| S.No. | Cancer<br>Vaccine Type | Drug Name | Biological Name                           | Developer                               | Current<br>Development<br>Phase | Additional<br>Information  | Start<br>Date | Completio<br>Date |
|-------|------------------------|-----------|---|---|---------------------------------|--|---------------|-------------------|
| 1     | Anal                   |           | PSMA/PRAME                                | MannKind<br>Corporation                 | l                               | Completed The present clinical trial is a dose comparison of a multi-component active immunotherapy designed to stimulate an immune reaction to specific tumor associated antigens which are highly expressed on a large number of solid cancers.            | 2007          | 2009              |
| 2     | Biliary Tract          |           | PSMA/PRAME                                | MannKind<br>Corporation                 | I                               | Completed The present clinical trial is a dose comparison of a multi-component active immunotherapy designed to stimulate an immune reaction to specific tumor associated antigens which are highly expressed on a large number of solid cancers.            | 2007          | 2009              |
| 3     | Bladder                |           | CDX-1310                                  | Celldex Therapeutics                    | I                               | Celldex Therapeutics, Inc. is testing a form of immune therapy (vaccine) to see if it can be used to make the immune system attack the cancer  | 2006          | 2009              |
| 4     | Bladder                |           | Lapuleucel-T                              | Dendreon                                | Preclinical                     |  |               |                   |
| 5     | Bladder                |           | NY-ESO-1 plasmid<br>DNA Cancer<br>Vaccine | Ludwig Institute for<br>Cancer Research | I                               | Completed: To estimate the safety of NY-ESO-1 Plasmid DNA (pPJV7611) Cancer Vaccine given by PMED in patients with tumor type known to express NY-ESO-1 or LAGE-1 using frequency, severity, and duration of treatment-related adverse effects as endpoints. | 2004          | 2007              |
| 6     | Bladder                |           | V934/V935                                 | Merck                                   | I                               | Completed. This is a two-part study to test the safety, tolerability, and immune response for V934/V935 vaccine using a new prime-boost regimen in participants with selected solid tumors.  | 2008          | 2011              |
| 7     | Bone                   |           | PSMA/PRAME                                |   | I                               |  | 2007          | 2009              |

|    |       |   |   | MannKind<br>Corporation                 |    | Completed The present clinical trial is a dose comparison of a multi-component active immunotherapy designed to stimulate an immune reaction to specific tumor associated antigens which are highly expressed on a large number of solid cancers.   |      |      |
|----|-------|---|---|---|----|---|------|------|
| 8  | Bone  |   | PSMA/PRAME                                | MannKind<br>Corporation                 | I  | Completed The present clinical trial is a dose comparison of a multi-component active immunotherapy designed to stimulate an immune reaction to specific tumor associated antigens which are highly expressed on a large number of solid cancers.   | 2007 | 2009 |
| 9  | Bone  |   | Trivalent ganglioside<br>vaccine, OPT-821 | MabVax<br>Therapeutics                  | =  | The trivalent vaccine is being developed to teach the patient?s immune system to recognize 3 types of sugars called GM2, GD2 and GD3 that are found primarily on the surface of sarcoma cells. If the trivalent vaccine can stimulate the patient?s immune system to develop antibodies which recognize and target the GM2, GD2 and GM3 sugars, then the patient?s antibodies could attack and kill any remaining sarcoma cells potentially preventing the recurrence of sarcoma. | 2010 | 2014 |
| 10 | Bone  |   | NY-ESO-1 plasmid<br>DNA Cancer<br>Vaccine | Ludwig Institute for<br>Cancer Research | I  | Completed: To estimate the safety of NY-ESO-1 Plasmid DNA (pPJV7611) Cancer Vaccine given by PMED in patients with tumor type known to express NY-ESO-1 or LAGE-1 using frequency, severity, and duration of treatment-related adverse effects as endpoints.  | 2004 | 2007 |
| 11 | Brain | CDX-110 with<br>GM-CSF,<br>temozolomide |   | Celldex Therapeutics                    | II | This study is designed to evaluate the clinical activity of CDX-110 vaccination when given with standard of care treatment (maintenance temozolomide therapy).  | 2007 | 2011 |

