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(54) Title: ANTIBODIES, VARIABLE DOMAINS & CHAINS TAILORED FOR HUMAN USE

Recombined BAC Vectors to add Polymorphic V-regions to the Mouse Genome

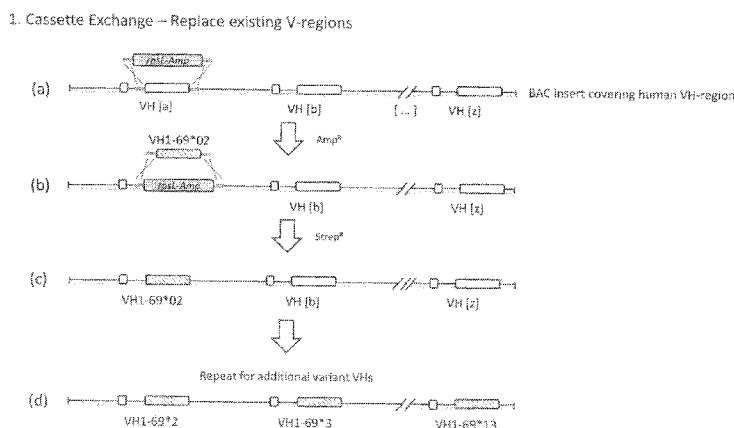


Figure 1

(57) Abstract: The invention relates to the provision of antibody therapeutics and prophylactics that are tailored specifically for human use. The present invention provides libraries, vertebrates and cells, such as transgenic mice or rats or transgenic mouse or rat cells. Furthermore, the invention relates to methods of using the vertebrates to isolate antibodies or nucleotide sequences encoding antibodies. Antibodies, heavy chains, polypeptides, nucleotide sequences, pharmaceutical compositions and uses are also provided by the invention.





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**ANTIBODIES, VARIABLE DOMAINS & CHAINS TAILORED FOR HUMAN USE****FIELD OF THE INVENTION**

The present invention relates to the provision of antibody therapeutics and prophylactics that are tailored specifically for human use.

The present invention provides libraries, vertebrates and cells, such as transgenic mice or rats or transgenic mouse or rat cells. Furthermore, the invention relates to methods of using the vertebrates to isolate antibodies or nucleotide sequences encoding antibodies. Antibodies, heavy chains, polypeptides, nucleotide sequences, pharmaceutical compositions and uses are also provided by the invention.

**BACKGROUND**

The state of the art provides non-human vertebrates (eg, mice and rats) and cells comprising transgenic immunoglobulin loci, such loci comprising human variable (V), diversity (D) and/or joining (J) segments, and optionally human constant regions. Alternatively, endogenous constant regions of the host vertebrate (eg, mouse or rat constant regions) are provided in the transgenic loci. Methods of constructing such transgenic vertebrates and use of these to generate antibodies and nucleic acids thereof following antigen immunisation are known in the art, eg, see US7501552 (Medarex), US5939598 (Abgenix), US6130364 (Abgenix), WO02/066630 (Regeneron), WO2011004192 (Genome Research Limited), WO2009076464, WO2009143472 and WO2010039900 (Ablexis), the disclosures of which are explicitly incorporated herein. Such transgenic loci in the art include varying amounts of the human V(D) J repertoire. Existing transgenic immunoglobulin loci are based on a single human DNA source. The potential diversity of human antibody variable regions in non-human vertebrates bearing such transgenic loci is thus confined.

The inventors considered that it would be desirable to tailor the genomes of these transgenic non-human vertebrates (and thus antibody and antibody chain products of these) to address the variability – and commonality – in the natural antibody gene usage of humans. The inventors wanted to do this in order to better address human use of antibody-based therapeutic and prophylactic drugs.

It would be desirable also to provide for novel and potentially expanded repertoire and diversity of human variable regions in transgenic immunoglobulin loci and non-human vertebrates harbouring these, as well as in antibodies produced following immunisation of such animals.

### **SUMMARY OF THE INVENTION**

The present invention has been developed from extensive bioinformatics analysis of natural antibody gene segment distributions across a myriad of different human populations and across more than two thousand samples from human individuals. The inventors have undertaken this huge task to more thoroughly understand and design non-human vertebrate systems and resultant antibodies to better address human medical therapeutics as a whole, as well as to enable rational design to address specific ethnic populations of humans. Using such rational design, the inventors have constructed transgenic non-human vertebrates and isolated antibodies, antibody chains and cells expressing these in a way that yields products that utilise gene segments that have been purposely included on the basis of the human bioinformatics analysis. The examples illustrate worked experiments where the inventors isolated many cells and antibodies to this effect.

The invention also relates to synthetically-extended & ethnically-diverse superhuman immunoglobulin gene repertoires. The present invention thus provides for novel and potentially expanded synthetic immunoglobulin diversities, thus providing a pool of diversity from which human antibody therapeutic leads can be selected. This expanded pool is useful when seeking to find antibodies with desirable characteristics, such as relatively high affinity to target antigen without the need for further affinity maturation (eg, using laborious *in vitro* techniques such as phage or ribosome display), or improved biophysical characteristics, or to address targets and new epitopes that have previously been difficult to address with antibodies are not reached by prior antibody binding sites.

The invention also provides for diversity that is potentially biased towards variable gene usage common to members of a specific human population, which is useful for generating antibodies for treating and/or preventing diseases or conditions within such population. This ability to bias the

antibody repertoire allows one to tailor antibody therapeutics with the aim of more effectively treating and/or preventing disease or medical conditions in specific human populations.

The present inventors realised the possibility of providing immunoglobulin gene segments from disparate sources in transgenic loci, in order to provide for novel and potentially-expanded antibody diversities from which antibody therapeutics (and antibody tool reagents) could be generated. This opens up the potential of transgenic human-mouse/rat technologies to the possibility of interrogating different and possibly larger antibody sequence-spaces than has hitherto been possible.

In rationally designing transgenic antibody loci, as well as antibodies and antibody chains, the inventors also realised that a relatively long HCDR3 length (at least 20 amino acids) is often desirable to address epitopes. For example, naturally-occurring antibodies have been isolated from humans infected with infectious disease pathogens, such antibodies having a long HCDR3 length. Neutralising antibodies have been found in this respect. A long HCDR3 length would be desirable to address other antigens (eg, receptor clefts or enzyme active sites), not just limited to infectious disease pathogens, and thus the inventors realised the general desirability of the possibility of engineering transgenic loci to be able to produce long HCDR3 antibodies and heavy chains. The inventors, through laborious execution of bioinformatics on in excess of 2000 human DNA samples *via* the 1000 Genomes project together with rational sequence choices, identified that the inclusion of the specific human gene segment variant JH6\*02 is desirable for producing long HCDR3 antibodies and chains.

Additional rational design and bioinformatics has led the inventors to realise that specific human constant region variants are conserved across many diverse human populations. The inventors realised that this opens up the possibility of making a choice to humanise antibodies, chains and variable domains by using such specific constant regions in products, rather than arbitrarily choosing the human constant region (or a synthetic version of a human constant region). This aspect of the invention also enables one to tailor antibody-based drugs to specific human ethnic populations, thereby more closely matching drug to patient (and thus disease setting) than has hitherto been performed. It can be a problem in the state of the art that antibodies are humanised with an arbitrary choice of human constant region (presumably derived from one (often unknown) ethnic

population or non-naturally occurring) that does not function as well in patients of a different human ethnic population. This is important, since the constant region has the major role in providing antibody effector functions, eg, for antibody recycling, cellular and complement recruitment and for cell killing.

To this end, in a first configuration of the invention, there is provided

#### First Configuration

A non-human vertebrate or vertebrate cell (optionally an ES cell or antibody-producing cell) comprising a genome having a superhuman immunoglobulin heavy chain human VH and/or D and/or J gene repertoire.

A non-human vertebrate or vertebrate cell (optionally an ES cell or antibody-producing cell) comprising a genome having a superhuman immunoglobulin light chain human VL gene repertoire; optionally wherein the vertebrate or cell is according to the first configuration.

A non-human vertebrate or vertebrate cell (optionally an ES cell or antibody-producing cell) whose genome comprises a transgenic immunoglobulin locus (eg, a heavy chain locus or a light chain locus), said locus comprising immunoglobulin gene segments according to the first and second human immunoglobulin gene segments (optionally V segments) as mentioned below operably connected upstream of an immunoglobulin constant region; optionally wherein the genome is homozygous for said transgenic immunoglobulin locus; optionally wherein the immunoglobulin locus comprises more than the natural human complement of functional V gene segments; and/or optionally wherein the immunoglobulin locus comprises more than the natural human complement of functional D gene segments; and/or optionally wherein the immunoglobulin locus comprises more than the natural human complement of functional J gene segments.

A transgenic non-human vertebrate (eg, a mouse or rat) or vertebrate cell (optionally an ES cell or antibody-producing cell) whose genome comprises a transgenic immunoglobulin locus comprising a plurality of human immunoglobulin gene segments operably connected upstream of a non-human vertebrate constant region for the production of a repertoire of chimaeric antibodies, or chimaeric light or heavy chains, having a non-human vertebrate constant region and a human variable region; wherein the transgenic locus comprises one or more human immunoglobulin V gene segments, one or more human J gene segments and optionally one or more human D gene segments, a first (optionally a V segment) of said gene segments and a second (optionally a V segment) of said gene segments being different and derived from the genomes of first and second human individuals respectively, wherein the individuals are different; and optionally not related; optionally wherein the immunoglobulin locus comprises more than the natural human complement of functional V gene segments; and/or optionally wherein the immunoglobulin locus comprises more than the natural human complement of functional D gene segments; and/or optionally wherein the immunoglobulin locus comprises more than the natural human complement of functional J gene segments.

A transgenic non-human vertebrate (eg, a mouse or rat) or vertebrate cell (optionally an ES cell or antibody-producing cell) whose genome comprises first and second transgenic immunoglobulin loci, each locus comprising a plurality of human immunoglobulin gene segments operably connected upstream of a non-human vertebrate constant region for the production of a repertoire of chimaeric antibodies, or chimaeric light or heavy chains, having a non-human vertebrate constant region and a human variable region;

wherein (i) the first transgenic locus comprises one or more human immunoglobulin V gene segments, one or more human J gene segments and optionally one or more human D gene segments, (ii) the second transgenic locus comprises one or more human immunoglobulin V gene segments, one or more human J gene segments and optionally one or more human D gene segments; and (iii) wherein a first (optionally a V) gene segment of said first locus and a second (optionally a V) gene segment of said second gene locus are different and derived from the genomes of first and second human individuals respectively, wherein the individuals are different; and optionally not related;

optionally wherein the first and second loci are on different chromosomes (optionally chromosomes with the same chromosome number) in said genome;

optionally wherein each immunoglobulin locus comprises more than the natural human complement of functional V gene segments; and/or

optionally wherein each immunoglobulin locus comprises more than the natural human complement of functional D gene segments; and/or

optionally wherein each immunoglobulin locus comprises more than the natural human complement of functional J gene segments.

A method of constructing a cell (eg, an ES cell) according to the invention, the method comprising

- (a) identifying functional V and J (and optionally D) gene segments of the genome sequence of a (or said) first human individual;
- (b) identifying one or more functional V and/or D and/or J gene segments of the genome sequence of a (or said) second human individual, wherein these additional gene segments are not found in the genome sequence of the first individual;
- (c) and constructing a transgenic immunoglobulin locus in the cell, wherein the gene segments of (a) and (b) are provided in the locus operably connected upstream of a constant region.

In one embodiment, the gene segment(s) in step (b) are identified from an immunoglobulin gene database selected from the 1000 Genomes, Ensembl, Genbank and IMGT databases.

Throughout this text, Genbank is a reference to Genbank release number 185.0 or 191.0; the 1000 Genomes database is Phase 1, release v3, 16<sup>th</sup> March 2012; the Ensembl database is assembly GRCh37.p8 (10/04/2012); the IMGT database is available at [www.imgt.org](http://www.imgt.org).

In one embodiment, the first and second human individuals are members of first and second ethnic populations respectively, wherein the populations are different, optionally wherein the human



immunoglobulin gene segment derived from the genome sequence of the second individual is low-frequency (optionally rare) within the second ethnic population.

This configuration of the invention also provides a method of making a transgenic non-human vertebrate (eg, a mouse or rat), the method comprising

- (a) constructing an ES cell (eg, a mouse C57BL/6N, C57BL/6J, 129S5 or 129Sv strain ES cell) by carrying out the method above;
- (b) injecting the ES cell into a donor non-human vertebrate blastocyst (eg, a mouse C57BL/6N, C57BL/6J, 129S5 or 129Sv strain blastocyst);
- (c) implanting the blastocyst into a foster non-human vertebrate mother (eg, a C57BL/6N, C57BL/6J, 129S5 or 129Sv strain mouse); and
- (d) obtaining a child from said mother, wherein the child genome comprises a transgenic immunoglobulin locus.

In one embodiment, the invention provides a method of isolating an antibody that binds a predetermined antigen (eg, a bacterial or viral pathogen antigen), the method comprising immunising a non-human vertebrate according to the invention.

### Second Configuration

A library of antibody-producing transgenic cells whose genomes collectively encode a repertoire of antibodies, wherein

- (a) a first transgenic cell expresses a first antibody having a chain encoded by a first immunoglobulin gene, the gene comprising a first variable domain nucleotide sequence produced following recombination of a first human unrearranged immunoglobulin gene segment;
- (b) a second transgenic cell expresses a second antibody having a chain encoded by a second immunoglobulin gene, the second gene comprising a second variable domain nucleotide sequence produced following recombination of a second human unrearranged immunoglobulin gene segment, the first and second antibodies being non-identical;
- (c) the first and second gene segments are different and derived from the genome sequences of first and second human individuals respectively, wherein the individuals are different; and optionally not related;

(d) wherein the cells are non-human vertebrate (eg, mouse or rat) cells.

In one embodiment, the first and second human individuals are members of first and second ethnic populations respectively, wherein the populations are different; optionally wherein the ethnic populations are selected from those identified in the 1000 Genomes database.

In another embodiment, the second human immunoglobulin gene segment is a polymorphic variant of the first human immunoglobulin gene segment; optionally wherein the second gene segment is selected from the group consisting of a gene segment in any of Tables 1 to 7 and 9 to 14 below (eg, selected from Table 13 or Table 14), eg, the second gene segment is a polymorphic variant of VH1-69.

### Third Configuration

An isolated antibody having

(a) a heavy chain encoded by a nucleotide sequence produced following recombination in a transgenic non-human vertebrate cell of an unrearranged human immunoglobulin V gene segment with a human D and human J segment, optionally with affinity maturation in said cell, wherein one of the gene segments is derived from the genome of an individual from a first human ethnic population; and the other two gene segments are derived from the genome of an individual from a second, different, human ethnic population, and wherein the antibody comprises heavy chain constant regions of said non-human vertebrate (eg, rodent, mouse or rat heavy chain constant regions); and/or

(b) a light chain encoded by a nucleotide sequence produced following recombination in a transgenic non-human vertebrate cell of an unrearranged human immunoglobulin V gene segment with a human J segment, optionally with affinity maturation in said cell, wherein one of the gene segments is derived from the genome of an individual from a first human ethnic population (optionally the same as the first population in (a)); and the other gene segment is derived from the genome of an individual from a second, different, human ethnic population (optionally the same as the second population in (a)), and wherein the antibody comprises light chain constant regions of said non-human vertebrate (eg, rodent, mouse or rat heavy light constant regions);

- (c) Optionally wherein each variable domain of the antibody is a human variable domain.
- (d) Optionally wherein the heavy chain constant regions are gamma-type constant regions.

The invention also provides an isolated nucleotide sequence encoding the antibody, optionally wherein the sequence is provided in an antibody expression vector, optionally in a host cell.

The invention also provides a method of producing a human antibody, the method comprising replacing the non-human vertebrate constant regions of the antibody of the third configuration with human antibody constant regions.

The invention also provides a pharmaceutical composition comprising an antibody according to the third configuration, or an antibody produced according to the method above and a diluent, excipient or carrier; optionally wherein the composition is provided in a container connected to an IV needle or syringe or in an IV bag.

The invention also provides an antibody-producing cell that expresses the second antibody recited in any one of the configurations.

In an alternative configuration, the invention contemplates the combination of nucleotide sequences of first and second immunoglobulin gene segments (eg, two or more polymorphic variants of a particular human germline VH or VL gene segment) to provide a synthetic gene segment. Such synthetic gene segment is used, in one embodiment, to build a transgenic immunoglobulin locus, wherein the synthetic gene segment is provided in combination with one or more human variable and J regions (and optionally one or more human D regions) operably connected upstream of a constant region. When provided in the genome of a non-human vertebrate or cell (eg, mouse or rat cell, eg, ES cell), the invention provides for superhuman gene segment diversity. The sequences to be combined can be selected from gene segments that have been observed to be commonly used in human antibodies raised against a particular antigen (eg, a flu antigen, such as haemagglutinin). By combining the sequences, the synthetic gene segment may

recombine *in vivo* to produce an antibody that is well suited to the treatment and/or prevention of a disease or condition (eg, influenza) mediated by said antigen.

#### Fourth Configuration

A non-human vertebrate (optionally a mouse or a rat) or vertebrate cell whose genome comprises an immunoglobulin heavy chain locus comprising human gene segment JH6\*02, one or more VH gene segments and one or more D gene segments upstream of a constant region; wherein the gene segments in the heavy chain locus are operably linked to the constant region thereof so that the mouse is capable of producing an antibody heavy chain produced by recombination of the human JH6\*02 with a D segment and a VH segment.

A non-human vertebrate cell (optionally a mouse cell or a rat cell) whose genome comprises an immunoglobulin heavy chain locus comprising human gene segment JH6\*02, one or more VH gene segments and one or more D gene segments upstream of a constant region; wherein the gene segments in the heavy chain locus are operably linked to the constant region thereof for producing (eg, in a subsequent progeny cell) an antibody heavy chain produced by recombination of the human JH6\*02 with a D segment and a VH segment.

A heavy chain (eg, comprised by an antibody) isolated from a vertebrate of the invention wherein the heavy chain comprises a HCDR3 of at least 20 amino acids.

A method for producing a heavy chain, VH domain or an antibody specific to a target antigen, the method comprising immunizing a non-human vertebrate according to the invention with the antigen and isolating the heavy chain, VH domain or an antibody specific to a target antigen or a cell producing the heavy chain, VH domain or an antibody, wherein the heavy chain, VH domain or an antibody comprises a HCDR3 that is derived from the recombination of human JH6\*02 with a VH gene segment and a D gene segment.

A heavy chain, VH domain or an antibody produced by the method.

A B-cell or hybridoma expressing a heavy chain VH domain that is identical to the VH domain of the heavy chain.

A nucleic acid encoding the VH domain of the heavy chain of claim 22, 23 or 28, or encoding the heavy chain.

A vector (eg, a CHO cell or HEK293 cell vector) comprising the nucleic acid; optionally wherein the vector is in a host cell (eg, a CHO cell or HEK293 cell).

A pharmaceutical composition comprising the antibody, heavy chain or VH domain (eg, comprised by an antibody), together with a pharmaceutically-acceptable excipient, diluent or a medicament (eg, a further antigen-specific variable domain, heavy chain or antibody).

The antibody, heavy chain or VH domain (eg, comprised by an antibody) as above for use in medicine.

The use of an antibody, heavy chain or VH domain (eg, comprised by an antibody) as above in the manufacture of a medicament for treating and/or preventing a medical condition in a human.

#### Fifth Configuration

A method of producing an antibody heavy chain, the method comprising

- (a) providing an antigen-specific heavy chain variable domain; and
- (b) combining the variable domain with a human heavy chain constant region to produce an antibody heavy chain comprising (in N- to C-terminal direction) the variable domain and the constant region;

wherein

the human heavy chain constant region is an IGHG1ref, IGHG2ref, IGHG2a, IGHG3ref, IGHG3a, IGHG3b, IGHG4ref or IGHG4a constant region.

An antibody comprising a human heavy chain, the heavy chain comprising a variable domain that is specific for an antigen and a constant region that is an IGHG1ref, IGHG2ref, IGHG2a, IGHG3ref, IGHG3a, IGHG3b, IGHG4ref or IGHG4a constant region . Optionally, the variable domain comprises mouse-pattern AID somatic mutations.

A polypeptide comprising (in N- to C- terminal direction) a leader sequence, a human variable domain that is specific for an antigen and a human constant region that is an IGHG1ref, IGHG2ref, IGHG2a, IGHG3ref, IGHG3a, IGHG3b, IGHG4ref or IGHG4a constant region wherein (i) the leader sequence is not the native human variable domain leader sequence; and/or (ii) the variable domain comprises mouse AID-pattern somatic mutations and/or mouse Terminal deoxynucleotidyl transferase (TdT)- pattern junctional mutations.

A nucleotide sequence encoding (in 5' to 3' direction) a leader sequence and a human antibody heavy chain, the heavy chain comprising a variable domain that is specific for an antigen and a constant region that is an IGHG1ref, IGHG2ref, IGHG2a, IGHG3ref, IGHG3a, IGHG3b, IGHG4ref or IGHG4a constant region; and the leader sequence being operable for expression of the heavy chain and wherein the leader sequence is not the native human variable domain leader sequence.

A nucleotide sequence encoding (in 5' to 3' direction) a promoter and a human antibody heavy chain, the heavy chain comprising a variable domain that is specific for an antigen and a constant region that is an IGHG1ref, IGHG2ref, IGHG2a, IGHG3ref, IGHG3a, IGHG3b, IGHG4ref or IGHG4a constant region ; and the promoter being operable for expression of the heavy chain and wherein the promoter is not the native human promoter.

A vector (eg, a CHO cell or HEK293 cell vector) comprising a IGHG1ref, IGHG2ref, IGHG2a, IGHG3ref, IGHG3a, IGHG3b, IGHG4ref or IGHG4a constant region nucleotide sequence that is 3' of a cloning site for the insertion of a human antibody heavy chain variable domain nucleotide sequence, such

that upon insertion of such a variable domain sequence the vector comprises (in 5' to 3' direction) a promoter, a leader sequence, the variable domain sequence and the constant region sequence so that the vector is capable of expressing a human antibody heavy chain when present in a host cell.

#### Sixth Configuration

A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising at least 3 human variable region gene segments of the same type (eg, at least 3 human VH6-1 gene segments, at least 3 human JH6 gene segments, at least 3 human V $\kappa$ 1-39 gene segments, at least 3 human D2-2 gene segments or at least 3 human J $\kappa$ 1 gene segments), wherein at least two of the human gene segments are variants that are not identical to each other.

A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising at least 2 different non-endogenous variable region gene segments of the same type (eg, at least 2 human VH6-1 gene segments, at least 3 human JH6 gene segments, at least 2 human V $\kappa$ 1-39 gene segments, at least 2 human D2-2 gene segments or at least 2 human J $\kappa$ 1 gene segments) *cis* at the same Ig locus.

A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising at least 2 different human variable region gene segments of the same type (eg, at least 2 human VH6-1 gene segments, at least 2 human JH6 gene segments, at least 2 human V $\kappa$ 1-39 gene segments, at least 2 human D2-2 gene segments or at least 2 human J $\kappa$ 1 gene segments) *trans* at the same Ig locus; and optionally a third human gene segment of the same type, wherein the third gene segment is *cis* with one of said 2 different gene segments.

A population of non-human vertebrates (eg, mice or rats) comprising a repertoire of human variable region gene segments, wherein the plurality comprises at least 2 human variable region gene segments of the same type (eg, at least 2 human VH6-1 gene segments, at least 2 human JH6 gene segments, at least 2 human V $\kappa$ 1-39 gene segments, at least 2 human D2-2 gene segments or at least 2 human J $\kappa$ 1 gene segments), a first of said different gene segments is provided in the genome of a

first vertebrate of the population, and a second of said different gene segments being provided in the genome of a second vertebrate of the population, wherein the genome of the first vertebrate does not comprise the second gene segment.

A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising at least 2 different non-endogenous variable region gene segments of the same type (eg, at least 2 human VH6-1 gene segments, at least 2 human JH6 gene segments, at least 2 human V $\kappa$ 1-39 gene segments, at least 2 human D2-2 gene segments or at least 2 human J $\kappa$ 1 gene segments), wherein the gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations.

A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 3 human variable region gene segments of the same type (eg, at least 3 human VH6-1 gene segments, at least 3 human JH6 gene segments, at least 3 human V $\kappa$ 1-39 gene segments, at least 3 human D2-2 gene segments or at least 3 human J $\kappa$ 1 gene segments), wherein at least two of the human gene segments are variants that are not identical to each other.

A method of enhancing the immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 2 different non-endogenous variable region gene segments of the same type (eg, at least 2 human VH6-1 gene segments, at least 2 human JH6 gene segments, at least 2 human V $\kappa$ 1-39 gene segments, at least 2 human D2-2 gene segments or at least 2 human J $\kappa$ 1 gene segments) *cis* at the same Ig locus.

A method of enhancing the immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 2 different human variable region gene segments of the same type (eg, at least 2 human VH6-1 gene segments, at least 2 human JH6 gene segments, at least 2 human V $\kappa$ 1-39 gene segments, at least 2 human D2-2 gene segments or at least 2 human J $\kappa$ 1 gene segments) *trans* at the same Ig locus; and



optionally a third human gene segment of the same type, wherein the third gene segment is *cis* with one of said 2 different gene segments.

A method of providing an enhanced human immunoglobulin variable region gene segment repertoire, the method comprising providing a population of non-human vertebrates (eg, a mouse or rat) comprising a repertoire of human variable region gene segments, wherein the method comprises providing at least 2 different human variable region gene segments of the same type (eg, at least 2 human VH6-1 gene segments, at least 2 human JH6 gene segments, at least 2 human Vk1-39 gene segments, at least 2 human D2-2 gene segments or at least 2 human Jk1 gene segments), wherein a first of said different gene segments is provided in the genome of a first vertebrate of the population, and a second of said different gene segments is provided in the genome of a second vertebrate of the population, wherein the genome of the first vertebrate does not comprise the second gene segment.

A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 2 different non-endogenous variable region gene segments of the same type (eg, at least 2 human VH6-1 gene segments, at least 2 human JH6 gene segments, at least 2 human Vk1-39 gene segments, at least 2 human D2-2 gene segments or at least 2 human Jk1 gene segments), wherein the gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations.

A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 2 human variable region gene segments of the same type (eg, at least 2 human VH6-1 gene segments, at least 2 human JH6 gene segments, at least 2 human Vk1-39 gene segments, at least 2 human D2-2 gene segments or at least 2 human Jk1 gene segments), wherein the gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations; optionally wherein at least 2 or 3 of said different gene segments are provided at the same Ig locus in said genome.

A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising first and second human Ig locus gene segments of the same type (eg, first and second human JH6 gene segments; or first and second IgG2 gene segments; or first and second human Jλ7 gene segments), wherein the first gene segment is a gene segment selected from any one of Tables 1 and 9 to 14 (eg, selected from Table 13 or Table 14) (eg, IGHJ6-a) and the second gene segment is the corresponding reference sequence.

A population of non-human vertebrates (eg, mice or rats) comprising first and second human Ig locus gene segments of the same type (eg, first and second human JH6 gene segments; or first and second IgG2 gene segments; or first and second human Jλ7 gene segments), wherein the first gene segment is a gene segment selected from any one of Tables 1 and 9 to 14 (eg, selected from Table 13 or Table 14) (eg, IGHJ6-a) and the second gene segment is the corresponding reference sequence, wherein the first gene segment is provided in the genome of a first vertebrate of the population, and the second gene segment is provided in the genome of a second vertebrate of the population.

A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising first and second human Ig locus gene segments of the same type (eg, first and second human JH6 gene segments; or first and second IgG2 gene segments; or first and second human Jλ7 gene segments), wherein the first gene segment is a gene segment selected from any one of Tables 1 and 9 to 14 (eg, selected from Table 13 or Table 14) (eg, IGHJ6-a) and the second gene segment is the corresponding reference sequence.

In one aspect of this configuration, the invention relates to human D gene segment variants as described further below.

In one aspect of this configuration, the invention relates to human V gene segment variants as described further below.

In one aspect of this configuration, the invention relates to human J gene segment variants as described further below.

### **BRIEF DESCRIPTION OF THE FIGURES**

Figures 1 to 3: Schematic illustrating a protocol for producing recombiner BAC vectors to add V gene segments into a mouse genome;

Figure 4: Schematic illustrating a protocol for adding V gene segments to a mouse genome using sequential recombinase mediated cassette exchange (sRMCE); and

Figure 5 (in 4 parts): Alignment of 13 IGHV1-69 variants showing the variable (V) coding region only. Nucleotides that differ from VH1-69 variant \*01 are indicated at the appropriate position whereas identical nucleotides are marked with a dash. Where nucleotide changes result in amino acid differences, the encoded amino acid is shown above the corresponding triplet. Boxed regions correspond to CDR1, CDR2 and CDR3 as indicated.

Figure 6 is a schematic illustrating gene segment diversity and the effect of including variant variants in *cis* according to the invention:-

- (a) Situation in a normal person: Recombination on the same chromosome limits combinations of variants, for instance the antibody gene V4-4 can only be recombined within variant 1 to form for instance for instance V4-4-D-J6 or V4-4-D-J2<sup>A</sup>. Similarly the variant V4-4<sup>A</sup> can't be recombined with either J6 or J2<sup>A</sup> from variant 1 and can only be joined with J-genes from variant 2 to form V4-4<sup>A</sup>-D-J6<sup>A</sup> and V4-4<sup>A</sup>-D-J2. V4-4-J2/J6 complexity = 4.
- (b) Situation in a transgenic mouse: Only one variant is provided so the genome is limited. V4-4-J6/J2 complexity = 2.
- (c) Supra mouse of the invention: The variants are added in *cis* and thus can be recombined in every combination, expanding the repertoire. For instance V4-4 can be combined with J6A, J6, J2A or J2 and similarly V4-4A can be recombined with these same J-genes. The V4-4-J6/J2 complexity = 8, which in this simple example is double that of a person and 4X that of a mouse with a single variant.

Figure 7: Alignment of human JH6\*02 variants. Nucleotides that differ from JH6 \*01 are indicated at the appropriate position whereas identical nucleotides are marked with a dash. Where nucleotide changes result in amino acid differences, the encoded amino acid is shown above. Accession numbers (eg, J00256) are shown to the left of the IMGT variant name.

Figure 8: Alignment of JH sequences from various species.

Figure 9: Codon Table

Figure 10: BAC database extract

### **BRIEF DESCRIPTION OF THE TABLES**

<b><u>Table 1:</u></b>	Human IgH V Polymorphic Variants
<b><u>Table 2:</u></b>	Human IgH D Polymorphic Variants
<b><u>Table 3:</u></b>	Human IgH J Polymorphic Variants
<b><u>Table 4:</u></b>	Human Ig Vk Polymorphic Variants
<b><u>Table 5:</u></b>	Human Ig Vλ Polymorphic Variants
<b><u>Table 6:</u></b>	Human IgH Jk Polymorphic Variants
<b><u>Table 7:</u></b>	Human IgH Jλ Polymorphic Variants
<b><u>Table 8:</u></b>	1000 Genomes Project Human Populations
<b><u>Table 9:</u></b>	Immunoglobulin Gene Usage in Human Antibody Responses to Infectious Disease Pathogens
<b><u>Table 10A:</u></b>	Human IgH JH5 Variant Occurrences
<b><u>Table 10B:</u></b>	Non-Synonymous Human IgH JH5 Variants
<b><u>Table 11A:</u></b>	Human IgH JH6 Variant Occurrences
<b><u>Table 11B:</u></b>	Non-Synonymous Human IgH JH6 Variants
<b><u>Table 12A:</u></b>	Human IgH JH2 Variant Occurrences
<b><u>Table 12B:</u></b>	Non-Synonymous Human IgH JH2 Variants
<b><u>Table 13:</u></b>	Variant Frequency Analyses & Human Population Distributions

<u>Table 14:</u>	Frequent Human Variant Distributions
<u>Table 15:</u>	Human Gene Segment Usage: Heavy Chain Repertoires From Naïve Non-Human Vertebrates
<u>Table 16:</u>	Human Gene Segment Usage: Heavy Chain Repertoires From Immunised Non-Human Vertebrates
<u>Table 17:</u>	Human Gene Segment Usage: Heavy Chain Repertoires From Antigen-Specific Hybridomas
<u>Table 18:</u>	Sequence Correlation Table
<u>Table 19:</u>	Summary Of Function Correlated With Human Gamma Constant Region Sub-Type
<u>Table 20:</u>	Gene Segments Prevalent In Few Human Populations
<u>Table 21:</u>	Genomic and sequence information

### **DETAILED DESCRIPTION OF THE INVENTION**

A suitable source of JH6\*02 and other human DNA sequences for use in the invention will be readily apparent to the skilled person. For example, it is possible to collect a DNA sample from a consenting human donor (eg, a cheek swab sample as per the Example herein) from which can be obtained suitable DNA sequences for use in constructing a locus of the invention. Other sources of human DNA are commercially available, as will be known to the skilled person. Alternatively, the skilled person is able to construct gene segment sequence by referring to one or more databases of human Ig gene segment sequences disclosed herein.

An example source for human V, D and J gene segments according to the invention are Bacterial Artificial Chromosomes (RPCI-11 BACs) obtained from Roswell Park Cancer Institute (RPCI)/Invitrogen. See <http://bacpac.chori.org/hmale11.htm>, which describes the BACs as follows:-

*“RPCI - 11 Human Male BAC Library*

*The RPCI-11 Human Male BAC Library (Osoegawa et al., 2001) was constructed using improved cloning techniques (Osoegawa et al., 1998) developed by Kazutoyo Osoegawa. The library was generated by Kazutoyo Osoegawa. Construction was funded by a grant from the National Human Genome Research Institute (NHGRI, NIH) (#1R01RG01165-03). This library was generated according to the new NHGRI/DOE "Guidance on Human Subjects in Large-Scale DNA Sequencing...*

*"Male blood was obtained via a double-blind selection protocol. Male blood DNA was isolated from one randomly chosen donor (out of 10 male donors)".*

- Osoegawa K, Mammoser AG, Wu C, Frengen E, Zeng C, Catanese JJ, de Jong PJ; Genome Res. 2001 Mar;11(3):483-96; "A bacterial artificial chromosome library for sequencing the complete human genome";
- Osoegawa, K., Woon, P.Y., Zhao, B., Frengen, E., Tateno, M., Catanese, J.J., and de Jong, P.J. (1998); "An Improved Approach for Construction of Bacterial Artificial Chromosome Libraries"; Genomics 52, 1-8.

### Superhuman Immunoglobulin Gene Repertoires

The invention relates to synthetically-extended & ethnically-diverse superhuman immunoglobulin gene repertoires. The human immunoglobulin repertoires are beyond those found in nature (ie, "Superhuman"), for example, they are more diverse than a natural human repertoire or they comprise combinations of human immunoglobulin gene segments from disparate sources in a way that is non-natural. Thus, the repertoires of the invention are "superhuman" immunoglobulin repertoires, and the invention relates to the application of these in transgenic cells and non-human vertebrates for utility in producing chimaeric antibodies (with the possibility of converting these into fully-human, isolated antibodies using recombinant DNA technology). The present invention thus provides for novel and potentially expanded synthetic immunoglobulin diversities, which provides for a pool of diversity from which antibody therapeutic leads (antibody therapeutics and antibody tool reagents) can be selected. This opens up the potential of transgenic human-mouse/rat technologies to the possibility of interrogating different and possibly larger antibody sequence-spaces than has hitherto been possible. To this end, in one embodiment, the invention provides a SUPERHUMAN MOUSE<sup>™</sup> (aka SUPRA-MOUSE<sup>™</sup>) and a SUPERHUMAN RAT<sup>™</sup> (aka SUPRA-RAT<sup>™</sup>)

In developing this thinking, the present inventors have realised the possibility of mining the huge genetics resources now available to the skilled person thanks to efforts such as the HapMap Project, 1000 Genomes Project and sundry other immunoglobulin gene databases (see below for more details). Thus, in some embodiments, the inventors realised the application of these genome sequencing developments in the present invention to generate synthetically-produced and ethnically-diverse artificial immunoglobulin gene repertoires. In one aspect, the inventors realised that such repertoires are useful for the production of antibodies having improved affinity and/or biophysical characteristics, and/or wherein the range of epitope specificities produced by means of such repertoire is novel, provides for antibodies to epitopes that have hitherto been intractable by prior transgenic immunoglobulin loci or difficult to address.

The present invention provides libraries, vertebrates and cells, such as transgenic mice or rats or transgenic mouse or rat cells. Furthermore, the invention relates to methods of using the vertebrates to isolate antibodies or nucleotide sequences encoding antibodies. Antibodies, nucleotide sequences, pharmaceutical compositions and uses are also provided by the invention.

### **Variation Analysis**

The present inventors have realized methods and antibody loci designs that harness the power of genetic variation analysis. The reference human genome provides a foundation for experimental work and genetic analysis of human samples. The reference human is a compilation of the genomes from a small number of individuals and for any one segment of the genome a high quality single reference genome for one of the two chromosomes is available. Because the reference genome was assembled from a series of very large insert clones, the identity of these clones is known. Accordingly, experimental work with human genomic DNA is usually conducted on the clones from which the reference sequence was derived.

Individual humans differ in their sequence and recently several individuals have had their genomes sequenced, for instance James Watson and Craig Venter. Comparison of the genome sequence of these individuals has revealed differences between their sequences and the reference genome in both coding and non-coding parts of the genome, approximately 1 in 1000 bases are different. Some variants will be significant and contribute to differences between individuals. In extreme cases

these will result in genetic disease. Variation can be implicated in differing responses to drugs administered to human patients, eg, yielding an undesirable lowering of patient response to treatment.

The 1000-Genomes Project has the objective of identifying the most frequent variations in the human genome. This public domain project involved sequencing the genomes of more than 1000 individuals from diverse ethnic groups, comparing these sequences to the reference and assembling a catalogue of variants. This has enabled the annotation of variants in coding regions, but because this sequence wasn't derived from large clones of DNA, the analysis of the sequence from diploid individuals can't discriminate the distribution of the variation between the maternal and paternally inherited chromosomes. Where more than one variant is identified in a protein coding gene, it is not possible to illuminate the distribution of the pattern of variants in each version of the protein. For example, if two variants are detected in different positions of the same protein in an individual, this could have resulted from one copy with two variants and none in the other or each copy could have just one variant. To illuminate the sequence of real proteins, the 1000-Genome Project has sequenced mother-father-child trios. This allows one to "phase" the sequence variants, in other words identify blocks of sequence that are inherited from one or other parent and deconvolute the variants.

To further understand the variation within the 1000-genome set a tool has been developed that can identify the significant variants (defined as non-synonymous amino acid changes) from a region of DNA from the phased data in the 1000-genome data set. This tool has been made available online <http://www.1000genomes.org/variation-pattern-finder>. This tool allows an investigator to download non-synonymous variation delimited between specific coordinates. The downloaded files are configured as individual genotypes, but the data is phased so the haplotype information and the frequencies of specific halotypes in different populations can be extracted.

The inventors' analysis of the 1000-genome data for the individual human coding segments of the C, V D and J genes from the heavy and light chains reveals that there is significant variation in these segments. Individuals will usually have two different heavy chain alleles and also different light chain alleles at both kappa and lambda loci. The repertoire of antibodies that can be generated



from each allele will be different. This variation will contribute to a better or differing immune response to certain antigens.

Humanized mice that have hitherto been generated with immunoglobulin heavy and light chain loci contain just one type of immunoglobulin locus. Even if these mice contain a full human heavy chain locus, the variation will be less than contained in a typical human because only one set of C, V, D and J genes are available, while a typical human would have two sets.

The inventors have devised ways to improve on this limitation when constructing transgenic non-human vertebrates and cells for human antibody and variable region production *in vivo*.

Mice can be generated with two different loci, each engineered to have a different repertoire of V, D and J segments. This could be in a single mouse or two or more separate mouse strains and would be analogous to or beyond the repertoire found in a normal human. The engineering of such a mouse would go beyond the repertoire described in humanized mice to date which only have one set of alleles.

However, the inventors also realized that this also has limitations, because the different loci would not normally interact to shuffle V, D and J variants between loci. This same limitation is also inherent in a human, thus this system does not utilize the advantage of recombining variants in all combinations.

To go beyond the normal repertoire in humans and take advantage of combinations of C, V, D and J variants the inventors decided, in one embodiment, to provide these on the same chromosome in *cis*. See figure 6. These loci would be characterized by having more than the normal number of J, D or V genes. For example n=6 for the J genes, but including one J6 variant and one J2 variant would increase this to n=8. This could be combined with additional variants for the D and V genes, for example. By detailed analysis of the 1000- Genomes database, the inventors have devised a collection of candidate polymorphic human variant gene segments, eg, JH gene segments (eg, see the examples), that can be built into the design of transgenic heavy and light chain loci in mice for

expressing increasingly diverse and new, synthetic repertoires of human variable regions. Moreover, by utilizing naturally-occurring human variant gene segments, as per embodiments of the invention, this addresses compatibility with human patients since the inventor's analysis has drawn out candidate variants that are naturally conserved and sometimes very prevalent amongst human ethnic populations. Additionally this enables one to tailor the configurations of the invention to provide for antibody-based drugs that better address specific human ethnic populations.

In an example according to any configuration of the invention, loci (and cells and vertebrates comprising these) are provided in which gene segments from different human populations are used. This is desirable to increase antibody gene diversity to better address more diverse human patients. In an example, the gene segments are from first and second different human populations respectively, and thus the second gene segment is found in the second human population, but not so (or rarely) in the first human population. Rarely means, for example, that the gene segment is found in 5, 4, 3, 2, or 1 or zero individuals in the first population in the 1000 Genomes database. For example, the first gene segment may be shown as present in a first population by reference to Table 13 or 14 herein, the second gene segment may be shown as present in the second population by reference to Table 13 and not in the first population. Optionally, the first gene segment may also be shown as being present in the second population by reference to Table 13 or 14.

In any configuration or aspect of the invention, where a V gene segment is used, this may be used optionally with the native leader sequence. For example, use of genomic DNA (eg, from BACs as in the examples) will mean that the native leader will be used for each V gene segment incorporated into the locus and genomes of the invention. In an alternative, the skilled person may wish to inert a non-native leader sequence together with one or more of the V gene segments. Similarly, in any configuration or aspect of the invention, where a V gene segment is used, this may be used optionally with the native 5' UTR sequence. For example, use of genomic DNA (eg, from BACs as in the examples) will mean that the native 5' UTR sequence will be used for each V gene segment incorporated into the locus and genomes of the invention. In an alternative, the skilled person may wish to exclude the native 5' UTR sequence.

The present invention provides, in a first configuration

(a) Superhuman heavy chain gene repertoires

A non-human vertebrate or vertebrate cell (optionally an ES cell or antibody-producing cell) comprising a genome having a superhuman immunoglobulin heavy chain human VH and/or D and/or J gene repertoire.

In one aspect the cell of the invention is an embryonic stem cell. For example, the ES cell is derived from the mouse C57BL/6N, C57BL/6J, 129S5 or 129Sv strain. In one aspect the non-human vertebrate is a rodent, suitably a mouse, and cells of the invention, are rodent cells or ES cells, suitably mouse ES cells. The ES cells of the present invention can be used to generate animals using techniques well known in the art, which comprise injection of the ES cell into a blastocyst followed by implantation of chimaeric blastocysts into females to produce offspring which can be bred and selected for homozygous recombinants having the required insertion. In one aspect the invention relates to a transgenic animal comprised of ES cell-derived tissue and host embryo derived tissue. In one aspect the invention relates to genetically-altered subsequent generation animals, which include animals having a homozygous recombinants for the VDJ and/or VJ regions.

The natural human immunoglobulin gene segment repertoire consists of (see eg, [www.imgt.org](http://www.imgt.org)):-

VH: total-125 ; functional-41

DH: total-27; functional-23

JH: total-8; functional-6

Vk: total-77; functional-38

Jk: total-5; functional-5

V lambda: total-75; functional-31

J lambda: total-7; functional-5

In one embodiment, the vertebrate or cell genome comprises a transgenic immunoglobulin heavy chain locus comprising a plurality of human immunoglobulin VH gene segments, one or more human D gene segments and one or more human J gene segments, wherein the plurality of VH gene segments consists of more than the natural human repertoire of functional VH gene segments; optionally wherein the genome is homozygous for said transgenic heavy chain locus.

In one embodiment of the vertebrate or cell, the VH gene repertoire consists of a plurality of VH gene segments derived from the genome sequence of a first human individual, supplemented with one or more different VH gene segments derived from the genome sequence of a second, different human individual. Optionally the D and J segments are derived from the genome sequence of the first human individual. Optionally the VH gene segments from the genome sequence of the second individual are selected from the VH gene segments listed in Table 1, 13 or 14. In this way, the locus provides a superhuman repertoire of D gene segments.

Optionally the individuals are not related. Individuals are "not related" in the context of any configuration or aspect of the invention, for example, if one of the individuals does not appear in a family tree of the other individual in the same generation or going back one, two, three or four generations. Alternatively, are not related, for example, if they do not share a common ancestor in the present generation or going back one, two, three or four generations.

In one embodiment of the vertebrate or cell, the transgenic locus comprises more than 41 functional human VH gene segment species, and thus more than the natural human functional repertoire. Optionally the locus comprises at least 42, 43, 44, 45, 46, 47, 48, 49 or 50 functional human VH gene segment species (eg, wherein the locus comprises the full functional VH repertoire of said first individual supplemented with one or more VH gene segments derived from the genome sequence of the second human individual and optionally with one or more VH gene segments derived from the genome sequence of a third human individual). In this way, the locus provides a superhuman repertoire of VH gene segments that is useful for generating a novel gene and antibody diversity for

use in therapeutic and tool antibody selection.

In one embodiment of the vertebrate or cell, the transgenic locus comprises a first VH gene segment derived from the genome sequence of the first individual and a second VH gene segment derived from the genome sequence of the second individual, wherein the second VH gene segment is a polymorphic variant of the first VH gene segment. For example, the VH gene segments are polymorphic variants of VH1-69 as illustrated in the examples below. Optionally the locus comprises a further polymorphic variant of the first VH gene segment (eg, a variant derived from the genome sequence of a third human individual). In this way, the locus provides a superhuman repertoire of VH gene segments.

In one embodiment of the vertebrate or cell, the genome (alternatively or additionally to the superhuman VH diversity) comprises a transgenic immunoglobulin heavy chain locus comprising a plurality of human immunoglobulin VH gene segments, a plurality of human D gene segments and one or more human J gene segments, wherein the plurality of D gene segments consists of more than the natural human repertoire of functional D gene segments. Optionally the genome is homozygous for said transgenic heavy chain locus.

In one embodiment of the vertebrate or cell, the D gene repertoire consists of a plurality of D gene segments derived from the genome sequence of a (or said) first human individual, supplemented with one or more different D gene segments derived from the genome sequence of a (or said) second, different human individual. Optionally the individuals are not related. Optionally the J segments are derived from the genome sequence of the first human individual. Optionally the D gene segments from the genome sequence of the second individual are selected from the D gene segments listed in Table 2, 13 or 14. In this way, the locus provides a superhuman repertoire of D gene segments.

In one embodiment of the vertebrate or cell, the transgenic locus comprises more than 23 functional human D gene segment species; optionally wherein the locus comprises at least 24, 25, 26, 27, 28, 29, 30 or 31 functional human D gene segment species (eg, wherein the locus comprises the full functional D repertoire of said first individual supplemented with one or more D gene segments derived from the genome sequence of the second human individual and optionally with one or more

D gene segments derived from the genome sequence of a third human individual). In this way, the locus provides a superhuman repertoire of D gene segments.

In one embodiment of the vertebrate or cell, the transgenic locus comprises a first D gene segment derived from the genome sequence of the first individual and a second D gene segment derived from the genome sequence of the second individual, wherein the second D gene segment is a polymorphic variant of the first D gene segment. Optionally the locus comprises a further polymorphic variant of the first D gene segment (eg, a variant derived from the genome sequence of a third human individual). In this way, the locus provides a superhuman repertoire of D gene segments.

In one embodiment of the vertebrate or cell (alternatively or additionally to the superhuman VH and/or JH diversity), the genome comprises a (or said) transgenic immunoglobulin heavy chain locus comprising a plurality of human immunoglobulin VH gene segments, one or more human D gene segments and a plurality of human JH gene segments, wherein the plurality of J gene segments consists of more than the natural human repertoire of functional J gene segments; optionally wherein the genome is homozygous for said transgenic heavy chain locus.

In one embodiment of the vertebrate or cell, the JH gene repertoire consists of a plurality of J gene segments derived from the genome sequence of a (or said) first human individual, supplemented with one or more different J gene segments derived from the genome sequence of a (or said) second, different human individual. Optionally the individuals are not related. Optionally D segments are derived from the genome sequence of the first human individual. Optionally the J gene segments from the genome sequence of the second individual are selected from the J gene segments listed in Table 3 13 or 14. In this way, the locus provides a superhuman repertoire of JH gene segments.

In one embodiment of the vertebrate or cell, the transgenic locus comprises more than 6 functional human JH gene segment segments. Optionally the locus comprises at least 7, 8, 9, 10, 11, 12, 13, 14, 15, or 16 functional human JH gene segments (eg, wherein the locus comprises the full functional JH repertoire of said first individual supplemented with one or more JH gene segments derived from

the genome sequence of the second human individual and optionally with one or more JH gene segments derived from the genome sequence of a third human individual). In this way, the locus provides a superhuman repertoire of JH gene segments.

In one embodiment of the vertebrate or cell, the transgenic locus comprises a first JH gene segment derived from the genome sequence of the first individual and a second JH gene segment derived from the genome sequence of the second individual, wherein the second JH gene segment is a polymorphic variant of the first JH gene segment. Optionally the locus comprises a further polymorphic variant of the first JH gene segment (eg, a variant derived from the genome sequence of a third human individual). In this way, the locus provides a superhuman repertoire of JH gene segments.

(b) Superhuman light chain gene repertoires

The first configuration of the invention also provides:-

A non-human vertebrate or vertebrate cell (optionally an ES cell or antibody-producing cell) comprising a genome having a superhuman immunoglobulin light chain human VL gene repertoire. Optionally the vertebrate or cell comprises a heavy chain transgene according to aspect (a) of the first configuration. Thus, superhuman diversity is provided in both the heavy and light chain immunoglobulin gene segments in the cell and vertebrate. For example, the genome of the cell or vertebrate is homozygous for the heavy and light chain transgenes and endogenous antibody expression is inactivated. Such a vertebrate is useful for immunisation with a predetermined antigen to produce one or more selected antibodies that bind the antigen and have human variable regions resulting from recombination within the superhuman gene segment repertoire. This provides potentially for a novel antibody and gene sequence space from which to select therapeutic, prophylactic and tool antibodies.

In one embodiment of aspect (b) of the first configuration, the vertebrate or cell genome comprises

- (i) a transgenic immunoglobulin kappa light chain locus comprising a plurality of human

immunoglobulin V $\kappa$  gene segments and one or more human J gene segments, wherein the plurality of V $\kappa$  gene segments consists of more than the natural human repertoire of functional V $\kappa$  gene segments; optionally wherein the genome is homozygous for said transgenic kappa light chain locus; and/or

(ii) a transgenic immunoglobulin lambda light chain locus comprising a plurality of human immunoglobulin V $\lambda$  gene segments and one or more human J gene segments, wherein the plurality of V $\lambda$  gene segments consists of more than the natural human repertoire of functional V $\lambda$  gene segments; optionally wherein the genome is homozygous for said transgenic lambda light chain locus.

In this way, the locus provides a superhuman repertoire of VL gene segments.

In one embodiment of the vertebrate or cell,

(i) the V $\kappa$  gene repertoire consists of a plurality of V $\kappa$  gene segments derived from the genome sequence of a first human individual, supplemented with one or more V $\kappa$  gene segments derived from the genome sequence of a second, different human individual; optionally wherein the individuals are not related; optionally wherein the J segments are derived from the genome sequence of the first human individual; and optionally wherein the V $\kappa$  gene segments from the genome sequence of the second individual are selected from the V $\kappa$  gene segments listed in Table 4, 13 or 14; and

(i) the V $\lambda$  gene repertoire consists of a plurality of V $\lambda$  gene segments derived from the genome sequence of a first human individual, supplemented with one or more V $\lambda$  gene segments derived from the genome sequence of a second, different human individual; optionally wherein the individuals are not related; optionally wherein the J segments are derived from the genome sequence of the first human individual; and optionally wherein the V $\lambda$  gene segments from the genome sequence of the second individual are selected from the V $\lambda$  gene segments listed in Table 5, 13 or 14.

In this way, the locus provides a superhuman repertoire of VL gene segments.



In one embodiment of the vertebrate or cell,

-the kappa light transgenic locus comprises more than 38 functional human  $V\kappa$  gene segment species; optionally wherein the locus comprises at least 39, 40, 41, 42, 43, 44, 45, 46, 47 or 48 functional human  $V\kappa$  gene segment species (eg, wherein the locus comprises the full functional  $V\kappa$  repertoire of said first individual supplemented with one or more  $V\kappa$  gene segments derived from the genome sequence of the second human individual and optionally with one or more  $V\kappa$  gene segments derived from the genome sequence of a third human individual); and

-the lambda light transgenic locus comprises more than 31 functional human  $V\lambda$  gene segment species; optionally wherein the locus comprises at least 32, 33, 34, 35, 36, 37, 38, 39, 40 or 41 functional human  $V\lambda$  gene segment species (eg, wherein the locus comprises the full functional  $V\lambda$  repertoire of said first individual supplemented with one or more  $V\lambda$  gene segments derived from the genome sequence of the second human individual and optionally with one or more  $V\lambda$  gene segments derived from the genome sequence of a third human individual).

In this way, the locus provides a superhuman repertoire of VL gene segments.

In one embodiment of the vertebrate or cell,

-the kappa light transgenic locus comprises a first  $V\kappa$  gene segment derived from the genome sequence of the first individual and a second  $V\kappa$  gene segment derived from the genome sequence of the second individual, wherein the second  $V\kappa$  gene segment is a polymorphic variant of the first  $V\kappa$  gene segment; optionally wherein the locus comprises a further polymorphic variant of the first  $V\kappa$  gene segment (eg, a variant derived from the genome sequence of a third human individual); and

-the lambda light transgenic locus comprises a first  $V\lambda$  gene segment derived from the genome sequence of the first individual and a second  $V\lambda$  gene segment derived from the genome sequence of the second individual, wherein the second  $V\lambda$  gene segment is a polymorphic variant of the first  $V\lambda$  gene segment; optionally wherein the locus comprises a further polymorphic variant of the first  $V\lambda$  gene segment (eg, a variant derived from the genome sequence of a third human individual).

In this way, the locus provides a superhuman repertoire of VL gene segments.

In one embodiment of the vertebrate or cell, the genome comprises a (or said) transgenic immunoglobulin light chain locus comprising a plurality of human immunoglobulin VL gene segments and a plurality of human JL gene segments, wherein the plurality of J gene segments consists of more than the natural human repertoire of functional J gene segments; optionally wherein the genome is homozygous for said transgenic heavy chain locus.

In one embodiment of the vertebrate or cell,

(i) the J<sub>k</sub> gene repertoire consists of a plurality of J<sub>k</sub> gene segments derived from the genome sequence of a (or said) first human individual, supplemented with one or more J<sub>k</sub> gene segments derived from the genome sequence of a (or said) second, different human individual; optionally wherein the individuals are not related; optionally wherein the V<sub>k</sub> segments are derived from the genome sequence of the first human individual; optionally wherein the J<sub>k</sub> gene segments from the genome sequence of the second individual are selected from the J<sub>k</sub> gene segments listed in Table 6, 13 or 14; and

(ii) the J<sub>k</sub> gene repertoire consists of a plurality of J<sub>λ</sub> gene segments derived from the genome sequence of a (or said) first human individual, supplemented with one or more J<sub>λ</sub> gene segments derived from the genome sequence of a (or said) second, different human individual; optionally wherein the individuals are not related; optionally wherein the V<sub>λ</sub> segments are derived from the genome sequence of the first human individual; optionally wherein the J<sub>λ</sub> gene segments from the genome sequence of the second individual are selected from the J<sub>λ</sub> gene segments listed in Table 7, 13 or 14.

In this way, the locus provides a superhuman repertoire of JL gene segments.

In one embodiment of the vertebrate or cell,

(i) the transgenic light chain locus comprises more than 5 functional human J<sub>k</sub> gene segment species; optionally wherein the locus comprises at least 6, 7, 8, 9, 10, 11, 12, 13, 14 or 15 functional human J<sub>k</sub> gene segment species (eg, wherein the locus comprises the full functional J<sub>k</sub> repertoire of said first individual supplemented with one or more J<sub>k</sub> gene segments derived from the genome sequence of the second human individual and optionally with one or more J<sub>k</sub> gene segments derived

from the genome sequence of a third human individual); and/or

(i) the transgenic light chain locus comprises more than 5 functional human J $\lambda$  gene segment species; optionally wherein the locus comprises at least 6, 7, 8, 9, 10, 11, 12, 13, 14 or 15 functional human J $\lambda$  gene segment species (eg, wherein the locus comprises the full functional J $\lambda$  repertoire of said first individual supplemented with one or more J $\lambda$  gene segments derived from the genome sequence of the second human individual and optionally with one or more J $\lambda$  gene segments derived from the genome sequence of a third human individual).

In this way, the locus provides a superhuman repertoire of JL gene segments.

In one embodiment of the vertebrate or cell,

(i) the kappa light transgenic locus comprises a first Jk gene segment derived from the genome sequence of the first individual and a second Jk gene segment derived from the genome sequence of the second individual, wherein the second Jk gene segment is a polymorphic variant of the first Jk gene segment; optionally wherein the locus comprises a further polymorphic variant of the first Jk gene segment (eg, a variant derived from the genome sequence of a third human individual); and

(ii) the lambda light transgenic locus comprises a first J $\lambda$  gene segment derived from the genome sequence of the first individual and a second J $\lambda$  gene segment derived from the genome sequence of the second individual, wherein the second Jk gene segment is a polymorphic variant of the first J $\lambda$  gene segment; optionally wherein the locus comprises a further polymorphic variant of the first J $\lambda$  gene segment (eg, a variant derived from the genome sequence of a third human individual).

In this way, the locus provides a superhuman repertoire of JL gene segments.

Further aspects of the first configuration are described below.

#### The present invention provides, in a second configuration

A library of antibody-producing transgenic cells whose genomes collectively encode a repertoire of antibodies, wherein

- (a) a first transgenic cell expresses a first antibody having a chain (eg, heavy chain) encoded by a first immunoglobulin gene, the gene comprising a first variable domain nucleotide sequence produced following recombination of a first human unrearranged immunoglobulin gene segment (eg, a VH);
- (b) a second transgenic cell expresses a second antibody having a chain (eg, a heavy chain) encoded by a second immunoglobulin gene, the second gene comprising a second variable domain nucleotide sequence produced following recombination of a second human unrearranged immunoglobulin gene segment (eg, a VH), the first and second antibodies being non-identical;
- (c) the first and second gene segments are different and derived from the genome sequences of first and second human individuals respectively, wherein the individuals are different; and optionally not related;
- (d) wherein the cells are non-human vertebrate (eg, mouse or rat) cells (eg, B-cells or hybridomas).

In one embodiment, the library is provided *in vitro*. In another embodiment, the library is provided *in vivo* by one or a plurality of transgenic non-human vertebrates. For example, the or each vertebrate is according to any aspect of the first configuration of the invention.

In one embodiment, the library encodes an antibody repertoire of from 10 to  $10^9$  antibodies, for example, 10, 20, 30, 40, 50, 100 or 1000 to  $10^8$ ; or 10, 20, 30, 40, 50, 100 or 1000 to  $10^7$ ; or 10, 20, 30, 40, 50, 100 or 1000 to  $10^6$ ; or 10, 20, 30, 40, 50, 100 or 1000 to  $10^5$ ; or 10, 20, 30, 40, 50, 100 or 1000 to  $10^4$  antibodies. In an example, library encodes an antibody repertoire of at least  $10^3$ ,  $10^4$ ,  $10^5$ ,  $10^6$ ,  $10^7$ ,  $10^8$ ,  $10^9$  or  $10^{10}$  antibodies.

The first variable domain nucleotide sequence is produced following recombination of the first human unrearranged immunoglobulin gene segment with one or more other immunoglobulin gene segments (for example, human immunoglobulin gene segments). For example, where the first gene segment is a VH, the first variable domain nucleotide sequence (a VH domain) is produced following recombination of the VH with a human D and JH segments *in vivo*, optionally with somatic hypermutation, in the first transgenic cell or an ancestor thereof. For example, where the first gene segment is a VL, the first variable domain nucleotide sequence (a VL domain) is produced following recombination of the VL with a human JL segment *in vivo*, optionally with somatic hypermutation, in

the first transgenic cell or an ancestor thereof.

The second variable domain nucleotide sequence is produced following recombination of the second human unrearranged immunoglobulin gene segment with one or more other immunoglobulin gene segments (for example, human immunoglobulin gene segments). For example, where the second gene segment is a VH, the second variable domain nucleotide sequence (a VH domain) is produced following recombination of the VH with a human D and JH segments *in vivo*, optionally with somatic hypermutation, in the second transgenic cell or an ancestor thereof. For example, where the second gene segment is a VL, the second variable domain nucleotide sequence (a VL domain) is produced following recombination of the VL with a human JL segment *in vivo*, optionally with somatic hypermutation, in the second transgenic cell or an ancestor thereof.

The first and second gene segments are respectively derived from genome sequences of first and second human individuals. In one example, such a gene segment is isolated or cloned from a sample cell taken from said individual using standard molecular biology techniques as known to the skilled person. The sequence of the gene segment may be mutated (eg, by the introduction of up to 5, 6, 7, 8, 9, 10, 11, 12, 13, 14 or 15 nucleotide changes) prior to use in the present invention. In another example, a gene segment is derived by identifying a candidate human immunoglobulin gene segment in a database (see guidance below) and a nucleotide sequence encoding a gene segment for use in the present invention is made by reference (eg, to be identical or a mutant with up to 5, 6, 7, 8, 9, 10, 11, 12, 13, 14 or 15 nucleotide changes to the reference sequence) to the database sequence. The skilled person will be aware of methods of obtaining nucleotide sequences by reference to databases or by obtaining from cellular samples.

In one embodiment of the vertebrate, cell or library of any configuration of the invention, the first and second human individuals are members of first and second ethnic populations respectively, wherein the populations are different. This, therefore, provides for superhuman gene diversity in transgenic loci, cells and vertebrates as per the invention.

#### Human Populations

Optionally the ethnic populations are selected from those identified in the 1000 Genomes Project of database. In this respect, see Table 8 which provides details of the ethnic populations on which the 1000 Genomes database is based.

N A Rosenberg *et al* (Science 20 December 2002: vol. 298 no. 5602 2342-2343) studied the genetic structure of human populations of differing geographical ancestry. In total, 52 populations were sampled, these being populations with:

*African ancestry*

(Mbuti Pygmies, Biaka Pygmies, San peoples, and speakers of Niger-Kordofanian languages (Bantu, Yoruba or Mandenka populations),

*Eurasian ancestry*

(European ancestry (Orkadian, Adygel, Basque, French, Russians, Italians, Sardinian, Tuscan), Middle Eastern ancestry (Mozabite, Bedouin, Druze, Palestinians), Central/South Asian ancestry (Balochi, Brahui, Makrani, Sindhi, Pathan, Burusho, Hazara, Uyghur, Kalash)),

*East Asian ancestry*

(Han, Dal, Daur, Hezhen, Lahu, Miao, Oroqen, She, Tujia, Tu, Xibo, Yi, Mongola, Naxi, Cambodian, Japanese, Yakut), Oceanic ancestry (Melanesian, Papuan); or

*Americas ancestry*

(Karitiana, Surui, Colombian, Maya, Pima).

The International HapMap Project, Nature, 2003 Dec 18;426(6968):789-96, discloses that goal of the HapMap Project: to determine the common patterns of DNA sequence variation in the human genome by determining the genotypes of one million or more sequence variants, their frequencies and the degree of association between them in DNA samples from populations with ancestry from parts of Africa, Asia and Europe. The relevant human populations of differing geographical ancestry include Yoruba, Japanese, Chinese, Northern European and Western European populations. More specifically:-

Utah population with Northern or Western European ancestry (samples collected in 1980 by the Centre d'Etude du Polymorphisme Humain (CEPH));  
population with ancestry of Yoruba people from Ibadan, Nigeria;  
population with Japanese ancestry; and  
population with ancestry of Han Chinese from China.

The authors, citing earlier publications, suggest that ancestral geography is a reasonable basis for sampling human populations.

A suitable sample of human populations from which the populations used in the present invention are selected is as follows:-

(a) European ancestry

- (b) Northern European ancestry; Western European ancestry; Toscani ancestry; British ancestry, Finnish ancestry or Iberian ancestry.
- (c) More specifically, population of Utah residents with Northern and/or Western European ancestry; Toscani population in Italia; British population in England and/or Scotland; Finnish population in Finland; or Iberian population in Spain.

(a) East Asian ancestry

- (b) Japanese ancestry; Chinese ancestry or Vietnamese ancestry.
- (c) More specifically, Japanese population in Toyko, Japan; Han Chinese population in Beijing, China; Chinese Dai population in Xishuangbanna; Kinh population in Ho Chi Minh City, Vietnam; or Chinese population in Denver, Colorado, USA.

(a) West African ancestry

- (b) Yoruba ancestry; Luhya ancestry; Gambian ancestry; or Malawian ancestry.

- (c) More specifically, Yoruba population in Ibadan, Nigeria; Luhya population in Webuye, Kenya; Gambian population in Western Division, The Gambia; or Malawian population in Blantyre, Malawi.

(a) Population of The Americas

- (b) Native American ancestry; Afro-Caribbean ancestry; Mexican ancestry; Puerto Rican ancestry; Columbian ancestry; or Peruvian ancestry.
- (c) More specifically, population of African Ancestry in Southwest US; population of African American in Jackson, MS; population of African Caribbean in Barbados; population of Mexican Ancestry in Los Angeles, CA; population of Puerto Rican in Puerto Rico; population of Colombian in Medellin, Colombia; or population of Peruvian in Lima, Peru.

(a) South Asian ancestry

- (b) Ahom ancestry; Kayadtha ancestry; Reddy ancestry; Maratha; or Punjabi ancestry.
- (c) More specifically, Ahom population in the State of Assam, India; Kayadtha population in Calcutta, India; Reddy population in Hyderabad, India; Maratha population in Bombay, India; or Punjabi population in Lahore, Pakistan.

In any configuration of the invention, in one embodiment, each human population is selected from a population marked "(a)" above.

In any configuration of the invention, in another embodiment, each human population is selected from a population marked "(b)" above.

In any configuration of the invention, in another embodiment, each human population is selected from a population marked "(c)" above.

In one embodiment of the library of the vertebrate, cell or library of the invention, the first and second ethnic populations are selected from the group consisting of an ethnic population with



European ancestry, an ethnic population with East Asian, an ethnic population with West African ancestry, an ethnic population with Americas ancestry and an ethnic population with South Asian ancestry.

In one embodiment of the library of the vertebrate, cell or library of the invention, the first and second ethnic populations are selected from the group consisting of an ethnic population with Northern European ancestry; or an ethnic population with Western European ancestry; or an ethnic population with Toscani ancestry; or an ethnic population with British ancestry; or an ethnic population with Icelandic ancestry; or an ethnic population with Finnish ancestry; or an ethnic population with Iberian ancestry; or an ethnic population with Japanese ancestry; or an ethnic population with Chinese ancestry; or an ethnic population Vietnamese ancestry; or an ethnic population with Yoruba ancestry; or an ethnic population with Luhya ancestry; or an ethnic population with Gambian ancestry; or an ethnic population with Malawian ancestry; or an ethnic population with Native American ancestry; or an ethnic population with Afro-Caribbean ancestry; or an ethnic population with Mexican ancestry; or an ethnic population with Puerto Rican ancestry; or an ethnic population with Columbian ancestry; or an ethnic population with Peruvian ancestry; or an ethnic population with Ahom ancestry; or an ethnic population with Kayadtha ancestry; or an ethnic population with Reddy ancestry; or an ethnic population with Maratha; or an ethnic population with Punjabi ancestry.

In one embodiment of any configuration of the vertebrate, cell or library of the invention, the human immunoglobulin gene segment derived from the genome sequence of the second individual is low-frequency (optionally rare) within the second ethnic population. Optionally human immunoglobulin gene segment has a Minor Allele Frequency (MAF) (cumulative frequency) of between 0.5% - 5%, optionally less than 0.5%, in the second human population, eg, as in the 1000 Genomes database.

In one embodiment of any configuration of the vertebrate, cell or library of the invention, the first variable region nucleotide sequence is produced by recombination of the first human immunoglobulin gene segment with a first J gene segment and optionally a first D gene segment, wherein the first human immunoglobulin gene segment is a V gene segment and the V, D and J segments are derived from the first human population, optionally from the genome of one individual

of the first human population.

In one embodiment of the library of the vertebrate, cell or library of the invention, the second variable region nucleotide sequence is produced by recombination of the second human immunoglobulin gene segment with a second J gene segment and optionally a second D gene segment, wherein the second human immunoglobulin gene segment is a V gene segment derived from the second population and the D and/or J segments are derived from the first human population, optionally the D and J gene segments being from the genome of one individual of the first human population.

In one embodiment of the library of the vertebrate, cell or library of the invention, all of the D and J segments that have been recombined with the first and second V gene segments are D and J segments derived from the first human population, optionally the D and J gene segments being from the genome of one individual of the first human population.

In one embodiment of the library, the second human immunoglobulin gene segment is a polymorphic variant of the first human immunoglobulin gene segment; optionally wherein the second gene segment is selected from the group consisting of a gene segment in any of Tables 1 to 7 and 9 to 14 (eg, selected from Table 13 or 14).

In one embodiment of the library, the first and second human immunoglobulin gene segments are both (i)  $V_H$  gene segments; (ii) D segments; (iii) J segments (optionally both  $J_H$  segments, both  $J_K$  segments or both  $J_\lambda$  segments); (iv) constant regions (optionally both a gamma constant region, optionally both a C gamma-1 constant region); (v) CH1 regions; (vi) CH2 regions; or (vii) CH3 regions.

The library is, for example, a naive and optionally has a library size of from 10 or  $10^2$  to  $10^9$  cells. For example, from 10, 20, 30, 40, 50, 100 or 1000 to  $10^8$ ; or 10, 20, 30, 40, 50, 100 or 1000 to  $10^7$ ; or 10, 20, 30, 40, 50, 100 or 1000 to  $10^6$ ; or 10, 20, 30, 40, 50, 100 or 1000 to  $10^5$ ; or 10, 20, 30, 40, 50, 100 or 1000 to  $10^4$  cells.

The library has, for example, been selected against a predetermined antigen and optionally has a library size of from 10 or  $10^2$  to  $10^9$  cells. For example, from 10, 20, 30, 40, 50, 100 or 1000 to  $10^8$ ; or 10, 20, 30, 40, 50, 100 or 1000 to  $10^7$ ; or 10, 20, 30, 40, 50, 100 or 1000 to  $10^6$ ; or 10, 20, 30, 40, 50, 100 or 1000 to  $10^5$ ; or 10, 20, 30, 40, 50, 100 or 1000 to  $10^4$  cells.

In one embodiment of the library of the invention, said first and second cells are progeny of first and second ancestor non-human vertebrate cells respectively, wherein the first ancestor cell comprises a genome comprising said first human immunoglobulin gene segment; and the second ancestor cell comprises a genome comprising said second human immunoglobulin gene segment.

The invention further provides a library of antibody-producing transgenic cells whose genomes collectively encode a repertoire of antibodies, wherein the library comprises the first and second ancestor cells described above.

The invention further provides a library of hybridoma cells produced by fusion of the library of the invention (eg, a B-cell library) with fusion partner cells and optionally has a library size of from 10 or  $10^2$  to  $10^9$  cells. For example, from 10, 20, 30, 40, 50, 100 or 1000 to  $10^8$ ; or 10, 20, 30, 40, 50, 100 or 1000 to  $10^7$ ; or 10, 20, 30, 40, 50, 100 or 1000 to  $10^6$ ; or 10, 20, 30, 40, 50, 100 or 1000 to  $10^5$ ; or 10, 20, 30, 40, 50, 100 or 1000 to  $10^4$  cells. Production of hybridomas is well known to the skilled person. Examples of fusion partners are SP2/0-g14 (obtainable from ECACC), P3X63-Ag8.653 (obtainable from LGC Standards; CRL-1580), NS1 and NS0 cells. PEG fusion or electrofusion can be carried out, as is conventional.

**The invention provides, in a third configuration:-**

An isolated antibody having

- (a) a heavy chain encoded by a nucleotide sequence produced following recombination in a transgenic non-human vertebrate cell of an unrearranged human immunoglobulin V gene segment with a human D and human J segment, optionally with affinity maturation in said cell, wherein one of the gene segments (eg, VH) is derived from the genome of an individual from a first human ethnic population; and the other two gene segments (eg, D and JH) are derived from the genome of an individual from a second (eg, a second and third respectively), different, human ethnic population,

and wherein the antibody comprises heavy chain constant regions (eg, C gamma) of said non-human vertebrate (eg, rodent, mouse or rat heavy chain constant regions); and/or

- (b) a light chain encoded by a nucleotide sequence produced following recombination in a transgenic non-human vertebrate cell of an unrearranged human immunoglobulin V gene segment with a human J segment, optionally with affinity maturation in said cell, wherein one of the gene segments (eg, VL) is derived from the genome of an individual from a first human ethnic population (optionally the same as the first population in (a)); and the other gene segment (eg, JL) is derived from the genome of an individual from a second, different, human ethnic population (optionally the same as the second population in (a)), and wherein the antibody comprises light chain constant regions of said non-human vertebrate (eg, rodent, mouse or rat heavy light constant regions);
- (c) Optionally wherein each variable domain of the antibody is a human variable domain.
- (d) Optionally wherein the heavy chain constant regions are mu- or gamma-type constant regions.

The invention also provides an isolated nucleotide sequence encoding the antibody of the third configuration, optionally wherein the sequence is provided in an antibody expression vector, optionally in a host cell. Suitable vectors are mammalian expression vectors (eg, CHO cell vectors or HEK293 cell vectors), yeast vectors (eg, a vector for expression in *Picchia pastoris*, or a bacterial expression vector, eg, a vector for *E. coli* expression).

The invention also provides a method of producing a human antibody, the method comprising replacing the non-human vertebrate constant regions of the antibody of the third configuration with human antibody constant regions (eg, a C variant disclosed in table 13 or 18). The skilled person will be aware of standard molecular biology techniques to do this. For example, see Harlow, E. & Lane, D. 1998, 5<sup>th</sup> edition, *Antibodies: A Laboratory Manual*, Cold Spring Harbor Lab. Press, Plainview, NY; and Pasqualini and Arap, *Proceedings of the National Academy of Sciences* (2004) 101:257-259 for standard immunisation. Joining of the variable regions of an antibody to a human constant region can be effected by techniques readily available in the art, such as using conventional recombinant DNA and RNA technology as will be apparent to the skilled person. See e.g. Sambrook, J and Russell, D. (2001, 3<sup>d</sup> edition) *Molecular Cloning: A Laboratory Manual* (Cold Spring Harbor Lab. Press, Plainview, NY).

In one embodiment, the method comprises further making a mutant or derivative of the antibody.

The invention also provides a pharmaceutical composition comprising an antibody according to the third configuration, or a human antibody of the invention and a diluent, excipient or carrier; optionally wherein the composition is provided in a container connected to an IV needle or syringe or in an IV bag.

The invention also provides an antibody-producing cell (eg, a mammalian cell, eg, CHO or HEK293; a yeast cell, eg, *P pastoris*; a bacterial cell, eg, *E coli*; a B-cell; or a hybridoma) that expresses the second antibody of the third configuration or the isolated antibody of the invention.

The first configuration of the invention also provides:-

A non-human vertebrate or vertebrate cell (optionally an ES cell or antibody-producing cell) whose genome comprises a transgenic immunoglobulin locus (eg, a heavy chain locus or a light chain locus), said locus comprising immunoglobulin gene segments according to the first and second human immunoglobulin gene segments (optionally V segments) described above in connection with the third configuration. The gene segments are operably connected upstream of an immunoglobulin constant region; optionally wherein the genome is homozygous for said transgenic immunoglobulin locus.

Optionally the immunoglobulin locus comprises more than the natural human complement of functional V gene segments; and/or

Optionally wherein the immunoglobulin locus comprises more than the natural human complement of functional D gene segments; and/or

Optionally wherein the immunoglobulin locus comprises more than the natural human complement of functional J gene segments.

In this way, a superhuman immunoglobulin gene repertoire is provided in a transgenic non-human vertebrate or vertebrate cell according to the invention.

The first configuration also provides:-

A transgenic non-human vertebrate (eg, a mouse or rat) or vertebrate cell (optionally an ES cell or antibody-producing cell) whose genome comprises a transgenic immunoglobulin locus comprising a plurality of human immunoglobulin gene segments operably connected upstream of a non-human vertebrate constant region for the production of a repertoire of chimaeric antibodies, or chimaeric light or heavy chains, having a non-human vertebrate constant region and a human variable region; wherein the transgenic locus comprises one or more human immunoglobulin V gene segments, one or more human J gene segments and optionally one or more human D gene segments, a first (optionally a V segment) of said gene segments and a second (optionally a V segment) of said gene segments being different and derived from the genomes of first and second human individuals respectively, wherein the individuals are different; and optionally not related;

optionally wherein the immunoglobulin locus comprises more than the natural human complement of functional V gene segments; and/or

optionally wherein the immunoglobulin locus comprises more than the natural human complement of functional D gene segments; and/or

optionally wherein the immunoglobulin locus comprises more than the natural human complement of functional J gene segments.

In this way, a superhuman immunoglobulin gene repertoire is provided in a transgenic non-human vertebrate or vertebrate cell according to the invention.

The first configuration also provides:-

A transgenic non-human vertebrate (eg, a mouse or rat) or vertebrate cell (optionally an ES cell or antibody-producing cell) whose genome comprises first and second transgenic immunoglobulin loci, each locus comprising a plurality of human immunoglobulin gene segments operably connected upstream of a non-human vertebrate constant region for the production of a repertoire of chimaeric antibodies, or chimaeric light or heavy chains, having a non-human vertebrate constant region and a human variable region;

wherein (i) the first transgenic locus comprises one or more human immunoglobulin V gene segments, one or more human J gene segments and optionally one or more human D gene segments, (ii) the second transgenic locus comprises one or more human immunoglobulin V gene segments, one or more human J gene segments and optionally one or more human D gene segments; and (iii) wherein a first (optionally a V) gene segment of said first locus and a second (optionally a V) gene segment of said second gene locus are different and derived from the genomes of first and second human individuals respectively, wherein the individuals are different; and optionally not related;

optionally wherein the first and second loci are on different chromosomes (optionally chromosomes with the same chromosome number) in said genome;

optionally wherein each immunoglobulin locus comprises more than the natural human complement of functional V gene segments; and/or

optionally wherein each immunoglobulin locus comprises more than the natural human complement of functional D gene segments; and/or

optionally wherein each immunoglobulin locus comprises more than the natural human complement of functional J gene segments.

In this way, a superhuman immunoglobulin gene repertoire is provided in a transgenic non-human vertebrate or vertebrate cell according to the invention.

In these embodiments of the first configuration, the immunoglobulin gene segments are optionally as described for the third configuration.

In these embodiments of the first configuration, the genome optionally comprises a third immunoglobulin gene segment (optionally a V segment), the third gene segment being derived from a human individual that is different from the individual from which the first (and optionally also the second) gene segment is derived; optionally wherein the first, second and third gene segments are polymorphic variants of a human immunoglobulin gene segment (eg, VH1-69 – see the examples for

further description).

In these embodiments of the first configuration, the genome of the vertebrate or cell is optionally homozygous for the first, second and optional third gene segment, wherein a copy of the first, second and optional third gene segments are provided together on the same chromosome operably connected upstream of a common non-human vertebrate constant region.

For example, each first, second and optional third gene segment is a V gene segment.

In one example, the library of the invention is provided by a collection of non-human vertebrates (optionally a collection of rodents, mice or rats); optionally, wherein a first member of said collection produces said first antibody but not said second antibody, and a second member of the collection produces said second antibody (but optionally not said first antibody). It is therefore contemplated to make non-human vertebrates where different human genomes have been used as a source for building the transgenic loci in the vertebrates. For example, a first vertebrate comprises a transgenic heavy chain locus having gene segments only from a first (and optionally a second) human population or individual; a second vertebrate comprises a transgenic heavy chain locus having gene segments only from a third (and optionally a fourth) human population or individual; and optionally third and more vertebrates can be built similarly based on unique or overlapping human population genomes. However, when provided as a mixed population of transgenic vertebrates, the mixed population provides a collective pool of human immunoglobulin genes that is greater than found in a natural human repertoire. This is useful to extend the antibody and gene sequence space beyond those possible with prior transgenic mice and rats bearing human immunoglobulin loci. As explained above, these have been based on a single human genome.

In one embodiment, the collection of non-human vertebrates bear human immunoglobulin genes confined to human populations that are together grouped under the same population genus "(a)" mentioned above. This provides for a gene repertoire that is biased to producing human antibody variable regions prevalent in the population genus (a) and thus useful for generating antibody therapeutics/prophylactics for members of said population. Alternatively, where gene segments from different human populations are provided in a single transgene according to the invention (not necessarily in a collection of vertebrates), the different human populations are for example together



grouped under the same population genus "(a)" mentioned above.

The invention also provides a repertoire of antibodies expressed from a library of cells according to the invention.

In the non-human vertebrate or cell of any configuration of the invention, the constant region of the transgenic locus is, in one example, an endogenous constant region of said vertebrate (eg, endogenous mouse or rat constant region, eg, from the same strain of mouse or rat as the non-human vertebrate itself).

The invention also provides a method of constructing a cell (eg, an ES cell) according to the invention, the method comprising

- (a) identifying functional V and J (and optionally D) gene segments of the genome sequence of a (or said) first human individual;
- (b) identifying one or more functional V and/or D and/or J gene segments of the genome sequence of a (or said) second human individual, wherein these additional gene segments are not found in the genome sequence of the first individual;
- (c) and constructing a transgenic immunoglobulin locus in the cell, wherein the gene segments of (a) and (b) are provided in the locus operably connected upstream of a constant region.

Optionally the cell comprises a heavy chain locus constructed according to steps (a) to (c) and/or a light chain locus (kappa and/or lambda loci) constructed according to steps (a) to (c).

Optionally the cell is homozygous for the or each transgenic locus; optionally wherein antibody expression from loci endogenous to said cell has been inactivated. This is useful for confining the functional antibody gene repertoire, and thus antibody production, to antibodies bearing human variable regions.

Optionally the gene segment(s) in step (b) are identified from an immunoglobulin gene database selected from the 1000 Genomes, Ensembl, Genbank and IMGT databases.

Optionally the first and second human individuals are members of first and second ethnic populations respectively, wherein the populations are different, optionally wherein the human immunoglobulin gene segment derived from the genome sequence of the second individual is low-frequency (optionally rare) within the second ethnic population.

The invention also provides a method of making a transgenic non-human vertebrate (eg, a mouse or rat), the method comprising

- (a) constructing an ES cell (eg, a mouse C57BL/6N, C57BL/6J, 129S5 or 129Sv strain ES cell) by carrying out the method above;
- (b) injecting the ES cell into a donor non-human vertebrate blastocyst (eg, a mouse C57BL/6N, C57BL/6J, 129S5 or 129Sv strain blastocyst);
- (c) implanting the blastocyst into a foster non-human vertebrate mother (eg, a C57BL/6N, C57BL/6J, 129S5 or 129Sv strain mouse); and
- (d) obtaining a child from said mother, wherein the child genome comprises a transgenic immunoglobulin locus.

The invention provides a transgenic non-human vertebrate (eg, a mouse or rat) made by the method or a progeny thereof. The invention also provides a population of such non-human vertebrates.

Microinjection of ES cells into blastocysts and generation of transgenic mice thereafter are conventional practices in the state of the art, and the skilled person is aware of techniques useful to effect this. C57BL/6N, C57BL/6J, 129S5 or 129Sv mouse strains and ES cells are readily and publicly available.

The invention also provides a method of isolating an antibody that binds a predetermined antigen (eg, a bacterial or viral pathogen antigen), the method comprising

- (a) providing a vertebrate (optionally a mouse or rat) according to the invention;
- (b) immunising (eg, using a standard prime-boost method) said vertebrate with said antigen (optionally wherein the antigen is an antigen of an infectious disease pathogen);
- (c) removing B lymphocytes from the vertebrate and selecting one or more B lymphocytes expressing antibodies that bind to the antigen;
- (d) optionally immortalising said selected B lymphocytes or progeny thereof, optionally by

producing hybridomas therefrom; and

(e) isolating an antibody (eg, and IgG-type antibody) expressed by the B lymphocytes; and

(f) optionally producing a derivative or variant of the antibody.

This method optionally further comprises after step (e) the step of isolating from said B lymphocytes nucleic acid encoding said antibody that binds said antigen; optionally exchanging the heavy chain constant region nucleotide sequence of the antibody with a nucleotide sequence encoding a human or humanised heavy chain constant region and optionally affinity maturing the variable region of said antibody; and optionally inserting said nucleic acid into an expression vector and optionally a host.

### **Bioinformatics Analysis & Selection of Immunoglobulin Gene Segments**

See also the discussion on variation analysis above.

The skilled person will know of sources of human antibody gene sequences, such as IMGT ([www.imgt.org](http://www.imgt.org)), GenBank ([www.ncbi.nlm.nih.gov/genbank](http://www.ncbi.nlm.nih.gov/genbank)) Bioinformatics tools for database manipulation are also readily available and known to the skilled person, eg, as publicly available from the 1000 Genomes Project/EBI ([www.1000genomes.org](http://www.1000genomes.org))

As a source of antibody gene segment sequences, the skilled person will also be aware of the following available databases and resources (including updates thereof):-

1.1. The Kabat Database (G. Johnson and T. T.Wu, 2002; <http://www.kabatdatabase.com>).

Created by E. A. Kabat and T. T. Wu in 1966, the Kabat database publishes aligned sequences of antibodies, T-cell receptors, major histocompatibility complex (MHC) class I and II molecules, and other proteins of immunological interest. A searchable interface is provided by the SeqhuntII

tool, and a range of utilities is available for sequence alignment, sequence subgroup classification, and the generation of variability plots. See also Kabat, E. A., Wu, T. T., Perry, H., Gottesman, K., and Foeller, C. (1991) *Sequences of Proteins of Immunological Interest*, 5th ed., NIH Publication No. 91-

3242, Bethesda, MD, which is incorporated herein by reference, in particular with reference to human gene segments for use in the present invention.

1.2. KabatMan (A. C. R. Martin, 2002; <http://www.bioinf.org.uk/abs/simkab.html>). This is a web interface to make simple queries to the Kabat sequence database.

1.3. *IMGT, the International ImMunoGeneTics Information System*<sup>®</sup>; M.-P. Lefranc, 2002; <http://imgt.cines.fr>. IMGT is an integrated information system that specializes in antibodies, T cell receptors, and MHC molecules of all vertebrate species. It provides a common portal to standardized data that include nucleotide and protein sequences, oligonucleotide primers, gene maps, genetic polymorphisms, specificities, and two-dimensional (2D) and three-dimensional (3D) structures. IMGT includes three sequence databases (*IMGT/LIGM-DB*, *IMGT/MHC-DB*, *IMGT/PRIMERDB*), one genome database (*IMGT/GENE-DB*), one 3D structure database (*IMGT/3Dstructure-DB*), and a range of web resources (“*IMGT Marie-Paule page*”) and interactive tools.

1.4. *V-BASE* (I. M. Tomlinson, 2002; <http://www.mrc-cpe.cam.ac.uk/vbase>). V-BASE is a comprehensive directory of all human antibody germline variable region sequences compiled from more than one thousand published sequences. It includes a version of the alignment software DNAPLOT (developed by Hans-Helmar Althaus and Werner Müller) that allows the assignment of rearranged antibody V genes to their closest germline gene segments.

1.5. *Antibodies—Structure and Sequence* (A. C. R. Martin, 2002; <http://www.bioinf.org.uk/abs>). This page summarizes useful information on antibody structure and sequence. It provides a query interface to the Kabat antibody sequence data, general information on antibodies, crystal structures, and links to other antibody-related information. It also distributes an automated summary of all antibody structures deposited in the Protein Databank (PDB). Of particular interest is a thorough description and comparison of the various numbering schemes for antibody variable regions.

1.6. *AAAAA—AHO’s Amazing Atlas of Antibody Anatomy* (A. Honegger, 2001; <http://www.unizh.ch/~antibody>). This resource includes tools for structural analysis, modeling, and engineering. It adopts a unifying scheme for comprehensive structural alignment of antibody and T-

cell-receptor sequences, and includes Excel macros for antibody analysis and graphical representation.

1.7. WAM—Web Antibody Modeling (N. Whitelegg and A. R. Rees, 2001; <http://antibody.bath.ac.uk>). Hosted by the Centre for Protein Analysis and Design at the University of Bath, United Kingdom. Based on the AbM package (formerly marketed by Oxford Molecular) to construct 3D models of antibody Fv sequences using a combination of established theoretical methods, this site also includes the latest antibody structural information.

1.8. Mike's Immunoglobulin Structure/Function Page (M. R. Clark, 2001; <http://www.path.cam.ac.uk/~mrc7/mikeimages.html>) These pages provide educational materials on immunoglobulin structure and function, and are illustrated by many colour images, models, and animations. Additional information is available on antibody humanization and Mike Clark's Therapeutic Antibody Human Homology Project, which aims to correlate clinical efficacy and anti-immunoglobulin responses with variable region sequences of therapeutic antibodies.

1.9. The Antibody Resource Page (The Antibody Resource Page, 2000; <http://www.antibodyresource.com>). This site describes itself as the "complete guide to antibody research and suppliers." Links to amino acid sequencing tools, nucleotide antibody sequencing tools, and hybridoma/cell-culture databases are provided.

1.9. Humanization by Design (J. Saldanha, 2000; <http://people.cryst.bbk.ac.uk/~ubca07s>). This resource provides an overview on antibody humanization technology. The most useful feature is a searchable database (by sequence and text) of more than 40 published humanized antibodies including information on design issues, framework choice, framework back-mutations, and binding affinity of the humanized constructs.

See also Antibody Engineering Methods and Protocols, Ed. Benny K C Lo, Methods in Molecular Biology™, Human Press. Also at <http://www.blogsua.com/pdf/antibody-engineering-methods-and-protocolsantibody-engineering-methods-and-protocols.pdf>

As a source of genomic sequence variation data, the skilled person will also be aware of the following available databases and resources (including updates thereof):-

1. HapMap (The International HapMap Consortium. 2003; <http://hapmap.ncbi.nlm.nih.gov/index.html.en>). The HapMap Project is an international project that aims to compare the genetic sequences of different individuals to identify chromosomal regions containing shared genetic variants. The HapMap www site provides tools to identify chromosomal regions and the variant therein, with options to drill down to population level frequency data.
2. 1000 Genomes ([The 1000 Genomes Project Consortium](http://www.1000genomes.org/) 2010; <http://www.1000genomes.org/>). This resource provides complete genomic sequence for 2500 unidentified individuals from one of 25 distinct population groups, with the aim of identifying genomic variants of >1%. The site provides the ability to interrogate data utilizing online tools (e.g. 'Variation Pattern Finder') and to download variant data for individual population groups.
3. Japanese SNP Database (H.Haga et al. 2002; <http://snp.ims.u-tokyo.ac.jp/index.html>). Based on a study identifying 190,562 human genetic variants this site catalogues genomic variants with useful features for searching and summarizing data.

It is possible to identify variants in immunoglobulin genes classed as low-frequency or rare variants that segregate with specific human ethnic populations. For the purpose of this analysis, a low-frequency immunoglobulin gene segment is classed as one with 'Minor Allele Frequency' (MAF) (cumulative frequency) of between 0.5% - 5%, rare variants are those classed as having a MAF of less than 0.5% in a particular human population.

The following bioinformatics protocol is envisaged to identify human immunoglobulin gene segments for use in the present invention:

- (a) Identify one or more genomic regions containing gene segments of interest ('target genomic regions') and calculate the genomic coordinates, using coordinates that match the sequence assembly build used by either the 1000 Genomes project or International HapMap project (or another selected human gene database of choice).
- (b) Identify genomic variants mapped to the genomic regions previously identified in (a). Retrieve variant frequencies for variants for each super population and preferably sub-population where such data is available. Tools readily available on the HapMap WWW site and the VWC tools for the 1000Genomes Project are useful for this step.
- (c) Filter list of genomic variants from target genomic regions to contain only variants classed as either 'Non-synonymous' single nucleotide polymorphisms (SNPs) or genomic 'insertions or deletions' (indels). Filter further to include those that are present in exonic sequences only.
- (d) Correlate population frequency data for each of the identified variants for each of the super populations (for example 'European Ancestry', 'East Asian ancestry', 'West African ancestry', 'Americas', and 'South Asian ancestry') to identify those variants that segregate with less than two super-populations. Further correlate all identified variants with each of the sub-populations (for example, 'European ancestry' super-population might be subdivided into groups such as 'CEU – Utah residents with Northern or Western European ancestry', 'TSI Toscani in Italia' and 'British from England and Scotland') and produce a second score for rarity of variants in within a super-population.
- (e) Collect one or more gene segments that show segregation to specific sub-populations for construction of synthetic loci according to the invention.

In one embodiment throughout the present text, “germline” refers to the canonical germline gene segment sequence.

By detailed analysis of the 1000 Genomes database, the inventors have devised a collection of candidate polymorphic antibody gene segment variants, eg, human variant JH gene segments (eg, see Example 4), that can be built into the design of transgenic heavy chain loci in mice for expressing increasingly diverse and new, synthetic repertoires of human variable regions. To this end, the invention provides the following embodiments.

The present invention provides in a fourth configuration:-

#### Selection of Human JH6\*02 Variant

#### Transgenic IgH Loci, Non-Human Vertebrates, Cells & Antibodies Based on Human JH6\*02

As explained above, in designing transgenic Ig heavy chain loci the present inventors have considered the huge amount of data available from the 1000 Genomes project (see [www.1000genomes.org](http://www.1000genomes.org)) that analyses gene distributions amongst many human populations, and in particular data on Ig gene segments. The inventors were also aware of human gene segments disclosed in the IMGT database (see [www.imgt.org](http://www.imgt.org)) and in Ensembl (see [www.ensembl.org](http://www.ensembl.org)). The inventors needed to make choices about which human gene segments to include amongst the large number of human gene segments presented in these databases and the other sources of human Ig gene segment information known in the art, including those other databases disclosed herein. When choosing human JH gene segments, the inventors were aware that human JH6 encodes a relatively long amino acid sequence, and thus the inventors thought it desirable to include this for increasing the chances of producing IgH chains with relatively long HCDR3 regions. Antibodies with long HCDR3 (at least 20 amino acids according to IMGT nomenclature) have been shown to neutralise a variety of pathogens effectively including HIV, Influenza virus, malaria and Africa trypanosomes. Reference is also made to naturally-occurring Camelid (eg, llama or camel) heavy chain-only antibodies which bear long HCDR3s for reaching relatively inaccessible epitopes (see, eg, EP0937140). Long HCDR3s can form unique stable subdomains with extended loop structure that towers above the antibody



surface to confer fine specificity. In some cases, the long HCDR3 itself is sufficient for epitope binding and neutralization (Liu, L *et al*; Journal of Virology. 2011. 85: 8467-8476, incorporated herein by reference). The unique structure of the long HCDR3 allows it to bind to cognate epitopes within inaccessible structure or extensive glycosylation on a pathogen surface. In human peripheral blood, there is around 3.5% of naïve B antibodies or 1.9% of memory B IgG antibodies containing the HCDR3s with lengths of more than 24 amino acids (PLoS One. 2012;7(5):e36750. Epub 2012 May 9; “Human peripheral blood antibodies with long HCDR3s are established primarily at original recombination using a limited subset of germline genes”; Briney BS *et al*, incorporated herein by reference) (Fig. 1). The usage analysis indicates that these antibodies have the preference to use human JH6 with human D2-2, D3-3 or D2-15 (Brinley, BS *et al*, Figs. 2-5). See also PLoS One. 2011 Mar 30;6(3):e16857; Comparison of antibody repertoires produced by HIV-1 infection, other chronic and acute infections, and systemic autoimmune disease”; Breden F *et al*, incorporated herein by reference. Around 20% of all HCDR3 of antibodies use JH6. However, in those antibodies with HCDR3 of more than 24 amino acids, 70% use JH6 (Brinley, BS *et al*, Fig.2).

There is a need in the art for genetically modified non-human vertebrates and cells that can make antibodies and heavy chains that have long human HCDR3s, as well as antibodies, chains and VH domains that can be selected from such vertebrates and cells wherein these can address target epitopes better accessed by long HCDR3s.

The inventors, therefore, chose in this configuration of the invention to include a human JH6 gene segment as a mandatory human gene segment in their IgH locus design. Several different naturally-occurring human JH6 variants are known (eg, JH6\*01 to \*04 as well as others; IMGT nomenclature). The inventors considered this when deciding upon which human JH6 variant should be included in the transgenic IgH locus design. An alignment of some human JH6 variants is shown in Figure 7 (from [www.imgt.org](http://www.imgt.org); dashes indicate identical nucleotides; nucleotide changes versus the \*01 variant are shown by underlined nucleotides and corresponding amino acid changes are shown by underlined amino acids; Genbank accession numbers (release 185.0) are shown prefixed by J, X, M or A). The inventors used sequencing of human genomic DNA samples, inspection of public IgH DNA databases as well as informed choices on the basis of variant sequences as means to arrive at a rational choice of which JH6 variant to use.

The 1000 Genomes database uses human JH6\*03 as the reference sequence, which would be a possible choice for the skilled person wishing to construct a transgenic IgH locus. The inventors noticed (eg, Figure 7 herein) that position 6 in JH6\*03 is a tyrosine (Y) encoded by a TAC codon, whereas some other naturally-occurring human variants have a glycine (G) encoded by a GGT codon (the glycine being present as a YYG motif, forming part of a larger YYGXDX motif). To understand the potential significance of this, the inventors carried out analysis of JH sequences from other vertebrate species. The inventors surprisingly noticed that YYG and YYGXDX motifs are conserved across many vertebrate species (see Figures 7 & 8). This suggested to the inventors, therefore, that preservation of this motif might be desirable, which could guide the choice of JH6 variant for use in the present invention.

Another pointer arose when the inventors considered the TAC codon versus the GGT codon encoding Y or G respectively. The inventors considered the impact of these nucleotide sequences on the action of activation-induced cytidine deaminase (AID). The inventors knew that activation-induced cytidine deaminase (AID) is believed to initiate Ig somatic hypermutation (SHM) in a multi-step mechanism and they addressed this activity when rationally designing the locus. AID catalyses the deamination of C to U in DNA, generating mutations at C bases. Cytidines located within hotspot motifs are preferentially deaminated. Certain motifs are hotspots for AID activity (DGYW, WRC, WRCY, WRCH, RGYW, AGY, TAC, WGCW, wherein W=A or T, Y=C or T, D=A, G or T, H=A or C or T, and R=A or G). The presence of a TAC codon encoding Y at position 6 in JH6\*03 creates AID mutation hotspots (the cytidine being the substrate of AID), these hotspots being the underlined motifs in the previous sentence. The inventors considered the impact of this and in doing so they considered possible mutants created by AID activity at the cytidine. Reference is made to Figure 9. The inventors noticed that a mutation at the third base of the TAC codon would yield 3 possible outcomes: Y, stop or stop. Thus, out of the three stop codons possible in the genetic code (the other being encoded by TGA – see Figure 9), two of them would be provided by mutation of the cytidine in the TAC codon encoding position 6 in JH6\*03. The inventors, therefore, considered that this might increase the chances of non-productive IgH variable region production in transgenic loci based on JH6\*03. Moreover, the inventors noticed that provision of a GGT codon instead (as per the other human JH6 variants) seemed preferable since mutation of the third base would never yield a stop codon (see Figure 9), and furthermore would retain coding, and thus conservation, of glycine at position 6, which the inventors also noticed was is in the YYG and YYGXDX motifs conserved across species.

Having decided against using JH6\*03, the inventors needed to make a choice from other possible human variants. The MDV motif is at the C-terminus of HCDR3 based on human JH6, the adjacent framework 4 (FW4) starting with the WGQ motif (with reference to the sequence shown encoded by JH6\*01; Figure 7). In making their choices for locus design, the inventors wished to maximise conservation of this HCDR3/FW4 junction in product IgH chains and antibodies including these. The inventors believed this to be desirable for heavy chain variable domain functionality and conformation. The inventors thought that this might in some cases be desirable to minimise immunogenicity (suitable for human pharmaceutical use). Consistent with these considerations, the inventors wanted to make a choice that would minimise mutation around the HCDR3/FW4 junction as a result of SHM *in vivo* to conserve junction configuration. See Rogozin & Diaz; "Cutting Edge: DGYW/WRCH Is a Better Predictor of Mutability at G:C Bases in Ig Hypermutation Than the Widely Accepted RGYW/WRCY Motif and Probably Reflects a Two-Step Activation-Induced Cytidine Deaminase-Triggered Process"; Journal of Immunology; March 15, 2004 vol. 172 no. 6 3382-3384. An example of a DGYW motif is GGCA. The inventors had this in mind when analysing the variant sequences.

With these considerations in mind, the inventors decided specifically to use human JH6\*02 as the mandatory human JH6 for their IgH locus design. JH6\*01 was rejected as the mandatory JH6 gene segment since the nucleotide sequence GGG CAA (encoding G and Q) contains a GGCA motif which is an AID recognition hotspot. The inventors realised that JH6\*04 also contains such a motif due to the presence of the sequence GGC AAA encoding G and K (positions 11 and 12 respectively). The inventors also realised that the \*02 variant has a C instead of a G that is in the \*01 variant, the C desirably being a synonymous change (ie, not changing the encoded amino acid sequence around the CDR3/FW4 junction) and also this does not provide a GGCA AID hotspot motif. The inventors, therefore, decided that the mandatory JH6 should have this C base and this too pointed them to using the human JH6\*02 variant.

In one example of any configuration of the invention herein, the only JH6 species included in the locus or genome is human JH6\*02.

The inventors obtained 9 anonymised DNA samples from cheek swabs of 9 consenting human adults. Sequencing was performed on IgH locus DNA to confirm natural JH6\*02 variant usage. It was found that the genome of all 9 humans contained a JH6\*02 variant gene segment. In 7 out of the 9 humans, the genome was homozygous for JH6\*02 (ie, each chromosome 14 had JH6\*02 as its JH6 gene segment in the IgH locus). The inventors also inspected the publicly-available sequence information from the genomes of well-known scientists Craig Venter and Jim Watson. Both of these genomes contain JH6\*02 too. This indicated to the inventors that this variant is common in humans.

So, the inventors made a choice of human JH6\*02 on the basis of

- (i) Containing the YYG and YYGXDX motifs that is conserved across several vertebrate species;
- (ii) Provision of one less TAC codon (an AID hotspot that risks stop codons) and a choice instead of a codon that preserves the YYG and YYGXDX motifs;
- (iii) Avoidance of a GGCA AID hotspot in the region of the HCDR3/FW4 junction; and
- (iv) Common occurrence (and thus conservation and acceptability) in humans of the JH6\*02 variant.

This rationale was tested by the inventors in laboratory examples, in order to see if human JH6\*02 could desirably participate in antibody gene segment recombination and heavy chain production in a foreign (non-human vertebrate) setting, and moreover to assess if long HCDR3s based on human JH6\*02 could be produced *in vivo* (in naïve and immunised settings) in such non-human systems. It was noted that in some non-human settings, such as a mouse, the YYG and YYGXDX motifs are not conserved, and thus the inventors decided that it was important to test whether or not JH6\*02 (having the YYG and YYGXDX motifs) could function properly in such a foreign setting to participate in VDJ recombination and selection against antigen.

Thus, as explained further in the examples, the inventors constructed transgenic JH6\*02-containing IgH loci in ES cells, generated transgenic non-human vertebrates from the ES cells (both naïve and immunised with a range of different target antigen types), isolated antibodies and heavy chain sequences based on JH6\*02 as well as B-cells expressing these and made hybridomas expressing antigen-specific antibodies that are based on the chosen JH6\*02 variant. The inventors found that the JH6\*02 variant was extensively used and could contribute to the production of HCDR3 of at least

20 amino acids in many different heavy chains (including antigen-specific heavy chains). The chosen variant was preferably used over other JH gene segments in all settings (naïve, immunised and antigen-specific) for the production of HCDR3 of at least 20 amino acids.

Thus, the present invention provides an IgH locus including human JH6\*02 (IMGT nomenclature) as a mandatory JH gene segment. In one embodiment, the locus comprises non-human vertebrate (eg, mouse or rat) constant region gene segments downstream (ie, 3' of) the human JH6\*02; and one or more VH gene segments (eg, a plurality of human VH gene segments) and one or more D gene segments (eg, a plurality of human D gene segments) upstream of (ie, 5' of) the human JH6\*02. For example, the locus is comprised by a vector (eg, a DNA vector, eg, a yeast artificial chromosome (YAC), BAC or PAC). Such a vector (eg, YAC) can be introduced into a non-human vertebrate (eg, mouse or rat) cell using standard techniques (eg, pronuclear injection) so that the locus is integrated into the cell genome for expression of IgH chains comprising at least one chain whose variable domain is a product of the recombination of human JH6\*02 with a VH and a D gene segment.

In another example, the locus (eg, with a completely human, rat or mouse constant region, or a human/mouse chimaeric constant region) can be provided in the genome of a non-human vertebrate (eg, mouse or rat) cell. For example, the cell is an ES cell or an antibody-producing cell (eg, an isolated B-cell, an iPS cell or a hybridoma).

In another example, the invention provides a non-human vertebrate (eg, a mouse or a rat) comprising an IgH locus of the invention which comprises a human JH6\*02 gene segment, wherein the locus can express an IgH chain whose variable domain is a product of the recombination of human JH6\*02 with a VH and a D gene segment. As shown in the examples, the inventors have successfully produced such mice which produce such IgH chains with VH domains based on human JH6\*02. The inventors isolated and sequenced IgH chains from the mice before (naïve) and after (immunised) exposure to a range of target antigens and confirmed by comparison to IMGT IgH gene segment sequences that the isolated chains (and antibodies containing these) were produced based on JH6\*02. Such chains were found in naïve mice, as well as in antigen-specific antibodies from immunised mice. B-cells were isolated from immunised mice, wherein the B-cells express antibodies based on JH6\*02 and hybridomas were generated from the B-cells, the hybridomas expressing antigen-specific antibodies based on JH6\*02. The inventors, therefore, provided the locus,

vertebrate, cell and hybridoma of the invention based on the use of human JH6\*02 and showed that antibodies based on JH6\*02 and B-cells expressing these can be successfully produced and isolated following immunisation of the vertebrates, corresponding hybridomas being a good source of antibodies whose VH domains are based on JH6\*02, eg for administration to a patient, eg, for human medicine. Furthermore, it was found possible to produce and isolated antigen-specific antibodies whose VH domains are based on JH6\*02 and which had a relatively long HCDR3 (eg, 20 amino acids).

Thus, the present invention provides embodiments as in the following clauses:-

1. A non-human vertebrate (optionally a mouse or a rat) or vertebrate cell whose genome comprises an immunoglobulin heavy chain locus comprising human gene segment JH6\*02, one or more VH gene segments and one or more D gene segments upstream of a constant region; wherein the gene segments in the heavy chain locus are operably linked to the constant region thereof so that the mouse is capable of producing an antibody heavy chain produced by recombination of the human JH6\*02 with a D segment and a VH segment.

In another example, the invention provides

A non-human vertebrate (optionally a mouse or a rat) or vertebrate cell whose genome comprises an immunoglobulin heavy chain locus comprising one, more or all of human IGHV gene segments selected from V3-21, V3-13, V3-7, V6-1, V1-8, V1-2, V7-4-1, V1-3, V1-18, V4-4, V3-9, V3-23, V3-11 and V3-20 (eg, one, more or all of V3-21\*03, V3-13\*01, V3-7\*01, V6-1\*01, V1-8\*01, V1-2\*02, V7-4-1\*01, V1-3\*01, V1-18\*01, V4-4\*01, V3-9\*01 and V3-23\*04). These segments were found in naive repertoires to be productive to produce HCDR3s of at least 20 amino acids in length. In an embodiment, the locus comprises a human JH6, eg, JH6\*02.

The invention also provides a HCDR3, VH domain, antibody heavy chain or antibody having a HCDR3 size of at least 20 amino acids. Optionally, the HCDR3 or VH domain (or VH domain of the heavy chain or antibody) comprises mouse AID-pattern somatic hypermutations and/or mouse dTd-pattern mutations. This can be provided, for example, wherein VH domain is produced in a mouse comprising mouse AID and/or mouse TdT (eg, endogenous AID or TdT). See also Annu. Rev. Biochem. 2007. 76:1–22; Javier M. Di Noia and Michael S. Neuberger, “Molecular Mechanisms of Antibody Somatic Hypermutation” (in particular figure 1 and

associated discussion on AID hotspots in mouse); and Curr Opin Immunol. 1995 Apr;7(2):248-54, "Somatic hypermutation", Neuberger MS and Milstein C (in particular, discussion on hotspots in mouse), the disclosures of which are incorporated herein by reference.

These segments were found in naive repertoires to be productive in recombination with human JH6\*02 to produce HCDR3s of at least 20 amino acids in length.

In an example, the vertebrate is naïve. In another embodiment, the vertebrate instead is immunised with a target antigen.

In an example, the vertebrate or cell mentioned below is capable of so producing an antibody heavy chain upon immunisation with a target antigen. In an example, the vertebrate is an immunised vertebrate that produces antibody heavy chains specific for a target antigen and wherein the variable domains of the heavy chains are the product of recombination between a VH, D and JH6\*02. For example, the D is selected from human D3-3, D2-15, D3-9; D4-17; D3-10; D2-2; D5-24; D6-19; D3-22; D6-13; D5-12; D1-26; D1-20; D5-18; D3-16; D2-21; D1-14; D7-27; D1-1; D6-25; D2-14 and D4-23 (eg, selected from D3-9\*01; D4-17\*01; D3-10\*01; D2-2\*02; D5-24\*01; D6-19\*01; D3-22\*01; D6-13\*01; D5-12\*01; D1-26\*01; D1-20\*01; D5-18\*01; D3-16\*02; D2-21\*02; D1-14\*01; D7-27\*02; D1-1\*01; D6-25\*01; D2-15\*01; and D4-23\*01). For example, the D is human D3-9 or D3-10. In an example, the HCDR3 length is at least 20 amino acids (eg, 20, 21, 23 or 24).

