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#### Abstract

In some embodiments, the invention relates to methods for creating a monoclonal antibody that specifically binds to antigen. The method may start from a polyclonal population of antibodies such as a non-specific polyclonal population or a polyclonal population of antibodies that specifically bind to the antigen. The method includes obtaining nucleic acid molecules encoding heavy and light immunoglobulin chains (or variable regions thereof) of multiple immunoglobulins from an animal; obtaining mass spectra information of peptide fragments of a population of polyclonal immunoglobulins that specifically bind to an antigen of choice; comparing and/or correlating the mass spectra information of the peptide fragments of the polyclonal immunoglobulins with predicted mass spectra information of predicted amino acid sequences encoded by the nucleic acid molecules, and then assembling the heavy and light chains to create an antibody (or variable region thereof) that specifically binds to the antigen.


# METHODS AND REAGENTS FOR CREATING MONOCLONAL ANTIBODIES 

## Cross Reference to Related Applications

This application claims priority from U.S. Provisional Application No. 61/450,922, filed March 9, 2011, U.S. Provisional Application No. 61/560,006, filed November 15, 2011, U.S. Provisional Application No. 61/566,876, filed December 5, 2011, and U.S. Provisional Application No. 61/594,729, filed February 3, 2012, the entire contents of each of which are incorporated herein by reference.

## Field of the Disclosure

This disclosure relates to biology, and more specifically, to molecular biology and immunology.

## Background of the Disclosure

Antibodies are biologically and commercially significant polypeptides that bind with great specificity and affinity to a particular target molecule, called an antigen. Antibodies are produced by immune cells of vertebrate animals, and all naturally-occurring antibodies share the same basic structure, namely two identical heavy chains covalently bonded to two identical light chains. The N -terminal regions of a single heavy chain and a single light chain form an antigen-binding site that is particular to each individual antibody. The Cterminal region of the heavy chains determines the particular isotype of the antibody, and the same antibody-producing cell can produce antibodies of different isotypes, where all the antibodies produced by the cell have the same antigen-binding site. The different isotypes typically perform different functions in the animal. For example, antibodies of the E isotype (i.e., IgE antibodies) are involved in the allergic response while antibodies of the A isotype (i.e., in IgA antibodies) can be found in mucosal membrane, saliva, and breast milk. The four-chain antibody molecule can exist by itself (e.g., an IgG antibody) or with additional monomers to form dimers (e.g., an IgA antibody) or even pentamers (e.g., an IgM antibody).

With the basic structure of an antibody well-understood, one can produce recombinant antibodies by manipulating the different regions of an antibody using standard
molecular biology techniques. For example, U.S. Patent Nos. 6,180,370 and 6,548,640 (herein incorporated by reference in their entirety) describe humanizing an antibody that naturally occurs in a non-human animal by manipulating various regions of the non-human antibody using molecular biology techniques. Other methods for manipulating or generating recombinant antibodies using standard molecular biology techniques are described (see, e.g., PCT Publication No. WO91/17271, PCT Publication No. WO92/01047; U.S. Patent Nos. $5,969,108,6,331,415,7,498,024$, and $7,485,291$, all of which are herein incorporated by reference in their entirety).

During an immune response, an animal will generate numerous different antibodies, each with a different antigen-binding specificity. This population of antibodies is called a polyclonal population of antibodies. If the immune response is directed toward a particular antigen, most (but not all) of the polyclonal antibodies made by the animal will specifically bind that antigen. However, with differences in binding affinity and binding sites on the antigen, some of the polyclonal antibodies are more favored than other polyclonal antibodies. In their Nobel Prize-winning discovery in 1975, Kohler and Milstein discovered a way to isolate and immortalize a single antibody-producing cell, which produces a monoclonal antibody that specifically binds to the antigen of interest, from a polyclonal antibody-producing animal (Kohler and Milstein, Nature 256: 495-497, 1975). This immortalization technology, which involves fusing the antibody-producing cells to an immortalized cell to produce a monoclonal antibody-producing hybridoma, has been the industry standard for making monoclonal antibodies for the past 35 years.

Despite its popularity and its longevity, the Kohler and Milstein hybridoma method has numerous drawbacks. For example, it is very time-consuming and labor-intensive. More relevantly, given how time-consuming and labor-intensive it is, only a small fraction of the antibody-producing cells of the animal are immortalized and screened for their ability to produce an antibody that specifically binds to the antigen. Finally, even once a hybridoma with the desired antigen specificity is isolated, obtaining the amino acid sequence of the antibody to facilitate further manipulation, such as humanization, of the antibody, is arduous and time-consuming.

There is a need for improved methods for creating monoclonal antibodies that
specifically bind to a desired antigen.

## Summary of the Disclosure

The various aspects and embodiments of the invention provide methods and systems to rapidly
5 and accurately create monoclonal antibodies that specifically bind to an antigen of interest. In further aspects and embodiments, the invention provides reagents and compositions for performing the various methods of the invention, and reagents and compositions resulting from the performance of the various methods of the invention. In some embodiments, the methods, reagents, and compositions disclosed herein are useful to create monocloncal antibodies from the circulation of a subject. In one aspect, the invention provides

A method for obtaining nucleotide sequences or amino acid sequences of heavy or light chains of immunoglobulins that specifically bind to an antigen of interest, comprising: (a) providing nucleic acid sequences encoding immunoglobulin chains of at least one animal, and deriving predicted mass spectra information from predicted amino acid sequences encoded by said nucleic acid sequences; (b) obtaining a population of polyclonal immunoglobulins, wherein said population of polyclonal immunoglobulins is a purified population of polyclonal immunoglobulins obtained by antigen affinity purification with said antigen of interest, obtaining peptide fragments by protease cleavage of said population, and obtaining mass spectra information of said peptide fragments; (c) correlating the mass spectra information of said peptide fragments obtained in step (b) with the predicted mass spectra information derived from the nucleic acid sequences in step (a), to identify nucleic acid sequences encoding immunoglobulin chains that comprise amino acid sequences corresponding to one or more of said peptide fragments; determining the variable regions and CDR3 regions of the immunoglobulin chains encoded by the identified nucleic acid sequences; and for the variable region and the CDR3 region encoded by each identified nucleic acid sequence, determining the amino acid sequence coverage of the variable region by said peptide fragments, and determining the amino acid sequence coverage of the CDR3 region by said peptide fragments; and (d) selecting from the identified nucleic acid sequences, nucleic acid sequences having an amino acid sequence coverage of the variable region of at least $60 \%$ and an amino acid sequence coverage of the CDR3 region of at least $60 \%$, to obtain nucleic acid sequences or amino acid sequences of heavy or light chains of immunoglobulins that specifically bind to said antigen of interest.

In some embodiments, a heavy immunoglobulin chain and a light immunoglobulin chain (or variable regions thereof) selected in step (d) are assembled to create an immunoglobulin (or variable regions thereof) that specifically binds to the antigen.

In some embodiments, the nucleotide sequences or amino acid sequences of the
immunoglobulin chain variable regions obtained in step (d) are synthesized by recombinant molecular biology techniques or gene synthesis techniques prior to assembly.

In some embodiments, the method further comprises: screening with an immunoassay the immunoglobulin (or variable regions thereof) created to confirm said immunoglobulin (or variable regions thereof) specifically binds to the antigen. In some embodiments, the immunoassay is selected from the group consisting of a flow cytometry assay, an enzyme-linked immunosorbent assay (ELISA), a Western blotting assay, an immunohistochemistry assay, an immunofluorescence assay, a radioimmunoassay, a neutralization assay, a binding assay, an affinity assay, or a protein or peptide immunoprecipitation assay.

In some embodiments, the selection of heavy immunoglobulin chains and light immunoglobulin chains in step (d) is made based on amino acid sequence coverage of a portion of the chains (e.g., the variable region or a complementarity determining region) by the peptide fragments.

In other embodiments, the selection of heavy or light immunoglobulin chains in step (d) is made based on the amino acid sequence coverage of the immunoglobulin chains or fragments thereof by the peptide fragments, in combination with at least one parameter selected from the group consisting of the number of unique peptides mapped, spectrum share, total peptide count, unique peptide count, frequency of the encoding nucleic acid sequences, and clonal relatedness.

In various embodiments, the nucleic acid sequences and information derived from the nucleic acid sequences (including, for example, the nucleotide sequences, the predicted amino acid sequences, and the predicted mass spectra) are located in a genetic material database.

In some embodiments, the animal from which the nucleic acid sequences are obtained is an animal exposed to the antigen.

In some embodiments, the predicted amino acid sequences encoded by said nucleic acid sequences encoding immunoglobulin chains (or variable regions thereof) of multiple immunoglobulins from the animal are obtained by: (1) isolating nucleic acid molecules from white blood cells from said animal; (2) amplifying immunoglobulin chain (or variable
region thereof)-encoding nucleic acid molecules using primers specific for polynucleotide sequences adjacent to said immunoglobulin chain (or variable region thereof)-encoding nucleic acid molecules; (3) obtaining nucleotide sequences of said amplified nucleic acid molecules encoding immunoglobulin chains (or variable regions thereof) of multiple immunoglobulins from the animal; and (4) using the genetic code to translate the nucleotide sequences into predicted amino acid sequences.

In some embodiments, the nucleic acid sequences are expressed nucleic acid sequences (e.g., transcribed into RNA and/or translated into protein in cells of the animal).

In some embodiments, the predicted amino acid sequences encoded by the nucleic acid molecules encoding immunoglobulin chains (or variable regions thereof) of multiple immunoglobulins from the animal are obtained by: (1) isolating nucleic acid molecules from white blood cells from said animal; (2) sequencing immunoglobulin chain (or variable region thereof)-encoding nucleic acid molecules using primers specific for polynucleotide sequences adjacent to said immunoglobulin chain (or variable region thereof)-encoding nucleic acid molecules to obtain the nucleotide sequences encoding immunoglobulin chains (or variable regions thereof) of multiple immunoglobulins from the animal; and (3) using the genetic code to translate the nucleic acid sequences into amino acid sequences. In some embodiments, the nucleic acid molecules are RNA molecules and said amplification step includes an initial reverse transcription step.

In some embodiments, the polynucleotide sequences adjacent to the immunoglobulin chain (or variable region thereof)-encoding nucleic acid molecules are selected from the group consisting of genomic DNA flanking immunoglobulin genes, immunoglobulin chain constant region-encoding polynucleotide sequences, and immunoglobulin chain framework region-encoding polynucleotide sequences.

In some embodiments, the predicted mass spectra information is obtained using a method comprising the steps of: (i) performing a theoretical digest of predicted amino acid sequences encoded by the nucleotide sequences of the nucleic acid molecules with one or more proteases and/or one or more chemical protein cleavage reagents to generate virtual peptide fragments; and (ii) creating predicted mass spectra of said virtual peptide fragments.

In some embodiments, the observed mass spectra information of the peptide
fragments are obtained using a method comprising the steps of: (i) isolating a population of polyclonal immunoglobulins that specifically bind to the antigen; (ii) digesting the population with one or more proteases and/or one or more chemical protein cleavage reagents to generate fragments; and (iii) obtaining mass spectra information of said peptide
5 fragments. In some embodiments, the population of polyclonal antibodies is isolated using a method comprising the steps of: (1) obtaining body fluid or a fraction thereof (e.g., blood, serum and/or plasma) from an animal; (2) passing the body fluid or a fraction thereof over the antigen under conditions whereby immunoglobulins that specifically bind to the antigen will become attached the antigen; and (3) collecting said immunoglobulins attached to said 10 antigen to obtain the population of polyclonal immunoglobulins that specifically bind to the antigen. In some embodiments, the antigen is attached to a solid support (e.g., the antigen is covalently or non-covalently bound to the solid support). In some embodiments, the solid support may be a bead (e.g., an agarose or a magnetic bead), a wall of a column, or a bottom of a plate (e.g., a tissue culture plate).

15 In some embodiments, the animal is an animal previously exposed to the antigen. In some embodiments, the animal previously exposed to the antigen is an animal previously immunized with the antigen.

In another aspect, the invention provides a method for obtaining nucleic acid sequences or amino acid sequences of an immunoglobulin chain variable region of an immunoglobulin that specifically binds to an antigen of interest comprising: (a) providing nucleic acid sequences encoding immunoglobulin variable regions of multiple immunoglobulins of at least one animal , and deriving predicted mass spectra information from predicted amino acid sequences encoded by said nucleic acid sequences; (b) obtaining a population of polyclonal immunoglobulins, wherein said population of polyclonal immunoglobulins is a purified population of polyclonal immunoglobulins obtained by antigen affinity purification with said antigen of interest, obtaining peptide fragments of immunoglobulin chain variable regions of said population by protease cleavage, and obtaining mass spectra information of said peptide fragments; (c) correlating the mass spectra information of the peptide fragments obtained in step (b) with the predicted mass spectra information derived from the nucleic acid sequences in step (a), to identify nucleic acid sequences encoding immunoglobulin chain variable regions that comprise amino acid sequences corresponding to one or more of said peptide fragments; determining the CDR3 regions of the immunoglobulin chains encoded by the identified nucleic acid sequences; and for the variable region and the CDR3 region encoded by each identified nucleic acid sequence, determining the amino acid sequence coverage of
the variable region by said peptide fragments, and determining the amino acid sequence coverage of the CDR3 region by said peptide fragments; and (d) selecting from the identified nucleotide sequences, nucleic acid sequences having an amino acid sequence coverage of the variable region of at least $60 \%$ and an amino acid sequence coverage of the CDR 3 region of at
5 least $60 \%$, to obtain nucleic acid sequences or amino acid sequences of variable regions of immunoglobulins that specifically bind to said antigen of interest.

In some embodiments, the method further comprises step (e) screening the amino acid sequences of said immunoglobulin chain variable regions with an immunoassay to isolate an immunoglobulin chain variable region of an immunoglobulin that specifically binds to the antigen. In some embodiments, the nucleotide sequences or amino acid sequences of the immunoglobulin chain variable regions obtained in step (d) are synthesized by recombinant molecular biology techniques or gene synthesis techniques prior to the step (e) screening step. In some embodiments, the immunoglobulin chain variable region produced in step (d) is assembled with a second immunoglobulin chain variable region to create an antibody binding domain of an immunoglobulin that specifically binds to the antigen. In some embodiments, the immunoassay is selected from the group consisting of a flow cytometry assay, an enzymelinked immunosorbent assay (ELISA), a Western blotting assay, and immunohistochemistry assay, an immunofluorescence assay, a radioimmunoassay, a neutralization assay, a binding assay, an affinity assay, or a protein or peptide immunoprecipitation assay.

In some embodiments, the immunoglobulin chain variable region is a heavy chain variable region or a light chain variable region.

In a further aspect, the invention provides a method for creating an antigen binding domain of an immunoglobulin that specifically binds to an antigen comprising: (a) providing nucleic acid sequences encoding immunoglobulin heavy chain variable regions and light chain variable regions of multiple immunoglobulins from an animal; (b) obtaining mass spectra information of peptide fragments of heavy immunoglobulin chains and light immunoglobulin chains of a population of polyclonal immunoglobulins that specifically bind to an antigen; (c) correlating mass spectra information of the peptide fragments with predicted mass spectra information of the nucleic acid sequences, said predicted mass spectra information derived from predicted amino acid sequences encoded by said nucleic acid sequences, to obtain nucleotide sequences or amino acid sequences of immunoglobulin chain variable regions comprising the peptide fragments; (d) selecting from the identified nucleotide sequences or amino acid sequences of immunoglobulin chain variable regions
based on the amino acid sequence coverage of the variable regions by the peptide fragments, to obtain nucleotide sequences or amino acid sequences of variable regions of immunoglobulins that specifically bind to an antigen; and (e) assembling a selected heavy immunoglobulin chain variable region with a selected light immunoglobulin chain variable region to create an antigen binding domain of an immunoglobulin that specifically binds to the antigen.

In various embodiments of all of the aspects of the invention, the animal is a vertebrate animal. In various embodiments, the animal is a mammal. In some embodiments, the animal is a human. In some embodiments, the animal is a rat, a rabbit or a mouse. In some embodiments, the animal is a bird, domesticated animal, a companion animal, a livestock animal, a rodent, or a primate. In some embodiments, the animal is a transgenic non-human animal, e.g., a transgenic non-human animal that expresses human antibody sequences and/or produces antibodies that are at least partly human.

In various aspects, the invention also provides an immunoglobulin (or variable region thereof), or an immunoglobulin chain variable region or an antigen binding domain of an immunoglobulin that specifically binds to an antigen isolated or created in accordance with the various non-limiting embodiments of the invention. In various embodiments, the immunoglobulin (or variable region thereof), or an immunoglobulin chain variable region or an antigen binding domain of an immunoglobulin that specifically binds to an antigen are isolated or recombinant. In various embodiments, the invention also provides a pharmaceutically acceptable carrier and an immunoglobulin (or variable region thereof), or an immunoglobulin chain variable region or an antigen binding domain of an immunoglobulin that specifically binds to an antigen isolated or created in accordance with the various non-limiting embodiments of the invention.

In a further aspect, the invention provides a method of treating an animal having or suspected of having a disease characterized by a disease antigen, wherein the method comprising administering an effective amount of a composition in accordance with various embodiments of the invention, wherein the antigen specifically bound by the immunoglobulin (or variable region thereof), or immunoglobulin chain variable region or an antigen binding domain of the composition and the disease antigen are the same. In some
embodiments, the animal is a human. In some embodiments, the animal is a rodent, a livestock animal, a domesticated animal, a companion animal, or a primate.

In a further aspect, the invention provides a method of reducing the likelihood of occurrence in an animal of a disease characterized by the presence in the animal of a disease in accordance with various embodiments of the invention, wherein the antigen specifically bound by the immunoglobulin (or variable region thereof), or immunoglobulin chain variable region or an antigen binding domain of the composition and the disease antigen are the same. In some embodiments, the composition further comprises an adjuvant. In some embodiments, the animal is a human. In some embodiments, the animal is a rodent, a livestock animal, a domesticated animal, a companion animal, or a primate.

## Brief Description of the Drawings

Figure 1 is a schematic diagram of an antibody comprising two heavy chains and two light chains. The two heavy chains are joined to each other by two disulfide bonds located in the hinge region of the antibody. Each light chain is joined to a heavy chain via a single disulfide bond. The antigen-binding site is created at the N -terminus of the heavy and light chains.

Figure 2 is a schematic diagram showing an example of a non-limiting method of various embodiments of the invention. In this example, samples comprising B lymphocytes (e.g., a blood sample or a tissue sample) and blood serum and/or plasma are collected from the same animal (e.g., a human, mouse, or rabbit). Nucleic acid molecules encoding immunoglobulin chains (or variable regions thereof) are sequenced and these nucleic acid sequences are used to generate theoretical or predicted mass spectra information based on the predicted amino acid sequences encoded by the nucleic acid sequences. Polyclonal antibodies from the blood sera are proteolytically digested or chemically fragmented and the resulting peptide fragments subjected to analysis by mass spectrometry. The information from the nucleic acid sequences (e.g., the mass spectra) is compared to the mass spectra information of the peptides fragments to identify the sequence of an immunoglobulin chain
(or variable domain thereof) of an antibody. This antibody can then be generated recombinantly according to standard methods.

Figure 3 is a schematic diagram showing another example of a non-limiting method of various embodiments of the invention. In this example, B lymphocytes and blood serum and/or plasma are collected from the same animal (in this case a rabbit). From the B lymphocytes, mRNA is extracted and subjected to sequencing using the Genome Sequencer FLX System machine commercially available from 454 Life Sciences using immunoglobulin gene-specific sequencing primers. This information is used to generate theoretical mass spectra based on the predicted amino acid sequences. From the blood serum and/or plasma, polyclonal antibodies are isolated and subjected to digestion with proteases and/or cleavage with chemical protein cleavage reagents. The resulting peptide fragments are separated by liquid chromatography, followed by mass spectrometry analysis (MS/MS). The mass spectra of the peptide fragments are correlated with the theoretical mass spectra of the nucleic acid sequences to obtain the amino acid sequences of the immunoglobulin chains that include the peptide fragments. A heavy and light chain can then be assembled to create a recombinant immunoglobulin by cloning nucleic acid sequences encoding the immunoglobulin chains into expression vector(s) and expressing the expression vectors in a cell. The expressed recombinant immunoglobulin is then further characterized.

Figure 4 is schematic diagram depicting another example of a non-limiting method of various embodiments of the invention. In this example, a non-limiting B cell source (e.g., splenocytes) and polyclonal antibodies are collected from the same animal (e.g., a human, mouse, or rabbit). Nucleic acid molecules are extracted from the B cell source and are subjected to next generation sequencing (NGS) using the Roche 454 machine using immunoglobulin gene-specific sequencing primers. This information, which can be put into a genetic material database, can be used to generate theoretical mass spectra based on the predict amino acid sequences encoded by the nucleic acid sequence. Also from the animal (e.g., a human, mouse, or rabbit), polyclonal antibodies (or peptide fragments thereof) are loaded into the mass spectrometer for analysis. The nucleic acid sequences are analyzed using Kabat rules to identify the sequences of the variable regions (e.g., one of the CDR
regions or FR regions) of the sequence. The sequences of the peptide fragments from the analyzed polyclonal antibodies are then screened to identify which peptides match all or part of the variable region from a predicted amino acid sequence.

Figure 5 is a table showing heavy and light chain NGS (i.e., next generation sequencing) sequences that had good mass spectrometry correlation and peptide over the variable region. Some of these peptides appeared quite frequently (see, e.g., light chain ref. no. G623FKB01A3GC7) and some had high nucleic acid sequence frequency count (see, e.g., light chain ref. no. G623FKB01AXJ1C). The rows in bold italics represent immunoglobulin chains that, upon testing, were found to contain sequences that specifically bound antigen (see testing results in Figure 6).

Figure 6 is a table showing the results of ELISA assays testing antibodies made using a non-limiting method of the invention screened against ELISA plates coated with pErk peptides. The different light chains and heavy chains shown in Fig. 5 were randomly combined with each other. As can be seen from Fig. 6, a number of pairings resulted in antibodies that were able to specifically bind to the p-ERK-coated plates (positive antibodies shown in shade).

Figure 7 is a photographic representation of an agarose gel showing the results of an RT-PCR reaction (i.e., reverse-transcriptase-polymerase chain reaction) of heavy chains and kappa and lambda light chains from cDNA generated from splenocytes of rabbits immunized with p -MET antigen.

Figure $\mathbf{8}$ is a table showing the sequences of the antibody chains after combining the theoretical (i.e., predicted) mass spectra derived from the nucleic acid sequences with LCMS/MS data from affinity purified antibody. Antibody chain abundance based on NGS frequency was also displayed. The chains depicted in italics were synthesized and assembled into antibody; and the bold italics chains are those which, upon testing with Western blotting analysis, were found to specifically bind the p-MET antigen.

Figure 9 is a photographic representation showing the results of a Western blotting experiment probing lysates prepared from Hela cells untreated (- lanes in all three blots) or treated with human growth factor (HGF) (+ lanes in all three blots) with two different rabbit antibodies generated using a non-limiting method of the invention (blots labeled 1 and 2)
and with a control antibody (left-most blots). Following positive results with Western blotting, antigen specific antibodies (heavy and light chain pairing) were then identified. As shown, the antibodies identified in both lanes 1 and 2 used the same heavy chain, but had different light chains. The amino acid sequences of the heavy and light chains of the two rabbit antibodies are shown below the Western blotting results, with the CDR3 regions of the heavy and light chains being underlined.

Figures 10a-e. Affinity purification of progesterone receptor-specific polyclonal rabbit $\operatorname{IgG}$. (a) Total IgG from the serum of the immunized rabbit was isolated with Protein A and further affinity purified on immobilized antigen peptides by gravity flow. After extensive washing to reduce non-specific IgG, a sequential elution with progressively acidic pH was used to fractionate the antigen-specific polyclonal IgG. Each fraction was tested for specific activity by Western blotting at matched antibody concentration $(21.5 \mathrm{ng} / \mathrm{ml})$ to detect PR A/B in lysates from T47D cells (+). Negative control lysates from HT1080 (-) were also tested. (b). The fraction with the highest specific activity, pH 1.8 , was processed with four proteases for LC-MS/MS analysis. (c). An MS/MS spectrum matched by SEQUEST to the V-region full tryptic peptide GFALWGPGTLVTVSSGQPK (SEQ ID NO: 305) containing CDRH3 (underlined) with an XCorr of 5.560 and a $\Delta \mathrm{M}$ (observed $m / z$ expected $\mathrm{m} / \mathrm{z}$ ) of 0.39 ppm . (d). MS/MS spectra were mapped to V -region peptides by SEQUEST and filtered to an FDR of $\leq 2 \%$. High confidence peptides were then remapped to the V-region database generated by NGS, taking into account the protease used for sample preparation and keeping track of the total number of peptides, the unique number of peptides, the spectrum share, and the amino acid coverage of the entire V-region. High coverage V-region sequences were selected, expressed as monoclonal antibodies, and screened for desired activity. (e). Heavy and light chain sequence identification coverage of clone F9. The depicted V-region sequences, when paired, specifically bind human PR A/B (see Figure 11a-e). Amino acids mapped by one or more peptides are shown in bold. To maximize V-region coverage and account for highly variable amino acid composition, complementary proteases were used (Chymotrypsin, Elastase, Pepsin, Trypsin.

Figures 11a-e. Identification and characterization of functional monoclonal antibodies against progesterone receptor A/B. (a). Combinatorial pairing of heavy and light
chains yielded 12 antigenspecific ELISA-reactive clones indicated in yellow. CDR3 sequence is used as an identifier: $\checkmark$ indicates Western blot-positive clones (See Fig.11b). (b). Six clones (F1 F9, H1, C1, F7, and H9) were specific for progesterone receptor A/B detection by Western blotting. Clones E6 (ELISA-negative, Western-negative) and H7
(ELISA-positive, Westernnegative) are shown as controls. +, T47D (PR A/B-positive); -, MDA-MB-231 (PR A/B-negative). All antibodies tested at $21.5 \mathrm{ng} / \mathrm{mL}$. (c). Comparison of specific activity of clone F9 to the affinity-purified polyclonal mixture by immunohistochemistry. $0.4 \mathrm{ug} / \mathrm{mL}$ of F9 specifically stained PR A/B-positive tissue or cell lines (T47D and MCF-7), but not a PR A/B-negative cell line (MDA-MB-231). $0.2 \mu \mathrm{~g} / \mathrm{mL}$ of polyclonal antibody was used as positive control. (d). Flow cytometry analysis. Blue, T47D cells (progesterone receptor A/B positive cell line); Black, MDA-MB-231 (progesterone receptor $\mathrm{A} / \mathrm{B}$ negative cell line). Polyclonal antibody signal/noise ratio, 1.69; concentration, $3.7 \mu \mathrm{~g} / \mathrm{mL}$. Monoclonal antibody F9 signal/noise ratio $=36.4$; concentration $0.5 \mu \mathrm{~g} / \mathrm{mL}$. (e). Confocal immunofluorescence microscopy analysis showed specific nuclear staining pattern on progesterone receptor $\mathrm{A} / \mathrm{B}$ positive cell line MCF-7 but not on MDA-MB-231 cells at $0.46 \mu \mathrm{~g} / \mathrm{mL}$. No primary antibody was included as background staining control. Polyclonal antibodies were also used as comparison at a concentration of $1.85 \mu \mathrm{~g} / \mathrm{mL}$.

Figures 12a-d. Characterization of clone C3 anti-Lin28A monoclonal antibody. (a) Combinatorial pairing of heavy and light chains yielded 5 antigen-specific ELISA-reactive clones indicated in shade. $\sqrt{ }$ indicates Western blot-positive clones. CDR3 sequence is used as an identifier. (b) Western blot analysis was performed using various Lin28A positive cell lysates, NCCIT, NTERA, mMES, and IGROV1. (c) Confocal immunofluorescence analysis was performed with Lin28A negative cells (HeLa) and Lin28A positive cells (NTERA). (d) Flow cytometry analysis of monoclonal antibody. Left peak, HeLa cells (Lin28A -); right peak, NTERA cells (Lin28A +). *V-regions had the same CDR3 sequence but not identical V region sequences.

Figures 13a-c. Identification and characterization of functional mouse monoclonal antibodies against phospho-Erk. (a) Purification of phospho-Erk polyclonal antibodies from the pooled sera of three mice. The pooled sera, protein G-purified total IgG from the pooled
sera, the unbound fraction from the protein G purification, and acid elution fractions of $\mathrm{pH} 3.5,2.7$ and 1.8 were assayed by Western blotting for binding specificity against phospho-Erk in Jurkat cell lysate. +, Jurkat cells stimulated with TPA; -, Jurkat cells treated with U0126. (b) Combinatorial pairing of heavy and light chains yielded 15 clones, indicated in shade, that are reactive by peptide antigen ELISA. $\sqrt{ }$ indicates Western blotpositive clones (See (c)). CDR3 sequence is used as an identifier. For the heavy chain sequences the underlined portion indicates the end of Frame Work Region 3. (c) Three clones (C10, F10 and M3) were specific for phospho-Erk detection by Western blotting. Clone C9 (ELISA positive, Western-negative) is shown as a control. All antibodies were tested at $100 \mathrm{ng} / \mathrm{mL}$.

## Detailed Description

This disclosure is directed to methods and systems for rapidly and accurately obtaining the amino acid sequences (and encoding nucleic acid sequences) of monoclonal antibodies that specifically bind to an antigen of interest. More specifically, the present methodology involves a direct, mass spectrometry-based proteomic investigation of circulating polyclonal antibodies from the serum of an animal, against a genetic material database which is comprised of nucleic acid molecules encoding full length immunoglobulin chains or variable regions. In specific embodiments, the genetic material database is generated from the B cell repertoire of an animal (e.g., the same animal whose serum was collected to obtain the polyclonal antibodies) by utilizing nucleic acid sequencing technologies. Thus, the present approach essentially involves correlating (i.e., crosscomparing or cross-referencing) the information from two sources: mass spectra information from the actual circulating polyclonal antibodies of an animal, and information (including, e.g., predicted mass spectra) from the genetic material database. A list of heavy and light chain DNA sequences can then be identified from the genetic material database that correspond to actual antibodies from the serum. Such heavy and light chains can be expressed in pairs to obtain functional monoclonal antibodies.

In some embodiments, the present methodology does not require B cell immortalization, single cell sorting and molecular cloning, or phage display, and does not
involve assembly of antibody sequences based on guesswork. By leveraging the strengths of both mass spectrometry technologies and nucleic acid sequencing technologies (such as Next Generation DNA Sequencing or NGS), the approach of this invention can significantly reduce the amount of time needed to isolate the sequences of antigen-specific monoclonal antibodies from a polyclonal population, thereby enabling a faster transition to recombinant antibodies such as fully human antibodies or humanized antibodies (e.g., humanized murine antibodies) that may be used therapeutically.

Furthermore, the present methodology is capable of identifying rare antibodies likely missed by existing technologies. The inventors have surprisingly found that individual antibodies with very selective specificity (e.g., an antibody that specifically binds to a phosphorylated tyrosine residue within a polypeptide) may occur very rarely within a polyclonal population. Methods that rely on the frequencies of antibody-encoding mRNAs and PCR amplification may miss these antibodies because their variable chains occur with low frequency. In contrast, the present methodology utilizes, for example, mass spectrometry based proteomics analysis of actual peptide fragments derived from a polyclonal antibody population, and therefore does not suffer from the errors of frequency following PCR amplification.

In addition, the present methodology allows for the rapid creation of novel antigenspecific antibodies that may not exist in the starting polyclonal population. For example, the created immunoglobulin molecule that has the highest desired qualities (e.g., highest binding affinity (or lowest KD) for the antigen or a desired isotype (e.g., IgG2a)) may be the result of a light chain from a first antibody in the polyclonal population assembled with a heavy chain of a second antibody (i.e., different from the first antibody) in the polyclonal population.

The methods described herein have applications in basic immunology and therapeutics. For example, the methods can provide the basis for understanding central questions in the field of immunology, including serum antibody diversity, dynamics, kinetics, clonality, and migration of B cells following antigen exposure. The methods can also be used to pursue therapeutically relevant human monoclonal antibodies from immunized, naturally infected, or diseased individuals.

As demonstrated herein, the present methodology has been successfully applied to several different antigens in both laboratory animal species and human, and has led to the isolation of monoclonal antibodies with antigen-specific activities that recapitulate or surpass those of the original affinity-purified polyclonal antibodies found in the serum of immunized subjects.

Accordingly, this disclosure further provides isolated recombinant monoclonal antibodies specific for an antigen, including therapeutic antibodies specific for a disease antigen, as well as therapeutic methods for treating a disease based on administration of therapeutic monoclonal antibodies.

The various aspects and embodiments of the invention are described in more detail below. The patents, published applications, and scientific literature referred to herein establish the knowledge of those with skill in the art and are hereby incorporated by reference in their entirety to the same extent as if each was specifically and individually indicated to be incorporated by reference. Any conflict between any reference cited herein and the specific teachings of this specification shall be resolved in favor of the latter. Likewise, any conflict between an art-understood definition of a word or phrase and a definition of the word or phrase as specifically taught in this specification shall be resolved in favor of the latter.

## Definitions

As used herein, the following terms have the meanings indicated. As used in this specification, the singular forms "a," "an" and "the" specifically also encompass the plural forms of the terms to which they refer, unless the content clearly dictates otherwise. The term "about" is used herein to mean approximately, in the region of, roughly, or around. When the term "about" is used in conjunction with a numerical range, it modifies that range by extending the boundaries above and below the numerical values set forth. In general, the term "about" is used herein to modify a numerical value above and below the stated value by a variance of $20 \%$.

By "peptide" or "peptide fragment" is meant a short polymer formed from the linking individual amino acid residues together, where the link between one amino acid residue and
the second amino acid residue is called an amide bond or a peptide bond. A peptide comprises at least two amino acid residues. A peptide is distinguished from a polypeptide in that it is shorter. At least two peptides, linked together by an amide bond or peptide bond between the $\mathrm{C}^{\prime}$ terminal amino acid residue of one peptide and the $\mathrm{N}^{\prime}$ terminal amino acid residue of the second peptide, form a polypeptide in accordance with various embodiments of the invention.

By "polypeptide" is meant a long polymer formed from the linking individual amino acid residue, where the link between one amino acid residue and the second amino acid residue is called an amide bond or a peptide bond. A polypeptide comprises at least four amino acid residues; however, multiple polypeptides can be linked together via amide or peptide bonds to form an even longer polypeptide.

By "nucleic acid molecule" is meant a polymer formed from linking individual nucleotides (e.g., deoxyribonucleotides or ribonucleotides) together, where the link between one nucleotide and the other nucleotide is a covalent bond including, for example, a phosphodiester bond. Thus, the term includes, without limitation, DNA, RNA, and DNARNA hybrids.

By "nucleic acid sequence" is meant a nucleic acid sequence (or nucleotide sequence complementary thereto) that includes nucleotides that encode all or part of an immunoglobulin chain (e.g., a heavy chain or a light chain). In some embodiments, the nucleic acid sequence is genomic DNA (e.g., exonic DNA with or without intronic DNA). In some embodiments, the nucleic acid sequence is cDNA or some form of RNA (e.g., hn RNA, mRNA, etc.). In some embodiments, the nucleic acid sequence is an expressed nucleic acid sequence that will be either transcribed into a nucleic acid molecule (e.g., DNA transcribed into RNA) or translated into a polypeptide in a cell containing that nucleic acid sequence. Accordingly, an expressed nucleic acid molecule includes, without limitation, hnRNA, mRNA, cDNA, and genomic exon sequences. By "complementary" in terms of nucleic acid molecules simply means that two single-stranded nucleic acid molecules contain nucleotides that will form standard Watson-Crick basepairs to form a doublestranded nucleic acid molecule, whether that double-stranded molecule is DNA, RNA, or a DNA-RNA hybrid.

As used herein, by "B lymphocyte" is meant any white blood cell in which gene recombination (or gene rearrangement) has begun to occur at a locus containing an immunoglobulin chain-encoding gene. For example, human immunoglobulin genes occur on chromosome 14 (heavy chain locus), chromosome 2 (kappa light chain locus), and chromosome 22 (lambda light chain locus). If a human white blood cell has undergone a gene rearrangement event in an immunoglobulin chain locus (e.g., on chromosome 14, chromosome 2, or chromosome 22), that cell is considered a B lymphocyte. Accordingly, B lymphocytes include, without limitation, B cells, pre-B cells, pro-B cells including early pro- $B$ cells (e.g., where the D and J regions of the heavy chain genes have undergone rearrangement but the light chain gene are germline (i.e., are not rearranged)) and late pro-B cells (e.g., where the $\mathrm{V}, \mathrm{D}$, and J regions of the heavy chain gene is rearranged but the light chain gene is still germline and where no immunoglobulin proteins are expressed on the cell surface), pre-B cells including large pre-B cells and small pre-B cells, immature $B$ cells, active B cells, germinal center B cells, plasma cells (including plasmablasts), and memory B cells.

Throughout the specification and the claims, the terms "antibody" and "immunoglobulin" are used interchangeably and are meant to include intact immunoglobulin polypeptide molecules of any isotype or sub-isotype (e.g., $\operatorname{IgG}, \operatorname{IgG} 1, \operatorname{IgG} 2, \operatorname{IgG} 2 \mathrm{a}, \operatorname{IgG} 2 \mathrm{~b}$, $\operatorname{IgG} 3, \operatorname{IgG} 4, \operatorname{IgM}, \operatorname{IgD}, \operatorname{IgE}, \operatorname{IgE} 1, \operatorname{IgE} 2, \operatorname{IgA})$ from any species of animal such as primates (e.g., human or chimpanzees), rodents (e.g., mice or rats), lagomorphs (e.g., rabbits or hares), livestock animals (e.g., cows, horses, goats, pigs, and sheep), fish (e.g., sharks), birds (e.g., chickens) or camelids (e.g., camels or llamas) or from transgenic non-human animals (e.g., rodents) genetically engineered to produce human antibodies (see, e.g., Lonberg et al., WO93/12227; U.S. Pat. No. 5,545,806; Kucherlapati, et al., WO91/10741; U.S. Pat. No. 6,150,584; US 2009/0098134; US 2010/0212035; US 2011/0236378; US 2011/0314563; WO2011/123708; WO2011/004192; WO2011/158009); antigen binding domain fragments thereof, such as Fab, Fab ', $\mathrm{F}\left(\mathrm{ab}^{\prime}\right)_{2}$; variants thereof such as $\mathrm{scFv}, \mathrm{Fv}, \mathrm{Fd}$, dAb, bispecific scFvs, diabodies, linear antibodies (see U.S. Pat. No. 5,641,870; Zapata et al., Protein Eng. 8 (10): 1057-1062.1995); single-chain antibody molecules; and multispecific antibodies formed from antibody fragments; and any polypeptide comprising a binding domain which
is, or is homologous to, an antibody binding domain (defined herein elsewhere). Nonlimiting antibodies of various embodiments of the invention include but are not limited to polyclonal, monoclonal, monospecific, polyspecific antibodies and fragments thereof, chimeric antibodies comprising an immunoglobulin binding domain fused to another polypeptide, and humanized antibodies such as a non-human antibody (e.g., a rabbit antibody) whose constant and/or FR domains have been replaced with constant and/or FR domains from a human antibody (see, e.g., U.S. Pat. Nos: 5,530,101; 5,585,089; 5,693,761; $5,693,762 ; 6,180,370$; and $6,548,640$ ). Transgenic non-human animals genetically engineered to produce human (e.g., at least partially human) antibodies are available from Harbour Antibodies (Rotterdam, The Netherlands), Ablexis (San Francisco, CA), Kymab Ltd (Cambridge, UK), OMT, Inc. (Palo Alto, CA), Amgen (Thousand Oaks, CA), Medarex (Princeton, NJ), and Regeneron (Tarrytown, NY).

Naturally-occurring intact antibodies are made up of two classes of polypeptide chains, light chains and heavy chains. A non-limiting antibody of various aspects of the invention can be an intact, four immunoglobulin chain antibody comprising two heavy chains and two light chains. The heavy chain of the antibody can be of any isotype including $\operatorname{IgM}$, $\operatorname{IgG}, \operatorname{IgE}, \operatorname{IgA}$ or $\operatorname{IgD}$ or sub-isotype including $\operatorname{IgG} 1, \operatorname{IgG} 2, \operatorname{IgG} 2 a, \operatorname{IgG} 2 b$, $\operatorname{IgG} 3, \mathrm{IgG} 4, \mathrm{IgE} 1, \mathrm{IgE} 2$, etc. The light chain can be a kappa light chain or a lambda light chain. For example, a single IgG naturally-occurring (or intact) antibody comprises two identical copies of a light chain and two identical copies of an IgG heavy chain. The heavy chains of all naturally-occurring antibodies, where each heavy chain contains one variable domain $\left(\mathrm{V}_{\mathrm{H}}\right)$ and one constant domain $\left(\mathrm{C}_{\mathrm{H}}\right.$, which itself comprises the CH 1 region, the hinge region, the CH 2 region, and the CH 3 region), bind to one another via multiple disulfide bonds within their constant domains to form the "stem" of the antibody. The light chains of all naturally-occurring antibodies, where each light chain contains one variable domain ( $\mathrm{V}_{\mathrm{L}}$ ) and one constant domain $\left(\mathrm{C}_{\mathrm{L}}\right)$, each bind through its constant domain to one heavy chain constant domain via disulfide binding. A schematic of a four immunoglobulin chain antibody (e.g., an IgG antibody) is shown in Figure 1. In Figure 1, the three CH domains are shown in light blue, the single VH domain is shown in dark blue, the single CL domain is shown in light pink and the single VL domain is shown in dark pink. As shown in Figure 1,
the VL and the VH domains of the light and heavy chains, respectively, come together to form the antibody binding domain.

In some embodiments, an intact immunoglobulin chain (e.g., a heavy chain or a light chain) may comprise in order from 5' to $3^{\prime}$ (for a nucleic acid sequence encoding the chain) or from the amino terminus to the carboxy terminus (for the amino acid sequence of the chain): a variable domain and a constant domain. The variable domain may comprise three complementarity determining regions (CDRs; also called hypervariable regions or HVs ), with interspersed framework (FR) regions. The variable domains of both the light chains and heavy chains contain three hypervariable regions sandwiched between four more conserved framework regions (FR), for a structure of $5^{\prime}$ (or $\mathrm{N}^{\prime}$ )- FR1, CDR1, FR2, CDR2, FR3, CDR3, FR4 3' (or C'), with the constant region 3' (or C') to the FR4 region. The CDRs form loops that comprise the principal antigen binding surface of the antibody (see Kabat, E.
A. et al., Sequences of Proteins of Immunological Interest, National Institutes of Health, Bethesda, Md., (1987) and Wu, T.T. and Kabat, E.A. (1970) J. Exp. Med. 132: 211-250 (1970)) with the four framework regions largely adopting a beta-sheet conformation and the CDRs forming loops connecting, and in some cases forming part of, the beta-sheet structure. The CDRs in each chain are held in close proximity by the framework regions and, with the CDRs from the other chain, contribute to the formation of the antigen binding domain.

