assessed using standard techniques, such as, but not limited to, SPR, HTRF, ELISA (which are described elsewhere herein). Other substitutions could be made to enhance the disruption of binding such as changing the charge on the side chain of antigen sequence amino acids (e.g. Lysine change to glutamic acid), switching polar and non-polar residues (e.g. Serine change to leucine). The alanine scan or other amino substitution method may be carried out either with recombinant soluble antigen, or where the target is a cell membrane target, directly on cells using transient or stable expression of the mutated versions.
- (154) In one embodiment, protein crystallography may be used to determine contact residues between antibody and antigen (i.e. to determine the epitope to which the antibody binds), crystallography allows the direct visualisation of contact residues involved in the antibody-antigen interaction. As well as standard X-ray crystallography, cryo-electro microscopy has been used to determine contact residues between antibodies and HIV capsid protein (see Lee, Jeong Hyun, et L. "Antibodies to a conformational epitope on gp41 neutralize HIV-1 by destabilizing the Env spike.", Nature communications, 6, (2015)).
- (155) In one embodiment, if the antibody recognises a linear epitope, short peptides based on the antigen sequence can be produced and binding of the antibody to these peptides can be assessed using standard techniques, such as, but not limited to, SPR, HTRF, ELISA (which are described elsewhere herein). Further investigation of the epitope could be provided by performing an Alanine scan on any peptides that show binding. Alternative to linear peptides, conformational scans could be carried out using Pepscan technology (http://www.pepscan.com/) using their chemical linkage of peptides onto scaffolds, which has been used to determine discontinuous epitopes on CD20 targeting antibodies (Niederfellner, Gerhard, et al. "Epitope characterization and crystal structure of GA101 provide insights into the molecular basis for type I/II distinction of CD20 antibodies.", Blood, 118.2, (2011), 358-367). (156) In one embodiment, limited proteolytic digestion and mass spectrophotometry can be used to identify binding epitopes. The antibody-antigen complex is digested by a protease, such as, but not limited to, trypsin. The digested complex peptides are compared to antibody-alone and antigen-alone digestion mass spectrophotometry to determine if a particular epitope is protected by the complexation. Further work involving amino acid substitution, competition binding, may then be employed to narrow down to individual amino acid residues involved in the interaction (see, for example, Suckau, Detley, et al. "Molecular epitope identification by limited proteolysis of an immobilized antigen-antibody complex and mass spectrometric peptide mapping.", Proceedings of the National Academy of Sciences, 87.24, (1990), 9848-9852).
- (157) Thus, in one embodiment, the contact residues of the epitope are identified with an unrelated amino acid scan (e.g. alanine scan). In another embodiment, an unrelated amino acid scan (e.g. alanine scan) is carried out using a technique selected from SPR, HTRF, ELISA, X-ray crystallography, cryo-electro microscopy and a combination of limited proteolytic digestion and mass spectrometry.
- (158) In one embodiment, the unrelated amino acid scan (e.g. alanine scan) is carried out using HTRF. In one embodiment, the unrelated amino acid scan (e.g. alanine scan) is carried out using ELISA. When the alanine scan is carried out with either ELISA or HTRF, an amino acid residue is identified as contributing to the epitope if the reduction in signal is at least 25%. In one embodiment, the reduction in signal is at least 30%. In one embodiment, the reduction in signal is at least 40%. In one embodiment, the reduction in signal is at least 50%. In one embodiment, the reduction in signal is at least 50%. In one embodiment, the reduction in signal is at least 60%. In one embodiment, the reduction in signal is at least 50%. In one embodiment, the reduction in signal is at least 60%.

- 75%. In one embodiment, the reduction in signal is at least 80%. In one embodiment, the reduction in signal is at least 85%. In one embodiment, the reduction in signal is at least 90%.
- (159) When the alanine scan is carried out with SPR, an amino acid residue is identified as contributing to the epitope if there is at least a 10-fold reduction in affinity. In one embodiment, the reduction in affinity is at least 15-fold. In one embodiment, the reduction in affinity is at least 30-fold. In one embodiment, the reduction in affinity is at least 40-fold. In one embodiment, the reduction in affinity is at least 50-fold. In one embodiment, the reduction in affinity is at least 100-fold.
- (160) In one embodiment, the contact residues of the epitope are identified by X-ray crystallography. In one embodiment, the contact residues of the epitope are identified by cryo-electro microscopy. In one embodiment, the contact residues of the epitope are identified by a combination of limited proteolytic digestion and mass spectrometry. 27. The antibody or fragment according to Clause 26, wherein the contact residues of the epitope are defined by a reduction in affinity of at least 10-fold in an unrelated amino acid scan, e.g. an alanine scan as determined by SPR.
- (161) In one embodiment, the reduction in affinity is at least 15-fold. In one embodiment, the reduction in affinity is at least 20-fold. In one embodiment, the reduction in affinity is at least 30-fold. In one embodiment, the reduction in affinity is at least 50-fold. In one embodiment, the reduction in affinity is at least 100-fold.
- (162) SPR may be carried out as described herein. 28. An antibody or fragment (optionally according to any preceding Clause) which competes for binding to human PCSK9 with the antibody of any preceding Clause. (163) Optionally, competition is determined by surface plasmon resonance (SPR) or ELISA. The skilled person will be familiar with these techniques and standard conditions, for example.
- (164) In one embodiment, the antibody or fragment competes (e.g. in a dose-dependent manner) with hPCSK9 (or a fusion protein thereof) for binding to cell surface-expressed hPCSK9. In one embodiment, the antibody or fragment competes (e.g. in a dose-dependent manner) with hPCSK9 (or a fusion protein thereof) for binding to soluble hPCSK9.
- (165) Optionally, the competition for binding to hPCSK9 is conducted using SPR. SPR may be carried out as described herein. 29. The antibody or fragment according to any preceding Clause which specifically binds to human PCSK9 comprising any one of SEQ ID NOs: 189-192; and/or a cynomolgus PCSK9 comprising SEQ ID NO: 193; and/or a mouse PCSK9 comprising SEQ ID NO: 194.
- (166) Optionally, the antibody or fragment of the invention specifically binds to the amino acid sequence of SEQ ID NO: 189. Optionally, the antibody or fragment of the invention specifically binds to the amino acid sequence of SEQ ID NO: 193. Optionally, the antibody or fragment of the invention specifically binds to the amino acid sequence of SEQ ID NO: 194.
- (167) In an example, PCSK9 herein is a human, mouse or cynomolgus monkey PCSK9.
- (168) In one embodiment, the antibody or fragment binds to cynomolgus PCSK9 with an affinity of less than 1 nM (e.g. from 1 nM to 0.01 pM or from 1 nM to 0.1 pM, or from 1 nM to 1 pM). In one embodiment, the antibody or fragment binds to cynomolgus PCSK9 with an affinity of less than 10 nM (e.g. from 10 nM to 0.01 pM or from 10 nM to 0.1 pM, or from 10 nM to 1 pM). In one embodiment, the antibody or fragment binds to cynomolgus PCSK9 with an affinity of less than 0.1 nM (e.g. from 0.1 nM to 0.01 pM or from 0.1 nM to 0.1 pM, or from 0.1 nM to 1 pM). In one embodiment, the antibody or fragment binds to cynomolgus PCSK9 with an affinity of less than 0.01 nM (e.g. from 0.011 nM to 0.01 pM or from 0.01 nM to 0.11 pM).
- (169) In one embodiment, the antibody or fragment binds to cynomolgus PCSK9 with an affinity of within 2-fold of the affinity to hPCSK9. In one embodiment, the antibody or fragment binds to cynomolgus PCSK9 with an affinity of within 4-fold of the affinity to hPCSK9. In one embodiment, the antibody or fragment binds to cynomolgus PCSK9 with an affinity of within 5-fold of the affinity to hPCSK9. In one embodiment, the antibody or fragment binds to cynomolgus PCSK9 with an affinity of within 8-fold of the affinity to hPCSK9. In one embodiment, the antibody or fragment binds to cynomolgus PCSK9 with an affinity of within 8-fold of the affinity to hPCSK9. In one embodiment, the antibody or fragment binds to cynomolgus PCSK9 with an affinity of within 10-fold of the affinity to hPCSK9. "hPCSK9" herein is a human PCSK9, eg, a human PCSK9 disclosed herein, eg, comprising SEQ ID NO: 562.
- (170) In one embodiment, the antibody or fragment does not detectably bind to cynomolgus PCSK9. In one embodiment, the antibody or fragment does not detectably bind to murine (eg, mouse and/or rat) PCSK9. (171) In one embodiment, the antibody or fragment binds to murine (eg, mouse and/or rat) PCSK9 with an affinity of less than 1 nM (e.g. from 1 nM to 0.01 pM or from 1 nM to 0.1 pM, or from 1 nM to 1 pM). In one embodiment, the antibody or fragment binds to murine PCSK9 with an affinity of less than 10 nM (e.g. from 10 nM to 0.01 pM or from 10 nM to 0.1 pM, or from 10 nM to 1 pM). In one embodiment, the antibody or fragment binds to murine PCSK9 with an affinity of less than 0.1 nM (e.g. from 0.1 nM to 0.01 pM or from 0.1 nM to 0.1 pM, or from 0.1 nM to 1 pM). In one embodiment, the antibody or fragment binds to murine PCSK9 with an affinity of less than 0.01

- nM (e.g. from 0.011 nM to 0.01 pM or from 0.01 nM to 0.1 pM).
- (172) Optionally, the antibody or fragment comprises an effector-enabled or effector-disabled constant region, such as a human constant region, for example an effector-null human constant region, e.g. an IgG4 constant region or an IgG1 constant region, optionally wherein the constant region is IgG4-PE, or a disabled IgG1. Optionally, the antibody or fragment comprises a murine (eg, mouse and/or rat) constant region. Optionally, the antibody or fragment comprises any of the heavy chain constant region sequences described herein.
- (173) Optionally, the constant region has CDC and/or ADCC activity. 30. The antibody or fragment according to any preceding Clause, wherein the antibody or fragment comprises a human constant region, e.g. an IgG4 constant region or an IgG1 constant region.
- (174) For example, the constant region comprises a heavy chain constant region disclosed herein.
- (175) In an example (optionally in addition to the heavy chain region as per the paragraph immediately above), the constant region comprises a light chain constant region, the light chain constant region comprising a light chain constant region amino acid sequence disclosed herein. 31. The antibody or fragment according to Clause 30, wherein the constant region is an IgG4-PE constant region.
- (176) The anti-PCSK9 antibody or fragment according to the invention may comprise a constant region, such as a human constant region, for example an effector-null human constant region, e.g. an IgG4 constant region or an IgG1 constant region, optionally wherein the constant region is IgG4-PE, or a disabled IgG1 as defined in the sequence table herein.
- (177) In other embodiments, the antibody or fragment is any of the isotypes or constant regions as defined herein. In one embodiment, the constant region is wild-type human IgG1. For example, the constant region is an effector-enabled IgG1 constant region, optionally having ADCC and/or CDC activity. In one embodiment, the constant region is engineered for enhanced ADCC and/or CDC and/or ADCP. In another embodiment, the constant region is engineered for enhanced effector function.
- (178) The IgG4 constant region may be any of the IgG4 constant region amino acid sequences or encoded by any of the nucleic acid sequences of the sequence table herein. A heavy chain constant region may be an IgG4 comprising both the Leu235Glu mutation and the Ser228Pro mutation. This "IgG4-PE" heavy chain constant region (see the sequence table for an example) is effector null.
- (179) An alternative effector null human constant region is a disabled IgG1 being an IgG1*01 allele comprising the L235A and/or G237A mutations (e.g. LAGA, see the sequence table). In one embodiment, the antibodies or antibody fragments disclosed herein comprise an IgG1 heavy chain constant region, wherein the sequence contains alanine at position 235 and/or 237 (EU index numbering).
- (180) The potency of Fc-mediated effects may be enhanced by engineering the Fc domain by any of the techniques as will be apparent to the skilled person. In another embodiment, the antibodies and fragments disclosed herein may comprise a triple mutation (M252Y/S254T/T256E) which enhances binding to FcRn. 32. The antibody or fragment according to any preceding Clause (eg, a bispecific antibody), further comprising an antigen-binding site that specifically binds another target antigen (eg, ANGPTL3, eg, human ANGPTL3).
- (181) In an example, the further binding site is an agonist binding site for said another antigen. In an example, the further binding site is an antagonist binding site for said another antigen.
- (182) In an example, the further binding site is an antibody binding site comprising a VH and a VL; a binding site comprised by a constant domain of the antibody (eg, an Fcab binding site) or a non-immunoglobulin binding site (eg, a fibronectin domain). Optionally, the antigen-binding site is any antigen-binding site disclosed herein.
- (183) For example, the antibody or fragment is a bispecific antibody or fragment. For example, the antibody or fragment is a dual binding antibody or fragment, or a fusion protein comprising an antibody or fragment thereof as defined in any preceding Clause. A dual binding antibody has the meaning as set out above.
- (184) In an example, the antibody or fragment comprises a bispecific format selected from DVD-Ig, mAb.sup.2, FIT-Ig, mAb-dAb, dock and lock, SEEDbody, scDiabody-Fc, diabody-Fc, tandem scFv-Fc, Fab-scFv-Fc, Fab-scFv, intrabody, BiTE, diabody, DART, TandAb, scDiabody, scDiabody-CH.sub.3, Diabody-CH.sub.3, minibody, knobs-in-holes, knobs-in-holes with common light chain, knobs-in-holes with common light chain and charge pairs, charge pairs with common light chain, in particular mAb.sup.2, knob-in-holes, knob-in-holes with common light chain, knobs-in-holes with common light chain and charge pairs and FIT-Ig, e.g. mAb.sup.2 and FIT-Ig. (185) In one embodiment, the bispecific format is selected from DVD-Ig, mAb.sup.2, FIT-Ig, mAb-dAb, dock and lock, Fab-arm exchange, SEEDbody, Triomab, LUZ-Y, Fcab, Kλ-body, orthogonal Fab, scDiabody-Fc, diabody-Fc, tandem scFv-Fc, Fab-scFv-Fc, intrabody, BiTE, diabody, DART, TandAb, scDiabody, scDiabody-CH.sub.3, Diabody-CH.sub.3, Triple body, Miniantibody, minibody, TriBi minibody, scFv-CH.sub.3 KIH, scFv-CH-CL-scFv, F(ab').sub.2-scFv, scFv-KIH, Fab-scFv-Fc, tetravalent HCab, ImmTAC, knobs-in-holes, knobs-in-holes with common light chain, knobs-in-holes with common light chain, DT-IgG, DutaMab, IgG(H)-scFv, scFv-(H)IgG, IgG(L)-scFv, scFv-(L)IgG, IgG(L,H)-Fv, IgG(H)-V, V(H)—IgG, IgG(L)-V, V(L)-IgG, KIH IgG-scFab, 2scFv-IgG, IgG-2scFv, scFv4-Ig and zybody.

(186) In one embodiment, the bispecific format is selected from DVD-Ig, FIT-Ig, mAb-dAb, dock and lock, Fab-arm exchange, SEEDbody, Triomab, LUZ-Y, Fcab, Kλ-body, orthogonal Fab, scDiabody-Fc, diabody-Fc, tandem scFv-Fc, Fab-scFv-Fc, Fab-scFv, intrabody, BiTE, diabody, DART, TandAb, scDiabody, scDiabody-CH.sub.3, Diabody-CH.sub.3, Triple body, Miniantibody, minibody, TriBi minibody, scFv-CH.sub.3 KIH, scFv-CH-CL-scFv, F(ab').sub.2-scFv, scFv-KIH, Fab-scFv-Fc, tetravalent HCab, ImmTAC, knobs-in-holes, knobs-in-holes with common light chain, knobs-in-holes with common light chain and charge pairs, charge pairs, charge pairs with common light chain, DT-IgG, DutaMab, IgG(H)-scFv, scFv-(H)IgG, IgG(L)-scFv, scFv-(L)IgG, IgG(L,H)-Fv, IgG(H)-V, V(H)—IgG, IgG(L)-V, V(L)-IgG, KIH IgG-scFab, 2scFv-IgG, IgG-2scFv, scFv4-Ig and zybody, for example DVD-Ig, FIT-Ig, mAb-dAb, dock and lock, SEEDbody, scDiabody-Fc, diabody-Fc, tandem scFy-Fc, FabscFv-Fc, Fab-scFv, intrabody, BiTE, diabody, DART, TandAb, scDiabody, scDiabody-CH.sub.3, Diabody-CH.sub.3, minibody, knobs-in-holes, knobs-in-holes with common light chain, knobs-in-holes with common light chain and charge pairs, charge pairs, charge pairs with common light chain, in particular knob-in-holes, knob-in-holes with common light chain, knobs-in-holes with common light chain and charge pairs and FIT-Ig, e.g. FIT-Ig. (187) In one embodiment, the bispecific format is selected from DVD-Ig, mAb.sup.2, mAb-dAb, dock and lock, Fab-arm exchange, SEEDbody, Triomab, LUZ-Y, Fcab, Kλ-body, orthogonal Fab, scDiabody-Fc, diabody-Fc, tandem scFv-Fc, Fab-scFv-Fc, Fab-scFv, intrabody, BiTE, diabody, DART, TandAb, scDiabody, scDiabody-CH.sub.3, Diabody-CH.sub.3, Triple body, Miniantibody, minibody, TriBi minibody, scFy-CH.sub.3 KIH, scFy-CH-CL-scFv, F(ab').sub.2-scFv, scFv-KIH, Fab-scFv-Fc, tetravalent HCab, ImmTAC, knobs-in-holes, knobs-in-holes with common light chain, knobs-in-holes with common light chain and charge pairs, charge pairs, charge pairs with common light chain, DT-IgG, DutaMab, IgG(H)-scFv, scFv-(H)IgG, IgG(L)-scFv, scFv-(L)IgG, IgG(L,H)-Fv, IgG(H)-V, V(H)—IgG, IgG(L)-V, V(L)-IgG, KIH IgG-scFab, 2scFv-IgG, IgG-2scFv, scFv4-Ig and zybody, for example DVD-Ig, mAb.sup.2, mAb-dAb, dock and lock, SEEDbody, scDiabody-Fc, diabody-Fc, tandem scFv-Fc, Fab-scFv-Fc, Fab-scFv, intrabody, BiTE, diabody, DART, TandAb, scDiabody, scDiabody-CH.sub.3, Diabody-CH.sub.3, minibody, knobs-in-holes, knobs-in-holes with common light chain, knobs-in-holes with common light chain and charge pairs, charge pairs, charge pairs with common light chain, in particular mAb.sup.2, knob-in-holes, knobs-in-holes with common light chain and charge pairs, and knob-in-holes with common light chain, e.g. mAb.sup.2.

(188) In one embodiment, the bispecific format is selected from DVD-Ig, mAb-dAb, dock and lock, Fab-arm exchange, SEEDbody, Triomab, LUZ-Y, Fcab, Kλ-body, orthogonal Fab, scDiabody-Fc, diabody-Fc, tandem scFv-Fc, Fab-scFv-Fc, Fab-scFv, intrabody, BiTE, diabody, DART, TandAb, scDiabody, scDiabody-CH.sub.3, Diabody-CH.sub.3, Triple body, Miniantibody, minibody, TriBi minibody, scFv-CH.sub.3 KIH, scFv-CH-CL-scFv, F(ab').sub.2-scFv, scFv-KIH, Fab-scFv-Fc, tetravalent HCab, ImmTAC, knobs-in-holes, knobs-in-holes with common light chain, knobs-in-holes with common light chain and charge pairs, charge pairs with common light chain, DT-IgG, DutaMab, IgG(H)-scFv, scFv-(H)IgG, IgG(L)-scFv, scFv-(L)IgG, IgG(L,H)-Fv, IgG(H)-V, V(H)—IgG, IgG(L)-V, V(L)-IgG, KIH IgG-scFab, 2scFv-IgG, IgG-2scFv, scFv4-Ig and zybody, for example DVD-Ig, mAb-dAb, dock and lock, SEEDbody, scDiabody-Fc, diabody-Fc, tandem scFv-Fc, Fab-scFv-Fc, Fab-scFv, intrabody, BiTE, diabody, DART, TandAb, scDiabody, scDiabody-CH.sub.3, Diabody-CH.sub.3, minibody, knobs-in-holes, knobs-in-holes with common light chain, knobs-in-holes with common light chain and charge pairs, charge pairs with common light chain, in particular knob-in-holes, knobs-in-holes with common light chain and charge pairs, and knob-in-holes with common light chain. 33. An anti-PCSK9 antibody or fragment as defined in any preceding Clause for treating or preventing a PCSK9-mediated disease or condition (eg, hyperlipidaemia or hypercholesterolaemia) in a subject.

(189) In an example, the subject is a human. In an alternative, the subject is a non-human animal. In an example, the subject is an adult human. In an example, the subject is a paediatric human. In an example, the subject is a human CKD patient on dialysis treatment. In an example, the subject is a human having end-stage renal disease. (190) In an example, the antibody or fragment herein is for treating or preventing a disease or condition selected from hypercholesterolemia, hyperlipidemia, hypercholesterolemia, dyslipidemia, cholestatic liver disease, nephrotic syndrome, hypothyroidism, obesity, diabetes, atherosclerosis or a cardiovascular disease. In an example, the disease or condition is selected from a lipid disorder, hyperlipoproteinemia, hyperlipidemia; dyslipidemia; hypercholesterolemia, a heart attack, a stroke, coronary heart disease, atherosclerosis, peripheral vascular disease, claudication, type II diabetes, high blood pressure, and a cardiovascular disease or condition. For example, the disease or condition is hyperlipidaemia. For example, the disease or condition is hypercholesterolaemia. For example, the disease or condition is diabetes. For example, the disease or condition is stroke. For example, the disease or condition is atherosclerosis. For example, the disease or condition is a CNS disorder. For example, the disease or condition is a neurological disorder. For example, the disease or condition is depression. (191) In an example, the disease or condition is in a human. In an example, the disease or condition is in an animal. (192) In an example, the PCSK9-mediated disease or condition is a neurodegenerative disease, disorder or condition, e.g. selected from Alzheimer's disease, amyotrophic lateral sclerosis, Parkinson's disease, Huntington's

pigmentosa; anterior ischemic optic neuropathy, glaucoma, uveitis, depression, trauma-associated stress or posttraumatic stress disorder, frontotemporal dementia, Lewy body dementias, mild cognitive impairments, posterior cortical atrophy, primary progressive aphasia and progressive supranuclear palsy or aged-related dementia, in particular, the neurodegenerative disease, disorder or condition is selected from Alzheimer's disease, amyotrophic lateral sclerosis, Parkinson's disease and Huntington's disease, for example, Alzheimer's disease. (193) In an example, the antibody, fragment, combination of the invention is administered intravenously to the subject; or is for administration intravenously to the subject. In an example, the antibody, fragment, combination of the invention is administered subcutaneously to the subject; or is for administration subcutaneously to the subject. 34. The antibody or fragment of Clause 33, wherein the antibody or fragment is administered to the subject simultaneously or sequentially with a statin. 35. A combination of an amount of an anti-PCSK9 antibody or fragment and an amount of a statin (eg, comprising multiple doses of said antibody and/or statin), wherein the antibody or fragment is according to any one of Clauses 1 to 34. (194) There is also provided: A medical kit comprising the combination, a first sterile container comprising said amount of antibody or fragment, and a second sterile container comprising said amount of statin, and optionally instructions for using the combination to treat hyperlipidaemia or hypercholesterolaemia in a subject. (195) There is also provided: A medical kit comprising the combination, a first sterile container comprising said amount of antibody or fragment, and a second sterile container comprising said amount of a ANGPTL3 inhibitor, and optionally instructions for using the combination to treat hyperlipidaemia or hypercholesterolaemia in a subject. (196) In an example, the combination is for treating or preventing hyperlipidaemia or hypercholesterolaemia in a subject, wherein over a 4 consecutive week period a total dose of the antibody and total dose of statin are administered to said subject in a ratio of X:Y, wherein X is from 10 to 2×10.sup.6 and Y=4, eg, X is from 10 to 2×10.sup.6 micrograms and Y=4 micrograms. 36. The antibody, fragment or combination according to the invention for use in a method of treating or preventing hypercholesterolaemia in a subject that has previously been on a statin treatment regime at a first dose, wherein the method comprises reducing the dose of statin that is administered to the subject or administering no statin to the subject, wherein the method comprises administering the antibody or fragment to the subject. 37. The combination of clause 35 or 36, wherein comprising statin at a daily dose of 10 to 20 mg (eg, 10 mg); or <60 mg (eg, 40 mg). 38. The antibody, fragment or combination of any preceding clause for administering to a human or animal subject suffering from elevated cholesterol, for lowering plasma low density lipoprotein cholesterol (LDL-C) level in the subject after the subject has received the anti-PCSK9 antibody or fragment. 39. Use of the antibody, fragment or combination as defined in any preceding Clause in the manufacture of a medicament for administration to a subject for treating or preventing a PCSK9-mediated disease or condition, e.g. hyperlipidaemia or hypercholesterolaemia. 40. A method of treating or preventing a PCSK9-mediated disease or condition in a subject (e.g. hyperlipidaemia or hypercholesterolaemia), the method comprising administering to said subject a therapeutically effective amount of an antibody, fragment or combination as defined in any one of Clauses 1 to 38, wherein the PCSK9-mediated disease or condition is thereby treated or prevented. (197) The disease or condition can be any disclosed herein. 41. The use according to Clause 39 or the method according to Clause 40, wherein the PCSK9-mediated disease or condition is hyperlipidaemia or hypercholesterolaemia, 42. The antibody, fragment, combination, use or the method according to any one of Clauses 33 to 41, further comprising administering to the subject a further therapy, for example a further therapeutic agent, optionally wherein the further therapeutic agent is selected from the group consisting of a: a. Statin; b. An ANGPTL3 inhibitor (eg, an anti-ANGPTL3 antibody, eg, evinacumab) c. Fibrate; d. Bile acid sequestrant; e. Nicotinic acid; and f. Niacin. (198) The disclosure includes generic versions of the branded drugs instead and the disclosure of these generic drugs

disease, primary progressive multiple sclerosis, secondary progressive multiple sclerosis, corticobasal degeneration,

Rett syndrome, a retinal degeneration disorder selected from age-related macular degeneration and retinitis

is included by reference herein for possible use in the invention, eg, as part of a combination. 43. A pharmaceutical composition comprising an antibody, fragment or combination as defined in any one of Clauses 1 to 38 and 42 and a pharmaceutically acceptable excipient, diluent or carrier and optionally in combination with a further therapeutic agent selected from those mentioned above (eg, in Clause 42). 44. The pharmaceutical composition according to Clause 43 for treating and/or preventing a PCSK9-mediated condition or disease, e.g. hyperlipidaemia or hypercholesterolaemia. 45. The pharmaceutical composition according to Clause 43 or 44 in combination with a label or instructions for use to treat and/or prevent said disease or condition in a human; optionally wherein the label or instructions comprise a marketing authorisation number (e.g., an FDA or EMA authorisation number); optionally wherein the kit comprises an IV or injection device that comprises the antibody or fragment. 46. A nucleic acid that encodes a VH domain and/or a VL domain of an antibody or fragment as defined in any one of Clauses 1 to 32. 47. A nucleic acid that encodes a VH domain comprising the amino acid sequence of a VH domain of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or an amino acid that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto.

(199) For example, the identity is at least 85%. For example, the identity is at least 90%. For example, the identity is at least 95%.

(200) Optionally, there is provided a nucleic acid that encodes a VH domain comprising the amino acid sequence of SEQ ID NO: 114, or an amino acid that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto. For example, the identity is at least 85%. For example, the identity is at least 90%. For example, the identity is at least 95%. 48. A nucleic acid that encodes a VL domain comprising the amino acid sequence of a VL domain of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or an amino acid that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto. (201) Optionally, the nucleic acid also encodes a VH domain comprising the amino acid sequence of a VH domain of the selected antibody; or an amino acid that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto. For example, the identity is at least 85%. For example, the identity is at least 90%. For example, the identity is at least 95%. 49. A nucleic acid (eg, in a host cell, eg, a CHO or HEK293 or Cos cell) comprising a nucleotide sequence that is at least 70% identical to the sequence of SEQ ID NO: 2 or 66.

(202) Herein in any instance where % identity is mentioned, in an example there is 100% identity. 50. A nucleic acid that encodes a heavy chain and/or a light chain of an antibody or fragment as defined in any one of Clauses 1 to 32. 51. A nucleic acid that encodes a heavy chain comprising a VH amino acid sequence that is at least 70% identical to SEQ ID NO: 22 or 66; and a C region amino acid sequence that is at least 70% identical to a IGHG4 sequence, optionally at least 70% identical to SEQ ID NO: 3 or 152. 52. A nucleic acid that encodes a light chain comprising a VL amino acid sequence that is at least 70% identical to SEQ ID NO: 93 or 95; and a C region amino acid sequence that is at least 70% identical to a IGKC sequence, optionally at least 70% identical to SEQ ID NO: 156. 53. A nucleic acid (eg, in a host cell, eg, a CHO or HEK293 or Cos cell) comprising a. a nucleotide sequence that is at least 70% identical to a heavy chain sequence of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); and/or b. a nucleotide sequence that is at least 70% identical to a light chain sequence of an antibody selected respectively from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT). 54. A vector comprising the nucleic acid(s) (eg, the nucleic acid(s) of any one of clauses 46 to 53); optionally wherein the vector is a CHO or HEK293 vector.

(203) All of the nucleic acids of the invention herein are expressible in a host cell, eg, a CHO or HEK293 or Cos cell, such as for expressing a variable domain or chain of an antibody or fragment of the invention. 55. A host cell comprising the nucleic acid(s) (eg, the nucleic acid(s) of any one of Clauses 46 to 53) or the vector of Clause 54. (204) In an example, the antibody or fragment comprises a HCDR3 length of 9, 10, 11 or 12 residues, eg, 10, eg, 11. In an example, the antibody or fragment comprises a LCDR3 length of 7, 8 or 9 residues, eg, 8, eg, 9. In an example, each V.sub.H domain of the antibody or fragment comprises from 1-11 non-germline residues, eg, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10 or 11 non-germline residues. In an example, each V.sub.L domain of the antibody or fragment comprises from 3-8 non-germline residues, eg, 3, 4, 5, 6, 7 or 8 non-germline residues.