In an example of the vertebrate or cell, the genome comprises additional human JH gene segments (eg, JH2, 3, 4 and 5 gene segments).

In an example of the vertebrate or cell, the genome comprises an immunoglobulin light chain locus comprising one or more human V gene segments and one or more human J gene segments upstream of a constant region (eg, a human or a mouse lambda or kappa constant region).

For rearrangement and expression of heavy chains, the locus comprises control elements, such as an E $\mu$  and S $\mu$  between the J gene segment(s) and the constant region as is known by the skilled person. In one example, a mouse E $\mu$  and S $\mu$  is included in the heavy chain locus between the JH6\*02 and the constant region (ie, in 5' to 3' order the locus comprises the JH6\*02, E $\mu$  and S $\mu$  and constant region). In an example, the E $\mu$  and S $\mu$  are E $\mu$  and S $\mu$  of a mouse 129-derived

genome (eg, a 129Sv-derived genome, eg, 129Sv/EV (such as 129S7Sv/Ev (such as from AB2.1 or AB2.2 cells obtainable from Baylor College of Medicine, Texas, USA) or 129S6Sv/Ev)); in another example, the E $\mu$  and S $\mu$  are E $\mu$  and S $\mu$  of a mouse C57BL/6 -derived genome. In this respect, the locus can be constructed in the IgH locus of the genome of a cell selected from AB2.1, AB2.2, VGF1, CJ7 and FH14. VGF1 cells were established and described in Auerbach W, Dunmore JH, Fairchild-Huntress V, *et al*; Establishment and chimera analysis of 129/SvEv- and C57BL/6-derived mouse embryonic stem cell lines. *Biotechniques* 2000; 29:1024–8, 30, 32, incorporated herein by reference.

Additionally or alternatively, the constant region (or at least a C $\mu$ ; or C $\mu$  and gamma constant regions thereof) is a constant region (or C $\mu$ ; or C $\mu$  and gamma constant regions thereof) is of a genome described in the paragraph immediately above.

A suitable source of JH6\*02 and other human DNA sequences will be readily apparent to the skilled person. For example, it is possible to collect a DNA sample from a consenting human donor (eg, a cheek swab sample as per the Example herein) from which can be obtained suitable DNA sequences for use in constructing a locus of the invention. Other sources of human DNA are commercially available, as will be known to the skilled person. Alternatively, the skilled person is able to construct gene segment sequence by referring to one or more databases of human Ig gene segment sequences disclosed herein.

2. The vertebrate of clause 1, wherein the vertebrate has been immunised with a target antigen and wherein the variable domain of the heavy chain is the product of recombination between a VH, D and JH6\*02 and wherein the HCDR3 length is at least 20 amino acids (eg, 20, 21, 23 or 24).

Optionally, the immunised vertebrate produces an antibody heavy chain specific for a target antigen and wherein the variable domain of the heavy chain is the product of recombination between a VH, D and JH6\*02 and wherein the HCDR3 length is at least 20 amino acids (eg, 20, 21, 23 or 24).

3. A non-human vertebrate cell (optionally a mouse cell or a rat cell) whose genome comprises an immunoglobulin heavy chain locus comprising human gene segment JH6\*02, one or more VH gene segments and one or more D gene segments upstream of a constant region; wherein the gene segments in the heavy chain locus are operably linked to the constant region thereof for



producing (eg, in a subsequent progeny cell) an antibody heavy chain produced by recombination of the human JH6\*02 with a D segment and a VH segment.

4. The cell of clause 3, which is an ES cell capable of differentiation into a progeny antibody-producing cell that expresses said heavy chain.
5. The vertebrate or cell of any preceding clause, wherein the heavy chain locus comprises a human JH6\*02 recombination signal sequence (RSS) operably connected 5' to the JH6\*02 gene segment.

For example, the native RSS-JH6\*02 sequence can be used to advantageously maintain the natural pairing between RSS and their JH gene segment. In this respect, the following sequence is used:-

ggtttttgtggggtgaggatggacattctgccattctggattactactactactacggtatggacgtctggggccaagggaccacggtcaccg  
tctcctcag (SEQ ID NO: 238)

RSSs have a common architecture: 9mer (eg, first underlined sequence above) followed by a 22bp spacer and then a 7mer (eg, second underlined sequence above). Spacers are 23bp +/- 1 normally, while the 9 and 7mer are more conserved.

6. The vertebrate or cell of clause 5, wherein the RSS is SEQ ID NO: 238 or a sequence having an identical 9mer and 7mer sequence flanking a sequence that is at least 70% identical to the 22mer sequence of SEQ ID NO: 238.
7. The vertebrate or cell of clause 6, wherein the RSS and JH6\*02 are provided as SEQ ID NO: 237.
8. The vertebrate or cell of any preceding clause, wherein the JH6\*02 is the only JH6-type gene segment in the genome.
9. The vertebrate or cell of any preceding clause, wherein the JH6\*02 is the closest JH gene segment to the constant region in the locus.

10. The vertebrate or cell of any preceding clause, wherein the locus comprises one, more or all human D gene segments D3-9; D4-17; D3-10; D2-2; D5-24; D6-19; D3-22; D6-13; D5-12; D1-26; D1-20; D5-18; D3-16; D2-21; D1-14; D7-27; D1-1; D6-25; D2-14; and D4-23.

For example, the locus comprises one, more or all of human D gene segments D3-9\*01; D4-17\*01; D3-10\*01; D2-2\*02; D5-24\*01; D6-19\*01; D3-22\*01; D6-13\*01; D5-12\*01; D1-26\*01; D1-20\*01; D5-18\*01; D3-16\*02; D2-21\*02; D1-14\*01; D7-27\*02; D1-1\*01; D6-25\*01; D2-15\*01; and D4-23\*01.

11. The vertebrate or cell of clause 10, wherein the locus comprises one, more or all human D gene segments D3-9, D3-10, D6-19, D4-17, D6-13, D3-22, D2-2, D2-25 and D3-3.

These D segments were found to be productive in recombination with human JH6\*02 to produce HCDR3s of at least 20 amino acids in length.

In an example, the locus comprises one, more or all human D gene segments D3-9, D3-10, D6-19, D4-17, D6-13 and D3-22 (for example one, more or all of D3-9\*01, D3-10\*01, D6-19\*01, D4-17\*01, D6-13\*01 and D3-22\*01). These D segments were found in naïve repertoires to be productive in recombination with human JH6\*02 to produce HCDR3s of at least 20 amino acids in length.

In an example, the locus comprises one, more or all human D gene segments D3-10, D6-19 and D1-26 (for example, one, more or all of D3-10\*01, D6-19\*01 and D1-26\*01). These D segments were found in immunised repertoires to be productive in recombination with human JH6\*02 to produce HCDR3s of at least 20 amino acids in length.

In an example, the locus comprises one, more or all human D gene segments D3-9 and D3-10 (for example, one, more or all of D3-9\*01 and D3-10\*01). These D segments were found in antigen-specific repertoires to be productive in recombination with human JH6\*02 to produce HCDR3s of at least 20 amino acids in length.

12. The vertebrate or cell of any preceding clause, wherein the locus comprises a plurality of human D gene segments and the JH6\*02 is in human germline configuration with respect to the 3'-most human D gene segment (or all of the human D segments comprised by the locus).

In an example, the 3'-most D gene segment is D7-27. In an example, the locus comprises all of human D gene segments from D1-1 to D7-27 as present in a germline human IgH locus (eg, as shown in the IMGT database).

Alternatively or additionally, the JH6\*02 is in human germline configuration with respect to one, more or all of the E $\mu$ , S $\mu$  and constant region (eg, C $\mu$ ).

13. The vertebrate or cell of any preceding clause, wherein the locus comprises one, more or all of IGHV gene segments selected from V3-21, V3-13, V3-7, V6-1, V1-8, V1-2, V7-4-1, V1-3, V1-18, V4-4, V3-9, V3-23, V3-11 and V3-20.

In an example, the locus comprises one, more or all human IGHV gene segments V3-21, V3-13, V3-7, V6-1, V1-8, V1-2, V7-4-1, V1-3, V1-18, V4-4, V3-9, V3-23 (for example, one, more or all of V3-21\*03, V3-13\*01, V3-7\*01, V6-1\*01, V1-8\*01, V1-2\*02, V7-4-1\*01, V1-3\*01, V1-18\*01, V4-4\*01, V3-9\*01 and V3-23\*04). These segments were found in naive repertoires to be productive in recombination with human JH6\*02 to produce HCDR3s of at least 20 amino acids in length.

In an example, the locus comprises one, more or all human IGHV gene segments V3-7, V3-11 and V4-4 (for example, one, more or all of V3-7\*01, V3-11\*01 and V4-4\*02). These segments were found in immunised repertoires to be productive in recombination with human JH6\*02 to produce HCDR3s of at least 20 amino acids in length.

In an example, the locus comprises one, more or all human IGHV gene segments V4-4, V1-8, V3-9, V3-11 and V3-20 (for example, one, more or all of V4-4\*02, V1-8\*01, V3-9\*01, V3-11\*01 and V3-20 (eg, \*d01)). These segments were found in antigen-specific repertoires to be productive in recombination with human JH6\*02 to produce HCDR3s of at least 20 amino acids in length.

14. The vertebrate or cell of any preceding clause, wherein the locus comprises one, more or all of human D3-9\*01, D3-10\*01, D6-19\*01, D6-13\*01, D1-26\*01, IGHV1-8\*01, IGHV4-61\*01, IGHV6-1\*01, IGHV4-4\*02, IGHV1-3\*01, IGHV3-66\*03, IGHV3-7\*01 and IGHV3-9\*01.

These are gene segments that very frequently combine with JH6\*02 to produce productive heavy chains and antibodies.

For example, the locus comprises one, more or all of human IGHV1-8\*01, D3-9\*01 and D3-10\*01. These gene segments were productive with JH6\*02 to produce HCDR3s of at least 20 amino acids in more than 10 antibodies.

15. An antibody-producing cell (eg, a B-cell) that is a progeny of the cell of any one of clauses 3 to 14, wherein the antibody-producing cell comprises a heavy chain locus comprising a rearranged variable region produced by recombination of human JH6\*02 with a D segment and a VH segment (eg, JH6\*02 with human VH3-11 (eg, VH3-11\*01) and D3-9; VH3-20 (eg, VH3-20\*01) and D3-10; VH4-4 (eg, VH4-4\*02) and D3-10; VH3-9 (eg, VH3-9\*01) and D3-10; or VH1-8 (eg, VH1-8\*01) and D310).

Such a variable region would be the product of *in vivo* somatic hypermutation in a non-human vertebrate or cell of the invention.

16. The cell of clause 15, which is a B-cell or hybridoma that expresses a target antigen-specific antibody comprising a heavy chain that comprises a rearranged variable region produced by recombination of human JH6\*02 with a D segment and a VH segment (eg, JH6\*02 with human VH3-11 (eg, VH3-11\*01) and D3-9; VH3-20 (eg, VH3-20\*01) and D3-10; VH4-4 (eg, VH4-4\*02) and D3-10; VH3-9 (eg, VH3-9\*01) and D3-10; or VH1-8 (eg, VH1-8\*01) and D310).

Such a variable region would be the product of *in vivo* somatic hypermutation in a non-human vertebrate or cell of the invention

17. The vertebrate or cell of any preceding clause, wherein the antibody heavy chain specifically binds a target antigen.
18. The vertebrate or cell of any preceding clause, wherein the antibody heavy chain has a HCDR3 length of at least 20 amino acids.

Optionally, the HCDR3 length is at least 21, 22, 23, 24, 25, 26, 27, 28, 29 or 30 amino acids. Additionally, in one example the length is no more than 35, 34, 33, 32 or 31 amino acids. For example, the HCDR3 length is 20, 21, 22, 23 or 24 amino acids.

19. The vertebrate or cell of any preceding clause, wherein the antibody heavy chain is a product of the recombination of JH6\*02 with a human VH gene segment recited in clause 13 or 14 and/or a D gene segment recited in clause 10, 11 or 14.
20. The vertebrate or cell of any preceding clause, wherein all endogenous non-human vertebrate heavy chain variable region gene segments have been inactivated in the genome (Eeg, by gene segment deletion or inversion).
21. The vertebrate or cell of any preceding clause, wherein the genome is homozygous for said heavy chain locus.
22. A heavy chain (eg, comprised by an antibody) isolated from a vertebrate of any one of clauses 1, 2, 5 to 14 and 17 to 21 wherein the heavy chain comprises a HCDR3 of at least 20 amino acids.
23. The heavy chain of clause 22, wherein the HCDR3 is the product of recombination of human JH6\*02 with a human VH gene segment recited in clause 13 or 14 and/or a D gene segment recited in clause 10, 11 or 14.

In an example, the heavy chain is chimaeric where the C region is non-human. In an example, the heavy chain is human where the C region is human.

24. A heavy chain (eg, comprised by an antibody) whose VH variable domain is identical to the VH variable domain of the heavy chain of clause 22 or 23, and which comprises a human constant region or a human-mouse chimaeric constant region (eg, CH1 is human and the other constant domains are mouse).
25. The heavy chain of clause 22, 23 or 24, whose VH variable domain is specific for a target antigen.
26. A method for producing a heavy chain, VH domain or an antibody specific to a target antigen, the method comprising immunizing a non-human vertebrate according to any one of clauses 1, 2, 5 to 14 and 17 to 21 with the antigen and isolating the heavy chain, VH domain or an antibody specific to a target antigen or a cell producing the heavy chain, VH domain or an antibody, wherein the heavy chain, VH domain or an antibody comprises a HCDR3 that is derived from the recombination of human JH6\*02 with a VH gene segment and a D gene segment.

27. A method for producing a human heavy chain or antibody comprising carrying out the method of clause 26, wherein the constant region of the locus is a non-human vertebrate (eg, mouse or rat) constant region, and then replacing the non-human constant region of the isolated heavy chain or antibody with a human constant region (eg, by engineering of the nucleic acid encoding the antibody).
28. A heavy chain, VH domain or an antibody produced by the method of clause 26 or 27.

Optionally the HCDR3 length is at least 20 amino acids as herein described.

29. A B-cell or hybridoma expressing a heavy chain VH domain that is identical to the VH domain of the heavy chain of clause 22, 23 or 28.
30. A nucleic acid encoding the VH domain of the heavy chain of clause 22, 23 or 28, or encoding the heavy chain of clause 22, 23, 24, 25 or 28.
31. A vector (eg, a CHO cell or HEK293 cell vector) comprising the nucleic acid of clause 30; optionally wherein the vector is in a host cell (eg, a CHO cell or HEK293 cell).
32. A pharmaceutical composition comprising the antibody, heavy chain or VH domain (eg, comprised by an antibody) of any one of clauses 22 to 25 and 28, together with a pharmaceutically-acceptable excipient, diluent or a medicament (eg, a further antigen-specific variable domain, heavy chain or antibody).
33. The antibody, heavy chain or VH domain (eg, comprised by an antibody) of any one of clauses 22 to 25 and 28 for use in medicine (eg, human medicine).

For example, the locus comprises the following human VH gene segments

IGHV6-1  
IGHV3-7  
IGHV1-8  
IGHV3-9  
IGHV3-11  
IGHV3-13  
IGHV1-18  
IGHV3-30  
IGHV4-31

IGHV4-39

IGHV4-59

Optionally also (i) and/or (ii)

(i)

IGHV1-2

IGHV2-5 and

IGHV3-21

(ii)

IGHV1-2

IGHV2-5

IGHV3-21

IGHV1-24

For example, the locus comprises the following human VH gene segment variants

IGHV6-1\*01

IGHV3-7\*01

IGHV1-8\*01

IGHV3-9\*01

IGHV3-11\*01

IGHV3-13\*01

IGHV1-18\*01

IGHV3-30\*18

IGHV4-31\*03

IGHV4-39\*01 and

IGHV4-59\*01;

Optionally also (iii) or (iv)

(ii)

IGHV1-2\*04

IGHV2-5\*10 and

IGHV3-21\*03

(iv)

IGHV1-2\*02

IGHV2-5\*01

IGHV3-21\*01 and

IGHV1-24\*01

For example, the locus comprises the following human JH gene segment variants

IGHJ2\*01

IGHJ3\*02

IGHJ4\*02  
IGHJ5\*02 and  
IGHJ6\*02

For example, the locus comprises the following human D gene segments

IGHD1-1  
IGHD2-2  
IGHD3-9  
IGHD3-10  
IGHD5-12  
IGHD6-13  
IGHD1-14  
IGHD2-15  
IGHD3-16  
IGHD4-17  
IGHD6-19  
IGHD2-21  
IGHD5-24  
IGHD1-26 and  
IGHD7-27

and optionally also (v) or (vi)

(v)  
IGHD3-3

(vi)  
IGHD3-3  
IGHD4-4  
IGHD5-5  
IGHD6-6  
IGHD1-7  
IGHD2-8 and  
IGHD2-8

The present invention provides in a fifth configuration:-

Constant Regions Tailored to Human Use & Antibody Humanisation



Additional rational design and bioinformatics has led the inventors to realise that specific human constant region variants are conserved across many diverse human populations. The inventors realised that this opens up the possibility of making a choice to humanise antibodies, chains and variable domains by using such specific constant regions in products, rather than arbitrarily choosing the human constant region (or a synthetic version of a human constant region). This aspect of the invention also enables one to tailor antibody-based drugs to specific human ethnic populations, thereby more closely matching drug to patient (and thus disease setting) than has hitherto been performed. It can be a problem in the state of the art that antibodies are humanised with an arbitrary choice of human constant region (presumably derived from one (often unknown) ethnic population or non-naturally occurring) that does not function as well in patients of a different human ethnic population. This is important, since the constant region has the major role in providing antibody effector functions, eg, for antibody recycling, cellular and complement recruitment and for cell killing.

As discussed further in WO2011066501, human IgG sub-types IgG1, IgG2, IgG3 and IgG4 exhibit differential capacity to recruit immune functions, such as antibody-dependent cellular cytotoxicity (ADCC, e.g., IgG1 and IgG3), antibody-dependent cellular phagocytosis (ADCP, e.g., IgG1, IgG2, IgG3 and IgG4), and complement dependent cytotoxicity (CDC, e.g., IgG1, IgG3). Sub-type-specific engagement of such immune functions is based on selectivity for Fc receptors on distinct immune cells and the ability to bind C1q and activate the assembly of a membrane attack complex (MAC). Among the various types, relative affinity for Fc $\gamma$  receptors (e.g., Fc $\gamma$ RI, Fc $\gamma$ RIIa/b/c, Fc $\gamma$ RIIIa/b) is high for IgG1 and IgG3, however, there is minimal affinity for IgG2 (restricted to the Fc $\gamma$ RIIIa 131H polymorphism), and IgG4 only has measurable affinity for Fc $\gamma$ RI. Using comparative sequence analysis and co-crystal structures, the key contact residues for receptor binding have been mapped to the amino acid residues spanning the lower hinge and CH2 region. Using standard protein engineering techniques, some success in enhancing or reducing the affinity of an antibody preparation for Fc receptors and the C1q component of complement has been achieved.

Among the isotypes, IgG2 is least capable of binding the family of Fc receptors. Using IgG2 as the starting point, efforts have been made to find a mutant with diminished effector functions but which retains FcRn binding, prolonged stability, and low immunogenicity. Improved mutants of this nature may provide improved antibody therapeutics with retained safety. Human IgG1 therapeutic

antibodies that bind to cell surface targets are able to engage effector cells that may mediate cell lysis of the target cell by antibody-dependent cellular cytotoxicity (ADCC) or complement dependent cytotoxicity (CDC). These mechanisms occur through interaction of the CH2 region of the antibody Fc domain to FcγR receptors on immune effector cells or with C1q, the first component of the complement cascade. Table 19 shows the activities of different human gamma sub-types. The skilled person may choose accordingly to promote or dampen-down activity depending upon the disease setting in humans of interest. For example, use of a human gamma-1 constant region is desirable when one wishes to isolated totally human heavy chains and antibodies that have relatively high complement activation activity by the classical pathway and FcγR1 recognition in human patients. See also Mol Immunol. 2003 Dec;40(9):585-93; "Differential binding to human FcγRIIIa and FcγRIIIb receptors by human IgG wild type and mutant antibodies"; Armour KL *et al*, which is incorporated herein by reference.

IgG2 constant regions are well suited to producing antibodies and heavy chains according to the invention for binding to cytokines or soluble targets in humans, since IgG2 is essentially FcγRI,III-silent, FcγRIIIa-active and has little Complement activity.

IgG1 constant regions have wide utility for human therapeutics, since IgG1 antibodies and heavy chains are FcγRI,II,III- active and have complement activity. This can be enhanced by using a human gamma-1 constant region that has been activated by engineering as is known in the art.

The work of the inventors has therefore identified a collection of human constant region of different isotypes from which an informed choice can be made when humanising chimaeric antibody chains (or conjugating V domains, such as dAbs or *Camelid* VHH, to constant regions). The collection was identified on the basis of bioinformatics analysis of the 1000 Genomes database, the inventors selecting constant region variants that are frequently occurring across several human ethnic populations, as well as those that appear with relatively high frequency within individual populations (as assessed by the number of individuals whose genomes comprise the variant). By sorting through the myriad possible sequences on this basis, the inventors have provided a collection of human constant region variants that are naturally-occurring and which can be used when rationally designing

antibodies, heavy chains and other antibody-based formats that bear a human constant region. In particular, this is useful when humanising chimaeric heavy chains to produce totally human chains in which both the variable and constant regions are human. This is useful for compatibility with human patients receiving antibody-based drugs.

To this end, the invention provides the following aspects:-

1. A method of producing an antibody heavy chain, the method comprising
  - (a) providing an antigen-specific heavy chain variable domain (eg, VH (such as a human VH or dAb) or VHH or a humanised heavy chain variable domain); and
  - (b) combining the variable domain with a human heavy chain constant region to produce an antibody heavy chain comprising (in N- to C-terminal direction) the variable domain and the constant region;

wherein

the human heavy chain constant region is an IGHAref, IGHA1a, IGHA2a, IGHA2b, IGHG1ref, IGHG2ref, IGHG2a, IGHG3ref, IGHG3a, IGHG3b, IGHG4ref, IGHG4a, IGHDref, IGHEref, IGHMref, IGHMa or IGHMb constant region.

Step (b) can be carried out, eg, using recombinant DNA technology using the corresponding nucleotide sequences.

For the constant region according to any aspect of this configuration, either genomic DNA or equivalent (ie, having introns and exons and optionally also 5' UTR sequences, eg, with native or a non-native leader sequence) can be used for the constant region. For example, any of the "GENOMIC" sequences disclosed as SEQ ID NO: 365 onwards herein. Alternatively, an intronless sequence can be used, for example any of the "CDS" sequences disclosed as SEQ ID NO: 365 onwards herein (eg, with native or a non-native leader sequence).

Optionally for any aspect of this configuration of the invention, the human heavy chain constant region is an IGHAref constant region.

Optionally for any aspect of this configuration of the invention, the human heavy chain constant region is an IGHA1a constant region.

Optionally for any aspect of this configuration of the invention, the human heavy chain constant region is an IGHA2a constant region.

Optionally for any aspect of this configuration of the invention, the human heavy chain constant region is an IGHA2b constant region.

Optionally for any aspect of this configuration of the invention, the human heavy chain constant region is IGHG1ref constant region.

Optionally for any aspect of this configuration of the invention, the human heavy chain constant region is an IGHG2ref constant region.

Optionally for any aspect of this configuration of the invention, the human heavy chain constant region is an IGHG2a constant region.

Optionally for any aspect of this configuration of the invention, the human heavy chain constant region is an IGHG3ref constant region.

Optionally for any aspect of this configuration of the invention, the human heavy chain constant region is an IGHG3a constant region.

Optionally for any aspect of this configuration of the invention, the human heavy chain constant region is an IGHG3b constant region.

Optionally for any aspect of this configuration of the invention, the human heavy chain constant region is an IGHG4ref constant region.

Optionally for any aspect of this configuration of the invention, the human heavy chain constant region is an IGHG4a constant region.

Optionally for any aspect of this configuration of the invention, the human heavy chain constant region is an IGHDref constant region.

Optionally for any aspect of this configuration of the invention, the human heavy chain constant region is an IGHEref constant region.

Optionally for any aspect of this configuration of the invention, the human heavy chain constant region is an IGHMref constant region.

Optionally for any aspect of this configuration of the invention, the human heavy chain constant region is an IGHMa constant region.

Optionally for any aspect of this configuration of the invention, the human heavy chain constant region is an IGHMb constant region.

Optionally, a derivative (eg, a mutant or conjugate) of the heavy chain or an antibody containing the heavy chain is produced. For example, a toxic payload can be conjugated (eg, for oncology applications). For example, one or more mutations can be introduced, as is known in the art, to inactivate or enhance Fc effector function.

2. The method of aspect 1, wherein the variable domain is a human variable domain.

A human variable domain is, for example, the product of recombination in a transgenic non-human vertebrate of human VH, D and JH gene segments. Alternatively, the variable domain is identified using *in vitro* display technology from a human VH library, eg, using phage display, ribosome display or yeast display, as is known in the art.

In another embodiment, the variable domain is a humanised variable domain, eg, comprising human frameworks with non-human (eg, mouse or rat) CDRs). Humanisation technology is conventional in the art, and will be readily known to the skilled person.

3. The method of any preceding aspect, wherein the variable domain has previously been selected from a non-human vertebrate that has been immunised with the antigen.

For example, the vertebrate (such as a mouse or rat) genome comprises a chimaeric heavy chain locus comprising a human variable region (human V, D and JH gene segments) operably connected upstream of a non-human vertebrate constant region so that the locus is able to rearrange for the expression of heavy chains comprising human variable domains and non-human vertebrate constant regions.

In alternative embodiments, the variable domain is selected using an *in vitro* technology such as phage display, ribosome display or yeast display. In this case the variable domain may be displayed with or without an constant region, provided that it is later combined with a human constant region as per the invention.

4. The method of any preceding aspect, comprising providing an expression vector (Eg, a mammalian expression vector, such as a CHO or HEK293 vector) comprising a nucleotide sequence encoding the constant region; inserting a nucleotide sequence encoding the variable domain into the vector 5' of the constant region sequence; inserting the vector into a host cell and expressing the heavy chain by the host cell; the method further comprising isolating a heavy chain (eg, as part of an antibody) comprising the variable domain and the human constant region.

The vector comprises regulatory elements sufficient to effect expression of the heavy chain when the vector is harboured by a host cell, eg, a CHO or HEK293 cell.

5. The method of any preceding aspect, further comprising obtaining a nucleotide sequence encoding the heavy chain.

6. An antibody comprising a human heavy chain, the heavy chain comprising a variable domain that is specific for an antigen and a constant region that is an IGHAref, IGHA1a, IGHA2a, IGHA2b, IGHG1ref, IGHG2ref, IGHG2a, IGHG3ref, IGHG3a, IGHG3b, IGHG4ref, IGHG4a, IGHDref, IGHEref, IGHMref, IGHMa or IGHMb constant region.
  
7. A polypeptide comprising (in N- to C- terminal direction) a leader sequence, a human variable domain that is specific for an antigen and a human constant region that is an IGHAref, IGHA1a, IGHA2a, IGHA2b, IGHG1ref, IGHG2ref, IGHG2a, IGHG3ref, IGHG3a, IGHG3b, IGHG4ref, IGHG4a, IGHDref, IGHEref, IGHMref, IGHMa or IGHMb constant region; wherein (i) the leader sequence is not the native human variable domain leader sequence (eg, the leader sequence is another human leader sequence or a non-human leader sequence); and/or (ii) the variable domain comprises mouse AID-pattern somatic mutations or mouse terminal deoxynucleotidyl transferase (TdT)- pattern junctional mutations.
  
8. A nucleotide sequence encoding (in 5' to 3' direction) a leader sequence and a human antibody heavy chain, the heavy chain comprising a variable domain that is specific for an antigen and a constant region that is an IGHAref, IGHA1a, IGHA2a, IGHA2b, IGHG1ref, IGHG2ref, IGHG2a, IGHG3ref, IGHG3a, IGHG3b, IGHG4ref, IGHG4a, IGHDref, IGHEref, IGHMref, IGHMa or IGHMb constant region; and the leader sequence being operable for expression (eg, in a mammalian CHO or HEK293 cell) of the heavy chain and wherein the leader sequence is not the native human variable domain leader sequence (eg, the leader sequence is another human leader sequence or a non-human leader sequence).

In an example, the leader sequence is

ATGGGCTGGTCCTGCATCATCCTGTTTCTGGTGGCCACCGCCACCGGCGTGACAGC

Which translates to

MGWSCILFLVATATGVHS

9. A nucleotide sequence encoding (in 5' to 3' direction) a promoter and a human antibody heavy chain, the heavy chain comprising a variable domain that is specific for an antigen and a constant region that is an IGHAref, IGHA1a, IGHA2a, IGHA2b, IGHG1ref, IGHG2ref, IGHG2a, IGHG3ref,

IGHG3a, IGHG3b, IGHG4ref, IGHG4a, IGHDref, IGHEref, IGHMref, IGHMa or IGHMb constant region; and the promoter being operable for expression (eg, in a mammalian CHO or HEK293 cell) of the heavy chain and wherein the promoter is not the native human promoter.

In one embodiment, the promoter sequence is a human IGK 3-15 promoter.

10. The antibody, polypeptide or nucleotide sequence of any one of aspects 6 to 9, wherein the variable domain comprises mouse AID-pattern somatic mutations and/or mouse terminal deoxynucleotidyl transferase (TdT)- pattern junctional mutations.

For example, one way, in any aspect of this configuration of the invention, to provide mouse AID-pattern somatic mutations and/or mouse terminal deoxynucleotidyl transferase (TdT)- pattern junctional mutations is to select a variable domain from a non-human vertebrate or cell. For example, a vertebrate or cell as disclosed herein.

11. A vector (eg, a CHO cell or HEK293 cell vector) comprising the nucleic acid of aspect 8, 9 or 10; optionally wherein the vector is in a host cell (eg, a CHO cell or HEK293 cell).
12. A pharmaceutical composition comprising the antibody or polypeptide of any one of aspects 6, 7 and 10, together with a pharmaceutically-acceptable excipient, diluent or a medicament (eg, a further antigen-specific variable domain, antibody chain or antibody).
13. The antibody or polypeptide of any one of aspects 6, 7 and 10 for use in treating and/or preventing a medical condition in a human patient.
14. Use of the antibody or polypeptide of any one of aspects 6, 7 and 10 for the manufacture of a medicament for treating and/or preventing a medical condition in a human patient.
15. The antibody, polypeptide or use of aspect 13 or 14, wherein the human is a member of a human population selected from population numbers 1-14, wherein the populations are numbered as follows (population labels being according to 1000 Genomes Project nomenclature)

1= ASW;



- 2= CEU;
- 3=CHB;
- 4=CHS;
- 5=CLM;
- 6=FIN;
- 7=GBR;
- 8=IBS;
- 9=JPT;
- 10=LWK;
- 11=MXL;
- 12=PUR;
- 13=TSI;
- 14=YRI.

16. The antibody, polypeptide or use of aspect 15, wherein the constant region is a

- (i) IGHA1a constant region and the human population is selected from any population number 1-14;
- (ii) IGHA2a constant region and the human population is selected from any population number 1-14;
- (iii) IGHA2b constant region and the human population is selected from any population number 1-14;
- (iv) IGHG2a constant region and the human population is selected from any population number 1-9 and 11-13;
- (v) IGHG3a constant region and the human population is selected from any population number 1-14;
- (vi) IGHG3b constant region and the human population is selected from any population number 1-8 and 11-13;
- (vii) IGHG4a constant region and the human population is selected from any population number 1-9 and 11-13;
- (viii) IGHMa constant region and the human population is selected from any population number 1-14; or
- (ix) IGHMb constant region and the human population is selected from any population number 1-14;

Wherein the populations are numbered as follows (population labels being according to 1000 Genomes Project nomenclature)

1= ASW;

2= CEU;

3=CHB;

4=CHS;

5=CLM;

6=FIN;

7=GBR;

8=IBS;

9=JPT;

10=LWK;

11=MXL;

12=PUR;

13=TSI;

14=YRI.

17. A vector (eg, a CHO cell or HEK293 cell vector) comprising a IGHG1ref, IGHG2ref, IGHG2a, IGHG3ref, IGHG3a, IGHG3b, IGHG4ref or IGHG4a constant region nucleotide sequence that is 3' of a cloning site for the insertion of a human antibody heavy chain variable domain nucleotide sequence, such that upon insertion of such a variable domain sequence the vector comprises (in 5' to 3' direction) a promoter, a leader sequence, the variable domain sequence and the constant region sequence so that the vector is capable of expressing a human antibody heavy chain when present in a host cell.

The present invention provides in a Sixth configuration:-

Multiple Variants in the Same Genome *Cis* or *Trans*

The inventors' analysis has revealed groupings of naturally-occurring human antibody gene segment variants as set out in Table 13 and Table 14. This revealed the possibility of producing transgenic genomes in non-human vertebrates and cells wherein the genomes contain more than the natural

human complement of specific human gene segments. In one example, this can be achieved by providing more than the natural human complement of a specific gene segment type on one or both of the respective Ig locus (eg, one or both chromosomes harbouring IgH in a mouse genome or mouse cell genome).

To this end, this configuration of the invention provides the following (as set out in numbered paragraphs):-

1. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising at least 3 human variable region gene segments of the same type (eg, at least 3 human VH6-1 gene segments, at least 3 human JH6 gene segments, at least 3 human Vk1-39 gene segments, at least 3 human D2-2 gene segments or at least 3 human Jk1 gene segments), wherein at least two of the human gene segments are variants that are not identical to each other.

For example, the genome comprises a variable region that comprises V, D and J gene segments (for the variable region of a heavy chain locus) or V and J gene segments (for the variable region of a light chain locus) upstream of a constant region for expression of heavy or light chains respectively.

In an alternative, the skilled person can choose to provide more than the wild type human complement of a specific gene segment type by providing several copies of one variant type of the human gene segment. Thus, there is provided

A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising at least 3 human variable region gene segments of the same type (eg, at least 3 human VH6-1 gene segments, at least 3 human JH6 gene segments, at least 3 human Vk1-39 gene segments, at least 3 human D2-2 gene segments or at least 3 human Jk1 gene segments), wherein the human gene segments are identical variants.

For example, the genome comprises a variable region that comprises V, D and J gene segments (for the variable region of a heavy chain locus) or V and J gene segments (for the variable region of a light chain locus) upstream of a constant region for expression of heavy or light chains respectively.

2. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising at least 2 different non-endogenous variable region gene segments of the same type (eg, at least 2 human VH6-1 gene segments, at least 3 human JH6 gene segments, at least 2 human V $\kappa$ 1-39 gene segments, at least 2 human D2-2 gene segments or at least 2 human J $\kappa$ 1 gene segments) *cis* at the same Ig locus.

In an alternative, the skilled person can choose to provide more than the wild type human complement of a specific gene segment type by providing several copies of one variant type of the human gene segment. Thus, there is provided

A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising at least 2 non-endogenous variable region gene segments of the same variant type (eg, at least 2 human JH6\*02 gene segments) *cis* at the same Ig locus.

3. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising at least 2 different human variable region gene segments of the same type (eg, at least 2 human VH6-1 gene segments, at least 2 human JH6 gene segments, at least 2 human V $\kappa$ 1-39 gene segments, at least 2 human D2-2 gene segments or at least 2 human J $\kappa$ 1 gene segments) *trans* at the same Ig locus; and optionally a third human gene segment of the same type, wherein the third gene segment is *cis* with one of said 2 different gene segments.

In an alternative, the skilled person can choose to provide more than the wild type human complement of a specific gene segment type by providing several copies of one variant type of the human gene segment. Thus, there is provided

A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising at least 2 different human variable region gene segments of the same variant type (eg, at least 2 human JH6\*02 gene segments) *trans* at the same Ig locus; and optionally a third human gene segment of the same variant type, wherein the third gene segment is *cis* with one of said 2 different gene segments.

4. A population of non-human vertebrates (eg, mice or rats) comprising a repertoire of human variable region gene segments, wherein the plurality comprises at least 2 human variable region gene segments of the same type (eg, at least 2 human VH6-1 gene segments, at least 2 human JH6 gene segments, at least 2 human V $\kappa$ 1-39 gene segments, at least 2 human D2-2 gene segments or at least 2 human J $\kappa$ 1 gene segments), a first of said different gene segments is provided in the genome of a first vertebrate of the population, and a second of said different gene segments being provided in the genome of a second vertebrate of the population, wherein the genome of the first vertebrate does not comprise the second gene segment.
5. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising at least 2 different non-endogenous variable region gene segments of the same type (eg, at least 2 human VH6-1 gene segments, at least 2 human JH6 gene segments, at least 2 human V $\kappa$ 1-39 gene segments, at least 2 human D2-2 gene segments or at least 2 human J $\kappa$ 1 gene segments), wherein the gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations.
6. A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 3 human variable region gene segments of the same type (eg, at least 3 human VH6-1 gene segments, at least 3 human JH6 gene segments, at least 3 human V $\kappa$ 1-39 gene segments, at least 3 human D2-2 gene segments or at least 3 human J $\kappa$ 1 gene segments), wherein at least two of the human gene segments are variants that are not identical to each other.
7. A method of enhancing the immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 2 different non-endogenous variable region gene segments of the same type (eg, at least 2 human VH6-1 gene segments, at least 2 human JH6 gene segments, at least 2 human V $\kappa$ 1-39 gene segments, at least 2 human D2-2 gene segments or at least 2 human J $\kappa$ 1 gene segments) *cis* at the same Ig locus.

8. A method of enhancing the immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 2 different human variable region gene segments of the same type (eg, at least 2 human VH6-1 gene segments, at least 2 human JH6 gene segments, at least 2 human Vk1-39 gene segments, at least 2 human D2-2 gene segments or at least 2 human Jk1 gene segments) *trans* at the same Ig locus; and optionally a third human gene segment of the same type, wherein the third gene segment is *cis* with one of said 2 different gene segments.
9. A method of providing an enhanced human immunoglobulin variable region gene segment repertoire, the method comprising providing a population of non-human vertebrates (eg, a mouse or rat) comprising a repertoire of human variable region gene segments, wherein the method comprises providing at least 2 different human variable region gene segments of the same type (eg, at least 2 human VH6-1 gene segments, at least 2 human JH6 gene segments, at least 2 human Vk1-39 gene segments, at least 2 human D2-2 gene segments or at least 2 human Jk1 gene segments), wherein a first of said different gene segments is provided in the genome of a first vertebrate of the population, and a second of said different gene segments is provided in the genome of a second vertebrate of the population, wherein the genome of the first vertebrate does not comprise the second gene segment.
10. A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 2 different non-endogenous variable region gene segments of the same type (eg, at least 2 human VH6-1 gene segments, at least 2 human JH6 gene segments, at least 2 human Vk1-39 gene segments, at least 2 human D2-2 gene segments or at least 2 human Jk1 gene segments), wherein the gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations.
11. The vertebrate, cell or method of any preceding paragraph, wherein at least 2 or 3 of said different gene segments are provided *cis* at the same Ig locus in said genome.
12. The vertebrate, cell or method of any preceding paragraph, wherein the gene segments are derived from the genome sequence of different human individuals that are not genetically

related over at least 3 generations.

13. The vertebrate, cell or method of any preceding paragraph, wherein the gene segments are derived from the genome sequence of two or more different human individuals; optionally wherein the different human individuals are from different human populations.
14. The vertebrate, cell or method of paragraph 13, wherein the individuals are not genetically related.
15. A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 2 human variable region gene segments of the same type (eg, at least 2 human VH6-1 gene segments, at least 2 human JH6 gene segments, at least 2 human Vk1-39 gene segments, at least 2 human D2-2 gene segments or at least 2 human Jk1 gene segments), wherein the gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations; optionally wherein at least 2 or 3 of said different gene segments are provided at the same Ig locus in said genome.
16. The method of paragraph 15, wherein the different human individuals are from different human populations.
17. The method of paragraph 15, wherein the individuals are not genetically related.
18. The vertebrate, cell or method of preceding paragraph, wherein at least one of the different segments is a synthetic mutant of a human germline gene segment.
19. The vertebrate, cell or method of any preceding paragraph, wherein each of said gene segments occurs in 10 or more different human populations.

20. The vertebrate, cell or method of preceding paragraph, wherein each of said gene segments has a human frequency of 5% or greater (eg, 10, 20, 25, 30, 35, 40, 45, 50, 55, 60, 65, 70, 75, 80, 85, 90, or 95% or greater).

In this respect, the skilled person can be guided by the information provided in Table 14.

Frequency can, for example, be cumulative frequency in the 1000 Genomes database.

21. The vertebrate, cell or method of paragraph 20, wherein each of said gene segments occurs in 10 or more different human populations.
22. The vertebrate, cell or method of any preceding paragraph, wherein each of said gene segments occurs in the 1000 Genomes database in more than 50 individuals.
23. The vertebrate, cell or method of preceding paragraph, wherein each of said gene segments (i) has a human frequency of 5% or greater (eg, 10, 20, 25, 30, 35, 40, 45, 50, 55, 60, 65, 70, 75, 80, 85, 90, or 95% or greater); and (ii) occurs in 10 or more different human populations.

In this respect, the skilled person can be guided by the information provided in Table 14.

Frequency can, for example, be cumulative frequency in the 1000 Genomes database.

24. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising first and second human Ig locus gene segments of the same type (eg, first and second human JH6 gene segments; or first and second IgG2 gene segments; or first and second human Jλ7 gene segments), wherein the first gene segment is a gene segment selected from Table 14 (eg, IGHJ6-a) and the second gene segment is the corresponding reference sequence (eg, IGHJ6 ref; SEQ ID NO: 244).

Table 14 lists commonly-occurring natural human variants. It can be seen that these occur across many human populations and thus usefully have wide applicability for human antibody-based drugs.



For example, the gene segments are provided as targeted insertions into an endogenous non-human vertebrate Ig locus. Alternatively, random integration (eg, using YACs) as is known in the art can be performed.

For example, the genome comprises a variable region that comprises V, D and J gene segments (for the variable region of a heavy chain locus) or V and J gene segments (for the variable region of a light chain locus) upstream of a constant region for expression of heavy or light chains respectively.

In another embodiment, the invention enables the skilled person to select two or more different naturally-occurring human gene segment variants for combination into the genome of a non-human vertebrate or cell. A reference sequence need not be included. It may be desirable to use one or more rare gene segments to increase diversity of the repertoire. Additionally or alternatively, it may be desirable to include a mixture of frequent and rare variants of the same type to provide repertoire diversity. The variants may be chosen additionally or alternatively to tailor the gene segment inclusion to one or more specific human populations as indicated by the information provided in Table 13 or Table 14.

Thus, the invention provides

A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising first and second human Ig locus gene segments of the same type (eg, first and second human JH6 gene segments; or first and second IgG2 gene segments; or first and second human Jλ7 gene segments), wherein the gene segments are gene segments selected from Table 13 or Table 14; and optionally wherein one or more of the gene segments appears in Table 14 (eg, IGHJ6-a) or is a reference sequence (eg, IGHJ6 ref; SEQ ID NO: 244).

25. The vertebrate or cell of paragraph 24, wherein the genome comprises a third human gene segment of said type, the third gene segment being different from the first and second gene segments.

26. The vertebrate or cell of paragraph 24 or 25, wherein the first and second gene segments are *cis* on the same chromosome; and optionally the third gene segment is also *cis* on said chromosome.

27. The vertebrate or cell of paragraph 26, wherein the gene segments are targeted insertions into an endogenous non-human Ig locus.

For example, the gene segments are heavy chain gene segments and the non-human locus is an IgH locus. For example, the gene segments are light chain ( $\kappa$  or  $\lambda$ ) gene segments and the non-human locus is an IgL locus.

28. The vertebrate or cell of paragraph 24 or 25, wherein the first and second gene segments are *trans* on different chromosomes.

Thus, the chromosomes are the same type (eg, both mouse chromosome 6 or rat chromosome 4).

29. The vertebrate or cell of any one of paragraphs 24 to 28, wherein the first gene segment is a gene segment selected from any one of Tables 1 to 7 and 9 to 14 (eg, selected from Table 13 or 14) and the second gene segment is the corresponding reference sequence.

30. A population of non-human vertebrates (eg, mice or rats) comprising first and second human Ig locus gene segments of the same type (eg, first and second human JH6 gene segments; or first and second IgG2 gene segments; or first and second human  $\lambda$ 7 gene segments), wherein the first gene segment is a gene segment selected from any one of Tables 1 to 7 and 9 to 14 (eg, Table 13 or 14) (eg, IGHJ6-a) and the second gene segment is the corresponding reference sequence (eg, SEQ ID NO: 7), wherein the first gene segment is provided in the genome of a first vertebrate of the population, and the second gene segment is provided in the genome of a second vertebrate of the population.

31. The population of paragraph 30, wherein the genome of the first vertebrate does not comprise the second gene segment.
32. The population of paragraph 30 or 31, wherein the population comprises a third human gene segment of said type, the third gene segment being different from the first and second gene segments and optionally wherein the first and third gene segments are present in the genome of the first vertebrate.
33. The population of paragraph 30, 31 or 32, wherein the gene segments are targeted insertions into an endogenous non-human Ig locus in the respective genome.

For example, the gene segments are heavy chain gene segments and the non-human locus is an IgH locus. For example, the gene segments are light chain (kappa or lambda) gene segments and the non-human locus is an IgL locus.

34. The population of of any one of paragraphs 30 to 33, wherein the first gene segment is a gene segment selected from any one of Tables 1 to 7 and 9 to 14 (eg, Table 13 or 14) and the second gene segment is the corresponding reference sequence.
35. A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising first and second human Ig locus gene segments of the same type (eg, first and second human JH6 gene segments; or first and second IgG2 gene segments; or first and second human J $\lambda$ 7 gene segments), wherein the first gene segment is a gene segment selected from any one of Tables 1 to 7 and 9 to 14 (eg, Table 13 or 14) (eg, IGHJ6-a) and the second gene segment is the corresponding reference sequence (eg, SEQ ID NO: 7).
36. A method of providing an enhanced human immunoglobulin gene segment repertoire, the method comprising providing a population according to any one of paragraphs 30 to 33.

#### Variants Prevalent in Few Populations

In another aspect, it is of note that certain human gene segment variants may appear relatively frequently in one or a small number of populations, but is not found prevalently across many different human populations. There is thinking that specific germline gene segment repertoires have evolved in individual human ethnic populations due to iterative exposure to antigens (eg, disease pathogen antigens) to which the population is often exposed. Repeated exposure and mutation may have lead to the evolution of gene segment variants that can provide an effective response to the antigen (pathogen) in the population, and this may explain the conservation of the gene segments in those populations (as opposed to other human ethnic populations that may not have frequently encountered the antigen). With this in mind, the inventors identified gene segment variants from their analysis that are relatively prevalent in a small number of human populations, and not across many populations. The inventors realized that inclusion of one or more of such gene segments in the configurations of the invention (eg, in transgenic Ig loci, vertebrates and cells) would be useful for producing antibodies, Ig chains and variable domains that can address antigens (eg, disease-causing antigens or pathogens) to which the small number of human populations may become exposed. Such products would be useful for treating and/or preventing disease or medical conditions in members of such a population. This aspect could also be useful for addressing infectious disease pathogens that may have been common in the small number of populations, but which in the future or relatively recently in evolution has become a more prevalent disease-causing pathogen in other human populations (ie, those not listed in Table 13 against the gene segment variant(s) in question). To this end, from the 1000 Genomes database the inventors have identified the gene segment variants listed in Table 20.

Thus, according to any configuration or aspect described herein, one, more or all of the gene segments used in the present invention can be a gene segment listed in Table 20A, 20B, 20C or 20D.

#### Multiple JH Gene Segment Variants

A specific application of this configuration is the provision of multiple human JH gene segments as follows.

A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising at least 3 human JH gene segments of the same type (JH1, JH2, JH3, JH4, JH5 or JH6), wherein at least two of the human JH gene segments are variants that are not

identical to each other.

In an example, any cell of the invention is an isolated cell. An "isolated" cell is one that has been identified, separated and/or recovered from a component of its production environment (eg, naturally or recombinantly). Preferably, the isolated cell is free of association with all other components from its production environment, eg, so that the cell can produce an antibody to an FDA-approvable or approved standard. Contaminant components of its production environment, such as that resulting from recombinant transfected cells, are materials that would typically interfere with research, diagnostic or therapeutic uses for the resultant antibody, and may include enzymes, hormones, and other proteinaceous or non-proteinaceous solutes. In preferred embodiments, the polypeptide will be purified: (1) to greater than 95% by weight of antibody as determined by, for example, the Lowry method, and in some embodiments, to greater than 99% by weight; (2) to a degree sufficient to obtain at least 15 residues of N-terminal or internal amino acid sequence by use of a spinning cup sequenator, or (3) to homogeneity by SDS-PAGE under non-reducing or reducing conditions using Coomassie blue or, preferably, silver stain. Ordinarily, however, an isolated cell will be prepared by at least one purification step.

A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising at least 2 different non-endogenous JH gene segments (eg, human gene segments) of the same type (JH1, JH2, JH3, JH4, JH5 or JH6) *cis* at the same Ig (eg, IgH, eg, endogenous IgH, eg, mouse or rat IgH) locus. In an example, the genome comprises a human VH, D and JH repertoire comprising said different JH gene segments. Optionally the non-endogenous JH gene segments are non-mouse or non-rat, eg, human JH gene segments. In an example one or more or all of the non-endogenous gene segments are synthetic.

A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising at least 2 different human JH gene segments of the same type (JH1, JH2, JH3, JH4, JH5 or JH6) *trans* at the same Ig (eg, IgH, eg, endogenous IgH, eg, mouse or rat IgH) locus; and optionally a third human JH gene segments of the same type, wherein the third JH is *cis* with one of said 2 different JH gene segments.

A population of non-human vertebrates (eg, mice or rats) comprising a repertoire of human JH gene segments, wherein the plurality comprises at least 2 different human JH gene segments of the same type (JH1, JH2, JH3, JH4, JH5 or JH6), a first of said different JH gene segments is provided in the genome of a first vertebrate of the population, and a second of said different JH gene segments being provided in the genome of a second vertebrate of the population, wherein the genome of the first vertebrate does not comprise the second JH gene segment.

A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising at least 2 different non-endogenous (eg, human) JH gene segments of the same type (JH1, JH2, JH3, JH4, JH5 or JH6), wherein the JH gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations (eg, 3, 4, 5 or 6 generations). Optionally the non-endogenous JH gene segments are human JH gene segments. In an example one or more or all of the non-endogenous gene segments are synthetic.

A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 3 human JH gene segments of the same type (JH1, JH2, JH3, JH4, JH5 or JH6), wherein at least two of the human JH gene segments are variants that are not identical to each other.

A method of enhancing the immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 2 different non-endogenous (eg, human) JH gene segments of the same type (JH1, JH2, JH3, JH4, JH5 or JH6) *cis* at the same Ig (eg, IgH, eg, endogenous IgH, eg, mouse or rat IgH) locus). Optionally the non-endogenous JH gene segments are non-mouse or non-rat, eg, human JH gene segments. In an example one or more or all of the non-endogenous gene segments are synthetic.

A method of enhancing the immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 2 different human JH gene segments of the same type (JH1, JH2, JH3, JH4, JH5 or JH6) *trans* at the same Ig (eg, IgH, eg, endogenous IgH, eg, mouse or rat IgH) locus; and optionally a third human JH

gene segments of the same type, wherein the third JH is *cis* with one of said 2 different JH gene segments.

A method of providing an enhanced human immunoglobulin JH gene segment repertoire, the method comprising providing a population of non-human vertebrates (eg, a mouse or rat) comprising a repertoire of human JH gene segments, wherein the method comprises providing at least 2 different human JH gene segments of the same type (JH1, JH2, JH3, JH4, JH5 or JH6), wherein a first of said different JH gene segments is provided in the genome of a first vertebrate of the population, and a second of said different JH gene segments is provided in the genome of a second vertebrate of the population, wherein the genome of the first vertebrate does not comprise the second JH gene segment.

A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 2 different non-endogenous (eg, human) JH gene segments of the same type (JH1, JH2, JH3, JH4, JH5 or JH6), wherein the JH gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations (eg, 3, 4, 5, or 6 generations). Optionally the non-endogenous JH gene segments are human JH gene segments. In an example one or more or all of the non-endogenous gene segments are synthetic.

In an example of the vertebrate or cell or the method of the invention at least 2 or 3 of said different gene segments are provided *cis* at the same Ig locus in said genome.

In an example of the vertebrate or cell or the method of the invention the JH gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations (eg, 3, 4, 5, or 6 generations).

In an example of the vertebrate or cell or the method of the invention the JH gene segments are derived from the genome sequence of two or more different human individuals; optionally wherein the different human individuals are from different human populations.

In an example of the vertebrate or cell or the method of the invention the individuals are not genetically related (eg, going back 3, 4, 5, or 6 generations).

In an example of the vertebrate or cell or the method of the invention at least one of the different JH segments is a synthetic mutant of a human germline JH gene segment.

The invention also provides a method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 2 human JH gene segments of the same type (JH1, JH2, JH3, JH4, JH5 or JH6), wherein the JH gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations (eg, 3, 4, 5, or 6 generations); optionally wherein at least 2 or 3 of said different gene segments are provided at the same IgH locus in said genome.

In an example of the vertebrate or cell or the method of this embodiment of the invention the genome comprises a substantially complete functional repertoire of human JH gene segment types supplemented with one, two or more human JH gene segments, wherein said substantially complete functional repertoire and the supplementary JH gene segments are not found together in the germline genome of a human individual.

In an example of the population of the invention, the population comprises a substantially complete functional repertoire of human JH gene segment types supplemented with one, two or more human JH gene segments, wherein said substantially complete functional repertoire and the supplementary JH gene segments are not found together in the germline genome of a human individual.

A non-human vertebrate (eg, a mouse or rat) or a non-human cell (eg, an ES cell or a B-cell) having a genome comprising a substantially complete functional repertoire of human JH gene segment types supplemented with one, two or more human JH gene segments, wherein said substantially complete functional repertoire and the supplementary JH gene segments are not found together in the germline genome of a human individual.



A population of non-human vertebrates (eg, mice or rats) comprising a substantially complete functional repertoire of human JH gene segment types supplemented with one, two or more human JH gene segments, wherein said substantially complete functional repertoire and the supplementary JH gene segments are not found together in the germline genome of a human individual.

In an example of the vertebrate or the population, at least one of said JH gene segments is SEQ ID NO: 1, 2, 3 or 4. For example, at least one of said JH gene segments is SEQ ID NO: 1 and at least one, two or more of said supplementary JH gene segments is a variant according to any example above. For example, at least one of said JH gene segments is SEQ ID NO: 2 and at least one, two or more of said supplementary JH gene segments is a variant according to any one of the examples above. For example, at least one of said JH gene segments is SEQ ID NO: 2 and at least one, two or more of said supplementary JH gene segments is a variant according to any one of the examples above.

In an embodiment, the non-human vertebrate or vertebrate cell of the invention comprises a genome that comprises VH, D and JH gene repertoires comprising human gene segments, the JH gene repertoire (eg, a human JH gene segment repertoire) comprising

a plurality of JH1 gene segments provided by at least 2 different JH1 gene segments in *cis* at the same Ig locus in said genome;

a plurality of JH2 gene segments provided by at least 2 different JH2 gene segments in *cis* at the same Ig locus in said genome;

a plurality of JH3 gene segments provided by at least 2 different JH3 gene segments in *cis* at the same Ig locus in said genome;

a plurality of JH4 gene segments provided by at least 2 different JH4 gene segments in *cis* at the same Ig locus in said genome;

a plurality of JH5 gene segments provided by at least 2 different JH5 gene segments in *cis* at the same Ig locus in said genome; and/or

a plurality of JH6 gene segments provided by at least 2 different JH6 gene segments in *cis* at the same Ig locus in said genome;

optionally wherein the JH gene segments are derived from the genome sequence of two or more different human individuals.

Optionally said at least 2 different JH gene segments are human gene segments or synthetic gene segments derived from human gene segments.

Optionally, the Ig locus is a IgH locus, eg, an endogenous locus, eg, a mouse or rat IgH locus.

In an embodiment, the non-human vertebrate or vertebrate cell of the invention comprises a genome that comprises VH, D and JH gene repertoires comprising human gene segments, the JH gene repertoire (eg, a human JH gene segment repertoire) comprising

- a plurality of JH1 gene segments provided by at least 3 different JH1 gene segments;
- a plurality of JH2 gene segments provided by at least 3 different JH2 gene segments;
- a plurality of JH3 gene segments provided by at least 3 different JH3 gene segments;
- a plurality of JH4 gene segments provided by at least 3 different JH4 gene segments;
- a plurality of JH5 gene segments provided by at least 3 different JH5 gene segments; and/or
- a plurality of JH6 gene segments provided by at least 3 different JH6 gene segments;

optionally wherein the JH gene segments are derived from the genome sequence of two or three different human individuals;

optionally wherein at least 2 or 3 of said different gene segments are provided in *cis* at the same Ig locus in said genome.

Optionally said at least 3 different JH gene segments are human gene segments or synthetic gene segments derived from human gene segments.

Optionally, the Ig locus is a IgH locus, eg, an endogenous locus, eg, a mouse or rat IgH locus.

Optionally in the vertebrate or cell the different human individuals are from different human populations.

Optionally in the vertebrate or cell the individuals are not genetically related (eg, Going back 3, 4, 5 or 6 generations).

Optionally in the vertebrate or cell at least one of the different JH segments is a synthetic mutant of a human germline JH gene segment.

In an embodiment of a non-human vertebrate or vertebrate cell (optionally an ES cell or B-cell) according to the invention, the vertebrate or cell genome comprises human VH, D and JH gene repertoires, the JH gene repertoire (eg, a human JH gene repertoire) comprising a plurality of JH1 gene segments provided by at least 2 different human JH1 gene segments, optionally in *cis* at the same Ig locus in said genome;  
a plurality of JH2 gene segments provided by at least 2 different human JH2 gene segments, optionally in *cis* at the same Ig locus in said genome;  
a plurality of JH3 gene segments provided by at least 2 different human JH3 gene segments, optionally in *cis* at the same Ig locus in said genome;  
a plurality of JH4 gene segments provided by at least 2 different human JH4 gene segments, optionally in *cis* at the same Ig locus in said genome;  
a plurality of JH5 gene segments provided by at least 2 different human JH5 gene segments, optionally in *cis* at the same Ig locus in said genome; and/or  
a plurality of JH6 gene segments provided by at least 2 different human JH6 gene segments, optionally in *cis* at the same Ig locus in said genome;  
wherein the JH gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations (eg, 3, 4, 5 or 6 generations).

Optionally said at least 2 different JH gene segments are human gene segments or synthetic gene segments derived from human gene segments.

Optionally, the Ig locus is a IgH locus, eg, an endogenous locus, eg, a mouse or rat IgH locus.

Optionally in the vertebrate or cell the human individuals are from different human populations.

### JH5

An embodiment provides a vertebrate, cell or population of the invention whose genome comprises a plurality of JH5 gene segments, wherein the plurality comprises a human JH5 gene variant of SEQ ID NO: 1, wherein the variant comprises a nucleotide mutation at one or more positions

corresponding to positions

106,330,024

106,330,027

106,330,032

106,330,041

106,330,044

106,330,045

106,330,062

106,330,063

106,330,065

106,330,066

106,330,067

106,330,068 and

106,330,071

on human chromosome 14.

In the vertebrate, cell or population optionally the plurality comprises a human JH5 gene variant of SEQ ID NO: 1, wherein the variant comprises a guanine at a position corresponding to position 106,330,067 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 1.

Optionally the variant comprises additionally a mutation at a position corresponding to (i) position 106,330,071 on human chromosome 14 (optionally the additional mutation being a guanine); (ii) position 106,330,066 on human chromosome 14 (optionally the additional mutation being a guanine); and/or (iii) position 106,330,068 on human chromosome 14 (optionally the additional mutation being a thymine).

Optionally the plurality comprises a human JH5 gene variant of SEQ ID NO: 1, wherein the variant comprises a guanine at a position corresponding to position 106,330,071 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 1.

Optionally the variant comprises additionally a mutation at a position corresponding to (i) position 106,330,063 on human chromosome 14 (optionally the additional mutation being an adenine); and/or (ii) position 106,330,067 on human chromosome 14 (optionally the additional mutation being a guanine).

Optionally the plurality comprises a human JH5 gene variant of SEQ ID NO: 1, wherein the variant comprises a cytosine at a position corresponding to position 106,330,045 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 1.

Optionally the plurality comprises a human JH5 gene variant of SEQ ID NO: 1, wherein the variant comprises an adenine at a position corresponding to position 106,330,044 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 1.

Optionally the variant comprises additionally a mutation at a position corresponding to (i) position 106,330,066 on human chromosome 14 (optionally the additional mutation being a guanine); and/or (ii) position 106,330,068 on human chromosome 14 (optionally the additional mutation being a thymine).

Optionally the plurality comprises a human JH5 gene variant of SEQ ID NO: 1, wherein the variant comprises a guanine at a position corresponding to position 106,330,066 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 1.

Optionally the variant comprises additionally a mutation at a position corresponding to (i) position 106,330,067 on human chromosome 14 (optionally the additional mutation being a guanine); and/or (ii) position 106,330,068 on human chromosome 14 (optionally the additional mutation being a thymine).

Optionally the plurality comprises a human JH5 gene variant of SEQ ID NO: 1, wherein the variant comprises a thymine at a position corresponding to position 106,330,068 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 1.

Optionally the variant comprises additionally a mutation at a position corresponding to (i) position 106,330,067 on human chromosome 14 (optionally the additional mutation being a guanine); and/or (ii) position 106,330,066 on human chromosome 14 (optionally the additional mutation being a guanine).

Optionally the plurality comprises a human JH5 gene variant of SEQ ID NO: 1, wherein the variant comprises a cytosine at a position corresponding to position 106,330,027 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 1.

Optionally the plurality comprises a human JH5 gene variant of SEQ ID NO: 1, wherein the variant comprises an adenine at a position corresponding to position 106,330,024 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 1.

Optionally the plurality comprises a human JH5 gene variant of SEQ ID NO: 1, wherein the variant comprises a thymine at a position corresponding to position 106,330,032 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 1.

Optionally the plurality comprises a human JH5 gene variant of SEQ ID NO: 1, wherein the variant comprises a thymine at a position corresponding to position 106,330,041 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 1.

Optionally the plurality comprises a human JH5 gene variant of SEQ ID NO: 1, wherein the variant comprises an adenine or thymine at a position corresponding to position 106,330,063 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 1.

Optionally the variant comprises additionally a mutation at a position corresponding to position 106,330,071 on human chromosome 14 (optionally the additional mutation being a guanine).

Optionally the plurality comprises a human JH5 gene variant of SEQ ID NO: 1, wherein the variant comprises a cytosine at a position corresponding to position 106,330,062 on human chromosome

14; and optionally no further mutation from the sequence of SEQ ID NO: 1.

Optionally the genome comprises SEQ ID NO:1; optionally in *cis* at the same Ig locus as one, two or more of the variants.

### JH6

An embodiment provides a vertebrate, cell or population of the invention whose genome comprises a plurality of JH6 gene segments, wherein the plurality comprises a human JH6 gene variant of SEQ ID NO: 2, wherein the variant comprises a nucleotide mutation at one or more positions corresponding to positions

106,329,411

106,329,413

106,329,414

106,329,417

106,329,419

106,329,426

106,329,434

106,329,435, and

106,329,468

on human chromosome 14.

Optionally the genome of the vertebrate, cell or population comprises a plurality of JH6 gene segments, wherein the plurality comprises a human JH6 gene variant of SEQ ID NO: 2, wherein the variant comprises a guanine at a position corresponding to position 106,329,435 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 2.

Optionally the variant comprises additionally a mutation at a position corresponding to (i) position 106,329,468 on human chromosome 14 (optionally the additional mutation being a guanine); (ii) position 106,329,419 on human chromosome 14 (optionally the additional mutation being an adenine); (iii) position 106,329,434 on human chromosome 14 (optionally the additional mutation

being a cytosine) and/or position 106,329,414 on human chromosome 14 (optionally the additional mutation being a guanine); (iv) position 106,329,426 on human chromosome 14 (optionally the additional mutation being an adenine); (v) position 106,329,413 on human chromosome 14 (optionally the additional mutation being an adenine); (vi) position 106,329,417 on human chromosome 14 (optionally the additional mutation being a thymine); (vii) position 106,329,411 on human chromosome 14 (optionally the additional mutation being a thymine); (viii) position 106,329,451 on human chromosome 14 (optionally the additional mutation being an adenine); (ix) position 106,329,452 on human chromosome 14 (optionally the additional mutation being a cytosine); and/or (x) position 106,329,453 on human chromosome 14 (optionally the additional mutation being a cytosine).

Optionally the variant comprises additionally mutations at positions corresponding to position 106,329,451 on human chromosome 14, the additional mutation being an adenine; position 106,329,452 on human chromosome 14, the additional mutation being a cytosine; and position 106,329,453 on human chromosome 14, the additional mutation being a cytosine.

The vertebrate, cell or population optionally comprises a plurality of JH6 gene segments,, wherein the plurality comprises a human JH6 gene variant of SEQ ID NO: 2, wherein the variant comprises a guanine at a position corresponding to position 106,329,468 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 2.

Optionally the variant comprises additionally a mutation at a position corresponding to position 106,329,435 on human chromosome 14 (optionally the additional mutation being a guanine).

Optionally the vertebrate, cell or population comprises a plurality of JH6 gene segments,, wherein the plurality comprises a human JH6 gene variant of SEQ ID NO: 2, wherein the variant comprises a thymine at a position corresponding to position 106,329,417 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 2.

Optionally the variant comprises additionally a mutation at a position corresponding to position 106,329,435 on human chromosome 14 (optionally the additional mutation being a guanine).

Optionally the vertebrate, cell or population comprises a plurality of JH6 gene segments,, wherein the plurality comprises a human JH6 gene variant of SEQ ID NO: 2, wherein the variant comprises a



cytosine at a position corresponding to position 106,329,434 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 2.

Optionally the variant comprises additionally a mutation at a position corresponding to (i) position 106,329,414 on human chromosome 14 (optionally the additional mutation being a guanine); and/or (ii) position 106,329,435 on human chromosome 14 (optionally the additional mutation being a guanine).

Optionally the vertebrate, cell or population comprises a plurality of JH6 gene segments, wherein the plurality comprises a human JH6 gene variant of SEQ ID NO: 2, wherein the variant comprises a thymine at a position corresponding to position 106,329,411 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 2.

Optionally the variant comprises additionally a mutation at a position corresponding to position 106,329,435 on human chromosome 14 (optionally the additional mutation being a guanine).

Optionally the vertebrate, cell or population comprises a plurality of JH6 gene segments, wherein the plurality comprises a human JH6 gene variant that is an antisense sequence of a variant described above.

Optionally the genome comprises SEQ ID NO:2; optionally *cis* at the same Ig locus as one, two or more of the JH6 variants.

## JH2

An embodiment provides a vertebrate, cell or population of the invention whose genome comprises a plurality of JH2 gene segments, wherein the plurality comprises a human JH2 gene variant of SEQ ID NO: 3, wherein the variant comprises a nucleotide mutation at one or more positions corresponding to positions

106,331,455

106,331,453, and

106,331,409

on human chromosome 14.

Optionally the vertebrate, cell or population comprises said plurality of JH2 gene segments, wherein the plurality comprises a human JH2 gene variant of SEQ ID NO: 3, wherein the variant comprises a guanine at a position corresponding to position 106,331,455 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 3.

Optionally the variant comprises additionally a mutation at a position corresponding to (i) position 106,331,453 on human chromosome 14 (optionally the additional mutation being an adenine); and/or (ii) position 106,331,409 on human chromosome 14 (optionally the additional mutation being an adenine); (iii) position 106,329,434 on human chromosome 14 (optionally the additional mutation being an adenine).

Optionally the vertebrate, cell or population comprises a plurality of JH2 gene segments, wherein the plurality comprises a human JH2 gene variant of SEQ ID NO: 3, wherein the variant comprises an adenine at a position corresponding to position 106,331,453 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 3.

Optionally the variant comprises additionally a mutation at a position corresponding to position 106,331,409 on human chromosome 14 (optionally the additional mutation being an adenine).

Optionally the vertebrate, cell or population comprises a plurality of JH2 gene segments, wherein the plurality comprises a human JH2 gene variant of SEQ ID NO: 3, wherein the variant comprises an adenine at a position corresponding to position 106,331,409 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 3.

Optionally the vertebrate, cell or population comprises a plurality of JH2 gene segments, wherein the plurality comprises a human JH2 gene variant that is an antisense sequence of a variant described above.

Optionally the genome comprises SEQ ID NO:3; optionally *cis* at the same Ig locus as one, two or more of the JH2 variants.

Optionally the vertebrate, cell or population genome comprises two or more different JH gene segments selected from SEQ ID NOs: 1 to 3 and variants described above; optionally wherein said JH gene segments are *cis* at the same immunoglobulin Ig locus.

#### Multiple Human D Gene Segment Variants

A specific application of this configuration is the provision of multiple human D gene segments as follows (as set out in numbered clauses, starting at clause number 154).

154. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising at least 3 human D gene segments of the same type (eg, D2-2 gene segments), wherein at least two of the human D gene segments are variants that are not identical to each other (eg, D2-2ref and D2-2a).

In an example of any aspect of the sixth configuration of the invention (V, D, J or C), one or more or all of the variants are naturally-occurring human gene segments.

In an example of any aspect of the sixth configuration of the invention (V, D, J or C), one or more of the variants may be a synthetic variant of a human gene segment.

155. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising at least 2 different non-endogenous D gene segments of the same type type (eg, D2-2ref and D2-2a) *cis* at the same Ig locus.

156. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising at least 2 different human D gene segments of the same type (eg, D2-2ref and D2-2a) *trans* at the same Ig locus; and optionally a third human D gene segment (eg, (eg, D2-2ref, D2-2a or D2-2b) of the same type, wherein the third D is *cis* with one of said 2 different D gene segments.

157. A population of non-human vertebrates (eg, mice or rats) comprising a repertoire of human D gene segments, wherein the plurality comprises at least 2 different human D gene segments of the same type (eg, D2-2 gene segments), a first of said different D gene segments (eg, D2-2ref) is provided in the genome of a first vertebrate of the population, and a second of said different D gene segment (eg, D2-2a) being provided in the genome of a second vertebrate of the

population, wherein the genome of the first vertebrate does not comprise the second D gene segment.

158. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising at least 2 different non-endogenous D gene segments of the same type (eg, human D2-2 gene segments), wherein the D gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations.
159. A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 3 human D gene segments of the same type (eg, D2-2 gene segments), wherein at least two of the human D gene segments are variants that are not identical to each other (eg, D2-2ref and D2-2a).
160. A method of enhancing the immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 2 different non-endogenous D gene segments of the same type (eg, human D2-2 gene segments) *cis* at the same Ig locus.
161. A method of enhancing the immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 2 different human D gene segments of the same type (eg, D2-2ref and D2-2a) *trans* at the same Ig locus; and optionally a third human D gene segment (eg, D2-2ref, D2-2a or D2-2b) of the same type, wherein the third D is *cis* with one of said 2 different D gene segments.
162. A method of providing an enhanced human immunoglobulin D gene segment repertoire, the method comprising providing a population of non-human vertebrates (eg, a mouse or rat) comprising a repertoire of human D gene segments, wherein the method comprises providing at least 2 different human D gene segments of the same type (eg, D2-2ref and D2-2a), wherein a first of said different D gene segments is provided in the genome of a first vertebrate of the population, and a second of said different D gene segments is provided in the genome of a second vertebrate of the population, wherein the genome of the first vertebrate does not

comprise the second D gene segment.

163. A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 2 different non-endogenous D gene segments of the same type (eg, D2-2ref and D2-2a), wherein the D gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations.
164. The vertebrate or cell of clause 154, 156 or 158, or the method of clause 159, 161 or 163, wherein at least 2 or 3 of said different gene segments are provided *cis* at the same Ig locus in said genome.
165. The vertebrate or cell of clause 154, 155 or 156, or the method of any one of clauses 159 to 162 and 164, wherein the D gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations.
166. The vertebrate or cell of any one of clauses 154 to 157, or the method of any one of clauses 159 to 162 and 165, wherein the D gene segments are derived from the genome sequence of two or more different human individuals; optionally wherein the different human individuals are from different human populations.
167. The vertebrate, cell or method of clause 166, wherein the individuals are not genetically related.
168. The vertebrate, cell or method of any one of clauses 154 to 167, wherein at least one of the different D segments is a synthetic mutant of a human germline D gene segment.
169. A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 2 human D gene segments of the same type (eg, D2-2ref and D2-2a), wherein the D gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations; optionally wherein at least 2 or 3 of said different gene segments are provided at the same IgH locus in said genome.

170. The vertebrate or cell of any one of clauses 154 to 158 and 164 to 168, wherein the genome comprises a substantially complete functional repertoire of human D gene segment types supplemented with one, two or more variant human D gene segments, wherein said substantially complete functional repertoire and the supplementary D gene segments are not found together in the germline genome of a human individual.
171. The population of clause 157, wherein the population comprises a substantially complete functional repertoire of human D gene segment types supplemented with one, two or more variant human D gene segments, wherein said substantially complete functional repertoire and the supplementary D gene segments are not found together in the germline genome of a human individual.
172. A non-human vertebrate (eg, a mouse or rat) or a non-human cell (eg, an ES cell or a B-cell) having a genome comprising a substantially complete functional repertoire of human D gene segment types supplemented with one, two or more variant human D gene segments, wherein said substantially complete functional repertoire and the supplementary D gene segments are not found together in the germline genome of a human individual.
173. A population of non-human vertebrates (eg, mice or rats) comprising a substantially complete functional repertoire of human JH gene segment types supplemented with one, two or more variant human D gene segments, wherein said substantially complete functional repertoire and the supplementary D gene segments are not found together in the germline genome of a human individual.
174. The vertebrate or cell of clause 172 or the population of clause 173, comprising first and second D gene segments selected from  
D2-2ref and D2-2a; or  
D2-21ref and D2-21a; or  
D3-10ref and D3-10a; or  
D3-16ref and D3-16a; or  
D2-8ref and D2-8a; or  
D3-3ref and D3-3a; or  
D4-23ref and D4-23a; or  
D6-13ref and D6-13a; or

D3-9ref and D3-9a; or

D4-4ref and D4-4a; or

D7-27ref and D7-27a;

Optionally wherein the first and/or second D gene segment is present in two or more copies.

For example, there are provided two or three copies of the first gene segment, optionally with one, two or three copies of the second gene segment. Copies can be arranged in *cis* or *trans*.

175. The vertebrate, cell or population of clause 174, comprising human gene segments D2-2ref and D2-2a; and D3-3ref and D3-3a; and optionally also D2-15.

In an example, the vertebrate, cell or population comprises one or more D segments selected from human D3-3, D2-15, D3-9; D4-17; D3-10; D2-2; D5-24; D6-19; D3-22; D6-13; D5-12; D1-26; D1-20; D5-18; D3-16; D2-21; D1-14; D7-27; D1-1; D6-25; D2-14 and D4-23 (eg, selected from D3-9\*01; D4-17\*01; D3-10\*01; D2-2\*02; D5-24\*01; D6-19\*01; D3-22\*01; D6-13\*01; D5-12\*01; D1-26\*01; D1-20\*01; D5-18\*01; D3-16\*02; D2-21\*02; D1-14\*01; D7-27\*02; D1-1\*01; D6-25\*01; D2-15\*01; and D4-23\*01), together with the reference sequence(s) of said selected segment(s). These were found in variable domains having a HCDR3 length of at least 20 amino acids (see examples herein).

176. A non-human vertebrate or vertebrate cell according to clause 155, comprising a genome that comprises VH, D and JH gene repertoires comprising human gene segments, the D gene repertoire comprising one or more of

a plurality of D2-2 gene segments provided by at least 2 different D2-2 gene segments in *cis* at the same Ig locus in said genome;

a plurality of D2-21 gene segments provided by at least 2 different D2-21 gene segments in *cis* at the same Ig locus in said genome;

a plurality of D3-10 gene segments provided by at least 2 different D3-10 gene segments in *cis* at the same Ig locus in said genome;

a plurality of D3-16 gene segments provided by at least 2 different D3-16 gene segments in *cis* at the same Ig locus in said genome;

a plurality of D2-8 gene segments provided by at least 2 different D2-8 gene segments in *cis* at the same Ig locus in said genome;

a plurality of D3-3 gene segments provided by at least 2 different D3-3 gene segments in *cis* at the same Ig locus in said genome;

a plurality of D4-23 gene segments provided by at least 2 different D4-23 gene segments in *cis* at the same Ig locus in said genome;

a plurality of D6-13 gene segments provided by at least 2 different D6-13 gene segments in *cis* at the same Ig locus in said genome;

a plurality of D3-9 gene segments provided by at least 2 different D3-9 gene segments in *cis* at the same Ig locus in said genome;

a plurality of D4-4 gene segments provided by at least 2 different D4-4 gene segments in *cis* at the same Ig locus in said genome; and

a plurality of D7-27 gene segments provided by at least 2 different D7-27 gene segments in *cis* at the same Ig locus in said genome;

optionally wherein the D gene segments are derived from the genome sequence of two or more different human individuals.

177. A non-human vertebrate or vertebrate cell according to clause 155, comprising a genome that comprises VH, D and JH gene repertoires comprising human gene segments, the D gene repertoire comprising one or more of

a plurality of D2-2 gene segments provided by at least 2 different D2-2 gene segments in *trans* in said genome;

a plurality of D2-21 gene segments provided by at least 2 different D2-21 gene segments in *trans* in said genome;

a plurality of D3-10 gene segments provided by at least 2 different D3-10 gene segments in *trans* in said genome;

a plurality of D3-16 gene segments provided by at least 2 different D3-16 gene segments in *trans* in said genome;

a plurality of D2-8 gene segments provided by at least 2 different D2-8 gene segments in *trans* in said genome;

a plurality of D3-3 gene segments provided by at least 2 different D3-3 gene segments in *trans* in said genome;

a plurality of D4-23 gene segments provided by at least 2 different D4-23 gene segments in *trans* in said genome;



a plurality of D6-13 gene segments provided by at least 2 different D6-13 gene segments in *trans* in said genome;

a plurality of D3-9 gene segments provided by at least 2 different D3-9 gene segments in *trans* in said genome;

a plurality of D4-4 gene segments provided by at least 2 different D4-4 gene segments in *trans* in said genome; and

a plurality of D7-27 gene segments provided by at least 2 different D7-27 gene segments in *trans* in said genome;

optionally wherein the D gene segments are derived from the genome sequence of two or more different human individuals.

178. A non-human vertebrate or vertebrate cell (optionally an ES cell or B-cell), according to clause 154, comprising a genome that comprises VH, D and JH gene repertoires comprising human gene segments, the D gene repertoire comprising one or more of

a plurality of D2-2 gene segments provided by at least 3 different D2-2 gene segments;

a plurality of D2-21 gene segments provided by at least 3 different D2-21 gene segments;

a plurality of D3-10 gene segments provided by at least 3 different D3-10 gene segments;

a plurality of D3-16 gene segments provided by at least 3 different D3-16 gene segments;

a plurality of D2-8 gene segments provided by at least 3 different D2-8 gene segments;

a plurality of D3-3 gene segments provided by at least 3 different D3-3 gene segments;

a plurality of D4-23 gene segments provided by at least 3 different D4-23 gene segments;

a plurality of D6-13 gene segments provided by at least 3 different D6-13 gene segments;

a plurality of D3-9 gene segments provided by at least 3 different D3-9 gene segments;

a plurality of D4-4 gene segments provided by at least 3 different D4-4 gene segments; and

a plurality of D7-27 gene segments provided by at least 3 different D7-27 gene segments;

optionally wherein the D gene segments are derived from the genome sequence of two or three different human individuals;

optionally wherein at least 2 or 3 of said different gene segments are provided in *cis* at the same

Ig locus in said genome.

179. The vertebrate or cell of clause 176, 177 or 178, wherein the different human individuals are from different human populations.
180. The vertebrate or cell of any one of clauses 176 to 179, wherein the individuals are not genetically related.
181. The vertebrate or cell of any one of clauses 176 to 180, wherein at least one of the different D segments is a synthetic mutant of a human germline D gene segment.
182. A non-human vertebrate or vertebrate cell (optionally an ES cell or B-cell) according to clause 158, comprising a genome comprising human VH, D and JH gene repertoires, the D gene repertoire comprising of one or more of
- a plurality of D2-2 gene segments provided by at least 2 different D2-2 gene ; optionally in *cis* in said genome;
  - a plurality of D2-21 gene segments provided by at least 2 different D2-21 gene ; optionally in *cis* in said genome;
  - a plurality of D3-10 gene segments provided by at least 2 different D3-10 gene ; optionally in *cis* in said genome;
  - a plurality of D3-16 gene segments provided by at least 2 different D3-16 gene ; optionally in *cis* in said genome;
  - a plurality of D2-8 gene segments provided by at least 2 different D2-8 gene ; optionally in *cis* in said genome;
  - a plurality of D3-3 gene segments provided by at least 2 different D3-3 gene ; optionally in *cis* in said genome;
  - a plurality of D4-23 gene segments provided by at least 2 different D4-23 gene ; optionally in *cis* in said genome;
  - a plurality of D6-13 gene segments provided by at least 2 different D6-13 gene ; optionally in *cis* in said genome;
  - a plurality of D3-9 gene segments provided by at least 2 different D3-9 gene ; optionally in *cis* in said genome;
  - a plurality of D4-4 gene segments provided by at least 2 different D4-4 gene ; optionally in *cis* in

said genome; and

a plurality of D7-27 gene segments provided by at least 2 different D7-27 gene ; optionally in *cis* in said genome;

wherein the D gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations.

183. The vertebrate or cell of clause 182, wherein the human individuals are from different human populations.

184. The vertebrate, cell or population of any one of clauses 154 to 183, wherein one or more of the D gene segments is a variant of a human germline D gene segment, wherein the variant gene segment encodes an amino acid sequence that differs by 1, 2 or 3 amino acids from the corresponding amino acid sequence encoded by the human germline D gene segment, provided in that said amino acid sequence encoded by the variant does not include a stop codon when said corresponding amino acid sequence does not include a stop codon.

Optionally, the variant and germline D gene segments encode the respective amino acid sequences in reading frame 2 (IMGT numbering). See Briney *et al* 2012.

185. The vertebrate, cell or population of clause 184, wherein said corresponding amino acid sequence encoded by the human germline D gene segment is a hydrophilic or hydrophobic sequence (according to J Mol Biol. 1997 Jul 25;270(4):587-97; Corbett SJ *et al*; Table 2).

186. The vertebrate, cell or population of clause 184 or 185, comprising said variant and said germline human D gene segments; optionally wherein the variant and germline human D gene segments are *cis* on the same chromosome.

187. The vertebrate, cell or population of any one of clauses 184 to 186, wherein germline human D gene segment is a D2, D3, D5 or D6 family gene segment; optionally a D2-2, D2-15, D3-3, D3-9, D3-10, D3-22, D5-5, D5-18, D6-6, D6-13, D6-19 gene segment.

These D segments are usable in all three reading frames.

Optionally a variant of 1, 2, 3, 4, 5, 6, 7, 8, 9, 10 or all of these human germline D gene segments is used.

188. The vertebrate, cell or population of any one of clauses 154 to 187, comprising a plurality of D2-2 gene segments, wherein the plurality comprises D2-2 gene segments that vary from each other at one or more nucleotide positions corresponding to positions

106,382,687 and

106,382,711

on human chromosome 14.

189. The vertebrate, cell or population of clause 188, wherein the plurality comprises a human D2-2 gene segment ((optionally two copies and/or in homozygous state) comprising a thymine at a position corresponding to position 106,382,687 on human chromosome 14; and optionally no further mutation from the sequence of D2-2ref.

190. The vertebrate, cell or population of clause 188 or 189, wherein the plurality comprises a human D2-2 gene segment comprising a cytosine at a position corresponding to position 106,382,687 on human chromosome 14; and optionally no further mutation from the sequence of D2-2a.

191. The vertebrate, cell or population of any one of clauses 188 to 190, wherein the plurality comprises a human D2-2 gene segment comprising an adenine at a position corresponding to position 106,382,711 on human chromosome 14; and optionally no further mutation from the sequence of D2-2b.

192. The vertebrate, cell or population of any one of clauses 188 to 191, wherein the plurality comprises a human D2-2 gene segment comprising an thymine at a position corresponding to position 106,382,711 on human chromosome 14; and optionally no further mutation from the sequence of D2-2ref.

193. The vertebrate, cell or population of any one of clauses 154 to 192, comprising a plurality of D7-27 gene segments, wherein the plurality comprises D7-27 gene segments that vary from each other at a nucleotide position corresponding to position 106,331,767 on human chromosome 14.

194. The vertebrate, cell or population of clause 193, wherein the plurality comprises a human D7-27 gene segment (optionally two copies and/or in homozygous state) comprising a cytosine at a position corresponding to position 106,331,767 on human chromosome 14; and optionally no further mutation from the sequence of D7-27ref.
195. The vertebrate, cell or population of clause 193 or 194, wherein the plurality comprises a human D7-27 gene segment comprising a guanine at a position corresponding to position 106,331,767 on human chromosome 14; and optionally no further mutation from the sequence of D7-27a.
196. The vertebrate, cell or population of any one of clauses 154 to 195, comprising a plurality of D4-23 gene segments, wherein the plurality comprises D4-23 gene segments that vary from each other at a nucleotide position corresponding to position 106,350,740 on human chromosome 14.
197. The vertebrate, cell or population of clause 196, wherein the plurality comprises a human D4-23 gene segment (optionally two copies and/or in homozygous state) comprising an adenine at a position corresponding to position 106,350,740 on human chromosome 14; and optionally no further mutation from the sequence of D4-23ref.
198. The vertebrate, cell or population of clause 196 or 197, wherein the plurality comprises a human D4-23 gene segment (optionally two copies and/or in homozygous state) comprising an guanine at a position corresponding to position 106,350,740 on human chromosome 14; and optionally no further mutation from the sequence of D4-23a.
199. The vertebrate, cell or population of any one of clauses 154 to 197, comprising a plurality of D2-21 gene segments, wherein the plurality comprises D2-21 gene segments that vary from each other at a nucleotide position corresponding to position 106,354,418 on human chromosome 14.
200. The vertebrate, cell or population of clause 199, wherein the plurality comprises a human D2-21 gene segment (optionally two copies and/or in homozygous state) comprising an adenine

at a position corresponding to position 106,354,418 on human chromosome 14; and optionally no further mutation from the sequence of D2-21ref.

201. The vertebrate, cell or population of clause 199 or 200, wherein the plurality comprises a human D2-21 gene segment (optionally two copies and/or in homozygous state) comprising a guanine at a position corresponding to position 106,354,418 on human chromosome 14; and optionally no further mutation from the sequence of D2-21a.
202. The vertebrate, cell or population of any one of clauses 154 to 201, comprising a plurality of D3-16 gene segments, wherein the plurality comprises D3-16 gene segments that vary from each other at a nucleotide position corresponding to position 106,354,418 on human chromosome 14.
203. The vertebrate, cell or population of clause 202, wherein the plurality comprises a human D3-16 gene segment (optionally two copies and/or in homozygous state) comprising a thymine at a position corresponding to position 106,361,515 on human chromosome 14; and optionally no further mutation from the sequence of D3-16ref.
204. The vertebrate, cell or population of clause 202 or 203, wherein the plurality comprises a human D3-16 gene segment (optionally two copies and/or in homozygous state) comprising a cytosine at a position corresponding to position 106,361,515 on human chromosome 14; and optionally no further mutation from the sequence of D3-16a.
205. The vertebrate, cell or population of any one of clauses 154 to 204, comprising a plurality of D6-13 gene segments, wherein the plurality comprises D6-13 gene segments that vary from each other at a nucleotide position corresponding to position 106,367,013 on human chromosome 14.
206. The vertebrate, cell or population of clause 205, wherein the plurality comprises a human D6-13 gene segment (optionally two copies and/or in homozygous state) comprising a thymine at a position corresponding to position 106,367,013 on human chromosome 14; and optionally no further mutation from the sequence of D6-13ref.

207. The vertebrate, cell or population of clause 205 or 206, wherein the plurality comprises a human D6-13 gene segment (optionally two copies and/or in homozygous state) comprising a cytosine at a position corresponding to position 106,367,013 on human chromosome 14; and optionally no further mutation from the sequence of D6-13a.
208. The vertebrate, cell or population of any one of clauses 154 to 207, comprising a plurality of D3-10 gene segments, wherein the plurality comprises D3-10 gene segments that vary from each other at one or more nucleotide positions corresponding to positions 106,370,370 and 106,370,371 on human chromosome 14.
209. The vertebrate, cell or population of clause 208, wherein the plurality comprises a human D3-10 gene segment (optionally two copies and/or in homozygous state) comprising a thymine at a position corresponding to position 106,370,370 on human chromosome 14; and optionally no further mutation from the sequence of D3-10ref.
210. The vertebrate, cell or population of clause 208 or 209, wherein the plurality comprises a human D3-10 gene segment (optionally two copies and/or in homozygous state) comprising a cytosine at a position corresponding to position 106,370,370 on human chromosome 14; and optionally no further mutation from the sequence of D3-10a.
211. The vertebrate, cell or population of clause 208, 209 or 210 wherein the plurality comprises a human D3-10 gene segment (optionally two copies and/or in homozygous state) comprising an adenine at a position corresponding to position 106,370,371 on human chromosome 14; and optionally no further mutation from the sequence of D3-10ref.
212. The vertebrate, cell or population of any one of clauses 208 to 211, wherein the plurality comprises a human D3-10 gene segment (optionally two copies and/or in homozygous state) comprising a guanine at a position corresponding to position 106,370,371 on human chromosome 14; and optionally no further mutation from the sequence of D3-10b.
213. The vertebrate, cell or population of any one of clauses 154 to 212, comprising a plurality of D3-9 gene segments, wherein the plurality comprises D3-9 gene segments that vary from each

other at a nucleotide position corresponding to position 106,370,567 on human chromosome 14.

214. The vertebrate, cell or population of clause 213, wherein the plurality comprises a human D3-9 gene segment (optionally two copies and/or in homozygous state) comprising an adenine at a position corresponding to position 106,370,567 on human chromosome 14; and optionally no further mutation from the sequence of D3-9ref.
215. The vertebrate, cell or population of clause 213 or 214, wherein the plurality comprises a human D3-9 gene segment (optionally two copies and/or in homozygous state) comprising a thymine at a position corresponding to position 106,370,567 on human chromosome 14; and optionally no further mutation from the sequence of D3-9a.
216. The vertebrate, cell or population of any one of clauses 154 to 215, comprising a plurality of D2-8 gene segments, wherein the plurality comprises D2-8 gene segments that vary from each other at one or more nucleotide positions corresponding to positions  
106,373,085;  
106,373,086 and  
106,373,089  
on human chromosome 14.
217. The vertebrate, cell or population of clause 216, wherein the plurality comprises a human D2-8 gene segment (optionally two copies and/or in homozygous state) comprising a cytosine at a position corresponding to position 106,373,085 on human chromosome 14.
218. The vertebrate, cell or population of clause 216 or 217, wherein the plurality comprises a human D2-8 gene segment (optionally two copies and/or in homozygous state) comprising a thymine at a position corresponding to position 106,373,085 on human chromosome 14; and optionally no further mutation from the sequence of D2-8b.
219. The vertebrate, cell or population of clause 216, 217 or 218 wherein the plurality comprises a human D2-8 gene segment (optionally two copies and/or in homozygous state) comprising a cytosine at a position corresponding to position 106,373,086 on human chromosome 14; and



optionally no further mutation from the sequence of D2-8ref.

220. The vertebrate, cell or population of any one of clauses 216 to 219, wherein the plurality comprises a human D2-8 gene segment comprising a thymine at a position corresponding to position 106,373,086 on human chromosome 14; and optionally no further mutation from the sequence of D2-8ref.
221. The vertebrate, cell or population of any one of clauses 154 to 220, comprising a plurality of D4-4 gene segments, wherein the plurality comprises D4-4 gene segments that vary from each other at one or more nucleotide positions corresponding to positions 106,379,086; and 106,379,089 on human chromosome 14.
222. The vertebrate, cell or population of clause 221, wherein the plurality comprises a D4-4 gene segment (optionally two copies and/or in homozygous state) comprising a cytosine at a position corresponding to position 106,379,086 on human chromosome 14; and optionally no further mutation from the sequence of D4-4ref.
223. The vertebrate, cell or population of clause 221 or 222, wherein the plurality comprises a human D4-4 gene segment (optionally two copies and/or in homozygous state) comprising a thymine at a position corresponding to position 106,379,086 on human chromosome 14; and optionally no further mutation from the sequence of D4-4a.
224. The vertebrate, cell or population of clause 221, 222 or 223 wherein the plurality comprises a human D4-4 gene segment (optionally two copies and/or in homozygous state) comprising a cytosine at a position corresponding to position 106,379,089 on human chromosome 14; and optionally no further mutation from the sequence of D4-4ref or a cytosine at a position corresponding to position 106,379,086 on human chromosome 14.
225. The vertebrate, cell or population of any one of clauses 221 to 224, wherein the plurality comprises a human D4-4 gene segment (optionally two copies and/or in homozygous state) comprising a thymine at a position corresponding to position 106,373,089 on human

chromosome 14; and optionally no further mutation from the sequence of D4-4a.

226. The vertebrate, cell or population of any one of clauses 154 to 225, comprising a plurality of D3-3 gene segments, wherein the plurality comprises D3-3 gene segments that vary from each other at one or more nucleotide positions corresponding to positions 106,380,241; and 106,380,246 on human chromosome 14.
227. The vertebrate, cell or population of clause 226, wherein the plurality comprises a D3-3 gene segment (optionally two copies and/or in homozygous state) comprising a thymine at a position corresponding to position 106,380,241 on human chromosome 14; and optionally no further mutation from the sequence of D3-3ref.
228. The vertebrate, cell or population of clause 226 or 227, wherein the plurality comprises a human D3-3 gene segment (optionally two copies and/or in homozygous state) comprising a cytosine at a position corresponding to position 106,380,241 on human chromosome 14; and optionally no further mutation from the sequence of D3-3a.
229. The vertebrate, cell or population of clause 226, 227 or 228 wherein the plurality comprises a human D3-3 gene segment (optionally two copies and/or in homozygous state) comprising an adenine at a position corresponding to position 106,380,246 on human chromosome 14; and optionally no further mutation from the sequence of D3-3ref.
230. The vertebrate, cell or population of any one of clauses 226 to 229, wherein the plurality comprises a human D3-3 gene segment (optionally two copies and/or in homozygous state) comprising a thymine at a position corresponding to position 106,380,246 on human chromosome 14; and optionally no further mutation from the sequence of D3-3a.

#### Multiple Human JL Gene Segment Variants

A specific application of this configuration is the provision of multiple human JL gene segments (Jk and/or Jλ) as follows (as set out in numbered paragraphs, starting at paragraph number 80).

80. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising at least 3 human JL gene segments of the same type (eg, Jk1), wherein at least two of the human JL gene segments are variants that are not identical to each other.
81. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising at least 2 different non-endogenous JL gene segments of the same type (eg, Jk1) *cis* at the same Ig locus.
82. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising at least 2 different human JL gene segments of the same type (eg, Jk1) *trans* at the same Ig locus; and optionally a third human JL gene segment of the same type, wherein the third JL is *cis* with one of said 2 different JL gene segments.
83. A population of non-human vertebrates (eg, mice or rats) comprising a repertoire of human JL gene segments, wherein the plurality comprises at least 2 different human JL gene segments of the same type (eg, Jk1), a first of said different JL gene segments is provided in the genome of a first vertebrate of the population, and a second of said different JL gene segments being provided in the genome of a second vertebrate of the population, wherein the genome of the first vertebrate does not comprise the second JL gene segment.
84. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising at least 2 different non-endogenous JL gene segments of the same type (eg, Jk1), wherein the JL gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations.
85. A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 3 human JL gene segments of the same type (eg, Jk1), wherein at least two of the human JL gene segments are variants that are not identical to each other.
86. A method of enhancing the immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 2 different non-endogenous JL gene segments of the same type (eg, Jk1) *cis* at the same Ig

locus.

87. A method of enhancing the immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 2 different human JL gene segments of the same type(eg, J $\kappa$ 1) *trans* at the same Ig locus; and optionally a third human JL gene segment of the same type, wherein the third JL is *cis* with one of said 2 different JL gene segments.
88. A method of providing an enhanced human immunoglobulin JL gene segment repertoire, the method comprising providing a population of non-human vertebrates (eg, a mouse or rat) comprising a repertoire of human JL gene segments, wherein the method comprises providing at least 2 different human JL gene segments of the same type (eg, J $\kappa$ 1), wherein a first of said different JL gene segments is provided in the genome of a first vertebrate of the population, and a second of said different JL gene segments is provided in the genome of a second vertebrate of the population, wherein the genome of the first vertebrate does not comprise the second JL gene segment.
89. A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 2 different non-endogenous JL gene segments of the same type (eg, J $\kappa$ 1), wherein the JL gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations.
90. The vertebrate or cell of paragraph 80, 82 or 84, or the method of paragraph 85, 82 or 89, wherein at least 2 or 3 of said different gene segments are provided *cis* at the same Ig locus in said genome.
91. The vertebrate or cell of paragraph 80, 81 or 82, or the method of paragraph 85, 86 or 87, wherein the JL gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations.
92. The vertebrate or cell of paragraph 80, 81 or 82, or the method of paragraph 85, 86 or 87, wherein the JL gene segments are derived from the genome sequence of two or more different human individuals; optionally wherein the different human individuals are from different human

populations.

93. The vertebrate, cell or method of paragraph 92, wherein the individuals are not genetically related.
94. The vertebrate, cell or method of any one of paragraphs 80 to 93, wherein at least one of the different JL segments is a synthetic mutant of a human germline JL gene segment.
95. A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 2 human JL gene segments of the same type (eg, Jk1), wherein the JL gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations; optionally wherein at least 2 or 3 of said different gene segments are provided at the same IgL locus in said genome.
96. The vertebrate or cell of any one of paragraphs paragraph 80 to 82 and 84, wherein the genome comprises a substantially complete functional repertoire of human Jk and/or Jλ gene segment types supplemented with one, two or more human Jk and/or Jλ gene segments respectively, wherein said substantially complete functional repertoire and the supplementary gene segments are not found together in the germline genome of a human individual.
97. The population of paragraph 83, wherein the population comprises a substantially complete functional repertoire of human JL gene segment types supplemented with one, two or more human Jk and/or Jλ gene segments respectively, wherein said substantially complete functional repertoire and the supplementary gene segments are not found together in the germline genome of a human individual.
98. A non-human vertebrate (eg, a mouse or rat) or a non-human cell (eg, an ES cell or a B-cell) having a genome comprising a substantially complete functional repertoire of human Jk and/or Jλ gene segment types supplemented with one, two or more human Jk and/or Jλ gene segments respectively, wherein said substantially complete functional repertoire and the supplementary gene segments are not found together in the germline genome of a human individual.

99. A population of non-human vertebrates (eg, mice or rats) comprising a substantially complete functional repertoire of human J $\kappa$  and/or J $\lambda$  gene segment types supplemented with one, two or more human J $\kappa$  and/or J $\lambda$  gene segments respectively, wherein said substantially complete functional repertoire and the supplementary gene segments are not found together in the germline genome of a human individual.
100. A non-human vertebrate or vertebrate cell according to paragraph 81, comprising a genome that comprises VL and JL gene repertoires comprising human gene segments, the JL gene repertoire comprising
- a plurality of human J $\kappa$ 1 gene segments provided by at least 2 different human J $\kappa$ 1 gene segments in *cis* at the same Ig locus in said genome;
  - a plurality of human J $\kappa$ 2 gene segments provided by at least 2 different human J $\kappa$ 1 gene segments in *cis* at the same Ig locus in said genome;
  - a plurality of human J $\kappa$ 3 gene segments provided by at least 2 different human J $\kappa$ 1 gene segments in *cis* at the same Ig locus in said genome;
  - a plurality of human J $\kappa$ 4 gene segments provided by at least 2 different human J $\kappa$ 1 gene segments in *cis* at the same Ig locus in said genome;
  - a plurality of human J $\kappa$ 5 gene segments provided by at least 2 different human J $\kappa$ 1 gene segments in *cis* at the same Ig locus in said genome;
  - a plurality of human J $\lambda$ 1 gene segments provided by at least 2 different human J $\lambda$ 1 gene segments in *cis* at the same Ig locus in said genome;
  - a plurality of human J $\lambda$ 2 gene segments provided by at least 2 different human J $\lambda$ 2 gene segments in *cis* at the same Ig locus in said genome;
  - a plurality of human J $\lambda$ 3 gene segments provided by at least 2 different human J $\lambda$ 3 gene segments in *cis* at the same Ig locus in said genome;
  - a plurality of human J $\lambda$ 4 gene segments provided by at least 2 different human J $\lambda$ 4 gene segments in *cis* at the same Ig locus in said genome;
  - a plurality of human J $\lambda$ 5 gene segments provided by at least 2 different human J $\lambda$ 5 gene segments in *cis* at the same Ig locus in said genome;
  - a plurality of human J $\lambda$ 6 gene segments provided by at least 2 different human J $\lambda$ 6 gene segments in *cis* at the same Ig locus in said genome; or
  - a plurality of human J $\lambda$ 7 gene segments provided by at least 2 different human J $\lambda$ 7 gene segments in *cis* at the same Ig locus in said genome;

optionally wherein the JL gene segments are derived from the genome sequence of two or more different human individuals.

101. A non-human vertebrate or vertebrate cell (optionally an ES cell or B-cell), according to paragraph 80, comprising a genome that comprises VL and JL gene repertoires comprising human gene segments, the JL gene repertoire comprising

a plurality of human J $\kappa$ 1 gene segments provided by at least 3 (eg, 3, 4, 5, 6, or 7) different human J $\kappa$ 1 gene segments;

a plurality of human J $\kappa$ 2 gene segments provided by at least 3 (eg, 3, 4, 5, 6, or 7) different human J $\kappa$ 2 gene segments;

a plurality of human J $\kappa$ 3 gene segments provided by at least 3 (eg, 3, 4, 5, 6, or 7) different human J $\kappa$ 3 gene segments;

a plurality of human J $\kappa$ 4 gene segments provided by at least 3 (eg, 3, 4, 5, 6, or 7) different human J $\kappa$ 4 gene segments;

a plurality of human J $\kappa$ 5 gene segments provided by at least 3 (eg, 3, 4, 5, 6, or 7) different human J $\kappa$ 5 gene segments;

a plurality of human J $\lambda$ 1 gene segments provided by at least 3 (eg, 3, 4, 5, 6, or 7) different human J $\lambda$ 1 gene segments;

a plurality of human J $\lambda$ 2 gene segments provided by at least 3 (eg, 3, 4, 5, 6, or 7) different human J $\lambda$ 2 gene segments;

a plurality of human J $\lambda$ 3 gene segments provided by at least 3 (eg, 3, 4, 5, 6, or 7) different human J $\lambda$ 3 gene segments;

a plurality of human J $\lambda$ 4 gene segments provided by at least 3 (eg, 3, 4, 5, 6, or 7) different human J $\lambda$ 4 gene segments;

a plurality of human J $\lambda$ 5 gene segments provided by at least 3 (eg, 3, 4, 5, 6, or 7) different human J $\lambda$ 5 gene segments;

a plurality of human J $\lambda$ 6 gene segments provided by at least 3 (eg, 3, 4, 5, 6, or 7) different human J $\lambda$ 6 gene segments; or

a plurality of human J $\lambda$ 7 gene segments provided by at least 3 (eg, 3, 4, 5, 6, or 7) different human J $\lambda$ 7 gene segments;

optionally wherein the JL gene segments are derived from the genome sequence of two or three

different human individuals;

optionally wherein at least 2 or 3 of said different gene segments are provided in *cis* at the same Ig locus in said genome.

102. The vertebrate or cell of paragraph 104 or 105, wherein the different human individuals are from different human populations.

103. The vertebrate or cell of any one of paragraphs 104 to 106, wherein the individuals are not genetically related.

104. The vertebrate or cell of any one of paragraphs 104 to 107, wherein at least one of the different JL segments is a synthetic mutant of a human germline JL gene segment.

105. A non-human vertebrate or vertebrate cell (optionally an ES cell or B-cell) according to paragraph 84, comprising a genome comprising human VL and JL gene repertoires, the JL gene repertoire comprising

a plurality of human J $\kappa$ 1 gene segments provided by at least 2 different human J $\kappa$ 1 gene segments, optionally in *cis* at the same Ig locus in said genome;

a plurality of human J $\kappa$ 2 gene segments provided by at least 2 different human J $\kappa$ 1 gene segments, optionally in *cis* at the same Ig locus in said genome;

a plurality of human J $\kappa$ 3 gene segments provided by at least 2 different human J $\kappa$ 1 gene segments, optionally in *cis* at the same Ig locus in said genome;

a plurality of human J $\kappa$ 4 gene segments provided by at least 2 different human J $\kappa$ 1 gene segments, optionally in *cis* at the same Ig locus in said genome;

a plurality of human J $\kappa$ 5 gene segments provided by at least 2 different human J $\kappa$ 1 gene segments, optionally in *cis* at the same Ig locus in said genome;

a plurality of human J $\lambda$ 1 gene segments provided by at least 2 different human J $\lambda$ 1 gene segments, optionally in *cis* at the same Ig locus in said genome;

a plurality of human J $\lambda$ 2 gene segments provided by at least 2 different human J $\lambda$ 2 gene segments, optionally in *cis* at the same Ig locus in said genome;

a plurality of human J $\lambda$ 3 gene segments provided by at least 2 different human J $\lambda$ 3 gene segments, optionally in *cis* at the same Ig locus in said genome;



a plurality of human Jλ4 gene segments provided by at least 2 different human Jλ4 gene segments, optionally in *cis* at the same Ig locus in said genome;

a plurality of human Jλ5 gene segments provided by at least 2 different human Jλ5 gene segments, optionally in *cis* at the same Ig locus in said genome;

a plurality of human Jλ6 gene segments provided by at least 2 different human Jλ6 gene segments, optionally in *cis* at the same Ig locus in said genome; or

a plurality of human Jλ7 gene segments provided by at least 2 different human Jλ7 gene segments, optionally in *cis* at the same Ig locus in said genome;

wherein the JL gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations.

106. The vertebrate or cell of paragraph 109, wherein the human individuals are from different human populations.

The skilled person will realise that standard molecular biology techniques can be used to provide vectors comprising synthetic combinations of immunoglobulin gene segments (eg, V, D and/or J) for use in the invention, such that the vectors can be used to build a transgenic immunoglobulin locus (eg, using homologous recombination and/or recombinase mediated cassette exchange as known in the art, eg, see US7501552 (Medarex), US5939598 (Abgenix), US6130364 (Abgenix), WO02/066630 (Regeneron), WO2011004192 (Genome Research Limited), WO2009076464, WO2009143472 and WO2010039900 (Ablexis), the disclosures of which are explicitly incorporated herein. For example, such synthetic combinations of gene segments can be made using standard recombineering techniques in E coli to construct BAC vectors harbouring the synthetic combination prior to insertion in embryonic stem cells using homologous recombination or RMCE (eg, using cre/lox site-specific recombination). Details of recombineering can be found at [www.genebridges.com](http://www.genebridges.com) and in EP1034260 and EP1204740 the disclosures of which are explicitly incorporated herein.

In one embodiment, it is useful to bias the immune response of the vertebrate (and thus resultant lead antibodies) to a predetermined gene segment, eg, one known to be commonly used in natural human immune responses to antigens, such as antigens of infectious disease pathogens. For example, VH1-69 is commonly used to produce antibodies in humans against Influenza virus; it is possible, therefore, to include two or more polymorphic DNA versions of the VH segment VH1-69 in

the locus of the invention. The examples below illustrate how such a transgenic locus can be constructed in which diversity is extended by extending the VH1-69 gene segment repertoire based on naturally-occurring VH1-69 polymorphic variants.

In one embodiment in any configuration of the invention, the genome has been modified to prevent or reduce the expression of fully-endogenous antibody. Examples of suitable techniques for doing this can be found in PCT/GB2010/051122, US7501552, US6673986, US6130364, WO2009/076464, EP1399559 and US6586251, the disclosures of which are incorporated herein by reference. In one embodiment, the non-human vertebrate VDJ region of the endogenous heavy chain immunoglobulin locus, and optionally VJ region of the endogenous light chain immunoglobulin loci (lambda and/or kappa loci), have been inactivated. For example, all or part of the non-human vertebrate VDJ region is inactivated by inversion in the endogenous heavy chain immunoglobulin locus of the mammal, optionally with the inverted region being moved upstream or downstream of the endogenous Ig locus (see, eg, WO2011004192, the disclosure of which is incorporated herein by reference). For example, all or part of the non-human vertebrate VJ region is inactivated by inversion in the endogenous kappa chain immunoglobulin locus of the mammal, optionally with the inverted region being moved upstream or downstream of the endogenous Ig locus. For example, all or part of the non-human vertebrate VJ region is inactivated by inversion in the endogenous lambda chain immunoglobulin locus of the mammal, optionally with the inverted region being moved upstream or downstream of the endogenous Ig locus. In one embodiment the endogenous heavy chain locus is inactivated in this way as is one or both of the endogenous kappa and lambda loci.

Additionally or alternatively, the vertebrate has been generated in a genetic background which prevents the production of mature host B and T lymphocytes, optionally a RAG-1-deficient and/or RAG-2 deficient background. See US5859301 for techniques of generating RAG-1 deficient animals.

Thus, in one embodiment of any configuration or aspect of the invention herein, endogenous heavy and light chain expression has been inactivated.

In one embodiment each said locus constant region is a heavy chain endogenous non-human vertebrate (optionally host mouse or rat) constant region.

In one embodiment each said locus constant region is a light chain endogenous non-human vertebrate (optionally host mouse or rat) constant region.

The invention provides a monoclonal or polyclonal antibody composition prepared by immunisation of at least one vertebrate (eg, mouse or rat) according to the invention, optionally wherein the antigen is an antigen of an infectious disease pathogen (eg, a bacterial or viral pathogen antigen), optionally wherein the same antigen is used to immunise all the vertebrates; optionally wherein the antibody or antibodies are IgG-type (eg, IgG1).

