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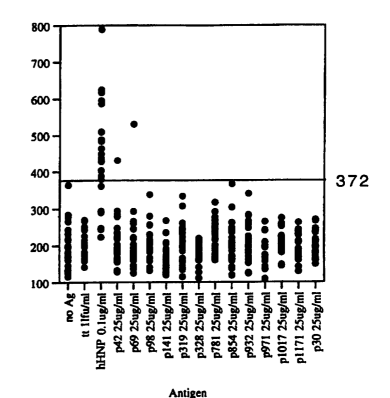
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(54) Title: INTRACELLULAR DOMAIN OF THE HER-2/NEU PROTEIN FOR PREVENTION OR TREATMENT OF MALIGNAN-CIES

(57) Abstract

Compounds and compositions for eliciting or enhancing immune reactivity to HER-2/neu protein are disclosed. The compounds include polypeptides and nucleic acid molecules encoding such peptides. The compounds may be used for the prevention or treatment of malignancies in which the HER-2/neu oncogene is associated.



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Description

INTRACELLULAR DOMAIN OF THE HER-2/NEU PROTEIN FOR PREVENTION OR TREATMENT OF MALIGNANCIES

Technical Field

The present invention is generally directed toward polypeptides, and nucleic acid molecules encoding such polypeptides, for eliciting or enhancing an immune response to HER-2/neu protein, including for use in the treatment of malignancies in which the HER-2/neu oncogene is associated.

15 Background of the Invention

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Despite enormous investments of financial and human resources, cancer remains one of the major causes of death. For example, cancer is the leading cause of death in women between the ages of 35 and 74. Breast cancer is the most common malignancy in women and the incidence for developing breast cancer is on the rise. will be diagnosed with the disease. approaches to cure breast cancer have centered around a combination of surgery, radiation and chemotherapy. approaches have resulted in some dramatic successes in certain malignancies. However, these approaches have not been successful for all malignancies and breast cancer is most often incurable when attempting to treat beyond a certain stage. Alternative approaches to prevention and therapy are necessary.

A common characteristic of malignancies is uncontrolled cell growth. Cancer cells appear to have undergone a process of transformation from the normal phenotype to a malignant phenotype capable of autonomous

growth. Amplification and overexpression of somatic cell genes is considered to be a common primary event that results in the transformation of normal cells to malignant cells. The malignant phenotypic characteristics encoded by the oncogenic genes are passed on during cell division to the progeny of the transformed cells.

Ongoing research involving oncogenes identified at least forty oncogenes operative in malignant cells and responsible for, or associated transformation. Oncogenes have been classified into 10 different groups based on the function or putative location of their gene products (such as the protein expressed by the oncogene).

are believed to be essential Oncogenes certain aspects of normal cellular physiology. 15 In this regard, the HER-2/neu oncogene is a member of the tyrosine protein kinase family of oncogenes and shares a high homology with the epidermal growth factor degree of ${\tt HER-2/\it neu}$ presumably plays a role in cell receptor. growth and/or differentiation. HER-2/neu appears induce malignancies through quantitative mechanisms that result from increased or deregulated expression of essentially normal gene product.

HER-2/neu (p185) is the protein product of the 25 HER-2/neu oncogene. The HER-2/neu gene is amplified and the HER-2/neu protein is overexpressed in a variety of cancers including breast, ovarian, colon, lung prostate cancer. HER-2/neu is related to malignant transformation. It is found in 50%-60% of ductal in situ 30 carcinoma and 20%-40% of all breast cancers, as well as a substantial fraction of adenocarcinomas arising in the ovaries, prostate, colon and lung. HER-2/neu intimately associated not only with the malignant phenotype, but also with the aggressiveness of the

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malignancy, being found in one-fourth of all invasive breast cancers. HER-2/neu overexpression is correlated with a poor prognosis in both breast and ovarian cancer. HER-2/neu is a transmembrane protein with a relative 5 molecular mass of 185 kd that is approximately 1255 amino acids (aa) in length. It has an extracellular binding domain (ECD) of approximately 645 aa, with 40% homology to epidermal growth factor receptor (EGFR), hydrophobic transmembrane anchor domain (TMD), and 10 carboxyterminal cytoplasmic domain (CD) of approximately 580 aa with 80% homology to EGFR.

Due to the difficulties in the current approaches to therapy of cancers in which the HER-2/neu oncogene is associated, there is a need in the art for improved compounds and compositions. The present invention fulfills this need, and further provides other related advantages.

Summary of the Invention

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20 Briefly stated, the present invention provides polypeptides, nucleic acid molecules (directing expression of such polypeptides) and viral vectors (directing the expression of such polypeptides) for use for the immunization, or the manufacture of a medicament 25 for immunization, of a warm-blooded animal against a malignancy in which the HER-2/neu oncogene is associated. A polypeptide or nucleic acid molecule according to this invention may be present in a composition that includes a pharmaceutically acceptable carrier or diluent. polypeptide, nucleic acid molecule, viral vector or pharmaceutical composition may be used for immunization on a one-time basis (e.g., when a malignancy is suspected) or on a periodic basis (e.g., for an individual with an elevated risk of acquiring or reacquiring a malignancy).

A medicament for immunization may be useful in the treatment of an existing tumor or to prevent tumor occurrence or reoccurrence.

In one embodiment, the present provides a polypeptide encoded by a DNA sequence selected (a) nucleotides 2026 through 3765 of SEQ ID NO:1; and (b) DNA sequences that hybridize to a nucleotide sequence complementary to nucleotides 2026 through 3765 of SEQ ID NO:1 under moderately stringent conditions, wherein 10 the DNA sequence encodes a polypeptide that produces an immune response to HER-2/neu protein. In a preferred embodiment, a polypeptide has the amino acid sequence of SEQ ID NO:2 from lysine, amino acid 676, through valine, amino acid 1255, or a variant thereof that produces at 15 least an equivalent immune response. A composition is provided that comprises a polypeptide of the present invention in combination with a pharmaceutically acceptable carrier or diluent.

In another embodiment, a polypeptide or composition of the present invention is provided for the immunization of a warm-blooded animal against a malignancy in which the HER-2/neu oncogene is associated. In another embodiment, such a polypeptide or composition is used for the manufacture of a medicament for immunization of a warm-blooded animal against a malignancy in which the HER-2/neu oncogene is associated.

In another embodiment, a nucleic acid molecule directing the expression of a polypeptide according to the present invention is provided for immunization by transfecting the cells of a warm-blooded animal with the nucleic acid molecule. In another embodiment, such a nucleic acid molecule is used for the manufacture of a medicament for immunization of a warm-blooded animal

against a malignancy in which the HER-2/neu oncogene is associated.

In another embodiment, a viral vector directing the expression of a polypeptide according to the present invention is provided for immunization by infecting the cells of a warm-blooded animal with the vector. another embodiment, such a viral vector is used for the manufacture of a medicament for immunization of a warmblooded animal against a malignancy in which the HER-2/neu oncogene is associated.

These and other aspects of the present invention will become evident upon reference to the following detailed description and attached drawings.

15 Brief Description of the Drawings

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Figure 1 shows the results of the priming of naive T lymphocytes to HER-2/neu polypeptide by dendritic Bone marrow-derived DC were generated with GM-CSF and IL6 from CD34+ stem cells. DC pulsed with HER-2/neu 20 polypeptide induced protein-specific proliferation autologous CD4+/CD45RA+ T lymphocytes after 7 days of culturing T cells with DC. Bone marrow-derived CD34+ stem cells cultured for one week in serum-free containing GM-CSF and IL-6 were used as APC. APC were plated into 96-well round-bottomed plates (Corning, Corning, NY, USA) at various concentrations and incubated for 16-18 hours with 20-25 μ g/ml of recombinant HER-2/neu polypeptide. CD4+ T lymphocytes were isolated from autologous peripheral blood mononuclear cells by positive selection using immunoaffinity columns (CellPro, Inc., Bothell, WA, USA). Antigen-pulsed APC were irradiated (10 Gy), and CD4+ T lymphocytes were added at 10^5 per well. Proliferative response of T cells was measured by the uptake of (3 H)thymidine (1μ Ci/well) added on day 7 for 1610

18 hours. Proliferation assays were performed in serumand cytokine-free medium in 5 well replicates. The symbols represent: —— DC + HER-2/neu polypeptide + CD4+/CD45RA+ T cells; —O— DC + CD4+/CD45RA+ T cells; and 5 —— DC + HER-2/neu polypeptide.

Figure 2 shows the response of CD4+ cells to HER-2/neu polypeptide. Using the priming assay described for Figure 1, CD4+ T cells from normal donors were tested for responses to recombinant human HER-2/neu polypeptide. The symbols represent: SC+CD4; and SC+CD4+HER-2/neu polypeptide. "SC" is stem cells.

Figure 3 shows that rats immunized with rat HER-2/neu polypeptide develop rat neu specific antibodies. Rats were immunized with recombinant rat HER-2/neu 15 polypeptide 25 ug in MPL or vaccel adjuvant. immunizations were given, each 20 days apart. Twenty days immunization rats were assessed after the final antibody responses to rat neu. Animals immunized with rat HER-2/neu polypeptide and the vaccel adjuvant showed high 20 titer rat *neu* specific responses. The control was an animal immunized with human HER-2/neu polypeptide (foreign protein). In separate experiments, rats immunized with 100 ug and 300 ug of purified whole rat neu did not develop detectable neu specific antibodies (data 25 Data represents the mean and standard deviation shown). polypeptide/MPL; ······●······ rat HER-2/neu polypeptide/vaccel; ---- MPL alone; ----O--- vaccel alone; and ---\$--- control. "MPL" and "vaccel" are adjuvants (Ribi, Bozeman, MT, USA). 30 "Neu" is HER-2/neu protein.

Figure 4 shows that breast cancer patients have preexistent immunity to HER-2/neu polypeptide. Patient PBMC were evaluated by tritiated thymidine incorporation

in 24 well replicates. Responsive wells are scored as greater than the mean and 3 standard deviations (372 cpm) of the control wells. This HER-2/neu positive-stage II breast cancer patient has a significant response to recombinant human HER-2/neu polypeptide. The symbols "p" represent peptides for HER-2/neu protein, "tt" represents tetanus toxoid, and "hHNP" represents recombinant human HER-2/neu polypeptide.

10 Detailed Description of the Invention

Prior to setting forth the invention, it may be helpful to an understanding thereof to set forth definitions of certain terms to be used hereinafter.

HER-2/neu polypeptide - as used herein, refers

to a portion of the HER-2/neu protein (the protein also known as p185 or c-erbB2) having the amino acid sequence of SEQ ID NO:2 from lysine, amino acid 676, through valine, amino acid 1255; and may be naturally derived, synthetically produced, genetically engineered, or a functionally equivalent variant thereof, e.g., where one or more amino acids are replaced by other amino acid(s) or non-amino acid(s) which do not substantially affect elicitation or enhancement of an immune response to HER-2/neu protein (e.g., variant stimulates a response by helper T cells or cytotoxic T cells).

Proliferation of T cells - as used herein, includes the multiplication of T cells as well as the stimulation of T cells leading to multiplication, i.e., the initiation of events leading to mitosis and mitosis 30 itself. Methods for detecting proliferation of T cells are discussed below.

As noted above, the present invention is directed toward compounds and compositions to elicit or enhance immunity to the protein product expressed by the

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HER-2/neu oncogene, including for malignancies in a warm-blooded animal wherein an amplified HER-2/neu gene is associated with the malignancies. Association of an amplified HER-2/neu gene with a malignancy does not require that the protein expression product of the gene be present on the tumor. For example, overexpression of the protein expression product may be involved with initiation of a tumor, but the protein expression may subsequently be lost. A use of the present invention is to elicit or enhance an effective autochthonous immune response to convert a HER-2/neu positive tumor to HER-2/neu negative.