By "antigen" is meant a target molecule (e.g., a polypeptide or a carbohydrate) that can be specifically bound by an antibody. The portion of an antigen that is specifically bound by the antibody is referred to as an "epitope". An "epitope" is smallest portion of a target molecule capable being specifically bound by the antigen binding domain of an antibody. The minimal size of an epitope may be about three to seven amino acids (e.g., five or six amino acids). There may be multiple epitopes on a single antigen, thus, a single antigen can be specifically bound by multiple different antibodies, all of which antibodies specifically bind the antigen (i.e., all of these antibodies are antigen-specific antibodies) even though each individual antibody specifically binds to a different epitope on the antigen.

By "disease antigen" is meant an antigen which arises in an animal during a disease state. For example, a viral antigen (e.g., an antigen encoded by a nucleic acid molecule of a virus's genetic material) is a disease antigen in animal infected with that virus. Similarly,
some diseases (e.g., cancer) are characterized by gene translocations which produce chimeric proteins (e.g., BCR-ABL). Thus, a BCR-ABL protein is a disease antigen. It should be understood that a disease antigen is not necessarily seen only in an animal suffering from that disease.

By "disease" is simply meant any abnormal condition affecting an animal. Nonlimiting examples of diseases include, without limitation, autoimmune disease (e.g., rheumatoid arthritis or type I diabetes), cancer (e.g., leukemia, colon cancer, or prostate cancer, etc.), viral infections (e.g., AIDS caused by infection of the HIV virus or chicken pox caused by infection of the varicella zoster virus), parasitic infection (e.g., schistosomiasis or scabies), and bacterial infection (e.g., tuberculosis or diptheria).

By "specifically bind" is meant that an immunoglobulin or antibody interacts with its antigen (i.e., its specific antigen), where the interaction is dependent upon the presence of a particular structure (e.g., an epitope) on the antigen; in other words, the antibody is recognizing and binding to a specific structure rather than to all molecules or structures in general. An antibody that specifically binds to the antigen may be referred to as an "antigen-specific antibody" or an "antibody specific for the antigen". In some embodiments, an antibody that specifically binds to antigen can immunoprecipitate that antigen from a solution containing the antigen as well as other molecules (e.g., a cell lysate). In some embodiments, an antibody that specifically binds to its antigen has a $K_{D}$ for its antigen of 1 x $10^{-6} \mathrm{M}$ or less. In some embodiments, an antibody that specifically binds to its antigen has a $K_{D}$ for its antigen of $1 \times 10^{-7} \mathrm{M}$ or less, or a $K_{D}$ of $1 \times 10^{-8} \mathrm{M}$ or less, or a $K_{D}$ of $1 \times 10^{-9} \mathrm{M}$ or less, or a $K_{D}$ of $1 \times 10^{-10} \mathrm{M}$ or less, of a $K_{D}$ of $1 \times 10^{-11} \mathrm{M}$ or less, of a $K_{D}$ of $1 \times 10^{-12} \mathrm{M}$ or less. In certain embodiments, the $\mathrm{K}_{\mathrm{D}}$ of an antibody that specifically binds to its antigen for its specific antigen is between 1 pM to 500 pM , or between 500 pM to $1 \mu \mathrm{M}$, or between 1 $\mu \mathrm{M}$ to 100 nM , or between 100 mM to 10 nM . As used herein, by the term " $\mathrm{K}_{\mathrm{D}}$ ", is intended to refer to the dissociation constant of an interaction between two molecules (e.g., the dissociation constant between an antibody and its specific antigen).

By "variable region of an immunoglobulin chain" or an "immunoglobulin chain variable region" is a polypeptide comprising at least a portion of the variable domain of a heavy (i.e., the VH domain) or a light chain (i.e., the VL domain) of an immunoglobulin,
where the portion of the VL and the VH domains form an antigen binding domain of an immunoglobulin (see Fig. 1). Thus, the variable region of an immunoglobulin may include, without limitation, a single CDR (e.g., CDR1), two CDRs interspersed with a single FR (e.g., CDR1, FR2, and CDR2), three CDRs interspersed with two FRs (e.g., CDR1, FR2, CDR2, FR3, and CDR3), or three CDRs flanked by either or both of FR1 and FR4 (e.g., FR1, CDR1, FR2, CDR2, FR3, CDR3, FR4). In some embodiments, the immunoglobulin chain variable region is the region on one of either the heavy or the light chain which, when combined with the immunoglobulin chain variable region of the other chain (i.e., the light or the heavy chain) of the intact immunoglobulin, forms the antigen binding domain.

By "antigen binding domain" is meant the region of a single heavy chain assembled with a single light chain in an immunoglobulin, which retains the specific binding activity of the intact antibody for its specific antigen. Thus, an intact IgG immunoglobulin, which comprises two heavy chains and two light chains, has two antigen binding domains. Likewise, fragmentation of an intact antibody which retains a covalent bond between the heavy chain and the light chain will also result in an immunoglobulin fragment having an antigen binding domain. For example, digestion of an immunoglobulin with the enzyme papain will generate $\mathrm{F}(\mathrm{ab})$ fragments, each of which has a single antigen binding domain. Of course the entire $\mathrm{F}(\mathrm{ab})$ is not the antigen binding domain; rather, only the portion of the $\mathrm{F}(\mathrm{ab})$ fragment which retains the ability to specifically bind the antigen is the antigen binding domain.

Technical and scientific terms used herein have the meaning commonly understood by one of skill in the art to which the present invention pertains, unless otherwise defined. Reference is made herein to various methodologies and materials known to those of skill in the art. Standard reference works setting forth the general principles of antibody and/or recombinant DNA technologies include Harlow and Lane, Antibodies: A Laboratory Manual, Cold Springs Harbor Laboratory Press, Cold Spring Harbor, New York (1988); Sambrook et al., Molecular Cloning: A Laboratory Manual, 2nd Ed., Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y. (1989); Coligan et al., Current Protocols in Immunology, John Wiley \& Sons, New York, NY (1991-2010); Ausubel et al., Current Protocols in Molecular Biology, John Wiley \& Sons, New York, NY (1987-2010); Kaufman
et al., Eds., Handbook of Molecular and Cellular Methods in Biology in Medicine, CRC Press, Boca Raton (1995); McPherson, Ed., Directed Mutagenesis: A Practical Approach, IRL Press, Oxford (1991); all of which are incorporated by reference in their entirety. Standard reference works setting forth the general principles of pharmacology include Companies Inc., New York (2006), which is incorporated by reference in its entirety.

## Methods for Obtaining Sequences of Antigen-Specific Immunoglobulins

In one aspect, this invention is directed to a method for obtaining the amino acid and/or nucleic acid sequences of immunoglobulin chains (or variable regions thereof) of a single immunoglobulin from a population of polyclonal antibodies.

According to the present method, a population of polyclonal antibodies of interest is obtained from an animal and fragmented to generate peptide fragments which are analyzed by mass spectrometry. The mass spectra information observed from the peptide fragments is then correlated with predicted mass spectra information derived from a genetic material database comprised of nucleic acid sequences that encode full-length immunoglobulin heavy and/or light chains (or variable regions thereof). As a result of such correlating, immunoglobulin heavy and/or light chains (or variable regions thereof) can be identified from the genetic material database that correspond to immunoglobulin heavy and/or light chains (or variable regions thereof) of immunoglobulin molecules within the starting polyclonal antibody population.

The various aspects of the present method are described in more detail below.

## The Starting Population of Polyclonal Antibodies

Immunoglobulins that specifically bind to an antigen of interest may be collected from an animal, which includes any mammal, such as human. Immunoglobulins can be collected from a body fluid sample of the animal including, for example, blood, serum or plasma of the blood, cerebrospinal fluid, synovial fluid, peritoneal fluid, mucosal secretions, tears, nasal secretions, saliva, milk, and genitourinary secretions.

In some embodiments, immunoglobulins need not come from a single individual
animal but, rather, may be a cocktail of different antibodies (monoclonal or polyclonal) taken from different individuals. In some embodiments, the immunoglobulins are collected from a transgenic non-human animal, e.g., a transgenic non-human animal that expresses human antibody sequences and/or produces antibodies that are at least partly human. either because the animal from whom the immunoglobulins are collected was previously immunized with the antigen, or because the animal from whom the immunoglobulins are collected was previously exposed to a condition whereby the animal was likely to generate antigen-specific antibodies. In an example of the latter case, the animal may have been infected with a virus (e.g., Epstein Barr Virus), where the antigen of interest is the EBNA1 protein, which is encoded by the genome of the Epstein Barr Virus.

In various embodiments, the animal whose immunoglobulins are collected (i.e., obtained) is of the same species as the animal whose B lymphocyte nucleic acid sequences are collected to create the reference database. In some embodiments, the animal whose immunoglobulins are collected for the peptide database and the animal whose B lymphocyte nucleic acids are collected for the reference database are the same animal.

As shown in Figure 2, where the animal is the same animal, blood taken from the animal can provide both the nucleic acid sequences (e.g., from the cells in the blood) and the polyclonal antibodies (e.g., from the sera or plasma of the blood).

The immunoglobulins collected from the animal form a polyclonal population of immunoglobulins, because different B lymphocytes produced members of the population. It should be noted that in such a polyclonal population, not all of the individual antibodies within that polyclonal population will specifically bind the same antigen. In fact, each of the antibodies within the population may bind a different antigen. However, this polyclonal population still is said to specifically bind a particular antigen if at least one individual antibody, preferably multiple antibodies, of the polyclonal population binds that antigen (see, e.g., Example 3 below). In another example, some antibodies in the polyclonal population may bind the antigen with low affinity. However, a polyclonal population is said to specifically bind an antigen if some (e.g., at least one or more) of the antibodies in that population specifically bind the antigen.

It should be noted that by the phrase "polyclonal antibody (or immunoglobulin) that specifically binds to an antigen" is meant that within the polyclonal population, at least one antibody specifically binds to the antigen, however that one antibody is not necessarily isolated from the other antibodies within the polyclonal population that do not specifically bind to the antigen. Of course in some embodiments, more than one different antibody within the polyclonal population specifically binds to the antigen.

It should also be noted that different antibody molecules are antibody molecules produced by a different B cell. For example, after collecting sera, a polyclonal population of 1000 antibody molecules may be isolated from the sera (e.g., using the antibodies' adherence to a protein A column to isolate the antibodies from the other sera components). Within that population of 1000 antibody molecules, 900 may be identical (i.e., secreted by the same B cell) and thus there are really only 101 different antibodies within that polyclonal population. Regarding a polyclonal population, if all 900 identical antibody molecules specifically bind the antigen, the polyclonal population of 1000 antibody molecules is a polyclonal antibody that specifically binds to the antigen. Similarly, if an additional 5 different antibody molecules of the remaining 100 different antibody molecules also specifically bind to the antigen, the polyclonal population of 1000 antibody molecules is likewise is a polyclonal antibody that specifically binds to the antigen.

The majority of antibody molecules within a polyclonal population need not specifically bind to an antigen for that population to be referred to as a "polyclonal antibody that specifically binds to the antigen". For example, if within a polyclonal population of 1000 antibody molecules, even if only 1 antibody molecule specifically binds to the antigen and 999 antibody molecules do not, that population of 1000 antibody molecules is still a "polyclonal antibody that specifically binds to the antigen" as the term is used herein.

Note also that all of the antibodies in a polyclonal antibody population need not bind the same epitope on the antigen. For example, a polyclonal population can be specific for the antigen where every different antibody within the population specifically binds a different epitope on the antigen.

In various embodiments of the non-limiting methods of the invention, the population of polyclonal immunoglobulins may have, for example, at least two different
immunoglobulins within the population, or at least three, or at least five, or at least ten, or at least twenty, or at least fifty, or at least one hundred or at least five hundred different immunoglobulins within the population.

The invention also contemplates collecting a polyclonal population of the nucleic acid sequences are collected from the B cells themselves). For example, a population of B cells may be collected from an animal that has been subjected to the Epstein Barr virus. The population can be expanded, e.g., to enrich B lymphocytes in the population as compared to other white blood cells. From this cultured media of these cells (into which the polyclonal antibodies are secreted by the cells), the polyclonal population of antibodies can be isolated.

The polyclonal population of immunoglobulins collected, either from an animal(s) or from tissue culture supernatants of B cells, can be first purified prior to digestion into peptide fragments. For example, the collected polyclonal antibodies can be subjected to a protein A or protein G sepharose column, which can separate antibodies from other blood sera proteins, for example. See, for example, Figure 2 and Figure 3. Alternatively or additionally, the collected polyclonal antibodies are subjected to antiegn affinity purification to enrich for antibodies with high specific activity. While not entirely necessary, a purification step, especially antigen affinity purification, can reduce the complexity of a polyclonal mixture and ultimately reduce the number of potential false positive or negative candidate immunoglobulins. The collected polyclonal antibodies may be concentrated or buffer exchanged or both, either before or after purification.

In one illustrative embodiment, to collect immunoglobulins that specifically bind to an antigen of interest from an animal, peripheral blood is drawn from the animal, and serum and/or plasma antibodies are collected according to standard methods (e.g., adherence of the antibodies to protein A). The serum and/or plasma antibodies are then purified or screened to enrich for immunoglobulins that specifically bind to the antigen. This screen can be, for example, by coating a solid-phase surface (e.g., a sepharose bead or bottom of a plastic well) with antigen and pass the serum and/or plasma over the antigen-coated surface under conditions where immunoglobulins that specifically bind to the antigen will bind. The
bound antibodies may be treated with a protease (e.g., papain) or a chemical protein cleavage reagent that specifically cuts near the hinge region of the immunoglobulin to remove the non-adherent Fc portions. After rinsing away non-binding serum and/or plasma proteins (including non-specific immunoglobulins), the antigen-specific immunoglobulins can be collected and their quantities thus enriched as compared to antibodies that do not specifically bind to the antigen.

## Observed Mass Spectra From the Collected Polyclonal Antibodies

To obtain observed (i.e., actual) mass spectra, the collected polyclonal antibodies (or fragments thereof) are analyzed by protein analysis methods (e.g., mass spectrometry, liquid chromatography, etc.).

In some embodiments, observed mass spectra information is obtained from peptide fragments which are generated from the polyclonal antibodies. The polyclonal antibodies can be fragmented, for example, with one or more proteases, and/or a chemical protein cleavage reagent, such as cyanogen bromide.

Certain proteases are known to cleave their substrates at specific sites. Table 1 provides a non-comprehensive list of commonly used proteases and their cleavage sites (in 3 letter amino acid code).

Table 1

| Protease | Cleavage Site |
| :--- | :--- |
| Trypsin | cleaves after (i.e., on the carboxyl side of) Arg or Lys, <br> unless followed by Pro |
| Chymotrypsin | cleaves after Phe, Trp, or Tyr, unless followed by Pro |
| Elastase | cleaves after Ala, Gly, Ser, or Val, unless followed by <br> Pro. |
| Endoproteinase Lys-C | cleaves after Lys |
| Pepsin | cleaves after Phe or Leu. |
| Thermolysin | cleaves before Ile, Met, Phe, Trp, Tyr, or Val, unless <br> preceded by Pro. |
| Endopeptidase V8 (alias Glu- <br> C) | cleaves after Glu. |

A more comprehensive listing of proteases that can be used to digest proteins to smaller fragments is given in Tables 11.1.1 and 11.1.3 of Riviere and Tempst (Riviere LR, Tempst P. Enzymatic digestion of proteins in solution. Curr Protoc Protein Sci. 2001 May; Chapter 11:Unit 11.1. PubMed PMID: 18429101; herein incorporated by reference in its entirety).

In specific embodiments, multiple (i.e., two or more) proteases are used (e.g., independently or together) to digest the polyclonal antibodies to maximize V -region coverage and account for highly variable amino acid compositions of immunoglobulins. For example, a combination of chymotrypsin, elastase, pepsin and trypsin can be used, as illustrated in Example 7 herein. In some embodiments, a protease or proteases are chosen on the basis that they do not cleave within predicted CDR3 regions based on analysis of the nucleic acid molecules in the genetic material database.

Proteins may be digested to smaller fragments that are amenable to mass spectrometry by treatment with particular chemical protein cleavage reagents rather than proteolytic enzymes. See for example chapter 3 of G. Allen. Sequencing of Proteins and Peptides, Laboratory Techniques in Biochemistry and Molecular Biology, Vol. 9. Elsevier 1989. Such chemical protein cleavage reagents include, without limitation, cyanogen bromide, BNPS-skatole, o-iodosobenzoic acid, dilute acid (e.g., dilute HCl ), and so forth. For example, proteins can be cleaved at Met residues with cyanogen bromide, at Cys residues after cyanylation, after Trp residues with BNPS-skatole or o-iodosobenzoic acid, etc. Protein fragments can also be generated by exposure to dilute acid, e.g., HCl . An example of the use of partial acid hydrolysis to determine protein sequences by mass spectrometry is given by Zhong et al. (Zhong H, et al., J. Am. Soc. Mass Spectrom. 16(4):471-81, 2005. PubMed PMID: 15792716, incorporated by reference in its entirety). Zhong et al., supra used microwave-assisted acid hydrolysis with $25 \%$ trifluoroacetic acid in water to fragment bacteriorhodopsin for sequencing by mass spectrometry. See also Wang N, and Li L., J. Am. Soc. Mass. Spectrom. 21(9):1573-87, 2010.PubMed PMID: 20547072 (herein incorporated by reference in its entirety).

Proteins can be fragmented to make them more amenable for mass spectrometry by treatment with one protease, by treatment with more than one protease in combination, by
treatment with a chemical cleavage reagent, by treatment with more than one chemical cleavage reagent in combination, or by treatment with a combination of proteases and chemical cleavage reagents. The reactions may occur at elevated temperatures or elevated pressures. See for example López-Ferrer D, et al., J. Proteome. Res. 7(8):3276-81, 2008.
reference in its entirety). The fragmentation can be allowed to go to completion so the protein is cleaved at all bonds that the digestion reagent is capable of cleaving; or the digest conditions can be adjusted so that fragmentation does not go to completion deliberately, to produce larger fragments that may be particularly helpful in deciphering antibody variable region sequences; or digest conditions may be adjusted so the protein is partially digested into domains, e.g., as is done with E. coli DNA polymerase I to make Klenow fragment. The conditions that may be varied to modulate digestion level include duration, temperature, pressure, pH , absence or presence of protein denaturing reagent, the specific protein denaturant (e.g., urea, guanidine HCl , detergent, acid-cleavable detergent, methanol, acetonitrile, other organic solvents), the concentration of denaturant, the amount or concentration of cleavage reagent or its weight ratio relative to the protein to be digested, among other things.

In some embodiments, the reagent (i.e., the protease or the chemical protein cleavage reagents) used to cleave the proteins is a completely non-specific reagent. Using such a reagent, no constraints are made may be made at the N -terminus of the peptide, the C terminus of the peptide, or both of the N - and C -termini. For example, a partially proteolyzed sequence that is constrained to have a tryptic cleavage site at one end of the peptide sequence or the other, but not both, may be used in the various methods described herein.

The resulting peptide fragments can be detected and analyzed using an HPLC coupled to a mass spectrometer from which observed mass spectra are generated. This method may be referred to as a "bottom up" proteomics approach, where proteome components are separated and identified after reducing the proteins to relatively small peptides, e.g., 3 to 45 residues in length.

In other embodiments, an alternative, "top down" proteomics approach can be
employed to obtain observed mass spectra, which involves mass spectrometry analysis of intact proteins or large protein fragments or protein domains or large polypeptides. For example, to identify the parts of the antibody variable regions that bestow specific antigen recognition to a particular polyclonal antibody molecule, it is helpful to sequence large portions of the variable regions to identify its CDRs, by direct analysis of fragments large enough that the CDRs remain linked together.

For a review describing both "bottom up" and "top down" strategies, see Han X, Aslanian A, Yates JR 3rd. Mass spectrometry for proteomics. Curr Opin Chem Biol. 2008 Oct;12(5):483-90. Review. PubMed PMID: 18718552; PubMed Central PMCID: PMC2642903 (incorporated by reference in its entirety). For a recent review of top-down proteomics applied to determining antibody sequences, see Zhang Z. et al., Mass Spectrom Rev. 2009 Jan-Feb;28(1):147-76. Review. PubMed PMID: 18720354 (incorporated by reference in its entirety). For a recent paper showing extensive sequencing of a monoclonal antibody by top-down proteomics, see Tsybin et al, Anal Chem. 2011 Oct 21. PubMed PMID: 22017162 (incorporated by reference in its entirety).

In some embodiments of the above non-limiting method, while the antigen-specific immunoglobulins are bound to the antigen-coated surface, the immunoglobulins can be digested with either papain or pepsin to generate $\mathrm{F}(\mathrm{ab})$ and $\mathrm{F}(\mathrm{ab})_{2}$ fragments, respectively. Since the entirety of an immunoglobulin chain variable region is located on a chain of an $\mathrm{F}(\mathrm{ab})$ fragment, this pre-treatment with papain and/or pepsin will enrich for immunoglobulin chain variable regions. After rinsing away the non-binding portions of the immunoglobulins, the immunoglobulin chain variable regions can be collected.

After passage of the immunoglobulin fragments through the mass spectrometer, numerous observed mass spectra will be generated. However, given the potentially large number of different immunoglobulins within a polyclonal population, each with a different amino acid sequence, that are analysed with the mass spectrometer, the resulting observed mass spectra will be difficult to assemble back into a functional immunoglobulin chain variable region. In the methods of various embodiments of the invention, because the underlying nucleic acid sequence is available, there is no need to assemble the observed mass spectra data. Instead, the observed mass spectrum of a single peptide fragment can be
correlated with the predicted mass spectra of the nucleic acid sequence to obtain the amino acid (and underlying nucleotide) sequence of the entire immunoglobulin chain (or variable region thereof) of an immunoglobulin that specifically binds to an antigen from a starting polyclonal immunoglobulin population. This correlating step is further described hereinbelow.

In addition to mass spectra information, additional information derived from the peptide fragments of the polyclonal antibodies is useful in various embodiments of the invention. This information includes, without limitation, the mass of each peptide, the length (in amino acid residues) of each peptide, the observed mass spectrum of each peptide (e.g., from tandem mass spectrometry such as the MS2 or MS3 spectrum), the mass to charge ratio of each peptide, the ionic charge of each peptide, the chromatographic profile of each peptide, and the amino acid sequence of each peptide.

## Mass Spectrometry Analysis

In the methods of this invention, mass spectra information can be obtained by mass spectrometry analysis of collected immunoglobulins or fragments generated therefrom. A mass spectrometer is an instrument capable of measuring the mass-to-charge $(\mathrm{m} / \mathrm{z})$ ratio of individual ionized molecules, allowing researchers to identify unknown compounds, to quantify known compounds, and to elucidate the structure and chemical properties of molecules. In some embodiments, one begins mass spectrometry analysis by isolating and loading a sample onto the instrument. Once loaded, the sample is vaporized and then ionized. Subsequently, the ions are separated according to their mass-to-charge ratio via exposure to a magnetic field. In some embodiments, a sector instrument is used, and the ions are quantified according to the magnitude of the deflection of the ion's trajectory as it passes through the instrument's electromagnetic field, which is directly correlated to the ions mass-to-charge ratio. In other embodiments, ion mass-to-charge ratios are measured as the ions pass through quadrupoles, or based on their motion in three dimensional or linear ion traps or Orbitrap, or in the magnetic field of a Fourier transform ion cyclotron resonance mass spectrometer. The instrument records the relative abundance of each ion, which is used to determine the chemical, molecular and/or isotopic composition of the original
sample. In some embodiments, a time-of-flight instrument is used, and an electric field is utilized to accelerate ions through the same potential, and measures the time it takes each ion to reach the detector. This approach depends on the charge of each ion being uniform so that the kinetic energy of each ion will be identical. The only variable influencing velocity in this scenario is mass, with lighter ions traveling at larger velocities and reaching the detector faster consequently. The resultant data is represented in a mass spectrum or a histogram, intensity vs. mass-to-charge ratio, with peaks representing ionized compounds or fragments.

To obtain mass spectra data of a protein sample, the sample is loaded onto the instrument and ionized. Ionization can be done by, e.g., electrospray ionization and matrixassisted laser desorption/ionization ("MALDI"). See, e.g., Zenobi, "Ion Formation in MALDI Mass Spectrometry", 17 Mass Spectrometry Review, 337 (1998). Protein characterization can be done in one of two ways, top-down or bottom-up. The top-down approach involves ionizing intact proteins or larger protein fragments. See, e.g., Allison Doerr, "Top-down Mass Spectrometry", 5 Nature Methods, 24 (2008). The bottom-up approach involves enzymatically or chemically digesting the protein into constituent peptides using a protease. See Biran Chait, "Mass Spectrometry: Bottom-Up or TopDown?", 6 Science 65 (2006). The resultant peptides are introduced into the instrument and ultimately identified by peptide mass fingerprinting or tandem mass spectrometry.

In some embodiments, mass spectrometry analysis may be combined with a chromatographic fractionation (e.g., liquid chromatography).

Mass spectra data useful in this invention can be obtained by peptide mass fingerprinting. Peptide mass fingerprinting involves inputting the observed mass from a spectrum of the mixture of peptides generated by proteolytic digestion into a database and correlating the observed masses with the predicted masses of fragments arising from digestions of known proteins in silico. Known masses corresponding to sample masses provide evidence that the known protein is present in the sample tested.

Mass spectra data can be obtained by tandem mass spectrometry. In some embodiments, tandem mass spectrometry typically utilizes collision-induced-dissociation, which causes peptide ions to collide with gas and to fragment (e.g., due to vibrational energy
imparted by the collision). The fragmentation process produces cleavage products that break at the peptide bonds at various sites along the protein. The observed fragments' masses may be matched with a database of predicted masses for one of many given peptide sequences, and the presence of a protein may be predicted. See, e.g., Eng, 5 An Approach to Database, JASMS, 976 (1994).

In another embodiment, tandem mass spectrometry is performed by higher-energy collision induced dissociation (HCD), which on some mass spectrometers shows fragment product ions closer to peptide termini than collision induced dissociation. See Olsen JV, Macek B, Lange O, Makarov A, Horning S, Mann M. Higher-energy C-trap dissociation for peptide modification analysis. Nat. Methods. 2007 Sep;4(9):709-12. Epub 2007 Aug 26. PubMed PMID: 17721543.

In another embodiment, tandem mass spectrometry is performed by electron transfer dissociation (ETD), which is based on ion-ion reactions where a distinct reagent chemical ion donates a radical to a peptide ion, which then promptly fragments to form product ions. See Mikesh LM, Ueberheide B, Chi A, Coon JJ, Syka JE, Shabanowitz J, Hunt DF. The utility of ETD mass spectrometry in proteomic analysis. Biochim Biophys Acta. 2006 Dec;1764(12):1811-22. Epub 2006 Oct 30. Review. PubMed PMID: 17118725; PubMed Central PMCID: PMC1853258. Certain fragmentation methods, such as ETD, are particularly well-suited to "top down" proteomics strategies. Other fragmentation mechanisms are specific to certain ionization mechanisms, for example, such as post-source decay (PSD) is compatible with matrix-assisted laser desorption ionization (MALDI), and is also well-suited to "top down" proteomics strategies.

## Genetic Material Database

In accordance with the present invention, the observed mass spectra information from the starting polyclonal immunoglobulin population is correlated with predicted mass spectra information derived from a genetic material database, in order to obtain the amino acid (and underlying nucleotide) sequences of immunoglobulin chains (or variable regions thereof) of immunoglobulins from the starting polyclonal immunoglobulin population.

As used herein, a genetic material database includes nucleic acid sequences encoding a plurality of immunoglobulin chains (or variable regions thereof). Thus, information which can be obtained or derived from such a genetic material database includes, for example, the nucleotide sequence information of each nucleic acid molecule, the length (in nucleotides) of each nucleic acid molecule, amino acid sequence information of the polypeptides or peptides encoded by each nucleic acid molecule, the mass of a polypeptide or peptide encoded by each nucleic acid molecule, the length (in amino acid residues) of a polypeptide or peptide encoded by each nucleic acid molecule, the mass spectra information of polypeptides or peptides encoded by each nucleic acid molecule (e.g., a predicted mass spectra information based on the amino acid sequence of the polypeptide or peptide), and the amino acid sequence of a polypeptide or peptide encoded by each nucleic acid molecule.

In some embodiments of the invention, the genetic material database contains genetic information of nucleic acid sequences encoding full length immunoglobulin chains (and not just the variable regions thereof). In some embodiments, the nucleic acid sequences are expressed (i.e., transcribed into RNA and/or translated into protein) by the cell from which said sequences are derived. In specific embodiments, the genetic material database includes expressed nucleic acid sequences encoding immunoglobulin chain variable regions of multiple immunoglobulins from an animal. In some embodiments, the genetic material database contains at least one hundred different expressed nucleic acid sequences. In other embodiments, the genetic material database contains at least one thousand different expressed nucleic acid sequences.

Nucleic acid molecules encoding immunoglobulin chains (or the variable regions thereof) are readily obtainable from a population of cells (e.g., peripheral white blood cells) containing B lymphocytes. In some embodiments, the nucleic acid molecules are obtained from splenocytes or mononuclear cells, such as peripheral blood mononuclear cells (PBMCs). In some embodiments, the B lymphocytes are from a naïve animal (e.g., an animal that has not been exposed to the antigen to which an antigen-specific antibody is sought). In some embodiments, the naïve animal has been exposed to very few antigens (e.g., an animal raised in sterile or pathogen-free environment). In some embodiments, the naïve animal is a typical animal that has been exposed to typical antigens, but has not been
exposed to the antigen of choice.
In some embodiments, the animal from which the nucleic acid molecules encoding immunoglobulin chains (or the variable regions thereof) are obtained is an animal that has been previously exposed to the antigen. For example, the animal may be an animal immunized with the antigen (e.g., the antigen mixed with an adjuvant or an antigen coupled to an immunogenic carrier such as keyhole limpet hemocyanin (KLH)), may be an animal infected with a pathogen comprising the antigen (e.g., an animal infected with HIV virus when the antigen of choice is the HIV p24 antigen), or may otherwise be previously exposed to the antigen. In some embodiments, the animal is a bird (e.g., a chicken or turkey) or a mammal, such as a primate (e.g., a human or a chimpanzee), a rodent (e.g., a mouse, hamster, or rat), a lagomorph (e.g., a rabbit or hare), a camelid (e.g., a camel or a llama), or a domesticated mammal such as a companion animal (e.g., a cat, a dog, or a horse), or a livestock animal (e.g., a goat, sheep, or a cow).

It shall be understood that the nucleic acid sequences of the various aspects and embodiments of the invention need not come from a single animal. For example, some of the nucleic acid sequences of various embodiments of the invention may come from an animal previously exposed to an antigen, and some of the nucleic acid sequences may come from naïve animal. In some embodiments of the invention, nucleic acid sequences are from animals of a single species. For example, where there are multiple animals from which nucleic acid sequences are obtained, all of those animals may be the same species (e.g., all are rabbits or all are humans). In some embodiments, the nucleic acid sequences are obtained from animals of a single species. In other embodiments, nucleic acid sequences from more than one species of animal may be obtained. For examples, nucleic acid sequences may be obtained from mice and rats, and predicted mass spectra based from these sequences can be used to correlate with and/or compare to the actual mass spectra information of peptide fragment of polyclonal antibodies to create an immunoglobulin (or variable region, antigen binding domain, or chain thereof) that specifically binds to the antigen. In some embodiments, the nucleic acid sequences are obtained from animals of a single gender (e.g., all animals are female).

The animal from whom the polyclonal antibodies are collected and the animal from
whom the nucleic acid sequences are collected may be the same animal, or the same species of animal, or syngenic animals (e.g., both are Balb/c mice), or from animals of the same gender (e.g., both are female animals). The MS2 spectra from the antigen-binding components of the polyclonal antibodies can thus be correlated to the theoretical MS2 the nucleic acid sequences that encode antigen-binding antibodies.

It shall also be understood that the nucleic acid sequences and the polyclonal antibodies can be collected from cells of an animal where the cells were cultured in vitro following removal from the animal and prior to collection of the polyclonal antibodies (e.g., from the supernatant or cultured media of the cultured cells) and collection of the nucleic acid sequences from the cells. This culturing step is useful, e.g., to expand or enrich B lymphocytes as compared to other blood or tissue cells (e.g., to enrich B lymphocytes over red blood cells or epithelial cells). The number of individual nucleic acid sequences used to create theoretical mass spectra in the various embodiments of the invention is limitless. For example, five or ten or fifty, or one hundred, or one thousand, or one million, or one billion, or one trillion or more different nucleic acid sequences can be obtained and used to create theoretical mass spectra. The nucleic acid sequences may come from any source, and may be from a combination of sources. For example, nucleic acid sequences can be obtained by sequencing expressed nucleic acid molecules encoding immunoglobulin chain variable regions (or the entire full length immunoglobulin chain including the variable regions and constant region) as described herein. Nucleic acid sequences can also be obtained from genomic DNA that may or may not have undergone full V(D)J recombination. Nucleic acid sequences can also be obtained from publicly available sources. For example, numerous amino acid sequences (and nucleotide sequences) of immunoglobulin chain variable regions (and polynucleotides encoding the same) from multiple species of animal are known (see, for example, the following US and PCT patent publications (including issued US patents), the entirety of each of which is hereby incorporated by reference: US 20100086538; WO 2010/097435; US 20100104573; US 7,887,805; US 7,887,801; US 7,846,691; US 7,833,755; US 7,829,092.

The B lymphocytes from which nucleic acid sequences are obtained can be from any
blood or tissue source including, without limitation, bone marrow, fetal blood, fetal liver, sites of inflammation (e.g. inflamed joints surrounding synovial fluid in rheumatoid patients), tumors (e.g., tumor-infiltrating lymphocytes), peripheral blood, in lymph nodes, in peyer's patches, in tonsils, and in the spleen or in any lymphoid organ. In some embodiments, the entire tissue (e.g., bone marrow or lymph node) can be processed (e.g., cells separated from one another and lysed), genetic material removed, and the nucleic acid molecules encoding immunoglobulin chains (or variable regions thereof) sequenced.

In some embodiments, B lymphocytes are enriched from tissues or a population of cells (e.g., peripheral blood) containing them prior to isolating genetic material from the B lymphocytes. In accordance with various embodiments of the invention, methods for enriching B lymphocytes from an animal are well known. B lymphocytes can be found in many organs and areas of the body including, without limitation, bone marrow, fetal blood, fetal liver, sites of inflammation (e.g. inflamed joints surrounding synovial fluid in rheumatoid patients), tumors (e.g., tumor-infiltrating lymphocytes), peripheral blood, in lymph nodes, and in the spleen. From these tissue samples (e.g., peripheral blood or the spleen of an animal), white blood cells may be isolated according to standard methods (e.g., using the Ficoll-Paque PLUS or Ficoll-Paque PREMIUM reagents commercially available from GE Healthcare, Piscataway, NJ, according to manufacturer's instructions). B lymphocytes themselves can then be further isolated from other white blood cells using, for example, cell surface markers found on B lymphocytes. B lymphocyte cell surface markers include, without limitation, cell surface expressed immunoglobulin chains (e.g., lambda light chain, kappa light chain, and heavy chain such as $\operatorname{IgM}$ or $\operatorname{IgG})$. Additional B lymphocyte cell surface markers include, without limitation, CD21, CD27, CD138, CD20, CD19, CD22, CD72, and CD79A. Yet additional B lymphocyte cell surface markers include, without limitation, CD38, CD78, CD80, CD83, DPP4, FCER2, IL2RA, TNFRSF8, CD24, CD37, CD40, CD74, CD79B, CR2, IL1R2, ITGA2, ITGA3, MS4A1, ST6GAL1, CD1C, CD138, and CHST10.

These B lymphocyte surface markers can be used sequentially to enrich for B lymphocytes. For example, antibodies specific to a B lymphocyte cell surface markers (e.g., CD19) can be coupled to magnetic beads (e.g., Dynabeads commercially available from

Invitrogen Corp., Carlsbad, CA), and cells adhering to the beads (e.g., CD19 positive cells) isolated from non-CD19 expressing cells. B lymphocytes can be further enriched from the CD19 positive cells by, for example, flow cytometry sorting of cells expressing immunoglobulin chains at their cell surface. These enriched B lymphocytes can thus be isolated for use in the methods of various embodiments of the invention.

Antigen specific B lymphocytes can also be purified directly using the desired antigen as bait to isolate B cells expressing the antigen specific B cell receptor (membrane immunoglobulin). For example, B cells can be added to a column to which is adhered the desired antigen. The antigen-specific B cells will flow through the column more slowly than non-specific B cells or other cells (e.g., red blood cells, macrophages, etc.). The antigenspecific B cells can thus be enriched using this method.

Enriched or non-enriched B lymphocytes from an animal (e.g., enriched by various methods) can also be subjected to in vitro cell culture for 1 or 2 or 3 or 4 or more days prior to nucleic acid extraction. Such culture in vitro may expand the number of B lymphocytes and thus enrich them over non-B lymphocyte cells. In one non-limiting example, CD27 isolated human B lymphocytes can be subjected to various cytokine and extracellular molecule cocktails (such as but not limited to activated T cell conditioned medium, or any combination of B cell growth, and/or differentiation factors) prior to nucleic acid extraction in order to stimulate growth and/or differentiation of the $B$ lymphocytes prior to nucleic acid extraction from the B lymphocytes. Other biological molecules can also be added to the tissue culture media during the in vitro culturing to assist in growth, differentiation, and/or in vitro immunization, and/or any combination of the above.

From these isolated, enriched, or stimulated B lymphocytes, nucleic acid sequences (e.g., genomic DNA, hnRNA, mRNA, etc.) can be extracted using standard methods (e.g., phenol: chloroform extraction; see Ausubel et al., supra). This nucleic acid can then be subjected to sequencing analysis using a variety of methods for sequencing.

In some embodiments, the nucleic acid sequences can be directly sequenced from the biological material (i.e., without being amplified prior to sequencing). Services and reagents for directly sequencing from nucleic acid sequences are commercially available, for example, from Helicos BioSicences Corp. (Cambridge, MA). For example, Helicos' True

Single Molecule Sequencing allows direct sequencing of DNA, cDNA, and RNA. See also U.S. Patent Nos. 7,645,596; 7,037,687, 7,169,560; and publications Harris et al., Science 320: 106-109, 2008; Bowers et al., Nat. Methods 6: 493-494, 2009; and Thompson and Milos, Genome Biology 12: 217, 2011 (all of which patents and publications are incorporated herein by reference in their entireties).

In other embodiments, the nucleic sequences are amplified (e.g., by polymerase chain reaction (PCR)) prior to obtaining sequence information.

In one non-limiting example, an oligo dT PCR primer is used for RT-PCR. In another non-limiting example, gene-specific RT-PCR is performed using the PCR primers described herein, such as the 454 specific fusion mouse primers, the 454 rabbit immunoglobulin chain fusion primers or the variable heavy and variable light region primers. In another example, PCR primers against heavy chain and light chain populations in a mouse have sequences set forth in PCT publication no. WO2010/097435, herein incorporated by reference.

With or without B cell enrichment, purified genetic materials (DNA or mRNA) can be amplified (e.g., by PCR or RT-PCR) following standard procedures (see, e.g., Ausubel et al., supra) to prepare a library before NGS sequencing.

Isolated B lymphocytes mentioned above by various means can also be subjected to single cell encapsulation by using method in the art such as oil emulsion encapsulation or by commercial instrument such as RainDance technology (RainDance Technologies, Inc., Lexington, MA). These encapsulated B lymphocytes can then be fused with an appropriate single cell RT-PCR reagent (e.g., the reagent sold by Qiagen, as Cat \# 210210) with the appropriate amplification primers to generate linked Heavy and Light chain PCR products from each single B cells. Ligation or overlap PCR is known in the field and is practiced routinely for various molecular biology applications to stitch 2 DNA pieces into one (see, e.g., Meijer P.J. et al., J. Mol. Biol. 358(3):764-72, 2006 for overlap PCR). This approach allows for cognate pairing preservation and identification during sequencing.

## DNA Sequencing Methods

Methods for DNA sequencing that are well known and generally available in the art
may be used to obtain the nucleic acid sequences of the various embodiments of the invention. The methods may employ such enzymes as the Klenow fragment of DNA polymerase I, SEQUENASE® (US Biochemical Corp, Cleveland, Ohio), Taq polymerase (Invitrogen), thermostable T7 polymerase (Amersham, Chicago, Ill.), DNA ligase (e.g., from T4) or combinations of recombinant polymerases and proofreading exonucleases such as the ELONGASE Amplification System marketed by Gibco BRL (Gaithersburg, Md.). The process may be automated with machines such as the Hamilton Micro Lab 2200 (Hamilton, Reno, Nev.), Peltier Thermal Cycler (PTC200; MJ Research, Watertown, Mass.) and the ABI 377 DNA sequencers (Applied Biosystems).

Non-limiting methods to sequence nucleic acid molecules and thus generate nucleic acid sequences (e.g., to populate a genetic material database) of various embodiments of the invention include the Sanger method (see, e.g., Sanger et al, Nature 24: 687-695, 1977), the Maxam-Gilbert method (see, e.g., Maxam and Gilbert, Proc. Natl. Acad. Sci. USA 74: 560564, 1977), and pyrosequencing (see, e.g., Ronaghi et al., Science 281 (5375): 363, 1998 and Ronaghi et al., Analytical Biochemistry 242 (1): 84, 1996). Pyrosequencing, another nonlimiting sequencing method that can be used to obtain polynucleotide sequences, uses luciferase to generate light for detection of the individual nucleotides (either dATP, dTTP, dGTP, or dCTP, collectively "dNTPs") added to the nascent DNA, and the combined data are used to generate sequence read-outs.

In some embodiments, the nucleic acid sequences are obtained using deep sequencing or next generation sequencing. One rate-limiting step in conventional DNA sequencing arises from the need to separate randomly terminated DNA polymers by gel electrophoresis. Next generation sequencing devices bypass this limitation, e.g., by physically arraying DNA molecules on solid surfaces and determining the DNA sequence in situ, without the need for gel separation. These high throughput sequencing techniques allow numerous nucleic acid molecules to be sequenced in parallel.

Thus, thousands or millions of different nucleic acid molecules can be sequenced simultaneously (see Church, G.M., Sci. Am. 294 (1): 46-54, 2006; Hall, N., J. Exp. Biol. 210(Pt. 9): 1518-1525, 2007; Schuster et al., Nature Methods 5(1): 16-18, 2008; and MacLean et al., Nature Reviews Microbiology 7: 287-296, 2009). A variety of different
methods and machines for performing next generation sequencing exist, any of which can be used to generate nucleic acid sequences. See Lin et al., Recent Patents on Biomedical Engineering 1:60-67, 2008 for an overview of numerous next generation sequencing technologies. Publication No. 20070087362, describe the polony next generation sequencing method which uses a ligation-based sequencing method (see also U.S. Patent No. 5750341). The SOLiD technology commercially available from Applied Biosystems (a LifeTechnolgies Corp. company, Carlsbad, CA) employs sequencing by ligation. Using the SOLiD technology, a library of DNA fragments to be sequenced are amplified by emulsion PCR, and of the multiple fragments in the library, a single fragment species will be attached to a single magnetic bead (so called clonal beads). The fragments attached to the magnetic beads will have a universal P1 adapter sequence attached so that the starting sequence of every fragment is both known and identical. Primers are then selected that hybridize to the P1 adapter sequence within the library template. A set of four fluorescently labeled di-base probes compete for ligation to the sequencing primer. Specificity of the di-base probe is achieved by interrogating every 1 st and 2 nd base in each ligation reaction.