- (205) In an embodiment, a CDR sequence herein is determined according to Kabat. In an alternative, the CDR sequence is determined according to IMGT.
- (206) In an example, the selected antibody is CL-148219. In an example, the selected antibody is CL-148489.
- (207) In an example, the selected antibody comprises the heavy chain of CL-148219 or CL-148489.
- (208) In an example, the heavy chain of the antibody or fragment of the invention is a human gamma-1, gamma-2, gamma-3, gamma-4, mu, delta, epsilon or alpha isotype, preferably a gamma isotype (eg, an IgG4 isotype). In an example, the light chain of the antibody or fragment of the invention comprises a human kappa constant region. Alternatively, in an example, the light chain of the antibody or fragment of the invention comprises a human lambda constant region.
- (209) Optionally, the antibody is a 4-chain antibody comprising a dimer of a heavy chain associated with a dimer of a light chain. In an example, the heavy chain comprises one or heavy chain CDRs or a CDR combination as disclosed herein and/or the light chain comprises one or heavy chain CDRs or a CDR combinations as disclosed herein, such as from the same selected antibody. In an example, the heavy chain comprises a VL as disclosed herein, such as from the same selected antibody. In an example, the heavy chain and the light chain are from the same selected antibody, eg, any antibody disclosed in the sequence table herein or the tables in the Examples herein.
- (210) In an example, the selected antibody comprises the light (and optionally the heavy) chain(s) of CL-148219 or CL-148489.
- (211) In an example, the selected antibody comprises the variable domains of CL-148219 or CL-148489.
- (212) In an example, the selected antibody comprises the VH domains of CL-148219 or CL-148489.
- (213) In an example, the selected antibody comprises the VH and VL domains of CL-148219 or CL-148489.
- (214) Optionally, the VH is encoded by a nucleotide sequence that is derived from the recombination of a human IGHV4-59 (eg, IGHV4-59*01) a D gene segment (eg, IGHD3-10, eg, IGHD3-10*01) and a IGHJ6 (eg, IGHJ6*02)

```
gene segment. Optionally additionally or alternatively the VL is encoded by a nucleotide sequence that is derived
from the recombination of a human IGKV2-28 (eg IGKV2-28*01) and IGKJ3 (eg, IGKJ3*01).
(215) Optionally, the VH is encoded by a nucleotide sequence that is derived from the recombination of a human
IGHV3-9 (eg, IGHV3-9*01) a D gene segment (eg, IGHD3-9, eg, IGHD3-9*01) and a IGHJ6 (eg, IGHJ6*02) gene
segment. Optionally additionally or alternatively the VL is encoded by a nucleotide sequence that is derived from
the recombination of a human IGKV2-29 (eg IGKV2-29*01 or IGKV2D-29*01) and IGKJ4 (eg, IGKJ4*01).
(216) In an example, the antibody or fragment comprises a HCDR3 length of 9-12 residues and/or the antibody or
fragment comprises a LCDR3 length of 7-9 residues. In an example, the antibody or fragment comprises a HCDR3
length of 9, 10, 11 or 12 residues, eg, 10, eg, 11. In an example, the antibody or fragment comprises a LCDR3 length
of 7, 8 or 9 residues, eg, 8, eg, 9. In an example, each VH domain of the antibody or fragment comprises from 1-11
non-germline residues, eg, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10 or 11 non-germline residues. In an example, each VL domain of
the antibody or fragment comprises from 3-8 non-germline residues, eg, 3, 4, 5, 6, 7 or 8 non-germline residues.
(217) Optionally, the antibody or fragment competes with CL-148219 (eg, CL-148219 in IgG format, eg, IgG4-PE)
for binding to PCSK9 (eg, human PCSK9) as determined by SPR.
(218) Optionally, the antibody or fragment competes with CL-148489 (eg, CL-148489 in IgG format, eg, IgG4-PE)
for binding to PCSK9 (eg, human PCSK9) as determined by SPR.
(219) Optionally, the amino acid substitutions are conservative amino acid substitutions, optionally wherein each
conservative substitution is from group (1) to (6): 1) Alanine (A), Serine (S), Threonine (T); 2) Aspartic acid (D),
Glutamic acid (E); 3) Asparagine (N), Glutamine (Q); 4) Arginine (R), Lysine (K); 5) Isoleucine (I), Leucine (L),
Methionine (M), Valine (V); and 6) Phenylalanine (F), Tyrosine (Y), Tryptophan (W).
(220) Any SPR herein is, for example, surface plasmon resonance (SPR) at 37° C. and pH 7.6.
(221) Optionally, any PCSK9 herein is (for example, in in vitro testing) human PCSK9.
(222) In an example, the antibody or fragment of the invention binds to human PCSK9 with a Ka of eg, 5×10.sup.6
M.sup.-1×s.sup.-1; or about 5×10.sup.6 M.sup.-1×s.sup.-1. In an example, the antibody or fragment of the
invention binds to human PCSK9 with a Kd of eg, 4 or 5 s.sup.-1; or about 4 or 5 s.sup.-1. In an example, the
antibody or fragment of the invention binds to human PCSK9 with a KD of eg. 0.07 or 0.14 nM; or about 0.07 or
0.14 nM. In an embodiment, the fragment is a Fab fragment. In an embodiment, the fragment is a scFv.
(223) As used herein, "inhibits", "inhibition", "inhibiting" and the like, as used herein refers to the ability of an
antagonist (e.g. an antibody or fragment thereof) to bind to an epitope (eg, of hPCSK9) which either partially or
completely prevents the binding of another antigen If the epitope to which the antagonist binds completely blocks
the binding site of the ligand, then ligand binding is completely prevented (which may be a physical blocking—in
the case of overlapping epitopes—or steric blocking—where the antagonist is large such that it prevents the ligand
binding to its distinct epitope), and the ligand is not removed from circulation. The concentration of circulating
ligand may therefore appear to be increased. If the epitope to which the antagonist binds partially blocks the binding
site of the ligand, the ligand may be able to bind, but only weakly (in the case of partial inhibition), or in a different
orientation to the natural binding interaction. In this case, some of the ligand may be removed from circulation, but
not as much as when the ligand binding site is completely free and available for binding. Inhibition thus refers to the
physical interaction of ligand and receptor. Inhibition can be measured by HTRF, which is described in more detail
elsewhere herein and in Mathis (1995) Clinical Chemistry 41(9), 1391-1397. Inhibition can also be measured by
flow cytometry, where receptor is expressed on cells, or by ELISA, where receptor is adsorbed onto plates.
(224) Optionally, the antibody of the invention has an affinity (KD) for binding PCSK9 of from 1 pM to 5 nM,
optionally wherein binding is determined by SPR using a Fab of said antibody at 37° C. at pH 7.6.
(225) Optionally, the antibody has off-rate (K.sub.off) for binding PCSK9 of from 1×10.sup. –5 to 1×10.sup.
-3S.sup.-1, optionally wherein binding is determined by SPR using a Fab of said antibody at 37° C. at pH 7.6.
(226) Optionally, the antibody has on-rate (K.sub.on) for binding PCSK9 of from 1×10.sup.5 to 1×10.sup.7 M.sup.
-1S.sup.-1, optionally wherein binding is determined by SPR using a Fab of said antibody at 37° C. at pH 7.6.
(227) In an example, the antibody (eg, as a Fab) or fragment has an affinity (KD) for binding PCSK9 (eg, human
PCSK9) of (a) from 2, 3, 4, 5 or 10 pM to 3, 4 or 5 nM; (b) from 1-10 pM to 5 nM; (c) from 10 pM to 3, 4 or 5 nM;
(d) from 50 or 80 pM to 200 nM; (e) from 50 or 80 pM to 150 nM; or (f) from 50 or 80 pM to 100 nM.
(228) In an example, the KD is (or is about) 5-15 pM (eg, 10 pM). In an example, the KD is (or is about) 2-5 nM
(eg, 3 nM). In an example, the KD is (or is about) 100-400 pM (eg, 140 or 390 pM).
(229) In an example, the antibody (eg, as a Fab) or fragment has an off-rate (K.sub.off) for binding PCSK9 (eg,
human PCSK9) of (a) from 1×10.sup.-5 to 5×10.sup.-4S.sup.-1; (b) from 1×10.sup.-5 to 6×10.sup.-4S.sup.-1; (c)
from 1×10.sup.-5 to 7×10.sup.-4 S.sup.-1; (d) from 1×10.sup.-5 to 8×10.sup.-4S.sup.-1; (e) from 2×10.sup.-5 to
1×10.sup.-3 S.sup.-1; (f) from 2×10.sup.-5 to 5×10.sup.-4 S.sup.-1; (g) from 2×10.sup.-5 to 6×10.sup.-4 S.sup.
-1; (h) from 2×10.sup.-5 to 7×10.sup.-4S.sup.-1; or (i) from 2×10.sup.-5 to 8×10.sup.-4S.sup.-1.
(230) In an example, the K.sub.off is (or is about) 5×10.sup.-4S.sup.-1 (eg, when the KD is (or is about) from 2 nM
to 400 pM; when the KD is (or is about) 2-5 nM (eg. 3 nM); or when the KD is (or is about) 100-400 pM (eg. 140 or
```

- 390 pM)). In an example, the K.sub.off is (or is about) 3×10.sup.-5 S.sup.-1 (eg, when the KD is (or is about) from 5-15 pM (eg, 10 pM)).
- (231) In an example, the antibody (eg, as a Fab) or fragment has an on-rate (K.sub.on) for binding PCSK9 (eg, human PCSK9) of (a) from $1\times10.\sup.5$ to $1\times10.\sup.6$ M.sup. -1S.sup. -1S.sup. -1S. -1S.sup. -1S.sup.
- -1S.sup.-1; (c) from 1×10.sup.5 to 3×10.sup.6 M.sup.-1S.sup.-1; (d) from 1×10.sup.5 to 4×10.sup.6 M.sup.
- -1S.sup.-1; (e) from 1×10.sup.5 to 5×10.sup.6 M.sup.-1S.sup.-1; (f) from 2×10.sup.5 to 5×10.sup.6 M.sup.
- -1S.sup.-1; (g) from 3×10.sup.5 to 5×10.sup.6 M.sup.-1S.sup.-1; (h) from 4×10.sup.5 to 5×10.sup.6 M.sup.
- -1S.sup.-1; (i) from 5×10.sup.5 to 5×10.sup.6 M.sup.-1S.sup.-1; or (j) from 6×10.sup.5 to 5×10.sup.6 M.sup. -1S.sup.-1.
- (232) In an example, the K.sub.on is (or is about) 1 or 2×10.sup.-5 M.sup.-1S.sup.-1 (eg, when the KD is 2-5 nM (eg, 3 nM)). In an example, the K.sub.on is (or is about) 1-4, 1, 2, 3 or 4×10 .sup.-6 M.sup.-1S.sup.-1 (eg, when the KD is (or is about) from 5-400 pM (eg, 140 or 390 pM) or 5-15 pM (eg, 10 pM)).
- (233) As provided in the Clauses or other aspects herein, an anti-PCSK9 antibody or fragment may bind to PCSK9, e.g. human PCSK9 with a K.sub.D of less than 50 nM, less than 40 nM, less than 30 nM as determined by surface plasmon resonance. Another embodiment, anti-PCSK9 antibody or fragment may bind to PCSK9, e.g. human PCSK9 with a K.sub.D of less than 20 nM, less than 15 nM, less than 10 nM as determined by surface plasmon resonance. The anti-PCSK9 antibody or fragment may bind to PCSK9, e.g. human PCSK9 with a K.sub.D of less than 8 nM, less than 5 nM, less than 4 nM, less than 3 nM, less than 2 nM or less than 1 nM as determined by surface plasmon resonance. The K.sub.D may be 0.9 nM or less, 0.8 nM or less, 0.7 nM or less, 0.6 nM or less, 0.5 nM or less, 0.4 nM or less, 0.3 nM or less, 0.2 nM or less, or 0.1 nM or less.
- (234) In another embodiment, the K.sub.D is within a range of 0.01 to 1 nM, or a range of 0.05 to 2 nM, or a range of 0.05 to 1 nM. The K.sub.D may be with regard to hPCSK9, cynomolugus monkey (ie, "cyno") PCSK9 and/or mouse PCSK9.
- (235) In another embodiment, the anti-PCSK9 antibodies described herein have a K.sub.ON rate (e.g. as measured by SPR, e.g. at 25° C. or at 37° C.) of approximately 0.5 to 10 μ M, for example approximately 1 to 8 μ M or approximately 1 to 7 uM. In another embodiment, the K.sub.ON rate is approximately 1 to 5 uM, e.g. approximately 1 μM, approximately 1.5 μM, approximately 2 μM, approximately 2.5 μM or approximately 3 μM. In another embodiment, the K.sub.ON rate is approximately 3.5 μM, approximately 4 μM, approximately 4.5 μM, approximately 5 μ M or approximately 5.5 μ M.
- (236) In another embodiment, the anti-PCSK9 antibodies described herein have a K.sub.OFF rate (e.g. as measured by SPR, e.g. at 25° C. or at 37° C.) of approximately 0.01 to 100 mM, for example approximately 0.1 to 50 mM or approximately 0.5 to 50 mM. In another embodiment, the K.sub.OFF rate is approximately 0.5 to 10 mM, or approximately 0.5 to 10 mM, e.g. approximately 1 mM, approximately 2 mM, approximately 3 mM, approximately 4 mM or approximately 5 mM. In another embodiment, the K.sub.OFF rate is approximately 0.6 mM, approximately 0.7 mM, approximately 0.8 mM or approximately 0.9 mM.
- (237) Optionally, the antibody of the invention comprises a human IgG4 constant region.
- (238) Preferably, an antibody or a fragment thereof that specifically binds to a hPCSK9 does not cross-react with other antigens (but may optionally cross-react with different PCSK9 species, e.g., rhesus, cynomolgus, or murine). An antibody or a fragment thereof that specifically binds to a PCSK9 antigen can be identified, for example, by immunoassays, BIAcore™, or other techniques known to those of skill in the art. An antibody or a fragment thereof binds specifically to a hPCSK9 antigen when it binds to a hPCSK9 antigen with higher affinity than to any crossreactive antigen as determined using experimental techniques, such as radioimmunoassays (RIA) and enzyme-linked immunosorbent assays (ELISAs). Typically, a specific or selective reaction will be at least twice background signal or noise and more typically more than 10 times background. See, e.g. Paul, ed., 1989, Fundamental Immunology Second Edition, Raven Press, New York at pages 332-336 for a discussion regarding antibody specificity. (239) Contact amino acid residues involved in the interaction of antibody and antigen, such as PCSK9, may be
- determined by various known methods to those skilled in the art.
- (240) In one embodiment, if the antibody recognises a linear epitope, short peptides based on the antigen sequence can be produced and binding of the antibody to these peptides can be assessed using standard techniques. (241) In one embodiment, limited proteolytic digestion and mass spectrophotometry can be used to identify binding
- epitopes.
- (242) In one embodiment, the contact residues of the epitope are identified by X-ray crystallography. In one embodiment, the contact residues of the epitope are identified by cryo-electro microscopy. In one embodiment, the contact residues of the epitope are identified by a combination of limited proteolytic digestion and mass spectrometry.
- (243) In another embodiment, the anti-PCSK9 antibodies (and fragments) described in herein provide improved transient expression levels over other anti-PCSK9 antibodies and fragments. Thus, in one embodiment, the anti-PCSK9 antibody (or fragment) is expressed in a HEK293 cell, e.g. a HEK293T cell, at an expression level of

approximately 100 μ g/mL, or in a range of approximately 100 to 350 μ g/mL. In another embodiment, the expression level is above approximately 350 μ g/mL.

- (244) In another embodiment, the anti-PCSK9 antibody (or fragment) is expressed in a CHO cell, e.g. an Expi-CHO cell, at an expression level of approximately 100 μ g/mL, or in a range of approximately 100 to 350 μ g/mL. In another embodiment, the expression level is above approximately 350 μ g/mL.
- (245) In another embodiment, the anti-PCSK9 antibody (or fragment) is expressed in a CHO cell, e.g. an Expi-CHO cell or a CHO-E7 EBNA cell, at an expression level of approximately 100 μ g/mL, or in a range of approximately 100 to 350 μ g/mL. In another embodiment, the expression level is above approximately 350 μ g/mL. The antibody for example, comprises the VH and VL domains of any one of CL-58838, formatted as a human IgG1 or human IgG4 (eg, IgG4-PE).
- (246) In any of these expression systems, the expression is carried out of a scale of between approximately 0.5 mL and 3 mL, for example between approximately 0.5 mL and 2 mL. In any of these expression systems, the anti-PCSK9 antibody (or fragment) may be expressed from a pTT5 vector. In any of these expression systems, the anti-PCSK9 antibody (or fragment) may be expressed in conjunction with a lipid transfection reagent, and may optionally be expressed in a CHO cell, e.g. an Expi-CHO cell. In any of these expression systems, the anti-PCSK9 antibody (or fragment) may be expressed in conjunction with a PEI transfection reagent, and may optionally be expressed in a CHO cell, e.g. an CHO-E7 EBNA cell. In any of these expression systems, the anti-PCSK9 antibody (or fragment) may be expressed in conjunction with a helper plasmid (e.g. an AKT helper plasmid), and may optionally be expressed in a CHO cell, e.g. an CHO-E7 EBNA cell.
- (247) In any of these expression systems, the expression level is between approximately 100 μ g/mL and approximately 1500 μ g/mL, for example between approximately 1000 μ g/mL and approximately 1000 μ g/mL and approximately 200 μ g/mL and approximately 1000 μ g/mL. In any of these expression systems, the lower limit of expression may be approximately 100 μ g/mL, approximately 200 μ g/mL, approximately 300 μ g/mL, or approximately 400 μ g/mL. In another embodiment, the lower limit of expression may be approximately 500 μ g/mL, approximately 600 μ g/mL, approximately 700 μ g/mL, or approximately 800 μ g/mL. In any of these expression systems, the upper limit of expression may be approximately 2000 μ g/mL, approximately 1800 μ g/mL, approximately 1600 μ g/mL, or approximately 1500 μ g/mL. In another embodiment, the upper limit of expression may be approximately 1250 μ g/mL, approximately 1000 μ g/mL, approximately 900 μ g/mL, or approximately 800 μ g/mL.
- (248) In another embodiment, the expression system is a Lonza expression system, e.g. Lonza X-Ceed® system. In the Lonza expression system, the expression may be carried out at a scale of approximately 30 mL to 2 L, for example 50 mL to 1 L, or 1 L to 2 L. In the Lonza expression system, the anti-PCSK9 antibody (or fragment) may be expressed in conjunction with electroporation, and optionally without any helper plasmids. In the Lonza expression system, the anti-PCSK9 antibody (or fragment) may be expressed at a level of approximately 1 g/L, or approximately 900 mg/L, or approximately 800 mg/L, or approximately 700 mg/L. In another embodiment, In the Lonza expression system, the anti-PCSK9 antibody (or fragment) may be expressed at a level of approximately 600 mg/L or approximately 500 mg/L or approximately 400 mg/L. In the Lonza expression system, the anti-PCSK9 antibody (or fragment) may be expressed at a level of between approximately 400 mg/L and approximately 2 g/L, for example between approximately 500 mg/L and approximately 1.5 g/L, or between approximately 500 mg/L and approximately 1 g/L. In another embodiment, the expression level is above 1 g/L. In another embodiment, the anti-PCSK9 antibodies provide improved half-life over other anti-PCSK9 antibodies.
- (249) In one embodiment, the antibody or fragment is a human antibody or fragment. In one embodiment, the antibody or fragment is a fully human antibody or fragment. In one embodiment, the antibody or fragment is a fully human monoclonal antibody or fragment.
- (250) in one embodiment, the antibody or fragment is a humanised antibody or fragment. In one embodiment, the antibody or fragment is a humanised monoclonal antibody or fragment.
- (251) Contact amino acid residues involved in the interaction of antibody and antigen may be determined by various known methods to those skilled in the art, such as alanine scanning, protein crystallography, mass spectrophotometry or any other technique as will be apparent to the skilled addressee.
- (252) In one embodiment, the recited CDR comprises one amino acid substitution, which may be a conservative amino acid substitution. In one embodiment, the recited CDR comprises two amino acid substitutions, which may be conservative amino acid substitutions. In one embodiment, the recited CDR comprises three amino acid substitutions, which may be conservative amino acid substitutions. In one embodiment, the recited CDR comprises four amino acid substitutions, which may be conservative amino acid substitutions. In one embodiment, the recited CDR comprises five amino acid substitutions, which may be conservative amino acid substitutions. In one embodiment, the recited CDR comprises six amino acid substitutions, which may be conservative amino acid substitutions.
- (253) Amino acid substitutions include alterations in which an amino acid is replaced with a different naturally-

occurring amino acid residue. Such substitutions may be classified as "conservative", in which case an amino acid residue contained in a polypeptide is replaced with another naturally occurring amino acid of similar character either in relation to polarity, side chain functionality or size. Such conservative substitutions are well known in the art. Substitutions encompassed by the present invention may also be "non-conservative", in which an amino acid residue which is present in a peptide is substituted with an amino acid having different properties, such as naturally-occurring amino acid from a different group (e.g. substituting a charged or hydrophobic amino; acid with alanine), or alternatively, in which a naturally-occurring amino acid is substituted with a non-conventional amino acid. (254) In one embodiment, the conservative amino acid substitutions are as described herein. For example, the substitution may be of Y with F, T with S or K, P with A, E with D or Q, N with D or G, R with K, G with N or A, T with S or K, D with N or E, I with L or V, F with Y, S with T or A, R with K, G with N or A, K with R, A with S, K or P. In another embodiment, the conservative amino acid substitutions may be wherein Y is substituted with F, T with A or S, I with L or V, W with Y, M with L, N with D, G with A, T with A or S, D with N, I with L or V, F with Y or L, S with A or T and A with S, G, T or V.

(255) In an embodiment, the present invention provides a pharmaceutical composition comprising an anti-PCSK9 antagonist (eg, an antibody, or PCSK9-binding fragment thereof) of the present invention, and an acceptable carrier, diluent, or excipient. More particularly, the compositions of the present invention further comprise one or more additional therapeutic agents, eg, a statin. As referred to herein, a "statin" (also known as HMG-CoA reductase inhibitors) are inhibitors of the enzyme HMG-coA reductase, which mediates cholesterol production in the liver. Statins, by competitively binding HMG-CoA reductase, prevent the binding of HMG-CoA to the enzyme and thereby inhibit the activity of the reductase (e.g. the production of mevalonate). Non-limiting examples of statins can include atorvastatin (LIPITORTM), fluvastatin (LESCOLTM), lovastatin (MEVACORTM, ALTOCORTM), pitavastatin (LIVALOTM), pravastatin (PRAVACHOLTM), rosuvastatin (CRESTORTM), and simvastatin (ZOCORTM). (256) Statins can be administered in combination with other agents, e.g. the combination of ezetimibe and simvastatin.

(257) In an example, the anti-PCSK9 antagonist, antibody or fragment binds to PCSK9 with a KD of less than about 1×10.sup.–8 M, preferably, less than about 1×10.sup.–9 M as determined by common methods known in the art, eg, by use of a surface plasmon resonance (SPR) biosensor at 37° C.

(258) "Effective amount" means the amount of an antagonist (eg, antibody) of the present invention or pharmaceutical composition of the present invention that will elicit the biological or medical response or desired therapeutic effect on a subject, mammal or human that is being sought by the researcher, medical doctor, or other clinician. An effective amount may vary according to factors such as the disease state, age, sex, and weight of the individual, and the ability of the antibody and/or statin to elicit a desired response in the individual. An effective amount is also one in which any toxic or detrimental effect is outweighed by the therapeutically beneficial effects. (259) An anti-PCSK9 antagonist antibody, or antigen-binding fragment thereof, of the present invention, combination or pharmaceutical composition comprising the same, may be administered by parenteral routes (eg, subcutaneous, intravenous, intraperitoneal, intramuscular, or transdermal). Administration may be to a subject alone or in combination with a pharmaceutically acceptable carrier and/or diluent in single or multiple doses. Pharmaceutical compositions, combinations or antagonists of the present invention can be prepared by methods well known in the art (e.g., Remington: *The Science and Practice of Pharmacy*, 19.sup.th ed. (1995), A. Gennaro et al., Mack Publishing Co.) and may comprise or be combined with one or more pharmaceutically acceptable carriers, diluents, or excipients.

(260) In an embodiment, the subject is a human male, eg, an adult or infant. In an embodiment, the subject is a human female, eg, an adult or infant, eg, a non-pregnant female or pregnant female. I an example, the human is a dialysis patient. The infant may be a human that is >1 month old. In an example, the subject is an adult human with established cardiovascular disease or with primary hyperlipidemia (eg, heterozygous familial hypercholesterolemia [HeFH]). In an example, the subject is an human with HoFH, eg, an adult human.

(261) A subject can be one who has been previously diagnosed with or identified as suffering from or having a condition in need of treatment or one or more complications related to such a condition, and optionally, have already undergone treatment for the condition or the one or more complications related to the condition. Alternatively, a subject can also be one who has not been previously diagnosed as having the condition or one or more complications related to the condition. For example, a subject can be one who exhibits one or more risk factors for the condition or one or more complications related to the condition or a subject who does not exhibit risk factors. (262) The invention may comprise simultaneously or sequentially administering the anti-PCSK9 antagonist and statin. In an example, antagonist and statin are administered no more than 1 month, 4 weeks, 3 weeks, 2 weeks, 1 week, 10 days, 9 days, 8 days, 7 days, 6 days, 5 days, 4 days, 3 days, 2 days or 1 day apart. As exemplified herein, administration of the antagonist and statin can be effective if no more than 7 days (eg, no more than one day) apart. In an example, the anti-PCSK9 antagonist and statin are administered to the subject no more than 10, 14, 21 or 28 days apart.

- (263) In an example, the statin is administered 2, 3 or 4 times weekly. In another example, the statin is administered 1, 2, 3 or 4 times monthly or in a 8 week period.
- (264) In an example, the statin and/or antibody or fragment is administered to the subject intraveneously or subcutaneously.
- (265) In an example, the hyperlipidaemia or hypercholesterolaemia is in a subject receiving or having received statin treatment, eg, a patient that is a low or non-responder to statin treatment.
- (266) In an embodiment, the present invention provides the use of an anti-PCSK9 antagonist and a statin for the manufacture of a medicament. In a further embodiment, the present invention provides the use of an anti-PCSK9 antagonist and a statin for the manufacture of a medicament for the treatment or prevention of hyperlipidaemia or hypercholesterolaemia, eg, moderate to severe hyperlipidaemia or hypercholesterolaemia, eg, in a subject that is a low or no-responder to statin treatment.
- (267) In an example the anti-PCSK9 antibody of the invention is an antibody that competes with a reference antibody in an HTRF assay. For example, wherein in the HTRF assay the antibody of the invention is a labelled antibody that is pre-incubated with human PCSK9 and subsequently combined with unlabelled reference antibody (according to part I or II), wherein competition between the antibodies is detected by the assay. In an example, the assay uses AlexaFluorTM 647 labelled antibody of the invention. In an alternative, the human PCSK9 is labelled (eg, with AlexaFluorTM 647, the test antibody is labelled with biotin for binding to Eu3+cryptate-streptavidin, and the reference antibody is unlabelled). Example reference antibodies are preferably CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT). Alternatives are alirocumab or evolocumab.
- (268) Optionally, the anti-PCSK9 antibody of the invention (test antibody) competes in an HTRF assay with the reference antibody for binding human PCSK9 (or binds the same epitope of human PCSK9 as the reference antibody), wherein the assay uses a directly or indirectly labelled test antibody directly or indirectly labelled with a donor (such as for example Eu3+cryptate) or an acceptor fluorophore (such as for example AlexaFluor™ 647) and a target PCSK9 labelled with either a donor or acceptor fluorophore to enable energy transfer between donor and acceptor, whereby a fluorescence signal is produced and detected. In an example, where AlexaFluor™ 647 labelling is used, competition is detected by a reduction in fluorescence signal at 665 nM of at least 20% when the test antibody is in the presence of the reference antibody versus signal without the reference antibody. Optionally, the reduction in signal at 665 nM is at least 20, 30, 40, 50, 60, 70, 80 or 90%.
- (269) In an example, the antibody or fragment is for reducing the total dose of administered over a 4 week period to a human or animal subject for treating or preventing hyperlipidaemia or hypercholesterolaemia or any other disease or condition disclosed herein.
- (270) In an example, the antibody or fragment is for reducing to ½ to ½ the total dose required in a control subject receiving identical treatment over the 4 week period except for administration of statin without administration of an anti-PCSK9 antagonist (eg, antibody or fragment) to a human or animal subject for treating or preventing hyperlipidaemia or hypercholesterolaemia or any other disease or condition disclosed herein.
- (271) In an example, the antibody or fragment is for sparing by ½ to ½ the administration of statin administered to a human or animal subject over a treatment period, eg, a 4 week period, for treating or preventing hyperlipidaemia or hypercholesterolaemia osteoporosis or any other disease or condition disclosed herein.
- (272) Optionally, the antibody is an IgG4 antibody.
- (273) Optionally, the dose of statin administered to the subject is not effective when administered in the absence of the anti-PCSK9 antagonist (eg, antibody or fragment of the invention).
- (274) In an example, the subject herein is refractory to a dose of statin, but is responsive for treatment of hyperlipidaemia or hypercholesterolaemia or another disease or condition when administered the antibody or fragment of the invention and the statin dose.
- (275) Thus, in an example, the antibody or fragment of the invention is for administration in combination with a dose of a statin to a human or animal subject for treating a PCSK9-related disease or condition in the subject, wherein the subject is treated with the combination, but is not treatable for the disease or condition by administration of the dose of statin in the absence of administration of said antibody or fragment.
- (276) The disease or condition may be hyperlipidaemia or hypercholesterolaemia or any other disease or condition disclosed herein, for example selected from a lipid disorder, hyperlipoproteinemia, hyperlipidemia; hypercholesterolemia, a heart attack, a stroke, coronary heart disease, atherosclerosis, peripheral vascular disease, claudication, type II diabetes, high blood pressure, and a cardiovascular disease or condition.
- (277) The invention provides a antibody or fragment which specifically binds to PCSK9. Optionally, the antibody or fragment comprises a CDRH1 sequence as disclosed herein or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto. Optionally, the antibody or fragment comprises a CDRH2 sequence as disclosed herein or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto. Optionally, the antibody or fragment comprises a CDRH3 sequence as disclosed herein or a sequence that is at least 70, 80, 85, 90, 95, 96, 97,

98 or 99% identical thereto. Optionally, the antibody or fragment comprises a CDRL1 sequence as disclosed herein or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto. Optionally, the antibody or fragment comprises a CDRL2 sequence as disclosed herein or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto. Optionally, the antibody or fragment comprises a CDRL3 sequence as disclosed herein or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto.

(278) Optionally, the antibody or fragment comprises the CDRH1 sequence of CL-148219 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto; the CDRH2 sequence of CL-148219 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto; and the CDRH3 sequence of CL-148219 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto. Optionally, the antibody or fragment comprises the CDRL1 sequence of CL-148219 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto; the CDRL2 sequence of CL-148219 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto; and the CDRL3 sequence of CL-148219 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto.

(279) Optionally, the antibody or fragment comprises the CDRH1 sequence of CL-148489 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto; the CDRH2 sequence of CL-148489 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto; and the CDRH3 sequence of CL-148489 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto. Optionally, the antibody or fragment comprises the CDRL1 sequence of CL-148489 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto; the CDRL2 sequence of CL-148489 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto; and the CDRL3 sequence of CL-148489 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto.

(280) Optionally, the antibody or fragment comprises the VH FR1 sequence of CL-148219 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto; the VH FR2 sequence of CL-148219 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto; the VH FR3 sequence of CL-148219 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto; and the VH FR4 sequence of CL-148219 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto. Additionally or alternatively, optionally the antibody or fragment comprises the VL FR1 sequence of CL-148219 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto; the VL FR2 sequence of CL-148219 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto; and the VL FR4 sequence of CL-148219 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto; and the VL FR4 sequence of CL-148219 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto; and the VL FR4 sequence of CL-148219 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto.

(281) Optionally, the antibody or fragment comprises the VH FR1 sequence of CL-148489 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto; the VH FR2 sequence of CL-148489 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto; the VH FR3 sequence of CL-148489 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto; and the VH FR4 sequence of CL-148489 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto. Additionally or alternatively, optionally the antibody or fragment comprises the VL FR1 sequence of CL-148489 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto; the VL FR2 sequence of CL-148489 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto; the VL FR3 sequence of CL-148489 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto; and the VL FR4 sequence of CL-148489 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto; and the VL FR4 sequence of CL-148489 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto.

(282) In an example, each CDR and FR1-4 herein is according to IMGT nomenclature. In an alternative, each CDR and FR1-4 herein is according to Kabat nomenclature.

