	Primary: •Determine the dose-limiting toxicity and the maximum tolerated dose of autologous dendritic cells pulsed with autologous tumor cell lysate in patients with stage III or IV melanoma. /•Determine the safety and tolerability of this therapy in these patients. Secondary: •Determine the immune response, in terms of the type and degree of T-cell proliferation and delayed-type hypersensitivity responses, in patients treated			
	with this therapy. Patients are followed at day 84 and then every 3 months thereafter	Autologica de adritio della (DC) are presente		
	Patients undergo leukapheresis for the collection of peripheral blood mononuclear cells (PBMC) on days -9, 19, and 47.			
	from autologous PBMC exposed to sargramostim (GM-CSF), interleukin-4, and tumor necrosis factor alpha and pulsed v	with autologous tumor cell lysate. Patients		
Completed	Evaluation of Influenza H1N1 Vaccine in Adults With Lymphoid Malignancies on Chemotherapy	Pandemic H1N1(2009) Influenza Vaccine in		
	Conditions: Lymphoma; Multiple Myeloma; Influenza A Virus, H1N1 Subtype	Adults With Lymphoid Malignancies on		
	Intervention: Biological: AS03-adjuvanted H1N1 pandemic influenza vaccine	Active Systemic Treatment or Post Stem		
Active, not	Broad Spectrum HPV (Human Papillomavirus) Vaccine in 16 to 26 Year Old Women (V505-001)			
recruiting	Conditions: Cervical Cancer; Vulvar Cancer; Vaginal Cancer; Genital Warts; Human Papillomavirus Infection	The purpose of this study is to evaluate		
	Biological: Comparator: V505 formulation 1; Drug: Comparator: V505 formulation 2; Biological: Comparator:	the safety and immunogenicity of V505		
	Interventions: V505 formulation 3; Biological: Comparator: Quadrivalent Human Papillomavirus (Types 6, 11, 16, 18)	in comparison to GARDASIL ™		
	Recombinant; Biological: Comparator: Placebo (unspecified)			
Recruiting	Combination of Chemoradiation Therapy and Epitope Peptide Vaccine Therapy in Treating Patients With Esophageal Cancer	Chemoradiation Therapy With Epitope		
	Condition: Esophageal Cancer 2008 慶応大学	Peptide Vaccine Therapy in Treating		
	Intervention: Biological: URLC10, TTK, KOC1, VEGFR1, VEGFR2, cisplatin, fluorouracil	Patients With Unresectable, Advanced or		
	Primary Outcome: •Safety(toxicities as assessed by NCI CTCAE version3) [3 months] Secondary Outcome: •Peptide specific CTL induction [3 months]. /•DTH to peptide [3 months]. /•Changes in levels of response rate as assessed by RECIST criteria [1 year]. /•Time to progression [1 year]. /•survival [1 year] Detailed Description: Up-regulated ling cancer 10 (URLC10), TTK protein kinase (TTK) and K homology domain contai (KOC1) were identified as new targets of tumor associated antigens using cDNA microarray technologies combined with cancer tissues. Furthermore, anti-angiogenic therapy is now considered to be one of promising approaches for treating or receptor 1 (VEGFR1) and vascular endothelial growth factor receptor 2 (VEGFR2) are essential targets for tumor angiog are able to induce cytotoxic T lymphocytes (CTL) restricted to HLA-A *2402 in vivo. On the other hand, chemotherapy (C to be a standard treatment for unresectable advanced esophageal cancer. In this clinical trial, we evaluate the safety and multiple peptides (URLC10, TTK, KOC1, VEGFR1, and VEGFR 2) emulsified with Montanide ISA 51 in combination with therapy in treating patients with unresectable, advanced or recurrent esophageal cancer.	ning protein over expressed in cancer the expression profiles of normal and cancer. Vascular endothelial growth factor enesis. Epitope peptides for these targets EDDP, 5-FU) plus radiation therapy has been immune responses of different doses of chemotherapy (CDDP, 5-FU) plus radiation		
	A Study of V503 in Preadolescents and Adolescents	the Immunogenicity, Tolerability, and		
recruiting	Conditions: Cervical Cancers; Vulvar Cancer; Vaginal Cancer; Genital Lesions; PAP Test Abnormalities; HPV	Manufacturing Consistency of V503 (A		
	Intervention: Biological: V503 2009	Multivalent Human Papillomavirus [HPV		
Completed		Lymphodepleting Conditioning Followed by		
	Condition: Melanoma (Skin) 2004	Infusion of Anti-MART-1 TCR-Gene		
Interventions: Biological: MART-1:27-35 peptide vaccine; Biological: aldesleukin; Biological: filgrastim; Biological: Engineered Lymphocytes and Subs				
	lincomplete Freund's adjuvant: Biological: therapeutic autologous lymphocytes: Biological: therapeutic tumor	Peptide Immunization		

	Primary Outcome: •Safety /•Tumor regression Secondary Outcome: •In vivo survival of transplanted cells /•Clinical response RATIONALE: Inserting a laboratory-treated gene into a person's white blood cells may make the body build an immune response to kill tumor cells. Giving cyclophosphamide and fludarabine before a white blood cell infusion may suppress the immune system and allow tumor cells to be killed. Vaccines may make the body build an immune response to kill tumor cells. Aldesleukin may stimulate a person's white blood cells to kill tumor cells. Combining white blood cell infusion with vaccine therapy and aldesleukin may cause a stronger immune response and kill more tumor cells.		
Active, not recruiting	Vaccine Therapy in Treating Patients With Stage IV Melanoma Condition: Melanoma (Skin) Interventions: Biological: autologous tumor cell vaccine; Biological: therapeutic autologous dendritic cells 2005	Vaccination With Mature, Autologous Monocyte-Derived Dendritic Cells Transfected With Unselected Autologous	
	Primary Outcome Measures: •Safety. /•Immunogenicity. /•Objective tumor response. /•Time to disease progression /•Progression-free interval /•Overall survival OBJECTIVES: •Determine the safety and tolerability of vaccine therapy comprising autologous dendritic cells (DC) transfected with autologous polymerase chain reaction-amplified tumor RNA in patients with stage IV cutaneous melanoma. /•Determine whether tumor RNA- or tumor antigen-specific T-cell responses are induced in patients treated with this vaccine. /•Determine whether there are major differences in the immunogenicity of DC transfected at immature stage or at mature stage in patients treated with this vaccine. •Determine objective tumor response in patients treated with this vaccine. /•Determine time to disease progression and progression-free interval in patients treated with this vaccine. •Determine overall survival of patients treated with this vaccine. Patients are followed periodically for up to 10 years.		
Active, not recruiting	Vaccine Therapy With Immune Adjuvant in Treating Patients With Stage IIB, Stage IIC, Stage III, or Stage IV Melanoma Condition: Melanoma (Skin) Interventions: Biological: gp100 antigen; Biological: sargramostim plasmid DNA melanoma vaccine adjuvant; Biological: tyrosinase peptide 2004	A Multi-Epitope Peptide Vaccine Using GM- CSF DNA As An Adjuvant: A Pilot Trial To Assess Safety And Immunity	
	Primary Outcome: •Immunological efficacy in terms of T-cell response as measured by enzyme-linked immunospot. Primary: •Determine the maximum tolerated dose and recommended dose of sargramostim (GM-CSF) plasmid DNA accomprising tyrosinase peptide and gp100 antigen in patients with stage IIB, IIC, III, or IV melanoma who are HLA-A2-pose in these patients. •Determine the pharmacokinetics of this regimen in these patients. /•Determine the dose-limiting toxic effects of this regimenes the immunogenicity of this regimen in these patients.	itive. /•Determine the safety of this regimen	
Completed	Peptide Vaccinations to Treat Patients With Low-Risk Myeloid Cancers Conditions: Myelodysplastic Syndrome (MDS); Acute Myeloid Leukemia (AML); Chronic Myeloid Leukemia (CML) Interventions: Biological: WT1:126-134 Peptide; Biological: PR1:169-177 Peptide; Drug: WT1 and PR1 Peptide Vaccines; Drug: GM-CSF (Sargramostim); Biological: WT1 and PR1 Peptide Vaccines 2007	WT1 and PRI Peptide Vaccination for Patients With Low Risk Myeloid Malignancies	

	Primary Outcome: •The efficacy and toxicity associated with 6 doses of a combination of WT-1:126-134 and PR1:169-177 peptide vaccines for myeloid malignancies. Secondary Outcome: •Changes in marrow blast cells, blood counts, transfusion dependence, time to disease progression, survival and response to booster vaccination. Therefore we propose this Phase II trial, the third in a series of planned peptide vaccine research protocols, which will evaluate the safety and efficacy associated with an immunotherapy approach using two peptide vaccines, namely PR 1 : 169- 177 and WT-1: 126-1 34 in Montanide adjuvant, administered concomitantly with			
Completed Has Results	GM-CSF (Sargramostim), every 2 weeks for 10 weeks (6 doses WT1 plus 6 doses PRI plus GM-CSF) in select patients Evaluation of Safety and Immunogenicity of Co-administering Human Papillomavirus (HPV) Vaccine With Other Vaccines in Conditions: Cervical Intraepithelial Neoplasia; Papillomavirus Vaccines; Human Papillomavirus Infection Interventions: Biological: Boostrix ® Polio; Biological: GSK Biologicals' HPV-16/18 L1 AS04 vaccine (Cervarix TM)	diagnosed with MDS, AML or CML. Subjects Evaluate the Immunogenicity and Safety of GSK Biologicals' HPV Vaccine (580299) Co-administered With Boostrix Polio (dTpa-IPV)		
Completed	Vaccine Plus Interleukin-2 in Treating Patients With Advanced Melanoma Condition: Melanoma (Skin) 2000 Interventions: Biological: aldesleukin; Biological: gp100 antigen; Biological: incomplete Freund's adjuvant OBJECTIVES: •Determine clinical response rates in patients with advanced melanoma treated with gp100:209-217(210) interleukin-2.	Melanoma Vaccine (NSC #683472/675756, IND #6123) and Low-Dose, Subcutaneous Interleukin-2 in Advanced Melanoma M) melanoma vaccine and low-dose		
Suspended	M-Vax + Low Dose Interleukin-2 Versus Placebo Vaccine in Metastatic Melanoma in Patients With Stage IV Melanoma Condition: Melanoma Intervention: Biological: M-Vax- autologous, hapten-modified melanoma vaccine 2007	M-Vax Plus Low Dose Interleukin-2 Versus Placebo Vaccine Plus Low Dose Interleukin- 2 for Stage IV Melanoma		
	Primary Outcome: •Best overall anti-tumor response. [Time Frame: 1 year] /•Survival - % patients surviving at two year Secondary Outcome: •Safety [Time Frame: 5 years] The primary endpoints of the study are: 1)Best overall anti-tumor response, and 2)Survival, measured by % surviving at tumor response by modified RECIST criteria between weeks 24 and 25 (i.e., 5-6 weeks after completion of IL2). At the 6 will receive an additional single booster dose of M-Vax or Placebo Vaccine mixed with BCG. This will be followed by four evaluations for anti-tumor response will take place at the 38-39 week (month 9) and one-year points. Then patients will be adverse events until evidence of tumor progression that requires new therapy. Patients who remain on-study will be followers.	two years. Patients will be evaluated for anti- month point patients who remain on study more courses of IL2. Two additional re regularly evaluated for tumor status and		
Active, not recruiting	Vaccine Therapy and Ganciclovir in Treating Patients With Mesothelioma Condition: Malignant Mesothelioma Biological: PA-1-STK ovarian carcinoma vaccine; Drug: ganciclovir 2000	Treatment of Malignant Pleural Mesothelioma With Gene Modified Cancer Cell Lines		
	OBJECTIVES: I. Determine the safety and side effects of intrapleurally administered PA-1-STK modified ovarian carcino stage I, II, or III malignant mesothelioma. II. Determine the maximum tolerated dose and dose limiting toxicities of this valimmunologic response to this treatment regimen in these patients. IV. Determine the intrapleural pharmakokinetics of ga OUTLINE: This is a dose escalation study of PA-1-STK modified ovarian carcinoma vaccine. Patients receive PA-1-STK intrapleurally on day 1 followed by ganciclovir IV over 1 hour for 7 days beginning on day 1. Patients in the first 2 cohorts subsequent cohorts, treatment repeats every 3 weeks for a total of 3 courses in the absence of disease progression or u receive escalating doses of PA-1-STK modified ovarian carcinoma vaccine until the maximum tolerated dose is determine 16 patients will be accrued for this study.	ccine in these patients. III. Determine the nciclovir in these patients. modified ovarian carcinoma vaccine receive 1 course of treatment only. In all nacceptable toxicity. Cohorts of 3 patients		