More specifically, the disclosure of the present invention, in one aspect, shows that a polypeptide based on a particular portion (HER-2/neu polypeptide) of the protein expression product of the HER-2/neu gene can be 15 recognized by thymus-dependent lymphocytes (hereinafter "T cells") and, therefore, the autochthonous immune T cell response can be utilized prophylactically or to treat malignancies in which such a protein is or has been 20 overexpressed. The disclosure of the present invention also shows, in another aspect, that nucleic acid molecules directing the expression of such a peptide may be used alone or in a viral vector for immunization.

general, CD4+ Τ cell populations are considered to function as helpers/inducers through the 25 release of lymphokines when stimulated by a specific antigen; however, a subset of CD4+ cells can act as cytotoxic T lymphocytes (CTL). Similarly, CD8⁺ T cells are considered to function by directly lysing antigenic 30 targets; however, under a variety of circumstances they can secrete lymphokines to provide helper or DTH function. Despite the potential of overlapping function, and CD8 markers are phenotypic CD4 linked recognition of peptides bound to class II or class I MHC

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antigens. The recognition of antigen in the context of class II or class I MHC mandates that $CD4^+$ and $CD8^+$ T cells respond to different antigens or the same antigen presented under different circumstances. The binding of immunogenic peptides to class II MHC antigens commonly for occurs antigens ingested by presenting cells. Therefore, CD4⁺ T cells generally recognize antigens that have been external to the tumor cells. By contrast, under normal circumstances, binding 10 of peptides to class I MHC occurs only for proteins present in the cytosol and synthesized by the target itself, proteins in the external environment are excluded. An exception to this is the binding of exogenous peptides with a precise class I binding motif which are present outside the cell in high concentration. Thus, $CD4^+$ and 15 CD8+ T cells have broadly different functions and tend to recognize different antigens as a reflection of where the antigens normally reside.

As disclosed within the present invention, a 20 polypeptide portion of the protein product expressed by HER-2/neu oncogene is recognized by T cells. Circulating HER-2/neu polypeptide is degraded to peptide fragments. Peptide fragments from the polypeptide bind to major histocompatibility complex (MHC) antigens. 25 display of a peptide bound to MHC antigen on the cell surface and recognition by host T cells of the combination of peptide plus self MHC antigen, HER-2/neu polypeptide (including that expressed on a malignant cell) will be immunogenic to T cells. The exquisite specificity of the 30 T cell receptor enables individual T cells to discriminate between peptides which differ by a single amino acid residue.

During the immune response to a peptide fragment from the polypeptide, T cells expressing a T cell receptor

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with high affinity binding of the peptide-MHC complex will the peptide-MHC complex and thereby become activated and induced to proliferate. In the encounter with a peptide, small numbers of immune T cells will secrete lymphokines, proliferate and differentiate into effector and memory T cells. The primary immune response will occur in vivo but has been difficult detect in vitro. Subsequent encounter with the same antigen by the memory T cell will lead to a faster and 10 more intense immune response. The secondary response will occur either in vivo or in vitro. The in vitro response is easily gauged by measuring the degree of proliferation, the degree of cytokine production, or the generation of cytolytic activity of the T cell population re-exposed in 15 the antigen. Substantial proliferation of the T cell population in response to a particular antigen considered to be indicative of prior exposure or priming to the antigen.

compounds of this invention 20 comprise HER-2/neu polypeptides or DNA molecules that direct the expression of such peptides, wherein the DNA molecules may be present in a viral vector. above, the polypeptides of the present invention include variants of the polypeptide of SEQ ID NO:2 from amino acid 676 through amino acid 1255, that retain the ability to 25 stimulate an immune response. Such variants include various structural forms of the native polypeptide. to the presence of ionizable amino and carboxyl groups, for example, a HER-2/neu polypeptide may be in the form of 30 an acidic or basic salt, or may be in neutral form. Individual amino acid residues may also be modified by oxidation or reduction.

Variants within the scope of this invention also include polypeptides in which the primary amino acid

structure native HER-2/neu polypeptide is modified by forming covalent or aggregative conjugates with other peptides or polypeptides, or chemical moieties such as glycosyl groups, lipids, phosphate, acetyl groups and the like. Covalent derivatives may be prepared, for example, by linking particular functional groups to amino acid side chains or at the N- or C-terminus.

The present invention also includes HER-2/neu polypeptides with or without glycosylation. Polypeptides 10 expressed in yeast or mammalian expression systems may be similar to or slightly different in molecular weight and glycosylation pattern than the native molecules, depending upon the expression system. For instance, expression of DNA encoding polypeptides in bacteria such as E. 15 typically provides non-glycosylated molecules. glycosylation sites of eukaryotic proteins characterized by the amino acid triplet $Asn-A_1-Z$, where A_1 is any amino acid except Pro, and Z is Ser or Variants of HER-2/neu polypeptides having inactivated N-20 glycosylation sites can be produced by techniques known to ordinary skill in the art, oligonucleotide synthesis and ligation or site-specific mutagenesis techniques, and are within the scope of this invention. Alternatively, N-linked glycosylation sites 25 can be added to a HER-2/new polypeptide.

The polypeptides of this invention also include variants of the SEQ ID NO:2 polypeptide (i.e., variants of a polypeptide having the amino acid sequence of SEQ ID NO:2 from amino acid 676 through amino acid 1255) that have an amino acid sequence different from this sequence because of one or more deletions, insertions, substitutions or other modifications. In one embodiment, such variants are substantially homologous to the native HER-2/new polypeptide and retain the ability to stimulate

immune response. an "Substantial homology," as used herein, refers to amino acid sequences that may be encoded by DNA sequences that are capable of hybridizing under moderately stringent conditions to a nucleotide sequence complimentary to a naturally occurring DNA encoding the specified polypeptide portion of SEQ ID NO:2herein (i.e., nucleotides 2026 through 3765 of SEQ ID NO:1). Suitable moderately stringent conditions include prewashing in a solution of 5 X SSC, 0.5% SDS, 1.0 mM EDTA (pH 8.0); hybridizing at 50°C-65°C, 5 X SSC, overnight; 10 followed by washing twice at 65°C for 20 minutes with each of 2X, 0.5X and 0.2X SSC (containing 0.1% SDS). hybridizing DNA sequences are also within the scope of this invention. The effect of any such modifications on the ability of a HER-2/neu polypeptide to produce 15 immune response may be readily determined (e.g., analyzing the ability of the mutated HER-2/neu polypeptide induce a T cell response using, for example, the methods described herein).

20 Generally, amino acid substitutions may be made in a variety of ways to provide other embodiments of within variants the present invention. First, for example, amino acid substitutions may be conservatively; i.e., a substitute amino acid replaces an amino acid that has similar properties, such that one 25 skilled in the art of peptide chemistry would expect the secondary structure and hydropathic nature polypeptide to be substantially unchanged. In general, the following groups of amino acids represent conservative changes: (1) ala, pro, gly, glu, asp, gln, asn, ser, thr; 30 (2) cys, ser, tyr, thr; (3) val, ile, leu, met, ala, phe; (4) lys, arg, his; and (5) phe, tyr, trp, his. An example of a non-conservative change is to replace an amino acid of one group with an amino acid from another group.

Another way to make amino acid substitutions to produce variants of the present invention is to identify and replace amino acids in T cell motifs with potential to bind to class II MHC molecules (for CD4+ T cell response) or class I MHC molecules (for CD8+ T cell response). Peptide segments (of a HER-2/neu polypeptide) with a motif with theoretical potential to bind to class II molecules may be identified by computer analysis. For example, a protein sequence analysis package, T Sites, 10 that incorporates several computer algorithms designed to distinguish potential sites for T cell recognition can be used (Feller and de la Cruz, Nature 349:720-721, searching algorithms are used: (1) the algorithm described by Margalit (Feller and de la Cruz, Nature 349:720-721, 1991; Margalit et al., J. 15 138:2213-2229, 1987) identifies epitope motifs according to alpha-helical periodicity and amphipathicity; (2) the Rothbard and Taylor algorithm identifies epitope motifs according to charge and polarity pattern (Rothbard and 20 Taylor, EMBO 7:93-100, 1988). Segments with both motifs most appropriate for binding to class CD8+ T cells recognize peptide bound to class molecules. MHC molecules. Falk et al. have determined that peptides binding to particular MHC molecules share 25 discernible sequence motifs (Falk et al., Nature 351:290-296, 1991). A peptide motif for binding in the groove of HLA-A2.1 has been defined by Edman degradation of peptides stripped from HLA-A2.1 molecules of a cultured cell line (Table 2, from Falk et al., supra). The method identified 30 the typical or average HLA-A2.1 binding peptide as being 9 amino acids in length with dominant anchor residues occurring at positions 2 (L) and 9 (V). Commonly occurring strong binding residues have been identified at positions 2 (M), 4 (E,K), 6 (V), and 8 (K).

identified motif represents the average of many binding peptides.

The HLA-A2.1 Restricted Motif

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	Amino Acid Position 1 2 3 4 5 6 7 8 9	Point Assignment
Dominant Binding Anchor Residue	L V	+3
Strong Binding Residue	M E V K K	+2
Weak Binding	I AGIIAEL	+1
Residue	L YPKLYS	
	F FDYTH	
	K PTN	
	M M G	
	Y S V	
	Н	

The derived peptide motif as currently defined is not particularly stringent. Some HLA-A2.1 binding peptides do not contain both dominant anchor residues and the amino acids flanking the dominant anchor residues play major roles in allowing or disallowing binding. Not every peptide with the current described binding motif will bind, and some peptides without the motif will bind. However, the current motif is valid enough to allow identification of some peptides capable of binding. Of note, the current HLA-A2.1 motif places 6 amino acids between the dominant anchor amino acids at residues 2 and 9.

Following identification of peptide motifs 20 within a HER-2/neu polypeptide, amino acid substitutions may be made conservatively or non-conservatively. The latter type of substitutions are intended to produce an improved polypeptide that is more potent and/or more

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broadly cross-reactive (MHC polymorphism). An example of a more potent polypeptide is one that binds with higher affinity to the same MHC molecule as natural polypeptide, without affecting recognition by T cells specific for 5 natural polypeptide. An example of a polypeptide with broader cross-reactivity is one that induces more broadly cross-reactive immune responses (i.e., binds to a greater range of MHC molecules) than natural polypeptide. Similarly, one or more amino acids residing between peptide motifs and having a spacer function (e.g., do not interact with a MHC molecule or T cell receptor) may be substituted conservatively or non-conservatively. It will be evident to those of ordinary skill in the art that polypeptides containing one or more amino acid 15 substitutions may be tested for beneficial or adverse immunological interactions by a variety of assays, including those described herein for the ability to stimulate T cell recognition.

Variants within the scope of this invention may 20 also, or alternatively, contain other modifications, including the deletion or addition of amino acids, that have minimal influence on the desired immunological properties of the polypeptide. It will be appreciated by those of ordinary skill in the art that truncated forms or 25 non-native extended forms of a HER-2/neu polypeptide may be used, provided the desired immunological properties are at least roughly equivalent to that of full length, native HER-2/neu polypeptide. Cysteine residues may be deleted or replaced with other amino acids to prevent formation of 30 incorrect intramolecular disulfide bridges upon renaturation. Other approaches to mutagenesis involve modification of adjacent dibasic amino acid residues to enhance expression in yeast systems in which KEX2 protease activity is present.

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HER-2/neu polypeptide may generally obtained using a genomic or cDNA clone encoding protein. A genomic sequence that encodes full length HER-2/neu is shown in SEQ ID NO:1, and the deduced amino acid sequence is presented in SEQ ID NO:2. Such clones may be isolated by screening an appropriate expression library for clones that express HER-2/neu protein. library preparation and screen may generally be performed using methods known to those of ordinary skill in the art, such as methods described in Sambrook et al., Molecular 10 Cloning: A Laboratory Manual, Cold Spring Laboratories, Cold Spring Harbor, N.Y., 1989, which is incorporated herein by reference. Briefly, bacteriophage expression library may be plated 15 transferred to filters. The filters may then be incubated with a detection reagent. In the context of this invention, a "detection reagent" is any compound capable binding to HER-2/neu protein, which may then detected by any of a variety of means known to those of 20 ordinary skill in the art. Typical detection reagents contain a "binding agent," such as Protein A, Protein G, IgG or a lectin, coupled to a reporter group. Preferred reporter groups include enzymes, substrates, cofactors, inhibitors, dyes, radionuclides, luminescent 25 fluorescent groups and biotin. More preferably, reporter group is horseradish peroxidase, which may detected by incubation with a substrate tetramethylbenzidine or 2,2'-azino-di-3-ethylbenzthiazoline sulfonic acid. Plaques containing genomic or 30 cDNA sequences that express HER-2/neu protein are isolated and purified by techniques known to those of ordinary skill in the art. Appropriate methods may be found, for Sambrook et al., Molecular Cloning: A example, in

Laboratory Manual, Cold Spring Harbor Laboratories, Cold Spring Harbor, N.Y., 1989.

Variants of the polypeptide that retain the ability to stimulate an immune response may generally be identified by modifying the sequence in one or more of the aspects described above and assaying the resulting polypeptide for the ability to stimulate an response, e.g., a T cell response. For example, such assays may generally be performed by contacting T cells 10 with the modified polypeptide and assaying the response. Naturally occurring variants of the polypeptide may also be isolated by, for example, screening an appropriate cDNA genomic library with a DNA sequence encoding the polypeptide or a variant thereof.