- (283) Optionally, the antibody or fragment comprises a human gamma-1 heavy chain constant region, eg, comprising a gamma-1 heavy chain constant region sequence disclosed herein or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto. For example, said C region comprises SEQ ID NO: 126 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto.
- (284) Optionally, the antibody or fragment comprises a human gamma-2 heavy chain constant region, eg, comprising a sequence gamma-2 heavy chain constant region sequence disclosed herein or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto. For example, said C region comprises SEQ ID NO: 136 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto.
- (285) Optionally, the antibody or fragment comprises a human gamma-3 heavy chain constant region.
- (286) Optionally, the antibody or fragment comprises a human gamma-4 heavy chain constant region, eg, comprising a sequence gamma-4 heavy chain constant region sequence disclosed herein or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto. For example, said C region comprises SEQ ID NO: 3, 144, 152 or 154 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto.
- (287) Optionally, the antibody or fragment comprises a human kappa light chain constant region, eg, comprising a kappa light chain constant region sequence disclosed herein or a sequence that is at least 70, 80, 85, 90, 95, 96, 97,

- 98 or 99% identical thereto. For example, said C region comprises SEQ ID NO: 156 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto. In an embodiment, the antibody or fragment comprises a human gamma-4 heavy chain constant region, eg, a IgG4-PE constant region.
- (288) Optionally, the antibody or fragment comprises a human lambda light chain constant region, eg, comprising a lambda light chain constant region sequence disclosed herein or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto. For example, said C region comprises SEQ ID NO: 166 or a sequence that is at least 70, 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto.
- (289) In an embodiment, the antibody or fragment comprises a human gamma-4 heavy chain constant region, eg, a IgG4-PE constant region.
- (290) Target binding ability, specificity and affinity (Kd, K.sub.off and/or K.sub.on) can be determined by any routine method in the art, eg, by surface plasmon resonance (SPR). The term "Kd", as used herein, is intended to refer to the equilibrium dissociation constant of a particular antibody-antigen interaction.
- (291) In one embodiment, the surface plasmon resonance (SPR) is carried out at 25° C. In another embodiment, the SPR is carried out at 37° C.
- (292) In one embodiment, the SPR is carried out at physiological pH, such as about pH7 or at pH7.6 (eg, using Hepes buffered saline at pH7.6 (also referred to as HBS-EP)).
- (293) In one embodiment, the SPR is carried out at a physiological salt level, eg, 150 mM NaCl.
- (294) In one embodiment, the SPR is carried out at a detergent level of no greater than 0.05% by volume, eg, in the presence of P20 (polysorbate 20; eg, Tween-20TM) at 0.05% and EDTA at 3 mM.
- (295) In one example, the SPR is carried out at 25° C. or 37° C. in a buffer at pH7.6, 150 mM NaCl, 0.05% detergent (eg, P20) and 3 mM EDTA. The buffer can contain 10 mM Hepes. In one example, the SPR is carried out at 25° C. or 37° C. in HBS-EP. HBS-EP is available from Teknova Inc (California; catalogue number H8022). (296) In an example, the affinity of the ligand (eg, antibody) is determined using SPR by 1. Coupling anti-mouse (or other relevant human, rat or non-human vertebrate antibody constant region species-matched) IgG (eg, Biacore™ BR-1008-38) to a biosensor chip (eg, GLM chip) such as by primary amine coupling; 2. Exposing the anti-mouse IgG (or other matched species antibody) to a test IgG antibody to capture test antibody on the chip: 3. Passing the test antigen over the chip's capture surface at 1024 nM, 256 nM, 64 nM, 16 nM, 4 nM with a OnM (i.e. buffer alone); and 4. And determining the affinity of binding of test antibody to test antigen using surface plasmon resonance, eg, under an SPR condition discussed above (eg, at 25° C. in physiological buffer). SPR can be carried out using any standard SPR apparatus, such as by Biacore™ or using the ProteOn XPR36™ (Bio-Rad®). (297) Regeneration of the capture surface can be carried out with 10 mM glycine at pH1.7. This removes the captured antibody and allows the surface to be used for another interaction. The binding data can be fitted to 1:1 model inherent using standard techniques, eg, using a model inherent to the ProteOn XPR36TM analysis software. (298) In an example, the antagonist (eg, antibody or fragment) of the invention is contained in a medical container, eg, a vial, syringe, IV container or an injection device (eg, an intraocular or intravitreal injection device). In an example, the antagonist is in vitro, eg, in a sterile container. In an example, the invention provides a kit comprising the antagonist of the invention, packaging and instructions for use in treating or preventing or diagnosing in a human a disease or condition mediated by PCSK9.
- (299) In an example, the instructions indicate that the human should be genotyped for a PCSK9 variant sequence of the invention before administering the antagonist to the human. In an example, the instructions indicate that the human should be phenotyped for a PCSK9 variant of the invention before administering the ligand to the human. In an example, the human is of Chinese (eg, Han) ethnicity and the instructions are in Chinese (eg, Mandarin). (300) In an example, the antagonist is (or has been determined as) a neutraliser of PCSK9. In an example, determination is carried out in a human (eg, in a clinical trial). In an example, determination is carried out in a non-human, eg, in a mouse, rat, rabbit, pig, dog, sheep or non-human primate (eg, Cynomolgous monkey, rhesus monkey or baboon).
- (301) Variants of PCSK9 can include the forms described in WO2015092393 as a, f, c, r, p, m, e h, aj, and q. Sequences of these variants are provided therein, see, e.g, SEQ ID NOs:1-27 and in Table 1, 2 or 6. The disclosure of this reference and these sequences are incorporated herein by reference for possible use in the present invention. In an example, the antagonist specifically binds to a PCSK9 variant disclosed in WO2015092393. In an example, additionally or alternatively the human subject expresses such a PCSK9 variant.
- (302) In an example, the antagonist specifically binds to PCSK9 variants comprising a E670G amino acid. In an example, additionally or alternatively the human subject expresses such a PCSK9 variant.
- (303) In an example, the antagonist specifically binds to PCSK9 variants comprising a 1474V amino acid. In an example, additionally or alternatively the human subject expresses such a PCSK9 variant.
- (304) Antagonists of the invention are useful, for instance, in specific binding assays, for genotyping or phenotyping humans, affinity purification of the PCSK9 and in screening assays to identify other antagonists of PCSK9 activity. Some of the antagonists of the invention are useful for inhibiting binding of PCSK9 to a cognate human receptor or

protein, or inhibiting PCSK9-mediated activities.

(305) The invention encompasses anti-PCSK9 (eg, PCSK9) antibody antagonists having a modified glycosylation pattern. In some applications, modification to remove undesirable glycosylation sites may be useful, or e.g., removal of a fucose moiety to increase antibody dependent cellular cytotoxicity (ADCC) function (see Shield et al. (2002) JBC 277:26733). In other applications, modification of galactosylation can be made in order to modify complement dependent cytotoxicity (CDC).

- (306) In an example, the invention features a pharmaceutical composition comprising a antagonist of the invention, wherein the antagonist is or comprises a recombinant human antibody or fragment thereof which specifically binds the PCSK9 (eg, a rare variant as described herein) and a pharmaceutically acceptable carrier. In one embodiment, the invention features a composition which is a combination of an antibody antagonist or antigen-binding fragment of an antibody of the invention, and a second therapeutic agent. The second therapeutic agent may be any of an anti-inflammatory agent, an anti-angiogenesis agent, a painkiller, a diuretic, a chemotherapeutic agent, an anti-neoplastic agent, a vasodilator, a vasoconstrictor, a statin, a beta blocker, a nutrient, an adjuvant, an anti-obesity agent and an anti-diabetes agent.
- (307) In an example, the invention features a method for inhibiting PCSK9 activity using the anti-PCSK9 antagonist of the invention (eg, an antibody or antigen-binding portion of the antibody of the invention), wherein the therapeutic method comprises administering a therapeutically effective amount of a pharmaceutical composition comprising the antagonist. The disorder treated is any disease or condition which is improved, ameliorated, inhibited or prevented by removal, inhibition or reduction of PCSK9 activity. (308) Paragraphs:
- (309) The invention provides the following aspects set out in Paragraphs 1 et seq.
- (310) 1. An antibody or fragment comprising a binding site which specifically binds to Proprotein Convertase Subtilisin Kexin type 9 (PCSK9), wherein the binding site comprises a VH domain that is encoded by a nucleotide sequence that is derived from the recombination of a human VH gene segment, DH gene segment and JH gene segment, wherein the VH gene segment is selected from IGHV4-31, IGHV4-59, IGHV4-4 and IGHV3-9.
- (311) 2. The antibody or fragment according to Paragraph 1, wherein (i) the VH gene segment is IGHV4-31 and the DH gene segment is human gene segment IGHD3-9; or (ii) the VH gene segment is IGHV4-59 and the DH gene segment is human gene segment IGHD3-10; or (iii) the VH gene segment is IGHV4-4 and the DH gene segment is human gene segment IGHD2-15; or (iv) the VH gene segment is IGHV3-9 and the DH gene segment is human gene segment IGHD3-9.
- (312) 3. The antibody or fragment according to Paragraph 1 or 2, wherein the JH gene segment is human gene segment IGHJ6.
- (313) 4. The antibody or fragment according to any preceding Paragraph, wherein the binding site comprises a CDRH3 sequence selected from SEQ ID NO: 271, 285, 15, 77, 29, 91, 211 and 225.
- (314) 5. The antibody or fragment according to any preceding Paragraph, wherein the binding site comprises a VH domain comprising SEQ ID NO: 259, 1, 65 or 199.
- (315) 6. The antibody or fragment according to any preceding Paragraph, wherein the binding site comprises a VH domain comprising SEQ ID NO: 259 paired with a VL domain comprising SEQ ID NO: 289.
- (316) 7. The antibody or fragment according to any one of Paragraphs 1 to 5, wherein the binding site comprises a VH domain comprising SEQ ID NO: 319 paired with a VL domain comprising SEQ ID NO: 349.
- (317) 8. An antibody or fragment which specifically binds to PCSK9 and comprises the CDRH3 sequence of an anti-PCSK9 antibody according to any preceding Paragraph, or said CDRH3 sequence comprising 3, 2 or 1 amino acid substitution(s).
- (318) 9. An antibody or fragment (optionally according to any preceding Paragraph) which specifically binds to PCSK9 and comprises a VH domain which comprises a CDRH3 sequence of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or said sequence comprising 3, 2 or 1 amino acid substitution(s).
- (319) 10. The antibody or fragment according to Paragraph 9, wherein the VH domain comprises (i) a CDRH3 sequence of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or said CDRH3 sequence comprising 3, 2 or 1 amino acid substitution(s); and (ii) a CDRH1 sequence of said selected antibody; or said CDRH1 sequence comprising 3, 2 or 1 amino acid substitution(s).
- (320) 11. The antibody or fragment according to Paragraph 9 or 10, wherein the VH domain comprises (iii) a CDRH3 sequence of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or said CDRH3 sequence comprising 3, 2 or 1 amino acid substitution(s); and (iv) a CDRH2 sequence of said selected antibody; or said CDRH2 sequence comprising 3, 2 or 1 amino acid substitution(s).
- (321) 12. An antibody or fragment (optionally according to any preceding Paragraph) comprising a binding site

- which specifically binds to PCSK9, wherein the binding site comprises a VH domain that comprises the amino acid sequence of a VH domain of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or an amino acid that is at least 70% identical thereto.
- (322) 13. The antibody or fragment according to Paragraph 9, 10, 11 or 12, wherein the selected antibody is CL-274711 or CL-148219QLT.
- (323) 14. The antibody or fragment according to any preceding Paragraph comprising first and second copies of said VH domain.
- (324) 15. An antibody or fragment (optionally according to any preceding Paragraph) comprising a binding site which specifically binds to PCSK9, wherein the binding site comprises a VL domain that is encoded by a nucleotide sequence that is derived from the recombination of a human VL gene segment and JL gene segment, wherein the VL gene segment is selected from IGKV3-11, IGKV2-28 and IGKV2-29.
- (325) 16. The antibody or fragment according to Paragraph 15, wherein the JL gene segment is a human gene segment selected from IGKJ4, IGJK3 and IGKJ1.
- (326) 17. An antibody or fragment which specifically binds to PCSK9 and comprises the CDRL3 sequence of an anti-PCSK9 antibody according to any preceding Paragraph, or said CDRL3 sequence comprising 3, 2 or 1 amino acid substitution(s).
- (327) 18. An antibody or fragment (optionally according to any preceding Paragraph) which specifically binds to PCSK9 and comprises a VL domain which comprises a CDRL3 sequence selected from SEQ ID NO: 301, 315, 47, 107, 61, 121, 241 and 255, or said selected CDRL3 sequence comprising 3, 2 or 1 amino acid substitution(s). (328) 19. An antibody or fragment (optionally according to any preceding Paragraph) which specifically binds to PCSK9 and comprises a VL domain which comprises a CDRL3 (and optionally a CDRH3) sequence of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or said sequence(s) each comprising 3, 2 or 1 amino acid substitution(s).
- (329) 20. The antibody or fragment according to Paragraph 19, wherein the VL domain comprises (i) a CDRL3 sequence (and optionally a CDRH3) of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or said CDR3 sequence(s) each comprising 3, 2 or 1 amino acid substitution(s); and (ii) a CDRL1 (and optionally a CDRH1) sequence of said selected antibody; or said CDR1 sequence(s) each comprising 3, 2 or 1 amino acid substitution(s).
- (330) 21. The antibody or fragment according to Paragraph 19 or 20, wherein the VL domain comprises (iii) a CDRL3 (and optionally a CDRH3) sequence of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or said CDR3 sequence(s) each comprising 3, 2 or 1 amino acid substitution(s); and (iv) a CDRL2 (and optionally a CDRH2) sequence of said selected antibody; or said CDR2 sequence(s) each comprising 3, 2 or 1 amino acid substitution(s).
- (331) 22. An antibody or fragment (optionally according to any preceding Paragraph) comprising a binding site which specifically binds to PCSK9, wherein the binding site comprises a VL domain that comprises the amino acid sequence of a VL domain of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or an amino acid that is at least 70% identical thereto.
- (332) 23. An antibody or fragment (optionally according to any preceding Paragraph) which specifically binds to PCSK9 and comprises the heavy chain amino acid sequence of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or an amino acid that is at least 70% identical thereto.
- (333) 24. An antibody or fragment (optionally according to any preceding Paragraph) which specifically binds to PCSK9 and comprises the light chain amino acid sequence of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or an amino acid that is at least 70% identical thereto.
- (334) 25. The antibody or fragment of Paragraph 23, comprising the light chain amino acid sequence of said selected antibody; or an amino acid that is at least 70% identical thereto.
- (335) 26. An antibody or fragment (optionally according to any preceding Paragraph) which specifically binds to a human PCSK9 epitope that is identical to an epitope to which the antibody of any preceding Paragraph binds. (336) 27. The antibody or fragment according to Paragraph 26, wherein the epitope is identified by unrelated amino acid scanning, or by X-ray crystallography.
- (337) 28. The antibody or fragment according to Paragraph 27, wherein the contact residues of the epitope are defined by a reduction in affinity of at least 10-fold in an unrelated amino acid scan, e.g. an alanine scan as determined by SPR.
- (338) 29. An antibody or fragment (optionally according to any preceding Paragraph) which competes for binding to human PCSK9 with the antibody of any preceding Paragraph.
- (339) 30. The antibody or fragment according to any preceding Paragraph which specifically binds to a PCSK9 comprising an amino acid sequence selected from SEQ ID NOs: 189-194.

- (340) 31. The antibody or fragment according to any preceding Paragraph, wherein the antibody or fragment comprises a human constant region, optionally an IgG4 constant region or an IgG1 constant region.
- (341) 32. The antibody or fragment according to Paragraph 31, wherein the constant region is an IgG4-PE constant region, optionally the constant region comprises the amino acid sequence of SEQ ID NO: 3 or 152.
- (342) 33. The antibody or fragment according to any preceding Paragraph further comprising an antigen-binding site that specifically binds another target antigen, optionally ANGPTL3.
- (343) 34. An anti-PCSK9 antibody or fragment as defined in any preceding Paragraph for treating or preventing a PCSK9-mediated disease or condition (optionally hypercholesterolaemia) in a subject.
- (344) 35. The anti-PCSK9 antibody or fragment of Paragraph 34, wherein the disease or condition is selected from hypercholesterolemia, hypercholesterolemia, dyslipidemia, cholestatic liver disease, nephrotic syndrome, hypothyroidism, obesity, diabetes, atherosclerosis or a cardiovascular disease.
- (345) 36. The antibody or fragment of Paragraph 34 or 35, wherein the antibody or fragment is administered to the subject simultaneously or sequentially with a statin.
- (346) 37. A combination of an amount of an anti-PCSK9 antibody or fragment and an amount of a statin (optionally comprising multiple doses of said antibody and/or statin), wherein the antibody or fragment is according to any one of Paragraphs 1 to 36.
- (347) 38. The antibody, fragment or combination according to any one of Paragraphs 1 to 37 for use in a method of treating or preventing hypercholesterolaemia in a subject that has previously been on a statin treatment regime at a first dose, wherein the method comprises reducing the dose of statin that is administered to the subject or administering no statin to the subject, wherein the method comprises administering the antibody or fragment to the subject.
- (348) 39. The combination of Paragraph 37 or 38, wherein comprising statin at a daily dose of 10 to 20 mg (eg, 10 mg); or <60 mg (eg, 40 mg).
- (349) 40. The antibody, fragment or combination of any preceding Paragraph for administering to a human or animal subject suffering from elevated cholesterol, for lowering plasma low density lipoprotein cholesterol (LDL-C) level in the subject after the subject has received the anti-PCSK9 antibody or fragment.
- (350) 41. Use of the antibody, fragment or combination as defined in any preceding Paragraph in the manufacture of a medicament for administration to a subject for treating or preventing a PCSK9-mediated disease or condition, optionally hypercholesterolaemia.
- (351) 42. A method of treating or preventing a PCSK9-mediated disease or condition in a subject (optionally hypercholesterolaemia), the method comprising administering to said subject a therapeutically effective amount of an antibody, fragment or combination as defined in any one of Paragraphs 1 to 40, wherein the PCSK9-mediated disease or condition is thereby treated or prevented.
- (352) 43. The use according to Paragraph 41 or the method according to Paragraph 42, wherein the PCSK9-mediated disease or condition is hypercholesterolaemia.
- (353) 44. The antibody, fragment, combination, use or the method according to any one of Paragraphs 34 to 43, further comprising administering to the subject a further therapy, for example a further therapeutic agent, optionally wherein the further therapeutic agent is selected from the group consisting of a: a. Statin; b. An ANGPTL3 inhibitor (eg, an anti-ANGPTL3 antibody, eg, evinacumab) c. Fibrate; d. Bile acid sequestrant; e. Nicotinic acid; and f. Niacin.
- (354) 45. A pharmaceutical composition comprising an antibody, fragment or combination as defined in any one of Paragraphs 1 to 40 and 44 and a pharmaceutically acceptable excipient, diluent or carrier and optionally in combination with a further therapeutic agent selected from an agent recited in Paragraph 44.
- (355) 46. The pharmaceutical composition according to Paragraph 45 for treating and/or preventing a PCSK9-mediated condition or disease, optionally hypercholesterolaemia.
- (356) 47. The pharmaceutical composition according to Paragraph 45 or 46 in combination with a label or instructions for use to treat and/or prevent said disease or condition in a human; optionally wherein the label or instructions comprise a marketing authorisation number (optionally an FDA or EMA authorisation number); optionally wherein the kit comprises an IV or injection device that comprises the antibody or fragment.
- (357) 48. A nucleic acid that encodes a VH domain and/or a VL domain of an antibody or fragment as defined in any one of Paragraphs 1 to 33.
- (358) 49. A nucleic acid that encodes a VH domain comprising the amino acid sequence of a VH domain of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or an amino acid that is at least 70% identical thereto.
- (359) 50. A nucleic acid that encodes a VL domain comprising the amino acid sequence of a VL domain of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); or an amino acid that is at least 70% identical thereto.
- (360) 51. A nucleic acid comprising a nucleotide sequence that is at least 70% identical to the sequence of SEQ ID

- NO: 260 or 320.
- (361) Alternatively:
- (362) A nucleic acid comprising a nucleotide sequence that is at least 70% identical to the sequence of SEQ ID NO: 2, 66 or 200.
- (363) 52. A nucleic acid that encodes a heavy chain and/or a light chain of an antibody or fragment as defined in any one of Paragraphs 1 to 33.
- (364) 53. A nucleic acid that encodes a heavy chain comprising a VH amino acid sequence that is at least 70% identical to SEQ ID NO: 259 or 319; and a C region amino acid sequence that is at least 70% identical to a IGHG4 sequence, optionally at least 70% identical to SEQ ID NO: 3 or 152.
- (365) Alternatively:
- (366) A nucleic acid that encodes a heavy chain comprising a VH amino acid sequence that is at least 70% identical to SEQ ID NO: 1, 65, and a C region amino acid sequence that is at least 70% identical to a IGHG4 sequence, optionally at least 70% identical to SEQ ID NO: 3 or 152.
- (367) 54. A nucleic acid that encodes a light chain comprising a VL amino acid sequence that is at least 70% identical to SEQ ID NO: 289 or 349; and a C region amino acid sequence that is at least 70% identical to a IGKC sequence, optionally at least 70% identical to SEQ ID NO: 156. (368) Alternatively:
- (369) A nucleic acid that encodes a light chain comprising a VL amino acid sequence that is at least 70% identical to SEQ ID NO: 33, 95 and 229; and a C region amino acid sequence that is at least 70% identical to a IGKC sequence, optionally at least 70% identical to SEQ ID NO: 156.
- (370) 55. A nucleic acid (eg, in a host cell, eg, a CHO or HEK293 or Cos cell) comprising a. a nucleotide sequence that is at least 70% identical to a heavy chain sequence of an antibody selected from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT); and/or b. a nucleotide sequence that is at least 70% identical to a light chain sequence of an antibody selected respectively from CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT).
- (371) 56. A vector comprising the nucleic acid(s) (eg, the nucleic acid(s) of any one of Paragraphs 48 to 55); optionally wherein the vector is a CHO or HEK293 vector.
- (372) 57. A host cell comprising the nucleic acid(s) (eg, the nucleic acid(s) of any one of Paragraphs 48 to 55) or the vector of Paragraph 56.
- (373) 58. An antibody, fragment, combination, vector, host cell, use or method as herein described.
- (374) In any embodiment herein, preferably the selected antibody is CL-274711.
- (375) Therapeutic Administration and Formulations
- (376) The invention provides therapeutic compositions comprising the anti-PCSK9 antagonist, eg, antibodies or antigen-binding fragments thereof, of the present invention. The administration of therapeutic compositions in accordance with the invention will be administered with suitable carriers, excipients, and other agents that are incorporated into formulations to provide improved transfer, delivery, tolerance, and the like. A multitude of appropriate formulations can be found in the formulary known to all pharmaceutical chemists: Remington's Pharmaceutical Sciences, Mack Publishing Company, Easton, Pa. These formulations include, for example, powders, pastes, ointments, jellies, waxes, oils, lipids, lipid (cationic or anionic) containing vesicles (such as LIPOFECTINT™), DNA conjugates, anhydrous absorption pastes, oil-in-water and water-in-oil emulsions, emulsions carbowax (polyethylene glycols of various molecular weights), semi-solid gels, and semi-solid mixtures containing carbowax. See also Powell et al. "Compendium of excipients for parenteral formulations" PDA (1998) J Pharm Sci Technol 52:238-311.
- (377) The dose may vary depending upon the age and the size of a subject to be administered, target disease, conditions, route of administration, and the like. When the antagonist, eg, antibody, of the present invention is used for treating various conditions and diseases associated with the PCSK9 in an adult patient, it is advantageous to intravenously administer the antibody of the present invention normally at a single dose of about 0.01 to about 20 mg/kg body weight, more preferably about 0.02 to about 7, about 0.03 to about 5, or about 0.05 to about 3 mg/kg body weight. Depending on the severity of the condition, the frequency and the duration of the treatment can be adjusted.
- (378) Various delivery systems are known and can be used to administer the antagonist or pharmaceutical composition of the invention, for example a antagonist provided by e.g., encapsulation in liposomes, microparticles, microcapsules, recombinant cells capable of expressing the mutant viruses, receptor mediated endocytosis (see, e.g., Wu et al. (1987) J. Biol. Chem. 262:4429-4432). Methods of introduction include, but are not limited to, intradermal, intramuscular, intraperitoneal, intravenous, subcutaneous, intranasal, epidural, and oral routes. The antagonist or composition may be administered by any convenient route, for example by infusion or bolus injection, by absorption through epithelial or mucocutaneous linings (e.g., oral mucosa, rectal and intestinal mucosa, etc.) and may be administered together with other biologically active agents. Administration can be systemic or local.

(379) The antagonist or pharmaceutical composition can be also delivered in a vesicle, in particular a liposome (see Langer (1990) Science 249:1527-1533; Treat et al. (1989) in Liposomes in the Therapy of Infectious Disease and Cancer, Lopez Berestein and Fidler (eds.), Liss, New York, pp. 353-365; Lopez-Berestein, ibid., pp. 317-327; see generally ibid.).

(380) In certain situations, the antagonist or pharmaceutical composition can be delivered in a controlled release

system. In one embodiment, a pump may be used (see Langer, supra; Sefton (1987) CRC Crit. Ref Biomed. Eng. 14:201). In another embodiment, polymeric materials can be used; see, Medical Applications of Controlled Release, Langer and Wise (eds.), CRC Pres., Boca Raton, Fla. (1974). In yet another embodiment, a controlled release system can be placed in proximity of the composition's target, thus requiring only a fraction of the systemic dose (see, e.g., Goodson, in Medical Applications of Controlled Release, supra, vol. 2, pp. 115-138, 1984). (381) The injectable preparations may include dosage forms for intravenous, subcutaneous, intracutaneous and intramuscular injections, drip infusions, etc. These injectable preparations may be prepared by methods publicly known. For example, the injectable preparations may be prepared, e.g., by dissolving, suspending or emulsifying the antibody or its salt described above in a sterile aqueous medium or an oily medium conventionally used for injections. As the aqueous medium for injections, there are, for example, physiological saline, an isotonic solution containing glucose and other auxiliary agents, etc., which may be used in combination with an appropriate solubilizing agent such as an alcohol (e.g., ethanol), a polyalcohol (e.g., propylene glycol, polyethylene glycol), a nonionic surfactant [e.g., polysorbate 80, HCO-50 (polyoxyethylene (50 mol) adduct of hydrogenated castor oil)], etc. As the oily medium, there are employed, e.g., sesame oil, soybean oil, etc., which may be used in combination with a solubilizing agent such as benzyl benzoate, benzyl alcohol, etc. The injection thus prepared is preferably filled in an appropriate ampoule. A pharmaceutical composition of the present invention can be delivered subcutaneously or intravenously with a standard needle and syringe. In addition, with respect to subcutaneous delivery, a pen delivery device readily has applications in delivering a pharmaceutical composition of the present invention. Such a pen delivery device can be reusable or disposable. A reusable pen delivery device generally utilizes a replaceable cartridge that contains a pharmaceutical composition. Once all of the pharmaceutical composition within the cartridge has been administered and the cartridge is empty, the empty cartridge can readily be discarded and replaced with a new cartridge that contains the pharmaceutical composition. The pen delivery device can then be reused. In a disposable pen delivery device, there is no replaceable cartridge. Rather, the disposable pen delivery device comes prefilled with the pharmaceutical composition held in a reservoir within the device. Once the reservoir is emptied of the pharmaceutical composition, the entire device is discarded. (382) Numerous reusable pen and autoinjector delivery devices have applications in the subcutaneous delivery of a antagonist or pharmaceutical composition of the present invention. Examples include, but certainly are not limited to AUTOPEN™ (Owen Mumford, Inc., Woodstock, UK), DISETRONIC™ pen (Disetronic Medical Systems, Burghdorf, Switzerland), HUMALOG MIX 75/25 pen, HUMALOG™ pen, HUMALIN 70/30™ pen (Eli Lilly and Co., Indianapolis, Ind.), NOVOPENT™ I, II and Ill (Novo Nordisk, Copenhagen, Denmark), NOVOPEN JUNIOR™ (Novo Nordisk, Copenhagen, Denmark), BD™ pen (Becton Dickinson, Franklin Lakes, N.J.), OPTIPENTTM, OPTIPEN PROTM, OPTIPEN STARLETTM, and OPTICLIKTTM (Sanofi-Aventis, Frankfurt, Germany), to name only a few. Examples of disposable pen delivery devices having applications in subcutaneous delivery of a pharmaceutical composition of the present invention include, but certainly are not limited to the SOLOSTAR™ pen (Sanofi-Aventis), the FLEXPEN™ (Novo Nordisk), and the KWIKPEN™ (Eli Lilly). (383) Advantageously, the pharmaceutical compositions for oral or parenteral use described above are prepared into dosage forms in a unit dose suited to fit a dose of the antagonist(s). Such dosage forms in a unit dose include, for example, tablets, pills, capsules, injections (ampoules), suppositories, etc. The amount of the aforesaid antibody contained is generally about 5 to about 500 mg per dosage form in a unit dose; especially in the form of injection, it is preferred that the aforesaid antibody is contained in about 5 to about 100 mg and in about 10 to about 250 mg for the other dosage forms. (384) For convenience, the meaning of some terms and phrases used in the specification, examples, and appended

claims, are provided below. Unless stated otherwise, or implicit from context, the following terms and phrases include the meanings provided below. The definitions are provided to aid in describing particular embodiments, and are not intended to limit the claimed invention, because the scope of the invention is limited only by the claims. Unless otherwise defined, all technical and scientific terms used herein have the same meaning as commonly understood by one of ordinary skill in the art to which this invention belongs. If there is an apparent discrepancy between the usage of a term in the art and its definition provided herein, the definition provided within the specification shall prevail.

(385) For convenience, certain terms employed herein, in the specification, examples and appended claims are collected here.

(386) The terms "decrease", "reduced", or "reduction" are all used herein to mean a decrease by a statistically significant amount. In some embodiments, "reduce," "reduction" or "decrease" typically means a decrease by at

least 10% as compared to a reference level (e.g. the absence of a given treatment) and can include, for example, a decrease by at least about 10%, at least about 20%, at least about 25%, at least about 30%, at least about 35%, at least about 40%, at least about 45%, at least about 55%, at least about 60%, at least about 65%, at least about 70%, at least about 75%, at least about 85%, at least about 90%, at least about 99%, or more. As used herein, "reduction" does not encompass a complete reduction as compared to a reference level. A decrease can be preferably down to a level accepted as within the range of normal for an individual without a given disorder. However, for example, for the purposes of lowering or reducing cholesterol level, for example, a reduction by about 5-10 points can be considered a "decrease" or "reduction."

(387) In certain aspects of all embodiments of the invention, the term "inhibition" is used. Inhibition refers and refers to decrease by at least 10% as compared to a reference level (e.g. the absence of a given treatment) and can include, for example, a decrease by at least about 10%, at least about 20%, at least about 25%, at least about 30%, at least about 35%, at least about 40%, at least about 45%, at least about 50%, at least about 55%, at least about 60%, at least about 65%, at least about 70%, at least about 75%, at least about 80%, at least about 85%, at least about 90%, at least about 95%, at least about 98%, at least about 99%, or more including 100% inhibition as compared to a reference level. "Complete inhibition" refers to a 100% inhibition as compared to a reference level. (388) The terms "increased", "increase", "enhance", or "activate" are all used herein to mean an increase by a statically significant amount. In some embodiments, the terms "increased", "increase", "enhance", or "activate" can mean an increase of at least 10% as compared to a reference level, for example an increase of at least about 20%, or at least about 30%, or at least about 40%, or at least about 50%, or at least about 60%, or at least about 70%, or at least about 80%, or at least about 90% or up to and including a 100% increase or any increase between 10-100% as compared to a reference level, or at least about a 2-fold, or at least about a 3-fold, or at least about a 4-fold, or at least about a 5-fold or at least about a 10-fold increase, or any increase between 2-fold and 10-fold or greater as compared to a reference level. In the context of a marker or symptom, an "increase" is a statistically significant increase in such level.

(389) As used herein, the term "substantially" refers to the qualitative condition of exhibiting total or near-total extent or degree of a characteristic or property of interest. One of ordinary skill in the biological arts will understand that biological and chemical phenomena rarely, if ever, go to completion and/or proceed to completeness or achieve or avoid an absolute result. The term "substantially" is therefore used herein to capture the potential lack of completeness inherent in many biological and chemical phenomena. For the removal of doubt, "substantially" can refer to at least a 90% extent or degree of a characteristic or property of interest, e.g. at least 90%, at least 92%, at least 95%, at least 98%, at least 99% or greater.

(390) All patents and other publications; including literature references, issued patents, published patent applications, and co-pending patent applications; cited throughout this application are expressly incorporated herein by reference for the purpose of describing and disclosing, for example, the methodologies described in such publications that might be used in connection with the technology described herein. These publications are provided solely for their disclosure prior to the filing date of the present application. Nothing in this regard should be construed as an admission that the inventors are not entitled to antedate such disclosure by virtue of prior invention or for any other reason. All statements as to the date or representation as to the contents of these documents is based on the information available to the applicants and does not constitute any admission as to the correctness of the dates or contents of these documents.

(391) The description of embodiments of the disclosure is not intended to be exhaustive or to limit the disclosure to the precise form disclosed. While specific embodiments of, and examples for, the disclosure are described herein for illustrative purposes, various equivalent modifications are possible within the scope of the disclosure, as those skilled in the relevant art will recognize. For example, while method steps or functions are presented in a given order, alternative embodiments may perform functions in a different order, or functions may be performed substantially concurrently. The teachings of the disclosure provided herein can be applied to other procedures or methods as appropriate. The various embodiments described herein can be combined to provide further embodiments. Aspects of the disclosure can be modified, if necessary, to employ the compositions, functions and concepts of the above references and application to provide yet further embodiments of the disclosure. Moreover, due to biological functional equivalency considerations, some changes can be made in protein structure without affecting the biological or chemical action in kind or amount. These and other changes can be made to the disclosure in light of the detailed description. All such modifications are intended to be included within the scope of the appended claims.

(392) Specific elements of any of the foregoing embodiments can be combined or substituted for elements in other embodiments. Furthermore, while advantages associated with certain embodiments of the disclosure have been described in the context of these embodiments, other embodiments may also exhibit such advantages, and not all embodiments need necessarily exhibit such advantages to fall within the scope of the disclosure.