Completed	Vaccine Therapy in Treating Patients With Metastatic Melanoma	Intradermally Administered MART-			
'	Conditions: Intraocular Melanoma; Melanoma (Skin)	1gp100/Tyrosinase Peptide-Pulsed Dendritic			
	Riclogical: MART-1 antigon: Riclogical: gp100:200-217(210M) partide veccine: Riclogical: they proutice	Cell Vaccine Matured With a Cytokine			
	Interventions: autologous dendritic cells; Biological: tyrosinase peptide 2007	Cocktail for Metastatic Melanoma			
	Primary Outcome: •Overall survival. /•Progression-free survival / •Time to progression /•Toxicity.				
	Primary: •Determine clinical response in HLA-A *0201-positive patients with metastatic melanoma treated with an intrade	ermally administered vaccine comprising			
	autologous dendritic cells pulsed with MART-1, gp100, and tyrosinase peptides and matured with a cytokine cocktail.	,			
	Secondary: •Determine immunologic response in patients treated with this regimen.				
Active, not	Extension Study of the Efficacy of the GSK 580299 Vaccine in Japanese Women Vaccinated in the Primary NCT00316693	the 580299 Vaccine in the Prevention of			
recruiting	Condition: Human Papillomavirus Infection	HPV-16 and/or HPV-18 Associated Cervical			
	Interventions: Procedure: Blood sampling; Procedure: Liquid-based cytology (LBC) sampling	Intraepithelial Neoplasia (CIN) in Japanese			
	A Study to Evaluate the Safety, Immune Response, and Efficacy of Gardasil (V501) in Women				
recruiting					
Has Results	Conditions: Healthy; Papillomavirus Infection				
	Interventional Biological: Quadrivalent Human Papillomavirus (Types 6, 11, 16, 18) Recombinant Vaccine; Biological:	-			
	Interventions: Comparator: Placebo				
Active, not	Vaccine Therapy in Treating Patients Who Have Received First-Line Therapy for Hodgkin's Lymphoma	KGEL Vaccine After Initial Therapy of			
recruiting	Condition: Lymphoma	1			
	Interventions: Biological: Hodgkin's antigens-GM-CSF-expressing cell vaccine; Procedure: adjuvant therapy 2007	Hodgkin's Lymphoma			
	Primary Outcome: •Immunologic response. /•Durability of immunologic response. /•Utility of Epstein-Barr virus reporter	system for monitoring cellular vaccine			
	responses. /•Safety and tolerability				
	•Determine immunologic responses in patients who have completed first-line therapy for Hodgkin's lymphoma treated with Hodgkin's antigens-GM-CSF-expressing				
	Icell vaccine.				
	•Determine the durability of these immunologic responses in these patients. /•Determine the utility of an Epstein-Barr virus reporter system for monitoring cellular				
	vaccine responses.				
	•Determine the safety and tolerability of this vaccine in these patients. /OUTLINE: Beginning 4-6 months after last chemotherapy, patients receive Hodgkin's				
	antigens-GM-CSF-expressing cell vaccine on day 1. Treatment repeats every 3 weeks for up to 4 courses. /Immunologic responses are serially monitored along with				
	disease status				
Completed	Immunogenicity and Safety of GlaxoSmithKline Biologicals' HPV Vaccine 580299 in Healthy Females 10 - 25 Years of Age.				
	Conditions: HPV-16/18 Infections; Papillomavirus Vaccines; Cervical Neoplasia				
	Interventions: Biological: CervarixTM; Biological: Placebo vaccine (Al(OH)3)				
Terminated	Vaccine Therapy Following Chemotherapy and Peripheral Stem Cell Transplantation in Treating Patients With Non-Hodgkin's	Evaluate Immune Response Using Idiotype			
Has Results	Condition: Lymphoma	Vaccines Following High-Dose			
	Riological: autologous tumor cell vaccine: Riological: keyhole limnet hemocyanin: Riological: sargramostim:	Chemotherapy and Hematopoietic Stem Cell			
	Interventions: Procedure: adjuvant therapy 2000	Transplantation for Follicular Lymphoma			

Primary Outcome: •Humoral and Cellular Immune Response. /evaluate the humoral immune responses and cellular immune responses to idiotype vaccine with KLH and GM-CSF adjuvant given to patients with follicular lymphoma following high-dose chemotherapy and autologous stem cell transplantation Secondary Outcome: •Safety. /To evaluate the safety and toxicity of idiotype vaccine with KLH and GM-CSF adjuvant in the post-transplant setting /•Toxicity. /To evaluate the safety and toxicity of idiotype vaccine with KLH and GM-CSF adjuvant in the post-transplant setting /•Changes in Quantitative Bcl-2 [Time Frame: 1 year]. To evaluate changes in quantitative bcl-2 of the blood and bone marrow prior to and at various time points following the series of idiotype vaccines. OBJECTIVES: •Determine the humoral and cellular immune responses in patients with follicular non-Hodgkin's lymphoma treated with autologous lymphomaderived idiotype vaccine with keyhole limpet hemocyanin plus sargramostim (GM-CSF). *Determine the safety and toxicity of this regimen in these patients in the post-transplant setting. •Determine the changes in quantitative bcl-2 in the blood and bone marrow of these patients before and at various times after the series of idiotype vaccines. Patients are followed every 3 months for 2 years, every 6 months for 2 years Recruiting Safety Study of DNA Vaccine Delivered by Intradermal Electroporation to Treat Colorectal Cancer Immunogenicity of Intradermal Condition: Colorectal Cancer Electroporation of tetwtCEA DNA in Patients Interventions: Biological: tetwtCEA DNA (wt CEA with tetanus toxoid Th epitope); Device: Derma Vax (electroporation With Colorectal Cancer device); Biological: GM-CSF; Drug: Cyclophosphamide 2010 Primary Outcome: •To evaluate the safety and immunogenicity of a DNA immunisation approach where tetwtCEA DNA will be administered in combination with electroporation. Secondary Outcome: •To assess the efficiency of priming immunological responses to CEA by intradermal administration of tetwtCEA DNA in combination with electroporation. •To assess the efficiency of boosting immunological responses to CEA by intradermal administration of tetwtCEA DNA in combination with electroporation in subjects already vaccinated with CEA DNA /•To compare effects (safety and immunogenicity) of additional adjuvance with GM-CSF. The purpose of this study is to evaluate the safety and immunogenicity of a CEA DNA immunisation approach in patients with colorectal cancer. The DNA plasmid, tetwtCEA, encodes wild type human CEA fused to a tetanus toxoid T helper epitope. The vaccine will be delivered using an intradermal electroporation device. Derma Vax (Cyto Pulse Sciences). The following will be assessed: Completed Human Papilloma Virus (HPV) Vaccine Immunogenicity and Safety Trial in Young and Adult Women With GSK Biologicals' HPV-Has Results Conditions: Cervical Intraepithelial Neoplasia: Human Papillomavirus Infection Intervention: Biological: Cervarix™ Suspended GM-CSF With or Without Vaccine Therapy After Combination Chemotherapy and Rituximab as First-Line Therapy in Treating Lymphoma Double-Blind, Randomized, Placebo-Condition: Controlled Trial of FavID® (Id/KLH) and GM-Drug: autologous immunoglobulin idiotype-KLH conjugate vaccine; Drug: cyclophosphamide; Drug: CSF Following CHOP/Rituximab as Firstdoxorubicin hydrochloride: Drug: prednisone: Drug: rituximab: Drug: sargramostim: Drug: vincristine: Line Therapy in Subjects With High-Procedure: Intervention/procedure; Procedure: antibody therapy; Procedure: biological therapy; Procedure: Interventions Intermediate and High-Risk Diffuse Large Bchemotherapy; Procedure: colony-stimulating factor therapy; Procedure: cytokine therapy; Procedure: Cell Lymphoma monoclonal antibody therapy; Procedure: non-specific immune-modulator therapy; Procedure: therapeutic procedure: Procedure: tumor cell derivative vaccine: Procedure: vaccine therapy