The above-described sequence modifications may be introduced using standard recombinant techniques or by automated synthesis of the modified polypeptide. For example, mutations can be introduced at particular loci by synthesizing oligonucleotides containing a mutant sequence, flanked by restriction sites enabling ligation to fragments of the native sequence. Following ligation, the resulting reconstructed sequence encodes an analogue having the desired amino acid insertion, substitution, or deletion.

Alternatively, oligonucleotide-directed sitespecific mutagenesis procedures can be employed to provide
a gene in which particular codons are altered according to
the substitution, deletion, or insertion required.
Exemplary methods of making the alterations set forth
above are disclosed by Walder et al., Gene 42:133, 1986;
Bauer et al., Gene 37:73, 1985; Craik, BioTechniques,
January 1985, 12-19; Smith et al., Genetic Engineering:
Principles and Methods, Plenum Press, 1981; and U.S.
Patent Nos. 4,518,584 and 4,737,462.

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Mutations in nucleotide sequences constructed for expression of such HER-2/neu polypeptides must, of course, preserve the reading frame of the coding sequences and preferably will not create complementary regions that could hybridize to produce secondary mRNA structures, such loops or hairpins, which would adversely affect translation of the mRNA. Although a mutation site may be predetermined, it is not necessary that the nature of the mutation per se be predetermined. For example, in order 10 to select for optimum characteristics of mutants at a given site, random mutagenesis may be conducted at the target codon and the expressed HER-2/neu polypeptide mutants screened for the desired activity.

Not all mutations in a nucleotide sequence which encodes a HER-2/neu polypeptide will be expressed in the final product. For example, nucleotide substitutions may be made to enhance expression, primarily to avoid secondary structure loops in the transcribed mRNA (see, e.g., European Patent Application 75,444A), or to provide codons that are more readily translated by the selected host, such as the well-known *E. coli* preference codons for *E. coli* expression.

The polypeptides of the present invention, both naturally occurring and modified, are preferably produced 25 by recombinant DNA methods. Such methods inserting a DNA sequence encoding a HER-2/neu polypeptide into a recombinant expression vector and expressing the DNA sequence in a recombinant microbial, mammalian or insect cell expression system under conditions promoting 30 expression. DNA sequences encoding the polypeptides provided by this invention can be assembled from cDNA fragments and short oligonucleotide linkers, or from a series of oligonucleotides, to provide a synthetic gene which is capable of being inserted in a recombinant

expression vector and expressed in a recombinant transcriptional unit.

Recombinant expression vectors contain a DNA sequence encoding a HER-2/neu polypeptide operably linked to suitable transcriptional or translational regulatory elements derived from mammalian, microbial, viral insect genes. Such regulatory elements include transcriptional promoter, an optional operator sequence to control transcription, a sequence encoding suitable mRNA 10 ribosomal binding sites, and sequences which control the termination of transcription and translation. An origin replication and a selectable marker to facilitate recognition of transformants may additionally incorporated.

15 DNA regions are operably linked when they are functionally related to each other. For example, DNA for a signal peptide (secretory leader) is operably linked to DNA for a polypeptide if it is expressed as a precursor which participates in the secretion of the polypeptide; a 20 promoter is operably linked to a coding sequence if it controls the transcription of the sequence; or a ribosome binding site is operably linked to a coding sequence if it is positioned so as to permit translation. Generally, operably linked means contiguous and, in the case of 25 secretory leaders, in reading frame. DNA encoding HER-2/neu polypeptides which are to be expressed in a microorganism will preferably contain no introns that could prematurely terminate transcription of DNA mRNA.

Expression vectors for bacterial use may comprise a selectable marker and bacterial origin of replication derived from commercially available plasmids comprising genetic elements of the well known cloning vector pBR322 (ATCC 37017). Such commercial vectors

include, for example, pKK223-3 (Pharmacia Fine Chemicals, Uppsala, Sweden) and pGEM1 (Promega Biotec, Madison, WI, USA). These pBR322 "backbone" sections are combined with an appropriate promoter and the structural sequence to be expressed. E. coli is typically transformed using derivatives of pBR322, a plasmid derived from an E. coli species (Bolivar et al., Gene 2:95, 1977). pBR322 contains genes for ampicillin and tetracycline resistance and thus provides simple means for identifying transformed cells.

10 Promoters commonly used in recombinant microbial expression vectors include the β -lactamase (penicillinase) and lactose promoter system (Chang et al., Nature 275:615, 1978; and Goeddel et al., Nature 281:544, 1979), tryptophan (trp) promoter system (Goeddel et al., Nucl. Acids Res. 8:4057, 1980; and European Patent Application 15 36,776) and the tac promoter (Maniatis, Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory, p.412, 1982). A particularly useful bacterial expression system employs the phage λ P_L promoter and cI857ts thermolabile 20 Plasmid vectors available from the American repressor. Type Culture Collection which incorporate derivatives of the λ P_L promoter include plasmid pHUB2, resident in E. coli strain JMB9 (ATCC 37092) and pPLc28, resident in E. coli RR1 (ATCC 53082).

25 Suitable promoter sequences in yeast vectors include the promoters for metallothionein, phosphoglycerate kinase (Hitzeman et al., J. Biol. Chem. 255:2073, 1980) or other glycolytic enzymes (Hess et al., J. Adv. Enzyme Reg. 7:149, 1968; and Holland et Biochem. 17:4900, 1978), such as enolase, glyceraldehyde-30 3-phosphate dehydrogenase, hexokinase, decarboxylase, phosphofructokinase, glucose-6-phosphate isomerase, 3-phosphoglycerate mutase, pyruvate

triosephosphate isomerase, phosphoglucose isomerase, and glucokinase. Suitable vectors and promoters for use in yeast expression are further described in R. Hitzeman et al., European Patent Application 73,657.

5 Preferred yeast vectors can be assembled using DNA sequences from pBR322 for selection and replication in E. coli (Ampr gene and origin of replication) and yeast DNA sequences including a glucose-repressible ADH2 promoter and α -factor secretion leader. The ADH2 promoter has been 10 described by Russell et al. (J. Biol. Chem. 258:2674, 1982) and Beier et al. (Nature 300:724, 1982). The yeast α -factor leader, which directs secretion of heterologous proteins, can be inserted between the promoter and the structural gene to be expressed (see, e.g., Kurjan et al., 15 Cell 30:933, 1982; and Bitter et al., Proc. Natl. Sci. USA 81:5330, 1984). The leader sequence may be modified to contain, near its 3' end, one or more useful restriction sites to facilitate fusion of the leader sequence to foreign genes. The transcriptional and 20 translational control sequences in expression vectors to be used in transforming vertebrate cells may be provided by viral sources. For example, commonly used promoters and enhancers are derived from polyoma, adenovirus simian virus 40 (SV40), and human cytomegalovirus. DNA sequences derived from the SV40 viral genome, for example, SV40 origin, early and late promoter, enhancer, splice, and polyadenylation sites may be used to provide the other genetic elements required for expression of a heterologous early and late promoters sequence. The particularly useful because both are obtained easily from 30 the virus as a fragment which also contains the SV40 viral origin of replication (Fiers et al., Nature 273:113, 1978). Smaller or larger SV40 fragments may also be used, provided the approximately 250 bp sequence extending from

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the Hind III site toward the Bgl II site located in the viral origin of replication is included. Further, viral genomic promoter, control and/or signal sequences may be utilized, provided such control sequences are compatible with the host cell chosen. Exemplary vectors can be constructed as disclosed by Okayama and Berg, Mol. Cell. Biol. 3:280, 1983.

A useful system for stable high level expression of mammalian receptor cDNAs in C127 murine epithelial cells can be constructed substantially described by Cosman et al. (Mol. Immunol. 23:935, 1986). A preferred eukaryotic vector for expression of LbeIF4A protein DNA is pDC406 (McMahan et al., EMBO J. 10:2821, and includes regulatory sequences derived from SV40, human immunodeficiency virus (HIV), and Epstein-Barr 15 virus (EBV). Other preferred vectors include pDC409 and pDC410, which are derived from pDC406. pDC410 was derived from pDC406 by substituting the EBV origin of replication with sequences encoding the SV40 large T antigen. pDC409 differs from pDC406 in that a Bgl II restriction site 20 outside of the multiple cloning site has been deleted, making the Bgl II site within the multiple cloning site unique.

A useful cell line that allows for episomal replication of expression vectors, such as pDC406 and pDC409, which contain the EBV origin of replication, is CV-1/EBNA (ATCC CRL 10478). The CV-L/EBNA cell line was derived by transfection of the CV-1 cell line with a gene encoding Epstein-Barr virus nuclear antigen-I (EBNA-1) and constitutively express EBNA-1 driven from human CMV immediate-early enhancer/promoter.

Transformed host cells are cells which have been transformed or transfected with expression vectors constructed using recombinant DNA techniques and which

contain sequences encoding a HER-2/neu polypeptide of the present invention. Transformed host cells may express the desired HER-2/neu polypeptide, but host cells transformed for purposes of cloning or amplifying HER-2/neu DNA do not 5 need to express the HER-2/neu polypeptide. polypeptides will preferably be secreted into the culture supernatant, depending on the DNA selected, but may also be deposited in the cell membrane.

Suitable host for expression cells of 10 recombinant proteins include prokaryotes, yeast or higher eukaryotic cells under the control of appropriate promoters. Prokaryotes include gram negative or gram positive organisms, for example E. coli or Bacilli. Higher eukaryotic cells include established cell lines of 15 insect or mammalian origin as described below. Cell-free translation systems could also be employed to produce HER-2/neu polypeptides using RNAs derived from DNA Appropriate cloning and expression vectors constructs. for use with bacterial, fungal, yeast, and mammalian 20 cellular hosts are described, for example, by Pouwels et al., Cloning Vectors: A Laboratory Manual, Elsevier, New York, 1985.

Prokaryotic expression hosts may be used for expression of HER-2/neu polypeptides that do not require 25 extensive proteolytic disulfide and processing. Prokaryotic expression vectors generally comprise one or more phenotypic selectable markers, for example a gene encoding proteins conferring antibiotic resistance supplying an autotrophic requirement, and an origin of replication recognized by the host to ensure amplification host. Suitable prokaryotic hosts within the transformation include E. coli, Bacillus subtilis, Salmonella typhimurium, and various species within the

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genera *Pseudomonas*, *Streptomyces*, and *Staphylococcus*, although other hosts may also be employed.

Recombinant HER-2/neu polypeptides may also be expressed in yeast hosts, preferably from the Saccharomyces species, such as S. cerevisiae. other genera, such as Pichia or Kluyveromyces may also be employed. Yeast vectors will generally contain an origin replication from the 2μ yeast plasmid autonomously replicating sequence (ARS), a promoter, DNA 10 encoding HER-2/neu polypeptide, sequences the polyadenylation and transcription termination selection gene. Preferably, yeast vectors will include an origin of replication and selectable marker permitting transformation of both yeast and E. coli, e.g., 15 ampicillin resistance gene of E. coli and the cerevisiae trpl gene, which provides a selection marker for a mutant strain of yeast lacking the ability to grow in tryptophan, and a promoter derived from a highly expressed yeast gene to induce transcription of 20 structural sequence downstream. The presence of the trpl lesion in the yeast host cell genome then provides effective environment for detecting transformation growth in the absence of tryptophan.

Suitable yeast transformation protocols 25 known to those of skill in the art. An exemplary technique described by Hind et al. (Proc. Natl. Acad. Sci. USA 75:1929, 1978), involves selecting for transformants in a selective medium consisting of 0.67%yeast nitrogen base, 0.5% casamino acids, 2% glucose, 10 30 mg/ml adenine and 20 mg/ml uracil. Host transformed by vectors comprising the ADH2 promoter may be grown for expression in a rich medium consisting of 1%yeast extract, 2% peptone, and 1% glucose supplemented with 80 mg/ml adenine and 80 mg/ml uracil. Derepression

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of the ADH2 promoter occurs upon exhaustion of medium glucose. Crude yeast supernatants are harvested by filtration and held at 4°C prior to further purification.

Various mammalian or insect (e.g., Spodoptera or 5 Trichoplusia) cell culture systems can also be employed to express recombinant polypeptide. Baculovirus systems for production of heterologous polypeptides in insect cells reviewed, for example, by Luckow and Bio/Technology 6:47, 1988. Examples of suitable mammalian host cell lines include the COS-7 lines of monkey kidney 10 cells, described by Gluzman (Cell 23:175, 1981), and other cell lines capable of expressing an appropriate vector including, for example, CV-1/EBNA (ATCC CRL 10478), L cells, C127, 3T3, Chinese hamster ovary (CHO), COS, NS-1, Mammalian expression vectors may 15 HeLa and BHK cell lines. comprise nontranscribed elements such as an origin replication, a suitable promoter and enhancer linked to the gene to be expressed, and other 5' or 3' flanking nontranscribed sequences, and 5' or 3' nontranslated sequences, such as necessary ribosome binding sites, a 20 polyadenylation site, splice donor and acceptor sites, and transcriptional termination sequences.