(393) It will be understood that particular configurations, aspects, examples, clauses and embodiments described herein are shown by way of illustration and not as limitations of the invention. The principal features of this invention can be employed in various embodiments without departing from the scope of the invention. Those skilled in the art will recognize, or be able to ascertain using no more than routine study, numerous equivalents to the specific procedures described herein. Such equivalents are considered to be within the scope of this invention and are covered by the claims. All publications and patent applications mentioned in the specification are indicative of the level of skill of those skilled in the art to which this invention pertains. All publications and patent applications are herein incorporated by reference to the same extent as if each individual publication or patent application was specifically and individually indicated to be incorporated by reference. The use of the word "a" or "an" when used in conjunction with the term "comprising" in the claims and/or the specification may mean "one," but it is also consistent with the meaning of "one or more," "at least one," and "one or more than one." The use of the term "or" in the claims is used to mean "and/or" unless explicitly indicated to refer to alternatives only or the alternatives are mutually exclusive, although the disclosure supports a definition that refers to only alternatives and "and/or." Throughout this application, the term "about" is used to indicate that a value includes the inherent variation of error for the device, the method being employed to determine the value, or the variation that exists among the study subjects.

(394) As used in this specification and claim(s), the words "comprising" (and any form of comprising, such as "comprise" and "comprises"), "having" (and any form of having, such as "have" and "has"), "including" (and any form of including, such as "includes" and "include") or "containing" (and any form of containing, such as "contains" and "contain") are inclusive or open-ended and do not exclude additional, unrecited elements or method steps

(395) Any part of this disclosure may be read in combination with any other part of the disclosure, unless otherwise apparent from the context.

(396) All of the compositions and/or methods disclosed and claimed herein can be made and executed without undue experimentation in light of the present disclosure. While the compositions and methods of this invention have been described in terms of preferred embodiments, it will be apparent to those of skill in the art that variations may be applied to the compositions and/or methods and in the steps or in the sequence of steps of the method described herein without departing from the concept, spirit and scope of the invention. All such similar substitutes and modifications apparent to those skilled in the art are deemed to be within the spirit, scope and concept of the invention as defined by the appended claims.

(397) The present invention is described in more detail in the following non-limiting Examples EXAMPLES

(398) Plasma LDL cholesterol (LDL-C) is a main risk factor of cardiovascular diseases, which cause 4 million deaths annually in Europe. Clearance of plasma LDL-C is primarily carried out in liver by LDL receptor (LDLR). LDLR constantly internalises LDL-C from plasma into hepatocytes and continues LDL-C uptake by recycling back to the cell surface. The surface level of LDLR is negatively regulated by PCSK9, which directly binds to LDLR and facilitates lysosome-mediated degradation of LDLR. Diminished surface exposure of LDLR to LDL-C reduces LDL-C internalisation in hepatocytes and thus increases plasma LDL-C, which may lead to plaque forming and clogging in blood stream. Recently targeting PCSK9 emerges as a promising strategy to control plasma LDL-C level.

(399) We discovered monoclonal antibodies that block PCSK9 function and have therapeutic potential. We first immunised KymiceTM (see, eg, WO2011/004192 and Lee et al, Nat Biotechnol, 2014 April; 32(4):356-63. doi: 10.1038/nbt.2825. Epub 2014 Mar. 16, "Complete humanization of the mouse immunoglobulin loci enables efficient therapeutic antibody discovery") with both human and mouse PCSK9 proteins to generate cross-reactive antibodies. To avoid the unwanted effect of a His tag, we made a cell line that actively secretes the native form of both human and mouse PCSK9 protein without His tag for immunisation. Following immunisation, antibody sequences were retrieved from antigen-specific B cells by Next Generation Sequencing and analyzed in silico to identify favourable antibodies to progress. After expression, antibodies were screened and selected based on cross-reactive binding, affinity, and function by a sequential in vitro screening cascade. Binding to human PCSK9 variants was confirmed to ensure efficacy in the majority of human population. Biophysical features of selected antibodies were characterised from the perspective of developability. In a mouse model of hyperlipidemia, two outstanding antibodies (CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT)) showed superior function in decreasing plasma non-HDL cholesterol without changing HDL cholesterol and sustained effect with longer duration compared to benchmark antibodies. In conclusion, we successfully elicited cross-reactive antibodies against PCSK9 and identified at least two superior fully human monoclonal antibodies to block PCSK9 function in vitro and in vivo.

Example 1: Identification of Antibodies by In Vitro Assays

(400) Five immunisation campaigns were conducted in Kymice™ and in vitro assays were conducted including the

- following selection criteria:— (a) Cross-reactive binders against human PCSK9 (hPCSK9), cynomolgus monkey PCSK9 (cynoPCSK9) and mouse PCSK9 (mPCSK9) were selected; (b) Antibody affinity against hPCSK9a and mPCSK9 were determined and antibodies with higher affinity were prioritized; (c) In a cell-based assay in vitro, neutralisation of antibodies was demonstrated to reduce internalisation of fluorescent hPCSK9 in a human liver cell line (HepG2); and (d) Simultaneously, functionality of antibodies to restore LDL uptake in HepG2 cells was determined in the same in vitro cell-based assay as in (c).
- (401) Two antibodies (CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT)) showed particularly promising function both in vitro and in vivo (Example 2). (402) CL-148219
- (403) Compared to the benchmarks, CL-148219 showed stronger neutralisation activity with higher capacity to restore LDL uptake in vitro, suggesting CL-148219 may directly blocks the interaction between PCSK9 and LDLR. In a mouse model of hyperlipidemia (E3L.CETP mice; de Knijff, Westerterp, Ason, van den Hoek and Kühnast), CL-148219 reduced plasma non-HDL cholesterol superior to benchmark Ab with longer duration of effect. See FIGS. **1-4**.

(404) CL-148489

- (405) In the in vitro cell-based assay, CL-148489 neutralised hPCSK9 moderately but contained significant capacity to restore LDL uptake, suggesting this antibody may indirectly interfere the binding of PCSK9 to LDLR, possibly via blocking HSPG binding to PCSK9. This antibody also reduced plasma non-HDL cholesterol in a mouse model of hyperlipidemia (E3L.CETP mice) better than benchmarks with similar duration of effect. See FIGS. **5-8**. (406) The benchmarks were: alirocumab hIgG4-PE (SEQ ID Nos: 195 and 196) and "Regeneron pH-D" (a pH-dependent (differential PCSK9 binding inside versus outside lysosome) anti-PCSK9 hIgG4-PE antibody) (SEQ ID Nos: 197 and 198).
- (407) The antibodies' VH and VL regions were derived from the recombination of the following gene segments:— (408) TABLE-US-00001 CL-148219 VH VH IGHV4-59*01 D IGHD3-10*01 JH IGHJ6*02 CL-148219 VL VL IGKV2-28*01 JL IGKJ3*01 CL-148489 VH VH IGHV3-9*01 D IGHD3-9*01 JH IGHJ6*02 CL-148489 VL VL IGKV2D-29*01 JL IGKJ4*01

REFERENCES

(409) de Knijff, P. et al, 1991, "Familial dysbetalipoproteinemia associated with apolipoprotein E3-Leiden in an extended multigeneration pedigree", J. Clin. Invest. 88: 643-655; Westerterp, M. et al, 2006, "Cholesteryl ester transfer protein decreases high-density lipoprotein and severely aggravates atherosclerosis in APOE*3-Leidenmice", *Arterioscler. Thromb. Vasc. Biol.* 26: 2552-2559; Ason, B. et al, 2011, "PCSK9 inhibition fails to alter hepatic LDLR, circulating cholesterol, and atherosclerosis in the absence of ApoE", J Lipid Res. 2011 April, 52(4): 679-687; van den Hoek AM. et al, 2014, "APOE*3Leiden.CETP transgenic mice as model for pharmaceutical treatment of the metabolic syndrome", Diabetes Obes Metab. 2014 June; 16(6):537-44; and Kühnast, S et al, 2014, "Alirocumab inhibits atherosclerosis, improves the plaque morphology, and enhances the effects of a statin", J Lipid Res. 2014 October, 55(10): 2103-2112.

Example 2: In Vivo Assays

- (410) In this study we used APOE*3Leiden.cholesteryl ester transfer protein (CETP) mice, which contain mouse ApoE, human mutant APOE3*Leiden, and a functional LDLR. These mice have shown excellent translational value with the increases in cholesterol seen in the model treatable by the same therapies which are effective in patients and showing a similar maximum reduction in LDL levels to those obtainable in the clinic.
- (411) CETP mice were fed a high fat diet for 4 weeks and then received a single dose of antibody, isotype control or standard of care (alirocumab, an FDA approved PCSK9 inhibitor) on day 0. The mice then received a second higher dose (half log), on day 18. Individual animal cholesterol levels were measured on day 0, 3, 7, 14, 21 and 28. A full lipid profile was carried out on a pooled group sample from the terminal bleed at day 28.
- (412) Of the 10 of our antibodies tested two (CL-148219 and CL148489) reduced cholesterol levels by more than the equivalent dose of alirocumab with one those CL-148219 showing a longer duration of effect. In summary we have identified two potentially best in class PCSK9 inhibitors.

Methods

(413) See FIG. **9** for the study design. One hundred and twenty-eight female, 8-14 weeks of age, APOE*3Leiden.CETP transgenic mice were put on Western-type diet (WTD) with 0.15% cholesterol and 15% saturated fat. After 4 weeks run-in period 24 low-responder mice were removed from the study and the animals placed into groups (n=6-8), matched on age, body weight, plasma cholesterol, triglycerides and HDL-cholesterol after 4 h fasting (t=0). The mice were dosed subcutaneously at t=0 with 3 mg/kg of isotype control, benchmark or our anti-PCSK9 antibodies. Four of our antibodies, identified as excellent from prior in vitro screening, had an additional dose group of 1 mg/kg at t=0. At t=18 days the mice received a 2nd subcutaneous dose of the same antibody at a %.sub.z log higher dose than the first dose (i.e. first dose 3 mg/kg, 2nd dose 10 mg/kg). (414) Body weight and food intake were measured twice weekly. On day 0, 3, 7, 14, 21 and 28 post the initial dose 4

- h fasted plasma samples were collected for the determination of plasma total cholesterol, HDL-cholesterol and triglycerides. Plasma non-HDL-cholesterol values were calculated. At t=28 days, after the last blood collection, mice were sacrificed, and plasma and liver tissue collected. Lipoprotein profiles were determined on FPLC fractions from group pooled plasma samples.
- (415) Statistics: Two way annova and Dunnetts post-hoc analysis—Graphpad Prismi™.
- (416) Results
- (417) Total Cholesterol (FIG. 10)
- (418) The isotype control group showed plasma total cholesterol levels of 16-22 mmol/L during the study.
- (419) Alirocumab significantly reduced total cholesterol at day 3 in the 3 mg/kg dose (-34.9%), and day 21 and 28 (respectively 3 and 10 days after the second injection) in the 10 mg/kg dose (-33.2% and -32.3%, respectively) compared to the isotype control antibody.
- (420) CL-148219 significantly reduced total cholesterol at day 3 in both the 1 and 3 mg/kg dose (-36.4% and -47.0%, respectively) and at day 7 in the 3 mg/kg dose (-32.6%). Following the second dose CL-148219 significantly reduced total cholesterol at day 21 and day in both the 3 mg/kg dose (-33.4% and -27.1%, respectively) and 10 mg/kg dose (-46.3% and -49.6%, respectively).
- (421) CL-148489 significantly reduced total cholesterol at day 3 and day 21 (3 days after the second injection) in the 3 mg/kg dose (-47.3% and -32.3%, respectively) and following the second dose on day 21 and day 28 in the 10 mg/kg dose (-42.9% and -56.0%, respectively).
- (422) Total Non-HDL Cholesterol (FIG. 11)
- (423) The isotype control group showed non-HDL-cholesterol levels of 15.2-21.3 mmol/L during the study.
- (424) Alirocumab significantly reduced non-HDL-cholesterol at day 3 in the 3 mg/kg dose (-36.5%), and day 21 and 28 (respectively 3 and 10 days after the second injection) in the 10 mg/kg dose (-35.5% and 34.3%, respectively) compared to the isotype control antibody.
- (425) CL-148219 significantly reduced non-HDL-cholesterol at day 3 in both the 1 and 3 mg/kg dose (-37.7% and -49.1%, respectively), and at day 7 in the 3 mg/kg dose (-33.5%). Following the second injection CL-148219 significantly reduced non-HDL-cholesterol at day 21 and day 28 in both the 3 mg/kg (-34.4% and -29.2%, respectively) and 10 mg/kg dose (-49.6% and -51.8%, respectively).
- (426) CL-148489 significantly reduced non-HDL-cholesterol at day 3 and day 21 (3 days after the second injection) in the 3 mg/kg dose (-49.4% and -35.0%, respectively) and at following the second injection at day 21 and day 28 in the 10 mg/kg dose (-45.5% and -58.5%, respectively).
- (427) HDL Cholesterol (FIG. 12)
- (428) None of the antibodies had an effect on HDL cholesterol.
- (429) Triglycerides (FIG. 13)
- (430) During the study, plasma triglyceride levels remained stable in the isotype control group. Whilst none of the antibodies significantly affected plasma triglycerides compared to the isotype control group CL-148219, CI-148489 and Alirocumab showed a trend towards lower triglyceride levels at day 28.
- (431) Phospholipid and Cholesterol Profile (FIG. 14)
- (432) Phospholipid (A) and Cholesterol (B) profiles for the pooled samples are shown below—fractions 4-8 are considered VLDL, 9-15 as LDL and 16-24 as HDL. No statistics were performed as the measurements are carried out on one pooled sample was used per group. The lipoprotein profiles confirm that alirocumab, CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT) reduced VLDL-cholesterol and LDL-cholesterol and had no effect on HDL-cholesterol.

CONCLUSIONS

(433) Our antibodies were well tolerated with the animals showing normal behavior and no signs of discomfort during the study; Sectioning of the tissues showed no gross pathology; Body weight and food intake were not significantly affected by any of the antibodies; Liver weight were not significantly affected by any of the antibodies; CL-148219, and CL-148489 significantly decreased plasma total cholesterol and plasma non-HDL-cholesterol in a dose-dependent fashion compared to the isotype control group. Lipoprotein profiles showed decreased plasma VLDL-cholesterol in mice treated with these antibodies; None of our antibodies significantly affected HDL-cholesterol compared to the isotype control group. None of the antibodies significantly affected plasma triglyceride levels, although Alirocumab, CL-274711, CL-148219QLT, CL-274698, CL-148219 and CL-148489 (eg, CL-274711 or CL-148219QLT) showed a trend towards decreased plasma triglyceride levels compared to the isotype control group.

Examples 3 & 4

(434) Mice

(435) For Example 3, female APOE*3Leiden.CETP transgenic mice were bred. Mice were housed in macrolon cages (3 or 4 mice per cage) in animal rooms; relative humidity 40-70%, temperature 20-24° C., light cycle 7 am to 7 pm. Mice were supplied with a food and sterilized tap water ad libitum. For Example 4, C57BL/6 mice weighing

- 18-22 g on the day of the experiment were used. The animals were be housed in groups of 2-4 in polysulfone cages (floor area=1500 cm2) under standard conditions: room temperature (22±2° C.), hygrometry (55±10%), light/dark cycle (12 h/12 h), air replacement (15-20 volumes/hour), water and food ad libitum.
- (436) Plasma Cholesterol, Lipoprotein and Triglyceride Analysis
- (437) Total plasma cholesterol was determined using the Cholesterol CHOD-PAP kit (Roche, Mannheim, Germany) and triglycerides was determined using the Triglycerides GPO-PAP kit (Roche, Mannheim, Germany). To determine plasma HDL-cholesterol ApoB-containing lipids were precipitated using PEG-6000/glycine followed by quantification using the Cholesterol CHOD-PAP kit (Roche, Mannheim, Germany). Lipoprotein profiles were measured by FPLC analysis using an AKTA apparatus (GE Healthcare). Analysis was performed on pooled group samples pooled. Fractions were collected based on retention time and cholesterol and phospholipids measured in the fractions using the Cholesterol CHOD-PAP kit from (Roche, Mannheim, Germany) and Phospholipids kit (Instruchemie, Delfzijl, Netherlands).
- (438) Serum Antibody Concentration
- (439) Serum antibody concentration were determined using a capture ELISA. Plates were coated overnight at 2-8° C. with an Anti-Human IgG (Fc specific) antibody. The following day the assay plate was washed three times with 300 μ L/well of PBS+0.1% Tween (PBS-T) using a plate washer. The plates were then blocked with 250 μ L/well 2.5% Milk per well for at least 1 hr at room temperature.
- (440) Standards and QC's were prepared in Eppendorf tubes. The standards, samples and QCs were then diluted to the 1 in 20 MRD in assay buffer in Eppendorf tubes. The plates were washed three times with 300 μ L/well of PBS-T using a plate washer. 100 μ L per well of standard curve, sample and QC was added to the assay plate as per plate map. The plate was incubated for 1 hr at RT, shaking at 300 RPM. Plates were washed five times with 300 μ L/well of PBS-T using a plate washer and 100 μ L of Anti-Human IgG (Fc specific)-Peroxidase diluted 1 in 5000 in PBS added to each well and the plate then incubated for 1 hr at RT, shaking at 300 RPM. The plate was then washed five times with 300 μ L of PBS-T using a plate washer. 100 μ L of TMB substrate was added to each well and the plate incubated for 15 minutes at room temperature in the dark. 100 μ L/well of stop solution (1M sulfuric acid) was then added to each well and the optical density determined using a microplate reader set to 450 nm with a reference read at 540 nm. The reference reading was then subtracted from the 450 nm reading using plate reader software. Softmax Pro was then used for analysis of the data using regression analysis and a 4PL curve fit. The concentrations were measured off the standard curve.

Example 3

- (441) Female, 8-14 weeks of age APOE*3Leiden.CETP transgenic mice were put on Western-type diet (WTD) containing 0.15% cholesterol and 15% saturated fat. After the 4 weeks run-in period low-responder mice were removed from the study. The remaining mice were matched for age, body weight, plasma cholesterol, and triglycerides. The animals were placed in groups of 6 (isotype control group) or 10.
- (442) Animals received a subcutaneous injection at day 0 (5 mL/kg). Antibodies (Isotype Control, Benchmark Antibody F, CL-148218 and CL-274711) were administered at 1, 3 or 10 mg/kg. On days 0, 3, 7, 14, 18, 21, and 28 post treatment blood samples were taken from the tail vein following a 4-hour fasting period, animals. Blood was collected using CB 300 K2E microvettes (Sarstedt, Nürnbrecht, Germany) containing EDTA-dipotassium salt for total cholesterol, triglycerides (day 0, 7 and 28), and HDL-cholesterol measurements. The tubes or capillaries were placed on ice immediately. EDTA plasma was obtained after centrifugation (10 min at 6000 rpm) at 4° C. Mice were sacrificed at day 28 by CO.sub.2 asphyxiation, directly after the last tail vein blood sampling point. Group lipoprotein profiles were ascertained on the day 28 samples.
- (443) The aim of this experiment was to evaluate the pharmacodynamic effects on total/non-HDL cholesterol and lipoprotein profiles of a dose response of antibodies CL-274711, CL-148219 QLT and Benchmark Antibody F in APOE*3Leiden.CETP transgenic mice feed a Western-type diet containing 0.15% cholesterol and 15% saturated fat. (444) All three antibodies reduced cholesterol levels at all doses tested (see FIGS. **15-17**) and increasing the dose increased the reduction in cholesterol seen and duration of the effect. At 1 and 3 mg/kg CL-148219 QLT was more effective than Benchmark Antibody F at reducing cholesterol levels and at all doses tested CL-148219 QLT had a longer duration of effect than Benchmark Antibody F (FIGS. **15-17**). Further analysis showed that CL-148219 QLT was as effective as a 3 times higher dose of Benchmark Antibody F at reducing cholesterol levels and had the same duration of effect as that higher dose of Benchmark Antibody F (FIGS. **18-19**). CL-274711 reduced cholesterol levels to the same levels as those seen with the same dose of Benchmark Antibody F but had a longer duration of effect (FIG. **16-17**).
- (445) The antibodies showed an equivalent reduction in non-HDL cholesterol levels at all doses tested (see FIG. **20**) with increasing doses resulting in incremental increases in the reduction in non-HDL cholesterol and the duration of the response. At 1 and 3 mg/kg CL-148219 QLT was more effective than Benchmark Antibody F at reducing non-HDL cholesterol levels and at all the doses tested CL-148219 QLT had a longer duration of effect than Benchmark Antibody F (FIG. **20**). Further analysis also showed that CL-148219 QLT was as effective as a 3 times higher dose

of Benchmark Antibody F at reducing non-HDL cholesterol levels and had the same duration of effect as that higher dose of Benchmark Antibody F (FIG. **21**). CL-274711 reduced cholesterol levels to the same levels as those seen with the same dose of Benchmark Antibody F but had a longer duration of effect (FIG. **21**). Evaluation of the phospholipid profiles at day 28 (FIG. **22**) showed that both CL-274711 and CL-148219 QLT reduced VLDL/LDL cholesterol and phospholipids in a dose dependent manner. In comparison with Benchmark Antibody F both CL-274711 and CL-148219 QLT at 10 mg/kg were more effective at reducing VLDL/LDL cholesterol and phospholipids

Example 4

- (446) Male C57BL/6 mice (JANVIER LABS, C.S. 4105, Saint-Berthevin F-53941, France), weighing 18-22 g on the day of the experiment were used for the study. Animals received a 3 mg/kg subcutaneous injection of CL-148219 QLT, CL-274711 or Benchmark F on day 0 (10 mL/kg). Blood sampling was performed on days 1, 3, 7, 14, 21 and 28 post-dose (n=3 per sampling point). Serum was then prepared from each blood sample which was then used to determine the antibody concentration at each time point.
- (447) The aim of the experiment was to evaluate the pharmacokinetics of CL-148219 QLT, CL-274711 and Benchmark Antibody F over the course of 28 days. Evaluation of the PK profiles showed that both CL-274711 and CL-148218 had a higher initial serum antibody concentration than Benchmark Antibody F and this higher serum antibody concentration was retained for the 28 days of the study (FIG. **23**). CL-148219 QLT initially had a higher serum antibody concentration than CL-274711 but CL-274711 had a slower clearance from the serum. (448) Further Data:
- (449) Reference is made to FIG. **24** onwards showing efficacy of antibodies of the invention. Reference is also made to Table 1 showing antibody affinity against human PCSK9 variants, cynomolgus monkey PCSK9, and mouse PCSK9 at neutral pH (7.6) versus acidic pH (5.8). KD was measured by ProteOn™ SPR analysis system. (450) We included six PCSK9 benchmark antibodies as controls for comparison. Three benchmark antibodies were versions of marketed antibodies (benchmark antibody A, B, and F) and three were selected from antibodies in clinical trials (benchmark antibody C, D, and E). Benchmark antibody A, B, C, D, and E were made in human IgG4PE format. Benchmark antibody F was in human IgG1 format.
- (451) Antigens used for immunisation and/or screening included 4 human PCSK9 variants, cynomolgus monkey PCSK9, and mouse PCSK9. Here we called 4 human PCSK9 variants as following: human PCSK9 reference (hPCSK9 Ref), human PCSK9 variant a (hPCSK9a), human PCSK9 variant b (hPCSK9b), and human PCSK9 D374Y gain-of-function mutant (hPCSK9 GOF). Relative to the human PCSK9 reference sequence, human variant a has V4741 and G670E amino acid substitution, and human variant b has A53V, V4741, and G670E amino acid substitution. We included these 4 human variants for assay screening in order to find an antibody with capability to neutralise all the 4 variants and the mutant in human population. (452) CL-148219
- (453) Compared to the benchmarks, CL-148219 showed stronger neutralisation activity with higher capacity to restore LDL uptake in vitro, suggesting CL-148219 may directly block the interaction between PCSK9 and LDLR. In a mouse model of hyperlipidemia (E3L.CETP mice; de Knijff, Westerterp, Ason, van den Hoek and Kühnast), CL-148219 reduced plasma non-HDL cholesterol superior to benchmark Ab with longer duration of effect. See FIGS. **24-27** and Table 1.

(454) CL-148219 QLT Mutant

- (455) By sequence analysis in silicon, a N-glycosylation motif was found in the heavy chain of CL-148219. The NLT motif, located in the framework 3 region of the heavy chain, may be glycosylated and may have adverse effect in developability. It was later confirmed by Mass Spectrometry that the NLT motif in CL-148219 was indeed N-glycosylated. We thus introduced mutation to the NLT motif by replacing N with Q residue, generating CL-148219 QLT mutant to disrupt glycosylation. Removal of glycosylation was further confirmed by Mass Spectrometry. CL-148219 QLT mutant showed similar affinity and functionality in vitro when compared to the parental CL-148219. Like parental CL-148219, the QLT mutant reduced plasma non-HDL cholesterol superior to benchmarks with longer duration of effect in a hyperlipidemia mouse model (E3L.CETP mice). See FIGS. **27-31** and Table 1. (456) CL-148489
- (457) In the in vitro cell-based assay, CL-148489 neutralised hPCSK9 moderately but contained significant capacity to restore LDL uptake, suggesting this antibody may indirectly interfere the binding of PCSK9 to LDLR, possibly via blocking HSPG binding to PCSK9. This antibody also reduced plasma non-HDL cholesterol in a mouse model of hyperlipidemia (E3L.CETP mice) better than benchmarks with similar duration of effect. See FIGS. **32-35** and Table 1.

(458) CL-274698

(459) Unlike CL-148219 and CL-148489 that were derived from a mouse with humanized antibody loci and without knock-out of the PCSK9 gene, CL-274698 was derived from a PCSK9 knockout mouse. In the absence of PCSK9, mice produced high affinity antibodies against immunogens including both human and mouse PCSK9. CL-274698

was thus generated and screened. CL-274698 not only showed high affinity against PCSK9 with KD in a sub-nM range, but also exhibited superior function to neutralise hPCSK9 and restore LDL uptake in vitro. In the in vivo efficacy study by using E3L.CETP mice, CL-274698 reduced plasma non-HDL cholesterol more than benchmarks within 3 days after Ab injection, with the fact that the level of non-HDL cholesterol was similar to benchmarks after day 7. See FIGS. **36-39** and Table 1.

(460) CL-274711

(461) Similar to CL-274698, CL-274711 was generated by immunizing a PCSK9 knockout mouse. In the in vitro cell-based assays, CL-274711 neutralised hPCSK9 and restored LDL uptake better than benchmarks. In the in vivo efficacy assay by using E3L.CETP mice, CL-274711 showed slightly superior function in reducing plasma non-HDL cholesterol with longer duration of effect compared to benchmarks. See FIGS. **40-43** and Table 1. Method

(462) Cell-Based Functional Assay In Vitro

(463) HepG2 cells were seeded at 6×10.sup.4 cells/well into 96-well clear flat-bottom cell culture plates (Costar) in complete culture medium, containing 10% FBS, 1× Penicillin/Streptomycin and 1× NEAA in MEMα. To allow attachment, cells were incubated in 5% CO.sub.2 incubator at 37° C. for 24 hours. The next day media were replaced by serum-free culture media (1× NEAA and 1× Penicillin/Streptomycin in MEMα), and cells were kept in incubator overnight (16-18 hours). On the following day, cells were treated with various concentrations of anti-PCSK9 monoclonal antibodies that were pre-incubated with 5 μg/ml AF647-labelled hPCSK9 GOF antigen. The antibodies were diluted from 60 μg/ml with 3 fold series dilutions. After 1 hour incubation with hPCSK9 GOF plus anti-PCSK9 antibodies, BODIPY LDL (Invitrogen) was added to cells at a final concentration of 10 μg/ml. LDL uptake was allowed by incubating cells for additional 3 hours. In the end of treatment, cells were detached by accutase (BD), and followed by fixation with 2% paraformaldehyde diluted in PBS (Alfa Aesar) overnight at 4° C. After washes, cells were resuspended in PBS containing 5 mM EDTA, and fluorescent cells were detected by CytoFlex flow cytometry.