	Primary: •Compare the 3-year disease-free survival of patients with high-intermediate- or high-risk bulky stage II or stage III or IV diffuse large B-cell lymphoma treated with sargramostim (GM-CSF) with or without autologous immunoglobulin idiotype-KLH conjugate vaccine (FavId®) after combination chemotherapy comprising cyclophosphamide, doxorubicin, vincristine, prednisone, and rituximab (CHOP-R).			
	Secondary: •Compare the 2-year disease-free survival, duration of response, time to progression, overall survival, and so	afety in patients treated with these		
	regimens.	arety in patiente areated than arese		
	•Estimate the rate of immune reactivity to FavId®. Patients are followed periodically for up to 2 years			
Active, not				
recruiting	Condition: Melanoma (Skin)	Peptide Based Vaccine Therapy in Patients		
	Interventions: Biological: MAGE-10.A2; Biological: MART-1 antigen; Biological: NY-ESO-1 peptide vaccine; Biological: sargramostim; Biological: tyrosinase peptide 2002	With High-Risk or Metastatic Melanoma		
	OBJECTIVES: •Compare the safety of melanoma peptide vaccine with or without sargramostim (GM-CSF) in patients with	th high-risk or metastatic melanoma.		
	•Compare changes in peptide-specific cellular and humoral immunologic profiles in patients treated with these regimens.			
	•Compare tumor response in patients treated with these regimens.			
	OUTLINE: This is a randomized, open-label study. Patients are randomized to 1 of 2 treatment arms.			
	•Arm I: Patients receive melanoma peptide vaccine comprising tyrosinase leader injected at 2 separate sites, Melan-A ELA injected at another site, NY-ESO-1a and			
	NY-ESO-1b combined and injected at one site, and MAGE-10.A2 injected at another site, intradermally once weekly on weeks 1-6.			
	•Arm II: Patients receive vaccine as in arm I. Patients also receive sargramostim (GM-CSF) subcutaneously daily beginning	ing 2 days before each vaccination and		
	continuing for 5 days.			
	Vaccine Therapy With or Without Interleukin-12 in Treating Patients With Stage III or Stage IV Melanoma	Vaccine Combining Tyrosinase/gp100		
recruiting	Conditions: Intraocular Melanoma; Melanoma (Skin)	Peptides Emulsified With Montanide ISA 51		
	Interventions: Biological: gp100 antigen; Biological: incomplete Freund's adjuvant; Biological: recombinant interleukin-12; Biological: tyrosinase peptide 1999	With and Without Interleukin-12 for Patients With Resected Stages III and IV Melanoma		
	Detailed Description: OBJECTIVES: I. Evaluate immune reactivity to tyrosinase and gp100 peptides emulsified with Montanide ISA-51 (ISA-51) with or without interleukin-12 following surgical resection in HLA-A2 positive patients with stage III or IV melanoma. OUTLINE: This is a randomized, parallel study. Patients are stratified by prior therapy (immunotherapy or chemotherapy vs surgery only). Patients are randomized to			
	receive 1 of 2 treatment arms: Arm I: Following surgery, patients receive tyrosinase and gp100 peptides emulsified with Montanide ISA-51 (ISA-51) subcutaneously			
	(SQ) once weekly during weeks 0, 2, 4, 6, 10, 14, 18, and 26 for a total of 8 vaccinations. Arm II: Following surgery, patients receive treatment as in Arm I followed			
	by interleukin-12 SQ once weekly during weeks 0, 2, 4, 6, 10, 14, 18, and 26 for a total of 8 vaccinations. Patients are fol 2 years after resection, then every 6 months for 3 years, and then yearly if without evidence of disease.	lowed at 2-4 weeks, then every 3 months to		
Recruiting	Imatinib Mesylate, Interferon Alfa, and GM-CSF Compared With Imatinib Mesylate and Vaccine Therapy in Treating Patients			
	Condition: Leukemia 2006	Interferon + GM-CSF Versus K562/GM-		
	Interventions: Biological: GM-K562 cell vaccine; Biological: recombinant interferon alfa; Biological: sargramostim	CSF Vaccination for CML		
	The following the first transfer the first transfer the first transfer the first transfer tra	1		

Primary Outcome: •Progression-free survival at 1 year. /•Rate of molecular complete remission. Secondary Outcome: •Time to Philadelphia chromosome (Ph) negativity as measured by polymerase chain reaction /•Disease-free survival. /•Percent molecular complete remission. /•Toxicity . /•Time to progression. Primary: •Compare clinical response, in terms of 1-year progression-free survival and rate of molecular complete remission, in patients with Philadelphia chromosome-positive chronic myelogenous leukemia (Ph+ CML) in chronic phase who have achieved a complete cytogenetic remission to single-agent imatinib mesylate treated with imatinib mesylate, interferon alfa, and sargramostim (GM-CSF) vs imatinib mesylate and GM-K562 cell vaccine. Secondary: •Compare time to Ph-negativity by polymerase chain reaction after randomization. /•Compare disease-free survival and percent molecular complete remissions. / Determine the toxicity of these treatment regimens in these patients. Patients are followed periodically for up to 1 year. Active, not Vaccine Therapy in Treating Patients With Stage IV Melanoma recruiting Condition: Melanoma (Skin) 1999 Phase I Trial of a Dendritic Cell Vaccine for Biological: dendritic cell-MART-1 peptide vaccine; Biological: gp100 antigen; Biological: therapeutic tumor Melanoma Interventions: infiltrating lymphocytes: Biological: tyrosinase peptide OBJECTIVES: I. Determine the dose-limiting toxicities, maximum tolerated dose, recommended phase II dose, and rate of sensitization of T cells at each dose level in patients with melanoma receiving dendritic cell vaccine. II. Determine the overall (complete and partial) response rate, duration of response, and optimal route of administration in this patient population. OUTLINE: This is a dose escalation study. Patients are randomized to one of three treatment arms. All patients undergo leukopheresis to obtain lymphocyte and myeloid origin mononuclear cell fractions for preparation of dendritic cell (DC) vaccine. In each arm, cohorts of up to 5 patients receive escalating doses of vaccine. The maximum tolerated dose (MTD) is defined as the dose preceding that at which 2 or more of 5 patients experience dose-limiting toxicity. Randomization ceases if the MTD has been reached in 2 arms, although accrual may continue. Treatment repeats every 2 weeks for a total of 4 doses. Arm I: Patients receive 3 different doses of peptide pulsed DC vaccine IV, each divided into 3 different peptide pulsed pools administered over 30 minutes. Arm II: Patients receive 3 different doses of peptide pulsed DC vaccine subcutaneously/intradermally to sites with no evidence of disease. At the lowest dose, patients receive 3 different peptide pulsed pools, each administered at a separate site. At the higher doses, patients receive 3 injections further subdivided into 6 and administered at 6 distinct sites. Arm III: Patients receive peptide pulsed DC vaccine intranodally in groin or ancillary lymph nodes at the lower 2 doses of the 3 administered to arms I and II. At the lower dose, patients receive 3 different peptide pulsed pools, each administered into a different node. At the higher dose, patients receive 3 injections further subdivided into 6 and administered at 6 distinct sites. Patients are followed at 2 weeks and then monthly for 3 months. Recruiting Evaluating the Safety and the Biological Effects of Intratumoral Interferon Gamma and a Peptide-Based Vaccine in Patients Intratumoral Injection of Interferon Gamma During Vaccination in Patients Condition: Melanoma 2009 Intervention: Biological: A combination of intratumoral IFN-gamma plus systemic vaccination with MELITAC 12.1 With Subcutaneous or Cutaneous

	Primary Outcome: *Safety: To determine the safety of administration of intratumoral interferon gamma with a peptide-based vaccine in patients with cutaneous or subcutaneous metastases of melanoma. [6 months]. /*Biologic effect: To evaluate the biological effects of vaccine plus IFN-gamma at the tumor site, to include expression of CXCR3 ligands (CXCL9, CXCL10 & CXCL11) and the magnitude of infiltration of CD8+ CXCR3+ T cells and vaccine-specific T cells. [6 months] Secondary Outcome: •To estimate the effects of vaccine on CXCR3 expression by circulating an interferon cD4 and CD8 T cells. [6 months]. /*To estimate the effects of vaccine plus IFN-gamma on changes in the percentage of FoxP3+ CD25hi CD4+ (putative regulatory T cells, Tregs) among tumor infiltrating T cells. [6 months]. /*To obtain preliminary data on the variability of immunologic parameters among multiple biopsies of subcutaneous or cutaneous metastases of melanoma. [6 months] /*To obtain preliminary data on the clinical response of cutaneous or subcutaneous metastases of melanoma to the proposed combination regimen. [6 months] It is generally agreed that one mechanism to improve the immunologic outcomes of vaccine therapy is to optimize T cell trafficking to the tumor site. CXCR3 is the chemokine receptor on T cells which directs them to sites of inflammation by following the chemokine gradient. The ligands for CXCR3 (CXCL9 (MIG), CXCL10 (IP 10) and CXCL11 (I-TAC)) are known to be induced by interferon gamma. This protocol proposes administering a peptide vaccine to activate tumor antigen-specific		
	CD8+ T cells expressing CXCR3, followed by intratumoral interferon gamma to increase CXCR3 ligands (CXCL9-11) at least	the tumor site and recruit the CXCR3+ T	
Recruiting	Rituximab and Cyclophosphamide Followed by Vaccine Therapy in Treating Patients With Relapsed Hodgkin Lymphoma Condition: Lymphoma Interventions: Biological: Hodgkin's antigens-GM-CSF-expressing cell vaccine; Biological: filgrastim; Biological: rituximab; Drug: cyclophosphamide	Rituximab, High Dose Cyclophosphamide, and GM-CSF Based Immunotherapy for Relapsed Hodgkin's Lymphoma	
	Primary: •Determine the safety and tolerability of rituximab and high-dose cyclophosphamide followed by vaccine the that expresses Hodgkin's tumor antigens and sargramostim (GM-CSF) (KGEL vaccine) as salvage therapy in patient Determine the immunologic response to this vaccine in these patients. Secondary: •Determine the 3-year relapse-free and overall survival of patients treated with this regimen./•Determine the in patients treated with this regimen. Patients receive rituximab IV on days -10 and -7 and then on days 29, 36, 43, and 50 (weeks 4-7) and high-dose (transpatients) to 0 without stem cell rescue. Patients receive filgrastim (G-CSF) subcutaneously once daily beginning on day 6 and contents also receive vaccine therapy comprising an allogeneic vaccine that expresses Hodgkin's tumor antigens and GM	patterns of cellular immune reconstitution asplant-dose) cyclophosphamide IV on days continuing until blood counts recover.	
Active, not recruiting	Vaccine Therapy and Imatinib Mesylate in Treating Patients With Chronic Phase Chronic Myelogenous Leukemia Condition: Leukemia Interventions: Biological: GM-K562 cell vaccine; Drug: imatinib mesylate	Vaccination for CML Patients With Persistent Disease on Imatinib Mesylate	
Active, not	Primary: •Determine the maximum tolerated dose of GM-K562 cell vaccine when administered with imatinib mesylate in chronic myelogenous leukemia in first hematologic response. /•Determine the safety and toxic effects of GM-K562 cell vaccine myelogenous leukemia in first hematologic response. /•Determine the safety and toxic effects of GM-K562 cell vaccine secondary: •Determine the disease response by serial BCR-ABL quantitative polymerase chain reaction measurements. Determine the development of tumor immunity in patients treated with this regimen. /OUTLINE: This is a dose-escalation Patients continue to receive oral imatinib mesylate at the same stable dose as before study entry. Patients receive GM-K43, 57, 85, 113, and 141 in the absence of disease progression or unacceptable toxicity. Cohorts of 10 patients receive escalating doses of GM-K562 until the maximum tolerated dose (MTD) is determined. The at which 3 of 10 patients experience dose-limiting toxicity. Patients are followed periodically for 20 years. Broad Spectrum HPV (Human Papillomavirus) Vaccine Study in 16-to 26-Year-Old Women (V503-001)	accination in patients. s in patients treated with this regimen. /• n study of GM-K562. 562 subcutaneously on days 1, 8, 15, 29,	
1		Toggie Dillided (Midi III Flodse Dillidillg/,	