Another next generation sequencing method that of Margulies et al., Nature 437: 376-380, 2005 and US Patent Nos. 7,211,390; 7,244,559; and 7,264,929, which describe a parallelized version of pyrosequencing which amplifies DNA inside water droplets in an oil solution (emulsion PCR), with each droplet containing a single DNA template attached to a single primer-coated bead. Using the sequencing machine (the Genome Sequencer FLX System machine commercially available from 454 Life Sciences, a Roche company, Branford, CT), oligonucleotide adaptors are ligated to fragmented nucleic acid molecules and are then immobilized to the surface of microscopic beads before PCR amplification in an oil-droplet emulsion. Beads are then isolated in multiple picolitre-volume wells, each containing a single bead, sequencing enzymes, and dNTPs. Incorporation of a dNTP into the complementary strand releases pyrophosphate, which produces ATP, which in turn generates light that can then be recorded as an image for analysis.
U.S. Patent No. 7,115,400 describes another technique for solid-phase amplification
of nucleic acid molecules. This allows a large number of different nucleic acid sequences to be arrayed and amplified simultaneously. This technology is embodied in the Genome Analyzer system commercially available from Solexa (Illumina, Inc.). In this technology, DNA molecules are first attached to primers on a slide and amplified so that local clonal extended utilizing a partial nucleotide sequence and employing various methods known in the art to detect upstream sequences such as promoters and regulatory elements. For example, one method that may be employed, "restriction-site" PCR, uses universal primers to retrieve unknown sequence adjacent to a known locus (Sarkar, G., PCR Methods Applic. 2: 318-322 (1993)). In particular, genomic DNA is first amplified in the presence of primer to linker sequence and a primer specific to the known region. Exemplary primers are those described in Example 4 herein. The amplified sequences are then subjected to a second round of PCR with the same linker primer and another specific primer internal to the first one. Products of each round of PCR are transcribed with an appropriate RNA polymerase and sequenced using reverse transcriptase.

Inverse PCR may also be used to amplify or extend sequences using divergent primers based on a known region (Triglia et al., Nucleic Acids Res. 16: 8186 (1988)). The primers may be designed using OLIGO 4.06 Primer Analysis software (National Biosciences Inc., Plymouth, Minn.), or another appropriate program, to be 22-30 nucleotides in length, to have a GC content of $50 \%$ or more, and to anneal to the target sequence at temperatures about $68-72{ }^{\circ} \mathrm{C}$. The method uses several restriction enzymes to generate a suitable fragment in the known region of a gene. The fragment is then circularized by intramolecular ligation and used as a PCR template.

Another method which may be used is capture PCR which involves PCR amplification of DNA fragments adjacent to a known sequence in human and yeast artificial
chromosome DNA (Lagerstrom et al., PCR Methods Applic. 1: 111-119 (1991)). In this method, multiple restriction enzyme digestions and ligations may also be used to place an engineered double-stranded sequence into an unknown portion of the DNA molecule before performing PCR. Another method which may be used to retrieve unknown sequences is that described in Parker et al., Nucleic Acids Res. 19: 3055-3060 (1991)). Additionally, one may use PCR, nested primers, and PROMOTERFINDER® libraries to walk in genomic DNA (Clontech, Palo Alto, Calif.). This process avoids the need to screen libraries and is useful in finding intron/exon junctions.

It shall be understood that the nucleic acid from B lymphocytes may be further screened for those nucleic acid molecules encoding immunoglobulins prior to sequencing. To do this, primers specific for immunoglobulin-encoding nucleic acid molecules (or specific for regions adjacent thereto) may be employed.

As used herein, by "primer" is meant a nucleic acid sequence that may be at least about 15 nucleotides, or at least about 20 nucleotides, or at least about 30 nucleotides, or at least about 40 nucleotides in length. A primer specific for a particular nucleic acid molecule is meant to include a primer that hybridizes to a portion of the nucleic acid molecule under PCR annealing conditions (e.g., $60^{\circ} \mathrm{C}$ for thirty seconds). In some embodiments, a primer specific for a particular nucleic acid molecule is one that is complementary to that nucleic acid molecule.

Primers used for sequencing the nucleic acid sequence may be referred to as Sequencing Primers. Primers used for amplification of a target nucleic acid sequence by the polymerase chain reaction (PCR) may also be referred to as PCR primers or amplification primers (see description of PCR, for example, in Sambrook et al., supra and Ausubel et al., supra) the entire disclosure of which is hereby incorporated herein by reference.

In one non-limiting example for obtaining nucleic acid sequences in accordance with various embodiments of the invention, total nucleic acid from B lymphocytes may be rendered single-stranded (e.g., by heating the nucleic acid to $94-98^{\circ} \mathrm{C}$ for at least one minute. The single-stranded nucleic acid may then be passed over a solid support (e.g., a column or gel) to which are adhered single-stranded primers that are specific for non-variant regions of immunoglobulin-encoding nucleic acid molecules or non-coding regions adjacent thereto
(e.g., immunoglobulin gene promoters, enhancers, and/or introns). Some non-limiting examples for these non-variant regions of immunoglobulins include the constant region of the heavy chain, and the constant region of the light chains, and the FR1 region of either the heavy chain or the light chain. The nucleic acid is allowed to hybridize to the solid-phase support-bound primers, and the non-hybridizing nucleic acid removed. After removal, the hybridized nucleic acid (which is enriched for immunoglobulin-encoding nucleic acid molecules) is released from the primers by, for example, addition of heat or increasing the concentration of EDTA in the buffer.

In another embodiment of the invention, regardless of whether the nucleic acid from the B lymphocytes is enriched for immunoglobulin-encoding nucleic acid molecules, the immunoglobulin-encoding nucleic acid molecules may be amplified to increase their copy number. This amplification can be performed, for example, by PCR amplification using primers specific for non-variant regions of immunoglobulin-encoding nucleic acid molecules or non-coding regions adjacent thereto.

In all of the above methods for obtaining nucleic acid sequences in accordance with the various embodiments of the invention, it will be understood that the primers (e.g., sequencing or PCR primers) used to generate the immunoglobulin chain variable regionencoding nucleic acid sequences may be universal (e.g., polyA tail) or may be specific to immunoglobulin-encoding sequences.

In some embodiments, the starting material from which the immunoglobulin geneencoding nucleic acid sequence information is obtained is genomic DNA. For example, if the immunoglobulin chain variable regions are from humans, primers (e.g., sequencing primers and/or PCR primers) may be selected to be identical to or hybridize to an immunoglobulin chain gene promoter. For example, the human genome sequence is known. Since the heavy chain-encoding gene occurs on chromosome 14 and the light chainencoding gene occurs on chromosome 22 (lambda light chain) and 2 (kappa light chain), it would be routine for the ordinarily skilled biologist to design primers that hybridize to regulatory elements of the heavy chain-encoding gene and the light chain-encoding gene. Such regulatory elements include, without limitation, promoters, enhancers, and introns.

Immunoglobulin variable region-specific primers can likewise be readily determined
for mice immunoglobulins since the murine kappa light chain gene is known to be located on chromosome 6 and the murine heavy chain gene is known to be located on chromosome 12.

In another non-limiting embodiment, the starting material from which the cDNA reversed translated from the mRNA. In this example, to obtain immunoglobulin variable region-encoding nucleic acid sequences, primers can be selected to be identical to or hybridize to the polyA tail of an mRNA or the complementary TTTT (SEQ ID NO: 306)rich sequence of the mRNA's corresponding cDNA. Alternatively, or in addition, primers can also be selected to be identical to or hybridize to the FR1-encoding nucleic acid sequences. Alternatively, or in addition, primers can also be selected to be identical to or hybridize to a portion of (or all of) one of the CH regions (i.e., $\mathrm{CH} 1, \mathrm{CH} 2$, or CH 3 ) and/or the VH region-encoding nucleic acid sequences.

Sequencing errors can arise from using universal degenerate primers to sequence nucleic acid molecules encoding immunoglobulins from hybridomas. For example, Essono et al, Protein Engineering, Design and Selection, pp. 1-8, 2009 describe a method combining sequencing with peptide mass spectrometry fingerprinting of the corresponding Ig chain to determine the correct sequence of a monoclonal antibody produced by a hybridoma clone. However, in the non-limiting methods of various embodiments of the invention, the presence of sequencing errors will merely increase the number of different nucleic acid sequences. Unlike Essono et al., supra, since the methods of various embodiments of the invention allow the creation of a single antibody (both heavy and light chains or variable regions thereof) from a starting polyclonal population of antibodies (where the created antibody may not actually occur within the starting polyclonal population of antibodies), having a large number of sequences in the genetic material database with which to correlate the observed mass spectra data of the peptide database is an asset.

## Predicted Mass Spectra Information From the Genetic Material Database

In accordance with various embodiments of the invention, once nucleotide sequences of the nucleic acid molecules are generated, additional information may be generated based
on the nucleotide sequence information alone. For example, the nucleotide sequence information can be translated into predicted amino acid sequences using the genetic code. Although the ordinarily skilled artisan can readily translate nucleotide sequences into amino acid sequences using the genetic code, several automated translation tools (which are publicly available) can be used, such as the ExPASy translate tool from the Swiss Institute of Bioinformatics or the EMBOSS Transeq translation tool from EMBL-EBI.

Similarly, predicted mass spectra information of the predicted amino acid sequences encoded by the nucleic acid sequences can be readily determined by the ordinarily skilled artisan. For example, following virtual (i.e., in silico) digestion of the predicted polypeptides encoded by the nucleic acid sequences, predicted mass spectra of the peptide fragments can be generated by using standard publicly available software algorithm tools including, without limitation, the Sequest software (from Thermo Fisher Scientific, Inc., West Palm Beach, FL), the Sequest 3G software (from Sage-N Research, Inc., Milpitas, CA), the Mascot software (from Matrix Science, Inc., Boston, MA; see also Electrophoresis, 20(18) 3551-67 (1999)), and the X!Tandem software (opensource from The Global Proteome Machine Organization, whose use is described in Baerenfaller K. et al., Science 320:938-41, 2008).

As used herein, the words "predicted," "theoretical," and "virtual" are used interchangeably to refer to nucleotide sequences, amino acid sequences or mass spectra that are derived from in silico (i.e., on a computer) transcription and/or translation (for the predicted nucleotide and amino acid sequences) or in silico digestion and/or mass spectrometry analysis (for the predicted mass spectra) of information from the nucleic acid sequences. For example, nucleic acid sequences are derived from genomic nucleic acid molecules obtained from B lymphocytes as described herein. The nucleotide sequence of, for example, mRNA derived from genomic DNA is predicted following in silico translation of the genomic DNA. This predicted mRNA (or cDNA) may then be translated in silico to produce predicted amino acid sequences. The predicted amino acid sequences may then be digested in silico with proteases (e.g., trypsin) and/or chemical protein cleavage reagents (e.g., cyanogen bromide) to produce predicted (or theoretical or virtual) peptide fragments. The virtual peptide fragments can be then analyzed in silico to produce predicted mass
spectra information. Thus, predicted mass spectra information, predicted peptide fragments, predicted amino acid sequences, and predicted mRNA or cDNA sequences can all be derived from the nucleic acid sequences collected from B lymphocytes (e.g., from an animal). predicted polypeptides to generate predicted peptides fragments and ultimately predicted mass spectra is the same protease(s) and/or reagent(s) used to digest the starting population of polyclonal antibodies, as described above.

## Correlating Observed Mass Spectra with Predicted Mass Spectra

As described above, passage of the fragments derived from the starting polyclonal population of antibodies through a mass spectrometer generates numerous observed mass spectra. Given the potentially large number of different immunoglobulins within a polyclonal population, each with a different amino acid sequence, that are analyzed with the mass spectrometer, the resulting observed mass spectra will be difficult to assemble back into a functional immunoglobulin chain variable region. In the methods of various embodiments of the invention, because the encoding nucleic acid sequences are available, there is no need to assemble the observed mass spectra data. Instead, the observed mass spectra are correlated with the predicted mass spectra derived from the nucleic acid sequences of the genetic material database to obtain the amino acid (and underlying nucleotide) sequences of full-length immunoglobulin chains (or variable regions thereof) of an immunoglobulin that specifically binds to an antigen from a starting polyclonal immunoglobulin population.

Also as described above, the genetic material database can be derived from nucleic acid molecules isolated from the B-cell repertoire of an immunized animal, including nucleic acid molecules encoding full length immunoglobulin heavy and light chains and variable regions thereof. Attempts to identify nucleic acids encoding antigen-specific immunoglobulins based solely on the information from the genetic material database (e.g., frequency rankings of variable region sequences) may miss those immunoglobulins that occur at low frequencies yet manifest superior antigen-specific activities. In accordance
with the various embodiments of this invention, however, by correlating the predicted mass spectra information from the genetic material database with the observed mass spectra information from the actual circulating polyclonal antibodies as disclosed herein, those immunoglobulin chains (or variable regions thereof) in the genetic material database can be selected that correspond to immunoglobulins within the circulating polyclonal antibodies.

By "correlating" it is meant that the observed mass spectra information derived from the starting polyclonal antibodies and the predicted mass spectra information derived from the genetic material database are cross-referenced and compared against each other, such that immunoglobulin heavy and/or light chains (or variable regions thereof) can be identified or selected from the genetic material database that correspond to immunoglobulin heavy and/or light chains (or variable regions thereof) of antigen-specific immunoglobulins in the starting polyclonal population.

In specific embodiments, the correlating process involves comparing the observed mass spectra information with the predicted spectra information to identify matches. For example, each of the observed spectra can be searched against the collection of predicted mass spectra derived from the genetic material database, with each predicted spectrum being identifiably associated with a peptide sequence from the genetic material database. Once a match is found, i.e., an observed mass spectrum is matched to a predicted mass spectrum, because each predicted mass spectrum is identifiably associated with a peptide sequence in the genetic material database, the observed mass spectrum is said to have found its matching peptide sequence - such match also referred to herein as "peptide spectrum match" or "PSM". Because of the large number of spectra to be searched and matched, this search and matching process can be performed by computer-executed functions and softwares, such as the SEQUEST algorithm (Sage-N Research, Inc., Milpitas, CA).

In some embodiments, the search and matching is directed to functional domains or fragments of immunoglobulins, such as variable region sequences, constant region sequences, and/or one or more CDR sequences. For example, the observed spectra are only searched against predicted mass spectra derived from V regions (and/or CDR3 sequences) of immunoglobulins to identify V-region (and/or CDR3) PSMs. In other embodiments, the search and matching is directed to full-immunoglobulin heavy or light chain sequences.

After the search and matching has been completed, immunoglobulin heavy or light chains in the genetic material database are analyzed and selected based on one or more of the following parameters: the number of unique peptides, the spectrum share, the amino acid sequence coverage, the count of peptides (either total peptide count or unique peptide count), frequency of the encoding nucleic acid sequences, and clonal relatedness.

The term "coverage" in referring to a sequence or region (e.g., a heavy or light chain sequence, a V-region sequence, or a CDR sequence) is defined as the total number of amino acids within the sequence that have been identified in peptides which map to the sequence or region and which have a matching observed spectrum, divided by the number of amino acids in the sequence or region. The higher the coverage, the more likely the sequence or region appears in the actual polyclonal population.

By "number of unique peptides" it is meant the number of distinct peptides observed mapping to a single protein sequence (e.g., a single immunoglobulin heavy or light chain or a variable region thereof). The higher the number, the more likely the immunoglobulin chain is present in the polyclonal population. In specific embodiments, selection of an immunoglobulin chain is made based on a number of unique peptides of at least 5, 6, 7, 8, 9, $10,11,12$ or more in the immunoglobulin chain or its variable region.
"Spectrum share" is determined by dividing the total number of peptides mapped to the sequence by the total number of confident PSMs mapped to the entire genetic database. Spectrum share provides a human readable count of peptides expressed as the percentage of PSMs that map to a specific V-region sequence.

The term "peptide count" in referring to a protein sequence (e.g., a CDR3 region or a variable region) means the number of times a peptide is identified from the observed mass spectra that matches the protein sequence. For example, the count of a CDR3 region means the number of times a peptide is identified from the observed mass spectra that matches the CDR3 region. The count of a variable region means the number of times a peptide is identified from the observed mass spectra that matches the variable region. "Total peptide count" in referring to a protein sequence means the number of times any peptide (unique or non-unique) is identified from the observed mass spectra that matches the protein sequence. "Unique peptide count" means the number of times a unique peptide is identified from the
observed mass spectra that matches the protein sequence. If the same peptide has been identified multiple times from the observed mass spectra, the total number of times this peptide is observed will be considered in determining the total peptide count, yet this peptide will be counted only once for determining the unique peptide count. on sequence coverage. In other embodiments, the selection is made based on a combination of of sequence coverage with one or more other parameters, including the number of unique peptides, spectrum share, total peptide count, unique peptide count, frequency of the encoding nucleic acid sequence, or clonal relatedness.

The above parameters can be independently determined with respect to a full-length heavy or light chain, or with respect to one or more portions of an immunoglobulin heavy or light chain, e.g., the variable region, and a CDR (e.g., CDR1, CDR2, or CDR3, especially CDR3). In certain embodiments, selection of immunoglobulin chains (or variable regions thereof) is made based on the V -region coverage and/or CDR coverage (e.g., CDR3 coverage).

The selection of immunoglobulin heavy or light chains (or variable regions thereof) can be made based on the absolute value of one or more parameters, or based on the ranking of absolute values for a relevant parameter. Where ranking for a particular parameter is considered, the top ranked $10,20,30,40,50,60,70,80,90,100$ or more sequences can be selected irrespective of the absolute values of that parameter. Where the value of a parameter is considered, e.g., the percentage of sequence coverage, in some embodiments, selection of immunoglobulin chains is made based on a CDR coverage (such as CDR3 coverage) of at least $10 \%, 15 \%, 20 \%, 25 \%, 30 \%, 35 \%, 40 \%, 45 \%, 50 \%, 55 \%, 60 \%, 65 \%$, $70 \%, 75 \%, 80 \%, 85 \%, 90 \%, 95 \%, 98 \%$ or higher; additionally or alternatively, based on a V-region coverage of at least $5 \%, 10 \%, 15 \%, 20 \%, 25 \%, 30 \%, 35 \%, 40 \%, 45 \%, 50 \%, 55 \%$, $60 \%, 65 \%, 70 \%, 75 \%, 80 \%, 85 \%, 90 \%$ or higher.

In some embodiments, a phylogenetic analysis is performed to determine clonal relatedness of the heavy chain variable region, light chain variable region, or one or more CDR's (e.g., CDRH3 or CDRL3). Changes or mutations of nucleic sequence of heavy and light chains compared to germline sequence can provide evidence of affinity maturation of
antibodies following antigen exposure. Clonal relatedness can be used as a factor in selection of antibody sequences. A phylogenetic analysis can be performed by methods known in the art, e.g., those described in Dereeper et al., 2008, Nucl. Acids Res., 36(Web Server issue):W456-459; Dereeper et al., 2010, BMC Evol. Biol., 10:8, and available online at www.phylogeny.friversion 2 cg//index.cgi. In some embodiments, the entire heavy or light chain variable regions are grouped by homology, then further grouped by CDR (e.g., CDR3) homology.

The selected heavy and light chain sequences can then be expressed in pairs to assemble into monoclonal antibodies which are analyzed to confirm antigen-specific functionality. The pairing of selected heavy and light chain sequences can be entirely random, or can take into consideration of one or more parameters described above, including sequence coverage, unique number of peptides, spectrum share, total peptide count, and unique peptide count.

In some embodiments, the abundance of a population of antibodies having a particular peptide sequence can be determined using a heavy isotope labeled (e.g., AQUA) peptide. See, e.g., WO 03/016861 and Gerber et al., 2003, 100:6940-45. These methods employ the introduction of a known quantity of at least one heavy-isotope labeled peptide standard (which has a unique signature detectable by LC-SRM chromatography) into a digested biological sample in order to determine, by comparison to the peptide standard, the absolute quantity of a peptide with the same sequence and protein modification in the biological sample. The peptide can be unique to one species of antibody or found in multiple (e.g., clonally-related) antibodies. In some embodiments, the peptide can include at least a portion of a CDR (e.g., CDR3). Quantitation of the abundance of antibody populations can be useful in methods of monitoring serum antibody composition, e.g., following vaccination of a subject.

It should be noted that the immunoglobulin that specifically binds to the antigen whose amino acid sequence (or nucleic acid sequence) created using the non-limiting methods of various embodiments of the invention need not actually be present within the starting polyclonal population of immunoglobulins. Rather, the non-limiting methods of various embodiments of the invention simply allow the rapid creation of an immunoglobulin
that specifically binds the antigen whether or not that immunoglobulin actually existed in the starting polyclonal population. For example, the created immunoglobulin that has the highest desired qualities (e.g., highest binding affinity (or lowest KD) for the antigen or a desired isotype (e.g., IgG2a)) may be the result of a light chain from a first antibody in the polyclonal population assembled with a heavy chain of a second antibody (i.e., different from the first antibody) in the polyclonal population. The resulting created immunoglobulin can be further characterized (e.g., binding affinity for the antigen or isotype) according to standard methods.

## Method of Making Recombinant Antibodies

Once the nucleotide sequence of an immunoglobulin chain (or variable region thereof) of an antibody that specifically binds to the antigen is elucidated, a nucleic acid molecule comprising that sequence can be generated.

For example, if the starting population from which the immunoglobulin chain (or variable region thereof)-encoding nucleic acid molecules is obtained is a cDNA library, the nucleic acid molecule comprising the elucidated sequence can be readily obtained from the library (e.g., by screening the library with a primer identical to or capable of hybridizing to a portion of the elucidated sequence) or by PCR amplifying the nucleic acid molecule from the library using primers designed to amplify the elucidated nucleic acid sequence.

Alternatively (or in addition), nucleic acid molecules comprising the elucidated nucleotide sequence can be generated by simply artificially generating the nucleic acid molecule using a standard DNA synthesis machine. Numerous DNA synthesis machines are commercially available including, without limitation, the MerMade series of synthesizers (e.g., MerMade 4, Mermade 6, MerMade 384, etc.) available from BioAutomation, Plano, TX; the various DNA/RNA synthesizers commercially available from Applied Biosystems (now part of Life Technologies, Corp., Carlsbad, CA). Several companies also offer DNA synthesis services (e.g., BioPioneer, Bio S\&R, Biomatik, Epoch BioLabs, etc.)

Methods to express nucleic acid encoding heavy and light chains of an immunoglobulin to produce recombinant immunoglobulins are known (see, e.g., U.S. Patent Publication Nos. 6,331,415; 5,969,108; 7,485,291; US 2011-0045534; and PCT Publ. No.

WO 2011/022077). Recombinant immunoglobulins can be made in a variety of cells including, without limitation, insect cells (e.g., SF9 cells), hamster cells (e.g., CHO cells), murine cells (e.g., NIH-3T3 cells), primate cells (e.g., COS cells), human cells (e.g., Hela cells), and prokaryotic cells (e.g., E. coli cells). In some embodiments, the cells expressing the recombinant immunoglobulins of various embodiments of the invention are able to add secondary modifications (e.g., glycosylation) to the recombinant immunoglobulin in a manner similar to that of the species from which the immunoglobulin was originally derived. For example, where the population of polyclonal antibodies whose fragments were used to generate the observed mass spectra data are collected from a human, human cells (or cells which glycosylate proteins similarly or identically to human cells) may be used.

To obtain expression of the nucleic acid sequences of a recombinant immunoglobulin (or antigen binding fragment thereof) that specifically binds to the antigen in a cell, the nucleic acid sequences may be ligated into a vector (e.g., a plasmid or a retroviral vector) containing appropriate regulatory sequences such that the inserted nucleic acid sequences are expressed in the cell into which the nucleic acid sequence are introduced. Such regulatory sequences include, for example, promoters, enhancers, intron acceptor elements, poly adenylation sites, etc. Any method can be employed to introduce the nucleic acid sequences of a recombinant immunoglobulin (or vector containing the same) into a cell including, without limitation, electroporation, transfection by chemical means (e.g., CaPO4, DEAE-dextran, polyethylenimine), infection, transduction, liposome fusion, etc. (see methods, e.g., in Ausubel et al., supra).

In accordance with some embodiments of the invention, the heavy immunoglobulin chain and the light immunoglobulin chain are randomly selected to be assembled into an immunoglobulin (or variable region or antibody binding domain thereof). For example, correlation of the actual mass spectra from a peptide fragment of the polyclonal antibodies with the predicted mass spectra of a predicted peptide encoded by the nucleic acid sequences will be used to obtain the nucleotide sequence or predicted amino acid sequence of an immunoglobulin chain comprising the peptide fragment. The obtained nucleotide sequence of the immunoglobulin chain can then be randomly co-expressed and/or with a second similarly obtained nucleotide sequence of an immunoglobulin chain, where the second
nucleotide sequence encodes the other chain of an intact antibody under conditions where the two encoded immunoglobulin chains will assemble into an intact antibody.

Conditions for co-expressing two nucleotide sequences (e.g., in cells) each encoding an immunoglobulin chain such that an intact immunoglobulin is assembled are known (see, e.g., U.S. Pat. Nos. $5,969,108 ; 6,331,415 ; 7,498,024 ; 7,485,291$; and US Pat. Pub. No. 20110045534, all herein incorporated by reference in their entireties). Because of the number of different nucleotide sequences that can be obtained using the methods described herein, the invention contemplates the use of robotics and high-throughput methods to screen the encoded immunoglobulins to create an immunoglobulin that specifically binds to the antigen.

As used herein, by "assembled" or "assembling" is meant that a light chain of an antibody (or a fragment thereof) and a heavy chain of an antibody (or a fragment thereof) are combined together in a manner in which the two chains join to create an antibody (or a fragment thereof). In some embodiments, in the assembled antibody (or fragment thereof), amino acid residues from both the heavy chain and light chain contribute to the antigen binding domain of the assembled antibody (or fragment thereof). In some embodiments, the assembled antibody (or fragment thereof) comprises a light chain (or fragment thereof) covalently bonded to a heavy chain (or fragment thereof). In some embodiments, the assembled antibody (or fragment thereof) comprises a light chain (or fragment thereof) noncovalently bonded to a heavy chain (or fragment thereof).

In some embodiments, the nucleotide sequences or amino acid sequences of the immunoglobulin chains (or variable regions thererof) identified in the proteomics analysis described above are synthesized by recombinant molecular biology techniques or gene synthesis techniques prior to assembly of recombinant antibodies. For example, the nucleotide or amino acid sequences may be synthesized on a nucleotide or peptide synthesis machine prior to assembly. Or, the nucleotide or amino acid sequences may be expressed recombinantly by cloning the nucleotide sequences into an expression vector (e.g., pCDNA3.1 from Invitrogen, Carlsbad, CA), and expressing the encoded polypeptide in a cell (e.g., HeLa cells, CHO cells, COS cells, etc.) transfected with the expression vector. In some embodiments, the assembly step occurs in the transfected cell (e.g., a single cell is
transfected with one or more expression vectors comprising nucleic acid sequences encoding one heavy and one light chain, where the heavy and light chain will be expressed as polypeptides in the transfected cell).

In various embodiments of the invention, the recombinant antibodies are isolated. As used herein, by "isolated" (or "purified") is meant an antibody is substantially free of other biological material with which it is naturally associated, or free from other biological materials derived, e.g., from a cell that has been genetically engineered to express the antibody of the invention. For example, an isolated recombinant antibody is one that is physically separated from other components of the host cell (e.g., the endoplasmic reticulum or cytoplasmic proteins and RNA). Likewise, a purified antibody from blood sera and/or plasma is an antibody that is isolated from other serum or plasma components (e.g., albumin or cells) (using, for example, adherence of the antibodies to protein A, where the nonantibody sera components will not adhere to protein A). Thus, an isolated antibody (or isolated immunoglobulin) of the present invention includes an antibody that is at least 70$100 \%$ pure, i.e., an antibody which is present in a composition wherein the antibody constitutes $70-100 \%$ by weight of the total composition. In some embodiments, the isolated antibody of the present invention is $75 \%-99 \%$ by weight pure, $80 \%-99 \%$ by weight pure, $90-99 \%$ by weight pure, or $95 \%$ to $99 \%$ by weight pure. The relative degree of purity of an antibody various non-limiting embodiments of the invention is easily determined by wellknown methods.

In some embodiments, the recombinant antibodies (or variable regions thereof) are further screened or analyzed in an immunoassay to confirm that the antibodies specifically bind to the antigen. In some embodiments, the immunoassay is a standard immunoassay such as a flow cytometry assay (e.g., a FACS scan), an enzyme-linked immunosorbent assay (ELISA), a Western blotting assay, an immunohistochemistry assay, an immunofluorescence assay, a radioimmunoassay, a neutralization assay, a binding assay, an affinity assay, or a protein or peptide immunoprecipitation assay. All of these immunoassays are well known standard assays and have been well described in standard methods books (see, e.g., Ausubel et al., supra; Coligan et al., supra; Harlow and Lane, supra).

## Therapeutic Antibodies

The various non-limiting embodiments and methods of the invention are useful, for example, in isolating antibodies that have therapeutic value. For example, in the course of a normal immune response in an animal to a pathogen, antibodies with the highest specificity to an antigen of the pathogen may take weeks to arise. This is because the $B$ lymphocyte producing the antibody must first be stimulated by the appropriate T lymphocyte that also recognize the antigen presented on an antigen presenting cell in context of the major histocompatibility complex expressed by every nucleated cell of an animal. B lymphocytes initially responding to the antigen produce antibodies that specifically bind to the antigen. However, the highest affinity antibodies are actually those that are produced by B lymphocytes that have bound their antigen (through cell surface expressed immunoglobulin complexed with other cell surface antigens to form the B cell receptor) and, upon stimulation through the B cell receptor and other cells (including T lymphocytes), undergo affinity maturation to produce antibodies with high affinity for their specific antigen. Such a $B$ lymphocyte that has undergone affinity maturation (or its progeny with the same antibody specificity) is available in the animal to quickly produce high affinity antibody should the animal encounter the same pathogen again.

This tight regulation of T lymphocytes and B lymphocytes responding to an antigen the first time that antigen is seen (e.g., the first time the animal is infected with a particular pathogen) is necessary to prevent autoimmune or inappropriate immune response. However, one drawback is that by the time an antigen-specific B lymphocyte is secreting antibody of the highest affinity and specificity for the antigen, a quickly growing pathogen may have grown within the animal to the extent that it can no longer be easily cleared. In some embodiments of the invention, the methods allow for the rapid development of an antigenspecific antibody that skips the time-consuming process of first isolating an antigen-specific B lymphocyte that is secreting the antibody and immortalizing that lymphocyte.

Thus, in another aspect, the invention provides a therapeutic composition comprising a recombinant antibody with a pharmaceutically acceptable carrier.

As used herein, "pharmaceutically acceptable carrier" includes any material which,
when combined with an active ingredient (e.g., a recombinant antibody made in accordance with various embodiments of the invention), allows the ingredient to retain biological activity and is non-reactive with the subject's immune system and non-toxic to the subject when delivered. Examples include, but are not limited to, any of the standard pharmaceutical carriers such as a phosphate buffered saline solution, water, emulsions such as oil/water emulsion, and various types of wetting agents. Non-limiting examples of diluents for aerosol or parenteral administration are phosphate buffered saline, normal ( $0.9 \%$ ) saline, Ringer's solution and dextrose solution. The pH of the solution may be from about 5 to about 8 , or from about 7 to about 7.5. Further carriers include sustained release preparations such as semipermeable matrices of solid hydrophobic polymers containing the antibody, which matrices are in the form of shaped articles, e.g., films, liposomes or microparticles. It will be apparent to those persons skilled in the art that certain carriers may be more preferable depending upon, for instance, the route of administration and concentration of antibody being administered. Compositions comprising such carriers are formulated by well known conventional methods (see, for example, Remington's Pharmaceutical Sciences, 18th edition, A. Gennaro, ed., Mack Publishing Co., Easton, Pa., 1990; and Remington, The Science and Practice of Pharmacy, 20th Ed. Mack Publishing, 2000).

While any suitable carrier known to those of ordinary skill in the art may be employed in the pharmaceutical compositions of this invention, the type of carrier will vary depending on the mode of administration. In various embodiments of the invention, numerous delivery techniques for the non-limiting pharmaceutical compositions described herein (e.g., containing a binding agent or a binding agent-encoding polynucleotide) are well known in the art, such as those described by Rolland, 1998, Crit. Rev. Therap. Drug Carrier Systems 15:143-198, and references cited therein.

## Methods of Treatment

In another aspect, the invention provides a method a treating an animal having or suspected of having disease a characterized by a disease antigen, wherein the method comprising administering an effective amount of a therapeutic composition comprising an immunoglobulin that specifically binds to an antigen made in accordance with the methods
of various embodiments of the invention, wherein the antigen specifically bound by the immunoglobulin of the therapeutic composition and the disease antigen are the same.

In some embodiments, the animal is a human or a domesticated animal (e.g., a dog, cat, cow, goat, sheep, chicken, turkey, llama, emu, elephant, or ostrich).

As used herein, the phrase "characterized by" with respect to a disease and indicated disease antigen (e.g., an HIVgp120 antigen from AIDS) is meant a disease in which the indicated disease antigen is present in an animal with that disease. In some embodiments, the disease antigen is encoded by nucleic acid from the disease's etiological agent (e.g., a virus). In some embodiments, the disease antigen is encoded by the animal's genome (e.g., the BCR-ABL fusion disease antigen encoded by the Philadelphia chromosome in patients with chronic myelogenous leukemia (CML).

By "treating" is meant halting, retarding, or inhibiting progression of a disease or preventing development of disease in an animal. Methods of detecting whether the treatment is successful are known. For example, where the disease is a solid tumor, progression of the disease is inhibited, halted, or retarded if there is a regression of the tumor, reduction in metastases, reduction in tumor size and/or reduction in tumor cell count following administration of the effective amount of a therapeutic composition comprising a recombinant immunoglobulin produced using the methods of various embodiments of the invention.

As used herein, by an "effective amount" is an amount or dosage sufficient to effect beneficial or desired results including halting, slowing, halting, retarding, or inhibiting progression of a disease in an animal or preventing development of disease in an animal. An effective amount will vary depending upon, e.g., an age and a body weight of a subject to which the therapeutic composition comprising the recombinant immunoglobulin is to be administered, a severity of symptoms and a route of administration, and thus administration is determined on an individual basis. In general, the daily adult dosage for oral administration is about 0.1 to 1000 mg , given as a single dose or in divided doses. For continuous intravenous administration, the compositions can be administered in the range of $0.01 \mathrm{ug} / \mathrm{kg} / \mathrm{min}$ to $1.0 \mathrm{ug} / \mathrm{kg} / \mathrm{min}$, desirably $0.025 \mathrm{ug} / \mathrm{kg} / \mathrm{min}$ to $0.1 \mathrm{ug} / \mathrm{kg} / \mathrm{min}$.

An effective amount can be administered in one or more administrations. By way of
example, an effective amount of a recombinant immunoglobulin produced using the methods of various embodiments of the invention, is an amount sufficient to ameliorate, stop, stabilize, reverse, slow and/or delay progression of a disease (e.g., a cancer) in an animal or is an amount sufficient to ameliorate, stop, stabilize, reverse, slow and/or delay growth of a diseased cell (e.g., a biospsied cancer cell) in vitro. As is understood in the art, an effective amount of a recombinant antibody of various embodiments of the invention may vary, depending on, inter alia, the animal's medical history as well as other factors such as the isotype (and/or dosage) of the recombinant antibody.

Effective amounts and schedules for administering the compositions comprising a non-limiting recombinant antibody of various embodiments of the invention may be determined empirically, and making such determinations is within the skill in the art. Those skilled in the art will understand that the dosage that must be administered will vary depending on, for example, the animal that will receive the compositions of various embodiments of the invention, the route of administration, the particular type of compositions used (e.g., the isotype of the recombinant antibody within the composition) and other drugs being administered to the animal. Where the animal (e.g., a human patient) is administered a composition comprising an antibody, guidance in selecting appropriate doses for antibody is found in the literature on therapeutic uses of antibodies, e.g., Handbook of Monoclonal Antibodies, Ferrone et al., eds., Noges Publications, Park Ridge, N.J., 1985, ch. 22 and pp. 303-357; Smith et al., Antibodies in Human Diagnosis and Therapy, Haber et al., eds., Raven Press, New York, 1977, pp. 365-389.

A typical daily dosage of an effective amount of an antibody used alone might range from about $1 \mathrm{ug} / \mathrm{kg}$ to up to $100 \mathrm{mg} / \mathrm{kg}$ of body weight or more per day, depending on the factors mentioned above. Generally, any of the following doses may be used: a dose of at least about $50 \mathrm{mg} / \mathrm{kg}$ body weight; at least about $10 \mathrm{mg} / \mathrm{kg}$ body weight; at least about 3 $\mathrm{mg} / \mathrm{kg}$ body weight; at least about $1 \mathrm{mg} / \mathrm{kg}$ body weight; at least about $750 \mathrm{ug} / \mathrm{kg}$ body weight; at least about $500 \mathrm{ug} / \mathrm{kg}$ body weight; at least about $250 \mathrm{ug} / \mathrm{kg}$ body weight; at least about $100 \mathrm{ug} / \mathrm{kg}$ body weight; at least about $50 \mathrm{ug} / \mathrm{kg}$ body weight; at least about $10 \mathrm{ug} / \mathrm{kg}$ body weight; at least about $1 \mathrm{ug} / \mathrm{kg}$ body weight, or more, is administered. In some embodiments, a dose of a binding agent (e.g., antibody) provided herein is between about
$0.01 \mathrm{mg} / \mathrm{kg}$ and about $50 \mathrm{mg} / \mathrm{kg}$, between about $0.05 \mathrm{mg} / \mathrm{kg}$ and about $40 \mathrm{mg} / \mathrm{kg}$, between about 0.1 mg and about $30 \mathrm{mg} / \mathrm{kg}$, between about 0.1 mg and about $20 \mathrm{mg} / \mathrm{kg}$, between about 0.5 mg and about 15 mg , or between about 1 mg and 10 mg . In some embodiments, the dose is between about 1 mg and 5 mg . In some alternative embodiments, the dose is between about 5 mg and 10 mg .

The methods described herein (including therapeutic methods) can be accomplished by a single direct injection at a single time point or multiple time points to a single or multiple sites. Administration can also be nearly simultaneous to multiple sites. Frequency of administration may be determined and adjusted over the course of therapy, and is base on accomplishing desired results. In some cases, sustained continuous release formulations of the recombinant immunoglobulins of various embodiments of the invention may be appropriate. Various formulations and devices for achieving sustained release are known in the art.

Compositions comprising the recombinant antibodies of present invention may be formulated for any appropriate manner of administration, including for example, systemic, topical, oral, nasal, intravenous, intracranial, intraperitoneal, subcutaneous or intramuscular administration, or by other methods, such as infusion, which ensure its delivery to the bloodstream in an effective form. The composition may also be administered by isolated perfusion techniques, such as isolated tissue perfusion, to exert local therapeutic effects. For parenteral administration, such as subcutaneous injection, the carrier preferably comprises water, saline, alcohol, a fat, a wax or a buffer. For oral administration, any of the above carriers or a solid carrier, such as mannitol, lactose, starch, magnesium stearate, sodium saccharine, talcum, cellulose, glucose, sucrose, and magnesium carbonate, may be employed. In some embodiments, for oral administration, the formulation of the compositions is resistant to decomposition in the digestive tract, for example, as microcapsules encapsulating the recombinant immunoglobulin of various embodiments of the invention within liposomes. Biodegradable microspheres (e.g., polylactate polyglycolate) may also be employed as carriers for the therapeutic compositions of this invention. Suitable biodegradable microspheres are disclosed, for example, in U.S. Pat. Nos. $4,897,268$ and $5,075,109$.

In some embodiments of the invention, compositions may also comprise buffers (e.g., neutral buffered saline or phosphate buffered saline), carbohydrates (e.g., glucose, mannose, sucrose or dextran), mannitol, proteins, polypeptides or amino acids such as glycine, antioxidants, chelating agents such as EDTA or glutathione, adjuvants (e.g., aluminum hydroxide) and/or preservatives. Alternatively, non-limiting compositions of various embodiments of the present invention may be formulated as a lyophilizate.

In some embodiments of the invention, the recombinant immunoglobulins also may be entrapped in microcapsules prepared, for example, by coacervation techniques or by interfacial polymerization (for example, hydroxymethylcellulose or gelatin-microcapsules and poly(methylmethacylate) microcapsules, respectively), in colloidal drug delivery systems (for example, liposomes, albumin microspheres, microemulsions, nano-particles and nanocapsules), or in macroemulsions. Such techniques are disclosed in Remington's Pharmaceutical Sciences, 18th edition, A. Gennaro, ed., Mack Publishing Co., Easton, Pa., 1990; and Remington, The Science and Practice of Pharmacy 20th Ed. Mack Publishing, 2000. To increase the serum half life of the recombinant immunoglobulin of various embodiments of the invention, one may incorporate a salvage receptor binding epitope into the antibody (especially an antibody fragment) as described in U.S. Pat. No. 5,739,277, for example. As used herein, the term "salvage receptor binding epitope" refers to an epitope of the Fc region of an $\operatorname{IgG}$ molecule (e.g., $\operatorname{IgG} 1, \operatorname{IgG} 2, \operatorname{IgG} 3$, and $\operatorname{IgG} 4$ ) that is responsible for increasing the in vivo serum half-life of the IgG molecule.

In some embodiments of the invention, the recombinant immunoglobulins may also be formulated as liposomes. Liposomes containing the recombinant immunoglobulins are prepared by methods known in the art, such as described in Epstein et al., 1985, Proc. Natl. Acad. Sci. USA 82:3688; Hwang et al., 1980, Proc. Natl Acad. Sci. USA 77:4030; and U.S. Pat. Nos. $4,485,045$ and 4,544,545. Liposomes with enhanced circulation time are disclosed in U.S. Pat. No. 5,013,556. Particularly useful liposomes can be generated by the reverse phase evaporation method with a lipid composition comprising phosphatidylcholine, cholesterol and PEG-derivatized phosphatidylethanolamine (PEG-PE). Liposomes are extruded through filters of defined pore size to yield liposomes with the desired diameter. In addition, antibodies of various embodiments of the invention (including antigen binding
domain fragments such as Fab' fragments) can be conjugated to the liposomes as described in Martin et al., 1982, J. Biol. Chem. 257:286-288, via a disulfide interchange reaction. Administration of the recombinant antibodies of various embodiments of the invention includes local or systemic administration, including injection, oral administration, particle gun or catheterized administration, and topical administration. One skilled in the art is familiar with administration of expression vectors to obtain expression of an exogenous protein in vivo. See, e.g., U.S. Pat. Nos. 6,436,908, 6,413,942, and 6,376,471.

In another aspect, the invention provides a method of reducing the likelihood of occurrence in an animal of a disease characterized by the presence in the animal of a disease antigen, wherein the method comprising administering an effective amount of a therapeutic composition comprising a recombinant immunoglobulin of various embodiments of the invention, wherein the antigen specifically bound by the immunoglobulin of the therapeutic composition and the disease antigen are the same.