(464) ProteOn SPR Analysis

(465) ProteOn™ XPR36 (Bio-Rad) was used to measure antibody binding affinity to antigen (human PCSK9 variants, cynomolgus monkey PCSK9 and mouse PCSK9) in neutral (HBS-EP+, pH7.6) and acidic buffer (30 mM sodium acetate, 150 mM NaCl, 0.05% P20, pH5.8) at 25° C. The SPR runs were performed using a GLC chip (Bio-Rad) immobilised with anti-human Fc antibodies. The analyte human antibodies were captured on the chip surface at 2 µg/ml. The PCSK9 proteins were injected at 100 nM, 25 nM, 6.25 nM, 1.56 nM, 0.39 nM for 180 seconds and dissociation was monitored for 600 seconds at a flow rate of 30 μL/min. The chip surface was regenerated using two injections of 10 mM Glycine pH 1.5 for 160 seconds at 30 μl/min. The sensorgrams were then fitted with a 1:1 model (Langmuir kinetic model) where ka and kd are fitted and KD was calculated by the ProteOn software. (466) TABLE-US-00002 TABLE 1 pH 7.6 pH 5.8 ka kd ka kd (1/Ms) (1/s) KD (M) (1/Ms) (1/s) KD (M) Benchmark Human PCSK9 var A 1.53E+06 3.00E-04 1.96E-10 1.68E+06 1.20E-04 7.13E-11 Ab B Human PCSK9 var B 1.57E+06 2.91E-04 1.86E-10 1.75E+06 1.10E-04 6.27E-11 Human PCSK9 Ref 1.50E+06 2.93E-04 1.95E-10 1.77E+06 1.06E-04 5.99E-11 Human PCSK9 GOF 1.32E+06 7.21E-04 5.48E-10 1.69E+06 3.27E-04 1.94E-10 Cyno PCSK9 1.21E+06 2.30E-04 1.90E-10 1.31E+06 6.95E-05 5.32E-11 Mouse PCSK9 1.16E+06 1.33E-03 1.15E-09 1.34E+06 4.96E-04 3.70E-10 CL-148219 Human PCSK9 var A 1.08E+06 2.34E-04 2.16E-10 9.08E+05 3.87E-04 4.26E-10 Human PCSK9 var B 1.05E+06 2.25E-04 2.15E-10 9.18E+05 3.78E-04 4.12E-10 Human PCSK9 Ref 1.13E+06 2.30E-04 2.05E-10 1.02E+06 3.74E-04 3.68E-10 Human PCSK9 GOF 9.49E+05 2.33E-04 2.45E-10 9.08E+05 3.91E-04 4.31E-10 Cyno PCSK9 9.58E+05 2.84E-04 2.96E-10 9.75E+05 3.36E-04 3.45E-10 Mouse PCSK9 7.91E+05 4.11E-04 5.20E-10 1.01E+06 7.01E-04 6.91E-10 CL-148219 Human PCSK9 var A 8.56E+05 2.29E-04 2.67E-10 6.02E+05 4.29E-04 7.12E-10 OLT Human PCSK9 var B 8.62E+05 2.21E-04 2.57E-10 6.07E+05 4.21E-04 6.94E-10 Human PCSK9 Ref 8.33E+05 2.26E-04 2.72E-10 6.22E+05 4.15E-04 6.67E-10 Human PCSK9 GOF 7.21E+05 2.19E-04 3.03E-10 6.06E+05 4.56E-04 7.53E-10 Cyno PCSK9 7.28E+05 2.56E-04 3.52E-10 5.84E+05 3.31E-04 5.68E-10 Mouse PCSK9 6.00E+05 3.52E-04 5.87E-10 4.72E+05 6.75E-04 1.43E-09 CL-148489 Human PCSK9 var A 8.94E+05 1.57E-04 1.75E-10 1.25E+06 1.00E-5* <8.0E-12** Human PCSK9 var B 8.48E+05 1.64E-04 1.93E-10 1.54E+06 1.00E-5* <6.49E-12** Human PCSK9 Ref 9.24E+05 1.63E-04 1.76E-10 1.50E+06 1.00E-5* <6.67E-12** Human PCSK9 GOF 9.47E+05 1.29E-04 1.36E-10 1.37E+06 1.00E-5* <7.30E-12** Cvno PCSK9 1.13E+06 5.73E-05 5.06E-11 1.31E+06 1.21E-04 9.24E-11 Mouse PCSK9 7.14E+05 1.92E-03 2.69E-09 1.38E+06 6.32E-04 4.60E-10 CL-274698 Human PCSK9 var A 1.37E+06 1.36E-04 9.98E-11 1.56E+06 4.81E-05 3.09E-11 Human PCSK9 var B 1.39E+06 1.27E-04 9.10E-11 1.60E+06 4.19E-05 2.62E-11 Human PCSK9 Ref 1.35E+06 1.25E-04 9.29E-11 1.60E+06 4.60E-05 2.86E-11 Human PCSK9 GOF 1.21E+06 8.36E-05 6.89E-11 1.53E+06 3.83E-05 2.50E-11 Cyno PCSK9 1.10E+06 1.08E-04 9.84E-11 1.26E+06 3.75E-05 2.97E-11 Mouse PCSK9 9.41E+05 2.57E-04 2.73E-10 1.33E+06 1.37E-04 1.03E-10 CL-274711 Human PCSK9 var A 2.95E+05 6.11E-05 2.07E-10 4.46E+05 6.99E-05 1.57E-10 Human PCSK9 var B

```
3.04E+05 5.44E-05 1.79E-10 4.42E+05 7.15E-05 1.62E-10 Human PCSK9 Ref 2.94E+05 6.04E-05 2.05E-10
4.33E+05 6.58E-05 1.52E-10 Human PCSK9 GOF 2.99E+05 5.29E-05 1.77E-10 4.86E+05 9.60E-05
1.98E-10 Cyno PCSK9 4.06E+05 5.87E-05 1.45E-10 5.45E+05 8.91E-05 1.64E-10 Mouse PCSK9 2.85E+05
2.53E-04 8.89E-10 5.08E+05 6.39E-04 1.26E-09 *1e-5 is the lower detection limits **KD cannot be accuratly
(467) TABLE-US-00003 VARIABLE REGION SEQUENCE SUMMARY SEQ DESCRIPTION ID CL- CL-
AMINO NO 148219 148489 ACID NUCLEOTIDE 1 X VH REGION X 65 X X 2 X X 66 X X 7 X HCDR1-IMGT
X 69 X X 8 X X 70 X X 11 X HCDR2-IMGT X 73 X X 12 X X 74 X X 15 X HCDR3-IMGT X 77 X X 16 X X 78
X X 21 X HCDR1-Kabat X 83 X X 22 X X 84 X X 25 X HCDR2-Kabat X 87 X X 26 X X 88 X X 29 X HCDR3-
Kabat X 91 X X 30 X X 92 X X 5 X VH FR1-IMGT X 67 X X 6 X X 68 X X 9 X VH FR2-IMGT X 71 X X 10 X
X 72 X X 13 X VH FR3-IMGT X 75 X X 14 X X 76 X X 17 X VH FR4-IMGT X 79 X X 18 X X 80 X X 19 X VH
FR1-Kabat X 81 X X 20 X X 82 X X 23 X VH FR2-Kabat X 85 X X 24 X X 86 X X 27 X VH FR3-Kabat X 89 X
X 28 X X 90 X X 31 X VH FR4-Kabat X 93 X X 32 X X 94 X X 33 X VL REGION X 95 X X 34 X X 96 X X 39
X LCDR1-IMGT X 99 X X 40 X X 100 X X 43 X LCDR2-IMGT X 103 X X 44 X X 104 X X 47 X LCDR3-IMGT
X 107 X X 48 X X 108 X X 53 X LCDR1-Kabat X 113 X X 54 X X 114 X X 57 X LCDR2-Kabat X 117 X X 58 X
X 118 X X 61 X LCDR3-Kabat X 121 X X 62 X X 122 X X 37 X VL FR1-IMGT X 97 X X 38 X X 98 X X 41 X
VL FR2-IMGT X 101 X X 42 X X 102 X X 45 X VL FR3-IMGT X 105 X X 46 X X 106 X X 49 X VL FR4-IMGT
X 109 X X 50 X X 110 X X 51 X VL FR1-Kabat X 111 X X 52 X X 112 X X 55 X VL FR2-Kabat X 115 X X 56 X
X 116 X X 59 X VL FR3-Kabat X 119 X X 60 X X 120 X X 63 X VL FR4-Kabat X 123 X X 64 X X 124 X X
(468) TABLE-US-00004 CONSTANT REGION SEQUENCE SUMMARY SEQ ID NO: AMINO ACID
NUCLEOTIDE 3 IgG4-PE X 4 X 125 IGHG1*01 X 126 X 127 IGHG1*02 or X 128 IGHG1*05 X 129 IGHG1*03
X 130 X 131 IGHG1*04 X 132 X 133 Disabled human IGHG1*01 X 134 X 135 IGHG2*01 or X 136 IGHG2*04 or
X IGHG2*05 137 IGHG2*02 X 138 X 139 IGHG2*04 X 140 X 141 IGHG2*06 X 142 X 143 IGHG4*01 or X 144
IGHG4*04 X 145 IGHG4*02 X 146 X 147 IGHG4*03 X 148 X 149 IGHG4-PE X 150 X 151 X 152 X 153
Inactivated X 154 IGHG4 X 155 IGKC*01 X 156 X 157 IGKC*02 X 158 X 159 IGKC*03 X 160 X 161 IGKC*04
X 162 X 163 IGKC*05 X 164 X 165 IGLC1*01 X 166 X 167 IGLC1*02 X 168 X 169 X 170 IGLC2*01 X 171 X
172 X 173 IGLC2*02 or X 174 IGLC2*03 X 175 IGLC3*01 X 176 X 177 IGLC3*02 X 178 X 179 IGLC3*03 X
180 X 181 IGLC3*04 X 182 X 183 IGLC6*01 X 184 X 185 IGLC7*01 or X 186 IGLC7*02 X 187 IGLC7*03 X
188 X
SEQUENCES
(469) CL-148219
(470) TABLE-US-00005 CL-148219 Heavy Chain Variable region amino acid sequence (SEQ ID
NO: 1)Q V H L Q E S G P G L V K P S E T L S LT C T V S G
G S Í S S Y Y W S W I R Q YP G K G L E W I G Y I S Y S
G S S N Y NP S L K R R V T I S R D T S K N Q F S LN L T
  V I A A D T A V Y Y C A R N L MI R G A Y G M D V W
     T T V T V S S Variable region nucleotide sequence (SEQ ID NO: 2)
caggtgcacctgcaggagtcgggcccaggactggtgaagccttcggagac cctgtccctcacgtgcactgtctctggtggctccatcagtagttactact
ggagctggatccggcagtacccaggaaagggactggagtggattggatat atctcttacagtgggagcagcaattataatcccccctcaagaggcgagtc
accatatcacgagacacgtccaagaaccagttctccctgaatctgacctc tgtaatcgctgcggacacggccgtttattactgtgcgagaaatcttatga
ttcggggagcctacggcatggacgtctggggccaagggaccacggtcacc gtctcctca FR1-IMGT amino acid sequence (SEQ
ID NO: 5)Q V H L Q E S G P G L V K P S E T L S L T C T V
FR1-IMGT nucleotide sequence (SEQ ID NO: 6)
CAGGTGCACCTGCAGGAGTCGGGCCCAGGACTGGTGAAGCCTTCGGAGAC
CCTGTCCCTCACGTGCACTGTCTCT CDR1-IMGT amino acid sequence (SEQ ID NO: 7) G G
S I S S Y Y CDR1-IMGT nucleotide sequence (SEO ID NO: 8)
GGTGGCTCCATCAGTAGTTACTAC FR2-IMGT amino acid sequence (SEQ ID NO: 9) W S W
I R Q Y P G K G L E W I G Y FR2-IMGT nucleotide sequence (SEQ ID NO: 10)
TGGAGCTGGATCCGGCAGTACCCAGGAAAGGGACTGGAGTGGATTGGATA T CDR2-IMGT amino
acid sequence (SEQ ID NO: 11) I S Y S G S S CDR2-IMGT nucleotide sequence (SEQ
ID NO: 12) ATCTCTTACAGTGGGAGCAGC FR3-IMGT amino acid sequence (SEQ ID NO: 13)
N Y N P S L K R R V T I S R D T S K N Q F S L N LT S V I
  A D T A V Y Y C FR3-IMGT nucleotide sequence (SEQ ID NO: 14)
AATTATAATCCCTCCCTCAAGAGGCGAGTCACCATATCACGAGACACGTC
CAAGAACCAGTTCTCCCTGAATCTGACCTCTGTAATCGCTGCGGACACGG CCGTTTATTACTGT CDR3-
IMGT amino acid sequence (SEQ ID NO: 15) A R N L M I R G A Y G M D V
CDR3-IMGT nucleotide sequence (SEQ ID NO: 16)
```

GCGAGAAATCTTATGATTCGGGGAGCCTACGGCATGGACGTC FR4-IMGT amino acid sequence

```
(SEQ ID NO: 17) W G Q G T T V T V S S FR4-IMGT nucleotide sequence (SEQ
ID NO: 18) TGGGGCCAAGGGACCACGGTCACCGTCTCCTCA FR1-KABAT amino acid sequence
(SEQ ID NO: 19) Q V H L Q E S G P G L V K P S E T L S L T C
T V S G G S I S FR1-KABAT nucleotide sequence (SEQ ID NO: 20)
CAGGTGCACCTGCAGGAGTCGGGCCCAGGACTGGTGAAGCCTTCGGAGAC
CCTGTCCCTCACGTGCACTGTCTCTGGTGGCTCCATCAGT CDR1-KABAT amino acid sequence
(SEQ ID NO: 21) S Y Y W S CDR1-KABAT nucleotide sequence (SEQ ID NO: 22)
AGTTACTACTGGAGC FR2-KABAT amino acid sequence (SEQ ID NO: 23) W I R Q Y P
G K G L E W I G FR2-KABAT nucleotide sequence (SEQ ID NO: 24)
TGGATCCGGCAGTACCCAGGAAAGGGACTGGAGTGGATTGGA CDR2-KABAT amino acid sequence
(SEQ ID NO: 25) Y I S Y S G S S N Y N P S L K R CDR2-KABAT
nucleotide sequence (SEQ ID NO: 26)
sequence (SEQ ID NO: 27) R V T I S R D T S K N Q F S L N L T S
I A A D TA V Y Y C A R FR3-KABAT nucleotide (SEQ ID NO: 28)
CGAGTCACCATATCACGAGACACGTCCAAGAACCAGTTCTCCCTGAATCT
GACCTCTGTAATCGCTGCGGACACGGCCGTTTATTACTGTGCGAGACDR3-KABAT amino acid
sequence (SEQ ID NO: 29) N L M I R G A Y G M D V CDR3-KABAT nucleotide
sequence (SEO ID NO: 30) AATCTTATGATTCGGGGAGCCTACGGCATGGACGTC FR4-KABAT
amino acid sequence (SEQ ID NO: 31) W G Q G T T V T V S S FR4-KABAT
nucleotide sequence (SEQ ID NO: 32) TGGGGCCAAGGGACCACGGTCACCGTCTCACL-
148219 Light Chain kappa Variable region amino acid sequence (SEQ ID NO: 33) D T V
MTQSPLSLPVTPGEPASISCRSSQSLLHS
N G Y N Y L D WY L Q K A G Q S P Q L L I Y L G S N R A
S
       P D R F S G S V S G T D F T L K IS R V E A E D V
          C M Q A L Q T PF T F G P G T K V D I K Variable region
nucleotide sequence (SEQ ID NO: 34) gatactgtgatgactcagtctccactctccctgcccgtcacccctggaga
gccggcctccatctcctgcaggtctagtcagagcctcctgcatagtaatg gatacaattatttggattggtacctgcagaaggcaggacagtctccacaa
ctcctgatctatttgggttctaatcgggcctccggggtccctgacaggtt\ cagtggcagtgtatcaggcacagatttcacactgaaaatcagcagagtgg
aggctgaggatgttgggatttattactgcatgcaagctctacaaactcca ttcactttcggccctgggaccaaagtggatatcaaa C-region amino acid
sequence (SEQ ID NO: 35) R T V A A P S V F I F P P S D E Q L K S
G T A S V V C L L N N F Y P R E A K V QW K V D N A L Q
  G N S Q E S V T E Q DS K D S T Y S L S S T L T L S K A
  Y EK H K V Y A C E V T H Q G L S S P V T KS F N R G
E C C-region nucleotide sequence (SEQ ID NO: 36) cgtacggtggccgctccctccgtgttcatcttcccaccttccgacgagca
gctgaagtccggcaccgcttctgtcgtgtgcctgctgaacaacttctacc cccgcgaggccaaggtgcagtggaaaggtggacaacgccctgcagtccggc
aactcccaggaatccgtgaccgagcaggactccaaggacagcacctactc cctgtcctccaccctgaccctgtccaaggccgactacgagaagcacaagg
tgtacgcctgcgaagtgacccaccagggcctgtctagccccgtgaccaag tctttcaaccggggcgagtgt FR1-IMGT amino acid sequence
(SEQ ID NO: 37) D T V M T Q S P L S L P V T P G E P A S I S
C R S S FR1-IMGT nucleotide sequence (SEQ ID NO: 38)
GATACTGTGATGACTCAGTCTCCACTCTCCCTGCCCGTCACCCCTGGAGA
GCCGGCCTCCATCTCCTGCAGGTCTAGT CDR1-IMGT amino acid sequence (SEQ ID NO: 39) Q
S L L H S N G Y N Y CDR1-IMGT nucleotide sequence (SEQ ID NO: 40)
CAGAGCCTCCTGCATAGTAATGGATACAATTAT FR2-IMGT amino acid sequence (SEQ ID NO:
41) L D W Y L Q K A G Q S P Q L L I YFR2-IMGT nucleotide sequence
(SEO ID NO: 42) TTGGATTGGTACCTGCAGAAGGCAGGACAGTCTCCACAACTCCTGATCTA T
CDR2-IMGT amino acid sequence (SEQ ID NO: 43) L G S CDR2-IMGT nucleotide sequence
(SEQ ID NO: 44) TTGGGTTCT FR3-IMGT amino acid sequence (SEQ ID NO: 45) N R A
    V P D R F S G S V S G T D F T L K I S RV E A E D V
    Y Y C FR3-IMGT nucleotide sequence (SEQ ID NO: 46)
AATCGGGCCTCCGGGGTCCCTGACAGGTTCAGTGGCAGTGTATCAGGCAC
AGATTTCACACTGAAAATCAGCAGAGTGGAGGCTGAGGATGTTGGGATTT ATTACTGC CDR3-IMGT
amino acid sequence (SEQ ID NO: 47) M Q A L Q T P F T CDR3-IMGT nucleotide
sequence (SEQ ID NO: 48) ATGCAAGCTCTACAAACTCCATTCACT FR4-IMGT amino acid
sequence (SEQ ID NO: 49) F G P G T K V D I K FR4-IMGT nucleotide sequence
(SEQ ID NO: 50) TTCGGCCCTGGGACCAAAGTGGATATCAAA FR1-KABAT amino acid sequence
(SEQ ID NO: 51) D T V M T Q S P L S L P V T P G E P A S I S C
FR1-KABAT nucleotide sequence (SEQ ID NO: 52)
```

```
GATACTGTGATGACTCAGTCTCCACTCTCCCTGCCCGTCACCCCTGGAGA GCCGGCCTCCATCTCCTGC
CDR1-KABAT amino acid sequence (SEQ ID NO: 53) R S S Q S L L H S N G Y
N Y L D CDR1-KABAT nucleotide sequence (SEQ ID NO: 54)
AGGTCTAGTCAGAGCCTCCTGCATAGTAATGGATACAATTATTTGGAT FR2-KABAT amino acid
sequence (SEQ ID NO: 55) W Y L Q K A G Q S P Q L L I Y FR2-KABAT
nucleotide sequence (SEQ ID NO: 56)
TGGTACCTGCAGAAGGCAGGACAGTCTCCACAACTCCTGATCTAT CDR2-KABAT amino acid
sequence (SEQ ID NO: 57) L G S N R A S CDR2-KABAT nucleotide sequence (SEQ ID
NO: 58) TTGGGTTCTAATCGGGCCTCC FR3-KABAT amino acid sequence (SEQ ID NO: 59) G
V P D R F S G S V S G T D F T L K I S R V E A ED V G I Y
Y C FR3-KABAT nucleotide sequence (SEQ ID NO: 60)
GGGGTCCCTGACAGGTTCAGTGGCAGTGTATCAGGCACAGATTTCACACT
GAAAATCAGCAGAGTGGAGGCTGAGGATGTTGGGATTTATTACTGC CDR3-KABAT amino acid
sequence (SEQ ID NO: 61) M Q A L Q T P F T CDR3-KABAT nucleotide sequence
(SEQ ID NO: 62) ATGCAAGCTCTACAAACTCCATTCACT FR4-KABAT amino acid sequence
(SEQ ID NO: 63) F G P G T K V D I K FR4-KABAT nucleotide sequence (SEQ ID
NO: 64) TTCGGCCCTGGGACCAAAGTGGATATCAAA
CL-148489
(471) TABLE-US-00006 CL-148489 Heavy Chain Variable region amino acid sequence (SEQ ID
NO: 65) E V Q L V E S G G G L V Q P G R S L R LS C
  T F A D Y V M H W V R Q TP G K G L E W V S G I S
  Y S I N YA D S V K G R F T I S R D N A Q N S L YL Q M
  S L R A E D T A L Y F C A K D IT Y D L L T G Y N Y N
  G L D V W G Q G T T V T V S S Variable region nucleotide sequence (SEQ
  NO: 66) gaagtgcagctggtagagtctggggggggggcttggtacagcctggcaggtccctgagactc
tcctgtacagcctctggattcacctttgctgattatgtcatgcactgggtccggcaaact
ccagggaagggcctggagtgggtctcaggtattagttggaatagttatagtataaattat
gcggactctgtgaagggccgattcaccatctccagagacaacgcccagaactccctgtat
ctgcaaatgaacagtctgagagctgaggacacggccttgtatttttgtgcaaaagatata
acttacgatcttttgactggttataactacaactacggtttagacgtctggggccaaggg accacggtcaccgtctcctca FR1-IMGT amino acid
sequence (SEQ ID NO: 67) E V Q L V E S G G G L V Q P G R S L R L
S C T A S FR1-IMGT nucleotide sequence (SEQ ID NO: 68)
GAAGTGCAGCTGGTAGAGTCTGGGGGGGGGGCTTGGTACAGCCTGGCAGGTCCCTGAGACTCTCCTGTACAGCC
TCT CDR1-IMGT amino acid sequence (SEQ ID NO: 69) G F T F A D Y V CDR1-
IMGT nucleotide sequence (SEQ ID NO: 70) GGATTCACCTTTGCTGATTATGTC FR2-IMGT
amino acid sequence (SEQ ID NO: 71) M H W V R Q T P G K G L E W V
S G FR2-IMGT nucleotide sequence (SEQ ID NO: 72)
ATGCACTGGGTCCGGCAAACTCCAGGGAAGGGCCTGGAGTGGGTCTCAGGT CDR2-IMGT amino
acid sequence (SEQ ID NO: 73) I S W N S Y S I CDR2-IMGT nucleotide sequence
(SEQ ID NO: 74) ATTAGTTGGAATAGTTATAGTATA FR3-IMGT amino acid sequence (SEQ ID
NO: 75) N Y A D S V K G R F T I S R D N A Q N S L Y L Q M
N S L R A E D T A L Y F C FR3-IMGT nucleotide sequence (SEQ ID NO: 76)
AATTATGCGGACTCTGTGAAGGGCCGATTCACCATCTCCAGAGACACGCCCAGAACTCCCTGTATCTGCAAA
TGAACAGTCTGAGAGCTGAGGACACGGCCTTGTATTTTTGT CDR3-IMGT amino acid sequence
(SEQ ID NO: 77) A K D I T Y D L L T G Y N Y N Y G L D VCDR3-
    nucleotide sequence (SEQ ID NO: 78)
GCAAAAGATATAACTTACGATCTTTTGACTGGTTATAACTACAACTACGGTTTAGACGTC FR4-IMGT
amino acid sequence (SEQ ID NO: 79) W G Q G T T V T V S S FR4-IMGT
nucleotide sequence (SEQ ID NO: 80) TGGGGCCAAGGGACCACGGTCACCGTCTCA FR1-
KABAT amino acid sequence (SEQ ID NO: 81) E V Q L V E S G G G L V Q
P G R S L R L S C T A S G F T F A FR1-KABAT nucleotide sequence (SEQ
ID NO: 82)
GAAGTGCAGCTGGTAGAGTCTGGGGGGGGGGCTTGGTACAGCCTGGCAGGTCCCTGAGACTCTCCTGTACAGCC
TCTGGATTCACCTTTGCT CDR1-KABAT amino acid sequence (SEQ ID NO: 83) D Y V M
CDR1-KABAT nucleotide sequence (SEQ ID NO: 84) GATTATGTCATGCAC FR2-KABAT amino
acid sequence (SEQ ID NO: 85) W V R Q T P G K G L E W V S FR2-KABAT
nucleotide sequence (SEQ ID NO: 86)
TGGGTCCGGCAAACTCCAGGGAAGGGCCTGGAGTGGGTCTCACDR2-KABAT amino acid sequence
```

```
(SEQ ID NO: 87) G I S W N S Y S I N Y A D S V K G CDR2-KABAT
nucleotide sequence (SEQ ID NO: 88)
GGTATTAGTTGGAATAGTTATAGTATAAATTATGCGGACTCTGTGAAGGGC FR3-KABAT amino acid
sequence (SEQ ID NO: 89) R F T I S R D N A Q N S L Y L Q M N S L
R A E D T A L Y F C A K FR3-KABAT nucleotide sequence (SEQ ID NO: 90)
CGATTCACCATCTCCAGAGACAACGCCCAGAACTCCCTGTATCTGCAAATGAACAGTCTGAGAGCTGAGGACA
CGGCCTTGTATTTTTGTGCAAAA CDR3-KABAT amino acid sequence (SEQ ID NO: 91) D I
T Y D L L T G Y N Y N Y G L D V CDR3-KABAT nucleotide sequence (SEQ
ID NO: 92) GATATAACTTACGATCTTTTGACTGGTTATAACTACAACTACGGTTTAGACGTC FR4-
KABAT amino acid sequence (SEQ ID NO: 93) W G Q G T T V T V S S FR4-
KABAT nucleotide sequence (SEQ ID NO: 94) TGGGGCCAAGGGACCACGGTCACCGTCTCCTCA
CL-148489 Light Chain kappa Variable region amino acid sequence (SEQ ID NO: 95) D I
          T P L S L S V T P G Q P A SI S C R S S Q S L L H
S
            Y L Y WY L Q K P G Q P P Q L L I Y E V S N R F
  G V P D R F S G S G S G T D F T L K IS R V E A E D V
    Y Y C M Q S I Q L PL T F G G G T K V E I K Variable region
nucleotide sequence (SEQ ID NO: 96) gatattgtgatgacccagactccactctctgtccgtcacccctggacagccggcctcc
atctcctgcaggtctagtcagagcctcctacatagtgatggaaagacctatttgtattgg
tacctgcagaagcccggccagcctccacagctcctgatctatgaagtttccaaccgtttc
tctggagtgccagataggttcagtggcagcgggtcagggacagatttcacactgaagatc\\
amino acid sequence (SEQ ID NO: 97) D I V M T Q T P L S L S V T P G
Q P A S I S C R S S FR1-IMGT nucleotide sequence (SEQ ID NO: 98)
GATATTGTGATGACCCAGACTCCACTCTCTGTCCGTCACCCCTGGACAGCCGGCCTCCATCTCCTGCAGGTC
TAGT CDR1-IMGT amino acid sequence (SEQ ID NO: 99) Q S L L H S D G K T Y
CDR1-IMGT nucleotide sequence (SEQ ID NO: 100)
CAGAGCCTCCTACATAGTGATGGAAAGACCTAT FR2-IMGT amino acid sequence (SEQ ID NO:
101) L Y W Y L Q K P G Q P P Q L L I Y FR2-IMGT nucleotide sequence
(SEQ ID NO: 102) TTGTATTGGTACCTGCAGAAGCCCGGCCAGCCTCCACAGCTCCTGATCTAT
CDR2-IMGT amino acid sequence (SEQ ID NO: 103) E V S CDR2-IMGT nucleotide sequence
(SEQ ID NO: 104) GAAGTTTCC FR3-IMGT amino acid sequence (SEQ ID NO: 105) N R
F S G V P D R F S G S G S G T D F T L K I S R V E A E D
V G L Y Y C FR3-IMGT nucleotide sequence (SEQ ID NO: 106)
AACCGTTTCTCTGGAGTGCCAGATAGGTTCAGTGGCAGCGGGTCAGGGACAGATTTCACACTGAAGATCAGC
CGGGTGGAGGCTGAAGATGTTGGCCTTTATTACTGC CDR3-IMGT amino acid sequence (SEQ ID
NO: 107) M Q S I Q L P L T CDR3-IMGT nucleotide sequence (SEQ ID NO: 108)
ATGCAAAGTATACAGCTTCCGCTCACT FR4-IMGT amino acid sequence (SEQ ID NO: 109) F
G G G T K V E I K FR4-IMGT nucleotide sequence (SEQ ID NO: 110)
TTCGGCGGAGGACCAAGGTAGAGATCAAA FR1-KABAT amino acid sequence (SEQ ID NO:
111) D I V M T Q T P L S L S V T P G Q P A S I S CFR1-KABAT
nucleotide sequence (SEQ ID NO: 112)
GATATTGTGATGACCCAGACTCCACTCTCTCTGTCCGTCACCCCTGGACAGCCGGCCTCCATCTCCTGC
CDR1-KABAT amino acid sequence (SEQ ID NO: 113) R S S Q S L L H S D G
K T Y L Y CDR1-KABAT nucleotide sequence (SEQ ID NO: 114)
AGGTCTAGTCAGAGCCTCCTACATAGTGATGGAAAGACCTATTTGTAT FR2-KABAT amino acid
sequence (SEQ ID NO: 115) W Y L Q K P G Q P P Q L L I Y FR2-KABAT
nucleotide sequence (SEQ ID NO: 116)
TGGTACCTGCAGAAGCCCGGCCAGCCTCCACAGCTCCTGATCTAT CDR2-KABAT amino acid
sequence (SEQ ID NO: 117) E V S N R F S CDR2-KABAT nucleotide sequence (SEQ ID
NO: 118) GAAGTTTCCAACCGTTTCTCT FR3-KABAT amino acid sequence (SEQ ID NO: 119)
G V P D R F S G S G S G T D F T L K I S R V E A E D V G
L Y Y C FR3-KABAT nucleotide sequence (SEQ ID NO: 120)
GGAGTGCCAGATAGGTTCAGTGGCAGCGGGTCAGGGACAGATTTCACACTGAAGATCAGCCGGGTGGAGGC
TGAAGATGTTGGCCTTTATTACTGC CDR3-KABAT amino acid sequence (SEQ ID NO: 121) M
Q S I Q L P L T CDR3-KABAT nucleotide sequence (SEQ ID NO: 122)
ATGCAAAGTATACAGCTTCCGCTCACT FR4-KABAT amino acid sequence (SEQ ID NO: 123) F
G G G T K V E I K FR4-KABAT nucleotide sequence (SEQ ID NO: 124)
```

TTCGGCGGAGGGACCAAGGTAGAGATCAAA

```
CL-274698
(472) TABLE-US-00007 CL-274698 Heavy Chain Variable region amino acid sequence (SEQ ID
NO: 199) QVQLQESGPG LVKPSGTLSL TCAVSGGSIS SSKWWSWVRQ PPGKGLEWIG
ETHYSGSTNY NPSLKSRVTI SVDKSKNOFS LKLRSVTAAD TAVYYCARVG ATENFWGOGT
LVTVSS Variable region nucleotide sequence (SEQ ID NO: 200) CAGGTGCAGC
TGCAGGAGTC GGGCCCAGGA CTGGTGAAGC CTTCGGGGAC CCTGTCCCTC ACCTGCGCTG
TCTCTGGTGG CTCCATCAGC AGTAGTAAAT GGTGGAGTTG GGTCCGCCAG CCCCCAGGGA
AGGGGCTGGA GTGGATTGGG GAAACCCATT ATAGTGGGAG CACCAACTAC AACCCGTCCC
TCAAGAGTCG AGTCACCATA TCAGTAGACA AGTCCAAGAA CCAGTTCTCC
CTGAAGCTGA GGTCTGTGAC CGCCGCGGAC ACGGCCGTTT ATTACTGTGC GAGAGTGGGT
GCTACTGAGA ACTTCTGGGG CCAGGGAACC CTGGTCACCG TCTCCTCA FR1-IMGT amino
acid sequence (SEQ ID NO: 201) QVQLQESGPG LVKPSGTLSL TCAVS FR1-IMGT nucleotide
sequence (SEQ ID NO: 202)
CAGGTGCAGCTGCAGGAGTCGGGCCCAGGACTGGTGAAGCCTTCGGGGACCCTGTCCCTCACCTGCGCTGTC
TCT CDR1-IMGT amino acid sequence (SEQ ID NO: 203) GGSIS SSKW CDR1-IMGT
nucleotide sequence (SEQ ID NO: 204) GGTGGCTCCATCAGCAGTAGTAAATGG FR2-IMGT amino
acid sequence (SEQ ID NO: 205) WSWVRQ PPGKGLEWIG E FR2-IMGT nucleotide sequence
(SEQ ID NO: 206) TGGAGTTGGGTCCGCCAGCCCCCAGGGAAGGGGCTGGAGTGGATTGGGGAA
CDR2-IMGT amino acid sequence (SEQ ID NO: 207) THYSGST CDR2-IMGT nucleotide sequence
(SEQ ID NO: 208) ACCCATTATAGTGGGAGCACC FR3-IMGT amino acid sequence (SEQ ID
NO: 209) NY NPSLKSRVTI SVDKSKNQFS LKLRSVTAAD TAVYYC FR3-IMGT nucleotide
sequence (SEQ ID NO: 210)
AACTACAACCCGTCCCTCAAGAGTCGAGTCACCATATCAGTAGACAAGTCCAAGAACCAGTTCTCCCTGAAGC
TGAGGTCTGTGACCGCCGGGACACGGCCGTTTATTACTGT CDR3-IMGT amino acid sequence
(SEQ ID NO: 211) ARVG ATENF CDR3-IMGT nucleotide sequence (SEQ ID NO: 212)
GCGAGAGTGGGTGCTACTGAGAACTTC FR4-IMGT amino acid sequence (SEQ ID NO: 213)
WGQGT LVTVSS FR4-IMGT nucleotide sequence (SEQ ID NO: 214)
TGGGGCCAGGGAACCCTGGTCACCGTCTCCTCAG FR1-KABAT amino acid sequence (SEQ ID
NO: 215) QVQLQESGPG LVKPSGTLSL TCAVSGGSIS FR1-KABAT nucleotide sequence (SEQ ID
CAGGTGCAGCTGCAGGAGTCGGGCCCAGGACTGGTGAAGCCTTCGGGGACCCTGTCCCTCACCTGCGCTGTC
TCTGGTGGCTCCATCAGC CDR1-KABAT amino acid sequence (SEQ ID NO: 217) SSKWWS
CDR1-KABAT nucleotide sequence (SEQ ID NO: 218) AGTAGTAAATGGTGGAGT FR2-KABAT
amino acid sequence (SEQ ID NO: 219) WVRQ PPGKGLEWIG FR2-KABAT nucleotide sequence
(SEQ ID NO: 220) TGGGTCCGCCAGCCCCCAGGGAAGGGGCTGGAGTGGATTGGG CDR2-KABAT
amino acid sequence (SEQ ID NO: 221) ETHYSGSTNY NPSLKS CDR2-KABAT nucleotide
sequence (SEQ ID NO: 222) GAAACCCATTATAGTGGGAGCACCAACTACAACCCGTCCCTCAAGAGT
FR3-KABAT amino acid sequence (SEQ ID NO: 223) RVTI SVDKSKNQFS LKLRSVTAAD
TAVYYCAR FR3-KABAT nucleotide sequence (SEQ ID NO: 224)
CGGCCGTTTATTACTGTGCGAGA CDR3-KABAT amino acid sequence (SEQ ID NO: 225) VG
ATENF CDR3-KABAT nucleotide sequence (SEQ ID NO: 226) GTGGGTGCTACTGAGAACTTC FR4-
KABAT amino acid sequence (SEQ ID NO: 227) WGQGT LVTVSS FR4-KABAT nucleotide
sequence (SEQ ID NO: 228) TGGGGCCAGGGAACCCTGGTCACCGTCTCCTCAG CL-274698 Light
Chain kappa Variable region amino acid sequence (SEQ ID NO: 229) EIVLTQSPAT
LSLSPGERAT LSCRASQSVF RYLAWYQQKP GQAPRLLIYD ASTRATDIPA RFSGSGSGTD
FTLTISSLEP EDFAVYYCQQ RSNWPPTFGQ GTKVEIK Variable region nucleotide sequence (SEQ
ID NO: 230) GAAATTGTGT TGACACAGTC TCCAGCCACC CTGTCTTTGT CTCCAGGGGA
AAGAGCCACC CTCTCCTGCA GGGCCAGTCA GAGTGTTTTC AGGTACTTAG CCTGGTACCA
ACAGAAACCT GGCCAGGCTC CCAGGCTCCT CATCTATGAT GCATCCACCA GGGCCACTGA
CATCCCAGCC AGGTTCAGTG GCAGTGGGTC TGGGACAGAT TTCACTCTCA CCATCAGCAG
CCTAGAGCCT GAAGATTTTG CAGTTTATTA CTGTCAGCAA CGTAGCAACT GGCCTCCGAC
GTTCGGCCAA GGGACCAAGG TGGAAATCAA A FR1-IMGT amino acid sequence (SEQ ID
NO: 231) EIVLTQSPAT LSLSPGERAT LSCRAS FR1-IMGT nucleotide sequence (SEQ ID NO:
GAAATTGTGTTGACACAGTCTCCAGCCACCCTGTCTTTGTCTCCAGGGGAAAGAGCCACCCTCTCCTGCAGGG
```