Purified HER-2/new polypeptides may be prepared by culturing suitable host/vector systems to express the recombinant translation products of the DNAs of the present invention, which are then purified from culture media or cell extracts. For example, supernatants from systems which secrete recombinant polypeptide into culture media may be first concentrated using a commercially available protein concentration filter, such as an Amicon or Millipore Pellicon ultrafiltration unit. Following the concentration step, the concentrate may be applied to a suitable purification matrix. For example, a suitable affinity matrix may comprise a counter structure protein

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(i.e., a protein to which a HER-2/new polypeptide binds in a specific interaction based on structure) or lectin or antibody molecule bound to а suitable Alternatively, an anion exchange resin can be employed, example, a matrix or substrate having diethylaminoethyl (DEAE) groups. The matrices can be acrylamide, agarose, dextran, cellulose or other types commonly employed in protein purification. Alternatively, a cation exchange step can be employed. Suitable cation exchangers include various insoluble matrices comprising 10 sulfopropyl or carboxymethyl groups. Sulfopropyl groups preferred. Gel filtration chromatography also provides a means of purifying a HER-2/neu.

Affinity chromatography is a preferred method of purifying HER-2/neu polypeptides. For example, monoclonal antibodies against the HER-2/neu polypeptide may also be useful in affinity chromatography purification, by utilizing methods that are well-known in the art.

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Finally, one or more reverse-phase 20 performance liquid chromatography (RP-HPLC) steps employing hydrophobic RP-HPLC media (e.g., silica gel having pendant methyl or other aliphatic groups) may be employed to further purify a HER-2/neu polypeptide composition. Some or all of the foregoing purification steps, in various combinations, can also be employed to 25 provide a homogeneous recombinant polypeptide.

Recombinant HER-2/neu polypeptide produced in bacterial culture is preferably isolated by initial extraction from cell pellets, followed by one or more concentration, salting-out, aqueous ion exchange or size exclusion chromatography steps. High performance liquid chromatography (HPLC) may be employed for final purification steps. Microbial cells employed in expression of recombinant LbeIF4A protein can be disrupted by any

convenient method, including freeze-thaw cycling, sonication, mechanical disruption, or use of cell lysing agents.

Fermentation of yeast which express HER-2/neu 5 polypeptide as a secreted protein greatly simplifies purification. Secreted recombinant protein resulting from a large-scale fermentation can be purified by methods analogous to those disclosed by Urdal et al. (*J. Chromatog. 296:*171, 1984). This reference describes two sequential, reverse-phase HPLC steps for purification of recombinant human GM-CSF on a preparative HPLC column.

Preparations of HER-2/neu polypeptides in recombinant culture may contain nonsynthesized HER-2/neu cell components, including proteins, in amounts and of a character which depend upon the purification steps taken to recover the HER-2/neu polypeptide from the culture. These components ordinarily will be of yeast, prokaryotic or non-human eukaryotic origin. preparations are typically free of other proteins which may be normally associated with the HER-2/neu protein as it is found in nature in its species of origin.

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Automated synthesis provides an alternate method preparing polypeptides of this invention. example, any of the commercially available solid-phase techniques may be employed, such as the Merrifield solid method, in which amino acids are phase synthesis sequentially added to a growing amino acid chain. Chem. Soc. 85:2149-2146, Merrifield, J. Am. Equipment for automated synthesis of polypeptides commercially available from suppliers such as Biosystems, Inc. of Foster City, CA, and may generally be operated according to the manufacturer's instructions.

Within one aspect of the present invention, use of a HER-2/new polypeptide (or a DNA molecule that directs

the expression of such a peptide) to generate an immune to the HER-2/neu protein response (including expressed on a malignancy in which a HER-2/neu oncogene is associated) may be detected. Representative examples of such malignancies include breast, ovarian, colon, lung and prostate cancers. An immune response to the HER-2/neu protein, once generated by a HER-2/neu polypeptide, can be long-lived and can be detected long after immunization, regardless of whether the protein is present or absent in the body at the time of testing. An immune response to 10 the HER-2/neu protein generated by reaction to a HER-2/neupolypeptide can be detected by examining for the presence or absence, or enhancement, of specific activation of $CD4^+$ or CD8 $^+$ T cells. More specifically, T cells isolated from an immunized individual by routine techniques (such as $\ensuremath{\text{b}} \ensuremath{\text{y}}$ 15 Ficoll/Hypaque density gradient centrifugation peripheral blood lymphocytes) are incubated with ${\tt HER-2/neu}$ protein. For example, T cells may be incubated in vitro for 2-9 days (typically 4 days) at 37°C with HER-2/neu protein (typically, 5 $\mu g/ml$ of whole protein or graded 20 numbers of cells synthesizing HER-2/neu protein). be desirable to incubate another aliquot of a T cell sample in the absence of ${\tt HER-2/neu}$ protein to serve as a control.

25 Specific activation of CD4+ or CD8+ T cells may be detected in a variety of ways. Methods for detecting specific T cell activation include detecting proliferation of T cells, the production of cytokines lymphokines), or the generation of cytolytic 30 activity (i.e., generation of cytotoxic T cells specific for HER-2/neu protein). For $CD4^+$ T cells, a preferred method for detecting specific T cell activation is the detection of the proliferation of T cells. T cells, a preferred method for detecting specific T cell activation is the detection of the generation of cytolytic activity.

Detection of the proliferation of T cells may be accomplished by a variety of known techniques. example, T cell proliferation can be detected by measuring the rate of DNA synthesis. T cells which have been stimulated to proliferate exhibit an increased rate of DNA synthesis. A typical way to measure the rate of DNA synthesis is, for example, by pulse-labeling cultures of T cells with tritiated thymidine, a nucleoside precursor 10 which is incorporated into newly synthesized DNA. amount of tritiated thymidine incorporated determined using a liquid scintillation spectrophotometer. proliferation ways to detect T cell 15 measuring increases in interleukin-2 (IL-2) production, Ca^{2+} flux, or dye uptake, such as 3-(4,5-dimethylthiazol-2yl)-2,5-diphenyl-tetrazolium. Alternatively, synthesis of lymphokines (such as interferon-gamma) can be measured or the relative number of T cells that can respond to intact p185HER-2/new protein may be quantified. 20

By use or expression of a HER-2/neu polypeptide, T cells which recognize the HER-2/neu protein can For example, a medicament proliferated in vivo. immunization with a HER-2/new peptide (i.e., as a vaccine) can induce continued expansion in the number of T cells necessary for therapeutic attack against a tumor in which Typically, about the HER-2/neu oncogene is associated. $0.01 \, \mu g/kg$ to about $100 \, mg/kg$ body weight will intradermal, subcutaneous administered by the intravenous route. A preferred dosage is about 1 μ g/kg to about 1 mg/kg, with about 5 μ g/kg to about 200 μ g/kg It will be evident to those particularly preferred. skilled in the art that the number and frequency of administration will be dependent upon the response of the

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patient. It may be desirable to administer the HER-2/neu polypeptide repetitively. It will be evident to those skilled in this art that more one HER-2/neu than polypeptide may be administered, either simultaneously or Preferred peptides for use in a medicament sequentially. for immunization are those that include the amino acid sequence of SEQ ID NO:2 beginning at about the lysine residue at amino acid position 676 and extending to about the valine residue at amino acid position 1255. 10 be appreciated by those in the art that the present invention contemplates the use of an intact HER-2/neu polypeptide as well as division of such a polypeptide into Neither intact p185HER-2/neu plurality of peptides. protein nor a peptide having the amino acid sequence of its entire extracellular domain (i.e., a peptide having an 15 amino acid sequence of SEQ ID NO:2 from amino position 1 up to amino acid position 650, plus or minus about one to five positions, and with or without the first 21 amino acid positions) are used alone for immunization.

20 A HER-2/neu polypeptide (or nucleic acid) is preferably formulated for use in the above methods as a pharmaceutical composition (e.g., Pharmaceutical compositions generally comprise one or more polypeptides in combination with a pharmaceutically 25 acceptable carrier, excipient or diluent. Such carriers be nontoxic to recipients at the dosages concentrations employed. The use of HER-2/neu polypeptide in conjunction with chemotherapeutic agents is also contemplated.

In addition to the HER-2/neu polypeptide (which functions as an antigen), it may be desirable to include other components in the vaccine, such as a vehicle for antigen delivery and immunostimulatory substances designed to enhance the protein's immunogenicity. Examples of

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vehicles for antigen delivery include aluminum salts, water-in-oil emulsions, biodegradable oil vehicles, oilin-water emulsions, biodegradable microcapsules, liposomes. Examples of immunostimulatory substances (adjuvants) include N-acetylmuramyl-L-alanine-Dlipopoly-saccharides (LPS), isoglutamine (MDP), IL-12, GM-CSF, gamma interferon and IL-15. It will be evident to those of ordinary skill in this art that a HERpolypeptide for a vaccine may be prepared synthetically or be naturally derived.

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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 15 whether a sustained release is desired. 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, 20 lactose, starch, magnesium stearate, sodium saccharine, talcum, cellulose, glucose, sucrose, and magnesium carbonate, may be employed. Biodegradable microspheres (e.g., polylactic galactide) may also be employed carriers for the pharmaceutical compositions of this 25 Suitable biodegradable microspheres invention. disclosed, for example, in U.S. Patent Nos. 4,897,268 and 5,075,109. A HER-2/neu polypeptide may be encapsulated within the biodegradable microsphere or associated with the surface of the microsphere. For example, in preferred embodiment, a polypeptide having the amino acid sequence of SEQ ID NO:2 from amino acid 676 through amino acid 1255 is encapsulated within a biodegradable microsphere. In this regard, it is preferable that the microsphere be larger than approximately 25 microns.

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Pharmaceutical compositions (including vaccines) may also contain diluents such as buffers, antioxidants such as ascorbic acid, low molecular weight (less than about 10 residues) polypeptides, proteins, amino acids, carbohydrates including glucose, sucrose or dextrins, chelating agents such as EDTA, glutathione and other stabilizers and excipients. Neutral buffered saline or saline mixed with nonspecific serum albumin are exemplary appropriate diluents. Preferably, product is formulated as a lyophilizate using appropriate excipient solutions (e.g., sucrose) as diluents.

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As alternative to an the presentation HER-2/neu polypeptides, the subject invention includes compositions capable of delivering nucleic acid molecules 15 encoding a HER-2/neu polypeptide. Such compositions include recombinant viral vectors (e.g., retroviruses (see WO 90/07936, WO 91/02805, WO 93/25234, WO 93/25698, and WO 94/03622), adenovirus (see Berkner, Biotechniques 6:616-627, 1988; Li et al., Hum. Gene Ther. 4:403-409, 1993; 20 Vincent et al., Nat. Genet. 5:130-134, 1993; and Kolls et al., Proc. Natl. Acad. Sci. USA 91:215-219, 1994), pox virus (see U.S. Patent No. 4,769,330; U.S. Patent No. 5,017,487; and WO 89/01973)), naked DNA (see WO 90/11092), nucleic acid molecule complexed to a polycationic molecule (see WO 93/03709), and nucleic acid associated with 25 liposomes (see Wang et al., Proc. Natl. Acad. Sci. 84:7851, 1987). In certain embodiments, the DNA may be linked to killed or inactivated adenovirus (see Curiel et al., Hum. Gene Ther. 3:147-154, 1992; Cotton et al., Proc. 30 Natl. Acad. Sci. USA 89:6094, 1992). Other suitable compositions include DNA-ligand (see Wu et al., J. Biol. Chem. 264:16985-16987, 1989) and lipid-DNA combinations (see Felgner et al., Proc. Natl. Acad. Sci. USA 84:7413-7417, 1989). In addition, the efficiency of naked DNA

uptake into cells may be increased by coating the DNA onto biodegradable beads.

In addition to direct in vivo procedures, vivo procedures may be used in which cells are removed from an animal, modified, and placed into the same or another animal. It will be evident that one can utilize any of the compositions noted above for introduction of HER-2/neu nucleic acid molecules into tissue cells in an ex vivo context. Protocols for viral, physical chemical methods of uptake are well known in the art.