Vaccine preparation is generally described in Vaccine Design ("The subunit and adjuvant approach" (eds Powell M. F. \& Newman M. J., (1995) Plenum Press New York).

In another aspect, the invention provides a kit for determining the amino acid sequence of an antibody from an animal comprising (a) a means for obtaining nucleic acid sequences encoding immunoglobulin chain variable regions of multiple immunoglobulins from an animal, and (b) instructions for correlating mass spectra information from an antibody analyzed by mass spectrometry with predicted mass spectra information derived from the nucleic acid sequences to determine the amino acid sequence of the antibody.

The methods disclosed herein can be used to monitor circulating antibodies over time, e.g., in a subject immunized with an antigen. In these embodiments, samples can be taken from the subject at a plurality of time points (e.g., before and after immunization) and the methods disclosed herein used to identify circulating antibodies at each time point. The composition of circulating antibodies can be compared at the plurality of time points to determine the efficacy and/or time course of the vaccination. This can be useful for monitoring immune responses in individual subjects and also in the development of vaccines.

The following examples are provided to illustrate, but not to limit, the various
aspects and embodiments of the invention.

## Example 1

## Identifying Individual Antibody Heavy Chains from a Polyclonal Population of Antibodies that Specifically Bind an Antigen.

In this example, multiple monoclonal antibodies were derived from a polyclonal population of antibodies that specifically bound an antigen. Using the methods of various embodiments of the invention, the information from the genetic material database generated from nucleic acid molecules from the animal whose sera comprised the starting polyclonal population were compared to peptide database information from analysis of the monoclonal antibodies.

The nucleic acid sequences were obtained from splenocytes from an animal immunized with the antigen according to the methods described herein using primers specific for rabbit immunoglobulin chain-encoding sequences (see, for example, the primer sequences in Example 6 below). The CDR3 regions from the heavy chains of the polyclonal antibodies were ranked based on the number of times they appeared in the database and the percentage of times each CDR3 appeared among all of the CDR3 regions in the database. Table 2 shows the top 25 CDR3 regions and their frequencies. These results show that the same CDR3 sequences were found in many different antibodies in the polyclonal mixture. This information shows that antibodies that specifically bind to the same antigen often share sequences in their CDR3 regions (and presumably in the other CDR regions). This information shows that the methods described herein will be able to identify and isolate those immunoglobulin chains (or fragments thereof) that will specifically bind to the antigen.

Table 2

| SEQ ID NO: | CDR3 | Count | Percent |
| :--- | :--- | :--- | :--- |
| 29 | GVKF | 582 | $7.90 \%$ |
| 30 | GVSTNV | 530 | $7.20 \%$ |
| 31 | DPYDDPTYRGYGMDL | 372 | $5.05 \%$ |
| 32 | NPAVNTYAS | 345 | $4.69 \%$ |


|  | GGL | 198 | 2.69\% |
| :---: | :---: | :---: | :---: |
| 33 | HLFLHF | 196 | 2.66\% |
| 34 | HLFLNL | 172 | 2.34\% |
|  | GNV | 169 | 2.30\% |
|  | GNI | 143 | 1.94\% |
| 35 | HLFLNF | 129 | 1.75\% |
| 36 | GLGYVGSSVYIVKYINL | 126 | 1.71\% |
| 37 | DLIRVAGDTFYDGAFNL | 113 | 1.53\% |
| 38 | GRYNGWGYSNDL | 113 | 1.53\% |
| 39 | GGGTTLYTYFDL | 111 | 1.51\% |
| 40 | GLGYVGSDVYIVKYINL | 105 | 1.43\% |
| 41 | GGYGYGYGNTDFNL | 93 | 1.26\% |
| 42 | DDGGVRVDFDL | 87 | 1.18\% |
| 43 | VDDSGWMPFKL | 85 | 1.15\% |
| 44 | NVGSSSHYNLNL | 76 | 1.03\% |
| 45 | DGTDHGFNIDL | 72 | 0.98\% |
| 46 | STFRNSYARLAL | 69 | 0.94\% |
| 47 | IPYGWYSGGGAAPYFDL | 65 | 0.88\% |
| 48 | NAAIL | 62 | 0.84\% |
| 49 | AVSDNGYGMYWFNL | 61 | 0.83\% |
| 50 | ELAGYDVGVEF | 59 | 0.80\% |

For the creation of the peptide database, the following methods were used.

## Proteolytic Digestion of Antibodies

Approximately 10ug of the polyclonal population of antibodies was concentrated and buffer exchanged by ultrafiltration ( 0.5 ml 10 K Amicon: Millipore). The initial volume was first concentrated, then exchanged by adding 400 ul of 200 mM Hepes at pH 8 . Samples were denatured by resuspending in 80 ul of 8 M urea in pH 8 Hepes for 15 min at room temperature. Antibodies were reduced in 10 mM DTT at room temperature for 40 min .

Alkylation was performed for 1 hour with 20mM IAA. Urea concentration was reduced to a final concentration of 2 M . Samples were then divided equally by five and digested separately overnight at 37C with Trypsin, Lys-C, Glu-C, Pepsin, or Chymotrypsin respectively. For Pepsin digests, samples were concentrated and exchanged with 3M acetic
purified using Sep-Pack cartridges (Waters). Cleaned samples were lyophilized and resuspended for analysis on an LTQ Orbitrap Velos mass spectrometer.

## Mass Spectrometry

Peptide mixtures produced by digesting the antibody fraction with the proteases LysC, trypsin, chymotrypsin, Pepsin, or Glu-C (i.e., peptides were produced by digesting the antibody fraction with each of these proteases individually) were analyzed by LC-MS/MS individually using the LTQ Orbitrap Velos (Thermo-Fisher) hybrid mass spectrometer. Samples were loaded for 15 min using a Famos autosampler (LC Packings) onto a handpoured fused silica capillary column ( 125 um internal diameter 18 cm ) packed with MagicC18aQ resin ( $5 \mathrm{~m}, 200 \AA$ ) using an Agilent 1100 series binary pump with an in-line flow splitter. Chromatography was developed using a binary gradient at $400 \mathrm{nl} / \mathrm{min}$ of $8-$ $30 \%$ solvent B for 35 min (Solvent A, $0.25 \%$ formic acid (FA); Solvent B, $0.1 \%$ FA, $97 \%$ acetonitrile). As peptides eluted from the liquid chromatography column into the mass spectrometer, they were ionized and the peptide ion mass-to-charge ratios were measured to generate an MS1 spectrum. The mass spectrometer then selected the 20 most abundant peptide ions eluting at that moment and that had not been subjected to MS2 spectrum acquisition in the past 35 seconds, then isolated and fragmented, in turn, each of those 20 precursor peptide ions to produce 20 MS 2 product ion spectra. An entire cycle of acquiring one MS1 spectrum of precursor ions followed by acquiring 20 MS 2 product ion spectra in a data-dependent manner was accomplished in about 1.6 seconds, and then repeated continuously as peptides eluted from the liquid chromatography column. Charge-state screening was used to reject singly charged species, and a threshold of 500 counts was required to trigger an MS/MS spectrum. When possible, the LTQ and Orbitrap were operated in parallel processing mode.

## Database searching and data processing.

MS/MS spectra were searched using the SEQUEST algorithm against a genetic database. Search parameters included full enzyme specificity for Chymotrypsin, Glu-C, Lys-C, and trypsin, and no enzyme specificity for pepsin with a parent mass tolerance of 50 p.p.m., a static modification of 57.0214 on cysteine and dynamic modifications of 15.9949 on methionine. HCD spectra were searched with a fragment ion tolerance of $\pm 0.02 \mathrm{Da}$, while CID spectra were searched with a fragment ion tolerance of $\pm 1 \mathrm{Da}$. Peptides were filtered to a $1 \%$ peptide FDR via the target-decoy approach, using a linear discriminant function to score each peptide based on parameters such as Xcorr, $\Delta \mathrm{Cn}$, and precursor mass error.

## Results

Figure 4 schematically depicts the method followed in this example. The nucleic acid sequences were analyzed using the Kabat rules (see Kabat, E. A. et al., Sequences of Proteins of Immunological Interest, National Institutes of Health, Bethesda, Md., (1987) and Wu, T.T. and Kabat, E.A. J. Exp. Med. 132: 211-250 (1970)) to determine where the variable and CDR3 region (and the sequences thereof) were located within the sequences. Next, the percent coverage of CDR3 regions of the heavy chain of multiple monoclonal antibodies identified by Mass spectrometry was elucidated. As shown below in Table 3, sixteen different peptide sequences from the MS-analyzed polyclonal antibody mixture were identified, where each of the sixteen peptides comprised the entirety (i.e., $100 \%$ ) of the CDR3 region of the corresponding sequence from the nucleic acid sequences collected from the animal.

## Table 3

| SEQ ID NO: | CDR3 | \% CDR3 coverage |
| :--- | :--- | ---: |
|  | GNL | 100 |
|  | GNV | 100 |
| 29 | GVKF | 100 |
| 30 | GVSTNV | 100 |
| 51 | SRSTSYYINL | 100 |
| 45 | DGTDHGFNIDL | 100 |
| 52 | DGSDHGFNIDL | 100 |
| 53 | GADSIYRIYFDL | 100 |

## NVGSSSYYNLNL

Of the peptides listed in Table 3, five of the most frequent-occurring observed peptides by mass spectrometry were also seen as theoretical mass spectra derived from the information from the nucleic acid sequences. Thus, this experiment proved that by comparing and correlating the predicted mass spectra (and underlying sequences) derived from the nucleic acid sequences with the observed mass spectra from the actual peptide fragments from the polyclonal antibodies, the sequences of multiple monoclonal antibodies (or at least the heavy chains thereof) were readily obtained.

## Example 2

## Development of an influenza antigen-specific recombinant human antibody

During the winter of 2009-2010, a strain of H1N1 influenza virus infected a large number of humans, causing death and permanent injury. Using the non-limiting methods of various embodiments of the invention, neutralizing antibodies may be cloned from humans previously exposed to a similar virus strain, and used as a composition to treat human patients currently suffering from the disease.

Accordingly, elderly individuals who were known to have been exposed to the influenza virus during the 1918 influenza epidemic are screened for the presence of serum antibodies that can neutralize the 1918 virus. To do this, the method described in Yu et al., Nature 455: 532-536, 2008 (and online supplement; article and supplement incorporated herein by reference in their entirety) is followed.

Patients whose blood serum and/or plasma contains virus-neutralizing antibodies are identified, and blood is taken from these patients and separated into cells and serum and/or plasma.

From the blood cells, B lymphocytes are isolated according to standard methods (see, for example, the methods described here) and nucleic acid molecules from the B lymphocytes are obtained. Immunoglobulin chain-encoding nucleic acid molecules are isolated from these cells by PCR amplifying genomic DNA using primers that hybridize to regions upstream and downstream of the human immunoglobulin heavy (VH)- and light (VL)-chain variable-region genes. Methods for making such primers are standard in the field of immunology (see, e.g., the methods described in Marks and Bradbury, "PCR Cloning of Human Immunoglobulin Genes" in Antibody Engineering: Methods and Protocols, 248: 117-134, 2003, incorporated herein by reference).

These nucleic acid molecules obtained by using these primers for PCR amplification are used to populate the genetic material database. Within the genetic database, the nucleic acid sequences are further manipulated using standard software packages to determine the amino acid sequence of the polypeptide encoded by each nucleic acid sequence, and the encoded polypeptides are virtually digested with trypsin, where the predicted resulting peptides generated from such digest are used to generate predicted mass spectra.

From the blood from the patients, serum and/or plasma is collected. Antibodies present in the serum and/or plasma are isolated by standard methods. For example, serum proteins are passed through a protein A sepharose column, to which immunoglobulins adhere and non-immunoglobulin proteins do not. Because the individuals whose blood is collected are not newly exposed to the 1918 influenza virus, their serum antibodies are further enriched for antibodies that specifically bind to a 1918 viral antigen by passing the serum antibodies over a second column coated with 1918 virus (e.g., attenuated virus or fragments thereof). The bound antibodies are next treated with a protease (e.g., papain) or chemical protein cleavage reagent that specifically cuts near the hinge region of the immunoglobulin, and the non-adherent Fc portions removed. Finally, the bound Fab or Fab2 fragments are treated with trypsin to generate peptide fragments, and all fragments are then fractionated using liquid chromatography, with the fragments then being analyzed by mass spectrometry. Using an algorithm such as the Sequest program, the observed tandem mass spectra of the peptides are correlated with the predicted mass spectra from the nucleic acid sequences extracted from the patients' B lymphocytes. Using this process, at least one
peptide found within the predicted amino acid sequence of a unique immunoglobulin chain of the genetic material database may be identified. The nucleic acid sequence encoding this immunoglobulin chain (or variable region thereof) is then retrieved from the genetic database and synthesized using standard DNA synthesis methods. The synthesized DNA sequences are then subcloned into expression vectors which are then transfected into CHO cells. The recombinant antibodies produced by the cells are next isolated and tested for the ability to bind to the 1918 virus (or fragments thereof).

Recombinant antibodies produced using this method are then combined with a pharmaceutically acceptable carrier and administered to patients suffering from H1N1 virus infection. Because these recombinant antibodies are wholly human in origin, it is not expected that they will be rejected by the patients' immune systems.

## Example 3

## Obtaining Nucleic Acid Sequences

This protocol uses next generation sequencing (NGS), and is based on 454 NGS platform (FLX+, FLX or junior; commercially available from 454 Life Sciences, a Roche company, Branford, CT). Slight modifications will be needed for other high throughput NGS platforms and will be based on NGS manufacturing's instructions.

Mice are immunized with antigen of interest (peptide(s), recombinant proteins, virus, toxin, etc) with standard immunization protocols (see, e.g., Coligan et al., supra). Immune responses are monitored by plasma immunoglobulins titer against the specific antigen. Blood, spleen, bone marrow, lymph nodes, or any lymphoid organs can be collected and processed to isolate B cells according to standard methods. This isolation procedure can also be reduced if material is limited and replaced with a direct RT-PCR procedure using immunoglobulin variable domain specific PCR primers against heavy and light chains populations from the animal.

Of course in some embodiments, the nucleic acid sequences can be directly sequenced straight from the biological material (i.e., without being amplified prior to sequencing). Services and reagents for directly sequencing from nucleic acid sequences are commercially available, for example, from Helicos BioSicences Corp. (Cambridge, MA).

For example, Helicos' True Single Molecule Sequencing allows direct sequencing of DNA, cDNA, and RNA. See also US Patent Nos. 7,645,596; 7,037,687, 7,169,560; and publications Harris et al., Science 320: 106-109, 2008; Bowers et al., Nat. Methods 6: 493494, 2009; and Thompson and Milos, Genome Biology 12: 217, 2011 (all of which patents and publications are incorporated herein by reference in their entireties).

In some embodiments, the nucleic sequences are amplified (e.g., by polymerase chain reaction) prior to obtaining sequence information.

In one non-limiting example, an oligo dT PCR primer is used for RT-PCR. In another non-limiting example, gene-specific RT-PCR is performed using the PCR primers described below are used. In another example, PCR primers against heavy chain and light chain populations in a mouse have sequences set forth in PCT publication no. WO2010/097435, herein incorporated by reference.

With or without B cell enrichment, purified genetic materials (DNA or mRNA) will then be subjected to RT-PCR following standard procedures (see, e.g., Ausubel et al., supra). This is the library preparation stage of the genetic materials before NGS sequencing run. Reverse transcription (RT) reaction can apply oligo dT or immunoglobulin specific primers to generate cDNAs. Polymerase chain reaction procedure will apply immunoglobulin specific primers to amplify variable region of (rearranged or/and expressed) heavy and light chains from the sample.

These methods are described in further details below.

## Library Preparation

Sample preparation example:
Blood, spleen, bone marrow, or lymph nodes are isolated after mice received final boost with antigen. Mononuclear cells are isolated by Ficoll separation as previously described above. Ficolled cells are then washed by PBS, counted, and snap frozen for total RNA preparation.

Total RNA is isolated from the cells using the Qiagen RNeasy kit (commercially available from Qiagen Inc., Hilden, Germany) according to manufacturer's instructions, and the total

RNA is stored at $-80^{\circ} \mathrm{C}$.

10uM CST mouse RT-Ig primer or Oligo dT
2.5ug total RNA (splenocytes)

10 mM dNTP
Sterile, distilled water

For gene-specific RT-PCR or standard RT-PCR (using oligo dT), the following protocol may be used.

Incubate mixture at $65^{\circ} \mathrm{C}$ for 5 minutes and then place on ice.
5x cDNA Synthesis Buffer 4ul
0.1M DTT 1ul

Invitrogen Thermoscript RT (15U/ul) 1 ul
Mix contents gently and incubate at $60^{\circ} \mathrm{C}$ for 60 mins Terminate reaction by heating at $85^{\circ} \mathrm{C}$ for 5 mins cDNA is ready for use in making library
cDNA will then be subjected to PCR using CST 454 specific fusion mouse primers for Heavy and Light chains. The primers will have the following sequences:

Mouse 454 amplicon primers
Heavy Chains (Forward and Reverse primers)
HV1 CCATCTCATCCCTGCGTGTCTCCGACTCAGACGAGTGCGTGATGTGAAGCTTCAGGAGTC (SEQ ID NO: 1)
HV2 CCATCTCATCCCTGCGTGTCTCCGACTCAGACGCTCGACACAGGTGCAGCTGAAGGAGTC (SEQ ID NO: 2)
HV3 CCATCTCATCCCTGCGTGTCTCCGACTCAGAGACGCACTCCAGGTGCAGCTGAAGCAGTC (SEQ ID NO: 3)
HV4 CCATCTCATCCCTGCGTGTCTCCGACTCAGAGCACTGTAGCAGTTACTCTGAAAAGAGTC (SEQ ID NO: 4)
HV5 CCATCTCATCCCTGCGTGTCTCCGACTCAGATCAGACACGGAGGTCCAGCTGCAACAATCT (SEQ ID NO: 5)
HV6 CCATCTCATCCCTGCGTGTCTCCGACTCAGATATCGCGAGGAGGTCCAGCTGCAGCAGTC (SEQ ID NO: 6)
HV7 CCATCTCATCCCTGCGTGTCTCCGACTCAGCGTGTCTCTACAGGTCCAACTGCAGCAGCCT (SEQ ID NO: 7)
HV8 CCATCTCATCCCTGCGTGTCTCCGACTCAGCTCGCGTGTCGAGGTGAAGCTGGTGGAGTC (SEQ ID NO: 8)
HV9 CCATCTCATCCCTGCGTGTCTCCGACTCAGTCTCTATGCGGAGGTGAAGCTGGTGGAATC (SEQ ID NO: 9)
HV 10 CCATCTCATCCCTGCGTGTCTCCGACTCAGTGATACGTCTGATGTGAACTTGGAAGTGTC

HVFOR1 CCTATCCCCTGTGTGCCTTGGCAGTCTCAGTGCAGAGACAGTGACCAGAGT (SEQ ID NO: 11)

HVFOR2 CCTATCCCCTGTGTGCCTTGGCAGTCTCAGTGAGGAGACTGTGAGAGTGGT (SEQ ID NO: 12)
HVFOR3 CCTATCCCCTGTGTGCCTTGGCAGTCTCAGTGAGGAGACGGTGACTGAGGT (SEQ ID NO: 13)
HVFOR4 CCTATCCCCTGTGTGCCTTGGCAGTCTCAGTGAGGAGACGGTGACCGTGGT (SEQ ID NO: 14)

Kappa chains (Forward and Reverse Primers)
KV1 CCATCTCATCCCTGCGTGTCTCCGACTCAGCATAGTAGTGGATGTTTTGATGACCCAAACT (SEQ ID NO: 15)
KV2 CCATCTCATCCCTGCGTGTCTCCGACTCAGCGAGAGATACGATATTGTGATGACGCAGGCT (SEQ ID NO: 16)
KV3 CCATCTCATCCCTGCGTGTCTCCGACTCAGATACGACGTAGATATTGTGATAACCCAG (SEQ ID NO: 17)
KV4 CCATCTCATCCCTGCGTGTCTCCGACTCAGTCACGTACTAGACATTGTGCTGACCCAATCT (SEQ ID NO: 18)
KV5 CCATCTCATCCCTGCGTGTCTCCGACTCAGCGTCTAGTACGACATTGTGATGACCCAGTCT (SEQ ID NO: 19)
KV6 CCATCTCATCCCTGCGTGTCTCCGACTCAGTCTACGTAGCGATATTGTGCTAACTCAGTCT (SEQ ID NO: 20)
KV7 CCATCTCATCCCTGCGTGTCTCCGACTCAGTGTACTACTCGATATCCAGATGACACAGACT (SEQ ID NO: 21)
KV8 CCATCTCATCCCTGCGTGTCTCCGACTCAGACGACTACAGGACATCCAGCTGACTCAGTCT (SEQ ID NO: 22)
KV9 CCATCTCATCCCTGCGTGTCTCCGACTCAGCGTAGACTAGCAAATTGTTCTCACCCAGTCT (SEQ ID NO: 23)
KVFOR1 CCTATCCCCTGTGTGCCTTGGCAGTCTCAGCCGTTTCAGCTCCAGCTTG (SEQ ID NO: 24)
KVFOR2 CCTATCCCCTGTGTGCCTTGGCAGTCTCAGCCGTTTTATTCCAGCTTGGT(SEQ ID NO: 25)
KVFOR3 CCTATCCCCTGTGTGCCTTGGCAGTCTCAGCCGTTTTATTTCCAACTTTG (SEQ ID NO: 26)
Lambda Chains (Forward and Reverse Primers)
LV CCATCTCATCCCTGCGTGTCTCCGACTCAGTACGAGTATGCAGGCTGTTGTGACTCAGGAA (SEQ ID NO: 27)
LVFOR CCTATCCCCTGTGTGCCTTGGCAGTCTCAGCTTGGGCTGACCTAGGACAGT (SEQ ID NO: 28)

In all of the above sequences, the underlined sequences are for the 454 sequencing, the bolded sequences are barcodes for multiplexing, and the regular font sequences are mousespecific sequences.

The primers are used to amplify the above-described libraries as follows:

Heavy chain PCR:

CST454 mouse heavy chain primers mix cDNA 2x Phusion Master Mix H2O

1ul
1ul
12.5ul
10.5ul

Light chain PCR:
CST454 mouse light chain primers mix cDNA
2x Phusion Master Mix
1ul
1ul
12.5ul
10.5 ul

The PCR condition cycle conditions may be as follows in Table 4:
Table 4

| Step | Temperature | Time (in minutes) |
| :---: | :---: | :---: |
| 1: Denaturing Step | $98^{\circ} \mathrm{C}$ | $01: 30$ |
| 2: Denaturing Step | $98^{\circ} \mathrm{C}$ | $00: 10$ |
| 3: Annealing Step | $60^{\circ} \mathrm{C}$ | $00: 30$ |
| 4: Extension step | $72^{\circ} \mathrm{C}$ | $00: 30$ |

20 cycles are applied of steps 2-4 are applied. PCR products will then be subjected to Agencourt Ampure DNA purification (commercially available from Beckman Coulter Genomics, Danvers, MA) 2 times, following manufacture's protocol (see, e.g., the protocols of Beckman Coulter Genomics' Agencourt AMPure XP system).

Once the PCR/genetic library is prepared, all subsequent steps will follow 454 manufacturing protocols for emPCR and sequencing reactions. See publications by 454 Life Sciences Corp., a Roche Company, Branford, CT 06405 entitled, "Sequencing Method Manual, GS Junior Titanium Series" (May 2010 (rev. June 2010)) and "emPCR Amplification Method Manual - Lib-L, GS junior Titanium Series (May 2010 (rev. June 2010)), both of which are hereby incorporated by reference in their entirety.

Multiple samples can be combined at this stage into a single sequencing run. They will be distinguished by a unique barcode (or MID from 454 platform). For example, a
barcode is incorporated into the PCR primer.
In some embodiments, the emPCR Amplification Method Manual - Lib-L, GS junior Titanium Series (May 2010 (rev. June 2010); 454 Life Sciences Corp.) is followed. In some embodiments, the Sequencing Method Manual, GS Junior Titanium Series" (May 2010 (rev. June 2010); 454 Life Sciences Corp.) is next followed.

Sequencing data can be produced as FASTA files (or any standard file formats) and stored in a genetic material database. These sequence data will be used to generate the predicted mass spectra database to analyze the observed peptide mass spectra generated from the same animal's serum and/or plasma immunoglobulins. Standard programs can be used to do this. In this example, the predicted mass spectra were generated by the Sequest software package.

## Example 4

## Identifying Individual Antibody Chains from a Polyclonal Population

The methods described herein were next used to identify the sequence of individual antibodies from several different polyclonal populations. The methods of this example are shown schematically in Figures 2 and 4.

Using the methods described above in Example 2, three different polyclonal populations of antibodies that specifically bind to three different antigens were made into three different libraries. Deep sequencing using the 454 sequencing methods described above were performed using primers specific for rabbit immunoglobulin chain-encoding sequences to obtain three different genetic material databases.

Correspondingly, the genetic material databases were used to generate three different protein databases using the methods described in Example 3 above.

The results for the first antigen are shown in Tables 5 (light chain) and 6 (heavy chain); the second antigen are shown in Tables 7 (light chain) and 8 (heavy chain) and the third antigen are shown in Tables 9 (light chain) and 10 (heavy chain).

Table 5

| CDR3 | CDR3 <br> count | CDR3 coverage | Total Peptides | Unique <br> Peptides | CDR3 peptide |
| :---: | :---: | :---: | :---: | :---: | :---: |
| QGEFSCRDFDCTV <br> (SEQ ID NO: 61) | 16 | 100 | 58 | 30 | CQGEFSCRDFDCTVF (SEQ ID NO: 62) |
| AGGYKSSGDTVS (SEQ ID NO: 63) | 15 | 100 | 48 | 24 | YCAGGYKSSGDTVSF (SEQ ID NO: 64) |
| AGGYKSTTDGSA (SEQ ID NO: 65) | 9 | 100 | 29 | 17 | CAGGYKSTTDGSAF (SEQ ID NO: 66) |
| QQGRRSVDVDNV (SEQ ID NO: 67) | 8 | 100 | 25 | 12 | CADAATYYCQQGRRSVDVDNVFGGGTE (SEQ ID NO: 68) |
| QGEFNCDGVGCTT (SEQ ID NO: 69) | 2 | 100 | 17 | 9 | YCQGEFNCDGVGCTTF (SEQ ID NO: 70) |

Table 6

| CDR3 | $\begin{gathered} \text { CDR3 } \\ \text { count } \end{gathered}$ | CDR3 coverage | Total Peptides | Unique <br> Peptides | CDR3 peptide |
| :---: | :---: | :---: | :---: | :---: | :---: |
| GVRDWGDALDL (SEQ ID NO: 71) | 5 | 100 | 42 | 22 | $\begin{aligned} & \text { GVRDWGDALDLWGQGTLVTVSSGQPK } \\ & \text { (SEQ ID NO: 72) } \end{aligned}$ |
| LYNSVVGDDI (SEQ ID NO: 73) | 10 | 100 | 38 | 20 | LYNSVVGDDIWGPGTLVTVSLGQPK <br> (SEQ ID NO: 74) |
| LYNSVVGDDM (SEQ ID NO: 75) | 4 | 100 | 37 | 21 | LYNSVVGDDMWGPGTLVTVSLGQPK <br> (SEQ ID NO: 76) |
| GMPGSTSGNSNI <br> (SEQ ID NO: 77) | 2 | 100 | 34 | 20 | GMPGSTSGNSNIWGPGTLVTVSLGQPK <br> (SEQ ID NO: 78) |
| LYNSLVGDDI <br> (SEQ ID NO: 79) | 2 | 100 | 30 | 15 | LYNSLVGDDIWGPGTLVTVSLGQPK <br> (SEQ ID NO: 80) |
| KGDPGHPNGLFFTM (SEQ ID NO: 81) | 3 | 100 | 22 | 19 | KGDPGHPNGLFFTMWGPGTLVTVSFGQPK <br> (SEQ ID NO: 82) |
| GGGSHSGSAIYDMDP (SEQ ID NO: 83) | 2 | 100 | 20 | 14 | GGGSHSGSAIYDMDPWGPGTLVTVSSGQPK <br> (SEQ ID NO: 84) |
| GTSRGSDYRLDL (SEQ ID NO: 85) | 2 | 100 | 15 | 11 | GTSRGSDYRLDLWGQGTLVTVSSGQPK <br> (SEQ ID NO: 86) |
| GMPASTSGNSNI (SEQ ID NO: 87) | 2 | 100 | 14 | 14 | GMPASTSGNSNIWGPGTLVTVSLGQPK <br> (SEQ ID NO: 88) |
| $\begin{aligned} & \text { DAIANI } \\ & \quad \text { (SEQ ID NO: 89) } \end{aligned}$ | 2 | 100 | 10 | 8 | DAIANIWGPGTLVTVSLGQPK <br> (SEQ ID NO: 90) |
| $\begin{aligned} & \text { DKWMVFGDLRL } \\ & \quad \text { (SEQ ID NO: 91) } \end{aligned}$ | 2 | 100 | 9 | 4 | DKWMVFGDLRLWGPGTLVTVSSGQPK (SEQ ID NO: 92) |

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| Table 7 |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
| CDR3 | $\begin{aligned} & \text { CDR3 } \\ & \text { count } \end{aligned}$ | CDR3 <br> coverage | Total Peptides | Unique <br> Peptides | CDR3 peptide |
| QQGRTYSDVANV (SEQ ID NO: 93) | 1 | 66.67 | 42 | 20 | TYSDVANVFGGGTEVVVK (SEQ ID NO: 94) |
| QQGYSSYNVDNA (SEQ ID NO: 95) | 2 | 41.67 | 75 | 20 | NVDNAFGGGTEVVVK <br> (SEQ ID NO: 96) |
| QQGYSSSNVDNA (SEQ ID NO: 97) | 2 | 41.67 | 41 | 19 | NVDNAFGGGTEVVVK <br> (SEQ ID NO: 98) |
| LGTYDCRSADCNA (SEQ ID NO: 99) | 2 | 46.15 | 33 | 18 | SADCNAFGGGTEVVVK <br> (SEQ ID NO: 100) |
| QHGYYSNVDNA (SEQ ID NO: 101) | 2 | 45.45 | 46 | 18 | NVDNAFGGGTEVVVK <br> (SEQ ID NO: 102) |
| QQGFSSRNVDNA (SEQ ID NO: 103) | 2 | 41.67 | 24 | 18 | NVDNAFGGGTEVVVK <br> (SEQ ID NO: 104) |
| QQGYSSVNVDNA <br> (SEQ ID NO: 105) | 2 | 41.67 | 26 | 18 | NVDNAFGGGTEVVVK <br> (SEQ ID NO: 106) |
| QQGYTYNNVDNA (SEQ ID NO: 107) | 2 | 41.67 | 27 | 16 | NVDNAFGGGTEVVVK <br> (SEQ ID NO: 108) |
| $\begin{gathered} \text { LGTYDCRSGDCNV } \\ \quad(\text { SEQ ID NO: } 109) \end{gathered}$ | 1 | 46.15 | 25 | 15 | SGDCNVFGGGTEVVVK <br> (SEQ ID NO: 110) |
| QQGYTSNVDNA <br> (SEQ ID NO: 111) | 2 | 45.45 | 26 | 15 | NVDNAFGGGTEVVVK <br> (SEQ ID NO: 112) |
| QQGQTPENVDNA (SEQ ID NO: 113) | 2 | 41.67 | 22 | 14 | NVDNAFGGGTEVVVK <br> (SEQ ID NO: 114) |
| QQGSTYSDVANV (SEQ ID NO: 115) | 1 | 66.67 | 29 | 14 | TYSDVANVFGGGTEVVVK (SEQ ID NO: 116) |
| QQGATYSDVANV (SEQ ID NO: 117) | 1 | 66.67 | 63 | 13 | TYSDVANVFGGGTEVVVK (SEQ ID NO: 118) |
| QQGTTYSDVANV (SEQ ID NO: 119) | 1 | 66.67 | 25 | 13 | TYSDVANVFGGGTEVVVK (SEQ ID NO: 120) |
| QQGYTRSNVDNA <br> (SEQ ID NO: 121) | 2 | 41.67 | 21 | 11 | NVDNAFGGGTEVVVK <br> (SEQ ID NO: 122) |
| AGYKSYGNADID (SEQ ID NO: 123) | 4 | 66.67 | 24 | 10 | SYGNADIDFGGGTEVVVK (SEQ ID NO: 124) |
| QQGYTSSNVDNA (SEQ ID NO: 125) | 2 | 41.67 | 17 | 9 | NVDNAFGGGTEVVVK <br> (SEQ ID NO: 126) |
| LVSYDCSSADCNA (SEQ ID NO: 127) | 2 | 46.15 | 51 | 8 | SADCNAFGGGTEVVVK (SEQ ID NO: 128) |
| QQAYTSSNVDNA (SEQ ID NO: 129) | 2 | 41.67 | 4 | 3 | NVDNAFGGGTEVVVK <br> (SEQ ID NO: 130) |

Table 8

| CDR3 |
| :---: | :--- | :--- | :--- | :--- | :--- |$\quad$| CDR3 |
| :--- |
| count | CDR3 | Total |
| :--- |
| coverage | Peptides | Unique |
| :--- |
| Peptides | CDR3 peptide

Table 9

| CDR3 | $\begin{aligned} & \text { CDR3 } \\ & \text { count } \end{aligned}$ | CDR3 coverage | Total <br> Peptides | Unique <br> Peptides | CDR3 peptide |
| :---: | :---: | :---: | :---: | :---: | :---: |
| LANYDCSSGDCSV <br> (SEQ ID NO: 141 ) | 1 | 100 | 28 | 18 | CLANYDCSSGDCSVF (SEQ ID NO: 142 ) |
| QGNFDCSSADCSA (SEQ ID NO: 143 ) | 2 | 100 | 37 | 21 | CQGNFDCSSADCSAF (SEQ ID NO: 144 ) |
| QGNFDCTSADCSA (SEQ ID NO: 145 ) | 2 | 100 | 37 | 21 | CQGNFDCTSADCSAF <br> (SEQ ID NO: 146 ) |

Table 10

| CDR3 | CDR3 count | CDR3 <br> coverage | Total <br> Peptides | Unique <br> Peptides | CDR3 Peptide |
| :---: | :---: | :---: | :---: | :---: | :---: |
| DGTDHGFNIDL <br> (SEQ ID NO: 45) | 8 | 100 | 38 | 22 | DGTDHGFNIDLWGPGTLVTVSSGQPK <br> (SEQ ID NO: 147) |
| GNV | 2 | 100 | 36 | 21 | TSTTVTLQMTSLTAADTATYFCASGNV WGPGTLVTVSSGQPK $\quad$ (SEQ ID NO: 148) |
| $\begin{gathered} \hline \text { GVSTNV } \\ \text { (SEQ ID NO: } 30 \text { ) } \end{gathered}$ | 6 | 100 | 29 | 19 | GVSTNVWGPGTLVTVSSGQPK (SEQ ID NO: 149) |
| GVKF (SEQ ID NO: 29) | 4 | 100 | 30 | 18 | FCTRGVKF (SEQ ID NO: 150) |
| $\begin{gathered} \text { DGSDHGFNIDL } \\ \text { (SEQ ID NO: 52) } \end{gathered}$ | 6 | 100 | 29 | 16 | DGSDHGFNIDLWGPGTLVTVSSGQPK <br> (SEQ ID NO: 151) |
| NAAIL (SEQ ID NO: 152) | 10 | 100 | 34 | 16 | NAAILWGPGTLVTVSSGQPK <br> (SEQ ID NO: 153) |
| $\begin{array}{\|} \hline \text { SRSTSYYINL } \\ \text { (SEQ ID NO: 154) } \\ \hline \end{array}$ | 12 | 100 | 33 | 15 | SRSTSYYINLWGPGTLVTVSSGQPK (SEQ ID NO: 155) |
| GGDAGYGSFDAFGP (SEQ ID NO: 56) | 6 | 100 | 30 | 14 | GGDAGYGSFDAFGPWGPGTLVTVSSGQPK (SEQ ID NO: 156) |
| GVSTDV <br> (SEQ ID NO: 157) | 2 | 100 | 25 | 14 | GVSTNVWGPGTLVTVSSGQPK <br> (SEQ ID NO: 158) |


| NVGSSSYYNLNL <br> (SEQ ID NO: 54) | 6 | 100 | 28 | 14 | NVGSSSYYNLNLWGPGTLVTVSSGQPK <br> (SEQ ID NO: 159) |
| :---: | :---: | :---: | :---: | :---: | :---: |
| $\begin{array}{\|l\|} \hline \text { GVSTSV } \\ \text { (SEQ ID NO: } 160 \text { ) } \end{array}$ | 2 | 100 | 24 | 13 | GVSTNVWGPGTLVTVSSGQPK (SEQ ID NO: 161) |
| GGYAGAGYFDAFNP <br> (SEQ ID NO:162) | 2 | 100 | 21 | 12 | GGYAGAGYFDAFNPWGPGTLVTVSSGQPK (SEQ ID NO: 163) |
| $\begin{array}{\|c\|} \hline \text { NYNL } \\ \text { (SEQ ID NO: 164) } \\ \hline \end{array}$ | 6 | 100 | 26 | 12 | NYNLWGPGTLVTVSSGQPK (SEQ ID NO: 165) |
| RDGFSTDRYFNL (SEQ ID NO: 166) | 7 | 91.67 | 25 | 12 | DGFSTDRYFNLWGPGTLVTVSSGQPK (SEQ ID NO: 167) |
| DRGTGSGDYTPFNL <br> (SEQ ID NO: 168) | 5 | 71.43 | 26 | 12 | GSGDYTPFNLWGPGTLVTVSSGQPK (SEQ ID NO: 169) |
| $\begin{array}{\|l\|} \hline \text { DAAIL } \\ \text { (SEQ ID NO: 170) } \end{array}$ | 8 | 100 | 27 | 11 | NAAILWGPGTLVTVSSGQPK <br> (SEQ ID NO: 171) |
| GPYVDSTYYNL (SEQ ID NO: 172) | 6 | 100 | 23 | 11 | GPYVDSTYYNLWGPGTLVTVSSGQPK (SEQ ID NO: 173) |
| GSGDYTPFNL (SEQ ID NO: 174) | 6 | 100 | 23 | 11 | GSGDYTPFNLWGPGTLVTVSSGQPK (SEQ ID NO: 175) |
| YYDGADYHTYNL (SEQ ID NO: 176) | 6 | 100 | 21 | 11 | YYDGADYHTYNLWGPGTLVTVSSGQPK (SEQ ID NO: 177) |
| EFGNNGWNIDL <br> (SEQ ID NO: 178) | 6 | 100 | 21 | 10 | EFGNNGWNIDLWGPGTLVTVSSGQPK (SEQ ID NO: 179) |
| VEYGNDWGNL (SEQ ID NO: 180) | 6 | 100 | 20 | 10 | VEYGNDWGNLWGPGTLVTVSSGQPK (SEQ ID NO: 181) |
| YFDGADYHTYNL (SEQ ID NO: 182) | 6 | 100 | 20 | 10 | YFDGADYHTYNLWGPGTLVTVSSGQPK (SEQ ID NO: 183) |
| RFSGGGYGYDL (SEQ ID NO: 184) | 5 | 90.91 | 25 | 10 | FSGGGYGYDLWGPGTLVTVSSGQPK (SEQ ID NO: 185) |
| DRDL (SEQ ID NO: 186) | 6 | 100 | 19 | 9 | DRDLWGPGTLVTVSSGQPK (SEQ ID NO: 187) |
| $\begin{array}{\|c\|} \hline \text { GLDL } \\ \quad \text { (SEQ ID NO: 188) } \\ \hline \end{array}$ | 5 | 100 | 19 | 9 | YGLDLWGPGTLVTVSSGQPK (SEQ ID NO: 189) |
| $\begin{array}{\|c\|} \hline \text { YDVDSVSAYDL } \\ \text { (SEQ ID NO: 190) } \end{array}$ | 6 | 100 | 24 | 9 | YDVDSVSAYDLWGPGTLVTVSSGQPK (SEQ ID NO: 191) |
| EVVGYDYSGDL <br> (SEQ ID NO: 192) | 6 | 100 | 18 | 8 | EVVGYDYSGDLWGPGTLVTVSSGQPK <br> (SEQ ID NO: 193) |
| DPYDDPTY <br> (SEQ ID NO: 194) | 2 | 100 | 10 | 6 | DPYDDPTYR $\quad$ (SEQ ID NO: 195) |
| GGL | 1 | 100 | 3 | 3 | GGLVKPGASLTL $\quad$ (SEQ ID NO: 196) |

Tables 5-10 show peptides identified with high confidence ( $>99 \%$ certainty) by mass spectrometry (CDR3 peptide) that correspond to sequences (specifically the CDR3 region) generated by deep sequencing from the antibody repertoire of the animal. CDR 3 count shows the number of times a peptide was identified from the polyclonal antibody mixture that matched the CDR3 region. CDR3 coverage indicated the percent of those amino acids
in the CDR 3 region (shown in the CDR 3 column) that appear in the peptides identified by mass spectrometry relative to the total amino acids of the CDR3 region. Total peptides represent the total number of peptides by sequence identified by mass spectrometry corresponding to the full length variable region sequence determined by deep sequencing. Unique peptides represent the number of unique peptides by sequence identified by mass spectrometry corresponding to the full length variable region sequence determined by deep sequencing.

## Example 5

In another example, the following protocols can be used to generate the nucleic acid sequences and the polyclonal antibodies. The results show success in generating an antigenspecific antibody using these methods.

In these protocols, mice were immunized with an immunogenic P-ERK antigen. The genetic material database and peptide database can be generated using the following methods.

## I. Genetic Material Database:

## Cell isolation.

Spleens from immunized mice were flushed 5 times with 5 mL of RPMI/ $10 \%$ FCS using a syringe and 21 G needle. Cells were frozen in $90 \%$ FCS/10\% DMSO. A total of 50$100 \times 10^{\wedge} 6$ cells were isolated from each spleen.

## RNA Isolation and cDNA Synthesis.

Total RNA was isolated from Splenocytes according to manufacturer's protocol using QIAshredder (Qiagen cat\#79654) and RNeasy mini kit (Qiagen, Hilden, Germany; cat\#74104). RNA was DNAse treated on column as per a standard next generation sequencing protocol. Total RNA concentration was measured using an ND-1000 spectrophotometer (NanoDrop; commercially available from Thermo Scientific, Wilmington, DE).

The isolated RNA was used for first-strand cDNA synthesis by reverse transcription using Thermoscript RT-PCR system (Invitrogen (part of Life Technologies), Carlsbad, CA cat\#11146-024). cDNA was synthesized using 1.5 ug of RNA and oligo dT primer according to manufacturer's protocol. sequences of the primers used are shown below.