CCAGT CDR1-IMGT amino acid sequence (SEQ ID NO: 233) QSVF RY CDR1-IMGT nucleotide sequence (SEQ ID NO: 234) CAGAGTGTTTTCAGGTAC FR2-IMGT amino acid

```
sequence (SEQ ID NO: 235) LAWYQQKP GQAPRLLIY FR2-IMGT nucleotide sequence (SEQ ID
NO: 236) TTAGCCTGGTACCAACAGAAACCTGGCCAGGCTCCCAGGCTCCTCATCTAT CDR2-IMGT
amino acid sequence (SEQ ID NO: 237) D AS CDR2-IMGT nucleotide sequence (SEQ ID
NO: 238) GATGCATCC FR3-IMGT amino acid sequence (SEQ ID NO: 239) TRATDIPA
RFSGSGSGTD FTLTISSLEP EDFAVYYC FR3-IMGT nucleotide sequence (SEQ ID NO: 240)
ACCAGGGCCACTGACATCCCAGCCAGGTTCAGTGGCAGTGGGTCTGGGACAGATTTCACTCTCACCATCAGCA
GCCTAGAGCCTGAAGATTTTGCAGTTTATTACTGT CDR3-IMGT amino acid sequence (SEQ ID
NO: 241) QQ RSNWPPT CDR3-IMGT nucleotide sequence (SEQ ID NO: 242)
CAGCAACGTAGCAACTGGCCTCCGACG FR4-IMGT amino acid sequence (SEQ ID NO: 243)
FGQ GTKVEIK FR4-IMGT nucleotide sequence (SEQ ID NO: 244)
TTCGGCCAAGGGACCAAGGTGGAAATCAAA FR1-KABAT amino acid sequence (SEQ ID NO:
245) EIVLTQSPAT LSLSPGERAT LSC FR1-KABAT nucleotide sequence (SEQ ID NO: 246)
GAAATTGTGTTGACACAGTCTCCAGCCACCCTGTCTTTGTCTCCAGGGGAAAGAGCCACCCTCTCCTGC
CDR1-KABAT amino acid sequence (SEQ ID NO: 247) RASQSVF RYLA CDR1-KABAT
nucleotide sequence (SEQ ID NO: 248) AGGGCCAGTCAGAGTGTTTTCAGGTACTTAGCC FR2-
KABAT amino acid sequence (SEQ ID NO: 249) WYQQKP GQAPRLLIY FR2-KABAT
nucleotide sequence (SEQ ID NO: 250)
TGGTACCAACAGAAACCTGGCCAGGCTCCCAGGCTCCTCATCTAT CDR2-KABAT amino acid
sequence (SEQ ID NO: 251) D ASTRAT CDR2-KABAT nucleotide sequence (SEQ ID NO: 252)
GATGCATCCACCAGGGCCACT FR3-KABAT amino acid sequence (SEQ ID NO: 253) DIPA
RFSGSGSGTD FTLTISSLEP EDFAVYYC FR3-KABAT nucleotide sequence (SEQ ID NO: 254)
GACATCCCAGCCAGGTTCAGTGGCAGTGGGTCTGGGACAGATTTCACTCTCACCATCAGCAGCCTAGAGCCTG
AAGATTTTGCAGTTTATTACTGT CDR3-KABAT amino acid sequence (SEQ ID NO: 255) QQ
RSNWPPT CDR3-KABAT nucleotide sequence (SEQ ID NO: 256)
CAGCAACGTAGCAACTGGCCTCCGACG FR4-KABAT amino acid sequence (SEQ ID NO: 257)
FGQ GTKVEIK FR4-KABAT nucleotide sequence (SEQ ID NO: 258)
TTCGGCCAAGGGACCAAGGTGGAAATCAAA
CL-274711
(473) TABLE-US-00008 CL-274711 Heavy Chain Variable region amino acid sequence (SEQ ID
NO: 259) QVQLQESGPG LVKPSQTLSL TCTVSGGSIS SGGYYWSWIR QHPGKGLEWI
GHIFYSGSTY YNPSLKSRVT ISVDTSKNQF SLKLNSVTAA DTAVYYCASE GGYYDIPDVW
GQGTTVTVSS Variable region nucleotide sequence (SEQ ID NO: 260) CAGGTGCAGC
TGCAGGAGTC GGGCCCAGGA CTGGTGAAGC CTTCACAGAC CCTGTCCCTC ACCTGCACTG
TCTCTGGTGG CTCCATCAGC AGTGGTGGTT ACTACTGGAG CTGGATCCGC CAGCACCCAG
GGAAGGGCCT GGAGTGGATT GGGCACATCT TTTACAGTGG GAGCACCTAC TACAACCCGT
CCCTCAAGAG TCGAGTTACC ATATCAGTTG ACACGTCTAA GAACCAGTTC TCCCTGAAGC
TGAACTCTGT GACTGCCGCG GACACGGCCG TGTATTACTG TGCGAGCGAG GGAGGGTATT
ACGATATTCC GGACGTCTGG GGCCAAGGGA CCACGGTCAC CGTCTCCTCA FR1-IMGT amino
acid sequence (SEQ ID NO: 261) QVQLQESGPG LVKPSQTLSL TCTVS FR1-IMGT nucleotide
sequence (SEQ ID NO: 262)
CT CDR1-IMGT amino acid sequence (SEQ ID NO: 263) GGSIS SGGYY CDR1-IMGT
nucleotide sequence (SEQ ID NO: 264) GGTGGCTCCATCAGCAGTGGTGGTTACTAC FR2-IMGT
amino acid sequence (SEO ID NO: 265) WSWIR OHPGKGLEWI GH FR2-IMGT nucleotide
sequence (SEQ ID NO: 266)
TGGAGCTGGATCCGCCAGCACCCAGGGAAGGGCCTGGAGTGGATTGGGCAC CDR2-IMGT amino
acid sequence (SEQ ID NO: 267) IFYSGST CDR2-IMGT nucleotide sequence (SEQ ID
ATCTTTTACAGTGGGAGCACC FR3-IMGT amino acid sequence (SEQ ID NO: 269) Y
YNPSLKSRVT ISVDTSKNQF SLKLNSVTAA DTAVYYC FR3-IMGT nucleotide sequence (SEQ
TACTACAACCCGTCCCTCAAGAGTCGAGTTACCATATCAGTTGACACGTCTAAGAACCAGTTCTCCCTGAAGCT
GAACTCTGTGACTGCCGCGGACACGGCCGTGTATTACTGT CDR3-IMGT amino acid sequence
(SEQ ID NO: 271) ASE GGYYDIPDV CDR3-IMGT nucleotide sequence (SEQ ID NO: 272)
GCGAGCGAGGGAGGGTATTACGATATTCCGGACGTC FR4-IMGT amino acid sequence (SEQ ID
NO: 273) W GQGTTVTVSS FR4-IMGT nucleotide sequence (SEQ ID NO: 274)
TGGGGCCAAGGGACCACGGTCACCGTCTCCTCA FR1-KABAT amino acid sequence (SEQ ID
NO: 275) QVQLQESGPG LVKPSQTLSL TCTVSGGSIS FR1-KABAT nucleotide sequence (SEQ ID
NO: 276)
```

```
\mathsf{CAGGTGCAGCTGCAGGAGTCGGGGCCCAGGACTGGTGAAGCCTTCACAGACCCTGTCCCTCACCTGCACTGTC
CTGGTGGCTCCATCAGC CDR1-KABAT amino acid sequence (SEQ ID NO: 277) SGGYYWS
CDR1-KABAT nucleotide sequence (SEQ ID NO: 278) AGTGGTGGTTACTACTGGAGC FR2-
KABAT amino acid sequence (SEQ ID NO: 279) WIR QHPGKGLEWI G FR2-KABAT
nucleotide sequence (SEQ ID NO: 280)
TGGATCCGCCAGCACCCAGGGAAGGGCCTGGAGTGGATTGGG CDR2-KABAT amino acid sequence
(SEQ ID NO: 281) HIFYSGSTY YNPSLKS CDR2-KABAT nucleotide sequence (SEQ ID NO:
282) CACATCTTTTACAGTGGGAGCACCTACTACAACCCGTCCCTCAAGAGT FR3-KABAT amino
acid sequence (SEQ ID NO: 283) RVT ISVDTSKNQF SLKLNSVTAA DTAVYYCAS FR3-
KABAT nucleotide sequence (SEQ ID NO: 284)
CGAGTTACCATATCAGTTGACACGTCTAAGAACCAGTTCTCCCTGAAGCTGAACTCTGTGACTGCCGCGGACA
CGGCCGTGTATTACTGTGCGAGC CDR3-KABAT amino acid sequence (SEQ ID NO: 285) E
GGYYDIPDV CDR3-KABAT nucleotide sequence (SEQ ID NO: 286)
GAGGGAGGTATTACGATATTCCGGACGTC FR4-KABAT amino acid sequence (SEQ ID NO: 287)
W GQGTTVTVSS FR4-KABAT nucleotide sequence (SEQ ID NO: 288)
TGGGGCCAAGGGACCACGGTCACCGTCTCCTCA CL-274711 Light Chain kappa Variable region
amino acid sequence (SEQ ID NO: 289) EIVLTQSPAT LSLSPGERAT LSCRASQSVS
NYLAWYQQKP GQAPRLLISD ASNRATGIPA RFSGSGSGTD FTLTISSLEP EDFAIYYCQQ
RSNWPLTFGG GTKVEIK Variable region nucleotide sequence (SEQ ID NO: 290)
GAAATTGTGT TGACACAGTC TCCAGCCACC CTGTCTTTGT CTCCAGGGGA AAGAGCCACC
CTCTCCTGCA GGGCCAGTCA GAGTGTTAGC AACTACTTAG CCTGGTACCA ACAGAAACCT
GGCCAGGCTC CCAGGCTCCT CATCTCTGAT GCATCCAACA GGGCCACTGG CATCCCAGCC
AGGTTCAGTG GCAGTGGGTC TGGGACAGAC TTCACTCTCA CCATCAGCAG CCTAGAGCCT
GAAGATTTTG CAATTTATTA CTGTCAGCAG CGTAGCAACT GGCCGCTCAC TTTCGGCGGA
GGGACCAAGG TGGAGATCAA A FR1-IMGT amino acid sequence (SEQ ID NO: 291)
EIVLTQSPAT LSLSPGERAT LSCRAS FR1-IMGT nucleotide sequence (SEQ ID NO: 292)
GAAATTGTGTTGACACAGTCTCCAGCCACCCTGTCTTTGTCTCCAGGGGAAAGAGCCACCCTCTCCTGCAGGG
CCAGT CDR1-IMGT amino acid sequence (SEQ ID NO: 293) QSVS NY CDR1-IMGT
nucleotide sequence (SEQ ID NO: 294) CAGAGTGTTAGCAACTAC FR2-IMGT amino acid
sequence (SEQ ID NO: 295) LAWYQQKP GQAPRLLIS FR2-IMGT nucleotide sequence (SEQ ID
NO: 236) TTAGCCTGGTACCAACAGAAACCTGGCCAGGCTCCCAGGCTCCTCATCTCT CDR2-IMGT
amino acid sequence (SEQ ID NO: 297) D AS CDR2-IMGT nucleotide sequence (SEQ ID
    298) GATGCATCC FR3-IMGT amino acid sequence (SEQ ID NO: 299) NRATGIPA
RFSGSGSGTD FTLTISSLEP EDFAIYYC FR3-IMGT nucleotide sequence (SEQ ID NO: 300)
AACAGGGCCACTGGCATCCCAGCCAGGTTCAGTGGCAGTGGGTCTGGGACAGACTTCACTCTCACCATCAGC
AGCCTAGAGCCTGAAGATTTTGCAATTTATTACTGT CDR3-IMGT amino acid sequence (SEQ ID
NO: 301) QQ RSNWPLT CDR3-IMGT nucleotide sequence (SEQ ID NO: 302)
CAGCAGCGTAGCAACTGGCCGCTCACT FR4-IMGT amino acid sequence (SEQ ID NO: 303)
FGG GTKVEIK FR4-IMGT nucleotide sequence (SEQ ID NO: 304)
TTCGGCGGAGGACCAAGGTGGAGATCAAA FR1-KABAT amino acid sequence (SEQ ID NO:
305) EIVLTQSPAT LSLSPGERAT LSC FR1-KABAT nucleotide sequence (SEQ ID NO: 306)
GAAATTGTGTTGACACAGTCTCCAGCCACCCTGTCTTTGTCTCCAGGGGAAAGAGCCACCCTCTCCTGC
CDR1-KABAT amino acid sequence (SEQ ID NO: 307) RASQSVS NYLA CDR1-KABAT
nucleotide sequence (SEQ ID NO: 308) AGGGCCAGTCAGAGTGTTAGCAACTACTTAGCC FR2-
KABAT amino acid sequence (SEQ ID NO: 309) WYQQKP GQAPRLLIS FR2-KABAT
nucleotide sequence (SEQ ID NO: 310)
TGGTACCAACAGAAACCTGGCCAGGCTCCCAGGCTCCTCATCTCT CDR2-KABAT amino acid
sequence (SEQ ID NO: 311) D ASNRAT CDR2-KABAT nucleotide sequence (SEQ ID NO: 312)
GATGCATCCAACAGGGCCACT FR3-KABAT amino acid sequence (SEQ ID NO: 313) GIPA
RFSGSGSGTD FTLTISSLEP EDFAIYYC FR3-KABAT nucleotide sequence (SEQ ID NO: 314)
GGCATCCCAGCCAGGTTCAGTGGCAGTGGGTCTGGGACAGACTTCACTCTCACCATCAGCAGCCTAGAGCCTC
AAGATTTTGCAATTTATTACTGT CDR3-KABAT amino acid sequence (SEQ ID NO: 315) QQ
RSNWPLT CDR3-KABAT nucleotide sequence (SEQ ID NO: 316)
CAGCAGCGTAGCAACTGGCCGCTCACT FR4-KABAT amino acid sequence (SEQ ID NO: 317)
FGG GTKVEIK FR4-KABAT nucleotide sequence (SEQ ID NO:
TTCGGCGGAGGGACCAAGGTGGAGATCAAA
CL-148219QLT
(474) TABLE-US-00009 CL-148219QLT Heavy Chain Variable region amino acid sequence (SEQ
```

```
ID NO: 319) QVHLQESGPG LVKPSETLSL TCTVSGGSIS SYYWSWIRQY PGKGLEWIGY
ISYSGSSNYN PSLKRRVTIS RDTSKNQFSL QLTSVIAADT AVYYCARNLM IRGAYGMDVW
GQGTTVTVSS Variable region nucleotide sequence (SEQ ID NO: 320) CAGGTGCACC
TGCAGGAGTC GGGCCCAGGA CTGGTGAAGC CTTCGGAGAC CCTGTCCCTC ACGTGCACTG
TCTCTGGTGG CTCCATCAGT AGTTACTACT GGAGCTGGAT CCGGCAGTAC CCAGGAAAGG
GACTGGAGTG GATTGGATAT ATCTCTTACA GTGGGAGCAG CAATTATAAT CCCTCCTCA
AGAGGCGAGT CACCATATCA CGAGACACGT CCAAGAACCA GTTCTCCCTG CAGCTGACCT
CTGTAATCGC TGCGGACACG GCCGTTTATT ACTGTGCGAG AAATCTTATG ATTCGGGGAG
CCTACGGCAT GGACGTCTGG GGCCAAGGGA CCACGGTCAC CGTCTCCTCA FR1-IMGT
acid sequence (SEQ ID NO: 321) QVHLQESGPG LVKPSETLSL TCTVS FR1-IMGT nucleotide
sequence (SEQ ID NO: 322)
CAGGTGCACCTGCAGGAGTCGGGCCCAGGACTGGTGAAGCCTTCGGAGACCCTGTCCCTCACGTGCACTGTC
TCT CDR1-IMGT amino acid sequence (SEQ ID NO: 323) GGSIS SYY CDR1-IMGT nucleotide
sequence (SEQ ID NO: 324) GGTGGCTCCATCAGTAGTTACTAC FR2-IMGT amino acid sequence
(SEO ID NO: 325) WSWIROY PGKGLEWIGY FR2-IMGT nucleotide sequence
TGGAGCTGGATCCGGCAGTACCCAGGAAAGGGACTGGAGTGGATTGGATAT CDR2-IMGT amino
acid sequence (SEQ ID NO: 327) ISYSGSS CDR2-IMGT nucleotide sequence (SEQ ID NO: 328)
ATCTCTTACAGTGGGAGCAGC FR3-IMGT amino acid sequence (SEQ ID NO: 329) NYN
PSLKRRVTIS RDTSKNQFSL QLTSVIAADT AVYYC FR3-IMGT nucleotide sequence (SEQ ID
NO: 330)
AATTATAATCCCTCCCTCAAGAGGCGAGTCACCATATCACGAGACACGTCCAAGAACCAGTTCTCCCTGCAGCT
GACCTCTGTAATCGCTGCGGACACGCCGTTTATTACTGT CDR3-IMGT amino acid sequence (SEQ
ID NO: 331) ARNLM IRGAYGMDV CDR3-IMGT nucleotide sequence (SEQ ID NO: 332)
GCGAGAAATCTTATGATTCGGGGAGCCTACGGCATGGACGTC FR4-IMGT amino acid sequence
(SEQ ID NO: 333) W GQGTTVTVSS FR4-IMGT nucleotide sequence (SEQ ID NO: 334)
TGGGGCCAAGGGACCACGGTCACCGTCTCCTCA FR1-KABAT amino acid sequence (SEQ ID
    335) QVHLQESGPG LVKPSETLSL TCTVSGGSI FR1-KABAT nucleotide sequence (SEQ ID
NO:
CAGGTGCACCTGCAGGAGTCGGGCCCAGGACTGGTGAAGCCTTCGGAGACCCTGTCCCTCACGTGCACTGTC
TCTGGTGGCTCCATC CDR1-KABAT amino acid sequence (SEQ ID NO: 337) S SYYWS CDR1-
KABAT nucleotide sequence (SEQ ID NO: 338) AGTAGTTACTACTGGAGC FR2-KABAT amino
    sequence (SEQ ID NO: 339) WIRQY PGKGLEWIG FR2-KABAT nucleotide sequence (SEQ
ID NO: 340) TGGATCCGGCAGTACCCAGGAAAGGGACTGGAGTGGATTGGA CDR2-KABAT amino
acid sequence (SEQ ID NO: 341) Y ISYSGSSNYN PSLKR CDR2-KABAT nucleotide sequence
KABAT amino acid sequence (SEQ ID NO: 343) RVTIS RDTSKNQFSL QLTSVIAADT
AVYYCAR FR3-KABAT nucleotide sequence (SEQ ID NO: 344)
CGAGTCACCATATCACGAGACACGTCCAAGAACCAGTTCTCCCTGCAGCTGACCTCTGTAATCGCTGCGGACA
CGGCCGTTTATTACTGTGCGAGA CDR3-KABAT amino acid sequence (SEQ ID NO: 345) NLM
IRGAYGMDV CDR3-KABAT nucleotide sequence (SEQ ID NO:
AATCTTATGATTCGGGGAGCCTACGGCATGGACGTC FR4-KABAT amino acid sequence (SEQ ID
NO: 347) W GQGTTVTVSS FR4-KABAT nucleotide sequence (SEQ ID NO: 348)
TGGGGCCAAGGGACCACGGTCACCGTCTCCTCA CL-148219QLT Light Chain kappa Variable
region amino acid sequence (SEQ ID NO: 349) DTVMTQSPLS LPVTPGEPAS ISCRSSQSLL
HSNGYNYLDW YLQKAGQSPQ LLIYLGSNRA SGVPDRFSGS VSGTDFTLKI SRVEAEDVGI
YYCMQALQTP FTFGPGTKVD IK Variable region nucleotide sequence (SEQ ID NO: 350)
GATACTGTGA TGACTCAGTC TCCACTCTCC CTGCCCGTCA CCCCTGGAGA GCCGGCCTCC
ATCTCCTGCA GGTCTAGTCA GAGCCTCCTG CATAGTAATG GATACAATTA TITGGATTGG
TACCTGCAGA AGGCAGGACA GTCTCCACAA CTCCTGATCT ATTTGGGTTC TAATCGGGCC
TCCGGGGTCC CTGACAGGTT CAGTGGCAGT GTATCAGGCA CAGATTTCAC ACTGAAAATC
AGCAGAGTGG AGGCTGAGGA TGTTGGGATT TATTACTGCATGCAAGCTCT ACAAACTCCA
TTCACTTTCG GCCCTGGGAC CAAAGTGGAT ATCAAA FR1-IMGT amino acid sequence (SEQ
ID NO: 351) DTVMTQSPLS LPVTPGEPAS ISCRSS FR1-IMGT nucleotide sequence (SEQ ID
NO: 352)
GATACTGTGATGACTCAGTCTCCACTCTCCCTGCCCGTCACCCCTGGAGAGCCGGCCTCCATCTCCTGCAGGTC
TAGT CDR1-IMGT amino acid sequence (SEQ ID NO: 353) QSLL HSNGYNY CDR1-IMGT
nucleotide sequence (SEQ ID NO: 354) CAGAGCCTCCTGCATAGTAATGGATACAATTAT FR2-IMGT
```

amino acid sequence (SEQ ID NO: 355) LDW YLQKAGQSPQ LLIY FR2-IMGT nucleotide

```
sequence (SEQ ID NO: 356)
TTGGATTGGTACCTGCAGAAGGCAGGACAGTCTCCACAACTCCTGATCTAT CDR2-IMGT amino
acid sequence (SEQ ID NO: 357) LGS CDR2-IMGT nucleotide sequence (SEQ ID NO:
TTGGGTTCT FR3-IMGT amino acid sequence (SEQ ID NO: 359) NRA SGVPDRFSGS
VSGTDFTLKI SRVEAEDVGI YYC FR3-IMGT nucleotide sequence (SEQ ID NO: 360)
AATCGGGCCTCCGGGGTCCCTGACAGGTTCAGTGGCAGTGTATCAGGCACAGATTTCACACTGAAAATCAGC
AGAGTGGAGGCTGAGGATGTTGGGATTTATTACTGC CDR3-IMGT amino acid sequence (SEQ ID
NO: 361) MQALQTP FT CDR3-IMGT nucleotide sequence (SEQ ID NO: 362)
ATGCAAGCTCTACAAACTCCATTCACT FR4-IMGT amino acid sequence (SEQ ID NO: 363)
FGPGTKVD IK FR4-IMGT nucleotide sequence (SEQ ID NO: 364)
TTCGGCCCTGGGACCAAAGTGGATATCAAA FR1-KABAT amino acid sequence DTVMTQSPLS
LPVTPGEPAS ISC FR1-KABAT nucleotide sequence (SEQ ID NO: 366)
GATACTGTGATGACTCAGTCTCCACTCTCCCTGCCCGTCACCCCTGGAGAGCCGGCCTCCATCTCCTGC
CDR1-KABAT amino acid sequence (SEQ ID NO: 367) RSSQSLL HSNGYNYLD CDR1-KABAT
nucleotide sequence (SEQ ID NO: 368)
AGGTCTAGTCAGAGCCTCCTGCATAGTAATGGATACAATTATTTGGAT FR2-KABAT amino acid
sequence (SEQ ID NO: 369) W YLQKAGQSPQ LLIY FR2-KABAT nucleotide sequence (SEQ
ID NO: 370) TGGTACCTGCAGAAGGCAGGACAGTCTCCACAACTCCTGATCTAT CDR2-KABAT
amino acid sequence (SEQ ID NO: 371) LGSNRA S CDR2-KABAT nucleotide sequence (SEQ
   NO: 372) TTGGGTTCTAATCGGGCCTCC FR3-KABAT amino acid sequence (SEQ ID NO:
373) GVPDRFSGS VSGTDFTLKI SRVEAEDVGI YYC FR3-KABAT nucleotide sequence (SEQ ID
NO: 374)
GGGGTCCCTGACAGGTTCAGTGGCAGTGTATCAGGCACAGATTTCACACTGAAAATCAGCAGAGTGGAGGCT
GAGGATGTTGGGATTTATTACTGC CDR3-KABAT amino acid sequence (SEQ ID NO: 375)
MQALQTP FT CDR3-KABAT nucleotide sequence (SEQ ID NO: 376)
ATGCAAGCTCTACAAACTCCATTCACT FR4-KABAT amino acid sequence (SEQ ID NO: 377)
FGPGTKVD IK FR4-KABAT nucleotide sequence (SEQ ID NO: 378)
TTCGGCCCTGGGACCAAAGTGGATATCAAA
(475) TABLE-US-00010 Constant Regions for Antibodies & Fragments of the Invention SEQ ID
NO: Human IGHG1*01 Human Heavy 125
gcctccaccaagggcccatcggtcttcccctggcaccctctccaagagcacctctgggggcacagcggccct IgG1 Chain Constant
gggctgcctggtcaaggactacttccccgaaccggtgacggtgtcgtggaactcaggcgccctgaccagcggc constant Region
gtgcacaccttcccggctgtcctacagtcctcaggactctactccctcagcagcgtggtgaccgtgccctccagc region (IGHG1*01)
agcttgggcacccagacctacatctgcaacgtgaatcacaagcccagcaacaccaaggtggacaagaaagtt Nucleotide
gagcccaaatcttgtgacaaaactcacacatgcccaccgtgcccagcacctgaactcctggggggaccgtcag Sequence
tcttcctcttcccccaaaacccaaggacaccctcatgatctcccggacccctgaggtcacatgcgtggtggtgg
acgtgagccacgaagaccctgaggtcaagttcaactggtacgtggacggcgtggaggtgcataatgccaaga
caaagccgcgggaggagcagtacaacagcacgtaccgggtggtcagcgtcctcaccgtcctgcaccaggact
ggctgaatggcaaggagtacaagtgcaaggtctccaacaaagccctcccagcccccatcgagaaaaccatct
ccaaagccaaagggcagccccgagaaccacaggtgtacaccctgcccccatcccgggatgagctgaccaaga
accaggtcagcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggagtgggagagcaatgg
gcagccggagaacaactacaagaccacgcctcccgtgctggactccgacggctccttcttcctctacagcaagc
tcaccgtggacaagagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgaggctctgcacaa
ccactacacgcagaagagcctctccctgtctccgggtaaa Human Heavy 126
ASTKGPSVFPLAPSSKSTSGGTAALGCLVKDYFPEPVTVSWNSGALTSGVHTFPAVLQSS Chain Constant
GLYSLSSVVTVPSSSLGTOTYICNVNHKPSNTKVDKKVEPKSCDKTHTCPPCPAPELLGGP Region
SVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNAKTKPREEQYN (IGHG1*01)
STYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTISKAKGQPREPQVYTLPPSRDE Protein
LTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKSRW Sequence
QQGNVFSCSVMHEALHNHYTQKSLSLSPGK (P01857) Human IGHG1*02 Human Heavy 127
gcctccaccaagggcccatcggtcttccccctggcaccctcctccaagagcacctctgggggcacagcggccct IgG1 or Chain Constant
gggctgcctggtcaaggactacttccccgaaccggtgacggtgtcgtggaactcaggcgccctgaccagcggc constant IGHG1*05 Region
gtgcacaccttcccggctgtcctacagtcctcaggactctactccctcagcagcgtggtgaccgtgccctccagc region (IGHG1*02 or
agcttgggcacccagacctacatctgcaacgtgaatcacaagcccagcaacaccaaggtggacaagaaagtt IGHG1*05)
gagcccaaatcttgtgacaaaactcacacatgcccaccgtgcccagcacctgaactcctggggggaccgtcag Nucleotide
tcttcctcttcccccaaaacccaaggacaccctcatgatctcccggacccctgaggtcacatgcgtggtggtgg Sequence
acgtgagccacgaagaccctgaggtcaagttcaactggtacgtggacggcgtggaggtgcataatgccaaga
caaagccgcgggaggagcagtacaacagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactg
```

```
gctgaatggcaaggagtacaagtgcaaggtctccaacaaagccctcccagcccccatcgagaaaaccatctcc
aaagccaaagggcagccccgagaaccacaggtgtacaccctgcccccatcccgggatgagctgaccaagaa
ccaggtcagcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggagtgggagagcaatggg
cagccggagaacaactacaagaccacgcctcccgtgctggactccgacggctccttcttcctctacagcaagct
caccgtggacaagagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgaggctctgcacaac
cactacacgcagaagagcctctccctgtctccgggtaaa Human Heavy 128
ASTKGPSVFPLAPSSKSTSGGTAALGCLVKDYFPEPVTVSWNSG Chain
                                                                 Constant
ALTSGVHTFPAVLQSSGLYSLSSVVTVPSSSLGTQTYICNVNHKP Region
SNTKVDKKVEPKSCDKTHTCPPCPAPELLGGPSVFLFPPKPKDT (IGHG1*02)
LMISRTPEVTCVVVDVSHEDPEVKENWYVDGVEVHNAKTKPRE Protein
EQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTIS Sequence
KAKGQPREPQVYTLPPSRDELTKNQVSLTCLVKGFYPSDIAVEW
ESNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKSRWQQGNVFS CSVMHEALHNHYTQKSLSLSPGK
Human IGHG1*03 Human Heavy 129
gcctccaccaagggcccatcggtcttccccctggcaccctcctccaagagcacctctgggggcacagcggccct IgG1 Chain Constant
gggctgcctggtcaaggactacttccccgaaccggtgacggtgtcgtggaactcaggcgccctgaccagcggc constant Region
gtgcacaccttcccggctgtcctacagtcctcaggactctactccctcagcagcgtggtgaccgtgccctccagc region (IGHG1*03)
agcttgggcacccagacctacatctgcaacgtgaatcacaagcccagcaacaccaaggtggacaagagagtt Nucleotide
gagcccaaatcttgtgacaaaactcacacatgcccaccgtgcccagcacctgaactcctggggggaccgtcag Sequence
tcttcctcttcccccaaaacccaaggacaccctcatgatctcccggacccctgaggtcacatgcgtggtggtgg (Y14737)
acgtgagccacgaagaccctgaggtcaagttcaactggtacgtggacggcgtggaggtgcataatgccaaga
caaagccgcgggaggagcagtacaacagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactg
gctgaatggcaaggagtacaagtgcaaggtctccaacaaagccctcccagccccatcgagaaaaccatctcc
aaagccaaagggcagccccgagaaccacaggtgtacaccctgccccatcccgggaggagatgaccaagaa
ccaggtcagcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggagtgggagagcaatggg
cagccggagaacaactacaagaccacgcctcccgtgctggactccgacggctccttcttcctctatagcaagct
caccgtggacaagagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgaggctctgcacaac
cactacacgcagaagagcctctccctgtccccgggtaaa Human Heavy 130
ASTKGPSVFPLAPSSKSTSGGTAALGCLVKDYFPEPVTVSWNSG Chain
                                                                 Constant
ALTSGVHTFPAVLQSSGLYSLSSVVTVPSSSLGTQTYICNVNHKP Region
SNTKVDKRVEPKSCDKTHTCPPCPAPELLGGPSVFLFPPKPKDT (IGHG1*03)
LMISRTPEVTCVVVDVSHEDPEVKENWYVDGVEVHNAKTKPRE Protein
EQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTIS Sequence
KAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE
WESNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKSRWQQGNVF SCSVMHEALHNHYTQKSLSLSPGK
Human IGHG1*04 Human 131 gcctccaccaagggcccatcggtcttccccctggcaccctcctccaagagcacctctgggggcacagcggccct
IgG1 Heavy Chain gggctgcctggtcaaggactacttccccgaaccggtgacggtgtcgtggaactcaggcgccctgaccagcggc constant
Constant gtgcacaccttcccggctgtcctacagtcctcaggactctactccctcagcagcgtggtgaccgtgccctccagc region Region
agcttgggcacccagacctacatctgcaacgtgaatcacaagcccagcaacaccaaggtggacaagaaagtt (IGHG1*04)
gagcccaaatcttgtgacaaaactcacacatgcccaccgtgcccagcacctgaactcctggggggaccgtcag Nucleotide
tcttcctcttccccccaaaacccaaggacaccctcatgatctcccggacccctgaggtcacatgcgtggtggtgg Sequence
acgtgagccacgaagaccctgaggtcaagttcaactggtacgtggacggcgtggaggtgcataatgccaaga
caaagccgcgggaggagcagtacaacagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactg
gctgaatggcaaggagtacaagtgcaaggtctccaacaaagccctcccagcccccatcgagaaaaccatctcc
aaagccaaagggcagccccgagaaccacaggtgtacaccctgcccccatcccgggatgagctgaccaagaa
ccaggtcagcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggagtgggagagcaatggg
cagccggagaacaactacaagaccacgcctcccgtgctggactccgacggctccttcttcctctacagcaagct
caccgtggacaagagcaggtggcagcaggggaacatcttctcatgctccgtgatgcatgaggctctgcacaac
cactacacgcagaagagcctctccctgtctccgggtaaa Human 132
ASTKGPSVFPLAPSSKSTSGGTAALGCLVKDYFPEPVTVSWNSGALTSGVHTFPAVLQSS Heavy Chain
GLYSLSSVVTVPSSSLGTQTYICNVNHKPSNTKVDKKVEPKSCDKTHTCPPCPAPELLGGP Constant
SVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNAKTKPREEQYN Region
STYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTISKAKGQPREPQVYTLPPSRDE (IGHG1*04)
LTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKSRW Protein
QQGNIFSCSVMHEALHNHYTQKSLSLSPGK Sequence Disabled Disabled 133
Gcctccaccaagggcccatcggtcttccccctggcaccctcctccaagagcacctctgggggcacagcggccct Human human Human
gggctgcctggtcaaggactacttccccgaaccggtgacggtgtcgtggaactcaggcgccctgaccagcggc IgG1 IGHG1*01 IGHG1*01
```