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Accordingly, the present invention is useful for enhancing or eliciting, in a patient or cell culture, a cellular immune response (e.g., the generation of antigenspecific cytolytic T cells). As used herein, the term "patient" refers to any warm-blooded animal, preferably a human. A patient may be afflicted with cancer, such as breast cancer, or may be normal (i.e., free of detectable disease and infection). A "cell culture" is T cells or isolated component cells preparation of 20 (including, but not limited to, macrophages, monocytes, B cells and dendritic cells). Such cells may be isolated by any of a variety of techniques well known to those of ordinary skill in the art (such as Ficoll-hypaque density centrifugation). The cells may (but need not) have been isolated from a patient afflicted with a HER-2/neu associated malignancy, and may be reintroduced into a patient after treatment.

The present invention also discloses that HER-2/neu polypeptide, in addition to being immunogenic to 30 T cells, appears to stimulate B-cells to antibodies capable of recognizing HER-2/neu polypeptide. Antibodies specific (i.e., which exhibit a binding affinity of about 10⁷ liters/mole or better) for HER-2/neu protein may be found in a variety of body fluids including

sera and ascites. Briefly, a body fluid sample is isolated from a warm-blooded animal, such as a human, for is desired to determine whether whom it antibodies specific for ${\tt HER-2/\it neu}$ polypeptide are present. The body incubated with HER-2/neu polypeptide conditions and for a time sufficient to permit immunocomplexes to form between the polypeptide antibodies specific for the protein. For example, a body fluid and HER-2/neu polypeptide may be incubated at 4°C for 24-48 hours. Following the incubation, the reaction 10 mixture is tested for the presence of immunocomplexes. Detection of one or more immunocomplexes formed between ${\tt HER-2/\it neu}$ polypeptide and antibodies specific for ${\tt HER-1}$ 2/neu polypeptide may be accomplished by a variety of known techniques, such as radioimmunoassays (RIA) and 15 enzyme linked immunosorbent assays (ELISA).

Suitable immunoassays include the double monoclonal antibody sandwich immunoassay technique of David et al. (U.S. Patent 4,376,110); monoclonal-20 polyclonal antibody sandwich assays (Wide et al., Kirkham and Hunter, eds., Radioimmunoassay Methods, E. and S. Livingstone, Edinburgh, 1970); the "western blot" method of Gordon et al. (U.S. Patent 4,452,901); immunoprecipitation of labeled ligand (Brown et al., 25 J. Biol. Chem. 255:4980-4983, 1980); enzyme-linked immunosorbent assays as described by, for example, Raines Chem. Ross (J. Biol. *257*:5154-5160, 1982); immunocytochemical techniques, including the use of fluorochromes (Brooks et al., Clin. Exp. Immunol. 39: 477, 1980); and neutralization of activity [Bowen-Pope et al., 30 Proc. Natl. Acad. Sci. USA 81:2396-2400 (1984)], all of which are hereby incorporated by reference. In addition to the immunoassays described above, a number of other immunoassays are available, including those described in U.S. Patent Nos.: 3,817,827; 3,850,752; 3,901,654; 3,935,074; 3,984,533; 3,996,345; 4,034,074; and 4,098,876, all of which are herein incorporated by reference.

For detection purposes, HER-2/neu polypeptide ("antigen") may either be labeled or unlabeled. unlabeled, the antigen finds use in agglutination assays. In addition, unlabeled antigen can be used in combination with labeled molecules that are reactive immunocomplexes, or in combination with labeled antibodies (second antibodies) that are reactive with the antibody 10 directed against HER-2/neu polypeptide, such as antibodies specific for immunoglobulin. Alternatively, the antigen can be directly labeled. Where it is labeled, reporter group can include radioisotopes, fluorophores, 15 enzymes, luminescers, or dye particles. These and other labels are well known in the art and are described, for example, in the following U.S. patents: 3,766,162; 3,791,932; 3,817,837; 3,996,345; and 4,233,402.

Typically in an ELISA assay, antigen is adsorbed 20 to the surface of a microtiter well. Residual proteinbinding sites on the surface are then blocked with an appropriate agent, such as bovine serum albumin (BSA), heat-inactivated normal goat serum (NGS), (buffered solution of nonfat dry milk which also contains 25 a preservative, salts, and an antifoaming agent). well is then incubated with a sample suspected containing specific antibody. The sample can be applied neat, or, more often, it can be diluted, usually in a buffered solution which contains a small amount (0.1%-5.0% 30 by weight) of protein, such as BSA, NGS, or BLOTTO. After incubating for a sufficient length of time to allow specific binding to occur, the well is washed to remove unbound protein and then incubated with an anti-species specific immunoglobulin antibody labeled with a reporter

group. The reporter group can be chosen from a variety of enzymes, including horseradish peroxidase, betagalactosidase, alkaline phosphatase, and glucose oxidase. Sufficient time is allowed for specific binding to occur, then the well is again washed to remove unbound conjugate, and the substrate for the enzyme is added. Color is allowed to develop and the optical density of the contents of the well is determined visually or instrumentally.

In one preferred embodiment of this aspect of the present invention, a reporter group is bound to HER-2/neu protein. The step of detecting immunocomplexes involves removing substantially any unbound HER-2/neu protein and then detecting the presence or absence of the reporter group.

15 In another preferred embodiment, а group is bound to a second antibody capable of binding to the antibodies specific for HER-2/neu protein. The step detecting immunocomplexes involves (a) removing substantially any unbound antibody, (b) adding the second 20 antibody, (c) removing substantially any unbound second antibody and then (d) detecting the presence or absence of the reporter group. Where the antibody specific for HER-2/neu protein is derived from a human, the second antibody is an anti-human antibody.

In a third preferred embodiment for detecting immunocomplexes, a reporter group is bound to a molecule capable of binding to the immunocomplexes. The step of detecting involves (a) adding the molecule, (b) removing substantially any unbound molecule, and then (c) detecting the presence or absence of the reporter group. An example of a molecule capable of binding to the immunocomplexes is protein A.

It will be evident to one skilled in the art that a variety of methods for detecting the

immunocomplexes may be employed within the present invention. Reporter groups suitable for use in any of the methods include radioisotopes, fluorophores, enzymes, luminescers, and dye particles.

5 In a related aspect of the present invention, detection of immunocomplexes formed between HER-2/neu polypeptide and antibodies in body fluid which are specific for HER-2/neu polypeptide may be used to monitor the effectiveness of cancer therapy, which involves a 10 HER-2/neu polypeptide, for a malignancy in which the HER-2/neu oncogene is associated. Samples of body fluid taken from an individual prior to and subsequent to initiation of therapy may be analyzed for the immunocomplexes by the methodologies described above. Briefly, the number of 15 immunocomplexes detected in both samples are compared. substantial change in the number of immunocomplexes in the second sample (post-therapy initiation) relative to the first sample (pre-therapy) reflects successful therapy.

The following examples are offered by way of 20 illustration and not by way of limitation.

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EXAMPLES

EXAMPLE 1

Expression and Purification of Recombinant Human HER-2/NEU Polypeptide

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The human ${\tt HER-2/neu}$ polypeptide was recovered by method (e.g., U.S. Patent Nos. 4,683,195; 4,683,202; 4,800,159) from a plasmid prepared according to Di Fiore et al. (King et al., Science 229:974-976, 1.0 Di Fiore et al., Science *237*:178-182, 1987) oligonucleotide primers that additionally introduced a BssHII restriction site and an enterokinase protease site on the 5' end and an EcoRI site on the 3' end. The primer for the 5'-end was

5'-TCTGGCGCGCTGGATGACGATGACAAGAACGACGGCAGCAGAAGATC-3' 15 ID NO:3) while the primer for the 3'-end was 5'-TGAATTCTCGAGTCATTACACTGGCACGTCCAGACCCAG-3' (SEO ΙD The resulting 1.8 kb PCR fragment was subcloned into the T-vector from Novagen (Madison, WI, USA) and the sequence of selected clones was determined on the ABI 373 automated DNA sequencer (Applied Biosystems Inc., Foster City, CA, USA) using overlapping sequencing primers. fragments with sequence that corresponded to the published DNA sequence for the human HER-2/neu cDNA (SEQ ID NO:1; Coussens et al., Science 230:1132, 1985; Yamamoto et al., 25 Nature 319:230, 1986) were then connected in the correct reading frame via the BssHII site to a modified E. coli thioredoxin reductase. A 6Xhistidine affinity employed in Ni-NTA affinity purification of the expressed 30 fusion protein was incorporated into the thioredoxin reductase fusion partner. This cDNA for the trxA-human HER-2/neu polypeptide fusion protein was subcloned into a

modified pET expression vector for expression in E. coli.

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While thioredoxin reductase has been reported to stabilize and solubilize other heterologous proteins expressed in E. coli, it did not appear to offer any significant advantage for human HER-2/neu polypeptide expression in E. coli. While a significant proportion of the trxA-HER-2/neu polypeptide fusion protein was soluble, a majority was expressed in inclusion bodies. subjected protein was also to degradation expression in E. coli. The presence of the thioredoxin 10 reductase fusion partner may, however, stabilize the protein during purification. The availability of monoclonal antibodies to thioredoxin reductase provides a convenient marker to follow during purification.

purification of the human 15 polypeptide with the thioredoxin reductase fusion partner containing the 6XHis affinity tag, the E. coli pellet was resuspended with protease inhibitors and lysozyme sonicated. The inclusion bodies were isolated by centrifugation, and are washed 3X with deoxycholate, the 20 last wash being overnight to remove LPS. The washed inclusion bodies solubilized in are GuHCl The Ni column was eluted with Imidazole in purification. urea and dialyzed against 10 mM Tris pH8. The recovery of HER-2/neu polypeptide using this protocol was from 80%-95% 25 pure full length protein with the main contaminant being degraded protein. From 500 ml of fermentation, 20 mg were recovered. It was >98% HER-2/neu polypeptide. techniques used herein are well known to those in the art and have been described, for example, in J. Sambrook et al., Molecular Cloning: A Laboratory Manual, 2nd Ed., Cold Spring Harbor Laboratory Press, 1989, Cold Spring Harbor, New York, USA.

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EXAMPLE 2

DENDRITIC CELLS CAN PRIME HUMAN HER-2/NEU POLYPEPTIDE

5 A. Generation of DC Cultures From Bone Marrow

DC cultures were generated from hematopoietic progenitor cells (HPC). CD34+ cells were purified from bone marrow of normal donors using the cell separation system Ceprate LC Kit (CellPro, Bothell, WA, Purity of recovered CD34+ cells was determined by 10 USA). flow cytometric analysis to be 80% to 95%. CD34+ cells cultured in serum-free medium (X-VIVO Biowhittaker, Inc., Walkersville, MD, USA) supplemented L-glutamine (584 $\mu g/l)$, penicillin (10 streptomycin (100 μ g/ml), 100 ng/ml human rGM-CSF and 50 15 ng/ml human rIL-6 (Immunex, Seattle, WA, USA). After 0 to 17 days of culture time, cells were harvested and used for phenotyping and T cell stimulation assays. GM-CSF alone and in combination with IL-4 or $\text{TNF}\alpha$ have been described to induce the in vitro growth of DC. In experiments using 20 KLH and OVA as antigens to prime naive T cells, GM-CSF IL-6 consistently qave a comparable stimulation, but with a lower background and thus a higher

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B. T Cell Priming Assay

Bone marrow derived CD34+ HPC cultured in serum-free medium containing GM-CSF and IL-6 were used as APC after a culture period of 0-17 days. Priming ability of DC was determined by culturing them with autologous, naive T lymphocytes in the presence or absence of the protein antigen recombinant human HER-2/neu polypeptide (hHNP) (10 µg/ml). CD4+ T lymphocytes were isolated from peripheral

stimulation index as compared to GM-CSF plus IL-4 or $\text{TNF}\alpha$.

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blood mononuclear cells by positive selection using immunoaffinity columns (CellPro, Inc., Bothell, WA, USA). CD4+ CD45RA+ (naive) T lymphocytes were selected from CD4+ T lymphocytes using an anti-CD45RA mAb directly conjugated (Immunotech, Westbrook, ME, USA) cytometric sorting. The CD4+ CD45RA+ T cells obtained were 99% pure. DC cultures were plated into 96-well round-bottomed plates (Corning, Corning, NY, USA) various concentrations and incubated for 16-18 hours with 10 hHNP 10 μ g/ml final concentration. Antigen-pulsed DC were irradiated (10 Gy), and autologous CD4+ CD45RA+ T lymphocytes were added (5 x 10^4 /well). Proliferative response of T cells was measured by the uptake of (^{3}H) thymidine (1 μ Ci/well) added on day 6 for 16-18 hours. Proliferation assays were performed in serum-free and 15 cytokine-free medium. The results are shown in Figure 1. Figure 2 shows the results of testing CD4+ T cells, from a normal donor, for responses to hHNP. Similar data was obtained with T cells from nine out of ten normal 20 individuals.