	Conditions:	Cervical Cancer; Vulvar Cancer; Vaginal Cancer; Genital Warts; Human Papillomavirus Infection	Controlled With GARDASIL, Dose-
	Interventions:	Biological: Comparator: GARDASIL(R); Biological: Comparator: V503	Ranging, Tolerability, Immunogenicity,
Withdrawn	DCVax-L Vacci	nation With CD3/CD28 Costimulated Autologous T-Cells for Recurrent Ovarian or Primary Peritoneal Cancer	Maintenance Vaccination Combined With
	Conditions:	Ovarian Cancer; Primary Peritoneal Cancer	Metronomic Cyclophosphamide w/wo Adoptive
		Biological: DCVax-L and T Cells 2008	Transfer of CD3/CD28-CoStimulated T-Cells for
	Intervention:		Recurrent Ovarian or Primary Peritoneal Cancer Previously Vaccinated DCVax-L
	DCVax-L, an a cyclophosphar autologous T c Primary Object blood T cells Phase II: Twe •ARM-IIA: main	recurrent epithelial ovarian carcinoma or primary peritoneal cancer, who have previously undergone vaccination autologous vaccine with DC loaded in vitro with autologous tumor lysate. Phase I Subjects enrolled in this mide/fludarabine-induced lymphodepletion; followed by adoptive transfer of ex vivo CD3/CD28-costimulate cells; followed by a single DCVax-L vaccination, to establish feasibility and safety of this approach. Cives of Phase I: To determine the feasibility and safety of administering vaccine-primed, ex vivo CD3/C in combination with DCVax-L vaccination, following lymphodepletion with high dose cyclophosphamical mity-two additional subjects will be randomized to receive either: Intenance DCVax-L vaccination, in combination with oral metronomic cyclophosphamide, or	study will receive leukapheresis; followed by d vaccine-primed peripheral blood D28-costimulated autologous peripheral
Action	vaccine-primed Primary Object oral metronom lymphodepletic Primary Outco Rates of diseas	apheresis, followed by cyclophosphamide/fludarabine-induced lymphodepletion, followed by adoptive tran diperipheral blood autologous T cells, followed by maintenance DCVax-L vaccination, plus oral metronomicative of Phase II: To assess the distribution of progression-free survival at 6 months for patients treated wite cyclophosphamide as well as patients treated with ex vivo CD3/CD28-costimulated vaccine-primed peripon with high dose cyclophosphamide / fludarabine, followed by DCVax-L boost vaccination and metronomic *Disease status will be assessed with CT (or MRI) of chest/abdomen/pelvis at enrollment, after vaccinate progression will be recorded at the time of study conclusion. [3 months after enrollment]	c cyclophosphamide. with maintenance DCVax-L vaccination plus pheral blood autologous T cells after c oral cyclophosphamide. The 2 and at the conclusion of the study.
	vaccine-primed Primary Object oral metronom lymphodepletic Primary Outco Rates of diseas	apheresis, followed by cyclophosphamide/fludarabine-induced lymphodepletion, followed by adoptive tran diperipheral blood autologous T cells, followed by maintenance DCVax-L vaccination, plus oral metronomicative of Phase II: To assess the distribution of progression-free survival at 6 months for patients treated with cyclophosphamide as well as patients treated with ex vivo CD3/CD28-costimulated vaccine-primed peripon with high dose cyclophosphamide / fludarabine, followed by DCVax-L boost vaccination and metronomic recorded at the time of study conclusion. [3 months after enrollment] 3 in Females 12–26 Years of Age Who Have Previously Received GARDASIL™ (V503–006)	c cyclophosphamide. with maintenance DCVax-L vaccination plus pheral blood autologous T cells after c oral cyclophosphamide. ne 2 and at the conclusion of the study. Placebo-Controlled, Double-Blind Clinical
Active, not recruiting	vaccine-primed Primary Object oral metronom lymphodepletic Primary Outco Rates of diseas	apheresis, followed by cyclophosphamide/fludarabine-induced lymphodepletion, followed by adoptive tran diperipheral blood autologous T cells, followed by maintenance DCVax-L vaccination, plus oral metronomicative of Phase II: To assess the distribution of progression-free survival at 6 months for patients treated wite cyclophosphamide as well as patients treated with ex vivo CD3/CD28-costimulated vaccine-primed peripon with high dose cyclophosphamide / fludarabine, followed by DCVax-L boost vaccination and metronomic *Disease status will be assessed with CT (or MRI) of chest/abdomen/pelvis at enrollment, after vaccinate progression will be recorded at the time of study conclusion. [3 months after enrollment]	c cyclophosphamide. vith maintenance DCVax-L vaccination plus pheral blood autologous T cells after c oral cyclophosphamide. ne 2 and at the conclusion of the study. Placebo-Controlled, Double-Blind Clinical
	vaccine-primed Primary Object oral metronom lymphodepletic Primary Outcoon Rates of disease A Study of V500 Conditions:	apheresis, followed by cyclophosphamide/fludarabine-induced lymphodepletion, followed by adoptive tran diperipheral blood autologous T cells, followed by maintenance DCVax-L vaccination, plus oral metronomicative of Phase II: To assess the distribution of progression-free survival at 6 months for patients treated with cyclophosphamide as well as patients treated with ex vivo CD3/CD28-costimulated vaccine-primed peripon with high dose cyclophosphamide / fludarabine, followed by DCVax-L boost vaccination and metronomic recorded at the time of study conclusion. [3 months after enrollment] 3 in Females 12–26 Years of Age Who Have Previously Received GARDASIL™ (V503–006)	c cyclophosphamide. with maintenance DCVax-L vaccination plus pheral blood autologous T cells after c oral cyclophosphamide. ne 2 and at the conclusion of the study. Placebo-Controlled, Double-Blind Clinical Trial to Study the Tolerability and
recruiting Not yet	vaccine-primed Primary Object oral metronom lymphodepletic Primary Outcook Rates of disease A Study of V500 Conditions: Interventions:	apheresis, followed by cyclophosphamide/fludarabine-induced lymphodepletion, followed by adoptive tran disperipheral blood autologous T cells, followed by maintenance DCVax-L vaccination, plus oral metronomic crive of Phase II: To assess the distribution of progression-free survival at 6 months for patients treated with cyclophosphamide as well as patients treated with ex vivo CD3/CD28-costimulated vaccine-primed peripon with high dose cyclophosphamide / fludarabine, followed by DCVax-L boost vaccination and metronomic me: •Disease status will be assessed with CT (or MRI) of chest/abdomen/pelvis at enrollment, after vaccing progression will be recorded at the time of study conclusion. [3 months after enrollment] 3 in Females 12–26 Years of Age Who Have Previously Received GARDASIL™ (V503–006) Cervical Cancers; Vulvar Cancers; Vaginal Cancers; Genital Warts; Human Papillomavirus (HPV) Infection	c cyclophosphamide. vith maintenance DCVax-L vaccination plus pheral blood autologous T cells after c oral cyclophosphamide. ne 2 and at the conclusion of the study. Placebo-Controlled, Double-Blind Clinical Trial to Study the Tolerability and Immunogenicity of V503, a Multivalent Human Papillomavirus
recruiting	vaccine-primed Primary Object oral metronom lymphodepletic Primary Outcook Rates of disease A Study of V500 Conditions: Interventions: Feasibility of Au	apheresis, followed by cyclophosphamide/fludarabine-induced lymphodepletion, followed by adoptive tran deperipheral blood autologous T cells, followed by maintenance DCVax-L vaccination, plus oral metronomic properties of Phase II: To assess the distribution of progression-free survival at 6 months for patients treated with ex vivo CD3/CD28-costimulated vaccine-primed periphon with high dose cyclophosphamide / fludarabine, followed by DCVax-L boost vaccination and metronomic me: •Disease status will be assessed with CT (or MRI) of chest/abdomen/pelvis at enrollment, after vaccing progression will be recorded at the time of study conclusion. [3 months after enrollment] 3 in Females 12–26 Years of Age Who Have Previously Received GARDASIL™ (V503–006) Cervical Cancers; Vulvar Cancers; Vaginal Cancers; Genital Warts; Human Papillomavirus (HPV) Infection Biological: V503; Biological: Comparator: Placebo to V503 2010	c cyclophosphamide. vith maintenance DCVax-L vaccination plus pheral blood autologous T cells after c oral cyclophosphamide. ne 2 and at the conclusion of the study. Placebo-Controlled, Double-Blind Clinical Trial to Study the Tolerability and Immunogenicity of V503, a Multivalent
recruiting Not yet	vaccine-primed Primary Object oral metronom lymphodepletic Primary Outco Rates of disease A Study of V50 Conditions: Interventions: Feasibility of Au Conditions:	apheresis, followed by cyclophosphamide/fludarabine-induced lymphodepletion, followed by adoptive tran disperipheral blood autologous T cells, followed by maintenance DCVax-L vaccination, plus oral metronomic ctive of Phase II: To assess the distribution of progression-free survival at 6 months for patients treated with cyclophosphamide as well as patients treated with ex vivo CD3/CD28-costimulated vaccine-primed peripon with high dose cyclophosphamide / fludarabine, followed by DCVax-L boost vaccination and metronomic receive setatus will be assessed with CT (or MRI) of chest/abdomen/pelvis at enrollment, after vaccing progression will be recorded at the time of study conclusion. [3 months after enrollment] 3 in Females 12−26 Years of Age Who Have Previously Received GARDASIL™ (V503−006) Cervical Cancers; Vulvar Cancers; Vaginal Cancers; Genital Warts; Human Papillomavirus (HPV) Infection Biological: V503; Biological: Comparator: Placebo to V503 2010 tologous Tumor Cell−TLR9 Agonist Vaccination for Metastatic Colorectal Cancer	c cyclophosphamide. with maintenance DCVax-L vaccination plus pheral blood autologous T cells after c oral cyclophosphamide. ne 2 and at the conclusion of the study. Placebo-Controlled, Double-Blind Clinical Trial to Study the Tolerability and Immunogenicity of V503, a Multivalent Human Papillomavirus Autologous Tumor Cell-TLR9 Agonist

Recently, the Strober lab developed a preclinical model that effectively treated colon cancer in mice by combining immunotherapy and autologous bone marrow transplantation in order to markedly augment the anti-tumor potency of immunotherapy. They used the CT26 colon cancer as the therapeutic target either as a single subcutaneous tumor nodule, as a disseminated tumor in the lungs and peritoneum, or as a metastatic tumor in the liver depending on the route of administration of the tumor cells in BALB/c mice. Mice were vaccinated mice with established primary tumors or disseminated/ metastatic disease with irradiated tumor cells mixed with the adjuvant CpG, and found that vaccination alone had no effect on tumor growth. Similarly radiation conditioning of tumor bearing hosts followed by transplantation of bone marrow and spleen cells or purified T cells and hematopoietic stem cells from unvaccinated donors of the same strain had no effect. In contrast, radiation conditioning of mice followed by transplantation of hematopoietic and immune cells from donors of the same strain vaccinated with tumor cells and CpG cured almost all subcutaneous primary as well as disseminated and metastatic tumors in the hosts. A similar result was obtained after autologous transplantation of hematopoietic and immune cells from tumor bearing mice that had been vaccinated after tumor establishment. Investigation of tumor infiltrating cells showed that the injected donor T cells do not accumulate in the tumors unless the host has been irradiated before injection.