First round Primers, universal tail is underlined
Heavy chain sense primers:
$15 \quad \mathrm{~V}_{\mathrm{H}} 1.1$ :
ACGAGCTACGCACGAACTGCAGGTRTCCACTCC (SEQ ID NO: 197)
ACGAGCTACGCACGAATAGCAGGTGTCCACTCC (SEQ ID NO: 198) ACGAGCTACGCACGARGTACAGGTGTCCACTCC (SEQ ID NO: 199) ACGAGCTACGCACGAGCYACAGMTGTCCACTCC (SEQ ID NO: 200)
ACGAGCTACGCACGAACTGCAGGTGTCCWMTCC (SEQ ID NO: 201)
$\mathrm{V}_{\mathrm{H}} 1$.2:
ACGAGCTACGCACGARCTRCAGGTGTKCACTCC (SEQ ID NO: 202)
ACGAGCTACGCACGAGCTAWMGGTGTCCACTCC (SEQ ID NO: 203)
$25 \quad$ ACGAGCTACGCACGACCTCAGGTGTCCACTCC (SEQ ID NO: 204)
ACGAGCTACGCACGACCTCAGGTGTCCACTCC (SEQ ID NO: 204)
ACGAGCTACGCACGAGCTACAGGTGCTCACTCC (SEQ ID NO: 205)
ACGAGCTACGCACGAACTGCAGGTGTCCTCTCT (SEQ ID NO: 206)
$\mathrm{V}_{\mathrm{H}} 1.3$ :
ACGAGCTACGCACGAAYTGCAGGTGTCCAYTGC (SEQ ID NO: 207)
ACGAGCTACGCACGAGCTAMMGGTGTCCACTTC (SEQ ID NO: 208)
ACGAGCTACGCACGACTCCTGTCAKTAACTKCAGGT (SEQ ID NO: 209)
ACGAGCTACGCACGAAACTGCAGGTGTCTCTCT (SEQ ID NO: 210)
ACGAGCTACGCACGARCTRCAGGYGTCCACTCT (SEQ ID NO: 211)
$\mathrm{V}_{\mathrm{H}} 2$ :
ACGAGCTACGCACGACCAAGCTGTATCCTTTCC (SEQ ID NO: 212)
ACGAGCTACGCACGACCAAGCTGTGTCCTRTCC (SEQ ID NO: 213)