EXAMPLE 3

Assay for Detecting Low Frequency Lymphocyte Precursors

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Three assays can be used for the detection of CD4⁺ responses: a standard proliferation assay, a screening method for low frequency events, and a limiting dilution assay (LDA). Conventional proliferative assays are capable of readily detecting primed responses. The proliferative response stimulation index provides a rough correlation with precursor frequency of antigen-reactive T cells. Any specific proliferative response detected from PBL is considered to be a primed response.

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To provide a more quantitative interpretation of CD4⁺ T cell responses, the assay system developed for detecting low lymphocyte precursor frequency responses (described below) is used. This assay is simple and costeffective. In circumstances in which more precision is needed, the precursor frequency is validated by limiting dilution assays (Bishop and Orosz, Transplantation 47:671-677, 1989).

Responses greater than detected in normal individuals are defined as a primed response and imply existent immunity. Low responses, detectable only by LDA conditions are considered to be unprimed responses. An absent response by LDA or a response lower than that defined by the normal population analysis is considered to be tolerance/anergy.

In general, primed CD4⁺ T cell responses can be detected in conventional proliferative assays, whereas unprimed responses are not detectable in the same assays. Detection of small numbers of unprimed T cells is limited by confounding background thymidine uptake including the autologous mixed lymphocyte response (AMLR) to self MHC antigen plus responses to processed self serum proteins and exogenously added serum proteins.

system for low frequency responses based on Poisson sampling statistics was used (In: Pinnacles, Chiron Corporation, 1:1-2, 1991). This type of analysis applies specifically to low frequency events in that, if the precursor frequency is less than the number of cells in one replicate culture, many replicates are required to detect a statistically significant number of positives. Theoretically, the analysis will correct for autologous responses by setting up a known positive control (such as PHA or tetanus toxoid) and known negative control (no

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antigen) and evaluating all data points from lowest to highest irrespective of the experimental group to which they belong. A cutoff value is calculated based on the equation cutoff = M + (F + SD), where M = arithmetic mean, 5 F = 3.29, a factor from tables of standardized normal distribution chosen so not more than 0.1% of the "true negatives" of a normally distributed background will be above the cutoff, and SD = standard deviation. In this screening assay, wells above the cutoff are considered true positives that potentially contain a lymphocyte that is specifically proliferating to the antigen of interest. Although estimations of lymphocyte precursor frequency is possible using this method, precise determination requires formal LDA analysis.

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EXAMPLE 4

HER-2/NEU POLYPEPTIDE BASED VACCINE ELICITS IMMUNITY TO HER-2/NEU PROTEIN

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A. Animals

Rats used in this study were Fischer strain 344 (CDF (F-344)/CrlBR) (Charles River Laboratories, Portage MI). Animals were maintained at the University of Washington Animal facilities under specific pathogen free conditions and routinely used for experimental studies between 3 and 4 months of age.

B. Immunization

Fischer rats were immunized with recombinant rat HER-2/neu polypeptide (rHNP) in a variety of adjuvants (MPL, Vaccel; Ribi, Bozeman, MT, USA). Animals received 50 μ g of rHNP mixed with adjuvant subcutaneously. Twenty days later the animals were boosted with a second

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immunization of 50 μ g of rHNP administered in the same fashion. Twenty days after the booster immunization animals were tested for the presence of antibodies directed against rat HER-2/neu protein (neu).

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C. Cell Lines

Two cell lines were used as a source of neu SKBR3, a human breast cancer cell line that is proteins. a marked overexpressor of HER-2/neu (American Type Culture 10 Collection, Rockville, MD), was maintained in culture in 10% fetal bovine serum (FBS) (Gemini Bioproducts, Inc., Calabasas, CA) and RPMI. DHFR-G8, an NIH/3T3 cell line cotransfected with cneu-p and pSV2-DHFR (American Type Culture Collection, Rockville, MD), was used as a source 15 of non-transforming rat new protein (Bernards et al., Proc. Natl. Acad. Sci. USA 84:6854-6858, 1987). This cell line was maintained in 10% FBS and Dulbecco's modified Eagle's medium with 4.5g/L glucose. DHFR-G8 cells were passaged through the same medium supplemented with 0.3 μM 20 methotrexate at every third passage to maintain the neu transfectant.

D. Preparation of Cell Lysates

Lysates of both SKBR3 and DHFR-G8 were prepared 25 and used as a source of neu protein. Briefly, a lysis buffer consisting of tris base, sodium chloride Triton-X (1%) pH 7.5 was prepared. Protease inhibitors were added; aprotinin (lµg/ml), benzamidine (lmM) and PMSF 1 ml of the lysis buffer was used to suspend 10 $^{\prime}$ (1mM). 30 cells. The cells were vortexed for 15 seconds every 10 minutes for an hour until disrupted. All procedures were performed on ice in a 4°C cold room. After disruption the cells were microfuged at 4°C for 20 minutes. Supernatant was removed from cell debris and stored in small aliquots at -70°C until used. Presence of human and rat neu in the lysates was documented by Western blot analysis.

E. ELISA for Rat new Antibody Responses

5 96 well Immulon 4 plates (Baxter SP, Redmond, WA: Dynatech Laboratories) were incubated overnight at 4°C with a rat new specific monoclonal antibody (Oncogene Science), 7.16.4, at a concentration of 10 μ g/ml diluted in carbonate buffer (equimolar concentrations of Na₂CO₃ and NaHCO₃ pH 9.6). After incubation, all wells were blocked 10 with PBS-1% BSA (Sigma Chemical, St. Louis, MO, USA), 100 μ l/well for 3 hours at room temperature. The plate was washed with PBS-0.5% Tween and lysates of DHFRG8, a murine cell line transfected with rat neu DNA (American Type 15 Culture Collection, Rockville, MD, USA); a source of rat neu protein, were added to alternating rows. The plate was incubated overnight at 4°C. The plate was then washed with PBS-0.5% Tween and experimental sera was added at the following dilutions: 1:25 to 1:200. The sera was diluted 20 in PBS-1% BSA-1% FBS-25 $\mu g/ml$ mouse IgG-0.01% NaN₃ and then serially into PBS-1% BSA. 50 μ l of diluted sera was added/well and incubated 1 hour at room temperature. Each experimental sera was added to a well with rat neu and a without rat neu. Sheep anti-rat Ιq F(ab')2 horseradish peroxidase (HRP) was added to the wells at a 1:5000 dilution in PBS-1% BSA and incubated for 45 minutes at room temperature (Amersham Co., Arlington Heights, IL, Following the final wash, TMB (Kirkegaard and Perry Laboratories, Gaithersburg, MD) developing reagent was added. Color reaction was read at an optical density of 30 The OD of each serum dilution was calculated as the OD of the rat new coated wells minus the OD of the PBS-1% BSA coated wells. Sera from animals immunized with the adjuvants alone and an animal immunized with hHNP (foreign protein) were also evaluated in a similar manner. The results are shown in Figure 3.

5 F. <u>T Cell Proliferation Assays</u>

For analysis of HER-2/neu polypeptide specific responses: Fresh spleen or lymph node cells are harvested by mechanical disruption and passage through wire mesh and washed. 2 x 10^5 spleen cells/well and 1 x 10^5 lymph node 10 cells/well are plated into 96-well round bottom microtiter plates (Corning, Corning, NY) with 6 replicates experimental group. The media consists of EHAA (Biofluids) with L-glutamine, penicillin/streptomycin, mercaptoethanol, and 5% FBS. Cells are incubated with 15 polypeptides. After 4 days, wells are pulsed with 1 μCi of [3H]thymidine for 6-8 hours and counted. expressed as a stimulation index (SI) which is defined as the mean of the experimental wells divided by the mean of the control wells (no antigen). For analysis of HER-2/neu protein specific responses: Spleen or lymph node cells are cultured for 3 in vitro stimulations. At the time of analysis 1×10^5 cultured spleen or lymph node T cells are plated into 96 well microtiter plates as described above. Cells are incubated with lpg/ml immunoaffinity column 25 purified rat neu (from DHFR-G8 cells as the source of rat After 4 days, wells were pulsed with 1 µCi of [3H] thymidine for 6-8 hours and counted. Data is expressed as a stimulation index which is defined as the mean of the experimental wells divided by the mean of the control 30 wells (no antigen).

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EXAMPLE 5

PRIMED RESPONSES TO HUMAN HER-2/NEU POLYPEPTIDE

CAN BE DETECTED IN PATIENTS WITH BREAST CANCER

Heparinized blood was obtained from a patient with stage II HER-2/neu overexpressing breast Peripheral blood mononuclear cells (PBMC) were separated by Ficoll Hypaque density centrifugation. PBMC were plated at a concentration of 2×10^5 /well into 96-well 10 round-bottomed plates (Corning, Corning, NY, USA). well replicates were performed for each experimental group. Antigens consisting of HER-2/neu derived peptides (15-20 amino acids in length with number of first amino acid in sequence listed) 25 μ g/ml, human HER-2/neu 15 polypeptide (hHNP) 1 μ g/ml, tetanus toxoid 1 μ g/ml, and p30 a peptide derived from tetanus 25 μ g/ml were added to each 24 well replicate. The assay was performed in media containing 10% human sera. Proliferative response of T cells was measured by the uptake of (3H)thymidine 20 (1 μ Ci/well) added on day 4 for 10 hours. Positive wells, antigen reactive wells, were scored as positive if the cpm was greater than the mean and 3 standard deviations of the no antigen wells. The results are shown in Figure 4. This stage II breast cancer patient has a significant 25 response to recombinant hHNP.

From the foregoing, it will be evident that, although specific embodiments of the invention have been described herein for purposes of illustration, various modifications may be made without deviating from the spirit and scope of the invention.

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Sequence Listing

- (1) GENERAL INFORMATION:
 - (i) APPLICANT: University of Washington
 - (ii) TITLE OF INVENTION: COMPOUNDS FOR ELICITING OR ENHANCING IMMUNE REACTIVITY TO HER-2/neu PROTEIN FOR PREVENTION OR TREATMENT OF MALIGNANCIES IN WHICH THE HER-2/neu ONCOGENE IS ASSOCIATED
 - (iii) NUMBER OF SEQUENCES: 4
 - (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: SEED and BERRY LLP
 - (B) STREET: 6300 Columbia Center, 701 Fifth Avenue
 - (C) CITY: Seattle
 - (D) STATE: Washington
 - (É) COUNTRY: USA
 - (F) ZIP: 98104-7092
 - (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.30
 - (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER:
 - (B) FILING DATE: 28-MAR-1996
 - (C) CLASSIFICATION:
 - (viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: Sharkey, Richard G.
 - (B) REGISTRATION NUMBER: 32,629
 - (C) REFERENCE/DOCKET NUMBER: 920010.448PC
 - (ix) TELECOMMUNICATION INFORMATION:
 - (A) TELEPHONE: (206) 622-4900
 - (B) TELEFAX: (206) 682-6031
- (2) INFORMATION FOR SEQ ID NO:1:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 3768 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(ix) FEATURE:

(A) NAME/KEY: CDS

(B) LOCATION: 1..3765

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

GAG G1u								48
CCC Pro								96
CGG Arg								144
TAC Tyr 50								192
CCC Pro							 	240
GGC Gly								288
AGG Arg								336
 CTG Leu							 	384

50

		115					120					125				
	ACA Thr 130															432
	ACA Thr															480
	TGC Cys															528
AAC Asn	CAG Gln	CTG Leu	GCT Ala 180	CTC Leu	ACA Thr	CTG Leu	ATA Ile	GAC Asp 185	ACC Thr	AAC Asn	CGC Arg	TCT Ser	CGG Arg 190	GCC Ala	TGC Cys	576
	CCC Pro															624
	GAG Glu 210															672
	CGC Arg															720
	GCC Ala															768
	TTC Phe															816
	TAC Tyr															864
	ACA Thr 290															912

ACG Thr										960
GTG Val									1	800
TGT Cys									1	056
AGG Arg									1	104
ATC Ile 370									1	152
GCC Ala									1	.200
ACT Thr									1	.248
AGC Ser									1	.296
CGA Arg									1	.344
ATC Ile 450									1	1392
GCC Ala					Phe	_		_	1	1440

		TGG Trp															1488
		AAC Asn															1536
		CTG Leu															1584
		AAC Asn 530															1632
,		GTA Val															1680
		CCG Pro															1728
		GGA Gly															1776
		CCC Pro															1824
		TAC Tyr 610															1872
		TGC Cys															1920
		TGC Cys															1968
(GCG	GTG	GTT	GGC	ATT	CTG	CTG	GTC	GTG	GTC	TTG	GGG	GTG	GTC	П	GGG	2016