The invention also provides a monoclonal or polyclonal antibody mixture produced by the method of the invention or a derivative antibody or mixture thereof, eg, where one or more constant region has been changed (eg, replaced with a different constant region such as a human constant region; or mutated to enhance or ablate Fc effector function). In an aspect of the invention, the monoclonal or polyclonal antibody mixture is provided for therapy and/or prophylaxis of a disease or condition in a human, eg, for the treatment and/or prevention of an infectious disease, wherein optionally wherein each antibody binds an antigen of an infectious disease pathogen, preferably the same antigen.

In an aspect of the invention, there is provided the use of an isolated, monoclonal or polyclonal antibody according to the invention, or a mutant or derivative antibody thereof in the manufacture of a medicament for the treatment and/or prevention of a disease or condition in a human, eg, an infectious disease, optionally wherein the infectious disease is a disease caused by a bacterial or viral pathogen.

An example of a mutant antibody is one that bears up to 15 or 10 amino acid mutations in its variable regions relative to an isolated antibody (eg, IgG-type, such as IgG1-type, antibody) obtainable or obtained by the method of the invention. An example of a derivative is one that has been modified to replace a constant region with a different constant region such as a human constant region; or mutated to enhance or ablate Fc effector function.

Examples of infectious diseases are diseases caused or mediated by a bacterial or viral pathogen. For example, the infectious disease is selected from the group consisting of a disease caused by a pathogen selected from the group consisting of Haemophilus influenza, E coli, Neisseria meningitidis, a herpes family virus, cytomegalovirus (CMV), HIV and influenza virus.

**Tailoring V(D)J Incorporation Into Immunoglobulin Loci For The Generation of Antibodies Against Infectious Disease**

The inventors realised that it would be desirable to provide for vertebrates, cells, methods etc for the production of therapeutic and/or prophylactic antibodies based on natural human immune responses to antigens, such as antigens of infectious disease pathogens. In this respect, the literature observes frequently used immunoglobulin gene segments to raise anti-infective responses in humans (Table 9).

In the various configurations, aspects, embodiments and examples above, the invention provides the skilled addressee with the possibility of choosing immunoglobulin gene segments in a way that tailors or biases the repertoire for application to generating antibodies to treat and/or prevent infectious diseases. The inventors have categorised the following groups of gene segments for use in the invention according to the desired application of resultant antibodies.

**List A:**

**Immunoglobulin gene segments for antibodies that bind an antigen expressed by a Pathogen**

- (a) a VL gene segment selected from the group consisting of a V<sub>λ</sub>II gene family member, V<sub>λ</sub>VII 4A, V<sub>λ</sub>II 2.1, V<sub>λ</sub>VII 4A, a V<sub>λ</sub>1 gene family member, a V<sub>λ</sub>3 gene family member, IGLV1S2, V<sub>λ</sub>3-cML70, Iah2, Ialvl, Ia3h3, Kv325, a V<sub>κ</sub>I gene family member, κI-15A (KL012), V<sub>κ</sub>II family member, a V<sub>κ</sub>III family member, a V<sub>κ</sub>I gene family member, κI-15A (KL012), V<sub>κ</sub>II A2 (optionally the A2a variant), V<sub>κ</sub>A27 (Humkv325) and a gene segment at least 80% identical thereto.
- (b) a V<sub>λ</sub> gene segment selected from a V<sub>λ</sub>II gene family member, V<sub>λ</sub>VII 4A, V<sub>λ</sub>II 2.1, V<sub>λ</sub>VII 4A, a V<sub>λ</sub>1 gene family member, a V<sub>λ</sub>3 gene family member, IGLV1S2, V<sub>λ</sub>3-cML70, Iah2, Ialvl, Ia3h3 and a gene segment at least 80% identical thereto.
- (c) a V<sub>κ</sub> gene segment selected from Kv325, a V<sub>κ</sub>I gene family member, κI-15A (KL012), V<sub>κ</sub>II family member, a V<sub>κ</sub>III family member, a V<sub>κ</sub>I gene family member, κI-15A (KL012), V<sub>κ</sub>II A2 (optionally the A2a variant), V<sub>κ</sub>A27 (Humkv325) and a gene segment at least 80% identical thereto.
- (d) a V<sub>H</sub> gene segment a V<sub>H</sub>III gene family member (optionally, a VHIIIa or VHIIIb family member), a V<sub>H</sub>IV gene family member, V<sub>H</sub>III 9.1 (VH3-15), V<sub>H</sub>III VH26 (VH3-23), V<sub>H</sub>3-21, LSG6.1, LSG12.1, DP77 (V3-21), V<sub>H</sub>H11, VH1GRR, ha3h2, V<sub>H</sub>I-ha1c1, V<sub>H</sub>III-VH2-1, VH4.18, ha4h3, Hv1051, 71-2, Hv1f10, VH4.11, 71-4, VH251, VH1-69 and a gene segment at least 80% identical thereto.
- (e) a J<sub>λ</sub> gene segment selected from J<sub>λ</sub>2, J<sub>λ</sub>3 and a gene segment at least 80% identical thereto.

- (f) a D gene segment selected from Dk1, Dxp'1, Dn4r, D2r and a gene segment at least 80% identical thereto.

List A1:

Immunoglobulin gene segments for antibodies that bind an antigen expressed by a Bacterial Pathogen

- (a) a  $V_{\lambda}$  gene segment selected from a  $V_{\lambda}$ II gene family member,  $V_{\lambda}$ VII 4A,  $V_{\lambda}$ II 2.1,  $V_{\lambda}$ VII 4A and a gene segment at least 80% identical thereto.
- (b) a  $V_{\kappa}$  gene segment selected from a  $V_{\kappa}$ I gene family member,  $\kappa$ I-15A (KL012),  $V_{\kappa}$ II family member, a  $V_{\kappa}$ III family member, a  $V_{\kappa}$ I gene family member,  $\kappa$ I-15A (KL012),  $V_{\kappa}$ II A2 (optionally the A2a variant),  $V_{\kappa}$  A27 (Humkv325) and a gene segment at least 80% identical thereto.
- (c) a  $V_H$  gene segment a VH3 gene family member (optionally, a VHIIIa or VHIIIb family member),  $V_H$ III 9.1 (VH3-15),  $V_H$ III VH26 (VH3-23),  $V_H$ 3-21, LSG6.1, LSG12.1, DP77 (V3-21),  $V_H$  H11 and a gene segment at least 80% identical thereto.
- (d) a  $J_{\lambda}$  gene segment selected from  $J_{\lambda}$ 2,  $J_{\lambda}$ 3 and a gene segment at least 80% identical thereto.
- (e) a  $J_H$  gene segment selected from  $J_H$ 2,  $J_H$ 3,  $J_H$ 4 and a gene segment at least 80% identical thereto.

List A1.1:

Immunoglobulin gene segments for antibodies that bind an antigen expressed by *H influenza*

- (a) a  $V_{\lambda}$  gene segment selected from a  $V_{\lambda}$ II gene family member,  $V_{\lambda}$ VII 4A,  $V_{\lambda}$ II 2.1,  $V_{\lambda}$ VII 4A and a gene segment at least 80% identical thereto.
- (b) a  $V_{\kappa}$  gene segment selected from a  $V_{\kappa}$ II family member, a  $V_{\kappa}$ III family member, a  $V_{\kappa}$ I gene family member,  $\kappa$ I-15A (KL012),  $V_{\kappa}$ II A2 (optionally the A2a variant),  $V_{\kappa}$  A27 (Humkv325) and a gene segment at least 80% identical thereto.
- (c) a  $V_H$  gene segment a VH3 gene family member (optionally, a VHIIIb family member),  $V_H$ III 9.1 (VH3-15),  $V_H$ III VH26 (VH3-23),  $V_H$ 3-21, LSG6.1, LSG12.1, DP77 (V3-21) and a gene segment at least 80% identical thereto.
- (d) a  $J_{\lambda}$  gene segment selected from  $J_{\lambda}$ 2,  $J_{\lambda}$ 3 and a gene segment at least 80% identical thereto.

List A1.2:

Immunoglobulin gene segments for antibodies that bind an antigen expressed by E Coli or Neisseria meningitidis

- (a) a V<sub>H</sub> gene segment a VH3 gene family member (optionally a VHIIIa or VHIIIb member), V<sub>H</sub>III 9.1 (VH3-15), V<sub>H</sub> H11, V<sub>H</sub>III VH26 (VH3-23) a gene segment at least 80% identical thereto, eg, V<sub>H</sub>III 9.1 + J<sub>H</sub>3; or V<sub>H</sub> H11 + J<sub>H</sub>4; or V<sub>H</sub>III VH26 + J<sub>H</sub>2.
- (b) a V<sub>K</sub> gene segment selected from a VκI gene family member, κI-15A (KL012) and a gene segment at least 80% identical thereto.
- (c) a V<sub>λ</sub> gene segment selected from a VλII gene family member, VλII 2.1 and a gene segment at least 80% identical thereto.
- (d) a J<sub>H</sub> gene segment selected from J<sub>H</sub>2, J<sub>H</sub>3, J<sub>H</sub>4 and a gene segment at least 80% identical thereto.

A2:Immunoglobulin gene segments for antibodies that bind an antigen expressed by a viral Pathogen

- (a) a V<sub>H</sub> gene segment selected from a V<sub>H</sub>III gene family member, a V<sub>H</sub>IV gene family member, V<sub>H</sub>III-VH26 (VH3-23), VH1GRR, ha3h2, V<sub>H</sub>I-ha1c1, V<sub>H</sub>III-VH2-1, VH4.18, ha4h3, Hv1051, 71-2, Hv1f10, VH4.11, 71-4, VH251, VH1-69 and a gene segment at least 80% identical thereto.
- (b) a V<sub>λ</sub> gene segment selected from a Vλ1 gene family member, a Vλ3 gene family member, IGLV1S2, Vλ3-cML70, lalh2, lalvl, la3h3 and a gene segment at least 80% identical thereto.
- (c) a V<sub>K</sub> gene segment selected from Kv325 and a gene segment at least 80% identical thereto.
- (d) a J<sub>H</sub> gene segment selected from J<sub>H</sub>3, J<sub>H</sub>5, J<sub>H</sub>6 and a gene segment at least 80% identical thereto.
- (e) a D gene segment selected from Dk1, Dxp'1, Dn4r, D2r and a gene segment at least 80% identical thereto.
- (f) a J<sub>λ</sub> gene segment selected from Jλ2, Jλ3 and a gene segment at least 80% identical thereto.

A2.1:Immunoglobulin gene segments for antibodies that bind an antigen expressed by Herpes Virus Family (eg, VZV or HSV)

- (a) a V<sub>H</sub> gene segment selected from a V<sub>H</sub>III gene family member, a V<sub>H</sub>IV gene family member, V<sub>H</sub>III-VH26 (VH3-23), VH1GRR, ha3h2, V<sub>H</sub>I-ha1c1, V<sub>H</sub>III-VH2-1, VH4.18, ha4h3, and a gene segment at least 80% identical thereto.

- (b) a  $V_{\lambda}$  gene segment selected from a  $V_{\lambda 1}$  gene family member, a  $V_{\lambda 3}$  gene family member, IGLV1S2,  $V_{\lambda 3}$ -cML70, Iah2, Ialvl, Ia3h3 and a gene segment at least 80% identical thereto.
- (c) a  $J_H$  gene segment selected from  $J_H3$ ,  $J_H5$ ,  $J_H6$  and a gene segment at least 80% identical thereto.
- (d) a D gene segment selected from Dk1, Dxp'1, Dn4r, D2r and a gene segment at least 80% identical thereto.
- (e) a  $J_{\lambda}$  gene segment selected from  $J_{\lambda 2}$ ,  $J_{\lambda 3}$  and a gene segment at least 80% identical thereto.

A2.2:Immunoglobulin gene segments for antibodies that bind an antigen expressed by CMV

- (a) a  $V_H$  gene segment selected from Hv1051 and a gene segment at least 80% identical thereto.
- (b) a  $V_k$  gene segment selected from Kv325 and a gene segment at least 80% identical thereto.

A2.3:Immunoglobulin gene segments for antibodies that bind an antigen expressed by HIV

- (a) a  $V_H$  gene segment selected from 71-2, Hv1f10, VH4.11, 71-4, VH251, VH1-69 and a gene segment at least 80% identical thereto.

A2.4:Immunoglobulin gene segments for antibodies that bind an antigen expressed by Influenza Virus

- (a) a  $V_H$  gene segment selected from VH1-69 and a gene segment at least 80% identical thereto.

Thus,

Where one wishes to generate an antibody or antibody mixture to treat and/or prevent an infectious disease, one or more V, D and/or all J gene segments used in any configuration, aspect, method, example or embodiment of the invention can be selected from List A1. Thus, for example in (a) of the first configuration of the invention, the recited heavy chain V gene segment is selected from the VH gene segments in List A, optionally with a D in that list.

Where one wishes to generate an antibody or antibody mixture to treat and/or prevent an infectious disease caused or mediated by a bacterial pathogen, one or more or all V, D and/or J gene

segments used in any configuration, aspect, method, example or embodiment of the invention can be selected from List A1.

Where one wishes to generate an antibody or antibody mixture to treat and/or prevent an infectious disease caused or mediated by a viral pathogen, one or more or all V, D and/or J gene segments used in any configuration, aspect, method, example or embodiment of the invention can be selected from List A2.

Where one wishes to generate an antibody or antibody mixture to treat and/or prevent an infectious disease caused or mediated by *H influenza*, one or more or all V, D and/or J gene segments used in any configuration, aspect, method, example or embodiment of the invention can be selected from List A1.1.

Where one wishes to generate an antibody or antibody mixture to treat and/or prevent an infectious disease caused or mediated by *E Coli* or *Neisseria meningitidis*, one or more or all V, D and/or J gene segments used in any configuration, aspect, method, example or embodiment of the invention can be selected from List A1.2.

Where one wishes to generate an antibody or antibody mixture to treat and/or prevent an infectious disease caused or mediated by Herpes Virus Family (eg, VZV or HSV), one or more or all V, D and/or J gene segments used in any configuration, aspect, method, example or embodiment of the invention can be selected from List A2.1.

Where one wishes to generate an antibody or antibody mixture to treat and/or prevent an infectious disease caused or mediated by CMV, one or more or all V, D and/or J gene segments used in any configuration, aspect, method, example or embodiment of the invention can be selected from List A2.2.

Where one wishes to generate an antibody or antibody mixture to treat and/or prevent an infectious disease caused or mediated by HIV, one or more or all V, D and/or J gene segments used in any configuration, aspect, method, example or embodiment of the invention can be selected from List A2.3.

Where one wishes to generate an antibody or antibody mixture to treat and/or prevent an infectious disease caused or mediated by Influenza Virus, one or more or all V, D and/or J gene



segments used in any configuration, aspect, method, example or embodiment of the invention can be selected from List A2.4.

Optionally each VH segment in the locus of the invention is selected from List A1, A2, A1.1, A1.2, A2.1, A2.2, A2.3 or A2.4.

Optionally each VL segment in the locus of the invention is selected from List A1, A2, A1.1, A1.2, A2.1, A2.2, A2.3 or A2.4

Optionally each D segment in the locus of the invention is selected from List A1, A2, A1.1, A1.2, A2.1, A2.2, A2.3 or A2.4.

Optionally each J<sub>L</sub> segment in the locus of the invention is selected from List A1, A2, A1.1, A1.2, A2.1, A2.2, A2.3 or A2.4.

#### Antibodies For Therapy & Prophylaxis of Patients of Specific Ancestry

The inventors, having undertaken the extensive Bioinformatics analysis exercise described herein, realised that the output of that analysis has made it possible to identify specific gene segments that are useful to produce antibody- and VH domain-based drugs that are tailored specifically to a patient's ancestry (ie, genotype). That is, antibodies can be selected on the basis that they are made *in vivo* in a transgenic non-human vertebrate (eg, mouse or rat with transgenic IgH loci) and particularly derived from gene segments that are relatively prevalent in members of the patient's population, ie, from individuals of the same human ancestry. Since variant distributions differ across different populations (see Table 13), this presumably reflects the effects of evolution, adaptation and conservation of useful variant gene types in those populations. Thus, by tailoring the antibody-based drugs according to the invention, it is possible to match the drug to the population gene biases, thus with the aim of making better drugs for that specific population of humans. Better can, for example, mean more efficacious, better neutralising, higher target antigen affinity, less immunogenic, less patient reactions to the drug etc. This can be determined empirically, as is standard in drug research and development processes.

Thus, the invention provides the following embodiments (numbered from clause 345 onwards):-

345. An isolated antibody for administration to a Chinese patient, the antibody comprising a human heavy chain, the heavy chain comprising a variable domain that is specific for an antigen

and a constant region, wherein the constant region is a human constant region selected from a constant region (eg, an IGHG constant region) in Table 13 found in a Chinese population and with a cumulative frequency of at least 1 or 5%, and wherein

(i) the variable domain is derived from the recombination of said human gene segments in a non-human vertebrate (eg, in a mouse or a rat); and/or (ii) the variable domain comprises non-human vertebrate (eg, mouse or rat) AID-pattern mutations and non-human vertebrate (eg, mouse or rat) terminal deoxynucleotidyl transferase (TdT)-pattern mutations.

In another embodiment, the invention provides

An isolated antibody for administration to a Chinese patient, the antibody comprising a human heavy chain, the heavy chain comprising a variable domain that is specific for an antigen and a constant region, wherein the constant region is a human constant region selected from a constant region (eg, an IGHG constant region) present in a Chinese population with a cumulative frequency of at least 5%, and wherein

(i) the variable domain is derived from the recombination of said human gene segments in a non-human vertebrate (eg, in a mouse or a rat); and/or (ii) the variable domain comprises non-human vertebrate (eg, mouse or rat) AID-pattern mutations and non-human vertebrate (eg, mouse or rat) terminal deoxynucleotidyl transferase (TdT)-pattern mutations.

In an example, the constant region is found in the 1000 Genomes database. In an example, the constant region is found in Table 13.

346. The antibody of clause 345 wherein the constant region is a IGHG1a, IGHG2a, IGHG3a, IGHG3b or IGHG4a constant region.

347. The antibody of clause 345 or 346, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the VH gene segment being selected from a VH in Table 13 found in a Chinese population and with a cumulative frequency of at least 5%.

In another embodiment, the invention provides

The antibody of clause 345 or 346, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the VH gene segment being selected from a VH present in a Chinese population with a cumulative frequency of at least 5%.

In an example, the gene segment is found in the 1000 Genomes database. In an example, the gene segment is found in Table 13.

348. The antibody of clause 345, 346 or 347, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the D gene segment being selected from a D in Table 13 found in a Chinese population and with a cumulative frequency of at least 5%.

In another embodiment, the invention provides

The antibody of clause 345, 346 or 347, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the D gene segment being selected from a D present in a Chinese population with a cumulative frequency of at least 5%.

In an example, the gene segment is found in the 1000 Genomes database. In an example, the gene segment is found in Table 13.

349. The antibody of clause 345, 346, 347 or 348 wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the JH gene segment being selected from a JH in Table 13 found in a Chinese population and with a cumulative frequency of at least 5%.

In another embodiment, the invention provides

The antibody of clause 345, 346, 347 or 348 wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the JH gene segment being selected from a JH present in a Chinese population with a cumulative frequency of at least 5%.

In an example, the gene segment is found in the 1000 Genomes database. In an example, the gene segment is found in Table 13.

350. An isolated VH domain identical to a variable domain as recited in any one of clauses 347 to 349, optionally fused at its C-terminus to a polypeptide (eg, an antibody Fc).

In an embodiment, there is provided an isolated VH domain identical to a variable domain as recited in any one of clauses 347 to 349 which is part of a conjugate, conjugated with a label (eg, for imaging in the patient) or a toxin (eg, a radioactive toxic payload, such as for cancer treatment in the patient) or a half-life-extending moiety (eg, PEG of human serum albumin).

351. A pharmaceutical composition comprising the antibody or variable domain of any one of clauses 345 to 350 together with a pharmaceutically-acceptable excipient, diluent or a medicament (eg, a further antigen-specific variable domain, antibody chain or antibody).

352. An isolated antibody for administration to a Chinese patient, the antibody comprising a human heavy chain, the heavy chain comprising a variable domain that is specific for an antigen

and a constant region, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the VH gene segment being selected from a VH in Table 13 found in a Chinese population and with a cumulative frequency of at least 5%; and wherein

(i) the variable domain is derived from the recombination of said human gene segments in a non-human vertebrate (eg, in a mouse or a rat); and/or (ii) the variable domain comprises non-human vertebrate (eg, mouse or rat) AID-pattern mutations and non-human vertebrate (eg, mouse or rat) terminal deoxynucleotidyl transferase (TdT)-pattern mutations.

In another embodiment, the invention provides

An isolated antibody for administration to a Chinese patient, the antibody comprising a human heavy chain, the heavy chain comprising a variable domain that is specific for an antigen and a constant region, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the VH gene segment being selected from a VH present in a Chinese population with a cumulative frequency of at least 5%; and wherein

(i) the variable domain is derived from the recombination of said human gene segments in a non-human vertebrate (eg, in a mouse or a rat); and/or (ii) the variable domain comprises non-human vertebrate (eg, mouse or rat) AID-pattern mutations and non-human vertebrate (eg, mouse or rat) terminal deoxynucleotidyl transferase (TdT)-pattern mutations.

353. The antibody of clause 352, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the D gene segment being selected from a D in Table 13 found in a Chinese population and with a cumulative frequency of at least 5%.

In another embodiment, the invention provides

The antibody of clause 352, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the D gene segment being selected from a D present in a Chinese population with a cumulative frequency of at least 5%.

In an example, the gene segment is found in the 1000 Genomes database. In an example, the gene segment is found in Table 13.

354. The antibody of clause 352 or 353, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the JH gene segment being selected from a JH in Table 13 found in a Chinese population and with a cumulative frequency of at least 5%.

In another embodiment, the invention provides

The antibody of clause 352 or 353, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the JH gene segment being selected from a JH present in a Chinese population with a cumulative frequency of at least 5%.

In an example, the gene segment is found in the 1000 Genomes database. In an example, the gene segment is found in Table 13.

355. An isolated VH domain identical to a variable domain as recited in any one of clauses 352 to 354, optionally fused at its C-terminus to a polypeptide (eg, an antibody Fc).

In an embodiment, there is provided a VH domain identical to a variable domain as recited in any one of clauses 352 to 354 which is part of a conjugate, conjugated with a label (eg, for imaging in the patient) or a toxin (eg, a radioactive toxic payload, such as for cancer treatment in the patient) or a half-life-extending moiety (eg, PEG of human serum albumin).

356. A pharmaceutical composition comprising the antibody or variable domain of any one of clauses 352 to 355 together with a pharmaceutically-acceptable excipient, diluent or a medicament (eg, a further antigen-specific variable domain, antibody chain or antibody).

357. An antibody heavy chain or VH domain (eg, provided as part of an antibody) for therapy and/or prophylaxis of a disease or medical condition in a Chinese patient, wherein the heavy chain is a heavy chain produced by the following steps (or is a copy of such a heavy chain):-

(a) Selection of an antigen-specific antibody heavy chain or VH domain from a non-human vertebrate (eg, a mouse or a rat), wherein the heavy chain or VH domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the VH gene segment being selected from a VH in Table 13 found in a Chinese population and with a cumulative frequency of at least 5%;

(b) Optional humanisation of the heavy chain by combining the variable domain of the heavy chain with a human constant region; or optional humanisation of the selected VH domain by combining with a human constant region.

In another embodiment, the invention provides

An antibody heavy chain or VH domain (eg, provided as part of an antibody) for therapy and/or prophylaxis of a disease or medical condition in a Chinese patient, wherein the heavy chain is a heavy chain produced by the following steps (or is a copy of such a heavy chain):-

(a) Selection of an antigen-specific antibody heavy chain or VH domain from a non-human vertebrate (eg, a mouse or a rat), wherein the heavy chain or VH domain is derived from the

recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the VH gene segment being selected from a VH present in a Chinese population with a cumulative frequency of at least 5%;

(b) Optional humanisation of the heavy chain by combining the variable domain of the heavy chain with a human constant region; or optional humanisation of the selected VH domain by combining with a human constant region.

In an example, the VH gene segment is found in the 1000 Genomes database. In an example, the gene segment is found in Table 13.

358. The antibody heavy chain or VH domain of clause 357, wherein the human constant region is as recited in clause 345 or 346.

359. An antibody heavy chain or VH domain as recited in clause 357 or 358 for use in a medicament for therapy and/or prophylaxis of a disease or medical condition in a Chinese patient.

360. A method of treating and/or preventing a disease or medical condition in a Chinese patient, the method comprising administering to the patient a therapeutically or prophylactically-effective amount of the antibody heavy chain or VH domain as recited in clause 357 or 358.

361. An isolated antibody for administration to a patient of European, East Asian, West African, South Asian or Americas ancestry, the antibody comprising a human heavy chain, the heavy chain comprising a variable domain that is specific for an antigen and a constant region, wherein the constant region is a human constant region selected from a constant region (eg, an IGHG constant region) in Table 13 found in a population of European, East Asian, West African, South Asian or Americas ancestry respectively and with a cumulative frequency of at least 1 or 5%, and wherein

(i) the variable domain is derived from the recombination of said human gene segments in a non-human vertebrate (eg, in a mouse or a rat); or (ii) the variable domain comprises non-human vertebrate (eg, mouse or rat) AID-pattern mutations and non-human vertebrate (eg, mouse or rat) terminal deoxynucleotidyl transferase (TdT)-pattern mutations.

In another embodiment, the invention provides

An isolated antibody for administration to a patient of European, East Asian, West African, South Asian or Americas ancestry, the antibody comprising a human heavy chain, the heavy chain comprising a variable domain that is specific for an antigen and a constant region, wherein the constant region is a human constant region selected from a constant region (eg, an IGHG constant region) present in a population of European, East Asian, West African, South Asian or Americas ancestry respectively with a cumulative frequency of at least 1 or 5%, and wherein

(i) the variable domain is derived from the recombination of said human gene segments in a

non-human vertebrate (eg, in a mouse or a rat); or (ii) the variable domain comprises non-human vertebrate (eg, mouse or rat) AID-pattern mutations and non-human vertebrate (eg, mouse or rat) terminal deoxynucleotidyl transferase (TdT)-pattern mutations.

In an example, the constant region is found in the 1000 Genomes database. In an example, the constant region is found in Table 13.

362. The antibody of clause 361 wherein the constant region is a IGHG1a, IGHG2a, IGHG3a, IGHG3b or IGHG4a constant region and the patient is of European ancestry.

363. The antibody of clause 361 or 362, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the VH gene segment being selected from a VH in Table 13 found in said population and with a cumulative frequency of at least 1 or 5%.

In another embodiment, the invention provides

The antibody of clause 361 or 362, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the VH gene segment being selected from a VH present in a Chinese population with a cumulative frequency of at least 5%.

In an example, the gene segment is found in the 1000 Genomes database. In an example, the gene segment is found in Table 13.

364. The antibody of clause 361, 362 or 363, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the D gene segment being selected from a D in Table 13 found in said population and with a cumulative frequency of at least 1 or 5%.

In another embodiment, the invention provides

The antibody of clause 361, 362 or 363, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the D gene segment being selected from a D present in a Chinese population with a cumulative frequency of at least 5%.

In an example, the gene segment is found in the 1000 Genomes database. In an example, the gene segment is found in Table 13.

365. The antibody of clause 361, 362, 363 or 364 wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the JH gene segment being selected from a JH in Table 13 found in said population and with a cumulative frequency of at least 1 or 5%.

In another embodiment, the invention provides

The antibody of clause 361, 362, 363 or 364 wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the JH gene segment being selected from a JH present in a Chinese population with a cumulative frequency of at least 5%.

In an example, the gene segment is found in the 1000 Genomes database. In an example, the gene segment is found in Table 13.

366. An isolated VH domain identical to a variable domain as recited in any one of clauses 363 to 365, optionally fused at its C-terminus to a polypeptide (eg, an antibody Fc).
367. A pharmaceutical composition comprising the antibody or variable domain of any one of clauses 361 to 366 together with a pharmaceutically-acceptable excipient, diluent or a medicament (eg, a further antigen-specific variable domain, antibody chain or antibody).
368. An isolated antibody for administration to a patient of European, East Asian, West African or Americas ancestry, the antibody comprising a human heavy chain, the heavy chain comprising a variable domain that is specific for an antigen and a constant region, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the VH gene segment being selected from a VH in Table 13 found in a population of European, East Asian, West African, South Asian or Americas ancestry respectively and with a cumulative frequency of at least 1 or 5%; and wherein
- (i) the variable domain is derived from the recombination of said human gene segments in a non-human vertebrate (eg, in a mouse or a rat); or (ii) the variable domain comprises non-human vertebrate (eg, mouse or rat) AID-pattern mutations and non-human vertebrate (eg, mouse or rat) terminal deoxynucleotidyl transferase (TdT)-pattern mutations.

In another embodiment the invention provides:-

An isolated antibody for administration to a patient of European, East Asian, West African or Americas ancestry, the antibody comprising a human heavy chain, the heavy chain comprising a variable domain that is specific for an antigen and a constant region, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the VH gene segment being selected from a VH present in a population of European, East Asian, West African, South Asian or Americas ancestry respectively with a cumulative frequency of at least 1 or 5%; and wherein

- (i) the variable domain is derived from the recombination of said human gene segments in a non-human vertebrate (eg, in a mouse or a rat); or (ii) the variable domain comprises non-human vertebrate (eg, mouse or rat) AID-pattern mutations and non-human vertebrate (eg,



mouse or rat) terminal deoxynucleotidyl transferase (TdT)-pattern mutations.

In an example, the VH gene segment is found in the 1000 Genomes database. In an example, the gene segment is found in Table 13.

369. The antibody of clause 368, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the D gene segment being selected from a D in Table 13 found in said population and with a cumulative frequency of at least 1 or 5%.

In another example there is provided

The antibody of clause 368, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the D gene segment being selected from a D present in said population with a cumulative frequency of at least 1 or 5%.

In an example, the D gene segment is found in the 1000 Genomes database. In an example, the gene segment is found in Table 13.

370. The antibody of clause 368 or 369, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the JH gene segment being selected from a JH in Table 13 found in said population and with a cumulative frequency of at least 1 or 5%.

In another example there is provided

The antibody of clause 368 or 369, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the JH gene segment being selected from a JH present in said population and with a cumulative frequency of at least 1 or 5%.

In an example, the JH gene segment is found in the 1000 Genomes database. In an example, the gene segment is found in Table 13.

371. An isolated VH domain identical to a variable domain as recited in any one of clauses 368 to 370, optionally fused at its C-terminus to a polypeptide (eg, an antibody Fc).

372. A pharmaceutical composition comprising the antibody or variable domain of any one of clauses 368 to 371 together with a pharmaceutically-acceptable excipient, diluent or a medicament (eg, a further antigen-specific variable domain, antibody chain or antibody).

373. An antibody heavy chain or VH domain (eg, provided as part of an antibody) for therapy and/or prophylaxis of a disease or medical condition in a patient of European, East Asian, West African, South Asian or Americas ancestry, wherein the heavy chain is a heavy chain produced by

the following steps (or is a copy of such a heavy chain):-

(a) Selection of an antigen-specific antibody heavy chain or VH domain from a non-human vertebrate (eg, a mouse or a rat), wherein the heavy chain or VH domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the VH gene segment being selected from a VH in Table 13 found in said population and with a cumulative frequency of at least 1 or 5%;

(b) Optional humanisation of the heavy chain by combining the variable domain of the heavy chain with a human constant region; or optional humanisation of the selected VH domain by combining with a human constant region.

In another embodiment, there is provided:-

An antibody heavy chain or VH domain (eg, provided as part of an antibody) for therapy and/or prophylaxis of a disease or medical condition in a patient of European, East Asian, West African, South Asian or Americas ancestry, wherein the heavy chain is a heavy chain produced by the following steps (or is a copy of such a heavy chain):-

(a) Selection of an antigen-specific antibody heavy chain or VH domain from a non-human vertebrate (eg, a mouse or a rat), wherein the heavy chain or VH domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the VH gene segment being selected from a VH present in said population with a cumulative frequency of at least 1 or 5%;

(b) Optional humanisation of the heavy chain by combining the variable domain of the heavy chain with a human constant region; or optional humanisation of the selected VH domain by combining with a human constant region.

In an example, the VH gene segment is found in the 1000 Genomes database. In an example, the gene segment is found in Table 13.

374. The antibody heavy chain or VH domain of clause 373, wherein the human constant region is as recited in clause 361 or 362.

375. An antibody heavy chain or VH domain as recited in clause 373 or 374 for use in a medicament for therapy and/or prophylaxis of a disease or medical condition in a patient of said ancestry.

376. A method of treating and/or preventing a disease or medical condition in a patient of European, East Asian, West African, South Asian or Americas ancestry, the method comprising administering to the patient a therapeutically or prophylactically-effective amount of the antibody heavy chain or VH domain as recited in clause 373 or 374.

In embodiments herein, a Chinese patient can be a Han Chinese patient.

In embodiments herein, a patient of European ancestry can be a patient of Northern or Western European ancestry, Italian ancestry, British or Scottish ancestry, Finnish ancestry or Iberian ancestry.

In embodiments herein, a patient of East Asian ancestry can be a patient of Han Chinese ancestry, Japanese ancestry Chinese Dai ancestry, Vietnamese ancestry or Kinh ancestry.

In embodiments herein, a patient of West African ancestry can be a patient of Yoruba ancestry, Luhya ancestry, Gambian ancestry or Malawian ancestry.

In embodiments herein, a patient of Americas ancestry can be a patient of African American ancestry, African Caribbean ancestry, Mexican ancestry, Puerto Rican ancestry, Colombian ancestry or Peruvian ancestry.

In embodiments herein, a patient of South Asian ancestry can be a patient of Ahom ancestry, Kayadtha ancestry, Reddy ancestry, Maratha ancestry, or Punjabi ancestry.

In an example of any aspect, the cumulative frequency is at least 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25, 30, 35, 40, 45, 50, 55, 60, 65, 70, 75, 80, 85, 90 or 95%.

It will be understood that particular 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.

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

The term "or combinations thereof" as used herein refers to all permutations and combinations of the listed items preceding the term. For example, "A, B, C, or combinations thereof" is intended to include at least one of: A, B, C, AB, AC, BC, or ABC, and if order is important in a particular context, also BA, CA, CB, CBA, BCA, ACB, BAC, or CAB. Continuing with this example, expressly included are combinations that contain repeats of one or more item or term, such as BB, AAA, MB, BBC, AAABCCCC, CBBAAA, CABABB, and so forth. The skilled artisan will understand that typically there is no limit on the number of items or terms in any combination, unless otherwise apparent from the context.

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

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.

The present invention is described in more detail in the following non limiting prophetic Examples.

## **EXAMPLES**

### **EXAMPLE 1**

“Recombineered BAC Vectors to add Polymorphic V-regions to the Mouse Genome”

Figure 1 through 3 depict recombineering methods (see references above) that can be used to introduce polymorphic V-gene regions into genomic DNA. In one embodiment, a genomic fragment from the human heavy chain region is inserted into a bacterial artificial chromosome (BAC) vector by standard techniques. Preferably, such a BAC, which can range in size from 20-kb to 200-kb or more, can be isolated from libraries of BACs by standard techniques including sequence searches of commercially available libraries or by hybridization to bacterial colonies containing BACs to identify those with a BAC of interest.

A BAC is chosen that has several VH gene segments; in Figure 1, these are generically identified as VH[a] through VH[z] for example. One skilled in the art will readily identify appropriate genomic fragments, for example, an approximately 120-kb fragment from human VH5-78 through VH1-68 which includes 5 endogenous active VH gene segments and 7 VH psuedogenes. Using recombineering techniques, the endogenous VH gene segments can be replaced by polymorphic VH or VL gene segments. In this example, two steps are required. The first step replaces the V-region coding exon of an endogenous VH gene segment with a positive-negative selection operon, in this example, an operon encoding an ampicillin resistance gene (*Amp*) and a streptomycin-sensitizing ribosomal protein (*rpsL*). Certain strains of bacteria can be selected for the absence of the *rpsL* gene by resistance to streptomycin. Short stretches of DNA homologous to sequences flanking the

endogenous VH gene exon are placed 5' and 3' of the *rpsL*-Amp operon. In the presence of appropriate recombination factors per standard recombineering techniques (see references above) recombination between the operon fragment and the BAC will result in replacement of the endogenous VH gene exon with the operon (Figure 1a) which are selected by resistance to ampicillin. The second step uses the same homologous sequences in order to replace the inserted operon with a desired polymorphic VH gene segment. In this example, a human VH1-69 gene is inserted (Figure 1b and 1c). In particular the \*02 variant of VH1-69 is used [ref IMGT and Figure 5]. Successful integrations of the polymorphic VH gene segment are selected in bacteria that become resistant to streptomycin due to the loss of the operon, specifically the *rpsL* portion.

In this example, the two step process as described can be repeated for each of the endogenous VH gene segments or for as many endogenous gene segments that one wishes to replace with polymorphic V gene segments (Figure 1d).

As is apparent, any polymorphic V gene segment can be inserted in this manner and any endogenous V gene segment can act as a target, including pseudogenes. V gene segments in each of the heavy chain and two light chain loci can be replaced using this technique with appropriate genomic fragments available as BAC inserts.

Figure 2 depicts another method for creating a genomic fragment encoding polymorphic V gene segments. In this example, polymorphic V gene segments are inserted into a region of genomic DNA devoid of other genes, control elements or other functions. Such 'desert' regions can be selected based on sequence analysis and corresponding DNA fragments cloned into BACs or identified in existing BAC libraries. Starting with such a genomic fragment, recombineering techniques can be used to insert polymorphic V gene segments at intervals of, for example, 10-kb. In this example, a 150-kb genomic fragment might accommodate insertion of up to 15 polymorphic V gene segments. Insertion of the segments is a two-step process. The first recombineering step inserts the *rpsL*-Amp operon at a specific site. Sequences homologous to a specific site are used to flank the operon. These are used by the recombineering system to insert the element specifically into the BAC genomic fragment and positive events are selected by resistance to ampicillin (Figure 2a). The second step replaces the operon in the genomic fragment with a polymorphic V gene segment by a similar recombineering step using the same sequence homology (Figure 2b). In this example, both

exons and promoter element of a polymorphic VH gene segment are inserted, resulting in replacement of the *rpsL-Amp* operon and therefore resistance to streptomycin (Figure 2c).

The two step technique for inserting polymorphic V gene segments into a specific site on the genomic fragment can be repeated multiple times resulting in a BAC genomic fragment with several polymorphic gene segments, including their promoter elements. It is apparent that the examples shown in Figures 1 and 2 can be combined wherein the technique for insertion can be used to add *extra* polymorphic V gene segments to a BAC genomic fragment as depicted in Figure 1. One might choose to add these extra segments to an IG genomic fragment since such a fragment would be more amenable to proper IG gene expression once inserted into a non-human mammal's genome. It is known that a genomic fragment can have elements such as enhancers or elements that contribute to certain chromatin conformations, both important in wild-type gene expression.

Figure 3 depicts an additional method to create genomic fragments with polymorphic V gene segments. This method depends upon the efficiency with which short (around 50 to 150 bases, preferably 100 bases) single stranded DNA fragments recombine with a homologous sequence using recombineering (Nat Rev Genet. 2001 Oct;2(10):769-79; Recombineering: a powerful new tool for mouse functional genomics; Copeland NG, Jenkins NA, Court DL). The recombinases used in recombineering preferentially bind and use such short single-stranded fragments of DNA as a substrate for initiating homologous recombination. The efficiency can be as high as 10<sup>-2</sup>, that is, a positive event can be found in approximately 100 randomly picked (not selected) clones resulting from recombineering. A positive event in this example occurring when one or more single nucleotide changes introduced into the single-stranded fragment get transferred to the BAC insert containing V gene segments and surrounding genomic DNA, said nucleotide change or changes occurring at a homologous sequence on the BAC.

Polymorphic V gene segments can differ from endogenous V gene segments by only 1 or 2, or up to 10 or 15 nucleotide changes, for example. An example of such nucleotide polymorphisms are depicted in Figure 5. Short single stranded regions that encompass the polymorphic nucleotide changes can be chemically synthesized using standard techniques. The resulting single stranded DNA fragments are introduced into bacteria and *via* recombineering techniques approximately 1 in 100 BAC fragments will have incorporated the polymorphic nucleotides *via* homologous incorporation of

the single stranded fragment (Figure 3a). BACs with the desired nucleotide change can be identified by screening for example several hundred individual clones by polymerase chain reaction (PCR) amplification and sequencing, both by standard techniques. In the example, two nucleotide changes will convert a VH1-69\*01 gene segment into a VH1-69\*02 gene segment (Figure 3b).

It is clear that this process can be repeated for multiple endogenous V gene segments contained on a single BAC genomic fragment. In addition, the techniques depicted in Figure 2 can be used to add additional polymorphic V gene segments by insertion into regions between existing V gene segments. As would be evident to one skilled in the art, a combination of these techniques can be used to create numerous variations of both polymorphic and endogenous human V gene segments. And it would be evident that several different genomic fragments with engineered polymorphic V gene segments and endogenous human V gene segments can be combined to create even more variations.

## EXAMPLE 2

### “Adding Polymorphic V-regions to the Genome using SRMCE of Modified BACs”

Modified BACs with polymorphic V gene segments created using the methods described in Example 1 can be used to alter the genome of non-human mammals. These alterations can result in an intact IG locus in which normal immunoglobulin region recombination results in VDJ or VJ combinations which includes the human V gene segments. An example of how such an animal can be created is by altering the genome of, for example, mouse embryonic stem (ES) cells using the strategy outlined in Figure 4.

One technique to integrate modified BACs with polymorphic V gene segments into a genome is sequential recombinase mediated cassette exchange (SRMCE). The technique is described in WO2011004192 (Genome Research Limited), which is incorporated here in its entirety by reference.



SRMCE provides for a locus modified with a 'landing pad' inserted at a specific location. This insertion can either be *de novo* via homologous recombination or as a consequence of a previous BAC insertion. In this example, the landing pad is inserted in the mouse IGH locus between the most 3' J gene segment and the C $\mu$  gene segment and a previous BAC insertion *via* SRMCE techniques have resulted in the addition of 5 human V gene segments and 2 V region pseudogenes. The landing pad has elements as shown in Figure 4 that will allow the selection of correct insertion of a second targeting BAC fragment. The specificity of this insertion is provided by cre recombinase-mediated exchange between permissive *lox* sites. A *lox* site is permissive for recombination only with a compatible *lox* site. In this example, the *loxP* site will only recombine with *loxP* and *lox2272* will only recombine with *lox2272*. This provides directionality to the insertion of the BAC fragment as depicted in Figure 4b and 4c.

ES cell clones with correct insertions are selected from a pool of clones without insertions or with non-productive insertions by resistance to puromycin. Resistance to puromycin results from the juxtaposition of an active promoter element, PGK, with the *puroTK* coding region. Correct insertions are verified by standard techniques including PCR of junctions, PCR of internal elements, Southern blotting, comparative genomic hybridization (CGH), sequencing and *etc.* In the example, correct *lox2272-lox2272* and *loxP-loxP* recombination also results in two intact sets of *piggyBac* elements that did not exist prior to insertion. An intact *piggyBac* element is comprised of a set of inverted repeats which are depicted in the figure by "PB5'" and "PB3'". An appropriated oriented set of *piggyBac* elements are the substrate of *piggyBac* transposase which can catalyse recombination between the elements, resulting in deletion of intervening sequences as well as both elements. The DNA remaining after a *piggyBac* transposition is left intact and is lacking any remnant of the *piggyBac* element. In the example, ES cell clones with successful *piggyBac* transposition are selected by loss of the active *puroTK* element which renders the cells resistant to the drug FIAU (Figure 4c and 4d).

The final product of the SRMCE method in this example is a IGH locus with several polymorphic V gene segments inserted along with a set of endogenous unmodified VH gene segments between sequences of the mouse genome on the 5' side and the mouse IGH constant region gene segments on the 3' side. The polymorphic V gene segments are positioned such that they can participate in the recombination events associated with B cell maturation yielding VDJ gene segments. These gene

segments can then be transcribed and spliced to the mouse constant region. Translation of these transcripts will result in the production of an antibody heavy chain encoded by the polymorphic V gene segment, a human DH gene segment, a human JH gene segment and a mouse constant heavy chain gene segment.

As is well known to those skilled in the art, an ES cell clone can be used to create a line of genetically modified mice *via* injection of said cells into a mouse blastocyst embryo, transferring the injected embryo to a suitable recipient and breeding the chimeric offspring that result. The modified gene locus can be propagated through breeding and made either heterozygous or homozygous depending on the genetic cross.

It is evident from the structure of the IGH locus provided in this example and by knowledge of the mechanisms involved in B cell receptor (BCR) and antibody gene rearrangements that a large set of different combinations of polymorphic V gene segments with various DH and JH gene segments will result and these can contribute to a large repertoire of functional antibody genes in a population of B cells in genetically modified animals. In this example, several different human VH1-69 polymorphs are incorporated to provide superhuman VH diversity. This particular VH gene segment is known to be prevalent in antibodies that bind infectious disease pathogens (such as influenza virus) and therefore the antibody repertoire of a mouse with the genetic modification of this example would be expected to produce antibodies with a bias in favour of those that bind infectious disease pathogens. The repertoire, in other words, would have a larger subset of antibodies with superior affinities for pathogen antigens. Examples of such pathogens include influenza virus, hepatitis C virus (HCV) and human immunodeficiency virus-1 (HIV-1) (see also table above).

### EXAMPLE 3

“Alignment of 13 VH1-69 Alleles”

Building a more diverse antibody repertoire by incorporating additional V gene segment polymorphs requires availability of polymorphic variants of V gene segments. One source of such variants include sequence databases. In this example, 13 distinct variants of the VH1-69 gene segment are provided.

These variant sequences and comparisons are drawn from the “IMmunoGeneTics” IMGT Information System ([www.imgt.com](http://www.imgt.com)) database. Figure 5 is a diagram of the alignment of variants \*02 through \*13 with the \*01 variant. The VH1-69\*01 nucleotide and amino acid sequence is provided at the top of the figure. Where the remaining variants are identical to the \*01 variant sequence a dash is inserted below the sequence. Nucleotide differences are noted alongside the appropriate variant and if the sequence change results in a protein coding change, the amino acid change is indicated above the triplet.

Figure 5 depicts between 1 and 4 amino acid changes for each variant in comparison to the \*01 variant. All of the amino acid changes occur in the part of the heavy chain protein encoding the complementarity determining regions (CDRs). These regions are responsible for antigen specificity and the affinity of the antibody for the antigen. It is evident that providing additional polymorphic CDRs in a repertoire of antibodies will increase the likelihood of there being an antibody with superior binding characteristics for various antigens. In several reports, it has been observed that the VH1-69-encoded variable region of the heavy chain is often found in antibodies that bind influenza virus, HCV and HIV-1 antigens (see table above). Therefore incorporating the polymorphic V gene segments of this example into a transgenic animal model using the methods of Examples 1 and 2 would likely result in an antibody repertoire in said transgenic animal with more antibodies that bind to antigens associated with these and other pathogens. And as is known in the art, a larger repertoire increases the probability of finding monoclonal antibodies using, for example, hybridoma technology, that bind with high affinity and specificity to a desired antigen.

This disclosure therefore describes in these examples a transgenic mouse model which can be immunized with pathogen or other antigens. Plasma B cells from such an immunized mouse can be used to make a hybridoma library that can be screened for production of antibodies that bind the pathogen antigens. This library will be superior to libraries from traditional transgenic mice for finding such antibodies given the addition of polymorphic VH1-69 gene segments to the IGH locus in said transgenic mouse.

These examples are not limiting to the human polymorphic V gene segments that can be chosen or to the methods used to introduce them into an animal model. The method can be used to construct a transgenic locus with immunoglobulin D and/or J segments. The V, D, J segments can be from a plurality of human sources (optionally more than one human ethnic population).

**EXAMPLE 4****Human IgH JH Gene Variants Selected from the 1000 Genomes Database**

Data is presented for human JH2, 5 and 6 variants. In Tables 10A, 11A and 12A samples from humans from various populations are listed where the sequence analysis of the inventors has revealed the presence of polymorphisms in one or both IgH JH alleles. The population codes are explained in Table 8 above. The polymorphisms are nucleotide variants from JH2, 5 and 6 reference sequences (SEQ ID NOs: 1, 2 and 3 respectively; see below). All references are sequences taken from the Ensembl database ([www.ensembl.org](http://www.ensembl.org)). The JH5 reference is human IgH J5-001 disclosed in that database. The JH6 reference is human IgH J6-001 disclosed in that database. The JH2 reference is human IgH J2-001 disclosed in that database.

The reference nucleotide and encoded amino acid sequences are shown on the next page.

Alignments with encoded amino acid sequences are also provided, including the corresponding position numbers on human chromosome 14.

Variant Frequencies are shown in Tables 10A, 11A and 12A and these relate to the frequency of the variants in the 1000 Genomes Database (release current at October 2011).

Tables 10B, 11B and 12B show the non-synonymous nucleotide polymorphisms in the human JH variants, as sorted by the present inventors from the 1000 Genomes database. Position numbers corresponding to nucleotide positions on human chromosome 14 are shown for variant positions (chromosome 14 being the chromosome bearing the IgH locus in humans). Thus, for example, the first entry in Table 11B is "14:106330027:A/C" which refers to a position in a variant JH5 sequence wherein the position corresponds to position 106,330,027 on human chromosome 14, such position being A (adenine) in the reference sequence. The "C" indicates that the present inventors observed a mutation to cytosine at this position in the variants found in the 1000 Genomes database. This change leads to a change at the amino acid level of the encoded sequence (ie, a "non-synonymous" change), in this case a change from a serine (found in the reference) to an alanine in the variant.

**Example 5:**

### **Human Antibody Gene Segment Variant Identification & Population Analysis**

The genomic coding region coordinates for each target gene for variant analysis were identified from the Ensembl WWW site ([www.ensembl.org](http://www.ensembl.org)) using coordinates from the GRCh.p8 Human Genome assembly ([www.ncbi.nlm.nih.gov/projects/genome/assembly/grc](http://www.ncbi.nlm.nih.gov/projects/genome/assembly/grc)). Using the collected gene location coordinates, variant data was extracted from the public ftp site of the 1000 Genomes Project using the Perl 'Variant Pattern Finder' (VPF - [www.1000genomes.org/variation-pattern-finder-api-documentation](http://www.1000genomes.org/variation-pattern-finder-api-documentation)).

Data extracted by VPF was post processed using software to extract all non-synonymous (NSS) variants with their associated genotype calls. Genotypes calls were assembled to form unique haplotypes, representing groups of NSS variants associated with 1000 Genome population groups and frequency of occurrence within those populations.

The output of the analysis results in tables such as in Table 13. The main body of the table describes each haplotype in turn giving a unique ID for that gene (in the range a-z,aa-zz), the population frequencies and occurrence in individuals and unique population groups; one or more subsequent columns describe the DNA base calls at each location that form the haplotype giving both the base from the reference sequence or the variant base call.

Table 13 was constructed in this manner. The table can be read as follows:

The first four columns (left to right) consist of (1) the haplotype ID letter ('ref' indicates reference - the DNA base call at each genomic location from the GRCh37 Human Reference Assembly) (2) the observed cumulative frequency of the haplotype among the different populations (3) the number of individuals in which a specific haplotype was observed (4) the number of unique population groups that the identified individuals belong to (the actual population group identifiers are displayed as a string of ID's in the most right hand column for each haplotype. For example haplotype 'a' has a population ID string of '3,4,9,13').

The populations are numbered as follows (population labels being according to 1000 Genomes Project nomenclature)

- 1= ASW;
- 2= CEU;
- 3=CHB;
- 4=CHS;
- 5=CLM;
- 6=FIN;
- 7=GBR;
- 8=IBS;

9=JPT;  
 10=LWK;  
 11=MXL;  
 12=PUR;  
 13=TSI;  
 14=YRI.

Subsequent columns detail a single point variant and have the following format (top to bottom) (1) the human genomic location of the variant (format [chromosome number]: [location] e.g. '14:106204113'); (2) The identifier for the point variant as defined in DbSNP ([www.ncbi.nlm.nih.gov/projects/SNP/](http://www.ncbi.nlm.nih.gov/projects/SNP/)); (3) One or additional rows show the amino acid change as result of the variant for a specific transcript (denoted by the Ensembl transcript ID in the most right-hand column for each row), the format is the amino acid in the reference sequence followed by '->' and the amino acid caused by the substitution of the variant in the reference sequence (e.g. 'Gly->Arg' means a that the translated reference sequence would result in a glycine at that location, whereas the substitution of the identified variant would result in translated protein containing arginine) using the IUPAC three letter amino acid codes (<http://pac.iupac.org/publications/pac/pdf/1972/pdf/3104x0639.pdf>). Subsequent rows (one per haplotype) show the DNA base at each location, bases matching the reference sequence are shown in black on white back ground, bases varying from the reference are shown as white text on a black background.

The most right-hand column contains the Ensembl transcript ID's (e.g. 'ENST00000390542') for each of the gene transcript and relates to the amino acid changes to the left of this column. Because the transcripts are differing lengths each variant position may or may not have an associated amino acid change at the that position.

### **Example 6:**

#### **Transgenic Mice, B-cells, Hybridomas, Antibodies & Heavy Chains Based on Human JH6\*02**

A functional human gene segment repertoire (from V<sub>H</sub>2-26 to J<sub>H</sub>6, see the IMGT database for the structure of the human IgH locus;

<http://www.imgt.org/IMGTrepertoire/index.php?section=LocusGenes&repertoire=locus&species=human&group=IGK>) was sectored by the inventors to produce two different transgenic heavy chain alleles (denoted S2F and S3F) and corresponding mice. The transgenic alleles were expressed in the mice and the heavy chain repertoires were assessed at the RNA transcript level. Deep sequence

analysis was carried out using Bioinformatics methods to assess V, D and JH gene usage, including in variable domain sequences having a HCDR3 length of at least 20 amino acids. Endogenous, mouse variable region gene segments were inactivated by inversion (as per the method described in WO2011004192, this disclosure being incorporated herein by reference).

#### Sequencing of Human Donor DNA Samples: Identification of Conserved JH6\*02 Variant

DNA samples from 9 anonymised consenting human donors were obtained by taking cheek swabs.

The samples were processed and the DNA Samples were extracted follow the protocol of QIAamp DNA Mini Kit (Cat.No.51304, Qiagen).

PCR reactions were set up to amplify the JH6 region and PCR products were sequenced (PCR Oligos sequence: Fwd. 5'-AGGCCAGCAGAGGGTTCATG-3' (SEQ ID NO: 444), Rev. 5'-GGCTCCAGATCCTCAAGGCAC-3' (SEQ ID NO: 445)).

Sequence analysis was carried out by comparing to the JH6 reference sequence from IMGT annotated database (<http://www.imgt.org/>), and this identified that all 9 donor genomes contained the human JH6\*02 variant, with this variant being in the homozygous state in 7 out of the 9 donors. The inventors also consulted the genomic sequences publicly available for Jim Watson and Craig Venter at Ensembl human genome database [<http://www.ensembl.org/>]. These too contained the human JH6\*02 variant. This confirmed to the inventors that human JH6\*02 is a common, conserved variant in humans, and thus a good candidate for construction of a transgenic IgH locus as per the invention

#### Identification of Suitable Human DNA Sequence BACs

A series of human bacterial artificial chromosome (BAC) clones were identified from Ensemble (<http://www.ensembl.org/index.html>) or UCSC (<http://genome.ucsc.edu/>) human database searches based on gene name (IGH) or location (chromosome 14: 106026574-107346185). Seven human RP11 BAC clones (see an extract of the UCSC databse in Figure 10, identified BACs being circled) were selected, RP11-1065N8 BAC carrying human JH6\*02. In total, the following BACs were

identified as sources of human IgH locus DNA: RP11-1065N8, RP11-659B19, RP11-141I7, RP-112H5, RP11-101G24, RP11-12F16 and RP11-47P23.

With a similar approach, different BAC clones (eg, different RP11 clone IDs or different sources from RP11) or genetically engineered BACs can be selected for insertion into the mouse IGH locus to provide different sets of human repertoires in the transgenic mouse.

#### Construction of Transgenic IgH Loci

Insertion of human heavy gene segments from a 1st IGH BAC (RP11-1065N8) into the IGH locus of mouse AB2.1 ES cells (Baylor College of Medicine) was performed to create a heavy chain allele denoted the S1 allele. The inserted human sequence corresponds to the sequence of human chromosome 14 from position 106494908 to position 106328951 and comprises functional heavy gene segments V<sub>H</sub>2-5, V<sub>H</sub>7-4-1, V<sub>H</sub>4-4, V<sub>H</sub>1-3, V<sub>H</sub>1-2, V<sub>H</sub>6-1, D1-1, D2-2, D3-9, D3-10, D4-11, D5-12, D6-13, D1-14, D2-15, D3-16, D4-17, D5-18, D6-19, D1-20, D2-21, D3-22, D4-23, D5-24, D6-25, D1-26, D7-27, J<sub>H</sub>1, J<sub>H</sub>2, J<sub>H</sub>3, J<sub>H</sub>4, J<sub>H</sub>5 and J<sub>H</sub>6 (in 5' to 3' order), wherein the J<sub>H</sub>6 was chosen to be the human J<sub>H</sub>6\*02 variant. The insertion was made between positions 114666435 and 114666436 on mouse chromosome 12, which is upstream of the mouse C<sub>μ</sub> region. The mouse V<sub>H</sub>, D and J<sub>H</sub> gene segments were retained in the locus, immediately upstream of (5' of) the inserted human heavy chain DNA.

A second allele, S2 was constructed in which more human functional V<sub>H</sub> gene segments were inserted upstream (5') of the 5'-most V<sub>H</sub> inserted in the S1 allele by the sequential insertion of human DNA from a second BAC (BAC2). The inserted human sequence from BAC2 corresponds to the sequence of human chromosome 14 from position 106601551 to position 106494909 and comprises functional heavy chain gene segments V<sub>H</sub>3-13, V<sub>H</sub>3-11, V<sub>H</sub>3-9, V<sub>H</sub>1-8, V<sub>H</sub>3-7. The mouse V<sub>H</sub>, D and J<sub>H</sub> gene segments were retained in the locus, immediately upstream of (5' of) the inserted human heavy chain DNA. In a subsequent step, these were inverted to inactivate them, thereby producing S2F mice in which only the human heavy chain variable region gene segments are active.

A third allele, S3 was constructed in which more human functional V<sub>H</sub> gene segments were inserted upstream (5') of the 5'-most V<sub>H</sub> inserted in the S2 allele by the sequential insertion of human DNA



from a third BAC (BAC3). The inserted sequence corresponds to the sequence of human chromosome 14 from position 106759988 to position 106609301, and comprises functional heavy chain gene segments, V<sub>H</sub>2-26, V<sub>H</sub>1-24, V<sub>H</sub>3-23, V<sub>H</sub>3-21, V<sub>H</sub>3-20, V<sub>H</sub>1-18, and V<sub>H</sub>3-15. The mouse V<sub>H</sub>, D and J<sub>H</sub> gene segments were retained in the locus, immediately upstream of (5' of) the inserted human heavy chain DNA. In a subsequent step, these were inverted to inactivate them, thereby producing S3F mice in which only the human heavy chain variable region gene segments are active.

Mice bearing either the S2F or S3F insertion into an endogenous heavy chain locus were generated from the ES cells using standard procedures. The other endogenous heavy chain locus was inactivated in the mice by insertion of an inactivating sequence comprising neo<sup>R</sup> into the mouse J<sub>H</sub>-C<sub>μ</sub> intron (to produce the "HA" allele).

#### Immunisation procedure

Transgenic mice of the S2F or S3F genotype were primed with 20-40ug recombinant proteins obtained commercially or produced in house with Antigen 1 (OVA (Sigma A7641); Antigen 2 (a human infectious disease pathogen antigen) and Antigen 3 (a human antigen) via the ip route in complete Freund's adjuvant (Sigma F 5881) and 10ug/animal CpG (CpG oligo; Invivogen, San Diego, California, USA) and then boosted twice in about two weekly intervals with about half the amount of antigen in incomplete Freund's adjuvant (Sigma F 5506) and 10ug/animal CpG. Final boosts were administered two weeks later iv without any adjuvant and contained 5-10 ug protein in PBS.

#### Hybridoma fusion procedure

Spleens were taken 3 days after the final boost and spleenocytes were treated with CpG (25 μm final concentration) for and left until the following day. Cells were then fused with SP0/2 Ag14 myeloma cells (HPA Cultures Cat No 85072401) using a BTX ECM2001 electrofusion instrument. Fused cells were left to recover for 20 minutes then seeded in a T75 flask until next morning. Then the cells were spun down and plated out by dilution series on 96-well culture plates and left for about 10 days before screening. Media was changed 1-3 times during this period.

#### Screening

### Screening

Culture supernatants of the hybridoma wells above were screened using homogenous time resolved fluorescence assay (htfrf) using Europium cryptate labelled anti-mouse IgG (Cisbio anti-mouse Ig Europium Cryptate) and a biotin tagged target antigen with a commercially available streptavidin conjugated donor (Cisbio; streptavidin conjugated D2) or by IgG-specific 384 well ELISA. Positive wells identified by htfrf were scaled to 24-well plates or immediately counterscreened using an IgG-specific detection ELISA method. Positives identified by primary ELISA screen were immediately expanded to 24-well plates. Once cultures were expanded to 24-well stage and reached confluency, supernatants were re-tested using htfrf or IgG-specific ELISA to confirm binding to target antigen. Supernatant of such confirmed cultures were then also analysed by surface plasmon resonance using a BioRad ProteOn XPR36 instrument. For this, antibody expressed in the hybridoma cultures was captured on a biosensor GLM chip (BioRad 176-512) which had an anti-mouse IgG (GE Healthcare BR-1008-38) covalently coupled the biosensor chip surface. The antigen was then used as the analyte and passed over the captured hybridoma antibody surface. For Antigen 2 and Antigen 3, concentrations of 256nM, 64nM, 16nM, 4nM and 1nM were typically used, for Antigen 1, concentrations of 1028nM, 256nM, 64nM, 16nM and 4nM were typically used, binding curves were double referenced using a 0nM injection (i.e. buffer alone). Kinetics and overall affinities were determined using the 1:1 model inherent to the BioRad ProteOn XPR36 analysis software.

Any clones with confirmed binding activity were used for preparing total RNA and followed by PCR to recover the heavy chain variable region sequences. Standard 5'-RACE was carried out to analyse RNA transcripts from the transgenic heavy chain loci in the S2F and S3F mice. Additionally, deep sequence analysis of almost 2000 sequences produced by the mice was carried out.

### Bionformatics Analysis

Sequences for analysis were obtained from two different methods:

- The first is from RNA extracted from the spleen: first cDNA strand was synthesized using an oligo based on the Cmu region of the mouse IGH locus as a PCR template. PCR was performed using this oligo with an oligo dT-anchor primer. Then PCR product was cloned into pDrive vector (Qiagen) and then sequenced.

- The second is from hybridomas generated through electro-fusion: total RNA was extracted from hybridoma lines of interest using standard Trizol methods and frozen at -80 °C for long term storage. cDNA was generated from 100ng total RNA using standard Superscript III reverse transcriptase and a gene-specific reverse primer binding to all mouse IgG isotypes for heavy chain and a mouse kappa constant region primer for the light chain amplification. 2-3 ul of cDNA were then used as template in a PCR reaction using Pfu DNA polymerase and a panel of degenerate forward primers annealing to the leader sequence of the human immunoglobulin variable domain as well as one mouse pan-IgG reverse primer. PCR products were run out of a 1% agarose gel and bands of approximately 350-450 basepairs extracted and purified. DNA was then sequenced.

The sequences from the first method can either be from IgM from Naïve mice or IgG from immunised mice. The samples from the second method are all from IgG from immunised mice, and specific to the immunising antigen. Almost 2000 sequences were analysed.

The sequences were obtained as a pair of forward and reverse reads. These were first trimmed to remove low-quality base calls from the ends of the reads (trimmed from both ends until a 19 nucleotide window had an average quality score of 25 or more). The reads were combined together by taking the reverse complement of the reverse read, and aligning it against the forward read. The alignment scoring was 5 for a match, -4 for a mismatch, a gap open penalty of 10 and a gap extension penalty of 1. A consensus sequence was then produced by stepping through the alignment and comparing bases. When there was a disagreement the base with the highest quality value from sequencing was used.

The BLAST+ (Basic Local Alignment Search Tool) (Camacho C., Coulouris G., Avagyan V., Ma N., Papadopoulos J., Bealer K., & Madden T.L. (2008) "BLAST+: architecture and applications." *BMC Bioinformatics* 10:421 <http://www.ncbi.nlm.nih.gov/pubmed/20003500>) program 'blastn' was then used to find the germline J and V segments used in each sequence. A wordsize of 30 was used for V matching, and 15 for J matching. The database searched against was constructed from the NGS sequencing of the BACs which were used to generate the Kymouse.

If a sequence matched both a V and a J segment, the sequence between the two was then compared to a database of germline D segments in the mouse using ‘blastn’ with a wordsize of 4 and the options ‘blastn-short’ and ‘ungapped’. This was used to assign a D segment, if possible. The CDR3 was identified by searching for the conserved “TATTACTGT” sequence in the V segment, and the “CTGGGG” in the J segment. If these motifs were not found, then up to 4 mismatches were allowed. The IMGT definition of CDR3 was used, so the CDR3 length is calculated from after the “TGT” in the V to before the “TGG” in the J. Sequences with an out of frame junction (those which do not have a CDR3 nucleotide length divisible by 3) or which contained a stop codon (“TAA”, “TAG” or “TGA”) were excluded.

The identity of the matching V, J and D segments as well as the CDR3 length from this assignment were then saved as a table for downstream analysis. The ratio of IGHJ6\*02 used increased from the naïve to immunised mice, as well as being enriched in the sub-population of sequences with a long HCDR3 (defined as consisting of 20 or more amino acids):

	All		HCDR3>20		% HCDR3>20
	JH6*02%	Total Count	JH6*02%	Total Count	
Naïve	22.31%	1340	91.11%	45	3.36%
Immunised	37.50%	256	66.67%	9	3.52%
Hybridoma	36.13%	119	63.64%	11	9.24%

This shows that the JH6\*02 gene segment is selected for by immunisation, as the proportion of JH6\*02 usage increases after immunisation. JH6\*02 is also used in the majority of antibodies with a long HCDR3 length, which is desirable for targets which are specifically bound by long HCDR3 length antibodies.





G G G A C C A C G G T C A C C T C S  
G T C A C T V C T C S

SEQ ID NO: 3 (JH2 Reference)

A T G A C C A T G A A G C T A G A C  
C C C G C A C G T G G A C A G T  
G A C A G A G A G T C

JH2 Alignment:

(top line=SEQ ID NO: 3, Middle line=SEQ ID NO:9, Bottom line=SEQ ID NO:10)

100-51 Hg  
133 601  
T G A C C G  
A C T G G W  
G A A G C T A G A G A C C C G G C C G G C C G T G G C A C C C T G G A C C A G T G A C A G A G G A G T C  
133 601  
A A G C T A G A G A C C C G G C C G T G G C A C C C T G G T C A C T G T G A C A G A G G A G T C A  
133 601  
A A G C T A G A G A C C C G G C C G T G G C A C C C T G G A C C A G T G A C A G A G G A G T C A

**TABLES**

In the tables, the notation is illustrated by the following example

IGLV1-40	V1-40*02	X53936	g9>c c10>g,L4>V
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5

Polymorphic variant IGV lambda V1-40\*02 has Genbank Accession No. X53936 and when compared to the \*01 variant, the V1-40\*02 variant has mutations at positions 9, 10 and 4. For example, at position 9, a "C" appears instead of a "G" that is present in the \*01 variant. The "|" is simply a notation separator, and does not indicate any mutation. For example the "g282 l" notation indicates no change (ie, position 282 is a g). "del#" means that the residue at that position is absent.

166

10

**Table 1: Human IgH V Polymorphic Variants**

All variant sequences appearing in Genbank under the listed Accession Numbers are incorporated herein in their entirety by reference as though explicitly recited herein and for possible inclusion in any of the claims herein.

15

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subgroup	IGHV	IGHV	Accession number	Description of mutations
1	IGHV1-2	IGHV1-2*01	X07448	SEQ ID NO: 11
		IGHV1-2*02	X62106	c163>t,R55>W   g233>t,t234>g,S78>M t299>c,V100>A
		IGHV1-2*03	X92208	c44>t,P15>L c163>t,R55>W   g233>t,t234>g,S78>M t299>c,V100>A
		IGHV1-2*04	Z12310	c163>t,R55>W a223>t,R75>W g233>t,t234>g,S78>M t299>c,V100>A
	IGHV1-3	IGHV1-3*01	X62109	SEQ ID NO: 12
				c6  t12  t167 ,I56  a208 ,K70  a291  c296 ,T99
		IGHV1-3*02	X62107	c6>t t12>g t167>g,I56>S a208>g,K70>E a291>g c296>t,T99>M
	IGHV1-8	IGHV1-8*01	M99637	SEQ ID NO: 13
		IGHV1-18	IGHV1-18*01	M99641
	IGHV1-18*02		X60503	g282>a

				SEQ ID NO: 14
				SEQ ID NO: 15
				g139 ,G47  t237
				t237>c
				g139>a,G47>R t237>c
				SEQ ID NO: 16
				c92 ,T31  g315
				c92>a,T31>N
				g315>t
				SEQ ID NO: 17
				g115 ,V39
				g115>a,V39>M
IGHV1-24	M99642	IGHV1-24*01		
IGHV1-45	X92209	IGHV1-45*01		
	AB019438	IGHV1-45*02		
	Z17391	IGHV1-45*03		
IGHV1-46	X92343	IGHV1-46*01		
	J00240	IGHV1-46*02		
	L06612	IGHV1-46*03		
IGHV1-58	M29809	IGHV1-58*01		
	AB019438	IGHV1-58*02		

IGHV1-69	IGHV1-69*01	L22582	SEQ ID NO: 18
	IGHV1-69*02	Z27506	g6>c   g18>a     g100>a,A34>T   g163>a,G55>R   t178>c,F60>L   c185>t,T62>I     g244>a,E82>K
	IGHV1-69*03	X92340	g291>t,E97>D
	IGHV1-69*04	M83132	g6>c     g163>a,G55>R   t178>c,F60>L   c185>t,T62>I     g244>a,E82>K
	IGHV1-69*05	X67905	g6>c     g238>a,A80>T
	IGHV1-69*06	L22583	g244>a,E82>K
	IGHV1-69*07	Z29978	g163>a,G55>R
	IGHV1-69*08	Z14309	g6>c   g18>a     g100>a,A34>T   g163>a,G55>R   t178>c,F60>L     g244>a,E82>K
	IGHV1-69*09	Z14307	g163>a,G55>R   t178>c,F60>L   c185>t,T62>I     g244>a,E82>K
	IGHV1-69*10	Z14300	g6>c     g54>a     t178>c,F60>L   c185>t,T62>I     g244>a,E82>K
	IGHV1-	Z14296	g6>c     g163>a,G55>R   t178>c,F60>L

					g6>c	
					g6>c   g54>a	
					SEQ ID NO: 19	
					SEQ ID NO: 20	
					c201	
					c201>t	
					SEQ ID NO: 21	
					a175>g,N59>D	
					a175>g,N59>D    c234>t	
					c299>g,A100>G  c314>t,A105>V  a317>g,c318>g,H106>R	
					a175>g,N59>D  a202>g,S68>G	
69*11		Z14301				
IGHV1-69*12		Z14214(st)				
IGHV1-69*13		Z18904				
IGHV1-c*01	IGHV1-c	Z12305				
IGHV1-f*01	IGHV1-f	Z29977				
IGHV1-f*02		X62111				
IGHV2-5*01	IGHV2-5	Z14072				
IGHV2-5*02		X93619				
IGHV2-5*03		L21963				
IGHV2-5*04		L21964				
IGHV2-						
	2					

5*05			
IGHV2-5*06	L21966		g39>a   a175>g,N59>D a202>g,S68>G
IGHV2-5*07	L21968		c314>t,A105>V   c299>g,A100>G
IGHV2-5*08	L21971		a4>g,l2>V a31>g,T11>A   g60>a g103>a,V35>M g106>c,G36>R g118>a,G40>S a175>g,N59>D
IGHV2-5*09	L21972		a4>g,l2>V   a175>g,N59>D a202>g,S68>G
IGHV2-5*10	X69690		a175>g,N59>D
IGHV2-26*01	M99648		a317>g,c318>g,H106>R  SEQ ID NO: 22
IGHV2-70*01	L21969		SEQ ID NO: 23 22 a2 ,Q1  g4 ,V2  g14 ,R5  g31 ,A11  a60  t67 ,C23  a70 ,T24  t106 ,C36  t116 ,V39  a138  c157 ,L53  t164 ,L55  a197 ,Y66  t210  g216 ,K72  a297  c299 ,A100  a301 ,c302 ,T101  g303  c309  c314 ,A105  g317 ,g318 ,R106  t320 ,I107
IGHV2-70*02	X92241		a297>g   a301>g,c302>t,T101>V

IGHV2-70*03	X92238	g14>a,R5>K   t164>g,L55>R a197>t,Y66>F  02>t,T101>V	t106>c,C36>R   a297>g   a301>g,c3
IGHV2-70*04	Z12330	g14>a,R5>K   t164>g,L55>R a197>t,Y66>F	t106>c,C36>R
IGHV2-70*05	Z27502	t164>g,L55>R a197>t,Y66>F	t106>c,C36>R t116>c,V39>A
IGHV2-70*06	X92239	g14>a,R5>K   t164>g,L55>R a197>t,Y66>F t210>c  g,c302>t,T101>V	t106>c,C36>R   a297>g   a301>
IGHV2-70*07	X92243	a297>g	a138>g   a301>g,c302>t,T101>V
IGHV2-70*08	X92245	a70>g,T24>A   a297>g	a301>g,c302>t,T101>V   t
IGHV2-70*09	L21962	g4>a,V2>  g14>a,R5>K g31>a,A11>T a60>g t67>c,C23>R  G   g303>a   c314>t,A105>V	g216>c,K72>N   c299>g,A100>
IGHV2-70*10	L21965	g14>a,R5>K  c157>a,L53>  t164>g,L55>R	t106>c,C36>R  
IGHV2-70*11	L21967	a2>g,Q1>R  64>g,L55>R	t1

3	IGHV2-70*12	L21970	g4>a,V2>  g14>a,R5>K g31>a,A11>T a60>g   g303>a
		IGHV2-70*13	AB019437  c309>t
	IGHV3-7*01	M99649	SEQ ID NO: 24
		IGHV3-7*02	X92288 g144>a
	IGHV3-9*01	M99651	SEQ ID NO: 25
		IGHV3-11*01	M99652  g13 ,V5  g32 ,G11  g47 ,G16  g178 ,G60  a184 ,c185 ,T62  t 188 ,I63  t196 ,Y66  a206 ,D69  g240  c243 ,D81  c296 ,g297 , T99  g301 ,g303 ,V101  t304 ,a305 ,Y102  t307 ,a308 ,c309 ,Y103  t 312  c314 ,A105
	IGHV3-11*02	M15496	9^10>ins^t   g32>del#,G11>del# g47>del#,G16>del#   a206>del#,D69>del#   c243>g,D81>E c296>del#,g297 >del#,T99>del# g301>t,g303>a,V101>L t304>c,a305>t,Y102>L t307>a,a30 8>c,c309>t,Y103>T t312>c c314>a,A105>E
	IGHV3-7		
	IGHV3-9		
	IGH3-11		

							Y t188>c,l63>T t196>a,Y66>N   g240>a	g13>t,V5>L   g178>a,G60>S a184>t,c185>a,T62>
IGHV3-13	IGHV3-11*03	X92287	IGHV3-13*01	X92217			SEQ ID NO: 27	
							g9 ,Q3  t52 ,S18  g77 ,S26  g95 ,S32  t165  t167 ,I56  c222  g224 ,R75	
IGHV3-15	IGHV3-13*02	M99653	IGHV3-13*03	U29582			g9>t,Q3>H t52>g,S18>A   g95>a,S32>N t165>c t167>a,I56>N c222>g	
							c77>g,S26>C   g224>a,R75>Q	
	IGHV3-15*01	X92216	IGHV3-15*02	M99654			SEQ ID NO: 28	
							g23 ,g24 ,G8  g32 ,G11  a40 ,K14  a81  t89 ,F30  g119 ,S40  t159  c163 ,R55  a169 ,K57  a178 ,T60  g181 ,D61  g196 ,D66  c210  a242 ,D81  a275 ,N92  c279  a297  a320	
IGHV3-15*03	IGHV3-15*02	M99654	IGHV3-15*03	M99408			g32>c,G11>A	
							g23>c,g24>c,G8>A g32>c,G11>A a40>c,K14>Q   t89>g,F30>C	
							a178>g,T60>A g181>a,D61>N   c210>t a242>t,D81>V	
	IGHV3-15*04	M99402				a169>g,K57>E		



					c279>t
					t159>c
					g196>a,D66>N
					a81>t   g119>a,S40>N t159>c
					g23>c ,G8>A   a40>c,K14>Q   t89>g,F30>C   c163>t, R55>C   a178>g,T60>A g181>a,D61>N   c210>t   a275 >t,N92>   a297>g a320>g
					SEQ ID NO: 29
					a6  a9
					a6>g a9>g
					SEQ ID NO: 30
					SEQ ID NO: 31
					SEQ ID NO: 32
					g9
					g9>a
IGHV3-15*05	M99403				
IGHV3-15*06	M99404				
IGHV3-15*07	M99406				
IGHV3-15*08	M99400				
IGHV3-16*01	M99655				
IGHV3-16*02	AB019440				
IGHV3-19*01	M99656				
IGHV3-20*01	M99657				
IGHV3-21*01	AB019439				
IGHV3-	M99658				

IGHV3-23	21*02				
	IGHV3-23*01	M99660			SEQ ID NO: 33 c164 ,A55  a169 ,g170 ,S57  g172 ,t174 ,G58  a175 ,S59  g181 ,G61  c201  c203 ,A68  c237  c243
	IGHV3-23*02	M35415			c203>g,A68>G c237>a
	IGHV3-23*03	U29481			c164>t,A55>V a169>t,g170>a,S57>Y g172>a,t174>c,G58>S a175>g,S59>G  g181>a,G61>S c201>t   c243>t
	IGHV3-23*04	AJ879486			t13>g
IGHV3-30	IGHV3-23*05	AY757302			a154>t,g155>a,S>Y g157>a,t159>a,G>S g166>a,G>S
	IGHV3-30*01	M83134			SEQ ID NO: 34 g18 ,E6  a27  a49 ,R17  c75  g80 ,G27  c101 ,t102 ,A34  t135  a138  a150  g163 ,V55  t169 ,c170 ,a171 ,S57  c201  g202 ,A68  a229 ,T77  c257 ,T86  g267  a275 ,N92  t288  a293 ,D98  g317 ,R106
	IGHV3-30*02	L26401			a49>g,R17>G c75>g   c101>g,t102>c,A34>G   a150 >g g163>t,V55>F t169>c,c170>g,a171>g,S57>R c201>t   g317>a,R106>K
	IGHV3-30*03	M99663			c101>g,t102>c,A34>G   a150>g   c201>t

					a150>g
IGHV3-30*04	L06615				
IGHV3-30*05	M77323			c101>g,t102>c,A34>G   a293>g,D98>G	
IGHV3-30*06	L06617			c75>g   c101>c,t102>c,A34>G	
IGHV3-30*07	L06614			t288>c	
IGHV3-30*08	M62737			g18>c,E6>D   g80>c,G27>A	
IGHV3-30*09	M77300			a229>g,T77>A   a150>g	
IGHV3-30*10	M77326			g202>a,A68>T	
IGHV3-30*11	M77331			c75>g	
IGHV3-30*12	M77338			a27>g   c75>g   c101>g,t102>c,A34>G   t288>c	
IGHV3-30*13	M77339			c101>g,t102>c,A34>G   c257>g,I86>R	

							a150>g
							g267>t
							a275>g,N92>S
							t135>c
							a138>g
							c101>g,t102>g,A34>G   a150>g
						106>K	c201>t   g317>a,R
							c75>g   c101>g,t102>g,A34>G   a150>g
							SEQ ID NO: 35
							c75  g317 ,R106
							c75>g g317>a,R106>K
							SEQ ID NO: 36
							g6  g150  g170 ,g171 ,W57  t177  t212 ,V71  t246  a251 ,K84  a263 ,Y88  g317 ,R106

				g6>a	t212>c,V71>A	a251>c,K84>T  a263>t,Y88>F
					t246>c	g317>a,R106>K
				g150>a	t177>c	
					g170>c,g171>a,W57>S	
				SEQ ID NO: 37		
				SEQ ID NO: 38		
				c302 ,A101		
				c302>t,A101>V		
				SEQ ID NO: 39		
				t32 ,V11  a100 ,T34  g138  t172 ,W58		
				t32>g,V11>G  a100>g,T34>A  g138>a  t172>g,W58>G		
				SEQ ID NO: 40		
				c58  c101 ,A34  t149 ,L50  t267 ,L89  t270 ,H90  t310		
IGHV3-33*02	M99665					
IGHV3-33*03	M77305					
IGHV3-33*04	M77335					
IGHV3-33*05	M77334					
IGHV3-35*01	M99666					
IGHV3-38*01	M99669					
IGHV3-38*02	AB019439					
IGHV3-43*01	M99672					
IGHV3-43*02	Z18901					
IGHV3-47*01	Z18900					
IGHV3-35						
IGHV3-38						
IGHV3-43						
IGHV3-47						

IGHV3-47*02	AB019438	c58>a c101>t,A34>V t149>c,L50>P   t270>a,H90>Q	
		IGHV3-47*03	M99674
IGHV3-48	M99675	SEQ ID NO: 41	g48  c96  a100 ,g101 ,c102 ,S34  a178 ,S60  t246  c287 ,A96  g303
	AB019438	c287>a,A96>D	
	U03893	g48>a c96>t a100>g,g101>a,c102>a,S34>E a178>g,S60>G t246>c g303>t	
IGHV3-49	M99676	SEQ ID NO: 42	g50 ,R17  t93  g94 ,D32  g100 ,A34  t124 ,F42  a202 ,T68  g245 ,G82
	M99401	g50>c,R17>P t93>g g94>t,D32>Y g100>c,A34>P t124>g,F42>V a202>g,T68>A g245>a,G82>D	
	AB019438	a202>g,T68>A g245>a,G82>D	
	AM940220	t124>g ,F42>V a202>g,T68>A g245>a,G82>D	

		IGHV3-49*05	AM940221	a202>g,T68>A g245>a,G82>D
IGHV3-53		IGHV3-53*01	M99679	t19 ,S7  g133 ,A45  c210  g315  a318
		IGHV3-53*02	Z12342	SEQ ID NO: 43 t19>a,S7>T
		IGHV3-53*03	J03617	g133>c,A45>P c210>t g315>t a318>g
		IGHV3-64*01	M99682	SEQ ID NO: 44 g1 ,E1  g26 ,G9  g70 ,A24  t198  t201  a205 ,N69  t210  c265 ,t267 ,L89  g274 ,g275 ,G92  c279  t296 ,M99  c314 ,A105  g317 ,R106
IGHV3-64		IGHV3-64*02	AB019437	g26>a,G9>E   a205>g,N69>D
		IGHV3-64*03	M77298	g70>t,A24>S t198>c t201>c a205>g,N69>D t210>a c265>g,t267>c,L89>V g274>a ,G92>S c279>t t296>c,M99>T c314>t,A105>V g317>a,R106>K
		IGHV3-64*04	M77299	g1>c,E1>Q   g70>t,A24>S t198>c t201>c a205>g,N69>D t210>a t267>g,  g274>a,g275>a,G92>N   t296>c,M99>T
		IGHV3-64*05	M77301	g70>t,A24>S t198>c t201>c a205>g,N69>D t210>a c265>g, ,L89>V g274>a ,G92>S c279>t t296>c,M99>T c314>t,A105>V g317>a,R106>K

IGHV3-66	IGHV3-66*01	X92218	SEQ ID NO: 45 g24   g37 ,V13  a81  g175 ,G59  a223  c288  g319
	IGHV3-66*02	Z27504	a223>c c288>t
	IGHV3-66*03	AB019437	g24>a g37>a,V13> a81>g g175>t,G59>C a223>c c288>t
	IGHV3-66*04	X70208	g319>c
IGHV3-72	IGHV3-72*01	X92206	SEQ ID NO: 46 t186
	IGHV3-72*02	Z29979	t186>c
IGHV3-73	IGHV3-73*01	X70197	SEQ ID NO: 47 t21
	IGHV3-73*02	AB019437	t21>c
IGHV3-74	IGHV3-74*01	L33851	SEQ ID NO: 48 c21  g197 ,c198 ,S66
	IGHV3-	Z17392	c21>t



							g197>c,c198>g,S66>T
							SEQ ID NO: 49
							SEQ ID NO: 50
							g303>t
							SEQ ID NO: 51
							c46>t,P16>S  >Y   g308>a,C103
							g308>a,C103>Y
							g73>a,V25>I  C103>Y   g308>a,
74*02							
IGHV3-74*03		J00239					
IGHV3-d*01	IGHV3-d	Z18898					
IGHV3-h*01	IGHV3-h	AJ879484					
IGHV3-h*02		AJ879485					
IGHV4-4*01	IGHV4-4	X05713					
IGHV4-4*02		X92232					
IGHV4-4*03		X92252					
IGHV4-4*04		X92253					
						4	

IGHV4-4*05	X92254	c20>t,S7>L   g308>a,C103>Y
IGHV4-4*06	Z75355	a234>g,I78>M a245>c,K82>T  g308>a,C103>Y
IGHV4-4*07	X62112	c46>t,P16>S g50>a,G17>E g70>a,A24>T   c93>t a97>t,g98>a,t99>c,S33>Y a100>t,N34>Y t120>c g124>a,V42> c129>g c136>g,a138>c,P46>A g147>a g163>c,a164>g,a165>t,E55>R c172>a,a173>c,t174>c,H58>T g207>c a234>g,I78>M a245>c,K82>T g308>a,C103>Y   a234>g,I78>M a245>c,K82>T g308>a,C103>Y
IGHV4-28*01	X05714	SEQ ID NO: 52 g48  g49 ,c51 ,D17  c185 ,T62  g297  c299 ,A100  a319
IGHV4-28*02	M83133	g48>a g49>c,c51>g,D17>Q c185>t,T62>
IGHV4-28*03	X92233	a319>g
IGHV4-28*04	X56358	g297>c c299>g,A100>G
IGHV4-28*05	X92260	c185>t,T62>

IGHV4-30-2	IGHV4-30-2*01	L10089	SEQ ID NO: 53 t163 ,a164 ,c165 ,Y55  c172 ,H58  a237  g245 ,R82  c288  g291  c300  c315  g319    c288>t   c315>g
	IGHV4-30-2*02	M95122	
	IGHV4-30-2*03	X92229	t163>a,a164>g,c165>t,Y55>S c172>t,H58>Y a237>c g245>c,R82>T c288>t g291>a c300>t c315>g g319>c
	IGHV4-30-2*04	Z75351	g245>c,R82>T   g291>a   c315>g
IGHV4-30-4	IGHV4-30-4*01	Z14238	SEQ ID NO: 54 g18 ,E6  a48  c49 ,g51 ,Q17  c134 ,P45  a166 ,I56  t228  c288  a291    a48>g c49>g,g51>c,Q17>D   c288>a
	IGHV4-30-4*02	Z14239	
	IGHV4-30-4*03	X92274	a291>g
	IGHV4-30-4*04	X92275	g18>c,E6>D   a166>t,I56>F
	IGHV4-30-4*05	Z75353	t228>c

			Z75360		c134>a,P45>H
IGHV4-31	IGHV4-31*01	IGHV4-31*01	L10098	SEQ ID NO: 55	a8 ,Q3  g37 ,V13  c69  c84  g100 ,G34  a164 ,Y55  t224 ,L75  a237  a244 ,c245 ,T82  t249  t285  t285-c287  a295 ,T99  t304 ,Y102
	IGHV4-31*02	IGHV4-31*02	M99683		c69>t   t224>g,L75>R
	IGHV4-31*03	IGHV4-31*03	Z14237		t224>g,L75>R
	IGHV4-31*04	IGHV4-31*04	M95120	a8>g,Q3>R	t224>g,L75>R
	IGHV4-31*05	IGHV4-31*05	M95121		t224>g,L75>R   t28 5-c287>del(3nt)# a295>g,T99>A
	IGHV4-31*06	IGHV4-31*06	X92270		g100>a,G34>S   t224>g,L75>R
	IGHV4-31*07	IGHV4-31*07	X92271		c84>a   t224>g,L75>R
	IGHV4-31*08	IGHV4-31*08	X92272		t224>g,L75>R a237>c   t249>c
	IGHV4-	IGHV4-	X92273		t224>g,L75>R    ,c245>a,T82>K t24

		31*09		9>c t285>c
		IGHV4-31*10	Z14235	g37>t,V13>L   a164>g,Y55>C t224>g,L75>R   a244>c, T82>P t249>c   t304>g,Y102>D
	IGHV4-34	IGHV4-34*01	AB019439	SEQ ID NO: 56 a12  g15  c16 ,Q6  g20 ,W7  g25 ,A9  t37 ,L13  g48  g49 ,E17  t85 ,S29  t88 ,F30  a118 ,S40  c129  c137 ,P46  g147  g163 ,a165 ,E55  a169 ,a170 ,t171 ,N57  c172 ,H58  a180  t199 ,Y67  g207  g226 ,t227 ,c228 ,V76  a234 ,I78  c249  c263 ,S88  g270 ,K90  a274 ,S92  c285  t300  a308 ,Y103
		IGHV4-34*02	M99684	g15>a
		IGHV4-34*03	X92255	t300>c
		IGHV4-34*04	X92236	V76>A   t199>a,Y67>N    ,t227>c ,
		IGHV4-34*05	X92237	c137>t,P46>L    ,t227>c ,V76>A   a118>t,S40>C   t199>a,Y67>N

IGHV4-34*06	X92256	a274>g,S92>G   t300>c
IGHV4-34*07	X92258	,t171>c    t300>c
IGHV4-34*08	M95113	t85>a,S29>T
IGHV4-34*09	Z14241	a12>g   c16>g,Q6>E g20>c,W7>S g25>c,A9>P t37>g,L13>V g48>a g49>c,E17>Q   g147>a   c228>t    c249>t   c285>t t300>c
IGHV4-34*10	Z14242	a12>g   c16>g,Q6>E g20>c,W7>S g25>c,A9>P t37>g,L13>V   g147>a   g226>a, ,V76> a234>g,I78>M   c263>a,S88>Y   t300>c
IGHV4-34*11	X05716	>g   g163>t,a165>t,E55>Y a169>t ,N57>Y c172>t,H58>Y a180>g t199>a,Y67>N g207>c  t227>c ,V76>A   g270>c,K90>N   t300>c a308>g,Y103>C   t88>g,F30>V   c129

				a170>t ,N57>
				g207>c
			SEQ ID NO: 57	
			a2 ,Q1  t56 ,L19  c237  g258 ,Q86  t262 ,S88  a291  t300  c319	
			g258>c,Q86>H	c319>g
			t300>c	
			a291>g	
			t56>c,L19>P	
			a2>g,Q1>R   c237>a   t262>c,S88>P a291>g t300>c	
			c237>a	a291>g t300>c  c319>g
IGHV4-34*12	X56591			
IGHV4-34*13	Z75356			
IGHV4-39*01	AB019439			
IGHV4-39*02	X05715			
IGHV4-39*03	X92259			
IGHV4-39*04	X92297			
IGHV4-39*05	M95116			
IGHV4-39*06	Z14236			
IGHV4-39*07	AM940222			
IGHV4-39				

	IGHV4-55	IGHV4-55*01	M99685	SEQ ID NO: 58 c44 ,P15  c237  a251 ,K84  c255 ,N85  a263 ,Y88  c288  c290 ,A97  g291  t319
		IGHV4-55*02	X92223	c237>a
		IGHV4-55*03	X92263	c237>a   a263>c,Y88>S
		IGHV4-55*04	X92265	c44>t,P15>L c237>a
		IGHV4-55*05	X92266	c44>t,P15>L
		IGHV4-55*06	X92267	c255>g,N85>K   c288>t
		IGHV4-55*07	X92268	a251>g,K84>R   a263>c,Y88>S   g291>a
		IGHV4-55*08	X92234	c237>a   t319>g
		IGHV4-55*09	X92235	a263>c,Y88>S   c290>t,A97>V   t319>a



	IGHV4-59	<p>IGHV4-59*01</p> <p>IGHV4-59*02</p> <p>IGHV4-59*03</p> <p>IGHV4-59*04</p> <p>IGHV4-59*05</p> <p>IGHV4-59*06</p> <p>IGHV4-59*07</p>	<p>AB019438</p> <p>M29812</p> <p>M95114</p> <p>M95117</p> <p>M95118</p> <p>M95119</p> <p>X56360</p>	<p>SEQ ID NO: 59</p> <p>g12  g16 ,E6  c20 ,S7  c25 ,P9  g37  g51 ,E17  a70 ,T24  t76 ,c77 ,S26  a88 ,I30  c135  c136 ,a138 ,P46  a147  t163 ,a164 ,t165 ,Y55  t172 ,a173 ,c174 ,Y58  a196 ,N66  c207  c228  a234 ,I78  a237  c249  g258  c285  t288  g291  c300  g319  a320  </p> <p> a88&gt;g,I30&gt;V </p> <p> g258&gt;a </p> <p>  ,c174&gt;t  a196&gt;t,N66&gt;Y c207&gt;g   a234&gt;g,I78&gt;M   t288&gt;c g291&gt;a c300&gt;t </p> <p> c135&gt;g  ,a138&gt;g   t163&gt;c,a164&gt;g, ,Y55&gt;R  ,c174&gt;t  a196&gt;t,N66&gt;Y c207&gt;g   a237&gt;c   t288&gt;c g291&gt;a c300&gt;t </p> <p> t76&gt;a ,S26&gt;T   c136&gt;g, a138&gt;t,P46&gt;A a147&gt;c  ,t165&gt;c,    a196&gt;t,N66&gt;Y c207&gt;g c228&gt;t   c249&gt;t   c285&gt;t t288&gt;c </p> <p> g51&gt;c,E17&gt;D </p>
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	IGHV7- 81	IGHV7- 81*01	AB019437	SEQ ID NO: 66
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**Table 2: Human IgH D Polymorphic Variants**

All variant sequences appearing in Genbank under the listed Accession Numbers are incorporated herein in their entirety by reference as though explicitly recited herein and for possible inclusion in any of the claims herein.

IGHD subgroup	IGHD name	Variant name	Genbank Accession number	Sequence
1	IGHD1-1	IGHD1-1*01	X97051	SEQ ID NO: 67
	IGHD1-7	IGHD1-7*01	X13972	SEQ ID NO: 68
	IGHD1-14	IGHD1-14*01	X13972	SEQ ID NO: 69
	IGHD1-20	IGHD1-20*01	X97051	SEQ ID NO: 70
	IGHD1-	IGHD1-	X97051	SEQ ID NO: 71

2	26	26*01	IGHD2- 2*01	J00232	SEQ ID NO: 72	AGGATATTGTTAGTAGTACCAGCTGCTATGCC			
					IGHD2- 2	IGHD2- 2*02	X97051	G>A	
						IGHD2- 2*03	M35648	A>T	
						IGHD2- 8	IGHD2- 8*01	X13972	SEQ ID NO: 73
					IGHD2- 8*02		J00233	AA>GG	
					IGHD2- 15	IGHD2- 15*01	J00234	SEQ ID NO: 74	
						IGHD2- 21	IGHD2- 21*01	J00235	SEQ ID NO: 75
							IGHD2- 21*01	X97051	I>C

3	IGHD3-3	21*02	X13972	SEQ ID NO: 76 GTATTACGATTTTGGAGTGGTTATTATACC
		IGHD3-3*01		
	IGHD3-9	IGHD3-3*02	X93618	CG>GC
		IGHD3-9*01	X13972	SEQ ID NO: 77
	IGHD3-10	IGHD3-10*01	X13972	SEQ ID NO: 78 GTATTACTATGGTTCGGGGGAGTTATTATAAC
		IGHD3-10*02	X93615	G>[OMITTED]
	IGHD3-16	IGHD3-16*01	X93614	SEQ ID NO: 79 GTATTATGATTACGTTTGGGGGAGTTATGCTTATACC
		IGHD3-16*02	X97051	GC>CG
	IGHD3-	IGHD3-	X93616	SEQ ID NO: 80



4	22	22*01			
	IGHD4-4	IGHD4-4*01	X13972	SEQ. ID NO: 81	
	IGHD4-11	D4-11*01	X13972	SEQ. ID NO: 82	
	IGHD4-17	IGHD4-17*01	X97051	SEQ. ID NO: 83	
5	IGHD4-23	IGHD4-23*01	X97051	SEQ. ID NO: 84	
	IGHD5-5	IGHD5-5*01	X13972	SEQ. ID NO: 85	
	IGHD5-12	IGHD5-12*01	X13972	SEQ. ID NO: 86	
	IGHD5-18	IGHD5-18*01	X97051</A<<B>	SEQ. ID NO: 87	
6	IGHD5-24	IGHD5-24*01	X97051	SEQ. ID NO: 88	
	IGHD6-6	IGHD6-6*01	X13972	SEQ. ID NO: 89	

	IGHD6-13	IGHD6-13*01	X13972	SEQ ID NO: 90
	IGHD6-19	IGHD6-19*01	X97051	SEQ ID NO: 91
	IGHD6-25	IGHD6-25*01	X97051	SEQ ID NO: 92
7	IGHD7-27	IGHD7-27*01	J00256	SEQ ID NO: 93

Underlined nucleotides in the \*01 variant are positions that are different in other variants, eg, GC>CG for IGHD3-16\*02 indicates that GC in the \*01 variant is instead CG in the \*02 variant. All other positions are identical to between the \*01 and other variant. Similar notation is used in other tables below to denote changes between variants of gene segments.

**Table 3: Human IgH J Polymorphic Variants**

All variant sequences appearing in Genbank under the listed Accession Numbers are incorporated herein in their entirety by reference as though explicitly recited herein and for possible inclusion in any of the claims herein.

IGHV name	Variant name	Genbank Accession number	Sequence
IGHJ1	J1*01	J00256	SEQ ID NO: 94
IGHJ2	J2*01	J00256	SEQ ID NO: 95
IGHJ3	J3*01	J00256	SEQ ID NO: 96 T GAT GCT TTT GAT <u>GTC</u> TGG GGC CAA GGG ACA ATG GTC ACC GTC TCT TCA G
	J3*02	X86355	<u>G</u> >A
IGHJ4	J4*01	J00256	SEQ ID NO: 97 <u>AC</u> TAC TTT GAC TAC TGG GGC <u>CAA</u> <u>GGA</u> ACC CTG GTC ACC GTC TCC TCA G

	J4*02	X86355	AA>AG
	J4*03	M25625	AC>GC GA>GG
IGHJ5	J5*01	J00256	SEQ ID NO: 98 AC AAC TGG TTC GAC <u>I</u> CC TGG GGC CAA GGA ACC CTG GTC ACC GTC TCC TCA G
	J5*02	X86355	<u>I</u> >C A>G
IGHJ6	J6*01	J00256	SEQ ID NO: 99 ATT TAC TAC TAC TAC <u>G</u> GT ATG GAC GTC TGG <u>G</u> GG <u>C</u> AA GGG ACC ACG GTC ACC GTC TCC TCA G
	J6*02	X86355	<u>G</u> G>GC
	J6*03	X86356	<u>G</u> GT>TAC <u>G</u> G>GC

				CA>AA
				GG>GC
				CA>AA
J6*04			AJ8794887	

**Table 4: Human Ig Vk Polymorphic Variants**

All variant sequences appearing in Genbank under the listed Accession Numbers are incorporated herein in their entirety by reference as though explicitly recited herein and for possible inclusion in any of the claims herein.

IGKV subgroup	IGKV name	Variant name	Accession number	Sequence
1	IGKV1-5	IGKV1-5*01	Z00001	SEQ ID NO: 100 GAC ATC CAG ATG ACC CAG TCT CCT TCC ACC CTG TCT GCA TCT GTA GGA GAC AGA GTC ACC ATC <u>ACT</u> TGC CGG GCC AGT CAG AGT ATT AGT AGC TGG ... .. TTG GCC TGG TAT CAG CAG AAA CCA GGG AAA GCC CCT AAG

				CTC CTG ATC TAT <u>GAT</u> GCC TCC ... .. AGT TTG GAA AGT GGG GTC CCA ... TCA AGG TTC AGC GGC AGT GGA ... .. TCT GGG ACA GAA TTC ACT CTC ACC ATC AGC AGC CTG CAG CCT GAT GAT TTT GCA ACT TAT TAC TGC CAA CAG TAT AAT AGT TAT TCT CC
	IGKV1-5*02	M23851		<u>AC</u> >AT
	IGKV1-5*03	X72813		<u>GAT</u> >AAG <u>GCC</u> >GCG <u>ICC</u> >TCT <u>TTG</u> >TTA
IGKV1-6	IGKV1-6*01	M64858		SEQ ID NO: 101
IGKV1-8	IGKV1-8*01	Z00014		SEQ ID NO: 102
IGKV1D-8	IGKV1D-8*01	Z00008		SEQ ID NO: 103
IGKV1-9	IGKV1-9*01	Z00013		SEQ ID NO: 104
IGKV1-12	IGKV1-	V01577		SEQ ID NO: 105

		12*01		<p>GAC ATC CAG ATG ACC CAG TCT CCA TCT <u>TCC</u> GTG TCT GCA TCT GTA GGA  GAC AGA GTC ACC ATC ACT TGT CGG GCG AGT CAG GGT ATT AGC AGC TGG  ..... TTA GCC TGG TAT CAG CAG AAA CCA GGG AAA GCC CCT AAG  CTC CTG ATC TAT GCT GCA TCC ..... AGT TTG CAA AGT GGG  GTC CCA ... TCA AGG TTC AGC GGC AGT GGA ..... TCT GGG ACA GAT TTC  ACT CTC <u>ACC</u> ATC AGC AGC CTG CAG CCT GAA GAT TTT GCA ACT TAC TAT  TGT CAA CAG GCT AAC AGT TTC CCT <u>CC</u></p>
IGKV1D-12	IGKV1D-12*01	X17263	<p><u>TCC</u>&gt;TCT  <u>ACC</u>&gt;ACT</p>	
IGKV1-12/IGKV1D-12 (1)	IGKV1-12*02/1D-12*02	V01576	<p><u>CC</u>&gt;TC</p>	
IGKV1-13	IGKV1-13*01	Z00010	<p>SEQ ID NO: 106  GCC ATC CAG TTG ACC CAG TCT CCA TCC TCC CTG TCT GCA TCT GTA GGA  GAC AGA GTC ACC ATC ACT TGC CGG GCA AGT CAG GGC ATT AGC AGT GCT  ..... TTA GCC <u>IGA</u> TAT CAG CAG AAA CCA GGG AAA GCT CCT AAG  CTC CTG ATC TAT GAT GCC TCC ..... AGT TTG GAA AGT GGG  GTC CCA ... TCA AGG TTC AGC GGC AGT GGA ..... TCT GGG ACA GAT TTC  ACT CTC ACC ATC AGC AGC CTG CAG CCT GAA GAT TTT GCA ACT TAT TAC  TGT CAA CAG TTT AAT <u>AAAT</u> TAC CCT CA</p>	

											<u>TGA&gt;TGG</u> <u>AAT&gt;AGT</u>	
											<u>TGA&gt;TGG</u>	
											SEQ ID NO: 107	
											GAC ATC CAG ATG ACC CAG TCT CCA TCC TCA CTG TCT GCA TCT GTA GGA GAC AGA GTC ACC ATC ACT TGT CGG GCG AGT CAG GGC ATT AGC AAT TAT ..... TTA GCC TGG TTT CAG CAG AAA CCA GGG AAA GCC CCT AAG TCC CTG ATC TAT GCT GCA TCC ..... AGT TTG CAA AGT GGG GTC CCA ... TCA <u>AGG</u> TTC AGC GGC AGT GGA ..... TCT GGG ACA GAT TTC ACT CTC ACC ATC AGC AGC CTG CAG CCT GAA GAT TTT GCA ACT TAT TAC TGC CAA CAG TAT AAT AGT TAC CCT CC	
											<u>AGG&gt;AAG</u>	
											SEQ ID NO: 108	
											GAC ATC CAG ATG ACC CAG TCT CCA TCC TCA CTG TCT GCA TCT GTA GGA GAC AGA GTC ACC ATC ACT TGT CGG GCG <u>AGI</u> CAG GGT ATT AGC AGC TGG ..... TTA GCC TGG TAT CAG CAG AAA CCA GAG AAA GCC CCT AAG TCC CTG ATC TAT GCT GCA TCC ..... AGT TTG CAA AGT GGG GTC CCA ... TCA AGG TTC AGC GGC AGT GGA ..... TCT GGG ACA GAT TTC ACT CTC ACC ATC AGC AGC CTG CAG CCT GAA GAT TTT GCA ACT TAT TAC	



				TGC CAA CAG TAT AAT AGT TAC CCT CC
			V00558	<u>AGT</u> >AGG
			X72808	SEQ ID NO: 109 GAC ATC CAG ATG ACC CAG TCT CCA TCC TCC CTG TCT GCA TCT GTA GGA GAC AGA GTC ACC ATC ACT TGC CGG GCA AGT CAG GGC ATT AGA AAT GAT ... .. TTA GGC TGG TAT CAG CAG AAA CCA GGG AAA GCC CCT AAG CGC CTG ATC TAT GCT GCA TCC ... .. AGT TTG CAA AGT GGG GTC CCA ... TCA AGG TTC AGC GGC AGT GGA ... .. TCT GGG ACA GAA TTC ACT CTC ACA ATC AGC <u>AGC</u> CTG CAG CCT GAA GAT TTT GCA ACT TAT TAC TGT CTA CAG CAT AAT AGT TAC CCT CC
			D88255	<u>AGC</u> > AAC
			X63392	SEQ ID NO: 110 <u>AAC</u> ATC CAG ATG ACC CAG TCT CCA TCT GCC ATG TCT GCA TCT GTA GGA GAC AGA GTC ACC ATC ACT TGT CGG GCG AGG CAG GGC ATT AGC AAT TAT ... .. TTA GCC TGG TTT CAG CAG AAA CCA GGG AAA GTC CCT AAG <u>CAC</u> CTG ATC TAT GCT GCA TCC ... .. AGT TTG CAA AGT GGG GTC CCA ... TCA AGG TTC AGC GGC AGT GGA ... .. TCT GGG ACA GAA TTC ACT CTC ACA ATC AGC AGC CTG CAG CCT GAA GAT TTT GCA ACT TAT TAC TGT CTA CAG CAT AAT AGT TAC CCT CC
			FM164407	<u>AAC</u> >GAC
IGKV1D-16*02	IGKV1D-17	IGKV1-17	IGKV1-16*02	
IGKV1D-17*01			IGKV1D-17*01	
IGKV1D-17*02			IGKV1D-17*02	
IGKV1D-17*01			IGKV1D-17*01	
IGKV1D-17*02			IGKV1D-17*02	

		17*02		<u>CAC</u> >CGC
IGKV1-27	IGKV1-27*01	IGKV1-27*01	X63398	SEQ ID NO: 111
IGKV1-33	IGKV1-33*01	IGKV1-33*01	M64856	SEQ ID NO: 112
IGKV1D-33	IGKV1D-33*01	IGKV1D-33*01	M64855	SEQ ID NO: 113
IGKV1-37	IGKV1-37*01	IGKV1-37*01	X59316	SEQ ID NO: 114
IGKV1D-37	IGKV1D-37*01	IGKV1D-37*01	X71893	SEQ ID NO: 115
IGKV1-39	IGKV1-39*01	IGKV1-39*01	X59315	SEQ ID NO: 116 GAC ATC CAG ATG ACC CAG TCT CCA TCC <u>ICC</u> CTG TCT GCA TCT GTA GGA GAC AGA GTC ACC ATC ACT TGC CGG GCA AGT CAG AGC ATT AGC AGC TAT ... .. TTA AAT TGG TAT CAG CAG AAA CCA GGG AAA GCC CCT AAG CTC CTG ATC TAT GCT GCA TCC ... .. AGT TTG CAA AGT GGG GTC CCA ... TCA AGG TTC AGT GGC AGT GGA ... .. TCT GGG ACA GAT TTC ACT CTC ACC ATC AGC AGT CTG CAA CCT GAA GAT TTT GCA ACT <u>TAC</u> TAC TGT <u>CAA</u> CAG <u>AGI</u> TAC AGT <u>ACC</u> CCT CC
	IGKV1-39*02	IGKV1-39*02	X59318	<u>TCC</u> >TTC <u>TAC</u> >TAT

				<p><u>CAA CAG AGT</u>&gt; CAG TGT GGT <u>ACC</u>&gt;ACA</p>	
IGKV1D-39	IGKV1D-39*01	X59312	SEQ ID NO: 117		
IGKV1D-42	IGKV1D-42*01	X72816	SEQ ID NO: 118		
IGKV1D-43	IGKV1D-43*01	X72817	SEQ ID NO: 119		
IGKV2-24	IGKV2-24*01	X12684	SEQ ID NO: 120		
IGKV2D-24	IGKV2D-24*01	X63401	SEQ ID NO: 121		
IGKV2-28	IGKV2-28*01	X63397	SEQ ID NO: 122		
IGKV2D-28	IGKV2D-28*01	X12691	SEQ ID NO: 123		
IGKV2-29	IGKV2-29*01	X63396	SEQ ID NO: 124	<p>GAT ATT GTG ATG ACC CAG ACT CCA CTC TCT CTG TCC GTC ACC CCT GGA CAG CCG GCC TCC ATC TCC TGC AAG TCT AGT CAG AGC CTC CTG CAT AGT GAT GGA AAG ACC TAT ... TTG TAT TGG TAC CTG CAG AAG CCA GGC CAG</p>	
2					

				TCT CCA CAG CTC CTG ATC TAT GAA GTT TCC ... .. AGC CGG TTC TCT GGA GTG CCA ... GAT AGG TTC AGT GGC AGC GGG ... .. TCA GGG ACA GAT TTC ACA CTG AAA ATC AGC CGG GTG GAG GCT GAG GAT GTT GGG GTT TAT TAC TGA ATG CAA GGT ATA CAC CTT CCT CC
	IGKV2-29*02	U41645		<u>CTG</u> > CTA <u>TGA</u> > TGC
	IGKV2-29*03	A1783437		<u>TGA</u> > TGC
IGKV2D-29	IGKV2D-29*01	M31952		SEQ ID NO: 125 GAT ATT GTG ATG ACC CAG ACT CCA CTC TCT CTG TCC GTC ACC CCT GGA CAG CCG GCC TCC ATC TCC TGC AAG TCT AGT CAG AGC CTC CTG CAT AGT GAT GGA AAG ACC TAT ... TTG TAT TGG TAC CTG CAG AAG CCA GGC CAG <u>CCT</u> CCA CAG CTC CTG ATC TAT GAA GTT TCC ... .. AAC CGG TTC TCT GGA GTG CCA ... GAT AGG TTC AGT GGC AGC GGG ... .. TCA GGG ACA GAT TTC ACA CTG AAA ATC AGC CGG GTG GAG GCT GAG GAT GTT GGG GTT TAT TAC TGC ATG CAA AGT ATA CAG CTT CCT CC
	IGKV2D-29*02	U41644		<u>CCT</u> >TCT
IGKV2-30	IGKV2-30*01	X63403		SEQ ID NO: 126 GAT GTT GTG ATG ACT CAG TCT CCA CTC TCC CTG CCC GTC ACC CTT GGA CAG CCG GCC TCC ATC TCC TGC AAG TCT AGT CAA AGC CTC GTA <u>IAC</u> AGT GAT GGA AAC ACC TAC ... TTG AAT TGG TTT CAG CAG AGG CCA GGC CAA

				<p>TCT CCA AGG CGC CTA ATT TAT AAG GTT TCT ... .. AAC CGG                  GAC TCT GGG GTC CCA ... GAC AGA TTC AGC GGC AGT GGG ... .. TCA GGC                  ACT GAT TTC ACA CTG AAA ATC AGC AGG GTG GAG GCT GAG GAT GTT GGG                  GTT TAT TAC TGC ATG CAA GGT ACA CAC TGG CCT CC</p> <p><u>TAC</u>&gt;CAC</p>
		IGKV2-30	FM164408	<p>SEQ ID NO: 127</p>
		IGKV2-40	X63402	<p>SEQ ID NO: 128</p> <p>GAT ATT GTG ATG ACC CAG ACT CCA CTC TCC CTG CCC GTC ACC CCT GGA                  GAG CCG GCC TCC ATC TCC TGC AGG TCT AGT CAG AGC CTC TTG GAT AGT                  GAT GAT GGA AAC ACC TAT TTG <u>GAC TGG</u> TAC CTG CAG AAG CCA GGG CAG                  TCT CCA CAG CTC CTG ATC TAT ACG CTT TCC ... .. TAT CGG GCC                  TCT GGA GTC CCA ... GAC AGG TTC AGT <u>GGC</u> AGT GGG ... .. TCA GGC ACT                  GAT TTC ACA CTG AAA ATC AGC AGG GTG GAG GCT GAG GAT GTT GGA GTT                  TAT TAC TGC ATG CAA CGT ATA GAG TTT CCT TC</p> <p><u>GAC TGG</u>&gt;GAT TGT  <u>GGC</u>&gt;GAC</p>
		IGKV2D-30	X59317	<p>SEQ ID NO: 129</p>
		IGKV2D-40	X59311	<p>SEQ ID NO: 129</p>

<p><b>3</b></p>	<p>IGKV3-7</p>	<p>IGKV3-7*01</p>	<p>X02725</p>	<p>SEQ ID NO: 130</p> <p>GAA ATT GTA ATG ACA CAG TCT CCA CCC ACC CTG TCT TTG TCT CCA GGG  GAA AGA GTC ACC CTC TCC TGC AGG GCC AGT CAG AGT GTT AGC AGC AGC  TAC ..... TTA ACC TGG TAT CAG CAG AAA CCT GGC CAG GCG CCC  AGG CTC CTC ATC TAT GGT GCA TCC ..... ACC AGG GCC ACT  AGC ATC CCA ... GCC AGG TTC AGT GGC AGT GGG ..... TCT GGG ACA GAC  TTC ACT CTC ACC ATC AGC AGC CTG CAG CCT GAA GAT TTT GCA GTT TAT  TAC TGT CAG CAG GAT CAT AAC TTA CCT CC</p>
		<p>IGKV3-7*02</p>	<p>X72812</p>	<p>ACC&gt;TCC  TAT&gt;TAC  GGC&gt;GGG  GCG&gt;GCT  AGC&gt;GGC  ACA&gt;AGA  CAT&gt;TAT</p>
		<p>IGKV3-7*03</p>	<p>K02769</p>	<p>SEQ ID NO: 131</p> <p>GAA ATT GTA ATG ACA CAG TCT CCA CCC ACC CTG TCT TTG TCT CCA GGG  GAA AGA GTC ACC CTC TCC TGC AGG GCC AGT CAG AGT GTT AGC AGC AGC  TAC ..... TTA ACC TGG TAT CAG CAG AAA CCT GGC CAG GCG CCC  AGG CTC CTC ATC TAT GGT GCA TCC ..... ACC AGG GCC ACT  AGC ATC CCA ... GCC AGG TTC AGT GGC AGT GGG ..... TCT GGG ACA GAC  TTC ACT CTC ACC ATC AGC AGC CTG CAG CCT GAA GAT TTT GCA GTT TAT</p>

		TAC TGT CAG CAG GAT CAT AAC TTA CCT CC	
	IGKV3-7*04	FM164409	<u>CAT</u> >TAT
IGKV3D-7	IGKV3D-7*01	X72820	SEQ ID NO: 132
IGKV3-11	IGKV3-11*01	X01668	SEQ ID NO: 133 GAA ATT GTG TTG ACA CAG TCT CCA GCC ACC CTG TCT TTG TCT CCA GGG GAA AGA GCC ACC CTC TCC TGC AGG GCC AGT CAG AGT GTT AGC AGC TAC ... .. TTA GCC TGG TAC CAA CAG AAA CCT GGC CAG GCT CCC AGG CTC CTC ATC TAT GAT GCA TCC ... .. AAC AGG GCC ACT GGC ATC CCA ... GCC AGG TTC AGT GGC AGT GGG ... .. TCT GGG <u>ACA</u> GAC TTC ACT CTC ACC ATC AGC AGC CTA GAG CCT GAA GAT TTT GCA GTT TAT TAC TGT CAG CAG CGT AGC AAC TGG CCT CC
	IGKV3-11*02	K02768	<u>ACA</u> >AGA
IGKV3D-11	IGKV3D-11*01	X17264	SEQ ID NO: 134
IGKV3-15	IGKV3-15*01	M23090	SEQ ID NO: 135
IGKV3D-15	IGKV3D-15	X72815	SEQ ID NO: 136

<p>15*01</p>	<p>GAA ATA GTG ATG <u>ACG</u> CAG TCT CCA GCC ACC CTG TCT GTG TCT CCA GGG          GAA AGA GCC ACC CTC TCC TGC AGG GCC AGT CAG AGT GTT AGC AGC AAC          ..... TTA GCC TGG TAC CAG AAA CCT GGC CAG GCT CCC AGG          CTC CTC ATC TAT GGT GCA TCC ..... ACC AGG GCC ACT GGC          ATC CCA ... GCC AGG TTC AGT GGC AGT GGG ..... TCT GGG ACA GAG TTC          ACT CTC ACC ATC AGC AGC CTG CAG TCT GAA GAT TTT GCA GTT TAT TAC          TGT CAG CAG TAT AAT AAC <u>IGG</u> CCT CC</p>	<p>M23091</p>	<p><u>ACG</u>&gt;ATG <u>TGG</u>&gt;TGA</p>
<p>IGKV3D-15*02</p>	<p>SEQ ID NO: 137</p>	<p>X12686</p>	<p>GAA ATT GTG TTG <u>ACG</u> CAG TCT CCA GGC ACC CTG TCT TTG TCT CCA GGG          GAA AGA GCC ACC CTC TCC TGC AGG GCC AGT CAG AGT GTT AGC AGC AGC          TAC ..... TTA GCC TGG TAC CAG AAA CCT GGC CAG GCT CCC          AGG CTC CTC ATC TAT GGT GCA TCC ..... AGC AGG GCC ACT          GGC ATC CCA ... <u>GAC</u> AGG TTC AGT GGC AGT GGG ..... TCT GGG ACA GAC          TTC ACT CTC ACC ATC AGC AGA CTG GAG CCT GAA GAT TTT GCA GTG TAT          TAC TGT CAG CAG TAT GGT AGC TCA CCT CC</p>
<p>IGKV3-20</p>	<p>IGKV3-20*01</p>	<p>X56593</p>	<p><u>ACC</u>&gt;CCC <u>CTG</u>&gt;TGT <u>ICI</u>&gt;CTT <u>TTG</u>&gt;TGT <u>TCT</u>&gt;CTC</p>
<p>IGKV3-20*01, Tou-kv325</p>	<p>IGKV3-20*01, Tou-kv325</p>	<p>X56593</p>	<p><u>ACC</u>&gt;CCC <u>CTG</u>&gt;TGT <u>ICI</u>&gt;CTT <u>TTG</u>&gt;TGT <u>TCT</u>&gt;CTC</p>





All variant sequences appearing in Genbank under the listed Accession Numbers are incorporated herein in their entirety by reference as though explicitly recited herein and for possible inclusion in any of the claims herein.