gtgcacaccttcccggctgtcctacagtcctcaggactctactccctcagcagcgtggtgaccgtgccctccagc heavy Heavy Chain

```
agcttgggcacccagacctacatctgcaacgtgaatcacaagcccagcaacaccaaggtggacaagaaagtg chain Constant
gagcccaaatcttgtgacaaaactcacacatgcccaccgtgcccagcacctgaactcgcgggggcaccgtcag constant Region
tcttcctcttccccccaaaacccaaggacaccctcatgatctcccggacccctgaggtcacatgcgtggtggtgg region Nucleotide
acgtgagccacgaagaccctgaggtcaagttcaactggtacgtggacggcgtggaggtgcataatgccaaga Sequence.
caaagccgcgggaggagcagtacaacagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactg
gctgaatggcaaggagtacaagtgcaaggtctccaacaaagccctcccagcccccatcgagaaaaccatctcc
aaagccaaagggcagccccgagaaccacaggtgtacaccctgcccccatcccgggatgagctgaccaagaa
ccaggtcagcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggagtgggagagcaatggg
cagccggagaacaactacaagaccacgcctcccgtgctggactccgacggctccttcttcctctacagcaagct
caccgtggacaagagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgaggctctgcacaac
cactacacgcagaagagcctctccctgtctccgggtaaa Disabled 134
ASTKGPSVFPLAPSSKSTSGGTAALGCLVKDYFPEPVTVSWNSGALTSGVHTFPAVLQSS Human
GLYSLSSVVTVPSSSLGTQTYICNVNHKPSNTKVDKKVEPKSCDKTHTCPPCPAPELAGA IGHG1*01
PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNAKTKPREEQY Heavy Chain
NSTYRVVSVLTVLHODWLNGKEYKCKVSNKALPAPIEKTISKAKGOPREPOVYTLPPSRD Constant
ELTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKSR Region
WQQGNVFSCSVMHEALHNHYTQKSLSLSPGK Amino Acid Sequence. Two residues that differ from
the wild-type sequence are identified in bold. Human IGHG2*01 Human 135
gcctccaccaagggcccatcggtcttccccctggcgccctgctccaggagcacctccgagagcacagccgccct IgG2 or Heavy Chain
gggctgcctggtcaaggactacttccccgaaccggtgacggtgtcgtggaactcaggcgctctgaccagcggc constant IGHG2*04 Constant
gtgcacaccttcccagctgtcctacagtcctcaggactctactccctcagcagcgtggtgaccgtgccctccagc region or Region
aacttcggcacccagacctacacctgcaacgtagatcacaagcccagcaacaccaaggtggacaagacagtt IGHG2*05 (IGHG2*01
gagcgcaaatgttgtgtcgagtgcccaccgtgcccagcaccacctgtggcaggaccgtcagtcttcctcttcccc or IGHG2*03
aagaccccgaggtccagttcaactggtacgtggacggcgtggaggtgcataatgccaagacaaagccacggg IGHG2*05)
aggagcagttcaacagcacgttccgtgtggtcagcgtcctcaccgttgtgcaccaggactggctgaacggcaa Nucleotide
ggagtacaagtgcaaggtctccaacaaaggcctcccagcccccatcgagaaaaccatctccaaaaccaaagg Sequence
gcagccccgagaaccacaggtgtacaccctgccccatcccgggaggagatgaccaagaaccaggtcagcct
gacctgcctggtcaaaggcttctaccccagcgacatcgccgtggagtgggagagcaatgggcagccggagaa
caactacaagaccacacctcccatgctggactccgacggctccttcttcctctacagcaagctcaccgtggaca
agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgaggctctgcacaaccactacacgca gaagagcctctccctgtctccgggtaaa Human
136 ASTKGPSVFPLAPCSRSTSESTAALGCLVKDYFPEPVTVSWNSGALTSGVHTFPAVLQSS Heavy
GLYSLSSVVTVPSSNFGTQTYTCNVDHKPSNTKVDKTVERKCCVECPPCPAPPVAGPSVF Constant
LFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVQFNWYVDGVEVHNAKTKPREEQFNST Region
FRVVSVLTVVHQDWLNGKEYKCKVSNKGLPAPIEKTISKTKGQPREPQVYTLPPSREEMT (IGHG2*01)
KNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTTPPMLDSDGSFFLYSKLTVDKSRW Protein
QQGNVFSCSVMHEALHNHYTQKSLSLSPGK Sequence Human IGHG2*02 Human Heavy 137
GCCTCCACCAAGGGCCCATCGGTCTTCCCCCTGGCGCCCTGCTCCAGGAGCACCTCC IgG2 Chain
Constant GAGAGCACAGCGGCCCTGGGCTGCCTGGTCAAGGACTACTTCCCCGAACCGGTGAC constant
region GGTGTCGTGGAACTCAGGCGCTCTGACCAGCGGCGTGCACACCTTCCCGGCTGTCCT Region
(IGHG2*02) ACAGTCCTCAGGACTCTACTCCCTCAGCAGCGTGGTGACCGTGACCTCCAGCAACTT
Nucleotide CGGCACCCAGACCTACACCTGCAACGTAGATCACAAGCCCAGCAACACCAAGGTGG
Sequence ACAAGACAGTTGAGCGCAAATGTTGTGTCGAGTGCCCACCGTGCCCAGCACCACCT
GTGGCAGGACCGTCAGTCTTCCTCTTCCCCCCAAAACCCAAGGACACCCTCATGATCT
CCCGGACCCCTGAGGTCACGTGCGTGGTGGTGGACGTGAGCCACGAAGACCCCGA
GGTCCAGTTCAACTGGTACGTGGACGCATGGAGGTGCATAATGCCAAGACAAAGC
CACGGGAGGAGCAGTTCAACAGCACGTTCCGTGTGGTCAGCGTCCTCACCGTCGTG
CACCAGGACTGGCTGAACGGCAAGGAGTACAAGTGCAAGGTCTCCAACAAAGGCCT
CCCAGCCCCATCGAGAAAACCATCTCCAAAACCAAAGGGCAGCCCCGAGAACCAC
AGGTGTACACCCTGCCCCCATCCCGGGAGGAGATGACCAAGAACCAGGTCAGCCTG
ACCTGCCTGGTCAAAGGCTTCTACCCCAGCGACATCGCCGTGGAGTGGGAGAGCAA
TGGGCAGCCGGAGAACAACTACAAGACCACACCTCCCATGCTGGACTCCGACGGCT
CCTTCTTCCTCTACAGCAAGCTCACCGTGGACAAGAGCAGGTGGCAGCAGGGGAAC
GTCTTCTCATGCTCCGTGATGCATGAGGCTCTGCACAACCACTACACACAGAAGAGC
CTCTCCCTGTCTCCGGGTAAA Human Heavy 138
ASTKGPSVFPLAPCSRSTSESTAALGCLVKDYFPEPVTVSWNSGALTSGVHTFPAVLQSS Chain Constant
GLYSLSSVVTVTSSNFGTQTYTCNVDHKPSNTKVDKTVERKCCVECPPCPAPPVAGPSVF Region
LFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVQFNWYVDGMEVHNAKTKPREEQFNST (IGHG2*02)
```

```
FRVVSVLTVVHQDWLNGKEYKCKVSNKGLPAPIEKTISKTKGQPREPQVYTLPPSREEMT Protein
KNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTTPPMLDSDGSFFLYSKLTVDKSRW Sequence
QQGNVFSCSVMHEALHNHYTQKSLSLSPGK Human IGHG2*04 Human Heavy 139
gcctccaccaagggcccatcggtcttccccctggcgccctgctccaggagcacctccgagagcacagcggccct IgG2 Chain Constant
gggctgcctggtcaaggactacttccccgaaccggtgacggtgtcgtggaactcaggcgctctgaccagcggc constant Region
gtgcacaccttcccagctgtcctacagtcctcaggactctactccctcagcagcgtggtgaccgtgccctccagc region (IGHG2*04)
agcttgggcacccagacctacacctgcaacgtagatcacaagcccagcaacaccaaggtggacaagacagtt Nucleotide
gagcgcaaatgttgtgtcgagtgcccaccgtgcccagcaccacctgtggcaggaccgtcagtcttcctcttcccc Sequence
aagaccccgaggtccagttcaactggtacgtggacggcgtggaggtgcataatgccaagacaaagccacggg
aggagcagttcaacagcacgttccgtgtggtcagcgtcctcaccgttgtgcaccaggactggctgaacggcaa
ggagtacaagtgcaaggtctccaacaaaggcctcccagccccatcgagaaaaccatctccaaaaccaaagg
gcagccccgagaaccacaggtgtacaccctgccccatcccgggaggagatgaccaagaaccaggtcagcct
gacctgcctggtcaaaggcttctaccccagcgacatcgccgtggagtgggagagcaatgggcagccggagaa
caactacaagaccacacctcccatgctggactccgacggctccttcttcctctacagcaagctcaccgtggaca
agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgaggctctgcacaaccactacacgca gaagagcctctccctgtctccgggtaaa Human
140 ASTKGPSVFPLAPCSRSTSESTAALGCLVKDYFPEPVTVSWNSGALTSGVHTFPAVLQSS Heavy
GLYSLSSVVTVPSSSLGTQTYTCNVDHKPSNTKVDKTVERKCCVECPPCPAPPVAGPSVF Constant
LFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVQFNWYVDGVEVHNAKTKPREEQFNST Region
FRVVSVLTVVHQDWLNGKEYKCKVSNKGLPAPIEKTISKTKGQPREPQVYTLPPSREEMT (IGHG2*04)
KNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTTPPMLDSDGSFFLYSKLTVDKSRW Protein
QQGNVFSCSVMHEALHNHYTQKSLSLSPGK Sequence Human IGHG2*06 Human 141
GCCTCCACCAAGGGCCCATCGGTCTTCCCCCTGGCGCCCTGCTCCAGGAGCACCTCC IgG2 Heavy
Chain GAGAGCACAGCGGCCCTGGGCTGCCTGGTCAAGGACTACTTCCCCGAACCGGTGAC constant
Constant GGTGTCGTGGAACTCAGGCGCTCTGACCAGCGGCGTGCACACCTTCCCGGCTGTCCT region
Region ACAGTCCTCAGGACTCTACTCCCTCAGCAGCGTGGTGACCGTGCCCTCCAGCAACTT
(IGHG2*06) CGGCACCCAGACCTACACCTGCAACGTAGATCACAAGCCCAGCAACACCAAGGTGG
Nucleotide ACAAGACAGTTGAGCGCAAATGTTGTGTCGAGTGCCCACCGTGCCCAGCACCACCT
Sequence GTGGCAGGACCGTCAGTCTTCCTCTTCCCCCAAAACCCAAGGACACCCTCATGATCT
CCCGGACCCCTGAGGTCACGTGCGTGGTGGTGGACGTGAGCCACGAAGACCCCGA
GGTCCAGTTCAACTGGTACGTGGACGGCGTGGAGGTGCATAATGCCAAGACAAAGC
CACGGGAGGAGCAGTTCAACAGCACGTTCCGTGTGGTCAGCGTCCTCACCGTCGTG
CACCAGGACTGGCTGAACGGCAAGGAGTACAAGTGCAAGGTCTCCAACAAAGGCCT
CCCAGCCCCATCGAGAAAACCATCTCCAAAAACCAAAGGGCAGCCCCGAGAACCAC
AGGTGTACACCCTGCCCCCATCCCGGGAGGAGATGACCAAGAACCAGGTCAGCCTG
ACCTGCCTGGTCAAAGGCTTCTACCCCAGCGACATCTCCGTGGAGTGGGAGAGCAA
TGGGCAGCCGGAGAACAACTACAAGACCACACCTCCCATGCTGGACTCCGACGGCT
CCTTCTTCCTCTACAGCAAGCTCACCGTGGACAAGAGCAGGTGGCAGCAGGGGAAC
GTCTTCTCATGCTCCGTGATGCATGAGGCTCTGCACAACCACTACACACAGAAGAGC
CTCTCCCTGTCTCCGGGTAAA Human Heavy 142
ASTKGPSVFPLAPCSRSTSESTAALGCLVKDYFPEPVTVSWNSGALTSGVHTFPAVLQSS Chain Constant
GLYSLSSVVTVPSSNFGTQTYTCNVDHKPSNTKVDKTVERKCCVECPPCPAPPVAGPSVF Region
LFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVQFNWYVDGVEVHNAKTKPREEQFNST (IGHG2*06)
FRVVSVLTVVHQDWLNGKEYKCKVSNKGLPAPIEKTISKTKGQPREPQVYTLPPSREEMT Protein
KNQVSLTCLVKGFYPSDISVEWESNGQPENNYKTTPPMLDSDGSFFLYSKLTVDKSRWQ Sequence
QGNVFSCSVMHEALHNHYTQKSLSLSPGK Human IGHG4*01 Human Heavy 143
gettecaceaagggeceateegtetteeeeetggegeeetgeteeaggageaceteegagageacageegeeet IgG4 or Chain Constant
gggctgcctggtcaaggactacttccccgaaccggtgacggtgtcgtggaactcaggcgccctgaccagcggc constant IGHG4*04 Region
gtgcacaccttcccggctgtcctacagtcctcaggactctactccctcagcagcgtggtgaccgtgccctccagc region (IGHG4*01 or
agcttgggcacgaagacctacacctgcaacgtagatcacaagcccagcaacaccaaggtggacaagagagtt IGHG4*04)
gagtccaaatatggtcccccatgcccatcatgcccagcacctgagttcctgggggggaccatcagtcttcctgttc Nucleotide
aggaagaccccgaggtccagttcaactggtacgtggatggcgtggaggtgcataatgccaagacaaagccgc
gggaggagcagttcaacagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggctgaacgg
caaggagtacaagtgcaaggtctccaacaaaggcctcccgtcctccatcgagaaaaccatctccaaagccaa
agggcagccccgagagccacaggtgtacaccctgccccatcccaggaggagatgaccaagaaccaggtca
gcctgacctgcctggtcaaaggcttctaccccagcgacatcgccgtggagtgggagagcaatgggcagccgga
gaacaactacaagaccacgcctcccgtgctggactccgacggctccttcttcctctacagcaggctaaccgtgg
```

```
acaagagcaggtggcaggaggggaatgtcttctcatgctccgtgatgcatgaggctctgcacaaccactacac acagaagagcctctccctgtctctgggtaaa
Human Heavy 144 ASTKGPSVFPLAPCSRSTSESTAALGCLVKDYFPEPVTVSWNSGALTSGVHTFPAVLQSS
      Constant GLYSLSSVVTVPSSSLGTKTYTCNVDHKPSNTKVDKRVESKYGPPCPSCPAPEFLGGPSVF
Region LFPPKPKDTLMISRTPEVTCVVVDVSQEDPEVQFNWYVDGVEVHNAKTKPREEQFNST (IGHG4*01)
YRVVSVLTVLHQDWLNGKEYKCKVSNKGLPSSIEKTISKAKGQPREPQVYTLPPSQEEMT Protein
KNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTTPPVLDSDGSFFLYSRLTVDKSRWQ Sequence
EGNVFSCSVMHEALHNHYTQKSLSLSLGK (P01861) Human IGHG4*02 Human Heavy 145
gettecaceaagggeceateegtetteceeetggegeeetgeteeaggageaceteeggaggacaegeegeeet IgG4 Chain Constant
gggctgcctggtcaaggactacttccccgaaccggtgacggtgtcgtggaactcaggcgccctgaccagcggc constant Region
gtgcacaccttcccggctgtcctacagtcctcaggactctactccctcagcagcgtggtgaccgtgccctccagc region (IGHG4*02)
agcttgggcacgaagacctacacctgcaacgtagatcacaagcccagcaacaccaaggtggacaagagagtt Nucleotide
gagtccaaatatggtccccgtgcccatcatgcccagcacctgagttcctggggggaccatcagtcttcctgttc Sequence
aggaagaccccgaggtccagttcaactggtacgtggatggcgtggaggtgcataatgccaagacaaagccgc
gggaggagcagttcaacagcacgtaccgtgtggtcagcgtcctcaccgtcgtgcaccaggactggctgaacgg
caaggagtacaagtgcaaggtctccaacaaaggcctcccgtcctccatcgagaaaaccatctccaaagccaa
agggcagccccgagagccacaggtgtacaccctgccccatcccaggaggagatgaccaagaaccaggtca
gcctgacctgcctggtcaaaggcttctaccccagcgacatcgccgtggagtgggagagcaatgggcagccgga
gaacaactacaagaccacgcctcccgtgctggactccgacggctccttcttcctctacagcaggctaaccgtgg
acaagagcaggtggcaggaggggaatgtcttctcatgctccgtgatgcatgaggctctgcacaaccactacac gcagaagagcctctccctgtctctgggtaaa
Human 146 ASTKGPSVFPLAPCSRSTSESTAALGCLVKDYFPEPVTVSWNSGALTSGVHTFPAVLQSS
Heavy Chain GLYSLSSVVTVPSSSLGTKTYTCNVDHKPSNTKVDKRVESKYGPPCPSCPAPEFLGGPSVF
Constant LFPPKPKDTLMISRTPEVTCVVVDVSQEDPEVQFNWYVDGVEVHNAKTKPREEQFNST Region
YRVVSVLTVVHQDWLNGKEYKCKVSNKGLPSSIEKTISKAKGQPREPQVYTLPPSQEEM (IGHG4*02)
TKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTTPPVLDSDGSFFLYSRLTVDKSRW Protein
QEGNVFSCSVMHEALHNHYTQKSLSLSLGK Sequence Human IGHG4*03 Human 147
gcttccaccaagggcccatccgtcttccccctggcgccctgctccaggagcacctccgagagcacagccgccct IgG4 Heavy Chain
gggctgcctggtcaaggactacttccccgaaccggtgacggtgtcgtggaactcaggcgccctgaccagcggc constant Constant
gtgcacaccttcccggctgtcctacagtcctcaggactctactccctcagcagcgtggtgaccgtgccctccagc region Region
agcttgggcacgaagacctacacctgcaacgtagatcacaagcccagcaacaccaaggtggacaagaggtt (IGHG4*03)
gagtccaaatatggtcccccatgcccatcatgcccagcacctgagttcctgggggggaccatcagtcttcctgttc Nucleotide
aggaagaccccgaggtccagttcaactggtacgtggatggcgtggaggtgcataatgccaagacaaagccgc
gggaggagcagttcaacagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggctgaacgg
caaggagtacaagtgcaaggtctccaacaaaggcctcccgtcctccatcgagaaaaccatctccaaagccaa
agggcagccccgagagccacaggtgtacaccctgcccccatcccaggaggagatgaccaagaaccaggtca
gcctgacctgcctggtcaaaggcttctaccccagcgacatcgccgtggagtgggagagcaatgggcagccgga
gaacaactacaagaccacgcctcccgtgctggactccgacggctccttcttcctctacagcaagctcaccgtgg
acaagagcaggtggcaggaggggaacgtcttctcatgctccgtgatgcatgaggctctgcacaaccactacac gcagaagagcctctccctgtctctgggtaaa
Human 148 ASTKGPSVFPLAPCSRSTSESTAALGCLVKDYFPEPVTVSWNSGALTSGVHTFPAVLQSS
Heavy Chain GLYSLSSVVTVPSSSLGTKTYTCNVDHKPSNTKVDKRVESKYGPPCPSCPAPEFLGGPSVF
Constant LFPPKPKDTLMISRTPEVTCVVVDVSQEDPEVQFNWYVDGVEVHNAKTKPREEQFNST Region
YRVVSVLTVLHQDWLNGKEYKCKVSNKGLPSSIEKTISKAKGQPREPQVYTLPPSQEEMT (IGHG4*03)
KNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKSRWQ Protein
EGNVFSCSVMHEALHNHYTQKSLSLSLGK Sequence Human IGHG4- Human 149
gcctccaccaagggcccatccgtcttccccctggcgccctgctccaggagcacctccgagagcacggccgccct IgG4PE PE Heavy
                                                                                 Chain
gggctgcctggtcaaggactacttccccgaaccagtgacggtgtcgtggaactcaggcgccctgaccagcggc constant Constant
gtgcacaccttcccggctgtcctacagtcctcaggactctactccctcagcagcgtggtgaccgtgccctccagc region Region
agcttgggcacgaagacctacacctgcaacgtagatcacaagcccagcaacaccaaggtggacaagagagtt (IGHG4-PE)
gagtccaaatatggtcccccatgcccaccatgcccagcgcctgaatttgaggggggaccatcagtcttcctgttc Nucleotide
cccccaaaacccaaggacactctcatgatctcccggacccctgaggtcacgtggtggtggtggacgtgagcc Sequence
aggaagaccccgaggtccagttcaactggtacgtggatggcgtggaggtgcataatgccaagacaaagccgc Version A
gggaggagcagttcaacagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggctgaacgg
agggcagccccgagagccacaggtgtacaccctgccccatcccaggaggagatgaccaagaaccaggtca
gcctgacctgcctggtcaaaggcttctaccccagcgacatcgccgtggagtgggagagcaatgggcagccgga
gaacaactacaagaccacgcctcccgtgctggactccgacggatccttcttcctctacagcaggctaaccgtgg
acaagagcaggtggcaggagggggaatgtcttctcatgctccgtgatgcatgaggctctgcacaaccactacac acagaagagcctctccctgtctctgggtaaa
```

```
Human 150 gcctccaccaagggacctagcgtgttccctctcgccccctgttccaggtccacaagcgagtccaccgctgccctc Heavy Chain
ggctgtctggtgaaagactactttcccgagcccgtgaccgtctcctggaatagcggagccctgacctccggcgt Constant
gcacacatttcccgccgtgctgcagagcagcggactgtatagcctgagcagcgtggtgaccgtgcccagctcca Region
gcctcggcaccaaaacctacacctgcaacgtggaccacaagccctccaacaccaaggtggacaagcggggg (IGHG4-PE)
ccccaaacccaaggacaccctgatgatctcccggacacccgaggtgacctgtgtggtcgtggacgtcagccag Sequence
gaggaccccgaggtgcagttcaactggtatgtggacggcgtggaggtgcacaatgccaaaaccaagcccagg Version B
gaggagcagttcaattccacctacagggtggtgagcgtgctgaccgtcctgcatcaggattggctgaacggca
aggagtacaagtgcaaggtgtccaacaagggactgcccagctccatcgagaagaccatcagcaaggctaag
ggccagccgagggagccccaggtgtataccctgcctcctagccaggaagagatgaccaagaaccaagtgtcc
ctgacctgcctggtgaagggattctacccctccgacatcgccgtggagtgggagagcaatggccagcccgaga
aagagcaggtggcaggagggcaacgtgttctcctgttccgtgatgcacgaggccctgcacaatcactacaccc agaagagcctctccctgtccctgggcaag
Human Heavy 151 gccagcaccaagggcccttccgtgttccccctggccccttgcagcaggagcacctccgaatccacagctgccct Chain
Constant gggctgtctggtgaaggactactttcccgagcccgtgaccgtgagctggaacagcggcgctctgacatccggcg Region
tccacacctttcctgccgtcctgcagtcctccggcctctactccctgtcctccgtggtgaccgtgcctagctcctcc PE) Nucleotide
ctcggcaccaagacctacacctgtaacgtggaccacaaaccctccaacaccacaaggtggacaaacgggtcgag Sequence
agcaagtacggccctccctgccctccttgtcctgcccccgagttcgaaggcggacccagcgtgttcctgttccct Version C
cctaagcccaaggacaccctcatgatcagccggacacccgaggtgacctgcgtggtggtggatgtgagccagg
aggaccctgaggtccagttcaactggtatgtggatggcgtggaggtgcacaacgccaagacaaagccccggg
aagagcagttcaactccacctacagggtggtcagcgtgctgaccgtgctgcatcaggactggctgaacggcaa
ggagtacaagtgcaaggtcagcaataagggactgcccagcagcatcgagaagaccatctccaaggctaaag
tgacctgcctggtgaagggattctacccttccgacatcgccgtggagtgggagtccaacggccagcccgagaa
gtccaggtggcaggaaggcaacgtgttcagctgctccgtgatgcacgaggccctgcacaatcactacacccag aagtccctgagcctgtccctgggaaag
Human Heavy 152 ASTKGPSVFPLAPCSRSTSESTAALGCLVKDYFPEPVTVSWNSGALTSGVHTFPAVLQSS
Chain Constant GLYSLSSVVTVPSSSLGTKTYTCNVDHKPSNTKVDKRVESKYGPPCPPCPAPEFEGGPSVF
Region (IGHG4- LFPPKPKDTLMISRTPEVTCVVVDVSQEDPEVQFNWYVDGVEVHNAKTKPREEQFNST
    Protein YRVVSVLTVLHQDWLNGKEYKCKVSNKGLPSSIEKTISKAKGQPREPQVYTLPPSQEEMT
Sequence (Amino KNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTTPPVLDSDGSFFLYSRLTVDKSRWQ
     substitution EGNVFSCSVMHEALHNHYTQKSLSLSLGK shown in BOLD) Inactivated Inactivated
Inactivated 153 gcctccaccaagggcccatccgtcttccccctggcgccctgctccaggagcacctccgagagcacggccgcct Human IGHG4
Human Heavy gggctgcctggtcaaggactacttccccgaaccagtgacggtgtcgtggaactcaggcgccctgaccagcggc IgG4 Chain
gtgcacaccttcccggctgtcctacagtcctcaggactctactccctcagcagcgtggtgaccgtgccctccagc constant Constant
agcttgggcacgaagacctacacctgcaacgtagatcacaagcccagcaacacccaaggtggacaagagagtt region Nucleotide
gagtccaaatatggtcccccatgcccaccatgcccagcgcctccagttgcggggggaccatcagtcttcctgttc Sequence
aggaagaccccgaggtccagttcaactggtacgtggatggcgtggaggtgcataatgccaagacaaagccgc
gggaggagcagttcaacagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggctgaacgg
agggcagccccgagagccacaggtgtacaccctgccccatcccaggaggagatgaccaagaaccaggtca
gcctgacctgcctggtcaaaggcttctaccccagcgacatcgccgtggagtgggagagcaatgggcagccgga
gaacaactacaagaccacgcctcccgtgctggactccgacggatccttcttcctctacagcaggctaaccgtgg
acaagagcaggtggcaggaggggaatgtcttctcatgctccgtgatgcatgaggctctgcacaaccactacac acagaagagcctctccctgtctctgggtaaa
Inactivated 154 ASTKGPSVFPLAPCSRSTSESTAALGCLVKDYFPEPVTVSWNSGALTSGVHTFPAVLQSS
      Heavy GLYSLSSVVTVPSSSLGTKTYTCNVDHKPSNTKVDKRVESKYGPPCPPCPAPPVAGGPSV
Chain FLFPPKPKDTLMISRTPEVTCVVVDVSQEDPEVQFNWYVDGVEVHNAKTKPREEQFNS Constant
TYRVVSVLTVLHQDWLNGKEYKCKVSNKGLPSSIEKTISKAKGQPREPQVYTLPPSQEEM Region
TKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTTPPVLDSDGSFFLYSRLTVDKSRW (IGHG4)
QEGNVFSCSVMHEALHNHYTQKSLSLSLGK Protein Sequence (inactivating mutations from human IgG4
shown in bold) Human IGKC*01 Human Cκ 155
cgtacggtggccgctccctccgtgttcatcttcccaccttccgacgagcagctgaagtccggcaccgcttctgtcg Ск Light Chain
tgtgcctgctgaacaacttctacccccgcgaggccaaggtgcagtggaaggtggacaacgccctgcagtccgg constant Constant
caactcccaggaatccgtgaccgagcaggactccaaggacagcacctactccctgtcctccaccctgaccctgt region Region
ccaaggccgactacgagaagcacaaggtgtacgcctgcgaagtgacccaccagggcctgtctagccccgtga (IGKC*01)
ccaagtctttcaaccggggcgagtgt Nucleotide Sequence Ск Light Chain 156
RTVAAPSVFIFPPSDEQLKSGTASVVCLLNNFYPREAKVQWKVDNALQSGNSQESVTEQ Constant
```

```
DSKDSTYSLSSTLTLSKADYEKHKVYACEVTHQGLSSPVTKSFNRGEC Region (IGKC*01) Amino Acid
Sequence Human IGKC*02 CK Light Chain 157
cgaactgtggctgcaccatctgtcttcatcttcccgccatctgatgagcagttgaaatctggaactgcctctgttg Ck Constant
tgtgcctgctgaataacttctatcccagagaggccaaagtacagtggaaggtggataacgccctccaatcgggt constant Region
aactcccaggagagtgtcacagagcaggagagcaaggacagcacctacagcctcagcagcaccctgacgct region (IGKC*02)
gagcaaagcagactacgagaaacacaaagtctacgccggcgaagtcacccatcagggcctgagctcgcccgt Nucleotide
cacaaagagcttcaacaggggagagtgt Sequence Ск Light Chain 158
RTVAAPSVFIFPPSDEQLKSGTASVVCLLNNFYPREAKVQWKVDNALQSGNSQESVTEQ Constant
ESKDSTYSLSSTLTLSKADYEKHKVYAGEVTHQGLSSPVTKSFNRGEC Region (IGKC*02) Amino
Sequence Human IGKC*03 CK Light Chain 159
cgaactgtggctgcaccatctgtcttcatcttcccgccatctgatgagcagttgaaatctggaactgcctctgttg Ck Constant
tgtgcctgctgaataacttctatcccagagaggccaaagtacagcggaaggtggataacgccctccaatcggg constant Region
taactcccaggagagtgtcacagagcaggagagcaaggacagcacctacagcctcagcagcaccctgacgct region (IGKC*03)
gagcaaagcagactacgagaaacacaaagtctacgcctgcgaagtcacccatcagggcctgagctcgcccgt Nucleotide
cacaaagagcttcaacaggggagagtgt Sequence Ck Light Chain 160
RTVAAPSVFIFPPSDEQLKSGTASVVCLLNNFYPREAKVQRKVDNALQSGNSQESVTEQ Constant
ESKDSTYSLSSTLTLSKADYEKHKVYACEVTHQGLSSPVTKSFNRGEC Region (IGKC*03) Amino
Sequence Human IGKC*04 Ck Light Chain 161
cgaactgtggctgcaccatctgtcttcatcttcccgccatctgatgagcagttgaaatctggaactgcctctgttg Ck Constant
tgtgcctgctgaataacttctatcccagagaggccaaagtacagtggaaggtggataacgccctccaatcgggt constant Region
aactcccaggagagtgtcacagagcaggacagcaaggacagcacctacagcctcagcagcaccctgacgctg region (IGKC*04)
agcaaagcagactacgagaaacacaaactctacgcctgcgaagtcacccatcagggcctgagctcgcccgtc Nucleotide
acaaagagcttcaacaggggagagtgt Sequence Ск Light 162
RTVAAPSVFIFPPSDEQLKSGTASVVCLLNNFYPREAKVQWKVDNALQSGNSQESVTEQ Chain
DSKDSTYSLSSTLTLSKADYEKHKLYACEVTHQGLSSPVTKSFNRGEC Constant Region (IGKC*04) Amino
Acid Sequence Human IGKC*05 Ck Light 163
cgaactgtggctgcaccatctgtcttcatcttcccgccatctgatgagcagttgaaatctggaactgcctctgttg Ck Chain
tgtgcctgctgaataacttctatcccagagaggccaaagtacagtggaaggtggataacgccctccaatcgggt constant Constant
aactcccaggagagtgtcacagagcaggacagcaaggacagcacctacagcctcagcaacaccctgacgctg region Region
agcaaagcagactacgagaaacacaaagtctacgcctgcgaagtcacccatcagggcctgagctcgcccgtc (IGKC*05)
acaaagagcttcaacaggggagagtgc Nucleotide Sequence Ск Light 164
RTVAAPSVFIFPPSDEQLKSGTASVVCLLNNFYPREAKVQWKVDNALQSGNSQESVTEQ Chain
DSKDSTYSLSNTLTLSKADYEKHKVYACEVTHQGLSSPVTKSFNRGEC Constant Region (IGKC*05)
Amino Acid Sequence Human IGLC1*01 Cλ Light 165
cccaaggccaaccccacggtcactctgttcccgccctcctctgaggagctccaagccaacaaggccacactagt Cλ Chain
gtgtctgatcagtgacttctacccgggagctgtgacagtggcttggaaggcagatggcagccccgtcaaggcg constant Constant
ggagtggagacgaccaaaccctccaaacagagcaacaacaagtacgcggccagcagctacctgagcctgac region Region
gcccgagcagtggaagtcccacagaagctacagctgccaggtcacgcatgaagggagcaccgtggagaaga (IGLC1*01)
cagtggcccctacagaatgttca Nucleotide Sequence (ENST000003 90321.2) Cλ Light 166
PKANPTVTLFPPSSEELQANKATLVCLISDFYPGAVTVAWKADGSPVKAGVETTKPSKQS Chain
NNKYAASSYLSLTPEQWKSHRSYSCQVTHEGSTVEKTVAPTECS Constant Region (IGLC1*01) Amino Acid
Sequence (A0A075B6K8) Human IGLC1*02 Cλ Light 167
ggtcagcccaaggccaaccccactgtcactctgttcccgccctcctctgaggagctccaagccaacaaggccac Cλ Chain
actagtgtgtctgatcagtgacttctacccgggagctgtgacagtggcctggaaggcagatggcagccccgtca constant Constant
aggcgggagtggagaccaccaaaccctccaaacagagcaacaacaagtacgcggccagcagctacctgagc region Region
ctgacgcccgagcagtggaagtcccacagaagctacagctgccaggtcacgcatgaagggagcaccgtggag (IGLC1*02)
aagacagtggcccctacagaatgttca Nucleotide Sequence Version A Cλ Light 168
ggtcagcccaaggccaaccccactgtcactctgttcccgccctcctctgaggagctccaagccaacaaggccac Chain
actagtgtgtctgatcagtgacttctacccgggagctgtgacagtggcctggaaggcagatggcagccccgtca Constant
aggcgggagtggagaccaccaaaccctccaaacagagcaacaacaagtacgcggccagcagctacctgagc Region
ctgacgcccgagcagtggaagtcccacagaagctacagctgccaggtcacgcatgaagggagcaccgtggag (IGLC1*02)
aagacagtggcccctacagaatgttca Nucleotide Sequence 169
GQPKANPTVTLFPPSSEELQANKATLVCLISDFYPGAVTVAWKADGSPVKAGVETTKPS Version B
KQSNNKYAASSYLSLTPEQWKSHRSYSCQVTHEGSTVEKTVAPTECS Cλ Light Chain Constant Region
(IGLC1*02) Amino Acid Sequence Human IGLC2*01 Cλ Light 170
ggccagcctaaggccgctccttctgtgaccctgttccccccatcctccgaggaactgcaggctaacaaggccac Cλ Chain
cctcgtgtgcctgatcagcgacttctaccctggcgccgtgaccgtggcctggaaggctgatagctctcctgtgaa constant Constant
ggccggcgtggaaaccaccacccttccaagcagtccaacaacaacaactacgccgcctcctcctacctgtccctga region Region
```

```
cccctgagcagtggaagtcccaccggtcctacagctgccaagtgacccacgagggctccaccgtggaaaaga (IGLC2*01)
ccgtggctcctaccgagtgctcc Nucleotide Sequence Version A Cλ Light 171
ggccagcctaaagctgcccccagcgtcaccctgtttcctcctccagcgaggagctccaggccaacaaggcca Chain
ccctcgtgtgcctgatctccgacttctatcccggcgctgtgaccgtggcttggaaagccgactccagccctgtca Constant
aagccggcgtggagaccaccaccaccctccaagcagtccaacaacaagtacgccgcctccagctatctctccct Region
gacccctgagcagtggaagtcccaccggtcctactcctgtcaggtgacccacgagggctccaccgtggaaaag (IGLC2*01)
accgtcgccccaccgagtgctcc Nucleotide Sequence Version B Cλ Light 172
GQPKAAPSVTLFPPSSEELQANKATLVCLISDFYPGAVTVAWKADSSPVKAGVETTTPSK Chain
QSNNKYAASSYLSLTPEQWKSHRSYSCQVTHEGSTVEKTVAPTECS Constant Region (IGLC1*02) Amino
Acid Sequence Human IGLC2*02 Cλ Light 173
ggtcagcccaaggctgcccctcggtcactctgttcccgccctcctctgaggagcttcaagccaacaaggccac Cλ or Chain
actggtgtgtctcataagtgacttctacccgggagccgtgacagtggcctggaaggcagatagcagccccgtca constant IGLC2*03 Constant
aggcgggagtggagaccaccacaccctccaaacaagcaacaacaagtacgcggccagcagctatctgagc region Region
ctgacgcctgagcagtggaagtcccacagaagctacagctgccaggtcacgcatgaagggagcaccgtggag (IGLC2*02
aagacagtggcccctacagaatgttca or IGLC2*03) Nucleotide Sequence Cλ Light 174
GQPKAAPSVTLFPPSSEELQANKATLVCLISDFYPGAVTVAWKADSSPVKAGVETTTPSK Chain
QSNNKYAASSYLSLTPEQWKSHRSYSCQVTHEGSTVEKTVAPTECS Constant Region (IGLC2*02) Amino
Acid Sequence Human IGLC3*01 Cλ Light 175
cccaaggctgcccctcggtcactctgttcccacctcctctgaggagcttcaagccaacaaggccacactggt Cλ Chain
gtgtctcataagtgacttctacccgggagccgtgacagttgcctggaaggcagatagcagccccgtcaaggcg constant Constant
ggggtggagaccaccacaccctccaaacaaagcaacaacaagtacgcggccagcagctacctgagcctgac region Region
gcctgagcagtggaagtcccacaaaagctacagctgccaggtcacgcatgaagggagcaccgtggagaaga (IGLC3*01) cagttgcccctacggaatgttca
Nucleotide Sequence Cλ Light 176
PKAAPSVTLFPPSSEELQANKATLVCLISDFYPGAVTVAWKADSSPVKAGVETTTPSKQS Chain
NNKYAASSYLSLTPEQWKSHKSYSCQVTHEGSTVEKTVAPTECS Constant Region (IGLC3*01) Amino
Sequence Human IGLC3*02 Cλ Light 177
ggtcagcccaaggctgcccctcggtcactctgttcccaccctcctctgaggagcttcaagccaacaaggccac Cλ Chain
actggtgtgtctcataagtgacttctacccgggggccagtgacagttgcctggaaggcagatagcagccccgtca constant Constant
ctgacgcctgagcagtggaagtcccacaaaagctacagctgccaggtcacgcatgaagggagcaccgtggag (IGLC3*02)
aagacagtggcccctacggaatgttca Nucleotide Sequence Cλ Light 178
GQPKAAPSVTLFPPSSEELQANKATLVCLISDFYPGPVTVAWKADSSPVKAGVETTTPSK Chain
QSNNKYAASSYLSLTPEQWKSHKSYSCQVTHEGSTVEKTVAPTECS Constant Region (IGLC1*02) Amino
Acid Sequence Human IGLC3*03 Cλ Light 179
ggtcagcccaaggctgcccctcggtcactctgttcccaccctctctgaggagcttcaagccaacaaggccac Cλ Chain
actggtgtgtctcataagtgacttctacccgggagccgtgacagtggcctggaaggcagatagcagccccgtca constant Constant
aggcgggagtggagaccaccaccaccctccaaacaagcaacaacaagtacgcggccagcagctacctgagc region Region
ctgacgcctgagcagtggaagtcccacaaaagctacagctgccaggtcacgcatgaagggagcaccgtggag (IGLC3*03)
aagacagtggcccctacagaatgttca Nucleotide Sequence Cλ Light 180
GQPKAAPSVTLFPPSSEELQANKATLVCLISDFYPGAVTVAWKADSSPVKAGVETTTPSK Chain
QSNNKYAASSYLSLTPEQWKSHKSYSCQVTHEGSTVEKTVAPTECS Constant Region (IGLC3*03) Amino
Acid Sequence Human IGLC3*04 Cλ Light 181
ggtcagcccaaggctgcccctcggtcactctgttcccgccctcctctgaggagcttcaagccaacaaggccac Cλ Chain
actggtgtgtctcataagtgacttctacccgggagccgtgacagtggcctggaaggcagatagcagccccgtca constant Constant
aggcgggagtggagaccaccacaccctccaaacaagcaacaacaagtacgcggccagcagctacctgagc region Region
ctgacgcctgagcagtggaagtcccacagaagctacagctgccaggtcacgcatgaagggagcaccgtggag (IGLC3*04)
aagacagtggcccctacagaatgttca Nucleotide Sequence Cλ Light 182
GQPKAAPSVTLFPPSSEELQANKATLVCLISDFYPGAVTVAWKADSSPVKAGVETTTPSK Chain
QSNNKYAASSYLSLTPEQWKSHRSYSCQVTHEGSTVEKTVAPTECS Constant Region (IGLC3*04) Amino
Acid Sequence Human IGLC6*01 Cλ Light 183
ggtcagcccaaggctgccccatcggtcactctgttcccgccctcctctgaggagcttcaagccaacaaggccac Cλ Chain
actggtgtgcctgatcagtgacttctacccgggagctgtgaaagtggcctggaaggcagatggcagccccgtc constant Constant
aacacgggagtggagaccaccacaccctccaaacagagcaacaacaagtacgcggccagcagctacctgag region Region
cctgacgcctgagcagtggaagtcccacagaagctacagctgccaggtcacgcatgaagggagcaccgtgga (IGLC6*01)
gaagacagtggcccctgcagaatgttca Nucleotide Sequence C\(\lambda\) Light 184
GQPKAAPSVTLFPPSSEELQANKATLVCLISDFYPGAVKVAWKADGSPVNTGVETTTPSK Chain
QSNNKYAASSYLSLTPEQWKSHRSYSCQVTHEGSTVEKTVAPAECS Constant Region (IGLC6*01) Amino
Acid Sequence Human IGLC7*01 Cλ Light 185
```

```
ggtcagcccaaggctgccccatcggtcactctgttcccaccctctctgaggagcttcaagccaacaaggccac Cλ or Chain
actggtgtgtctcgtaagtgacttctacccgggagccgtgacagtggcctggaaggcagatggcagccccgtca constant IGLC7*02 Constant
aggtgggagtggagaccaccaaaccctccaaacaaagcaacaagtatgcggccagcagctacctgagc region Region
ctgacgcccgagcagtggaagtcccacagaagctacagctgccgggtcacgcatgaagggagcaccgtggag (IGLC7*01
aagacagtggcccctgcagaatgctct or IGLC7*02) Nucleotide Sequence Cλ Light 186
GQPKAAPSVTLFPPSSEELQANKATLVCLVSDFYPGAVTVAWKADGSPVKVGVETTKPS Chain
KQSNNKYAASSYLSLTPEQWKSHRSYSCRVTHEGSTVEKTVAPAECS Constant Region (IGLC7*01) Amino
Acid Sequence Human IGLC7*03 Cλ Light 187
GGTCAGCCCAAGGCTGCCCCCTCGGTCACTCTGTTCCCACCCTCCTGAGGAGCTTC Cλ Chain
AAGCCAACAAGGCCACACTGGTGTCTCGTAAGTGACTTCAACCCGGGAGCCGTG constant Constant
ACAGTGGCCTGGAAGGCAGATGGCAGCCCCGTCAAGGTGGGAGTGGAGACCACCA region Region
AACCCTCCAAACAAGCAACAACAAGTATGCGGCCAGCAGCTACCTGAGCCTGACG (IGLC7*03)
CCCGAGCAGTGGAAGTCCCACAGAAGCTACAGCTGCCGGGTCACGCATGAAGGGA Nucleotide
GCACCGTGGAGAAGACAGTGGCCCCTGCAGAATGCTCT Sequence C\u03b1 Light 188
GQPKAAPSVTLFPPSSEELQANKATLVCLVSDFNPGAVTVAWKADGSPVKVGVETTKPS Chain
KQSNNKYAASSYLSLTPEQWKSHRSYSCRVTHEGSTVEKTVAPAECS Constant Region (IGLC7*03) Amino
Acid Sequence
(476) TABLE-US-00011 C region amino
                                acid sequence
                                             (SEQ ID NO:
                                                          3) A S T
                                                                          P
                                                                            E
S
  V F P L A P C S R S
                             Τ
                                S
                                   ES T A A L G
                                                      C
                                                       L V K D
           SW N S G A L
                             Τ
                                S G V
                                        H T F P A V L Q
                                                               S
                                 G
L
                  P S
                        S
                          S
                             L
                                   Τ
                                      K
                                         TY T C N
                                                     V D H K
     D K R V E SK Y G P
K
                                P
                                   C
                                      P
                                         P
                                           C P A P E F
                                                           Ε
                                                               G
                                                                 G
             P K D T L M I S
                                  R
                                      T P E V TC V V D
                                                                V
           Q F N W Y V DG V E V H N A K T K P R E E Q F
D
S
  Τ
            V S V L
                       T V L H Q D W L N G K E Y
                                                             KC
             S I E K T I S K A KG Q P R E P Q V Y T
K
  G
                                            G F Y P S D I A V EW E
Q
  E
             KN Q V S L T C L V K
             N N Y K T T P P V L D SD G S F F L Y S R L
Ν
  G Q P
V
           R W Q E GN V F S C S V M H E A L H N H Y
S L S L S L G K C region nucleotide sequence
                                             (SEQ ID
                                                     NO:
gccagcaccaagggcccttccgtgttccccctggccccttgcagcaggagcacctccgaa
tccacagctgccctgggctgtctggtgaaggactactttcccgagcccgtgaccgtgagc
tggaacagcggcgctctgacatccggcgtccacacctttcctgccgtcctgcagtcctcc
tacacctgtaacgtggaccacaaaccctccaacaccaaggtggacaaacgggtcgagagc
aagtacggccctccctgccctcttgtcctgccccgagttcgaaggcggacccagcgtg
ttcctgttccctcctaagcccaaggacaccctcatgatcagccggacacccgaggtgacc
tgcgtggtggtggatgtgagccaggaggaccctgaggtccagttcaactggtatgtggat
ggcgtggaggtgcacaacgccaagacaaagccccgggaagagcagttcaactccacctac
agggtggtcagcgtgctgaccgtgctgcatcaggactggctgaacggcaaggagtacaag
tgcaaggtcagcaataagggactgcccagcagcatcgagaagaccatctccaaggctaaa
aaccaggtgagcctgacctgcctggtgaagggattctacccttccgacatcgccgtggag
tgggagtccaacggccagcccgagaacaattataagaccacccctcccgtcctcgacagc
gacggatccttctttctgtactccaggctgaccgtggataagtccaggtggcaggaaggc
aacgtgttcagctgctccgtgatgcacgaggccctgcacaatcactacacccagaagtcc ctgagcctgtccctgggaaag >PCSK9 reference
       (SEQ ID NO: 189) QEDEDGDYEELVLALRSEEDGLAEAPEHGT
TATFHRCAKDPWRLPGTYVVVLKEETHLSQSERTARRLQAQAARRGYLTKILHVFHGLLP
GFLVKMSGDLLELALKLPHVDYIEEDSSVFAQSIPWNLERITPPRYRADEYQPPDGGSLV
EVYLLDTSIQSDHREIEGRVMVTDFENVPEEDGTRFHRQASKCDSHGTHLAGVVSGRDAG
VAKGASMRSLRVLNCQGKGTVSGTLIGLEFIRKSQLVQPVGPLVVLLPLAGGYSRVLNAA
CQRLARAGVVLVTAAGNFRDDACLYSPASAPEVITVGATNAQDQPVTLGTLGTNFGRCVD
LFAPGEDIIGASSDCSTCFVSQSGTSQAAAHVAGIAAMMLSAEPELTLAELRQRLIHFSA
KDVINEAWFPEDQRVLTPNLVAALPPSTHGAGWQLFCRTVWSAHSGPTRMATAVARCAPD
EELLSCSSFSRSGKRRGERMEAQGGKLVCRAHNAFGGEGVYAIARCCLLPQANCSVHTAP
PAEASMGTRVHCHQQGHVLTGCSSHWEVEDLGTHKPPVLRPRGQPNQCVGHREASIHASC
CHAPGLECKVKEHGIPAPQEQVTVACEEGWTLTGCSALPGTSHVLGAYAVDNTCVVRSRD
VSTTGSTSEGAVTAVAICCRSRHLAQASQELQ > PCSK9 variant a (SEQ ID NO: 190)
```

```
TATFHRCAKDPWRLPGTYVVVLKEETHLSQSERTARRLQAQAARRGYLTKILHVFHGLLP
GFLVKMSGDLLELALKLPHVDYIEEDSSVFAQSIPWNLERITPPRYRADEYQPPDGGSLV
EVYLLDTSIOSDHREIEGRVMVTDFENVPEEDGTRFHROASKCDSHGTHLAGVVSGRDAG
VAKGASMRSLRVLNCQGKGTVSGTLIGLEFIRKSQLVQPVGPLVVLLPLAGGYSRVLNAA
CORLARAGVVLVTAAGNFRDDACLYSPASAPEVITVGATNAQDQPVTLGTLGTNFGRCVD
LFAPGEDIIGASSDCSTCFVSQSGTSQAAAHVAGIAAMMLSAEPELTLAELRQRLIHFSA
KDVINEAWFPEDQRVLTPNLVAALPPSTHGAGWQLFCRTVWSAHSGPTRMATAIARCAPD
EELLSCSSFSRSGKRRGERMEAQGGKLVCRAHNAFGGEGVYAIARCCLLPQANCSVHTAP
PAEASMGTRVHCHQQGHVLTGCSSHWEVEDLGTHKPPVLRPRGQPNQCVGHREASIHASC
CHAPGLECKVKEHGIPAPQEQVTVACEEGWTLTGCSALPGTSHVLGAYAVDNTCVVRSRD
VSTTGSTSEEAVTAVAICCRSRHLAQASQELQ > PCSK9 variant b (SEQ ID NO: 191)
QEDEDGDYEELVLA
LRSEEDGLVEAPEHGTTATFHRCAKDPWRLPGTYVVVLKEETHLSQSERTARRLQAQA
ARRGYLTKILHVFHGLLPGFLVKMSGDLLELALKLPHVDYIEEDSSVFAOSIPWNLER
ITPPRYRADEYQPPDGGSLVEVYLLDTSIQSDHREIEGRVMVTDFENVPEEDGTRFHR
OASKCDSHGTHLAGVVSGRDAGVAKGASMRSLRVLNCOGKGTVSGTLIGLEFIRKSOL
VQPVGPLVVLLPLAGGYSRVLNAACQRLARAGVVLVTAAGNFRDDACLYSPASAPEVI
TVGATNAQDQPVTLGTLGTNFGRCVDLFAPGEDIIGASSDCSTCFVSQSGTSQAAAHV
AGIAAMMLSAEPELTLAELRQRLIHFSAKDVINEAWFPEDQRVLTPNLVAALPPSTHG
AGWQLFCRTVWSAHSGPTRMATAIARCAPDEELLSCSSFSRSGKRRGERMEAQGGKLV
CRAHNAFGGEGVYAIARCCLLPQANCSVHTAPPAEASMGTRVHCHQQGHVLTGCSSHW
EVEDLGTHKPPVLRPRGQPNQCVGHREASIHASCCHAPGLECKVKEHGIPAPQEQVTV
ACEEGWTLTGCSALPGTSHVLGAYAVDNTCVVRSRDVSTTGSTSEEAVTAVAICCRSR HLAQASQELQ
>PCSK9 GOF mutant (SEQ ID NO: 192) QEDEDGDYEELVLA
LRSEEDGLAEAPEHGTTATFHRCAKDPWRLPGTYVVVLKEETHLSQSERTARRLQAQA
ARRGYLTKILHVFHGLLPGFLVKMSGDLLELALKLPHVDYIEEDSSVFAQSIPWNLER
ITPPRYRADEYOPPDGGSLVEVYLLDTSIOSDHREIEGRVMVTDFENVPEEDGTRFHR
QASKCDSHGTHLAGVVSGRDAGVAKGASMRSLRVLNCQGKGTVSGTLIGLEFIRKSQL
VQPVGPLVVLLPLAGGYSRVLNAACQRLARAGVVLVTAAGNFRDDACLYSPASAPEVI
TVGATNAQDQPVTLGTLGTNFGRCVDLFAPGEDIIGASSYCSTCFVSQSGTSQAAAHV
AGIAAMMLSAEPELTLAELRQRLIHFSAKDVINEAWFPEDQRVLTPNLVAALPPSTHG
AGWQLFCRTVWSAHSGPTRMATAVARCAPDEELLSCSSFSRSGKRRGERMEAQGGKLV
CRAHNAFGGEGVYAIARCCLLPQANCSVHTAPPAEASMGTRVHCHQQGHVLTGCSSHW
EVEDLGTHKPPVLRPRGQPNQCVGHREASIHASCCHAPGLECKVKEHGIPAPQEQVTV
ACEEGWTLTGCSALPGTSHVLGAYAVDNTCVVRSRDVSTTGSTSEGAVTAVAICCRSR HLAQASQELQ
                  (SEQ ID NO: 193) QEDEDGDYEELVLALRSEEDGLADAPEHGA
>PCSK9 cyno monkey
TATFHRCAKDPWRLPGTYVVVLKEETHRSQSERTARRLQAQAARRGYLTKILHVFHHLLP
GFLVKMSGDLLELALKLPHVDYIEEDSSVFAQSIPWNLERITPARYRADEYQPPKGGSLV
EVYLLDTSIQSDHREIEGRVMVTDFESVPEEDGTRFHRQASKCDSHGTHLAGVVSGRDAG
VAKGAGLRSLRVLNCQGKGTVSGTLIGLEFIRKSQLVQPVGPLVVLLPLAGGYSRVFNAA
CQRLARAGVVLVTAAGNFRDDACLYSPASAPEVITVGATNAQDQPVTLGTLGTNFGRCVD
LFAPGEDIIGASSDCSTCFVSRSGTSQAAAHVAGIAAMMLSAEPELTLAELRQRLIHFSA
KDVINEAWFPEDQRVLTPNLVAALPPSTHRAGWQLFCRTVWSAHSGPTRMATAVARCAQD
EELLSCSSFSRSGKRRGERIEAQGGKRVCRAHNAFGGEGVYAIARCCLLPQVNCSVHTAP
PAGASMGTRVHCHOOGHVLTGCSSHWEVEDLGTHKPPVLRPRGOPNOCVGHREASIHASC
CHAPGLECKVKEHGIPAPQEQVIVACEDGWTLTGCSALPGTSHVLGAYAVDNTCVVRSRD
VSTTGSTSEEAVAAVAICCRSRHLVQASQELQ >PCSK9 mouse (SEQ ID NO: 194)
QDEDGDYEELMLALPSQEDGLADEAA
HVATATFRRCSKEAWRLPGTYIVVLMEETQRLQIEQTAHRLQTRAARRGYVIKVLHIFYD
LFPGFLVKMSSDLLGLALKLPHVEYIEEDSFVFAQSIPWNLERIIPAWHQTEEDRSPDGS
SQVEVYLLDTSIQGAHREIEGRVTITDFNSVPEEDGTRFHRQASKCDSHGTHLAGVVSGR
DAGVAKGTSLHSLRVLNCQGKGTVSGTLIGLEFIRKSQLIQPSGPLVVLLPLAGGYSRIL
NAACRHLARTGVVLVAAAGNFRDDACLYSPASAPEVITVGATNAQDQPVTLGTLGTNFGR
CVDLFAPGKDIIGASSDCSTCFMSQSGTSQAAAHVAGIVARMLSREPTLTLAELRQRLIH
FSTKDVINMAWFPEDQQVLTPNLVATLPPSTHETGGQLLCRTVWSAHSGPTRTATATARC
APEEELLSCSSFSRSGRRRGDWIEAIGGQQVCKALNAFGGEGVYAVARCCLVPRANCSIH
NTPAARAGLETHVHCHQKDHVLTGCSFHWEVEDLSVRRQPALRSRRQPGQCVGHQAASVY
```