Ala	Val	Val	Gly 660	Ile	Leu	Leu	Val	Val 665	Val	Leu	Gly	Val	Val 670	Phe	Gly	
	CTC Leu															2064
	CTG Leu 690															2112
	ATG Met															2160
	AAG Lys															2208
	ATC Ile															2256
	GTG Val															2304
	GAA Glu 770															2352
	CTG Leu															2400
	CCC Pro															2448
	GGC Gly					-										2496
	AGC												TTG Leu			2544

835		840	845
			TC AAA ATT ACA GAC TTC 2592 al Lys Ile Thr Asp Phe 860
			CA GAG TAC CAT GCA GAT 2640 or Glu Tyr His Ala Asp 880
			G GAG TCC ATT CTC CGC 2688 eu Glu Ser Ile Leu Arg 895
			T TAT GGT GTG ACT GTG 2736 r Tyr Gly Val Thr Val 910
	Met Thr Phe Gly		r Asp Gly Ile Pro Ala 925
			G CGG CTG CCC CAG CCC 2832 u Arg Leu Pro Gln Pro 940
			G GTC AAA TGT TGG ATG 2880 t Val Lys Cys Trp Met 5 960
			G TTG GTG TCT GAA TTC 2928 u Leu Val Ser Glu Phe 975
			G GTC ATC CAG AAT GAG 2976 1 Val Ile Gln Asn Glu 990
	Pro Ala Ser Pro		C TTC TAC CGC TCA CTG 3024 r Phe Tyr Arg Ser Leu 1005
		Asp Leu Val Asp	T GCT GAG GAG TAT CTG 3072 p Ala Glu Glu Tyr Leu 1020

GTA CCC CAG Val Pro Glr 1025	CAG GGC TT Gln Gly Ph 10	e Phe Cys						3120
GGC ATG GTC Gly Met Val				Ser Thr			Gly	3168
GGG GAC CTG Gly Asp Leu						Pro		3216
TCT CCA CTG Ser Pro Leu 107	Ala Pro Se		Ala Gly					3264
GAC CTG GGA Asp Leu Gly 1090					Leu Pro			3312
GAC CCC AGC Asp Pro Ser 1105		n Arg Tyr						3360
CCC TCT GAG Pro Ser Glu				Leu Thr			Gln	3408
CCT GAA TAT Pro Glu Tyr						Ser		3456
CGA GAG GGC Arg Glu Gly 115	Pro Leu Pr		Arg Pro					3504
AGG CCC AAG Arg Pro Lys 1170					Val Lys			3552
TTT GCC TTT Phe Ala Phe 1185	GGG GGT GC Gly Gly Al 11	a Val Glu						3600

		GCT Ala			Gln					Pro					Ala	36	548
		AAC Asn		Tyr					Asp					Gly		36	596
		AGC Ser 1235	Thr					Pro					Pro			37	'44
		CTG Leu)														37	'68
(2)		ORMAT	SEQUE (A) (B)	ENCE LEN		RACTE 125	ERIST 55 am act	ΓICS: nino id		is							
	(1	ii) M	10LEC	ULE	TYPE	: pr	otei	n									
	()	ki) S	SEQUE	NCE	DESC	CRIPT	ION:	SEC) ID	NO:2	2:						
Met 1	Glu	Leu	Ala	Ala 5	Leu	Cys	Arg	Trp	Gly 10	Leu	Leu	Leu	Ala	Leu 15	Leu		
Pro	Pro	Gly	Ala 20	Ala	Ser	Thr	Gln	Val 25	Cys	Thr	Gly	Thr	Asp 30	Met	Lys		
Leu	Arg	Leu 35	Pro	Ala	Ser	Pro	G1u 40	Thr	His	Leu	Asp	Met 45	Leu	Arg	His		
Leu	Tyr 50	Gln	Gly	Cys	Gln	Val 55	Val	Gln	Gly	Asn	Leu 60	Glu	Leu	Thr	Tyr		
Leu 65	Pro	Thr	Asn	Ala	Ser 70	Leu	Ser	Phe	Leu	G1n 75	Asp	Ile	Gln	Glu	Val 80		
Gln	GIV	Tyr	۱۵۱	برما	בוז	ΛΊэ	Hic	Acn	Gln	Val	Ara	Cln	Val	Dro	Lou		

Gln	Arg	Leu	Arg 100	Ile	Val	Arg	Gly	Thr 105	Gln	Leu	Phe	Glu	Asp 110	Asn	Tyr
Ala	Leu	Ala 115	Val	Leu	Asp	Asn	Gly 120	Asp	Pro	Leu	Asn	Asn 125	Thr	Thr	Pro
Val	Thr 130	Gly	Ala	Ser	Pro	Gly 135	Gly	Leu	Arg	Glu	Leu 140	Gln	Leu	Arg	Ser
Leu 145	Thr	Glu	Ile	Leu	Lys 150	Gly	Gly	Val	Leu	Ile 155	Gln	Arg	Asn	Pro	Gln 160
Leu	Cys	Tyr	Gln	Asp 165	Thr	Ile	Leu	Trp	Lys 170	Asp	Ile	Phe	His	Lys 175	Asn
Asn	Gln	Leu	Ala 180	Leu	Thr	Leu	Ile	Asp 185	Thr	Asn	Arg	Ser	Arg 190	Ala	Cys
His	Pro	Cys 195	Ser	Pro	Met	Cys	Lys 200	Gly	Ser	Arg	Cys	Trp 205	Gly	Glu	Ser
Ser	Glu 210	Asp	Cys	Gln	Ser	Leu 215	Thr	Arg	Thr	Val	Cys 220	Ala	Gly	Gly	Cys
A1a 225	Arg	Cys	Lys	Gly	Pro 230	Leu	Pro	Thr	Asp	Cys 235	Cys	His	Glu	Gln	Cys 240
Ala	Ala	Gly	Cys	Thr 245	Gly	Pro	Lys	His	Ser 250	Asp	Cys	Leu	Ala	Cys 255	Leu
His	Phe	Asn	His 260	Ser	Gly	Ile		G1u 265		His	Cys	Pro	A1a 270	Leu	Val
Thr	Tyr	Asn 275	Thr	Asp	Thr	Phe	G1u 280	Ser	Met	Pro	Asn	Pro 285	Glu	Gly	Arg
Tyr	Thr 290	Phe	Gly	Ala	Ser	Cys 295	Val	Thr	Ala	Cys	Pro 300	Tyr	Asn	Tyr	Leu
Ser 305	Thr	Asp	Val	Gly	Ser 310	Cys	Thr	Leu	Val	Cys 315	Pro	Leu	His	Asn	G1n 320
Glu	Val	Thr	Ala	G1u 325	Asp	Gly	Thr	Gln	Arg 330	Cys	Glu	Lys	Cys	Ser 335	Lys

Pro	Cys	Ala	Arg 340	Val	Cys	Tyr	Gly	Leu 345		Met	Glu	His	Leu 350	Arg	GΊι
Val	Arg	Ala 355	Val	Thr	Ser	Ala	Asn 360	Ile	Gln	Glu	Phe	A1a 365	Gly	Cys	Lys
Lys	Ile 370	Phe	Gly	Ser	Leu	A1a 375	Phe	Leu	Pro	Glu	Ser 380	Phe	Asp	Gly	Asp
Pro 385	Ala	Ser	Asn	Thr	Ala 390	Pro	Leu	Gln	Pro	G1u 395	Gln	Leu	Gln	Val	Phe 400
Glu	Thr	Leu	Glu	G1u 405	Пe	Thr	Gly	Tyr	Leu 410	Tyr	Ile	Ser	Ala	Trp 415	Pro
Asp	Ser	Leu	Pro 420	Asp	Leu	Ser	Val	Phe 425	Gln	Asn	Leu	Gln	Val 430	Ile	Arg
Gly	Arg	Ile 435	Leu	His	Asn	Gly	Ala 440	Tyr	Ser	Leu	Thr	Leu 445	Gln	Gly	Leu
Gly	Ile 450	Ser	Trp	Leu	Gly	Leu 455	Arg	Ser	Leu	Arg	Glu 460	Leu	Gly	Ser	Gly
Leu 465	Ala	Leu	Ile	His	His 470	Asn	Thr	His	Leu	Cys 475	Phe	Val	His	Thr	Va1 480
Pro	Trp	Asp	Gln	Leu 485	Phe	Arg	Asn	Pro	His 490	Gln	Ala	Leu	Leu	His 495	Thr
Ala	Asn	Arg	Pro 500	Glu	Asp	Glu	Cys	Va1 505	Gly	Glu	Gly	Leu	Ala 510	Cys	His
Gln	Leu	Cys 515	Ala	Arg	Gly	His	Cys 520	Trp	Gly	Pro	Gly	Pro 525	Thr	Gln	Cys
Val	Asn 530	Cys	Ser	Gln	Phe	Leu 535	Arg	Gly	G1n	Glu	Cys 540	Val	G1u	Glu	Cys
Arg 545	Val	Leu	Gln	Gly	Leu 550	Pro	Arg	Glu	Tyr	Val 555	Asn	Ala	Arg	His	Cys 560
Leu	Pro	Cys	His	Pro 565	Glu	Cys	Gln	Pro	G1n 570	Asn	Gly	Ser	Val	Thr 575	Cys

Phe	Gly	Pro	G1u 580	Ala	Asp	Gln	Cys	Va1 585	Ala	Cys	Ala	His	Tyr 590	Lys	Asp
Pro	Pro	Phe 595	Cys	Val	Ala	Arg	Cys 600	Pro	Ser	Gly	Val	Lys 605	Pro	Asp	Leu
Ser	Tyr 610	Met	Pro	Ile	Trp	Lys 615	Phe	Pro	Asp	Glu	Glu 620	Gly	Ala	Cys	Gln
Pro 625	Cys	Pro	Ile	Asn	Cys 630	Thr	His	Ser	Cys	Val 635	Asp	Leu	Asp	Asp	Lys 640
Gly	Cys	Pro	Ala	G1u 645	Gln	Arg	Ala	Ser	Pro 650	Leu	Thr	Ser	Ile	Ile 655	Ser
Ala	Val	Val	Gly 660	Ile	Leu	Leu	Val	Val 665	Val	Leu	Gly	Val	Val 670	Phe	Gly
Ile	Leu	Ile 675	Lys	Arg	Arg	Gln	Gln 680	Lys	Пе	Arg	Lys	Tyr 685	Thr	Met	Arg
Arg	Leu 690	Leu	Gln	Glu	Thr	G1u 695	Leu	Val	Glu	Pro	Leu 700	Thr	Pro	Ser	Gly
A1a 705	Met	Pro	Asn	Gln	Ala 710	Gln	Met	Arg	Ile	Leu 715	Lys	Glu	Thr	Glu	Leu 720
Arg	Lys	Val	Lys	Val 725	Leu	Gly	Ser	Gly	Ala 730	Phe	Gly	Thr	Val	Tyr 735	Lys
Gly	Пe	Trp	Ile 740	Pro	Asp	Gly		Asn 745		Lys	Ile	Pro	Va1 750	Ala	Ile
Lys	Val	Leu 755	Arg	Glu	Asn	Thr	Ser 760	Pro	Lys	Ala	Asn	Lys 765	Glu	Пе	Leu
Asp	G1u 770	Ala	Tyr	Val	Met	Ala 775	Gly	Val	Gly	Ser	Pro 780	Tyr	Val	Ser	Arg
Leu 785	Leu	Gly	Ile	Cys	Leu 790	Thr	Ser	Thr	Val	G1n 795	Leu	Val	Thr	Gln	Leu 800
Met	Pro	Tyr	Gly	Cys 805	Leu	Leu	Asp	His	Val 810	Arg	Glu	Asn	Arg	Gly 815	Arg