IGLV subgroup	IGLV Gene name	IGLV variant name	Accession number	Sequences & Description of mutations
1	IGLV1-36	V1-36*01	Z73653	SEQ ID NO: 144
	IGLV1-40	V1-40*01	M94116	SEQ ID NO: 145
		V1-40*02	X53936	g9>c c10>g,L4>V
		V1-40*03	Z22192	g9>c c10>g,L4>V a253>g,T85>A
	IGLV1-41	V1-41*01	M94118	SEQ ID NO: 146
		V1-41*02	D87010	g295>t,E99>* c332>t,P111>L

		V1-44*01	Z73654	SEQ ID NO: 147			
IGLV1-44		V1-47*01	Z73663	SEQ ID NO: 148			
		V1-47*02	D87016	g168>t,R56>S			
IGLV1-50		V1-50*01	M94112	SEQ ID NO: 149			
		V1-51*01	Z73661	SEQ ID NO: 150			
IGLV1-51		V1-51*02	M30446	t162>c c168>a,D56>E			
		V2-8*01	X97462	SEQ ID NO: 151			
IGLV2-8	2	V2-8*02	L27695	g37>a,G13>R			

		V2-8*03	Y12418	c230>t,S77>F
IGLV2-11		V2-11*01	Z73657	SEQ ID NO: 152
		V2-11*02	Z22198	t96>g
		V2-11*03	Y12415	g132>a
IGLV2-14		V2-14*01	Z73664	SEQ ID NO: 153
		V2-14*02	L27822	c87>t t93>g g94>a,G32>S t103>c,a104>t,Y35>L t170>g,V57>G t198>g,N66>K
		V2-14*03	Y12412	g132>a g168>t,E56>D
IGLV2-18		V2-14*04	Y12413	g168>t,E56>D
		V2-18*01	Z73642	SEQ ID NO: 154

18	V2-18*02	L27697	t317>c,L106>S
	V2-18*03	L27694	t272>c,I91>T t317>c,L106>S
	V2-18*04	L27692	t227>c,F76>S t249>c   t317>c,L106>S
	V2-23*01	X14616	SEQ ID NO: 155
IGLV2- 23	V2-23*02	Z73665	g170>t,G57>V a339>c,L113>F
	V2-23*03	D86994	a339>c,L113>F
	V2-33*01	Z73643	SEQ ID NO: 156
IGLV2- 33	V2-33*02	L27823	a3>g

		V2-33*03	L27691	t96>c a256>g,M86>V
3	IGLV3-1	V3-1*01	X57826	SEQ ID NO: 157
		V3-9*01	X97473	a52 ,T18  a82 ,I28  a88 ,g89 ,S30  a95 ,N32  g173 ,S58
			X74288	a52>g,T18>A a82>c,I28>L a88>t,g89>a,S30>Y a95>g ,N32>S g173>a,S58>N
	IGLV3-9	V3-9*02	X51754	a52>g,T18>A a82>c,I28>L a88>t,g89>a,S30>Y a95>del#,N32>del# g173>a,S58>N
		V3-10*01	X97464	SEQ ID NO: 158
			L29166	g166>a,E56>K c207>a t270>c c299>a,A100>D a319>g,T107>A a325>t,g326>a,S109>Y
	IGLV3-10	V3-12*01	Z73658	SEQ ID NO: 159
		V3-12*02	D86998	a259>g,T87>A
12	IGLV3-12			

IGLV3-16	V3-16*01	X97471	SEQ ID NO: 160		
IGLV3-19	V3-19*01	X56178	SEQ ID NO: 161		
IGLV3-21	V3-21*01	X71966	SEQ ID NO: 162		
	V3-21*02	D87007	a27>g a49>c,K17>Q a160>g,I54>V t166>g,Y56>D c324>t		
IGLV3-22	V3-21*03	M94115	a27>g   a160>g,I54>V t166>g,Y56>D c324>t		
	V3-22*01	Z73666	SEQ ID NO: 163		
IGLV3-25	V3-22*02	X71967	c53>a,T18>K g88>a,E30>K g140>del#,G47>del# c148>t,c149>g,t150>a,P50>* 150^151>ins^tatacf,50^51>ins^Y# g322>a,D108>N c330>t		
	V3-25*01	X97474	SEQ ID NO: 164		

		V3-25*02	D86994	t14>c,g15>a,M5>T
		V3-25*03	L29165	t14>c,g15>a,M5>T t294>c
	IGLV3-27	V3-27*01	D86994	SEQ ID NO: 165
	IGLV3-32	V3-32*01	Z73645	SEQ ID NO: 166
4	IGLV4-3	V4-3*01	X57828	SEQ ID NO: 167
	IGLV4-60	V4-60*01	Z73667	SEQ ID NO: 168
		V4-60*02	D87000	a288>t,L96>F
		V4-60*03	AF073885	t287>c,a288>t,L96>S
	IGLV4-	V4-69*01	Z73648	SEQ ID NO: 169



			V4-69*02	U03868					c198>t
			V5-37*01	Z73672					SEQ ID NO: 170
			V5-39*01	Z73668					t81 ,g135>t
			V5-39*02	AF216776					t81>c,g135>t
			V5-45*01	Z73670					SEQ ID NO: 171
			V5-45*02	Z73671					g22>t,A8>S g75>a
			V5-45*03	D86999					g22>t,A8>S
			V5-48*01	Z73649					SEQ ID NO: 172
69									
	5	IGLV5-37							
		IGLV5-39							
		IGLV5-45							
		IGLV5-48							

	IGLV5-52	V5-52*01	Z73669	SEQ ID NO: 173
6	IGLV6-57	V6-57*01	Z73673	SEQ ID NO: 174
7	IGLV7-43	V7-43*01	X14614	SEQ ID NO: 175
	IGLV7-46	V7-46*01	Z73674	SEQ ID NO: 176
		V7-46*02	D86999	c275>t,S92>L
8	IGLV8-61	V7-46*03	Z22210	c271>del# L91>del#
		V8-61*01	Z73650	SEQ ID NO: 177
		V8-61*02	U03637	c223>t,R75>C
		V8-61*03	AF266511	g210>c

9	IGLV9-49	V9-49*01	Z73675	SEQ ID NO: 178
		V9-49*02	Z73675	g291>a
		V9-49*03	U03869	g153>a
10	IGLV10-54	V10-54*01	Z73676	SEQ ID NO: 179
		V10-54*02	D86996	a86>t,N29>    a228>c,L76>F g320>t,W107>L
		V10-54*03	S70116	g33>c   t125>c,g126>t,L42>P c127>g,Q43>E
11	IGLV11-55	V11-55*01	D86996	SEQ ID NO: 180


**Table 6: Human IgH.Jk Polymorphic Variants**

All variant sequences appearing in Genbank under the listed Accession Numbers are incorporated herein in their entirety by reference as though explicitly recited herein and for possible inclusion in any of the claims herein.

IGKJ gene name	Variant name	Accession number	Sequence
IGKJ1	IGKJ1*01	J00242	SEQ ID NO: 181
	IGKJ2*01	J00242	SEQ ID NO: 182
IGKJ2	IGKJ2*01	Z70260 #c	TG TAC <u>ACT</u> TTT GGC CAG GGG ACC AAG CTG GAG ATC AAA C
	IGKJ2*02		<u>TAC</u> >TGC
	IGKJ2*03	U95246 #c	<u>ACT</u> >AGT

	IGK12*04	L40735 #c	<u>TAC</u> >TGC <u>ACT</u> >AGT
<b>IGK13</b>	IGK13*01	J00242	SEQ ID NO: 183
<b>IGK14</b>	IGK14*01	J00242	SEQ ID NO: 184 G CTC <u>ACT</u> TTC GGC GGA GGG ACC AAG GTG GAG ATC AAA C
	IGK14*02	AF103571 #c	<u>ACT</u> >ACG
<b>IGK15</b>	IGK15*01	J00242	SEQ ID NO: 185

**Table 7: Human IgH Jλ Polymorphic Variants**

All variant sequences appearing in Genbank under the listed Accession Numbers are incorporated herein in their entirety by reference as though explicitly recited herein and for possible inclusion in any of the claims herein.

IGLJ gene name	Variant name	Accession number	Sequence
IGLJ1	J1*01	X04457	SEQ ID NO: 186
IGLJ2	J2*01	M15641	SEQ ID NO: 187
IGLJ3	J3*01	M15642	SEQ ID NO: 188 T <u>GTG</u> <u>GTA</u> TTC GGC GGA GGG ACC AAG CTG ACC GTC CTA G
	J3*02	D87023(1)	<u>GTG</u> > TGG <u>GTA</u> > GTG
IGLJ4	J4*01	X51755	SEQ ID NO: 189
IGLJ5	J5*01	X51755	SEQ ID NO: 190 C TGG GTG TTT GGT GAG GGG <u>ACC</u> GAG CTG ACC GTC CTA G
	J5*02	D87017	<u>ACC</u> >ACG

IGLJ6	J6*01	M18338	SEQ ID NO: 191
IGLJ7	J7*01	X51755	SEQ ID NO: 192 T GCT GTG TTC GGA GGA GGC ACC CAG CTG ACC <u>GTC</u> CTC G
	J7*02	D87017	<u>GTC</u> >GCC

**TABLE 8: 1000 GENOMES PROJECT HUMAN POPULATIONS**

Below is a summary of the ethnic population origin of samples that the 1000 Genomes Project sequences.

**Population****European ancestry**

Utah residents (CEPH) with Northern and Western European ancestry (CEU)

Toscani in Italia (TSI)

British from England and Scotland (GBR)

Finnish from Finland (FIN)

Iberian populations in Spain (IBS)

**East Asian ancestry**

Han Chinese in Beijing, China (CHB)

Japanese in Toyko, Japan (JPT)

Han Chinese South (CHS)

Chinese Dai in Xishuangbanna (CDX)

Kinh in Ho Chi Minh City, Vietnam (KHV)

Chinese in Denver, Colorado (CHD) (pilot 3 only)

**West African ancestry**

Yoruba in Ibadan, Nigeria (YRI)

Luhya in Webuye, Kenya (LWK)

Gambian in Western Division, The Gambia (GWD)

Malawian in Blantyre, Malawi (MAB)

K00004-1 WO



**Population**

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West African Population (TBD)

**Americas**

African Ancestry in Southwest US (ASW)

African American in Jackson, MS (AJM)

African Caribbean in Barbados (ACB)

Mexican Ancestry in Los Angeles, CA (MXL)

Puerto Rican in Puerto Rico (PUR)

Colombian in Medellin, Colombia (CLM)

Peruvian in Lima, Peru (PEL)

**South Asian ancestry**

Ahom in the State of Assam, India

Kayadtha in Calcutta, India

Reddy in Hyderabad, India

Maratha in Bombay, India

Punjabi in Lahore, Pakistan

**Table 9: Immunoglobulin Gene Usage in Human Antibody Responses to Infectious Disease Pathogens**

		<p><b><u>KAPPA V GENES</u></b></p> <ul style="list-style-type: none"> <li>• Vk II germline gene A2 + JK3</li> <li>• Vk II family gene + JK4</li> <li>• 94% identical to the A27 (Humkv325) germ line gene</li> <li>• a VkI gene family member; κI-15A (KL012)</li> </ul> <p><b><u>LAMBDA V GENES</u></b></p> <ul style="list-style-type: none"> <li>• Four Vλ VII family members that are 96-98% identical to each other</li> <li>• Vλ II family members (82, 89 and 91% homologous to Vλ2.1 gene) + VHIII segments closely homologous to germline gene 9.1</li> <li>• Vλ VII 4A</li> <li>• All with Jλ homologous to germline Jλ2 and Jλ3</li> </ul> <p><b><u>VH GENES</u></b></p> <ul style="list-style-type: none"> <li>• VH 96% identical to the VH germ line gene segment DP77 (V3-21)</li> </ul>	<p>Haemophilus influenzae type b polysaccharide (Hib PS)</p>	<p>Haemophilus influenzae</p> <ol style="list-style-type: none"> <li>1. Lonberg, Nat Biotech 2005; [human PBMCs]</li> <li>2. Adderson et al, J Clin Invest 1992; [Human PBLs]</li> <li>3. Chung et al, J Immunol 1993</li> <li>4. Nadel et al, J Immunol 1998</li> <li>5. Feeney et al, J Clin Invest 1996</li> <li>6. Lucas et al, Infect Immun 1994; [Human PBLs]</li> <li>7. Adderson et al, J Clin Invest 1993; [Human PBLs]</li> <li>8. Granoff et al, J Clin Invest 1993; [human PBLs]</li> <li>9. Azmi et al, Infect Immun 1994; [human tonsil cells]</li> </ol>
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<ul style="list-style-type: none"> <li>• LSG6.1, LSG12.1, V<sub>H</sub>III VH26, V<sub>H</sub>III 9.1</li> <li>• <u>VH and VL COMBINATIONS</u></li> <li>• V<sub>H</sub>III 9.1 + V<sub>λ</sub>VII 4A</li> <li>• V<sub>H</sub>III 9.1 + V<sub>λ</sub>II 2.1</li> <li>• V<sub>H</sub>III 9.1 + V<sub>λ</sub>II A2</li> <li>• V<sub>H</sub>III VH26 + V<sub>λ</sub>II 2.1</li> </ul>			
<ul style="list-style-type: none"> <li>• V<sub>H</sub>III 9.1; V<sub>H</sub>III H11; V<sub>H</sub>III VH26</li> <li>• κI 15A</li> <li>• Vλ2.1</li> </ul>	<ul style="list-style-type: none"> <li>• Polysaccharide capsule of E coli K1</li> <li>• Meningococcal B polysaccharide; Poly[α(2→8)-N-acetylneuramic acid</li> </ul>	<ul style="list-style-type: none"> <li>• E coli K1</li> <li>• Neisseria meningitidis Group B</li> </ul>	<p>9. Azmi et al, Infect Immun 1994</p>
<ul style="list-style-type: none"> <li>• VHIII or VHIV family member</li> <li>• VλI or Vλ3 member</li> <li>• VH26 + Dk1 + JH6 with IGLV1S2 + Jλ2</li> <li>• VH4.18</li> <li>• VH2-1 (VH3) + D region Dxp'1 + JH5 with Vλ3 cML70 + Jλ3</li> <li>• VH1GRR+ JH3 + Dn4r or D2r with IGLV1S2 + Jλ2</li> </ul>	<ul style="list-style-type: none"> <li>• HSV 120-kD glycoprotein</li> <li>• 116-, 105-, 64-kD glycoproteins of VZV</li> </ul>	<ul style="list-style-type: none"> <li>• Herpes family virus</li> <li>• Herpes simplex virus (HSV); HSV-1; HSV-2</li> <li>• Varicella zoster virus (VZV)</li> </ul>	<p>10. Huang et al, J Clin Invest 1992; [human tonsils]</p>

<ul style="list-style-type: none"> <li>• For VZV Abs: ha3h2 (VH3) with la1h2 (VA); or ha1c1 (VH1) with la1v1 (VA1)</li> <li>• For VZV Abs: ha4h3 (VH4) with la3h3 (VA3)</li> </ul>			
<ul style="list-style-type: none"> <li>• Hv1051 (VH)</li> <li>• Kv325 (Vk)</li> </ul>		<ul style="list-style-type: none"> <li>• Cytomegalovirus (CMV)</li> </ul>	<p>10. Huang et al, J Clin Invest 1992;</p>
<ul style="list-style-type: none"> <li>• 71-2 (VH)</li> <li>• Hv1f10 (VH)</li> <li>• VH4.11</li> <li>• 71-4 (VH)</li> <li>• VH251</li> <li>• VH1-69</li> </ul>		<ul style="list-style-type: none"> <li>• HIV</li> </ul>	<p>10. Huang et al, J Clin Invest 1992; 11. Wang &amp; Palese, Science 2011</p>
<ul style="list-style-type: none"> <li>• VH1-69</li> </ul>	<p>Haemagglutinin (HA)</p>	<ul style="list-style-type: none"> <li>• Influenza virus, eg, Group 1 and/or Group 2 Influenza A virus; eg, H1N1, H2N2, or H3N2 or H7N2 or H7N7 influenza virus</li> </ul>	<p>12. Ekiert et al, Science 2009 13. Throsby et al, PLoS One 2008 14. Sui et al, Nat Struct Mol Biol 2009 15. Ekiert et al, Science 2011</p>

Table 10A: Human IgH JH5 Variant Occurrences

HUMAN POPULATION												
NA19835	NA12814; NA12763 and 1 other(s)	HG00683; HG00590	HG01359	HG00189	HG00261; HG00136	NA18987	NA19448; NA19331	HG00740	NA20799			0.017
	NA11994	HG00595; HG00584		HG00267; HG00276	HG00137		NA19456; NA19332		NA20542	NA18934; NA19147 and 1 other(s)		0.011
			HG01494	HG00351	HG00234	NA19005		NA19719				0.005
NA19921	NA12815	HG00650		HG00177						NA18517		0.005
	NA12829; NA12890 and 1 other(s)			HG00358					NA20503	NA19093		0.005





Table 10B: Non-Synonymous Human IgH JH5 Variants

			TYT NA RAV															
	1	1	67488400000T5 IE	A/S GM DDCS LO M NOMS _ NDN														
	1	1	67488400000T5 IE	D/V GM DDCS LO M NOMS _ NDN														
	1	1	67488400000T5 IE	Q/L GM DDCS LO M NOMS _ NDN														
	1	1	67488400000T5 IE	I/T GM DDCS LO M NOMS _ NDN														
	1	1	67488400000T5 IE	A/T GM DDCS LO M NOMS _ NDN														
	1	1	67488400000T5 IE	Q/D GM DDCS LO M NOMS _ NDN														
	1	1	67488400000T5 IE	H/D GM DDCS LO M NOMS _ NDN														
	1	1	67488400000T5 IE	Q/F GM DDCS LO M NOMS _ NDN														
	1	1	67488400000T5 IE	L/F GM DDCS LO M NOMS _ NDN														
	1	1	67488400000T5 IE	Q/WGM DDCS LO M NOMS _ NDN														
	1	1	67488400000T5 IE	*S/WDE MAG POS														
	1	1	67488400000T5 IE	T/M GM DDCS LO M NOMS _ NDN														







Table 11A: Human IgH JH6 Variant Occurrences

HUMAN POPULATION													H9	
NA20289; NA20294 and 19 other(s)	NA12546; NA12046 and 41 other(s)	NA18532; NA18562 and 58 other(s)	HG00628; HG00690 and 57 other(s)	HG01456; HG01275 and 33 other(s)	HG00278; HG00313 and 57 other(s)	HG00259; HG00129 and 53 other(s)	HG01619; HG01626 and 7 other(s)	NA19083; NA18986 and 48 other(s)	NA19395; NA19321 and 20 other(s)	NA19777; NA19795 and 48 other(s)	HG00640; HG01167 and 22 other(s)	NA20798; NA20513 and 58 other(s)	NA19172; NA18909 and 8 other(s)	0.509
NA20127; NA20414 and 11 other(s)	NA12348; NA10851 and 17 other(s)	NA18558; NA18579 and 9 other(s)	HG00464; HG00683 and 16 other(s)	HG01495; HG01550 and 7 other(s)	HG00358; HG00362 and 12 other(s)	HG00135; HG00245 and 6 other(s)	HG01620; HG01625 and 1 other(s)	NA18942; NA19004 and 11 other(s)	NA19038; NA19451 and 17 other(s)	NA19725; NA19773 and 6 other(s)	HG01108; HG01072 and 12 other(s)	NA20787; NA20754 and 14 other(s)	NA19116; NA18517 and 22 other(s)	0.173
NA20314; NA20317 and 11 other(s)	NA06989; NA12044 and 10 other(s)	NA18608; NA18624 and 16 other(s)	HG00436; HG00404 and 13 other(s)	HG01357; HG01461 and 10 other(s)	HG00188; HG00282 and 10 other(s)	HG00116; HG00102 and 16 other(s)	HG01518	NA18974; NA18978 and 17 other(s)	NA19372; NA19474 and 16 other(s)	NA19758; NA19749 and 4 other(s)	HG01191; HG01048 and 7 other(s)	NA20522; NA20815 and 14 other(s)	NA18520; NA19209 and 16 other(s)	0.171
	NA12827		HG00584		HG00336									0.003
									NA19471; NA19401				NA18868	0.003
													NA19121; NA19175	0.002
	NA07346								NA19376					0.002
													NA18934; NA18856	0.002
														0.001



**Table 11B:**  
**(i) Non-Synonymous Human IgH JH6 Variants**

Variant	Position	Reference	Variant	Position	Reference	Variant	Position	Reference	Variant	Position	Reference	Variant	Position	Reference	Variant	Position	Reference
	1			1			1			1			1			1	
	1			1			1			1			1			1	
	1			1			1			1			1			1	
	1			1			1			1			1			1	
	1			1			1			1			1			1	
	1			1			1			1			1			1	
	1			1			1			1			1			1	
	1			1			1			1			1			1	
	1			1			1			1			1			1	
	1			1			1			1			1			1	









Table 12A: Human IgH JH2 Variant Occurrences

HUMAN POPULATION											2H	AV	EPYT	
	NA12812					NA19351					NA18502; NA19172		0.004	
						NA19443							0.002	
													0.002	
													0.002	
													0.001	
													0.001	
													0.001	
NA19922													0.001	

Table 12B: Non-Synonymous Human IgH JH2 Variants

Variant	W/GN DOC_S UO M NONMS	Y GM DOC_S UO M NONMS	X/X NODC_LA	RAV	4630930000TS	JE	EPYT TMA RAV	ZH
	1	1	1					
	1	1	1					
	1	1	1					

1	2	3
4	5	6
7	8	9
10	11	12

Table 13.1 - IGHA1

h.type	cum.freq	# indivs	pops
ref	0.5884	-	-
a	0.31385	556	14
b	0.05755	115	8
c	0.01455	31	8
e	0.0072	16	4
g	0.00405	9	4
d	0.00315	7	3
h	0.0018	4	3
j	0.00135	3	2
f	0.00135	3	2
i	0.00135	3	1
n	0.0009	2	2
m	0.0009	2	2
p	0.0009	2	1
q	0.0009	2	2
r	0.00045	1	1
k	0.00045	1	1
l	0.00045	1	1
o	0.00045	1	1

14:106173524	14:106173573	14:106173749	14:106174188	14:106174191	14:106174211	14:106174221	14:106174261
rs1130877	rs11546792	rs182183329	rs150528881	rs1140166	rs185815870	rs117775520	rs1407
Val->Ala	Leu->Met	Arg->His	Thr->Ala	Lys->Glu	Ala->Val	Pro->Ser	Glu->Asp
A	A	C	T	T	G	G	C
A	A	C	T	T	G	G	G
A	A	T	T	T	G	G	C
A	A	C	T	T	G	G	G
A	A	C	C	T	G	G	C
A	T	C	C	T	G	G	C
A	A	C	T	T	G	G	C
A	A	C	T	T	G	G	C
G	T	C	T	C	G	G	G
A	T	T	T	T	G	G	C
G	A	C	T	T	G	G	C
A	A	C	T	T	G	G	C
A	A	C	T	T	G	G	C
A	T	T	C	T	G	G	C
A	T	C	T	T	G	G	C
A	A	T	T	C	G	G	C
G	A	T	T	T	G	G	C
G	A	T	T	T	G	G	C
A	A	C	T	T	G	G	C

Table 13.1 - IGHA1

h.type	cum.fr	# indivs	pops	14:106174368 rs188231401 Ser->Thr	14:106174744 rs193173336 Val->Met	14:106174984 rs112706225 Lys->Glu	ENST00000390547
ref	0.5884	-	-	A	C	T	
a	0.31385	556	14	A	C	T	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.05755	115	8	A	C	T	1,5,7,10,11,12,13,14
c	0.01455	31	8	A	C	T	1,2,5,6,7,8,11,13
e	0.0072	16	4	A	C	T	1,10,12,14
g	0.00405	9	4	A	C	T	4,5,10,14
d	0.00315	7	3	A	C	C	1,10,14
h	0.0018	4	3	A	C	T	2,6,7
j	0.00135	3	2	A	C	C	10,14
f	0.00135	3	2	A	C	T	1,10
i	0.00135	3	1	A	C	C	14
n	0.0009	2	2	A	T	T	9,10
m	0.0009	2	2	A	C	T	6,9
p	0.0009	2	1	A	C	T	10
q	0.0009	2	2	A	C	C	10,12
r	0.00045	1	1	A	C	T	1
k	0.00045	1	1	A	C	T	10
l	0.00045	1	1	T	C	T	10
o	0.00045	1	1	A	C	T	4

Table 13.2 - IGHA2

h.type	cum.frq	# indivs	pops	14:106053293 rs111788346 Ala->Val	14:106053409 rs11628783 Glu->Asp	14:106053441 rs192043389 Arg->Cys	14:106053461 rs11970 Tyr->Phe	14:106053491 rs138896525 Arg->Gln	14:106053930 rs139611644 Glu->Lys	14:106053975 rs72713340 Pro->Ser	14:106053978 rs142827225 Gln->Glu
ref	0.2023	-	-	G	C	G	T	C	C	G	G
a	0.35655	528	14		G	G		C	C	G	G
b	0.1647	331	14	G	C	G	T	C	C	G	G
c	0.07815	133	3	G	C	G	T	T	C		G
g	0.03195	65	7	G	G	G	T	C	C	G	G
f	0.02645	58	13	G	C	G		C	C	G	G
i	0.0228	45	7	G	C	G		C	C	G	G
d	0.0227	48	11	G	G	G		C	C	G	G
j	0.01865	41	14	G	G	G	T	C	C	G	G
e	0.01465	31	10		C	G		C	C	G	G
k	0.01045	23	11		C	G	T	C	C	G	G
h	0.00955	20	9		G	G	T	C	C	G	G
m	0.00955	21	11	G	C	G	T	C	C	G	G
l	0.0041	9	3	G	C	G	T	T	C	G	G
p	0.0036	8	5	G	G	G	T	C	C	G	G
o	0.00315	7	3		G	G		C	C		G
n	0.0027	6	4	G	C	G		C	C		G
q	0.0018	4	2	G	C	G	T	T	C	G	G
r	0.0018	4	4		C	G	T	C	C	G	G
ai	0.0018	4	1	G	G	G	T	T	T	G	G
ag	0.00135	3	3	G	C	G	T	C	C		G
s	0.00135	3	1	G	C	G		C	C	G	G
y	0.0009	2	2		G	G	T	C	C	G	G
v	0.0009	2	2		G	G		C	C	G	G
ab	0.0009	2	2	G	C	G	T	C	C	G	G
ak	0.0009	2	2	G	C	G	T	C	C	G	G
ad	0.0009	2	2	G	C	G	T	C	C	G	G
ac	0.0009	2	2	G	C	G	T	C	T	G	G
ah	0.00045	1	1	G	C	G		C	C		G
u	0.00045	1	1	G	C	G		C	C	G	G

Table 13.2 - IGHA2

aj	0.00045	G	G	T	T	G	G	G	T	C	G	G
aa	0.00045	G	C	■	■	G	C	C	C	T	G	G
t	0.00045	■	C	■	■	G	C	C	C	C	■	G
z	0.00045	■	C	T	■	G	C	C	T	C	■	G
w	0.00045	G	G	■	■	G	C	C	C	C	■	G
af	0.00045	G	C	T	■	■	C	C	C	C	G	G
x	0.00045	■	G	■	■	G	C	C	C	C	G	C
ae	0.00045	■	G	T	■	G	C	C	C	C	■	G

Table 13.2 - IGHA2

h.type	cum.fre	# indiv	pops	14:106054166 rs182368844 Arg->Gln	14:106054428 rs9972103 Arg->ProArg->Pro	14:106054456 rs61984162 Ser->Pro	ENST00000390539
ref	0.2023	-	-	C	C	A	
a	0.35655	528	14	C	G	G	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.1647	331	14	C	G	G	1,2,3,4,5,6,7,8,9,10,11,12,13,14
c	0.07815	133	3	C	C	A	3,4,9
g	0.03195	65	7	C	C	A	1,3,5,10,11,12,14
f	0.02645	58	13	C	G	G	1,2,3,4,5,6,7,9,10,11,12,13,14
i	0.0228	45	7	C	C	A	1,4,5,9,10,12,14
d	0.0227	48	11	C	G	G	2,3,4,5,6,7,9,11,12,13,14
j	0.01865	41	14	C	G	G	1,2,3,4,5,6,7,8,9,10,11,12,13,14
e	0.01465	31	10	C	G	G	2,3,5,6,7,8,9,11,12,13
k	0.01045	23	11	C	G	G	2,3,5,6,7,8,9,10,11,12,13
h	0.00955	20	9	C	G	G	1,3,5,6,7,9,11,13,14
m	0.00955	21	11	C	C	G	1,2,3,4,7,9,10,11,12,13,14
l	0.0041	9	3	C	C	A	3,4,9
p	0.0036	8	5	C	C	G	6,7,10,11,14
o	0.00315	7	3	C	G	G	2,7,13
n	0.0027	6	4	C	G	G	2,5,6,13
q	0.0018	4	2	C	G	G	4,9
r	0.0018	4	4	C	C	A	2,5,11,13
ai	0.0018	4	1	C	C	A	14
ag	0.00135	3	3	C	G	G	1,5,7
s	0.00135	3	1	C	C	G	1
y	0.0009	2	2	C	C	A	10,14
v	0.0009	2	2	C	C	G	1,9
ab	0.0009	2	2	C	G	A	4,10
ak	0.0009	2	2	T	C	A	10,12
ad	0.0009	2	2	C	C	A	7,10
ac	0.0009	2	2	C	C	A	1,9
ah	0.00045	1	1	C	C	A	5
u	0.00045	1	1	C	G	A	5



Table 13.2 - IGHA2

aj	0.00045	1	C	C	C	A	3
aa	0.00045	1	C	C	C	A	5
t	0.00045	1	C	G	G	G	11
z	0.00045	1	T	G	G	G	3
w	0.00045	1	C	G	G	G	13
af	0.00045	1	C	C	C	A	14
x	0.00045	1	C	G	G	G	13
ae	0.00045	1	C	G	G	G	2

Table 13.3 - IGHD2-2

h.type	cum.freq	# indivs	pops	14:106382687 rs2857339	14:106382711 rs191409821	ENST00000390591
ref	0.6095	-	-	Cys->Tyr	Ile->Lys	
a	0.3896	590	14	C	A	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.0009	2	2	T	A	5,13
				C	T	

Table 13.4 - IGHD2-8

h.type	cum.freq	# indivs	pops	14:106373089 rs181619638	ENST00000390585
ref	0.99955	-	-	Tyr->Phe	
a	0.00045	1	1	T	
					2

Table 13.5 - IGHD3-10

h.type	cum.freq	# indivs	pops	rs	ENST
ref	0.9011	-	-	14:106370370	
a	0.0989	173	14	rs141575023	ENST00000390583
				Gly->Arg	1,2,3,4,5,6,7,8,9,10,11,12,13,14
				C	
				T	

Table 13.6 - IGHD3-16

h.type					14:106361515 rs143676405	
					Arg->His	ENST00000390577
ref	0.9061	-	-	14	C	
a	0.0939	155	14		T	1,2,3,4,5,6,7,8,9,10,11,12,13,14

Table 13.7 - IGHD3-3

	14:106380241				
	rs112802424				
	Arg->Gln				ENST00000390590
<b>h.type</b>		0.9766	-	6	
<b>ref</b>		0.0234	40	6	
<b>a</b>					1,7,10,12,13,14

Table 13.8 - IGHD4-23

<b>h.type</b>					14:106350740 rs145157618	ENST00000437320
<b>cum.freq</b>	0.9767	-			Arg->Trp	
<b># indivs</b>	0.0233	41	10		G	
<b>pops</b>						
<b>ref</b>						
<b>a</b>						1,2,3,5,7,10,11,12,13,14

Table 13.9 - IGHD4-4

h.type	cum.freq	# indivs	pops	14:106379089 rs190861327	Gln->Arg	ENST000000414852
ref	0.9973	-	-	T		
a	0.0027	5	4	<b>C</b>		1,4,9,10



Table 13.10 - IGHD6-13

h.type	cum.freq	# indvs	pops	14:106367013	rs185385671	ENST00000390580
ref	0.9982	-	-	Ser->Asn		
a	0.0018	4	4	C		2,3,5,6
				T		

Table 13.11 - IGH D7-27

<b>h.type</b>						
ref	0.99955	-	-	Thr->Ser	14:106331767	
a	0.00045	1	1	G	rs187067877	ENST00000439842
				<b>C</b>		

Table 13.12 - IGHD

h.type	cum.freq	# indivs	pops	14:106303836 rs146595719	14:106303846 rs1005582	14:106306841 rs709589	14:106307012 rs184889907	14:106307300 rs139530086	14:106307432 rs111902251	14:106311758 rs188179725	14:106311961 rs149054178
ref	0.003	-	-	Thr->Met	Ala->Thr	Gly->Arg	Val->Ile	Val->Ile	Thr->Ser	Val->Glu	His->Gln
a	0.735	951	14	G	C	C	C	C	T	A	G
b	0.225	369	14	G	T	C	C	C	T	A	G
e	0.017	32	5	G	T	T	C	C	T	A	G
c	0.013	24	4	G	T	C	C	C	■	A	G
d	0.005	11	3	G	T	C	C	T	T	A	G
f	0.002	5	3	G	T	C	T	C	T	A	G
g	5E-04	1	1	■	T	T	C	C	T	A	G
h	5E-04	1	1	G	T	T	C	C	T	T	G

Table 13.12 - IGHD

h.type	cum.freq	# indivs	pops	
ref	0.003	-	-	ENST00000390556
a	0.735	951	14	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.225	369	14	1,2,3,4,5,6,7,8,9,10,11,12,13,14
e	0.017	32	5	1,5,10,12,14
c	0.013	24	4	1,5,11,12
d	0.005	11	3	3,4,9
f	0.002	5	3	2,6,7
g	5E-04	1	1	13
h	5E-04	1	1	6

Table 13.13 - IGHEP1

				14:106188166	
				rs142072061	
				Val->Ile	ENST00000558023
ref	0.99775	-		C	
a	0.00225	5	4	T	2,6,7,9
<b>h.type</b>					
<b>cum.freq</b>					
<b># indivs</b>					
<b>pops</b>					

Table 13.14 - IGHE

h.type	cum.freq	# indivs	pops	14:106066524 rs189365488	14:106066688 rs148738013	14:106066690 rs61675149	14:106066701 rs57724657	14:106067096 rs182957708	14:106067118 rs150823665	14:106067815 rs144154566	
ref	0.87655	-	-	Arg->His	Arg->His	Trp->Gly	Ala->Val	Pro->Leu	Ala->Thr	Thr->Ile	ENST00000390541
a	0.08375	144		C	C	A	G	G	C	G	3,4,5,9,10,11,14
b	0.03205	66		C	C	C	G	G	C	G	1,5,10,12,14
d	0.00225	5		C	C	C	■	G	T	G	10,14
c	0.00135	3		T	C	A	G	G	C	G	1,10
h	0.00135	3		C	C	A	G	G	T	G	14
g	0.0009	2		C	T	A	G	G	C	G	7,13
e	0.0009	2		C	C	C	G	■	C	G	3,4
f	0.0009	2		C	C	A	G	G	C	■	6,13

Table 13.15 - IGHG1

h.type	cum.freq	# indivs	pops	14:106204113	14:106204131	14:106208326	14:106208411	ENST00000390542
ref	0.89185	-	-	rs117518546	rs138262373	rs1043109	rs184594692	ENST00000390548
a	0.09825	158	4	Gly->Arg	Asp->Asn	Leu->Val	Asp->Glu	ENST00000390549
b	0.0063	14	6	T	T	G	G	
d	0.0018	4	2	C	C	C	G	
c	0.0018	4	1	C	C	G	T	

3,4,9,13  
2,5,6,7,11,13  
3,9  
14

Table 13.16 - IGHG2

h.type	cum.freq	# indivs	pops	14:106109830 rs185308159	14:106110017 rs182705016	14:106110056 rs113678609	14:106110137 rs8009156	14:106110209 rs190914111	14:106110914 rs11627594	14:106110926 rs139934867	
ref	0.6966	-	-	Pro->Thr	Lys->Glu	Val->Leu	Val->Met	Glu->Lys	Pro->Thr	Val->Met	ENST00000390545
a	0.2868	499	12	G	T	C	C	C	G	C	
b	0.0045	9	1	G	T	C	T	C	T	C	1,2,3,4,5,6,7,8,9,11,12,13
f	0.0036	8	3	T	T	C	C	C	G	T	10
h	0.0027	6	6	G	T	C	C	C	T	C	7,11,13
d	0.0023	5	3	G	T	G	C	C	G	C	1,6,7,9,10,13
e	0.0018	4	3	G	T	C	C	T	G	C	3,4,9
c	0.0009	2	2	T	T	C	T	C	G	C	2,4,5
g	0.0005	1	1	G	C	C	C	C	G	C	1,10
i	0.0005	1	1	G	T	C	C	T	T	C	10
											4



Table 13.17 - IGHG3

h.type	cum.fre	# indivs	pop	14:106235611	14:106235663	14:106235692	14:106235729	14:106235742	14:106235783	14:106236000	14:106236128
ref				rs1051112	rs190174000	rs187517394	rs139413052	rs149653267	rs113169458	rs141959627	rs12890621
				Phe->Tyr	Gln->Glu	Lys->Arg	Met->Val	Asn->Lys	Val->Met	Thr->Ala	Tyr->Phe
a	0.2495	-	-	A	G	T	T	G	C	T	T
b	0.2813	479	14	T	G	T	T	G	C	T	T
j	0.19955	345	11	A	G	T	T	G	C	T	■
d	0.02345	50	8	A	G	T	T	C	C	T	T
c	0.02095	44	6	T	G	T	C	C	T	T	T
i	0.0209	46	10	T	G	T	T	G	C	T	■
g	0.019	41	10	T	G	T	T	G	C	T	T
f	0.01735	38	10	T	G	T	C	C	C	T	T
h	0.01725	36	7	T	G	T	T	G	C	C	T
e	0.0164	36	11	T	G	T	T	C	C	T	T
k	0.015	33	9	A	G	T	T	G	C	T	T
t	0.01315	27	5	A	C	C	C	G	C	T	T
q	0.0126	28	6	T	G	T	T	C	C	T	T
v	0.0126	27	3	A	G	T	C	G	C	T	T
o	0.0072	16	8	T	G	T	C	G	C	T	T
r	0.00675	15	7	A	G	T	T	C	C	T	■
m	0.0063	14	5	T	G	T	C	C	C	T	T
n	0.00585	13	5	A	C	T	T	C	C	T	T
am	0.00495	11	4	T	G	T	T	C	T	T	T
l	0.0045	10	5	T	G	T	C	C	T	T	T
p	0.0036	8	6	T	G	T	T	C	C	T	■
u	0.0036	8	2	A	C	T	T	G	C	T	T
x	0.0036	8	4	T	G	T	C	C	C	T	■
ae	0.00315	7	3	T	C	T	C	C	C	T	■
ai	0.00225	5	3	T	C	C	C	G	C	T	T
aj	0.00225	5	2	T	G	T	T	C	T	T	T
w	0.0018	4	2	A	G	T	C	C	C	T	T
bd	0.0018	4	1	T	G	T	T	G	C	T	T
al	0.00135	3	1	T	G	T	C	C	C	T	T
	0.00135	3	2	A	G	T	C	C	T	T	T

Table 13.17 - IGHG3

s	0.00135	3	T	C	T	T	T	T	G	C	T	T	■
ab	0.00135	3	A	G	T	T	T	T	G	C	T	T	T
ak	0.00135	3	A	G	T	T	T	T	C	T	T	T	T
ag	0.0009	2	T	G	T	T	T	T	G	C	T	T	T
y	0.0009	2	A	G	T	T	T	T	G	C	T	■	■
az	0.0009	2	T	G	T	T	T	T	G	T	T	■	■
aa	0.0009	2	A	G	T	T	T	T	C	C	T	T	■
ap	0.0009	2	A	C	T	T	T	T	G	C	T	T	■
av	0.0009	2	T	G	T	T	T	T	C	C	T	T	T
af	0.0009	2	T	C	C	C	C	C	C	C	T	T	T
at	0.0009	2	T	G	C	C	C	C	G	C	T	T	T
aw	0.0009	2	T	G	C	C	C	T	C	C	C	T	T
ay	0.0009	2	A	G	T	T	T	T	G	C	T	■	■
ar	0.0009	2	A	G	C	C	C	C	G	C	T	T	T
an	0.0009	2	T	G	T	T	T	T	C	C	T	T	T
ah	0.00045	1	A	G	T	T	T	T	G	C	T	T	T
ba	0.00045	1	A	G	T	T	T	T	G	C	C	■	■
ax	0.00045	1	A	C	T	T	T	T	C	C	T	T	T
bb	0.00045	1	T	G	C	C	C	C	C	C	T	T	T
aq	0.00045	1	A	G	T	T	T	T	G	C	C	T	T
be	0.00045	1	T	G	C	C	C	T	G	C	T	T	T
z	0.00045	1	A	G	T	T	T	T	G	C	T	■	■
ao	0.00045	1	T	C	C	C	C	T	C	C	T	T	T
au	0.00045	1	T	G	T	T	T	T	G	T	T	T	T
ad	0.00045	1	T	C	T	T	T	T	C	T	T	T	T
bc	0.00045	1	A	G	T	T	T	T	C	T	T	T	T
ac	0.00045	1	T	C	T	T	T	T	C	T	T	■	■
as	0.00045	1	A	G	T	T	T	T	G	T	T	T	T

Table 13.17 - IGHG3

h.type	cum.fre	# indivs	pops	14:106236141 rs60746425 Arg->Trp	14:106236143 rs74093865 Pro->Leu	14:106236174 rs184265224 Gly->Ser	14:106236187 rs141238286 Lys->Asn	14:106236195 rs145035200 Gln->Lys	14:106236315 rs189074626 Leu->Phe	14:106237516 rs186135943 Leu->Phe
ref	0.2495	-	-	G	G	C	C	G	G	C
a	0.2813	479	14	G		C	C	G	G	C
b	0.19955	345	11	G	G	C	C	G	G	C
j	0.02345	50	8	G	G	C	C	G	G	C
d	0.02095	44	6		G	C	C	G	G	C
c	0.0209	46	10	G	G	C	C	G	G	C
i	0.019	41	10	G	G	C	C	G	G	C
g	0.01735	38	10	G		C	C	G	G	C
f	0.01725	36	7	G		C	C	G	G	C
h	0.0164	36	11	G		C	C	G	G	C
e	0.015	33	9	G		C	C	G	G	C
k	0.01315	27	5	G	G	C	C	G	G	C
t	0.0126	28	6	G	G	C	C	G	G	C
q	0.0126	27	3	G	G	C	C	G	G	C
v	0.0072	16	8	G		C	C	G	G	C
o	0.00675	15	7	G	G	C	C	G	G	C
r	0.0063	14	5	G	G	C	C	G	G	C
m	0.00585	13	5	G	G	C	C	G	G	C
n	0.00495	11	4		G	C	C	G	G	C
am	0.0045	10	5	G	G	C	C	G	G	G
l	0.0036	8	6	G	G	C	C	G	G	C
p	0.0036	8	2	G	G	C	C	G	G	C
u	0.0036	8	4	G	G	C	C	G	G	C
x	0.00315	7	3	G	G	C	C	G	G	C
ae	0.00225	5	3	G	G	C	C	G	G	C
ai	0.00225	5	2	G	G	C	C	G	G	G
aj	0.0018	4	2	G	G	C	C	G	G	C
w	0.0018	4	1	G		C	C	T	G	C
bd	0.00135	3	1	G		C	C	T	G	C
al	0.00135	3	2	G	G	C	C	G	G	G

Table 13.17 - IGHG3

s	0.00135	3	G	G	C	C	C	C	G	C	G	C	C
ab	0.00135	3	G	G	C	C	C	C	G	G	G	G	C
ak	0.00135	3	G	G	C	C	C	C	G	G	G	G	C
ag	0.0009	2	G	G	C	C	C	C	G	G	G	G	C
y	0.0009	2	G	G	T	C	C	C	G	G	G	G	C
az	0.0009	2	G	G	C	C	C	C	G	G	G	G	C
aa	0.0009	2	G	G	C	C	C	C	G	G	G	G	C
ap	0.0009	2	G	G	C	C	C	C	G	G	G	G	C
av	0.0009	2	G	G	C	C	C	C	G	T	G	G	C
af	0.0009	2	G	G	C	C	C	C	G	G	G	G	C
at	0.0009	2	G	G	C	C	C	C	G	G	G	G	C
aw	0.0009	2	G	G	C	C	C	C	G	G	G	G	C
ay	0.0009	2	G	G	C	C	C	C	G	G	G	G	C
ar	0.0009	2	G	G	C	C	C	C	G	G	G	G	C
an	0.0009	2	G	G	C	C	C	C	G	G	G	G	C
ah	0.00045	1	G	G	C	C	C	C	G	G	G	G	C
ba	0.00045	1	G	G	C	C	C	C	G	G	G	G	C
ax	0.00045	1	G	G	C	C	C	C	G	G	G	G	C
bb	0.00045	1	G	G	C	C	C	C	G	G	G	G	C
aq	0.00045	1	G	G	C	C	C	C	G	G	G	G	C
be	0.00045	1	G	G	C	C	C	C	G	G	G	G	C
z	0.00045	1	G	G	C	C	C	C	G	G	G	G	C
ao	0.00045	1	G	G	C	C	C	C	G	G	G	G	C
au	0.00045	1	G	G	C	C	C	C	G	G	G	G	C
ad	0.00045	1	G	G	C	C	C	C	G	G	G	G	C
bc	0.00045	1	G	G	C	C	C	C	G	G	G	G	C
ac	0.00045	1	G	G	C	C	C	C	G	G	G	G	C
as	0.00045	1	G	G	C	C	C	C	G	G	G	G	C

Table 13.17 - IGHG3

h.type	cum.fir	# indivs	pops	ref
	0.2495	-	-	ENST00000390551
a	0.2813	479	14	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.19955	345	11	1,2,3,4,5,6,7,8,11,12,13
j	0.02345	50	8	1,2,3,4,5,10,12,14
d	0.02095	44	6	1,3,4,8,9,11
c	0.0209	46	10	1,2,3,5,6,7,8,11,12,13
i	0.019	41	10	1,2,3,4,7,8,10,12,13,14
g	0.01735	38	10	2,3,4,5,6,7,9,11,12,13
f	0.01725	36	7	1,2,5,6,7,11,13
h	0.0164	36	11	2,3,4,5,6,7,8,9,11,12,13
e	0.015	33	9	2,4,5,6,7,9,11,12,13
k	0.01315	27	5	1,5,10,12,14
t	0.0126	28	6	1,2,5,10,13,14
q	0.0126	27	3	1,10,14
v	0.0072	16	8	2,3,5,6,7,9,11,12
o	0.00675	15	7	2,5,7,8,11,12,13
r	0.0063	14	5	1,4,10,12,14
m	0.00585	13	5	1,5,10,12,14
n	0.00495	11	4	3,4,9,11
am	0.0045	10	5	1,5,10,12,14
l	0.0036	8	6	2,6,7,8,12,13
p	0.0036	8	2	1,10
u	0.0036	8	4	1,2,6,11
x	0.00315	7	3	2,7,13
ae	0.00225	5	3	1,10,14
ai	0.00225	5	2	1,14
aj	0.0018	4	2	10,14
w	0.0018	4	1	4
bd	0.00135	3	1	9
al	0.00135	3	2	1,14

**Table 13.17 - IGHG3**

s	0.00135	3	2 7,11
ab	0.00135	3	2 1,14
ak	0.00135	3	2 10,14
ag	0.0009	2	2 1,10
y	0.0009	2	2 2,7
az	0.0009	2	2 2,13
aa	0.0009	2	2 2,6
ap	0.0009	2	2 10,14
av	0.0009	2	1 9
af	0.0009	2	1 14
at	0.0009	2	2 10,14
aw	0.0009	2	2 5,9
ay	0.0009	2	1 11
ar	0.0009	2	2 5,14
an	0.0009	2	1 9
ah	0.00045	1	1 10
ba	0.00045	1	1 13
ax	0.00045	1	1 10
bb	0.00045	1	1 1
aq	0.00045	1	1 5
be	0.00045	1	1 7
z	0.00045	1	1 7
ao	0.00045	1	1 14
au	0.00045	1	1 5
ad	0.00045	1	1 1
bc	0.00045	1	1 1
ac	0.00045	1	1 13
as	0.00045	1	1 4

Table 13.18 - IGHG4

h.type	cum.freq	# indivs	pops	14:106090959 rs190355823	14:106091083 rs146589796	14:106091281 rs112773794	14:106091329 rs8015545	14:106091332 rs138186657	14:106091341 rs143666134	14:106091347 rs145943426	14:106091450 rs138777601
ref	0.77355	-	-	Ser->Ala	Met->Ile	Asn->His	Leu->Val	Val->Ile	Val->Ile	Val->Ile	Gln->His
a	0.19825	389	12	A	C	T	G	C	C	C	C
b	0.01005	22	4	A	C	T	C	C	C	C	C
c	0.0055	12	4	A	T	T	G	C	C	C	C
d	0.0041	8	4	A	C	G	G	T	C	C	C
f	0.0027	6	3	A	C	T	G	C	T	C	C
e	0.00225	4	2	A	C	T	G	C	C	C	G
g	0.0018	4	2	A	C	T	G	C	C	C	C
j	0.00045	1	1	A	C	G	G	C	C	C	C
k	0.00045	1	1	C	C	T	G	C	C	C	C
h	0.00045	1	1	A	C	T	G	C	C	T	C
i	0.00045	1	1	A	C	T	C	C	C	C	G

Table 13.18 - IGHG4

h.type	cum.freq	# indivs	pops	14:106091714 rs139754373	14:106092363 rs183713766	ENST00000390543
ref	0.77355	-	-	A	C	
a	0.19825	389	12	A	C	1,2,3,4,5,6,7,8,9,11,12,13
b	0.01005	22	4	<b>G</b>	C	1,10,12,14
c	0.0055	12	4	A	C	1,5,10,14
d	0.0041	8	4	A	C	1,5,10,14
f	0.0027	6	3	A	C	1,10,14
e	0.00225	4	2	A	C	10,11
g	0.0018	4	2	A	<b>T</b>	1,14
j	0.00045	1	1	A	C	1
k	0.00045	1	1	A	C	1
h	0.00045	1	1	A	C	10
i	0.00045	1	1	A	C	4



Table 13.19 - IGHJ1

h.type	cum.fre	# indivs	pops	14:106331628 rs181379404 Thr->Ser	ENST00000390565
ref	0.9982	-	-	G	
a	0.0018	3	3	<b>C</b>	2,3,13

Table 13.20 - IGHJ2

h.type	cum.fre	# indivs	pops	14:106331453 rs181475274	14:106331455 rs185806009	ENST00000390564
ref	0.9936	-	-	Tyr->Phe T	Trp->Cys C	
a	0.00415	9	6	T	<b>G</b>	1,3,4,7,10,13
b	0.00225	4	3	■	C	1,10,14

Table 13.21 - IGHJ3

h.type	cum.freq	# indivs	pops	14:106330799 rs190137342	14:106330814 rs181332275	14:106330815 rs185836312	14:106330833 rs190698203	14:106330841 rs183030857	
ref	0.95755	-	-	Ser->Leu	Met->Thr	Met->Val	Ile->Phe	Ala->Val	ENST00000463911
a	0.01645	34	12	G	A	T	T	G	1,2,3,4,5,6,7,9,10,12,13,14
b	0.0114	23	10	G	A	T	T	■	2,3,4,6,7,9,10,11,13,14
c	0.0064	14	11	G	A	<b>C</b>	T	G	1,2,5,6,7,8,9,10,11,13,14
d	0.0046	9	7	G	A	T	■	G	5,7,9,10,11,13,14
e	0.00315	6	5	■	<b>G</b>	T	T	G	1,2,3,5,10
f	0.00045	1	1	G	A	<b>C</b>	T	■	5

Table 13.22 - IGHJ4

h.type	cum.freq	# indivs	pops	14:106330436 rs145372686	14:106330442 rs183645858	14:106330460 rs150237274	14:106330467 rs189135736	
ref	0.9758	-	-	Thr->Ser G	Leu->Arg A	Tyr->Phe T	Phe->Ile A	ENST00000461719
a	0.01785	39	11	G	A	■	A	2,3,4,6,7,9,10,11,12,13,14
b	0.0032	7	5	G	A	T	T	1,2,9,11,13
c	0.0018	4	4	G	<b>C</b>	T	A	3,4,13,14
d	0.00135	3	3	<b>C</b>	A	T	A	2,4,10

Table 13.23 - IGHJ5

h.type	cum.freq	# indivs	pops	14:106330027 rs147551384	14:106330035 rs141950820	14:106330067 rs78872461	ENST00000488476
ref	0.9497	-	-	Ser->Ala A	Thr->Ile G	Trp->Cys C	
a	0.0357	78	13	A	G	G	1,2,3,4,5,6,7,9,10,11,12,13,14
b	0.011	24	10	A		C	1,2,3,5,6,7,8,9,10,11
c	0.0027	5	4	C	G	C	2,3,7,13
d	0.0009	2	2	A		G	4,14

Table 13.24 - IGHJ6

h.type	cum.freq	# indivs	pops	14:106329414 rs111729691	14:106329417 rs142191790	14:106329434 rs140411866	14:106329435 rs1950943	ENST00000390560
ref	0.3083	-	-	Ser->Pro A	Val->Ile C	Lys->Arg T	Lys->Gln T	
a	0.6845	945	14	A	C	T	G	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.00315	7	2	A	T	T	T	10,14
e	0.0009	2	2	A	C	C	G	4,13
h	0.0009	2	2	G	C	T	G	2,6
f	0.0009	2	2	A	C	C	T	3,10
d	0.00045	1	1	A	T	T	G	9
g	0.00045	1	1	G	C	C	G	4
c	0.00045	1	1	G	C	T	T	10

Table 13.25 - IGHM

h.type	cum.freq	# indivs	pops	14:106320732	14:106321313	14:106321330	14:106321590	14:106321663	14:106322093
ref	0.0249	-	-	Asp->Asn	Ala->Thr	Thr->Ile	Gly->Val	Ser->Gly	Asp->Glu
a	0.7527	974	14	C	C	G	C	T	G
b	0.15245	287	14	C	C	G	■	T	G
d	0.03795	74	9	C	C	G	C	C	G
c	0.02705	54	3	C	C	■	■	C	C
e	0.0018	4	2	C	C	G	C	C	G
g	0.00135	3	3	C	T	G	C	T	G
f	0.0009	2	1	C	C	G	C	T	C
h	0.00045	1	1	C	T	G	■	T	G
i	0.00045	1	1	T	C	G	C	C	C
									ENST00000390559
									1,2,3,4,5,6,7,8,9,10,11,12,13,14
									1,2,3,4,5,6,7,8,9,10,11,12,13,14
									1,2,5,7,10,11,12,13,14
									3,4,9
									10,14
									5,12,14
									11
									5
									10

Table 13.26 - IGHV1-18

h.type	cum.freq	# indivs	pops	14:106641643 rs181586608	14:106641660 rs146311654	14:106641668 rs184544516	14:106641688 rs114910155	14:106641769 rs139305030	14:106641801 rs188509150	14:106641832 rs181324183	14:106641981 rs184535212
				Thr->Ala	Gly->Asp	Lys->Asn	Asn->Asp	Thr->Pro	Lys->Arg	Ala->Pro	Ser->Asn
ref	0.97595	-	-	T	C	C	T	T	T	C	C
a	0.01185	23	6	T	C	C	C	T	T	C	C
c	0.0045	7	4	T	T	C	T	T	T	C	C
b	0.00275	6	5	T	C	G	T	T	T	C	C
d	0.00135	3	3	T	C	C	T	T	T	C	C
e	0.00135	3	3	T	C	C	T	G	T	C	C
f	0.0009	2	2	T	C	C	T	T	T	C	T
g	0.00045	1	1	C	T	C	T	T	T	C	C
h	0.00045	1	1	T	C	C	T	T	T	G	C
i	0.00045	1	1	C	C	C	T	T	T	C	C



Table 13.26 - IGHV1-18

h.type	cum.freq	# indivs	pops	ENST00000390605
ref	0.97595	-	-	
a	0.01185	23	6	1,6,9,10,11,14
c	0.0045	7	4	1,10,11,14
b	0.00275	6	5	6,10,11,13,14
d	0.00135	3	3	4,11,14
e	0.00135	3	3	2,6,11
f	0.0009	2	2	8,9
g	0.00045	1	1	13
h	0.00045	1	1	10
i	0.00045	1	1	10

Table 13.27 - IGHV1-24

h.type	cum.freq	# indivs	pops	14:106733154 rs141629050	14:106733158 rs150510876	14:106733166 rs139845206	14:106733192 rs141655646	14:106733263 rs183704154	14:106733269 rs114334184	14:106733274 rs187836940	14:106733336 rs139283249
ref	0.97175	-	-	Tyr->Phe	Tyr->His	Thr->Met	Glu->Asp	Ile->Phe	Glu->Lys	Asp->Gly	Met->Ile
a	0.01185	24		T	A	G	C	T	C	T	C
b	0.0082	18		T	A	G	C	T	C	T	C
c	0.00415	9		T	A	G	<b>G</b>	T	C	T	C
g	0.0009	2		T	A	G	C	T	C	T	<b>T</b>
d	0.00045	1		T	<b>G</b>	■	C	T	C	T	C
j	0.00045	1		T	A	G	C	T	C	T	C
k	0.00045	1		T	A	G	C	T	C	<b>C</b>	C
e	0.00045	1		T	A	G	C	■	C	T	C
h	0.00045	1		T	A	G	C	T	<b>T</b>	T	C
f	0.00045	1		T	A	G	C	T	C	T	C
i	0.00045	1		■	A	G	C	T	C	T	C

Table 13.27 - IGHV1-24

h.type	cum.freq	# indivs	pops	14:106733337 rs8012805	14:106733341 rs186604864	14:106733359 rs184694183	14:106733446 rs147700118	14:106733570 rs192824042	ENST00000390610
ref	0.97175	-	-	Met->Arg A	Ser->Pro A	Tyr->Asn A	Thr->AlaThr->Ala T	Thr->Ala T	
a	0.01185	24	4	<b>C</b>	A	A	T	T	1,10,11,14
b	0.0082	18	4	A	A	A	<b>C</b>	T	1,10,12,14
c	0.00415	9	3	A	A	A	T	T	1,10,14
g	0.0009	2	1	A	A	A	T	T	2
d	0.00045	1	1	A	A	A	T	T	14
j	0.00045	1	1	A	A	A	T	<b>C</b>	5
k	0.00045	1	1	A	<b>G</b>	<b>T</b>	<b>C</b>	T	5
e	0.00045	1	1	A	<b>G</b>	A	T	T	5
h	0.00045	1	1	A	A	A	T	T	14
f	0.00045	1	1	A	A	<b>T</b>	T	T	13
i	0.00045	1	1	A	A	A	T	T	3

Table 13.28 - IGHV1-2

h.type	cum.freq	# indivs	pops	14:106452690 rs12588974	14:106452735 rs188093056	14:106452766 rs112806369	14:106452784 rs180944534	14:106452793 rs186314384	14:106452799 rs189402450	14:106452817 rs1065059	14:106452868 rs192043188
ref	0.43185	-	-	Ala->Val	Ser->Asn	Arg->Trp	Ala->Ser	Thr->Ser	Gly->Ser	Trp->Arg	Tyr->Asn
a	0.3268	511	14	G	C	T	C	T	C	A	A
b	0.1304	228	14	G	C	■	C	T	C	A	A
c	0.0797	158	14	■	C	T	C	T	C	G	A
d	0.01135	25	10	G	C	T	C	T	C	A	A
e	0.0073	16	6	G	C	T	C	T	C	A	A
g	0.0018	4	3	G	T	T	C	T	C	A	A
h	0.00135	3	2	■	C	■	C	T	C	A	A
f	0.00135	3	3	G	C	■	C	T	C	A	A
i	0.0009	2	2	■	C	T	C	T	C	G	A
m	0.0009	2	2	G	C	T	C	T	C	G	A
l	0.0009	2	2	G	C	T	C	T	C	A	A
j	0.00045	1	1	G	C	T	■	T	C	A	A
u	0.00045	1	1	G	T	T	C	T	C	A	A
k	0.00045	1	1	G	C	T	C	T	C	A	A
t	0.00045	1	1	G	C	T	C	T	C	G	A
v	0.00045	1	1	G	C	T	C	T	C	A	T
s	0.00045	1	1	G	T	T	C	■	C	A	A
q	0.00045	1	1	G	C	T	C	T	T	A	A
w	0.00045	1	1	G	C	■	C	T	C	A	A
r	0.00045	1	1	G	C	T	C	T	C	A	A
n	0.00045	1	1	G	C	T	C	T	C	A	A
p	0.00045	1	1	G	T	T	C	T	C	G	A
o	0.00045	1	1	G	C	T	C	T	C	A	A

Table 13.28 - IGHV1-2

h.type	cum.freq	# indivs	pops	14:106452873 rs1065058	14:106452882 rs11553010	14:106452897 rs185487229	14:106452903 rs189604828	14:106452956 rs182256840	14:106452972 rs186783545	
ref	0.43185	-	-	Gly->Asp	Thr->Ser	Lys->Met	Ser->Cys	Gln->His	Ala->Gly	ENST00000390594
a	0.3268	511	14	C	G	T	G	C	G	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.1304	228	14	C	G	T	G	C	G	1,2,3,4,5,6,7,8,9,10,11,12,13,14
c	0.0797	158	14	C	G	T	G	C	G	1,2,3,4,5,6,7,8,9,10,11,12,13,14
d	0.01135	25	10	C	G	T	G	C	G	1,2,3,4,5,9,10,11,13,14
e	0.0073	16	6	T	G	T	G	C	G	1,2,5,10,12,14
g	0.0018	4	3	C	G	T	G	C	G	6,12,14
h	0.00135	3	2	C	G	T	G	C	G	3,4
f	0.00135	3	3	T	G	T	G	C	G	2,6,7
i	0.0009	2	2	T	G	T	G	C	G	11,13
m	0.0009	2	2	T	G	T	G	C	G	3,6
l	0.0009	2	2	C	C	T	G	C	G	9,10
j	0.00045	1	1	C	G	■	G	C	G	2
u	0.00045	1	1	T	G	T	G	C	G	2
k	0.00045	1	1	C	C	T	G	C	G	2
t	0.00045	1	1	C	G	T	G	C	G	2
v	0.00045	1	1	T	G	■	G	C	G	10
s	0.00045	1	1	C	G	T	G	C	G	2
q	0.00045	1	1	C	G	T	G	C	G	10
w	0.00045	1	1	C	G	T	C	C	G	11
r	0.00045	1	1	C	G	T	G	C	C	2
n	0.00045	1	1	T	G	T	G	G	G	1
p	0.00045	1	1	C	G	T	G	C	G	1
o	0.00045	1	1	C	G	T	G	G	G	10

Table 13.29 - IGHV1-3

h.type	cum.freq	# indivs	pops	14:106471253 rs190814253	14:106471268 rs1143505	14:106471280 rs141136074	14:106471296 rs113948114	14:106471310 rs191691305	14:106471353 rs34069216	14:106471361 rs187612074	14:106471365 rs191777242
ref	0.34155	-	-	Cys->Tyr	Met->Thr	Arg->Thr	Glu->Gln	Ser->Asn	Glu->Lys	Tyr->Cys	Lys->Gln
a	0.3564	555	14	C	A	C	C	C	C	T	T
b	0.0724	155	13	C	G	C	C	C	T	T	T
d	0.0439	97	13	T	A	G	C	C	C	T	T
c	0.02865	63	12	C	G	C	C	C	T	T	T
h	0.019	42	12	C	A	C	C	C	T	T	T
i	0.0113	25	11	C	A	C	C	C	T	T	T
g	0.0105	23	8	C	G	C	G	C	T	T	T
f	0.00995	22	9	C	A	C	C	C	T	T	T
j	0.0077	16	8	C	A	C	C	C	C	T	T
e	0.0073	16	4	T	A	C	C	C	C	T	T
n	0.0072	16	10	C	G	C	C	C	C	T	T
t	0.00675	15	8	T	A	G	C	C	T	T	T
r	0.00585	13	9	C	G	C	C	C	T	T	T
m	0.00585	13	6	T	A	G	C	C	T	T	T
l	0.00585	13	8	C	A	C	C	C	C	T	T
o	0.00495	11	5	C	G	C	C	C	T	T	T
p	0.0045	10	4	T	A	G	C	C	C	T	T
ab	0.00405	9	6	C	A	C	C	C	T	T	T
k	0.0036	8	5	C	G	C	C	C	T	T	T
x	0.00315	7	6	C	A	C	G	C	C	T	T
ah	0.0027	6	5	C	A	C	G	C	T	T	T
u	0.00225	5	2	C	G	C	C	C	T	T	T
v	0.00225	5	2	T	A	G	C	C	T	T	T
ap	0.00225	4	3	C	G	C	C	C	T	T	T
ac	0.00225	5	4	T	A	G	C	C	C	T	T
y	0.0018	4	3	C	A	G	C	C	C	T	T
ar	0.0018	4	3	C	G	C	C	C	T	T	T
ai	0.0018	4	3	C	A	C	G	C	T	T	T

Table 13.29 - IGHV1-3

aa	0.00135	3	C	G	C	C	C	C	T	T	T	T
ak	0.00135	3	C	A	C	C	C	C	C	C	T	T
w	0.00135	3	C	G	C	C	C	C	C	C	C	T
at	0.00135	3	T	A	G	C	C	C	C	T	T	T
ag	0.0009	2	C	A	G	C	C	C	C	T	T	T
aj	0.0009	2	T	G	C	C	C	C	C	T	T	T
aq	0.0009	2	T	A	G	C	C	C	C	T	T	T
s	0.0009	2	T	A	G	C	C	C	C	T	T	T
q	0.0009	2	C	G	C	C	C	C	C	T	T	T
z	0.0009	2	C	G	C	C	C	C	C	T	T	T
ad	0.0009	2	C	G	C	C	C	C	C	T	T	T
aw	0.0009	2	C	A	C	C	C	C	C	T	T	T
ae	0.0009	2	C	G	C	C	C	C	C	T	T	T
an	0.0009	2	T	A	C	C	C	C	C	T	T	T
bd	0.00045	1	C	G	C	C	C	C	C	T	T	T
ba	0.00045	1	C	G	C	C	C	C	C	T	T	T
ax	0.00045	1	C	A	C	C	C	C	C	T	T	G
bb	0.00045	1	C	G	C	C	C	C	C	T	T	G
az	0.00045	1	C	G	C	C	C	C	C	T	T	T
al	0.00045	1	T	A	G	C	C	C	C	T	T	T
be	0.00045	1	C	G	C	C	C	C	C	T	T	T
bf	0.00045	1	C	A	C	C	C	C	C	T	T	T
av	0.00045	1	C	A	C	C	C	C	C	C	C	G
ao	0.00045	1	T	A	G	C	C	C	C	T	T	T
af	0.00045	1	C	A	C	C	C	C	C	T	T	T
au	0.00045	1	C	G	C	C	C	C	C	T	T	T
bh	0.00045	1	T	A	C	C	C	C	C	T	T	T
am	0.00045	1	C	A	C	C	C	C	T	T	T	T
bc	0.00045	1	C	G	C	C	C	C	C	T	T	T
bg	0.00045	1	T	A	G	C	C	C	C	T	T	G
ay	0.00045	1	C	G	C	C	C	C	T	T	T	T
as	0.00045	1	T	A	G	C	C	C	C	T	T	G

Table 13.29 - IGHV1-3

h.type	cum.freq	# indvs	pops	14:106471383 rs184320740	14:106471388 rs34874585	14:106471421 rs74091631	14:106471503 rs186640498	14:106471508 rs192340916	14:106471547 rs1143496	
				Ala->Thr	Ser->Ile	Ala->Asp	Lys->Gln	Val->Glu	Val->Ala	
ref	0.34155	-	-	C	C	G	T	A	A	ENST00000390595
a	0.3564	555	14	C		G	T	A	A	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.0724	155	13	C	C	G	T	A	G	1,2,3,4,5,6,7,9,10,11,12,13,14
d	0.0439	97	13	C		G	T	A	A	1,2,3,4,5,6,7,9,10,11,12,13,14
c	0.02865	63	12	C		G	T	A	G	1,2,3,4,5,6,7,9,11,12,13,14
h	0.019	42	12	C	C	G	T	A	A	1,2,3,4,5,6,7,9,10,11,13,14
i	0.0113	25	11	C		G	T	A	G	1,2,3,5,6,7,10,11,12,13,14
g	0.0105	23	8	C		G	T	A	G	2,3,4,6,7,9,11,13
f	0.00995	22	9	C		G	T	A	A	2,3,5,7,8,9,11,13,14
j	0.0077	16	8	C		G	T	A	A	1,4,5,6,7,10,12,14
e	0.0073	16	4	C	C	G	T	A	A	1,10,12,14
n	0.0072	16	10	C	C	G	T	A	A	2,4,5,6,7,9,10,12,13,14
t	0.00675	15	8	C	C	G	T	A	A	3,4,6,9,10,11,13,14
r	0.00585	13	9	C	C	G	T	A	A	3,4,5,6,7,10,12,13,14
m	0.00585	13	6	C		G	T	A	A	1,2,10,11,13,14
l	0.00585	13	8	C	C	G	T	A	G	1,3,5,6,7,9,10,13
o	0.00495	11	5	C	C	G	T	A	G	2,3,6,7,13
p	0.0045	10	4	C		G	T	A	A	1,9,10,14
ab	0.00405	9	6	C	C	G	T	A	G	4,6,8,9,10,13
k	0.0036	8	5	C		G	T	T	G	2,6,7,12,13
x	0.00315	7	6	C	C	G	T	A	A	3,4,7,9,10,11
ah	0.0027	6	5	C		G	T	A	G	3,5,6,9,13
u	0.00225	5	2	C		T	T	A	G	1,14
v	0.00225	5	2	C		T	T	A	G	1,14
ap	0.00225	4	3	C		G	G	A	G	3,4,9
ac	0.00225	5	4	C		G	T	A	G	1,10,12,14
y	0.0018	4	3	C	C	G	T	A	A	1,3,12
ar	0.0018	4	3	T		G	T	A	G	1,10,13
ai	0.0018	4	3	C	C	G	T	A	G	2,4,7



Table 13.29 - IGHV1-3

aa	0.00135	C		T	T	A	A	14
ak	0.00135	C		G	T	A	G	1,9,10
w	0.00135	C		G	T	A	A	9
at	0.00135	C	C	G	T	A	G	6,10
ag	0.0009	C		G	T	A	A	10,14
aj	0.0009	T	C	G	T	A	G	3,4
aq	0.0009	C	C	G	T	A	A	10
s	0.0009	C	C	G	T	A	A	7,10
q	0.0009	T	C	G	T	A	A	2
z	0.0009	C		G	T	A	G	5
ad	0.0009	C	C	G	T	A	A	5,9
aw	0.0009	C	C	G	T	A	A	7,13
ae	0.0009	C		G	T	A	G	5,10
an	0.0009	C	C	T	T	A	A	12,14
bd	0.00045	C	C	G	G	A	G	10
ba	0.00045	C		G	T	A	G	3
ax	0.00045	C		G	T	A	G	5
bb	0.00045	C	C	G	T	A	A	8
az	0.00045	C	C	G	T	A	G	4
al	0.00045	T	C	G	T	A	A	10
be	0.00045	C		G	T	A	A	14
bf	0.00045	C	C	G	T	A	A	2
av	0.00045	C		G	T	A	A	13
ao	0.00045	T	C	G	T	A	A	1
af	0.00045	C	C	T	T	A	A	13
au	0.00045	C		G	T	A	A	14
bh	0.00045	C	C	G	T	A	A	14
am	0.00045	T		G	T	A	A	10
bc	0.00045	C		G	T	A	A	10
bg	0.00045	C		G	T	A	A	10
ay	0.00045	C		G	T	A	G	4
as	0.00045	C		G	T	A	G	10

Table 13.30 - IGHV1-45

h.type	cum.freq	# indivs	pops	14:106962947 rs186738824	14:106962984 rs145326338	14:106963101 rs7141669	14:106963167 rs182373995	14:106963326 rs12884037	
ref	0.75745	-	-	Met->Thr	Met->Leu	Gly->Arg	Val->Phe	Ala->Val	ENST00000390621
a	0.1286	251	12	A	T	C	C	G	
b	0.1126	220	12	A	T	T	C	G	1,2,3,4,5,6,7,9,10,11,12,13
d	0.0045	1	1	G	T	C	C	■	1,2,3,5,6,7,8,10,11,12,13,14
e	0.0045	1	1	A	G	C	C	G	1
c	0.0045	1	1	A	T	C	■	G	9
									3

Table 13.31 - IGHV1-46

h.type	cum.frq	# indivs	pops	14:106967122	14:106967151	14:106967155	14:106967171	14:106967173	14:106967177	14:106967241	14:106967249
ref	0.9778	-	-	rs147211698	rs55801711	rs149338091	rs181189514	rs185595166	rs190309173	rs144704015	rs182132309
a	0.0082	16	6	Thr->Met	Phe->Leu	Lys->Arg	Thr->Ala	Ser->Thr	Gly->Ser	Met->Ile	Tyr->Asn
b	0.0032	7	3	G	G	T	T	C	C	C	A
c	0.00225	5	3	G	G	T	T	C	C	C	A
d	0.0018	4	4	G	G	T	T	C	C	T	A
k	0.0009	2	2	G	G	C	T	C	C	C	A
e	0.0009	2	2	G	G	T	T	G	C	C	A
n	0.0009	2	2	G	G	T	T	C	C	C	A
j	0.00045	1	1	G	G	T	T	C	C	C	A
g	0.00045	1	1	G	G	T	T	G	C	C	T
h	0.00045	1	1	G	G	T	T	C	C	C	A
f	0.00045	1	1	G	G	T	T	C	C	C	A
i	0.00045	1	1	G	G	T	T	C	C	C	T
m	0.00045	1	1	G	G	C	T	C	T	C	A
l	0.00045	1	1	G	G	T	C	C	C	C	A
p	0.00045	1	1	G	G	T	T	C	C	C	A
o	0.00045	1	1	G	G	T	T	G	C	T	A

Table 13.31 - IGHV1-46

h.type	cum.frq	# indivs	pops	14:106967264 rs187613260 Tyr->Asp	14:106967273 rs192285778 Ala->Ser	14:106967275 rs61995748 Lys->Met	14:106967285 rs61747196 Val->>Ile	14:106967306 rs188566927 Lys->Gln	14:106967451 rs191694446 Ala->Thr	14:106967453 rs143810901 Leu->Pro	ENST00000390622
ref	0.9778	-	-	A	C	T	C	T	C	A	
a	0.0082	16	6	A	C	T	C	T	C	A	1,5,7,8,12,13
b	0.0032	7	3	A	C	T	C	T	C	A	3,10,14
c	0.00225	5	3	A	C	T	C	T	C	A	1,10,14
d	0.0018	4	4	A	C	T	C	T	C	A	2,10,11,14
k	0.0009	2	2	A	C	T	C	T	C	A	1,10
e	0.0009	2	2	A	C	T	C	T	C	A	6,10
n	0.0009	2	2	A	C	T	T	T	C	A	4,6
j	0.00045	1	1	A	C	T	C	T	C	A	1
g	0.00045	1	1	A	C	T	C	T	C	A	5
h	0.00045	1	1	A	C	T	T	G	C	A	5
f	0.00045	1	1	A	C	T	T	T	T	A	10
i	0.00045	1	1	A	C	T	C	T	C	A	9
m	0.00045	1	1	A	C	T	C	T	C	A	3
l	0.00045	1	1	A	C	T	C	T	C	A	3
p	0.00045	1	1	A	C	T	C	G	C	A	4
o	0.00045	1	1	C	C	T	C	T	C	G	10

Table 13.32 - IGHV1-58

h.type	cum.freq	# indivs	pops	14:107078380 rs189473179	14:107078395 rs141236708	14:107078567 rs2516904	14:107078570 rs145028255	14:107078578 rs189734868	14:107078591 rs146830036	14:107078609 rs113248643	14:107078661 rs180889109
ref	0.16105	-	-	Cys->Tyr	Thr->Met	Met->Val	Ala->Pro	Thr->Ser	Gly->Arg	Val->Phe	Met->Ile
a	0.4503	718	14	C	G	T	C	G	C	C	C
b	0.3144	568	14	C	G	C	C	G	C	C	C
c	0.05525	105	6	C	G	T	C	G	C	■	C
d	0.0068	15	6	C	■	T	C	G	C	C	C
e	0.0068	15	4	C	G	C	C	G	C	C	C
f	0.00225	5	3	T	G	T	C	G	C	C	C
g	0.0009	2	1	C	G	C	C	G	C	■	C
j	0.00045	1	1	C	G	C	C	C	C	C	C
k	0.00045	1	1	C	G	T	C	C	T	C	C
h	0.00045	1	1	C	G	T	C	G	C	C	■
i	0.00045	1	1	C	G	C	C	G	C	C	C
l	0.00045	1	1	C	G	T	G	G	C	C	C

Table 13.32 - IGHV1-58

h.type	cum.freq	# indivs	pops	14:107078674 rs186300171	14:107078775 rs148981028	14:107078790 rs1858692	
ref	0.16105	-	-	Ala->Val	Gly->Arg	Val->Ile	ENST00000390628
a	0.4503	718	14	G	C	C	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.3144	568	14	G	C	T	1,2,3,4,5,6,7,8,9,10,11,12,13,14
c	0.05525	105	6	G	C	T	1,5,10,12,13,14
d	0.0068	15	6	G	C	C	3,4,5,9,11,12
e	0.0068	15	4	G	T	T	1,10,13,14
f	0.00225	5	3	G	C	T	3,4,9
g	0.0009	2	1	G	C	T	10
j	0.00045	1	1	G	C	T	10
k	0.00045	1	1	G	C	C	13
h	0.00045	1	1	■	C	C	10
i	0.00045	1	1	■	C	T	1
l	0.00045	1	1	G	C	C	13

Table 13.33 - IGHV1-69

h.type	cum.freq	# indivs	pops	14:107169948 rs190613018	14:107169992 rs183124973	14:107169995 rs1064565	14:107170004 rs191670994	14:107170011 rs74089691	14:107170014 rs11557979	14:107170019 rs192735419	14:107170025 rs185522094
ref	0.49795	-	-	Val->Leu	Thr->Arg	Ser->Thr	Lys->Arg	Ala->Thr	Thr->Ser	Thr->Lys	Arg->Thr
a	0.1689	323	14	C	G	C	T	C	T	G	C
b	0.1543	312	14	C	G	C	T	C	T	G	C
c	0.0561	118	13	C	G	C	T	C	T	G	C
g	0.0348	76	7	C	G	C	T	T	T	G	C
e	0.014	31	4	C	G	C	T	T	T	G	C
l	0.0108	24	6	C	G	C	T	C	T	G	C
d	0.00995	22	3	C	G	C	T	C	T	G	C
h	0.00765	17	10	C	G	C	T	C	T	G	C
f	0.00685	15	8	C	G	C	T	C	T	G	C
i	0.00675	15	6	C	G	C	T	C	T	G	C
m	0.0027	6	3	C	G	C	T	C	T	G	C
j	0.00225	5	3	C	G	C	T	C	T	G	C
u	0.00225	5	4	C	G	C	T	C	T	G	C
s	0.0018	4	3	C	G	C	T	C	T	G	C
k	0.00135	3	1	C	G	C	T	T	T	G	C
ac	0.00135	3	3	C	G	C	T	C	T	G	C
ah	0.0009	2	2	C	G	G	T	C	T	G	C
t	0.0009	2	2	C	G	C	T	C	T	G	C
ao	0.0009	2	1	C	G	C	T	T	T	G	C
af	0.0009	2	2	C	G	G	T	C	T	G	C
ad	0.0009	2	2	C	G	C	T	C	T	G	C
n	0.0009	2	2	C	G	C	T	C	T	G	C
an	0.0009	2	2	C	G	C	T	T	T	G	C
o	0.0009	2	2	C	G	C	T	C	T	G	C
ba	0.00045	1	1	C	G	C	T	C	T	G	C
ax	0.00045	1	1	C	G	C	T	T	T	G	C
ag	0.00045	1	1	C	G	C	T	C	T	G	C
y	0.00045	1	1	C	G	C	T	C	T	G	C





Table 13.33 - IGHV1-69

h.type	cum.freq	# indivs	pops	14:107170028 rs11557995	14:107170034 rs181467093	14:107170055 rs8009570	14:107170056 rs184214708	14:107170062 rs55891010	14:107170077 rs11845244	14:107170091 rs189019973	14:107170121 rs1064564
ref	0.49795	-	-	Gly->Ala	Phe->Ser	Thr->Ile	Thr->Ala	Phe->Leu	Gly->Arg	Leu->Pro	Ser->Asn
a	0.1689	323	14	C	A	G	T	A	C	A	C
b	0.1543	312	14	C	A	■	T	G	T	A	C
c	0.0561	118	13	C	A	■	T	G	C	A	C
g	0.0348	76	7	C	A	G	T	A	C	A	C
e	0.014	31	4	C	A	G	T	A	T	A	C
l	0.0108	24	6	C	A	G	T	A	T	A	C
d	0.00995	22	3	C	A	G	T	A	T	A	C
h	0.00765	17	10	C	A	■	T	A	C	A	C
f	0.00685	15	8	C	A	■	T	A	T	A	C
i	0.00675	15	6	C	A	■	T	A	T	A	C
m	0.0027	6	3	C	A	G	T	A	C	A	C
j	0.00225	5	3	C	A	G	T	G	T	A	C
u	0.00225	5	4	C	A	■	T	G	C	A	C
s	0.0018	4	3	C	A	G	T	A	C	A	T
k	0.00135	3	1	C	A	■	T	G	T	A	C
ac	0.00135	3	3	C	A	G	T	A	C	A	C
ah	0.0009	2	2	C	A	G	T	A	C	A	C
t	0.0009	2	2	C	A	G	T	A	C	A	C
ao	0.0009	2	1	C	A	■	T	G	C	A	C
af	0.0009	2	2	C	A	■	T	G	C	A	C
ad	0.0009	2	2	C	A	G	T	A	C	A	C
n	0.0009	2	2	C	A	■	T	G	T	A	T
an	0.0009	2	2	C	A	■	T	G	T	A	C
o	0.0009	2	2	C	A	■	T	G	T	A	C
ba	0.00045	1	1	C	A	G	T	A	C	A	C
ax	0.00045	1	1	C	A	G	T	A	C	A	C
ag	0.00045	1	1	C	A	■	T	G	T	A	C
y	0.00045	1	1	C	A	■	T	G	T	G	C

Table 13.33 - IGHV1-69

1	0.00045	az	C	A	G	T	A	T	A	C
1	0.00045	aj	C	A	G	C	A	C	A	T
1	0.00045	aa	C	A	■	T	A	C	A	C
1	0.00045	aq	C	A	G	C	A	C	A	C
1	0.00045	v	C	A	■	T	G	T	A	C
1	0.00045	al	C	A	■	T	G	T	A	C
1	0.00045	ab	C	A	■	T	G	C	A	T
1	0.00045	ak	C	A	G	T	A	C	A	C
1	0.00045	q	C	A	G	T	A	C	A	C
1	0.00045	ap	C	A	G	T	A	C	A	C
1	0.00045	z	C	A	G	T	G	C	A	C
1	0.00045	w	C	A	■	T	G	T	A	T
1	0.00045	av	C	A	■	T	G	T	A	C
1	0.00045	r	C	A	■	T	G	T	A	C
1	0.00045	x	C	A	■	T	G	T	A	C
1	0.00045	au	C	A	■	T	G	T	A	C
1	0.00045	am	C	A	G	T	A	C	A	C
1	0.00045	at	C	A	■	T	G	T	A	C
1	0.00045	aw	C	G	■	T	G	T	A	T
1	0.00045	ay	C	A	G	C	A	C	A	T
1	0.00045	ar	C	A	■	T	G	T	A	C
1	0.00045	p	C	A	■	T	G	T	A	C
1	0.00045	as	G	A	■	T	G	C	A	C
1	0.00045	ae	C	A	G	T	A	C	A	C
1	0.00045	ai	C	A	■	T	G	T	A	C

Table 13.33 - IGHV1-69

h.type	cum.freq	# indivs	pops	14:107170127 rs11557984	14:107170128 rs11558005	14:107170130 rs140392997	14:107170133 rs138194290	14:107170139 rs11558018	14:107170143 rs186595111	14:107170145 rs189122962	14:107170173 rs150196675
ref	0.49795	-	-	Ala->Val	Ala->Thr	Tyr->Phe	Ser->Asn	Phe->Ser	Thr->Pro	Gly->Asp	Val->Leu
a	0.1689	323	14	G	C	T	C	A	T	C	C
b	0.1543	312	14	G	T	T	C	A	T	C	C
c	0.0561	118	13	G	C	T	C	A	T	C	C
g	0.0348	76	7	G	C	T	C	A	T	C	C
e	0.014	31	4	G	C	T	C	A	T	C	C
l	0.0108	24	6	G	C	T	C	A	T	C	C
d	0.00995	22	3	G	T	T	C	A	T	C	C
h	0.00765	17	10	G	C	T	C	A	T	C	C
f	0.00685	15	8	G	T	T	C	A	T	C	C
i	0.00675	15	6	G	C	T	C	A	T	C	C
m	0.0027	6	3	G	C	T	C	A	T	C	C
j	0.00225	5	3	G	C	T	C	A	T	C	C
u	0.00225	5	4	G	T	T	C	A	T	C	C
s	0.0018	4	3	G	C	T	C	A	T	C	C
k	0.00135	3	1	G	T	T	C	A	T	C	C
ac	0.00135	3	3	G	T	T	C	A	T	C	C
ah	0.0009	2	2	G	C	T	C	A	T	C	C
t	0.0009	2	2	█	C	T	C	A	T	C	C
ao	0.0009	2	1	G	C	T	C	A	T	C	C
af	0.0009	2	2	G	C	T	C	A	T	C	C
ad	0.0009	2	2	G	C	T	T	A	T	C	C
n	0.0009	2	2	G	C	T	C	A	T	C	C
an	0.0009	2	2	G	C	T	C	A	T	C	C
o	0.0009	2	2	█	T	T	C	A	T	C	C
ba	0.00045	1	1	G	C	T	C	A	T	C	C
ax	0.00045	1	1	G	C	T	C	A	T	C	C
ag	0.00045	1	1	G	T	█	T	A	T	C	C
y	0.00045	1	1	G	C	T	C	A	T	C	C

Table 13.33 - IGHV1-69

1	0.00045	az	G	C	T	T	T	A	T	C	C
1	0.00045	aj	G	C	T	T	T	A	T	C	C
1	0.00045	aa	G	T	T	T	T	A	T	C	C
1	0.00045	aq	G	C	T	T	T	A	T	C	C
1	0.00045	v	G	C	T	T	T	A	G	C	C
1	0.00045	al	G	C	T	T	T	A	T	C	C
1	0.00045	ab	G	C	T	T	T	A	T	C	C
1	0.00045	ak	G	C	T	T	T	A	T	C	C
1	0.00045	q	G	C	T	T	T	G	T	C	C
1	0.00045	ap	G	C	T	T	T	A	T	C	C
1	0.00045	z	G	C	T	T	T	A	T	C	C
1	0.00045	w	G	T	T	T	T	A	T	C	C
1	0.00045	av	G	C	T	T	T	A	T	C	C
1	0.00045	r	G	C	T	T	T	A	T	C	C
1	0.00045	x	G	C	T	T	T	A	T	C	C
1	0.00045	au	G	C	T	T	T	A	T	C	C
1	0.00045	am	G	T	T	T	T	A	T	C	C
1	0.00045	at	G	C	T	T	T	A	T	C	C
1	0.00045	aw	G	C	T	T	T	A	T	C	C
1	0.00045	ay	G	C	T	T	T	A	T	C	C
1	0.00045	ar	G	C	T	T	T	A	T	C	C
1	0.00045	p	G	T	T	T	T	A	T	C	C
1	0.00045	as	G	C	T	T	T	A	T	C	C
1	0.00045	ae	G	C	T	T	T	A	T	C	C
1	0.00045	ai	G	T	T	T	T	A	T	C	C

Table 13.33 - IGHV1-69

h.type	cum.freq	# indivs	pops	14:107170185 rs181067247	14:107170326 .	14:107170343 rs190658595	ENST00000390633
ref	0.49795	-	-	Pro->Thr	->Ala	Phe->Val	
a	0.1689	323	14	G	AGCT	A	
b	0.1543	312	14	G	AGCT	A	1,2,3,4,5,6,7,8,9,10,11,12,13,14
c	0.0561	118	13	G	AGCT	A	1,2,3,4,5,6,7,8,9,10,11,12,13,14
g	0.0348	76	7	G	AGCT	A	1,2,3,4,5,6,7,9,10,11,12,13,14
e	0.014	31	4	G	AGCT	A	1,5,10,11,12,13,14
l	0.0108	24	6	G	AGCT	A	1,10,12,14
d	0.00995	22	3	G	AGCT	A	1,3,8,10,12,14
h	0.00765	17	10	G	AGCT	A	1,10,14
f	0.00685	15	8	G	AGCT	A	1,2,3,4,6,7,10,11,12,13
i	0.00675	15	6	G	AGCT	A	1,3,5,6,9,10,11,14
m	0.0027	6	3	G	■	A	3,4,5,9,10,12
j	0.00225	5	3	G	AGCT	A	10,12,14
u	0.00225	5	4	G	AGCT	A	5,10,14
s	0.0018	4	3	G	AGCT	A	1,2,8,14
k	0.00135	3	1	G	AGCT	A	6,8,10
ac	0.00135	3	3	G	AGCT	A	14
ah	0.0009	2	2	G	AGCT	A	2,10,14
t	0.0009	2	2	G	AGCT	A	11,13
ao	0.0009	2	1	G	AGCT	A	2,9
af	0.0009	2	2	G	AGCT	A	10
ad	0.0009	2	2	G	AGCT	A	10,14
n	0.0009	2	2	G	AGCT	A	2,7
an	0.0009	2	2	G	AGCT	A	2,3
o	0.0009	2	2	G	AGCT	A	10,14
ba	0.00045	1	1	T	AGCT	A	3,10
ax	0.00045	1	1	G	AGCT	A	13
ag	0.00045	1	1	G	AGCT	A	10
y	0.00045	1	1	G	AGCT	A	13
							10

Table 13.33 - IGHV1-69

az	0.00045	1	G	AGCT	A	10
aj	0.00045	1	G	AGCT	A	7
aa	0.00045	1	G	AGCT	A	2
aq	0.00045	1	G	AGCT	A	9
v	0.00045	1	G	AGCT	A	11
al	0.00045	1	G	■	A	10
ab	0.00045	1	G	AGCT	A	14
ak	0.00045	1	G	AGCT	A	4
q	0.00045	1	G	AGCT	A	4
ap	0.00045	1	G	AGCT	A	13
z	0.00045	1	G	AGCT	A	14
w	0.00045	1	G	AGCT	A	14
av	0.00045	1	G	AGCT	A	4
r	0.00045	1	G	AGCT	A	8
x	0.00045	1	G	AGCT	A	2
au	0.00045	1	G	AGCT	A	2
am	0.00045	1	G	AGCT	A	1
at	0.00045	1	G	AGCT	A	3
aw	0.00045	1	G	AGCT	A	2
ay	0.00045	1	G	AGCT	A	10
ar	0.00045	1	G	AGCT	<b>C</b>	3
p	0.00045	1	G	AGCT	A	13
as	0.00045	1	G	AGCT	A	12
ae	0.00045	1	G	AGCT	A	13
ai	0.00045	1	G	AGCT	A	11

Table 13.34 - IGHV1-8

h.type	cum.fre	# indivs	pops	14:106539147 rs148477632 Ile->Val	14:106539163 rs73371428 Met->Ile	14:106539171 rs187649961 Val->Ile	14:106539278 rs184294529 Tyr->Phe	14:106539284 rs151035485 Thr->Ile	14:106539291 rs187522932 Thr->Pro	14:106539476 rs189071216 Ala->Thr	14:106539497 rs180817521 Ile->Phe
ref	0.8844	-	-	T	C	C	T	G	T	C	T
a	0.1016	160	9	T		C	T	G	T	C	T
c	0.00405	8	2	<b>C</b>	C	C	T	G	T	C	T
b	0.0032	6	6	T	C	C	T	G	T	C	
d	0.0018	4	2	T		C	T		T	C	
e	0.00135	3	2	T		C	T		T	C	
f	0.00135	3	2	T	C	<b>T</b>	T	G	T	C	
h	0.0009	2	2	T	C	C	<b></b>	G	T	C	
j	0.00045	1	1	T	C	C	T	G	<b>G</b>	C	
g	0.00045	1	1	T		C	T	G	T	<b>T</b>	
i	0.00045	1	1	T	C	C	T		T	C	

Table 13.34 - IGHV1-8

h.type	cum.frq	# indivs	pops	ENST00000390599
ref	0.8844	-	-	
a	0.1016	160	9	1,5,7,8,10,11,12,13,14
c	0.00405	8	2	10,13
b	0.0032	6	6	2,5,6,11,12,13
d	0.0018	4	2	1,14
e	0.00135	3	2	2,11
f	0.00135	3	2	7,11
h	0.0009	2	2	3,5
j	0.00045	1	1	2
g	0.00045	1	1	14
i	0.00045	1	1	13



Table 13.35 - IGHV2-26

h.type	cum. freq	# indivs	pops	14:106757655 rs12586893	14:106757706 rs185534639	14:106757798 rs181476099	14:106757831 rs11177969	14:106757865 rs185965403	14:106757880 rs146488407	14:106758074 rs186573875	
ref	0.9245	-	-	G	G	G	C	G	Val->Ile C	Thr->Ala T	ENST00000390611
a	0.06135	126	13	■	G	G	C	G	C	T	1,3,4,5,6,7,8,9,10,11,1 2,13,14
b	0.0087	15	5	G	G	G	T	G	C	T	1,6,10,11,14
c	0.0032	7	2	G	G	T	C	C	C	T	10,14
f	0.0009	2	2	G	G	G	C	C	T	T	13,14
d	0.00045	1	1	G	■	G	C	G	C	T	1
g	0.00045	1	1	G	G	G	C	C	C	T	5
e	0.00045	1	1	G	G	G	C	G	C	C	10

Table 13.36 - IGHV2-5

h.type	ref	cum. fre	# indivs	pops	14:106494234	14:106494245	14:106494248	14:106494251	14:106494269	14:106494325	14:106494346	14:106494404
					rs1065554	rs1065552	rs181774040	rs141264800	rs12895651	rs150364725	rs185589849	rs190103562
					Ser->Arg	Ser->Thr	Pro->Thr	Ser->Gly	Asn->Asp	Gly->Ala	Ser->Thr	Leu->Val
		0.3179	-	-	G	A	G	T	T	C	C	G
a		0.6657	872	14	G	A	G	T	C	C	C	G
b		0.01055	18	3	G	A	G	C	C	C	C	G
d		0.0018	4	2	G	A	T	T	C	C	C	G
c		0.0009	2	2	G	A	G	T	C	G	C	G
j		0.00045	1	1	G	T	T	T	C	C	C	G
k		0.00045	1	1	G	A	G	T	T	C	G	G
g		0.00045	1	1	C	A	G	T	C	G	C	G
e		0.00045	1	1	G	A	G	T	C	C	G	G
h		0.00045	1	1	C	A	T	T	C	C	C	G
f		0.00045	1	1	G	A	G	T	C	C	C	C
i		0.00045	1	1	C	A	G	T	C	C	C	G

ENST00000390597

1,2,3,4,5,6,7,8,9,10,11,1  
2,13,14  
1,10,14  
10,14  
4,13  
6  
1  
9  
3  
5  
10  
12

Table 13.37 - IGHV2-70

h.type	cum.frec	# indivs	pops	14:107178820 rs139383686	14:107178837 rs144976346	14:107178849 rs185317654	14:107178875 rs188731942	14:107178878 rs192620545	14:107178890 rs138836314	14:107178899 rs11265724	14:107178911 rs188243654	14:107178938 rs17113976
ref	0.36095	-	-	Ile->Met	Tyr->Asp	Asp->Tyr	Leu->Pro	Val->Ala	Lys->Arg	Asp->Ala	Thr->Ile	Tyr->Cys
a	0.31	584	14	T	A	C	A	A	T	T	G	T
b	0.16595	353	14	T	A	C	A	A	T	T	G	T
d	0.06035	132	13	T	A	C	A	A	T	G	G	C
c	0.04295	94	11	T	A	C	A	A	T	T	G	T
e	0.0319	70	6	T	A	C	A	A	T	T	G	C
f	0.00405	9	5	T	A	C	A	A	T	T	G	C
l	0.00225	5	4	T	A	C	G	A	T	T	G	T
y	0.00135	3	3	T	A	C	A	A	T	T	G	T
z	0.00135	3	3	C	A	C	A	A	T	T	G	T
af	0.00135	3	2	T	A	C	A	A	T	G	G	C
h	0.00135	3	3	T	A	C	A	A	T	G	G	C
g	0.0009	2	2	T	A	C	A	A	T	T	G	T
s	0.0009	2	2	T	A	C	A	A	T	T	G	T
ab	0.0009	2	2	T	A	C	A	A	C	T	G	T
ah	0.00045	1	1	C	C	C	A	A	T	T	G	T
ag	0.00045	1	1	T	A	C	A	A	T	G	G	C
j	0.00045	1	1	T	C	C	A	A	T	T	G	T
u	0.00045	1	1	T	A	C	A	A	T	T	G	C
k	0.00045	1	1	T	A	C	A	A	T	T	G	T
aj	0.00045	1	1	C	A	C	A	A	T	T	G	T
aa	0.00045	1	1	T	A	C	A	A	T	T	G	C
t	0.00045	1	1	T	A	C	A	A	T	T	G	C
aq	0.00045	1	1	T	A	C	A	A	T	G	G	C
v	0.00045	1	1	C	A	C	A	A	T	G	G	T
al	0.00045	1	1	T	A	C	A	A	C	T	G	T
ak	0.00045	1	1	T	A	C	A	A	T	T	G	C
q	0.00045	1	1	T	A	C	A	A	T	T	G	T
ap	0.00045	1	1	T	A	C	A	A	T	G	G	C

Table 13.37 - IGHV2-70

w	0.00045	1	T	C	A	A	A	T	T	G	G	T
ao	0.00045	1	T	A	A	A	A	T	T	G	G	C
r	0.00045	1	T	A	A	A	A	T	T	G	G	T
x	0.00045	1	C	A	A	A	A	T	T	T	■	T
ad	0.00045	1	T	A	A	A	G	T	T	G	G	C
am	0.00045	1	T	A	A	A	A	C	C	T	G	C
i	0.00045	1	T	A	A	A	A	T	T	T	G	T
n	0.00045	1	T	A	A	A	A	T	T	T	G	T
ac	0.00045	1	C	A	A	A	A	T	T	G	G	C
m	0.00045	1	T	A	A	A	A	T	T	T	■	T
ar	0.00045	1	T	A	A	A	A	T	T	T	G	T
p	0.00045	1	T	A	A	A	A	T	T	T	G	T
ae	0.00045	1	T	A	A	A	A	T	T	T	■	T
ai	0.00045	1	T	A	A	A	A	T	T	T	G	C
an	0.00045	1	T	A	A	A	A	T	T	T	G	T
o	0.00045	1	T	C	A	A	A	T	T	G	G	C

Table 13.37 - IGHV2-70

h.type	cum.frec	# indivs	pops	14:107178959 rs184814603	14:107178965 rs2073669	14:107178976 rs148856487	14:107179098 rs191880827	14:107179103 rs185635448	14:107179106 rs10144703	14:107179115 rs182292946	14:107179123 rs187704106	14:107179243 rs61734101
ref	0.36095	-	-	Asp->Gly	Leu->Arg	Glu->Asp	Gly->Ser	Glu->Val	Arg->Lys	Val->Gly	Leu->Phe	Thr->Met
a	0.31	584	14	T	A	C	C	T	C	A	T	G
b	0.16595	353	14	T	C	C	C	T	C	A	T	G
d	0.06035	132	13	T	C	C	C	T	T	A	T	G
c	0.04295	94	11	T	A	C	C	T	T	A	T	G
e	0.0319	70	6	T	C	C	C	T	T	A	T	G
f	0.00405	9	5	T	C	C	C	T	T	A	T	G
l	0.00225	5	4	T	C	C	C	T	C	A	T	G
y	0.00135	3	3	C	C	C	C	T	C	A	T	G
z	0.00135	3	3	T	C	C	C	T	T	A	T	G
af	0.00135	3	2	T	C	C	C	T	T	A	T	G
h	0.00135	3	3	T	C	C	C	T	T	A	T	G
g	0.0009	2	2	T	A	G	C	T	C	A	T	G
s	0.0009	2	2	T	C	C	T	T	C	A	T	G
ab	0.0009	2	2	T	A	C	C	T	C	A	T	G
ah	0.00045	1	1	T	C	C	C	T	C	A	T	G
ag	0.00045	1	1	C	C	C	C	T	T	A	T	G
j	0.00045	1	1	T	C	C	C	T	C	A	T	G
u	0.00045	1	1	T	A	C	C	T	C	A	T	G
k	0.00045	1	1	T	A	G	C	T	T	A	T	G
aj	0.00045	1	1	T	C	C	C	T	C	A	T	G
aa	0.00045	1	1	T	C	C	C	T	T	A	T	G
t	0.00045	1	1	T	A	C	C	T	T	A	T	G
aq	0.00045	1	1	T	C	C	C	T	T	A	T	G
v	0.00045	1	1	T	A	C	C	T	T	A	T	G
al	0.00045	1	1	T	C	C	C	T	C	A	T	G
ak	0.00045	1	1	T	C	C	C	T	T	A	T	G
q	0.00045	1	1	T	C	C	C	T	C	A	T	G
ap	0.00045	1	1	T	C	C	C	T	C	A	T	G