## $\mathbf{V}_{\mathbf{H}}$ and $\mathbf{V}_{\mathbf{L}}$ amplification.

A two-step PCR reaction was used to amplify the $\mathrm{V}_{\mathrm{H}}$ and $\mathrm{V}_{\mathrm{L}}$ genes. A mix of degenerate sense and anti-sense primers was used for the first round of PCR and a set of universal primers was used for the second round of PCR. Due to the large number of sense degenerate primers the heavy chain PCR is divided up into 8 separate reactions. The
$\begin{aligned} 40 & \mathrm{~V}_{\mathrm{H}} 3: \\ & \text { ACGAGCTACGCACGATGTTGACAGYCVTTCCKGGT (SEQ ID NO: 214) } \\ & \text { ACGAGCTACGCACGATGTTCACAGCCTTTCCTGGT (SEQ ID NO: 215) }\end{aligned}$

```
    V 44:
    ACGAGCTACGCACGATTTAAAAGGGGTCCAGTGT (SEQ ID NO: 216)
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$5 \quad \mathrm{~V}_{\mathrm{H}} 5$ :
ACGAGCTACGCACGATAYTTTAAAARGTGTCMAGTGT (SEQ ID NO: 217)
ACGAGCTACGCACGAGTTTTAAAAGGTGTCCTGTG (SEQ ID NO: 218)
$\mathrm{V}_{\mathrm{H}} 6-8$ :
10 ACGAGCTACGCACGACTYTTAAAAGGKGTCCAGWG (SEQ ID NO: 219)
ACGAGCTACGCACGACYTTTAMATGGTATCCAGTGT (SEQ ID NO: 220)
ACGAGCTACGCACGACTTTTACATGGTTTCAAGTGT (SEQ ID NO: 221)
ACGAGCTACGCACGAYTGTCCCTGCATATGTCYT (SEQ ID NO: 222)
$15 \quad \mathrm{~V}_{\mathrm{H}} 9-15$ :
ACGAGCTACGCACGAATGGCAGCWGCYCCAAG (SEQ ID NO: 223)
ACGAGCTACGCACGATTTATCAAGGTGTGCATTGT (SEQ ID NO: 224)
ACGAGCTACGCACGACTTTTAAAAGWTGTCCAGKGT (SEQ ID NO: 225)
ACGAGCTACGCACGAGTGACAGTCCTTCCTGGTAG (SEQ ID NO: 226)
ACGAGCTACGCACGACTTCCTGATGGCAGTGGTT (SEQ ID NO: 227)
ACGAGCTACGCACGAAGCTACAGGTATCCAATCC (SEQ ID NO: 228)
Heavy chain anti-sense primers:
IgG1:
25 CACTGGTGTGAGTCAATGCAGACAGATGGGGGTGTCG (SEQ ID NO: 229)
IgG2a:
CACTGGTGTGAGTCAAGACCGATGGGGCTGTTGTT (SEQ ID NO: 230)
IgG2b:
CACTGGTGTGAGTCAACAGACTGATGGGGGTGTTGTT (SEQ ID NO: 231)
30 IgG3:
CACTGGTGTGAGTCAAGACAGATGGGGCTGTTGTT (SEQ ID NO: 232)
Kappa chain sense primer:
35 ACGAGCTACGCACGAGACATYWWGATGACCCAGTCTCC (SEQ ID NO: 233)
Kappa chain anti-sense primer:
CACTGGTGTGAGTCACAGTTGGTGCAGCATCAGCCCG (SEQ ID NO: 234)
40 Second round Primers, universal tail is underlined
Heavy or Light chain sense primer:
CCTATCCCCTGTGTGCCTTGGCAGTCACGAGCTACGCACGA (SEQ ID NO: 235)
Heavy chain anti-sense primers:
MID97:
CCATCTCATCCCTGCGTGTCTCCGACTCAGctagtcactcCACTGGTGTGAGTCA
(SEQ ID NO: 236)

MID81:
CCATCTCATCCCTGCGTGTCTCCGACTCAGAGAGCGTCACCACTGGTGTGAGTCA (SEQ ID NO: 237)
MID24:
CCATCTCATCCCTGCGTGTCTCCGACTCAGTAGAGACGAGCACTGGTGTGAGTCA (SEQ ID NO: 238)

Light chain anti-sense primers:
MID34:
CCATCTCATCCCTGCGTGTCTCCGACTCAGcacgctacgtCACTGGTGTGAGTCA (SEQ ID NO: 239)
MID66:
CCATCTCATCCCTGCGTGTCTCCGACTCAGTCACGCGAGACACTGGTGTGAGTCA (SEQ ID NO: 240)
MID57:
CCATCTCATCCCTGCGTGTCTCCGACTCAGCGCGTATACACACTGGTGTGAGTCA (SEQ ID NO: 241)

In the above sequences, the underline and italic sequences are for 2 step PCR amplification, the underline sequences are for the 454 sequencing, the bolded sequences are the 454 key, the lower case sequences are barcode for multiplexing and the regular font capitalized sequences are mouse-specific sequences.

The PCR reactions were set up using the above-primers as outlined in Table 11.
Table 11 (First round Heavy chain PCR set-up)

| Sample | Sense primers | Anti-sense primers |
| :---: | :---: | :--- |
| 1 | $\mathrm{~V}_{\mathrm{H}} 1.1$ | IgG1, IgG2a, IgG2b, IgG3 |
| 2 | $\mathrm{~V}_{\mathrm{H}} 1.2$ | IgG1, IgG2a, IgG2b, IgG3 |
| 3 | $\mathrm{~V}_{\mathrm{H}} 1.3$ | IgG1, IgG2a, IgG2b, IgG3 |
| 4 | $\mathrm{~V}_{\mathrm{H}} 2$ | IgG1, IgG2a, IgG2b, IgG3 |
| 5 | $\mathrm{~V}_{\mathrm{H}} 3$ | IgG1, IgG2a, IgG2b, IgG3 |
| 6 | $\mathrm{~V}_{\mathrm{H}} 5$ | IgG1, IgG2a, IgG2b, IgG3 |
| 7 | $\mathrm{~V}_{\mathrm{H}} 4 \& \mathrm{~V}_{\mathrm{H}} 6-8$ | IgG1, IgG2a, IgG2b, IgG3 |
| 8 | $\mathrm{~V}_{\mathrm{H}} 9-15$ | IgG1, IgG2a, IgG2b, IgG3 |

For the first round, a $50 \mu \mathrm{~L}$ heavy chain PCR reaction contained $0.2 \mu \mathrm{M}$ of each sense primer ( 5 sense primers per reaction) and $0.2 \mu \mathrm{M}$ of each anti-sense primer (4 antisense primers per reaction), $10 \mu \mathrm{~L}$ of 5 x Phusion HF reaction buffer (Finnzymes (part of

Thermos Scientific), cat\#F-518), $1 \mu \mathrm{~L}$ of cDNA, $0.2 \mu \mathrm{M} \mathrm{dNTP}$ (NEB, cat\#N0447), $1 \mu \mathrm{~L}$ of Phusion Hot Start II DNA polymerase (Finnzymes, cat\#F-549L) and $28 \mu$ L RT-PCR Grade water (Ambion (a Life Technologies company), Austin, TX, cat\#AM9935). For the first round, a $50 \mu \mathrm{~L}$ light chain PCR reaction contained $0.2 \mu \mathrm{M}$ of the sense primer and $0.2 \mu \mathrm{M}$ of the anti-sense primer, $10 \mu \mathrm{~L}$ of 5x Phusion HF reaction buffer (Finnzymes, cat\#F-518), 1 $\mu \mathrm{L}$ of cDNA, $0.2 \mu \mathrm{M}$ dNTP (NEB, cat\#N0447), $1 \mu \mathrm{~L}$ of Phusion Hot Start II DNA polymerase (Finnzymes, cat\#F-549L) and $35 \mu \mathrm{~L}$ RT-PCR Grade water (Ambion, cat\#AM9935). The PCR thermocycle program was as follows: $98^{\circ} \mathrm{C}$ for $2 \mathrm{~min} ; 15$ cycles $\left(98{ }^{\circ} \mathrm{C}\right.$ for $0.5 \mathrm{~min}, 55^{\circ} \mathrm{C}$ for $0.5 \mathrm{~min}, 72^{\circ} \mathrm{C}$ for 1 min ); $72{ }^{\circ} \mathrm{C}$ for $5 \mathrm{~min} ; 4^{\circ} \mathrm{C}$ storage. PCR products were purified according to manufacturer's protocol using DNA clean and Concentrator -5 kit (Zymo Research Co., Irvine, CA, cat\#DR014).

For the second round, a $50 \mu \mathrm{~L}$ heavy chain PCR reaction contained $0.2 \mu \mathrm{M}$ of universal sense and universal anti-sense primer $10 \mu \mathrm{~L}$ of 5 x Phusion HF reaction buffer (Finnzymes, cat\#F-518), $10 \mu \mathrm{~L}$ of the purified first round PCR product, $0.2 \mu \mathrm{MdNTP}$ (NEB, cat\#N0447), $1 \mu \mathrm{~L}$ of Phusion Hot Start II DNA polymerase (Finnzymes, cat\#F-549L) and $19 \mu \mathrm{~L}$ RT-PCR Grade water (Ambion, cat\#AM9935). The PCR thermocycle program was: $98^{\circ} \mathrm{C}$ for $2 \mathrm{~min} ; 10$ cycles $\left(98^{\circ} \mathrm{C}\right.$ for $0.5 \mathrm{~min}, 55^{\circ} \mathrm{C}$ for $0.5 \mathrm{~min}, 72{ }^{\circ} \mathrm{C}$ for 1 min$) ; 72$ ${ }^{\circ} \mathrm{C}$ for $5 \mathrm{~min} ; 4^{\circ} \mathrm{C}$ storage. For the second round a $50 \mu \mathrm{~L}$ light chain PCR reaction contained $0.2 \mu \mathrm{M}$ of universal sense and universal anti-sense primer $10 \mu \mathrm{~L}$ of 5 x Phusion HF reaction buffer (Finnzymes, cat\#F-518), $10 \mu \mathrm{~L}$ of the purified first round PCR product, $0.2 \mu \mathrm{M}$ dNTP (NEB, cat\#N0447), $1 \mu \mathrm{~L}$ of Phusion Hot Start II DNA polymerase (Finnzymes, cat\#F-549L) and $19 \mu \mathrm{~L}$ RT-PCR Grade water (Ambion, cat\#AM9935). The PCR thermocycle program was: $98^{\circ} \mathrm{C}$ for $2 \mathrm{~min} ; 8$ cycles $\left(98^{\circ} \mathrm{C}\right.$ for $0.5 \mathrm{~min}, 55^{\circ} \mathrm{C}$ for 0.5 $\min , 72{ }^{\circ} \mathrm{C}$ for 1 min ); $72{ }^{\circ} \mathrm{C}$ for $5 \mathrm{~min} ; 4^{\circ} \mathrm{C}$ storage. PCR products were purified according to manufacturer's protocol using AMPure XP (Agencourt; Beckman Coulter Genomics, Brea, CA, cat\#A63881) and analyzed using an Agilent 2100 BioAnalyzer.

The sequences of the PCR products are then translated into predicted amino acid sequences which are then theoretically digested (e.g., with a protease and/or a chemical protein cleavage reagent) to produce virtual peptide fragments. These virtual peptide
fragments are then used to generate predicted mass spectra.

## II. Generation of Actual Mass Spectra from Peptide Fragments of Polyclonal

## Antibodies:

Polyclonal antibodies are purified from the sera and/or plasma of an animal (e.g., from the sera and/or plasma of the animal from whom the nucleic acid sequences are obtained). To purify the antibodies, the following methods are used:

## Protein-G Purification:

1 mL of magnetic protein-G beads (Millipore (Billerica, MA), cat\# LSKMAGG10) were added to each of four 15 mL conical tubes (Falcon (BD Biosciences, Franklin Lake, NJ), cat\#352097). The beads in each tube were washed twice with 10 mL of phosphate buffered saline $\mathrm{pH} 7.4,0.05 \%$ Tween-20 (PBST) and three times with 10 mL of PBS. Sera from three mice (ID 1262-2, 1262-4, 1263-4) were pooled together and diluted ten-fold to a final volume of 6 ml in PBS. 1.5 ml of the combined, diluted sera was then added to each tube of beads and incubated overnight at $4^{\circ} \mathrm{C}$. The flow through was collected and put through the purification process another two times. After the flow through was collected each tube was washed two times with 10 mL PBST and three times with 10 mL of PBS. Each tube was then incubated at $4^{\circ} \mathrm{C}$ for 30 minutes with 0.5 mL of 0.1 M pH 2.7 glycine to elute the IgG. The elution was repeated five times. All eluates were neutralized with 1 M Tris pH 8.5 , dialyzed overnight against PBS and protein concentration was measured with an ND-1000 spectrophotometer (Nanodrop). In total 2.5 mg of $\operatorname{IgG}$ was purified.

## Antigen column preparation:

5.0 mL of fresh streptavidin (SA) magnetic beads (Pierce, cat\#88817) were washed three times with 10 mL PBS, and incubated overnight at $4^{\circ} \mathrm{C}$ with 105 uL of a $20 \mathrm{mg} / \mathrm{ml}$ stock of biotin p-ERK peptide (a biotinylated form of Catalog No. 1150 commercially available from Cell Signaling Technology, Inc., Danvers, MA.) diluted in 5.0 mL of PBS. Flow through was discarded and beads were washed three times with 10 mL of PBS and aliquoted
into 10 low binding 1.7 mL tubes (Axygen (Union City, CA), cat\# MCT-175-L-C). Aliquoted beads were placed on a magnetic rack (Invitrogen, DynaMag) and PBS was removed prior to adding the dilute seras.

## Antigen-specific purification:

Protein-G purified IgG from above was added to the SA-magnetic beads coupled with biotin P-Erk peptide. After overnight incubation at $4^{\circ} \mathrm{C}$ the flow through was collected and the beads were washed with PBS-containing buffers.

IgG was then eluted with 5 fractions of 1.5 mL 0.1 M Glycine pH 3.5 , then 5 fractions of 1.5 mL 0.1 M Glycine pH 2.7 , then 5 fractions of 1.5 mL 0.1 M Glycine pH 1.8 and neutralized with 1 M Tris pH 8.5 . Eluates were assayed for P-ERK (i.e., phosphorylated ERK kinase, the antigen used to immunize the mice) reactivity using 96 -well plates coated with p-ERK -BSA peptide. Fractions with activity were quantitated by ELISA (Thermo, cat\#23300) and assayed for p-ERK reactivity by Western blot using lysates from Jurkat T cells (e.g., commercially available from the American Type Culture Collection or ATCC, Manassas, VA) treated with either 20uM U0126 for 1 hour or 200nM Tetradecanoyl-Phorbol-Myristic Acid (TPA) for 15 minutes. The fractions with the cleanest p-ERKreactivity were analyzed by mass spectrometry.

## Mass spectrometry analysis

The antibody-containing fractions were digested with at least one protease (e.g., trypsin) and/or at least one chemical protein cleavage reaction, and the resulting peptides subjected to analysis using mass spectrometry. The mass spectrometry analysis methods used to analyze the peptides are standard and have been described before in detail. (see, e.g., US Patent No. 7,300,753; Geiger et al., Nature Methods 7: 383-385, 2010; Elias and Gygi, Nature Methods 4: 207-214, 2007; Keshishian et al., Molecular and Cellular Proteomics 6: 2212-2229, 2007, all of which are hereby incorporated by reference in their entireties).

As described above (see, e.g., Example 3), the mass spectra were analyzed using as a reference the information in the genetic material database. To do this, MS2 spectra are
collected and then correlated one-by-one to predicted MS2 spectra from reference sequences (i.e., from the genetic material database) using a standard computational program that finds a match for every MS2 spectrum, even when it is not a good quality spectrum or a good match. Such programs are commercially available. For example, the Sequest software can $\mathrm{CA})$. The spectra that are identified as being good quality spectra or good matches to the genetic material database are mapped onto the reference sequences from the genetic material database. If a peptide MS2 can be mapped to more than one distinct component of the genetic database, it is unclear which component was present in the antigen-binding polyclonal antibody fraction as it could be one or more of those identified components. Thus, the process is repeated, and with repetition, evidence can be collected to show that some components correlate with collected MS spectra better than others. In other words, much of their variable region sequences are observed as MS2 spectra after enrichment by antigen binding. These elements are assumed to encode true antigen binding antibodies, and thus their sequences are constructed (e.g., on a synthetic oligonucleotide generator), cloned into an expression plasmid (e.g., pcDNA3.1 from Invitrogen), expressed in cells, and tested for antigen binding.

## Results

As shown in Fig. 5, the correlation of the actual mass spectrometry results from the peptide fragments with the theoretical mass spectrometry information from the nucleic acid sequences allowed the identification the sequences of heavy and light chain fragments. Those peptides that had the highest degree of confidence as far as mass spectrometry coverage is concerned and correlation to the nucleic acid sequences are shown. The nucleic acid sequence encoding the full length chain comprising the actual peptide fragments was synthesized and cloned into a recombinant expression vector. By random pairing, heavy and light chains were combined and expressed together in a cell to produce (i.e., create) recombinant antibodies (see, e.g., method of US patent nos. 4,816,397; 4,816,567; and US patent application no. 20110045534). Figure 6 is a table showing the results of an ELISA experiment using pERK-coated plates. As can be seen, several pairings of the chains
identified in Fig. 5 resulted in antibodies that were able to specifically bind to the p-ERKcoated plates (positive antibodies shown in yellow in Fig. 6, and the positive peptides are shown in red in Fig. 5).

Surprisingly, these results showed that neither frequency of peptide occurrence alone nor frequency of CDR3 count alone predicted usage of a particular antibody chain that specifically bound to the antigen. For example, light chain nucleic acid sequence ref. no. G623FKB01A3GC7 matched to 235 peptides from LC-MS/MS (i.e., liquid chromatography, tandem mass spectrometry) analysis and light chain nucleic acid sequence ref. no.

G623FKB01AXJ1C had a sequence that appeared in 1068 times in a single NGS run (see Fig. 5, lower table). However, neither of these, when combined with a heavy chain, was actually able to form an antibody that could specifically bind to the pERK antigen. This result is very surprising and showed that method of Reddy et al., Nature Biotechnology 28(9):965-969, 2010, which relied solely on nucleic acid sequence frequency from the NGS analysis, would have missed the true antigen-binding sequence. Thus, the methods described herein can be used reliably to identify and isolate an antibody that specifically binds to a chosen antigen.

## Example 6

An antigen-specific rabbit antibody was generated in accordance with the methods described herein. To do this, the following protocols were followed.

## Rabbit splenocyte RNA purification

The p-MET antigen (Cell Signaling Technology, Inc., Danvers, MA Catalog \# 1645) was used to immunize rabbits using standard methods. Immunized rabbits who had antigenspecific sera (i.e., sera containing polyclonal antibodies that specifically bound to the immunizing antigen) were sacrificed after a final antigen injection (boost). 50 ml of blood was collected and spleen or other lymphoid organs was collected. 10 million splenocytes were used for RNA purification. Serum and/or plasma from the 50 ml collected blood was also set aside for antigen specific antibody affinity purification.

RNA was purified from splenocytes using Qiagen's RNeasy kit (Qiagen cat\# 74104)
following the manufacturer's protocol. On column DNase I-treatment was conducted to eliminate contaminating genomic DNA by incorporating a DNase I digest step. After the RW1 buffer wash, DNase I (Qiagen cat\# 79254) diluted in RDD buffer was applied to the RNA purification column and incubated for 20 minutes at room temperature. The column was then washed once more with RW1 buffer, followed by two washes with RPE buffer, and the RNA was eluted with either 30 or $50 \mu \mathrm{l}$ water. The concentration of the RNA was determined by absorbance measured on a Nanodrop spectrophotometer (Thermo Scientific) at wavelength 450 nm .

## cDNA synthesis and generation of amplicons by PCR

RNA isolated from rabbit splenocytes was first reverse transcribed using Invitrogen's Thermoscript reverse transcriptase (Invitrogen cat\#12236-022) as shown below:

| DNase treated RNA: | 5 uL |
| :--- | :--- |
| Oligo dT primer $(50 \mathrm{uM}):$ | 1 uL |
| dNTP's $(10 \mathrm{mM}):$ | 2 uL |
| dI H2O: | 4 uL |

Incubate at $65^{\circ} \mathrm{C}$ for 5 min , place on ice for 2 minutes, then add the following:

| 5X cDNA buffer: |  |
| :--- | :--- |
| 0.1mM DTT: | 4 uL |
| RNAse OUT: | 1 uL |
| dI H2O: | 1 uL |
| ThermoScript: |  |
|  | 1 uL |

The mixture was incubated at $50^{\circ} \mathrm{C}$ for 1 hour, followed by a heat-inactivation step at $85^{\circ} \mathrm{C}$ for 5 minutes. Finally, the complementary RNA strand was eliminated from the cDNA by adding $1 \mu 1$ of RNase H (Invitrogen (Carlsbad, CA), cat\#18021-071) and incubating at $37^{\circ} \mathrm{C}$ for 20 minutes.

Amplicons of heavy, kappa and lambda chain variable regions for sequencing were generated by PCR as follows.

## Kappa Chain Fusion Primers

| Meverse |  |
| :--- | :--- |
| MID16 | CCATCTCATCCCTGCGTGTCTCCGACTCAGtcacgtactaGAAGAGGAGGACAGWAGGCGC |
|  | (SEQ ID NO: 244) |

Forward

## CCTATCCCCTGTGTGCCTTGGCAGTCTCAGATGGACATGAGGGCCCCC (SEQ ID NO: 245)

Lambda Chain Fusion Primers
Reverse

| MID39 | CCATCTCATCCCTGCGTGTCTCCGACTCAGtacagatcgtCTTGTTGTCCTTGAGTTCCTCAGAGGA |
| :--- | :--- |
|  | (SEQ ID NO: 246) |

Forward
CCTATCCCCTGTGTGCCTTGGCAGTCTCAGATGGCCTGCACCCCG
(SEQ ID NO: 247)

In the above sequences, the underline sequences are for the 454 sequencing, the bolded sequences are 454 key, the lower case sequences are barcode for multiplexing and the regular font capitalized sequences are rabbit-specific sequences.

PCR amplification was done using Finnzyme's Phusion Hot Start II polymerase (Thermo Scientific cat\# F-540S) where the reaction mix and conditions were set up as shown below:

Reaction mixture:
cDNA:
5X Buffer GC:
10 mM dNTP mix:
Phusion HotStart II:
2.5 uL
0.25 uL

Primers (forward+reverse) $30 \mathrm{uM}: \quad 0.25 \mathrm{uL}$
Water: $\quad 16.75 \mathrm{uL}$

## PCR program:

Step $198^{\circ} \mathrm{C}-1.5$ minutes
Step $298^{\circ} \mathrm{C}-10$ seconds

Step $360^{\circ} \mathrm{C}-30$ seconds
Step $472^{\circ} \mathrm{C}-30$ seconds
Step 5 Repeat steps 2 through 4, 20 times
Step $672^{\circ} \mathrm{C}-2$ minutes

Step 7 - hold
To ensure the absence of any false amplification from contaminating template in any of the reagents, duplicate reactions were set up for each mixture (4 separate reactions for heavy chain, and one for each light chain) where the cDNA template was substituted with water. These negative control reactions with no template were run at the same time as the samples containing template. Upon completion of the PCR program, $3 \mu \mathrm{l}$ of each reaction (including the negative controls) were analyzed by electrophoresis on a $1.5 \%$ TAE agarose gel for the presence of the amplicons when template was added to the reaction but not in the absence of cDNA. Figure 7 shows the results of these electrophoresis gels.

## Amplicon purification, analysis, quantitation, and preparation for $\mathbf{4 5 4}$ sequencing

In order to eliminate excess primers and/or primer dimmers in the PCR samples, amplicons were purified using Agentcourt Ampure magnetic beads (Beckman Coulter cat\#A63881) following the manufacturer's protocol (000387v001). The eluted amplicons after Ampure purification were then analyzed for purity and absence of any contaminating DNA species on the Agilent 2100 Bioanalyzer using the high sensitivity DNA chip (Agilent Technologies cat\# 5067-4626) by following the manufacturer's protocol.

Once the purity of amplicons was verified, the concentration of the DNA was quantified on a fluorometer using the Quant-iT PicoGreen dsDNA Assay Kit (Invitrogen cat\#P7589) as described in the manufacturer's protocol. The Lambda DNA provided in the kit was used as a concentration standard with which a standard curve was generated from $100 \mathrm{ng} /$ well to $1.56 \mathrm{ng} /$ well. The fluorescence of each amplicon diluted 100 -fold in TE buffer was measured in duplicate, and the concentration of DNA was determined according to the linear portion of the standard curve. All fluorescence measurements were done in black 96well plates. If the value of fluorescence was out of the linear range of the standard curve, the samples were remeasured with either larger or smaller dilutions in order to capture fluorescence values that fall within the linear range. Using the approximate size in base

Each amplicon was normalized to $1 \times 10^{7}$ molecules $/ \mu \mathrm{l}$, then mixed at a ratio of heavy chain: kappa chain: lambda chain at $3: 3: 1$ by volume, vortexed, and finally diluted 1:10 to obtain a final concentration of the mixture at $1 \times 10^{6}$ molecules $/ \mu \mathrm{l}$.

10 Emulsion PCR amplification, bead enrichment, bead counting and sequencing
Emulsion PCR was conducted following the 454 published protocol: "emPCR Amplification Method Manual - Lib-L" (Edition: May 2010 (Rev. April 2011, herein incorporated by reference in its entirety) with the following modifications: Section 3.1.3 Step 2)

| Reagent | Volume $(\mu \mathrm{l})$ |
| :--- | :--- |
| Mol. Bio. Grade water | 458 |
| Additive | 515 |
| Amp Mix | 270 |
| Amp Primer | 32 |
| Enzyme Mix | 70 |
| PPiase | 2 |
|  |  |
| Total | 1347 |

Aperture: $\quad<100 \mu \mathrm{~m}>$
Aperture Kd: $<60.04>$
Set Upper cutoff: $<30.00 \mu \mathrm{~m}>$
Set Lower cutoff: $<10.00 \mu \mathrm{~m}>$
Count Mode:
<between>
Once the sequencing beads were enriched, from step 3.7 of "emPCR Amplification Method Manual - Lib-L", the beads were counted on Beckman Coulter's Z2 Particle Counter with the following settings:
Metered Volume:
$<0.5 \mathrm{ml}>$

Resolution:
<256>
The concentration of beads was calculated as:
Concentration of beads $=[$ Avg. reading from particle counter * 4] beads $/ \mu \mathrm{l}$

The enriched beads from the emulsion PCR were sequenced on the 454 Sequencer (Roche) following the 454 sequencing protocol for GS FLX+ or GS Junior.

The peptide fragments of the polyclonal antibodies collected from the sera of immunized rabbits were generated as described above for mice (see, for example, Example 6). Briefly, the following protocol was used.

## Peptide-affinity purification of rabbit IgG

1. Re-suspend the peptide-affinity resin and take 0.4 ml of the slurry into a new column (Bio-rad, $731-1550,0.8 \times 4 \mathrm{~cm}$ ), and this should make 0.2 ml settled purification resin. If necessary, make a control column of either blank resin or an un-related peptideaffinity resin of equal volume. The blank resin was made with no peptide in the conjugation process.
2. Wash the column with 10 ml PBS, and let it drain completely.
3. Load the Protein-A purified total IgG. Cap the bottom first and wrap with paraffin. Add $3-5 \mathrm{ml}$ of total IgG . Cap the top and wrap with paraffin.
4. Rotate on a roller for 15 min at RT.
5. Collect the flow through. Un-cap the top first, then the bottom, let the column drain completely.
6. Wash with $10 \mathrm{ml} \mathrm{PBS}, 3$ times (wash the column wall to make sure that all the resin is packed at the bottom).
7. Wash with 10 ml 1 x RIPA.
8. Wash with $10 \mathrm{ml} 20 \%$ Acetonitrile in PBS pH 7.4 .
9. Wash with $10 \mathrm{ml} 60 \%$ Ethylene glycol in PBS pH7.4.
10. Wash with 10 ml 2.0 M NaCl in $\mathrm{PBS}, \mathrm{pH} 7.4$.
11. Elute with 5 ml 0.1 M Glycine pH 3.5 , neutralized immediately with 70ul 1M Tris pH8.5.
12. Elute with 5 ml 0.1 M Glycine pH 2.7 , neutralized immediately with 300ul 1 M Tris pH8.5.
13. Elute with 5 ml 0.1 M Glycine pH 1.8 , neutralized immediately with 800 ul 1 M Tris pH8.5.
14. All or the fractions of interest are measured for IgG concentration using Rabbit IgG ELISA plates (provided by Molecular assay/ELISA group).
15. The antigen-specific activity can be assessed using ELISA and/or Western blot. The specific activity can also be assessed after normalizing all fractions to the same concentration.
16. Purified antibody materials are ready to be processed for LC-MS/MS

Liquid chromatography-tandem mass spectrometry (LC-MS/MS) analysis was performed on peptides from the purified antibodies (i.e., the purified antibodies were digested and the peptides analyzed) as described above. The resulting mass spectra were correlated with the theoretical mass spectrometry data based on information in the genetic material database.

As shown in the table set forth in Figure 8, a number of heavy and light chain peptides were identified by correlating the actual (i.e., observed) mass spectrometry of the peptides with the theoretical mass spectrometry data from the nucleic acid sequences. The frequency of occurrence of these peptides is shown in the right-most lane of the table. These chains were chosen based on their coverage of CDR3 (in most cases $100 \%$ ), and the underlying nucleotide sequences retrieved from the genetic material database and synthesized. Six heavy chain was randomly combined with five light chain (shown in red in Fig. 8), and the resulting antibodies tested using ELISA (with antigen-coated plates) and Western blotting analysis (against Hela cells untreated (- lanes) or treated with Human Growth Factor (+ lanes), where the HGF-treated cells are known to express the p-MET antigen. The results of the Western blotting analysis are shown in Fig. 9. A p-MET specific antibody (commercially available from Cell Signaling Technology, Inc., Danvers, MA, catalog no. 3126) was used as a control. The antibodies generated in accordance with the methods described herein that showed high specific binding to the antigen in the cell lysates are shown in bold red in Fig. 8 (i.e., heavy chain ref nos. GXRYQP201BIQD2 and GXRYQP201A97DZ and light chain ref nos. GXRYQP201A291T and GXRYQP201BRIWK and GXRYQP201ALDF5). Note that Fig. 9 shows only two of the 6 different antibodies that specifically bound to antigen generated in this example (i.e., Fig. 9
shows only the two antibodies that use the GXRYQP201BIQD2 heavy chain coupled with the GXRYQP201A291T light chain and the GXRYQP201BRIWK light chain.

Again, as observed with the mouse antibody, the chains with the highest frequency did not result in formation of an antigen-specific antibody (compare heavy chain

Table 12. Functionally relevant monsoclonal matibodies against multiple targets ideutified by the
NGS:LC-MSMS plationm tested by ELISA and Westem blot (WB).

New Zealand white rabbits were immunized with human Progesterone Receptor A/B specific (PR A/B) peptides conjugated to keyhole limpet hemocyanin (KLH). Antigenspecific antibody activity in the crude serum of each animal was screened to select the rabbit with the highest ELISA and Western blot signals to PR A/B. Serum from this animal was collected from 20 mL of blood, and RNA was obtained from splenic B cells. Total $\gamma$ immunoglobulin ( IgG ) was isolated from the serum using a protein A sepharose column,
and antigen-specific polyclonal antibodies were purified by affinity chromatography using a custom column consisting of antigen-specific peptide conjugated to sepharose beads. Bound IgGs were washed extensively with PBS then subjected to sequential elutions with progressively acidic buffers ( $\mathrm{pH} 3.5, \mathrm{pH} 2.7$ and pH 1.8 ) (Fig. 10a). Fractions from each elution were collected, neutralized, and screened by antigen specific ELISA and Western blotting of lysate from the PR A/B expressing cell line T47D and the PR A/B negative cell line HT1080 (Fig. 10a). It was found that PR A/B Western blot specific activity was greatly enriched in the pH 1.8 fraction, to a lesser extent in the pH 2.7 fraction, and was undetectable in the pH 3.5 fraction when the polyclonal fraction was concentration matched. The pH 1.8 fraction was therefore used for LC-MS/MS analysis.

To generate a custom database of Ig V-region sequences by NGS, RNA was isolated from total splenocytes collected from the same animal that showed strong specific activity to PR A/B. Ig heavy and light chain variable region amplicons were generated using rabbit Igspecific $\gamma$ and $\kappa$ chain primers to amplify the entire V-region. Primers contained barcodes and followed the specific requirements for 454 titanium fusion primer design for the Roche 454 NGS platform. To increase the number of V-region sequences collected, we combined three 454 GS Junior sequencing runs consisting of $\gamma$ and $\kappa$ chains that resulted in a total of 80,000 passed filter reads, of which 44,363 contained the entire V-region and provided the basis for the proteomic approach described below. Sequences collected included 5,279 unique $\gamma$ chain complementarity determining region 3 (CDR3) sequences and 11,681 unique $\kappa$ chain CDR3 sequences of varying length that followed a Gaussian distribution. Consistent with previous data, this rabbit preferentially used VH1 (V1S69+V1S40 $>64 \%$ ) followed by VH4 (V1S44+ V1S45 ~30\%) in heavy chain VDJ rearrangement (Becker et al., Eur J Immunol 20: 397-402, 1990, Knight, Annu Rev Immunol 10: 593-616, 1992, Mage et al., Dev Comp Immunol 30: 137-153, 2006).

Next, the pH 1.8 fraction was examined by LC-MS/MS based on its previous activity (Fig. 10a). To maximize sequence coverage, 5 ug of polyclonal antibody was divided evenly and digested separately by chymotrypsin, elastase, pepsin and trypsin. A total of four LCMS/MS runs using a 45 -minute gradient were collected using an Orbitrap Velos (Thermo Fisher), producing an average of 10,000 spectra per run (Fig. 10b). To estimate the false-
discovery rate (FDR), the target/decoy approach was used by generating a composite database of forward and reverse-oriented sequences (Elias et al., Nat Methods 4: 207-214, 2007), and each LC-MS/MS run was searched using the SEQUEST (Yates et al., Anal Chem 67: 1426-1436, 1995) program. Peptide spectral matches (PSMs) were filtered to a final FDR of $\leq 2 \%$ using a linear discriminant analysis (Huttlin et al., Cell 143: 1174-1189, 2010) taking into account enzyme specificity when possible (chymotrypsin/trypsin). An example of a high confidence heavy chain CDR3 peptide identified using this method is shown in Fig. 10c. Individual runs were combined and a total of $2,356 \mathrm{~V}$-region PSMs were identified with a FDR of $1.8 \%$.

A database of antibody V-region sequences is analogous to a database of protein isoforms. As a result, traditional approaches using shotgun sequencing by LC-MS/MS in which only a few peptides are often used to confidently identify a protein are insufficient for identifying an antibody V-region sequence in a polyclonal antibody mixture. In addition, since antibody V-region sequences can vary by as little as one amino acid, high mass accuracy helped provide additional confidence in PSMs. Each V-region PSM with a mass error $\geq-5$ and $\leq 5 \mathrm{ppm}$ as determined by SEQUEST was mapped back to the entire V-region database to address PSM redundancy and coverage across the dataset (Fig. 10d). After remapping, the total number of peptides, the unique number of peptides, spectrum share (total peptides mapping to sequence / total V-region PSMs), total V-region sequence coverage, and CDR3 coverage were determined for each V-region sequence. In order to identify V-region sequences with high confidence that are likely to be enriched from the polyclonal mixture, empirically stringent criteria were applied in the proteomics analysis including: a) overall high coverage ( $\geq 65 \%$ ), b) at least 12 unique peptides due to high degree of homology of V -region sequences, and c) high hyper-variable region coverage, specifically, $\geq 95 \%$ coverage of CDR3. Although V-region sequences could be identified using one protease alone, it was found that because of the high degree of variability in Vregion sequences along with the unpredictable complexity of a polyclonal mixture, it was advantageous to use multiple proteases to increase V-region coverage. For example, as shown in Fig. 10e, multiple overlapping peptide fragments from different proteases contributed to the identification of the entire CDR3 of both heavy and light chain sequences.

Identifying unique PSMs across multiple runs from multiple proteases that map to the same V-region sequence increased spectral counts and coverage across the entire V-region sequence, provided higher confidence that specific V-region sequences were present in the polyclonal mixture, and further increased confidence in the NGS sequence quality (Kircher et al., Bioessays 32: 524-536, 2010). Using the filtering criteria described above, a total of ten $\gamma$ and eight $\kappa$ chain sequences of high confidence were identified from the pH 1.8 elution fraction (Table 13).
$100 \%$ CDR3 Coverage and $\geq 65 \%$ V－region Coverage

| NGS Ref．\＃ | Total <br> Paptide Cosunt | 5s variable Region Covernge | Cor3 Sequence | $\begin{aligned} & \text { SEO } \\ & \text { ID } \\ & \text { NO: } \end{aligned}$ | SGS rank by cars frequency | Germbine v（0） |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| kx乡⿰幺幺丱M\％\％\％\％ | \％3 | ¢\％๕． | M\％\％ | 346． | \％\％ |  |
|  | 93 | 92303 | GF3 | 397 | \％ | 164V15859＊14 |
| 《isk Moysweky | W． | צ... | Wूy\％ | W＊N | W．．． |  |
|  | 70 | 88.31 | blgme | 399 | 361 | GLEVEAS5，54． 5.34 |
|  | k． | （4y\％ | W\％ |  | K． |  |
|  | 59 | $\$ 772$ | ¢7\％ | 400 | 337 |  |
| k | Kis |  | kMxiky | Win | Wi. |  |
|  | 50 | 8.34 | 6F3L | 402 | 109 | 164423569．＊34 |
|  | 4． | $\text { \#. . } \mathrm{k} \text { \% }$ |  |  | Wikm |  |
|  | 35 | as 34 | G\％fl． | 404 | 843？ | cisfunses．＊， 14 |



Table 13．Identification of bigh confidence heswy and high chains．Heary ard hight ckains with low


 expression of combinmanial antibodies for chamoterization．Whas mak indicates the frequency making of
 gene wa be adenumed．

Despite providing evidence for the existence of high confidence V-region sequences present in affinity purified serum, direct information on cognate heavy and light chain pairing is absent from LC-MS/MS data due to proteolysis and the reduction of disulfide bonds during sample preparation. As a result, all possible combinations of heavy and light chain pairings were expressed ( $8 \times 10$ matrix for a total of 80 antibodies, in one 96 -well plate transfection) in addition to the highest rank heavy and light sequences observed by NGS frequency and screened for antigen-specific binding activity to PR A/B peptide by ELISA. A total of 12 heavy and light chain pairs were positive by antigen-specific ELISA (Fig. 11a). Each antigen-specific ELISA-positive clone was then tested by Western blot for specificity against endogenously expressed PR A/B in cell lysates (Fig. 11b). Six clones were found that specifically bound to PR A/B (Fig. 11b); two clones showed a much stronger signal compared to the original polyclonal mixture when assayed at the same antibody concentration. Antigen-specific clones positive by Western blot were further characterized in additional assays. One monoclonal antibody, clone F9 and clone C1, exhibited superior signal and specificity in Western blotting and immunohistochemistry (IHC) (Fig. 11b-c) and also reacted specifically in flow cytometry (FC) and immunofluorescence (IF) assays where the polyclonal mixture failed (Fig. 11d-e). In contrast, $\gamma$ and $\kappa$ chains selected by virtue of their highest NGS rank did not yield antigen-specific antibodies. CDR3 containing peptides were not observed from the highest NGS rank $\gamma$ and $\kappa$ chains, and none of the CDR3 sequences from the 30 highest rank $\gamma$ or $\kappa$ chains was identified with high confidence by our proteomics approach. It could not be ruled out that the absence of activity may be due to a lack of cognate pairing, but the fact that none of these chains was observed by LC-MS/MS suggests none of the highest rank NGS chains was specific against the antigen. Thus, in these experiments antigen-specific antibodies could not be identified relying on NGS rank alone.

In order to visualize clonal diversity, phylogenetic analysis (Dereeper et al., Nucleic Acids Res 36: W465-469, 2008) was performed on high confidence heavy and light chain Vregion sequences shown in Table 13. Closely related sequences for either heavy or light chain clustered into discrete groups. Interestingly, all PR A/B-specific monoclonal antibodies discovered in this report clustered closely together in the phylogenetic tree, most
likely due to clonal expansion from closely related B cells during immunization. Germline usage also supported this observation (Table 13). Similar observations were made in an independent experiment with a different antigen (Lin28A, Figure 12).

The methods used in the experiments described in this Example are as follows.
immunized by intradermal injection with four separate doses, each 3 weeks apart, with a mixture of keyhole limpet hemocyanin-conjugated peptides derived from the amino acid sequence of different regions of each human protein antigen. Peptides were conjugated to Imject maleimide-activated KLH (Thermo-Pierce). Mouse immunizations were carried out in the same manner, except the route of immunization was intraperitoneal and the injections were 2 weeks apart. Blood was drawn at 3 days after the final boost. Whole spleen from each animal was harvested at time of euthanasia following confirmation of desired polyclonal activity.

## Next Generation DNA sequencing of rabbit and mouse B cell repertoires.

 Splenocytes from hyperimmunized rabbits and mice were harvested and lysed for total RNA purification using Qiagen's RNeasy kit following the manufacturer's protocol. The RNA was on-column treated with DNase I (Qiagen cat\# 79254) to eliminate genomic DNA using the provided protocol. To generate heavy and light chain amplicon libraries from this material to be sequenced with 454 Life Sciences platform (Roche), RT-PCR was carried out as follows. cDNA was generated from the splenocyte total RNA as template using Thermoscript reverse transcriptase (Invitrogen cat\# 12236014) with oligo dT as primer. For rabbit IgG sequencing, variable regions of $\gamma, \kappa 1, \kappa 2$, and $\lambda$ chains were amplified with sequence specific 454 fusion primers (hybridizing to the leader on the $5^{\prime}$ end and containing sequences on the $3^{\prime}$ end required for identification and bar-coding in the Lib-L format of 454 sequencing platoform) using Phusion® Hot Start II High-Fidelity DNA Polymerase (Finnzymes Oy, Finland) with the following steps: denaturation- $98^{\circ} \mathrm{C}$ for 90 seconds; 20 cycles of [denaturation $-98^{\circ} \mathrm{C}$ for 10 seconds; annealing $-60^{\circ} \mathrm{C}$ for 30 seconds; extension$72^{\circ} \mathrm{C}$ for 30 seconds]. For mouse IgG sequencing, heavy and light chain amplicons were generated by a two-step PCR process. In the first step $\gamma$ or $\kappa$ chain variable regions were amplified ( 15 cycles with the same conditions as described above for rabbit) with a mixtureof gene family-specific degenerate oligonucleotides as sense primers, and anti-sense primers that hybridize to a highly conserved region at the start of the constant region, each sense and antisense primer containing distinct adaptor sequences at its 5' end. Each reaction from the first round was column-purified with a commercial kit (Qiagen cat\#28104) then further amplified by an additional 10 ( $\gamma$ chain) and 8 cycles ( $\kappa$ chain) in the second step using adaptor sequence-specific primers that contain sequences on the 3 ' end required for identification and bar-coding in the Lib-L format of 454 sequencing platform. For either species all light chain amplification reactions for each animal were pooled. Excess primers for heavy and light chain samples were eliminated using Agencourt AMPure XP DNA purification system following the provided protocol. The quality and purity of the amplicon pool after primer elimination was verified on Agilent Bioanalyzer 2100 (Agilent Technologies), and the concentration of the DNA was accurately quantified on a fluorometer using Quant-iT PicoGreen dsDNA Assay Kit (Invitrogen). Following the Lib-L LV, GS FLX Titanium Series protocol from 454 Life Sciences, emulsion PCR and bead enrichment was carried out. Bead number was counted on Beckman Coulter Z2 Particle Counter, and the library was sequenced on 454 GS Junior (Roche).

Affinity purification of antigen-specific IgG. Total IgG from the serum of the hyperimmunized rabbits (New Zealand white) was purified using Protein A sepharose beads (GE Healthcare), then was incubated rotating for 15 minutes in a column with the immunogen peptide covalently coupled to sepharose beads. By gravity flow, the unbound fraction was drained, and the column was washed extensively with 1x phosphate-buffered saline (PBS) to eliminate non-specific IgG. Antigen-specific polyclonal IgG pool was eluted sequentially with 0.1 M glycine $/ \mathrm{HCl}$ buffer at pH 3.5 , followed by pH 2.7 , and finally pH 1.8. Each elution was immediately neutralized with 1 M Tris buffer ( pH 8.5 ). Total $\operatorname{IgG}$ from the serum of the hyperimmunized mice was purified using Protein-G magnetic beads (Millipore, cat\# LSKMAGG10), then incubated rotating overnight at $4^{\circ} \mathrm{C}$ with immunogen peptide immobilized on magnetic beads (Pierce, cat\#88817). Using a magnetic tube rack (Invitrogen, cat\# 12321D) beads were extensively washed with PBS, then antibody bound to the column was sequentially eluted with progressively acidic pH as described for the rabbit IgG purification.

Protease Digestion of affinity-purified antibody. Polyclonal antibody was denatured in 8 M urea in 20mM HEPES pH 8 then reduced in 10 mM DTT for 1 hour at 55 ${ }^{\circ} \mathrm{C}$. Reduced polyclonal was cooled to room temperature (RT) and alkylation was performed in the presence of 20 mM iodoacetamide for 1 hour. Chymotrypsin, elastase, and overnight at $37^{\circ} \mathrm{C}$ at an enzyme to substrate ratio of $1: 50$. Pepsin digestion was performed in the presence of 3 M acetic acid at RT overnight at an enzyme to substrate ratio of 1:50. Digested peptides were desalted by STAGE-TIPS as published previously (Rappsilber et al., Anal Chem 75: 663-670, 2003), and analyzed by LC-MS/MS.

Mass spectrometry. LC-MS/MS was performed using the LTQ Orbitrap Velos (Thermo-Fisher) mass spectrometer. The samples were loaded for 7 min using a Famos autosampler (LC Packings) onto a hand-poured fused silica capillary column ( $125 \mu \mathrm{~m}$ internal diameter X 20 cm ) packed with Magic C18aQ resin ( $5 \mu \mathrm{~m}, 200 \AA$ ) using an Agilent 1100 series binary pump with an in-line flow splitter. Chromatography was developed using a binary gradient at $400 \mathrm{nl} / \mathrm{min}$ of $5-30 \%$ solvent B for 45 min (Solvent A, $0.25 \%$ formic acid (FA); Solvent B, $0.1 \%$ FA, $97 \%$ acetonitrile). Twenty MS/MS spectra were acquired in a data-dependent fashion from a preceding master spectrum in the Orbitrap ( $300-1,500 \mathrm{~m} / \mathrm{z}$ at a resolution setting of $6 \times 10^{4}$ ) with an automatic gain control (AGC) target of $10^{6}$. Charge-state screening was used to reject singly charged species, and a threshold of 500 counts was required to trigger an MS/MS spectrum. When possible, the LTQ and Orbitrap were operated in parallel processing mode.

Database searching and data processing. MS/MS spectra were searched using the SEQUEST algorithm (version 28 rev 12) (Yates et al., Anal Chem 67: 1426-1436, 1995) against a custom hybrid database composed of 21,932 full length gamma and 22,431 full length kappa $V$-region sequences and gamma and kappa constant region sequences concatenated to 6,358 yeast proteins ( $S$. cerevisiae, NCBI) and 42 common contaminants, including several human keratins, trypsin and chymotrypsin. Since V-region sequences are highly related, the yeast proteome artificially contributed more diverse sequences to the reference database (Beausoleil et al., Nat Biotechnol 24: 1285-1292, 2006) and provided another source of confidence after filtering the final dataset since filtered data should not
include peptides identified from yeast. Search parameters included partial specificity for chymotrypsin and trypsin and no specificity for elastase and pepsin, a mass tolerance of $\pm 50$ ppm, a static modification of 57.0214 on cysteine, and dynamic modification of 15.9949 on methionine. False discovery rate in the dataset was estimated using the target/decoy approach (Elias et al., Nat Methods 4: 207-214, 2007). Datasets were filtered to an FDR of $\leq 2 \%$ using a linear discriminant analysis (Huttlin et al., Cell 143: 1174-1189, 2010). Although the mass accuracy of the Orbitrap greatly exceeds 50 ppm , when searched with a wider precursor ion tolerance, correct peptide identifications result in small precursor mass errors ( $\pm 1 \mathrm{ppm}$ ), while incorrect peptide identifications distribute across the entire 50 ppm window. As a result, stringent precursor mass filters selectively remove many incorrect PSMs from the dataset.

Post acquisition analysis was performed as described in the text. Briefly, passing peptides derived from V-region sequences were re-mapped to the NGS Ig database. For peptides that arose from chymotryptic and tryptic digests, matches were limited to those arising from expected cleavages (KR for trypsin, YWFLMA for chymotrypsin). CDR coverage was determined by identifying CDRs using the rules defined by Kabat (Wu et al., $J$ Exp Med 132: 211-250, 1970). In all cases, coverage was defined as the total number of amino acids identified from high confidence peptides divided by the number of amino acids in the mature V-region sequence.

## Cloning, expression and characterization of identified immunoglobulin chains.

 $\gamma$ and $\kappa$ chains identified through the mass spectrometry analysis of the affinity-purified polyclonal IgG pool were cloned and expressed as follows. For each identified chain, the nucleic acid sequence encoding the entire variable domain from FWR1 through FWR4 were synthesized (Integrated DNA Technologies, Coralville Iowa). Using overlap PCR, each heavy-light chain combination permutation was expressed with a viral 2A sequence that uses a ribosomal skip mechanism to generate two polypeptides from a single open reading frame (Doronina et al., Mol Cell Biol 28: 4227-4239, 2008, Donnelly et al., J Gen Virol 82: 1027-1041, 2001). A single open reading frame cassette of, in order from $5^{\prime}$ to 3 ', light chain variable and constant regions, 2A peptide sequence from Thosea asigna virus, and heavy chain variable domain was cloned into a CMV-promoter driven mammalianexpression plasmid containing in-frame rabbit $\gamma$ chain leader sequence and rabbit $\gamma$ chain constant regions, $5^{\prime}$ and $3^{\prime}$ of the cloning site, respectively. HEK293 were transfected with plasmid preps encoding each light-heavy chain combination assembled in this manner using polyethylenimine (Boussif et al., Proc Natl Acad Sci USA 92: 7297-7301, 1995). The supernatant was screened 2 to 5 days post-transfection for secretion of antigen-specific antibody by ELISA using the immunogen peptide as the coating antigen, and light-heavy chain permutations that showed reactivity were further characterized. For mouse antibody expression, constant regions were of mouse $\operatorname{IgG} 2 \mathrm{a}$.

Characterization of polyclonal and monoclonal antibodies by ELISA, Western blotting, flow cytometry, immunofluorescence and immunohistochemistry. Detailed protocols of ELISA, Western blotting, flow cytometry, immunofluorescence and immunohistochemistry can be found online at the web site of Cell Signaling Technology Inc. Costar cat\#3369 certified high binding polystyrene 96 -well plates were used for ELISA. Antigens used for ELISA analysis for each target were the same peptides used for immunizations. For Progesterone Receptor antibodies, Western blotting was performed on T47D (PR+), MDA-MB-231 cells (PR-) and HT-1080 (PR-) cell lysate, flow cytometry analysis on T47D (PR+) and MDA-MB-231 cells (PR-), confocal immunofluorescence analysis on MCF-7 cells (PR+) compared with MDA-MB-231 cells (PR-), and immunohistochemical analysis on paraffin-embedded primary human breast carcinoma sections, T47D and paraffin-embedded MCF-7 cells (PR + ) compared with MDA-MB-231 cells (PR-). For phospho-p44/42 MAPK mouse antibodies, Western blotting was performed on lysate from Jurkat cells treated with either U1026 (Cell Signaling Technology, Inc. cat\# 9903) or 12-O-Tetradecanoylphorbol-13-Acetate (TPA) (Cell Signaling Technology, Inc. cat\# 4174). For Lin28A antibodies, Western blotting was performed on total lysate from NCCIT, NTERTA, MES and IGROV1 cell lines, confocal immunofluorescence and flow cytometry analyses on NTERA (Lin28A+) and HeLa (Lin28A-) cells. For phospho-Met ( pMet ) antibodies, lysates from MKN45 cells untreated ( $\mathrm{pMet}+$ ) and treated ( pMet ) with SU11274 Met kinase inhibitor were used. For Sox 1 antibodies, mouse brain extract (Sox1+) and lysate from NIH-3T3 (Sox 1-) cells were used.

## Example 8

In this Example, human monoclonal antibodies specific for the Hepatitis B virus small surface antigen ( HBsAg ) were generated in accordance with the methods described herein. To do this, the following protocol was followed to generate the genetic material database. Polyclonal antibodies were purified as described below and were analyzed following the mass spectrometry analysis as described above for mouse and rabbit.

## I. Generation of the Nucleic Acid Sequences.

## Antigen-specific, memory and total B-cell isolation and RNA purification

Peripheral blood mononuclear cells (PBMC) were isolated from fresh, whole human blood collected in heparin vacuum tubes, using a Ficoll gradient. In a Greiner Leucocep 50 ml conical tube (Sigma Aldrich cat\# Z642843) containing 20ml of Histopaque 1077 (Sigma Aldrich cat\#10771) at the bottom, up to 25 ml of blood was applied on top, then the tubes were centrifuged for 20 minutes at 1500 xg at room temperature. The leukocytes (buffy coat) were collected using a sterile pipette, washed in RPMI medium twice by resupending the cells in 50 ml of RPMI, then centrifuged at 300 xg for 10 minutes at $4^{\circ} \mathrm{C}$. After the washes, the PBMC were either cryopreserved in $20 \%$ DMSO in fetal bovine serum or processed immediately for B -cell isolation.

For B-cell isolation, a negative selection method was used to eliminate all non-Bcells from the PBMC using Invitrogen's Dynabeads Untouched B-cell Isolation kit (Invitrogen cat\#113-51D) following the manufacturer's protocol. The resulting unlabeled Bcell population was further processed to isolate either antigen-specific or memory B-cells.

For antigen-specific B-cell isolation, total unlabeled B-cells were incubated with biotinylated antigen that is immobilized on streptavidin magnetic beads (Pierce-Thermo Scientific cat\#88816) on a rotator at room temperature for 20 minutes. The beads containing any antigen-binding B-cells were then washed twice with $1 x P B S$. The washed beads were then resuspended in Qiagen's RNeasy kit RLT lysis buffer (supplemented with $1 \% \beta$ mercaptoethanol) for RNA isolation.

unlabeled B-cells using Miltenyi's MACS kits for $\mathrm{CD}^{2} 7^{+}$and surface $\mathrm{IgG}^{+}$cell isolation (Miltenyi Biotec (Auburn, CA) cat\#130-051-601 and 130-047-501). In order to simultaneously isolate $\mathrm{CD} 27^{+}$and $\mathrm{sIgG}{ }^{+}$B-cells, magnetic bead-conjugated antibodies to both cell surface markers were added at the same time during the incubation step. Upon purification, memory B-cells were spun down at 300xg for 10 minutes, and then lysed in RLT buffer for RNA as described above for RNA isolation.

RNA was purified from selected cells using Qiagen's RNeasy kit (Qiagen cat\# 74104) following the manufacturer's protocol. On column DNase I-treatment was conducted to eliminate contaminating genomic DNA by incorporating a DNase I digest step. After the RW1 buffer wash, DNase I (Qiagen cat\# 79254) diluted in RDD buffer was applied to the RNA purification column and incubated for 20 minutes at room temperature. The column was then washed once more with RW1 buffer, followed by two washes with RPE buffer, and the RNA was eluted with either 30 or $50 \mu$ l water. The concentration of the RNA was determined by absorbance measured on a Nanodrop spectrophotometer (Thermo Scientific) at wavelength 450 nm .

## cDNA synthesis and generation of amplicons by PCR

RNA isolated from memory or antigen-specific B-cells was first reverse transcribed using Invitrogen's Thermoscript reverse transcriptase (Invitrogen cat\#12236-022) as shown below:

| DNase treated RNA: | 5 uL |
| :--- | :--- |
| Oligo dT primer $(50 \mathrm{uM}):$ | 1 uL |
| dNTP's $(10 \mathrm{mM}):$ | 2 uL |
| dI H2O: | 4 uL |

Incubated at $65^{\circ} \mathrm{C}$ for 5 min , placed on ice for 2 minutes, then added the following:

| 5X cDNA buffer: | 4 uL |
| :--- | :--- |
| 0.1mM DTT: | 1 uL |
| RNAse OUT: | 1 uL |
| dI H2O: | 1 uL |
| ThermoScript: | 1 uL |

The mixture was incubated at $50^{\circ} \mathrm{C}$ for 1 hour, followed by a heat-inactivation step at $85^{\circ} \mathrm{C}$ for 5 minutes. Finally, the complementary RNA strand was eliminated from the cDNA
by adding $1 \mu \mathrm{l}$ of RNase H (Invitrogen cat\#18021-071) and incubating at $37^{\circ} \mathrm{C}$ for 20 minutes.

Amplicons of heavy, kappa and lambda chain variable regions for sequencing were generated by PCR as follows. For amplification of heavy chain, 4 independent reactions (each one specific to gene families of $\mathrm{V}_{\mathrm{H}} 1$ and $7 ; \mathrm{V}_{\mathrm{H}} 2,5$ and $6 ; \mathrm{V}_{\mathrm{H}} 3$; and $\mathrm{V}_{\mathrm{H}} 4$ ) were run for each cDNA sample using the below listed primers in order to preserve the natural distribution of $V_{H}$ gene transcript frequency in the pool of B-cells. For kappa and lambda chain amplification, single reaction for each chain was run for each cDNA sample. For each reaction, an equimolar mixture of forward primers was used with the same concentration of reverse primer(s) as indicated below. Amplification was performed with fusion primers compatible for 454 Sequencing (Roche) by the Lib-L platform. Reverse primers were designed to hybridize to the 5 ' end of the constant region of each chain. These primers contain the Lib-L primer B and MID sequences so that sequencing reads would begin from the extreme $5^{\prime}$ end of each constant region (in reverse sense) and into the 3 ' end of the variable region. For heavy and kappa chains, a single reverse primer was used for each MID, whereas for lambda chain, two distinct reverse primers were required for each MID.

## Heavy Chain Fusion Primers:

Reverse

| Reverse | CCA TCT CAT CCC TGC GTG TCT CCG AC tcag CTGTGCGTCGCA gaa gac <br> Sga TGG GCC CTT GGT GGA (SEQ ID NO: 248) | MID136 |
| :--- | :--- | :--- |
| oli551 | CCA TCT CAT CCC TGC GTG TCT CCG AC tcag ACGCGAGTAT gaa gac Sga <br> TGG GCC CTT GGT GGA (SEQ ID NO: 249) | MID27 |
| oli602 | CCA TCT CAT CCC TGC GTG TCT CCG AC tcag CACGCTACGT gaa gac Sga <br> tgg gcc ctt ggt gga (SEQ ID NO: 250) | MID34 |
| oli606 | CCA TCT CAT CCC TGC GTG TCT CCG AC tcag TGAGTCAGTAT gaa gac sga <br> tgg gcc ctt ggt gga (SEQ ID NO: 251) | MID70 |
| oli670 | CCA TCT CAT CCC TGC GTG TCT CCG AC tcag ATCTACTGACatgat gaa gac <br> Sga tgg gcc ctt ggt gga (SEQ ID NO: 252) | MID88 |
| oli671 | CCA TCT CAT CCC TGC GTG TCT CCG AC tcag AGTAGTGATCtcaca gaa gac <br> Sga tgg gcc ctt ggt gga (SEQ ID NO: 253) | MID83 |

Forward
VH1/7

| oli621 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GAC TGG ACC TGG AGV ATC <br> (SEQ ID NO: 254) |
| :--- | :--- |
| oli622 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GAC TGG ATT TGG AGG RTC <br> (SEQ ID NO: 255) |
| oli623 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GAC TGC ACC TGG AGG ATC <br> (SEQ ID NO: 256) |


| oli624 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GAC TGG ACC TGG AGG KTC <br> (SEQ ID NO: 257) |
| :--- | :--- |

VH2/5/6

| oli618 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GAC ATA CTT TGT TCC ACG C <br> (SEQ ID NO: 258) |
| :--- | :--- |
| oli619 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GAC ACA CTT TGC TAC ACA C <br> (SEQ ID NO: 259) |
| oli620 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG TCT GTC TCC TTC CTC ATC T <br> (SEQ ID NO: 260) |
| oli629 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GGG TCA ACC GCC ATC CTC <br> (SEQ ID NO: 261) |

VH3

| oli625 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GAG TTK GGR CTG AGC TGG <br> (SEQ ID NO: 262) |
| :--- | :--- |
| oli626 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GAG TTT KGG CTK AGC TGG <br> (SEQ ID NO: 263) |
| oli627 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GAA CTG GGG CTC CGC TGG <br> (SEQ ID NO: 264) |
| oli628 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GAR TTG GGG CTG WGC TGG <br> (SEQ ID NO: 265) |
| CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG AAR CAY CTG TGG TTC TTC CT <br> (SEQ ID NO: 266)  |  |$.$| oli617 |
| :--- |

Kappa Chain Fusion Primers
Reverse

| oli552 | CCA TCT CAT CCC TGC GTG TCT CCG AC tcag ACGACAGCTC gaa gat gaa <br> gac aga tgg tgc agc cac (SEQ ID NO: 267) | MID77 |
| :--- | :--- | :--- |
| oli556 | CCA TCT CAT CCC TGC GTG TCT CCG AC tcag TCGATCACGT gaa gat gaa <br> gac aga tgg tgc agc cac (SEQ ID NO: 268) | MID42 |
| oli603 | CCA TCT CAT CCC TGC GTG TCT CCG AC tcag TACACACACT GAA GAT <br> GAA GAC AGA TGG TGC AGC cac (SEQ ID NO: 269) |  |
| oli607 | CCA TCT CAT CCC TGC GTG TCT CCG AC tcag TGTAGTGTGAT GAA GAT <br> GAA GAC AGA TGG TGC AGC cac (SEQ ID NO: 270) | MID37 |


| Forward | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GAC ATG AGG GTS CCY GCT CAG <br> oli630 <br> CTC (SEQ ID NO: 271) |
| :--- | :--- |
| oli631 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GAC ATG AGR GTC CTC GCT CAG <br> CTC (SEQ ID NO: 272) |
| oli632 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GAA GCC CCA GCD CAG CTT CTC <br> (SEQ ID NO: 273) |
| oli633 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GAA ACC CCA GCG CAG CTT CTC <br> (SEQ ID NO: 274) |
| oli634 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GTG TTG CAG ACC CAG GTC TTC <br> (SEQ ID NO: 275) |
| oli635 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GGG TCC CAG GTT CAC CTC CTC <br> (SEQ ID NO: 276) |
| oli636 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG AGG CTC CYT GCT CAG CTC CTG <br> (SEQ ID NO: 277) |


| Lambda Chain Fusion Primers Reverse |  |  |
| :---: | :---: | :---: |
| oli604 | CCA TCT CAT CCC TGC GTG TCT CCG AC tcag CGTAGACTAG AGG GCG GGA ACA GAG TGA CMG TGG (SEQ ID NO: 278) | MID21 |
| oli605 | CCA TCT CAT CCC TGC GTG TCT CCG AC tcag CGTAGACTAG AGG GYG GGA ACA GAG TGA CCG AKG (SEQ ID NO: 279) | MID21 |
| oli608 | CCA TCT CAT CCC TGC GTG TCT CCG AC tcag TCTAGCGACTAT AGG GCG GGA ACA GAG TGA CMG TGG (SEQ ID NO: 280) | MID45 |
| oli609 | CCA TCT CAT CCC TGC GTG TCT CCG AC tcag TCTAGCGACTAT AGG GYG GGA ACA GAG TGA CCG AKG (SEQ ID NO: 281) | MID45 |
| oli553 | CCA TCT CAT CCC TGC GTG TCT CCG AC tcag TAGCGCGCGCT agg gcg gga aca gag tga $\mathrm{cMg} \operatorname{tgg}$ (SEQ ID NO: 282) | MID101 |
| oli554 | CCA TCT CAT CCC TGC GTG TCT CCG AC tcag TAGCGCGCGCT agg gYg gga aca gag tga ccg aKg (SEQ ID NO: 283) | MID101 |
| oli557 | CCA TCT CAT CCC TGC GTG TCT CCG AC tcag ATAGAGTACT agg gcg gga aca gag tga $\mathrm{cMg} \operatorname{tgg}$ (SEQ ID NO: 284) | MID33 |
| oli558 | CCA TCT CAT CCC TGC GTG TCT CCG AC tcag ATAGAGTACT agg $g \mathrm{Yg}$ gga aca gag tga ccg aKg (SEQ ID NO: 285) | MID33 |
| Forward |  |  |
| oli637 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG ACC TGC TCC CCT CTC CTC CTC A(SEQ ID NO: 286) |  |
| oli638 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GCC GGC TTC CCT CTC CTC CTC A (SEQ ID NO: 287) |  |
| oli639 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GCC TGG TCT CCT CTC CTC CTC A (SEQ ID NO: 288) |  |
| oli640 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GCC TGG ACY CCT CTC CTC CTC M (SEQ ID NO: 289) |  |
| oli641 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG CCC TGG GCT CTG CTS CTC CTS A (SEQ ID NO: 290) |  |
| oli642 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG CCC TGG GTC ATG CTC CTC CTG A (SEQ ID NO: 291) |  |
| oli643 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GCC TGG ACT CCT CTC TTT CTG T (SEQ ID NO: 292) |  |
| oli644 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GAG AAG AAG AGG AGA CCT GGG G (SEQ ID NO: 293) |  |
| oli645 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GCC TGG ACC GCT CTC CTT CTG A (SEQ ID NO: 294) |  |
| oli646 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GCC TGG ACC GTT CTC CTC CTC G (SEQ ID NO: 295) |  |
| oli647 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GCA TGG ATC CCT CTC TTC CTC G (SEQ ID NO: 296) |  |
| oli648 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GCC TGG ATC CCT CTA CTT CTC C (SEQ ID NO: 297) |  |
| oli649 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GCC TGG AYC CCT CTC CTG CTC C <br> (SEQ ID NO: 298) |  |
| oli650 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GCA TGG GCC ACA CTC CTG CTC C (SEQ ID NO: 299) |  |
| oli651 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GCC TGG ACC CCT CTC TGG CTC A (SEQ ID NO: 300) |  |
| oli652 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GCC TGG GTC TCC TTC TAC CTA C (SEQ ID NO: 301) |  |
| oli653 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GCC TGG ACC CCA CTC CTC CTC C (SEQ ID NO: 302) |  |


| oli654 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GCC TGG GCT CCT CTG CTC CTC A <br> (SEQ ID NO: 303) |
| :--- | :--- |
| oli655 | CCT ATC CCC TGT GTG CCT TGG CAG TC tcag ATG GCC TGG GCT CCA CTA CTT CTC A <br> (SEQ ID NO: 304) |

PCR amplification was done using Finnzyme's Phusion Hot Start II polymerase (Thermo Scientific cat\# F-540S) where the reaction mix and conditions were set up as

## Reaction mixture:

cDNA: $\quad 2.5 \mathrm{uL}$
5X Buffer GC: 5 uL
10 mM dNTP mix:
0.25 uL

Phusion HotStart II:
0.25 uL

Primers (forward+reverse) $30 \mathrm{uM}: \quad 0.25 \mathrm{uL}$
Water:
16.75 uL

PCR program:
Step $198^{\circ} \mathrm{C}-2$ minutes
Step $298^{\circ} \mathrm{C}-10$ seconds
Step $360^{\circ} \mathrm{C}-30$ seconds
Step $472^{\circ} \mathrm{C}-30$ seconds
Step 5 Repeat steps 2 through 4
Step $672^{\circ} \mathrm{C}-2$ minutes
Step 7 - hold
For heavy chain amplification, 25 or 30 cycles (step 5 repeated either 24 or 29 times), and for kappa and lambda chains, 20 or 30 cycles were run when amplifying cDNA template generated from either memory B-cells or from antigen-specific B-cells, respectively, as 5 extra cycles were required for sufficient amplification from antigenspecific B-cell cDNA for each chain. To ensure the absence of any false amplification from contaminating template in any of the reagents, duplicate reactions were set up for each mixture ( 4 separate reactions for heavy chain, and one for each light chain) where the cDNA template was substituted with water. These negative control reactions with no template were run at the same time as the samples containing template. Upon completion of the PCR program, $3 \mu \mathrm{l}$ of each reaction (including the negative controls) were analyzed by electrophoresis on a $1.5 \%$ TAE agarose gel for the presence of the amplicons
(approximately 540bp for heavy chain, approximately 485bp for kappa chain and approximately 510bp for lambda chain) when template was added to the reaction but not in the absence of cDNA.

## Amplicon purification, analysis, quantitation, and preparation for 454 sequencing

In order to eliminate excess primers and/or primer dimmers in the PCR samples, amplicons were purified using Agentcourt Ampure magnetic beads (Beckman Coulter cat\#A63881) following the manufacturer's protocol (000387v001). For heavy chain, all four reactions (VH1/7, VH2/5/6, VH3, VH4) were pooled and purified as one sample, thus a total of 3 amplicon samples (heavy, kappa and lambda chains) were purified for each cDNA amplification. The protocol for ampure purification was modified in that purifications were done in single 1.5 ml microtubes using a generic magnetic rack that is suitable for 1.5 ml tubes instead of in a 96 -well plate format. All volumes and other procedures were as described in the protocol. The eluted amplicons after Ampure purification were then analyzed for purity and absence of any contaminating DNA species on the Agilent 2100 Bioanalyzer using the high sensitivity DNA chip (Agilent Technologies cat\# 5067-4626) by following the manufacturer's protocol.

Once the purity of amplicons was verified, the concentration of the DNA was quantified on a fluorometer using the Quant-iT PicoGreen dsDNA Assay Kit (Invitrogen cat\#P7589) as described in the manufacturer's protocol. The Lambda DNA provided in the kit was used as a concentration standard with which a standard curve was generated from $100 \mathrm{ng} /$ well to $1.56 \mathrm{ng} /$ well. The fluorescence of each amplicon diluted 100 -fold in TE buffer was measured in duplicate, and the concentration of DNA was determined according to the linear portion of the standard curve. All fluorescence measurements were done in black 96well plates. If the value of fluorescence was out of the linear range of the standard curve, the samples were remeasured with either larger or smaller dilutions in order to capture fluorescence values that fall within the linear range. Using the approximate size in base pairs of each chain type (heavy-540bp, kappa-485bp and lambda-510bp), the following formula was used to determine the concentration: Concentration of each amplicon (molecules $/ \mu \mathrm{l})=$ [sample conc $\left.(\mathrm{ng} / \mu \mathrm{l}) * 6.022 \times 10^{23}\right] /\left[656.6 \times 10^{9} *\right.$ amplicon length (bp)]

Each amplicon was normalized to $1 \times 10^{7}$ molecules $/ \mu$, then mixed at a ratio of $\mathrm{Hc}: \mathrm{Kc}:$ Lc at $3: 3: 1$ by volume, vortexed, and finally diluted $1: 10$ to obtain a final concentration of the mixture at $1 \times 10^{6}$ molecules $/ \mu \mathrm{I}$.

## Emulsion PCR amplification, bead enrichment, bead counting and sequencing

Emulsion PCR was conducted following the 454 published protocol: "emPCR
Amplification Method Manual - Lib-L" (Edition: May 2010 (Rev. April 2011)) with the following modifications:

10 Section 3.1.3 Step 2)

| Reagent | Volume $(\mu \mathrm{l})$ |
| :--- | :--- |
| Mol. Bio. Grade water | 458 |
| Additive | 515 |
| Amp Mix | 270 |
| Amp Primer | 32 |
| Enzyme Mix | 70 |
| PPiase | 2 |
|  |  |
| Total | 1347 |

Once the sequencing beads were enriched, from step 3.7 of "emPCR Amplification 15 Method Manual - Lib-L", the beads were counted on Beckman Coulter's Z2 Particle Counter with the following settings:

Aperture: $\quad<100 \mu \mathrm{~m}>$
Aperture Kd: $<60.04>$
Set Upper cutoff: $<30.00 \mu \mathrm{~m}>$

Metered Volume: $<0.5 \mathrm{ml}>$
Resolution: <256>

The concentration of beads was calculated as:
Concentration of beads $=[$ Avg. reading from particle counter * 4] beads $/ \mu \mathrm{l}$
The enriched beads from the emulsion PCR were sequenced on the 454 Sequencer
(Roche) following the 454 sequencing protocol: "Sequencing Method Manual - GS Junior Titanium Series" - May 2010 (Rev. June 2010), herein incorporated by reference in its entirety.

## Purification Of Antigen-Specific IgG From Human Donor Plasma

Donor plasma isolation and screening for reactivity to specific antigens.