Gly	Met	Val	His	His 1045	-	His	Arg	Ser	Ser 1050		Thr	Arg	Ser	Gly 1055	-
Val 1025	Pro	Gln	Gln	Gly	Phe 1030		Cys	Pro	Asp	Pro 1035		Pro	Gly	Ala	Gly 1040
Leu	Glu 1010		Asp	Asp	Met	Gly 1015	•	Leu	Val	Asp	Ala 1020		Glu	Tyr	Leu
Asp	Leu	Gly 995	Pro	Ala	Ser	Pro	Leu 1000	•	Ser	Thr	Phe	Tyr 1005	_	Ser	Leu
Ser	Arg	Met	Ala 980	Arg	Asp	Pro	Gln	Arg 985	Phe	Val	Val	Ile	G1n 990	Asn	G1u
Пе	Asp	Ser	Glu	Cys 965	Arg	Pro	Arg	Phe	Arg 970	Glu	Leu	Val	Ser	G1u 975	Phe
Pro 945	Пе	Cys	Thr	Пе	Asp 950	Val	Tyr	Met	Ile	Met 955	Val	Lys	Cys	Trp	Met 960
Arg	G1u 930	Ile	Pro	Asp	Leu	Leu 935	Glu	Lys	Gly	Glu	Arg 94 0	Leu	Pro	Gln	Pro
Trp	Glu	Leu 915	Met	Thr	Phe	Gly	Ala 920	Lys	Pro	Tyr	Asp	Gly 925	Ile	Pro	Ala
Arg	Arg	Phe	Thr 900	His	Gln	Ser	Asp	Val 905	Trp	Ser	Tyr	Gly	Val 910	Thr	Val
Gly	Gly	Lys	Val	Pro 885	Ile	Lys	Trp	Met	Ala 890	Leu	Glu	Ser	Пе	Leu 895	Arg
G1y 865	Leu	Ala	Arg	Leu	Leu 870	Asp	Ilе	Asp	Glu	Thr 875	Glu	Tyr	His	Ala	Asp 880
Arg	Asn 850	Val	Leu	Val	Lys	Ser 855	Pro	Asn	His	Val	Lys 860	Пе	Thr	Asp	Phe
Met	Ser	Tyr 835	Leu	Glu	Asp	Val	Arg 840	Leu	Val	His	Arg	Asp 845	Leu	Ala	Ala
Leu	Gly	Ser	G1n 820	Asp	Leu	Leu	Asn	Trp 825	Cys	Met	Gln	Ile	A1a 830	Lys	Gly

- Gly Asp Leu Thr Leu Gly Leu Glu Pro Ser Glu Glu Glu Ala Pro Arg 1060 1065 1070
- Ser Pro Leu Ala Pro Ser Glu Gly Ala Gly Ser Asp Val Phe Asp Gly 1075 1080 1085
- Asp Leu Gly Met Gly Ala Ala Lys Gly Leu Gln Ser Leu Pro Thr His 1090 1095 1100
- Asp Pro Ser Pro Leu Gln Arg Tyr Ser Glu Asp Pro Thr Val Pro Leu 1105 1110 1115 1120
- Pro Ser Glu Thr Asp Gly Tyr Val Ala Pro Leu Thr Cys Ser Pro Gln 1125 1130 1135
- Pro Glu Tyr Val Asn Gln Pro Asp Val Arg Pro Gln Pro Pro Ser Pro 1140 1145 1150
- Arg Glu Gly Pro Leu Pro Ala Ala Arg Pro Ala Gly Ala Thr Leu Glu 1155 1160 1165
- Arg Pro Lys Thr Leu Ser Pro Gly Lys Asn Gly Val Val Lys Asp Val 1170 1175 1180
- Phe Ala Phe Gly Gly Ala Val Glu Asn Pro Glu Tyr Leu Thr Pro Gln 1185 1190 1195 1200
- Gly Gly Ala Ala Pro Gln Pro His Pro Pro Pro Ala Phe Ser Pro Ala 1205 1210 1215
- Phe Asp Asn Leu Tyr Tyr Trp Asp Gln Asp Pro Pro Glu Arg Gly Ala 1220 1225 1230
- Pro Pro Ser Thr Phe Lys Gly Thr Pro Thr Ala Glu Asn Pro Glu Tyr 1235 1240 1245
- Leu Gly Leu Asp Val Pro Val 1250 1255
- (2) INFORMATION FOR SEQ ID NO:3:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 48 base pairs
 - (B) TYPE: nucleic acid

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(C)	STRANDEDNESS	: single
(D)	TOPOLOGY: lin	near

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

TCTGGCGCGC TGGATGACGA TGACAAGAAA CGACGGCAGC AGAAGATC

48

(2) INFORMATION FOR SEQ ID NO:4:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 39 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

TGAATTCTCG AGTCATTACA CTGGCACGTC CAGACCCAG

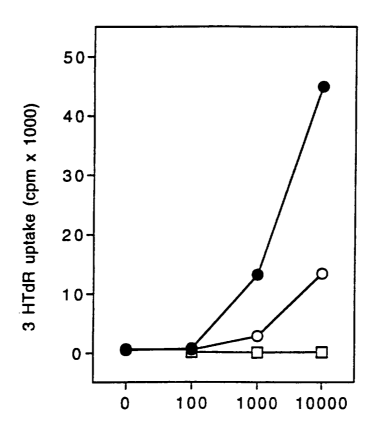
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Claims

- 1. A polypeptide encoded by a DNA sequence selected from:
- (a) nucleotides 2026 through 3765 of SEQ ID NO:1; and
- (b) DNA sequences that hybridize to a nucleotide sequence complementary to nucleotides 2026 through 3765 of SEQ ID NO:1 under moderately stringent conditions, wherein the DNA sequence encodes a polypeptide that produces an immune response to HER-2/neu protein.
- 2. A polypeptide having the amino acid sequence of SEQ ID NO:2 from lysine, amino acid 676, through valine, amino acid 1255, or a variant thereof that produces at least an equivalent immune response.
- 3. A polypeptide according to claim 2 having the amino acid sequence of SEQ ID NO:2 from amino acid 676 through amino acid 1255.
- 4. A composition comprising a polypeptide according to any one of claims 1, 2 or 3, in combination with a pharmaceutically acceptable carrier or diluent.
- 5. A polypeptide according to any one of claims 1, 2 or 3, or a composition according to claim 4, for the immunization of a warm-blooded animal against a malignancy in which the HER-2/neu oncogene is associated.
- 6. Use of a polypeptide according to any one of claims 1, 2 or 3, or a composition according to claim 4, for the manufacture of a medicament for immunization of a warm-

blooded animal against a malignancy in which the $\mbox{HER-2/neu}$ oncogene is associated.

- 7. A nucleic acid molecule directing the expression of a polypeptide according to any one of claims 1, 2 or 3 for immunization by transfecting the cells of a warmblooded animal with the nucleic acid molecule.
- 8. A nucleic acid molecule according to claim 7 wherein the cells are transfected $ex\ vivo$ and subsequently delivered to the animal.
- 9. Use of a nucleic acid molecule directing the expression of a polypeptide according to any one of claims 1, 2 or 3, for the manufacture of a medicament for immunization of a warm-blooded animal against a malignancy in which the HER-2/neu oncogene is associated.
- 10. A viral vector directing the expression of a polypeptide according to any one of claims 1, 2 or 3 for immunization by infecting the cells of a warm-blooded animal with the vector.
- 11. A viral vector according to claim 10 wherein the cells are infected ex vivo and subsequently delivered to the animal.
- 12. Use of a viral vector directing the expression of a polypeptide according to any one of claims 1, 2 or 3, for the manufacture of a medicament for immunization of a warmblooded animal against a malignancy in which the HER-2/neu oncogene is associated.



Number of stimulator cells per well

Fig. 1

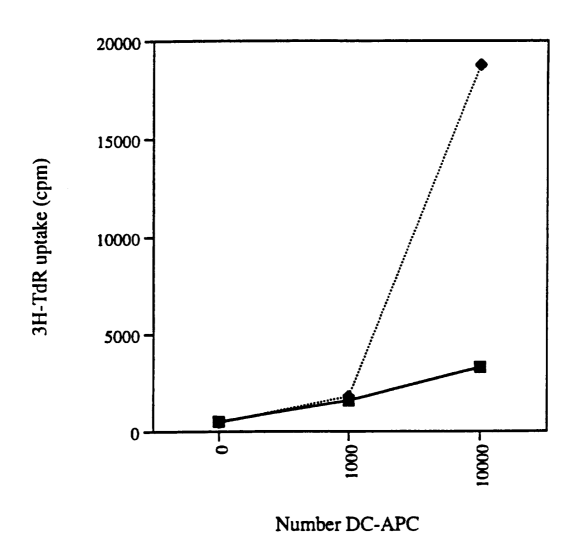


Fig. 2

SUBSTITUTE SHEET (RULE 26)

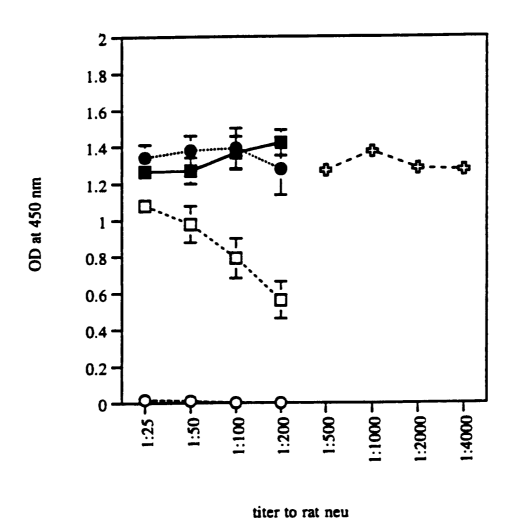


Fig. 3

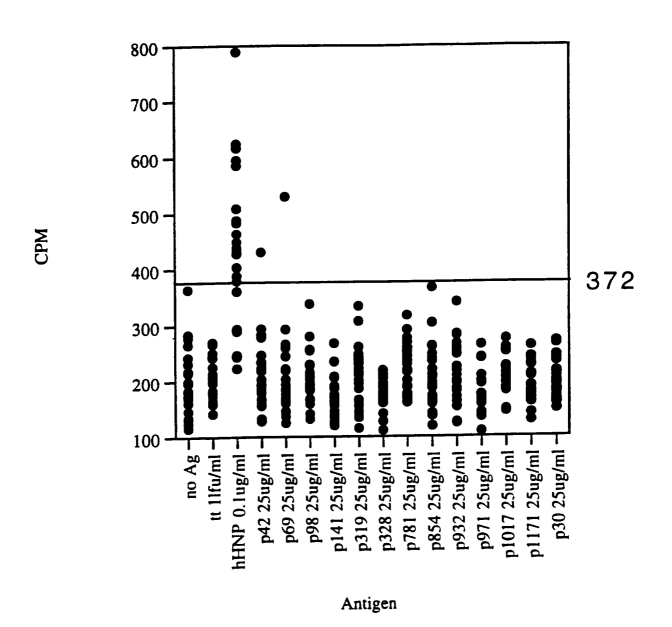


Fig. 4

SUBSTITUTE SHEET (RULE 26)

INTERNATIONAL SEARCH REPORT

Inte ional Application No PCI/US 96/01689

A. CLASS IPC 6	IFICATION OF SUBJECT MATTER C12N15/12 C07K14/71 C12N15/	86								
According to International Patent Classification (IPC) or to both national classification and IPC										
B. FIELDS	S SEARCHED									
Minimum d	locumentation searched (classification system followed by classification C12N C07K A61K	ution symbols)								
Documenta	tion searched other than minimum documentation to the extent that	such documents are included in the fields s	searched							
Electronic data base consulted during the international search (name of data base and, where practical, search terms used)										
C. DOCUM	IENTS CONSIDERED TO BE RELEVANT									
Category *	Citation of document, with indication, where appropriate, of the	relevant passages	Relevant to claim No.							
Х	CANCER RESEARCH, vol. 54, no. 1, 1 January 1994, pages 16-20, XP002010444 M.L.DISIS ET AL.: "Existent T-c antibody immunity to HER-2/neu p patients with breast cancer" see page 17, column 1, paragraph 3	1-12								
Α	WO,A,91 02062 (TRITON BIOSCIENCE February 1991 see the whole document	1-12								
Furt	her documents are listed in the continuation of box C.	X Patent family members are listed	in annex.							
'A' document defining the general state of the art which is not considered to be of particular relevance 'E' earlier document but published on or after the international filing date 'L' document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) 'O' document referring to an oral disclosure, use, exhibition or other means 'P' document published prior to the international filing date but later than the priority date claimed		"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention." "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone. "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. "&" document member of the same patent family.								
Date of the actual completion of the international search 8 August 1996		Date of mailing of the international search report 1 4. 08. 96								
Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+ 31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+ 31-70) 340-3016		Authorized officer Cupido, M								

INTERNATIONAL SEARCH REPORT Interional Application No

anormation on patent family members

PCI/US 96/01689

Patent document cited in search report	Publication date	Patent family member(s)		Publication date 27-01-94 11-03-91 05-02-91 04-09-91
WO-A-9102062	21-02-91	AU-B- 6413590 CA-A- 2042064 EP-A- 0444181		
		JP-T-	4503012	04-06-92