Table 13.37 - IGHV2-70

w	0.00045	1	T	A	C	C	T	C	T	A	T	G
ao	0.00045	1	T	C	C	C	T	T	T	A	T	G
r	0.00045	1	T	A	C	C	T	T	T	A	T	G
x	0.00045	1	T	C	C	C	T	C	T	A	T	G
ad	0.00045	1	T	C	C	C	T	T	T	A	T	G
am	0.00045	1	T	C	C	C	T	T	T	A	T	G
i	0.00045	1	T	A	C	C	T	C	T	C	T	G
n	0.00045	1	T	A	C	C	T	C	T	A	T	G
ac	0.00045	1	T	C	C	C	T	T	T	A	T	G
m	0.00045	1	T	C	C	C	T	T	T	A	T	G
ar	0.00045	1	C	C	G	C	T	C	T	A	T	G
p	0.00045	1	T	C	C	C	T	C	T	A	T	G
ae	0.00045	1	C	C	C	C	T	T	T	A	T	G
ai	0.00045	1	T	C	C	C	T	C	T	A	T	G
an	0.00045	1	C	A	C	C	T	C	T	A	T	G
o	0.00045	1	T	C	C	C	T	T	T	A	T	G

Table 13.37 - IGHV2-70

h.type	cum.frec	# indvs	pops	14:107179254	rs61734098	ENST00000390634
ref	0.36095	-	-		Ile->Met	
a	0.31	584	14		T	
b	0.16595	353	14		T	1,2,3,4,5,6,7,8,9,10,11,12,13,14
d	0.06035	132	13		C	1,2,3,4,5,6,7,8,9,10,11,12,13,14
c	0.04295	94	11		T	1,2,3,4,5,6,7,9,10,11,12,13,14
e	0.0319	70	6		C	1,2,5,6,7,8,10,11,12,13,14
f	0.00405	9	5		T	1,5,10,12,13,14
l	0.00225	5	4		T	1,5,9,10,14
y	0.00135	3	3		T	2,3,4,10
z	0.00135	3	3		T	7,9,12
af	0.00135	3	2		T	10,13,14
h	0.00135	3	3		T	6,10
g	0.0009	2	2		T	2,10,13
s	0.0009	2	2		T	4,6
ab	0.0009	2	2		T	5,6
ah	0.00045	1	1		T	2,5
ag	0.00045	1	1		C	2
j	0.00045	1	1		T	10
u	0.00045	1	1		T	10
k	0.00045	1	1		T	13
aj	0.00045	1	1		T	10
aa	0.00045	1	1		T	14
t	0.00045	1	1		T	10
aq	0.00045	1	1		C	4
v	0.00045	1	1		T	13
al	0.00045	1	1		T	14
ak	0.00045	1	1		C	10
q	0.00045	1	1		T	1
ap	0.00045	1	1		C	10



Table 13.37 - IGHV2-70

w	0.00045	1	1	T	13
ao	0.00045	1	1	C	1
r	0.00045	1	1	T	2
x	0.00045	1	1	T	4
ad	0.00045	1	1	C	12
am	0.00045	1	1	C	1
i	0.00045	1	1	T	4
n	0.00045	1	1	T	5
ac	0.00045	1	1	T	13
m	0.00045	1	1	T	12
ar	0.00045	1	1	T	11
p	0.00045	1	1	T	4
ae	0.00045	1	1	T	13
ai	0.00045	1	1	T	9
an	0.00045	1	1	T	13
o	0.00045	1	1	C	14

Table 13.38 - IGHV3-11

h.type	cum.freq	# indvs	pops	14:106573345 rs186167416	14:106573637 rs191020138	14:106573662 rs74207678	ENST00000390601
ref	0.9817	-	-	Ala->Val G	Lys->ArgLys->Arg T	Trp->Arg A	
a	0.01695	36		G	T	<b>G</b>	3,4,9
b	0.0009	2			T	A	3,5
c	0.00045	1		G	<b>C</b>	A	3

Table 13.39 - IGHV3-13

h.type	cum.freq	# indivs	pops	14:106586159 rs192397984	14:106586165 rs140926945	14:106586168 rs184992069	14:106586231 rs10151633	14:106586259 rs11625174	14:106586267 rs189564282	14:106586354 rs10131161	14:106586389 rs181545751
ref	0.69725	-	-	Thr->Met	Gly->Glu	Ala->Gly	Arg->Gln	Thr->Pro	Ala->Val	Ser->Cys	Gln->His
a	0.15245	276	13	G	C	G	C	T	G	G	C
b	0.1011	167	9	G	C	G	T	G	G	C	C
c	0.02925	59	10	G	T	G	C	T	G	G	C
d	0.0105	19	4	■	C	G	T	T	G	C	C
e	0.00225	5	3	G	C	G	C	T	G	C	C
j	0.0018	4	2	G	C	G	T	T	G	G	C
g	0.00135	3	2	G	C	G	C	T	G	G	■
f	0.0009	2	2	G	C	G	T	T	■	C	C
k	0.00045	1	1	G	T	G	C	G	G	G	C
h	0.00045	1	1	■	C	G	C	T	G	G	C
i	0.00045	1	1	■	C	G	T	T	G	C	■
n	0.00045	1	1	G	C	G	C	G	G	G	C
m	0.00045	1	1	G	C	G	C	T	G	G	C
l	0.00045	1	1	G	C	C	C	T	G	G	C
o	0.00045	1	1	G	T	G	T	T	G	C	C

Table 13.39 - IGHV3-13

h.type	cum.freq	# indivs	pops	14:106586557	14:106586563	rs112340466	rs137975618	ENST00000390602
ref	0.69725	-	-	C	A	Val->Ile	Phe->Leu	
a	0.15245	276	13	C	A			1,2,3,4,5,6,7,8,9,10,11,12,13
b	0.1011	167	9	C	A			1,5,7,8,10,11,12,13,14
c	0.02925	59	10	C	A			1,2,5,6,7,10,11,12,13,14
d	0.0105	19	4	C	A			1,10,12,14
e	0.00225	5	3	C	A			1,10,14
j	0.0018	4	2	C	A			1,10
g	0.00135	3	2	C	A			6,14
f	0.0009	2	2	C	A			10,12
k	0.00045	1	1	C	A			2
h	0.00045	1	1	C	A			1
i	0.00045	1	1	C	A			14
n	0.00045	1	1	C	G			11
m	0.00045	1	1	T	G			14
l	0.00045	1	1	C	A			3
o	0.00045	1	1	C	A			10

Table 13.40 - IGHV3-15

h.type	cum.freq	# indivs	pops	14:106610377	14:106610467	14:106610509	14:106610570	14:106610736	
ref	0.91495	-	-	Asn->Thr	Gly->Ala	Ser->Asn	Gly->Arg	Ile->Phe	ENST00000390603
a	0.0572	104	11	T	C	C	C	T	
b	0.0265	56	5	T	C	T	C	T	1,2,3,5,6,7,10,11,12,13,14
d	0.00045	1	1	T	C	C	T	T	1,3,4,9,13
e	0.00045	1	1	G	C	C	C	T	6
c	0.00045	1	1	T	G	C	C	T	6
									9

Table 13.41 - IGHV3-16

h.type	cum.fre	# indivs	pops	14:106622052 rs12323556	14:106622073 rs142354696	14:106622076 rs12433709	14:106622132 rs12323559	ENST00000390604
ref	0.8039	-	-	Glu->Lys C	Lys->Glu T	Arg->Cys G	Arg->Ile C	
a	0.11075	200	13	C	T	■	C	1,2,3,4,5,6,7,8,9,11,12,13,14
b	0.07085	122	8	T	T	G	■	1,5,7,10,11,12,13,14
c	0.01	21	4	C	C	G	■	1,10,11,14
d	0.00315	7	3	T	T	G	C	10,11,14
e	0.00135	3	2	C	C	G	C	10,12

Table 13.42 - IGHV3-20

h.type	cum.freq	# indvs	pops	14:106667592 rs3751514	14:106667756 rs190163710	14:106667785 rs181860901	14:106667786 rs3751513	14:106667810 rs112170273	14:106668011 rs180685266	
ref	0.5693	-	-	His->Tyr G	Ala->Val G	Asp->Glu A	Asp->Gly T	Cys->Phe C	Val->Ala A	ENST00000390606
a	0.2829	470	14		G	A	T	C	A	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.1209	221	13	G	G	A	T		A	1,2,3,4,5,6,7,8,9,10,11,12,13
c	0.02285	50	6		G	A	C	C	A	1,3,4,9,10,14
f	0.00135	3	3	G	G	A	C	C	A	9,11,14
d	0.0009	2	2		G	A	T		A	5,11
g	0.00045	1	1		G	A	T	C	G	10
e	0.00045	1	1	G	G	A	T		G	2
h	0.00045	1	1	G		A	T	C	A	14
i	0.00045	1	1	G		T	T	C	A	1

Table 13.43 - IGHV3-21

h.type	cum.freq	# indivs	pops	14:106691762	14:106691807	14:106691824	14:106691833	14:106691849	14:106691865	14:106691867	14:106691879
				rs183314182	rs188058888	rs143336904	rs192148312	rs184550642	rs192949902	rs184209619	rs180974163
				Thr->Ala	Ser->Gly	Val->Ala	Leu->Pro	Ala->Thr	Met->Ile	Met->Leu	Ser->Gly
ref	0.9892	-	-	T	T	A	A	C	C	T	T
a	0.00225	5	-	T	<b>C</b>	A	A	C	C	T	T
d	0.0018	4	3	T	T	A	A	C	C	T	T
c	0.0009	2	2	T	T	<b>G</b>	A	C	C	T	T
b	0.0009	1	1	T	T	A	A	C	C	T	T
h	0.0009	2	2	<b>C</b>	T	A	A	C	C	T	T
j	0.00045	1	1	T	T	A	<b>G</b>	C	C	T	T
k	0.00045	1	1	T	T	A	A	C	C	T	T
g	0.00045	1	1	T	<b>C</b>	A	A	C	C	T	T
e	0.00045	1	1	T	T	A	A	C	C	■	T
f	0.00045	1	1	T	T	A	A	C	<b>G</b>	■	<b>C</b>
i	0.00045	1	1	T	T	A	A	<b>T</b>	C	T	T
n	0.00045	1	1	T	<b>C</b>	A	A	<b>T</b>	C	T	T
m	0.00045	1	1	T	T	A	A	C	C	T	T
l	0.00045	1	1	T	T	A	A	C	C	T	T



Table 13.43 - IGHV3-21

h.type	cum.freq	# indivs	pops	14:106691884 rs184687518	14:106691896 rs187020320	14:106691899 rs191067794	14:106691909 rs181216108	ENST00000390607
ref	0.9892	-	-	Thr->Ile	Ala->Gly	Ala->Val	Leu->Phe	
a	0.00225	5	5	G	G	G	G	
d	0.0018	4	3	G	G	G	G	7,9,11,13,14
c	0.0009	2	2	G	G	G	G	1,5,14
b	0.0009	1	1	G	C	G	G	10,14
h	0.0009	2	2	G	G	G	G	1
j	0.00045	1	1	G	G	G	G	1,13
k	0.00045	1	1	G	G	G	G	2
g	0.00045	1	1	G	G	G	G	1
e	0.00045	1	1	G	G	G	G	5
f	0.00045	1	1	G	C	G	G	2
i	0.00045	1	1	G	G	G	G	4
n	0.00045	1	1	G	G	G	G	14
m	0.00045	1	1	G	C	G	G	3
l	0.00045	1	1	G	C	G	G	11
								1

Table 13.44 - IGHV3-23

h.type	cum.freq	# indivs	pops	14:106725320 rs189958807	14:106725328 rs1064091	14:106725329 rs1064090	14:106725346 rs1055799	14:106725357 rs143624256	14:106725397 rs61752504	14:106725400 rs183319819	14:106725433 rs187384491
ref	0.83265	-	-	Tyr->His	Gly->Asp	Gly->Ser	Ala->Val	Glu->Asp	Ala->Gly	Tyr->Ser	Ser->Phe
a	0.09825	190	12	A	C	C	G	C	G	T	G
b	0.02695	58	9	A	C	C	G	C	G	T	G
c	0.01365	29	9	A	C	T	■	C	G	T	G
d	0.00865	18	4	A	C	T	G	C	C	T	G
e	0.00455	10	4	A	C	T	■	C	G	T	G
f	0.00225	5	3	A	C	C	G	C	C	T	G
k	0.00225	5	5	A	T	C	G	C	G	T	G
g	0.00225	5	3	A	C	C	■	C	G	T	G
n	0.0018	4	3	A	C	C	G	C	C	T	G
j	0.0009	2	1	A	C	C	■	C	G	T	G
h	0.0009	2	2	A	C	C	G	C	G	T	G
u	0.00045	1	1	A	T	C	G	C	G	G	G
t	0.00045	1	1	A	T	C	G	G	G	T	G
v	0.00045	1	1	A	C	C	G	G	G	T	G
s	0.00045	1	1	A	T	C	G	C	C	T	G
q	0.00045	1	1	A	C	C	G	C	C	T	G
r	0.00045	1	1	A	C	T	G	C	G	T	G
i	0.00045	1	1	A	C	C	G	C	G	T	G
m	0.00045	1	1	A	C	C	G	C	G	T	G
l	0.00045	1	1	G	C	C	G	C	G	T	G
p	0.00045	1	1	A	T	C	G	C	G	G	G
o	0.00045	1	1	A	C	C	G	C	G	T	■

Table 13.44 - IGHV3-23

h.type	cum.freq	# indvs	pops	Ser->Thr	Leu->Val	Val->Gly	Glu->Gly	Ala->Pro	Phe->Val	Trp->Cys	ENST00000390609
ref	0.83265	-	-	A	A	A	T	C	A	Trp->Cys	
a	0.09825	190	12	A	C	A	T	C	A	C	1,2,3,5,6,7,8,10,11,12,13,14
b	0.02695	58	9	A	A	A	T	C	A	C	1,3,4,5,9,10,11,12,14
c	0.01365	29	9	A	A	A	T	C	A	C	1,3,4,5,6,9,10,11,14
d	0.00865	18	4	T	C	A	T	C	A	C	1,5,10,14
e	0.00455	10	4	A	C	A	T	C	A	C	1,10,11,14
f	0.00225	5	3	T	A	A	T	C	A	C	1,7,10
k	0.00225	5	5	A	A	A	T	C	A	C	1,5,9,10,11
g	0.00225	5	3	A	A	A	T	C	A	C	1,5,11
n	0.0018	4	3	A	A	A	T	C	A	C	2,9,14
j	0.0009	2	1	A	C	A	T	C	A	C	14
h	0.0009	2	2	A	A	A	T	C	A	C	6,10
u	0.00045	1	1	A	C	A	T	C	A	C	14
t	0.00045	1	1	A	A	A	T	C	A	C	13
v	0.00045	1	1	A	A	A	C	G	A	C	8
s	0.00045	1	1	T	C	A	T	C	A	C	10
q	0.00045	1	1	A	A	A	T	C	A	C	14
r	0.00045	1	1	A	C	A	T	C	A	C	14
i	0.00045	1	1	A	A	A	T	C	C	C	9
m	0.00045	1	1	A	A	C	T	C	A	C	6
l	0.00045	1	1	A	A	A	T	C	A	C	10
p	0.00045	1	1	A	A	A	T	C	A	C	2
o	0.00045	1	1	A	A	A	T	C	A	C	2

Table 13.45 - IGHV3-30

h.type	cum.freq	# indivs	pops	14:106791117 rs138563161	14:106791127 rs190763974	14:106791150 rs183005459	14:106791154 rs187117835	14:106791216 rs192185012	
ref	0.97985	-	-	Ala->Val G	Lys->Glu T	Val->Ala A	Ala->Thr C	Thr->Ser G	ENST00000390613
a	0.0147	28	-		T	A	C	G	2,3,4,5,9,10,11
b	0.0023	5	4	G	T	A	C	<b>C</b>	1,2,9,13
d	0.00135	3	2	G	T	<b>G</b>	C	G	10,13
c	0.00135	3	3	G	<b>C</b>	A	C	G	3,9,11
e	0.00045	1	1	G	T	A	<b>T</b>	G	13

Table 13.46 - IGHV3-33

h.type	cum.fre	# indivs	pops	14:106815855 rs183940317	14:106815858 rs187379689	14:106815862 rs112373679	14:106815868 rs111734923	14:106815924 rs192341521	14:106815933 rs187304239	14:106815948 rs192234669	14:106815970 rs138488134
ref	0.82905	-	-	Asp->Gly T	Tyr->Ser T	Trp->Arg A	Val->Phe C	Ser->Asn C	Thr->Ile G	Ala->Val G	Arg->Gly T
a	0.145	243	12	T	T	<b>G</b>	<b>G</b>	C	G	G	<b>C</b>
b	0.00965	21	10	T	T	<b>G</b>	<b>G</b>	C	G	G	T
c	0.0082	18	6	T	T	A	C	C	G	G	<b>C</b>
e	0.0027	6	3	<b>C</b>	T	A	C	C	G	G	T
d	0.0018	4	3	T	T	A	C	C	<b>G</b>	G	T
h	0.00135	3	3	T	T	A	C	T	G	G	T
f	0.0009	2	2	T	<b>G</b>	A	C	C	G	G	T
j	0.00045	1	1	<b>C</b>	T	A	C	T	G	<b>G</b>	T
g	0.00045	1	1	<b>C</b>	T	<b>G</b>	<b>G</b>	C	G	G	<b>C</b>
i	0.00045	1	1	T	T	<b>G</b>	<b>G</b>	T	G	G	<b>C</b>

ENST00000390  
615

1,2,4,5,6,7,8,10,  
11,12,13,14  
1,2,5,6,7,10,11,  
12,13,14  
1,2,7,10,12,14  
1,10,14  
2,13,14  
1,6,10  
5,8  
10  
10  
1

Table 13.47 - IGHV3-35

h.type	cum.frec	# indivs	pops	14:106845417	14:106845487	14:106845753
ref	0.78085	-	-	rs112759551	rs74514616	rs145022218
a	0.11765	227	14	Arg->Gln	Gly->Arg	Val->Leu
b	0.09695	190	10	C	C	C
c	0.0041	9	5	T	G	G
d	0.00045	1	1	C	C	C

ENST00000390617

1,2,3,4,5,6,7,8,9,10,11,12,13,14

1,3,4,5,6,7,9,10,12,14

1,2,5,7,8

14

Table 13.48 - IGHV3-38

h.type	cum.freq	# indivs	pops	14:106866449 rs78850219	14:106866451 rs144366955	14:106866664 rs150283006	14:106866828 rs112981911	ENST00000390618
ref	0.8889	-	-	Asn->Ser T	Asn->Lys G	Leu->Phe C	Leu->Pro A	
a	0.0723	158	14	<b>C</b>	G	C	A	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.02515	52	6	<b>C</b>	<b>C</b>	C	A	1,5,10,11,12,14
c	0.00725	16	5	T	G	<b>G</b>	A	1,10,11,12,14
d	0.0064	13	3	T	G	C	<b>G</b>	2,7,13

Table 13.49 - IGHV3-43

h.type	cum.freq	# indivs	pops	14:106926211 rs116899367	14:106926223 rs2467912	14:106926231 rs182842897	14:106926323 rs186760979	14:106926328 rs61999676	14:106926388 rs2467910	14:106926435 rs182650845	14:106926454 rs188914311
ref	0.57745	-	-	Ala->Thr	Thr->Ala	Ser->Asn	Asp->Glu	Trp->Gly	Thr->Ala	Ser->Phe	Val->Met
a	0.2428	512	14	C	T	C	A	A	T	G	C
b	0.08365	164	13	C	C	C	A	A	C	G	C
c	0.0302	66	10	C	T	C	A	C	T	G	C
e	0.0182	40	11	C	T	C	A	A	C	G	C
d	0.01455	30	4	T	T	C	A	A	T	G	C
g	0.00995	22	7	C	T	C	A	C	T	G	C
h	0.0064	13	5	C	C	C	A	A	C	G	C
f	0.0055	12	5	C	T	C	A	A	T	■	C
i	0.00365	8	3	C	C	C	A	A	C	■	C
j	0.00225	5	4	C	T	C	A	C	C	G	C
k	0.00135	3	2	C	T	T	A	A	T	G	C
m	0.00135	3	3	C	C	C	A	A	T	G	C
p	0.0009	2	2	C	C	C	A	A	C	G	T
q	0.00045	1	1	C	T	C	T	A	T	G	C
n	0.00045	1	1	C	T	C	A	A	C	■	C
l	0.00045	1	1	C	T	C	A	A	C	G	C
o	0.00045	1	1	C	C	C	A	A	C	G	C



Table 13.49 - IGHV3-43

h.type	cum.freq	# indivs	pops	14:106926456 rs79008247	14:106926481 rs111739001	ENST00000434710
ref	0.57745	-	-	Val->Gly A	Val->Met C	
a	0.2428	512	14	A	C	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.08365	164	13	C	C	1,2,3,4,5,6,7,9,10,11,12,13,14
c	0.0302	66	10	C	C	1,3,4,5,6,7,9,11,12,13
e	0.0182	40	11	A	C	1,3,4,5,6,8,10,11,12,13,14
d	0.01455	30	4	A	C	2,5,6,7
g	0.00995	22	7	A	C	2,3,4,6,9,10,11
h	0.0064	13	5	A	T	1,2,6,7,13
f	0.0055	12	5	A	C	1,7,10,12,14
i	0.00365	8	3	A	C	1,10,14
j	0.00225	5	4	A	C	3,4,9,10
k	0.00135	3	2	A	C	10,14
m	0.00135	3	3	A	C	2,7,13
p	0.0009	2	2	A	C	1,10
q	0.00045	1	1	A	C	13
n	0.00045	1	1	A	C	10
l	0.00045	1	1	C	C	10
o	0.00045	1	1	C	C	9

Table 13.50 - IGHV3-48

h.type	cum.freq	# indivs	pops	14:106993845	14:106993938	14:106993945	14:106993996	14:106994231	14:106994237	14:106994262
ref	0.3919	-	-	rs7148408	.	rs7148607	rs183130922	rs192884486	rs184894692	rs188639638
				Asp->Ala	->Thr	Ser->Gly	Arg->Cys	Ile->Val	Val->Leu	Glu->Asp
a	0.3355	589	14	G	GTAC	T	G	T	C	C
b	0.20455	404	14	G	GTAC	C	G	T	C	C
c	0.027	59	11	G	G	T	G	T	C	C
d	0.01595	35	11	T	GTAC	C	G	T	C	C
f	0.0128	28	9	G	G	C	G	T	C	C
e	0.0105	23	10	T	G	T	G	T	C	C
j	0.00045	1	1	T	GTAC	T	■	T	C	C
g	0.00045	1	1	T	GTAC	T	G	T	G	C
h	0.00045	1	1	G	GTAC	T	G	C	C	C
i	0.00045	1	1	G	GTAC	T	G	T	C	G

ENST00000390624  
 1,2,3,4,5,6,7,8,9,10,11,12,13,14  
 1,2,3,4,5,6,7,8,9,10,11,12,13,14  
 1,2,4,5,6,7,10,11,12,13,14  
 1,2,3,4,5,6,7,8,9,10,13  
 1,3,4,6,7,9,10,11,14  
 2,3,4,5,6,7,10,11,13,14  
 4  
 11  
 5  
 2  
 1

Table 13.51 - IGHV3-49

h.type	cum.frec	# indivs	pops	14:107012994 rs140678750	14:107013065 rs140027079	14:107013072 rs191590980	14:107013093 rs187070171	14:107013129 rs2073674	14:107013146 rs184357967	14:107013171 rs193139111	14:107013201 rs2073673
				Tyr->His	Gly->Glu	Tyr->Asp	Gly->Ser	Phe->Val	Asp->Ala	Thr->Ala	Gln->Lys
ref	0.322	-	-	A	C	A	C	A	T	T	G
a	0.38705	663	14	A	C	A	C	C	T	T	G
b	0.28555	499	13	A	C	A	C	A	T	T	T
c	0.0027	6	3	G	C	A	C	A	T	T	G
d	0.00045	1	1	A	C	C	T	C	G	T	G
g	0.00045	1	1	A	C	A	C	A	T	C	G
e	0.00045	1	1	G	C	A	C	C	T	T	G
h	0.00045	1	1	A	C	A	C	C	T	C	G
f	0.00045	1	1	A	C	A	C	C	T	T	G
i	0.00045	1	1	A	T	A	C	A	T	T	G

Table 13.51 - IGHV3-49

h.type	cum.frec	# indivs	pops	14:107013209 rs184512071	14:107013210 rs189168917	
ref	0.322	-	-	Gly->Asp	Gly->Ser	ENST00000390625
a	0.38705	663	14	C	C	
b	0.28555	499	13	C	C	1,2,3,4,5,6,7,8,9,10,11,12,13,14
c	0.0027	6	3	C	C	1,2,3,4,5,6,7,8,9,10,11,12,13
d	0.00045	1	1	C	C	7,11,13
g	0.00045	1	1	C	C	2
e	0.00045	1	1	C	C	1
h	0.00045	1	1	C	C	4
f	0.00045	1	1	T	T	1
i	0.00045	1	1	C	C	2
						10

Table 13.52 - IGHV3-53

h.type	cum.freq	# indivs	pops	14:107048717 rs141949273	14:107048749 rs11627238	14:107048795	14:107048796 rs182065680	14:107048929 rs35608607	14:107048944 rs35850880	14:107049080 rs2731152	
ref	0.4053	-	-	Met->Ile	Asp->His	Ser->	Ser->Asn	Ile->Val	Ser->Thr	Leu->Ser	ENST00000390627
a	0.2535	483	14	C	C	G	C	T	A	A	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.2264	399	13	C	G	G	C	C	A	A	1,2,3,4,5,6,7,8,10,11,12,13,14
c	0.0453	99	13	C	C	G	C	T	T	G	1,2,3,4,5,6,7,9,10,11,12,13,14
d	0.0247	54	6	T	C	G	C	T	A	A	1,5,10,12,13,14
e	0.015	33	9	C	C	G	C	C	T	G	1,2,5,7,8,10,11,12,14
f	0.0092	20	6	C	C	G	C	T	A	A	3,4,9,10,11,12
g	0.0046	10	5	C	C	G	C	C	A	G	7,10,11,13,14
k	0.0041	9	8	C	G	G	C	T	A	A	3,6,7,9,11,12,13,14
h	0.0036	8	2	T	C	G	C	C	A	A	10,14
j	0.0023	5	4	C	G	G	C	C	A	A	1,3,4,9
i	0.0018	4	4	C	C	G	C	T	A	G	8,11,13,14
n	0.0014	3	2	C	C	G	C	T	T	G	2,10
m	0.0014	3	3	C	C	G	C	C	A	A	5,9,10
q	0.0005	1	1	C	C	G	C	C	T	A	14
l	0.0005	1	1	C	G	G	T	C	A	A	3
p	0.0005	1	1	C	G	G	C	T	A	A	4
o	0.0005	1	1	T	C	G	C	C	T	G	14

Table 13.53 - IGHV3-64

h.type	cum.frec	# indivs	pops	14:107113770 rs187347757	14:107113851 rs2072045	14:107113877 rs190575084	14:107113902 rs138428007	14:107114009 rs11846079	14:107114176 rs182438044	14:107114189 rs139161493	14:107114193 rs2073670
				Glu->Gln	Asp->Asn	Ser->Asn	Leu->Val	Glu->Gly	Trp->Gly	Glu->Asp	Met->Thr
ref	0.4221	-	-	C	C	C	G	T	A	C	A
a	0.47965	793	14	C	T	C	G	C	A	C	G
b	0.06225	131	13	C	C	C	G	C	A	C	A
c	0.02785	61	14	C	C	C	G	C	A	C	G
d	0.00275	6	4	C	T	C	G	T	A	C	G
e	0.0018	4	2	C	T	C	G	C	C	C	G
f	0.00135	3	1	C	T	C	C	C	A	C	G
h	0.0009	2	1	C	T	C	G	C	A	C	A
j	0.00045	1	1	C	C	C	G	T	A	G	A
g	0.00045	1	1	G	C	C	G	T	A	C	A
i	0.00045	1	1	C	T	T	G	T	A	C	A

Table 13.53 - IGHV3-64

h.type	cum. freq	# indivs	pops	
ref	0.4221	-	-	ENST00000454421
a	0.47965	793	14	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.06225	131	13	1,2,3,4,5,6,7,9,10,11,12,13,14
c	0.02785	61	14	1,2,3,4,5,6,7,8,9,10,11,12,13,14
d	0.00275	6	4	2,4,13,14
e	0.0018	4	2	1,10
f	0.00135	3	1	10
h	0.0009	2	1	12
j	0.00045	1	1	2
g	0.00045	1	1	3
i	0.00045	1	1	8

Table 13.54 - IGHV3-66

h.type	cum.freq	# indivs	pops	14:107131089 rs77593173	14:107131164 rs6423677	14:107131223 rs183864069	14:107131290 rs149638514	ENST00000390632
ref	0.43755	-	-	Tyr->His	Cys->Gly	Met->Lys	Ile->Val	
a	0.27865	502	14	A	A	A	T	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.12575	252	14	A	C	A	C	1,2,3,4,5,6,7,8,9,10,11,12,13,14
c	0.0649	140	13	G	A	A	C	1,2,3,4,5,6,7,9,10,11,12,13,14
d	0.03975	87	13	G	C	A	C	1,2,3,4,5,6,7,9,10,11,12,13,14
e	0.02745	60	14	G	A	A	T	1,2,3,4,5,6,7,8,9,10,11,12,13,14
f	0.021	46	14	A	C	A	T	1,2,3,4,5,6,7,8,9,10,11,12,13,14
g	0.0027	6	3	G	C	A	T	10,13,14
h	0.0009	2	2	G	C	T	C	3,4
j	0.00045	1	1	A	A	T	T	5
k	0.00045	1	1	A	C	T	C	4
i	0.00045	1	1	A	A	T	C	3



Table 13.55 - IGHV3-72

h.type	cum.frec	# indivs	pops	14:107198966 rs193196315	14:107199062 rs184752334	14:107199136 rs139291045	14:107199164 rs191373077	14:107199222 rs188380722	14:107199355 rs192371094	ENST00000433072
ref	0.9973	-	-	Lys->Arg T	Ser->Asn C	His->Gln G	Ala->Gly G	Leu->Val G	Ile->Leu T	
a	0.00045	1	1	C	T	G	G	G	T	5
d	0.00045	1	1	T	C	C	C	G	T	2
e	0.00045	1	1	C	C	G	G	G	T	11
c	0.00045	1	1	T	C	G	G	C	T	2
b	0.00045	1	1	T	C	G	C	G	T	1
f	0.00045	1	1	T	C	G	G	G	G	10

Table 13.56 - IGHV3-73

h.type	cum.freq	# indivs	pops	14:107211000 rs142927366	14:107211135 rs184224767	14:107211176 rs61750833	14:107211199 rs188470998	14:107211371 rs61752554	
ref	0.91625	-	-	Lys->Glu T	Ala->Pro C	Lys->Arg T	Leu->Phe C	Trp->Cys C	ENST00000390636
a	0.05265	96	6	T	C	T	C	<b>G</b>	1,5,10,12,13,14
b	0.0293	60	5	T	C	<b>C</b>	C	C	1,7,9,10,14
c	0.00135	3	3	T	C	T	<b>■</b>	C	1,10,14
d	0.00045	1	1	<b>C</b>	<b>G</b>	T	C	C	2

Table 13.57 - IGHV3-74

h.type	cum.freq	# indivs	pops	14:107218737 rs184901617	14:107218809 rs183213717	14:107218812 rs140462183	14:107218858 rs191764284	14:107218878 rs183293744	14:107218908 rs149597490	
				Thr->Met	Asp->Gly	Ser->Thr	Arg->Cys	Ser->Asn	Ser->Cys	
ref	0.98635	-	-	G	T	C	G	C	G	ENST000000424969
a	0.0091	20	-	G	T	<b>G</b>	G	C	G	1,9,10,12,13,14
b	0.0023	5	-	■	T	C	G	C	G	2,3,5,11
c	0.0009	2	-	G	T	C	G	<b>T</b>	G	10,14
d	0.00045	1	-	G	T	C	■	C	G	1
e	0.00045	1	-	G	<b>C</b>	C	G	C	G	11
f	0.00045	1	-	G	T	C	G	C	<b>C</b>	10

Table 13.58 - IGHV3-7

h.type	cum.fre	# indivs	pops	14:106518545 rs185279906 Asn->Ser	14:106518602 rs185538237 Ser->Asn	14:106518686 rs180765266 Gln->Arg	14:106518840 rs190666315 Leu->Arg	ENST00000390598
ref	0.9808	-	-	T	C	T	A	
a	0.016	29	5	T	C	<b>C</b>	A	1,5,10,11,14
b	0.0023	5	4	T	<b>T</b>	T	A	5,9,13,14
d	0.00045	1	1	<b>C</b>	C	T	A	4
c	0.00045	1	1	T	C	T	<b>C</b>	13

Table 13.59 - IGHV3-9

h.type	cum.fr	# indivs	pops	14:106552299 rs188663119 Tyr->His	14:106552310 rs8020204 Thr->Met	14:106552409 rs140101036 Ile->Thr	14:106552412 rs10141136 Ser->Asn	14:106552415 rs185242411 Gly->Val	14:106552416 rs149492969 Gly->Ser	14:106552436 rs192281729 Ser->Leu	14:106552463 rs182143003 Ala->Val
ref											
a	0.84305	-	-	A	G	A	C	C	C	G	G
b	0.10975	167	9	A		A	C	C	C	G	G
c	0.0228	46	7	A	G	A	C	C	C	G	G
d	0.0064	12	5	A	G	<b>G</b>	C	C	C	G	G
e	0.00405	6	4	A	G	A	C	C	C	G	G
f	0.00315	7	7	A	G	A	T	C	C	G	G
g	0.00225	5	4	A	G	A	C	C	C	G	
h	0.0018	4	3	A	G	A	C	C	C	G	G
i	0.00135	2	1	A	G	A	C	C	C		G
j	0.0009	1	1	A	G	A	T	C	T	G	G
k	0.0009	2	1	<b>G</b>	G	A	C	C	C	G	G
l	0.00045	1	1	A		A	T	C	C	G	G
m	0.00045	1	1	A		A	C	C	C	G	G
n	0.00045	1	1	A	G	A	C	C	C	G	G
o	0.00045	1	1	A		A	C	C	T	G	G
p	0.00045	1	1	A	G	A	C	C	C	G	G
q	0.00045	1	1	A		A	C	C	T	G	G
r	0.00045	1	1	A		A	C	C	C	G	G
s	0.00045	1	1	A	G	A	C	C	C	G	G
t	0.00045	1	1	A		A	C	C	T	G	G
u	0.00045	1	1	A		A	C	C	T	G	G
v	0.00045	1	1	A		A	C	C	T	G	G
w	0.00045	1	1	A		A	C	C	T	G	G
x	0.00045	1	1	A		A	C	C	T	G	G
y	0.00045	1	1	A		A	C	C	T	G	G
z	0.00045	1	1	A		A	C	C	T	G	G

Table 13.59 - IGHV3-9

h.type	cum.fr	# indivs	pops	14:106552464 rs186818749 Ala->Pro	14:106552472 rs191640600 Val->Ala	14:106552491 rs182569652 Asp->Asn	14:106552493 rs141844186 Asp->Gly	14:106552713 rs116484141 Ser->Asn	ENST000000390600
ref	0.84305	-	-	C	A	C	T	C	
a	0.10975	167	9	C	A	C	T	C	1,5,7,8,10,11,12,13,14
b	0.0228	46	7	C	A	C	<b>C</b>	C	3,4,6,7,9,11,13
c	0.0064	12	5	C	A	C	T	C	1,7,10,12,14
e	0.00405	6	4	C	A	<b>T</b>	T	C	1,3,12,14
d	0.00315	7	7	C	A	C	T	C	1,2,3,5,6,8,10
f	0.00225	5	4	C	A	C	T	C	6,7,9,13
g	0.0018	4	3	<b>G</b>	A	C	T	C	2,10,14
h	0.00135	2	1	C	A	C	T	C	1
r	0.0009	1	1	C	A	C	T	C	2
o	0.0009	2	1	C	A	C	T	C	13
j	0.00045	1	1	C	A	C	T	C	10
k	0.00045	1	1	<b>G</b>	A	C	T	C	10
q	0.00045	1	1	C	A	C	T	<b>T</b>	2
i	0.00045	1	1	C	A	C	T	C	14
n	0.00045	1	1	C	<b>G</b>	C	T	C	6
m	0.00045	1	1	C	A	C	T	C	11
l	0.00045	1	1	C	A	C	T	C	14
p	0.00045	1	1	C	A	C	<b>C</b>	C	3

Table 13.60 - IGHV4-28

h.type	cum.fr	# indiv	pops	14:106780542 rs113785338 Val->Leu	14:106780632 rs112792995 Tyr->Asn	14:106780634 rs34629512 Thr->Ile	14:106780667 rs8010702 Glu->Ala	14:106780759 rs61994196 Asp->Glu	14:106780760 rs181692209 Asp->Gly	14:106780797 rs186483759 Leu->Val	14:106780817 rs8009554 Leu->TrpLeu->Trp
ref	0.22145	-	-	C	A	G	T	G	T	G	A
a	0.41435	691	14	C	A	G	T	G	T	G	C
b	0.13715	250	14	C	A		T	G	T	G	C
c	0.07385	161	14	C	A	G	T	C	T	G	C
d	0.06485	136	13	C	A		T	C	T	G	C
f	0.0195	42	5		T	G	T	G	T	G	C
e	0.01685	37	10	C	A	G	T	C	T	G	A
g	0.01355	29	4		T	G	T	C	T	G	C
i	0.01185	22	6	C	A	G	G	G	T	G	C
h	0.01175	25	11	C	A		T	G	T	G	A
j	0.00405	9	5	C	A		T	C	T	G	A
k	0.00315	7	3		T	G	T	C	C	G	C
p	0.00225	5	3	C	A	G	G	C	T	G	C
s	0.0009	2	2	C	A	G	T	G	C	G	C
n	0.0009	2	2		A	G	T	C	T	G	C
u	0.00045	1	1	C	T	G	T	G	T	G	C
t	0.00045	1	1		A	G	T	C	T	G	A
v	0.00045	1	1	C	A	G	T	G	T	C	C
q	0.00045	1	1		T	G	T	C	C	G	A
r	0.00045	1	1		T	G	T	C	C	G	C
m	0.00045	1	1		T	G	T	C	T	G	C
l	0.00045	1	1		T	G	T	C	T	G	A
o	0.00045	1	1		T	G	T	G	T	G	A

Table 13.60 - IGHV4-28

h.type	cum. fre	# indivs	pops	ENST00000390612
ref	0.22145	-	-	
a	0.41435	691	14	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.13715	250	14	1,2,3,4,5,6,7,8,9,10,11,12,13,14
c	0.07385	161	14	1,2,3,4,5,6,7,8,9,10,11,12,13,14
d	0.06485	136	13	1,2,3,5,6,7,8,9,10,11,12,13,14
f	0.0195	42	5	1,10,11,12,14
e	0.01685	37	10	1,2,5,6,7,10,11,12,13,14
g	0.01355	29	4	1,5,10,14
i	0.01185	22	6	1,8,10,11,13,14
h	0.01175	25	11	1,2,3,5,7,8,9,10,11,13,14
j	0.00405	9	5	2,6,7,10,13
k	0.00315	7	3	1,10,14
p	0.00225	5	3	1,10,14
s	0.0009	2	2	1,11
n	0.0009	2	2	10,14
u	0.00045	1	1	10
t	0.00045	1	1	11
v	0.00045	1	1	13
q	0.00045	1	1	11
r	0.00045	1	1	1
m	0.00045	1	1	10
l	0.00045	1	1	10
o	0.00045	1	1	14



Table 13.61 - IGHV4-31

h.type	cum.fr	# indivs	pops	14:106805289 rs188101179 Val->Leu	14:106805303 rs145562667 Arg->Leu	14:106805321 rs192498675 Asn->Ser	14:106805345 rs185638910 Tyr->Ser	14:106805358 rs190512954 Ile->Leu	14:106805381 rs77489245 His->Pro	14:106805388 rs149858616 Arg->Cys	14:106805408 rs61995642 Gly->Asp
ref	0.01175	-	-	C	T	T	T	T	T	G	C
a	0.5475	886	14	C	T	T	T	T	T	G	C
b	0.2035	420	14	C	T	T	T	T	G	G	T
c	0.1554	303	14	C	T	T	T	T	G	G	C
d	0.04565	83	11	C	T	T	T	T	T	G	C
e	0.00865	18	11	C	T	T	T	T	T	G	T
f	0.00865	19	7	C	T	T	T	T	G	G	T
i	0.0054	10	3	C	T	T	T	T	T	■	C
g	0.00315	7	4	C	T	T	T	T	T	G	C
j	0.00225	5	3	C	T	T	T	T	G	G	C
h	0.00135	3	2	C	T	T	T	T	T	■	C
t	0.0009	2	2	C	T	T	G	T	T	G	C
m	0.0009	2	2	C	T	T	T	T	G	G	C
l	0.0009	2	2	C	T	T	T	T	G	G	T
u	0.00045	1	1	C	T	T	T	G	T	G	C
k	0.00045	1	1	C	C	C	T	T	T	G	C
v	0.00045	1	1	C	T	T	G	T	G	G	T
s	0.00045	1	1	C	T	T	T	T	G	G	T
q	0.00045	1	1	G	T	T	T	T	T	G	C
r	0.00045	1	1	C	T	T	T	T	G	G	C
n	0.00045	1	1	C	T	T	T	T	T	G	C
p	0.00045	1	1	C	C	C	G	T	G	G	C
o	0.00045	1	1	G	T	T	T	T	T	G	C

Table 13.61 - IGHV4-31

h.type	cum.fr	# indivs	pops	14:106805411 rs185736587 Gly->Asp	14:106805421 rs188982921 Ile->Phe	14:106805497 rs181045179 Gln->His	14:106805508 rs4462488 Pro->Ser	ENST00000438142
ref	0.01175	-	-	C	T	C	G	
a	0.5475	886	14	C	T	C		1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.2035	420	14	C	T	C		1,2,3,4,5,6,7,8,9,10,11,12,13,14
c	0.1554	303	14	C	T	C		1,2,3,4,5,6,7,8,9,10,11,12,13,14
d	0.04565	83	11	C	T	C		1,2,5,6,7,8,10,11,12,13,14
e	0.00865	18	11	C	T	C		1,2,3,4,6,9,10,11,12,13,14
f	0.00865	19	7	C	T	C	G	1,3,6,10,11,12,14
i	0.0054	10	3	T		C		1,10,14
g	0.00315	7	4	C	T	C	G	1,2,5,7
j	0.00225	5	3	C	T	C	G	2,10,14
h	0.00135	3	2	C	T	C		1,10
t	0.0009	2	2	C	T	C		4,14
m	0.0009	2	2	C	T	C		1,10
l	0.0009	2	2	C	T	C		1,10
u	0.00045	1	1	C	T	C		4
k	0.00045	1	1	C	T	C		13
v	0.00045	1	1	C	T	C		1
s	0.00045	1	1	C	T	G		5
q	0.00045	1	1	C	T	C		5
r	0.00045	1	1	C	T	G		2
n	0.00045	1	1	C	T	C		7
p	0.00045	1	1	C	T	C		10
o	0.00045	1	1	C	T	C		5

Table 13.62 - IGHV4-34

h.type	cum.freq	# indivs	pops	14:106829649 rs183232225	14:106829717 rs11546808	14:106829767 rs188280613	14:106829775 rs191592421	14:106829793 rs11546811	14:106829817 rs1064183	14:106829874 rs181039074	ENST00000390616
ref	0.9955	-	-	Ser->Phe	Ser->Arg	Pro->Ser	Ile->Ser	Gly->Ala	Ala->Asp	Leu->Pro	
d	0.0009	2		G	G	G	A	C	G	A	
d	0.0009	2		G	G	G	A	<b>G</b>	G	A	2,14
g	0.0009	2		G	<b>T</b>	G	A	C	G	A	11,14
b	0.0009	2		G	G	G	A	C	G	<b>G</b>	3,10
a	0.00045	1		G	G	G	<b>C</b>	C	G	A	6
e	0.00045	1		G	G	<b>■</b>	A	C	G	A	5
c	0.00045	1		<b>■</b>	G	G	A	C	G	A	7
f	0.00045	1		G	G	G	A	C	<b>T</b>	A	2

Table 13.63 - IGHV4-39

h.type	cum. freq	# indivs	pops	14:106877626 rs188639181	14:106877648 rs191739990	14:106877705 rs187288552	14:106877730 rs191860623	14:106877753 rs35281264	14:106877755 rs186830319	14:106877761 rs115523671	14:106877831 rs190205925
ref	0.6445	-	-	Cys->Tyr	Ala->Thr	Ile->Leu	Asn->Lys	Tyr->His	Tyr->Ser	Ser->Asn	Ile->Phe
a	0.10295	220	14	C	C	T	G	A	T	C	T
b	0.0918	197	14	C	C	T	G	G	T	C	T
d	0.04425	93	13	C	C	T	G	A	T	C	T
c	0.02555	56	10	C	C	T	G	G	T	C	T
e	0.01595	35	12	C	C	T	G	A	T	T	T
f	0.0132	29	10	C	C	T	G	G	T	C	T
g	0.0105	23	9	C	C	T	G	G	T	C	T
h	0.00595	13	8	C	C	T	G	G	T	C	T
i	0.0059	13	7	C	C	T	G	A	T	C	T
j	0.0055	12	8	C	C	T	G	A	T	C	T
m	0.00415	9	5	C	C	T	G	G	T	C	T
k	0.00365	8	5	C	C	T	G	G	T	C	T
l	0.00275	6	6	C	C	T	G	G	T	T	T
y	0.0018	4	3	C	C	T	G	A	T	C	T
n	0.0018	4	3	C	C	T	G	G	T	C	T
o	0.0018	4	3	C	C	T	G	G	T	C	T
u	0.00135	3	3	C	C	T	G	A	T	C	T
t	0.00135	3	2	C	C	T	G	G	T	C	T
q	0.00135	3	3	C	C	T	G	A	T	C	T
aa	0.0009	2	2	C	C	T	G	G	T	C	T
s	0.0009	2	2	C	C	T	G	G	T	C	T
x	0.0009	2	2	C	C	T	G	G	T	C	T
p	0.0009	2	2	C	C	T	G	A	T	T	T
ah	0.00045	1	1	C	C	T	G	A	T	C	T
ag	0.00045	1	1	C	C	T	G	A	T	T	T
aj	0.00045	1	1	C	C	T	G	G	T	T	T
aq	0.00045	1	1	C	T	T	G	A	T	C	T
v	0.00045	1	1	C	C	T	G	G	T	C	T
al	0.00045	1	1	C	C	T	C	A	T	C	T



Table 13.63 - IGHV4-39

h.type	cum. freq	# indivs	pops	14:106877836 rs143740707	14:106877837 rs145759866	14:106877848 rs181859654	14:106877877 rs181228803	14:106877888 rs190955722	14:106877907 rs144078801	14:106877912 rs74093494	14:106878019 rs140461651
ref	0.6445	-	-	C	C	G	C	C	C	G	Ala->Thr
a	0.10295	220	14	C	C	G	C	C	C	C	C
b	0.0918	197	14	T	C	G	C	C	C	C	C
d	0.04425	93	13	C	C	G	C	C	C	G	C
c	0.02555	56	10	C	C	G	C	C	C	C	C
e	0.01595	35	12	C	C	G	C	C	C	G	C
f	0.0132	29	10	C	C	G	C	C	C	G	C
g	0.0105	23	9	C	C	G	C	C	C	G	C
h	0.00595	13	8	C	C	G	C	C	C	C	C
i	0.0059	13	7	C	C	G	C	C	C	C	C
j	0.0055	12	8	C	C	G	C	C	C	C	C
m	0.00415	9	5	C	C	G	C	C	C	C	C
k	0.00365	8	5	T	C	G	C	C	C	G	C
l	0.00275	6	6	C	C	G	C	C	C	C	C
y	0.0018	4	3	T	C	G	C	C	C	G	C
n	0.0018	4	3	C	C	G	C	C	C	G	C
o	0.0018	4	3	T	C	G	C	C	C	C	C
u	0.00135	3	3	C	C	G	C	C	C	G	C
t	0.00135	3	2	T	C	G	C	C	C	G	C
q	0.00135	3	3	C	C	G	C	C	C	G	C
aa	0.0009	2	2	C	C	G	C	C	C	G	C
s	0.0009	2	2	C	C	G	C	C	C	C	C
x	0.0009	2	2	T	C	G	C	C	C	C	C
p	0.0009	2	2	C	C	G	C	C	C	C	C
ah	0.00045	1	1	C	C	G	C	C	C	G	C
ag	0.00045	1	1	C	C	G	C	C	C	C	C
aj	0.00045	1	1	T	C	G	C	C	C	C	C
aq	0.00045	1	1	C	C	G	C	C	C	G	C
v	0.00045	1	1	T	C	G	C	C	C	G	C
al	0.00045	1	1	C	C	G	C	T	C	G	C

Table 13.63 - IGHV4-39

ab	0.00045	C	C	G	C	C	C	C	C	C	C	C	C	C	C	C	C	C
ak	0.00045	C	C	G	C	C	C	C	C	C	C	C	C	C	C	C	C	C
ap	0.00045	C	C	G	C	C	C	C	C	C	C	C	C	C	C	C	C	C
z	0.00045	C	C	G	C	C	C	C	C	C	C	C	C	C	C	C	C	C
w	0.00045	T	C	G	C	C	C	C	C	C	C	C	C	C	C	C	C	C
ao	0.00045	T	C	G	C	C	C	C	C	C	C	C	C	C	C	C	C	C
r	0.00045	C	C	G	C	C	C	C	C	C	C	C	C	C	C	C	C	C
af	0.00045	C	C	G	C	C	C	C	C	C	C	C	C	C	C	C	C	C
ad	0.00045	T	C	G	C	C	C	C	C	C	C	C	C	C	C	C	C	C
am	0.00045	C	C	G	C	C	C	C	C	C	C	C	C	C	C	C	C	C
at	0.00045	C	C	C	C	C	C	C	C	C	C	C	C	C	C	C	C	C
ac	0.00045	C	C	C	C	C	C	C	C	C	C	C	C	C	C	C	C	C
ar	0.00045	C	C	G	C	C	C	C	C	C	C	C	C	C	C	C	C	C
as	0.00045	T	C	G	C	C	C	C	C	C	C	C	C	C	C	C	C	T
ae	0.00045	T	C	G	C	C	C	C	C	C	C	C	C	C	C	C	C	C
ai	0.00045	C	C	G	C	C	C	C	C	C	C	C	C	C	C	C	C	C
an	0.00045	T	C	G	C	C	C	C	C	C	C	C	C	C	C	C	C	C

Table 13.63 - IGHV4-39

h.type	cum.freq	# indivs	pops	14:106878048 rs187579450	14:106878052 rs138364008	14:106878056 rs4774113	14:106878057 rs190089302	14:106878067 rs4774114	ENST00000390619
ref	0.6445	-	-	His->Arg	Lys->Gln	Lys->Asn	Lys->Thr	Met->Leu	
a	0.10295	220	14	T	T	G	T	T	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.0918	197	14	T	T	G	T	G	1,2,3,4,5,6,7,8,9,10,11,12,13,14
d	0.04425	93	13	T	T	G	T	G	1,2,3,4,5,6,7,9,10,11,12,13,14
c	0.02555	56	10	T	T	T	T	T	1,2,3,5,6,7,10,11,13,14
e	0.01595	35	12	T	T	T	T	T	1,2,3,4,5,6,9,10,11,12,13,14
f	0.0132	29	10	T	T	T	T	T	1,2,3,4,5,6,7,10,11,14
g	0.0105	23	9	T	T	G	T	G	1,2,4,5,10,11,12,13,14
h	0.00595	13	8	T	T	T	T	G	1,2,6,7,8,10,13,14
i	0.0059	13	7	T	T	G	T	G	1,2,4,8,10,13,14
j	0.0055	12	8	T	T	T	T	T	1,2,3,4,5,9,11,12
m	0.00415	9	5	T	T	G	T	T	2,3,7,10,14
k	0.00365	8	5	T	T	G	T	G	2,3,4,9,13
l	0.00275	6	6	T	T	G	T	G	1,2,5,7,12,13
y	0.0018	4	3	T	T	T	T	T	2,5,13
n	0.0018	4	3	T	T	G	T	T	3,13,14
o	0.0018	4	3	T	T	T	T	T	4,6,13
u	0.00135	3	3	T	T	T	T	G	1,4,14
t	0.00135	3	2	T	T	T	T	T	3,4
q	0.00135	3	3	T	T	G	T	T	2,3,6
aa	0.0009	2	2	T	T	T	T	G	1,6
s	0.0009	2	2	T	T	G	T	G	7,8
x	0.0009	2	2	T	T	G	T	G	3,4
p	0.0009	2	2	T	T	G	T	G	5,14
ah	0.00045	1	1	C	T	T	T	T	13
ag	0.00045	1	1	T	T	T	T	T	11
aj	0.00045	1	1	T	T	T	T	G	3
aq	0.00045	1	1	T	T	G	T	G	2
v	0.00045	1	1	T	T	T	T	T	7
al	0.00045	1	1	T	T	G	T	G	14



Table 13.63 - IGHV4-39

1	0.00045	T	T	T	T	T	T	4
1	0.00045	T	T	T	T	T	T	13
1	0.00045	T	T	T	T	T	T	13
1	0.00045	T	T	T	T	G	T	10
1	0.00045	T	T	T	T	G	T	10
1	0.00045	T	T	T	T	T	T	12
1	0.00045	T	T	T	T	T	T	3
1	0.00045	T	T	T	T	T	T	12
1	0.00045	T	T	T	T	T	T	5
1	0.00045	T	T	T	T	T	G	13
1	0.00045	T	T	T	T	T	T	1
1	0.00045	T	T	T	T	T	T	13
1	0.00045	T	T	T	T	T	T	6
1	0.00045	T	T	T	T	G	T	2
1	0.00045	T	T	T	T	G	T	9
1	0.00045	T	T	T	G	T	T	13
1	0.00045	T	T	T	T	G	T	2

Table 13.64 - IGHV4-4

h.type	cum.freq	# indivs	pops	14:106478120 rs150123115	14:106478183 rs77628366	14:106478194 rs76899215	14:106478252 rs73365408	14:106478311	14:106478354 rs188216744	
				Tyr->Cys	Thr->Lys	Met->Ile	Arg->His	Ser->	Glu->Gly	
ref	0.34	-	-	T	G	C	C	A		ENST00000390596
a	0.1136	222	14	T	G	C	C		C	1,2,3,4,5,6,7,8,9,10,11,12,13,14
d	0.0817	166	14	T	G	T	C		T	1,2,3,4,5,6,7,8,9,10,11,12,13,14
f	0.06675	138	14	T	G	T	C		C	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.06385	140	13	T	G	T	C	A	T	1,2,3,4,5,6,7,9,10,11,12,13,14
c	0.0599	127	14	T	G	C	C		T	1,2,3,4,5,6,7,8,9,10,11,12,13,14
e	0.05215	110	12	T	T	T	C		C	1,2,3,4,5,6,7,9,11,12,13,14
g	0.03255	65	10	C	T	T	C		C	1,2,3,4,5,9,10,11,12,14
m	0.0267	58	12	T	T	T	T		C	1,2,3,4,5,6,7,9,10,12,13,14
i	0.0262	58	14	T	T	T	C		T	1,2,3,4,5,6,7,8,9,10,11,12,13,14
j	0.01715	38	11	T	T	T	C	A	T	1,2,4,5,6,7,9,10,11,12,14
h	0.01535	33	11	T	G	C	C	A	C	2,3,4,5,6,7,9,10,11,12,13
k	0.0113	25	10	C	G	C	C		C	2,3,4,5,6,8,9,11,12,13
o	0.00855	19	9	T	T	C	C		C	1,2,3,4,5,6,7,8,13
q	0.00765	17	7	C	G	C	C	A	T	1,3,9,10,11,12,14
n	0.00765	16	5	T	G	T	T	A	T	1,5,10,13,14
p	0.00765	16	5	T	G	T	T		T	1,5,9,10,14
l	0.00685	15	4	C	G	C	C	A	C	3,4,9,10
s	0.00585	13	4	T	G	C	T	A	T	1,10,12,14
aj	0.0054	12	8	T	G	T	C	A	C	2,3,4,5,9,12,13,14
ah	0.00495	11	7	T	T	T	T		T	1,2,3,9,10,13,14
t	0.0045	10	6	C	G	T	C		C	3,4,6,8,9,11
w	0.00405	9	7	C	T	T	C		T	3,4,7,9,11,13,14
r	0.0036	8	7	T	G	T	T		C	2,4,9,11,12,13,14
v	0.00315	7	5	C	G	C	C		T	7,9,11,13,14
ag	0.0027	6	5	T	T	T	C	A	C	2,5,6,10,14
y	0.0027	6	4	C	T	C	C		C	3,4,5,9
u	0.0027	6	3	T	T	C	T	A	T	1,10,14
z	0.0027	6	4	C	T	T	T		C	2,4,9,10

Table 13.64 - IGHV4-4

ak	0.0018	4	T	G	C	T	T	T	T	T	3,4,5,10
ao	0.00135	3	C	G	T	C	C	C	T	T	3,9,11
ac	0.00135	3	T	G	C	T	T	T	C	C	10
ae	0.00135	3	T	T	C	C	C	A	C	C	2,4
aa	0.0009	2	C	T	T	T	C	A	T	T	9,14
af	0.0009	2	T	T	C	T	T		C	C	6,9
x	0.0009	2	C	G	T	C	C	A	T	T	9,12
ad	0.0009	2	T	T	C	C	C		T	T	3,7
ai	0.0009	2	T	T	C	C	C	A	T	T	7
al	0.00045	1	C	G	C	T	T		C	C	4
ab	0.00045	1	C	T	C	T	T		C	C	10
am	0.00045	1	C	G	T	T	T	A	C	C	9
an	0.00045	1	C	G	C	T	T	A	C	C	1

Table 13.65 - IGHV4-59

h.type	cum.freq	# indvs	pops	14:107082722	14:107083453	14:107083457
ref	0.95005	-	-	rs181388142	rs112607697	rs190037133
a	0.04905	102		Ser->Leu	Tyr->His	Ser->Arg
c	0.00045	1		G	Tyr->His	Ser->Arg
b	0.00045	1		G	G	A
				G	A	A
				█	A	A

ENST000000390629
ENST000000455737
1,2,5,10,12,14
1
14

Table 13.66 - IGHV4-61

h.type	cum.freq	# indivs	pops	14:107095319 rs11864170	14:107095325 rs186969565	14:107095326 rs1064309	14:107095338 rs2072046	ENST00000390630
ref	0.0109	-	-	Tyr->Ser	Gly->Asp	Gly->Ser	Val->Ile	
a	0.5701	806	14	T	C	C	C	
b	0.18115	339	14	T	C	T	T	1,2,3,4,5,6,7,8,9,10,11,12,13,14
c	0.13535	275	13	<b>G</b>	C	C	T	1,2,3,4,5,6,7,8,9,10,11,12,13,14
f	0.05855	120	14	T	C	C	T	1,2,3,4,5,6,7,8,9,10,11,12,13,14
e	0.0191	42	12	<b>G</b>	C	C	C	2,3,4,5,6,7,9,10,11,12,13,14
d	0.0109	24	10	<b>G</b>	C	T	C	1,2,3,6,7,9,10,11,12,13
g	0.00675	15	4	T	T	C	T	1,9,10,14
h	0.00585	13	7	<b>G</b>	C	T	T	2,3,6,7,9,10,14
j	0.00045	1	1	T	T	C	C	7
k	0.00045	1	1	T	T	T	T	10
i	0.00045	1	1	<b>G</b>	T	C	T	14

Table 13.67 - IGHV5-51

h.type	cum.freq	# indivs	pops	14:107034771 rs180672762	14:107034793 rs181977500	14:107034796 rs186695491	14:107034813 rs191700780	14:107034850 rs142572296	14:107034851 rs186314656	14:107034868 rs139665108	14:107034910 rs117410356
				Ser->Arg	Ser->Asn	Ile->Ser	Ile->Met	Thr->Ser	Thr->Ala	Tyr->Phe	Arg->His
ref	0.97045	-	-	G	C	A	G	G	T	T	C
a	0.0164	35	3	G	C	A	G	G	T	T	T
b	0.0055	11	3	G	C	A	G	G	T	T	C
g	0.00135	3	3	G	T	A	G	G	T	T	C
d	0.0009	2	2	G	C	A	G	C	T	T	C
k	0.0009	2	2	G	C	A	G	G	T	T	C
c	0.0009	2	2	G	T	A	G	G	T	T	C
j	0.00045	1	1	G	C	A	G	G	T	T	C
e	0.00045	1	1	G	C	A	C	G	T	T	C
h	0.00045	1	1	G	C	A	G	G	T	T	C
f	0.00045	1	1	G	C	A	G	G	T	T	C
i	0.00045	1	1	G	C	A	G	G	T	T	C
n	0.00045	1	1	G	C	C	G	G	T	T	C
m	0.00045	1	1	T	C	A	C	G	C	T	C
l	0.00045	1	1	G	C	A	G	G	T	T	C

Table 13.67 - IGHV5-51

h.type	cum.freq	# indivs	pops	14:107034931 rs113154616	14:107034940 rs113988349	14:107034998 rs183518874	14:107035123 rs116300992	14:107035141 rs147695427	ENST00000390626
ref	0.97045	-	-	Ser->Thr	Ser->Asn	Ala->Pro	Leu->Ile	Ala->Thr	
a	0.0164	35	3	C	C	C	G	C	3,4,9
b	0.0055	11	3	C	C	C	T	C	5,10,14
g	0.00135	3	3	C	C	C	G	C	6,7,13
d	0.0009	2	2	C	C	C	G	C	2,13
k	0.0009	2	2	G	C	C	G	C	6,10
c	0.0009	2	2	C	T	C	G	C	2,10
j	0.00045	1	1	C	C	C	G	C	13
e	0.00045	1	1	C	C	C	G	C	10
h	0.00045	1	1	C	C	C	G	T	7
f	0.00045	1	1	G	T	C	G	C	10
i	0.00045	1	1	C	T	C	G	C	10
n	0.00045	1	1	C	C	C	G	C	4
m	0.00045	1	1	G	C	C	G	C	4
l	0.00045	1	1	C	C	G	G	C	6

Table 13.68 - IGHV6-1

h.type	cum.freq	# indvs	pops	14:106405816 rs186349163	14:106405877 rs72715486	14:106405910 rs188294175	ENST00000390593
ref	0.99775	-	-	Ser->Asn C	Lys->Glu T	Val->Ile C	
a	0.0009	2	2	C	C	C	3,8
c	0.0009	2	1	C	T	T	4
b	0.00045	1	1	T	T	C	8



Table 13.69 - IGHV7-81

h.type	cum.frec	# indivs	pops	14:107282845 rs61741324	14:107282853 rs61741323	14:107282887 rs61741308	14:107283009 rs11845591	14:107283018 rs144292230	14:107283073 rs148758470	
ref	0.9127	-	-	Leu->Met G	Thr->Ile G	Arg->Trp G	Gly->Val C	Lys->Met T	Val->Leu C	ENST00000390639
a	0.0732	136	6	T		G	C	T	C	1,5,10,12,13,14
b	0.01005	21	3	G	G	G		T	C	1,10,14
c	0.0018	4	3	G	G		C	T	C	1,5,14
e	0.00135	3	1	G	G	G	C		C	10
d	0.0009	2	2	G	G	G	C	T		1,10

Table 13.70 - IGKJ2

h.type	cum.freq	# indivs	pops	2:89161040 rs76129343	2:89161044 rs75187701	2:89161045 rs146019185	2:89161050 rs181828141	2:89161055 rs183845604	2:89161067 rs150749817	2:89161069 rs113458930	2:89161072 rs113539064	
ref	0.921	-	-	Lys->Glu	Glu->Asp	Glu->Gly	Lys->Asn	Thr->Pro	Phe->Val	Thr->Ser	Tyr->Cys	ENST00000390241
a	0.0449	86	13	T	C	T	C	T	A	G	T	1,2,3,4,5,6,7,9,10,11,12,13,14
b	0.0243	48	10	T	C	T	C	T	A	C	T	2,4,5,6,7,8,10,11,12,13
d	0.0032	6	3	T	C	T	C	T	A	G	C	2,6,7
e	0.0018	4	3	T	C	C	C	T	A	G	T	2,10,13
c	0.0018	4	4	T	C	T	G	T	A	G	T	1,2,7,9
f	0.0014	3	2	T	C	T	C	T	C	G	T	2,10
g	0.0009	2	2	T	C	T	C	T	A	G	T	11,14
h	0.0005	1	1	C	C	T	C	T	C	G	C	9
i	0.0005	1	1	T	C	T	C	G	A	G	T	10

Table 13.71 - IGKJ3

h.type	cum.frec	# indivs	pops	2:89160742 rs188533155	2:89160748 rs181878952	2:89160762 rs190475170	2:89160769 rs139877244	ENST00000390240
ref	0.99045	-	-	Asp->His	Lys->Glu	Phe->Cys	Phe->Leu	
a	0.00685	15	10	C	T	A	A	
d	0.00135	3	3	C	T	A	<b>G</b>	2,4,5,6,9,10,11,12,13,14
b	0.0009	2	2	<b>G</b>	<b>C</b>	A	A	5,11,12
c	0.00045	1	1	C	T	<b>C</b>	A	2,3
								5

Table 13.72 - IGKJ4

h.type	cum.fre	# indivs	pops	2:89160399 rs78964890	2:89160404 rs139461207	2:89160405 rs180910647	2:89160406 rs185131270	2:89160407 rs145002796	2:89160409 rs191518742	2:89160412 rs147540982	2:89160415 rs139402168	2:89160427 rs186462161
ref	0.9883	-	-	Lys->Asn	Ile->Leu	Glu->Asp	Glu->Gly	Glu->Gln	Val->Ala	Lys->Arg	Thr->Ile	Phe->Ser
b	0.00225	5	4	T	T	C	T	C	A	T	G	A
a	0.0018	4	4	T	T	C	T	C	A	<b>C</b>	G	A
g	0.00135	3	3	■	T	C	T	C	A	T	G	A
j	0.0009	2	1	T	T	C	<b>C</b>	C	A	T	G	A
k	0.0009	2	2	T	<b>G</b>	C	T	C	A	T	G	A
c	0.0009	2	2	T	T	C	T	C	A	T	G	A
h	0.0009	2	2	T	T	C	T	<b>G</b>	A	T	G	A
d	0.00045	1	1	T	T	C	T	C	A	T	■	A
e	0.00045	1	1	T	T	C	T	C	A	T	G	<b>G</b>
f	0.00045	1	1	T	T	C	T	C	<b>G</b>	T	G	A
i	0.00045	1	1	T	T	<b>G</b>	T	C	A	T	G	A
m	0.00045	1	1	T	T	C	T	C	A	<b>C</b>	G	A
l	0.00045	1	1	■	T	C	T	C	A	T	G	A

Table 13.72 - IGKJ4

h.type	cum.fre	# indivs	pops	2:89160428 rs191712435	2:89160430 rs187905157	ENST00000390239
ref	0.9883	-	-	Phe->Leu	Thr->Ile	
b	0.00225	5	4	A	G	
a	0.0018	4	4	A	G	2,5,13,14
g	0.00135	3	3	A	■	2,5,6,10
j	0.0009	2	1	A	G	3,4,10
k	0.0009	2	2	A	G	10
c	0.0009	2	2	G	G	9,10
h	0.0009	2	2	A	G	2,14
d	0.00045	1	1	A	G	2,10
e	0.00045	1	1	A	G	5
f	0.00045	1	1	A	G	6
i	0.00045	1	1	A	G	14
m	0.00045	1	1	G	G	9
l	0.00045	1	1	G	G	9
						13

Table 13.73 - IGKJ5

h.type	cum.fre	# indvs	pops	2:89160082 rs185692630	2:89160087 rs142632901	2:89160106 rs190985433	2:89160109 rs140799895	2:89160110 rs114105515	2:89160113 rs188703994	ENST00000390238
ref	0.9928	-	-	Lys->Arg T	Glu->Asp C	Gly->Ala C	Phe->Cys A	Phe->Leu A	Thr->Ser T	
a	0.00225	5	4	C	C	C	A	A	T	4,5,6,14
c	0.00135	3	3	T	C	C	A	A	T	9,10,11
b	0.00135	3	3	T	G	C	A	G	T	3,5,9
d	0.00045	1	1	T	G	C	A	G	T	10
g	0.00045	1	1	T	C	C	A	A	■	10
e	0.00045	1	1	T	C	G	A	A	T	10
h	0.00045	1	1	C	G	C	A	A	T	14
f	0.00045	1	1	T	C	C	C	A	T	3

Table 13.74 - IGKV1-16

h.type	cum.frec	# indivs	pops	2:89399455 rs193208674	2:89399488 rs184390186	2:89399509 rs116145759	2:89399515 rs2848410	2:89399809 rs188494311	ENST00000479981
ref	0.97625	-	-	Lys->Arg T	Ala->Val G	Ala->Val G	Gly->Glu C	Ala->Thr C	
a	0.01145	25	8	C	G	G	T	C	1,4,6,7,8,9,11,13
b	0.00595	13	10	C	G	G	C	C	2,4,5,6,7,8,9,10,13,14
c	0.00455	10	7	T	G	G	T	C	2,5,6,9,11,13,14
d	0.0009	2	2	T	G	■	C	C	1,14
e	0.00045	1	1	T	G	G	C	T	9
f	0.00045	1	1	T	■	G	C	C	7

Table 13.75 - IGKV1-17

h.type	cum.freq	# indivs	pops	2:89416951 rs185657549	2:89417000 rs188985676	2:89417011 rs191430092	2:89417047 rs188555641	2:89417087 rs181054320	
ref	0.83295	-	-	Ser->Thr C	Pro->Ser G	Tyr->Phe T	Arg->Gln C	Leu->Met G	ENST00000490686
a	0.08655	170	13	C	G	T	C	T	1,2,3,4,5,6,7,9,10,11,12,13,14
b	0.0545	111	12	C	G	■	C	T	1,2,3,4,5,7,9,10,11,12,13,14
c	0.0224	47	11	C	G	■	C	G	1,2,4,5,6,7,9,10,12,13,14
d	0.00225	4	1	C	■	T	C	G	10
e	0.0009	2	2	G	G	T	C	G	2,10
f	0.00045	1	1	C	G	T	T	G	7



Table 13.76 - IGKV1-5

h.type	cum.freq	# indivs	pops	2:89246874 rs11546105	2:89246904 rs182050897	2:89246946 rs11546098	2:89247012 rs146302149	2:89247045 rs191761898	2:89247051 rs182340840	2:89247106 rs189714283	ENST00000496168
ref	0.9823	-	-	Ser->Asn C	Ser->Tyr G	Ser->Asn C	Ser->Thr C	Thr->Ser G	Arg->Thr C	Cys->Gly A	
a	0.0073	16	8	C	G	C	G	G	C	A	1,6,8,9,11,12,13,14
b	0.005	11	8	C	G	T	C	G	C	A	1,2,3,6,7,11,12,14
f	0.0018	4	4	T	G	C	C	G	C	A	7,10,13,14
c	0.00135	3	3	C	G	T	G	G	C	A	10,12,13
d	0.0009	2	1	C	G	C	C	G	C	C	10
g	0.00045	1	1	C	G	C	C	G	G	A	11
e	0.00045	1	1	C	G	C	C	C	C	A	11
h	0.00045	1	1	C	T	C	C	G	C	A	14

Table 13.77 - IGKV1-6

<b>h.type</b>										
ref	0.9955	-	-	-						
a	0.0045	1	1	1						10
<b>cum.fre</b>										
<b># indvs</b>										
<b>pops</b>										
	2:89265959									
	rs188876010									
	Tyr->Cys									
	T									
	C									
										ENST00000464162

Table 13.78 - IGKV1-8

h.type	cum.frec	# indivs	pops	2:89291968 rs187771921	2:89292080 rs190853230	2:89292130 rs183046670	2:89292149 rs186879210	2:89292182 rs59175543	2:89292188 rs192180011	2:89292201 rs35262290	
ref	0.60715	-	-	Asp->Ala	Lys->Glu	Gly->Asp	Thr->Ser	Phe->Leu	Ser->Pro	Met->Ile	ENST00000495489
a	0.20315	348	14	T	T	C	T	A	A	C	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.1334	235	14	T	T	C	T	<b>G</b>	A	<b>T</b>	1,2,3,4,5,6,7,8,9,10,11,12,13,14
c	0.03325	70	14	T	<b>C</b>	C	T	A	A	C	1,2,3,4,5,6,7,8,9,10,11,12,13,14
e	0.01035	23	8	T	<b>C</b>	C	T	<b>G</b>	A	C	1,2,3,4,5,6,7,8,9,10,11,12,13,14
d	0.00955	21	8	T	<b>C</b>	C	T	<b>G</b>	A	<b>T</b>	1,6,7,10,11,12,13,14
h	0.0009	2	2	T	T	<b>T</b>	T	<b>G</b>	A	<b>T</b>	1,3,4,5,7,9,10,14
j	0.00045	1	1	<b>G</b>	T	C	T	A	A	C	9,10
k	0.00045	1	1	T	<b>C</b>	<b>T</b>	T	<b>G</b>	A	C	7
g	0.00045	1	1	T	T	C	<b>■</b>	<b>G</b>	A	C	1
f	0.00045	1	1	T	T	C	T	<b>G</b>	A	C	14
i	0.00045	1	1	T	T	<b>T</b>	T	<b>G</b>	<b>G</b>	<b>T</b>	5
				T	T	<b>T</b>	T	A	A	C	11

Table 13.79 - IGKV1-9

h.type	cum.freq	# indvs	pops	2:89309554 rs147109427	2:89309694 rs185997823	2:89309735 rs80322626	2:89309762 rs77800356	ENST00000493819
ref	0.7266	-	-	Glu->Asp T	Arg->Trp G	Phe->Ser A	Asp->Ala T	
a	0.1854	317	14		G	G	G	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.02735	57	10	T	G	A	G	1,3,4,5,7,9,10,11,13,14
c	0.0197	43	13		G	A	G	1,2,3,4,5,6,7,9,10,11,12,13,14
d	0.01825	40	13		G	A	T	1,2,3,4,5,7,8,9,10,11,12,13,14
e	0.01095	24	11	T		A	T	2,4,5,6,7,8,9,10,11,12,13
f	0.00545	12	7	T	G	G	G	1,3,4,5,7,10,11
g	0.0018	4	4	T	G	G	T	1,10,11,14
h	0.0018	4	4	T		A	G	3,4,10,11
j	0.00135	3	3		G	G	T	5,12,13
i	0.0009	2	2			G	G	4,14
k	0.00045	1	1			A	G	4

Table 13.80 - IGKV1D-12

h.type	cum.freq	# indivs	pops	2:90198951 rs189385596	2:90199107 rs180730188	ENST00000390276
ref	0.9955	-	-	Gly->Arg G	Gly->Arg G	
b	0.0027	6	4	G		1,7,10,12
a	0.0018	4	4		G	2,8,13,14

Table 13.81 - IGKV1D-16

h.type	cum.freq	# indivs	pops	2:90139373 rs6760987	2:90139377 rs184550323	2:90139422 rs189369337	ENST00000492446
ref	0.96165	-	-	Ser->Arg T	Gly->Ser G	Ala->Thr G	
a	0.03655	71	13	<b>G</b>	G	G	1,2,3,4,5,6,7,8,10,11,12,13,14
c	0.0009	2	1	T	G	<b>■</b>	1
b	0.0009	2	1	T	<b>■</b>	G	10

Table 13.82 - IGKV1D-17

h.type	cum.fre	# indvs	pops	2:90121919 rs182773099	2:90121961 rs191806633	2:90121999 rs182889841	2:90122063 rs193008187	ENST00000483379
ref	0.99315	-	-	Arg->Gln	Gln->Arg	Ala->Ser	Thr->Ile	
a	0.0037	8	4	G	A	G	C	
c	0.00135	3	2	G	A	T	C	1,5,10,14
b	0.00135	3	2	■	G	G	C	7,12
d	0.00045	1	1	G	A	G	C	1,10
							T	6

Table 13.83 - IGKV1D-42

h.type	cum.freq	# indivs	pops	2:90229083 rs193096486	2:90229247 rs139321345	2:90229260 rs842173	2:90229265 rs145538558	2:90229292 rs147724164	2:90229314 rs192596253	2:90229471 rs184070665	2:90229477 rs188365115	2:90229491 rs181576535
ref	0.7584	-	-	Ala->Thr	Asp->Tyr	Ile->Thr	Ser->Pro	Gly->Arg	Cys->Phe	Ile->Met	Ser->Arg	Asp->Val
a	0.22345	383	14	G	G	T	T	G	G	C	C	A
b	0.01005	21	3	G	G	C	T	G	G	C	C	A
c	0.0027	6	1	G	G	T	T	G	G	C	C	A
d	0.00135	3	2	G	G	T	T	G	G	C	C	A
e	0.00135	3	1	G	T	T	T	G	G	C	C	A
g	0.0009	2	2	G	G	T	T	G	G	G	C	A
f	0.0009	2	1	G	G	T	T	G	T	C	C	A
h	0.00045	1	1	G	G	C	T	G	G	C	■	A
i	0.00045	1	1	G	G	T	T	G	G	C	C	T



Table 13.83 - IGKV1D-42

h.type	cum.freq	# indivs	pops	ENST00000390278
ref	0.7584	-	-	
a	0.22345	383	14	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.01005	21	3	1,10,14
c	0.0027	6	1	9
d	0.00135	3	2	10,14
e	0.00135	3	1	13
g	0.0009	2	2	12,14
f	0.0009	2	1	14
h	0.00045	1	1	13
i	0.00045	1	1	3

Table 13.84 - IGKV1D-43

h.type	cum.freq	# indivs	pops	2:90248957	2:90249120	2:90249147	2:90249151	2:90249237	2:90249247	2:90249348	2:90249372
ref	0.98095	-	-		rs188075270	rs184607123	rs138753337	rs142796670	rs192607496	rs184789392	rs188397844
b	0.00685	14	3	Leu->Gln	Met->Val	Ala->Thr	Ser->Phe	Ala->Thr	Leu->Arg	Pro->Ser	Cys->Arg
a	0.00635	12	4	T	A	G	C	G	T	C	
c	0.0027	5	3	T	A	G	C	G	T	C	ENST00000468879
d	0.0009	2	2	T	A	G	C	G	T	C	ENST00000560950
e	0.0009	2	1	T	A	G	C	G	T	T	
g	0.00045	1	1	T	G	G	C	G	T	T	1,10,14
h	0.00045	1	1	T	A	G	C	G	T	C	1,10,11,14
f	0.00045	1	1	T	A	G	C	G	T	C	1,6,14
											1,14
											11
											2
											2
											14

Table 13.85 - IGKV1D-8

h.type	cum.freq	# indivs	pops	2:90259956 rs190239413	2:90259965 rs842156	2:90260022 rs17699428	2:90260033 rs183140988	2:90260085 rs186043133	2:90260091 rs190501712	2:90260178 rs138677672
ref	0.7813	-	-	Ala->Val	Val->Ala	Thr->Ile	Arg->Trp	Gly->Glu	Ala->Val	Thr->Ile
a	0.1872	319	14	C	T	C	C	G	C	C
b	0.01785	36	3	C	C	C	C	G	C	C
c	0.0105	22	6	C	T	T	C	G	C	T
d	0.0009	2	1	C	C	C	C	G	T	C
f	0.0009	2	2	T	T	C	C	G	C	C
g	0.00045	1	1	C	T	C	C	G	C	C
e	0.00045	1	1	C	T	C	T	G	C	C
h	0.00045	1	1	C	T	C	C	G	T	T

ENST00000471857

1,2,3,4,5,6,7,8,9,10,11,12,13,14

3,4,9

2,5,6,7,12,13

1

2,12

1

1

9

Table 13.86 - IGKV2-24

h.type	cum.frec	# indivs	pops	2:89475831 rs149900655	2:89476044 rs183417517	ENST00000484817
ref	0.8082	-	-	Met->Thr A	Cys->Phe C	
a	0.1708	293	13	G	C	1,2,3,4,5,6,7,9,10,11,12,13,14
b	0.02055	44	13	G		1,2,3,4,5,6,7,8,9,10,12,13,14
c	0.00045	1	1	A		8

Table 13.87 - IGKV2-30

h.type	cum.fre	# indvs	pops	2:89544389	2:89544392	2:89544473	2:89545042
ref	0.75005	-	-	rs141762272	rs182126464	rs147079411	rs186725977
a	0.24085	437	13	Arg->Trp	Asn->Asp	Tyr->His	Leu->Arg
b	0.0082	18	7	G	T	A	A
d	0.00045	1	1	G	T	G	A
c	0.00045	1	1	G	C	A	A
							ENST00000468494
							1,2,3,4,5,6,7,9,10,11,12,13,14
							2,4,6,7,9,12,13
							11
							11

Table 13.88 - IGKV2D-24

h.type	cum.frec	# indivs	pops	2:90044207	2:90044426	ENST00000462693
ref	0.9863	-	-	rs150942262	rs190167237	
a	0.01235	27	12	Phe->Cys T	Ala->Val C	1,2,3,4,6,7,8,9,11,12,13,14
b	0.00135	3	2	G T	C T	5,10

Table 13.89 - IGKV2D-26

h.type	cum.freq	# indivs	pops	2:90025217 rs187527217	2:90025275 rs147187346	2:90025397 rs192055191	2:90025483 rs184819753	
ref	0.6664	-	-	Ile->Asn	Met->Ile	Gly->Glu	Tyr->Asn	ENST00000390268
a	0.325	556	14	T	G	G	T	
b	0.00455	9	3	T	C	G	T	1,2,3,4,5,6,7,8,9,10,11,12,13,14
c	0.0018	4	1	█	G	G	█	2,11,12
d	0.0009	2	2	T	C	G	█	10
f	0.0009	2	2	T	G	█	T	3,4
e	0.00045	1	1	█	C	G	T	1,10

Table 13.90 - IGKV2D-29

h.type	cum.frec	# indivs	pops	2:89986777	2:89986852	2:89986921	ENST00000491977
ref	0.6969	-	-	Ala->Thr	Ser->Pro	Pro->Ser	
a	0.30175	481	14	G	T	C	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.0009	2	1	G	T	T	13
c	0.00045	1	1	G	C	T	12



Table 13.91 - IGKV2D-30

h.type	cum.freq	# indivs	pops	2:89975711	2:89976341	2:89976353	2:89976364
ref	0.9689	-	-	rs185844838	rs148937425	rs184425514	rs147056147
a	0.02335	51	14	Leu->His	Arg->His	Lys->Thr	Trp->Arg
b	0.00595	13	4	->	Arg->His	Lys->Thr	Trp->Arg
c	0.00135	3	2	T	G	A	T
d	0.00045	1	1	T	G	C	T

ENST00000474213	ENST00000558962
1,2,3,4,5,6,7,8,9,10,11,12,13,14	1,2,3,4,5,6,7,8,9,10,11,12,13,14
1,5,10,14	1,5,10,14
1,14	1,14
14	14

Table 13.92 - IGKV3-11

h.type	cum.fr	# indiv	pops	2:89326669 rs182958807 Pro->His	2:89326681 rs185920904 Arg->His	2:89326754 rs191612627 Ser->Pro	2:89326858 rs182380932 Tyr->Ser	2:89326861 rs191953548 Ser->Thr	2:89326865 rs183365521 Ser->Gly	2:89326872 rs189180740 Gln->His
ref	0.94435	-	-	G	C	A	T	C	T	C
a	0.02835	60	14	T	C	G	T	C	T	C
b	0.0096	21	11	G	C	G	T	C	T	C
d	0.0073	14	3	G	T	A	T	C	T	C
c	0.00545	12	7	T	C	A	T	C	T	C
e	0.0018	4	4	G	C	A	T	G	T	C
g	0.0009	2	1	G	T	G	T	C	T	C
f	0.0009	2	2	G	C	A	T	C	T	G
j	0.00045	1	1	G	C	A	G	C	C	C
h	0.00045	1	1	G	C	A	T	C	C	C
i	0.00045	1	1	G	C	A	G	C	T	C

ENST00000483158  
 1,2,3,4,5,6,7,8,9,10,11,12,13,14  
 2,3,4,5,6,7,8,9,11,13,14  
 1,10,14  
 1,6,7,9,11,13,14  
 2,5,7,9  
 14  
 9,13  
 14  
 13  
 9

Table 13.93 - IGV3-20

h.type	cum.fr	# indivs	pops	2:89442183 rs1065533 Ser->Arg	2:89442191 rs190424065 Ala->Thr	2:89442193 rs181820749 Gly->Ala	2:89442196 rs185401231 Tyr->Phe	2:89442253 rs180810535 Ser->Asn	2:89442255 rs11546074 Ser->Arg	2:89442262 rs191417237 Ser->Thr	2:89442295 rs183453470 Glu->Ala	2:89442342 rs190965861 Glu->Asp
ref			0.9847	G	C	C	T	C	G	C	T	T
a		5	0.00225	G	C	C	T	C	G	G	T	T
b		5	0.00225	G	C	C	T	T	G	C	T	T
d		3	0.00135	G	C	C	T	C	G	C	T	T
f		3	0.00135	G	C	C	T	C	G	C	T	T
j		2	0.0009	G	C	C	T	C	C	C	T	T
e		2	0.0009	G	C	C	T	C	C	C	T	T
c		2	0.0009	G	C	G	T	C	G	C	T	T
i		2	0.0009	G	C	C	T	C	G	C	T	T
k		1	0.00045	C	T	C	T	C	G	G	T	T
g		1	0.00045	G	C	C	■	C	G	C	T	T
q		1	0.00045	G	T	C	T	C	G	C	T	T
r		1	0.00045	G	C	C	T	C	G	C	G	T
h		1	0.00045	G	C	C	■	C	G	C	G	T
n		1	0.00045	G	C	C	T	C	G	C	T	T
m		1	0.00045	G	C	C	T	C	G	C	T	T
l		1	0.00045	G	C	G	T	C	G	G	T	T
p		1	0.00045	G	C	C	T	T	C	C	T	T
o		1	0.00045	G	C	C	T	C	G	C	T	■

Table 13.93 - IGKV3-20

h.type	cum.fr	# indivs	pops	2:89442344 rs183373822	2:89442350 rs187696503	2:89442352 rs182188819	2:89442354 rs187989142	2:89442355 rs192831943	ENST00000492167
ref	0.9847	-	-	Glu->Gln	Thr->Ser	Thr->Ile	Asp->Glu/Asp->Glu	Asp->Ala/Asp->Ala	
a	0.00225	5	3	C	T	G	A	T	1,9,14
b	0.00225	5	4	C	T	G	A	T	5,7,10,14
d	0.00135	3	3	C	T	■	A	T	2,6,14
f	0.00135	3	2	C	■	G	A	T	1,14
j	0.0009	2	2	C	T	■	A	T	1,10
e	0.0009	2	2	C	T	G	A	T	9,13
c	0.0009	2	2	C	T	G	A	T	10,11
i	0.0009	2	2	C	T	G	T	T	9,14
k	0.00045	1	1	C	T	G	A	T	11
g	0.00045	1	1	G	T	G	A	T	14
q	0.00045	1	1	C	T	G	A	T	13
r	0.00045	1	1	C	T	G	A	T	9
h	0.00045	1	1	C	T	G	A	T	14
n	0.00045	1	1	C	T	G	A	G	10
m	0.00045	1	1	G	T	G	A	T	13
l	0.00045	1	1	C	T	G	A	T	10
p	0.00045	1	1	C	T	G	A	T	2
o	0.00045	1	1	C	T	G	A	T	4

Table 13.94 - IGKV3-7

h.type	cum.fre	# indivs	pops	2:89278103 rs144594870 Ser->Gly	2:89278106 rs189211039 Thr->Ser	2:89278219 rs2458828 Val->Ala	2:89278262 rs181802934 Thr->Ala	
ref	0.78325	-	-	T	T	A	T	ENST00000390247
a	0.2018	349	14	C	T	A	T	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.01	19	3	T	T	G	T	1,10,14
c	0.00225	5	3	T	T	A	T	1,10,14
d	0.00225	5	3	T	T	G	C	1,10,14
e	0.00045	1	1	C	T	G	T	9

Table 13.95 - IGKV3D-11

h.type	cum.fr	# indivs	pops	2:90212050 rs58164053 Gly->Ser	2:90212051 rs188947989 Gly->Val	2:90212126 rs7578124 Asn->Thr	2:90212138 rs191678148 Gly->Val	2:90212167 rs185410352 Pro->Ser	2:90212198 rs189915304 Ser->Asn	2:90212240 rs117046277 Arg->His	2:90212243 rs181647122 Ser->Ile
ref	0.635	-	-	G	G	A	G	C	G	G	G
a	0.3511	561	14	G	G	A	G	C	G	G	G
c	0.0036	8	5	G	G	A	G	T	G	G	G
b	0.0023	5	3	G	G	A	G	C	G	G	G
g	0.0018	3	2	G	G	A	G	C	G	G	T
f	0.0018	4	3	G	G	A	G	C	G	G	G
d	0.0014	3	2	G	G	A	G	C	G	G	G
h	0.0009	2	1	G	G	C	G	C	G	G	G
i	0.0009	2	1	G	G	A	T	C	G	G	G
j	0.0005	1	1	G	T	A	G	T	G	G	G
k	0.0005	1	1	G	T	A	G	C	G	G	G
e	0.0005	1	1	G	G	A	G	T	G	G	G

ENST000000390277  
1,2,3,4,5,6,7,8,9,10,11,12,13,14  
4,6,7,8,9  
3,4,9  
1,10  
1,3,4  
10,14  
4  
10  
14  
9  
1



Table 13.97 - IGKV3D-20

h.type	cum.freq	# indivs	pops	2:90077781 rs183765055	2:90078037 rs189772344	2:90078079 rs182769673	2:90078093 rs2555982	ENST00000390270
ref	0.9845	-	-	Glu->Gly A	Thr->Met C	Ala->Gly C	Gly->Arg G	
a	0.0073	16	9	A	C	C		1,2,4,6,7,8,9,10,11
b	0.0064	12	4	<b>G</b>	C	C	G	1,10,11,14
d	0.0009	2	2	A	<b>T</b>	C		7,12
e	0.00045	1	1	A	<b>T</b>	C	G	11
c	0.00045	1	1	A	C	<b>G</b>	G	13



Table 13.98 - IGV4-1

h.type	cum.freq	# indivs	pops	2:89185362 rs180851255	2:89185392 rs116328981	2:89185400 rs148071875	2:89185431 rs141793384	2:89185450 rs189675840	2:89185451 rs138603647	2:89185455 rs181816666	2:89185457 rs141709704	2:89185461 rs78280375
ref				Tyr->Phe	Asp->Gly	Ala->Thr	Asn->Ser	Ser->Arg	Val->Ile	Leu->Ser	Tyr->His	Ser->Asn
a	0.9516	-	-	A	A	G	A	T	G	T	T	G
b	0.01185	26	11	A	A	G	A	T	G	T	T	G
c	0.00545	11	9	A	A	G	A	T	G	T	T	G
d	0.00365	8	7	A	A	G	A	T	G	T	T	G
k	0.00315	7	6	A	A	G	A	T	G	T	T	G
w	0.0018	4	4	A	A	G	A	T	G	T	T	G
h	0.0018	3	3	A	A	G	A	T	G	T	T	G
f	0.0018	4	4	T	A	G	A	T	G	T	C	G
e	0.00135	3	3	A	A	G	A	T	G	T	T	G
r	0.00135	3	3	A	A	G	A	T	G	T	T	G
x	0.00135	3	2	A	A	G	A	T	G	T	T	G
o	0.00135	3	2	A	A	G	A	T	G	T	T	G
y	0.0009	2	2	A	A	G	A	T	G	T	T	G
g	0.0009	2	2	A	A	G	A	T	G	T	T	G
t	0.0009	2	2	A	A	G	A	T	G	T	T	G
i	0.0009	2	2	A	A	G	A	T	G	T	T	G
n	0.0009	2	2	A	A	G	A	T	G	T	T	G
l	0.0009	2	2	A	A	G	A	T	G	T	T	G
j	0.00045	1	1	A	A	G	G	T	G	T	T	G
u	0.00045	1	1	A	A	G	A	T	G	T	T	G
aa	0.00045	1	1	A	A	G	A	T	G	C	T	G
v	0.00045	1	1	A	A	G	A	T	G	T	T	G
s	0.00045	1	1	A	A	G	A	T	G	T	T	G
ab	0.00045	1	1	A	A	G	A	T	G	T	T	G
q	0.00045	1	1	T	A	G	A	T	G	T	T	G
z	0.00045	1	1	A	A	G	A	T	G	T	T	G
af	0.00045	1	1	A	A	G	A	T	G	T	T	G
ad	0.00045	1	1	A	A	G	A	T	G	T	T	G

Table 13.98 - IGV4-1

ac	0.00045	1	A	A	G	A	T	G	T	T	T	G
m	0.00045	1	A	A	G	A	T	G	T	T	T	G
p	0.00045	1	A	G	G	A	T	G	T	T	T	G
ae	0.00045	1	A	A	G	A	T	G	T	T	T	G

Table 13.98 - IGKV4-1

h.type	cum.freq	# indivs	pops	2:89185464 rs139176691	2:89185466 rs186837913	2:89185474 rs142402063	2:89185476 rs191518466	2:89185482 rs143877691	2:89185484 rs148599313	2:89185485 rs142986022	2:89185491 rs140954172	2:89185511 rs186251122
				Ser->Phe	Asn->His	Lys->Asn	Asn->Ser	Leu->Ser	Ala->Pro	Ala->Val	Tyr->Phe	Pro->Thr
ref	0.9516	-	-	C	A	G	A	T	G	C	A	C
a	0.01185	26	11	C	A	G	A	T	G	C	A	C
b	0.00545	11	9	C	A	G	A	T	G	C	A	C
c	0.00365	8	7	C	A	G	A	T	G	T	A	C
d	0.00315	7	6	C	A	G	A	T	G	C	A	C
k	0.0018	4	4	C	A	G	A	T	G	C	A	C
w	0.0018	3	3	C	A	G	A	T	C	C	A	C
h	0.0018	4	4	C	A	G	A	T	G	C	A	C
f	0.0018	4	4	C	A	G	A	T	G	C	A	C
e	0.00135	3	3	C	A	G	A	T	G	C	T	C
r	0.00135	3	3	C	A	G	A	T	G	C	A	C
x	0.00135	3	2	C	A	G	A	T	G	C	A	C
o	0.00135	3	2	C	A	G	G	T	G	C	A	C
y	0.0009	2	2	C	A	C	A	T	C	C	A	C
g	0.0009	2	2	C	A	G	A	T	G	C	A	C
t	0.0009	2	2	C	A	C	A	T	G	C	A	C
i	0.0009	2	2	C	A	G	A	T	G	C	A	C
n	0.0009	2	2	C	A	G	A	T	G	C	A	C
l	0.0009	2	2	C	A	G	A	T	G	C	A	C
j	0.00045	1	1	C	A	G	A	T	G	C	A	C
u	0.00045	1	1	C	A	C	A	T	G	C	A	C
aa	0.00045	1	1	C	A	G	A	T	G	C	A	C
v	0.00045	1	1	C	A	G	A	T	G	C	A	C
s	0.00045	1	1	C	A	G	A	T	G	C	A	C
ab	0.00045	1	1	C	A	G	A	T	G	T	A	C
q	0.00045	1	1	C	C	G	A	T	G	C	A	C
z	0.00045	1	1	C	A	G	A	T	G	C	A	C
af	0.00045	1	1	C	A	G	A	C	G	C	A	C
ad	0.00045	1	1	C	A	G	A	T	G	C	A	C

Table 13.98 - IGKV4-1

ac 0.00045  
 m 0.00045  
 p 0.00045  
 ae 0.00045

C	A	G	A	T	G	C	A	C
C	A	G	A	T	G	C	T	C
C	A	G	A	T	G	C	A	C
T	A	C	A	T	G	C	A	C

Table 13.98 - IGKV4-1

h.type	cum.freq	# indivs	pops	2:89185518 rs141640067	2:89185529 rs145197543	2:89185530 rs188057788	2:89185563 rs192839829	2:89185577 rs138751137	2:89185586 rs141777505	2:89185607 rs182919936	2:89185622 rs188769218	2:89185637 rs145449257
ref	0.9516	-	-	Lys->Arg	Tyr->Asp	Tyr->Cys	Asp->Gly	Ser->Arg	Gly->Arg	Ile->Phe	Ala->Thr	Val->Leu
a	0.01185	26	11	A	T	A	A	A	G	A	G	G
b	0.00545	11	9	A	T	A	A	A	G	A	■	G
c	0.00365	8	7	A	T	A	A	A	G	A	G	G
d	0.00315	7	6	A	T	A	A	A	G	A	G	G
k	0.0018	4	4	A	T	A	A	A	G	A	G	G
w	0.0018	3	3	A	T	A	A	A	G	A	G	G
h	0.0018	4	4	A	T	A	A	A	G	A	G	G
f	0.0018	4	4	A	T	A	A	A	G	A	G	G
e	0.00135	3	3	A	T	A	A	A	G	A	G	G
r	0.00135	3	3	A	T	A	A	A	G	A	G	G
x	0.00135	3	2	A	T	A	A	A	G	A	G	G
o	0.00135	3	2	A	T	A	A	A	G	A	G	G
y	0.0009	2	2	A	T	A	A	A	G	A	G	G
g	0.0009	2	2	A	T	<b>G</b>	A	A	G	A	G	G
t	0.0009	2	2	A	T	A	A	A	G	A	G	G
i	0.0009	2	2	A	<b>G</b>	A	A	A	G	A	G	G
n	0.0009	2	2	<b>G</b>	T	A	A	A	G	A	G	G
l	0.0009	2	2	A	T	A	A	A	G	A	G	G
j	0.00045	1	1	A	T	A	A	A	G	A	G	G
u	0.00045	1	1	A	T	A	A	A	G	A	G	G
aa	0.00045	1	1	A	T	A	A	<b>C</b>	G	A	G	G
v	0.00045	1	1	A	T	A	A	A	G	A	G	G
s	0.00045	1	1	A	T	A	A	A	G	A	G	G
ab	0.00045	1	1	A	T	A	A	A	G	A	■	G
q	0.00045	1	1	A	T	A	A	A	G	A	G	G
z	0.00045	1	1	A	T	A	A	A	G	<b>T</b>	G	G
af	0.00045	1	1	A	T	A	A	A	G	A	G	G
ad	0.00045	1	1	A	T	A	<b>G</b>	A	G	A	G	G

Table 13.98 - IGKV4-1

ac 1  
 m 1  
 p 1  
 ae 1

A	T	A	A	A	A	A	A	A	A	A	A	A	G
A	T	A	A	A	A	A	A	A	A	A	A	A	G
A	T	A	A	A	A	A	A	A	A	A	A	A	G
A	T	A	A	A	A	A	A	A	A	A	A	A	G

Table 13.98 - IGKV4-1

h.type	cum.freq	# indivs	pops	2:89185664 rs192141151	ENST00000390243
ref	0.9516	-	-	Thr->Ser	
a	0.01185	26	11	A	1,2,3,4,5,7,9,10,11,13,14
b	0.00545	11	9	A	2,3,4,5,6,10,12,13,14
c	0.00365	8	7	A	2,3,5,6,7,11,13
d	0.00315	7	6	A	2,5,6,7,9,11
k	0.0018	4	4	A	2,5,7,11
w	0.0018	3	3	A	2,5,14
h	0.0018	4	4	A	6,7,11,14
f	0.0018	4	4	A	4,5,6,14
e	0.00135	3	3	A	5,9,13
r	0.00135	3	3	T	5,7,8
x	0.00135	3	2	A	1,4
o	0.00135	3	2	A	9,11
y	0.0009	2	2	A	11,13
g	0.0009	2	2	A	5,6
t	0.0009	2	2	A	13,14
i	0.0009	2	2	A	9,14
n	0.0009	2	2	A	1,2
l	0.0009	2	2	A	2,11
j	0.00045	1	1	A	13
u	0.00045	1	1	A	13
aa	0.00045	1	1	A	7
v	0.00045	1	1	A	10
s	0.00045	1	1	A	11
ab	0.00045	1	1	A	13
q	0.00045	1	1	A	5
z	0.00045	1	1	A	11
af	0.00045	1	1	A	3
ad	0.00045	1	1	A	2

Table 13.98 - IGKV4-1

ac	0.00045	1	A	1
m	0.00045	1	A	12
p	0.00045	1	A	14
ae	0.00045	1	A	9



Table 13.99 - IGKV5-2

h.type	cum.freq	# indivs	pops	2:89197020	2:89197105	2:89197190	
ref	0.6714	-	-	rs183891175	rs189117859	rs55661410	ENST00000390244
a	0.32315	501	14	Thr->Arg C	Asp->Glu T	Pro->Ser C	
b	0.005	11	6	C	T	T	1,2,3,4,5,6,7,8,9,10,11,12,13,14
c	0.00045	1	1	G	G	T	2,5,6,7,11,13
						C	11

Table 13.100 - IGKV6-21

h.type	cum.frec	# indivs	pops	2:89459266 rs181879031	2:89459267 rs395483	ENST00000390256
ref	0.9638	-	-	Thr->Met	Thr->Ala	
a	0.03575	78	14	G	T	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.00045	1	1	G	C	14

Table 13.101 - IGKV6D-21

h.type	cum.freq	# indivs	pops	2:90060955	2:90060957	2:90060963	
ref	0.9735	-	-	rs190780280	rs141599034	rs150509294	
a	0.0192	42	13	Asp->Gly	Ala->Thr	Ala->Thr	ENST00000436451
b	0.00685	14	4	Asp->Gly	Ala->Thr	Ala->Thr	ENST00000557853
c	0.00045	1	1	Asp->Gly	Ala->Thr	Ala->Thr	ENST00000558079
				A	G	G	
				A	G		1,2,3,4,5,6,7,8,9,10,11,12,14
				A		G	1,5,10,14
				<b>G</b>	G	G	3

Table 13.102 - IGKV6D-41

h.type	cum.freq	# indivs	pops	2:90108534 rs191316044	2:90108797	2:90109001 rs192662836	ENST00000390271
ref	0.99865	-	-	Val->Ala T	->Asp ATGT	Thr->Ile C	
a	0.00045	1	1	C	ATGT	C	3
c	0.00045	1	1	T	ATGT	T	6
b	0.00045	1	1	T	ATGT	C	9

Table 13.103 - IGLJ1

h.type	cum_freq	# indivs	pops	22:23235930
ref	0.9982	-	-	rs182236201
a	0.0018	4	4	Arg->Gln
				ENST00000390320
				Arg->Gln
				ENST00000526893
				Arg->Gln
				ENST00000532223
				G
				3.4

Table 13.104 - IGLJ2

h.type	cum.freq	# indivs	pops	22:23241713	22:23241772	22:23241773	22:23241777	22:23241802	22:23241802
ref	0.90495	-	-	Ala->Gly	Phe->Leu	Phe->Tyr	Leu->Phe	->Val	Val->Ile
a	0.0741	160	13	<b>G</b>	T	T	G	GTAT	G
b	0.0178	38	12	C	<b>C</b>	T	G	GTAT	G
e	0.0009	2	2	<b>G</b>	<b>C</b>	T	G	GTAT	G
c	0.0009	2	2	C	T	T	<b>C</b>	GTAT	G
d	0.00045	1	1	C	T	T	G	GTAT	■
g	0.00045	1	1	C	T	■	G	GTAT	G
f	0.00045	1	1	C	T	T	G	<b>G</b>	G

ENST00000390322

1,2,3,4,5,6,7,9,10,11,12,13,14

1,2,3,4,5,7,9,10,11,12,13,14

4,9

1,10

2

13

2



Table 13.106 - IGLJ5

h.type	cum.freq	# indivs	pops	22:23256422 rs189343862	22:23256465 rs192396255	ENST00000390327
ref	0.99545	-	-	Leu->Phe G	Glu->Lys G	
a	0.00365	8	5	T	G	1,6,7,12,13
b	0.0009	2	1	G		14



Table 13.107 - IGLJ6

h.type	cum.freq	# indivs	pops	22:23260329 rs76121504	22:23260345 rs114500104	22:23260367 rs184683407	ENST00000390328
ref	0.9895	-	-	Ser->Leu C	Phe->Leu C	Val->Ile G	
a	0.00595	13	3	T	C	G	1,12,14
b	0.0032	7	3	C	G	G	1,10,14
c	0.00135	3	2	C	C	■	1,10

Table 13.108 - IGLJ7

h.type	cum.frec	# indivs	pops	22:23263601	22:23263602	rs150162743	rs373405	Ala->Thr	Ala->Val	ENST00000390330
ref	0.0664	-	-	G	C	G	C			
a	0.93225	1084	14	G	T	G	T			1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.00135	3	3	■	T	■	T			1,10,14

Table 13.109 - IGLV10-54

h.type	cum.fr	# indivs	pops	22:22569242	22:22569357	22:22569442	22:22569448	22:22569451	22:22569468	22:22569554	22:22569559
ref				Val->LeuVal->Leu	Ser->ThrSer->Thr	Ser->Thr	Ile->Asn	Val->Ala	Ala->Thr	Phe->Leu	Ala->Val
a	0.3126	-	-	G	T	G	T	T	G	C	C
b	0.50725	812	14	G	T	G		T	G		C
c	0.09425	196	14	G	T	G		T	G	C	C
d	0.05525	109	9	G	T	G	T	T	G	C	C
e	0.01265	27	5	G	T	G		C	G	C	C
f	0.0036	8	3	G	T	G		T	G		C
g	0.00315	7	2	G	T	G		T	G		C
h	0.0027	6	2	G	T	G		T	G		C
i	0.0027	6	2	G	T	G		T	G		C
k	0.0018	4	2	G	T	G		T	G		C
l	0.00135	3	1	G		G		T	G		T
m	0.0009	2	2	G		G		T	G		C
n	0.0009	2	2	G		G		T	G		C
o	0.00045	1	1	C		C		T	G		C
p	0.00045	1	1	G	T	G	T	T	G	C	C

Table 13.109 - IGLV10-54

h.type	cum.fr	# indivs	pops	22:22569562	22:22569609	22:22569618	22:22569640	22:22569649
ref	0.3126	-	-	Ser->Cys	Glu->Lys	Ala->Pro	Leu->Trp	Ser->Asn
a	0.50725	812	14	C	G	G	T	G
b	0.09425	196	14	C	G	G	G	G
c	0.05525	109	9	C	G	G	G	G
d	0.01265	27	5	C	G	G	G	G
e	0.0036	8	3	C	G	G	G	G
f	0.00315	7	2	C	G	G	G	G
h	0.0027	6	2	C	G	G	G	G
i	0.0027	6	2	C	G	G	G	G
k	0.0018	4	2	C	G	G	G	G
g	0.00135	3	1	C	G	G	G	G
j	0.0009	2	2	C	G	G	T	G
n	0.0009	2	2	C	G	G	G	G
m	0.00045	1	1	C	G	G	G	G
l	0.00045	1	1	G	G	G	T	G

ENST00000390287  
 1,2,3,4,5,6,7,8,9,10,11,12,13,14  
 1,2,3,4,5,6,7,8,9,10,11,12,13,14  
 1,3,5,6,9,10,11,12,14  
 1,5,10,12,14  
 10,11,14  
 10,14  
 10,14  
 1,14  
 10,14  
 10  
 11,13  
 2,14  
 13  
 11

Table 13.110 - IGLV11-55

h.type	cum.freq	# indivs	pops	22:22556076	22:22556114	22:22556368	22:22556443	22:22556479	22:22556517	22:22556577
ref	0.4994	-	-	rs117609231	rs61749483	rs9306341	rs188903924	rs61744363	rs151095711	rs113498536
a	0.4463	728	14	Leu->Gln	Gly->ArgGly->Arg	Pro->Leu	Arg->Gln	Ala->Val	Ala->Pro	Val->Phe
b	0.0325	71	12	T	G	C	G	C	G	G
c	0.0096	21	6	T	C	T	G	C	G	G
d	0.00545	12	6	T	C	C	G	T	G	G
g	0.00225	5	3	T	G	T	G	T	G	T
e	0.0018	4	3	■	G	T	G	C	G	G
h	0.00135	3	2	T	G	C	G	C	C	G
f	0.00135	3	2	T	G	T	■	C	G	G

ENST00000390286

1,2,3,4,5,6,7,8,9,10,11,12,13,14  
 1,2,3,5,6,7,8,9,10,11,13,14  
 3,4,6,9,12,13  
 2,4,9,10,13,14  
 1,10,14  
 2,5,13  
 1,10  
 4,9

Table 13.111 - IGLV1-36

h.type	cum.freq	# indivs	pops	22:22786604	22:22786673	22:22786674	22:22786692	22:22786709	22:22786750	22:22786792	22:22786797
ref	0.8795	-	-	Ala->Thr	Pro->Thr	Pro->Leu	Arg->Gln	Ser->Pro		Ser->Arg	Asn->Ser
a	0.0942	185	14	G	C	C	G	T	TGAG	C	A
b	0.00685	15	5	G	C	C	G	T	TGAG	G	A
d	0.0068	15	4	G	C	T	G	T	TGAG	C	A
c	0.00455	10	2	G	C	C	G	T	TGAG	C	G
f	0.0036	8	2	G	■	C	G	C	TGAG	C	A
g	0.0027	6	3	G	C	C	G	T	T	C	A
e	0.0018	4	3	G	C	C	■	T	TGAG	C	A
				rs61735471	rs190488947	rs118006119	rs182633696	rs139155569	rs59686180	rs57337788	rs61735468