Whole blood from human volunteers was collected following IRB guidelines in heparin tubes. During ficoll-gradient separation of PBMC (as described above), plasma samples were collected simultaneously and stored at $-80^{\circ} \mathrm{C}$. Reactivity of plasma $\operatorname{IgG}$ to various antigens was tested by ELISA. Briefly, high-binding 96-well plates (Costar cat\#) were coated $100 \mu \mathrm{l} /$ well of antigen at $2 \mu \mathrm{~g} / \mathrm{ml}$ dissolved in carbonate buffer at $37^{\circ} \mathrm{C}$ for two hours or $4^{\circ} \mathrm{C}$ overnight. The plates were rinsed three times with PBS-Tween $(0.1 \%)$, then blocked with $300 \mu \mathrm{l} /$ well of $5 \%$ non-fat dry milk in PBS-Tween at $37^{\circ} \mathrm{C}$ for 1 hour. Plasma samples were diluted at $1 / 100,1 / 500$ and $1 / 1000$ and $1 / 2000$ in $5 \%$ milk PBS-Tween, and $100 \mu \mathrm{l}$ of each dilution was added in duplicates of blocked wells of the 96 -well plate and incubated for 2 hours at $37^{\circ} \mathrm{C}$. The plates were washed 3 times with $1 x$ PBS-Tween, and horseradish peroxidase-conjugated anti-human IgG antibody (Southern Biotech 2040-05) diluted $1 / 4000$ in PBS-Tween was added to each well $(100 \mu \mathrm{l})$ and incubated at $37^{\circ} \mathrm{C}$ for one hour. The plates were washed 6 times with PBS-Tween and developed by addition of $50 \mu 1$ TMB substrate solution (BioFX cat\#TMBW-1000-01), followed by $50 \mu \mathrm{l}$ of stop solution (BioFX cat\# STPR1000-01). The signals were measured at optical density of 450 nm . Donors whose plasma showed significant signal at $1 / 500$ or greater dilution were selected for screening by NG-XMT procedure.

Hepatitis B virus small surface antigen (HBsAg) adw subtype was purchased from Prospec (Rehovot, Israel, cat\# HBS-872).

## Purification of antigen-specific IgG from total plasma IgG

## Protein G purification

1. 5 ml of bead slurry ( 2.5 ml bead bed volume) of Protein G Sepharose 4 Fast Flow (GE

Healthcare cat\#17-0618-05) were applied to a gravity flow column and washed with 1xPBS twice.
2. 5 ml of human plasma diluted with 1 xPBS to 15 ml was applied to the column with beads, and the column was incubated on a rotator overnight at 4 C , or room temperature for 2 hours.
3. The column was washed 4 times with 20 ml of 1 xPBS .
4. IgG was eluted with 20 ml of pH 2.70 .1 M glycine $/ \mathrm{HCl}$ buffer and collected in a tube containing 1.2 ml of 1 M Tris pH 8.5 for neutralization.
5. 10 ml of $1 \times \mathrm{xPBS}(\mathrm{pH} 7.4)$ was added to the neutralized eluate to minimize precipitation due to high concentration of IgG.
6. Purified IgG was dialyzed twice against 4 liters of 1 xPBS in a 10 kDa cut-off dialysis cassette (Pierce cat\#66456).
7. IgG concentration was determined by measuring the absorbance at 280 nm on a Nanodrop photospectrometer (Thermo Scientific).

## Affinity purification

1. HBsAg was conjugated with biotin (Pierce Cat \#20217) following the manufacturer's protocols. The conjugated antigen was dialyzed extensively in $1 \times \mathrm{PBS}$.
2. 2 mg of biotin-conjugated antigen was incubated with 5 ml of magnetic streptavidin beads (Thermo Scientific cat\#8816) overnight at $4^{\circ} \mathrm{C}$ or for two hours at room temperature on a rotator. The beads were rinsed with 1xPBS twice, then divided into nine 1.5 ml tubes.
3. The efficiency of immobilization of antigen to beads was evaluated by HBsAg Elisa and consistently showed greater than $80 \%$ binding.
4. To each tube containing immobilized antigen, 1 mg of protein G-purified IgG from a single donor was added, the beads were resuspended fully by vortexing and incubated rotating at room temperature for 15 minutes.
5. The tubes were placed in a magnetic rack, the supernatant was removed, and the beads were washed 5 times with 1 ml 1 xPBS.
6. After the last wash step, 0.9 ml of 0.1 M glycine -HCl buffer at pH 1.8 was applied to one tube, vortexed and incubated at room temperature for 5 minutes. After 5 minutes, the
first tube was placed on the magnetic rack, then the acidic buffer in the tube was removed and placed into a second tube. This procedure was repeated until all nine tubes were incubated with the acidic buffer. Eluted IgG was finally collected in a tube containing 0.14 ml of 1 M Tris pH 8.5 for neutralization. restarting the purification from the step where 1 mg of protein G-purified IgG was added to the beads. The procedure was repeated multiple times to generate sufficient material for protease treatment prior to MS analysis.

## III. Mass Spectrometry

Mass spectrometry analysis was performed as described above. Briefly, following digestion with a protease (e.g., trypsin) and/or a chemical protein cleavage reagent (e.g., cyanogen bromide), mass spectrometry analysis was performed on the peptides. The resulting MS2 spectra was correlated to the theoretical MS2 spectra derived from the information in the genetic material database, in order to identify the genetic sequences that encode antibodies that specifically bind to the Hepatitis B virus small surface antigen.

## IV. Expression and Identification of Monoclonal Antibodies

24 distinct heavy (gamma) chain variable region clones, 20 distinct kappa chain variable region clones and 10 distinct lambda chain variable region clones were expressed in a combinatorial format and screened for antigen-specific binding activity (See Tables 14-15, where gamma chain clones are indicated in the left most vertical column and light chain clones are indicated in the top horizontal row). Each gamma chain was paired with every light (kappa and lambda) chain to express antibodies by transient transfection of HEK293E cells in standard 96 -well tissue culture plates.

Antibody that was secreted from the transfected cells in each well was screened for binding to purified, recombinant hepatitis B surface antigen (HBsAg-adw subtype purchased from Prospec, Ness-Ziona, ISRAEL) by enzyme-linked immunosorbant assay (ELISA). High binding 96-well ELISA plates (Costar-3369) were coated with $50 \mu \mathrm{l} /$ well of HBsAg diluted in carbonate buffer at $2 \mu \mathrm{~g} / \mathrm{ml}$ by incubating at $37^{\circ} \mathrm{C}$ for two hours then blocked with
$300 \mu \mathrm{l} /$ well of $5 \%$ powdered milk in phosphate-buffered saline (PBS) by incubating at $37^{\circ} \mathrm{C}$ for one hour. The supernatant from the transiently transfected HEK293E cells were diluted five-fold in $5 \%$ milk in PBS with $0.05 \%$ Tween 20 (PBS-T), then $50 \mu \mathrm{l}$ of the diluted supernatant was applied to each well of HBsAg-coated ELISA plates. To assess nonspecific binding, the same supernatant was also applied to plates coated only with $5 \%$ milk in PBS. After addition of supernatant, the ELISA plates were incubated at $37^{\circ} \mathrm{C}$ for 2 hours followed by 3 washes with $250 \mu \mathrm{l} /$ well of PBS-T. To detect any binding of antibody, $50 \mu \mathrm{l} /$ well of horse radish peroxidase (HRP)-conjugated anti-human $\operatorname{IgG}$ (Southern Biotech) diluted 4000 -fold in PBS-T was added to each well and incubated at $37^{\circ} \mathrm{C}$ for one hour. The plates were washed 6 times as described above, and then $50 \mu 1 /$ well of a chromogenic substrate for HRP, 3,3',5,5'-Tetramethylbenzidine, was added, which was neutralized with $50 \mu \mathrm{l} /$ well of acid approximately 10 minutes later. The signal from the chromogenic substrate neutralized with acid was measured by absorbance (optical density) at 450 nm .

Tables 14-15 show the values obtained from the absorbance of HBsAg plates from which the absorbance of the milk only plates in each well was subtracted. The following supernatant samples were used as controls (values are averages of two independent wells in each case): positive = supernatant from transfection of anti-HBsAg human antibody heavy and light chain; negative $=$ supernatant from cells transfected with PEI only. Wells with signal greater than the negative control signal by 10-20-fold, 20-40-fold and higher than 40fold are indicated in increasing shades of grey. 30 heavy-light permutations exhibited strong reactivity to HBsAg , greater than 40 -fold over background in two or more out of four wells, 26 were between 20 to 40 -fold over background, and 18 were between 10 to 20 -fold over background (one of the 18, expressed as a combination of EVUGG gamma chain, shown with * on the table, and AKUOL lambda chain was later found to be non-reactive). Thus, out of 24 distinct variable region gamma chain clones tested, 17 expressed HBsAg-specific antibody when paired with at least one of the 30 light chain clones tested.

Table 14
Kappa Chain

|  |  | GRAWH |  | CHUZK\% |  | EDLIN |  | 1P54A |  | B04 58 |  | HMPSS |  | 1018 X |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| HSP82, |  | 2.86 | 2.94 | 2.72 | -0.04 | -0.04 | -0.06 | -0.04 | 0.01 | 2.58 | 2.80 | 1.89 | 2.60 | 0.04 | 0.20 |
|  |  | 2.59 | 2.71 | 0.68 | 0.00 | 0.09 | 0.04 | -0.03 | 0.08 | 2.62 | 2.63 | 2.38 | 0.62 | 0.05 | 0.13 |
|  | BIRWB | 0.92 | 0.41 | 0.06 | 0.03 | 2.89 | 3.01 | 0.06 | 0.16 | 0.71 | 0.22 | 0.22 | -0.01 | 0.05 | 0.06 |
|  |  | 0.07 | 0.05 | 0.03 | 0.01 | 2.83 | 2.85 | 0.08 | 0.09 | 0.04 | 0.02 | 0.00 | 0.33 | 0.03 | 0.05 |
|  | 80001. | 0.20 | 0.09 | 1.84 | 1.97 | 0.77 | 0.82 . | 1.85 | 2.05 | 0.05 | 0.00 | 1.52 | 1.92 | 1.05 | 1.38 |
|  |  | 0.18 | 0.15 | 1.89 | 2.11 | 0.85 | 0.83 | 1.91 | 2.16 | 0.06 | 0.03 | 1.72 | 1.87 | 1.06 | 1.43 |
|  | H20\% | 0.01 | -0.01 | 0.03 | 0.05 | 0.06 | 0.05 | 0.03 | -0.02 | 0.03 | 0.02 | 0.02 | 0.00 | 0.14 | 0.10 |
|  |  | 0.02 | -0.01 | -0.02 | 0.04 | 0.05 | 0.10 | 0.06 | 0.09 | -0.03 | 0.00 | 0.05 | 0.06 | 0.09 | 0.06 |
|  | CP4Y8, | 0.02 | 0.02 | 0.02 | 0.02 | 0.05 | 0.04 | 0.04 | 0.03 | 0.05 | 0.02 | 0.03 | 0.00 | 2.68 | 2.75 |
|  |  | 0.04 | 0.06 | 0.04 | 0.04 | 0.06 | 0.06 | 0.05 | 0.07 | 0.04 | 0.04 | 0.02 | 0.03 | 2.71 | 2.60 |
|  | EBOGO | 0.06 | 0.04 | 0.06 | 0.04 | 0.05 | 0.03 | 0.06 | 0.05 | 0.06 | 0.05 | 0.04 | 0.05 | 0.08 | 0.08 |
|  |  | 0.13 | 0.06 | 0.04 | 0.07 | 0.05 | 0.08 | 0.08 | 0.06 | 0.05 | 0.05 | 0.06 | 0.04 | 0.03 | 0.17 |
|  | 13ESP | 0.15 | 0.05 | 0.10 | 0.12 | 0.04 | 0.10 | 0.07 | 0.15 | 0.06 | 0.07 | 0.14 | 0.08 | 0.05 | 0.10 |
|  |  | 0.05 | 0.05 | 0.05 | 0.06 | 0.08 | 0.05 | 0.08 | 0.06 | 0.05 | 0.05 | 0.05 | 0.05 | 0.14 | 0.05 |
|  | APABY, | -0.06 | 0.04 | 0.03 | 0.05 | 0.03 | 0.03 | 0.04 | 0.04 | 0.04 | 0.04 | 0.10 | 0.03 | 0.05 | 0.00 |
|  |  | 0.05 | 0.08 | 0.09 | 0.04 | 0.08 | 0.05 | 0.04 | 0.04 | 0.05 | 0.05 | 0.07 | 0.13 | 0.04 | 0.04 |
|  | 19136\% | 0.75 | 0.71 | 2.23 | 2.59 | 0.97 | 1.27 | 2.55 | 2.63 | 0.67 | 0.74 | 2.57 | 2.63 | 0.21 | -0.32 |
|  |  | 0.50 | 0.95 | 2.46 | 2.48 . | 1.50 | 1.53 | 2.64 | 2,66 | 1.32 | 1.31 | 2.57 | 2.64 | 0.21 | -0.42 |
|  | EZQNT | 0.06 | 0.05 | 0.18 | 0.13 | 0.84 | 0.72 | 0.17 | 0.31 | 2.08 | 1.76 | 0.09 | 0.09 | -0.38 | -0.66 |
| 들 |  | 0.28 | 0.07 | 0.13 | 0.10 | 0.80 | 1.05 | 0.14 | 0.18 | 2.06 | 1.97 | 0.17 | 0.18 | -0.69 | -0.56 |
| \% | A9F90 | 2.60 | 2.47 | 0.23 | 0.10 | 0.05 | 0.07 | 0.11 | 0.25 | 3.01. | 2.90 | 0.08 | 0.17 | -0.46 | -0.69 |
| E |  | 2.57 | 2.62 | 0.15 | 0.05 | 0.06 | 0.04 | 0.06 | 0.29 | 2.72 | 2.79 | 0.06 | 0.13 | -0.42 | -0.80 |
| 0 | IXGB9, | 0.11 | 0.24 | 0.05 | 0.05 | 0.03 | 0.07 | 0.06 | 0.05 | 0.24 | 0.43 | 0.06 | 0.01 | -1.11 | -0.99 |
|  |  | -0.04 | 0.03 | 0.01 | 0.02 | 0.02 | 0.03 | 0.03 | 0.03 | 0.06 | 0.05 | 0.06 | 0.03 | -0.25 | -0.80 |
| E | 103139 | -0.17 | 0.02 | 0.02 | 0.01 | 0.02 | 0.02 | 0.02 | 0.02 | 0.01 | 0.00 | 0.01 | -0.02 | -0.17 | -0.01 |
|  |  | 0.01 | 0.03 | 0.04 | 0.02 | 0.05 | 0.03 | 0.02 | 0.03 | 0.03 | 0.02 | 0.00 | 0.00 | -0.07 | 0.01 |
| E | A6TR9 | 0.01 | 0.03 | 0.03 | 0.03 | 0.04 | 0.02 | 0.03 | 0.04 | 0.04 | 0.05 | 0.02 | 0.05 | -0.02 | 0.02 |
| 1 |  | 0.03 | 0.02 | 0.02 | 0.03 | 0.05 | 0.03 | 0.03 | 0.04 | 0.03 | 0.04 | 0.02 | 0.01 | 0.01 | 0.02 |
| 1 | EO222 | 0.05 | 0.04 | 0.03 | 0.03 | 0.05 | 0.04 | 0.04 | 0.05 | 0.06 | 0.04 | 0.05 | 0.01 | 0.01 | 0.03 |
|  |  | 0.06 | 0.06 | 0.04 | 0.04 | 0.05 | 0.06 | 0.05 | 0.04 | 0.03 | 0.05 | 0.05 | 0.03 | -0.04 | 0.03 |
|  | ASNDH: | 0.05 | 0.06 | 0.04 | 0.04 | 0.04 | 0.05 | 0.05 | 0.05 | 0.05 | 0.03 | 0.05 | 0.02 | 0.00 | 0.02 |
|  |  | 0.15 | 0.04 | 0.03 | 0.06 | 0.05 | 0.14 | 0.03 | 0.05 | 0.08 | 0.04 | 0.08 | 0.05 | 0.02 | 0.02 |
|  | GFMQM | 0.16 | 0.11 | 0.01 | 0.02 | 0.02 | 0.04 | 0.02 | 0.03 | 0.18 | 0.24 | 0.00 | -0.01 | 0.02 | 0.10 |
|  |  | 0.22 | 0.24 | 0.04 | 0.06 | 0.08 | 0.07 | 0.05 | 0.06 | 0.78 | 0.63 | 0.02 | 0.00 | -0.03 | 0.06 |
|  | $130 \% 6$ | 0.02 | 0.04 | 0.04 | 0.04 | 0.05 | 0.05 | 0.03 | 0.03 | 0.05 | 0.05 | 0.04 | 0.01 | 0.05 | 0.06 |
|  |  | 0.05 | 0.04 | 0.03 | 0.04 | 0.05 | 0.04 | 0.03 | 0.04 | 0.03 | 0.04 | 0.03 | 0.02 | -0.01 | 0.01 |
|  | $\mathrm{D} 4+72$ | 0.06 | 0.06 | 0.03 | 0.03 | 0.04 | 0.04 | 0.05 | 0.03 | 0.04 | 0.05 | 0.03 | -0.01 | 0.00 | 0.06 |
|  |  | 0.04 | 0.04 | 0.03 | 0.04 | 0.04 | 0.05 | 0.07 | 0.05 | 0.05 | 0.06 | 0.04 | 0.02 | -0.05 | 0.02 |
|  | EVIGG\% | 0.06 | 0.05 | 0.04 | 0.06 | 0.03 | 0.04 | 0.04 | 0.05 | 0.05 | 0.05 | 0.04 | 0.03 | 0.03 | 0.17 |
|  |  | 0.03 | 0.00 | 0.04 | 0.03 | 0.03 | 0.02 | 0.05 | 0.02 | 0.04 | 0.04 | 0.06 | 0.04 | 0.03 | 0.01 |
|  | DUFFV, | -0.01 | 0.04 | 0.01 | 0.04 | 0.05 | 0.03 | 0.04 | 0.02 | 0.03 | -0.01 | -0.01 | -0.04 | -0.04 | -0.02 |
|  |  | 0.01 | 0.03 | 0.03 | 0.03 | 0.05 | -0.03 | 0.02 | 0.03 | 0.02 | 0.01 | 0.00 | -0.01 | 0.00 | 0.04 |
|  | OTF5S | 0.03 | 0.05 | 0.04 | 0.04 | 0.26 | 0.20 | 0.03 | 0.05 | 0.04 | 0.05 | 0.04 | 0.04 | 0.02 | 0.04 |
|  |  | 0.03 | 0.03 | 0.03 | 0.03 | 0.25 | 0.20 | 0.03 | 0.03 | 0.04 | 0.04 | 0.03 | 0.00 | 0.02 | 0.04 |
|  | 623\%4C. | 0.04 | 0.05 | 0.02 | 0.04 | 0.04 | 0.04 | 0.05 | 0.03 | 0.04 | 0.04 | 0.05 | 0.01 | -0.02 | 0.05 |
|  |  | 0.10 | 0.03 | 0.02 | 0.03 | 0.03 | 0.05 | 0.05 | 0.04 | 0.03 | 0.04 | 0.05 | -0.02 | 0.06 | 0.08 |
|  | CHZNS | 0.04 | 0.04 | 0.03 | 0.03 | 0.02 | 0.04 | 0.05 | 0.05 | 0.04 | 0.05 | 0.01 | 0.02 | 0.02 | 0.05 |
|  |  | 0.05 | 0.05 | 0.03 | 0.04 | 0.05 | 0.05 | 0.03 | 0.04 | 0.07 | 0.07 | 0.08 | -0.01 | 0.01 | 0.03 |
|  |  | GRAWI |  | ChUZN |  | EOLIN |  | JPSTA |  |  |  | HKQES |  | 3018X |  |

Kappa Chain

Table 14 continued

## Kappa Chain

|  |  | HRONB |  | BAH2A |  | HSW2E |  | MGRCL |  | FPWBB |  | MHPL |  | CRY 2 A |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | 15P82 | 0.04 | 0.04 | 0.03 | 0.03 | 0.04 | 0.03 | 0.02 | 0.01 | 0.00 | -0.03 | 0.10 | 0.03 | 0.09 | 0.07 |
|  |  | 0.04 | 0.03 | 0.03 | 0.03 | 0.04 | 0.03 | 0.02 | 0.02 | 0.01 | -0.06 | 0.13 | 0.04 | 0.09 | 0.06 |
|  | BHRWB | 0.04 | 0.03 | 0.04 | 0.02 | 0.02 | 0.03 | 0.04 | 0.04 | 0.02 | 0.02 | 0.01 | 0.04 | 0.04 | 0.06 |
|  |  | 0.03 | 0.02 | 0.03 | 0.03 | 0.03 | 0.03 | 0.09 | 0.07 | 0.03 | -0.02 | -0.02 | 0.03 | 0.04 | 0.05 |
|  | 8000 ${ }^{\text {a }}$ | 1.18. | 1.06 | 0.81 | 1.04 | 1.52 | 1.46 | 2.58 | 2.53 | 0.11 | 0.05 | 0.61 | 1.91 | 1.63 | 1.86 |
|  |  | 2.06 | 1.51 | 0.59 | 0.95 | 2.01 | 2.05. | 2.65 | 2.68 | 0.15 | 0.00 | 0.57 | 1.79 | 1.98 | 2.07 |
|  | H20\% | 2.68 | 2.64 | 0.04 | 0.04 | 0.04 | 0.04 | 0.04 | 0.03 | 0.02 | 0.00 | 0.01 | 0.01 | 0.02 | 0.05 |
|  |  | 2,46. | 2.39 | 0.04 | 0.03 | 0.03 | 0.03 | 0.02 | 0.01 | 0.01 | 0.01 | 0.00 | 0.01 | 0.01 | 0.01 |
|  | Cofyra | 0.09 | 0.04 | 0.02 | 0.04 | 0.03 | 0.04 | 0.02 | 0.00 | 0.01 | -0.01 | 0.01 | 0.03 | 0.03 | 0.04 |
|  |  | 0.19 | 0.14 | 0.03 | 0.03 | 0.03 | 0.02 | 0.03 | 0.03 | 0.01 | 0.00 | 0.03 | 0.02 | 0.01 | 0.04 |
|  | GBOEAO | 2.23 | 2.36 | 0.05 | 0.03 | 0.03 | 0.03 | 0.04 | 0.04 | 0.03 | 0.00 | 0.03 | 0.02 | 0.05 | 0.05 |
|  |  | 0.21 | 0.16 | 0.03 | 0.03 | 0.03 | 0.05 | 0.02 | 0.03 | 0.02 | -0.01 | 0.01 | 0.01 | 0.03 | 0.05 |
|  | IIESP | 0.07 | 0.08 | 0.03 | 0.02 | 0.03 | 0.03 | 0.05 | 0.06 | 0.03 | 0.00 | 0.03 | 0.06 | 1,96 | 1.97 |
|  |  | 0.04 | 0.03 | 0.03 | 0.04 | 0.04 | 0.05 | 0.04 | 0.03 | 0.05 | 0.01 | 0.01 | 0.05 | 2,44 | 2.50 |
|  | APABY, | 0.03 | 0.03 | 0.02 | 0.04 | 0.04 | 0.04 | 0.03 | 0.02 | 0.03 | -0.03 | 0.01 | 0.09 | 0.15 | 0.12 |
|  |  | 0.04 | 0.03 | 0.03 | 0.04 | 0.03 | 0.04 | 0.03 | 0.01 | 0.03 | 0.03 | -0.03 | 0.01 | 0.01 | 0.07 |
|  | 1943G | -0.33 | 0.38 | -0.11 | 0.20 | 0.37 | 0.35 | 2.34 | 2.44 | 0.27 | 0.16 | 0.19 | 0.27 | 0.12 | 0.15 |
|  |  | 0.75 | 0.75 | 0.38 | 0.73 | 0.84 | 0.95 | 2.41 | 2.38 | 0.46 | 0.40 | 0.25 | 0.58 | 0.40 | 0.47 |
|  | EZSAT | 0.33 | 0.33 | -0.06 | 0.05 | 0.05 | 0.03 | 0.35 | 0.30 | 0.01 | 0.00 | 0.01 | 0.03 | 1.21 | 0.85 |
|  |  | 0.44 | 0.73 | 0.02 | 0.04 | 0.03 | 0.02 | 0.20 | 0.26 | 0.02 | -0.01 | -0.01 | 0.04 | 1.38 | 1.07 |
|  | A9F9\% | -0.30 | -0.18 | 0.03 | 0.02 | 0.03 | 0.03 | 0.07 | 0.07 | 0.00 | 0.01 | 0.04 | 0.04 | 0.02 | 0.06 |
|  |  | -0.38 | -0.01 | 0.02 | 0.03 | 0.04 | 0.04 | 0.03 | 0.02 | 0.03 | -0.01 | 0.08 | 0.05 | 0.04 | 0.04 |
|  | 3XGB9 | -0.94 | 0.01 | -0.05 | 0.02 | 0.03 | 0.03 | 0.03 | 0.04 | 0.00 | -0.01 | -0.01 | 0.02 | 0.00 | 0.04 |
|  |  | -0.02 | 0.03 | 0.02 | 0.01 | 0.03 | 0.02 | 0.02 | 0.01 | 0.01 | -0.08 | -0.04 | 0.00 | 0.01 | -0.01 |
|  | 131119 | 0.01 | 0.00 | 0.01 | 0.02 | 0.01 | 0.02 | 0.01 | 0.02 | -0.02 | -0.05 | 0.03 | 0.00 | 0.03 | 0.04 |
|  |  | -0.03 | 0.02 | 0.02 | 0.02 | 0.02 | 0.02 | 0.01 | 0.02 | -0.01 | -0.04 | -0.01 | 0.01 | 0.03 | 0.04 |
|  | A6TR9 | 0.03 | 0.03 | 0.02 | 0.02 | 0.02 | 0.03 | 0.04 | 0.03 | 0.01 | 0.01 | 0.03 | 0.02 | 0.03 | 0.04 |
|  |  | 0.02 | 0.02 | 0.02 | 0.02 | 0.01 | 0.01 | 0.01 | 0.01 | 0.01 | 0.01 | 0.01 | 0.00 | 0.03 | 0.03 |
|  | EO22F | 0.01 | 0.01 | 0.01 | 0.01 | 0.00 | 0.01 | 0.01 | 0.01 | 0.01 | -0.02 | 0.02 | 0.04 | 0.03 | 0.06 |
|  |  | 0.03 | 0.01 | 0.01 | 0.02 | 0.02 | 0.02 | 0.00 | 0.02 | 0.01 | 0.01 | 0.04 | 0.04 | 0.02 | 0.02 |
|  | ASNDH. | 0.01 | 0.00 | 0.01 | 0.01 | 0.01 | 0.00 | 0.02 | 0.02 | 0.00 | -0.04 | 0.03 | 0.02 | 0.02 | 0.04 |
|  |  | -0.02 | -0.04 | 0.00 | -0.16 | -0.18 | -0.26 | -0.08 | 0.02 | -0.02 | -0.04 | -0.02 | 0.01 | 0.00 | 0.03 |
|  | G7M9M | 0.03 | 0.05 | 0.11 | 0.09 | 0.61 | 0.45 | 0.18 | 0.28 | 0.03 | 0.03 | 0.02 | 0.03 | 0.04 | 0.04 |
|  |  | 0.05 | 0.07 | 0.13 | 0.25 | 0.89 | 0.52 | 0.59 | 0.60 | 0.02 | 0.03 | 0.04 | 0.08 | 0.03 | 0.04 |
|  | H10R6 | 0.06 | 0.06 | 0.03 | 0.04 | 0.01 | 0.02 | 0.05 | 0.05 | 0.04 | 0.04 | 0.03 | 0.03 | 0.03 | 0.04 |
|  |  | -0.05 | 0.00 | -0.01 | -0.01 | 0.01 | -0.01 | 0.04 | 0.00 | -0.02 | 0.04 | 0.00 | 0.06 | 0.05 | 0.02 |
|  | 04102 | 0.00 | 0.04 | 0.03 | 0.04 | -0.02 | 0.00 | 0.04 | 0.03 | 0.02 | 0.00 | 0.00 | 0.06 | 0.03 | 0.05 |
|  |  | 0.00 | 0.01 | 0.05 | 0.00 | 0.04 | 0.05 | 0.05 | 0.06 | 0.06 | 0.06 | 0.02 | 0.03 | 0.02 | 0.03 |
|  | EVUGG: | 0.04 | 0.06 | 0.04 | 0.07 | 0.04 | 0.05 | 0.06 | 0.06 | 0.06 | 0.06 | -0.01 | -0.02 | 0.00 | 0.05 |
|  |  | 0.00 | -0.01 | 0.03 | -0.04 | 0.03 | 0.00 | 0.02 | 0.01 | 0.04 | 0.03 | -0.04 | 0.00 | 0.01 | 0.03 |
|  | Dutiy | 0.03 | 0.03 | 0.03 | 0.03 | 0.05 | 0.04 | 0.05 | 0.05 | 0.08 | 0.09 | -0.03 | 0.02 | 0.03 | 0.04 |
|  |  | 0.04 | 0.05 | 0.06 | 0.08 | 0.05 | 0.05 | 0.05 | 0.06 | 0.04 | 0.06 | 0.00 | 0.01 | 0.02 | 0.04 |
|  | DTF5S | 0.05 | 0.04 | 0.04 | 0.04 | 0.05 | 0.05 | 0.07 | 0.06 | 0.06 | 0.04 | 0.02 | 0.06 | 0.10 | 0.03 |
|  |  | 0.04 | 0.05 | 0.05 | 0.03 | 0.04 | 0.07 | 0.05 | 0.06 | 0.04 | 0.03 | -0.02 | 0.01 | 0.06 | 0.04 |
|  | D23Y4. | 0.03 | 0.04 | 0.00 | 0.04 | 0.05 | 0.06 | 0.06 | 0.05 | 0.05 | 0.05 | 0.02 | 0.08 | 0.04 | 0.05 |
|  |  | 0.03 | 0.03 | 0.03 | 0.04 | 0.05 | 0.04 | 0.06 | 0.06 | 0.04 | 0.04 | -0.01 | 0.01 | 0.04 | 0.04 |
|  | ClZNS | 0.02 | 0.04 | 0.05 | 0.05 | 0.05 | 0.04 | 0.05 | 0.05 | 0.06 | 0.05 | -0.01 | 0.00 | 0.01 | 0.04 |
|  |  | 0.03 | 0.02 | 0.02 | 0.05 | 0.03 | 0.03 | 0.03 | 0.02 | 0.06 | 0.11 | -0.03 | 0.01 | 0.02 | 0.02 |
|  |  | HRONE |  | BAltzA. |  | MSHzE |  | H6RCl |  | PPYBB |  | H14\%Lu. |  | CRY2A. |  |

Kappa Chain

Table 14 continued

## Kappa Chain

|  |  | ARW3R |  | GRBLO |  | IAVS8 |  | BLP31\% |  | AEITS |  | 0002\% |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 15P82 |  | 0.05 | 0.04 | 0.06 | 0.05 | 0.09 | 0.05 | 0.02 | 0.01 | -0.02 | 0.01 | 0.00 | 0.00 |
|  |  | 0.06 | 0.00 | 0.06 | 0.07 | 0.06 | 0.07 | 0.05 | 0.08 | 0.02 | 0.00 | 0.02 | 0.04 |
| B1RW8 |  | 0.03 | 0.06 | 0.06 | 0.07 | 0.07 | 0.08 | 0.06 | 0.10 | -0.02 | 0.02 | 0.21 | 0.18 |
|  |  | 0.03 | 0.08 | 0.09 | 0.17 | 0.06 | 0.07 | 0.06 | 0.08 | 0.01 | 0.12 | 0.08 | 0.18 |
| B800, |  | 2.11. | 2.38 | 2.59. | 2.69 | 2.50 | 2.58 | 0.66 | 0.38 | 0.21 | 0.86 | 0.95 | 0.60 |
|  |  | 1.95 | 2.12. | 2.53. | 2.61 | 2.53 | 2.38. | 0.49 | 0.42 | 0.14 | 0.63 . | 0.91 | 0.35 |
| H420K. |  | 0.07 | 0.07 | 0.07 | 0.06 | 0.05 | 0.04 | 0.05 | 0.05 | -0.01 | 0.00 | 0.13 | 0.03 |
|  |  | 0.03 | 0.05 | 0.03 | 0.03 | 0.03 | 0.04 | 0.04 | 0.05 | -0.11 | 0.00 | 0.01 | 0.01 |
| CPAY8 |  | 0.05 | 0.01 | 0.06 | 0.05 | 0.06 | 0.03 | 0.04 | 0.04 | -0.02 | -0.02 | 0.00 | 0.01 |
|  |  | 0.04 | 0.05 | 0.08 | 0.05 | 0.05 | 0.06 | 0.00 | 0.00 | -0.01 | 0.00 | 0.01 | 0.01 |
| GBOGO. |  | 0.06 | 0.00 | 0.06 | 0.07 | 0.08 | 0.08 | 0.06 | 0.06 | -0.03 | 0.00 | 0.00 | 0.01 |
|  |  | 0.03 | 0.05 | 0.07 | 0.08 | 0.06 | 0.07 | 0.06 | 0.06 | 0.00 | 0.03 | 0.01 | 0.01 |
| 13ESP |  | 0.04 | 0.06 | 0.08 | 0.08 | 0.08 | 0.07 | 0.08 | 0.05 | -0.01 | 0.04 | 0.01 | 0.04 |
|  |  | 0.08 | 0.07 | 0.06 | 0.06 | 0.07 | 0.07 | 0.06 | 0.03 | -0.01 | 0.02 | 0.12 | 0.03 |
| APABY |  | 0.05 | 0.07 | 0.06 | 0.05 | 0.01 | 0.06 | 0.07 | 0.03 | 0.00 | -0.01 | 0.00 | 0.04 |
|  |  | 0.04 | 0.02 | 0.00 | 0.02 | 0.00 | -0.01 | 0.01 | 0.04 | 0.03 | -0.04 | -0.03 | 0.01 |
| 19436. |  | 0.30 | 0.58 | 2.22 . | 2.39 | 0.36 | 0.51 | 0.04 | -0.05 | 0.09 | 0.17 | 0.20 | 0.28 |
|  |  | 0.80 | 1.11, | 2.41. | 2.65. | 1.01 | 1.14. | 0.08 | 0.05 | 0.16 | 0.14 | 0.49 | 0.44 |
| EZKNT |  | 0.04 | 0.05 | 0.25 | 0.61 | 0.07 | 0.09 | 0.07 | 0.31 | 0.07 | 0.08 | 0.07 | 0.07 |
|  |  | 0.03 | 0.03 | 0.30 | 0.42 | 0.05 | 0.06 | 0.04 | 0.05 | 0.23 | 0.20 | 0.16 | 0.17 |
| A9F9\% |  | 0.04 | 0.05 | 0.12 | 0.14 | 0.05 | 0.09 | 0.06 | 0.05 | 0.13 | 0.09 | 0.09 | 0.10 |
|  |  | 0.06 | 0.05 | 0.05 | 0.06 | 0.04 | 0.05 | 0.05 | 0.03 | 0.16 | 0.12 | 0.12 | 0.11 |
| 14cers |  | 0.04 | 0.03 | 0.05 | 0.02 | 0.03 | 0.04 | 0.04 | 0.04 | 0.19 | 0.10 | 0.09 | 0.22 |
|  |  | 0.00 | 0.02 | 0.03 | 0.02 | 0.02 | 0.03 | 0.04 | 0.07 | 0.13 | 0.06 | 0.07 | 0.08 |
| 10119 |  | 0.03 | 0.05 | 0.05 | 0.32 | 0.07 | 0.06 | 0.04 | 0.02 | 0.01 | 0.05 | 0.02 | -0.02 |
|  |  | 0.03 | 0.03 | 0.06 | 0.40 | 0.02 | 0.06 | 0.03 | 0.01 | 0.03 | 0.04 | 0.04 | 0.04 |
| A6TR9\% |  | 0.03 | 0.03 | 0.04 | 0.05 | 0.07 | 0.12 | 0.05 | 0.03 | 0.17 | 0.04 | 0.05 | 0.01 |
|  |  | 0.02 | 0.01 | 0.05 | 0.06 | 0.05 | 0.07 | 0.05 | 0.04 | 0.04 | 0.06 | 0.05 | 0.07 |
| EOZ2F |  | 0.05 | 0.05 | 0.06 | 0.04 | 0.05 | 0.06 | 0.06 | 0.06 | 0.06 | 0.05 | 0.04 | 0.06 |
|  |  | 0.04 | 0.05 | 0.06 | 0.05 | 0.05 | 0.05 | 0.05 | 0.01 | -0.02 | 0.03 | 0.04 | 0.05 |
| 45NOH. |  | 0.04 | 0.04 | 0.07 | 0.29 | 0.04 | 0.06 | 0.07 | 0.07 | 0.04 | 0.03 | 0.08 | 0.06 |
|  |  | 0.02 | 0.02 | 0.03 | 0.05 | 0.01 | 0.01 | 0.04 | 0.05 | 0.00 | -0.03 | 0.22 | 0.54 |
| 67M9M4 |  | 0.06 | 0.12 | 0.06 | 0.07 | 0.07 | 0.07 | 0.04 | 0.02 | 0.04 | 0.10 | 2.56 | 2.50 |
|  |  | 0.05 | 0.09 | 0.04 | 0.04 | 0.06 | 0.07 | 0.03 | 0.00 | 0.09 | 0.62 | 2.49 | 2.88 |
| 15086 |  | 0.04 | 0.05 | 0.05 | 0.04 | 0.07 | 0.08 | 0.03 | 0.02 | 0.03 | 0.03 | 0.09 | 0.15 |
|  |  | 0.03 | 0.07 | 0.05 | 0.05 | 0.05 | 0.05 | 0.05 | 0.02 | 0.03 | 0.04 | 0.05 | 0.08 |
| 104 12 |  | 0.05 | 0.06 | 0.07 | 0.05 | 0.05 | 0.05 | 0.05 | 0.05 | 0.04 | 0.04 | 0.04 | 0.04 |
|  |  | 0.04 | 0.04 | 0.05 | 0.04 | 0.06 | 0.05 | 0.04 | 0.04 | 0.02 | 0.04 | 0.03 | 0.04 |
| EVUGG |  | 0.03 | 0.05 | 0.04 | 0.04 | 0.04 | 0.04 | 0.04 | 0.03 | 0.04 | 0.05 | 0.04 | 0.04 |
|  |  | 0.01 | -0.01 | -0.01 | 0.01 | 0.03 | 0.02 | 0.03 | 0.04 | 0.04 | 0.05 | 0.05 | 0.05 |
| butry |  | 0.03 | 0.06 | 0.04 | 0.05 | 0.04 | 0.04 | 0.02 | 0.02 | 0.00 | 0.03 | 0.03 | 0.03 |
|  |  | 0.03 | 0.04 | 0.05 | 0.04 | 0.05 | 0.06 | 0.05 | 0.01 | 0.02 | 0.04 | 0.02 | 0.02 |
| DTF5S |  | 0.04 | 0.04 | 0.05 | 0.04 | 0.06 | 0.03 | 0.04 | 0.02 | 0.01 | 0.05 | 0.03 | 0.04 |
|  |  | 0.05 | 0.05 | 0.06 | 0.06 | 0.05 | 0.06 | 0.05 | 0.04 | 0.03 | 0.03 | 0.04 | 0.03 |
| D23YC. |  | 0.03 | 0.04 | 0.06 | 0.05 | 0.05 | 0.06 | 0.04 | 0.04 | 0.03 | 0.05 | 0.06 | 0.05 |
|  |  | 0.03 | 0.04 | 0.03 | 0.05 | 0.04 | 0.05 | 0.04 | 0.02 | 0.02 | 0.04 | 0.04 | 0.04 |
| ClZNS |  | 0.01 | 0.03 | 0.05 | 0.04 | 0.03 | 0.02 | 0.03 | 0.04 | 0.02 | 0.04 | 0.03 | 0.03 |
|  |  | 0.02 | 0.02 | 0.05 | 0.02 | 0.03 | 0.04 | 0.03 | 0.03 | 0.04 | 0.04 | 0.04 | 0.03 |
|  |  | ARW3R |  | GRBEO |  | 3AVSR |  | BEPSKK |  | AESTS |  | GOOZ55. |  |

Kappa Chain


| Controls |  |
| :---: | :---: |
| positive | 3.176 |
| negative | 0.059 |

## Example 9

In this Example, a human subject is administered a vaccine comprising an antigen of interest, and blood samples are taken before vaccination (week 0 ) and then at weeks 1 and 2. Subsequent samples are taken at 4-week intervals up to week 52. PBMC are isolated as

## I. Generation of the Nucleic Acid Sequences.

## Antigen-specific, memory and total B-cell isolation and RNA purification

For B-cell isolation, a negative selection method is used to eliminate all non-B-cells from the PBMC using Invitrogen's Dynabeads Untouched B-cell Isolation kit (Invitrogen cat\#113-51D) following the manufacturer's protocol. The resulting unlabeled B-cell population is further processed to isolate either antigen-specific or memory B-cells.

For antigen-specific B-cell isolation, total unlabeled B-cells are incubated with biotinylated antigen that is immobilized on streptavidin magnetic beads (Pierce-Thermo Scientific cat\#88816) on a rotator at room temperature for 20 minutes. The beads containing any antigen-binding B-cells are then washed twice with 1xPBS. The washed beads are then resuspended in Qiagen's RNeasy kit RLT lysis buffer (supplemented with $1 \% \beta$ mercaptoethanol) for RNA isolation.

For memory B-cell isolation, $\mathrm{CD}_{2} 7^{+}$and surface $\mathrm{IgG}^{+}$cells are isolated from total unlabeled B-cells using Miltenyi's MACS kits for $\mathrm{CD}_{2} 7^{+}$and surface $\mathrm{IgG}^{+}$cell isolation (Miltenyi Biotec (Auburn, CA) cat\#130-051-601 and 130-047-501). To simultaneously isolate $\mathrm{CD} 27^{+}$and sIgG ${ }^{+}$B-cells, magnetic bead-conjugated antibodies to both cell surface markers are added at the same time during the incubation step. Upon purification, memory B-cells are spun down at 300x g for 10 minutes, and then lysed in RLT buffer for RNA as described above for RNA isolation.

RNA is purified from selected cells using Qiagen's RNeasy kit (Qiagen cat\# 74104) following the manufacturer's protocol. On-column DNase I-treatment is conducted to eliminate contaminating genomic DNA by incorporating a DNase I digest step. After the RW1 buffer wash, DNase I (Qiagen cat\# 79254) diluted in RDD buffer is applied to the RNA purification column and incubated for 20 minutes at room temperature. The column is then washed once more with RW1 buffer, followed by two washes with RPE buffer, and the RNA is eluted with either 30 or $50 \mu \mathrm{l}$ water. The concentration of the RNA is determined by absorbance measured on a Nanodrop spectrophotometer (Thermo Scientific) at wavelength 450 nm .

## cDNA synthesis and generation of amplicons by PCR

RNA isolated from memory or antigen-specific B-cells is first reverse transcribed as described in Example 8. Amplicons of heavy, kappa and lambda chain variable regions for sequencing are generated by PCR as follows. For amplification of heavy chain, four independent reactions (each one specific to gene families of $\mathrm{V}_{\mathrm{H}} 1$ and $7 ; \mathrm{V}_{\mathrm{H}} 2,5$ and $6 ; \mathrm{V}_{\mathrm{H}} 3$; and $V_{H} 4$ ) are run for each cDNA sample using the primers described in Example 8 to preserve the natural distribution of $\mathrm{V}_{\mathrm{H}}$ gene transcript frequency in the pool of B -cells. For kappa and lambda chain amplification, a single reaction for each chain is run for each cDNA sample. For each reaction, an equimolar mixture of forward primers is used with the same concentration of reverse primer(s). Amplification is performed with fusion primers compatible for 454 Sequencing (Roche) by the Lib-L platform. Reverse primers are designed to hybridize to the 5 ' end of the constant region of each chain. These primers contain the Lib-L primer B and MID sequences so that sequencing reads begin from the extreme $5^{\prime}$ end of each constant region (in reverse sense) and into the $3^{\prime}$ end of the variable region. For heavy and kappa chains, a single reverse primer is used for each MID, whereas for lambda chain, two distinct reverse primers were required for each MID.

PCR amplification is performed using Finnzyme's Phusion Hot Start II polymerase (Thermo Scientific cat\# F-540S) where the reaction mix and conditions are set up as described in Example 8.

To ensure the absence of any false amplification from contaminating template in any
of the reagents, duplicate reactions are set up for each mixture (four separate reactions for heavy chain, and one for each light chain) where the cDNA template is substituted with water. These negative control reactions with no template are run at the same time as the samples containing template. Upon completion of the PCR program, $3 \mu \mathrm{l}$ of each reaction (including the negative controls) is analyzed by electrophoresis on a $1.5 \%$ TAE agarose gel for the presence of the amplicons (approximately 540bp for heavy chain, approximately 485bp for kappa chain and approximately 510 bp for lambda chain) when template is added to the reaction but not in the absence of cDNA.

To preserve cognate pairing of antibody chains during sequencing, the isolated $B$ cells are subjected to single cell encapsulation using single-cell microdroplet encapsulation (Raindance Technologies, Inc., Lexington, MA). The encapsulated B cells are then fused with a single cell RT-PCR reagent (the reagent sold by Qiagen, as Cat \# 210210) with amplification primers to generate linked heavy and light chain PCR products from each single B cell. Overlap PCR (Meijer P.J. et al., J. Mol. Biol. 358(3):764-72, 2006) is used to stitch the heavy and light chain PCR products into one DNA for preservation of antibody chain pairs through downstream sequencing.

## Amplicon purification, analysis, quantitation, and preparation for 454 sequencing

To eliminate excess primers and/or primer dimers in the PCR samples, amplicons are purified using Agentcourt Ampure magnetic beads (Beckman Coulter cat\#A63881) following the manufacturer's protocol (000387v001). For heavy chain, all four reactions (VH1/7, VH2/5/6, VH3, VH4) are pooled and purified as one sample, thus a total of three amplicon samples (heavy, kappa and lambda chains) are purified for each cDNA amplification. The protocol for Ampure purification is modified in that purifications are done in single 1.5 ml microtubes using a generic magnetic rack that is suitable for 1.5 ml tubes instead of in a 96-well plate format. All volumes and other procedures are as described in the protocol. The eluted amplicons after Ampure purification are then analyzed for purity and absence of any contaminating DNA species on the Agilent 2100 Bioanalyzer using the high sensitivity DNA chip (Agilent Technologies cat\# 5067-4626) by following the manufacturer's protocol.

Once the purity of amplicons is verified, the concentration of the DNA is quantified on a fluorometer using the Quant-iT PicoGreen dsDNA Assay Kit (Invitrogen cat\#P7589) as described in the manufacturer's protocol. The Lambda DNA provided in the kit is used as a concentration standard with which a standard curve was generated from $100 \mathrm{ng} /$ well to 1.56

## II. Generation of Peptide Fragments:

## Purification Of Antigen-Specific IgG From Human Donor Plasma

## Screening for reactivity to antigen.

Reactivity of plasma IgG to the antigen(s) of interest is tested by ELISA. Briefly, high-binding 96 -well plates (Costar cat\#) are coated $100 \mu \mathrm{l} /$ well of antigen at $2 \mu \mathrm{~g} / \mathrm{ml}$ dissolved in carbonate buffer at $37^{\circ} \mathrm{C}$ for two hours or $4^{\circ} \mathrm{C}$ overnight. The plates are rinsed in PBS-Tween at $37^{\circ} \mathrm{C}$ for 1 hour. Plasma samples are diluted at $1 / 100,1 / 500$ and $1 / 1000$ and $1 / 2000$ in $5 \%$ milk PBS-Tween, and $100 \mu \mathrm{l}$ of each dilution is added in duplicates of blocked wells of the 96 -well plate and incubated for 2 hours at $37^{\circ} \mathrm{C}$. The plates are washed three times with 1x PBS-TWEEN and horseradish peroxidase-conjugated anti-human IgG antibody (Southern Biotech 2040-05) diluted 1/4000 in PBS-Tween is added to each well $(100 \mu \mathrm{l})$ and incubated at $37^{\circ} \mathrm{C}$ for one hour. The plates are washed 6 times with PBSTween and developed by addition of $50 \mu \mathrm{l}$ TMB substrate solution (BioFX cat\#TMBW-$1000-01$ ), followed by $50 \mu \mathrm{l}$ of stop solution (BioFX cat\# STPR1000-01). The signals are measured at optical density of 450 nm . Serum titers are observed to generally increase with time following vaccination.

## Purification of antigen-specific IgG from total plasma IgG

Total IgG are purified from each serum sample using Protein $G$ as described in Example 8. The purified IgG are dialyzed twice against 4 liters of 1 xPBS in a 10 kDa cut-off dialysis cassette (Pierce cat\#66456), and the IgG concentration is determined by measuring the absorbance at 280 nm on a Nanodrop photospectrometer (Thermo Scientific). The Protein G-purified IgG are then affinity purified using beads bound to the antigen as described in Example 8. The affinity-purified antibodies from each sample are collected for mass spectrometry analysis.

## III. Mass Spectrometry

Mass spectrometry analysis is performed as described above. Briefly, following digestion with a protease (e.g., trypsin) and/or a chemical protein cleavage reagent (e.g., cyanogen bromide), mass spectrometry analysis is performed on the peptides. The resulting MS2 spectra are correlated to the theoretical MS2 spectra derived from the information in
the genetic material database, in order to identify the genetic sequences that encode antibodies that specifically bind to the antigen of interest. By determining the sequences of the antibodies in the samples, the composition of the antigen-specific antibody population in the subject at multiple points in time following vaccination is determined.

## Example 10

This Example describes the production of antigen-specific human antibodies using a transgenic animal that expresses human antibody genes.

XENOMOUSE strain XMG1-KL mice (Amgen, Thousand Oaks, CA) have their endogenous mouse antibody machinery inactivated and contain human immunoglobulin heavy and light chain loci (Jakobovits et al., 2007, Nature Biotechnol., 25:1134-43). These mice produce fully human $\operatorname{IgG} 1 \kappa$ and $\operatorname{IgG} 1 \lambda$ antibodies. The mice are immunized with a human antigen of interest, and a genetic material database and peptide database are generated using the following methods.

## I. Genetic Material Database:

## Cell isolation.

Spleens from immunized mice are flushed 5 times with 5 mL of RPMI/10\%FCS using a syringe and 21 G needle. Cells are frozen in $90 \%$ FCS/10\% DMSO. A total of 50$100 \times 10^{\wedge} 6$ cells are isolated from each spleen.

## RNA Isolation and cDNA Synthesis.

Total RNA is isolated from Splenocytes according to manufacturer's protocol using QIAshredder (Qiagen cat\#79654) and RNeasy mini kit (Qiagen, Hilden, Germany; cat\#74104). RNA is DNAse treated on column as per a standard next generation sequencing protocol. Total RNA concentration is measured using an ND-1000 spectrophotometer (NanoDrop; commercially available from Thermo Scientific, Wilmington, DE).

The isolated RNA is used for first-strand cDNA synthesis by reverse transcription using Thermoscript RT-PCR system (Invitrogen (part of Life Technologies), Carlsbad, CA
cat\#11146-024). cDNA is synthesized using 1.5 ug of RNA and oligo dT primer according to manufacturer's protocol.

## $V_{H}$ and $V_{L}$ amplification.

Amplicons of heavy, kappa and lambda chain variable regions for sequencing are generated by PCR as follows using primers specific for human antibody sequences as described in Example 8. For amplification of heavy chain, four independent reactions (each one specific to gene families of $\mathrm{V}_{\mathrm{H}} 1$ and $7 ; \mathrm{V}_{\mathrm{H}} 2,5$ and $6 ; \mathrm{V}_{\mathrm{H}} 3$; and $\mathrm{V}_{\mathrm{H}} 4$ ) are run for each cDNA sample to preserve the natural distribution of $\mathrm{V}_{\mathrm{H}}$ gene transcript frequency in the pool of B-cells. For kappa and lambda chain amplification, a single reaction for each chain is run for each cDNA sample. For each reaction, an equimolar mixture of forward primers is used with the same concentration of reverse primer(s). Amplification is performed with fusion primers compatible for 454 Sequencing (Roche) by the Lib-L platform. Reverse primers are designed to hybridize to the $5^{\prime}$ end of the constant region of each chain. These primers contain the Lib-L primer B and MID sequences so that sequencing reads begin from the extreme $5^{\prime}$ end of each constant region (in reverse sense) and into the 3 ' end of the variable region. For heavy and kappa chains, a single reverse primer is used for each MID, whereas for lambda chain, two distinct reverse primers were required for each MID.

PCR amplification is performed using Finnzyme's Phusion Hot Start II polymerase (Thermo Scientific cat\# F-540S) where the reaction mix and conditions are set up as described in Example 8.

To ensure the absence of any false amplification from contaminating template in any of the reagents, duplicate reactions are set up for each mixture (four separate reactions for heavy chain, and one for each light chain) where the cDNA template is substituted with water. These negative control reactions with no template are run at the same time as the samples containing template. Upon completion of the PCR program, $3 \mu \mathrm{l}$ of each reaction (including the negative controls) is analyzed by electrophoresis on a $1.5 \%$ TAE agarose gel for the presence of the amplicons (approximately 540bp for heavy chain, approximately 485bp for kappa chain and approximately 510bp for lambda chain) when template is added to the reaction but not in the absence of cDNA. PCR products are purified according to
manufacturer's protocol using AMPure XP (Agencourt; Beckman Coulter Genomics, Brea, CA, cat\#A63881) and analyzed using an Agilent 2100 BioAnalyzer.

The sequences of the PCR products are then translated into predicted amino acid sequences, which are then theoretically digested (e.g., with a protease and/or a chemical

## Antigen column preparation:

5.0 mL of fresh streptavidin (SA) magnetic beads (Pierce, cat\#88817) are washed three times with 10 mL PBS, and incubated overnight at $4^{\circ} \mathrm{C}$ with 105 uL of a $20 \mathrm{mg} / \mathrm{ml}$
stock of the antigen of interest conjugated to biotin diluted in 5.0 mL of PBS. Flow through is discarded, and beads are washed three times with 10 mL of PBS and aliquoted into ten low binding 1.7 mL tubes (Axygen (Union City, CA), cat\# MCT-175-L-C). Aliquoted beads are placed on a magnetic rack (Invitrogen, DynaMag), and PBS is removed prior to adding the dilute sera.

## Antigen-specific purification:

Protein-G purified IgG from above is added to the SA-magnetic beads coupled with biotinylated antigen. After overnight incubation at $4^{\circ} \mathrm{C}$, the flow through is collected and the beads are washed with a total of 10 mL of each of the following buffers, in series:

## PBS

RIPA buffer (i.e., radioimmunoprecipitation assay buffer; Alcaraz et al., J. Vet. Diagn. Invest. 2(3): 191-196, 1990; Ngoka, L.C., Proteome Sci. 6(1): 30, 2008)
20\% Acetonitrile in PBS
60\% Ethylene glycol in PBS
0.5 M NaCl in PBS

PBS (i.e., phosphate buffered saline)
IgG is then eluted with 5 fractions of 1.5 mL 0.1 M Glycine pH 3.5 , then 5 fractions of 1.5 mL 0.1 M Glycine pH 2.7 , then 5 fractions of 1.5 mL 0.1 M Glycine pH 1.8 and neutralized with 1 M Tris pH 8.5 . Eluates are assayed for reactivity to the antigen of interest using 96 -well plates coated with the antigen. Fractions with activity are quantitated by ELISA (Thermo, cat\#23300) and assayed for antigen reactivity by western blot. The fractions with the cleanest reactivity are analyzed by mass spectrometry.

## Mass Spectrometry

Mass spectrometry analysis is performed as described above. Briefly, following digestion with a protease (e.g., trypsin) and/or a chemical protein cleavage reagent (e.g., cyanogen bromide), mass spectrometry analysis is performed on the peptides. The resulting MS2 spectra are correlated to the theoretical MS2 spectra derived from the information in the genetic material database, in order to identify the genetic sequences that encode antibodies that specifically bind to the antigen of interest.

## Expression and Identification of Monoclonal Antibodies

Distinct heavy (gamma) chain variable region clones, kappa chain variable region clones and lambda chain variable region clones are expressed in a combinatorial format and screened for antigen-specific binding activity. Each gamma chain is paired with every light (kappa and lambda) chain to express antibodies by transient transfection of HEK293E cells in standard 96 -well tissue culture plates.

Antibody that is secreted from the transfected cells in each well is screened for binding to purified, recombinant antigen by enzyme-linked immunosorbant assay (ELISA), as described above. Several pairings of the heavy and light chains result in antibodies that specifically bind to the antigen-coated plates. These heavy and light chain pairs are selected, resulting in the production of fully human antibodies that specifically bind to the human antigen of interest.

## Equivalents

Those skilled in the art will recognize, or be able to ascertain, using no more than routine experimentation, numerous equivalents to the specific embodiments described specifically herein. Such equivalents are intended to be encompassed in the scope of the following claims.

## CLAIMS:

1. A method for obtaining nucleotide sequences or amino acid sequences of heavy or light chains of immunoglobulins that specifically bind to an antigen of interest, comprising:
(a) providing nucleic acid sequences encoding immunoglobulin chains of at least one animal, and deriving predicted mass spectra information from predicted amino acid sequences encoded by said nucleic acid sequences;
(b) obtaining a population of polyclonal immunoglobulins, wherein said population of polyclonal immunoglobulins is a purified population of polyclonal immunoglobulins obtained by antigen affinity purification with said antigen of interest, obtaining peptide fragments by protease cleavage of said population, and obtaining mass spectra information of said peptide fragments;
(c) correlating the mass spectra information of said peptide fragments obtained in step (b) with the predicted mass spectra information derived from the nucleic acid sequences in step (a), to identify nucleic acid sequences encoding immunoglobulin chains that comprise amino acid sequences corresponding to one or more of said peptide fragments; determining the variable regions and CDR3 regions of the immunoglobulin chains encoded by the identified nucleic acid sequences; and for the variable region and the CDR3 region encoded by each identified nucleic acid sequence, determining the amino acid sequence coverage of the variable region by said peptide fragments, and determining the amino acid sequence coverage of the CDR3 region by said peptide fragments; and
(d) selecting from the identified nucleic acid sequences, nucleic acid sequences having an amino acid sequence coverage of the variable region of at least $60 \%$ and an amino acid sequence coverage of the CDR3 region of at least $60 \%$, to obtain nucleic acid sequences or amino acid sequences of heavy or light chains of immunoglobulins that specifically bind to said antigen of interest.
2. A method for obtaining nucleic acid sequences or amino acid sequences of an immunoglobulin chain variable region of an immunoglobulin that specifically binds to an antigen of interest comprising:
(a) providing nucleic acid sequences encoding immunoglobulin variable regions of multiple immunoglobulins of at least one animal, and deriving predicted mass spectra information from predicted amino acid sequences encoded by said nucleic acid sequences;
(b) obtaining a population of polyclonal immunoglobulins, wherein said population of polyclonal immunoglobulins is a purified population of polyclonal immunoglobulins obtained by antigen affinity purification with said antigen of interest, obtaining peptide fragments of immunoglobulin chain variable regions of said population by protease cleavage, and obtaining mass spectra information of said peptide fragments;
(c) correlating the mass spectra information of the peptide fragments obtained in step (b) with the predicted mass spectra information derived from the nucleic acid sequences in step (a), to identify nucleic acid sequences encoding immunoglobulin chain variable regions that comprise amino acid sequences corresponding to one or more of said peptide fragments; determining the CDR3 regions of the immunoglobulin chains encoded by the identified nucleic acid sequences; and for the variable region and the CDR3 region encoded by each identified nucleic acid sequence, determining the amino acid sequence coverage of the variable region by said peptide fragments, and determining the amino acid sequence coverage of the CDR3 region by said peptide fragments; and
(d) selecting from the identified nucleotide sequences, nucleic acid sequences having an amino acid sequence coverage of the variable region of at least $60 \%$ and an amino acid sequence coverage of the CDR 3 region of at least $60 \%$, to obtain nucleic acid sequences or amino acid sequences of variable regions of immunoglobulins that specifically bind to said antigen of interest.
3. The method of claim 2, wherein variable regions are heavy chain variable regions.
4. The method of claim 2, wherein the variable regions are light chain variable regions.
5. The method of claim 1 or 2 , wherein said at least one animal of step (a) is an animal exposed to said antigen of interest.
6. The method of claim 1 or 2 , wherein the nucleic acid sequences encoding immunoglobulin chains are obtained from said at least one animal by:
(1) isolating nucleic acid molecules from white blood cells from said at least one animal; and
(2) amplifying immunoglobulin chain encoding nucleic acid molecules using primers specific for polynucleotide sequences adjacent to said immunoglobulin chain encoding nucleic acid molecules, and
(3) obtaining nucleic acid sequences of said amplified nucleic acid molecules encoding immunoglobulin chains.
7. The method of claim 6, wherein (i) the immunoglobulin chain encoding nucleic acid molecules are RNA molecules and said amplification step includes an initial reverse transcription step; or (ii) said polynucleotide sequences adjacent to said immunoglobulin chain encoding nucleic acid molecules are selected from the group consisting of genomic DNA flanking immunoglobulin genes, immunoglobulin chain constant region-encoding polynucleotide sequences, and immunoglobulin chain framework region-encoding polynucleotide sequences.
8. The method of claim 1 or 2 , wherein the predicted mass spectra information is obtained using a method comprising the steps of:
(i) performing a theoretical digest of predicted amino acid sequences encoded by the nucleic acid sequences with one or more proteases and/or one or more chemical protein cleavage reagents to generate virtual peptide fragments; and
(ii) creating predicted mass spectra of said virtual peptide fragments.
9. The method of claim 1 or 2 , wherein the nucleic acid sequences of step (a), predicted amino acid sequences, and predicted mass spectra derived from the predicted amino acid sequences, are located within a genetic material database.
10. The method of claim 1 or 2 , wherein said population of polyclonal immunoglobulins of step (b) is obtained from a body fluid sample or fraction thereof of an animal previously exposed to said antigen of interest, wherein optionally said body fluid
is selected from the group consisting of blood, cerebrospinal fluid, synovial fluid, peritoneal fluid, mucosal secretions, tears, nasal secretions, saliva, milk, and genitourinary secretions.

5 11. The method of claim 10, wherein the animal from which the population of polyclonal immunoglobulins of step (b) is obtained is of the same animal species as said at least one animal in step (a).
12. The method of claim 10 , wherein the animal from which the population of polyclonal immunoglobulins of step (b) is obtained is the same as said at least one animal in step (a).
13. The method of claim 1 or 2 , wherein the affinity purification in step (b) is accomplished by washing with $20 \%$ Acetonitrile in PBS pH7.4, washing with $60 \%$ Ethylene glycol in PBS pH7.4, and washing with 2.0 M NaCl in PBS pH 7.4 wherein molecules that are separated from said antigen of interest by said washing steps are discarded, and obtaining said purified population of polyclonal immunoglobulins from molecules that remain bound to said antigen of interest after said washing steps.
14. The method of claim 1 or 2 , wherein said population of polyclonal immunoglobulins of step (b) is obtained from the medium of cultured white blood cells in vitro.
15. The method of claim 1 or 2 , wherein the peptide fragments of step (b) are obtained from the population of polyclonal immunoglobulins by digesting the population with multiple proteases.
16. The method of claim 1 or 2 , wherein the selection in step (d) is additionally based on at least one parameter selected from the group consisting of the number of unique peptides mapped, spectrum share, total peptide count, unique peptide count, frequency of the encoding nucleic acid sequences, and clonal relatedness.
17. The method of claim 1 or 2 , wherein the nucleic acid sequences selected in step (d) have an amino acid sequence coverage of the variable region of at least $60 \%$ and an amino acid sequence coverage of the CDR 3 region of at least $75 \%$.

5 18. The method of claim 1, further comprising:
(e) making a heavy chain and a light chain based on the obtained nucleic acid sequences or amino acid sequences by recombinant molecular biology techniques or gene synthesis techniques, and
(f) assembling said heavy chain with said light chain to create an

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P40417AU01
(b) Antigen-binding Antigen-binding

${ }^{\theta} \operatorname{OOC}_{\mathrm{COO}^{-}}$
Figure 1

An individual (e.g., a Human, Rabbit or Mouse)


Sequence Analysis of Immunoglobulin Chain-encoding nucleic acid molecules


Genetic Material database

H \& L Frequency analysis Preliminary $\mathrm{H}+\mathrm{L}$ pairing


Polyclonal Antibody
Protease Digest or
Chemical Cleavage


Mass Spectrometry Analysis


Peptide Database
§
Polyclonal deconvolution:
Peptide Coverage
Frequency re-ranking
Final $\mathrm{H}+\mathrm{L}$ pairing

Identification of monoclonal leads


Obtain nucleic acid molecules comprising sequences
Encoding heavy and light chains (or variable regions thereof)
$\downarrow$
Cloning into expression vectors
$\stackrel{y}{3}$
Expression and Antibody Validation

Figure 2

## A Rabbit immunized with an antigen

B-cells (spleen, PBLs, lymphoid organs)

mRNA extraction
cDNA seq. template prep.


NG sequence analysis

H\& L chains rabbit database

Identification of monoclonal leads


4new
Polyclonal deconvolution: Peptide Coverage ranking Frequency ranking

PCR of variable $H$ \& L lead chains
Or gene synthesis of variable H \& L chains
Cloning into expression vectors
Final $\mathrm{H}+\mathrm{L}$ chain pairing


Expression and Antibody Validation
Figure 3
201724539526 Oct 2017
Genetic Sequence
Database

Functional Validation

$\xrightarrow[\substack{\text { CDNA } \\ \text { libraries }}]{ }$

Figure 4

Functionally Validated
Poly Clonal Ab

> G623FKB01AF34N G623FKBO1BSNEU G623FKB01A2ZY4 G623FKB01AOMXT G623FKB01AXQN3 G623FKB01AC4H8 G623FKB01AB7YA G623FKB01AUXQW G623FKB01BO141 G623FKB01BBIXE G623FKB01AL5DX

Light ChainsG623FKB01A3GC7
G623FKB01AXJ1C G623FKB01ATC8J G623FKB01BCUK2 G623FKB01BY43F G623FKB01BAZ8V G623FKB01A580V G623FKB01A312C G623FKB01BADIC G623FKB01ADDWX G623FKB01A2982

G623FKBO1BPZWW Bold Italics: antibody Bold Italics: antibody chains that are reactive to the antigen

| Lc ID | Hc ID plate 1 | BSNEU |  | BO141 |  | AOMXT |  | BBIXE |  | AXQN3 |  | AB7YA |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | A | B | A | B | A | B | A | B | A | B | A | B |
| BY43F | A | 0.1991 | 0.229 | 0.0995 | 0.0932 | 0.0921 | 0.0895 | 0.1193 | 0.102 | 0.1548 | 0.1429 | 0.1796 | 0.1863 |
|  | B | 0.9938 | 1.2748 | 0.2205 | 0.1463 | 0.1588 | 0.1899 | 0.9913 | 2.3349 | 0.0819 | 0.183 | 0.1681 | 0.1131 |
| A3GC7 | A | 0.6476 | 0.4431 | 0.0954 | 0.0882 | 0.155 | 0.1128 | 0.1514 | 0.1829 | 0.1717 | 0.1595 | 0.3717 | 0.192 |
|  | B | 0.571 | 0.5798 | 0.2424 | 0.0955 | 0.1599 | 0.1177 | 0.1499 | 0.1755 | 0.1035 | 0.0986 | 0.6849 | 0.1495 |
| BAZ8V | A | 0.6732 | 0.7344 | 0.0882 | 0.0744 | 0.1114 | 0.1048 | 0.4041 | 0.8467 | 0.1023 | 0.0671 | 0.1646 | 0.1409 |
|  | B | 0.7187 | 0.6232 | 0.1125 | 0.1171 | 0.1907 | 0.1074 | 0.1949 | 0.2536 | 0.1141 | 0.0937 | 0.1411 | 0.0746 |
| A312C | A | 0.0961 | 0.0897 | 0.0814 | 0.0959 | 0.0869 | 0.0849 | 0.2561 | 0.1673 | 0.1375 | 0.1143 | 0.1255 | 0.1243 |
|  | B | 0.1122 | 0.0985 | 0.2757 | 0.1145 | 0.1526 | 0.1026 | 0.2335 | 0.1849 | 0.1215 | 0.1104 | 0.1061 | 0.1562 |
|  | plate 3 |  |  |  |  |  |  |  |  |  |  |  |  |
| A580V | A | 3.47 | 3.0952 | 0.0851 | 0.0929 | 1.2682 | 1.7622 | 0.2069 | 0.2057 | 0.1099 | 0.1162 | 0.1188 | 0.175 |
|  |  | 1. 5397 | 1.3374 | 0.085 | 0.107 | 1.0411 | 18156 | 0.2299 | 0.2735 | 0.0855 | 0.1234 | 0.0968 | 0.1281 |
| BPZWW | A | 0.1257 | 0.1044 | 0.1008 | 0.1091 | 0.0894 | 0.0842 | 1.1936 | 1.2218 | 0.1372 | 0.1115 | 0.1123 | 0.124 |
|  | B | 0.162 | 0.1011 | 0.2664 | 0.1034 | 0.176 | 0.1706 | 1.4229 | 0.9012 | 0.0883 | 0.1091 | 0.0803 | 0.1166 |
| ADDWX |  | 0.092 | 0.0975 | 0.0775 | 0.1021 | 0.2347 | 0.1038 | 0.3545 | 0.2561 | 0.1478 | 0.1201 | 0.1084 | 0.1312 |
|  | B | 0.1022 | 0.0911 | 0.0684 | 0.0896 | 0.1448 | 0.1014 | 0.2455 | 0.5667 | 0.1025 | 0.1033 | 0.1026 | 0.0754 |
| AXJ1C | A | 0.0929 | 0.0835 | 0.0838 | 0.0994 | 0.0983 | 0.1008 | 0.2138 | 0.3213 | 0.1731 | 0.1144 | 0.113 | 0.1104 |
|  | B | 0.104 | 0.1181 | 0.1993 | 0.2078 | 0.1369 | 0.3932 | 0.1526 | 0.3997 | 0.2355 | 0.1435 | 0.1263 | 0.1915 |
|  | plate 5 |  |  |  |  |  |  |  |  |  |  |  |  |
| BCUK2 | A | 0.1451 | 0.1077 | 0.1041 | 0.1548 | 0.1076 | 0.1561 | 0.353 | 0.2947 | 0.1246 | 0.1243 | 0.1462 | 0.1731 |
|  | B | 0.1074 | 0.0872 | 0.1016 | 0.0952 | 0.1507 | 0.1442 | 0.3181 | 0.3213 | 0.1058 | 0.1341 | 0.0987 | 0.1175 |
| ATC8J |  | 0.0832 | 0.0987 | 0.0996 | 0.1377 | 0.1399 | 0.1315 | 0.3191 | 0.2757 | 0.1496 | 0.096 | 0.1022 | 0.1378 |
|  | B | 0.1029 | 0.0872 | 0.2011 | 0.0915 | 0.1117 | 0.1118 | 0.2807 | 0.2755 | 0.1078 | 0.0796 | 0.0789 | 0.1487 |
| BADIC | A | 3.6013 | 2.0746 | 0.0854 | 0.1078 | 3.0399 | 2.3754 | 0.187 | 0.3191 | 0.1331 | 0.0846 | 0.1008 | 0.1468 |
|  | B | 1.0088 | 1.6872 | 0.0883 | 0.1117 | 1.7897 | 1. 2741 | 0.1918 | 0.2086 | 0.0892 | 0.1111 | 0.1075 | 0.0772 |
| A2982 | A | 0.1187 | 0.1052 | 0.1109 | 0.1093 | 0.1003 | 0.0997 | 0.2348 | 0.4317 | 0.139 | 0.1361 | 0.12 | 0.0998 |
|  | B | 0.1547 | 0.0901 | 0.112 | 0.1175 | 0.1186 | 0.1065 | 0.1875 | 0.3154 | 0.1201 | 0.1146 | 0.1422 | 0.1878 |
|  | plateAB |  |  |  |  |  |  |  |  |  |  |  |  |
| AS297 |  | 0.1615 | 0.1045 | 0.121 | 0.1232 | 0.1307 | 0.1299 | 0.4285 | 0.4476 | 0.143 | 0.1361 | 0.169 | 0.1675 |
|  |  | 0.1197 | 0.084 | 0.0982 | 0.101 | 0.1362 | 0.1288 | 0.3966 | 0.3443 | 0.1103 | 0.1664 | 0.1162 | 0.1344 |
| 9106 control 2ndary |  | 1.6003 | 1.0512 | 1.4815 | 1.46830.0959 |  |  |  |  |  |  |  |  |
|  |  |  |  |  |  |  |  |  |  |  |  |  |
|  |  | 0.1702 |  |  |  | 0.1136 | 0.1081 | 0.0882 | 0.0993 | 0.0822 | 0.1045 | 0.0924 | . 1662 |

[^0]| LC ID | Hc ID plate 1 | A2ZY4 |  | AC4H8 |  | BJOL7 |  | AL5DX |  | AUXQW |  | AF34N |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | A | B | A | B | A | B | A | B | A | B | A | B |
| BY43F | A | 0.1422 | 0.0906 | 0.0956 | 0.0965 | 0.1142 | 0.0951 | 0.125 | 0.1081 | 0.124 | 0.1346 | 0.1486 | 0.148 |
|  | B | 0.3013 | 0.6843 | 0.1651 | 0.1841 | 0.124 | 0.1012 | 0.1195 | 0.0961 | 0.0982 | 0.1439 | 0.1278 | 0.1596 |
| A3GC7 | A | 0.0958 | 0.194 | 0.2652 | 0.4371 | 0.4326 | 0.4052 | 0.2398 | 0.233 | 0.1992 | 0.1144 | 0.2087 | 0.1812 |
|  | B | 0.0953 | 0.089 | 0.5074 | 0.4602 | 0.44 | 0.4557 | 0.2844 | 0.2952 | 0.3203 | 0.1075 | 0.1821 | 0.1748 |
| BAZ8V | A | 0.0966 | 0.084 | 0.2187 | 0.3583 | 0.5084 | 0.3416 | 0.4487 | 0.4084 | 0.3246 | 0.1517 | 0.1331 | 0.1267 |
|  | B | 0.0937 | 0.0759 | 0.1215 | 0.3744 | 0.4007 | 0.3472 | 0.3713 | 0.4181 | 0.2366 | 0.0809 | 0.1075 | 0.0982 |
| A312C | A | 0.0973 | 0.0973 | 0.1037 | 0.0944 | 0.1731 | 0.1139 | 0.2151 | 0.1307 | 0.1041 | 0.1071 | 0.102 | 0.1016 |
|  | B | 0.0899 | 0.1232 | 0.194 | 0.1527 | 0.1216 | 0.1237 | 0.1262 | 0.0902 | 0.1538 | 0.1891 | 0.1267 | 0.145 |
|  | plate 3 |  |  |  |  |  |  |  |  |  |  |  |  |
| A580V | A | 0.1256 | 0.1403 | 0.1009 | 0.1162 | 0.1183 | 0.0843 | 0.091 | 0.0967 | 0.1121 | 0.1369 | 1. 61.13 | 4.0514 |
|  | B | 0.0965 | 0.0923 | 0.0925 | 0.1119 | 0.151 | 0.116 | 0.1443 | 0.095 | 0.078 | 0.1416 | 4.4636 | 4.7646 |
| BPZWW | A | 0.1009 | 0.084 | 0.1061 | 0.1346 | 0.1929 | 0.1668 | 0.1219 | 0.1021 | 0.147 | 0.1143 | 0.1262 | 0.1294 |
|  | B | 0.1098 | 0.1563 | 0.1862 | 0.1062 | 0.1875 | 0.1935 | 0.1004 | 0.1742 | 0.1604 | 0.1031 | 0.0813 | 0.1238 |
| ADDWX | A | 0.0917 | 0.0898 | 0.0737 | 0.1389 | 0.2034 | 0.1038 | 0.1136 | 0.1091 | 0.1574 | 0.08 | 0.1027 | 0.1345 |
|  | B | 0.0941 | 0.0766 | 0.0868 | 0.1144 | 0.1279 | 0.0779 | 0.1884 | 0.1056 | 0.0974 | 0.0851 | 0.1016 | 0.0867 |
| AXJ1C | A | 0.0996 | 0.0985 | 0.088 | 0.1245 | 0.1463 | 0.1464 | 0.153 | 0.1366 | 0.1708 | 0.1065 | 0.1297 | 0.1081 |
|  | B | 0.1232 | 0.0857 | 0.127 | 0.1102 | 0.1855 | 0.139 | 0.1621 | 0.1028 | 0.1296 | 0.1331 | 0.1085 | 0.1783 |
|  | plate 5 |  |  |  |  |  |  |  |  |  |  |  |  |
| BCUK2 | A | 0.1191 | 0.1299 | 0.1049 | 0.0994 | 0.1008 | 0.1397 | 0.1856 | 0.1667 | 0.1412 | 0.1217 | 0.1951 | 0.1944 |
|  | B | 0.1017 | 0.0901 | 0.0908 | 0.09 | 0.1329 | 0.1093 | 0.1191 | 0.1134 | 0.0902 | 0.1369 | 0.1233 | 0.1771 |
| ATC8J | A | 0.1139 | 0.1023 | 0.1015 | 0.1093 | 0.1077 | 0.1066 | 0.0904 | 0.1016 | 0.104 | 0.1135 | 0.1038 | 0.1455 |
|  | B | 0.116 | 0.0864 | 0.1698 | 0.0999 | 0.1153 | 0.1032 | 0.0848 | 0.1091 | 0.0921 | 0.0871 | 0.0887 | 0.1607 |
| BADIC | A | 0.118 | 0.0916 | 0.0875 | 0.0922 | 0.0869 | 0.0862 | 0.0886 | 0.1082 | 0.1101 | 0.0988 | 4.4611 | 4 4264 |
|  | B | 0.1211 | 0.1007 | 0.1083 | 0.0994 | 0.1223 | 0.1087 | 0.0998 | 0.1298 | 0.1193 | 0.11 | 3.7921. | 4049. |
| A2982 | A | 0.108 | 0.0961 | 0.0886 | 0.1053 | 0.1523 | 0.1132 | 0.1099 | 0.1466 | 0.2152 | 0.1184 | 0.1058 | 0.1696 |
|  | B | 0.1896 | 0.1141 | 0.1493 | 0.1354 | 0.1649 | 0.1358 | 0.1706 | 0.1357 | 0.193 | 0.1744 | 0.1836 | 0.3142 |
|  | plate 7 |  |  |  |  |  |  |  |  |  |  |  |  |
| AS297 | A | 0.2515 | 0.0969 | 0.1444 | 0.1882 | 0.1441 | 0.1392 | 0.1377 | 0.1882 | 0.2288 | 0.1873 | 0.2154 | 0.2061 |
|  | B | 0.1416 | 0.11 | 0.1125 | 0.1112 | 0.1225 | 0.1437 | 0.1244 | 0.1287 | 0.1193 | 0.1486 | 0.1216 | 0.1323 |
| 2ndary |  | 1.6764 | 1.4255 | 1.9828 | 1.6362 |  |  |  |  |  |  |  |  |
|  |  | 0.1395 | 0.109 | 0.256 | 0.1528 | 0.1541 | 0.1128 | 0.1023 | 0.1168 | 0.0921 | 0.1133 | 0.1056 | 0.1895 |

Figure 6 continued...
$2017245395 \quad 26$ Oct 2017
RT-PCR of Rabbit Splenocytes

## Kappa and Lambda Chains <br> Heavy Chains



Figure 7

| 3126(pMet) |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: |
| NGS Sequence ID | Chain | CDR3 sequence | SEQ ID NO: | NGS freq (max 9.12\%) |
| GXRYQP201BFGRR | heavy chain | IPSYASSRGYYLIPDRLDL | 331 | 0.03\% |
| GXRYQP201BIQD2 | heavy chain | IPSYVSGRGVYIIPDRFDL | 332 | 0.19\% |
| GXRYQP201AGL7B | heavy chain | ASDYDSSRGHWLVYNRLDL | 333 | 0.08\% |
| GXRYQP201A97DZ | heavy chain | KGDPGHPNGLFFTM | 334 | 0.05\% |
| GXRYQP201ARCKH | heavy chain | IPSYVSSRGYYLIPDGLDL | 335 | 0.02\% |
| GXRYQP201A9YV7 | heavy chain | IPSYVSSRGYYLVPDGLDL | 336 | 0.01\% |
| GXRYQP201BRYT5 | heavy chain | IPSYVSSRGYYLIPDRLDL | 337 | 0.05\% |
| GXRYQP201AKBWL | heavy chain | ISSYVSSRGYWLIPDGLDL | 338 | 0.13\% |
| GXRYQP201ANGRW | heavy chain | ISSYVSSRGYYLIPDGLDL | 339 | 0.01\% |
| GXRYQP201BGVA8 | heavy chain | LYNSVVGDDM | 340 | 0.55\% |
| GXRYQP201AGFK4 | heavy chain | ASDYDSSRGHWLVYDRLDL | 341 | 0.01\% |
| GXRYQP201BJROT | heavy chain | LYNSVVGDDI | 342 | 0.03\% |
| GXRYQP201A8DBE | heavy chain | LYNSLVGDDI | 343 | 0.22\% |
| GXRYQP201B2QN3 | heavy chain | GMPGSTSGNSNI | 344 | 0.59\% |
| GXRYQP201A0CZK | heavy chain | GMPASTSGNSNI | 345 | 0.17\% |
| GXRYQP201AZOWE | heavy chain | GMPGSTSGNSNI | 346 | 0.59\% |
| GXRYQP201A1C3B | heavy chain | GVPTNRDAM | 347 | 9.12\% |
| 3126(pMet) |  |  |  |  |
| NGS Sequence ID | Chain | CDR3 sequence |  | NGS freq (max 3.69\%) |
| GXRYQP201AAKYU | light chain | AGGYKSSGDTVS | 348 | 0.14\% |
| GXRYQP201AG5FC | light chain | QGEFSCRDFDCTV | 349 | 0.11\% |
| GXRYQP201A291T | light chain | LGGYKTTTDGSI | 350 | 0.78\% |
| GXRYQP201A09ZW | light chain | QSYYHNSGTSYIT | 351 | 0.89\% |
| GXRYQP201A3ZHW | light chain | AGGYKSTTDGSA | 352 | 0.07\% |
| GXRYQP201AJ2IR | light chain | QSYYHNSGNSYIT | 353 | 0.19\% |
| GXRYQP201A5GBW | light chain | QSYYYGSGTSYIT | 354 | 0.04\% |
| GXRYQP201BP3WS | light chain | AGGYKSSGDTFT | 355 | 0.01\% |
| GXRYQP201AY70W | light chain | LGGYKKTIDGSA | 356 | 0.05\% |
| GXRYQP201BRIWK | light chain | AGGYKSASDGSA | 357 | 0.30\% |
| GXRYQP201AYFKS | light chain | QGEFSCDAGVCTL | 358 | 0.07\% |
| GXRYQP201ALDF5 | light chain | QGEFSCRSYDCTV | 359 | 0.18\% |
| GXRYQP201AATVT | light chain | LQDWSPSYADVA | 360 | 0.06\% |
| GXRYQP201APNW9 | light chain | QQGRRSVDVDNV | 361 | 0.05\% |
| GXRYQP201AT2TB | light chain | AGGYKTTTDGSI | 362 | 0.02\% |
| GXRYQP201AHXJZ | light chain | QQGYTYSNVDNV | 363 | 3.69\% |

Bold Italics are the antigen specific antibody chains
(Total of 6 heavy chains (italics) and 5 light chains (bold italics) were synthesized, including the highest abundance antibody chains based on NGS frequency)

Figure 8
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Figure 9
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Figure 10c


Figure 10d
201724539526 Oct 2017

Kappa Chain: Coverage $=65 \%$ Total Peptides $=24$
now

 (SEQ ID NO: 368)
Sigure 10 e

- Trypsin $=$ CDR

Fign. <br> $$
\begin{array}{llll} 
& \text { Heavy Chain } \\
\text { Figure 11a } & \\
& &
\end{array}
$$ <br> \title{

Heavy Chain <br> \title{
Heavy Chain <br> <br> ELISAt
Western Blot+ <br> <br> ELISAt
Western Blot+ <br> <br> Figure 11a
} <br> <br> Figure 11a
}
26 Oct 2017


$$
\text { , } 1611111
$$

Figure 11b

$$
\begin{aligned}
& \text { 鿊 + } \\
& \text { ㅍ́ } \\
& \text { Clone: } \\
& \begin{array}{l}
\text { PR-B } \longrightarrow \\
\text { PR-A } \longrightarrow
\end{array}
\end{aligned}
$$

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Figure 11d

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Figure 11e

Figures 12a-d

## Oct 2017 <br> $\stackrel{\rightharpoonup}{\sim}$



2

Figures 13a-c

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<110> Polakiewicz, Roberto
    Cheung, Wan Cheung
    Rush, II, John Edward
    Beausoleil, Sean Andre
<120> METHODS AND REAGENTS FOR CREATING MONOCLONAL ANTIBODIES
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<150> 61/566,876
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|  | Leu val Thr $\underset{20}{\text { val }}$ Ser Ser Gly Gln |

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oligonucleotide

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<210> 248

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<210> 258

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<220>

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<211> 68
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<212> DNA
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<210> 283

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gtgg 64
<210> 285
<211> }6
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<223> Description of Artificial Sequence: Synthetic
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<400> 292
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<210> 293
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<210> 297
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oligonucleotide
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<210> 298

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2017245395 12 Oct 2017
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Ile Pro Ser Tyr Ala Ser Ser Arg Gly Tyr Tyr Leu Ile Pro Asp Arg
Leu Asp Leu
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Phe Asp Leu
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Leu Asp Leu
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N Leu Asp Leu (28446_CST314_SEQ 
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Leu Tyr Asn Ser val val Gly Asp Asp Met
1 5 10
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Gly Val Pro Thr Asn Arg Asp Ala Met
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1 5 10
<210> 349
<211> }1
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1 5 10
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<211> 12
<212> PRT
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Leu Gly Gly Tyr Lys Thr Thr Thr Asp Gly Ser Ile
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Gln Ser Tyr Tyr His Asn Ser Gly Thr Ser Tyr Ile Thr
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peptide
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Gln Ser Tyr Tyr His Asn Ser Gly Asn Ser Tyr Ile Thr

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<210> 359
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Gln Gln Gly Arg 
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<400> 364
\(\underset{1}{\text { Glu Gln Leu Lys }} \underset{5}{\text { Glu }}\) Ser Gly Gly Gly \(\underset{10}{\text { Leu Val }}\) Thr Pro Gly \(\underset{15}{\text { Gly }}\) Thr
Leu Thr Leu \(\underset{20}{\text { Thr }}\) Cys Thr val Ser \(\underset{25}{\text { Gly }}\) Phe Ser Leu Ser Ser Val Ala
Met Ile \(\underset{35}{\operatorname{Trp}}\) Val \(\underset{40}{\text { Arg Gln Ala }} \underset{40}{\text { Pro Gly }}\) Lys Gly Leu \(\underset{45}{\text { Glu }}\) Tyr Ile Gly

\(\underset{65}{\text { Arg }}\) Phe Thr val Ser \(\underset{70}{\text { Lys }}\) Thr Ser Thr Thr \(\underset{75}{\text { val }}\) Asp Leu Lys Met \(\underset{80}{\text { Thr }}\)
Ser Leu Thr Thr \(\underset{85}{\text { Glu }}\) Asp Thr Ala Thr \(\underset{90}{\text { Tyr }}\) Phe Cys Ala Arg \(\underset{95}{\text { Ile }}\) Pro
Ser Tyr val Ser Gly Arg Gly val \(\begin{aligned} & \text { Tyr Ile Ile Pro Asp } \\ & 100 \\ & 100\end{aligned} \quad \begin{aligned} & \text { Arg } \\ & 110\end{aligned}\)
Leu Trp Gly Gln Gly Thr Leu Val \(\begin{aligned} & 115 \\ & 120\end{aligned}\) Thr val Ser Ser Gly
<210> 365
<211> 116
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\({ }_{1}^{\text {Ala }}\) Thr Phe Ala Ala Val Leu Thr Gln \({ }_{5}\) Thr Pro Ser Pro Val \({ }_{10}\) Ser Ala
Ala val Gly \(\underset{20}{\text { Gly }}\) Thr val Thr Ile \(\underset{25}{\text { Ser Cys Gln Ser Ser }} \underset{30}{\text { Glu }}\) Ser val
Tyr Lys \(\underset{35}{\text { Asn }}\) Asn Tyr Leu Ser \(\underset{40}{\operatorname{Trp}}\) Tyr Gln Gln Lys \(\underset{45}{\text { Pro }}\) Gly His Ser

Tyr Lys Ser Ala Ser Asp Gly Ser Ala Phe Gly Gly Gly
100 \(\begin{aligned} & \text { Thr Glu Val } \\ & 110\end{aligned}\)
Val Val Lys Gly
<210> 366
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<212> PRT
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Ala Thr Phe Ala Ala Val Leu Thr Gln Thr Pro Ser Pro Val Ser Ala
Ala Val Gly \(\underset{20}{\text { Gly }}\) Thr Val Thr Ile Ser Cys Gln Ser Ser \(\underset{25}{ }{ }_{30}\) Ger Val
Tyr Lys Asn Asn Tyr Leu Ser \(\underset{35}{ } \operatorname{Trp}_{40}\) Tyr Gln Gln Lys \(\underset{45}{ } \quad\) Gro Gly His Ser

Ser
65 Arg Phe Lys Gly Ser Gly Ser Gly Thr Gln \(\underset{70}{75}\) Phe Thr Leu Thr Ile
Ser Asp Leu Glu Cys Ala Asp Ala Ala
85 \(\underset{90}{ } \begin{gathered}\text { Thr } \\ 90\end{gathered} \quad\) Tyr Cys Leu Gly Gly

Val Val Lys Gly
    115
<210> 367
<211> 120
<212> PRT
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Gln Ser Val Glu Glu Ser Gly Gly Arg Leu val Pro Pro Gly Thr Pro
Leu Thr Leu Thr Cys Thr val Phe Gly Phe Asp Val Ser Ser His Ile
Met Ser }\mp@subsup{\operatorname{Trp}}{35}{\rp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Ile Gly
Leu Val Asp Ile Gly Lys Ser Ile Lys Trp Tyr Ala Ser Trp Ala Lys
G7y Arg Phe Thr Ile Ser Lys Thr Ser Ser Thr Ile Tyr Leu Lys Leu