|    |                                |                                 | •                             |                                     |    |   |      |      |
|----|--------------------------------|---------------------------------|-------------------------------|-------------------------------------|----|---|------|------|
| 12 | Brain                          | Dendritic cell<br>immunotherapy |                               | Northwest<br>Biotherapeutics        | II | In US only The purpose of the study is to determine the safety and efficacy of an investigational therapy called DCVax(R)-L in patients with newly diagnosed GBM for whom surgery is indicated.   | 2006 | 2012 |
| 13 | Brain                          |                                 | GliaAtak                      | Advantagene                         | II |   |      |      |
| 14 | Brain                          |                                 | Glionix                       | NovaRx                              | I  | completed   |      |      |
| 15 | Brain                          |                                 | ICT-107 , Placebo<br>DC       | ImmunoCellular<br>Therapeutics      | Ш  | The goal is for the ICT-107 vaccine to stimulate the patient?s immune response to kill the remaining GBM tumor cells after surgery and chemotherapy.  | 2011 | 2014 |
| 16 | Brain                          |                                 | ICT-121                       | ImmunoCellular<br>Therapeutics      | 1  |   |      |      |
| 17 | Brain                          | Cyclophosphamide,<br>Imiquimod  | IMA950 plus<br>GM-CSF, IMA950 | Immatics<br>Biotechnologies<br>GmbH | I  | The primary objective of this study is to determine the safety and tolerability of IMA950 when given with cyclophosphamide, granulocyte macrophage-colony stimulating factor (GM-CSF) and imiquimod in patients with glioblastoma and to determine if IMA950 shows sufficient immunogenicity in these patients. | 2011 | 2013 |
| 18 | Carcinoma of<br>Unknown Origin |                                 | PSMA/PRAME                    | MannKind<br>Corporation             | I  | Completed The present clinical trial is a dose comparison of a multi-component active immunotherapy designed to stimulate an immune reaction to specific tumor associated antigens which are highly expressed on a large number of solid cancers.   | 2007 | 2009 |
| 19 | Esophageal                     |                                 | PSMA/PRAME                    | MannKind<br>Corporation             | I  | The present clinical trial is a dose comparison of a multi-component active immunotherapy designed to stimulate an immune reaction to specific tumor associated antigens which are highly expressed on a large number of solid cancers.   | 2007 | 2009 |
| 20 | Esophageal                     |                                 | PSMA/PRAME                    | MannKind<br>Corporation             | I  | Completed The present clinical trial is a dose comparison of a multi-component active   | 2007 | 2009 |
|    |                                |                                 |                               |                                     |    |   |      |      |

|    |                           |                                |   |  |          | immunotherapy<br>designed to<br>stimulate an<br>immune reaction to<br>specific tumor<br>associated antigens<br>which are highly<br>expressed on a<br>large number of<br>solid cancers.   |      |      |
|----|---------------------------|--------------------------------|---|--|----------|--|------|------|
| 21 | Esophageal                |                                | NY-ESO-1 plasmid<br>DNA Cancer<br>Vaccine                   | Ludwig Institute for<br>Cancer Research                | I        | Completed: To estimate the safety of NY-ESO-1 Plasmid DNA (pPJV7611) Cancer Vaccine given by PMED in patients with tumor type known to express NY-ESO-1 or LAGE-1 using frequency, severity, and duration of treatment-related adverse effects as endpoints.                 | 2004 | 2007 |
| 22 | Esophageal                | Celecoxib,<br>cyclophosphamide | K562 (Allogeneic<br>Tumor Cell Vaccine)                     | National Cancer<br>Institute (NCI)                     | 1/11     | To evaluate the safety and effectiveness of tumor cell vaccines in combination with cyclophosphamide and celecoxib in patients with cancers involving the chest.   | 2010 | 2011 |
| 23 | Extrahepatic<br>Bile Duct |                                | carcinoembryonic<br>antigen RNA-pulsed<br>DC cancer vaccine | Duke University,<br>National Cancer<br>Institute (NCI) | I        | Phase I trial to study<br>the effectiveness of<br>biological therapy in<br>treating patients<br>who have metastatic<br>cancer that has not<br>responded to<br>previous treatment.  | 2000 | 2009 |
| 24 | Fallopian Tube            |                                | DPX-Survivac<br>with low dose<br>cyclophosphamide           | ImmunoVaccine<br>Technologies, Inc.                    | 1/11     | This is a phase 1-2 study to determine the safety and immunogenicity profiles of DPX-Survivac, a therapeutic vaccine co-administered with a regimen of low dose oral cyclophosphamide. DPX-Survivac is for the treatment of ovarian, fallopian tube, and peritoneal cancers. |      |      |
| 25 | Gallbladder               |                                | PSMA/PRAME  | MannKind<br>Corporation                                | ļ        | Completed The present clinical trial is a dose comparison of a multi-component active immunotherapy designed to stimulate an immune reaction to specific tumor associated antigens which are highly expressed on a large number of solid cancers.                            | 2007 | 2009 |
| 26 | Gallbladder               |                                | carcinoembryonic<br>antigen RNA-pulsed<br>DC cancer vaccine | Duke University,<br>National Cancer<br>Institute (NCI) | 1        | Phase I trial to study<br>the effectiveness of<br>biological therapy in<br>treating patients<br>who have metastatic<br>cancer that has not<br>responded to   | 2000 | 2009 |
| •  | 1                         |                                |   |  | <b>.</b> | <b>.</b>   |      | 1    |