Based on this model, we have assembled a team of Stanford University faculty members with expertise in gastrointestinal cancers, immunotherapy, radiation oncology, and bone marrow transplantation in the Departments of Medicine and Pathology to translate the preclinical findings into a Phase I safety and feasibility clinical study for the treatment of 10 patients with metastatic colorectal cancer. Resected tumor cells will be irradiated and mixed with CpG to create a vaccine. Patients will receive subcutaneous vaccination at weeks 1 and 2 after resection. Six weeks later, immune T cells and then G-CSF "mobilized" purified blood progenitor cells will be harvested from the blood and cryopreserved. If needed patients will receive chemotherapy for tumor reduction. When disease is controlled off chemotherapy, patients will receive a conditioning regimen of fludarabine (30mg/m2 daily x 3 days) followed by intensive fractionated total body irradiation. The dose of fTBI will be escalated using a 3+3 design to ensure safety and will range from 400 to 800 gray. The patient will then undergo hematopoietic and immune cell rescue. They will undergo a third vaccination within 7-14 days after transplant. Thereafter, serial monitoring of tumor burden will continue.

Immune monitoring will occur before and after vaccination as well as after transplantation. Tests will include in vitro anti-tumor immune responses of T cells (proliferation, cytotoxicity, cytokine secretion etc.) to stimulation with whole tumor cells and tumor cell lysates pulsed on to antigen presenting cells, anti-tumor

antibody responses, and immune reconstitution after transplantation.

Primary Outcome: •To assess the feasibility of using an autologous tumor cell vaccine in combination with standard chemotherapy followed by investigational autologous hematopoietic and immune cell rescue in terms of acceptable clinical toxicity.

Secondary Outcome: •Preliminary efficacy in terms of response and time to progression. /•Ex vivo analysis of immune response

Completed Human Papillomavirus (HPV) Vaccine Consistency and Non-inferiority Trial in Young Adult Women.					
Intervention: Biological: Cervarix™ Completed Immunogenicity and Safety of GlaxoSmithKline Biologicals' Huma Papillomavirus (HPV) Vaccine 580299 in Healthy Females 15 − Has Results Condition: HPV-16/18 Infections and Cervical Neoplasia Intervention: Biological: Cervarix TM Completed A Study to Evaluate the Immunogenicity and Safety of GSK Biologicals' HPV Vaccine in Healthy Women Aged 18-35 Years. Conditions: Human Papillomavirus (HPV) Infection; Associated Cervical Neoplasia Intervention: Biological: HPV-16/18 L1 VLP AS04	Completed	Human Papillomavirus (HPV) Vaccine Consistency and Non-inferiority Trial in Young Adult Women.			
Completed Immunogenicity and Safety of GlaxoSmithKline Biologicals' Huma Papillomavirus (HPV) Vaccine 580299 in Healthy Females 15 – Has Results Condition: HPV-16/18 Infections and Cervical Neoplasia Intervention: Biological: Cervarix TM Completed A Study to Evaluate the Immunogenicity and Safety of GSK Biologicals' HPV Vaccine in Healthy Women Aged 18-35 Years. Conditions: Human Papillomavirus (HPV) Infection; Associated Cervical Neoplasia Intervention: Biological: HPV-16/18 L1 VLP AS04	Has Results	Conditions: Papillomavirus Type 16/18 Infection; Cervical Intraepithelial Neoplasia			
Has Results Condition: HPV-16/18 Infections and Cervical Neoplasia Intervention: Biological: Cervarix TM Completed A Study to Evaluate the Immunogenicity and Safety of GSK Biologicals' HPV Vaccine in Healthy Women Aged 18-35 Years. Conditions: Human Papillomavirus (HPV) Infection; Associated Cervical Neoplasia Intervention: Biological: HPV-16/18 L1 VLP AS04		Intervention: Biological: Cervarix™			
Has Results Condition: HPV-16/18 Infections and Cervical Neoplasia Intervention: Biological: Cervarix TM Completed A Study to Evaluate the Immunogenicity and Safety of GSK Biologicals' HPV Vaccine in Healthy Women Aged 18-35 Years. Conditions: Human Papillomavirus (HPV) Infection; Associated Cervical Neoplasia Intervention: Biological: HPV-16/18 L1 VLP AS04					
Intervention: Biological: Cervarix TM Completed A Study to Evaluate the Immunogenicity and Safety of GSK Biologicals' HPV Vaccine in Healthy Women Aged 18-35 Years. Conditions: Human Papillomavirus (HPV) Infection; Associated Cervical Neoplasia Intervention: Biological: HPV-16/18 L1 VLP AS04	Completed	Immunogenicity and Safety of GlaxoSmithKline Biologicals' Huma Papillomavirus (HPV) Vaccine 580299 in Healthy Females 15 -			
Completed A Study to Evaluate the Immunogenicity and Safety of GSK Biologicals' HPV Vaccine in Healthy Women Aged 18-35 Years. Conditions: Human Papillomavirus (HPV) Infection; Associated Cervical Neoplasia Intervention: Biological: HPV-16/18 L1 VLP AS04	Has Results	Condition: HPV-16/18 Infections and Cervical Neoplasia			
Conditions: Human Papillomavirus (HPV) Infection; Associated Cervical Neoplasia Intervention: Biological: HPV-16/18 L1 VLP AS04		Intervention: Biological: Cervarix TM			
Conditions: Human Papillomavirus (HPV) Infection; Associated Cervical Neoplasia Intervention: Biological: HPV-16/18 L1 VLP AS04					
Intervention: Biological: HPV-16/18 L1 VLP AS04	Completed	A Study to Evaluate the Immunogenicity and Safety of GSK Biologicals' HPV Vaccine in Healthy Women Aged 18-35 Years.			
		Conditions: Human Papillomavirus (HPV) Infection; Associated Cervical Neoplasia			
		Intervention: Biological: HPV-16/18 L1 VLP AS04			
Recruiting RNActive®-Derived Therapeutic Vaccine RNActive®-Derived Therapeutic Vaccine	Recruiting	RNActive®-Derived Therapeutic Vaccine	RNActive®-Derived Therapeutic Vaccine in		

	Condition: Hormone Refractory Prostate Cancer Intervention: Biological: CV9103 2006	Advanced or Metastatic Hormone Refractory Prostate Cancer
	The Phase I part of the study consists of a staggered inclusion of subjects in two cohorts of 3, to confirm the safety of the with a lower dose to be considered in case of dose-limiting toxicity (DLT) being reported in greater than or equal to 2 out recommended dose (RD) for the Phase IIa part of the study will be established. In the Phase IIa part of the study, additio confirm the safety and explore the activity of that dose. Primary Outcome: •Phase I: Assessment of safety and tolerability of the trial regimen [9 Weeks]. /•Phase I: Evaluation of CV9103 is an mRNA-based vaccine for the treatment of human prostate cancer that is based on CureVac's RNActive® to specific antigens. Because these antigens are present in prostate cancer cells, they are appropriate targets for intervention correlate frequently with the progression of prostate cancer, and are known to be immunogenic in humans, where they in expansion. As an RNA-based vaccine, CV9103 features several advantages over other approaches: it is highly specific, genotype, and it does not need to cross the nuclear membrane to be active. Finally, in the absence of reverse transcripting genome.	of 3-6 subjects; in this way, the nal subjects will be included at the RD, to finduction of immune response echnology. CV9103 encodes for 4 prostate on. These antigens have been shown to duce antigen specific T-cell or B cell there is no restriction to the patient's MHC
Recruiting	Trial for Vaccine Therapy With Dendritic Cells - Transfected With hTERT-, Survivin- and Tumor Cell Derived mRNA + ex Vivo Condition: Metastatic Malignant Melanoma Interventions: Biological: Dendritic cells - transfected with hTERT-, survivin- and tumor cell derived mRNA + ex vivo T cell expansion and reinfusion; Drug: Temozolomide 2010 Primary Outcome: *Safety and toxicity of vaccination with DC transfected h-TERT mRNA, survivin mRNA and tumor cell expansion and reinfusion in patients with metastatic malignant melanoma. [followed up for two years after start of values and toxicity of vaccination with metastatic malignant melanoma.	
Secondary Outcome: •Evaluation of immunological responses, time to disease progression and survival time. [5 years of follow-up.] The investigators have also included hTERT and survivin mRNA in the vaccine. Finally, the investigators want to introduce ex vivo T cell expansion lymphodepletion for the patients who show an immune response.		
Completed	A Safety and Immunology Study of a Modified Vaccinia Vaccine for HER-2(+) Metastatic Breast Cancer Condition: Breast Cancer Intervention: Biological: MVA-BN-HER2 2010	MVA-BN-HER2 Following 1st- or 2nd-Line Chemotherapy for HER-2-Positive Metastatic Breast Cancer

MVA-BN®-HER2 is a candidate breast cancer immunotherapy product comprised of a highly attenuated non-replicating vaccinia virus, MVA-BN®, engineered to encode a modified form of the Her-2 protein.

MVA-BN® is a well-characterized, clonal strain of modified vaccinia virus Ankara (MVA) being developed as a smallpox vaccine, suitable for use in high-risk (e.g., immunocompromised) individuals. MVA-BN®-derived vectors encoding heterologous antigens are being developed for use as vaccines for infectious diseases such as HIV, and for the treatment of cancer. A large database exists from safety evaluations in animals and in humans for MVA-BN®, and MVA-BN®-derived vectors. Her-2 is overexpressed in 20-30% of human breast cancers. It is an oncogene/growth factor receptor critical for malignant phenotype of Her-2 expressing tumors. It is an immunogenic target, and immune responses to this protein have been shown to mediate potent anti-tumor activity in multiple animal models. Means to stimulate anti-Her-2 reactivity are now being studied clinically. Sponsor, collaborators, and others have used both Protein and DNA vaccine forms of Her-2, and a safety database is developed and no significant adverse events have resulted from Her-2 directed vaccination.

MVA-BN®-HER2 encodes a modified form of the Her-2 protein, hereinafter referred to as HER2. HER2 contains the extracellular domain of Her-2 but lacks the intracellular, cell signaling domain. In addition, HER2 includes two universal T-cell epitopes from tetanus toxin to facilitate the stimulation of an immune response to Her-2, a self-protein.

The current trial, BNIT-BR-002, will evaluate the safety and biological activity of a fixed dose of MVA-BN®-HER2, with and without Herceptin, following 1st- or 2nd-line chemotherapy in patients with metastatic breast cancers which overexpress Her-2.