```
Thr Arg Pro Thr \(\begin{gathered}\text { Thr Gly Asp Thr Ala } \\ 85\end{gathered} \underset{90}{\text { Thr }} \begin{gathered}\text { Tyr Phe Cys Ser } \\ 95\end{gathered} \underset{95}{\operatorname{Arg}}\) Gly
Phe Ala Leu \(\begin{array}{r}\operatorname{Trp} \text { Gly Pro Gly Thr Leu Val Thr Val Ser Ser Gly Gln } \\ 100\end{array}\)
Pro Lys Ala Pro Ser Val Phe Pro
115
<210> 368
<211> 120
<212> PRT
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<220>
<223> Description of Artificial Sequence: Synthetic
<400> 368

Gly Thr Val Thr Ile Asn Cys Gln Ala Ser Pro Ser Val Tyr Gly Asn \(\begin{gathered}20 \\ 20\end{gathered}\)
Tyr Leu Ser \(\underset{35}{ } \operatorname{Trp}\) Phe Gln Gln Lys Pro Gly Gln Pro \(\underset{40}{ } \underset{45}{ } \quad\) Lys Leu Leu
Ile Leu Asn Ala Ser Thr Leu Pro Ser Gly Val Ser Ser Arg Phe Lys
50
50
Gly
65 Ser Gly Ser Gly \(\begin{aligned} & \text { Thr } \\ & 70\end{aligned}\) His Phe Thr Leu \(\begin{gathered}\text { Thr } \\ 75\end{gathered}\)
Cys Asp Asp Ala Ala
85
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Ser Asp Asp Cys Asn Val Phe Gly Gly Gly Thr Glu Val Val Val Lys
Gly Asp Pro Val Ala Pro Thr Val
1 1 5
120
<210> 369
<211> 14
<212> PRT
<213> Artificial Sequence
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<400> 369
Ala Gly Thr Asn Tyr Trp Ala Ile Tyr Tyr Gly Met Asp Leu
<210> 370
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<212> PRT
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peptide
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Gln Gln Gly Tyr Lys Ile Thr Asn Ile Glu Asn Val
<210> 371
<211> 9
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Cys Ala Arg Gly Ser Ala Phe Ala Tyr
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Cys Ala Arg Arg Gly Tyr Asp Gly Ser Tyr Tyr Phe Asp Tyr
<210> 373
<211> 12

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<400> 373
Cys Ala Arg His Glu Pro Leu Asn Trp Phe Pro Tyr
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Clys Ala Arg Arg Gly Tyr Tyr Ser Asn Thr Thr Tyr val Asp Tyr
<210> 375
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Cys Ala Arg Arg Gly Tyr Tyr Ala Asp Thr Thr Tyr val Asp Tyr
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1 Ala Arg Glo
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28446_CST314_SEQ
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peptide
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Cys Ala Arg Ser fr Arg Thr Gly Ile Phe Asp Tyr
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peptide
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Cys Val Arg Ser fry Arg Ser Gly Ile Phe Asp Tyr
<210> 381
<211> 13
<212> PRT
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peptide
<400> 381
Cys Ala Arg Tyr Tyr Arg Asn Tyr Gly Gly Phe Asp Tyr
<210> 382
<211> 10
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<223> Description of Artificial Sequence: Synthetic
peptide
<400> 382
Cys Ala Arg Leu Thr Ala Tyr Phe Asp Tyr
1 5 Ala

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<210> 383
<211> 9
<212> PRT
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<400> 383
Phe Gln Gly Ser His Val Pro Phe Thr
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<210> 384
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                                    peptide
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Leu Gln Tyr Ala ser Tyr Pro Trp Thr
1
<210> 385
<211> 9
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    peptide
<400> 385
Gln Gln His Phe Ser Thr Pro Pro Thr
1
<210> 386
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    peptide
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Gln Gln His Tyr Ser Thr Pro Phe Thr
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<210> 387
<211> 9
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<223> Description of Artificial Sequence: Synthetic
    peptide
<400> 387
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| :---: | :---: |
| O | $\begin{aligned} & <210>388 \\ & <211>9 \\ & <212>\text { PRT } \\ & <213>\text { Artificial sequence } \end{aligned}$ |
|  | <220> <br> <223> Description of Artificial Sequence: Synthetic peptide |
| $\begin{aligned} & n \\ & n \\ & n \end{aligned}$ | $<400>388$ <br> Gln Gln His Tyr Ser Thr Pro Pro Thr 1 |
| $\stackrel{\bigcirc}{\mathrm{N}}$ | $\begin{aligned} & <210>389 \\ & <211>9 \\ & <212>\text { PRT } \\ & <213>\text { Artificial sequence } \end{aligned}$ |
|  | <220> <br> <223> Description of Artificial Sequence: Synthetic peptide |
|  | $<400>389$ <br> Gln Gln Arg Ser Ser Tyr Pro Phe Thr 1 |
|  | $\begin{aligned} & <210>390 \\ & <211>9 \\ & <212>\text { PRT } \\ & <213>\text { Artificial sequence } \end{aligned}$ |
|  | <220> <br> <223> Description of Artificial Sequence: Synthetic peptide |
|  | $<400>390$ <br> Gln Gln Ser Lys ${ }_{5}$ Glu Val Pro Leu Thr 1 |
|  | $\begin{aligned} & <210>391 \\ & <211>9 \\ & <212>\text { PRT } \\ & <213>\text { Artificial sequence } \end{aligned}$ |
|  | <220> <br> <223> Description of Artificial Sequence: Synthetic peptide |
|  | <400> 391 <br> Gln Gln Ser Asn Glu Asp Pro Arg Thr 1 5 |
|  | $\begin{aligned} & <210>392 \\ & <211>9 \\ & <21 \gg \text { PRT } \\ & <213>\text { Artificial sequence } \end{aligned}$ |
|  | <220> <br> <223> Description of Artificial Sequence: Synthetic peptide |

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Gln His Phe Trp Gly Thr Pro Trp Thr
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<211> 9
<212> PRT
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peptide
<400> 393
Gln Gln Tyr Tyr ser Tyr Pro Arg Thr
<210> 394
<211> 9
<212> PRT
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<223> Description of Artificial Sequence: Synthetic
peptide
<400> 394
Trp Gln Gly Thr His Phe Pro Gln Thr
1 5
<210> 395
<211> 9
<212> PRT
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<223> Description of Artificial Sequence: Synthetic
peptide
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Ser Gln Ser Thr His Val Pro Trp Thr
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<223> Description of Artificial Sequence: Synthetic
peptide
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Lys Leu Gly Leu
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<210> 397
<211>
<212> PRT
<213> Artificial Sequence
<220>

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<400> 397
Gly Phe Ser Leu
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<210> 398
<211> 5
<212> PRT
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peptide
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[^0]:    Figure 6