|  | 1 1   |      |
|--|---|------|
| previous trea  | atment.   |      |
| Completed The present trial is a dos comparison multi-compo active immunother.  PSMA/PRAME  MannKind Corporation  I designed to stimulate an immune rear specific tume associated a which are hid expressed of large number solid cancer.  | se of a onent rapy 2007 laction to lacting antigens ighly on a                          | 2009 |
| Allogeneic whole epithelial tumor cells, DNP-conjugated and irradiated  Allogeneic whole epithelial tumor cells, DNP-conjugated and irradiated  Hadassah Medical Organization  I/II  This study is on the finding tumor cells to grown in the laboratory or modified in sway that, who injected to the patient, they stimulate his immune respectively and irradiated  I/II  This study is on the finding tumor cells to grown in the laboratory or modified in sway that, who injected to the patient, they stimulate his immune respectively and irradiated. | ng that that are ele such a hen he y will syher ponse. hch will d in h gastric, hast or |      |
| 29 Gastric Carcinoembryonic antigen RNA-pulsed DC cancer vaccine Duke University, National Cancer Institute (NCI)  Phase I trial the effective biological the treating paties who have many cancer that I responded to previous treating paties.   | eness of lerapy in lents lents actastatic has not lo                                    | 2009 |
| 30 Gastrointestinal GVAX BioSante Pharmaceuticals  |   |      |
| 31 Gastrointestinal ANZ-100 Aduro BioTech I completed  |   |      |
| 32 Gastrointestinal DCVax-Liver Northwest Biotherapeutics  |   |      |
| 33 Gastrointestinal PancAtak Advantagene Preclinical   |   |      |
| 34 Gastrointestinal DCVax-Pancreas Northwest Biotherapeutics Preclinical   |   |      |
| 35 Genital Warts V503 Merck III Expected re late 2011  | esults in 2010  | 2011 |
| TVI-Brain-1 experimenta treatment th advantage of fact that you can produce immune cell ?killer? whit cells that ha ability to kill numbers of   | al nat takes of the ur body e ls, called te blood ave the large the                     | 2014 |
| Multiforme    TVAX Biomedical   II   | our strain-1 is rge those te blood deliver into your t they                             |      |

|    | Glioblastoma<br>Multiforme | Trivax,<br>Temozolomide,<br>Surgery,<br>Radiotherapy,<br>Temozolomide |  | Trimed Biotech<br>GmbH                                 |           | A randomised, open-label, 2-arm, multi-centre, phase II clinical study with one group receiving standard therapy with Temozolomide, radiotherapy, and Trivax; and a control group receiving standard therapy with Temozolomide and radiotherapy only; after tumour resection of at least 70% in both groups. The hypothesis is based on the assumption that time to progression will be doubled in the treatment group.      |      |      |
|----|----------------------------|---|--|--|-----------|--|------|------|
| 38 | Grade IV<br>Astrocytoma    |   | Cancer vaccine plus<br>immune adjuvant,<br>plus activated white<br>blood cells | TVAX Biomedical  | II        | TVI-Brain-1 is an experimental treatment that takes advantage of the fact that your body can produce immune cells, called ?killer? white blood cells that have the ability to kill large numbers of the cancer cells that are present in your body. TVI-Brain-1 is designed to generate large numbers of those ?killer? white blood cells and to deliver those cells into your body so that they can kill your cancer cells. | 2011 | 2014 |
| 39 | Grade IV<br>Glioma         |   | Cancer vaccine plus<br>immune adjuvant,<br>plus activated white<br>blood cells | TVAX Biomedical  | <b>II</b> | TVI-Brain-1 is an experimental treatment that takes advantage of the fact that your body can produce immune cells, called ?killer? white blood cells that have the ability to kill large numbers of the cancer cells that are present in your body. TVI-Brain-1 is designed to generate large numbers of those ?killer? white blood cells and to deliver those cells into your body so that they can kill your cancer cells. | 2011 | 2014 |
| 40 | Head and Neck              |   | PV701  | University of Chicago                                  | I         | Phase I trial to study the effectiveness of intratumoral (in the tumor) PV701 in treating patients who have advanced or recurrent unresectable squamous cell carcinoma (cancer) of the head and neck.  |      |      |
| 41 | Head and Neck              |   | carcinoembryonic<br>antigen RNA-pulsed<br>DC cancer vaccine                    | Duke University,<br>National Cancer<br>Institute (NCI) | I         | Phase I trial to study<br>the effectiveness of<br>biological therapy in<br>treating patients<br>who have metastatic<br>cancer that has not   | 2000 | 2009 |