ENST00000390301

1,2,3,4,5,6,7,8,9,10,11,12,13,14  
2,6,7,12,13  
1,10,12,14  
10,14  
10,14  
1,4,10  
2,6,7

Table 13.112 - IGLV1-40

h.type	cum.freq	# indivs	pops	22:22764185 rs146878832	22:22764350 rs186944811	22:22764375 rs183685135	22:22764392 rs143677402	22:22764400 rs187953594	22:22764410 rs141905358	22:22764421 rs146585623	22:22764436 rs185509404	22:22764455 rs190358290
ref	0.94895	-	-	His->Tyr	Gly->Glu	Ile->Met	Ser->Thr	Ile->Phe	Gly->Asp	His->Asp	Leu->Phe	Lys->Arg
a	0.0302	66	10	C	G	C	G	A	G	C	C	A
b	0.0046	10	5	C	G	C	G	T	G	C	C	A
d	0.0036	7	7	C	G	C	G	A	G	C	C	A
c	0.00275	6	2	C	G	C	G	A	G	C	C	A
f	0.0018	4	3	T	G	C	G	A	G	C	C	A
g	0.00135	3	3	C	■	C	G	A	G	C	C	A
e	0.0009	2	2	C	G	C	G	A	G	C	T	A
h	0.0009	2	2	C	G	C	G	A	■	C	C	A
j	0.00045	1	1	C	G	C	C	A	G	C	C	G
k	0.00045	1	1	C	G	C	G	A	G	C	T	A
s	0.00045	1	1	C	G	C	G	A	G	C	C	A
q	0.00045	1	1	C	G	G	G	A	G	C	C	G
r	0.00045	1	1	C	G	C	G	A	G	G	C	A
i	0.00045	1	1	C	G	C	C	A	G	C	C	A
n	0.00045	1	1	C	G	C	G	A	G	C	T	A
m	0.00045	1	1	C	G	C	G	A	G	C	C	A
l	0.00045	1	1	C	G	C	C	A	G	C	C	A
p	0.00045	1	1	C	G	C	C	A	■	C	C	A
o	0.00045	1	1	C	G	C	G	A	G	C	C	G



Table 13.112 - IGLV1-40

h.type	cum.freq	# indivs	pops	22:22764472 rs181685335	22:22764473 rs185969886	22:22764476 rs111666735	22:22764487 rs138088831	22:22764512 rs181089093	22:22764541 rs190749695	22:22764548 rs143634461	ENST00000390299
ref	0.94895	-	-	A	A	G	T	G	G	Thr->Ser	
a	0.0302	66	10	A	A	G	T	G	G	C	1,2,3,4,5,6,7,10,13,14
b	0.0046	10	5	A	A	G	T	G	G	C	2,6,7,12,13
d	0.0036	7	7	A	A	G	T	G	G	C	1,2,5,10,11,13,14
c	0.00275	6	2	A	A	G	G	G	G	C	10,14
f	0.0018	4	3	A	A	G	T	G	G	C	1,10,14
g	0.00135	3	3	A	A	G	T	G	G	C	2,7,12
e	0.0009	2	2	A	A	G	T	G	G	C	10,13
h	0.0009	2	2	A	A	G	T	G	G	C	2,3
j	0.00045	1	1	A	A	G	T	G	G	C	1
k	0.00045	1	1	A	G	G	T	G	G	C	1
s	0.00045	1	1	T	A	G	T	G	G	C	13
q	0.00045	1	1	A	A	G	T	G	G	C	10
r	0.00045	1	1	A	A	G	T	G	G	C	10
i	0.00045	1	1	A	A	G	T	G	G	C	10
n	0.00045	1	1	A	A	G	T	G	G	C	2
m	0.00045	1	1	A	A	G	T	C	G	C	2
l	0.00045	1	1	A	A	G	T	G	G	C	10
p	0.00045	1	1	A	A	G	T	G	G	C	1
o	0.00045	1	1	A	A	G	T	G	G	C	5

Table 13.113 - IGLV1-44

h.type	cum.fre	# indivs	pops	22:22735255 rs183817639	22:22735448 rs180819320	22:22735517 rs61731361	22:22735551 rs190708617	22:22735573 rs111733579	22:22735661 rs6002086	22:22735678 rs181994204	22:22735710 rs149125249	ENST00000390297
ref	0.9531	-	-	Ser->Gly A	Ala->Thr G	Thr->Ala A	Ala->Gly C	Ser->Arg T	Ser->Ala T	Asp->Glu T	Asn->Ser A	
a	0.0292	60	12	A	G	A	C	G	T	T	A	1,2,3,4,5,6,7,9,10,11,13,14
b	0.00775	16	2	A	G	A	C	G	G	T	A	10,14
c	0.00365	8	4	A	G	A	G	T	T	T	A	6,7,11,13
d	0.0018	4	3	G	G	A	C	T	T	T	A	4,11,13
e	0.0018	4	3	A	G	A	C	T	T	■	A	7,13,14
f	0.00135	3	2	A	G	A	C	T	T	T	G	1,14
g	0.00045	1	1	A	G	G	C	T	T	T	A	9
h	0.00045	1	1	A	■	A	C	T	T	T	A	9
i	0.00045	1	1	A	G	A	C	G	T	T	G	3

Table 13.114 - IGLV1-47

h.type	cum.fr	# indivs	pops	22:22712158 rs12160269 Leu->Ile	22:22712333 rs185378771 Pro->Ala	22:22712403 rs181574862 Gly->Glu	22:22712410 rs185842267 Asn->Lys	22:22712444 rs8142074 Ala->Thr	22:22712467 rs5757973 Ser->Arg	22:22712552 rs61231611 Arg->Trp	22:22712562 rs184225865 Asp->Ala	ENST00000390294
ref	0.11455	-	-	C	C	G	T	G	T	C	A	
a	0.8413	1059	14	C	C	G	T	G	G	C	A	1,2,3,4,5,6,7,8,9,10,11, 12,13,14
b	0.0182	40	4	■	C	G	T	G	G	C	A	1,10,12,14
c	0.01695	33	4	C	C	G	T	■	G	C	A	1,5,10,14
d	0.00405	9	3	C	C	G	T	G	G	T	A	1,10,14
g	0.0018	4	2	C	C	G	G	G	G	C	A	10,14
f	0.00135	3	2	C	C	■	T	G	G	C	A	8,14
e	0.0009	2	2	C	C	G	T	■	T	C	A	1,14
h	0.00045	1	1	C	G	G	T	G	G	C	A	13
i	0.00045	1	1	C	C	G	T	G	G	C	C	10

Table 13.115 - IGLV1-50

h.type	cum.freq	# indivs	pops
ref	0.7922	-	-
a	0.19325	373	14
b	0.0105	23	5
c	0.00225	5	3
d	0.0018	4	2

22:22681721	22:22681967	22:22682039	22:22682063
rs12484322	rs140594791	rs150490781	rs73880621
Ser->Pro	Ala->Val	Asn->Ser	Gln->Arg
T	C	A	A
C	C	A	A
T	C	A	G
T	C	G	A
T	T	A	A

EINST00000390291
1,2,3,4,5,6,7,8,9,10,11,12,13,14
1,10,11,12,14
10,12,13
3,9

Table 13.116 - IGLV1-51

h.type	cum.freq	# indivs	pops	Asp->Glu	Gly->Asp	Asp->Asn	Gly->Val	Asp->Gly	Ser->Arg	Ser->Arg	ENST00000390290
ref	0.8737	-	-	C	G	G	G	A	A	A	
a	0.1177	221	11		G	G	G	A	A	A	
b	0.00275	6	3	C	G		G	A	A	A	1,2,5,6,7,8,10,11,12,13,14
d	0.0018	4	2		G	G	G	A	A	C	2,5,12
c	0.0018	4	3	C	G	G	G	A	A	C	3,4
g	0.0009	2	2	C	G	G	G	A	A	A	3,4,9
e	0.00045	1	1	C	G	G	G	A	A	A	6,10
h	0.00045	1	1	C	G	G	T	A	C	A	6
f	0.00045	1	1	C	G	G	G	G	A	A	13
											6

Table 13.117 - IGLV2-11

h.type	cum.freq	# indivs	pops	22:23135047	22:23135342	22:23135358	22:23135360	22:23135376	22:23135379	22:23135382	22:23135466
				rs185062160	rs187815581	rs193000225	rs185237664	rs189822640	rs181371196	rs184627661	rs142176771
ref	0.9745	-	-	Ser->Asn	Met->Ile	Lys->Gln	Lys->Asn	Pro->Ser	Asp->Asn	Arg->Cys	Cys->Ser
a	0.00775	17	3	G	G	A	G	C	G	C	T
b	0.00455	10	7	G	G	A	G	C	G	C	■
d	0.0037	8	5	G	G	A	G	T	G	C	T
c	0.0032	7	5	G	G	A	T	T	■	C	T
g	0.00225	5	3	G	G	A	G	C	■	C	T
j	0.0009	2	2	G	G	A	T	C	G	C	■
e	0.0009	2	1	■	G	A	G	C	G	C	T
f	0.0009	2	2	G	G	C	G	C	G	C	T
i	0.0009	2	2	G	G	A	G	C	G	T	T
h	0.00045	1	1	G	C	A	G	C	G	C	T

EINST00000390314  
 3,4,9  
 2,6,7,10,11,12,13  
 1,5,10,11,14  
 2,3,9,12,14  
 2,10,14  
 3,9  
 6  
 6,13  
 2,11  
 14

Table 13.118 - IGLV2-14

h.type	cum.freq	# indivs	pops	Leu->Phe	Thr->Asn	Ser->Thr	22:23101480 rs188164895	22:23101494 rs191388769	22:23101497 rs182533580	22:23101559 rs4134484	22:23101568 rs28561583	22:23101600 rs184972365	22:23101615 rs188991923
ref	0.26855	-	-	C	C	G	22:23101480 rs188164895	22:23101494 rs191388769	22:23101497 rs182533580	22:23101559 rs4134484	22:23101568 rs28561583	22:23101600 rs184972365	22:23101615 rs188991923
a	0.48365	750	14	T	C	G	rs188164895	rs191388769	rs182533580	G	T	G	Asn->Ser
c	0.08855	191	14	C	C	G	rs188164895	rs191388769	rs182533580	T	T	G	A
b	0.0804	159	10	C	C	G	rs188164895	rs191388769	rs182533580	T	T	G	A
d	0.02365	51	9	C	C	G	rs188164895	rs191388769	rs182533580	T	T	G	A
g	0.0104	22	3	C	C	G	rs188164895	rs191388769	rs182533580	T	T	G	A
e	0.00995	19	3	C	C	G	rs188164895	rs191388769	rs182533580	T	T	G	A
h	0.0068	15	4	C	C	G	rs188164895	rs191388769	rs182533580	T	T	G	A
f	0.0055	12	6	C	C	G	rs188164895	rs191388769	rs182533580	T	T	G	A
i	0.0036	8	4	T	C	G	rs188164895	rs191388769	rs182533580	T	T	G	A
j	0.0027	6	5	T	C	G	rs188164895	rs191388769	rs182533580	T	T	G	A
m	0.0027	6	5	T	C	G	rs188164895	rs191388769	rs182533580	T	T	G	A
n	0.00225	5	3	C	C	G	rs188164895	rs191388769	rs182533580	T	T	G	A
i	0.00185	4	2	T	C	G	rs188164895	rs191388769	rs182533580	T	T	G	A
k	0.0018	4	4	T	C	G	rs188164895	rs191388769	rs182533580	T	T	G	A
r	0.0018	4	1	T	C	G	rs188164895	rs191388769	rs182533580	T	T	G	A
q	0.0009	2	1	C	C	G	rs188164895	rs191388769	rs182533580	T	T	G	A
p	0.0009	2	2	C	C	G	rs188164895	rs191388769	rs182533580	T	T	G	A
o	0.0009	2	1	C	C	G	rs188164895	rs191388769	rs182533580	T	T	G	A
y	0.00045	1	1	T	C	G	rs188164895	rs191388769	rs182533580	T	T	G	A
u	0.00045	1	1	C	C	G	rs188164895	rs191388769	rs182533580	T	T	G	A
t	0.00045	1	1	C	C	G	rs188164895	rs191388769	rs182533580	T	T	G	A
v	0.00045	1	1	C	C	G	rs188164895	rs191388769	rs182533580	T	T	G	A
s	0.00045	1	1	C	C	G	rs188164895	rs191388769	rs182533580	T	T	G	A
w	0.00045	1	1	C	C	G	rs188164895	rs191388769	rs182533580	T	T	G	A
x	0.00045	1	1	T	C	G	rs188164895	rs191388769	rs182533580	T	T	G	A

Table 13.118 - IGLV2-14

h.type	cum.freq	# indivs	pops	22:23101680 rs183884653	22:23101690 rs188224577	22:23101698 rs16989343	ENST00000390312
ref	0.26855	-	-	Tyr->Asn	Ser->Thr	Leu->Phe	
a	0.48365	750	14	T	G	C	1,2,3,4,5,6,7,8,9,10,11,12,13,14
c	0.08855	191	14	T	G	C	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.0804	159	10	T	G	T	1,3,4,5,6,9,10,11,12,14
d	0.02365	51	9	T	G	T	1,2,3,4,5,6,9,10,14
g	0.0104	22	3	T	G	C	3,4,9
e	0.00995	19	3	T	G	C	5,10,11
h	0.0068	15	4	T	G	T	3,4,9,10
f	0.0055	12	6	T	G	T	1,9,10,12,13,14
l	0.0036	8	4	T	G	C	2,4,11,12
j	0.0027	6	5	T	G	C	1,9,10,12,13
m	0.0027	6	5	T	G	T	3,4,6,9,10
n	0.00225	5	3	T	G	C	2,5,12
i	0.00185	4	2	T	G	C	2,9
k	0.0018	4	4	T	G	C	6,9,11,12
r	0.0018	4	1	T	G	C	14
q	0.0009	2	1	T	G	C	10
p	0.0009	2	2	T	G	C	2,13
o	0.0009	2	1	T	G	C	9
y	0.00045	1	1	T	G	C	10
u	0.00045	1	1	T	G	C	6
t	0.00045	1	1	T	G	C	6
v	0.00045	1	1	T	C	C	1
s	0.00045	1	1	T	G	C	1
w	0.00045	1	1	T	G	T	9
x	0.00045	1	1	T	G	C	2



Table 13.119 - IGLV2-18

h.type	cum.frec	# indivs	pops	22:23077436	22:23077555	ENST00000390310
ref	0.2279	-	-	rs149656926	rs4822296	
a	0.763	998	14	Glu->Asp	Leu->Ser	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.0091	19	6	G	T	1,3,4,7,10,14
				G	C	
				T	C	

Table 13.120 - IGLV2-23

h.type	cum.freq	# indivs	pops	Thr->Asn	Val->Met	Val->Glu	Ser->Cys	Gly->Arg	Ser->Arg	Gln->Glu	Glu->Asp	Gly->Val
ref	0.58265	-	-	C	G	T	A	G	A	C	G	G
a	0.34655	593	14	C	G	T	A	G	A	C	G	T
b	0.0405	82	12		G	T	A	G	A	C	T	T
c	0.01545	31	5	C	G	T	A	G	A	C	G	T
d	0.00315	7	5	C	G		A	G	A	C	G	G
g	0.00315	7	5	C	G	T	A	G	A	C	T	T
f	0.00225	5	2		G	T	A	G	A	C	T	G
e	0.00135	3	3	C	G	T	A	G	A	C	G	G
k	0.0009	2	2		G	T	A	G	A	C	G	G
o	0.0009	2	2	C	G	T	A	G	A	C	T	G
j	0.00045	1	1	C	G	T	T	G	A	C	G	G
h	0.00045	1	1	C		T	A	G	A	C	G	G
i	0.00045	1	1	C	G	T	A		A	C	G	G
n	0.00045	1	1	C	G	T	A	G	A	G	G	G
m	0.00045	1	1	C		T	A	G	A	C	G	T
l	0.00045	1	1	C	G	T	A	G	C	C	G	G
p	0.00045	1	1	C	G	T	A	G	A	C	T	G

Table 13.120 - IGLV2-23

h.type	cum.freq	# indivs	pops	22:23040887	rs5759376	Tyr->Cys	ENST00000390306
ref	0.58265	-	-	A			
a	0.34655	593	14	A			1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.0405	82	12	A			1,2,3,4,5,7,9,10,11,12,13,14
c	0.01545	31	5	<b>G</b>			3,4,5,9,11
d	0.00315	7	5	A			2,5,6,7,12
g	0.00315	7	5	A			1,2,9,10,14
f	0.00225	5	2	A			10,13
e	0.00135	3	3	<b>G</b>			3,9,11
k	0.0009	2	2	A			8,10
o	0.0009	2	2	A			3,10
j	0.00045	1	1	A			9
h	0.00045	1	1	A			3
i	0.00045	1	1	A			9
n	0.00045	1	1	A			14
m	0.00045	1	1	A			13
l	0.00045	1	1	A			5
p	0.00045	1	1	<b>G</b>			11

Table 13.121 - IGLV2-33

h.type	cum.freq	# indivs	pops	22:22930684 rs11914178	22:22930896 rs143638085	22:22931059 rs185353299	ENST00000390302
ref	0.98955	-	-	Met->Val A	Gly->Arg G	Gly->Val G	
a	0.00775	16	4	<b>G</b>	G	G	10,11,12,14
b	0.0018	4	3	A	G	<b>T</b>	2,5,13
c	0.0009	2	1	A	<b>A</b>	G	14

Table 13.122 - IGLV2-8

h.type	cum.freq	# indivs	pops	22:23165340	22:23165526	22:23165610	22:23165680	22:23165701	22:23165757
ref	0.8422	-	-	rs5996397	rs191286893	rs185026152	rs149347098	rs144694239	rs11558656
a	0.1451	267	11	Thr->Ser	Pro->Ser	Pro->Ser	Ser->Phe	Thr->Met	Ser->Cys
b	0.0046	10	7	G	C	C	C	C	A
c	0.00315	7	3	C	T	C	T	C	A
d	0.0027	6	3	G	C	C	C	C	T
e	0.0018	4	2	C	C	C	C	T	A
f	0.00045	1	1	C	C	T	C	C	A
									ENST00000390317
									1,2,5,6,7,8,10,11,12,13,14
									2,5,6,7,11,12,13
									3,4,9
									1,10,14
									4,7
									2

Table 13.123 - IGLV3-10

h.type	cum.freq	# indivs	pops	22:23154549 rs182282578	22:23154570 rs192547759	22:23154580 rs183865794	22:23154582 rs1065464	22:23154583 rs188439978	22:23154592 rs184893196	22:23154646 rs141938356	22:23154659 rs188537134	22:23154667 rs191450268
				Thr->Ala	Pro->Thr	Tyr->Cys	Ala->Pro	Ala->Val	Tyr->Cys	Arg->Gln	Ile->Met	Arg->Lys
ref	0.92545	-	-	A	C	A	G	C	A	G	C	G
a	0.049	97	10	A	C	A	C	C	A	G	C	G
b	0.0206	41	4	A	C	A	G	C	A	G	C	G
c	0.00315	6	2	A	C	A	G	C	A	G	C	G
d	0.00045	1	1	A	C	G	G	C	A	G	C	G
g	0.00045	1	1	A	C	A	G	C	A	G	G	G
e	0.00045	1	1	G	C	A	G	C	G	G	C	G
f	0.00045	1	1	A	C	A	G	T	A	G	C	G

Table 13.123 - IGLV3-10

h.type	cum.freq	# indivs	pops	22:23154736	rs80223369	Ala->Asp	ENST00000390315
ref	0.92545	-	-			C	
a	0.049	97	10			C	1,2,5,6,7,8,11,12,13,14
b	0.0206	41	4				1,10,12,14
c	0.00315	6	2			C	1,10
d	0.00045	1	1			C	2
g	0.00045	1	1			C	11
e	0.00045	1	1			C	2
f	0.00045	1	1			C	11





Table 13.125 - IGLV3-16

h.type	cum.fre	# indivs	pops	22:23090155 rs191152677	22:23090222 rs186816034	22:23090242 rs11090180	22:23090243 rs11090181	22:23090248 rs191325224	22:23090260 rs149984128	22:23090327 rs145232556	22:23090381 rs149168687	
ref	0.9395	-	-	Ser->Thr	Tyr->Phe	Phe->Val	Phe->Ser	Val->Met	Tyr->Asn	Val->Asp	Leu->Gln	ENST00000390311
a	0.04295	82	6	T	A	T	T	G	T	T	T	1,5,10,11,12,14
b	0.0081	18	9	T	A	G	C	G	T	T	T	1,2,5,7,9,10,12,13,14
c	0.0045	10	5	T	A	T	T	G	T	T	T	1,9,10,11,14
h	0.0018	4	3	T	A	G	C	G	T	T	T	1,10,14
d	0.00135	3	3	T	A	T	T	G	T	T	T	2,5,9
g	0.00045	1	1	T	A	T	T	G	T	T	T	5
e	0.00045	1	1	T	T	T	T	G	T	T	T	10
f	0.00045	1	1	T	A	T	T	G	T	T	T	13
i	0.00045	1	1	T	T	G	C	G	T	T	T	10

Table 13.126 - IGLV3-19

h.type	cum.freq	# indivs	pops	22:23063178 rs12158984	22:23063348 rs117660534	22:23063367 rs144678284	22:23063406 rs193091235	22:23063411 rs144208459	22:23063421 rs191985323	22:23063436 rs117237784	22:23063448 rs76281239	22:23063480 rs188492127
ref	0.86895	-	-	Leu->Phe	Val->Ile	Gln->Pro	Arg->Thr	Thr->Ala	Gly->Glu	Ser->Asn	Ser->Asn	Val->Ile
a	0.10015	190	12	T	G	A	G	A	G	G	G	G
b	0.01785	38	7	C	G	A	G	A	G	G	G	G
c	0.00225	5	2	C	G	A	G	A	G	G	G	G
d	0.00135	3	1	C	G	A	G	A	G	G	G	G
l	0.00135	3	2	C	G	A	G	A	G	G	G	G
j	0.0009	2	2	C	G	A	G	A	G	G	G	G
k	0.0009	2	2	C	G	A	G	A	G	G	G	G
h	0.0009	2	1	C	G	A	G	A	G	G	G	G
i	0.0009	2	1	C	G	A	G	A	G	G	G	G
p	0.0009	2	2	C	G	A	G	A	G	G	G	G
g	0.00045	1	1	C	G	A	G	A	G	G	G	G
e	0.00045	1	1	C	G	A	G	A	G	G	G	G
q	0.00045	1	1	C	G	C	G	A	G	G	G	G
r	0.00045	1	1	C	G	A	G	A	G	G	G	G
f	0.00045	1	1	C	G	A	G	A	G	G	G	G
n	0.00045	1	1	C	G	A	G	A	G	G	G	G
m	0.00045	1	1	C	G	A	G	G	G	G	G	G
o	0.00045	1	1	C	G	A	C	A	G	G	G	G

Table 13.126 - IGLV3-19

h.type	cum.freq	# indivs	pops	Leu->Arg	Phe->Ile	Gly->Glu	Asn->Asp	Arg->Trp	ENST00000390309
ref	0.86895	-	-	T	T	G	A	C	
a	0.10015	190	12	T	T	G	A	C	
b	0.01785	38	7	<b>G</b>	T	G	A	<b>T</b>	1,2,3,4,5,6,9,10,11,12,13,14
c	0.00225	5	2	T	T	G	A	C	1,3,4,9,10,12,14
d	0.00135	3	1	T	T	G	A	<b>T</b>	1,14
l	0.00135	3	2	T	T	G	A	C	10
j	0.0009	2	2	T	T	G	A	C	10,14
k	0.0009	2	2	T	T	G	A	C	2,14
h	0.0009	2	1	T	T	G	A	C	4,13
i	0.0009	2	1	T	T	G	A	C	9
p	0.0009	2	2	T	T	G	A	C	10
g	0.00045	1	1	<b>G</b>	T	G	A	<b>T</b>	2,6
e	0.00045	1	1	T	T	■	A	C	9
q	0.00045	1	1	T	T	G	A	C	4
r	0.00045	1	1	<b>G</b>	T	G	A	C	14
f	0.00045	1	1	T	T	G	<b>G</b>	C	4
n	0.00045	1	1	T	■	G	A	C	4
m	0.00045	1	1	T	T	G	A	C	13
o	0.00045	1	1	T	T	G	A	C	13
				T	T	G	A	C	10

Table 13.127 - IGLV3-1

h.type	cum.fre	# indivs	pops	22:23223277 rs144850905	22:23223280 rs184648956	22:23223283 rs61736464	22:23223292 rs145382269	22:23223297 rs181792449	22:23223301 rs186574328	22:23223306 rs61736463	22:23223330 rs145228815
ref	0.9255	-	-	Gly->Ala	Ser->Cys	Val->Glu	Tyr->Ser	Leu->Val	Thr->Ile	Pro->Ala	Gly->Arg
a	0.01865	40	13	G	C	T	A	C	C	C	G
c	0.00585	13	8	G	C	T	A	C	C	C	G
b	0.005	11	8	G	C	T	A	C	C	C	G
h	0.00315	7	7	G	C	T	A	C	C	C	G
o	0.00315	7	4	G	C	T	A	C	C	C	G
e	0.0027	6	5	G	C	T	<b>C</b>	C	C	C	G
t	0.00225	5	5	G	C	T	A	C	C	C	G
v	0.00225	5	5	G	C	T	A	C	C	C	G
f	0.00225	5	4	G	C	T	A	<b>G</b>	C	C	G
i	0.0018	4	4	G	C	T	A	C	C	C	G
n	0.0018	4	4	G	C	T	A	C	C	C	G
ag	0.00135	3	3	G	C	T	A	C	C	C	G
y	0.00135	3	3	G	C	T	A	C	C	C	G
s	0.00135	3	3	G	C	T	A	C	C	<b>G</b>	G
x	0.00135	3	3	G	C	T	A	C	<b>T</b>	C	G
ad	0.00135	3	2	G	C	T	A	C	C	C	G
l	0.00135	3	3	G	C	T	A	C	C	C	G
p	0.00135	3	2	G	C	T	A	C	C	C	G
ah	0.0009	2	2	G	C	T	A	C	C	C	G
d	0.0009	2	2	G	C	T	A	C	C	C	G
j	0.0009	2	1	G	C	T	A	C	C	C	G
g	0.0009	2	2	G	C	T	A	C	C	C	G
z	0.0009	2	2	G	C	T	A	C	C	C	G
ac	0.0009	2	2	G	C	T	A	C	C	C	G
ae	0.0009	2	2	G	C	T	A	C	C	C	G
u	0.00045	1	1	G	C	T	A	<b>G</b>	C	C	G
k	0.00045	1	1	G	C	T	A	C	C	C	G
aj	0.00045	1	1	G	C	T	A	C	C	C	<b>C</b>

Table 13.127 - IGLV3-1

1	0.00045	aa	C	C	T	A	C	C	C	C	C	C	C	G
1	0.00045	aq	G	C	■	A	C	C	C	C	C	C	C	G
1	0.00045	al	G	C	T	A	C	C	C	C	C	C	C	G
1	0.00045	ab	C	C	T	A	C	C	C	C	C	C	C	G
1	0.00045	ak	G	C	T	A	C	C	C	C	C	C	C	G
1	0.00045	q	G	C	T	A	C	C	C	C	C	C	C	G
1	0.00045	ap	G	C	T	A	C	C	C	C	C	C	C	G
1	0.00045	w	G	C	T	A	C	C	C	C	C	C	C	G
1	0.00045	ao	G	C	T	A	C	C	C	C	C	C	C	G
1	0.00045	r	G	C	T	A	C	C	C	C	C	C	C	G
1	0.00045	af	G	C	T	A	C	C	C	C	C	C	C	G
1	0.00045	au	G	C	T	A	C	C	C	C	C	C	C	G
1	0.00045	am	G	C	T	A	C	C	C	C	C	C	C	G
1	0.00045	at	G	C	T	A	C	C	C	C	C	C	C	G
1	0.00045	m	G	C	T	A	C	C	C	C	C	C	C	G
1	0.00045	ar	G	C	T	A	C	C	C	C	C	C	C	G
1	0.00045	as	G	C	T	A	C	C	C	C	C	C	C	G
1	0.00045	ai	G	C	T	A	C	C	C	C	C	C	C	G
1	0.00045	an	G	C	T	A	C	C	C	C	C	C	C	G

Table 13.127 - IGLV3-1

h.type	cum.fre	# indivs	pops	22:23223348 rs189772470	22:23223355 rs183495921	22:23223363 rs187892119	22:23223364 rs191755097	22:23223369 rs188197731	22:23223375 rs192957059	22:23223376 rs61731694	22:23223382 rs138660033	22:23223385 rs61731693
ref				Thr->Pro	Ser->Cys	Lys->Glu	Lys->Ile	Gly->Arg	Lys->Glu	Lys->Thr	Ala->Val	Cys->Tyr
a	0.9255	-	-	A	C	A	A	G	A	A	C	G
c	0.01865	40	13	A	C	A	A	G	A	A	C	G
b	0.00585	13	8	A	C	A	A	G	A	A	T	G
h	0.005	11	8	A	C	A	A	G	A	A	C	■
o	0.00315	7	7	A	C	A	A	G	A	A	C	G
e	0.00315	7	4	A	C	A	A	G	A	A	C	G
t	0.0027	6	5	A	C	A	A	G	A	A	C	G
v	0.00225	5	5	A	C	A	A	G	A	A	C	G
f	0.00225	5	5	A	C	A	A	G	A	A	C	G
i	0.00225	5	4	A	C	A	A	G	A	A	C	G
n	0.0018	4	4	C	C	A	A	G	A	A	C	G
ag	0.0018	4	4	A	C	A	A	G	A	A	C	G
y	0.00135	3	3	A	C	A	A	G	A	A	C	G
s	0.00135	3	3	A	C	A	A	G	A	C	C	G
x	0.00135	3	3	A	C	A	A	G	A	A	C	G
ad	0.00135	3	2	A	C	A	A	G	A	A	C	G
l	0.00135	3	3	A	C	A	A	G	A	A	C	■
p	0.00135	3	2	A	C	A	A	G	A	A	C	G
ah	0.0009	2	2	A	C	G	A	G	A	A	C	G
d	0.0009	2	2	A	C	A	A	G	A	A	C	G
j	0.0009	2	1	A	C	A	A	G	A	A	C	G
g	0.0009	2	2	A	C	A	A	G	A	A	C	G
z	0.0009	2	2	A	C	A	A	G	A	A	C	G
ac	0.0009	2	2	A	C	A	A	G	A	A	C	G
ae	0.0009	2	2	A	C	A	A	G	A	A	C	G
u	0.00045	1	1	A	C	A	A	G	A	A	C	G
k	0.00045	1	1	A	C	A	A	G	G	A	C	G
aj	0.00045	1	1	A	C	A	A	G	A	A	C	G

Table 13.127 - IGLV3-1

1	0.00045	aa	A	C	A	A	A	G	A	A	A	C	G
1	0.00045	aq	A	C	A	A	A	G	A	A	A	C	G
1	0.00045	al	A	C	A	A	A	G	A	A	A	C	G
1	0.00045	ab	A	C	A	A	A	G	<b>G</b>	A	A	C	G
1	0.00045	ak	A	C	A	A	A	G	A	A	A	C	G
1	0.00045	q	<b>C</b>	C	A	A	A	G	A	A	A	<b>T</b>	G
1	0.00045	ap	A	C	A	A	A	G	<b>G</b>	A	A	C	G
1	0.00045	w	A	<b>G</b>	A	A	A	G	A	A	A	C	G
1	0.00045	ao	A	C	A	A	A	G	A	A	A	C	G
1	0.00045	r	A	C	A	A	<b>T</b>	G	A	A	A	C	G
1	0.00045	af	A	C	A	A	A	G	A	A	A	C	G
1	0.00045	au	A	C	A	A	A	G	A	A	A	<b>T</b>	G
1	0.00045	am	A	C	A	A	A	G	A	A	A	C	G
1	0.00045	at	A	C	A	A	<b>T</b>	G	A	A	A	C	G
1	0.00045	m	A	C	A	A	A	G	<b>G</b>	A	A	<b>T</b>	G
1	0.00045	ar	A	C	A	A	A	G	A	A	A	C	G
1	0.00045	as	A	C	A	A	A	<b>■</b>	A	A	A	C	G
1	0.00045	ai	A	C	A	A	A	G	A	A	A	C	G
1	0.00045	an	A	C	A	A	A	G	A	A	A	C	G

Table 13.127 - IGLV3-1

h.type	cum.fre	# indivs	pops	22:23223386	22:23223387	22:23223400	22:23223402	22:23223417	22:23223432	22:23223438	22:23223439	22:23223525
ref				rs61735526	rs18677494	rs61731688	rs61731687	rs190667156	rs73164939	rs143830449	rs61731685	rs61731681
				Cys->Trp	Trp->Arg	Lys->Arg	Pro->Ala	Val->Met	Gln->Glu	Ser->Gly	Ser->Asn	Met->Leu
a	0.9255	-	-	C	T	A	C	G	C	A	G	A
c	0.01865	40	13	C	T	A	C	G	C	A		A
b	0.00585	13	8	C	T	A	C	G	C	A	G	A
h	0.005	11	8	C	T	A	C	G	C	A	G	A
o	0.00315	7	7	C	T	A	C		C	A	G	A
e	0.00315	7	4	C	T	<b>G</b>	C	G	C	A	G	A
t	0.0027	6	5	C	T	A	C	G	C	A	G	A
v	0.00225	5	5	C	T	A	<b>G</b>	G	C	A	G	A
f	0.00225	5	5	C	T	A	C	G	<b>G</b>	A	G	A
i	0.00225	5	4	C	T	A	C	G	C	A	G	A
n	0.0018	4	4	C	T	A	C	G	C	A	G	A
ag	0.0018	4	4	C	T	A	C	G	C	<b>G</b>	G	A
y	0.00135	3	3	C	T	A	C	G	C	<b>G</b>		A
s	0.00135	3	3	C	T	A	C	G	C	A	G	A
x	0.00135	3	3	C	T	A	C	G	C	A	G	A
ad	0.00135	3	3	C	T	A	C	G	C	A	G	A
l	0.00135	3	3	C	T	A	C	G	C	A		A
p	0.00135	3	2	C	T	A	C	G	C	A	G	A
ah	0.0009	2	2	C	T	A	C	G	C	A	G	A
d	0.0009	2	2	C	T	A	C	G	C	A	G	<b>C</b>
j	0.0009	2	1	C	T	A	C	G	C	A	G	A
g	0.0009	2	2	<b>G</b>	T	A	C	G	C	A		A
z	0.0009	2	2	C	T	A	C	G	C	A	G	A
ac	0.0009	2	2	<b>G</b>	T	A	C	G	C	A	G	A
ae	0.0009	2	2	C	T	A	C	G	C	A	G	A
u	0.00045	1	1	C	T	A	C	G	C	<b>G</b>	G	A
k	0.00045	1	1	C	T	<b>G</b>	C	G	C	A	G	A
aj	0.00045	1	1	C	T	A	C	G	C	A	G	A



Table 13.127 - IGLV3-1

1	0.00045	aa	C	T	A	C	G	C	A	G	A	G	A
1	0.00045	aq	C	T	A	C	G	C	A	G	A	G	A
1	0.00045	al	C	T	A	C	G	C	A	G	A	G	A
1	0.00045	ab	C	T	A	C	G	C	A	G	A	G	A
1	0.00045	ak	C	T	A	C	G	C	A	G	A	G	A
1	0.00045	q	G	T	A	C	G	C	A	G	A	G	A
1	0.00045	ap	C	T	A	C	G	C	A	G	A	G	A
1	0.00045	w	C	T	A	C	G	C	A	G	A	G	A
1	0.00045	ao	C	T	A	C	G	C	A	G	A	G	A
1	0.00045	r	C	T	A	C	G	C	A	G	A	G	A
1	0.00045	af	C	T	A	C	G	C	A	G	A	G	A
1	0.00045	au	C	T	A	C	G	C	A	G	A	G	A
1	0.00045	am	G	T	A	C	G	C	A	G	A	G	A
1	0.00045	at	C	T	A	C	G	C	A	G	A	G	A
1	0.00045	m	C	T	A	C	G	C	A	G	A	G	A
1	0.00045	ar	C	T	A	C	G	C	A	G	A	G	A
1	0.00045	as	C	T	A	C	G	C	A	G	A	G	A
1	0.00045	ai	C	T	A	C	G	C	A	G	A	G	A
1	0.00045	an	C	T	A	C	G	C	A	G	A	G	A

Table 13.127 - IGLV3-1

h.type	cum.fre	# indivs	pops	22:23223531 rs190043877	22:23223535 rs145774047	22:23223540 rs180853064	22:23223541 rs142180628	22:23223544 rs185803370	22:23223550 rs192317432	22:23223573 rs145570740	ENST00000390319
ref	0.9255	-	-	Glu->Lys	Ala->Gly	Tyr->Asp	Tyr->Ser	Tyr->Ser	Gln->Pro	His->Tyr	
a	0.01865	40	13	G	C	T	A	A	A	C	2,3,4,5,6,7,8,9,10,11,12,13,14
c	0.00585	13	8	G	C	T	A	A	A	C	2,5,6,7,9,10,13,14
b	0.005	11	8	G	C	T	A	A	A	C	1,4,5,6,10,11,13,14
h	0.00315	7	7	G	C	T	A	A	A	C	1,3,4,5,9,10,14
o	0.00315	7	4	G	C	T	A	A	A	C	3,7,10,14
e	0.0027	6	5	G	C	T	A	A	A	C	1,2,3,10,11
t	0.00225	5	5	G	C	T	A	A	A	C	2,3,5,10,14
v	0.00225	5	5	G	C	T	A	A	A	C	1,5,6,7,9
f	0.00225	5	4	G	C	T	A	A	A	C	2,4,10,13
i	0.0018	4	4	G	C	T	A	A	A	C	1,7,10,11
n	0.0018	4	4	G	C	T	A	A	A	C	6,12,13,14
ag	0.00135	3	3	G	C	T	A	A	A	C	2,5,13
y	0.00135	3	3	G	C	T	A	A	A	C	5,10,12
s	0.00135	3	3	G	C	T	A	A	A	C	2,9,10
x	0.00135	3	3	G	C	T	A	A	A	C	3,10,14
ad	0.00135	3	2	G	C	T	A	A	A	T	2,13
l	0.00135	3	3	G	C	T	A	A	A	C	2,6,13
p	0.00135	3	2	G	C	T	A	C	A	C	5,9
ah	0.0009	2	2	G	C	T	A	A	A	C	9,14
d	0.0009	2	2	G	C	T	A	A	A	C	2,13
j	0.0009	2	1	G	C	T	C	A	A	C	2
g	0.0009	2	2	G	C	T	A	A	A	C	6,12
z	0.0009	2	2	G	C	T	A	A	C	C	1,4
ac	0.0009	2	2	G	C	T	A	A	C	C	6,11
ae	0.0009	2	2	G	G	T	A	A	A	C	13,14
u	0.00045	1	1	G	C	T	A	A	A	C	14
k	0.00045	1	1	G	C	T	A	A	A	C	11
aj	0.00045	1	1	G	C	T	A	A	A	C	2

Table 13.127 - IGLV3-1

1	0.00045	aa	G	C	T	A	A	A	A	C
1	0.00045	aq	G	C	T	A	A	A	A	C
1	0.00045	al	G	C	T	C	A	A	A	C
1	0.00045	ab	G	C	T	A	A	A	A	C
1	0.00045	ak	G	C	G	A	A	A	A	C
1	0.00045	q	G	C	T	A	A	A	A	C
1	0.00045	ap	G	C	T	A	A	A	A	C
1	0.00045	w	G	C	T	A	A	A	A	C
1	0.00045	ao	G	C	T	A	A	A	A	C
1	0.00045	r	G	C	T	A	A	A	A	C
1	0.00045	af	G	C	T	A	A	A	A	C
1	0.00045	au	G	C	T	A	A	A	A	C
1	0.00045	am	G	C	T	A	A	A	A	C
1	0.00045	at	G	C	T	A	A	A	A	C
1	0.00045	m	G	C	T	A	A	A	A	C
1	0.00045	ar	G	C	T	C	A	A	A	C
1	0.00045	as	G	C	T	A	A	A	A	C
1	0.00045	ai	G	C	T	A	A	A	A	C
1	0.00045	an	G	C	T	A	A	A	A	C

Table 13.128 - IGLV3-21

h.type	cum.fre	# indivs	pops	22:23054916 rs5759408	22:23054919 rs141888672	22:23054937 rs115492210	22:23055389 rs190621299	22:23055407 rs183242778	22:23055440 rs1008910	22:23055497 rs145459282	22:23055522 rs192152129	22:23055527 rs184274943
ref	0.2714	-	-	Val->Phe	Leu->Ile	Ser->Ala	Val->Met	Thr->Ala	Gln->Lys	Tyr->Asp	Pro->Leu	Leu->Val
a	0.3195	556	14	G	C	T	G	A	C	T	C	C
b	0.178	338	14	G	C	T	G	A		T	C	C
c	0.1613	296	14	T	C	T	G	A		T	C	C
f	0.02165	45	9	G	C	T	G	A		T	C	C
d	0.0173	36	10	T	C	T	G	A		T	C	C
g	0.01045	21	5	G	C	G	G	A	C	T	C	C
e	0.00995	22	5	G	C	T	G	A		G	C	C
h	0.0045	10	7	G	C	T	G	A	C	T	C	C
i	0.0027	6	4	T	C	T	G	A	C	T	C	C
j	0.0018	4	3	G	C	T	G	A	C	T	C	C
s	0.0018	4	4	T	C	T	G	A	C	T	C	C
t	0.00135	3	2	G	C	G	G	A		T	C	C
k	0.0009	2	2	G	C	T	G	A	C	T	T	C
r	0.0009	2	1	G	C	G	G	A	C	T	C	C
i	0.0009	2	2	G	C	G	G	A		T	C	C
u	0.00045	1	1	G	C	T		A		T	C	C
v	0.00045	1	1	G		T	G	A		T	C	C
q	0.00045	1	1	G	C	T	G	A		T	C	C
n	0.00045	1	1	G	C	T	G	A		T	C	C
m	0.00045	1	1	G	C	T	G	A		T	C	G
p	0.00045	1	1	G	C	T	G	A		T	C	C
o	0.00045	1	1	G	C	T	G	A		T	C	C

Table 13.128 - IGLV3-21

h.type	cum.fre	# indivs	pops	22:23055333 rs4446155	22:23055339 rs1985791	22:2305543 rs188081944	22:23055611 rs192901531	22:23055674 rs189479110	ENST00000390308
ref	0.2714	-	-	G	G	A	A	A	
a	0.31195	556	14	T	T	A	A	A	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.178	338	14	G	G	A	A	A	1,2,3,4,5,6,7,8,9,10,11,12,13,14
c	0.1613	296	14	T	T	A	A	A	1,2,3,4,5,6,7,8,9,10,11,12,13,14
f	0.02165	45	9	G	G	A	A	A	1,2,5,7,8,10,12,13,14
d	0.0173	36	10	G	G	A	A	A	1,3,4,6,7,9,10,12,13,14
g	0.01045	21	5	G	G	A	A	A	1,10,11,12,14
e	0.00995	22	5	T	T	A	A	A	2,4,7,11,13
h	0.0045	10	7	T	T	A	A	A	2,4,6,9,10,13,14
i	0.0027	6	4	G	G	A	A	A	2,5,11,14
j	0.0018	4	3	G	G	A	A	A	1,2,10
s	0.0018	4	4	T	T	A	A	A	2,6,9,13
t	0.00135	3	2	G	G	A	A	A	1,14
k	0.0009	2	2	G	G	A	A	A	7,12
r	0.0009	2	1	G	G	A	A	A	12
i	0.0009	2	2	T	T	A	A	A	1,10
u	0.00045	1	1	G	G	A	A	A	2
v	0.00045	1	1	T	T	A	A	A	13
q	0.00045	1	1	G	G	A	T	A	4
n	0.00045	1	1	G	G	A	A	A	6
m	0.00045	1	1	T	T	G	A	A	6
p	0.00045	1	1	T	T	A	A	A	13
o	0.00045	1	1	G	G	A	A	T	13

Table 13.129 - IGLV3-22

h.type	cum.freq	# indvs	pops	22:23046829	22:23047053	22:23047069	22:23047094	22:23047104	22:23047177	22:23047198	22:23047244	22:23047290
ref				Leu->Phe	Val->Leu	Thr->Lys	Asp->Glu	Glu->Lys	Arg->Gln	Arg->Gln	Ser->Arg	Asp->Asn
a	0.6639	-	-	C	G	C	T	G	G	G	C	G
b	0.2962	529	14	C	G	■	T	■	G	G	C	■
c	0.01545	33	7	C	T	■	T	■	G	G	C	■
d	0.00825	18	5	C	G	C	T	G	G	G	C	■
e	0.0027	6	1	C	T	■	T	■	G	G	C	G
f	0.0027	6	3	C	G	C	T	■	G	G	C	G
g	0.0018	4	3	C	G	■	T	■	G	G	C	G
h	0.0018	4	2	T	G	C	T	G	G	G	C	G
i	0.00135	3	3	C	G	C	T	■	G	G	C	■
j	0.00135	3	1	C	T	■	T	G	G	■	C	■
k	0.0009	2	1	C	G	C	G	G	G	G	C	G
l	0.0009	2	2	C	G	C	G	G	G	■	C	G
m	0.0009	2	2	C	T	■	T	G	G	■	C	G
n	0.0009	2	2	C	T	■	T	G	G	■	C	G
o	0.0009	2	2	C	T	■	T	G	G	G	C	G
p	0.00045	1	1	C	G	■	T	■	G	G	G	■
q	0.00045	1	1	C	G	■	T	■	■	G	C	■

Table 13.129 - IGLV3-22

h.type	cum.freq	# indivs	pops	ENST00000390307
ref	0.6639	-	-	
a	0.2962	529	14	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.01545	33	7	1,3,4,9,10,12,14
c	0.00825	18	5	3,4,6,9,14
e	0.0027	6	1	10
d	0.0027	6	3	1,10,14
g	0.0018	4	3	3,4,10
h	0.0018	4	2	10,14
f	0.00135	3	3	2,10,12
i	0.00135	3	1	10
j	0.0009	2	1	4
n	0.0009	2	2	10,12
m	0.0009	2	2	1,14
o	0.0009	2	2	10,14
k	0.00045	1	1	11
l	0.00045	1	1	14

Table 13.130 - IGLV3-25

h.type	cum.freq	# indivs	pops	22:23029258 rs12628782	22:23029263 rs443102	22:23029478 rs190375718	22:23029526 rs3205084	22:23029537 rs182727015	22:23029586 rs188154997	22:23029588 rs5751507	22:23029595 rs182651541	22:23029597 rs188941516
				Pro->His	Phe->Leu	Pro->His	Gly->Ala	Pro->Ser	Val->Glu	Leu->Val	Ile->Lys	Tyr->His
ref	0.1086	-	-	C	T	C	G	C	T	C	T	T
a	0.6581	943	14	C	C	C	G	C	T	C	T	T
b	0.19965	367	14	C	C	C	C	C	T	C	T	T
c	0.02105	41	6	C	C	C	G	C	T	C	T	T
d	0.00405	9	5	C	C	C	G	C	T	C	T	T
e	0.00135	3	3	C	C	C	G	C	T	C	T	T
g	0.0009	2	1	C	C	C	G	C	T	C	T	T
l	0.0009	2	2	C	C	C	G	C	T	C	T	C
j	0.00045	1	1	C	C	C	G	C	T	C	T	T
k	0.00045	1	1	C	C	C	C	C	T	C	T	C
s	0.00045	1	1	C	T	C	G	T	T	G	T	T
q	0.00045	1	1	C	C	C	G	C	T	G	T	C
r	0.00045	1	1	C	C	C	C	C	T	C	T	T
h	0.00045	1	1	C	C	C	G	C	T	C	T	T
f	0.00045	1	1	C	C	C	G	C	T	C	T	T
i	0.00045	1	1	C	C	C	G	C	T	C	T	T
n	0.00045	1	1	C	C	C	C	C	T	C	T	T
m	0.00045	1	1	C	C	C	C	T	T	C	T	T
p	0.00045	1	1	C	T	C	G	C	T	C	T	T
o	0.00045	1	1	C	C	C	G	C	T	C	T	T



Table 13.130 - IGLV3-25

h.type	cum.freq	# indivs	pops	Tyr->Ser	Ser->Ile	Thr->Arg	Gly->Ala	Ala->Ser	Gln->His	Ala->Thr
ref	0.1086	-	-	A	G	C	G	G	A	G
a	0.6581	943	14	A	G	C	G	G	A	G
b	0.19965	367	14	A	G	C	G	G	A	G
c	0.02105	41	6	A	G	C	G	G	A	G
d	0.00405	9	5	A	G	C	G	G	A	G
e	0.00135	3	3	A	T	C	G	G	A	G
g	0.0009	2	1	A	G	C	G	G	A	G
i	0.0009	2	2	A	G	C	G	G	A	G
j	0.00045	1	1	A	G	G	G	G	A	G
k	0.00045	1	1	A	G	C	G	G	A	G
s	0.00045	1	1	A	G	C	G	G	A	G
q	0.00045	1	1	A	G	C	G	G	C	G
r	0.00045	1	1	A	G	C	G	G	A	G
h	0.00045	1	1	A	G	C	G	T	A	G
f	0.00045	1	1	A	T	C	G	G	A	G
i	0.00045	1	1	A	G	C	C	G	A	G
n	0.00045	1	1	A	G	C	C	G	A	G
m	0.00045	1	1	A	G	C	G	G	A	G
p	0.00045	1	1	C	G	C	G	G	A	G
o	0.00045	1	1	A	G	C	G	G	A	G

Table 13.130 - IGLV3-25

h.type	cum.freq	# indivs	pops
ref	0.1086	-	ENST00000390305
a	0.6581	943	14 1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.19965	367	14 1,2,3,4,5,6,7,8,9,10,11,12,13,14
c	0.02105	41	6 2,3,4,5,9,11
d	0.00405	9	5 1,2,5,6,10
e	0.00135	3	3 2,5,14
g	0.0009	2	1 2
l	0.0009	2	2 3,5
j	0.00045	1	1 6
k	0.00045	1	1 5
s	0.00045	1	1 10
q	0.00045	1	1 2
r	0.00045	1	1 7
h	0.00045	1	1 10
f	0.00045	1	1 4
i	0.00045	1	1 6
n	0.00045	1	1 14
m	0.00045	1	1 8
p	0.00045	1	1 10
o	0.00045	1	1 10

Table 13.131 - IGLV3-27

h.type	cum.freq	# indivs	pops	22:23011202	22:23011253	ENST00000390304
ref	0.99865	-	-	rs185761809	rs147724316	
a	0.0009	2	2	Val->Leu	Cys->Arg	
b	0.00045	1	1	G	T	
				G	C	3,9
				C	T	8



Table 13.133 - IGLV3-9

h.type	cum.fre	# indivs	pops	22:23162047 rs187387271	22:23162116 rs111894401	ENST00000390316
ref	0.9982	-	-	Gly->Glu	Ser->Asn	
a	0.0009	2	2	G	G	1,13
b	0.0009	2	2	G	G	3,13

Table 13.134 - IGLV4-3

h.type	cum.freq	# indivs	pops	22:23213939 rs191969136	22:23213987 rs183930410	22:23214006 rs188425943	22:23214144 rs141025913	22:23214190 rs149834282	ENST00000390318
ref	0.9872	-	-	Leu->Phe	Ser->Arg	Tyr->His	Leu->Phe	Thr->Lys	
a	0.01055	22	4	G	C	T	C	C	
d	0.0009	2	2	G	C	T	C	■	1,10,12,14
e	0.00045	1	1	G	C	C	C	C	7,14
c	0.00045	1	1	G	C	T	T	C	2
b	0.00045	1	1	C	C	T	C	C	14
				G	■	T	C	C	3

Table 13.135 - IGLV4-60

h.type	cum.frec	# indivs	pops	22:22516624	22:22516785	22:22516825	22:22516832	22:22516864	22:22516869	22:22516922	22:22516929	22:22516933
ref	0.0851	-	-		Leu->Val	Ser->Leu	Lys->Asn	Ser->Asn	Ile->Phe	Lys->Asn	Gly->Arg	Ser->Asn
a	0.4868	785	14	TCTC	C	C	G	G	A	G	G	G
b	0.3607	628	14	TCTC	C	C	G	G	A	G	G	G
c	0.03615	74	5	TCTC	C	C	G	G	A	G	G	G
d	0.015	32	4	TCTC	C	T	G	G	A	G	G	G
e	0.00635	14	4	TCTC	C	C	G	G	A	G	G	■
f	0.00225	5	4	T	C	C	G	G	A	G	G	G
g	0.0009	2	2	TCTC	C	C	G	G	A	G	G	G
n	0.0009	2	2	TCTC	C	C	G	G	A	G	G	G
o	0.0009	2	2	TCTC	C	C	G	G	A	G	C	G
j	0.00045	1	1	TCTC	C	C	G	G	A	G	G	G
k	0.00045	1	1	TCTC	G	C	G	G	A	G	G	G
t	0.00045	1	1	TCTC	C	C	G	G	A	C	G	G
s	0.00045	1	1	TCTC	C	C	G	G	A	G	G	G
q	0.00045	1	1	TCTC	C	C	G	G	A	G	G	G
r	0.00045	1	1	TCTC	C	C	G	G	T	G	C	G
h	0.00045	1	1	TCTC	C	C	G	■	A	G	G	G
i	0.00045	1	1	TCTC	C	C	G	■	A	G	G	G
m	0.00045	1	1	TCTC	C	T	G	■	A	G	G	G
l	0.00045	1	1	TCTC	C	C	G	G	A	G	G	G
p	0.00045	1	1	TCTC	C	C	C	G	A	G	G	G

Table 13.135 - IGLV4-60

h.type	cum.frec	# indivs	pops	22:22516948 rs182062284	22:22516984 rs187948661	22:22516990 rs115153265	22:22516998 rs2073453	22:22517026 rs738885	22:22517065 rs192463945	22:22517071 rs183795322
ref	0.0851	-	-	Lys->Thr	Ser->Asn	Gly->Val	Arg->Cys	Phe->Ser	Ser->Asn	Thr->Ile
a	0.4868	785	14	A	G	G	C	T	G	C
b	0.3607	628	14	A	G	G	C	C	G	C
c	0.03615	74	5	A	G	T	T	C	G	C
d	0.015	32	4	A	G	G	C	C	G	C
e	0.00635	14	4	A	G	G	C	C	G	C
f	0.00225	5	4	A	G	G	C	C	G	C
g	0.0009	2	2	A	G	G	C	C	G	T
n	0.0009	2	2	A	G	G	C	C	G	C
o	0.0009	2	2	A	G	G	C	C	G	C
j	0.00045	1	1	A	G	G	C	C	G	C
k	0.00045	1	1	A	G	G	C	C	G	C
t	0.00045	1	1	A	G	G	C	C	G	C
s	0.00045	1	1	C	G	G	T	C	G	C
q	0.00045	1	1	A	G	G	T	C	G	C
r	0.00045	1	1	A	G	G	T	C	G	C
h	0.00045	1	1	A	G	G	C	C	G	C
i	0.00045	1	1	A	G	G	T	C	G	C
m	0.00045	1	1	A	G	G	C	C	G	C
l	0.00045	1	1	A	G	G	C	T	G	T
p	0.00045	1	1	A	G	G	C	C	G	C



Table 13.135 - IGLV4-60

h.type	cum.frec	# indivs	pops	ENST00000390284
ref	0.0851	-	-	
a	0.4868	785	14	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.3607	628	14	1,2,3,4,5,6,7,8,9,10,11,12,13,14
c	0.03615	74	5	1,5,10,12,14
d	0.015	32	4	1,5,10,14
e	0.00635	14	4	1,10,11,14
f	0.00225	5	4	1,3,10,14
g	0.0009	2	2	5,11
n	0.0009	2	2	2,9
o	0.0009	2	2	5,10
j	0.00045	1	1	10
k	0.00045	1	1	5
t	0.00045	1	1	6
s	0.00045	1	1	13
q	0.00045	1	1	13
r	0.00045	1	1	13
h	0.00045	1	1	14
i	0.00045	1	1	10
m	0.00045	1	1	10
l	0.00045	1	1	5
p	0.00045	1	1	10

Table 13.136 - IGLV4-69

h.type	cum.freq	# indivs	pops	22:22385618 rs150132122	22:22385671 rs185429538	22:22385747 rs140354453	22:22385851 rs188851121	ENST00000390282
ref	0.98945	-	-	Ala->Asp	Ala->Pro	Gly->Glu	Thr->Pro	
a	0.0092	20	4	C	G	G	A	
d	0.00045	1	1	C	C	G	A	1,8,10,14
c	0.00045	1	1	C	G	G	C	5
b	0.00045	1	1	C	G	G	A	10
								2

Table 13.137 - IGLV5-37

h.type	cum.freq	# indivs	pops	22:22782150 rs56060925	22:22782240 rs181085584	22:22782258 rs111480336	
ref	0.86435	-	-	Gly->Ser	Gly->Cys	Arg->Cys	ENST00000390300
a	0.1266	251	13	G	G	C	
b	0.0068	13	7	G	G	T	1,2,3,4,5,6,7,8,9,10,11,12,13
c	0.00225	4	2	G	T	C	2,6,10,11,12,13,14 10,14

Table 13.138 - IGLV5-45

h.type	cum.freq	# indivs	pops	22:22730582 rs12157664	22:22730584 rs1985918	22:22730635 rs61731372	22:22730636 rs149965450	22:22730663 rs191752433	22:22730725 rs192713801	22:22730732 rs117029072	22:22730786 rs14764645	
ref	0.8017	-	-	Pro->Leu C	Ser->Ala T	Arg->Cys C	Arg->His G	Arg->Lys G	Asp->His G	Asp->Gly A	Ser->Leu C	ENST00000390296
a	0.1706	312	12	C	<b>G</b>	C	G	G	G	A	C	1,2,3,5,6,7,8,10,11,12,13,14
b	0.0164	36	14	C	T	<b>T</b>	G	G	G	A	C	1,2,3,4,5,6,7,8,9,10,11,12,13,14
c	0.00545	12	2	<b>T</b>	<b>G</b>	C	G	G	G	A	C	10,14
d	0.0036	8	3	C	T	C	G	G	G	A	<b>T</b>	3,4,9
g	0.0009	2	2	C	T	C	<b>■</b>	G	G	A	C	2,13
e	0.00045	1	1	C	T	C	G	<b>■</b>	G	<b>G</b>	C	2
h	0.00045	1	1	C	<b>G</b>	C	G	G	<b>C</b>	A	C	10
f	0.00045	1	1	C	T	<b>T</b>	G	<b>■</b>	G	A	C	9

Table 13.139 - IGLV5-48

h.type	cum.frec	# indivs	pops	22:22707473	22:22707555	22:22707576	22:22707621	22:22707698	22:22707713	22:22707773	ENST00000390293
ref	0.79525	-	-	Pro->Ala	Asn->Ser	Phe->Tyr	Ser->Asn	Asn->Tyr	Val->Phe	Ser->Gly	
a	0.0873	174	14	G	G	G	G	A	T	A	1,2,3,4,5,6,7,8,9,10,11,12,13,14
c	0.0301	66	13	G	G	T	G	A	G	A	1,2,3,4,5,6,7,8,9,10,11,13,14
b	0.02875	63	14	G	G	T	G	A	T	A	1,2,3,4,5,6,7,8,9,10,11,12,13,14
d	0.0123	27	10	G	G	G	G	A	T	A	1,2,4,5,6,9,10,12,13,14
e	0.0082	18	9	C	G	G	G	A	T	A	2,4,5,6,7,10,11,13,14
i	0.00595	13	7	C	G	G	G	A	T	A	2,3,4,7,9,11,13
f	0.0055	12	9	G	G	T	G	A	T	A	2,3,4,6,7,9,10,11,14
h	0.00545	12	8	C	A	T	G	A	T	A	1,2,3,6,10,12,13,14
g	0.0041	9	5	G	G	G	G	A	G	A	3,6,10,12,13
o	0.00315	7	6	C	A	G	G	A	T	A	1,2,7,8,11,13
k	0.00225	5	3	C	G	T	G	A	G	A	2,6,13
j	0.0018	4	3	C	A	G	G	A	G	A	4,10,12
t	0.00135	3	2	G	A	T	G	A	T	A	3,4
s	0.00135	3	3	G	A	T	G	A	G	A	2,5,13
u	0.0009	2	2	C	A	G	G	A	T	A	6,9
q	0.0009	2	2	C	A	T	G	A	G	A	5,12
r	0.0009	2	2	G	G	G	G	T	T	A	5,11
n	0.0009	2	2	C	G	T	G	A	T	A	2,10
m	0.0009	2	1	G	A	T	G	A	T	A	6
l	0.0009	2	2	C	G	T	G	A	T	A	12,13
v	0.00045	1	1	C	G	G	G	A	G	A	10
w	0.00045	1	1	G	G	T	G	A	G	A	2
x	0.00045	1	1	G	G	G	G	A	G	A	10
p	0.00045	1	1	C	A	T	G	A	G	G	3

Table 13.140 - IGLV5-52

h.type	cum.freq	# indivs	pops	22:22673101	22:22673305	22:22673463	22:22673551	ENST00000390289
ref	0.9914	-	-	TCTC	C	C	C	
a	0.005	11		T	C	C	G	
d	0.00135	3		TCTC	T	C	G	5,7,8,10,13
c	0.00135	3		TCTC	C	T	G	10,13
b	0.0009	2		TCTC	C	C	G	9
								6

Table 13.141 - IGLV6-57

h.type	cum.freq	# indivs	pops	22:22550370 rs181723895	22:22550381 rs186024609	22:22550387 rs191529840	22:22550399 rs183090103	22:22550405 rs186363981	22:22550415 rs190816426	22:22550436 rs149903410	22:22550450 rs2073447
ref	0.2701	-	-	Gly->Ala	Asn->Asp	Met->Leu	Pro->Ser	Ser->Ala	Glu->Ala	Thr->Ser	Arg->Gly
a	0.3734	650	14	G	A	A	C	T	A	C	C
b	0.2802	500	14	G	A	A	C	T	A	C	C
c	0.0325	71	12	G	A	A	T	T	A	C	G
d	0.02015	44	11	G	A	A	T	T	A	C	C
e	0.0187	41	12	G	A	A	T	T	A	C	G
j	0.00045	1	1	G	A	A	T	T	A	C	C
k	0.00045	1	1	G	G	A	T	T	A	C	C
g	0.00045	1	1	G	A	T	C	T	A	C	C
h	0.00045	1	1	C	A	T	C	T	A	C	C
f	0.00045	1	1	G	A	A	C	T	A	C	C
i	0.00045	1	1	G	A	A	C	T	A	C	C
n	0.00045	1	1	G	A	A	C	T	A	C	C
m	0.00045	1	1	G	A	A	C	T	A	C	C
l	0.00045	1	1	G	A	A	C	T	A	G	C
p	0.00045	1	1	G	A	A	C	G	C	C	C
o	0.00045	1	1	G	A	A	C	T	A	C	C

Table 13.141 - IGLV6-57

h.type	cum.freq	# indivs	pops	22:22550466 rs144967736	22:22550490 rs182196598	22:22550510 rs2073448	22:22550525 rs187919298	22:22550565 rs192435787	22:22550670 rs188470766	ENST00000390285
ref	0.2701	-	-	Ile->Asn	Tyr->Phe	Ser->Ala	Ile->Leu	Arg->Gln	Ser->Asn	
a	0.3734	650	14	T	A	T	A	G	G	1,2,3,4,5,6,7,8,9,10,11,12,13,14
b	0.2802	500	14	T	A	G	A	G	G	1,2,3,4,5,6,7,8,9,10,11,12,13,14
c	0.0325	71	12	T	A	G	A	G	G	1,2,3,4,5,6,7,9,10,11,13,14
d	0.02015	44	11	T	A	G	A	G	G	1,2,3,4,6,7,9,10,11,13,14
e	0.0187	41	12	T	A	T	A	G	G	1,2,3,4,5,6,7,9,10,11,13,14
j	0.00045	1	1	T	A	T	A	G	■	14
k	0.00045	1	1	T	A	T	A	G	G	4
g	0.00045	1	1	T	A	G	A	G	G	10
h	0.00045	1	1	T	A	T	A	G	G	11
f	0.00045	1	1	T	T	G	C	G	G	10
i	0.00045	1	1	T	A	T	C	G	G	11
n	0.00045	1	1	■	A	T	A	G	G	13
m	0.00045	1	1	T	A	G	A	G	■	1
l	0.00045	1	1	T	A	T	A	G	G	3
p	0.00045	1	1	T	T	T	A	G	G	2
o	0.00045	1	1	T	A	T	A	■	G	10



Table 13.142 - IGLV7-43

h.type	cum.freq	# indivs	pops	22:22749426	22:22749555	22:22749576	22:22749580	22:22749591	22:22749676	22:22749693	22:22749704
ref	0.97765	-	-	Cys->Arg	Pro->Thr	Gly->Arg	Gly->Glu	Leu->Val	Ala->Val	Ser->Gly	His->Gln
a	0.01005	22		T	C	G	G	C	C	A	C
b	0.0087	19		T	C	■	G	C	C	A	C
c	0.00135	3		T	C	G	G	C	T	A	C
d	0.00045	1		T	■	G	G	C	C	A	<b>G</b>
g	0.00045	1		T	C	G	G	C	C	A	C
e	0.00045	1		T	C	G	G	<b>G</b>	C	A	C
h	0.00045	1		<b>C</b>	C	G	G	C	C	<b>G</b>	C
f	0.00045	1		T	C	G	■	C	C	A	C
											ENST00000390298
											2,3,4,6,7,9,10,11,14
											1,5,10,14
											4,13
											1
											6
											1
											1
											1

Table 13.143 - IGLV7-46

h.type	cum.freq	# indivs	pops	22:22724244 rs188705617	22:22724306 rs141014218	22:22724313 rs193180910	22:22724332 rs185250547	22:22724394 rs7292000	22:22724424 rs191717127	ENST00000390295
ref	0.0807	-	-	Ala->Gly	Leu->Met	Tyr->Cys	His->Gln	Leu->Ser	Tyr->Phe	
a	0.9166	1080	14	C	C	A	C	T	A	
c	0.0009	2	2	G	C	A	C	C	A	1,2,3,4,5,6,7,8,9,10,11,12,13,14
d	0.00045	1	1	C	C	A	C	C	A	5,10
e	0.00045	1	1	C	C	G	C	C	A	10
b	0.00045	1	1	C	■	A	C	C	T	9
f	0.00045	1	1	C	C	A	G	T	A	13
										2

Table 13.144 - IGLV8-61

h.type	cum.freq	# indivs	pops	Gly->Arg	Ser->Gly	Thr->Met	Ser->Ala	Arg->Cys	Ala->Thr	ENST00000390283
ref	0.7574	-	-	G	A	C	T	C	Ala->Thr	
a	0.2193	407	12	G	A	C	T	T	G	
b	0.02195	48	8	G	A	T	T	C	G	1,2,3,4,5,6,7,8,9,11,12,13
d	0.00045	1	1	G	G	C	T	T	G	1,3,4,5,6,7,9,11
e	0.00045	1	1	G	A	C	G	C	G	2
c	0.00045	1	1	■	A	C	T	T	■	2
									G	5

Table 13.145 - IGLV9-49

h.type	cum.fre	# indivs	pops	22:22697585 rs146805069	22:22697782 rs140612446	22:22697788 rs186017534	22:22697825 rs190876413	22:22697923 rs187756755	22:22697939 rs190665606	22:22697983 rs192733353	22:22697998 rs184707881	22:22698031 rs14889698
ref	0.98455	-	-	Ala->Pro	Thr->Ala	Pro->Ser	Thr->Lys	Thr->Ala	Gly->Glu	Ser->Pro	Tyr->His	Glu->Lys
a	0.00865	18	3	G	A	C	C	A	G	T	T	G
b	0.0032	7	2	G	<b>G</b>	C	C	A	G	T	T	G
c	0.0009	2	2	G	A	<b>T</b>	C	A	G	T	T	<b>■</b>
d	0.00045	1	1	G	A	C	<b>■</b>	A	G	T	T	G
g	0.00045	1	1	<b>C</b>	A	C	C	A	G	T	T	G
e	0.00045	1	1	G	A	C	C	A	G	<b>C</b>	T	G
h	0.00045	1	1	G	A	C	C	<b>G</b>	G	T	T	G
f	0.00045	1	1	G	A	C	C	A	G	T	<b>C</b>	G
i	0.00045	1	1	G	A	C	C	A	<b>■</b>	T	T	G

Table 13.145 - IGLV9-49

h.type	cum.fre	# indivs	pops	22:22698059	rs187595925	ENST00000427632
ref	0.98455	-	-	A		
a	0.00865	18	3	A		3,4,9
b	0.0032	7	2	A		6,13
c	0.0009	2	2	A		7,9
d	0.00045	1	1	A		1
g	0.00045	1	1	A		3
e	0.00045	1	1	T		13
h	0.00045	1	1	A		2
f	0.00045	1	1	A		13
i	0.00045	1	1	A		2

**Table 14A: Variants Having a Cumulative Frequency Greater than 5% or 10%**

<u>GENE SEGMENT</u>	<u>VARIANT NAME</u>	<u>CUMULATIVE FREQUENCY</u>	<u>CUMULATIVE FREQUENCY &gt;10%</u>	<u>NO UNIQUE POPULATIONS</u>
<b><u>Heavy C</u></b>				
<u>IGHA</u>	<u>IGHA1-a</u>	<u>0.31385</u>	<u>Yes</u>	<u>14</u>
<u>IGHA</u>	<u>IGHA2-a</u>	<u>0.35655</u>	<u>Yes</u>	<u>14</u>
<u>IGHA</u>	<u>IGHA2-b</u>	<u>0.1647</u>	<u>Yes</u>	<u>14</u>
<u>IGHG</u>	<u>IGHG2-a</u>	<u>0.28675</u>	<u>Yes</u>	<u>12</u>
<u>IGHG</u>	<u>IGHG3-a</u>	<u>0.2813</u>	<u>Yes</u>	<u>14</u>
<u>IGHG</u>	<u>IGHG3-b</u>	<u>0.19955</u>	<u>Yes</u>	<u>11</u>
<u>IGHG</u>	<u>IGHG4-a</u>	<u>0.19825</u>	<u>Yes</u>	<u>12</u>
<u>IGHM</u>	<u>IGHM-a</u>	<u>0.7527</u>	<u>Yes</u>	<u>14</u>
<u>IGHM</u>	<u>IGHM-b</u>	<u>0.15245</u>	<u>Yes</u>	<u>14</u>
<b><u>Heavy J</u></b>				
<u>IGHJ</u>	<u>IGHJ6-a</u>	<u>0.6845</u>	<u>Yes</u>	<u>14</u>
<b><u>Heavy V</u></b>				
<u>IGHV</u>	<u>IGHV1-2-a</u>	<u>0.3268</u>	<u>Yes</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV1-2-b</u>	<u>0.1304</u>	<u>Yes</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV1-2-c</u>	<u>0.0797</u>	<u>No</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV1-3-a</u>	<u>0.3564</u>	<u>Yes</u>	<u>14</u>

<u>IGHV</u>	<u>IGHV1-3-b</u>	<u>0.0724</u>	<u>No</u>	<u>13</u>
<u>IGHV</u>	<u>IGHV1-45-a</u>	<u>0.1286</u>	<u>Yes</u>	<u>12</u>
<u>IGHV</u>	<u>IGHV1-45-b</u>	<u>0.1126</u>	<u>Yes</u>	<u>12</u>
<u>IGHV</u>	<u>IGHV1-58-a</u>	<u>0.4503</u>	<u>Yes</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV1-58-b</u>	<u>0.3144</u>	<u>Yes</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV1-69-a</u>	<u>0.1689</u>	<u>Yes</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV1-69-b</u>	<u>0.1543</u>	<u>Yes</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV1-69-c</u>	<u>0.0561</u>	<u>No</u>	<u>13</u>
<u>IGHV</u>	<u>IGHV2-26-a</u>	<u>0.06135</u>	<u>No</u>	<u>13</u>
<u>IGHV</u>	<u>IGHV2-5-a</u>	<u>0.6657</u>	<u>Yes</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV2-70-a</u>	<u>0.31</u>	<u>Yes</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV2-70-b</u>	<u>0.16595</u>	<u>Yes</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV2-70-d</u>	<u>0.06035</u>	<u>No</u>	<u>13</u>
<u>IGHV</u>	<u>IGHV3-13-a</u>	<u>0.15245</u>	<u>Yes</u>	<u>13</u>
<u>IGHV</u>	<u>IGHV3-15-a</u>	<u>0.0572</u>	<u>No</u>	<u>11</u>
<u>IGHV</u>	<u>IGHV3-16-a</u>	<u>0.11075</u>	<u>Yes</u>	<u>13</u>
<u>IGHV</u>	<u>IGHV3-20-a</u>	<u>0.2829</u>	<u>Yes</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV3-20-b</u>	<u>0.1209</u>	<u>Yes</u>	<u>13</u>
<u>IGHV</u>	<u>IGHV3-23-a</u>	<u>0.09825</u>	<u>No</u>	<u>12</u>
<u>IGHV</u>	<u>IGHV3-33-a</u>	<u>0.145</u>	<u>Yes</u>	<u>12</u>
<u>IGHV</u>	<u>IGHV3-35-a</u>	<u>0.11765</u>	<u>Yes</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV3-35-b</u>	<u>0.09695</u>	<u>No</u>	<u>10</u>
<u>IGHV</u>	<u>IGHV3-38-a</u>	<u>0.0723</u>	<u>No</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV3-43-a</u>	<u>0.2428</u>	<u>Yes</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV3-43-b</u>	<u>0.08365</u>	<u>No</u>	<u>13</u>
<u>IGHV</u>	<u>IGHV3-48-a</u>	<u>0.3355</u>	<u>Yes</u>	<u>14</u>

<u>IGHV</u>	<u>IGHV3-48-b</u>	<u>0.20455</u>	<u>Yes</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV3-49-a</u>	<u>0.38705</u>	<u>Yes</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV3-49-b</u>	<u>0.28555</u>	<u>Yes</u>	<u>13</u>
<u>IGHV</u>	<u>IGHV3-53-a</u>	<u>0.2535</u>	<u>Yes</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV3-53-b</u>	<u>0.2264</u>	<u>Yes</u>	<u>13</u>
<u>IGHV</u>	<u>IGHV3-64-a</u>	<u>0.47965</u>	<u>Yes</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV3-64-b</u>	<u>0.06225</u>	<u>No</u>	<u>13</u>
<u>IGHV</u>	<u>IGHV3-66-a</u>	<u>0.27865</u>	<u>Yes</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV3-66-b</u>	<u>0.12575</u>	<u>Yes</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV3-66-c</u>	<u>0.0649</u>	<u>No</u>	<u>13</u>
<u>IGHV</u>	<u>IGHV4-28-a</u>	<u>0.41435</u>	<u>Yes</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV4-28-b</u>	<u>0.13715</u>	<u>Yes</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV4-28-c</u>	<u>0.07385</u>	<u>No</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV4-28-d</u>	<u>0.06485</u>	<u>No</u>	<u>13</u>
<u>IGHV</u>	<u>IGHV4-31-a</u>	<u>0.5475</u>	<u>Yes</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV4-31-b</u>	<u>0.2035</u>	<u>Yes</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV4-31-c</u>	<u>0.1554</u>	<u>Yes</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV4-39-a</u>	<u>0.10295</u>	<u>Yes</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV4-39-b</u>	<u>0.0918</u>	<u>No</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV4-4-a</u>	<u>0.1136</u>	<u>Yes</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV4-4-d</u>	<u>0.0817</u>	<u>No</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV4-4-f</u>	<u>0.06675</u>	<u>No</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV4-4-b</u>	<u>0.06385</u>	<u>No</u>	<u>13</u>
<u>IGHV</u>	<u>IGHV4-4-c</u>	<u>0.0599</u>	<u>No</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV4-4-e</u>	<u>0.05215</u>	<u>No</u>	<u>12</u>
<u>IGHV</u>	<u>IGHV4-61-a</u>	<u>0.5701</u>	<u>Yes</u>	<u>14</u>



<u>IGHV</u>	<u>IGHV4-61-b</u>	<u>0.18115</u>	<u>Yes</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV4-61-c</u>	<u>0.13535</u>	<u>Yes</u>	<u>13</u>
<u>IGHV</u>	<u>IGHV4-61-f</u>	<u>0.05855</u>	<u>No</u>	<u>14</u>
<b><u>Kappa C</u></b>				
<u>IGKC</u>	<u>IGKC-a</u>	<u>0.2427</u>	<u>Yes</u>	<u>14</u>
<b><u>Kappa V</u></b>				
<u>IGKV</u>	<u>IGKV1-17-a</u>	<u>0.08655</u>	<u>No</u>	<u>13</u>
<u>IGKV</u>	<u>IGKV1-17-b</u>	<u>0.0545</u>	<u>No</u>	<u>12</u>
<u>IGKV</u>	<u>IGKV1-8-a</u>	<u>0.20315</u>	<u>Yes</u>	<u>14</u>
<u>IGKV</u>	<u>IGKV1-8-b</u>	<u>0.1334</u>	<u>Yes</u>	<u>14</u>
<u>IGKV</u>	<u>IGKV1-9-a</u>	<u>0.1854</u>	<u>Yes</u>	<u>14</u>
<u>IGKV</u>	<u>IGKV1D-42-a</u>	<u>0.22345</u>	<u>Yes</u>	<u>14</u>
<u>IGKV</u>	<u>IGKV1D-8-a</u>	<u>0.1872</u>	<u>Yes</u>	<u>14</u>
<u>IGKV</u>	<u>IGKV2-24-a</u>	<u>0.1708</u>	<u>Yes</u>	<u>13</u>
<u>IGKV</u>	<u>IGKV2-30-a</u>	<u>0.24085</u>	<u>Yes</u>	<u>13</u>
<u>IGKV</u>	<u>IGKV2D-26-a</u>	<u>0.325</u>	<u>Yes</u>	<u>14</u>
<u>IGKV</u>	<u>IGKV2D-29-a</u>	<u>0.30175</u>	<u>Yes</u>	<u>14</u>
<u>IGKV</u>	<u>IGKV3-7-a</u>	<u>0.2018</u>	<u>Yes</u>	<u>14</u>
<u>IGKV</u>	<u>IGKV3D-11-a</u>	<u>0.35105</u>	<u>Yes</u>	<u>14</u>
<u>IGKV</u>	<u>IGKV3D-15-a</u>	<u>0.2061</u>	<u>Yes</u>	<u>14</u>
<u>IGKV</u>	<u>IGKV5-2-a</u>	<u>0.32315</u>	<u>Yes</u>	<u>14</u>
<b><u>Lambda C</u></b>				
<u>IGLC</u>	<u>IGLC7-a</u>	<u>0.0751</u>	<u>No</u>	<u>14</u>

<b><u>Lambda J</u></b>				
<u>IGLJ</u>	<u>IGLJ2-a</u>	<u>0.0741</u>	<u>No</u>	<u>13</u>
<u>IGLJ</u>	<u>IGLJ7-a</u>	<u>0.93225</u>	<u>Yes</u>	<u>14</u>
<b><u>Lambda V</u></b>				
<u>IGLV</u>	<u>IGLV10-54-a</u>	<u>0.50725</u>	<u>Yes</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV10-54-b</u>	<u>0.09425</u>	<u>No</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV11-55-a</u>	<u>0.4463</u>	<u>Yes</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV1-36-a</u>	<u>0.0942</u>	<u>No</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV1-47-a</u>	<u>0.8413</u>	<u>Yes</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV1-50-a</u>	<u>0.19325</u>	<u>Yes</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV1-51-a</u>	<u>0.1177</u>	<u>Yes</u>	<u>11</u>
<u>IGLV</u>	<u>IGLV2-14-a</u>	<u>0.48365</u>	<u>Yes</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV2-14-c</u>	<u>0.08855</u>	<u>No</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV2-14-b</u>	<u>0.0804</u>	<u>No</u>	<u>10</u>
<u>IGLV</u>	<u>IGLV2-18-a</u>	<u>0.763</u>	<u>Yes</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV2-23-a</u>	<u>0.34655</u>	<u>Yes</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV2-8-a</u>	<u>0.1451</u>	<u>Yes</u>	<u>11</u>
<u>IGLV</u>	<u>IGLV3-12-a</u>	<u>0.38715</u>	<u>Yes</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV3-12-b</u>	<u>0.1116</u>	<u>Yes</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV3-12-c</u>	<u>0.0774</u>	<u>No</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV3-19-a</u>	<u>0.10015</u>	<u>Yes</u>	<u>12</u>
<u>IGLV</u>	<u>IGLV3-21-a</u>	<u>0.31195</u>	<u>Yes</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV3-21-b</u>	<u>0.178</u>	<u>Yes</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV3-21-c</u>	<u>0.1613</u>	<u>Yes</u>	<u>14</u>

<u>IGLV</u>	<u>IGLV3-22-a</u>	<u>0.2962</u>	<u>Yes</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV3-25-a</u>	<u>0.6581</u>	<u>Yes</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV3-25-b</u>	<u>0.19965</u>	<u>Yes</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV4-60-a</u>	<u>0.4868</u>	<u>Yes</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV4-60-b</u>	<u>0.3607</u>	<u>Yes</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV5-37-a</u>	<u>0.1266</u>	<u>Yes</u>	<u>13</u>
<u>IGLV</u>	<u>IGLV5-45-a</u>	<u>0.1706</u>	<u>Yes</u>	<u>12</u>
<u>IGLV</u>	<u>IGLV5-48-a</u>	<u>0.0873</u>	<u>No</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV6-57-a</u>	<u>0.3734</u>	<u>Yes</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV6-57-b</u>	<u>0.2802</u>	<u>Yes</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV7-46-a</u>	<u>0.9166</u>	<u>Yes</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV8-61-a</u>	<u>0.2193</u>	<u>Yes</u>	<u>12</u>

**Table 14B: Variants Having a Cumulative Frequency Greater than 20%**

<u>GENE SEGMENT</u>	<u>VARIANT NAME</u>	<u>CUMULATIVE FREQUENCY</u>	<u>NO UNIQUE POPULATIONS</u>
<b><u>Heavy C</u></b>			
<u>IGHA</u>	<u>IGHA1-a</u>	<u>0.31385</u>	<u>14</u>
<u>IGHA</u>	<u>IGHA2-a</u>	<u>0.35655</u>	<u>14</u>
<u>IGHG</u>	<u>IGHG2-a</u>	<u>0.28675</u>	<u>12</u>
<u>IGHG</u>	<u>IGHG3-a</u>	<u>0.2813</u>	<u>14</u>
<u>IGHM</u>	<u>IGHM-a</u>	<u>0.7527</u>	<u>14</u>
<b><u>Heavy J</u></b>			
<u>IGHJ</u>	<u>IGHJ6-a</u>	<u>0.6845</u>	<u>14</u>

<b><u>Heavy V</u></b>			
<u>IGHV</u>	<u>IGHV1-2-a</u>	<u>0.3268</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV1-3-a</u>	<u>0.3564</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV1-58-a</u>	<u>0.4503</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV1-58-b</u>	<u>0.3144</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV2-5-a</u>	<u>0.6657</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV2-70-a</u>	<u>0.31</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV3-20-a</u>	<u>0.2829</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV3-43-a</u>	<u>0.2428</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV3-48-a</u>	<u>0.3355</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV3-48-b</u>	<u>0.20455</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV3-49-a</u>	<u>0.38705</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV3-49-b</u>	<u>0.28555</u>	<u>13</u>
<u>IGHV</u>	<u>IGHV3-53-a</u>	<u>0.2535</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV3-53-b</u>	<u>0.2264</u>	<u>13</u>
<u>IGHV</u>	<u>IGHV3-64-a</u>	<u>0.47965</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV3-66-a</u>	<u>0.27865</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV4-28-a</u>	<u>0.41435</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV4-31-a</u>	<u>0.5475</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV4-31-b</u>	<u>0.2035</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV4-61-a</u>	<u>0.5701</u>	<u>14</u>
<b><u>Kappa C</u></b>			
<u>IGKC</u>	<u>IGKC-a</u>	<u>0.2427</u>	<u>14</u>
	<u>IGKV1-8-a</u>	<u>0.20315</u>	<u>14</u>

<b><u>Kappa V</u></b>			
<u>IGKV</u>			
<u>IGKV</u>	<u>IGKV1D-42-a</u>	<u>0.22345</u>	<u>14</u>
<u>IGKV</u>	<u>IGKV2-30-a</u>	<u>0.24085</u>	<u>13</u>
<u>IGKV</u>	<u>IGKV2D-26-a</u>	<u>0.325</u>	<u>14</u>
<u>IGKV</u>	<u>IGKV2D-29-a</u>	<u>0.30175</u>	<u>14</u>
<u>IGKV</u>	<u>IGKV3-7-a</u>	<u>0.2018</u>	<u>14</u>
<u>IGKV</u>	<u>IGKV3D-11-a</u>	<u>0.35105</u>	<u>14</u>
<u>IGKV</u>	<u>IGKV3D-15-a</u>	<u>0.2061</u>	<u>14</u>
<u>IGKV</u>	<u>IGKV5-2-a</u>	<u>0.32315</u>	<u>14</u>
<b><u>Lambda J</u></b>			
<u>IGLJ</u>	<u>IGLJ7-a</u>	<u>0.93225</u>	<u>14</u>
<b><u>Lambda V</u></b>			
<u>IGLV</u>	<u>IGLV10-54-a</u>	<u>0.50725</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV11-55-a</u>	<u>0.4463</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV1-47-a</u>	<u>0.8413</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV2-14-a</u>	<u>0.48365</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV2-18-a</u>	<u>0.763</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV2-23-a</u>	<u>0.34655</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV3-12-a</u>	<u>0.38715</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV3-21-a</u>	<u>0.31195</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV3-22-a</u>	<u>0.2962</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV3-25-a</u>	<u>0.6581</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV4-60-a</u>	<u>0.4868</u>	<u>14</u>

<u>IGLV</u>	<u>IGLV4-60-b</u>	<u>0.3607</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV6-57-a</u>	<u>0.3734</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV6-57-b</u>	<u>0.2802</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV7-46-a</u>	<u>0.9166</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV8-61-a</u>	<u>0.2193</u>	<u>12</u>

**Table 14C: Variants Having a Cumulative Frequency Greater than 30%**

<u>GENE SEGMENT</u>	<u>VARIANT NAME</u>	<u>CUMULATIVE FREQUENCY</u>	<u>NO UNIQUE POPULATIONS</u>
<b><u>Heavy C</u></b>			
<u>IGHA</u>	<u>IGHA1-a</u>	<u>0.31385</u>	<u>14</u>
<u>IGHA</u>	<u>IGHA2-a</u>	<u>0.35655</u>	<u>14</u>
<u>IGHM</u>	<u>IGHM-a</u>	<u>0.7527</u>	<u>14</u>
<b><u>Heavy J</u></b>			
<u>IGHJ</u>	<u>IGHJ6-a</u>	<u>0.6845</u>	<u>14</u>
<b><u>Heavy V</u></b>			
<u>IGHV</u>	<u>IGHV1-2-a</u>	<u>0.3268</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV1-3-a</u>	<u>0.3564</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV1-58-a</u>	<u>0.4503</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV1-58-b</u>	<u>0.3144</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV2-5-a</u>	<u>0.6657</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV2-70-a</u>	<u>0.31</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV3-48-a</u>	<u>0.3355</u>	<u>14</u>

<u>IGHV</u>	<u>IGHV3-49-a</u>	<u>0.38705</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV3-64-a</u>	<u>0.47965</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV4-28-a</u>	<u>0.41435</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV4-31-a</u>	<u>0.5475</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV4-61-a</u>	<u>0.5701</u>	<u>14</u>
<u>IGKV</u>	<u>IGKV2D-26-a</u>	<u>0.325</u>	<u>14</u>
<u>IGKV</u>	<u>IGKV2D-29-a</u>	<u>0.30175</u>	<u>14</u>
<u>IGKV</u>	<u>IGKV3D-11-a</u>	<u>0.35105</u>	<u>14</u>
<u>IGKV</u>	<u>IGKV5-2-a</u>	<u>0.32315</u>	<u>14</u>
<b><u>Lambda J</u></b>			
<u>IGLJ</u>	<u>IGLJ7-a</u>	<u>0.93225</u>	<u>14</u>
<b><u>Lambda V</u></b>			
<u>IGLV</u>	<u>IGLV10-54-a</u>	<u>0.50725</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV11-55-a</u>	<u>0.4463</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV1-47-a</u>	<u>0.8413</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV2-14-a</u>	<u>0.48365</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV2-18-a</u>	<u>0.763</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV2-23-a</u>	<u>0.34655</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV3-12-a</u>	<u>0.38715</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV3-21-a</u>	<u>0.31195</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV3-25-a</u>	<u>0.6581</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV4-60-a</u>	<u>0.4868</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV4-60-b</u>	<u>0.3607</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV6-57-a</u>	<u>0.3734</u>	<u>14</u>

<u>IGLV</u>	<u>IGLV7-46-a</u>	<u>0.9166</u>	<u>14</u>
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**Table 14D: Variants Having a Cumulative Frequency Greater than 40%**

<u>GENE SEGMENT</u>	<u>VARIANT NAME</u>	<u>CUMULATIVE FREQUENCY</u>	<u>NO UNIQUE POPULATIONS</u>
<b><u>Heavy C</u></b>			
<u>IGHM</u>	<u>IGHM-a</u>	<u>0.7527</u>	<u>14</u>
<b><u>Heavy J</u></b>			
<u>IGHJ</u>	<u>IGHJ6-a</u>	<u>0.6845</u>	<u>14</u>
<b><u>Heavy V</u></b>			
<u>IGHV</u>	<u>IGHV1-58-a</u>	<u>0.4503</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV2-5-a</u>	<u>0.6657</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV3-64-a</u>	<u>0.47965</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV4-28-a</u>	<u>0.41435</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV4-31-a</u>	<u>0.5475</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV4-61-a</u>	<u>0.5701</u>	<u>14</u>
<b><u>Lambda J</u></b>			
<u>IGLJ</u>	<u>IGLJ7-a</u>	<u>0.93225</u>	<u>14</u>
<b><u>Lambda V</u></b>			
<u>IGLV</u>	<u>IGLV10-54-a</u>	<u>0.50725</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV11-55-a</u>	<u>0.4463</u>	<u>14</u>



<u>IGLV</u>	<u>IGLV1-47-a</u>	<u>0.8413</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV2-14-a</u>	<u>0.48365</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV2-18-a</u>	<u>0.763</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV3-25-a</u>	<u>0.6581</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV4-60-a</u>	<u>0.4868</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV7-46-a</u>	<u>0.9166</u>	<u>14</u>

**Table 14E: Variants Having a Cumulative Frequency Greater than 50%**

<u>GENE SEGMENT</u>	<u>VARIANT NAME</u>	<u>CUMULATIVE FREQUENCY</u>	<u>NO UNIQUE POPULATIONS</u>
<b><u>Heavy C</u></b>			
<u>IGHM</u>	<u>IGHM-a</u>	<u>0.7527</u>	<u>14</u>
<b><u>Heavy J</u></b>			
<u>IGHJ</u>	<u>IGHJ6-a</u>	<u>0.6845</u>	<u>14</u>
<b><u>Heavy V</u></b>			
<u>IGHV</u>	<u>IGHV2-5-a</u>	<u>0.6657</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV4-31-a</u>	<u>0.5475</u>	<u>14</u>
<u>IGHV</u>	<u>IGHV4-61-a</u>	<u>0.5701</u>	<u>14</u>
<b><u>Lambda J</u></b>			
<u>IGLJ</u>	<u>IGLJ7-a</u>	<u>0.93225</u>	<u>14</u>
<b><u>Lambda V</u></b>			
	<u>IGLV10-54-a</u>	<u>0.50725</u>	<u>14</u>

<u>IGLV</u>			
<u>IGLV</u>	<u>IGLV1-47-a</u>	<u>0.8413</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV2-18-a</u>	<u>0.763</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV3-25-a</u>	<u>0.6581</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV7-46-a</u>	<u>0.9166</u>	<u>14</u>

**Table 14F: Variants Having a Cumulative Frequency Greater than 60%**

<u>GENE SEGMENT</u>	<u>VARIANT NAME</u>	<u>CUMULATIVE FREQUENCY</u>	<u>NO UNIQUE POPULATIONS</u>
<b><u>Heavy C</u></b>			
<u>IGHM</u>	<u>IGHM-a</u>	<u>0.7527</u>	<u>14</u>
<b><u>Heavy J</u></b>			
<u>IGHJ</u>	<u>IGHJ6-a</u>	<u>0.6845</u>	<u>14</u>
<b><u>Heavy V</u></b>			
<u>IGHV</u>	<u>IGHV2-5-a</u>	<u>0.6657</u>	<u>14</u>
<b><u>Lambda J</u></b>			
<u>IGLJ</u>	<u>IGLJ7-a</u>	<u>0.93225</u>	<u>14</u>
<b><u>Lambda V</u></b>			
<u>IGLV</u>	<u>IGLV1-47-a</u>	<u>0.8413</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV2-18-a</u>	<u>0.763</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV3-25-a</u>	<u>0.6581</u>	<u>14</u>

<u>IGLV</u>	<u>IGLV7-46-a</u>	<u>0.9166</u>	<u>14</u>
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**Table 14G: Variants Having a Cumulative Frequency Greater than 70%**

<u>GENE SEGMENT</u>	<u>VARIANT NAME</u>	<u>CUMULATIVE FREQUENCY</u>	<u>NO UNIQUE POPULATIONS</u>
<b><u>Heavy C</u></b>			
<u>IGHM</u>	<u>IGHM-a</u>	<u>0.7527</u>	<u>14</u>
<b><u>Lambda J</u></b>			
<u>IGLJ</u>	<u>IGLJ7-a</u>	<u>0.93225</u>	<u>14</u>
<b><u>Lambda V</u></b>			
<u>IGLV</u>	<u>IGLV1-47-a</u>	<u>0.8413</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV2-18-a</u>	<u>0.763</u>	<u>14</u>
<u>IGLV</u>	<u>IGLV7-46-a</u>	<u>0.9166</u>	<u>14</u>

**Table 14H: Variants Having a Cumulative Frequency Greater than 80%**

<u>GENE SEGMENT</u>	<u>VARIANT NAME</u>	<u>CUMULATIVE FREQUENCY</u>	<u>NO UNIQUE POPULATIONS</u>
<b><u>Lambda J</u></b>			
<u>IGLJ</u>	<u>IGLJ7-a</u>	<u>0.93225</u>	<u>14</u>
<b><u>Lambda V</u></b>			
<u>IGLV</u>	<u>IGLV1-47-a</u>	<u>0.8413</u>	<u>14</u>

<u>IGLV</u>			
<u>IGLV</u>	<u>IGLV7-46-a</u>	<u>0.9166</u>	<u>14</u>

**Table 14I: Variants Having a Cumulative Frequency Greater than 90%**

<u>GENE SEGMENT</u>	<u>VARIANT NAME</u>	<u>CUMULATIVE FREQUENCY</u>	<u>NO UNIQUE POPULATIONS</u>
<b><u>Lambda J</u></b>			
<u>IGLJ</u>	<u>IGLJ7-a</u>	<u>0.93225</u>	<u>14</u>
<b><u>Lambda V</u></b>			
<u>IGLV</u>	<u>IGLV7-46-a</u>	<u>0.9166</u>	<u>14</u>

**Table 15: NAÏVE HEAVY CHAIN REPERTOIRES**

<b><u>TABLE 15A:</u></b>			
<b><u>All Naïve – J Usage</u></b>			
<u>J</u>	<u>Average HCDR3 Length</u>	<u>Count</u>	<u>%</u>
IGHJ1*01	11	12	0.90%
IGHJ2*01	12	47	3.51%
IGHJ3*02	12	205	15.30%
IGHJ4*02	11	689	51.42%
IGHJ5*02	13	88	6.57%
<b><u>IGHJ6*02</u></b>	<b><u>16</u></b>	<b><u>299</u></b>	<b><u>22.31%</u></b>

<b><u>Naïve HCDR3&gt;=20</u></b>			
<u>J</u>	<u>Average HCDR3 Length</u>	<u>Count</u>	<u>%</u>

IGHJ3*02	20	1	2.22%
IGHJ4*02	20	1	2.22%
IGHJ5*02	20	2	4.44%
<b>IGHJ6*02</b>	<b>21</b>	<b>41</b>	<b>91.11%</b>

**Naïve – Long HCDR3 using IGHJ6\*02**

<u>HCDR3Length</u>	<u>Count</u>
20	19
21	10
22	7
23	3
24	1
26	1

<b><u>TABLE 15B:</u></b>			
<b><u>All Naïve – J &amp; D Usage</u></b>			
<u>J</u>	<u>D</u>	<u>Average HCDR3 Length</u>	<u>Count</u>
IGHJ6*02	IGHD3-9*01	19	21
IGHJ6*02	IGHD4-17*01	18	7
IGHJ6*02	IGHD3-10*01	17	98
IGHJ6*02	IGHD2-2*02	17	4
IGHJ6*02	IGHD5-24*01	17	1
IGHJ6*02	IGHD6-19*01	16	30
IGHJ6*02	IGHD3-22*01	16	5
IGHJ6*02	IGHD6-13*01	16	33
IGHJ6*02	IGHD5-12*01	15	7
IGHJ6*02	IGHD1-26*01	15	25
IGHJ6*02	IGHD1-20*01	14	6
IGHJ6*02	IGHD5-18*01	14	3
IGHJ6*02	IGHD3-16*02	14	6
IGHJ6*02	IGHD2-21*02	13	5
IGHJ6*02	IGHD1-14*01	13	4
IGHJ6*02	IGHD7-27*02	12	8
IGHJ6*02	IGHD1-1*01	12	8
IGHJ6*02	IGHD6-25*01	12	3
IGHJ6*02	IGHD4-23*01	11	2

<b>Naive HCDR3&gt;=20</b>			
<u>J</u>	<u>D</u>	<u>Average HCDR3 Length</u>	<u>Count</u>
IGHJ6*02	IGHD6-19*01	23	1
IGHJ6*02	IGHD4-17*01	22	2
IGHJ6*02	IGHD3-9*01	21	11
IGHJ6*02	IGHD3-10*01	21	25
IGHJ6*02	IGHD6-13*01	20	1
IGHJ6*02	IGHD3-22*01	20	1

<b>TABLE 15C: All Naive – V &amp; J Usage</b>			
<u>V</u>	<u>J</u>	<u>Average HCDR3 Length</u>	<u>Count</u>
IGHV3-21*03	IGHJ6*02	20	5
IGHV1-18*01	IGHJ6*02	20	10
IGHV1-2*02	IGHJ6*02	19	5
IGHV3-13*01	IGHJ6*02	19	3
IGHV3-7*01	IGHJ6*02	19	6
IGHV1-8*01	IGHJ6*02	19	21
IGHV7-4-1*01	IGHJ6*02	19	9
IGHV3-9*01	IGHJ6*02	18	13
IGHV4-61*01	IGHJ6*02	17	20
IGHV3-23*04	IGHJ6*02	17	6
IGHV4-4*02	IGHJ6*02	16	34
IGHV1-3*01	IGHJ6*02	16	39
IGHV3-20*d01	IGHJ6*02	16	1
IGHV6-1*01	IGHJ6*02	14	69
IGHV2-5*10	IGHJ6*02	14	8
IGHV3-11*01	IGHJ6*02	14	3
IGHV3-66*03	IGHJ6*02	13	21
IGHV3-15*01	IGHJ6*02	12	3

<b>Naive HCDR3&gt;=20</b>			
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<u>V</u>	<u>J</u>	<u>Average HCDR3 Length</u>	<u>Count</u>
IGHV3-21*03	IGHJ6*02	23	3
IGHV3-13*01	IGHJ6*02	22	1
IGHV3-7*01	IGHJ6*02	22	3
IGHV6-1*01	IGHJ6*02	21	3
IGHV1-8*01	IGHJ6*02	21	10
IGHV1-2*02	IGHJ6*02	21	3
IGHV7-4-1*01	IGHJ6*02	21	3
IGHV1-3*01	IGHJ6*02	21	4
IGHV1-18*01	IGHJ6*02	21	5
IGHV4-4*02	IGHJ6*02	20	3
IGHV3-9*01	IGHJ6*02	20	2
IGHV3-23*04	IGHJ6*02	20	1

**TABLE 16: IMMUNISED HEAVY CHAIN REPERTOIRES**

<b><u>TABLE 16A:</u></b>			
<b><u>All Immunised – J Usage</u></b>			
<u>J</u>	<u>HCDR3 Length</u>	<u>Count</u>	<u>%</u>
IGHJ1*01	11	2	0.78%
IGHJ2*01	14	7	2.73%
IGHJ3*02	15	12	4.69%
IGHJ4*02	15	120	46.88%
IGHJ5*02	15	19	7.42%
<b>IGHJ6*02</b>	<b>17</b>	<b>96</b>	<b>37.50%</b>

<b><u>HCDR3&gt;20</u></b>			
<u>J</u>	<u>HCDR3 Length</u>	<u>Count</u>	<u>%</u>
IGHJ4*02	22	2	22.22%
IGHJ5*02	25	1	11.11%
<b>IGHJ6*02</b>	<b>21</b>	<b>6</b>	<b>66.67%</b>

**Immunised – Long HCDR3 using IGHJ6\*02**

<u>HCDR3Length</u>	<u>Count</u>
20	4

21	1
24	1

**TABLE 16B:**  
**All Immunised – J & D Usage**

<u>J</u>	<u>D</u>	<u>HCDR3 Length</u>	<u>Count</u>
IGHJ6*02	IGHD4-17*01	18	1
IGHJ6*02	IGHD1-26*01	18	5
IGHJ6*02	IGHD6-19*01	17	27
IGHJ6*02	IGHD3-10*01	17	42
IGHJ6*02	IGHD6-13*01	16	2
IGHJ6*02	IGHD5-18*01	15	2
IGHJ6*02	IGHD4-23*01	15	5
IGHJ6*02	IGHD5-12*01	14	3
IGHJ6*02	IGHD3-16*02	14	3
IGHJ6*02	IGHD3-22*01	13	3
IGHJ6*02	IGHD3-9*01	12	1
IGHJ6*02	IGHD1-20*01	11	1

<u>HCDR3&gt;20</u>			
<u>J</u>	<u>D</u>	<u>HCDR3 Length</u>	<u>Count</u>
IGHJ6*02	IGHD3-10*01	21	4
IGHJ6*02	IGHD6-19*01	20	1
IGHJ6*02	IGHD1-26*01	20	1

**TABLE 16C:**  
**All Immunised – V & J Usage**

<u>V</u>	<u>J</u>	<u>HCDR3 Length</u>	<u>Count</u>
IGHV3-7*01	IGHJ6*02	18	33
IGHV1-8*01	IGHJ6*02	17	3
IGHV3-9*01	IGHJ6*02	17	22
IGHV3-13*01	IGHJ6*02	17	5
IGHV1-2*02	IGHJ6*02	16	8
IGHV3-11*01	IGHJ6*02	16	3
IGHV4-4*02	IGHJ6*02	15	9
IGHV6-1*01	IGHJ6*02	15	6
IGHV1-3*01	IGHJ6*02	15	6



<b>HCDR3&gt;20</b>			
<u>V</u>	<u>J</u>	<u>HCDR3 Length</u>	<u>Count</u>
IGHV3-7*01	IGHJ6*02	21	4
IGHV3-11*01	IGHJ6*02	21	1
IGHV4-4*02	IGHJ6*02	20	1

**TABLE 17: ANTIGEN-SPECIFIC HEAVY CHAIN REPERTOIRES**

<b><u>TABLE 17A:</u></b>			
<b><u>All Antigen-Specific – J Usage</u></b>			
<u>J</u>	<u>HCDR3 Length</u>	<u>Count</u>	<u>%</u>
IGHJ1*01	12	2	1.68%
IGHJ3*02	17	4	3.36%
IGHJ4*02	13	64	53.78%
IGHJ5*02	19	6	5.04%
<b>IGHJ6*02</b>	<b>17</b>	<b>43</b>	<b>36.13%</b>
<b><u>HCDR3&gt;20</u></b>			
<u>J</u>	<u>HCDR3 Length</u>	<u>Count</u>	<u>%</u>
IGHJ4*02	22	1	9.09%
IGHJ5*02	21	3	27.27%
<b>IGHJ6*02</b>	<b>21</b>	<b>7</b>	<b>63.64%</b>

**Immunised – Long HCDR3 using IGHJ6\*02**

<u>HCDR3Length</u>	<u>Count</u>
20	4
21	1
22	1
24	1

<b>TABLE 17B:</b> <b>All Antigen-Specific – J &amp; D Usage</b>			
<u>J</u>	<u>D</u>	<u>HCDR3 Length</u>	<u>Count</u>
IGHJ6*02	IGHD3-9*01	19	6
IGHJ6*02	IGHD3-10*01	19	12
IGHJ6*02	IGHD6-19*01	17	13
IGHJ6*02	IGHD5-12*01	16	2
IGHJ6*02	IGHD2-15*01	16	1
IGHJ6*02	IGHD1-26*01	15	7
IGHJ6*02	IGHD3-16*02	12	1
IGHJ6*02	IGHD5-18*01	11	1
<b>HCDR3&gt;20</b>			
<u>J</u>	<u>D</u>	<u>HCDR3 Length</u>	<u>Count</u>
IGHJ6*02	IGHD3-9*01	21	1
IGHJ6*02	IGHD3-10*01	21	6

<b>TABLE 17C:</b> <b>All Antigen-Specific – V &amp; J Usage</b>			
<u>V</u>	<u>J</u>	<u>HCDR3 Length</u>	<u>Count</u>
IGHV1-3*01	IGHJ6*02	18	1
IGHV3-7*01	IGHJ6*02	18	6
IGHV3-9*01	IGHJ6*02	18	6
IGHV1-8*01	IGHJ6*02	17	6
IGHV3-20*d01	IGHJ6*02	17	3
IGHV3-11*01	IGHJ6*02	17	8
IGHV4-4*02	IGHJ6*02	17	5
IGHV3-15*01	IGHJ6*02	15	5
IGHV3-13*01	IGHJ6*02	15	3
<b>HCDR3&gt;20</b>			
<u>V</u>	<u>J</u>	<u>HCDR3 Length</u>	<u>Count</u>
IGHV4-4*02	IGHJ6*02	20	2

IGHV1-8*01	IGHJ6*02	22	2
IGHV3-9*01	IGHJ6*02	20	1
IGHV3-11*01	IGHJ6*02	21	1
IGHV3-20*d01	IGHJ6*02	22	1

**Table 18: Sequence Correlation Table**

SEQ ID NO:	DESCRIPTION:
1	JH5 reference (nucleotide sequence)
2	JH6*03=JH6 reference (nucleotide sequence)
3	JH2 reference (nucleotide sequence)
4	JH6*02 (nucleotide sequence)
5	JH5 reference (nucleotide sequence)
6	JH5 reference (amino acid sequence)
7	JH6*03=JH6 reference (nucleotide sequence)
8	JH6*03=JH6 reference (amino acid sequence)

9	JH2 reference (nucleotide sequence)
10	JH2 reference (amino acid sequence)
11	IGHV1-2*01
12	IGHV1-3*01
13	IGHV1-8*01
14	IGHV1-24*01
15	IGHV1-45*01
16	IGHV1-46*01
17	IGHV1-58*01
18	IGHV1-69*01
19	IGHV1-c*01

20	IGHV1-f*01
21	IGHV2-5*01
22	IGHV2-26*01
23	IGHV2-70*01
24	IGHV3-7*01
25	IGHV3-9*01
26	IGHV3-11*01
27	IGHV3-13*01
28	IGHV3-15*01
29	IGHV3-16*01
30	IGHV3-19*01
31	IGHV3-20*01
32	IGHV3-21*01
33	IGHV3-23*01
34	IGHV3-30*01

35	IGHV3-30-3*01
36	IGHV3-33*01
37	IGHV3-35*01
38	IGHV3-38*01
39	IGHV3-43*01
40	IGHV3-47*01
41	IGHV3-48*01
42	IGHV3-49*01
43	IGHV3-53*01
44	IGHV3-64*01
45	IGHV3-66*01
46	IGHV3-72*01
47	IGHV3-73*01
48	IGHV3-74*01
49	IGHV3-d*01

50	IGHV3-h*01
51	IGHV4-4*01
52	IGHV4-28*01
53	IGHV4-30-2*01
54	IGHV4-30-4*01
55	IGHV4-31*01
56	IGHV4-34*01
57	IGHV4-39*01
58	IGHV4-55*01
59	IGHV4-59*01
60	IGHV4-61*01
61	IGHV4-b*01
62	IGHV5-51*01
63	IGHV5-a*01

64	IGHV6-1*01
65	IGHV7-4-1*01
66	IGHV7-81*01
67	IGHD1-1*01
68	IGHD1-7*01
69	IGHD1-14*01
70	IGHD1-20*01
71	IGHD1-26*01
72	IGHD2-2*01
73	IGHD2-8*01
74	IGHD2-15*01
75	IGHD2-21*01
76	IGHD3-3*01
77	IGHD3-9*01
78	IGHD3-10*01
79	IGHD3-16*01
80	IGHD3-22*01
81	IGHD4-4*01
82	IGHD4-11*01
83	IGHD4-17*01
84	IGHD4-23*01
85	IGHD5-5*01
86	IGHD5-12*01

87	IGHD5-18*01
88	IGHD5-24*01
89	IGHD6-6*01
90	IGHD6-13*01
91	IGHD6-19*01
92	IGHD6-25*01
93	IGHD6-25*01
94	JH1*01
95	JH2*01
96	JH3*01
97	JH4*01
98	JH5*01
99	JH6*01
100	IGKV1-5*01
101	IGKV1-6*01
102	IGKV1-8*01
103	IGKV1D-8*01
104	IGKV1-9*01
105	IGKV1-12*01
106	IGKV1-13*01

107	IGKV1-16*01
108	IGKV1D-16*01
109	IGKV1-17*01
110	IGKV1D-17*01
111	IGKV1-27*01
112	IGKV1-33*01
113	IGKV1D-33*01
114	IGKV1-37*01
115	IGKV1D-37*01
116	IGKV1-39*01
117	IGKV1D-39
118	IGKV1D-42*01
119	IGKV1D-43*01
120	IGKV2-24*01
121	IGKV2D-24*01