Patients will receive 3 subcutaneous vaccinations at 3 week intervals and have tumor followed by CT/MRI imaging and blood drawn for immune function analysis

Completed

ed	Safety and Imm	unogenicity Study of the New dHER2 Vaccine to Treat HER2-positive Metastatic Breast Cancer	the dHER2 Recombinant Protein Combined With
	Condition:	Metastatic Dieast Calicei	Immunological Adjuvant AS15 in Patients With
		Biological: UNK Biologicals / 19125 ZUU5	Metastatic Breast Cancer Overexpressing HER2/Neu

Primary Outcome; •Vaccine-related Grade 3 or 4 toxicity (other than skin toxicity and influenza-like symptoms) according to the Common Terminology Criteria for Adverse Events version 3.0. /•Objective clinical response (CR or PR)

Secondary Outcome: •Stable disease. /•Mixed response /•Time to disease progression. /•Time to onset of response, defined as time from first vaccination to the initial response. /•The duration of overall response is measured from the time that measurement criteria are met for complete response or partial response until the first date that recurrent or progressive disease is objectively documented. /•Anti-dHER2, anti-HER2 ECD and anti-HER2 ICD seropositivity. /•Functional activity in vitro. /•Frequency of cellular immune response in vitro to dHER2, HER2 ECD and HER2 ICD. /•Adverse events of Grades 3 and 4. /•Adverse events related to potential cardiotoxicity. /•Solicited local and general signs and symptoms (recorded by the patients on diary cards) [four days following each administration]. /• Unsolicited adverse events (serious and non-serious). /•Unsolicited serious adverse events. /•Any documented toxicity. /•Left ventricular ejection fraction. /• Laboratory values: hematological and biochemical variables (including coagulation). /•Vital signs. /•Electrocardiographic results [Time Frame: at the end of cycle 1 and cycle 2 and at first follow-up visit]. /•Results of physical examination.

Active, not recruiting

ot	Safety and Immunogenicity of a Melanoma DNA Vaccine Delivered by Electroporation	C-f-+
g	I Conditions: IMelanoma (Skin): Intraocular Melanoma 2007	Safety and Immunogenicity of a Xenogeneic Tyrosinase DNA Vaccine Melanoma
	Interventions: Biological: Xenogeneic Tyrosinase DNA Vaccine; Device: TriGrid Delivery System for Intramuscular	Tyrosinase DNA vaccine melanoma

Primary Outcome: •Evaluate the safety and feasibility of electroporation mediated intramuscular delivery of a mouse tyrosinase plasmid DNA vaccine in patients with stage IIB, IIC, III, or IV melanoma. [one year], /•Assess the magnitude and frequency of tyrosinase specific immunologic responses in the immunized patients Secondary Outcome. /•Assess patients with measurable tumor for evidence of anti-tumor response following immunization. This study is designed to evaluate administration of a xenogeneic DNA vaccine encoding the melanosomal antigen tyrosinase by in vivo electroporation in patients with malignant melanoma. The objectives of the study are to characterize the safety and immunogenicity of a DNA vaccine encoding the murine tyrosinase gene delivered administered intramuscularly using the electroporation based TriGrid Delivery System (Ichor Medical Systems). We will assess the nature, frequency, and severity of any toxicity associated with vaccination at escalating pINGmuTyr doses and then expand enrollment at then expand enrollment at the Maximum Tolerated Dose to assess immunologic responses to the tyrosinase antigen. Completed Lymphocyte-Depleting Nonmyeloablative Preparative Chemotherapy Followed By Autologous Lymphocyte Infusion, Peptide Metastatic Melanoma Using Condition: Melanoma (Skin) Nonmyeloablative But Lymphocyte Depleting Biological: NY-ESO-1 peptide vaccine; Biological: aldesleukin; Biological: filgrastim; Biological: incomplete Regimen Followed By The Administration Of In Vitro Sensitized Lymphocytes Reactive Interventions: Freund's adjuvant; Biological: therapeutic autologous lymphocytes; Drug: cyclophosphamide; Drug: With ESO-1 Antigen fludarabine phosphate 2004 Primary: •Determine the clinical tumor regression in patients with metastatic melanoma treated with a lymphocyte-depleting nonmyeloablative preparative chemotherapy regimen followed by autologous lymphocyte infusion, ESO-1 peptide vaccination comprising ESO-1:157-165 (165V) and Montanide ISA-51, and L-2. Secondary: •Determine the survival of the infused lymphocytes in patients treated with this regimen./•Determine the long-term immune status of patients treated with this regimen, /OUTLINE: Patients are stratified according to type of lymphocyte infusion (ESO-1-reactive tumor-infiltrating lymphocytes [TIL] vs ESO-1 reactive peripheral blood lymphocytes [PBL]). /•Autologous lymphocyte collection and expansion: Autologous PBL or TIL are collected from patients during leukapheresis or biopsy. The cells are sensitized in vitro with ESO-1:157-165 (165V) melanoma antigen and expanded. /•Lymphocyte-depleting nonmyeloablative preparative chemotherapy: Patients receive lymphocyte-depleting nonmyeloablative preparative chemotherapy comprising cyclophosphamide IV over 1 hour on days -7 and -6 and fludarabine IV over 15-30 minutes on days -5 to -1. /•Autologous lymphocyte infusion: Autologous PBL or TIL are reinfused on day 0*. Patients also receive filgrastim (G-CSF) subcutaneously (SC) once daily beginning on day 1 and continuing until blood counts recover /•ESO-1 peptide vaccination: Patients receive ESO-1 peptide vaccination comprising ESO-1:157-165 (165V) peptide emulsified in Montanide ISA-51 SC on days 0*-4, 11, 18, and 25. /•Interleukin therapy: Patients receive interleukin-2 IV over 15 minutes 3 times daily on days 0*-4. /NOTE: *Day 0 is 1-4 days after the last dose of fludarabine. Patients are followed at 4-5 weeks, every 3-4 months for 2 years, and then annually thereafter. Active, not Vaccination With Autologous Breast Cancer Cells Engineered to Secrete Granulocyte-Macrophage Colony-Stimulating Factor Autologous, Lethally Irradiated Breast Cancer recruiting Condition: Breast Cancer Cells Engineered by Adenoviral Mediated Gene

Intervention: Biological: Autologous, Lethally Irradiated Breast Cancer Cells 2004

Transfer to Secrete GM-CSF

	the cancer. The Primary Outco CSF that can b Secondary Ou These cells will and #5, the pat study treatmen	this trial is to test the safety of a vaccine made from a patient's own breast cancer cells, and determine if a vaccine is made by genetically modifying a patient's own tumor cells to secrete GM-CSF to activate the same: •To determine the doses of lethally irradiated, autologous breast cancer cells engineered by adenoving the manufactured for metastatic breast cancer [3 years]. /•to determine the safety and biologic activity of the same in the time to progression and overall survival of metastatic breast cancer patients treatly to measure how the patient's immune system is reacting to the tumor cells. This is called Delayed-Trient will also receive a DTH injection. Two to three days after the vaccine and DTH injection, skin biopsies to the time to progression and chest, abdomen, and pelvic CT scan to determine if the vaccine model.	immune response. ral mediated gene transfer to secrete GM- nis vaccination in metastatic breast cancer eated with this vaccine type Hypersensitivity (DTH). With vaccine #1 s will be taken of both sites. At week 10 in the accine therapy has had an effect on their	
Completed		, Montanide ISA 51 and ISA 51 VG, and CpG 7909 in Treating Patients With Resected Stage IIC, Stage III, or	Vaccine Combining Multiple Class I Peptides	
		Intraocular Melanoma; Melanoma (Skin) 2004 Biological: gp100 antigen; Biological: incomplete Freund's adjuvant; Biological: recombinant MAGE-3.1	With Montanide ISA 51 and ISA 51 VG and CpG Adjuvant 7909 for Resected Stages	
	interventions.	antigen; Biological: tyrosinase peptide; Drug: agatolimod sodium; Procedure: adjuvant therapy	IIC/III and IV Melanoma	
	1	me: •Immunological response as measured by ELISPOT assy, tetramer assay, and chromium release as	say	
	, -	tcome: •Toxicity [Designated as safety issue: Yes] •Time to relapse	oine Mentenide ISA 51 and ISA 51 VC and	
•Determine the safety and tolerability of a multipeptide (gp100 antigen, MAGE-3, and tyrosinase peptide) melanoma vaccine, Montanide CpG 7909 in patients with resected stage IIC, III, or IV melanoma. /•Determine the immune reactivity of this regimen in these patients.				
		is a pilot study. Patients are stratified according to class I haplotype (HLA-A1 vs HLA-A3/A11).	panerne.	
	Patients receive gp100 antigen, MAGE-3, tyrosinase peptide, Montanide ISA 51 and ISA 51 VG, and CpG 7909 subcutaneously on weeks 0, 2, 4, 6, 8, 10, 14, 18,			
	22, 26, 38, 50,	and then every 6 months for 2 years (for a total of 16 vaccinations) in the absence of unacceptable toxicit	у.	
		afety and Immunogenicity of GSK Biologicals' HPV Vaccine GSK580299 (Cervarix TM) Administered in Healthy		
recruiting	Conditions:	Human Papillomavirus (HPV) Infection; Papillomavirus Vaccines		
	Interventions:	Biological: GSK Biologicals' HPV Vaccine GSK580299; Biological: Engerix−B™		
Completed	01	Description of Fludenships and Oracles bearing Fallowed By White Blood Call Inferior Vession Theorems and		
Completed		Consisting of Fludarabine and Cyclophosphamide Followed By White Blood Cell Infusion, Vaccine Therapy, and Melanoma (Skin)	Metastatic Melanoma Using Lymphocytes Reactive With the GP100 Antigen With	
	L		Immunization Using a Recombinant RF-	
	Interventions:	Biological: aldesleukin; Biological: filgrastim; Biological: fowlpox virus vaccine vector; Biological: gp100 antigen; Biological: therapeutic autologous lymphocytes; Biological: therapeutic tumor infiltrating lymphocytes; Drug: cyclophosphamide; Drug: fludarabine phosphate	GP100P209 Virus Encoding a GP100 Peptide Following a Nonmyeloblative Lymphocyte	
L	J	tymphocytes, Drug, cyclophosphaniac, Drug, hadarabine phosphate		

recruiting

Active, not Vaccine Therapy and Sargramostim in Treating Adults With Metastatic Cancer