|    |          | <b>i</b> 1 |   |   |             |   |      |         |
|----|----------|------------|---|---|-------------|---|------|---------|
|    |          |            |   |   |             | responded to previous treatment.  |      |         |
| 42 | Kidney   |            | Serum and urinary<br>CA9 level  | Centre Hospitalier<br>Universitaire de Saint<br>Etienne                             | Preclinical | It was demonstrated that the level of expression of CA9 in tumor tissue can be used as a predictive marker of response to immunotherapy.  | 2009 | 2013    |
| 43 | Leukemia |            | GRNVAC1   | Geron   | II          | This is a phase II study to evaluate the safety, feasibility and efficacy of immunotherapy with GRNVAC1 in patients with AML.   | 2007 | 2012    |
| 44 | Leukemia |            | PV327   | Wellstat Biologics  | Preclinical |   |      |         |
| 45 | Leukemia |            | Tumor Vaccine:<br>CD40 LIGAND AND<br>IL-2 GENE<br>MODIFIED<br>AUTOLOGOUS<br>SKIN<br>FIBROBLASTS<br>AND TUMOR<br>CELLS | Baylor College of<br>Medicine   | I           | This research study is to determine the safety and dosage of special cells that may make the patients own immune system fight the leukemia. To do this we will put special genes into cells called fibroblasts that we have grown in the laboratory from a skin sample. The genes we put in these fibroblasts make them produce substances called CD40 Ligand (CD40L) and interleukin-2 (IL-2). These are natural substances that may help the immune system kill leukemia cells. | 1999 | 2010    |
| 46 | Leukemia |            | ISF35   | University of<br>California, San Diego  | II          | This is a Phase II, open label, fixed dose, repeat injection, single institution study. Eligible subjects will receive up to six doses of Ad-ISF35 injected directly into a selected lymph node under ultrasound guidance. The primary goal is to determine and monitor clinical and biological responses in patients treated with repeat intranodal injections of Ad-ISF35.  | 2009 | 2011    |
| 47 | Leukemia |            | Ras peptide cancer vaccine, sargramostim  | Memorial<br>Sloan-Kettering<br>Cancer Center,<br>National Cancer<br>Institute (NCI) | I           | Phase I trial to study<br>the effectiveness of<br>vaccine therapy plus<br>sargramostim in<br>treating patients<br>who have<br>myelodysplastic<br>syndrome.  | 1999 | Ongoing |
| 48 | Liver    |            | PSMA/PRAME  | MannKind<br>Corporation   | 1           | Completed The present clinical trial is a dose comparison of a multi-component active immunotherapy designed to   | 2007 | 2009    |
|    |          |            |   |   |             |   |      |         |