QEDEDGDYEELVLALRSEEDGLAEAPEHGT

```
ASCCHAPGLECKIKEHGISGPSEQVTVACEAGWTLTGCNVLPGASLTLGAYSVDNLCVAR
VHDTARADRTSGEATVAAAICCRSRPSAKASWVQ Alirocumab HC (human IgG4) (SEQ ID
195) EVQLVESGGGLVQPGGSLRLSCAASGFTFNNYAMNWVRQ
APGKGLDWVSTISGSGGTTNYADSVKGRFIISRDSSKHTLYLOMNSLRAEDTAVYYCAKD
SNWGNFDLWGRGTLVTVSSASTKGPSVFPLAPCSRSTSESTAALGCLVKDYFPEPVTVSW
NSGALTSGVHTFPAVLOSSGLYSLSSVVTVPSSSLGTKTYTCNVDHKPSNTKVDKRVESKYG
PPCPPCPAPEFEGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSQEDPEVQFNWYVDGVE
VHNAKTKPREEOFNSTYRVVSVLTVLHODWLNGKEYKCKVSNKGLPSSIEKTISKAKGOPR
EPQVYTLPPSQEEMTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTTPPVLDSDGSFF
LYSRLTVDKSRWQEGNVFSCSVMHEALHNHYTQKSLSLSLGK Alirocumab LC (human
(SEQ ID NO: 196) DIVMTQSPDSLAVSLGERATINCKSSQSVLYRSNNRNFLGWY
QQKPGQPPNLLIYWASTRESGVPDRFSGSGSGTDFTLTISSLQAEDVAVYYCQQYYTTPYTF
GQGTKLEIKRTVAAPSVFIFPPSDEQLKSGTASVVCLLNNFYPREAKVQWKVDNALQSGNS
QESVTEQDSKDSTYSLSSTLTLSKADYEKHKVYACEVTHQGLSSPVTKSFNRGEC pH-Dependent
Antibody HC (human IgG4) (SEQ ID NO: 197)
EMQLVESGGGLVQPGGSLRLSCAASGFTFSSHWMKWV
ROAPGKGLEWVANINODGSEKYYVDSVKGRFTISRDNAKNSLFLOMNSLRAEDTAVYYCA
RDIVLMVYHMDYYYYGMDVWGQGTTVTVSSASTKGPSVFPLAPCSRSTSESTAALGCLVK
DYFPEPVTVSWNSGALTSGVHTFPAVLOSSGLYSLSSVVTVPSSSLGTKTYTCNVDHKPS
NTKVDKRVESKYGPPCPPCPAPEFEGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSQED
PEVQFNWYVDGVEVHNAKTKPREEQFNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKGLPS
SIEKTISKAKGQPREPQVYTLPPSQEEMTKNQVSLTCLVKGFYPSDIAVEWESNGQPENN
YKTTPPVLDSDGSFFLYSRLTVDKSRWQEGNVFSCSVMHEALHNHYTQKSLSLSLGK pH-dependent
Antibody LC (human kappa) (SEQ ID NO: 198)
DIVMTQSPLSLPVTPGEPASISCRSSQSLHHSNGNNY
LDWYLOKPGOSPOLLIYLGSNRASGVPDRFSGSGSGTDFTLKISRVEAEDVGVYYCMOTL
QTPLTFGGGTKVEIKRTVAAPSVFIFPPSDEQLKSGTASVVCLLNNFYPREAKVQWKVDN
ALOSGNSOESVTEODSKDSTYSLSSTLTLSKADYEKHKVYACEVTHOGLSSPVTKSFNRG EC
```

Claims

1. An antibody or a fragment thereof comprising a binding site which specifically binds to Proprotein Convertase Subtilisin Kexin type 9 (PCSK9), wherein the binding site of the antibody or the fragment thereof comprises: (i) a VH domain comprising a CDRH1 comprising SEQ ID NO: 263 or 277, a CDRH2 comprising SEQ ID NO: 267 or 281, and a CDRH3 comprising SEQ ID NO: 271 or 285, respectively; and a VL domain comprising a CDRL1 comprising SEQ ID NO: 293 or 307, a CDRL2 comprising SEQ ID NO: 297 or 311, and a CDRL3 comprising SEQ ID NO: 301 or 315, respectively; or (ii) a VH domain comprising a CDRH1 comprising SEQ ID NO: 323 or 337, a CDRH2 comprising SEQ ID NO: 327 or 341, and a CDRH3 comprising SEQ ID NO: 331 or 345, respectively; and a VL domain comprising a CDRL1 comprising SEQ ID NO: 353 or 367, a CDRL2 comprising SEQ ID NO: 357 or 371, and a CDRL3 comprising SEQ ID NO: 361 or 375, respectively; or (iii) a VH domain comprising a CDRH1 comprising SEQ ID NO: 7 or 21, a CDRH2 comprising SEQ ID NO: 11 or 25, and a CDRH3 comprising SEQ ID NO: 15 or 29, respectively; and a VL domain comprising a CDRL1 comprising SEQ ID NO: 39 or 53, a CDRL2 comprising SEQ ID NO: 43 or 57, and a CDRL3 comprising SEQ ID NO: 47 or 61, respectively; or (iv) a VH domain comprising a CDRH1 comprising SEQ ID NO: 69 or 83, a CDRH2 comprising SEQ ID NO: 73 or 87, and a CDRH3 comprising SEQ ID NO: 77 or 91, respectively; and a VL domain comprising a CDRL1 comprising SEQ ID NO: 99 or 113, a CDRL2 comprising SEQ ID NO: 103 or 117, and a CDRL3 comprising SEQ ID NO: 107 or 121, respectively, or (v) a VH domain comprising a CDRH1 comprising SEQ ID NO: 203 or 217, a CDRH2 comprising SEQ ID NO: 207 or 221, and a CDRH3 comprising SEQ ID NO: 211 or 225, respectively; and a VL domain comprising a CDRL1 comprising SEQ ID NO: 233 or 247, a CDRL2 comprising SEQ ID NO: 237 or 251, and a CDRL3 comprising SEQ ID NO: 241 or 255, respectively. 2. The antibody or the fragment thereof according to claim 1, wherein the binding site comprises: (i) a VH domain comprising SEQ ID NO: 259 or an amino acid sequence that is at least 70% identical thereto; and a VL domain

comprising SEQ ID NO: 289, or an amino acid sequence that is at least 70% identical thereto; or (ii) a VH domain comprising SEQ ID NO: 319, or an amino acid sequence that is at least 70% identical thereto; and a VL domain comprising SEQ ID NO: 349, or an amino acid sequence that is at least 70% identical thereto; or (iii) a VH domain comprising SEQ ID NO: 1, or an amino acid sequence that is at least 70% identical thereto; and a VL domain comprising SEQ ID NO: 33, or an amino acid sequence that is at least 70% identical thereto; or (iv) a VH domain comprising SEQ ID NO: 65, or an amino acid sequence that is at least 70% identical thereto; and a VL domain

comprising SEQ ID NO: 95, or an amino acid sequence that is at least 70% identical thereto; or (v) a VH domain comprising SEQ ID NO: 199, or an amino acid sequence that is at least 70% identical thereto; and a VL domain comprising SEQ ID NO: 229, or an amino acid sequence that is at least 70% identical thereto.

- 3. The antibody or the fragment thereof according claim 1, wherein the binding site comprises a VH domain comprising SEQ ID NO: 319 and a VL domain comprising SEQ ID NO: 349.
- 4. A method of treating hypercholesterolemia or hyperlipidemia in a subject, the method comprising: administering a therapeutically effective amount of the antibody or the fragment thereof according to claim 1 to the subject.
- 5. A nucleic acid that encodes the antibody or the fragment thereof according to claim 1.
- 6. A nucleic acid that encodes the antibody or the fragment thereof according to claim 3.
- 7. A vector comprising the nucleic acid of claim 5.
- 8. An isolated host cell comprising the nucleic acid of claim 5.
- 9. The antibody or the fragment thereof according to claim 1, wherein the binding site comprises a VH domain comprising SEQ ID NO: 259 and a VL domain comprising SEQ ID NO: 289.