Primary: •Determine complete clinical tumor regression in patients with recurrent or refractory metastatic melanoma treated with lymphocyte-depleting nonmyeloablative preparative chemotherapy comprising fludarabine and cyclophosphamide followed by autologous lymphocyte infusion, recombinant fowlpox virus encoding gp100 peptide, and aldesleukin. **Secondary:** •Determine the survival of patients treated with this regimen. •Determine the safety of this regimen in these patients. OUTLINE: Patients are stratified according to the availability of suitable reactive cells (peripheral blood lymphocytes [PBL] vs tumor-infiltrating lymphocytes [TIL]). •Autologous lymphocyte activation and expansion: Autologous PBL or TIL are activated in vitro with qp100:209-217 (210M) antigen (qp100) and expanded. •Lymphocyte-depleting nonmyeloablative preparative regimen: Patients receive cyclophosphamide IV on days -7 and -6 and fludarabine IV over 15-30 minutes on davs -5 to -1. •Autologous lymphocyte infusion: Autologous PBL or TIL are reinfused over 20-30 minutes on day 0*. Patients also receive filgrastim (G-CSF) subcutaneously (SC) once daily beginning on day 1 or 2 and continuing until blood counts recover. •Fowlpox vaccine administration: Patients receive recombinant fowlpox virus encoding gp100 peptide IV over 1-2 minutes on days 2 and 28 (if treated with high-dose aldesleukin [IL-2], as below) OR days 2 and 43 (if treated with low-dose IL-2, as below). •IL-2 therapy: Patients receive high-dose IL-2 IV over 15 minutes every 8 hours on days 0*-4 (beginning within 24 hours after lymphocyte infusion) and 28-32 OR low-dose IL-2 SC on days 0*-4 (beginning within 24 hours after lymphocyte infusion), 7-11, 14-18, 21-25, 28-32, 35-39, 50-54, 57-61, 64-68, 71-75, 78-82, and 85-NOTE: *Day 0 is 1-4 days after the last dose of fludarabine. Patients are evaluated between days 72-86 (if treated with high-dose IL-2) OR days 98-123 (if treated with low-dose IL-2). Patients with stable disease or a minor, mixed, or partial response may receive up to 2 retreatment courses as above. Patients with progressive disease after IV lymphocyte infusion may be retreated with intra-arterial lymphocytes along with all other agents outlined above. After completion of study treatment, patients are followed at 2-4 weeks (if treated with high-dose IL-2) OR at 3 weeks (if treated with low-dose IL-2) and then annually thereafter. PROJECTED ACCRUAL: A total of 68 will be accrued for this study Completed Study of Combination Immunotherapy for the Generation of HER-2/Neu Specific Cytotoxic T Cells Condition: Breast Cancer Intervention: Biological: HER2 CTL vaccine (plus trastuzumab) This will be a single arm phase I-II single institution clinical trial in patients with HER2 overexpressing Stage IV breast and ovarian cancer who are on maintenance trastuzumab alone after being treated with chemotherapy and trastuzumab or trastuzumab alone to NED or stable disease. Patients will receive a monthly vaccination for 6 months with a HER2 CTL peptide-based vaccine. Phase I-II Study of Combination Immunotherapy for the Generation of HER-2/Neu (HER2) Specific Cytotoxic T Cells (CTL) in Vivo Primary Outcome Measures: •Safety [Time Frame: 5 years] •Immune response [Time Frame: 1.5 years] Secondary Outcome Measures: •Overall survival [Time Frame: At least 5 years] [Designated as safety issue: No] Biological: HER2 CTL vaccine (plus trastuzumab) HER2 CTL peptide-based vaccine; administered intradermally every month for 6 total doses Active, not GARDASIL™ Vaccine Impact in Population Study Condition: Human Papillomavirus Infections Intervention:

recruiting	Conditions: Breast Cancer; Colorectal Cancer; Ovarian Cancer; Unspecified Adult Solid Tumor, Protocol Specific	!	
	Interventions: Biological: falimarev; Biological: inalimarev; Biological: sargramostim		
	Background: •Many cancers produce two proteins, carcinoembryonic antigen (CEA) and mucin-1 (MUC-1).		
	•The PANVAC-V priming vaccine and PANVAC-F boosting vaccine contain human genes that cause production of CEA an	nd MUC-1, which can be used as a target	
	for the immune system to attack the cancer. The vaccines also contain genes that cause production of other proteins that e		
	•Sargramostim is a protein that boosts the immune system.	ř	
	Objectives: •To evaluate the safety and effectiveness of PANVAC-V and PANVAC-F in patients with advanced cancer.		
	•To document the immune response to the vaccines and any anti-tumor responses that may occur.		
	Eligibility: Patients 18 years of age and older with advanced cancer whose tumors produce CEA or MUC-1 protein		
	Design: •This trial has four arms: the first arm includes 10 patients with advanced colorectal cancer; the second arm includes 10 to 15 patients with any ad		
	non-colorectal cancer that produces either EA or MCU-1; the third arm includes about 12 patients with advanced breast cal	ncer; the fourth arm includes about 12	
	patients with advanced ovarian cancer.		
	•All patients receive PANVAC-V on study day 1, followed by PANVAC-F on days 15, 29 and 43. The vaccines are given by	injection under the skin. Sargramostim is	
	injected at the vaccination site on the day of each vaccination and for the next 3 days following vaccination.		
	•Patients whose disease has not worsened after the last boosting vaccination may receive up to 12 additional monthly boost	•	
	vaccinations, patients may receive vaccine every 3 months. Patients whose scans show that their disease has progressed,	, but who are otherwise clinically stable	
	may revert back to monthly injections.		
	•Patients undergo apheresis to collect white blood cells (lymphocytes) on day 1 and day 71 of the study to measure the imr		
	collected through a needle placed in one arm and directed through a cell separator machine where the lymphocytes are ex	tracted. The rest of the blood components	
	are returned to the patient through the same needle.	to traction and	
	Patients are monitored with frequent blood tests and periodic imaging tests (scans) to monitor for safety and the response	to treatment.	
	Drug: PANVAC-V [Recombinant-Vaccinia-CEA (D609)/MUC-1(L93)/TRICOM] Drug: PANVAC-F [Recombinant-Fowlpox-CEA (D609)/MUC-1(L93)/TRICOM]		
	Drug: Leukine (Sargramostim)		
Active not			
recruiting	Vaccine Therapy in Treating Patients With Myelodysplastic Syndromes Condition IM relative Syndromes		
	Condition: Myelodysplastic Syndromes Interventions: Biological: GM-K562 cell vaccine; Genetic: cytogenetic analysis; Genetic: fluorescence in situ hybridization;		
	Interventions: Other: flow cytometry; Other: immunoenzyme technique; Other: laboratory biomarker analysis		
L	1 Totalet, now cytometry, Other, iniminioenzyme teorinique, Other, laboratory biolitainer analysis		

RATIONALE: Vaccines made from cancer cells may help the body build an effective immune response to kill abnormal cells. PURPOSE: This clinical trial is studying how well vaccine therapy works in treating patients with myelodysplastic syndromes (MDS). Primary Outcome Measures: •Safety, •Hematologic response, defined as achieving a major response in ≥ 1 lineage as described by an erythroid increase > 2 a/dL, platelet increase of 30,000/mm³, or neutrophil increase by 100%, •Cytogenetic response, defined as normalization of pretreatment cytogenetic abnormalities [Designated as safety issue: No 1 Secondary Outcome Measures: •Immune response to common myeloid antigens (e.g., Wilms' tumor-1 [WT-1], survivin, or proteinase-3) as measured by Elispot assay, •Correlation of immune response with clinical response (hematologic response, resolution of cytogenetic abnormalities, or decrease in other parameters, such as WT-1 mRNA levels) **OUTLINE: This is an open-label study.** Patients receive GM-K562 cell vaccine subcutaneously once in weeks 0, 3, 6, 9, and 17 in the absence of disease progression or unacceptable toxicity. Blood and tissue samples are collected periodically for correlative and biomarker studies. Samples are analyzed by cytogenetic studies, fluorescent in situ hybridization (FISH), and flow cytometry. Elispot is used to quantify cellular cytotoxic T-cell response to Wilms' tumor-1 (WT-1), survivin, and proteinase 3. After completion of study treatment, patients are followed every 3 months for 1 year. PROJECTED ACCRUAL: A total of 15 patients will be accrued for this study. Completed Vaccine Therapy in Treating Patients With Stage I or Stage II Pancreatic Cancer Condition: Pancreatic Cancer Intervention: Biological: vitespen RATIONALE: Vaccines made from a person's cancer cells may make the body build an immune response to and kill tumor cells. Combining vaccine therapy with surgery may be an effective treatment for pancreatic cancer. PURPOSE: Phase I trial to study the effectiveness of vaccine therapy in treating patients with stage I or stage II pancreatic cancer that has been surgically removed. A Phase I Pilot Trial of Immunotherapy With Autologous Tumor-Derived gp96 Heat Shock Protein - Peptide Complex (HSPPC-96) in Patients With Resected Pancreatic Adenocarcinoma OBJECTIVES: I. Study the safety of autologous tumor derived gp96 heat shock protein peptide complex (HSPPC-96) in patients with resected pancreatic adenocarcinoma. II. Examine the immune response to HSPPC-96 in this group of patients. OUTLINE: This is a dose escalation study. Six weeks after surgery patients are given autologous tumor derived gp96 heat shock protein peptide complex (HSPPC-96) subcutaneously once a week for 4 weeks. Five patients are initially enrolled at each of two dose levels. An additional three patients may be enrolled at each dose level to determine the optimal dose of HSPPC-96. Patients are followed at weeks 1, 4, and 12 after treatment. PROJECTED ACCRUAL: A maximum of 16 patients will be accrued for this study. Study of EMD531444 in Subjects With Stage III Unresectable Non-small Cell Lung Cancer (NSCLC) Following Primary Recruiting EMD531444(L-BLP25 or BLP25 Liposome Condition: Non-small Cell Lung Cancer メルクセローノが日本でも治験を実施 Vaccine) in Subjects With Stage III Interventions: Biological: cyclophosphamide + EMD531444(LBLP25 or BLP25リポゾームワクチン) + BSC; Biological: Saline + Unresectable Non-small Cell Lung Cancer Placebo + BSC 糖蛋白ワクチンの一つであるL-BLP25は、リポペプチドを生成する25のアミノ酸から構成される Following Primary Chemoradiotherapy Primary Outcome Measures: •Overall survival time [Time Frame: The date of randomization; week 1, 2, 3, 4, 5, 6, 7, 8, 9, 14; every 6 weeks after week 14; 6 weeks after last vaccination; 12 weeks after last vaccination. Additional inquires on survival until death.] [Designated as safety issue: No] Detailed Description: Phase I part is designed to evaluate the safety of EMD531444 1000mcg dose to be used in phase II. Phase II part is designed to be conducted as randomized, double blind, placebo controlled study to compare overall survival time in all randomized subjects.