| , i |                                      | į i                            | i   | Ī  | 1      | I  | Ĭ    | 1       |
|-----|--------------------------------------|--------------------------------|---|--|--------|--|------|---------|
|     |                                      |                                |   |  |        | stimulate an immune reaction to specific tumor associated antigens which are highly expressed on a large number of solid cancers.  |      |         |
| 49  | Liver                                |                                | carcinoembryonic<br>antigen RNA-pulsed<br>DC cancer vaccine | Duke University,<br>National Cancer<br>Institute (NCI) | I      | Phase I trial to study<br>the effectiveness of<br>biological therapy in<br>treating patients<br>who have metastatic<br>cancer that has not<br>responded to<br>previous treatment.  | 2000 | 2009    |
| 50  | Lymphoma                             |                                | tumor specific<br>immune response,<br>control vaccine       | Biovest International                                  | III    | Fast-Track Phase III<br>completed; Pending<br>U.S. and<br>European regulatory<br>applications  | 2000 | 2009    |
| 51  | Lymphoma                             |                                | MyVax   | Genitope<br>Corporation                                | 11/111 | This is a multi-center, open-label, single arm Phase 1/2 study evaluating the feasibility, safety, and tolerability of a series of 16 immunizations of Id-KLH with GM-CSF in patients with previously untreated B-CLL.   | 2006 | Ongoing |
| 52  | Lymphoma                             |                                | Ad-ISF36  | University of<br>California, San Diego                 | 11     | This is a Phase II, open label, fixed dose, repeat injection, single institution study. Eligible subjects will receive up to six doses of Ad-ISF35 injected directly into a selected lymph node under ultrasound guidance. The primary goal is to determine and monitor clinical and biological responses in patients treated with repeat intranodal injections of Ad-ISF35. |      |         |
| 53  | Lymphoma                             |                                | ISF35   | University of<br>California, San<br>Diego; Memgen, LLC | II     | This is a Phase II, open label, fixed dose, repeat injection, single institution study. Eligible subjects will receive up to six doses of Ad-ISF35 injected directly into a selected lymph node under ultrasound guidance. The primary goal is to determine and monitor clinical and biological responses in patients treated with repeat intranodal injections of Ad-ISF35. | 2009 | 2011    |
| 54  | Malignant<br>Pleural<br>Mesothelioma | Celecoxib,<br>cyclophosphamide | K562 (Allogeneic<br>Tumor Cell Vaccine)                     | National Cancer<br>Institute (NCI)                     | 1/11   | To evaluate the safety and effectiveness of tumor cell vaccines in combination with cyclophosphamide and celecoxib in  | 2010 | 2011    |
|     |                                      |                                |   |  |        |  |      |         |

|    |              |            |  |   |      | patients with cancers involving the chest.   |      |      |
|----|--------------|------------|--|---|------|--|------|------|
| 55 | Mesothelioma |            | PSMA/PRAME   | MannKind<br>Corporation   | I    | Completed The present clinical trial is a dose comparison of a multi-component active immunotherapy designed to stimulate an immune reaction to specific tumor associated antigens which are highly expressed on a large number of solid cancers.  | 2007 | 2009 |
| 56 | Metastatic   |            | PG13-MAGE-A3<br>TCR9W11<br>(anti-MAGE-A3/12<br>TCR) Transduced<br>Autologous<br>Peripheral Blood<br>Lymphocytes,<br>Aldesleukin,<br>Cyclophosphamide,<br>Fludarabine | GlaxoSmithKline/NCI   | II   | To evaluate the safety and effectiveness of anti-MAGE-A3/12 lymphocytes as a treatment for metastatic cancers that have not responded to standard treatment  | 2010 | 2012 |
| 57 | Metastatic   |            | Dendritic Cell<br>Vaccination  | Quantum<br>Immunologics   | 1/11 | The study uses a molecule or particle that is found only on cancer cells and is unique to cancer cells, as it is not detected on normal tissue   | 2008 | 2015 |
| 58 | Metastatic   | AlloStim-7 | AlloStim8 or<br>AlloStim-9   | Immunovative<br>Therapies, Ltd.   | 1/11 | This is a Phase I/II study to investigate the feasibility of creating a personalized therapeutic cancer vaccine within the body. A vaccine contains a source of tumor antigen and an adjuvant. In this study, tumor antigen is generated by freezing a tumor by a minimally invasive percutaneous (through the skin) cryoablation procedure. The study drug, AlloStim, is injected into the ablated tumor to promote development of an anti-tumor immune response. | 2009 | 2011 |
| 59 | Metastatic   |            | DC/tumor fusion vaccine  | Beth Israel<br>Deaconess Medical<br>Center, Dana-Farber<br>Cancer Institute | 1/11 | This study aims to determine if the vacccine can be used safely in patients with advanced melanoma (cancer of the pigment cells) and whether the cells in this vaccine are capabale of producing immune responses against your own cancer.   | 2000 | 2008 |
| 60 | Metastatic   |            | carcinoembryonic<br>antigen RNA-pulsed<br>DC cancer vaccine  | Duke University,<br>National Cancer<br>Institute (NCI)                      | I    | Phase I trial to study<br>the effectiveness of<br>biological therapy in<br>treating patients<br>who have metastatic<br>cancer that has not   | 2000 | 2009 |