122	IGKV2-28*01
123	IGKV2D-28*01
124	IGKV2-29*01
125	IGKV2D-29*01
126	IGKV2-30*01
127	IGKV2D-30*01
128	IGKV2-40*01
129	IGKV2D-40*01
130	IGKV3-7*01
131	IGKV3-7*03
132	IGKV3D-7*01
133	IGKV3-11*01
134	IGKV3D-11*01
135	IGKV3-15*01
136	IGKV3D-15*01

137	IGKV3-20*01
138	IGKV3D-20*01
139	IGKV4-1*01
140	IGKV5-2*01
141	IGKV6-21*01
142	IGKV6D-21*01
143	IGKV6D-41*01
144	IGLV1-36*01
145	IGLV1-40*01
146	IGLV1-41*01
147	IGLV1-44*01
148	IGLV1-47*01
149	IGLV1-50*01
150	IGLV1-51*01
151	IGLV2-8*01

152	IGLV2-11*01
153	IGLV2-14*01
154	IGLV2-18*01
155	IGLV2-23*01
156	IGLV2-33*01
157	IGLV3-1*01
158	IGLV3-10*01
159	IGLV3-12*01
160	IGLV3-16*01
161	IGLV3-19*01
162	IGLV3-21*01
163	IGLV3-22*01
164	IGLV3-25*01
165	IGLV3-27*01
166	IGLV3-32*01

167	IGLV4-3*01
168	IGLV4-60*01
169	IGLV4-69*01
170	IGLV5-37*01
171	IGLV5-45*01
172	IGLV5-48*01
173	IGLV5-52*01
174	IGLV6-57*01
175	IGLV7-43*01
176	IGLV7-46*01
177	IGLV8-61*01
178	IGLV9-49*01
179	IGLV10-54*01



180	IGLV11-55*01
181	IGKJ1*01
182	IGKJ2*01
183	IGKJ3*01
184	IGKJ4*01
185	IGKJ5*01
186	IGLJ1*01
187	IGLJ2*01
188	IGLJ3*01
189	IGLJ4*01
190	IGLJ5*01
191	IGLJ6*01
192	IGLJ7*01
193	IGKV2D-26*01
194	IGLV1-36*01
195	IGLV1-47*01
196	IGLV1-50*01
197	IGLV1-51*01
198	IGLV10-54*01

199	IGLV11-55*01
200	IGLV2-14*01
201	IGLV2-18*01
202	IGLV2-23*01
203	IGLV2-8*01
204	IGLV3-1*01
205	IGLV3-12*01
206	IGLV3-19*01
207	IGLV3-21*01
208	IGLV3-22*01
209	IGLV3-25*01
210	IGLV4-60*01
211	IGLV5-37*01
212	IGLV5-45*01

213	IGLV5-48*01
214	IGLV6-57*01
215	IGLV7-46*01
216	IGLV8-61*01
217	IGHG3
218	IGHA2
219	IGHG2

220	IGHE
221	IGHE
222	IGHA1
223	IGHG1
224	IGHG1

225	IGHM
226	IGHG1
227	IGHD

228	IGHG4
229	IGHE
230	IGKC
231	IGLC2
232	IGLC1
233	IGLC3
234	IGLC7
235	JH6*02 (nucleotide sequence)
236	JH6*01 & JH6*02(amino acid sequence)
237	Human RSS-JH6*02
238	Human RSS
239	IGHJ1 ref

240	IGHJ2 ref
241	IGHJ3 ref
242	IGHJ4 ref
243	IGHJ5 ref
244	IGHJ6 ref
245	IGHV1-18 ref
246	IGHV1-2 ref
247	IGHV1-24 ref
248	IGHV1-3 ref
249	IGHV1-45 ref
250	IGHV1-46 ref
251	IGHV1-58 ref
252	IGHV1-69 ref
253	IGHV1-8 ref
254	IGHV2-26 ref
255	IGHV2-5 ref
256	IGHV2-70 ref
257	IGHV3-11 ref
258	IGHV3-13 ref
259	IGHV3-15 ref

260	IGHV3-16 ref
261	IGHV3-20 ref
262	IGHV3-21 ref
263	IGHV3-23 ref
264	IGHV3-30 ref
265	IGHV3-33 ref
266	IGHV3-35 ref
267	IGHV3-38 ref
268	IGHV3-43 ref
269	IGHV3-48 ref
270	IGHV3-49 ref
271	IGHV3-53 ref
272	IGHV3-64 ref
273	IGHV3-66 ref
274	IGHV3-7 ref

275	IGHV3-72 ref
276	IGHV3-73 ref
277	IGHV3-74 ref
278	IGHV3-9 ref
279	IGHV4-28 ref
280	IGHV4-31 ref
281	IGHV4-34 ref
282	IGHV4-39 ref
283	IGHV4-4 ref
284	IGHV4-59 ref
285	IGHV4-61 ref
286	IGHV5-51 ref
287	IGHV6-1 ref
288	IGHV7-81 ref
289	IGKJ2 ref
290	IGKJ3 ref
291	IGKJ4 ref

292	IGKJ5 ref
293	IGKV1-16 ref
294	IGKV1-17 ref
295	IGKV1-5 ref
296	IGKV1-6 ref
297	IGKV1-8 ref
298	IGKV1-9 ref
299	IGKV1D-12 ref
300	IGKV1D-16 ref
301	IGKV1D-17 ref
302	IGKV1D-42 ref
303	IGKV1D-43 ref
304	IGKV1D-8 ref
305	IGKV2-24 ref
306	IGKV2-30 ref
307	IGKV2D-24 ref

308	IGKV2D-26 ref
309	IGKV2D-29 ref
310	IGKV2D-30 ref
311	IGKV3-11 ref
312	IGKV3-20 ref
313	IGKV3-7 ref
314	IGKV3D-11 ref
315	IGKV3D-15 ref
316	IGKV3D-20 ref
317	IGKV4-1 ref
318	IGKV5-2 ref
319	IGKV6-21 ref
320	IGKV6D-21 ref
321	IGKV6D-41 ref
322	IGLJ1 ref

323	IGLJ2 ref
324	IGLJ3 ref
325	IGLJ5 ref
326	IGLJ6 ref
327	IGLJ7 ref
328	IGLV10-54 ref
329	IGLV11-55 ref
330	IGLV1-36 ref
331	IGLV1-40 ref
332	IGLV1-44 ref
333	IGLV1-47 ref
334	IGLV1-50 ref
335	IGLV1-51 ref
336	IGLV2-11 ref
337	IGLV2-14 ref
338	IGLV2-18 ref
339	IGLV2-23 ref
340	IGLV2-33 ref

341	IGLV2-8 ref
342	IGLV3-1 ref
343	IGLV3-10 ref
344	IGLV3-12 ref
345	IGLV3-16 ref
346	IGLV3-19 ref
347	IGLV3-21 ref
348	IGLV3-22 ref
349	IGLV3-25 ref
350	IGLV3-27 ref
351	IGLV3-32 ref
352	IGLV3-9 ref
353	IGLV4-3 ref
354	IGLV4-60 ref

355	IGLV4-69 ref
356	IGLV5-37 ref
357	IGLV5-45 ref
358	IGLV5-48 ref
359	IGLV5-52 ref
360	IGLV6-57 ref
361	IGLV7-43 ref
362	IGLV7-46 ref
363	IGLV8-61 ref
364	IGLV9-49 ref

365	IGHG1 ref GENOMIC
366	IGHG1 ref - CDS (ensembl transcript ID ENST00000390542)
367	IGHG1 ref - CDS (Ensembl transcript ID ENST00000390549)
368	IGHG1 ref - CDS (ensembl transcript ID ENST00000390548)

369	IGHG2 ref - GENOMIC
370	IGHG2 ref - CDS (ensembl transcript ID ENST00000390545)
371	IGHG2-a CDS
372	IGHG2-a GENOMIC
373	IGHG3 ref - GENOMIC
374	IGHG3 ref - CDS (ensembl transcript ID ENST00000390551)
375	IGHG3-a CDS



376	IGHG3-a GENOMIC
377	IGHG3-b CDS
378	IGHG3-b GENOMIC
379	IGHG4 ref - GENOMIC
380	IGHG4 ref - CDS (ensembl transcript ID ENST00000390543)
381	IGHG4-a CDS

382	IGHG4-a GENOMIC
383	IGHA1 ref GENOMIC
384	IGHA1 ref- CDS (ensembl transcript ID ENST00000390547)
385	IGHA1-a GENOMIC
386	IGHA1-a CDS
387	IGHA2 ref GENOMIC
388	IGHA2 ref- CDS (ensembl transcript ID ENST00000390539)

389	IGHA2-a GENOMIC
390	IGHA2-a CDS
391	IGHA2-b GENOMIC
392	IGHA2-b CDS

393	IGHD ref GENOMIC
394	IGHD ref- CDS (ensembl transcript ID ENST00000390556)

395	IGHE ref GENOMIC
396	IGHE ref- CDS (ensembl transcript ID ENST00000390541)
397	IGHE ref- CDS (ensembl transcript ID ENST00000576077)
398	IGHE ref- CDS (ensembl transcript ID ENST00000577108)
399	IGHM ref GENOMIC
400	IGHM ref- CDS (ensembl transcript ID ENST00000390559)

401	IGHM-a GENOMIC
402	IGHM-a CDS
403	IGHM-b GENOMIC
404	IGHM-b CDS
405	IGHD7-27 GENOMIC
406	IGHD2-15 GENOMIC
407	IGHD3-16 GENOMIC
408	IGHD6-6 ref GENOMIC
409	IGHD5-18 ref GENOMIC
410	IGHD2-2 ref GENOMIC
411	IGHD4-11 ref GENOMIC
412	IGHD5-12 ref

	GENOMIC
413	IGHD3-3 ref GENOMIC
414	IGHD2-8 ref GENOMIC
415	IGHD4-4 ref GENOMIC
416	IGHD4-23 ref GENOMIC
417	IGHD1-14 ref GENOMIC
418	IGHD3-10 ref GENOMIC
419	IGHD1-26 ref GENOMIC
420	IGHD3-9 ref GENOMIC
421	IGHD1-1 ref GENOMIC
422	IGHD6-25 ref GENOMIC
423	IGHD5-24 ref GENOMIC
424	IGHD2-21 ref GENOMIC
425	IGHD1-20 ref GENOMIC
426	IGHD6-13 ref GENOMIC
427	IGHD4-17 ref GENOMIC

428	IGHD3-22 ref GENOMIC
429	IGHD5-5 ref GENOMIC
430	IGHD6-19 ref GENOMIC
431	IGHD1-7 ref GENOMIC
432	Rabbit JH6 (amino acid)
433	Rabbit JH6 (nucleotide)
434	Sheep JH6 (amino acid)
435	Sheep JH6 (nucleotide)
436	Bovine JH6 (amino acid)
437	Bovine JH6 (nucleotide)
438	Dog JH3 (amino acid)
439	Dog JH3 (nucleotide)
440	Human JH6*02 (amino acid)
441	Human JH6*02 (nucleotide)
442	Leader Sequence (nucleotide)
443	Leader Sequence (amino acid)
444	PRIMER
445	PRIMER

**Table 19**

	IgG1	IgG2	IgG3	IgG4
Complement activation				
Classical pathway	+++	+	+++	-
Alternative pathway	-	+	-	-
Fc receptor recognition				
FcγRI	+++	-	+++	++
FcγRIIa, 131R/R	++	-	++	-
FcγRIIa, 131H/H	+	+	++	-
FcγRIIb	++	-	++	+
FcγRIII	+	+/-	+	+/-

**Table 20****Table 20A: Variants Found In No More Than 3 Human Populations**

TYPE	VARIANT	C.FREQ	C.FREQ (%)	#INDIVIDUALS	#POPULATIONS
IGKV	IGKV1D-8-b	0.01785	1.79%	36	3
IGHV	IGHV3-11-a	0.01695	1.70%	36	3
IGHV	IGHV5-51-a	0.0164	1.64%	35	3
IGHV	IGHV2-5-b	0.01055	1.06%	18	3
IGLV	IGLV2-14-g	0.0104	1.04%	22	3
IGHV	IGHV7-81-b	0.01005	1.01%	21	3
IGKV	IGKV1D-42-b	0.01005	1.01%	21	3
IGKV	IGKV3-7-b	0.01	1.00%	19	3
IGHV	IGHV1-69-d	0.00995	1.00%	22	3
IGLV	IGLV2-14-e	0.00995	1.00%	19	3
IGLV	IGLV9-49-a	0.00865	0.87%	18	3
IGLV	IGLV1-44-b	0.00775	0.78%	16	2
IGLV	IGLV2-11-a	0.00775	0.78%	17	3
IGKV	IGKV3-11-d	0.0073	0.73%	14	3
IGKV	IGKV1D-43-b	0.00685	0.69%	14	3
IGHV	IGHV3-38-d	0.0064	0.64%	13	3

IGLJ	IGLJ6-a	0.00595	0.60%	13	3
IGHV	IGHV5-51-b	0.0055	0.55%	11	3
IGLV	IGLV5-45-c	0.00545	0.55%	12	2
IGHV	IGHV4-31-i	0.0054	0.54%	10	3
IGKV	IGKV2D-26-b	0.00455	0.46%	9	3
IGLV	IGLV1-36-c	0.00455	0.46%	10	2
IGHV	IGHV1-24-c	0.00415	0.42%	9	3
IGHV	IGHV1-8-c	0.00405	0.41%	8	2
IGLV	IGLV1-47-d	0.00405	0.41%	9	3
IGHV	IGHV3-43-i	0.00365	0.37%	8	3
IGHV	IGHV3-53-h	0.0036	0.36%	8	2
IGLV	IGLV10-54-e	0.0036	0.36%	8	3
IGLV	IGLV1-36-f	0.0036	0.36%	8	2
IGLV	IGLV5-45-d	0.0036	0.36%	8	3
IGHV	IGHV1-46-b	0.0032	0.32%	7	3
IGHV	IGHV2-26-c	0.0032	0.32%	7	2
IGLJ	IGLJ6-b	0.0032	0.32%	7	3
IGLV	IGLV3-32-a	0.0032	0.32%	7	3
IGLV	IGLV9-49-b	0.0032	0.32%	7	2
IGHJ	IGHJ6-b	0.00315	0.32%	7	2
IGHV	IGHV3-16-d	0.00315	0.32%	7	3
IGHV	IGHV4-28-k	0.00315	0.32%	7	3
IGKJ	IGKJ2-d	0.00315	0.32%	6	3
IGLV	IGLV10-54-f	0.00315	0.32%	7	2
IGLV	IGLV2-8-c	0.00315	0.32%	7	3
IGLV	IGLV3-10-c	0.00315	0.32%	6	2
IGLV	IGLV1-40-c	0.00275	0.28%	6	2
IGLV	IGLV1-51-b	0.00275	0.28%	6	3
IGHV	IGHV1-69-m	0.0027	0.27%	6	3
IGHV	IGHV3-33-e	0.0027	0.27%	6	3
IGHV	IGHV3-49-c	0.0027	0.27%	6	3
IGHV	IGHV3-66-g	0.0027	0.27%	6	3
IGHV	IGHV4-4-u	0.0027	0.27%	6	3
IGKV	IGKV1D-42-c	0.0027	0.27%	6	1
IGKV	IGKV1D-43-c	0.0027	0.27%	5	3
IGLV	IGLV10-54-h	0.0027	0.27%	6	2
IGLV	IGLV10-54-i	0.0027	0.27%	6	2
IGLV	IGLV1-36-g	0.0027	0.27%	6	3
IGLV	IGLV2-8-d	0.0027	0.27%	6	3
IGLV	IGLV3-22-e	0.0027	0.27%	6	1
IGLV	IGLV3-22-d	0.0027	0.27%	6	3
IGHJ	IGHJ2-b	0.00225	0.23%	4	3
IGHV	IGHV1-3-u	0.00225	0.23%	5	2
IGHV	IGHV1-3-v	0.00225	0.23%	5	2
IGHV	IGHV1-3-ap	0.00225	0.23%	4	3
IGHV	IGHV1-46-c	0.00225	0.23%	5	3

IGHV	IGHV1-58-f	0.00225	0.23%	5	3
IGHV	IGHV1-69-j	0.00225	0.23%	5	3
IGHV	IGHV3-13-e	0.00225	0.23%	5	3
IGHV	IGHV3-23-f	0.00225	0.23%	5	3
IGHV	IGHV3-23-g	0.00225	0.23%	5	3
IGHV	IGHV4-28-p	0.00225	0.23%	5	3
IGHV	IGHV4-31-j	0.00225	0.23%	5	3
IGKV	IGKV1-17-d	0.00225	0.23%	4	1
IGKV	IGKV3-20-a	0.00225	0.23%	5	3
IGKV	IGKV3-7-c	0.00225	0.23%	5	3
IGKV	IGKV3-7-d	0.00225	0.23%	5	3
IGKV	IGKV3D-11-b	0.00225	0.23%	5	3
IGLJ	IGLJ3-a	0.00225	0.23%	4	2
IGLV	IGLV11-55-g	0.00225	0.23%	5	3
IGLV	IGLV1-50-c	0.00225	0.23%	5	3
IGLV	IGLV2-11-g	0.00225	0.23%	5	3
IGLV	IGLV2-14-n	0.00225	0.23%	5	3
IGLV	IGLV2-23-f	0.00225	0.23%	5	2
IGLV	IGLV3-19-c	0.00225	0.23%	5	2
IGLV	IGLV5-37-c	0.00225	0.23%	4	2
IGLV	IGLV5-48-k	0.00225	0.23%	5	3
IGLV	IGLV2-14-i	0.00185	0.19%	4	2
IGHJ	IGHJ1-a	0.0018	0.18%	3	3
IGHV	IGHV1-2-g	0.0018	0.18%	4	3
IGHV	IGHV1-3-y	0.0018	0.18%	4	3
IGHV	IGHV1-3-ar	0.0018	0.18%	4	3
IGHV	IGHV1-3-ai	0.0018	0.18%	4	3
IGHV	IGHV1-69-s	0.0018	0.18%	4	3
IGHV	IGHV1-8-d	0.0018	0.18%	4	2
IGHV	IGHV2-5-d	0.0018	0.18%	4	2
IGHV	IGHV3-13-j	0.0018	0.18%	4	2
IGHV	IGHV3-21-d	0.0018	0.18%	4	3
IGHV	IGHV3-23-n	0.0018	0.18%	4	3
IGHV	IGHV3-33-d	0.0018	0.18%	4	3
IGHV	IGHV3-64-e	0.0018	0.18%	4	2
IGHV	IGHV3-9-g	0.0018	0.18%	4	3
IGHV	IGHV4-39-y	0.0018	0.18%	4	3
IGHV	IGHV4-39-n	0.0018	0.18%	4	3
IGHV	IGHV4-39-o	0.0018	0.18%	4	3
IGHV	IGHV7-81-c	0.0018	0.18%	4	3
IGKJ	IGKJ2-e	0.0018	0.18%	4	3
IGKV	IGKV2D-26-c	0.0018	0.18%	4	1
IGKV	IGKV3D-11-g	0.0018	0.18%	3	2
IGKV	IGKV3D-11-f	0.0018	0.18%	4	3
IGKV	IGKV4-1-w	0.0018	0.18%	3	3
IGLJ	IGLJ1-a	0.0018	0.18%	4	2

IGLV	IGLV10-54-k	0.0018	0.18%	4	2
IGLV	IGLV11-55-e	0.0018	0.18%	4	3
IGLV	IGLV1-36-e	0.0018	0.18%	4	3
IGLV	IGLV1-40-f	0.0018	0.18%	4	3
IGLV	IGLV1-44-d	0.0018	0.18%	4	3
IGLV	IGLV1-44-e	0.0018	0.18%	4	3
IGLV	IGLV1-47-g	0.0018	0.18%	4	2
IGLV	IGLV1-50-d	0.0018	0.18%	4	2
IGLV	IGLV1-51-d	0.0018	0.18%	4	2
IGLV	IGLV1-51-c	0.0018	0.18%	4	3
IGLV	IGLV2-14-r	0.0018	0.18%	4	1
IGLV	IGLV2-33-b	0.0018	0.18%	4	3
IGLV	IGLV2-8-e	0.0018	0.18%	4	2
IGLV	IGLV3-16-h	0.0018	0.18%	4	3
IGLV	IGLV3-21-j	0.0018	0.18%	4	3
IGLV	IGLV3-22-g	0.0018	0.18%	4	3
IGLV	IGLV3-22-h	0.0018	0.18%	4	2
IGLV	IGLV5-48-j	0.0018	0.18%	4	3
IGHJ	IGHJ4-d	0.00135	0.14%	3	3
IGHV	IGHV1-18-d	0.00135	0.14%	3	3
IGHV	IGHV1-18-e	0.00135	0.14%	3	3
IGHV	IGHV1-2-h	0.00135	0.14%	3	2
IGHV	IGHV1-2-f	0.00135	0.14%	3	3
IGHV	IGHV1-3-aa	0.00135	0.14%	3	1
IGHV	IGHV1-3-ak	0.00135	0.14%	3	3
IGHV	IGHV1-3-w	0.00135	0.14%	3	1
IGHV	IGHV1-3-at	0.00135	0.14%	3	2
IGHV	IGHV1-69-k	0.00135	0.14%	3	1
IGHV	IGHV1-69-ac	0.00135	0.14%	3	3
IGHV	IGHV1-8-e	0.00135	0.14%	3	2
IGHV	IGHV1-8-f	0.00135	0.14%	3	2
IGHV	IGHV2-70-y	0.00135	0.14%	3	3
IGHV	IGHV2-70-z	0.00135	0.14%	3	3
IGHV	IGHV2-70-af	0.00135	0.14%	3	2
IGHV	IGHV2-70-h	0.00135	0.14%	3	3
IGHV	IGHV3-13-g	0.00135	0.14%	3	2
IGHV	IGHV3-16-e	0.00135	0.14%	3	2
IGHV	IGHV3-20-f	0.00135	0.14%	3	3
IGHV	IGHV3-30-d	0.00135	0.14%	3	2
IGHV	IGHV3-30-c	0.00135	0.14%	3	3
IGHV	IGHV3-33-h	0.00135	0.14%	3	3
IGHV	IGHV3-43-k	0.00135	0.14%	3	2
IGHV	IGHV3-43-m	0.00135	0.14%	3	3
IGHV	IGHV3-53-n	0.00135	0.14%	3	2
IGHV	IGHV3-53-m	0.00135	0.14%	3	3
IGHV	IGHV3-64-f	0.00135	0.14%	3	1



IGHV	IGHV3-73-c	0.00135	0.14%	3	3
IGHV	IGHV3-9-h	0.00135	0.14%	2	1
IGHV	IGHV4-31-h	0.00135	0.14%	3	2
IGHV	IGHV4-39-u	0.00135	0.14%	3	3
IGHV	IGHV4-39-t	0.00135	0.14%	3	2
IGHV	IGHV4-39-q	0.00135	0.14%	3	3
IGHV	IGHV4-4-ao	0.00135	0.14%	3	3
IGHV	IGHV4-4-ac	0.00135	0.14%	3	1
IGHV	IGHV4-4-ae	0.00135	0.14%	3	2
IGHV	IGHV5-51-g	0.00135	0.14%	3	3
IGHV	IGHV7-81-e	0.00135	0.14%	3	1
IGKJ	IGKJ2-f	0.00135	0.14%	3	2
IGKJ	IGKJ3-d	0.00135	0.14%	3	3
IGKJ	IGKJ4-g	0.00135	0.14%	3	3
IGKJ	IGKJ5-c	0.00135	0.14%	3	3
IGKJ	IGKJ5-b	0.00135	0.14%	3	3
IGKV	IGKV1-5-c	0.00135	0.14%	3	3
IGKV	IGKV1-9-j	0.00135	0.14%	3	3
IGKV	IGKV1D-17-c	0.00135	0.14%	3	2
IGKV	IGKV1D-17-b	0.00135	0.14%	3	2
IGKV	IGKV1D-42-e	0.00135	0.14%	3	1
IGKV	IGKV1D-42-d	0.00135	0.14%	3	2
IGKV	IGKV2D-24-b	0.00135	0.14%	3	2
IGKV	IGKV2D-30-c	0.00135	0.14%	3	2
IGKV	IGKV3-20-d	0.00135	0.14%	3	3
IGKV	IGKV3-20-f	0.00135	0.14%	3	2
IGKV	IGKV3D-11-d	0.00135	0.14%	3	2
IGKV	IGKV4-1-r	0.00135	0.14%	3	3
IGKV	IGKV4-1-x	0.00135	0.14%	3	2
IGKV	IGKV4-1-e	0.00135	0.14%	3	3
IGKV	IGKV4-1-o	0.00135	0.14%	3	2
IGLJ	IGLJ6-c	0.00135	0.14%	3	2
IGLJ	IGLJ7-b	0.00135	0.14%	3	3
IGLV	IGLV10-54-g	0.00135	0.14%	3	1
IGLV	IGLV11-55-h	0.00135	0.14%	3	2
IGLV	IGLV11-55-f	0.00135	0.14%	3	2
IGLV	IGLV1-40-g	0.00135	0.14%	3	3
IGLV	IGLV1-44-f	0.00135	0.14%	3	2
IGLV	IGLV1-47-f	0.00135	0.14%	3	2
IGLV	IGLV2-23-e	0.00135	0.14%	3	3
IGLV	IGLV3-12-d	0.00135	0.14%	3	3
IGLV	IGLV3-16-d	0.00135	0.14%	3	3
IGLV	IGLV3-19-d	0.00135	0.14%	3	1
IGLV	IGLV3-19-l	0.00135	0.14%	3	2
IGLV	IGLV3-1-ag	0.00135	0.14%	3	3
IGLV	IGLV3-1-y	0.00135	0.14%	3	3

IGLV	IGLV3-1-s	0.00135	0.14%	3	3
IGLV	IGLV3-1-x	0.00135	0.14%	3	3
IGLV	IGLV3-1-ad	0.00135	0.14%	3	2
IGLV	IGLV3-1-l	0.00135	0.14%	3	3
IGLV	IGLV3-1-p	0.00135	0.14%	3	2
IGLV	IGLV3-21-t	0.00135	0.14%	3	2
IGLV	IGLV3-22-f	0.00135	0.14%	3	3
IGLV	IGLV3-22-i	0.00135	0.14%	3	1
IGLV	IGLV3-25-e	0.00135	0.14%	3	3
IGLV	IGLV5-48-t	0.00135	0.14%	3	2
IGLV	IGLV5-48-s	0.00135	0.14%	3	3
IGLV	IGLV5-52-d	0.00135	0.14%	3	2
IGLV	IGLV5-52-c	0.00135	0.14%	3	1
IGLV	IGLV7-43-c	0.00135	0.14%	3	2
IGHJ	IGHJ5-d	0.0009	0.09%	2	2
IGHJ	IGHJ6-e	0.0009	0.09%	2	2
IGHJ	IGHJ6-h	0.0009	0.09%	2	2
IGHJ	IGHJ6-f	0.0009	0.09%	2	2
IGHV	IGHV1-18-f	0.0009	0.09%	2	2
IGHV	IGHV1-24-g	0.0009	0.09%	2	1
IGHV	IGHV1-2-i	0.0009	0.09%	2	2
IGHV	IGHV1-2-m	0.0009	0.09%	2	2
IGHV	IGHV1-2-l	0.0009	0.09%	2	2
IGHV	IGHV1-3-ag	0.0009	0.09%	2	2
IGHV	IGHV1-3-aj	0.0009	0.09%	2	2
IGHV	IGHV1-3-aq	0.0009	0.09%	2	1
IGHV	IGHV1-3-s	0.0009	0.09%	2	2
IGHV	IGHV1-3-q	0.0009	0.09%	2	1
IGHV	IGHV1-3-z	0.0009	0.09%	2	1
IGHV	IGHV1-3-ad	0.0009	0.09%	2	2
IGHV	IGHV1-3-aw	0.0009	0.09%	2	2
IGHV	IGHV1-3-ae	0.0009	0.09%	2	2
IGHV	IGHV1-3-an	0.0009	0.09%	2	2
IGHV	IGHV1-46-k	0.0009	0.09%	2	2
IGHV	IGHV1-46-e	0.0009	0.09%	2	2
IGHV	IGHV1-46-n	0.0009	0.09%	2	2
IGHV	IGHV1-58-g	0.0009	0.09%	2	1
IGHV	IGHV1-69-ah	0.0009	0.09%	2	2
IGHV	IGHV1-69-t	0.0009	0.09%	2	2
IGHV	IGHV1-69-ao	0.0009	0.09%	2	1
IGHV	IGHV1-69-af	0.0009	0.09%	2	2
IGHV	IGHV1-69-ad	0.0009	0.09%	2	2
IGHV	IGHV1-69-n	0.0009	0.09%	2	2
IGHV	IGHV1-69-an	0.0009	0.09%	2	2
IGHV	IGHV1-69-o	0.0009	0.09%	2	2
IGHV	IGHV1-8-h	0.0009	0.09%	2	2

IGHV	IGHV2-26-f	0.0009	0.09%	2	2
IGHV	IGHV2-5-c	0.0009	0.09%	2	2
IGHV	IGHV2-70-g	0.0009	0.09%	2	2
IGHV	IGHV2-70-s	0.0009	0.09%	2	2
IGHV	IGHV2-70-ab	0.0009	0.09%	2	2
IGHV	IGHV3-11-b	0.0009	0.09%	2	2
IGHV	IGHV3-13-f	0.0009	0.09%	2	2
IGHV	IGHV3-20-d	0.0009	0.09%	2	2
IGHV	IGHV3-21-c	0.0009	0.09%	2	2
IGHV	IGHV3-21-h	0.0009	0.09%	2	2
IGHV	IGHV3-23-j	0.0009	0.09%	2	1
IGHV	IGHV3-23-h	0.0009	0.09%	2	2
IGHV	IGHV3-33-f	0.0009	0.09%	2	2
IGHV	IGHV3-43-p	0.0009	0.09%	2	2
IGHV	IGHV3-64-h	0.0009	0.09%	2	1
IGHV	IGHV3-66-h	0.0009	0.09%	2	2
IGHV	IGHV3-74-c	0.0009	0.09%	2	2
IGHV	IGHV3-9-o	0.0009	0.09%	2	1
IGHV	IGHV4-28-s	0.0009	0.09%	2	2
IGHV	IGHV4-28-n	0.0009	0.09%	2	2
IGHV	IGHV4-31-t	0.0009	0.09%	2	2
IGHV	IGHV4-31-m	0.0009	0.09%	2	2
IGHV	IGHV4-31-l	0.0009	0.09%	2	2
IGHV	IGHV4-34-d	0.0009	0.09%	2	2
IGHV	IGHV4-34-g	0.0009	0.09%	2	2
IGHV	IGHV4-34-b	0.0009	0.09%	2	2
IGHV	IGHV4-39-aa	0.0009	0.09%	2	2
IGHV	IGHV4-39-s	0.0009	0.09%	2	2
IGHV	IGHV4-39-x	0.0009	0.09%	2	2
IGHV	IGHV4-39-p	0.0009	0.09%	2	2
IGHV	IGHV4-4-aa	0.0009	0.09%	2	2
IGHV	IGHV4-4-af	0.0009	0.09%	2	2
IGHV	IGHV4-4-x	0.0009	0.09%	2	2
IGHV	IGHV4-4-ad	0.0009	0.09%	2	2
IGHV	IGHV4-4-ai	0.0009	0.09%	2	1
IGHV	IGHV5-51-d	0.0009	0.09%	2	2
IGHV	IGHV5-51-k	0.0009	0.09%	2	2
IGHV	IGHV5-51-c	0.0009	0.09%	2	2
IGHV	IGHV6-1-a	0.0009	0.09%	2	2
IGHV	IGHV6-1-c	0.0009	0.09%	2	1
IGHV	IGHV7-81-d	0.0009	0.09%	2	2
IGKJ	IGKJ2-g	0.0009	0.09%	2	2
IGKJ	IGKJ3-b	0.0009	0.09%	2	2
IGKJ	IGKJ4-j	0.0009	0.09%	2	1
IGKJ	IGKJ4-c	0.0009	0.09%	2	2
IGKJ	IGKJ4-k	0.0009	0.09%	2	2

IGKJ	IGKJ4-h	0.0009	0.09%	2	2
IGKV	IGKV1-16-d	0.0009	0.09%	2	2
IGKV	IGKV1-17-e	0.0009	0.09%	2	2
IGKV	IGKV1-5-d	0.0009	0.09%	2	1
IGKV	IGKV1-8-h	0.0009	0.09%	2	2
IGKV	IGKV1-9-i	0.0009	0.09%	2	2
IGKV	IGKV1D-16-c	0.0009	0.09%	2	1
IGKV	IGKV1D-16-b	0.0009	0.09%	2	1
IGKV	IGKV1D-42-g	0.0009	0.09%	2	2
IGKV	IGKV1D-42-f	0.0009	0.09%	2	1
IGKV	IGKV1D-43-e	0.0009	0.09%	2	1
IGKV	IGKV1D-43-d	0.0009	0.09%	2	2
IGKV	IGKV1D-8-d	0.0009	0.09%	2	1
IGKV	IGKV1D-8-f	0.0009	0.09%	2	2
IGKV	IGKV2D-26-d	0.0009	0.09%	2	2
IGKV	IGKV2D-26-f	0.0009	0.09%	2	2
IGKV	IGKV2D-29-b	0.0009	0.09%	2	1
IGKV	IGKV3-11-g	0.0009	0.09%	2	1
IGKV	IGKV3-11-f	0.0009	0.09%	2	2
IGKV	IGKV3-20-j	0.0009	0.09%	2	2
IGKV	IGKV3-20-i	0.0009	0.09%	2	2
IGKV	IGKV3-20-e	0.0009	0.09%	2	2
IGKV	IGKV3-20-c	0.0009	0.09%	2	2
IGKV	IGKV3D-11-h	0.0009	0.09%	2	1
IGKV	IGKV3D-11-i	0.0009	0.09%	2	1
IGKV	IGKV3D-20-d	0.0009	0.09%	2	2
IGKV	IGKV4-1-y	0.0009	0.09%	2	2
IGKV	IGKV4-1-g	0.0009	0.09%	2	2
IGKV	IGKV4-1-t	0.0009	0.09%	2	2
IGKV	IGKV4-1-i	0.0009	0.09%	2	2
IGKV	IGKV4-1-n	0.0009	0.09%	2	2
IGKV	IGKV4-1-l	0.0009	0.09%	2	2
IGLJ	IGLJ2-e	0.0009	0.09%	2	2
IGLJ	IGLJ2-c	0.0009	0.09%	2	2
IGLJ	IGLJ5-b	0.0009	0.09%	2	1
IGLV	IGLV10-54-n	0.0009	0.09%	2	2
IGLV	IGLV10-54-j	0.0009	0.09%	2	2
IGLV	IGLV1-40-h	0.0009	0.09%	2	2
IGLV	IGLV1-40-e	0.0009	0.09%	2	2
IGLV	IGLV1-47-e	0.0009	0.09%	2	2
IGLV	IGLV1-51-g	0.0009	0.09%	2	2
IGLV	IGLV2-11-j	0.0009	0.09%	2	2
IGLV	IGLV2-11-f	0.0009	0.09%	2	2
IGLV	IGLV2-11-i	0.0009	0.09%	2	2
IGLV	IGLV2-11-e	0.0009	0.09%	2	1
IGLV	IGLV2-14-p	0.0009	0.09%	2	2

IGLV	IGLV2-14-q	0.0009	0.09%	2	1
IGLV	IGLV2-14-o	0.0009	0.09%	2	1
IGLV	IGLV2-23-k	0.0009	0.09%	2	2
IGLV	IGLV2-23-o	0.0009	0.09%	2	2
IGLV	IGLV2-33-c	0.0009	0.09%	2	1
IGLV	IGLV3-19-j	0.0009	0.09%	2	2
IGLV	IGLV3-19-k	0.0009	0.09%	2	2
IGLV	IGLV3-19-h	0.0009	0.09%	2	1
IGLV	IGLV3-19-i	0.0009	0.09%	2	1
IGLV	IGLV3-19-p	0.0009	0.09%	2	2
IGLV	IGLV3-1-ah	0.0009	0.09%	2	2
IGLV	IGLV3-1-d	0.0009	0.09%	2	2
IGLV	IGLV3-1-j	0.0009	0.09%	2	1
IGLV	IGLV3-1-g	0.0009	0.09%	2	2
IGLV	IGLV3-1-z	0.0009	0.09%	2	2
IGLV	IGLV3-1-ac	0.0009	0.09%	2	2
IGLV	IGLV3-1-ae	0.0009	0.09%	2	2
IGLV	IGLV3-21-k	0.0009	0.09%	2	2
IGLV	IGLV3-21-r	0.0009	0.09%	2	1
IGLV	IGLV3-21-i	0.0009	0.09%	2	2
IGLV	IGLV3-22-j	0.0009	0.09%	2	1
IGLV	IGLV3-22-n	0.0009	0.09%	2	2
IGLV	IGLV3-22-m	0.0009	0.09%	2	2
IGLV	IGLV3-22-o	0.0009	0.09%	2	2
IGLV	IGLV3-25-g	0.0009	0.09%	2	1
IGLV	IGLV3-25-l	0.0009	0.09%	2	2
IGLV	IGLV3-27-a	0.0009	0.09%	2	2
IGLV	IGLV3-32-c	0.0009	0.09%	2	2
IGLV	IGLV3-9-a	0.0009	0.09%	2	2
IGLV	IGLV3-9-b	0.0009	0.09%	2	2
IGLV	IGLV4-3-d	0.0009	0.09%	2	2
IGLV	IGLV4-60-g	0.0009	0.09%	2	2
IGLV	IGLV4-60-n	0.0009	0.09%	2	2
IGLV	IGLV4-60-o	0.0009	0.09%	2	2
IGLV	IGLV5-45-g	0.0009	0.09%	2	2
IGLV	IGLV5-48-u	0.0009	0.09%	2	2
IGLV	IGLV5-48-q	0.0009	0.09%	2	2
IGLV	IGLV5-48-r	0.0009	0.09%	2	2
IGLV	IGLV5-48-n	0.0009	0.09%	2	2
IGLV	IGLV5-48-m	0.0009	0.09%	2	1
IGLV	IGLV5-48-l	0.0009	0.09%	2	2
IGLV	IGLV5-52-b	0.0009	0.09%	2	1
IGLV	IGLV7-46-c	0.0009	0.09%	2	2
IGLV	IGLV9-49-c	0.0009	0.09%	2	2

**Table 20B: Variants Found In No More Than 3 Human Populations & In At Least 10 Individuals**

<b>TYPE</b>	<b>VARIANT</b>	<b>C.FREQ</b>	<b>C.FREQ (%)</b>	<b>#INDIVIDUALS</b>	<b>#POPULATIONS</b>
IGKV	IGKV1D-8-b	0.01785	1.79%	36	3
IGHV	IGHV3-11-a	0.01695	1.70%	36	3
IGHV	IGHV5-51-a	0.0164	1.64%	35	3
IGHV	IGHV2-5-b	0.01055	1.06%	18	3
IGLV	IGLV2-14-g	0.0104	1.04%	22	3
IGHV	IGHV7-81-b	0.01005	1.01%	21	3
IGKV	IGKV1D-42-b	0.01005	1.01%	21	3
IGKV	IGKV3-7-b	0.01	1.00%	19	3
IGHV	IGHV1-69-d	0.00995	1.00%	22	3
IGLV	IGLV2-14-e	0.00995	1.00%	19	3
IGLV	IGLV9-49-a	0.00865	0.87%	18	3
IGLV	IGLV1-44-b	0.00775	0.78%	16	2
IGLV	IGLV2-11-a	0.00775	0.78%	17	3
IGKV	IGKV3-11-d	0.0073	0.73%	14	3
IGKV	IGKV1D-43-b	0.00685	0.69%	14	3
IGHV	IGHV3-38-d	0.0064	0.64%	13	3
IGLJ	IGLJ6-a	0.00595	0.60%	13	3
IGHV	IGHV5-51-b	0.0055	0.55%	11	3
IGLV	IGLV5-45-c	0.00545	0.55%	12	2
IGHV	IGHV4-31-i	0.0054	0.54%	10	3
IGLV	IGLV1-36-c	0.00455	0.46%	10	2

**Table 20C: Variants Found In No More Than 3 Human Populations & In At Least 20 Individuals**

<b>TYPE</b>	<b>VARIANT</b>	<b>C.FREQ</b>	<b>C.FREQ (%)</b>	<b>#INDIVIDUALS</b>	<b>#POPULATIONS</b>
IGKV	IGKV1D-8-b	0.01785	1.79%	36	3
IGHV	IGHV3-11-a	0.01695	1.70%	36	3
IGHV	IGHV5-51-a	0.0164	1.64%	35	3
IGLV	IGLV2-14-g	0.0104	1.04%	22	3
IGHV	IGHV7-81-b	0.01005	1.01%	21	3
IGKV	IGKV1D-42-b	0.01005	1.01%	21	3
IGHV	IGHV1-69-d	0.00995	1.00%	22	3

**Table 20D: Variants Found In No More Than 3 Human Populations & In At Least 30 Individuals**

<b>TYPE</b>	<b>VARIANT</b>	<b>C.FREQ</b>	<b>C.FREQ (%)</b>	<b>#INDIVIDUALS</b>	<b>#POPULATIONS</b>
IGKV	IGKV1D-8-b	0.01785	1.79%	36	3
IGHV	IGHV3-11-a	0.01695	1.70%	36	3
IGHV	IGHV5-51-a	0.0164	1.64%	35	3



**Table 21**

<u>Seq ID #</u>	<u>Associated Gene Name</u>	<u>Ensembl Gene ID</u>	<u>Gene Location</u> <u>[Chromosome Name:Gene Chr Start (bp)-GeneChr End (bp)]</u>	<u>Ensembl Transcript ID</u>	<u>Exon Positions</u> <u>[Chromosome Name:Exon Chr Start (bp)-Exon Chr End (bp)]</u>
<u>239</u>	<u>IGHJ1</u>	<u>ENSG00000211905</u>	<u>14:106331617-106331668</u>	<u>ENST00000390565</u>	<u>14:106331617-106331668</u>
<u>240</u>	<u>IGHJ2</u>	<u>ENSG00000211904</u>	<u>14:106331409-106331460</u>	<u>ENST00000390564</u>	<u>14:106331409-106331460</u>
<u>241</u>	<u>IGHJ3</u>	<u>ENSG00000242887</u>	<u>14:106330797-106330845</u>	<u>ENST00000463911</u>	<u>14:106330797-106330845</u>
<u>242</u>	<u>IGHJ4</u>	<u>ENSG00000240041</u>	<u>14:106330425-106330470</u>	<u>ENST00000461719</u>	<u>14:106330425-106330470</u>
<u>243</u>	<u>IGHJ5</u>	<u>ENSG00000242472</u>	<u>14:106330024-106330072</u>	<u>ENST00000488476</u>	<u>14:106330024-106330072</u>
<u>244</u>	<u>IGHJ6</u>	<u>ENSG00000211900</u>	<u>14:106329408-106329468</u>	<u>ENST00000390560</u>	<u>14:106329408-106329468</u>
<u>245</u>	<u>IGHV1-18</u>	<u>ENSG00000211945</u>	<u>14:106641563-106642056</u>	<u>ENST00000390605</u>	<u>14:106641952-106642056 14:106641563-106641867</u>
<u>246</u>	<u>IGHV1-2</u>	<u>ENSG00000211934</u>	<u>14:106452671-106453170</u>	<u>ENST00000390594</u>	<u>14:106453061-106453170 14:106452671-106452975</u>
<u>247</u>	<u>IGHV1-24</u>	<u>ENSG00000211950</u>	<u>14:106733144-106733639</u>	<u>ENST00000390610</u>	<u>14:106733534-106733639 14:106733144-106733448</u>
<u>248</u>	<u>IGHV1-3</u>	<u>ENSG00000211935</u>	<u>14:106471246-106471723</u>	<u>ENST00000390595</u>	<u>14:106471636-106471723 14:106471246-106471550</u>
<u>249</u>	<u>IGHV1-45</u>	<u>ENSG00000211961</u>	<u>14:106962931-106963424</u>	<u>ENST00000390621</u>	<u>14:106963321-106963424 14:106962931-106963235</u>
<u>250</u>	<u>IGHV1-46</u>	<u>ENSG00000211962</u>	<u>14:106967049-106967788</u>	<u>ENST00000390622</u>	<u>14:106967439-106967788 14:106967049-106967353</u>
<u>251</u>	<u>IGHV1-58</u>	<u>ENSG00000211968</u>	<u>14:107078373-107078869</u>	<u>ENST00000390628</u>	<u>14:107078763-107078869 14:107078373-107078677</u>



<u>252</u>	<u>IGHV1-69</u>	<u>ENSG00000211973</u>	<u>14:107169931-107170428</u>	<u>ENST00000390633</u>	<u>14:107170322-107170428 14:107169931-107170235</u>
<u>253</u>	<u>IGHV1-8</u>	<u>ENSG00000211939</u>	<u>14:106539079-106539577</u>	<u>ENST00000390599</u>	<u>14:106539470-106539577 14:106539079-106539383</u>
<u>254</u>	<u>IGHV2-26</u>	<u>ENSG00000211951</u>	<u>14:106757650-106758116</u>	<u>ENST00000390611</u>	<u>14:106758047-106758116 14:106757650-106757960</u>
<u>255</u>	<u>IGHV2-5</u>	<u>ENSG00000211937</u>	<u>14:106494135-106494597</u>	<u>ENST00000390597</u>	<u>14:106494532-106494597 14:106494135-106494445</u>
<u>256</u>	<u>IGHV2-70</u>	<u>ENSG00000211974</u>	<u>14:107178820-107179338</u>	<u>ENST00000390634</u>	<u>14:107179217-107179338 14:107178820-107179130</u>
<u>257</u>	<u>IGHV3-11</u>	<u>ENSG00000211941</u>	<u>14:106573233-106573800</u>	<u>ENST00000390601</u>	<u>14:106573635-106573800 14:106573233-106573537</u>
<u>258</u>	<u>IGHV3-13</u>	<u>ENSG00000211942</u>	<u>14:106586137-106586667</u>	<u>ENST00000390602</u>	<u>14:106586542-106586667 14:106586137-106586438</u>
<u>259</u>	<u>IGHV3-15</u>	<u>ENSG00000211943</u>	<u>14:1066610313-1066610852</u>	<u>ENST00000390603</u>	<u>14:1066610727-1066610852 14:1066610313-1066610623</u>
<u>260</u>	<u>IGHV3-16</u>	<u>ENSG00000211944</u>	<u>14:106621894-106622419</u>	<u>ENST00000390604</u>	<u>14:106622300-106622419 14:106621894-106622198</u>
<u>261</u>	<u>IGHV3-20</u>	<u>ENSG00000211946</u>	<u>14:106667581-106668095</u>	<u>ENST00000390606</u>	<u>14:106667988-106668095 14:106667581-1066667885</u>
<u>262</u>	<u>IGHV3-21</u>	<u>ENSG00000211947</u>	<u>14:1066691673-106692203</u>	<u>ENST00000390607</u>	<u>14:106692079-106692203 14:106691673-106691977</u>
<u>263</u>	<u>IGHV3-23</u>	<u>ENSG00000211949</u>	<u>14:106725201-106725733</u>	<u>ENST00000390609</u>	<u>14:106725609-106725733 14:106725201-106725505</u>
<u>264</u>	<u>IGHV3-30</u>	<u>ENSG00000211953</u>	<u>14:106791005-106791536</u>	<u>ENST00000390613</u>	<u>14:106791411-106791536 14:106791005-106791309</u>
<u>265</u>	<u>IGHV3-33</u>	<u>ENSG00000211955</u>	<u>14:106815722-106816253</u>	<u>ENST00000390615</u>	<u>14:106816128-106816253 14:106815722-106816026</u>
<u>266</u>	<u>IGHV3-35</u>	<u>ENSG00000211957</u>	<u>14:106845323-106845789</u>	<u>ENST00000390617</u>	<u>14:106845729-106845789 14:106845323-106845627</u>
<u>267</u>	<u>IGHV3-38</u>	<u>ENSG00000211958</u>	<u>14:106866406-106866934</u>	<u>ENST00000390618</u>	<u>14:106866811-106866934 14:106866406-106866707</u>
<u>268</u>	<u>IGHV3-43</u>	<u>ENSG00000232216</u>	<u>14:106926188-106926724</u>	<u>ENST00000434710</u>	<u>14:106926599-106926724 14:106926188-106926495</u>

<u>269</u>	<u>IGHV3-48</u>	<u>ENSG00000211964</u>	<u>14:106993814-106994346</u>	<u>ENST00000390624</u>	<u>14:106994222-106994346 14:106993814-106994118</u>
<u>270</u>	<u>IGHV3-49</u>	<u>ENSG00000211965</u>	<u>14:107012938-107013477</u>	<u>ENST00000390625</u>	<u>14:107013352-107013477 14:107012938-107013248</u>
<u>271</u>	<u>IGHV3-53</u>	<u>ENSG00000211967</u>	<u>14:107048672-107049341</u>	<u>ENST00000390627</u>	<u>14:107049075-107049341 14:107048672-107048973</u>
<u>272</u>	<u>IGHV3-64</u>	<u>ENSG00000223648</u>	<u>14:107113741-107114274</u>	<u>ENST00000454421</u>	<u>14:107114149-107114274 14:107113741-107114045</u>
<u>273</u>	<u>IGHV3-66</u>	<u>ENSG00000211972</u>	<u>14:107131033-107131560</u>	<u>ENST00000390632</u>	<u>14:107131436-107131560 14:107131033-107131334</u>
<u>274</u>	<u>IGHV3-7</u>	<u>ENSG00000211938</u>	<u>14:106518400-106518932</u>	<u>ENST00000390598</u>	<u>14:106518808-106518932 14:106518400-106518704</u>
<u>275</u>	<u>IGHV3-72</u>	<u>ENSG00000225698</u>	<u>14:107198932-107199471</u>	<u>ENST00000433072</u>	<u>14:107199346-107199471 14:107198932-107199242</u>
<u>276</u>	<u>IGHV3-73</u>	<u>ENSG00000211976</u>	<u>14:107210932-107211471</u>	<u>ENST00000390636</u>	<u>14:107211346-107211471 14:107210932-107211242</u>
<u>277</u>	<u>IGHV3-74</u>	<u>ENSG00000224650</u>	<u>14:107218676-107219365</u>	<u>ENST00000424969</u>	<u>14:107219084-107219365 14:107218676-107218980</u>
<u>278</u>	<u>IGHV3-9</u>	<u>ENSG00000211940</u>	<u>14:106552285-106552809</u>	<u>ENST00000390600</u>	<u>14:106552684-106552809 14:106552285-106552592</u>
<u>279</u>	<u>IGHV4-28</u>	<u>ENSG00000211952</u>	<u>14:106780513-106781017</u>	<u>ENST00000390612</u>	<u>14:106780900-106781017 14:106780513-106780817</u>
<u>280</u>	<u>IGHV4-31</u>	<u>ENSG00000231475</u>	<u>14:106805209-106805716</u>	<u>ENST00000438142</u>	<u>14:106805599-106805716 14:106805209-106805516</u>
<u>281</u>	<u>IGHV4-34</u>	<u>ENSG00000211956</u>	<u>14:106829594-106830076</u>	<u>ENST00000390616</u>	<u>14:106829979-106830076 14:106829594-106829895</u>
<u>282</u>	<u>IGHV4-39</u>	<u>ENSG00000211959</u>	<u>14:106877619-106878126</u>	<u>ENST00000390619</u>	<u>14:106878010-106878126 14:106877619-106877926</u>
<u>283</u>	<u>IGHV4-4</u>	<u>ENSG00000211936</u>	<u>14:106478110-106478603</u>	<u>ENST00000390596</u>	<u>14:106478494-106478603 14:106478110-106478411</u>
<u>284</u>	<u>IGHV4-59</u>	<u>ENSG00000224373</u>	<u>14:107081806-107083830</u>	<u>ENST00000390629</u>	<u>14:107083640-107083830 14:107083256-107083557</u>
<u>284</u>	<u>IGHV4-59</u>	<u>ENSG00000224373</u>	<u>14:107081806-107083830</u>	<u>ENST00000455737</u>	<u>14:107083640-107083725 14:107083240-107083557</u>

<u>285</u>	<u>IGHV4-61</u>	<u>ENSG00000211970</u>	<u>14:107095126-107095662</u>	<u>ENST00000390630</u>	<u>14:107081806-107082728</u>
<u>286</u>	<u>IGHV5-51</u>	<u>ENSG00000211966</u>	<u>14:107034729-107035221</u>	<u>ENST00000390626</u>	<u>14:107095516-107095662 14:107095126-107095433</u>
<u>287</u>	<u>IGHV6-1</u>	<u>ENSG00000211933</u>	<u>14:106405611-106406108</u>	<u>ENST00000390593</u>	<u>14:107035117-107035221 14:107034729-107035033</u>
<u>288</u>	<u>IGHV7-81</u>	<u>ENSG00000211979</u>	<u>14:107282792-107283280</u>	<u>ENST00000390639</u>	<u>14:106406008-106406108 14:106405611-106405924</u>
<u>289</u>	<u>IGKJ2</u>	<u>ENSG00000211596</u>	<u>2:89161037-89161074</u>	<u>ENST00000390241</u>	<u>14:107283181-107283280 14:107282792-107283096</u>
<u>290</u>	<u>IGKI3</u>	<u>ENSG00000211595</u>	<u>2:89160733-89160770</u>	<u>ENST00000390240</u>	<u>2:89161037-89161074</u>
<u>291</u>	<u>IGKI4</u>	<u>ENSG00000211594</u>	<u>2:89160398-89160434</u>	<u>ENST00000390239</u>	<u>2:89160733-89160770</u>
<u>292</u>	<u>IGKI5</u>	<u>ENSG00000211593</u>	<u>2:89160080-89160117</u>	<u>ENST00000390238</u>	<u>2:89160398-89160434</u>
<u>293</u>	<u>IGKV1-16</u>	<u>ENSG00000240864</u>	<u>2:89399352-89399854</u>	<u>ENST00000479981</u>	<u>2:89160080-89160117</u>
<u>294</u>	<u>IGKV1-17</u>	<u>ENSG00000240382</u>	<u>2:89416833-89417335</u>	<u>ENST00000490686</u>	<u>2:89399773-89399854 2:89399352-89399647</u>
<u>295</u>	<u>IGKV1-5</u>	<u>ENSG00000243466</u>	<u>2:89246819-89247475</u>	<u>ENST00000496168</u>	<u>2:89417254-89417335 2:89416833-89417128</u>
<u>296</u>	<u>IGKV1-6</u>	<u>ENSG00000239855</u>	<u>2:89265781-89266286</u>	<u>ENST00000464162</u>	<u>2:89247240-89247475 2:89246819-89247114</u>
<u>297</u>	<u>IGKV1-8</u>	<u>ENSG00000240671</u>	<u>2:89291928-89292450</u>	<u>ENST00000495489</u>	<u>2:89266203-89266286 2:89265781-89266076</u>
<u>298</u>	<u>IGKV1-9</u>	<u>ENSG00000241755</u>	<u>2:89309479-89310012</u>	<u>ENST00000493819</u>	<u>2:89292349-89292450 2:89291928-89292223</u>
<u>299</u>	<u>IGKV1D-12</u>	<u>ENSG00000240834</u>	<u>2:90198535-90199190</u>	<u>ENST00000390276</u>	<u>2:89309900-89310012 2:89309479-89309774</u>
<u>300</u>	<u>IGKV1D-16</u>	<u>ENSG00000241244</u>	<u>2:90139078-90139580</u>	<u>ENST00000492446</u>	<u>2:90198535-90198770 2:90198895-90199190</u>

<u>301</u>	<u>IGKV1D-17</u>	<u>ENSG00000242766</u>	<u>2:90121477-90122133</u>	<u>ENST00000483379</u>	<u>2:90121477-90121712 2:90121838-90122133</u>
<u>302</u>	<u>IGKV1D-42</u>	<u>ENSG00000211633</u>	<u>2:90229045-90229531</u>	<u>ENST00000390278</u>	<u>2:90229045-90229119 2:90229236-90229531</u>
<u>303</u>	<u>IGKV1D-43</u>	<u>ENSG00000242580</u>	<u>2:90248739-90249395</u>	<u>ENST00000468879</u>	<u>2:90248739-90248974 2:90249100-90249395</u>
<u>304</u>	<u>IGKV1D-8</u>	<u>ENSG00000239819</u>	<u>2:90259593-90260248</u>	<u>ENST00000471857</u>	<u>2:90259593-90259828 2:90259953-90260248</u>
<u>305</u>	<u>IGKV2-24</u>	<u>ENSG00000241294</u>	<u>2:89475812-89476644</u>	<u>ENST00000484817</u>	<u>2:89476566-89476644 2:89475812-89476122</u>
<u>306</u>	<u>IGKV2-30</u>	<u>ENSG00000243238</u>	<u>2:89544264-89545079</u>	<u>ENST00000468494</u>	<u>2:89545001-89545079 2:89544264-89544574</u>
<u>307</u>	<u>IGKV2D-24</u>	<u>ENSG00000241566</u>	<u>2:90043607-90044439</u>	<u>ENST00000462693</u>	<u>2:90043607-90043685 2:90044129-90044439</u>
<u>308</u>	<u>IGKV2D-26</u>	<u>ENSG00000211623</u>	<u>2:90024732-90025512</u>	<u>ENST00000390268</u>	<u>2:90024732-90024810 2:90025202-90025512</u>
<u>309</u>	<u>IGKV2D-29</u>	<u>ENSG00000243264</u>	<u>2:89986322-89987079</u>	<u>ENST00000491977</u>	<u>2:89986322-89986400 2:89986769-89987079</u>
<u>310</u>	<u>IGKV2D-30</u>	<u>ENSG00000239571</u>	<u>2:89975669-89976489</u>	<u>ENST00000474213</u>	<u>2:89975669-89975752 2:89976179-89976489</u>
<u>311</u>	<u>IGKV3-11</u>	<u>ENSG00000241351</u>	<u>2:89326668-89327228</u>	<u>ENST00000483158</u>	<u>2:89327133-89327228 2:89326668-89326963</u>
<u>312</u>	<u>IGKV3-20</u>	<u>ENSG00000239951</u>	<u>2:89442057-89442643</u>	<u>ENST00000492167</u>	<u>2:89442543-89442643 2:89442057-89442355</u>
<u>313</u>	<u>IGKV3-7</u>	<u>ENSG00000243063</u>	<u>2:89277987-89278600</u>	<u>ENST00000390247</u>	<u>2:89278455-89278600 2:89277987-89278285</u>
<u>314</u>	<u>IGKV3D-11</u>	<u>ENSG00000211632</u>	<u>2:90211643-90212253</u>	<u>ENST00000390277</u>	<u>2:90211643-90211788 2:90211958-90212253</u>
<u>315</u>	<u>IGKV3D-15</u>	<u>ENSG00000224041</u>	<u>2:90153696-90154258</u>	<u>ENST00000417279</u>	<u>2:90153696-90153793 2:90153963-90154258</u>
<u>316</u>	<u>IGKV3D-20</u>	<u>ENSG00000211625</u>	<u>2:90077680-90078311</u>	<u>ENST00000390270</u>	<u>2:90077680-90077825 2:90078013-90078311</u>
<u>317</u>	<u>IGKV4-1</u>	<u>ENSG00000211598</u>	<u>2:89184913-89185669</u>	<u>ENST00000390243</u>	<u>2:89184913-89185136 2:89185356-89185669</u>

<u>318</u>	<u>IGKV5-2</u>	<u>ENSG00000211599</u>	<u>2:89196748-89197300</u>	<u>ENST00000390244</u>	<u>2:89196748-89196859 2:89197005-89197300</u>
<u>319</u>	<u>IGKV6-21</u>	<u>ENSG00000211611</u>	<u>2:89459235-89459850</u>	<u>ENST00000390256</u>	<u>2:89459741-89459850 2:89459235-89459530</u>
<u>320</u>	<u>IGKV6D-21</u>	<u>ENSG00000225523</u>	<u>2:90060377-90060995</u>	<u>ENST00000436451</u>	<u>2:90060377-90060489 2:90060700-90060995</u>
<u>321</u>	<u>IGKV6D-41</u>	<u>ENSG00000211626</u>	<u>2:90108504-90109080</u>	<u>ENST00000390271</u>	<u>2:90108504-90108578 2:90108785-90109080</u>
<u>322</u>	<u>IGLJ1</u>	<u>ENSG00000211674</u>	<u>22:23235872-23235998</u>	<u>ENST00000390320</u>	<u>22:23235872-23235998</u>
<u>323</u>	<u>IGLJ2</u>	<u>ENSG00000211676</u>	<u>22:23241661-23241835</u>	<u>ENST00000390322</u>	<u>22:23241661-23241835</u>
<u>324</u>	<u>IGLJ3</u>	<u>ENSG00000211678</u>	<u>22:23247030-23247205</u>	<u>ENST00000390324</u>	<u>22:23247030-23247205</u>
<u>325</u>	<u>IGLJ5</u>	<u>ENSG00000211681</u>	<u>22:23256408-23256479</u>	<u>ENST00000390327</u>	<u>22:23256408-23256479</u>
<u>326</u>	<u>IGLJ6</u>	<u>ENSG00000211682</u>	<u>22:23260304-23260373</u>	<u>ENST00000390328</u>	<u>22:23260304-23260373</u>
<u>327</u>	<u>IGLJ7</u>	<u>ENSG00000211684</u>	<u>22:23263562-23263607</u>	<u>ENST00000390330</u>	<u>22:23263562-23263607</u>
<u>328</u>	<u>IGLV10-54</u>	<u>ENSG00000211642</u>	<u>22:22569184-22569660</u>	<u>ENST00000390287</u>	<u>22:22569184-22569242 22:22569355-22569660</u>
<u>329</u>	<u>IGLV11-55</u>	<u>ENSG00000211641</u>	<u>22:22556057-22556600</u>	<u>ENST00000390286</u>	<u>22:22556057-22556114 22:22556233-22556600</u>
<u>330</u>	<u>IGLV1-36</u>	<u>ENSG00000211655</u>	<u>22:22786296-22786802</u>	<u>ENST00000390301</u>	<u>22:22786296-22786381 22:22786497-22786802</u>
<u>331</u>	<u>IGLV1-40</u>	<u>ENSG00000211653</u>	<u>22:22764098-22764614</u>	<u>ENST00000390299</u>	<u>22:22764098-22764194 22:22764305-22764614</u>
<u>332</u>	<u>IGLV1-44</u>	<u>ENSG00000211651</u>	<u>22:22735135-22735715</u>	<u>ENST00000390297</u>	<u>22:22735135-22735294 22:22735410-22735715</u>
<u>333</u>	<u>IGLV1-47</u>	<u>ENSG00000211648</u>	<u>22:22712087-22712608</u>	<u>ENST00000390294</u>	<u>22:22712087-22712188 22:22712304-22712608</u>
<u>334</u>	<u>IGLV1-50</u>	<u>ENSG00000211645</u>	<u>22:22681658-22682172</u>	<u>ENST00000390291</u>	<u>22:22681658-22681754 22:22681865-22682172</u>

<u>335</u>	<u>IGLV1-51</u>	<u>ENSG00000211644</u>	<u>22:22676828-22677336</u>	<u>ENST00000390290</u>	<u>22:22676828-22676909 22:22677019-22677336</u>
<u>336</u>	<u>IGLV2-11</u>	<u>ENSG00000211668</u>	<u>22:23134980-23135496</u>	<u>ENST00000390314</u>	<u>22:23134980-23135067 22:23135185-23135496</u>
<u>337</u>	<u>IGLV2-14</u>	<u>ENSG00000211666</u>	<u>22:23101189-23101707</u>	<u>ENST00000390312</u>	<u>22:23101189-23101275 22:23101393-23101707</u>
<u>338</u>	<u>IGLV2-18</u>	<u>ENSG00000211664</u>	<u>22:23077095-23077584</u>	<u>ENST00000390310</u>	<u>22:23077095-23077153 22:23077270-23077584</u>
<u>339</u>	<u>IGLV2-23</u>	<u>ENSG00000211660</u>	<u>22:23040274-23040892</u>	<u>ENST00000390306</u>	<u>22:23040274-23040481 22:23040599-23040892</u>
<u>340</u>	<u>IGLV2-33</u>	<u>ENSG00000211656</u>	<u>22:22930626-22931145</u>	<u>ENST00000390302</u>	<u>22:22930626-22930729 22:22930852-22931145</u>
<u>341</u>	<u>IGLV2-8</u>	<u>ENSG00000211671</u>	<u>22:23165153-23165787</u>	<u>ENST00000390317</u>	<u>22:23165153-23165360 22:23165476-23165787</u>
<u>342</u>	<u>IGLV3-1</u>	<u>ENSG00000211673</u>	<u>22:23222886-23223576</u>	<u>ENST00000390319</u>	<u>22:23222886-23222983 22:23223277-23223576</u>
<u>343</u>	<u>IGLV3-10</u>	<u>ENSG00000211669</u>	<u>22:23154244-23154782</u>	<u>ENST00000390315</u>	<u>22:23154244-23154324 22:23154478-23154782</u>
<u>344</u>	<u>IGLV3-12</u>	<u>ENSG00000211667</u>	<u>22:23114317-23115079</u>	<u>ENST00000390313</u>	<u>22:23114317-23114371 22:23114775-23115079</u>
<u>345</u>	<u>IGLV3-16</u>	<u>ENSG00000211665</u>	<u>22:23089870-23090398</u>	<u>ENST00000390311</u>	<u>22:23089870-23089953 22:23090108-23090398</u>
<u>346</u>	<u>IGLV3-19</u>	<u>ENSG00000211663</u>	<u>22:23063108-23063630</u>	<u>ENST00000390309</u>	<u>22:23063108-23063193 22:23063340-23063630</u>
<u>347</u>	<u>IGLV3-21</u>	<u>ENSG00000211662</u>	<u>22:23054174-23055688</u>	<u>ENST00000390308</u>	<u>22:23054174-23054290 22:23054770-23054949</u> <u>22:23055384-23055688</u>
<u>348</u>	<u>IGLV3-22</u>	<u>ENSG00000211661</u>	<u>22:23046750-23047307</u>	<u>ENST00000390307</u>	<u>22:23046750-23046859 22:23047009-23047307</u>
<u>349</u>	<u>IGLV3-25</u>	<u>ENSG00000211659</u>	<u>22:23029190-23029735</u>	<u>ENST00000390305</u>	<u>22:23029190-23029278 22:23029445-23029735</u>
<u>350</u>	<u>IGLV3-27</u>	<u>ENSG00000211658</u>	<u>22:23010758-23011276</u>	<u>ENST00000390304</u>	<u>22:23010758-23010841 22:23010984-23011276</u>

<u>351</u>	<u>IGLV3-32</u>	<u>ENSG00000211657</u>	<u>22:22936998-22937501</u>	<u>ENST00000390303</u>	<u>22:22936998-22937079 22:22937226-22937501</u>
<u>352</u>	<u>IGLV3-9</u>	<u>ENSG00000211670</u>	<u>22:23161507-23162253</u>	<u>ENST00000390316</u>	<u>22:23161507-23161667 22:23161954-23162253</u>
<u>353</u>	<u>IGLV4-3</u>	<u>ENSG00000211672</u>	<u>22:23213686-23214214</u>	<u>ENST00000390318</u>	<u>22:23213686-23213763 22:23213890-23214214</u>
<u>354</u>	<u>IGLV4-60</u>	<u>ENSG00000211639</u>	<u>22:22516592-22517074</u>	<u>ENST00000390284</u>	<u>22:22516592-22516643 22:22516765-22517074</u>
<u>355</u>	<u>IGLV4-69</u>	<u>ENSG00000211637</u>	<u>22:22385332-22385870</u>	<u>ENST00000390282</u>	<u>22:22385332-22385440 22:22385561-22385870</u>
<u>356</u>	<u>IGLV5-37</u>	<u>ENSG00000211654</u>	<u>22:22781876-22782371</u>	<u>ENST00000390300</u>	<u>22:22781876-22781926 22:22782049-22782371</u>
<u>357</u>	<u>IGLV5-45</u>	<u>ENSG00000211650</u>	<u>22:22730355-22730874</u>	<u>ENST00000390296</u>	<u>22:22730355-22730428 22:22730552-22730874</u>
<u>358</u>	<u>IGLV5-48</u>	<u>ENSG00000211647</u>	<u>22:22707289-22707781</u>	<u>ENST00000390293</u>	<u>22:22707289-22707334 22:22707459-22707781</u>
<u>359</u>	<u>IGLV5-52</u>	<u>ENSG00000211643</u>	<u>22:22673082-22673581</u>	<u>ENST00000390289</u>	<u>22:22673082-22673132 22:22673254-22673581</u>
<u>360</u>	<u>IGLV6-57</u>	<u>ENSG00000211640</u>	<u>22:22550113-22550686</u>	<u>ENST00000390285</u>	<u>22:22550113-22550244 22:22550370-22550686</u>
<u>361</u>	<u>IGLV7-43</u>	<u>ENSG00000211652</u>	<u>22:22749356-22749827</u>	<u>ENST00000390298</u>	<u>22:22749356-22749435 22:22749523-22749827</u>
<u>362</u>	<u>IGLV7-46</u>	<u>ENSG00000211649</u>	<u>22:22723982-22724454</u>	<u>ENST00000390295</u>	<u>22:22723982-22724061 22:22724151-22724454</u>
<u>363</u>	<u>IGLV8-61</u>	<u>ENSG00000211638</u>	<u>22:22453110-22453622</u>	<u>ENST00000390283</u>	<u>22:22453110-22453216 22:22453316-22453622</u>
<u>364</u>	<u>IGLV9-49</u>	<u>ENSG00000223350</u>	<u>22:22697539-22698084</u>	<u>ENST00000427632</u>	<u>22:22697539-22697621 22:22697759-22698084</u>

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**CLAIMS:**

1. A non-human vertebrate (optionally a mouse or a rat) or vertebrate cell whose genome comprises an immunoglobulin heavy chain locus comprising human gene segment JH6\*02,  
5 one or more VH gene segments and one or more D gene segments upstream of a constant region; wherein the gene segments in the heavy chain locus are operably linked to the constant region thereof so that the mouse is capable of producing an antibody heavy chain produced by recombination of the human JH6\*02 with a D segment and a VH segment.
- 10 2. The vertebrate of claim 1, wherein the vertebrate has been immunised with a target antigen and wherein the variable domain of the heavy chain is the product of recombination between a VH, D and JH6\*02 and wherein the HCDR3 length is at least 20 amino acids.
- 15 3. A non-human vertebrate cell (optionally a mouse cell or a rat cell) whose genome comprises an immunoglobulin heavy chain locus comprising human gene segment JH6\*02, one or more VH gene segments and one or more D gene segments upstream of a constant region; wherein the gene segments in the heavy chain locus are operably linked to the constant region thereof for producing (eg, in a subsequent progeny cell) an antibody heavy chain produced by recombination of the human JH6\*02 with a D segment and a VH segment.
- 20 4. The cell of claim 3, which is an ES cell capable of differentiation into a progeny antibody-producing cell that expresses said heavy chain.
- 25 5. The vertebrate or cell of any preceding claim, wherein the heavy chain locus comprises a human JH6\*02 recombination signal sequence (RSS) operably connected 5' to the JH6\*02 gene segment.
- 30 6. The vertebrate or cell of claim 5, wherein the RSS is SEQ ID NO: 238 or a sequence having an identical 9mer and 7mer sequence flanking a sequence that is at least 70% identical to the 22mer sequence of SEQ ID NO: 238.
7. The vertebrate or cell of claim 6, wherein the RSS and JH6\*02 are provided as SEQ ID NO: 237.

8. The vertebrate or cell of any preceding claim, wherein the JH6\*02 is the only JH6-type gene segment in the genome.
- 5 9. The vertebrate or cell of any preceding claim, wherein the JH6\*02 is the closest JH gene segment to the constant region in the locus.
10. The vertebrate or cell of any preceding claim, wherein the locus comprises one, more or all human D gene segments D3-9; D4-17; D3-10; D2-2; D5-24; D6-19; D3-22; D6-13; D5-12; D1-26; D1-20; D5-18; D3-16; D2-21; D1-14; D7-27; D1-1; D6-25; D2-14; and D4-23.
- 10 11. The vertebrate or cell of claim 10, wherein the locus comprises one, more or all human D gene segments D3-9, D3-10, D6-19, D4-17, D6-13, D3-22, D2-2, D2-25 and D3-3.
12. The vertebrate or cell of any preceding claim, wherein the locus comprises a plurality of human D gene segments and the JH6\*02 is in human germline configuration with respect to the 3'-most human D gene segment.
- 15 13. The vertebrate or cell of any preceding claim, wherein the locus comprises one, more or all of IGHV gene segments selected from V3-21, V3-13, V3-7, V6-1, V1-8, V1-2, V7-4-1, V1-3, V1-18, V4-4, V3-9, V3-23, V3-11 and V3-20.
- 20 14. The vertebrate or cell of any preceding claim, wherein the locus comprises one, more or all of human D3-9\*01, D3-10\*01, D6-19\*01, D6-13\*01, D1-26\*01, IGHV1-8\*01, IGHV4-61\*01, IGHV6-1\*01, IGHV4-4\*02, IGHV1-3\*01, IGHV3-66\*03, IGHV3-7\*01 and IGHV3-9\*01.
- 25 15. An antibody-producing cell (eg, a B-cell) that is a progeny of the cell of any one of claims 3 to 14, wherein the antibody-producing cell comprises a heavy chain locus comprising a rearranged variable region produced by recombination of human JH6\*02 with a D segment and a VH segment.
- 30 16. The cell of claim 15, which is a B-cell or hybridoma that expresses a target antigen-specific antibody comprising a heavy chain that comprises a rearranged variable region produced by recombination of human JH6\*02 with a D segment and a VH segment.

17. The vertebrate or cell of any preceding claim, wherein the antibody heavy chain specifically binds a target antigen.
- 5 18. The vertebrate or cell of any preceding claim, wherein the antibody heavy chain has a HCDR3 length of at least 20 amino acids.
- 10 19. The vertebrate or cell of any preceding claim, wherein the antibody heavy chain is a product of the recombination of JH6\*02 with a human VH gene segment recited in claim 13 or 14 and/or a D gene segment recited in claim 10, 11 or 14.
20. The vertebrate or cell of any preceding claim, wherein all endogenous non-human vertebrate heavy chain variable region gene segments have been inactivated in the genome.
- 15 21. The vertebrate or cell of any preceding claim, wherein the genome is homozygous for said heavy chain locus.
- 20 22. A heavy chain (eg, comprised by an antibody) isolated from a vertebrate of any one of claims 1, 2, 5 to 14 and 17 to 21 wherein the heavy chain comprises a HCDR3 of at least 20 amino acids.
23. The heavy chain of claim 22, wherein the HCDR3 is the product of recombination of human JH6\*02 with a human VH gene segment recited in claim 13 or 14 and/or a D gene segment recited in claim 10, 11 or 14.
- 25 24. A heavy chain (eg, comprised by an antibody) whose VH variable domain is identical to the VH variable domain of the heavy chain of claim 22 or 23, and which comprises a human constant region or a human-mouse chimaeric constant region.
- 30 25. The heavy chain of claim 22, 23 or 24, whose VH variable domain is specific for a target antigen.
26. A method for producing a heavy chain, VH domain or an antibody specific to a target antigen, the method comprising immunizing a non-human vertebrate according to any one

- 5 of claims 1, 2, 5 to 14 and 17 to 21 with the antigen and isolating the heavy chain, VH domain or an antibody specific to a target antigen or a cell producing the heavy chain, VH domain or an antibody, wherein the heavy chain, VH domain or an antibody comprises a HCDR3 that is derived from the recombination of human JH6\*02 with a VH gene segment and a D gene segment.
- 10 27. A method for producing a human heavy chain or antibody comprising carrying out the method of claim 26, wherein the constant region of the locus is a non-human vertebrate (eg, mouse or rat) constant region, and then replacing the non-human constant region of the isolated heavy chain or antibody with a human constant region.
28. A heavy chain, VH domain or an antibody produced by the method of claim 26 or 27.
- 15 29. A B-cell or hybridoma expressing a heavy chain VH domain that is identical to the VH domain of the heavy chain of claim 22, 23 or 28.
30. A nucleic acid encoding the VH domain of the heavy chain of claim 22, 23 or 28, or encoding the heavy chain of claim 22, 23, 24, 25 or 28.
- 20 31. A vector (eg, a CHO cell or HEK293 cell vector) comprising the nucleic acid of claim 30; optionally wherein the vector is in a host cell (eg, a CHO cell or HEK293 cell).
- 25 32. A pharmaceutical composition comprising the antibody, heavy chain or VH domain (eg, comprised by an antibody) of any one of claims 22 to 25 and 28, together with a pharmaceutically-acceptable excipient, diluent or a medicament (eg, a further antigen-specific variable domain, heavy chain or antibody).
- 30 33. The antibody, heavy chain or VH domain (eg, comprised by an antibody) of any one of claims 22 to 25 and 28 for use in medicine.
34. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising at least 3 human variable region gene segments of the same type (eg, at least 3 human VH6-1 gene segments, at least 3 human JH6 gene segments, at least 3 human Vk1-39 gene segments, at least 3 human D2-2 gene segments or

at least 3 human Jk1 gene segments), wherein at least two of the human gene segments are variants that are not identical to each other.

- 5 35. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising at least 2 different non-endogenous variable region gene segments of the same type (eg, at least 2 human VH6-1 gene segments, at least 3 human JH6 gene segments, at least 2 human Vk1-39 gene segments, at least 2 human D2-2 gene segments or at least 2 human Jk1 gene segments) *cis* at the same Ig locus.
- 10 36. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising at least 2 different human variable region gene segments of the same type (eg, at least 2 human VH6-1 gene segments, at least 2 human JH6 gene segments, at least 2 human Vk1-39 gene segments, at least 2 human D2-2 gene segments or at least 2 human Jk1 gene segments) *trans* at the same Ig locus; and optionally  
15 a third human gene segment of the same type, wherein the third gene segment is *cis* with one of said 2 different gene segments.
- 20 37. A population of non-human vertebrates (eg, mice or rats) comprising a repertoire of human variable region gene segments, wherein the plurality comprises at least 2 human variable region gene segments of the same type (eg, at least 2 human VH6-1 gene segments, at least 2 human JH6 gene segments, at least 2 human Vk1-39 gene segments, at least 2 human D2-2 gene segments or at least 2 human Jk1 gene segments), a first of said different gene segments is provided in the genome of a first vertebrate of the population, and a second of said different gene segments being provided in the genome of a second vertebrate of the  
25 population, wherein the genome of the first vertebrate does not comprise the second gene segment.
- 30 38. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising at least 2 different non-endogenous variable region gene segments of the same type (eg, at least 2 human VH6-1 gene segments, at least 2 human JH6 gene segments, at least 2 human Vk1-39 gene segments, at least 2 human D2-2 gene segments or at least 2 human Jk1 gene segments), wherein the gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations.
- 35 39. A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 3 human variable region gene segments of the same type (eg, at

least 3 human VH6-1 gene segments, at least 3 human JH6 gene segments, at least 3 human V $\kappa$ 1-39 gene segments, at least 3 human D2-2 gene segments or at least 3 human Jk1 gene segments), wherein at least two of the human gene segments are variants that are not identical to each other.

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40. A method of enhancing the immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 2 different non-endogenous variable region gene segments of the same type (eg, at least 2 human VH6-1 gene segments, at least 2 human JH6 gene segments, at least 2 human V $\kappa$ 1-39 gene segments, at least 2 human D2-2 gene segments or at least 2 human Jk1 gene segments) *cis* at the same Ig locus.

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41. A method of enhancing the immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 2 different human variable region gene segments of the same type (eg, at least 2 human VH6-1 gene segments, at least 2 human JH6 gene segments, at least 2 human V $\kappa$ 1-39 gene segments, at least 2 human D2-2 gene segments or at least 2 human Jk1 gene segments) *trans* at the same Ig locus; and optionally a third human gene segment of the same type, wherein the third gene segment is *cis* with one of said 2 different gene segments.

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42. A method of providing an enhanced human immunoglobulin variable region gene segment repertoire, the method comprising providing a population of non-human vertebrates (eg, a mouse or rat) comprising a repertoire of human variable region gene segments, wherein the method comprises providing at least 2 different human variable region gene segments of the same type (eg, at least 2 human VH6-1 gene segments, at least 2 human JH6 gene segments, at least 2 human V $\kappa$ 1-39 gene segments, at least 2 human D2-2 gene segments or at least 2 human Jk1 gene segments), wherein a first of said different gene segments is provided in the genome of a first vertebrate of the population, and a second of said different gene segments is provided in the genome of a second vertebrate of the population, wherein the genome of the first vertebrate does not comprise the second gene segment.

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43. A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 2 different non-endogenous variable region gene segments of the same type (eg, at least 2 human VH6-1 gene segments, at least 2 human JH6 gene segments, at least 2 human V $\kappa$ 1-39 gene segments, at least 2 human D2-2 gene segments or at least 2 human Jk1 gene segments), wherein the gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations.

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44. The vertebrate, cell or method of any one of claims 34 to 43, wherein at least 2 or 3 of said different gene segments are provided *cis* at the same Ig locus in said genome.
- 5 45. The vertebrate, cell or method of any one of claims 34 to 44, wherein the gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations.
- 10 46. The vertebrate, cell or method of any one of claims 34 to 45, wherein the gene segments are derived from the genome sequence of two or more different human individuals; optionally wherein the different human individuals are from different human populations.
47. The vertebrate, cell or method of claim 46, wherein the individuals are not genetically related.
- 15 48. A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 2 human variable region gene segments of the same type (eg, at least 2 human VH6-1 gene segments, at least 2 human JH6 gene segments, at least 2 human V $\kappa$ 1-39 gene segments, at least 2 human D2-2 gene segments or at least 2 human J $\kappa$ 1 gene segments), wherein the gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations; optionally wherein at least 2 or 3 of said different gene segments are provided at the same Ig locus in said genome.
- 20 49. The method of claim 48, wherein the different human individuals are from different human populations.
- 25 50. The method of claim 48, wherein the individuals are not genetically related.
- 30 51. The vertebrate, cell or method of preceding claim, wherein at least one of the different segments is a synthetic mutant of a human germline gene segment.



52. The vertebrate, cell or method of any one of claims 34 to 51, wherein each of said gene segments occurs in 10 or more different human populations.
53. The vertebrate, cell or method of preceding claim, wherein each of said gene segments has a human frequency of 5% or greater.
54. The vertebrate, cell or method of claim 53, wherein each of said gene segments occurs in 10 or more different human populations.
55. The vertebrate, cell or method of any one of claims 34 to 54, wherein each of said gene segments occurs in the 1000 Genomes database in more than 50 individuals.
56. The vertebrate, cell or method of preceding claim, wherein each of said gene segments (i) has a human frequency of 5% or greater; and (ii) occurs in 10 or more different human populations.
57. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) having a genome comprising first and second human Ig locus gene segments of the same type (eg, first and second human JH6 gene segments; or first and second IgG2 gene segments; or first and second human J $\lambda$ 7 gene segments), wherein the first gene segment is a gene segment selected from any one of Tables 1 to 7 and 9 to 14 (eg, IGHJ6-a) and the second gene segment is the corresponding reference sequence (eg, IGHJ6 ref; SEQ ID NO: 244).
58. The vertebrate or cell of claim 57, wherein the genome comprises a third human gene segment of said type, the third gene segment being different from the first and second gene segments.
59. The vertebrate or cell of claim 57 or 58, wherein the first and second gene segments are *cis* on the same chromosome; and optionally the third gene segment is also *cis* on said chromosome.
60. The vertebrate or cell of claim 59, wherein the gene segments are targeted insertions into an endogenous non-human Ig locus.

61. The vertebrate or cell of claim 57 or 58, wherein the first and second gene segments are *trans* on different chromosomes.
- 5 62. The vertebrate or cell of any one of claims 57 to 61, wherein the first gene segment is a gene segment selected any one of Tables 1 to 7 and 9 to 14 and the second gene segment is the corresponding reference sequence.
- 10 63. A population of non-human vertebrates (eg, mice or rats) comprising first and second human Ig locus gene segments of the same type (eg, first and second human JH6 gene segments; or first and second IgG2 gene segments; or first and second human Jλ7 gene segments), wherein the first gene segment is a gene segment selected from any one of Tables 1 to 7 and 9 to 14 (eg, IGHJ6-a) and the second gene segment is the corresponding reference sequence (eg, SEQ ID NO: 244), wherein the first gene segment is provided in the genome of a first vertebrate of the population, and the second gene segment is provided in the genome of a second vertebrate of the population.
- 15 64. The population of claim 63, wherein the genome of the first vertebrate does not comprise the second gene segment.
- 20 65. The population of claim 63 or 64, wherein the population comprises a third human gene segment of said type, the third gene segment being different from the first and second gene segments and optionally wherein the first and third gene segments are present in the genome of the first vertebrate.
- 25 66. The population of claim 63, 64 or 65, wherein the gene segments are targeted insertions into an endogenous non-human Ig locus in the respective genome.
- 30 67. The population of any one of claims 63 to 66, wherein the first gene segment is a gene segment selected from any one of Tables 1 to 7 and 9 to 14 and the second gene segment is the corresponding reference sequence.
- 35 68. A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising first and second human Ig locus gene segments of the same type (eg,

first and second human JH6 gene segments; or first and second IgG2 gene segments; or first and second human J $\lambda$ 7 gene segments), wherein the first gene segment is a gene segment selected from any one of Tables 1 to 7 and 9 to 14 (eg, IGHJ6-a) and the second gene segment is the corresponding reference sequence (eg, SEQ ID NO: 244).

5

69. A method of providing an enhanced human immunoglobulin gene segment repertoire, the method comprising providing a population according to any one of claims 63 to 66.

70. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) according to claim 34, having a genome comprising at least 3 human D gene segments of the same type (eg, D2-2 gene segments), wherein at least two of the human D gene segments are variants that are not identical to each other (eg, D2-2ref and D2-2a).

71. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) according to claim 35, having a genome comprising at least 2 different non-endogenous D gene segments of the same type type (eg, D2-2ref and D2-2a) *cis* at the same Ig locus.

72. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) according to claim 36, having a genome comprising at least 2 different human D gene segments of the same type (eg, D2-2ref and D2-2a) *trans* at the same Ig locus; and optionally a third human D gene segment (eg, (eg, D2-2ref, D2-2a or D2-2b) of the same type, wherein the third D is *cis* with one of said 2 different D gene segments.

73. A population of non-human vertebrates (eg, mice or rats) according to claim 37, comprising a repertoire of human D gene segments, wherein the plurality comprises at least 2 different human D gene segments of the same type (eg, D2-2 gene segments), a first of said different D gene segments (eg, D2-2ref) is provided in the genome of a first vertebrate of the population, and a second of said different D gene segment (eg, D2-2a) being provided in the genome of a second vertebrate of the population, wherein the genome of the first vertebrate does not comprise the second D gene segment.

74. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) according to claim 38, having a genome comprising at least 2 different non-endogenous D gene segments of the same type (eg, human D2-2 gene segments), wherein the D gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations.

75. A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat) according to claim 39, the method comprising providing the vertebrate with

a genome comprising at least 3 human D gene segments of the same type (eg, D2-2 gene segments), wherein at least two of the human D gene segments are variants that are not identical to each other (eg, D2-2ref and D2-2a).

- 5 76. A method of enhancing the immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat) according to claim 40, the method comprising providing the vertebrate with a genome comprising at least 2 different non-endogenous D gene segments of the same type (eg, human D2-2 gene segments) *cis* at the same Ig locus.
- 10 77. A method of enhancing the immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat) according to claim 41, the method comprising providing the vertebrate with a genome comprising at least 2 different human D gene segments of the same type (eg, D2-2ref and D2-2a) *trans* at the same Ig locus; and optionally a third human D gene segment (eg, D2-2ref, D2-2a or D2-2b) of the same type, wherein the third D is *cis* with one of said 2 different D gene segments.
- 15
78. A method of providing an enhanced human immunoglobulin D gene segment repertoire according to claim 42, the method comprising providing a population of non-human vertebrates (eg, a mouse or rat) comprising a repertoire of human D gene segments, wherein the method comprises providing at least 2 different human D gene segments of the same type (eg, D2-2ref and D2-2a), wherein a first of said different D gene segments is provided in the genome of a first vertebrate of the population, and a second of said different D gene segments is provided in the genome of a second vertebrate of the population, wherein the genome of the first vertebrate does not comprise the second D gene segment.
- 20
- 25
79. A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat) according to claim 43, the method comprising providing the vertebrate with a genome comprising at least 2 different non-endogenous D gene segments of the same type (eg, D2-2ref and D2-2a), wherein the D gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations.
- 30
80. The vertebrate or cell of claim 70, 71 or 74, or the method of claim 75, 77 or 79, wherein at least 2 or 3 of said different gene segments are provided *cis* at the same Ig locus in said genome.
- 35
81. The vertebrate or cell of claim 70, 71 or 71, or the method of any one of claims 75 to 78 and 80, wherein the D gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations.
- 40
82. The vertebrate or cell of any one of claims 70 to 73, or the method of any one of claims 75 to 78 and 81, wherein the D gene segments are derived from the genome sequence of two or more

different human individuals; optionally wherein the different human individuals are from different human populations.

- 5
83. The vertebrate, cell or method of claim 82, wherein the individuals are not genetically related.
84. The vertebrate, cell or method of any one of claims 70 to 83, wherein at least one of the different D segments is a synthetic mutant of a human germline D gene segment.
- 10
85. A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat) according to claim 48, the method comprising providing the vertebrate with a genome comprising at least 2 human D gene segments (eg, D2-2ref and D2-2a), wherein the D gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations; optionally wherein at least 2 or 3 of said different gene segments are provided at the same IgH locus in said genome.
- 15
86. The vertebrate or cell of any one of claims 70 to 74 and 80 to 84, wherein the genome comprises a substantially complete functional repertoire of human D gene segment types supplemented with one, two or more variant human D gene segments, wherein said substantially complete functional repertoire and the supplementary D gene segments are not found together in the germline genome of a human individual.
- 20
87. The population of claim 73, wherein the population comprises a substantially complete functional repertoire of human D gene segment types supplemented with one, two or more variant human D gene segments, wherein said substantially complete functional repertoire and the supplementary D gene segments are not found together in the germline genome of a human individual.
- 25
88. A non-human vertebrate (eg, a mouse or rat) or a non-human cell (eg, an ES cell or a B-cell) having a genome comprising a substantially complete functional repertoire of human D gene segment types supplemented with one, two or more variant human D gene segments, wherein said substantially complete functional repertoire and the supplementary D gene segments are not found together in the germline genome of a human individual.
- 30
89. A population of non-human vertebrates (eg, mice or rats) comprising a substantially complete functional repertoire of human JH gene segment types supplemented with one, two or more variant human D gene segments, wherein said substantially complete functional repertoire and the supplementary D gene segments are not found together in the germline genome of a human individual.
- 35

90. The vertebrate or cell of claim 88 or the population of claim 89, comprising first and second D gene segments selected from  
D2-2ref and D2-2a; or  
D2-21ref and D2-21a; or  
5 D3-10ref and D3-10a; or  
D3-16ref and D3-16a; or  
D2-8ref and D2-8a; or  
D3-3ref and D3-3a; or  
D4-23ref and D4-23a; or  
10 D6-13ref and D6-13a; or  
D3-9ref and D3-9a; or  
D4-4ref and D4-4a; or  
D7-27ref and D7-27a;
- 15 optionally wherein the first and/or second D gene segment is present in two or more copies.
91. The vertebrate, cell or population of claim 90, comprising human gene segments D2-2ref and D2-2a; and D3-3ref and D3-3a; and optionally also D2-15.
- 20
92. A non-human vertebrate or vertebrate cell according to claim 71, comprising a genome that comprises VH, D and JH gene repertoires comprising human gene segments, the D gene repertoire comprising one or more of
- 25 a plurality of D2-2 gene segments provided by at least 2 different D2-2 gene segments in *cis* at the same Ig locus in said genome;  
a plurality of D2-21 gene segments provided by at least 2 different D2-21 gene segments in *cis* at the same Ig locus in said genome;  
a plurality of D3-10 gene segments provided by at least 2 different D3-10 gene segments in *cis* at  
30 the same Ig locus in said genome;  
a plurality of D3-16 gene segments provided by at least 2 different D3-16 gene segments in *cis* at the same Ig locus in said genome;  
a plurality of D2-8 gene segments provided by at least 2 different D2-8 gene segments in *cis* at the same Ig locus in said genome;  
35 a plurality of D3-3 gene segments provided by at least 2 different D3-3 gene segments in *cis* at the same Ig locus in said genome;  
a plurality of D4-23 gene segments provided by at least 2 different D4-23 gene segments in *cis* at the same Ig locus in said genome;  
a plurality of D6-13 gene segments provided by at least 2 different D6-13 gene segments in *cis* at  
40 the same Ig locus in said genome;  
a plurality of D3-9 gene segments provided by at least 2 different D3-9 gene segments in *cis* at the same Ig locus in said genome;  
a plurality of D4-4 gene segments provided by at least 2 different D4-4 gene segments in *cis* at

the same Ig locus in said genome; and  
a plurality of D7-27 gene segments provided by at least 2 different D7-27 gene segments in *cis* at  
the same Ig locus in said genome;

5 optionally wherein the D gene segments are derived from the genome sequence of two or more  
different human individuals.

93. A non-human vertebrate or vertebrate cell according to claim 71, comprising a genome that  
comprises VH, D and JH gene repertoires comprising human gene segments, the D gene  
10 repertoire comprising one or more of

a plurality of D2-2 gene segments provided by at least 2 different D2-2 gene segments in *trans* in  
said genome;

15 a plurality of D2-21 gene segments provided by at least 2 different D2-21 gene segments in *trans*  
in said genome;

a plurality of D3-10 gene segments provided by at least 2 different D3-10 gene segments in *trans*  
in said genome;

20 a plurality of D3-16 gene segments provided by at least 2 different D3-16 gene segments in *trans*  
in said genome;

a plurality of D2-8 gene segments provided by at least 2 different D2-8 gene segments in *trans* in  
said genome;

25 a plurality of D3-3 gene segments provided by at least 2 different D3-3 gene segments in *trans* in  
said genome;

a plurality of D4-23 gene segments provided by at least 2 different D4-23 gene segments in *trans*  
in said genome;

30 a plurality of D6-13 gene segments provided by at least 2 different D6-13 gene segments in *trans*  
in said genome;

a plurality of D3-9 gene segments provided by at least 2 different D3-9 gene segments in *trans* in  
said genome;

35 a plurality of D4-4 gene segments provided by at least 2 different D4-4 gene segments in *trans* in  
said genome; and

a plurality of D7-27 gene segments provided by at least 2 different D7-27 gene segments in *trans*  
in said genome;

optionally wherein the D gene segments are derived from the genome sequence of two or more  
different human individuals.

94. A non-human vertebrate or vertebrate cell (optionally an ES cell or B-cell), according to claim 70,  
40 comprising a genome that comprises VH, D and JH gene repertoires comprising human gene  
segments, the D gene repertoire comprising one or more of

a plurality of D2-2 gene segments provided by at least 3 different D2-2 gene segments;

- a plurality of D2-21 gene segments provided by at least 3 different D2-21 gene segments;  
a plurality of D3-10 gene segments provided by at least 3 different D3-10 gene segments;  
a plurality of D3-16 gene segments provided by at least 3 different D3-16 gene segments;  
a plurality of D2-8 gene segments provided by at least 3 different D2-8 gene segments;  
5 a plurality of D3-3 gene segments provided by at least 3 different D3-3 gene segments;  
a plurality of D4-23 gene segments provided by at least 3 different D4-23 gene segments;  
a plurality of D6-13 gene segments provided by at least 3 different D6-13 gene segments;  
a plurality of D3-9 gene segments provided by at least 3 different D3-9 gene segments;  
10 a plurality of D4-4 gene segments provided by at least 3 different D4-4 gene segments; and  
a plurality of D7-27 gene segments provided by at least 3 different D7-27 gene segments;
- optionally wherein the D gene segments are derived from the genome sequence of two or three  
different human individuals;
- 15 optionally wherein at least 2 or 3 of said different gene segments are provided in *cis* at the same  
Ig locus in said genome.
95. The vertebrate or cell of claim 92, 93 or 94, wherein the different human individuals are from  
different human populations.  
20
96. The vertebrate or cell of any one of claims 92 to 95, wherein the individuals are not genetically  
related.
97. The vertebrate or cell of any one of claims 92 to 96, wherein at least one of the different D  
25 segments is a synthetic mutant of a human germline D gene segment.
98. A non-human vertebrate or vertebrate cell (optionally an ES cell or B-cell) according to claim 74,  
comprising a genome comprising human VH, D and JH gene repertoires, the D gene repertoire  
comprising of one or more of  
30 a plurality of D2-2 gene segments provided by at least 2 different D2-2 gene ; optionally in *cis* in  
said genome;  
a plurality of D2-21 gene segments provided by at least 2 different D2-21 gene ; optionally in *cis*  
in said genome;  
35 a plurality of D3-10 gene segments provided by at least 2 different D3-10 gene ; optionally in *cis*  
in said genome;  
a plurality of D3-16 gene segments provided by at least 2 different D3-16 gene ; optionally in *cis*  
in said genome;  
40 a plurality of D2-8 gene segments provided by at least 2 different D2-8 gene ; optionally in *cis* in  
said genome;  
a plurality of D3-3 gene segments provided by at least 2 different D3-3 gene ; optionally in *cis* in  
said genome;



a plurality of D4-23 gene segments provided by at least 2 different D4-23 gene ; optionally in *cis* in said genome;

a plurality of D6-13 gene segments provided by at least 2 different D6-13 gene ; optionally in *cis* in said genome;

5 a plurality of D3-9 gene segments provided by at least 2 different D3-9 gene ; optionally in *cis* in said genome;

a plurality of D4-4 gene segments provided by at least 2 different D4-4 gene ; optionally in *cis* in said genome; and

10 a plurality of D7-27 gene segments provided by at least 2 different D7-27 gene ; optionally in *cis* in said genome;

wherein the D gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations.

15 99. The vertebrate or cell of claim 98, wherein the human individuals are from different human populations.

100. The vertebrate, cell or population of any one of claims 70 to 99, wherein one or more of the D gene segments is a variant of a human germline D gene segment, wherein the variant gene  
20 segment encodes an amino acid sequence that differs by 1, 2 or 3 amino acids from the corresponding amino acid sequence encoded by the human germline D gene segment, provided in that said amino acid sequence encoded by the variant does not include a stop codon when said corresponding amino acid sequence does not include a stop codon.

25 101. The vertebrate, cell or population of claim 100, wherein said corresponding amino acid sequence encoded by the human germline D gene segment is a hydrophilic or hydrophobic sequence (according to J Mol Biol. 1997 Jul 25;270(4):587-97; Corbett SJ *et al*; Table 2).

30 102. The vertebrate, cell or population of claim 100 or 101, comprising said variant and said germline human D gene segments; optionally wherein the variant and germline human D gene segments are *cis* on the same chromosome.

103. The vertebrate, cell or population of any one of claims 100 to 102, wherein germline human D gene segment is a D2, D3, D5 or D6 family gene segment; optionally a D2-2, D2-15, D3-3, D3-9,  
35 D3-10, D3-22, D5-5, D5-18, D6-6, D6-13, D6-19 gene segment.

104. The vertebrate, cell or population of any one of claims 70 to 103, comprising a plurality of D2-2 gene segments, wherein the plurality comprises D2-2 gene segments that vary from each other at one or more nucleotide positions corresponding to positions  
40

106,382,687 and

106,382,711  
on human chromosome 14.

- 5 105. The vertebrate, cell or population of claim 104, wherein the plurality comprises a human D2-2 gene segment comprising a thymine at a position corresponding to position 106,382,687 on human chromosome 14; and optionally no further mutation from the sequence of D2-2ref.
- 10 106. The vertebrate, cell or population of claim 104 or 105, wherein the plurality comprises a human D2-2 gene segment comprising a cytosine at a position corresponding to position 106,382,687 on human chromosome 14; and optionally no further mutation from the sequence of D2-2a.
- 15 107. The vertebrate, cell or population of any one of claims 104 to 106, wherein the plurality comprises a human D2-2 gene segment comprising an adenine at a position corresponding to position 106,382,711 on human chromosome 14; and optionally no further mutation from the sequence of D2-2b.
- 20 108. The vertebrate, cell or population of any one of claims 104 to 107, wherein the plurality comprises a human D2-2 gene segment comprising an thymine at a position corresponding to position 106,382,711 on human chromosome 14; and optionally no further mutation from the sequence of D2-2ref.
- 25 109. The vertebrate, cell or population of any one of claims 70 to 108, comprising a plurality of D7-27 gene segments, wherein the plurality comprises D7-27 gene segments that vary from each other at a nucleotide position corresponding to position 106,331,767 on human chromosome 14.
- 30 110. The vertebrate, cell or population of claim 109, wherein the plurality comprises a human D7-27 gene segment comprising a cytosine at a position corresponding to position 106,331,767 on human chromosome 14; and optionally no further mutation from the sequence of D7-27ref.
- 35 111. The vertebrate, cell or population of claim 109 or 110, wherein the plurality comprises a human D7-27 gene segment comprising a guanine at a position corresponding to position 106,331,767 on human chromosome 14; and optionally no further mutation from the sequence of D7-27a.
112. The vertebrate, cell or population of any one of claims 70 to 111, comprising a plurality of D4-23 gene segments, wherein the plurality comprises D4-23 gene segments that vary from each other at a nucleotide position corresponding to position 106,350,740 on human chromosome 14.

- 5 113. The vertebrate, cell or population of claim 112, wherein the plurality comprises a human D4-23 gene segment comprising an adenine at a position corresponding to position 106,350,740 on human chromosome 14; and optionally no further mutation from the sequence of D4-23ref.
- 10 114. The vertebrate, cell or population of claim 112 or 113, wherein the plurality comprises a human D4-23 gene segment comprising an guanine at a position corresponding to position 106,350,740 on human chromosome 14; and optionally no further mutation from the sequence of D4-23a.
- 15 115. The vertebrate, cell or population of any one of claims 70 to 113, comprising a plurality of D2-21 gene segments, wherein the plurality comprises D2-21 gene segments that vary from each other at a nucleotide position corresponding to position 106,354,418 on human chromosome 14.
- 20 116. The vertebrate, cell or population of claim 115, wherein the plurality comprises a human D2-21 gene segment comprising an adenine at a position corresponding to position 106,354,418 on human chromosome 14; and optionally no further mutation from the sequence of D2-21ref.
- 25 117. The vertebrate, cell or population of claim 115 or 116, wherein the plurality comprises a human D2-21 gene segment comprising a guanine at a position corresponding to position 106,354,418 on human chromosome 14; and optionally no further mutation from the sequence of D2-21a.
- 30 118. The vertebrate, cell or population of any one of claims 70 to 117, comprising a plurality of D3-16 gene segments, wherein the plurality comprises D3-16 gene segments that vary from each other at a nucleotide position corresponding to position 106,354,418 on human chromosome 14.
- 35 119. The vertebrate, cell or population of claim 118, wherein the plurality comprises a human D3-16 gene segment comprising a thymine at a position corresponding to position 106,361,515 on human chromosome 14; and optionally no further mutation from the sequence of D3-16ref.
120. The vertebrate, cell or population of claim 118 or 119, wherein the plurality comprises a human D3-16 gene segment comprising a cytosine at a position corresponding to position 106,361,515 on human chromosome 14; and optionally no further mutation from the sequence of D3-16a.

121. The vertebrate, cell or population of any one of claims 70 to 120, comprising a plurality of D6-13 gene segments, wherein the plurality comprises D6-13 gene segments that vary from each other at a nucleotide position corresponding to position 106,367,013 on human chromosome 14.
- 5
122. The vertebrate, cell or population of claim 121, wherein the plurality comprises a human D6-13 gene segment comprising a thymine at a position corresponding to position 106,367,013 on human chromosome 14; and optionally no further mutation from the sequence of D6-13ref.
- 10 123. The vertebrate, cell or population of claim 121 or 122, wherein the plurality comprises a human D6-13 gene segment comprising a cytosine at a position corresponding to position 106,367,013 on human chromosome 14; and optionally no further mutation from the sequence of D6-13a.
- 15 124. The vertebrate, cell or population of any one of claims 70 to 123, comprising a plurality of D3-10 gene segments, wherein the plurality comprises D3-10 gene segments that vary from each other at one or more nucleotide positions corresponding to positions 106,370,370 and 106,370,371
- 20 on human chromosome 14.
125. The vertebrate, cell or population of claim 124, wherein the plurality comprises a human D3-10 gene segment comprising a thymine at a position corresponding to position 106,370,370 on human chromosome 14; and optionally no further mutation from the sequence of D3-10ref.
- 25
126. The vertebrate, cell or population of claim 124 or 125, wherein the plurality comprises a human D3-10 gene segment comprising a cytosine at a position corresponding to position 106,370,370 on human chromosome 14; and optionally no further mutation from the sequence of D3-10a.
- 30
127. The vertebrate, cell or population of claim 124, 125 or 126 wherein the plurality comprises a human D3-10 gene segment comprising an adenine at a position corresponding to position 106,370,371 on human chromosome 14; and optionally no further mutation from the sequence of D3-10ref.
- 35
128. The vertebrate, cell or population of any one of claims 124 to 127, wherein the plurality comprises a human D3-10 gene segment comprising a guanine at a position corresponding to position 106,370,371 on human chromosome 14; and optionally no further mutation from the

sequence of D3-10b.

129. The vertebrate, cell or population of any one of claims 70 to 128, comprising a plurality of D3-9 gene segments, wherein the plurality comprises D3-9 gene segments that vary from each other at a nucleotide position corresponding to position 106,370,567 on human chromosome 14.
130. The vertebrate, cell or population of claim 129, wherein the plurality comprises a human D3-9 gene segment comprising an adenine at a position corresponding to position 106,370,567 on human chromosome 14; and optionally no further mutation from the sequence of D3-9ref.
131. The vertebrate, cell or population of claim 129 or 130, wherein the plurality comprises a human D3-9 gene segment comprising a thymine at a position corresponding to position 106,370,567 on human chromosome 14; and optionally no further mutation from the sequence of D3-9a.
132. The vertebrate, cell or population of any one of claims 70 to 131, comprising a plurality of D2-8 gene segments, wherein the plurality comprises D2-8 gene segments that vary from each other at one or more nucleotide positions corresponding to positions 106,373,085; 106,373,086 and 106,373,089 on human chromosome 14.
133. The vertebrate, cell or population of claim 132, wherein the plurality comprises a human D2-8 gene segment comprising a cytosine at a position corresponding to position 106,373,085 on human chromosome 14.
134. The vertebrate, cell or population of claim 132 or 133, wherein the plurality comprises a human D2-8 gene segment comprising a thymine at a position corresponding to position 106,373,085 on human chromosome 14; and optionally no further mutation from the sequence of D2-8b.
135. The vertebrate, cell or population of claim 132, 133 or 134 wherein the plurality comprises a human D2-8 gene segment comprising a cytosine at a position corresponding to position 106,373,086 on human chromosome 14; and optionally no further mutation from the sequence of D2-8ref.

136. The vertebrate, cell or population of any one of claims 132 to 135, wherein the plurality comprises a human D2-8 gene segment comprising a thymine at a position corresponding to position 106,373,086 on human chromosome 14; and optionally no further mutation from the sequence of D2-8ref.
- 5
137. The vertebrate, cell or population of any one of claims 70 to 136, comprising a plurality of D4-4 gene segments, wherein the plurality comprises D4-4 gene segments that vary from each other at one or more nucleotide positions corresponding to positions 106,379,086; and
- 10 106,379,089  
on human chromosome 14.
138. The vertebrate, cell or population of claim 137, wherein the plurality comprises a D4-4 gene segment comprising a cytosine at a position corresponding to position 106,379,086 on human chromosome 14; and optionally no further mutation from the sequence of D4-4ref.
- 15
139. The vertebrate, cell or population of claim 137 or 138, wherein the plurality comprises a human D4-4 gene segment comprising a thymine at a position corresponding to position 106,379,086 on human chromosome 14; and optionally no further mutation from the sequence of D4-4a.
- 20
140. The vertebrate, cell or population of claim 137, 138 or 139 wherein the plurality comprises a human D4-4 gene segment comprising a cytosine at a position corresponding to position 106,379,089 on human chromosome 14; and optionally no further mutation from the sequence of D4-4ref or a cytosine at a position corresponding to position 106,379,086 on human chromosome 14.
- 25
141. The vertebrate, cell or population of any one of claims 137 to 140, wherein the plurality comprises a human D4-4 gene segment comprising a thymine at a position corresponding to position 106,373,089 on human chromosome 14; and optionally no further mutation from the sequence of D4-4a.
- 30
142. The vertebrate, cell or population of any one of claims 70 to 141, comprising a plurality of D3-3 gene segments, wherein the plurality comprises D3-3 gene segments that vary from each other at one or more nucleotide positions corresponding to positions 106,380,241; and 106,380,246 on human chromosome 14.
- 35

143. The vertebrate, cell or population of claim 142, wherein the plurality comprises a D3-3 gene segment comprising a thymine at a position corresponding to position 106,380,241 on human chromosome 14; and optionally no further mutation from the sequence of D3-3ref.
- 5 144. The vertebrate, cell or population of claim 142 or 143, wherein the plurality comprises a human D3-3 gene segment comprising a cytosine at a position corresponding to position 106,380,241 on human chromosome 14; and optionally no further mutation from the sequence of D3-3a.
- 10 145. The vertebrate, cell or population of claim 142, 143 or 144 wherein the plurality comprises a human D3-3 gene segment comprising an adenine at a position corresponding to position 106,380,246 on human chromosome 14; and optionally no further mutation from the sequence of D3-3ref.
- 15 146. The vertebrate, cell or population of any one of claims 142 to 145, wherein the plurality comprises a human D3-3 gene segment comprising a thymine at a position corresponding to position 106,380,246 on human chromosome 14; and optionally no further mutation from the sequence of D3-3a.
- 20 147. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) according to claim 34, having a genome comprising at least 3 human JH gene segments of the same type (JH1, JH2, JH3, JH4, JH5 or JH6), wherein at least two of the human JH gene segments are variants that are not identical to each other.
- 25 148. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) according to claim 35, having a genome comprising at least 2 different non-endogenous JH gene segments of the same type (JH1, JH2, JH3, JH4, JH5 or JH6) *cis* at the same Ig locus.
- 30 149. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) according to claim 36, having a genome comprising at least 2 different human JH gene segments of the same type (JH1, JH2, JH3, JH4, JH5 or JH6) *trans* at the same Ig locus; and optionally a third human JH gene segments of the same type, wherein the third JH is *cis* with one of said 2 different JH gene segments.
- 35 150. A population of non-human vertebrates (eg, mice or rats) comprising a repertoire of human JH gene segments according to claim 37, wherein the plurality comprises at least 2 different human JH gene segments of the same type (JH1, JH2, JH3, JH4, JH5 or JH6), a first of said different JH gene segments is provided in the genome of a first vertebrate of the population, and
- 40 a second of said different JH gene segments being provided in the genome of a second vertebrate of the population, wherein the genome of the first vertebrate does not comprise the second JH gene segment.
- 45 151. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) according to claim 38, having a genome comprising at least 2 different non-

endogenous JH gene segments of the same type (JH1, JH2, JH3, JH4, JH5 or JH6), wherein the JH gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations.