|    |                              |  |   |      | responded to previous treatment.  |      |      |
|----|------------------------------|--|---|------|---|------|------|
| 61 | Metastatic                   | carcinoembryonic<br>antigen RNA-pulsed<br>DC cancer vaccine                    | Duke University,<br>National Cancer<br>Institute (NCI)                              | 1/11 | Phase I/II trial to study the effectiveness of immunotherapy with CEA-treated white blood cells in treating patients with resected liver metastases from colon cancer.  | 1999 | 2009 |
| 62 | Myelodysplastic<br>Syndromes | Ras peptide cancer vaccine, sargramostim                                       | Memorial<br>Sloan-Kettering<br>Cancer Center,<br>National Cancer<br>Institute (NCI) | I    | Phase I trial to study<br>the effectiveness of<br>vaccine therapy plus<br>sargramostim in<br>treating patients<br>who have<br>myelodysplastic<br>syndrome.  | 1999 | 2001 |
| 63 | Myeloma                      | Dendritic Cell Tumor<br>Fusion   | Beth Israel<br>Deaconess Medical<br>Center, Dana-Farber<br>Cancer Institute         | I    | The main purpose of this study is to test the safety and determine the type and severity of any side effects of the Dendritic Cell Fusion Vaccine given in combination with an autologous transplant for patients with multiple myeloma.  | 2007 | 2011 |
| 64 | Myeloma                      | Telomerase (hTERT vaccine + pneumococcal conjugate vaccine (PCV)), PCV vaccine | University of<br>Pennsylvania   | 1/11 | To evaluate the safety of activated T cell infusions and immunization with hTERT multi-peptide vaccine in the post-transplant setting and whether the combination can delay hematopoietic recovery or induce other autoimmune events.  To determine whether the strategy of infusing vaccine-primed T-cells early after transplant in conjunction with post-transplant boosters leads to the induction of cellular immune responses to hTERT. | 2008 | 2011 |
| 65 | Neoplasm                     | NY-ESO-1b peptide<br>plus CpG 7909 and<br>Montanide ISA-51                     | Ludwig Institute for<br>Cancer Research   | I    | This cancer vaccine research study involves the injection of the NY-ESO-1b peptide along with 2 other agents to help stimulate the immune system.   | 2003 | 2005 |
| 66 | Neuroendocrine               | PSMA/PRAME   | MannKind<br>Corporation   |      | Completed The present clinical trial is a dose comparison of a multi-component active immunotherapy designed to stimulate an immune reaction to specific tumor associated antigens which are highly expressed on a large number of  | 2007 | 2009 |

|    |                            |                            |   |                                       |      | solid cancers.   |      |      |
|----|----------------------------|----------------------------|---|---------------------------------------|------|--|------|------|
| 67 | Peritoneal                 |                            | DPX-Survivac<br>with low dose<br>cyclophosphamide   | ImmunoVaccine<br>Technologies, Inc.   | I/II | This is a phase 1-2 study to determine the safety and immunogenicity profiles of DPX-Survivac, a therapeutic vaccine co-administered with a regimen of low dose oral cyclophosphamide. DPX-Survivac is for the treatment of ovarian, fallopian tube, and peritoneal cancers.                                       |      |      |
| 68 | Peritoneal                 | carboplatin,<br>paclitaxel | MAGE-A1,<br>Her-2/neu, FBP<br>peptides ovarian<br>cancer vaccine;<br>tetanus toxoid<br>helper peptide | University of Virginia;<br>NCI        | II   | This phase II trial is studying how well giving vaccine therapy together with paclitaxel and carboplatin works in treating patients who are undergoing surgery for stage III or stage IV ovarian cancer, primary peritoneal cancer, or fallopian tube cancer.  |      |      |
| 69 | Peritoneal                 |                            | oregovomab;<br>cyclophosphamide   | Gynecologic<br>Oncology Group,<br>NCI | 1/11 | This randomized clinical trial is studying the side effects of oregovomab and to see how well it works with or without cyclophosphamide in treating patients with stage III or stage IV ovarian epithelial cancer, fallopian tube cancer, or primary peritoneal cancer that responded to second-line chemotherapy. |      |      |
| 70 | Pulmonary                  |                            | HSPPC-97  | Antigenics                            | 11   | Antigenics is enrolling patients in a Phase II study testing the feasibility to derive an autologous investigational vaccine (HSPPC-96) from the tumor tissue of patients with resectable non-small cell lung cancer.  | 2003 | 2007 |
| 71 | Squamous Cell<br>Carcinoma |                            | MAGE-A3 HPV-16 vaccine  | University of<br>Maryland             | I    | This study is being done to test the safety of experimental cancer vaccines made of MAGE-A3 and HPV-16 antigens. We also hope to learn what doses of the vaccine will best stimulate the immune system.  | 2009 | 2012 |
| 72 | Testicular                 |                            | PSMA/PRAME  | MannKind<br>Corporation               | I    | Completed The present clinical trial is a dose comparison of a multi-component active immunotherapy  | 2007 | 2009 |