- 5 152. A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat) according to claim 39, the method comprising providing the vertebrate with a genome comprising at least 3 human JH gene segments of the same type (JH1, JH2, JH3, JH4, JH5 or JH6), wherein at least two of the human JH gene segments are variants that are not identical to each other.
- 10 153. A method of enhancing the immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat) according to claim 40, the method comprising providing the vertebrate with a genome comprising at least 2 different non-endogenous JH gene segments of the same type (JH1, JH2, JH3, JH4, JH5 or JH6) *cis* at the same Ig locus.
- 15 154. A method of enhancing the immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat) according to claim 41, the method comprising providing the vertebrate with a genome comprising at least 2 different human JH gene segments of the same type (JH1, JH2, JH3, JH4, JH5 or JH6) *trans* at the same Ig locus; and optionally a third human JH gene segments of the same type, wherein the third JH is *cis* with one of said 2 different JH gene segments.
- 20 155. A method of providing an enhanced human immunoglobulin JH gene segment repertoire according to claim 42, the method comprising providing a population of non-human vertebrates (eg, a mouse or rat) comprising a repertoire of human JH gene segments, wherein the method comprises providing at least 2 different human JH gene segments of the same type (JH1, JH2, JH3, JH4, JH5 or JH6), wherein a first of said different JH gene segments is provided in the genome of a first vertebrate of the population, and a second of said different JH gene segments is provided in the genome of a second vertebrate of the population, wherein the genome of the first vertebrate does not comprise the second JH gene segment.
- 25 30 156. A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat) according to claim 43, the method comprising providing the vertebrate with a genome comprising at least 2 different non-endogenous JH gene segments of the same type (JH1, JH2, JH3, JH4, JH5 or JH6), wherein the JH gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations.
- 35 157. The vertebrate or cell of claim 147, 149 or 151, or the method of claim 152, 154, 155 or 156, wherein at least 2 or 3 of said different gene segments are provided *cis* at the same Ig locus in said genome.
- 40 158. The vertebrate or cell of claim 147, 148 or 149, or the method of claim 152, 153, 154 or 155, wherein the JH gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations.
- 45 159. The vertebrate or cell of claim 147, 148 or 149, or the method of claim 152, 153, 154 or 155, wherein the JH gene segments are derived from the genome sequence of two or more different human individuals; optionally wherein the different human individuals are from different human populations.
- 50



160. The vertebrate, cell or method of claim 159, wherein the individuals are not genetically related.
- 5 161. The vertebrate, cell or method of any one of claims 147 to 160, wherein at least one of the different JH segments is a synthetic mutant of a human germline JH gene segment.
- 10 162. A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 2 human JH gene segments of the same type (JH1, JH2, JH3, JH4, JH5 or JH6), wherein the JH gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations; optionally wherein at least 2 or 3 of said different gene segments are provided at the same IgH locus in said genome.
- 15 163. The vertebrate or cell of any one of claims 147 to 149 and 151, wherein the genome comprises a substantially complete functional repertoire of human JH gene segment types supplemented with one, two or more human JH gene segments, wherein said substantially complete functional repertoire and the supplementary JH gene segments are not found together in the germline genome of a human individual.
- 20 164. The population of claim 150, wherein the population comprises a substantially complete functional repertoire of human JH gene segment types supplemented with one, two or more human JH gene segments, wherein said substantially complete functional repertoire and the supplementary JH gene segments are not found together in the germline genome of a human individual.
- 25 165. A non-human vertebrate (eg, a mouse or rat) or a non-human cell (eg, an ES cell or a B-cell) having a genome comprising a substantially complete functional repertoire of human JH gene segment types supplemented with one, two or more human JH gene segments, wherein said substantially complete functional repertoire and the supplementary JH gene segments are not found together in the germline genome of a human individual.
- 30 166. A population of non-human vertebrates (eg, mice or rats) comprising a substantially complete functional repertoire of human JH gene segment types supplemented with one, two or more human JH gene segments, wherein said substantially complete functional repertoire and the supplementary JH gene segments are not found together in the germline genome of a human individual.
- 35 167. The vertebrate of claim 165 or the population of claim 166, wherein at least one of said JH gene segments is SEQ ID NO: 1, 2, 3 or 4.
- 40 168. The vertebrate or population of claim 167, wherein at least one of said JH gene segments is SEQ ID NO: 1 and at least one, two or more of said supplementary JH gene segments is a variant according to any one of claims 178 to 196.
- 45 169. The vertebrate or population of claim 167 or 168, wherein at least one of said JH gene segments is SEQ ID NO: 2 and at least one, two or more of said supplementary JH gene segments is a variant according to any one of claims 198 to 209.
- 50 170. The vertebrate or population of claim 167, 168 or 169, wherein at least one of said JH gene segments is SEQ ID NO: 2 and at least one, two or more of said supplementary JH gene segments

is a variant according to any one of claims 212 to 217.

5 171. A non-human vertebrate or vertebrate cell according to claim 148, comprising a genome that comprises VH, D and JH gene repertoires comprising human gene segments, the JH gene repertoire comprising

a plurality of JH1 gene segments provided by at least 2 different JH1 gene segments in *cis* at the same Ig locus in said genome;

10 a plurality of JH2 gene segments provided by at least 2 different JH2 gene segments in *cis* at the same Ig locus in said genome;

a plurality of JH3 gene segments provided by at least 2 different JH3 gene segments in *cis* at the same Ig locus in said genome;

a plurality of JH4 gene segments provided by at least 2 different JH4 gene segments in *cis* at the same Ig locus in said genome;

15 a plurality of JH5 gene segments provided by at least 2 different JH5 gene segments in *cis* at the same Ig locus in said genome; or

a plurality of JH6 gene segments provided by at least 2 different JH6 gene segments in *cis* at the same Ig locus in said genome;

20 optionally wherein the JH gene segments are derived from the genome sequence of two or more different human individuals.

25 172. A non-human vertebrate or vertebrate cell (optionally an ES cell or B-cell), according to claim 147, comprising a genome that comprises VH, D and JH gene repertoires comprising human gene segments, the JH gene repertoire comprising

a plurality of JH1 gene segments provided by at least 3 different JH1 gene segments;

a plurality of JH2 gene segments provided by at least 3 different JH2 gene segments;

30 a plurality of JH3 gene segments provided by at least 3 different JH3 gene segments;

a plurality of JH4 gene segments provided by at least 3 different JH4 gene segments;

a plurality of JH5 gene segments provided by at least 3 different JH5 gene segments; or

a plurality of JH6 gene segments provided by at least 3 different JH6 gene segments;

35 optionally wherein the JH gene segments are derived from the genome sequence of two or three different human individuals;

optionally wherein at least 2 or 3 of said different gene segments are provided in *cis* at the same Ig locus in said genome.

40 173. The vertebrate or cell of claim 171 or 172, wherein the different human individuals are from different human populations.

45 174. The vertebrate or cell of any one of claims 171 to 173, wherein the individuals are not genetically related.

175. The vertebrate or cell of any one of claims 171 to 174, wherein at least one of the different JH segments is a synthetic mutant of a human germline JH gene segment.

50 176. A non-human vertebrate or vertebrate cell (optionally an ES cell or B-cell) according to claim 151, comprising a genome comprising human VH, D and JH gene repertoires, the JH gene repertoire comprising

- 5 a plurality of JH1 gene segments provided by at least 2 different human JH1 gene segments, optionally in *cis* at the same Ig locus in said genome;
- 10 a plurality of JH2 gene segments provided by at least 2 different human JH2 gene segments, optionally in *cis* at the same Ig locus in said genome;
- 15 a plurality of JH3 gene segments provided by at least 2 different human JH3 gene segments, optionally in *cis* at the same Ig locus in said genome;
- 20 a plurality of JH4 gene segments provided by at least 2 different human JH4 gene segments, optionally in *cis* at the same Ig locus in said genome;
- 25 a plurality of JH5 gene segments provided by at least 2 different human JH5 gene segments, optionally in *cis* at the same Ig locus in said genome; or
- 30 a plurality of JH6 gene segments provided by at least 2 different human JH6 gene segments, optionally in *cis* at the same Ig locus in said genome;
- 35 wherein the JH gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations.

177. The vertebrate or cell of claim 176, wherein the human individuals are from different human populations.

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178. The vertebrate, cell or population of any one of claims 147 to 151, 157 to 161 and 163 to 177, comprising a plurality of JH5 gene segments, wherein the plurality comprises a human JH5 gene variant of SEQ ID NO: 1, wherein the variant comprises a nucleotide mutation at one or more positions corresponding to positions

25

106,330,024

106,330,027

106,330,032

106,330,041

30

106,330,044

106,330,045

106,330,062

106,330,063

106,330,065

35

106,330,066

106,330,067

106,330,068 and

106,330,071

40

on human chromosome 14.

179. The vertebrate, cell or population of claim 178, wherein the plurality comprises a human JH5 gene variant of SEQ ID NO: 1, wherein the variant comprises a guanine at a position corresponding to position 106,330,067 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 1.

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180. The vertebrate, cell or population of claim 179, wherein the variant comprises additionally a mutation at a position corresponding to (i) position 106,330,071 on human chromosome 14 (optionally the additional mutation being a guanine); (ii) position 106,330,066 on human

chromosome 14 (optionally the additional mutation being a guanine); and/or (iii) position 106,330,068 on human chromosome 14 (optionally the additional mutation being a thymine).

- 5 181. The vertebrate, cell or population of claim 178, 179 or 180, wherein the plurality comprises a human JH5 gene variant of SEQ ID NO: 1, wherein the variant comprises a guanine at a position corresponding to position 106,330,071 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 1.
- 10 182. The vertebrate, cell or population of claim 181, wherein the variant comprises additionally a mutation at a position corresponding to (i) position 106,330,063 on human chromosome 14 (optionally the additional mutation being an adenine); and/or (ii) position 106,330,067 on human chromosome 14 (optionally the additional mutation being a guanine).
- 15 183. The vertebrate, cell or population of claim 178 to 182, wherein the plurality comprises a human JH5 gene variant of SEQ ID NO: 1, wherein the variant comprises a cytosine at a position corresponding to position 106,330,045 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 1.
- 20 184. The vertebrate, cell or population of claim 178 to 183, wherein the plurality comprises a human JH5 gene variant of SEQ ID NO: 1, wherein the variant comprises an adenine at a position corresponding to position 106,330,044 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 1.
- 25 185. The vertebrate, cell or population of claim 184, wherein the variant comprises additionally a mutation at a position corresponding to (i) position 106,330,066 on human chromosome 14 (optionally the additional mutation being a guanine); and/or (ii) position 106,330,068 on human chromosome 14 (optionally the additional mutation being a thymine).
- 30 186. The vertebrate, cell or population of claim 178 to 185, wherein the plurality comprises a human JH5 gene variant of SEQ ID NO: 1, wherein the variant comprises a guanine at a position corresponding to position 106,330,066 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 1.
- 35 187. The vertebrate, cell or population of claim 186, wherein the variant comprises additionally a mutation at a position corresponding to (i) position 106,330,067 on human chromosome 14 (optionally the additional mutation being a guanine); and/or (ii) position 106,330,068 on human chromosome 14 (optionally the additional mutation being a thymine).
- 40 188. The vertebrate, cell or population of claim 178 to 185, wherein the plurality comprises a human JH5 gene variant of SEQ ID NO: 1, wherein the variant comprises a thymine at a position corresponding to position 106,330,068 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 1.
- 45 189. The vertebrate, cell or population of claim 188, wherein the variant comprises additionally a mutation at a position corresponding to (i) position 106,330,067 on human chromosome 14 (optionally the additional mutation being a guanine); and/or (ii) position 106,330,066 on human chromosome 14 (optionally the additional mutation being a guanine).
- 50 190. The vertebrate, cell or population of claim 178 to 189, wherein the plurality comprises a human JH5 gene variant of SEQ ID NO: 1, wherein the variant comprises a cytosine at a position corresponding to position 106,330,027 on human chromosome 14; and optionally no further

mutation from the sequence of SEQ ID NO: 1.

- 5 191. The vertebrate, cell or population of claim 178 to 190, wherein the plurality comprises a human JH5 gene variant of SEQ ID NO: 1, wherein the variant comprises an adenine at a position corresponding to position 106,330,024 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 1.
- 10 192. The vertebrate, cell or population of claim 178 to 191, wherein the plurality comprises a human JH5 gene variant of SEQ ID NO: 1, wherein the variant comprises a thymine at a position corresponding to position 106,330,032 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 1.
- 15 193. The vertebrate, cell or population of claim 178 to 192, wherein the plurality comprises a human JH5 gene variant of SEQ ID NO: 1, wherein the variant comprises a thymine at a position corresponding to position 106,330,041 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 1.
- 20 194. The vertebrate, cell or population of claim 178 to 193, wherein the plurality comprises a human JH5 gene variant of SEQ ID NO: 1, wherein the variant comprises an adenine or thymine at a position corresponding to position 106,330,063 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 1.
- 25 195. The vertebrate, cell or population of claim 194, wherein the variant comprises additionally a mutation at a position corresponding to position 106,330,071 on human chromosome 14 (optionally the additional mutation being a guanine).
- 30 196. The vertebrate, cell or population of claim 178 to 195, wherein the plurality comprises a human JH5 gene variant of SEQ ID NO: 1, wherein the variant comprises a cytosine at a position corresponding to position 106,330,062 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 1.
- 35 197. The vertebrate, cell or population of claim 178 to 196, wherein the genome comprises SEQ ID NO:1; optionally in *cis* at the same Ig locus as one, two or more of the variants.
- 40 198. The vertebrate, cell or population of any one of claims 147 to 151, 157 to 161 and 96 to 130, comprising a plurality of JH6 gene segments, wherein the plurality comprises a human JH6 gene variant of SEQ ID NO: 2, wherein the variant comprises a nucleotide mutation at one or more positions corresponding to positions
- 45 106,329,411  
106,329,413  
106,329,414  
106,329,417  
106,329,419  
106,329,426  
106,329,434  
106,329,435, and  
106,329,468

on human chromosome 14.

- 5 199. The vertebrate, cell or population of claim 198, comprising a plurality of JH6 gene segments, wherein the plurality comprises a human JH6 gene variant of SEQ ID NO: 2, wherein the variant comprises a guanine at a position corresponding to position 106,329,435 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 2.
- 10 200. The vertebrate, cell or population of claim 199, wherein the variant comprises additionally a mutation at a position corresponding to (i) position 106,329,468 on human chromosome 14 (optionally the additional mutation being a guanine); (ii) position 106,329,419 on human chromosome 14 (optionally the additional mutation being an adenine); (iii) position 106,329,434 on human chromosome 14 (optionally the additional mutation being a cytosine) and/or position 106,329,414 on human chromosome 14 (optionally the additional mutation being a guanine); (iv) position 106,329,426 on human chromosome 14 (optionally the additional mutation being an adenine); (v) position 106,329,413 on human chromosome 14 (optionally the additional mutation being an adenine); (vi) position 106,329,417 on human chromosome 14 (optionally the additional mutation being a thymine); (vii) position 106,329,411 on human chromosome 14 (optionally the additional mutation being a thymine); (viii) position 106,329,451 on human chromosome 14 (optionally the additional mutation being an adenine); (ix) position 106,329,452 on human chromosome 14 (optionally the additional mutation being a cytosine); and/or (x) position 106,329,453 on human chromosome 14 (optionally the additional mutation being a cytosine).
- 15 201. The vertebrate, cell or population of claim 200, wherein the variant comprises additionally mutations at positions corresponding to position 106,329,451 on human chromosome 14, the additional mutation being an adenine; position 106,329,452 on human chromosome 14, the additional mutation being a cytosine; and position 106,329,453 on human chromosome 14, the additional mutation being a cytosine.
- 20 202. The vertebrate, cell or population of any one of claims 147 to 151, 157 to 161 and 163 to 201, comprising a plurality of JH6 gene segments,, wherein the plurality comprises a human JH6 gene variant of SEQ ID NO: 2, wherein the variant comprises a guanine at a position corresponding to position 106,329,468 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 2.
- 25 203. The vertebrate, cell or population of claim 202, wherein the variant comprises additionally a mutation at a position corresponding to position 106,329,435 on human chromosome 14 (optionally the additional mutation being a guanine).
- 30 204. The vertebrate, cell or population of any one of claims 147 to 151, 157 to 161 and 163 to 203, comprising a plurality of JH6 gene segments,, wherein the plurality comprises a human JH6 gene variant of SEQ ID NO: 2, wherein the variant comprises a thymine at a position corresponding to position 106,329,417 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 2.
- 35 205. The vertebrate, cell or population of claim 204, wherein the variant comprises additionally a mutation at a position corresponding to position 106,329,435 on human chromosome 14 (optionally the additional mutation being a guanine).
- 40
- 45

206. The vertebrate, cell or population of any one of claims 147 to 151, 157 to 161 and 163 to 205, comprising a plurality of JH6 gene segments,, wherein the plurality comprises a human JH6 gene variant of SEQ ID NO: 2, wherein the variant comprises a cytosine at a position corresponding to position 106,329,434 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 2.
207. The vertebrate, cell or population of claim 206, wherein the variant comprises additionally a mutation at a position corresponding to (i) position 106,329,414 on human chromosome 14 (optionally the additional mutation being a guanine); and/or (ii) position 106,329,435 on human chromosome 14 (optionally the additional mutation being a guanine).
208. The vertebrate, cell or population of any one of claims 147 to 151, 157 to 161 and 163 to 207, comprising a plurality of JH6 gene segments, wherein the plurality comprises a human JH6 gene variant of SEQ ID NO: 2, wherein the variant comprises a thymine at a position corresponding to position 106,329,411 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 2.
209. The vertebrate, cell or population of claim 208, wherein the variant comprises additionally a mutation at a position corresponding to position 106,329,435 on human chromosome 14 (optionally the additional mutation being a guanine).
210. The vertebrate, cell or population of any one of claims 147 to 151, 157 to 161 and 163 to 209, comprising a plurality of JH6 gene segments, wherein the plurality comprises a human JH6 gene variant that is an antisense sequence of the variant of any one of claims 107 to 118.
211. The vertebrate, cell or population of any one of claims 198 to 210, wherein the genome comprises SEQ ID NO:2; optionally *cis* at the same Ig locus as one, two or more of the JH6 variants.
212. The vertebrate, cell or population of any one of claims 147 to 151, 157 to 161 and 163 to 211, comprising a plurality of JH2 gene segments, wherein the plurality comprises a human JH2 gene variant of SEQ ID NO: 3, wherein the variant comprises a nucleotide mutation at one or more positions corresponding to positions  
106,331,455  
106,331,453, and  
106,331,409  
on human chromosome 14.
213. The vertebrate, cell or population of claim 212, comprising said plurality of JH2 gene segments, wherein the plurality comprises a human JH2 gene variant of SEQ ID NO: 3, wherein the variant comprises a guanine at a position corresponding to position 106,331,455 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 3.

214. The vertebrate, cell or population of claim 213, wherein the variant comprises additionally a mutation at a position corresponding to (i) position 106,331,453 on human chromosome 14 (optionally the additional mutation being an adenine); and/or (ii) position 106,331,409 on human chromosome 14 (optionally the additional mutation being an adenine); (iii) position 106,329,434 on human chromosome 14 (optionally the additional mutation being an adenine).
215. The vertebrate, cell or population of any one of claims 147 to 151, 157 to 161 and 163 to 214, comprising a plurality of JH2 gene segments, wherein the plurality comprises a human JH2 gene variant of SEQ ID NO: 3, wherein the variant comprises an adenine at a position corresponding to position 106,331,453 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 3.
216. The vertebrate, cell or population of claim 215, wherein the variant comprises additionally a mutation at a position corresponding to position 106,331,409 on human chromosome 14 (optionally the additional mutation being an adenine).
217. The vertebrate, cell or population of any one of claims 147 to 151, 157 to 161 and 163 to 216, comprising a plurality of JH2 gene segments, wherein the plurality comprises a human JH2 gene variant of SEQ ID NO: 3, wherein the variant comprises an adenine at a position corresponding to position 106,331,409 on human chromosome 14; and optionally no further mutation from the sequence of SEQ ID NO: 3.
218. The vertebrate, cell or population of any one of claims 147 to 151, 157 to 161 and 163 to 217, comprising a plurality of JH2 gene segments, wherein the plurality comprises a human JH2 gene variant that is an antisense sequence of the variant of any one of claims 188 to 193.
219. The vertebrate, cell or population of any one of claims 212 to 218, wherein the genome comprises SEQ ID NO:3; optionally *cis* at the same Ig locus as one, two or more of the JH2 variants.
220. The vertebrate, cell or population of any one of claims 147 to 151, 157 to 161 and 163 to 219, wherein the genome comprises two or more different JH gene segments selected from SEQ ID NOs: 1 to 3 and variants according to any one of claims 178 to 219; optionally wherein said JH gene segments are *cis* at the same immunoglobulin Ig locus.
221. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) according to claim 34, having a genome comprising at least 3 human JL gene segments of the same type (eg, Jk1), wherein at least two of the human JL gene segments are variants that are not identical to each other.
222. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) according to claim 35, having a genome comprising at least 2 different non-endogenous JL gene segments of the same type (eg, Jk1) *cis* at the same Ig locus.
223. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) according to claim 36, having a genome comprising at least 2 different human JL gene segments of the same type (eg, Jk1) *trans* at the same Ig locus; and optionally a third human JL gene segment of the same type, wherein the third JL is *cis* with one of said 2 different JL gene segments.



224. A population of non-human vertebrates (eg, mice or rats) comprising a repertoire of human JL gene segments according to claim 37, wherein the plurality comprises at least 2 different human JL gene segments of the same type (eg, Jk1), a first of said different JL gene segments is provided in the genome of a first vertebrate of the population, and a second of said different JL gene segments being provided in the genome of a second vertebrate of the population, wherein the genome of the first vertebrate does not comprise the second JL gene segment.
225. A non-human vertebrate (eg, a mouse or rat) or a non-human vertebrate cell (eg, an ES cell or a B-cell) according to claim 38, having a genome comprising at least 2 different non-endogenous JL gene segments of the same type (eg, Jk1), wherein the JL gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations.
226. A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat) according to claim 39, the method comprising providing the vertebrate with a genome comprising at least 3 human JL gene segments of the same type (eg, Jk1), wherein at least two of the human JL gene segments are variants that are not identical to each other.
227. A method of enhancing the immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat) according to claim 40, the method comprising providing the vertebrate with a genome comprising at least 2 different non-endogenous JL gene segments of the same type (eg, Jk1) *cis* at the same Ig locus.
228. A method of enhancing the immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat) according to claim 41, the method comprising providing the vertebrate with a genome comprising at least 2 different human JL gene segments of the same type (eg, Jk1) *trans* at the same Ig locus; and optionally a third human JL gene segment of the same type, wherein the third JL is *cis* with one of said 2 different JL gene segments.
229. A method of providing an enhanced human immunoglobulin JL gene segment repertoire according to claim 42, the method comprising providing a population of non-human vertebrates (eg, a mouse or rat) comprising a repertoire of human JL gene segments, wherein the method comprises providing at least 2 different human JL gene segments of the same type (eg, Jk1), wherein a first of said different JL gene segments is provided in the genome of a first vertebrate of the population, and a second of said different JL gene segments is provided in the genome of a second vertebrate of the population, wherein the genome of the first vertebrate does not comprise the second JL gene segment.
230. A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat) according to claim 43, the method comprising providing the vertebrate with a genome comprising at least 2 different non-endogenous JL gene segments of the same type (eg, Jk1), wherein the JL gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations.
231. The vertebrate or cell of claim 221, 223 or 225, or the method of claim 226, 228, 229 or 230, wherein at least 2 or 3 of said different gene segments are provided *cis* at the same Ig locus in said genome.
232. The vertebrate or cell of claim 221, 222 or 223, or the method of claim 226, 227, 228 or 229, wherein the JL gene segments are derived from the genome sequence of different human

individuals that are not genetically related over at least 3 generations.

233. The vertebrate or cell of claim 221, 222 or 223, or the method of claim 226, 227, 228 or 229, wherein the JL gene segments are derived from the genome sequence of two or more different human individuals; optionally wherein the different human individuals are from different human populations.

234. The vertebrate, cell or method of claim 233, wherein the individuals are not genetically related.

235. The vertebrate, cell or method of any one of claims 221 to 234, wherein at least one of the different JL segments is a synthetic mutant of a human germline JL gene segment.

236. A method of enhancing the human immunoglobulin gene diversity of a non-human vertebrate (eg, a mouse or rat), the method comprising providing the vertebrate with a genome comprising at least 2 human JL gene segments of the same type (eg, Jk1), wherein the JL gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations; optionally wherein at least 2 or 3 of said different gene segments are provided at the same IgL locus in said genome.

237. The vertebrate or cell of any one of claims claim 221 to 223 and 225, wherein the genome comprises a substantially complete functional repertoire of human Jk and/or Jλ gene segment types supplemented with one, two or more human Jk and/or Jλ gene segments respectively, wherein said substantially complete functional repertoire and the supplementary gene segments are not found together in the germline genome of a human individual.

238. The population of claim 224, wherein the population comprises a substantially complete functional repertoire of human JL gene segment types supplemented with one, two or more human Jk and/or Jλ gene segments respectively, wherein said substantially complete functional repertoire and the supplementary gene segments are not found together in the germline genome of a human individual.

239. A non-human vertebrate (eg, a mouse or rat) or a non-human cell (eg, an ES cell or a B-cell) having a genome comprising a substantially complete functional repertoire of human Jk and/or Jλ gene segment types supplemented with one, two or more human Jk and/or Jλ gene segments respectively, wherein said substantially complete functional repertoire and the supplementary gene segments are not found together in the germline genome of a human individual.

240. A population of non-human vertebrates (eg, mice or rats) comprising a substantially complete functional repertoire of human Jk and/or Jλ gene segment types supplemented with one, two or more human Jk and/or Jλ gene segments respectively, wherein said substantially complete functional repertoire and the supplementary gene segments are not found together in the germline genome of a human individual.

241. A non-human vertebrate or vertebrate cell according to claim 222, comprising a genome that comprises VL and JL gene repertoires comprising human gene segments, the JL gene repertoire comprising

a plurality of human Jk1 gene segments provided by at least 2 different human Jk1 gene segments in *cis* at the same Ig locus in said genome;  
a plurality of human Jk2 gene segments provided by at least 2 different human Jk1 gene

segments in *cis* at the same Ig locus in said genome;  
 a plurality of human J $\kappa$ 3 gene segments provided by at least 2 different human J $\kappa$ 1 gene  
 segments in *cis* at the same Ig locus in said genome;  
 5 a plurality of human J $\kappa$ 4 gene segments provided by at least 2 different human J $\kappa$ 1 gene  
 segments in *cis* at the same Ig locus in said genome;  
 a plurality of human J $\kappa$ 5 gene segments provided by at least 2 different human J $\kappa$ 1 gene  
 segments in *cis* at the same Ig locus in said genome;  
 a plurality of human J $\lambda$ 1 gene segments provided by at least 2 different human J $\lambda$ 1 gene  
 segments in *cis* at the same Ig locus in said genome;  
 10 a plurality of human J $\lambda$ 2 gene segments provided by at least 2 different human J $\lambda$ 2 gene  
 segments in *cis* at the same Ig locus in said genome;  
 a plurality of human J $\lambda$ 3 gene segments provided by at least 2 different human J $\lambda$ 3 gene  
 segments in *cis* at the same Ig locus in said genome;  
 a plurality of human J $\lambda$ 4 gene segments provided by at least 2 different human J $\lambda$ 4 gene  
 15 segments in *cis* at the same Ig locus in said genome;  
 a plurality of human J $\lambda$ 5 gene segments provided by at least 2 different human J $\lambda$ 5 gene  
 segments in *cis* at the same Ig locus in said genome;  
 a plurality of human J $\lambda$ 6 gene segments provided by at least 2 different human J $\lambda$ 6 gene  
 segments in *cis* at the same Ig locus in said genome; or  
 20 a plurality of human J $\lambda$ 7 gene segments provided by at least 2 different human J $\lambda$ 7 gene  
 segments in *cis* at the same Ig locus in said genome;

optionally wherein the JL gene segments are derived from the genome sequence of two or more  
 different human individuals.

25

242. A non-human vertebrate or vertebrate cell (optionally an ES cell or B-cell), according to claim  
 221, comprising a genome that comprises VL and JL gene repertoires comprising human gene  
 segments, the JL gene repertoire comprising

30 a plurality of human J $\kappa$ 1 gene segments provided by at least 3 different human J $\kappa$ 1 gene  
 segments;  
 a plurality of human J $\kappa$ 2 gene segments provided by at least 3 different human J $\kappa$ 1 gene  
 segments;  
 a plurality of human J $\kappa$ 3 gene segments provided by at least 3 different human J $\kappa$ 1 gene  
 35 segments;  
 a plurality of human J $\kappa$ 4 gene segments provided by at least 3 different human J $\kappa$ 1 gene  
 segments;  
 a plurality of human J $\kappa$ 5 gene segments provided by at least 3 different human J $\kappa$ 1 gene  
 segments;  
 40 a plurality of human J $\lambda$ 1 gene segments provided by at least 3 different human J $\lambda$ 1 gene  
 segments;  
 a plurality of human J $\lambda$ 2 gene segments provided by at least 3 different human J $\lambda$ 2 gene  
 segments;  
 a plurality of human J $\lambda$ 3 gene segments provided by at least 3 different human J $\lambda$ 3 gene  
 45 segments;  
 a plurality of human J $\lambda$ 4 gene segments provided by at least 3 different human J $\lambda$ 4 gene  
 segments;  
 a plurality of human J $\lambda$ 5 gene segments provided by at least 3 different human J $\lambda$ 5 gene  
 segments;  
 50 a plurality of human J $\lambda$ 6 gene segments provided by at least 3 different human J $\lambda$ 6 gene  
 segments; or

a plurality of human  $\lambda$ 7 gene segments provided by at least 3 different human  $\lambda$ 7 gene segments;

5 optionally wherein the JL gene segments are derived from the genome sequence of two or three different human individuals;

optionally wherein at least 2 or 3 of said different gene segments are provided in *cis* at the same Ig locus in said genome.

10 243. The vertebrate or cell of claim 241 or 242, wherein the different human individuals are from different human populations.

15 244. The vertebrate or cell of any one of claims 241 to 243, wherein the individuals are not genetically related.

245. The vertebrate or cell of any one of claims 241 to 244, wherein at least one of the different JL segments is a synthetic mutant of a human germline JL gene segment.

20 246. A non-human vertebrate or vertebrate cell (optionally an ES cell or B-cell) according to claim 225, comprising a genome comprising human VL and JL gene repertoires, the JL gene repertoire comprising

a plurality of human  $\kappa$ 1 gene segments provided by at least 2 different human  $\kappa$ 1 gene segments, optionally in *cis* at the same Ig locus in said genome;

25 a plurality of human  $\kappa$ 2 gene segments provided by at least 2 different human  $\kappa$ 1 gene segments, optionally in *cis* at the same Ig locus in said genome;

a plurality of human  $\kappa$ 3 gene segments provided by at least 2 different human  $\kappa$ 1 gene segments, optionally in *cis* at the same Ig locus in said genome;

30 a plurality of human  $\kappa$ 4 gene segments provided by at least 2 different human  $\kappa$ 1 gene segments, optionally in *cis* at the same Ig locus in said genome;

a plurality of human  $\kappa$ 5 gene segments provided by at least 2 different human  $\kappa$ 1 gene segments, optionally in *cis* at the same Ig locus in said genome;

a plurality of human  $\lambda$ 1 gene segments provided by at least 2 different human  $\lambda$ 1 gene segments, optionally in *cis* at the same Ig locus in said genome;

35 a plurality of human  $\lambda$ 2 gene segments provided by at least 2 different human  $\lambda$ 2 gene segments, optionally in *cis* at the same Ig locus in said genome;

a plurality of human  $\lambda$ 3 gene segments provided by at least 2 different human  $\lambda$ 3 gene segments, optionally in *cis* at the same Ig locus in said genome;

40 a plurality of human  $\lambda$ 4 gene segments provided by at least 2 different human  $\lambda$ 4 gene segments, optionally in *cis* at the same Ig locus in said genome;

a plurality of human  $\lambda$ 5 gene segments provided by at least 2 different human  $\lambda$ 5 gene segments, optionally in *cis* at the same Ig locus in said genome;

a plurality of human  $\lambda$ 6 gene segments provided by at least 2 different human  $\lambda$ 6 gene segments, optionally in *cis* at the same Ig locus in said genome; or

45 a plurality of human  $\lambda$ 7 gene segments provided by at least 2 different human  $\lambda$ 7 gene segments, optionally in *cis* at the same Ig locus in said genome;

50 wherein the JL gene segments are derived from the genome sequence of different human individuals that are not genetically related over at least 3 generations.

247. The vertebrate or cell of claim 246, wherein the human individuals are from different human populations.

248. A method of producing an antibody heavy chain, the method comprising

- 5
- (a) providing an antigen-specific heavy chain variable domain; and
  - (b) combining the variable domain with a human heavy chain constant region to produce an antibody heavy chain comprising (in N- to C-terminal direction) the variable domain and the constant region;
- 10

wherein

the human heavy chain constant region is an IGHAref, IGHA1a, IGHA2a, IGHA2b, IGHG1ref, IGHG2ref, IGHG2a, IGHG3ref, IGHG3a, IGHG3b, IGHG4ref, IGHG4a, IGHDref, IGHEref, IGHMref, IGHMa or IGHMb constant region.

15

249. The method of claim 248, wherein the variable domain is a human variable domain.

250. The method of any claim 248 or 249, wherein the variable domain has previously been selected from a non-human vertebrate that has been immunised with the antigen.

20

251. The method of any one of claims 248 to 250, comprising providing an expression vector comprising a nucleotide sequence encoding the constant region; inserting a nucleotide sequence encoding the variable domain into the vector 5' of the constant region sequence; inserting the vector into a host cell and expressing the heavy chain by the host cell; the method further comprising isolating a heavy chain (eg, as part of an antibody) comprising the variable domain and the human constant region.

25

252. The method of any one of claims 248 to 251, further comprising obtaining a nucleotide sequence encoding the heavy chain.

30

253. An antibody comprising a human heavy chain, the heavy chain comprising a variable domain that is specific for an antigen and a constant region that is an IGHAref, IGHA1a, IGHA2a, IGHA2b, IGHG1ref, IGHG2ref, IGHG2a, IGHG3ref, IGHG3a, IGHG3b, IGHG4ref, IGHG4a, IGHDref, IGHEref, IGHMref, IGHMa or IGHMb constant region.

35

254. A polypeptide comprising (in N- to C- terminal direction) a leader sequence, a human variable domain that is specific for an antigen and a human constant region that is an IGHAref, IGHA1a, IGHA2a, IGHA2b, IGHG1ref, IGHG2ref, IGHG2a, IGHG3ref, IGHG3a, IGHG3b, IGHG4ref, IGHG4a, IGHDref, IGHEref, IGHMref, IGHMa or IGHMb constant region; wherein (i) the leader sequence is not the native human variable domain leader sequence; and/or (ii) the variable domain comprises mouse AID-pattern somatic mutations or mouse terminal deoxynucleotidyl transferase (TdT)- pattern junctional mutations.

40

255. A nucleotide sequence encoding (in 5' to 3' direction) a leader sequence and a human antibody heavy chain, the heavy chain comprising a variable domain that is specific for an antigen and a constant region that is an IGHAref, IGHA1a, IGHA2a, IGHA2b, IGHG1ref, IGHG2ref, IGHG2a, IGHG3ref, IGHG3a, IGHG3b, IGHG4ref, IGHG4a, IGHDref, IGHEref, IGHMref, IGHMa or IGHMb constant region; and the leader sequence being operable for expression of the heavy chain and wherein the leader sequence is not the native human variable domain leader sequence.
256. A nucleotide sequence encoding (in 5' to 3' direction) a promoter and a human antibody heavy chain, the heavy chain comprising a variable domain that is specific for an antigen and a constant region that is an IGHAref, IGHA1a, IGHA2a, IGHA2b, IGHG1ref, IGHG2ref, IGHG2a, IGHG3ref, IGHG3a, IGHG3b, IGHG4ref, IGHG4a, IGHDref, IGHEref, IGHMref, IGHMa or IGHMb constant region; and the promoter being operable for expression of the heavy chain and wherein the promoter is not the native human promoter.
257. The antibody, polypeptide or nucleotide sequence of any one of claims 253 to 256, wherein the variable domain comprises mouse AID-pattern somatic mutations or mouse terminal deoxynucleotidyl transferase (TdT)- pattern junctional mutations.
258. A vector (eg, a CHO cell or HEK293 cell vector) comprising the nucleic acid of claim 255, 256 or 257; optionally wherein the vector is in a host cell (eg, a CHO cell or HEK293 cell).
259. A pharmaceutical composition comprising the antibody or polypeptide of any one of claims 253, 254 and 257, together with a pharmaceutically-acceptable excipient, diluent or a medicament (eg, a further antigen-specific variable domain, antibody chain or antibody).
260. The antibody or polypeptide of any one of claims 253, 254 and 257 for use in treating and/or preventing a medical condition in a human patient.
261. Use of the antibody or polypeptide of any one of claims 253, 254 and 257 for the manufacture of a medicament for treating and/or preventing a medical condition in a human patient.
262. The antibody, polypeptide or use of claim 260 or 261, wherein the human is a member of a human population selected from population numbers 1-14, wherein the populations are numbered as follows (population labels being according to 1000 Genomes Project nomenclature)
- 1= ASW;
  - 2= CEU;
  - 3=CHB;
  - 4=CHS;
  - 5=CLM;

6=FIN;  
 7=GBR;  
 8=IBS;  
 9=JPT;  
 10=LWK;  
 11=MXL;  
 12=PUR;  
 13=TSI;  
 14=YRI.

5

10

263. The antibody, polypeptide or use of claim 262, wherein the constant region is a

(i) IGHA1a constant region and the human population is selected from any population number 1-14;

15

(ii) IGHA2a constant region and the human population is selected from any population number 1-14;

(iii) IGHA2b constant region and the human population is selected from any population number 1-14;

20

(iv) IGHG2a constant region and the human population is selected from any population number 1-9 and 11-13;

(v) IGHG3a constant region and the human population is selected from any population number 1-14;

(vi) IGHG3b constant region and the human population is selected from any population number 1-8 and 11-13;

25

(vii) IGHG4a constant region and the human population is selected from any population number 1-9 and 11-13;

(viii)IGHMa constant region and the human population is selected from any population number 1-14; or

30

(ix) IGHMb constant region and the human population is selected from any population number 1-14;

Wherein the populations are numbered as follows (population labels being according to 1000 Genomes Project nomenclature)

1= ASW;

35

2= CEU;

3=CHB;

4=CHS;

5=CLM;

6=FIN;

40

7=GBR;

8=IBS;

9=JPT;

10=LWK;

11=MXL;

45

12=PUR;

13=TSI;

14=YRI.

5 265. A vector (eg, a CHO cell or HEK293 cell vector) comprising a IGHG1ref, IGHG2ref, IGHG2a,  
IGHG3ref, IGHG3a, IGHG3b, IGHG4ref or IGHG4a constant region nucleotide sequence that is 3'  
of a cloning site for the insertion of a human antibody heavy chain variable domain nucleotide  
sequence, such that upon insertion of such a variable domain sequence the vector comprises (in  
5' to 3' direction) a promoter, a leader sequence, the variable domain sequence and the  
10 constant region sequence so that the vector is capable of expressing a human antibody heavy  
chain when present in a host cell.

15 A non-human vertebrate or vertebrate cell (optionally an ES cell or antibody-producing cell)  
comprising a genome having a superhuman immunoglobulin heavy chain human VH and/or D  
and/or J gene repertoire.

20 266. The vertebrate or cell of claim 265, wherein the genome comprises a transgenic  
immunoglobulin heavy chain locus comprising a plurality of human immunoglobulin VH gene  
segments, one or more human D gene segments and one or more human J gene segments,  
wherein the plurality of VH gene segments consists of more than the natural human repertoire  
of functional VH gene segments; optionally wherein the genome is homozygous for said  
transgenic heavy chain locus.

25 267. The vertebrate or cell of claim 266, wherein VH gene repertoire consists of a plurality of VH  
gene segments derived from the genome sequence of a first human individual, supplemented  
with one or more different VH gene segments derived from the genome sequence of a second,  
different human individual; optionally wherein the individuals are not related; optionally  
wherein the D and J segments are derived from the genome sequence of the first human  
individual; and optionally wherein the VH gene segments from the genome sequence of the  
30 second individual are selected from the VH gene segments listed in Table 1.

35 268. The vertebrate or cell of claim 267, wherein the transgenic locus comprises more than 26  
functional human VH gene segments; optionally wherein the locus comprises at least  
27 or 28 functional human VH gene segments (eg, wherein the locus comprises the full functional  
VH repertoire of said first individual supplemented with one or more VH gene segments derived  
from the genome sequence of the second human individual and optionally with one or more VH  
gene segments derived from the genome sequence of a third human individual).

40 269. The vertebrate of cell of claim 267 or 268, wherein the transgenic locus comprises a first VH  
gene segment derived from the genome sequence of the first individual and a second VH gene  
segment derived from the genome sequence of the second individual, wherein the second VH  
gene segment is a polymorphic variant of the first VH gene segment; optionally wherein the  
locus comprises a further polymorphic variant of the first VH gene segment (eg, a variant derived  
from the genome sequence of a third human individual).



270. The vertebrate or cell of any one of claims 265 to 269, wherein the genome comprises a (or said) transgenic immunoglobulin heavy chain locus comprising a plurality of human immunoglobulin VH gene segments, a plurality of human D gene segments and one or more human J gene segments, wherein the plurality of D gene segments consists of more than the natural human repertoire of functional D gene segments; optionally wherein the genome is homozygous for said transgenic heavy chain locus.
271. The vertebrate or cell of claim 270, wherein D gene repertoire consists of a plurality of D gene segments derived from the genome sequence of a (or said) first human individual, supplemented with one or more different D gene segments derived from the genome sequence of a (or said) second, different human individual; optionally wherein the individuals are not related; optionally wherein J segments are derived from the genome sequence of the first human individual; and optionally wherein the D gene segments from the genome sequence of the second individual are selected from the D gene segments listed in Table 2.
272. The vertebrate or cell of claim 271, wherein the transgenic locus comprises more than 26 functional human D gene segments; optionally wherein the locus comprises at least 27 or 28 functional human D gene segments (eg, wherein the locus comprises the full functional D repertoire of said first individual supplemented with one or more D gene segments derived from the genome sequence of the second human individual and optionally with one or more D gene segments derived from the genome sequence of a third human individual).
273. The vertebrate or cell of claim 271 or 272, wherein the transgenic locus comprises a first D gene segment derived from the genome sequence of the first individual and a second D gene segment derived from the genome sequence of the second individual, wherein the second D gene segment is a polymorphic variant of the first D gene segment; optionally wherein the locus comprises a further polymorphic variant of the first D gene segment (eg, a variant derived from the genome sequence of a third human individual).
274. The vertebrate or cell of any one of claims 265 to 273, wherein the genome comprises a (or said) transgenic immunoglobulin heavy chain locus comprising a plurality of human immunoglobulin VH gene segments, one or more human D gene segments and a plurality of human JH gene segments, wherein the plurality of J gene segments consists of more than the natural human repertoire of functional J gene segments; optionally wherein the genome is homozygous for said transgenic heavy chain locus.
275. The vertebrate or cell of claim 274, wherein the J gene repertoire consists of a plurality of J gene segments derived from the genome sequence of a (or said) first human individual, supplemented with one or more different J gene segments derived from the genome sequence of a (or said) second, different human individual; optionally wherein the individuals are not related; optionally wherein D segments are derived from the genome sequence of the first human individual; and optionally wherein the J gene segments from the genome sequence of the second individual are selected from the J gene segments listed in Table 3.

276. The vertebrate or cell of claim 275, wherein the transgenic locus comprises more than 26 functional human J gene segments; optionally wherein the locus comprises at least 27 or 28 functional human J gene segments (eg, wherein the locus comprises the full functional J repertoire of said first individual supplemented with one or more J gene segments derived from the genome sequence of the second human individual and optionally with one or more J gene segments derived from the genome sequence of a third human individual).
277. The vertebrate of cell of claim 275 or 276, wherein the transgenic locus comprises a first J gene segment derived from the genome sequence of the first individual and a second J gene segment derived from the genome sequence of the second individual, wherein the second J gene segment is a polymorphic variant of the first J gene segment; optionally wherein the locus comprises a further polymorphic variant of the first J gene segment (eg, a variant derived from the genome sequence of a third human individual).
278. A non-human vertebrate or vertebrate cell (optionally an ES cell or antibody-producing cell) comprising a genome having a superhuman immunoglobulin light chain human VL gene repertoire; optionally wherein the vertebrate or cell is according to any one of claims 265 to 277.
279. The vertebrate or cell of claim 278, wherein the genome comprises
- (i) a transgenic immunoglobulin kappa light chain locus comprising a plurality of human immunoglobulin Vk gene segments and one or more human J gene segments, wherein the plurality of Vk gene segments consists of more than the natural human repertoire of functional Vk gene segments; optionally wherein the genome is homozygous for said transgenic kappa light chain locus; and/or
  - (ii) a transgenic immunoglobulin lambda light chain locus comprising a plurality of human immunoglobulin Vλ gene segments and one or more human J gene segments, wherein the plurality of Vλ gene segments consists of more than the natural human repertoire of functional Vλ gene segments; optionally wherein the genome is homozygous for said transgenic lambda light chain locus.
280. The vertebrate or cell of claim 279, wherein
- (i) the Vk gene repertoire consists of a plurality of Vk gene segments derived from the genome sequence of a first human individual, supplemented with one or more Vk gene segments derived from the genome sequence of a second, different human individual; optionally wherein the individuals are not related; optionally wherein the J segments are derived from the genome sequence of the first human individual; and optionally wherein the Vk gene segments from the genome sequence of the second individual are selected from the Vk gene segments listed in Table 4; and
  - (i) the Vλ gene repertoire consists of a plurality of Vλ gene segments derived from the genome sequence of a first human individual, supplemented with one or more Vλ gene segments derived from the genome sequence of a second, different human individual; optionally wherein the

individuals are not related; optionally wherein the J segments are derived from the genome sequence of the first human individual; and optionally wherein the V $\lambda$  gene segments from the genome sequence of the second individual are selected from the V $\lambda$  gene segments listed in Table 5.

5

281. The vertebrate or cell of claim 280, wherein

the kappa light transgenic locus comprises more than 26 functional human V $\kappa$  gene segments; optionally wherein the locus comprises at least 27 or 28 functional human V $\kappa$  gene segments (eg, wherein the locus comprises the full functional V $\kappa$  repertoire of said first individual supplemented with one or more V $\kappa$  gene segments derived from the genome sequence of the second human individual and optionally with one or more V $\kappa$  gene segments derived from the genome sequence of a third human individual); and

10

the lambda light transgenic locus comprises more than 26 functional human V $\lambda$  gene segments; optionally wherein the locus comprises at least 27 or 28 functional human V $\lambda$  gene segments (eg, wherein the locus comprises the full functional V $\lambda$  repertoire of said first individual supplemented with one or more V $\lambda$  gene segments derived from the genome sequence of the second human individual and optionally with one or more V $\lambda$  gene segments derived from the genome sequence of a third human individual);

15

20

282. The vertebrate of cell of claim 280 or 281, wherein

the kappa light transgenic locus comprises a first V $\kappa$  gene segment derived from the genome sequence of the first individual and a second V $\kappa$  gene segment derived from the genome sequence of the second individual, wherein the second V $\kappa$  gene segment is a polymorphic variant of the first V $\kappa$  gene segment; optionally wherein the locus comprises a further polymorphic variant of the first V $\kappa$  gene segment (eg, a variant derived from the genome sequence of a third human individual); and

25

30

the lambda light transgenic locus comprises a first V $\lambda$  gene segment derived from the genome sequence of the first individual and a second V $\lambda$  gene segment derived from the genome sequence of the second individual, wherein the second V $\lambda$  gene segment is a polymorphic variant of the first V $\lambda$  gene segment; optionally wherein the locus comprises a further polymorphic variant of the first V $\lambda$  gene segment (eg, a variant derived from the genome sequence of a third human individual).

35

283. The vertebrate or cell of any one of claims 278 to 282, wherein the genome comprises a (or said) transgenic immunoglobulin light chain locus comprising a plurality of human immunoglobulin VL gene segments and a plurality of human JL gene segments, wherein the plurality of J gene segments consists of more than the natural human repertoire of functional J gene segments; optionally wherein the genome is homozygous for said transgenic heavy chain locus.

40

284. The vertebrate or cell of claim 283, wherein

5 (i) the J<sub>k</sub> gene repertoire consists of a plurality of J<sub>k</sub> gene segments derived from the genome sequence of a (or said) first human individual, supplemented with one or more J<sub>k</sub> gene segments derived from the genome sequence of a (or said) second, different human individual; optionally wherein the individuals are not related; optionally wherein the V<sub>k</sub> segments are derived from the genome sequence of the first human individual; optionally wherein the J<sub>k</sub> gene segments from the genome sequence of the second individual are selected from the J<sub>k</sub> gene segments listed in Table 6; and

10 (ii) the J<sub>k</sub> gene repertoire consists of a plurality of J<sub>λ</sub> gene segments derived from the genome sequence of a (or said) first human individual, supplemented with one or more J<sub>λ</sub> gene segments derived from the genome sequence of a (or said) second, different human individual; optionally wherein the individuals are not related; optionally wherein the V<sub>λ</sub> segments are derived from the genome sequence of the first human individual; optionally wherein the J<sub>λ</sub> gene segments from the genome sequence of the second individual are selected from the J<sub>λ</sub> gene segments listed in Table 7.

285. The vertebrate or cell of claim 284, wherein

20 (i) the transgenic light chain locus comprises more than 26 functional human J<sub>k</sub> gene segments; optionally wherein the locus comprises at least 27 or 28 functional human J<sub>k</sub> gene segments (eg, wherein the locus comprises the full functional J<sub>k</sub> repertoire of said first individual supplemented with one or more J<sub>k</sub> gene segments derived from the genome sequence of the second human individual and optionally with one or more J<sub>k</sub> gene segments derived from the genome sequence of a third human individual); and/or

30 (i) the transgenic light chain locus comprises more than 26 functional human J<sub>λ</sub> gene segments; optionally wherein the locus comprises at least 27 or 28 functional human J<sub>λ</sub> gene segments (eg, wherein the locus comprises the full functional J<sub>λ</sub> repertoire of said first individual supplemented with one or more J<sub>λ</sub> gene segments derived from the genome sequence of the second human individual and optionally with one or more J<sub>λ</sub> gene segments derived from the genome sequence of a third human individual).

35 286. The vertebrate or cell of claim 284 or 285, wherein

40 (i) the kappa light transgenic locus comprises a first J<sub>k</sub> gene segment derived from the genome sequence of the first individual and a second J<sub>k</sub> gene segment derived from the genome sequence of the second individual, wherein the second J<sub>k</sub> gene segment is a polymorphic variant of the first J<sub>k</sub> gene segment; optionally wherein the locus comprises a further polymorphic variant of the first J<sub>k</sub> gene segment (eg, a variant derived from the genome sequence of a third human individual); and

45 (ii) the lambda light transgenic locus comprises a first J<sub>λ</sub> gene segment derived from the genome sequence of the first individual and a second J<sub>λ</sub> gene segment derived from the genome

sequence of the second individual, wherein the second Jk gene segment is a polymorphic variant of the first Jλ gene segment; optionally wherein the locus comprises a further polymorphic variant of the first Jλ gene segment (eg, a variant derived from the genome sequence of a third human individual).

5

287. A library of antibody-producing transgenic cells whose genomes collectively encode a repertoire of antibodies, wherein

10

(a) a first transgenic cell expresses a first antibody having a chain encoded by a first immunoglobulin gene, the gene comprising a first variable domain nucleotide sequence produced following recombination of a first human unrearranged immunoglobulin gene segment;

15

(b) a second transgenic cell expresses a second antibody having a chain encoded by a second immunoglobulin gene, the second gene comprising a second variable domain nucleotide sequence produced following recombination of a second human unrearranged immunoglobulin gene segment, the first and second antibodies being non-identical;

20

(c) the first and second gene segments are different and derived from the genome sequences of first and second human individuals respectively, wherein the individuals are different; and optionally not related;

(d) wherein the cells are non-human vertebrate (eg, mouse or rat) cells.

25

288. The library of claim 23, wherein in step (c) the individuals are not related.

30

289. The vertebrate, cell or library of any one of claims 265 to 288, wherein the first and second human individuals are members of first and second ethnic populations respectively, wherein the populations are different; optionally wherein the ethnic populations are selected from those identified in the 1000 genomes database.

35

290. The vertebrate, cell or library of claim 289, wherein the human immunoglobulin gene segment derived from the genome sequence of the second individual is low-frequency (optionally rare) within the second ethnic population.

40

291. The library of claim 289 or 290, wherein the first variable region nucleotide sequence is produced by recombination of the first human immunoglobulin gene segment with a first J gene segment and optionally a first D gene segment, wherein the first human immunoglobulin gene segment is a V gene segment and the V, D and J segments are derived from the first human population, optionally from the genome of one individual of the first human population.

45

292. The library of claim 291, wherein the second variable region nucleotide sequence is produced by recombination of the second human immunoglobulin gene segment with a second J gene segment and optionally a second D gene segment, wherein the second human immunoglobulin gene segment is a V gene segment derived from the second population and the

D and/or J segments are derived from the first human population, optionally the D and J gene segments being from the genome of one individual of the first human population.

- 5 293. The library of claim 292, wherein all of the D and J segments that have been recombined with the first and second V gene segments are D and J segments derived from the first human population, optionally the D and J gene segments being from the genome of one individual of the first human population.
- 10 294. The vertebrate, cell or library of any one of claims 289 to 293, wherein the first and second ethnic populations are selected from the group consisting of an ethnic population with European ancestry, an ethnic population with East Asian, an ethnic population with West African ancestry, an ethnic population with Americas ancestry and an ethnic population with South Asian ancestry.
- 15 295. The vertebrate, cell or library of claim 294, wherein the first and second ethnic populations are selected from the group consisting of an ethnic population with Northern European ancestry; or an ethnic population with Western European ancestry; or an ethnic population with Toscani ancestry; or an ethnic population with British ancestry; or an ethnic population with Icelandic ancestry; or an ethnic population with Finnish ancestry; or an ethnic population with Iberian ancestry; or an ethnic population with Japanese ancestry; or an ethnic population with Chinese ancestry; or an ethnic population Vietnamese ancestry; or an ethnic population with Yoruba ancestry; or an ethnic population with Luhya ancestry; or an ethnic population with Gambian ancestry; or an ethnic population with Malawian ancestry; or an ethnic population with Native American ancestry; or an ethnic population with Afro-Caribbean ancestry; or an ethnic population with Mexican ancestry; or an ethnic population with Puerto Rican ancestry; or an ethnic population with Columbian ancestry; or an ethnic population with Peruvian ancestry; or an ethnic population with Ahom ancestry; or an ethnic population with Kayadtha ancestry; or an ethnic population with Reddy ancestry; or an ethnic population with Maratha; or an ethnic population with Punjabi ancestry.
- 20 296. The library of claim 287, wherein the second human immunoglobulin gene segment is a polymorphic variant of the first human immunoglobulin gene segment; optionally wherein the second gene segment is selected from the group consisting of a gene segment in any of Tables 1 to 7 and 9 to 14, eg, the second gene segment is a polymorphic variant of VH1-69.
- 25 297. The library of any one of claims 287 to 296, wherein the first and second human immunoglobulin gene segments are both (i) V<sub>H</sub> gene segments; (ii) D segments; (iii) J segments (optionally both J<sub>H</sub> segments, both J<sub>K</sub> segments or both J<sub>λ</sub> segments); (iv) constant regions (optionally both a gamma constant region, optionally both a C gamma-1 constant region); (v) CH1 regions; (vi) CH2 regions; or (vii) CH3 regions.
- 30 298. The library of any one of claims 287 to 297, wherein the library is naive and optionally has a library size of from 10<sup>2</sup> to 10<sup>9</sup> cells.

299. The library of any one of claims 287 to 298, wherein the library has been selected against a predetermined antigen optionally has a library size of from  $10^2$  to  $10^9$  cells.
- 5 300. The library of any one of claims 287 to 299, wherein said first and second cells are progeny of first and second ancestor non-human vertebrate cells respectively, wherein the first ancestor cell comprises a genome comprising said first human immunoglobulin gene segment; and the second ancestor cell comprises a genome comprising said second human immunoglobulin gene segment.
- 10 301. A library of antibody-producing transgenic cells whose genomes collectively encode a repertoire of antibodies, wherein the library comprises the first and second ancestor cells recited in claim 300.
- 15 302. A library of hybridoma cells produced by fusion of the library of any one of claims 265 to 301 with fusion partner cells; and optionally the hybridoma library has a size of from  $10^2$  to  $10^9$  cells
303. An isolated antibody having
- 20 (a) a heavy chain encoded by a nucleotide sequence produced following recombination in a transgenic non-human vertebrate cell of an unrearranged human immunoglobulin V gene segment with a human D and human J segment, optionally with affinity maturation in said cell, wherein one of the gene segments is derived from the genome of an individual from a first human ethnic population; and the other two gene segments are derived from the genome of an individual from a second, different, human ethnic population, and wherein the antibody
- 25 comprises heavy chain constant regions of said non-human vertebrate (eg, rodent, mouse or rat heavy chain constant regions); and/or
- 30 (b) a light chain encoded by a nucleotide sequence produced following recombination in a transgenic non-human vertebrate cell of an unrearranged human immunoglobulin V gene segment with a human J segment, optionally with affinity maturation in said cell, wherein one of the gene segments is derived from the genome of an individual from a first human ethnic population (optionally the same as the first population in (a)); and the other gene segment is derived from the genome of an individual from a second, different, human ethnic population (optionally the same as the second population in (a)), and wherein the antibody comprises light
- 35 chain constant regions of said non-human vertebrate (eg, rodent, mouse or rat heavy light constant regions);
- (c) Optionally wherein each variable domain of the antibody is a human variable domain.
- 40 (d) Optionally wherein the heavy chain constant regions are gamma-type constant regions.
304. An isolated nucleotide sequence encoding the antibody of claim 303, optionally wherein the sequence is provided in an antibody expression vector, optionally in a host cell.

305. A method of producing a human antibody, the method comprising replacing the non-human vertebrate constant regions of the antibody of claim 303 with human antibody constant regions.
306. A pharmaceutical composition comprising an antibody according to claim 303, or an antibody produced according to the method of claim 305 and a diluent, excipient or carrier; optionally wherein the composition is provided in a container connected to an IV needle or syringe or in an IV bag.
307. An antibody-producing cell that expresses the second antibody recited in any one of claims 287 to 302 or the isolated antibody of claim 303.
308. A non-human vertebrate or vertebrate cell (optionally an ES cell or antibody-producing cell) whose genome comprises a transgenic immunoglobulin locus (eg, a heavy chain locus or a light chain locus), said locus comprising immunoglobulin gene segments according to the first and second human immunoglobulin gene segments (optionally V segments) recited in any one of claims 287 to 302 operably connected upstream of an immunoglobulin constant region; optionally wherein the genome is homozygous for said transgenic immunoglobulin locus; optionally wherein the immunoglobulin locus comprises more than the natural human complement of functional V gene segments; and/or optionally wherein the immunoglobulin locus comprises more than the natural human complement of functional D gene segments; and/or optionally wherein the immunoglobulin locus comprises more than the natural human complement of functional J gene segments.
309. A transgenic non-human vertebrate (eg, a mouse or rat) or vertebrate cell (optionally an ES cell or antibody-producing cell) whose genome comprises a transgenic immunoglobulin locus comprising a plurality of human immunoglobulin gene segments operably connected upstream of a non-human vertebrate constant region for the production of a repertoire of chimaeric antibodies, or chimaeric light or heavy chains, having a non-human vertebrate constant region and a human variable region; wherein the transgenic locus comprises one or more human immunoglobulin V gene segments, one or more human J gene segments and optionally one or more human D gene segments, a first (optionally a V segment) of said gene segments and a second (optionally a V segment) of said gene segments being different and derived from the genomes of first and second human individuals respectively, wherein the individuals are different; and optionally not related; optionally wherein the immunoglobulin locus comprises more than the natural human complement of functional V gene segments; and/or optionally wherein the immunoglobulin locus comprises more than the natural human complement of functional D gene segments; and/or



optionally wherein the immunoglobulin locus comprises more than the natural human complement of functional J gene segments.

5

310. A transgenic non-human vertebrate (eg, a mouse or rat) or vertebrate cell (optionally an ES cell or antibody-producing cell) whose genome comprises first and second transgenic immunoglobulin loci, each locus comprising a plurality of human immunoglobulin gene segments operably connected upstream of a non-human vertebrate constant region for the production of a repertoire of chimaeric antibodies, or chimaeric light or heavy chains, having a non-human vertebrate constant region and a human variable region;

10

wherein (i) the first transgenic locus comprises one or more human immunoglobulin V gene segments, one or more human J gene segments and optionally one or more human D gene segments, (ii) the second transgenic locus comprises one or more human immunoglobulin V gene segments, one or more human J gene segments and optionally one or more human D gene segments; and (iii) wherein a first (optionally a V) gene segment of said first locus and a second (optionally a V) gene segment of said second gene locus are different and derived from the genomes of first and second human individuals respectively, wherein the individuals are different; and optionally not related

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optionally wherein the first and second loci are on different chromosomes (optionally chromosomes with the same chromosome number) in said genome;

25

optionally wherein each immunoglobulin locus comprises more than the natural human complement of functional V gene segments; and/or

optionally wherein each immunoglobulin locus comprises more than the natural human complement of functional D gene segments; and/or

30

optionally wherein each immunoglobulin locus comprises more than the natural human complement of functional J gene segments.

35

311. The non-human vertebrate or cell of claim 309 or 310, wherein the immunoglobulin gene segments are as recited in any one of claims 23 to 37.

40

312. The non-human vertebrate or cell of claim 308, 309, 310 or 311, wherein the genome comprises a third immunoglobulin gene segment (optionally a V segment), the third gene segment being derived from a human individual that is different from the individual from which the first (and optionally also the second) gene segment is derived; optionally wherein the first, second and third gene segments are polymorphic variants of a human immunoglobulin gene segment.

45

313. The non-human vertebrate or cell of any one of claims 308 to 312, wherein the genome of the vertebrate or cell is homozygous for the first, second and optional third gene segment,

wherein a copy of the first, second and optional third gene segments are provided together on the same chromosome operably connected upstream of a common non-human vertebrate constant region.

- 5 314. The non-human vertebrate or cell of any one of claims 308 to 313, wherein each first, second and optional third gene segment is a V gene segment.
315. The library of any one of claims 287 to 302, wherein the library is provided by a collection of non-human vertebrates (optionally a collection of rodents, mice or rats); optionally, wherein a first member of said collection produces said first antibody but not said second antibody, and a second member of the collection produces said second antibody (but optionally not said first antibody).
- 10 316. A repertoire of antibodies expressed from a library of cells according to any one of claims 287 to 302 and 315.
- 15 317. A method of constructing a cell (eg, an ES cell) according to any one of claims 265 to 286, the method comprising
- 20 (a) identifying functional V and J (and optionally D) gene segments of the genome sequence of a (or said) first human individual;
- (b) identifying one or more functional V and/or D and/or J gene segments of the genome sequence of a (or said) second human individual, wherein these additional gene segments are not found in the genome sequence of the first individual;
- 25 (c) and constructing a transgenic immunoglobulin locus in the cell, wherein the gene segments of (a) and (b) are provided in the locus operably connected upstream of a constant region.
318. The method of claim 317, wherein the cell comprises a heavy chain locus constructed according to steps (a) to (c) and/or a light chain locus (kappa and/or lambda loci) constructed according to steps (a) to (c).
- 30 319. The method of claim 317 or 318, wherein the cell is homozygous for the or each transgenic locus; optionally wherein antibody expression from loci endogenous to said cell has been inactivated.
- 35 320. The method of any one of claims 317 to 319, wherein the gene segment(s) in step (b) are identified from an immunoglobulin gene database selected from the 1000 genomes, Ensembl, Genbank and IMGT databases.
- 40 321. The method of any one of claims 317 to 320, wherein the first and second human individuals are members of first and second ethnic populations respectively, wherein the populations are different, optionally wherein the human immunoglobulin gene segment derived from the genome sequence of the second individual is low-frequency (optionally rare) within the second ethnic population.

322. A method of making a transgenic non-human vertebrate (eg, a mouse or rat), the method comprising
- (a) constructing an ES cell (eg, a mouse C57BL/6N, C57BL/6J, 129S5 or 129Sv strain ES cell) by carrying out the method of any one of claims 317 to 321;
  - 5 (b) injecting the ES cell into a donor non-human vertebrate blastocyst (eg, a mouse C57BL/6N, C57BL/6J, 129S5 or 129Sv strain blastocyst);
  - (c) implanting the blastocyst into a foster non-human vertebrate mother (eg, a C57BL/6N, C57BL/6J, 129S5 or 129Sv strain mouse); and
  - 10 (d) obtaining a child from said mother, wherein the child genome comprises a transgenic immunoglobulin locus.
323. A transgenic non-human vertebrate (eg, a mouse or rat) made by the method of claim 322 or a progeny thereof.
- 15 324. A population of non-human vertebrates according to claim 323.
325. A method of isolating an antibody that binds a predetermined antigen (eg, a bacterial or viral pathogen antigen), the method comprising
- 20 (a) providing a vertebrate (optionally a mouse or rat) according to any one of claims 265 to 286, 289, 290, 294, 295, 308 to 314 and 323;
  - (b) immunising (eg, using a standard prime-boost method) said vertebrate with said antigen (optionally wherein the antigen is an antigen of an infectious disease pathogen);
  - (c) removing B lymphocytes from the vertebrate and selecting one or more B lymphocytes expressing antibodies that bind to the antigen;
  - 25 (d) optionally immortalising said selected B lymphocytes or progeny thereof, optionally by producing hybridomas therefrom; and
  - (e) isolating an antibody (eg, and IgG-type antibody) expressed by the B lymphocytes; and
  - (f) optionally producing a derivative or variant of the antibody.
- 30 326. The method of claim 325, further comprising after step (e) the step of isolating from said B lymphocytes nucleic acid encoding said antibody that binds said antigen; optionally exchanging the heavy chain constant region nucleotide sequence of the antibody with a nucleotide sequence encoding a human or humanised heavy chain constant region and optionally affinity maturing the variable region of said antibody; and optionally inserting said nucleic acid into an expression vector and optionally a host.
- 35 327. A non-human vertebrate (eg, a mouse or rat) or cell (eg, a mouse cell or rat cell) comprising a genome that comprises a transgenic heavy chain immunoglobulin locus, wherein the locus comprises at least 42 (optionally at least 43, 44, 45, 46, 47, 48, 49, 50) functional human VH gene segments, one or more functional human D gene segments, one or more functional human JH gene segments operably connected upstream of a non-human vertebrate constant region (eg, a mouse constant region, eg, a Cmu and/or a C gamma).
- 40 328. The vertebrate or cell of claim 327, wherein the locus comprises at least 23 functional human D gene segments and/or at least 6 functional human JH gene segments, optionally at
- 45

least a full human repertoire of functional VH, D and JH segments (optionally derived from the genome sequence of the same human individual).

- 5 329. A non-human vertebrate (eg, a mouse or rat) or cell (eg, a mouse cell or rat cell) comprising a genome that comprises a transgenic heavy chain immunoglobulin locus, wherein the locus comprises a full human repertoire of functional VH, D and JH segments derived from the genome sequence of the same human individual, supplemented with one or more additional functional human VH, D and/or JH gene segment that is not found in the genome sequence of said human individual.
- 10 330. The vertebrate or cell of claim 329, wherein each additional VH, D and JH segment is selected from the group consisting of (i) a gene segment derived from a second, different individual, optionally wherein the first and second individuals are members of different ethnic populations; (ii) a mutant of a human gene segment (optionally a human germline gene segment having up to 5, 10 or 15 mutations); and (iii) a hybrid human gene segment comprising a first and second nucleotide sequences derived from genome sequences of different human individuals or different polymorphic variants of a human immunoglobulin gene segment (eg, variants of a germline human VH, D or JH gene segment).
- 15 331. The vertebrate or cell of any one of claims 328 to 330, wherein the genome is homozygous for said heavy chain transgene and optionally endogenous immunoglobulin heavy chain expression has been inactivated.
- 20 332. A non-human vertebrate (eg, a mouse or rat) or cell (eg, a mouse cell or rat cell) comprising a genome that comprises a transgenic kappa light chain immunoglobulin locus, wherein the locus comprises at least 39 functional human Vk gene segments and one or more functional human Jk gene segments operably connected upstream of a non-human vertebrate constant region (eg, a mouse constant region).
- 25 333. The vertebrate or cell of claim 332, wherein the locus comprises at least a full human repertoire of functional Vk and Jk segments (optionally derived from the genome sequence of the same human individual).
- 30 334. A non-human vertebrate (eg, a mouse or rat) or cell (eg, a mouse cell or rat cell) comprising a genome that comprises a transgenic kappa light chain immunoglobulin locus, wherein the locus comprises a full human repertoire of functional Vk and Jk segments derived from the genome sequence of the same human individual, supplemented with one or more additional functional human Vk and/or Jk gene segment that is not found in the genome sequence of said human individual.
- 35 335. The vertebrate or cell of claim 334, wherein each additional Vk and Jk segment is selected from the group consisting of (i) a gene segment derived from a second, different individual, optionally wherein the first and second individuals are members of different ethnic populations; (ii) a mutant of a human gene segment (optionally a human germline gene segment having up to 5, 10 or 15 mutations); and (iii) a hybrid human gene segment comprising a first and second nucleotide sequences derived from: genome sequences of different human individuals or
- 40 45

different polymorphic variants of a human immunoglobulin gene segment (eg, variants of a germline human Vk or Jk gene segment).

- 5 336. The vertebrate or cell of any one of claims 332 to 335, wherein the genome is homozygous for said kappa light chain transgene and optionally endogenous immunoglobulin kappa light chain expression has been inactivated.
- 10 337. A non-human vertebrate (eg, a mouse or rat) or cell (eg, a mouse cell or rat cell) comprising a genome that comprises a transgenic lambda light chain immunoglobulin locus, wherein the locus comprises at least 32 functional human V lambda gene segments and one or more functional human J lambda gene segments operably connected upstream of a non-human vertebrate constant region (eg, a mouse constant region).
- 15 338. The vertebrate or cell of claim 337, wherein the locus comprises at least a full human repertoire of functional V lambda and J lambda segments (optionally derived from the genome sequence of the same human individual).
- 20 339. A non-human vertebrate (eg, a mouse or rat) or cell (eg, a mouse cell or rat cell) comprising a genome that comprises a transgenic lambda light chain immunoglobulin locus, wherein the locus comprises a full human repertoire of functional V lambda and J lambda segments derived from the genome sequence of the same human individual, supplemented with one or more additional functional human V lambda and/or J lambda gene segment that is not found in the genome sequence of said human individual.
- 25 340. The vertebrate or cell of claim 339, wherein each additional V lambda and J lambda segment is selected from the group consisting of (i) a gene segment derived from a second, different individual, optionally wherein the first and second individuals are members of different ethnic populations; (ii) a mutant of a human gene segment (optionally a human germline gene segment having up to 5, 10 or 15 mutations); and (iii) a hybrid human gene segment comprising a first and second nucleotide sequences derived from genome sequences of different human individuals or different polymorphic variants of a human immunoglobulin gene segment (eg, variants of a germline human V lambda or J lambda gene segment).
- 30 341. The vertebrate or cell of any one of claims 337 to 340, wherein the genome is homozygous for said lambda light chain transgene and optionally endogenous immunoglobulin lambda light chain expression has been inactivated.
- 35 342. A transgenic immunoglobulin locus comprising a synthetic immunoglobulin gene haplotype, the haplotype comprising first and second human gene segments (each being a V, D or J), a switch region and a constant region, wherein
- 40 a) the second gene segment is a polymorphic variant of the first gene segment; or  
b) the first and second gene segments are derived respectively from genome sequence of individuals from different, first and second, ethnic populations (eg, according to the 1000 Genomes database) and the second gene segment is not found in the first population (eg,
- 45

according to the 1000 Genomes database), optionally wherein the second gene segment occurs at low-frequency (or is rare) in the second population; and

5           Wherein the constant region, and optionally the switch region, are non-human vertebrate (eg, mouse or rat) constant and switch regions (eg, mouse or rat Cmu; mouse or rat Smu; mouse or rat C gamma; or mouse or rat S gamma).

10       343.     The transgenic immunoglobulin locus of claim 342 wherein the transgene comprises a third human immunoglobulin gene segment (a V, D or J) which is (a) a polymorphic variant of the first and second gene segments; or (b) derived from a genome sequence of the second population (and optionally is a low-frequency or rare gene in that population) or is from a third human ethnic population, wherein the third gene segment is not found in the first population (eg, according to the 1000 Genomes database).

15       344.     A non-human vertebrate or cell comprising the transgene of claim 342 or 343; optionally wherein the vertebrate or cell is homozygous for said transgene.

20       345.     An isolated antibody for administration to a Chinese patient, the antibody comprising a human heavy chain, the heavy chain comprising a variable domain that is specific for an antigen and a constant region, wherein the constant region is a human constant region selected from a constant region (eg, an IGHG constant region) in Table 13 found in a Chinese population and with a cumulative frequency of at least 5%;, and wherein

25           (i) the variable domain is derived from the recombination of said human gene segments in a non-human vertebrate (eg, in a mouse or a rat); or (ii) the variable domain comprises non-human vertebrate (eg, mouse or rat) AID-pattern mutations and non-human vertebrate (eg, mouse or rat) terminal deoxynucleotidyl transferase (TdT)-pattern mutations.

30       346.     The antibody of claim 345 wherein the constant region is a IGHG1a, IGHG2a, IGHG3a, IGHG3b or IGHG4a constant region.

35       347.     The antibody of claim 345 or 346, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the VH gene segment being selected from a VH in Table 13 found in a Chinese population and with a cumulative frequency of at least 5%.

40       348.     The antibody of claim 345, 346 or 347, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the D gene segment being selected from a D in Table 13 found in a Chinese population and with a cumulative frequency of at least 5%.

45       349.     The antibody of claim 345, 346, 347 or 348 wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the JH gene segment being selected from a JH in Table 13 found in

a Chinese population and with a cumulative frequency of at least 5%.

350. An isolated VH domain identical to a variable domain as recited in any one of claims 347 to 349, optionally fused at its C-terminus to a polypeptide (eg, an antibody Fc).

5

351. A pharmaceutical composition comprising the antibody or variable domain of any one of claims 345 to 350 together with a pharmaceutically-acceptable excipient, diluent or a medicament (eg, a further antigen-specific variable domain, antibody chain or antibody).

10

352. An isolated antibody for administration to a Chinese patient, the antibody comprising a human heavy chain, the heavy chain comprising a variable domain that is specific for an antigen and a constant region, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the VH gene segment being selected from a VH in Table 13 found in a Chinese population and with a cumulative frequency of at least 5%; and wherein

15

(i) the variable domain is derived from the recombination of said human gene segments in a non-human vertebrate (eg, in a mouse or a rat); or (ii) the variable domain comprises non-human vertebrate (eg, mouse or rat) AID-pattern mutations and non-human vertebrate (eg, mouse or rat) terminal deoxynucleotidyl transferase (TdT)-pattern mutations.

20

353. The antibody of claim 352, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the D gene segment being selected from a D in Table 13 found in a Chinese population and with a cumulative frequency of at least 5%.

25

354. The antibody of claim 352 or 353, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the JH gene segment being selected from a JH in Table 13 found in a Chinese population and with a cumulative frequency of at least 5%.

30

355. An isolated VH domain identical to a variable domain as recited in any one of claims 352 to 354, optionally fused at its C-terminus to a polypeptide (eg, an antibody Fc).

35

356. A pharmaceutical composition comprising the antibody or variable domain of any one of claims 352 to 355 together with a pharmaceutically-acceptable excipient, diluent or a medicament (eg, a further antigen-specific variable domain, antibody chain or antibody).

40

357. An antibody heavy chain or VH domain (eg, provided as part of an antibody) for therapy and/or prophylaxis of a disease or medical condition in a Chinese patient, wherein the heavy chain is a heavy chain produced by the following steps (or is a copy of such a heavy chain):-

45

(a) Selection of an antigen-specific antibody heavy chain or VH domain from a non-human vertebrate (eg, a mouse or a rat), wherein the heavy chain or VH domain is derived

from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment , the VH gene segment being selected from a VH in Table 13 found in a Chinese population and with a cumulative frequency of at least 5%;

5 (b) Optional humanisation of the heavy chain by combining the variable domain of the heavy chain with a human constant region; or optional humanisation of the selected VH domain by combining with a human constant region.

10 358. The antibody heavy chain or VH domain of claim 357, wherein the human constant region is as recited in claim 345 or 346.

15 359. An antibody heavy chain or VH domain as recited in claim 357 or 358 for use in a medicament for therapy and/or prophylaxis of a disease or medical condition in a Chinese patient.

20 360. A method of treating and/or preventing a disease or medical condition in a Chinese patient, the method comprising administering to the patient a therapeutically or prophylactically-effective amount of the antibody heavy chain or VH domain as recited in claim 357 or 358.

25 361. An isolated antibody for administration to a patient of European, East Asian, West African, South Asian or Americas ancestry, the antibody comprising a human heavy chain, the heavy chain comprising a variable domain that is specific for an antigen and a constant region, wherein the constant region is a human constant region selected from a constant region (eg, an IGHG constant region) in Table 13 found in a population of European, East Asian, West African, South Asian or Americas ancestry respectively and with a cumulative frequency of at least 5%;, and wherein

30 (i) the variable domain is derived from the recombination of said human gene segments in a non-human vertebrate (eg, in a mouse or a rat); or ( ii) the variable domain comprises non-human vertebrate (eg, mouse or rat) AID-pattern mutations and non-human vertebrate (eg, mouse or rat) terminal deoxynucleotidyl transferase (TdT)-pattern mutations.

35 362. The antibody of claim 361 wherein the constant region is a IGHG1a, IGHG2a, IGHG3a, IGHG3b or IGHG4a constant region and the patient is of European ancestry.

40 363. The antibody of claim 361 or 362, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the VH gene segment being selected from a VH in Table 13 found in said population and with a cumulative frequency of at least 5%.

364. The antibody of claim 361, 362 or 363, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment , the D gene segment being selected from a D in Table 13 found in



said population and with a cumulative frequency of at least 5%.

365. The antibody of claim 361, 362, 363 or 364 wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the JH gene segment being selected from a JH in Table 13 found in said population and with a cumulative frequency of at least 5%.

366. An isolated VH domain identical to a variable domain as recited in any one of claims 363 to 365, optionally fused at its C-terminus to a polypeptide (eg, an antibody Fc).

367. A pharmaceutical composition comprising the antibody or variable domain of any one of claims 361 to 366 together with a pharmaceutically-acceptable excipient, diluent or a medicament (eg, a further antigen-specific variable domain, antibody chain or antibody).

368. An isolated antibody for administration to a patient of European, East Asian, West African or Americas ancestry, the antibody comprising a human heavy chain, the heavy chain comprising a variable domain that is specific for an antigen and a constant region, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the VH gene segment being selected from a VH in Table 13 found in a population of European, East Asian, West African, South Asian or Americas ancestry respectively and with a cumulative frequency of at least 5%; and wherein

(i) the variable domain is derived from the recombination of said human gene segments in a non-human vertebrate (eg, in a mouse or a rat); or (ii) the variable domain comprises non-human vertebrate (eg, mouse or rat) AID-pattern mutations and non-human vertebrate (eg, mouse or rat) terminal deoxynucleotidyl transferase (TdT)-pattern mutations.

369. The antibody of claim 368, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the D gene segment being selected from a D in Table 13 found in said population and with a cumulative frequency of at least 5%.

370. The antibody of claim 368 or 369, wherein the variable domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the JH gene segment being selected from a JH in Table 13 found in said population and with a cumulative frequency of at least 5%.

371. An isolated VH domain identical to a variable domain as recited in any one of claims 368 to 370, optionally fused at its C-terminus to a polypeptide (eg, an antibody Fc).

372. A pharmaceutical composition comprising the antibody or variable domain of any one of claims 368 to 371 together with a pharmaceutically-acceptable excipient, diluent or a medicament (eg, a further antigen-specific variable domain, antibody chain or antibody).

373. An antibody heavy chain or VH domain (eg, provided as part of an antibody) for therapy and/or prophylaxis of a disease or medical condition in a patient of European, East Asian, West African, South Asian or Americas ancestry, wherein the heavy chain is a heavy chain produced by the following steps (or is a copy of such a heavy chain):-

5

(a) Selection of an antigen-specific antibody heavy chain or VH domain from a non-human vertebrate (eg, a mouse or a rat), wherein the heavy chain or VH domain is derived from the recombination of a human VH gene segment with a human D gene segment and a human JH gene segment, the VH gene segment being selected from a VH in Table 13 found in said population and with a cumulative frequency of at least 5%;

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(b) Optional humanisation of the heavy chain by combining the variable domain of the heavy chain with a human constant region; or optional humanisation of the selected VH domain by combining with a human constant region.

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374. The antibody heavy chain or VH domain of claim 373, wherein the human constant region is as recited in claim 361 or 362.

375. An antibody heavy chain or VH domain as recited in claim 373 or 374 for use in a medicament for therapy and/or prophylaxis of a disease or medical condition in a patient of said ancestry.

20

376. A method of treating and/or preventing a disease or medical condition in a patient of European, East Asian, West African, South Asian or Americas ancestry, the method comprising administering to the patient a therapeutically or prophylactically-effective amount of the antibody heavy chain or VH domain as recited in claim 373 or 374.

25

# Recombineered BAC Vectors to add Polymorphic V-regions to the Mouse Genome

## 1. Cassette Exchange – Replace existing V-regions

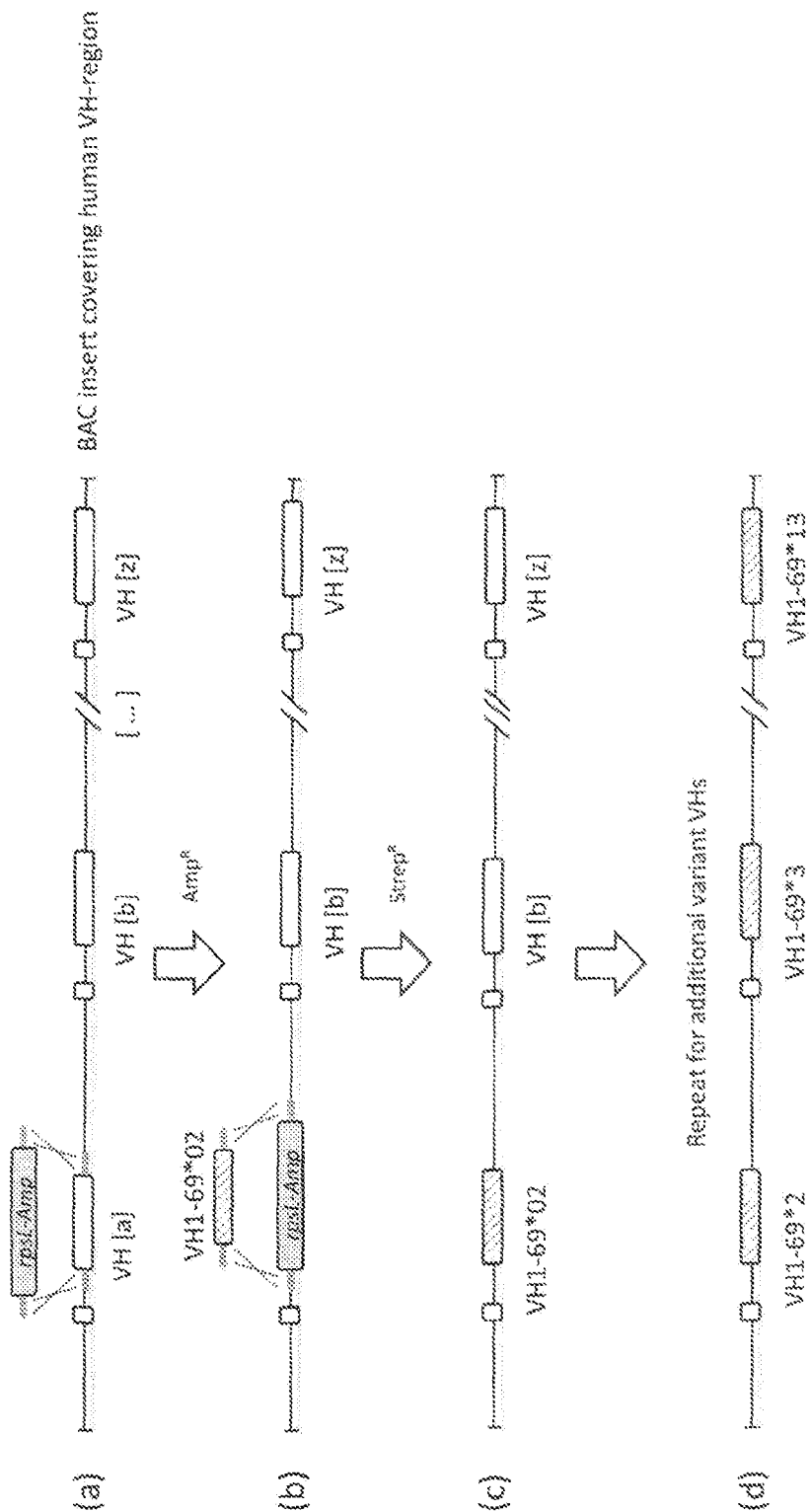


Figure 1

# Recombineered BAC Vectors to add Heterogeneous V-regions to the Mouse Genome

## 2. Insertion of V-regions into Heterologous Genomic DNA

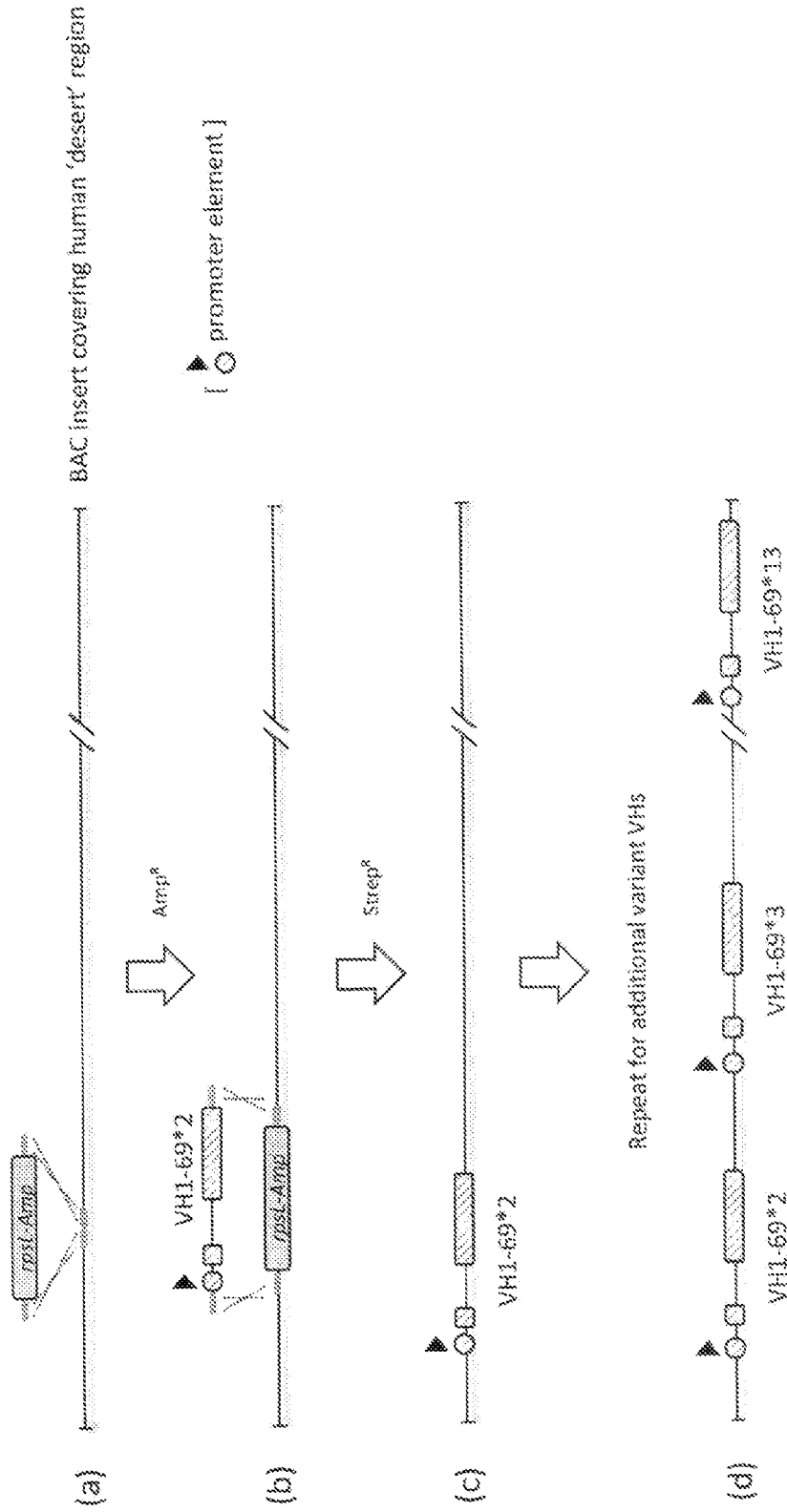


Figure 2

# Recombineered BAC Vectors to add Heterogeneous V-regions to the Mouse Genome

## 3. Single Strand Nucleotide Changes

[ Single Stranded targeting fragment with 2 nucleotide changes ]

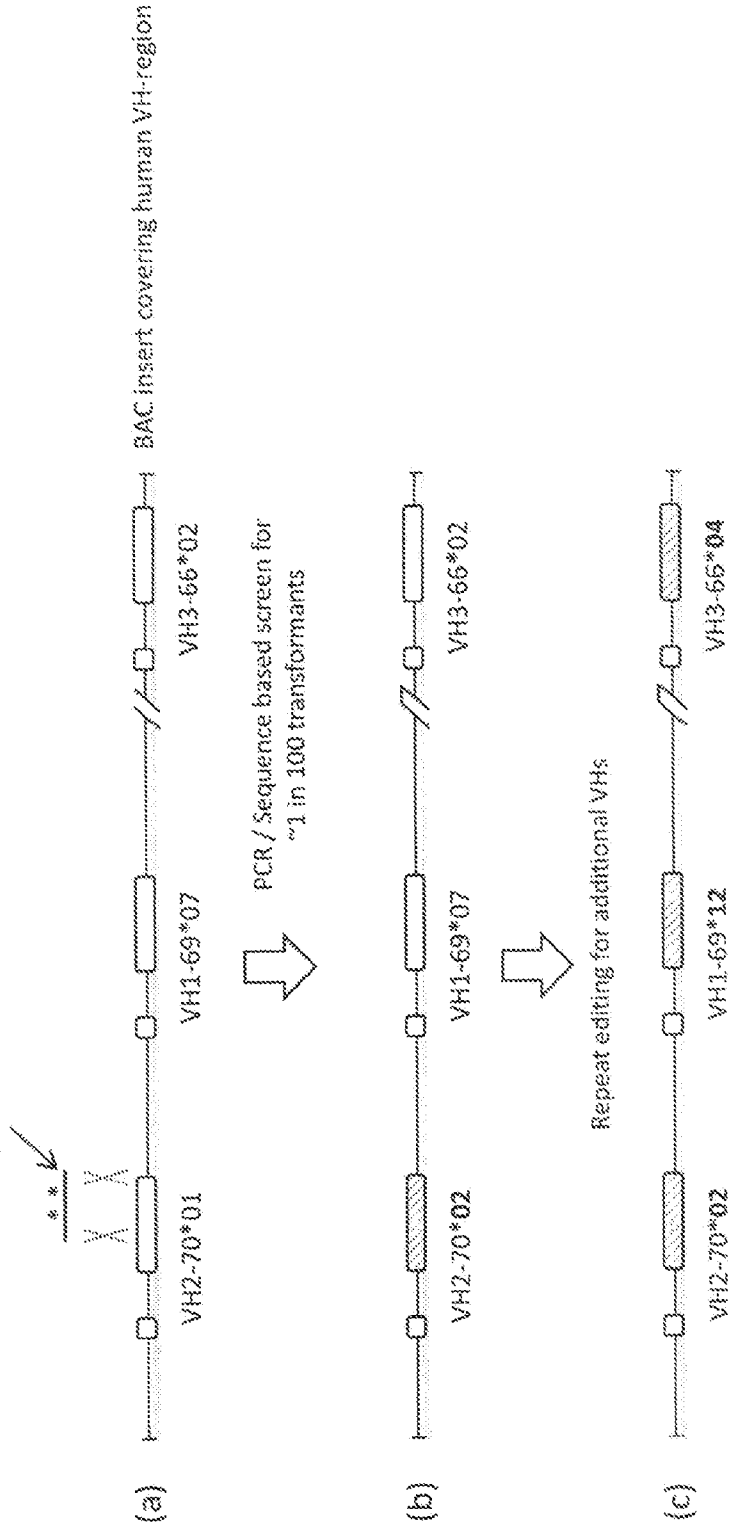


Figure 3

# Adding Polymorphic V-regions to the Genome using SRMCE of Modified BACs

1. Insertion at a 'Landing Pad' site during SRMCE

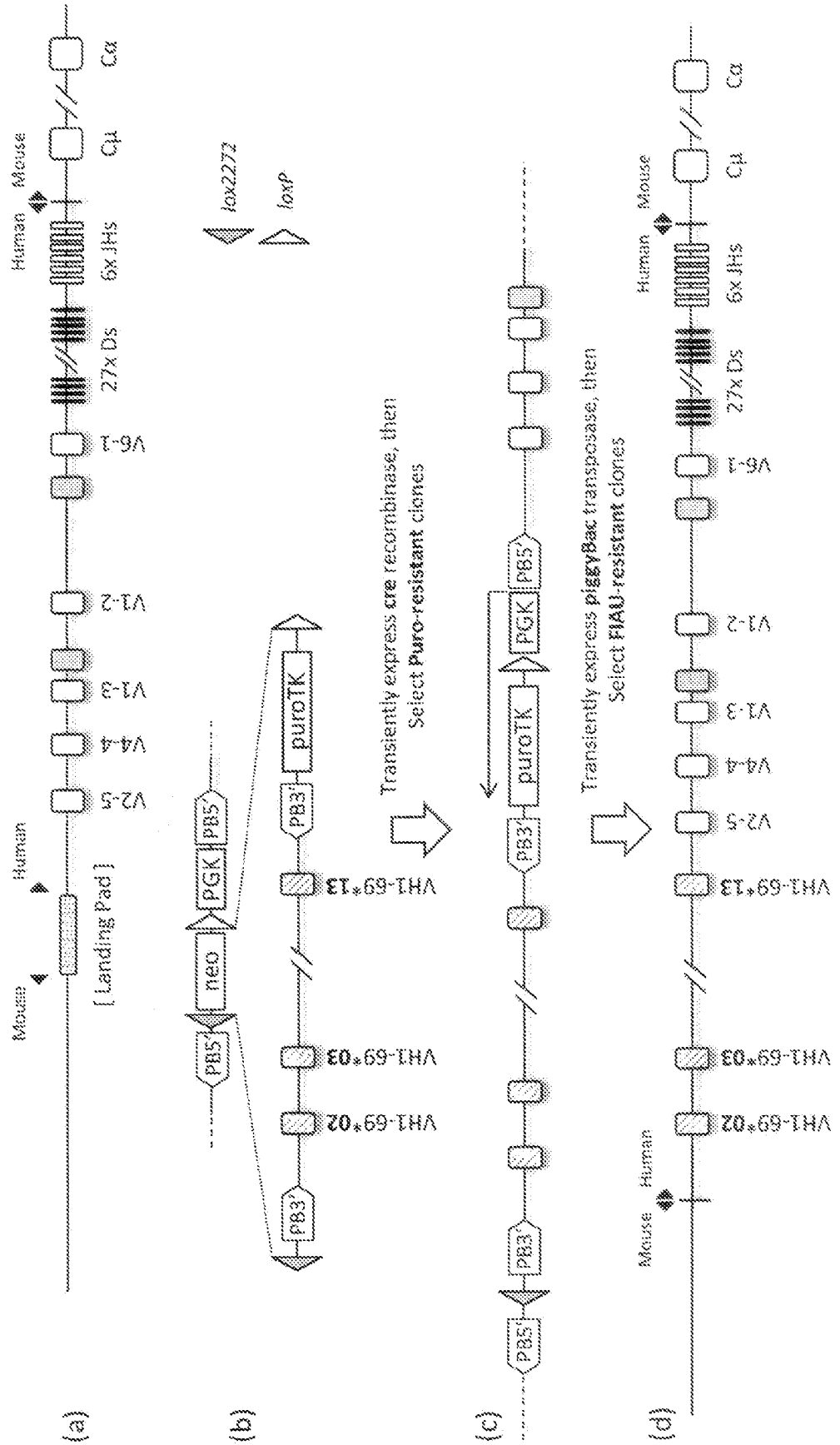


Figure 4

```

1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24 25 26
Q V Q L V Q E G A E V K R F G S S Q Y K V S C K A S
CAG GTG CAG CTG GTG CAG TCT GGG GCT ... GAG GTG AAG AAG OCT GGG TCC TCC GTG AAG GAC TTC TCC AAG GCG TCT
217505 ,IGHV1-69*02, Y18R6(YA67) ...C ...A
218240 ,IGHV1-69*03, 57G248
218312 ,IGHV1-69*04, hv1265
218405 ,IGHV1-69*05, RR.VH1.2
218553 ,IGHV1-69*06, hv1351K
218978 ,IGHV1-69*07, DA-2
219309 ,IGHV1-69*08
219307 ,IGHV1-69*09
219300 ,IGHV1-69*10
219296 ,IGHV1-69*11
219301 ,IGHV1-69*12
219219 ,IGHV1-69*13

```

Alignment of 13 IGHV1-69 alleles showing the variable (V) coding region only. Nucleotides that differ from VH1-69 allele \*01 are indicated at the appropriate position whereas identical nucleotides are marked with a dash. Where nucleotide changes result in amino acid differences, the encoded amino acid is shown above the corresponding triplet. Boxed regions correspond to CDR1, CDR2 and CDR3 as indicated.

Figure 5 (part 1 of 4)

	27	28	29	30	31	32	33	34	35	36	37	38	39	40	41	42	43	44	45	46	47	48	49	50	51	52	53
	G	G	T	F	S	S	V	A					I	S	W	Y	R	Q	A	P	G	Q	G	L	E	W	R
	GGA	GGC	ACC	TTC	AGC	AGC	TAT	GCT					ATC	AGC	TGG	GTC	CGA	CAG	GCC	GCT	GGA	CAA	GAG	GCT	GAG	TGG	ATG
L22582																											
227506								A																			
X92340																											
M83131																											
X67905																											
L22563																											
Z26979																											
Z14309								T																			
Z14307																											
Z14300																											
Z14296																											
Z14301																											
Z14214																											

Alignment of 13 IGHV1-69 alleles showing the variable (V) coding region only. Nucleotides that differ from VH1-69 allele. \*Q1 are indicated at the appropriate position whereas identical nucleotides are marked with a dash. Where nucleotide changes result in amino acid differences, the encoded amino acid is shown above the corresponding triplet. Boxed regions correspond to CDR1, CDR2 and CDR3 as indicated.

Figure 5 (part 2 of 4)



	54	55	56	57	58	59	60	61	62	63	64	65	66	67	68	69	70	71	72	73	74	75	76	77	78	79	80	
	G	G	I	I	P	I	F	I	F	G	T	A	R	Y	A	G	K	F	Q	G	G	R	V	I	I	I	A	
122582	CGA	GGG	ATC	ATC	CCT	ATC	ATC	TTT	GGT	ACA	GCA	...	...	...	...	...	...	...	...	...	...	...	...	...	...	...	...	...
	R				I			I																				
217566	A				C																							
832340																												
883132	A																											
867905																												
122563	R																											
229978	A																											
214309	R																											
214307	R																											
214300	R																											
214296	A																											
214301																												
214314																												

Alignment of 13 IGHV1-69 alleles showing the variable (V) coding region only. Nucleotides that differ from VH1-69 allele \*01 are indicated at the appropriate position whereas identical nucleotides are marked with a dash. Where nucleotide changes result in amino acid differences, the encoded amino acid is shown above the corresponding triplet. Boxed regions correspond to CDR1, CDR2 and CDR3 as indicated.

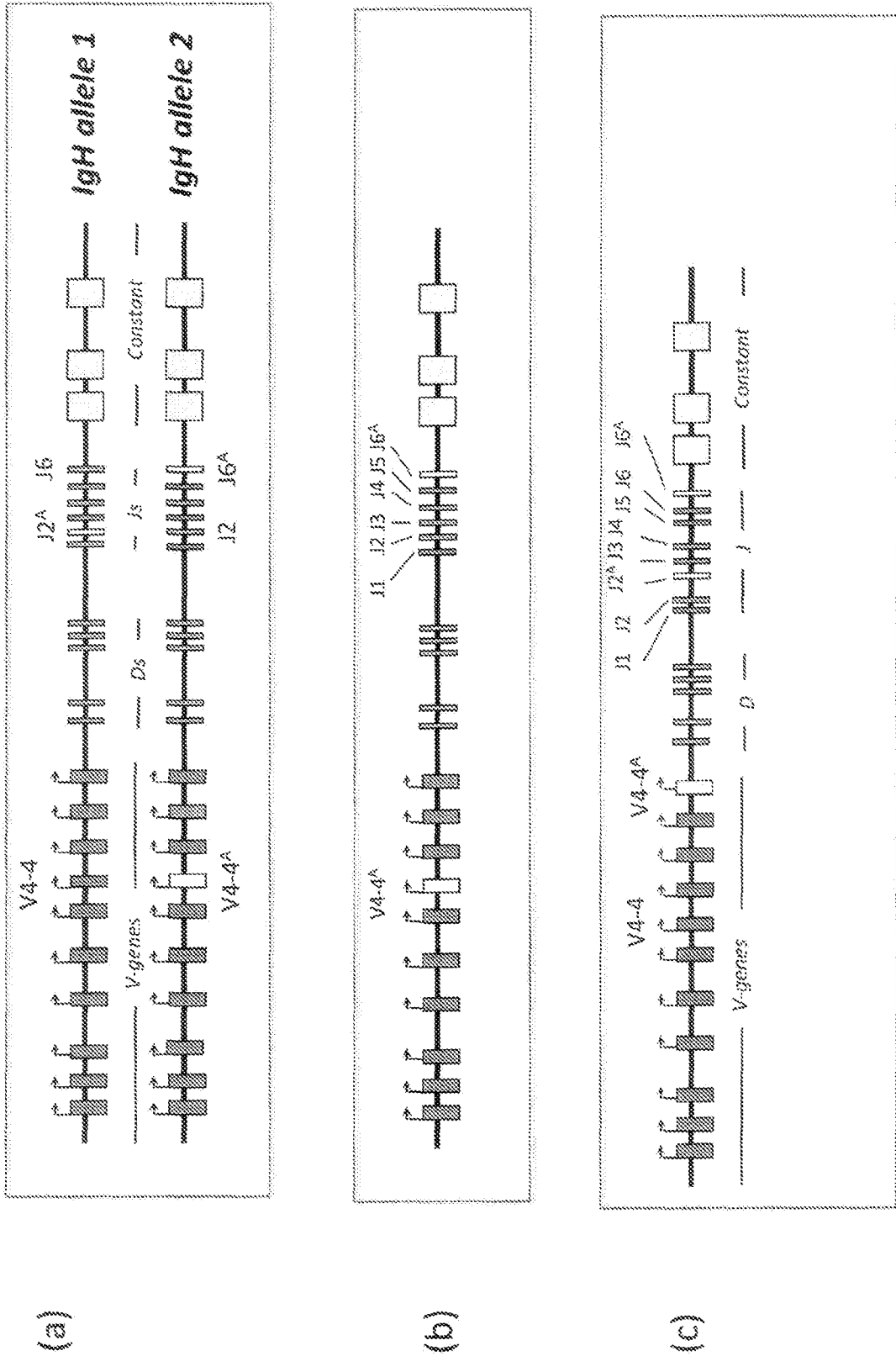
Figure 5 (part 3 of 4)

	81	82	83	84	85	86	87	88	89	90	91	92	93	94	95	96	97	98	99	100	101	102	103	104	105	106	CDR3
L22582 ,IGHV1-69*01, hV1051	D	E	S	T	S	T	A	T	M	E	L	S	S	L	K	S	E	D	T	A	V	Y	Y	C	A	K	
	GAC	GAA	TCC	ACG	ABC	ACA	GCC	TAC	ATG	GAG	CTG	AGC	AGC	CTG	AGA	TCT	GAG	GAC	ACG	GCC	GTC	GAT	TAC	TGT	GGG	AGA	GA
	K																										
L27506 ,IGHV1-69*02, yIGH6(YAC7)	A																										
X92340 ,IGHV1-69*03, S7GTA8	K																										
893332 ,IGHV1-69*04, hV1263																											
X67905 ,IGHV1-69*05, ER.VH1.2	K																										
L22583 ,IGHV1-69*06, hV1051K	A																										
Z09978 ,IGHV1-69*07, EA-2	K																										
Z14309 ,IGHV1-69*08	A																										
Z14307 ,IGHV1-69*09	K																										
Z14300 ,IGHV1-69*10	A																										
Z14296 ,IGHV1-69*11	K																										
Z14301 ,IGHV1-69*12	A																										
Z14214 ,IGHV1-69*13																											

Alignment of 13 IGHV1-69 alleles showing the variable (V) coding region only. Nucleotides that differ from VH1-69 allele \*01 are indicated at the appropriate position whereas identical nucleotides are marked with a dash. Where nucleotide changes result in amino acid differences, the encoded amino acid is shown above the corresponding triplet. Boxed regions correspond to CDR1, CDR2 and CDR3 as indicated.

Figure 5 (part 4 of 4)

Figure 6



**FIGURE 7**

	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	
J00256, IGHJ6*01	AT	TAC	TAC	TAC	TAC	GGT	ATG	GAC	GTC	TGG	GGG	CAA	GGG	ACC	ACG	GTC	ACC	GTC	TCC	TCA	G
X86355, IGHJ6*02	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---
X86357, IGHJ6*02	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---
X86358, IGHJ6*02	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---
M63031, IGHJ6*02	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---
X97051, IGHJ6*02	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---
M25625, IGHJ6*02	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---
X86356, IGHJ6*03	---	---	---	---	---	<u>Y</u>	<u>TAC</u>	---	---	---	---	<u>K</u>	---	---	---	---	---	---	---	---	---
X86359, IGHJ6*03	---	---	---	---	---	<u>Y</u>	<u>TAC</u>	---	---	---	---	<u>K</u>	---	---	---	---	---	---	---	---	---
M63030, IGHJ6*03	---	---	---	---	---	<u>Y</u>	<u>TAC</u>	---	---	---	---	<u>K</u>	---	---	---	---	---	---	---	---	---
AJ879487, IGHJ6*04	---	---	---	---	---	---	---	---	---	---	---	<u>K</u>	---	---	---	---	---	---	---	---	---

**FIGURE 8**

Rabbit JH6	<b>Y</b>	<b>Y</b>	<b>G</b>	M	D	L
	at	tac	tac	ggc	atg	gac ctc
Sheep JH6	<b>Y</b>	<b>Y</b>	<b>G</b>	V	D	V
	at	tac	tac	ggt	gta	gat gtc
Bovine JH6	<b>Y</b>	<b>Y</b>	<b>G</b>	V	D	V
	at	tac	tac	ggt	gta	gat gtc
Dog JH3	<b>Y</b>	<b>Y</b>	<b>G</b>	M	D	Y
	at	tac	tat	ggt	atg	gac tac
Human JH6*02	<b>Y</b>	<b>Y</b>	<b>Y</b>	<b>Y</b>	<b>G</b>	M D V
	<b>Y</b>	tac	tac	tac	tac	ggt atg gac gtc

**FIGURE 9**

	T	C	A	G
T	TTT Phe F	TCT Ser S	TAT Tyr Y	TGT Cys C
	TTC Phe F	TCC Ser S	TAC Tyr Y	TGC Cys C
	TTA Leu L	TCA Ser S	TAA stop *	TGA stop *
	TTG Leu L	TCG Ser S	TAG stop *	TGG Trp W
C	CTT Leu L	CCT Pro P	CAT His H	CGT Arg R
	CTC Leu L	CCC Pro P	CAC His H	CGC Arg R
	CTA Leu L	CCA Pro P	CAA Gln Q	CGA Arg R
	CTG Leu L	CCG Pro P	CAG Gln Q	CGG Arg R
A	ATT Ile I	ACT Thr T	AAT Asn N	AGT Ser S
	ATC Ile I	ACC Thr T	AAC Asn N	AGC Ser S
	ATA Ile I	ACA Thr T	AAA Lys K	AGA Arg R
	ATG Met M	ACG Thr T	AAG Lys K	AGG Arg R
G	GTT Val V	GCT Ala A	GAT Asp D	GGT Gly G
	GTC Val V	GCC Ala A	GAC Asp D	GGC Gly G
	GTA Val V	GCA Ala A	GAA Glu E	GGA Gly G
	GTG Val V	GCG Ala A	GAG Glu E	GGG Gly G

Figure 10

chr14:106,026,574-107,346,185 1,319,612 bp. enter position, gene symbol or search terms