|    |                             |                                |   |  |      | designed to<br>stimulate an<br>immune reaction to<br>specific tumor<br>associated antigens<br>which are highly<br>expressed on a<br>large number of<br>solid cancers.  |      |      |
|----|-----------------------------|--------------------------------|---|--|------|--|------|------|
| 73 | Testicular                  |                                | carcinoembryonic<br>antigen RNA-pulsed<br>DC cancer vaccine                       | Duke University,<br>National Cancer<br>Institute (NCI) | I    | Phase I trial to study<br>the effectiveness of<br>biological therapy in<br>treating patients<br>who have metastatic<br>cancer that has not<br>responded to<br>previous treatment.  | 2000 | 2009 |
| 74 | Thymic<br>Carcinoma         | Celecoxib,<br>cyclophosphamide | K562 (Allogeneic<br>Tumor Cell Vaccine)   | National Cancer<br>Institute (NCI)                     | I/II | To evaluate the safety and effectiveness of tumor cell vaccines in combination with cyclophosphamide and celecoxib in patients with cancers involving the chest.   | 2010 | 2011 |
| 75 | Thymoma                     | Celecoxib,<br>cyclophosphamide | K562 (Allogeneic<br>Tumor Cell Vaccine)   | National Cancer<br>Institute (NCI)                     | I/II | To evaluate the safety and effectiveness of tumor cell vaccines in combination with cyclophosphamide and celecoxib in patients with cancers involving the chest.   | 2010 | 2011 |
| 76 | Thyroid                     |                                | PSMA/PRAME  | MannKind<br>Corporation                                | I    | Completed The present clinical trial is a dose comparison of a multi-component active immunotherapy designed to stimulate an immune reaction to specific tumor associated antigens which are highly expressed on a large number of solid cancers.  | 2007 | 2009 |
| 77 | Tumors                      |                                | NY-ESO-I protein<br>with immune<br>adjuvants CpG 7909<br>and Montanide®<br>ISA-51 | Ludwig Institute for<br>Cancer Research                |      | This is a phase I, open-label, randomized study of NY-ESO-I protein with immune adjuvants CpG 7909 and Montanide® ISA-51 and NY-ESO-I protein 400µg with immune adjuvants CpG 7909 and Montanide® ISA-51 in patients with tumors that often express NY-ESO-1. The primary objective of the study is to define the safety. Secondarily, the study will evaluate whether patients develop a specific immunologic response to the NY-ESO-1 protein. | 2006 | 2006 |
| 78 | Upper GI Tract<br>Carcinoma |                                | V934/V935   | Merck  | I    | Completed.<br>This is a two-part<br>study to test the<br>safety, tolerability,   | 2008 | 2011 |
| •  |                             |                                | ı   |  |      | ı  |      | '    |

|    |         |      |       |   | and immune<br>response for<br>V934/V935 vaccine<br>using a new<br>prime-boost regimen<br>in participants with<br>selected solid<br>tumors. |      |      |
|----|---------|------|-------|---|--|------|------|
| 79 | Vaginal | V503 | Merck | Ш | Expected results in late 2011  | 2010 | 2011 |
| 80 | Vulvar  | V503 | Merck | Ш | Expected results in late 2011  | 2010 | 2011 |

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