Completed Vaccine Therapy Plus Chemotherapy in Treating Patients With Metastatic or Locally Recurrent Stomach Cancer or Esophageal Conditions: Esophageal Cancer; Gastric Cancer Interventions: Biological: G17DT Immunogen; Drug: cisplatin; Drug: fluorouracil RATIONALE: Vaccines may make the body build an immune response to kill tumor cells. Drugs used in chemotherapy use different ways to stop tumor cells.			
Interventions: Biological: G17DT Immunogen; Drug: cisplatin; Drug: fluorouracil			
	ells from		
dividing so they stop growing or die. Combining vaccine therapy with chemotherapy may kill more tumor cells.			
PURPOSE: Phase II trial to study the effectiveness of combining vaccine therapy and chemotherapy in treating patients who have metastatic or locally re	current		
stomach cancer or esophageal cancer.			
Primary Outcome Measures: •To determine whether a concomitant G17DT-chemotherapy regimen affects tumor response in subjects with gastric or			
gastroesophageal cancer. [Time Frame: 6 months to 1 year] [Designated as safety issue: No]			
Secondary Outcome Measures: •Time to disease progression, best overall response, and survival will be evaluated in the intent-to-treat population and the			
evaluable population. [Time Frame: 6 months to 1 year]			
BJECTIVES: I. Determine a safe and immunogenic combination of G17DT with cisplatin and fluorouracil in patients with chemotherapy-naive metastatic or locally recurrent gastric or gastroesophageal cancer. II. Determine the safety profile and tolerability of this regimen in these patients. III. Determine the tumor response rate,			
disease stabilization, best overall response, time to progression, time to treatment failure, and overall survival in patients treated with this regimen. IV. Determine the			
correlation of immunological response with clinical efficacy and benefit in patients treated with this regimen. V. Determine the pharmacokinetics and			
pharmacodynamics of this regimen in these patients.			
OUTLINE: This is a multicenter study. Patients are assigned to one of four treatment regimens. Regimen A: Patients receive high-dose G17DT intramusc	ılarly (IM)		
on days 7, 35, and 63. Patients also receive cisplatin IV over 1-3 hours on day 1 followed by fluorouracil IV continuously over days 1-5 every 4 weeks in the			
of disease progression or unacceptable toxicity. If inadequate immune response is seen on Regimen A, subsequent patients are treated on Regimen B. It			
unacceptable toxicity is seen on Regimen A, subsequent patients are treated on Regimen C. If inadequate immune response and unacceptable toxicity a			
Regimen A, or if unacceptable toxicity is seen on Regimen B or inadequate immune response is seen on Regimen C, then subsequent patients are treated on			
Regimen D. Regimen B: Patients receive high-dose G17D1 lift on days 1, 28, and 56. Patients also receive displatin IV over 1-3 hours on day 35 followed fluorouracil IV continuously over days 35-39 every four weeks in the absence of disease progression or unacceptable toxicity. Regimen C: Patients received.	Regimen D. Regimen B: Patients receive high-dose G17DT IM on days 1, 28, and 56. Patients also receive cisplatin IV over 1-3 hours on day 35 followed by		
G17DT IM on days 7, 35, and 63 with chemotherapy as in regimen A. Regimen D: Patients receive low-dose G17DT IM on days 1, 28, and 56 with chemotherapy			
in regimen B. Quality of life is assessed at baseline, on day 7, every 2 weeks for 10 weeks, and then every 4 weeks thereafter.	thorapy as		
PROJECTED ACCRUAL: A total of 15-75 patients will be accrued for this study within 5-30 months.			
Recruiting Melanoma Vaccine in Treating Patients With Stage III Melanoma After Surgery to Remove Lymph Nodes			
Condition: Melanoma (Skin)			
Biological: HLA-A1-binding MAGE-1/MAGE-3 multipeptide-pulsed autologous dendritic cell vaccine;			
Biological: HLA-A2-binding TYR/MART-1/gp100 multipeptide-pulsed autologous dendritic cell vaccine;			
Interventions: Biological: autologous melanoma lysate-pulsed autologous dendritic cell vaccine; Biological: autologous melanoma lysate/KLH-pulsed autologous dendritic cell vaccine; Biological: dendritic cell-idiotype-keyhole			
limpet hemocyanin vaccine: Other: flow cytometry: Procedure: adiuvant therapy			

Primary Outcome Measures: •Immune response, •Disease-free survival, •Overall survival, •Adverse events **OBJECTIVES:** •Determine the feasibility of adjuvant melanoma vaccine comprising autologous dendritic cells pulsed with tumor antigen peptides in patients with stage III melanoma following lymphadenectomy. •Determine the immune response (skin test of delayed-type hypersensitivity and flow cytometric enumeration of peripheral blood CD8+ lymphocytes producing IFNy) to this regimen in these patients. •Determine clinical outcome (disease-free survival, overall survival, and adverse events) in patients treated with this regimen. OUTLINE: Patients undergo leukapheresis for collection of peripheral blood mononuclear cells (PBMCs) and bone marrow mononuclear cells. Autologous dendritic cells (DCs) prepared from PBMCs and bone marrow mononuclear cells are exposed to various antigens and peptides, and autologous tumor cell lysate, if available. Patients receive autologous DCs pulsed with melanoma-associated antigen peptides, and autologous DCs pulsed with tumor lysates (if available), subcutaneously in weeks 0, 2, 5, 8, 12, 16, 20, 26, 31, 50, and 102. Patients with no evidence of disease may receive another booster injection 5 years after the start of vaccination. Blood samples are examined via flow cytometry and skin testing is performed to evaluate immune response. Recruiting Trial of Activated Marrow Infiltrating Lymphocytes Alone or in Conjunction With an Allogeneic Granulocyte Macrophage Condition: Multiple Myeloma Biological: aMILs; Biological: Allogeneic Myeloma Vaccine Interventions: Primary Outcome Measures: •Response rate utilizing Blade' criteria [Designated as safety issue: No] Secondary Outcome Measures: •Progression-free and overall survival [Designated as safety issue: Yes] •Anti-tumor immune response [Designated as safety issue: No] •The effect of aMILs on osteoclastogenesis [Designated as safety issue: No] •Effect of Marrow Infiltrating Lymphocytes on clonogenic myeloma precursors [Designated as safety issue: No] Estimated Enrollment: 32 Study Start Date: December 2009 Estimated Primary Completion Date: December 2012 (Final data collection date for primary outcome measure) Exp.1 Biological: aMILs: Activated marrow infiltrating lymphocytes Exp.2 Biological: aMILs Activated marrow infiltrating lymphocytes VER. Biological: Allogeneic Myeloma Vaccine Allogeneic granulocyte macrophage colony-stimulating factor (GM-CSF)-based myeloma cellular vaccine Completed Trial of Autologous, Hapten-Modified Vaccine in Patients With Stage III or IV Melanoma Condition: Melanoma Interventions: Biological: Autologous, DNP-modified vaccine (M-Vax); Biological: Autologous, DNP-Modified Melanoma Vaccine: Biological: Autologous, DNP-Modified Vaccine The purpose of this study is to determine whether a vaccine composed of patients' own melanoma cells treated with the chemical, dinitrophenyl (DNP)(called a hapten), is safe and stimulates an immune response to patients' own cancer cells. Primary Outcome Measures: •Immune response to patients' own melanoma cells [Time Frame: 2 months] [Designated as safety issue: No] Secondary Outcome Measures: •Safety [Time Frame: 9 months] [Designated as safety issue: Yes] M-Vax: A Feasibility and Bio-Equivalence Study Using a DNP-Modified Autologous Melanoma Tumor Cell Vaccine as Therapy in Patients With Stage III or IV Vaccine Therapy in Treating Patients With Metastatic Prostate Cancer That Has Not Responded to Hormone Therapy Completed Has Results Condition: Prostate Cancer Interventions: Biological: sipuleucel-T; Biological: Placebo

Primary Outcome Measures: •Time to Objective Disease Progression [Time Frame: 36 months from randomization] [Designated as safety issue: Yes] The time to objective disease progression in patients with asymptomatic metastatic hormone-refractory prostate cancer treated with APC8015 (sipuleucel-T). Secondary Outcome Measures: •Overall Survival [Time Frame: From randomization to 36 months] [Designated as safety issue: Yes] Overall Survival Biological: sipuleucel-T: Autologous peripheral blood mononuclear cells, including antigen presenting cells, that have been activated in vitro with a recombinant fusion protein, PAP-GM-CSF. Treatment consist of 3 doses administered approximately 2 weeks apart. Other Name: APC8015, Provenge Biological: Placebo: Approximately one-third of the autologous guiescent antigen presenting cells (APCs) prepared from a single leukapheresis procedure. A course of therapy consists of 3 complete doses given at approximately 2-week intervals. Interferon-gamma or Aldesleukin and Vaccine Therapy in Treating Patients With Multiple Myeloma Active, not recruitina Condition: Multiple Myeloma and Plasma Cell Neoplasm Biological: aldesleukin; Biological: idiotype-pulsed autologous dendritic cell vaccine APC8020; Biological: Interventions: recombinant interferon gamma: Genetic: polymerase chain reaction: Genetic: reverse transcriptasepolymerase chain reaction; Other: flow cytometry; Other: laboratory biomarker analysis Primary •To assess the clinical benefit in patients with plateau phase multiple myeloma treated with interferon-gamma vs aldesleukin in combination with idiotypepulsed autologous dendritic cell vaccine APC8020. •To describe response rates in patients who are in plateau phase status post-chemotherapy or status postperipheral blood cell transplantation treated with this regimen. Secondary: •To obtain data regarding the ability of this approach to produce an anti-idiotypic immunologic response. •To obtain information about the effects of interferon-gamma and aldesleukin on the number, function, and activation state of immune effector-cells including T-cells and B-cells. •To perform detailed analyses of lymphocyte phenotypes and T-cell repertoires before and after idiotype-pulsed autologous dendritic cell vaccine APC8020. OUTLINE: Patients are stratified according to gender (male vs female) and prior treatment (post-chemotherapy vs post-peripheral blood stem cell transplantation). Patients are randomized to 1 of 2 arms. In both arms, patients undergo apheresis for collection of peripheral blood mononuclear cells for generation of dendritic cells (DC) on days 0, 14, and 28. APC8020 is generated by loading DC with immunoglobulin idiotype prepared from the patient's serum. •Arm I: Patients receive interferon-gamma subcutaneously (SC) once daily on days 1-5, 15-20, and 29-34 and idiotype-pulsed autologous dendritic cell vaccine APC8020 IV over 30-minutes on days 2, 16, and 30. •Arm II: Patients receive aldesleukin SC once daily days 1-5, 15-20, and 29-34 and idiotype-pulsed autologous dendritic cell vaccine APC8020 as in arm I. In both arms, treatment continues in the absence of disease progression. Peripheral blood samples are collected at baseline and on day 5 of courses 1 and 4 for cytokine immunomodulatory studies, including immunophenotyping for lymphocyte phenotypic markers (CD69, CD40L, CD25, CD30, CD71, CDW137, CD134, and HLADR) by flow cytometry and immunofluorescence; T-cell spectratyping by PCR and RT-PCR; T-cell proliferation to idiotype protein; and CTL and T-helper response by flow cytometry. After completion of study treatment, patients are followed every 3 months for 2 years and then every 6 months thereafter. Completed Vaccine Therapy in Treating Patients With High-Risk Stage III or Completely Resected Metastatic Melanoma Conditions: Stage IV Melanoma; Stage III Melanoma; Recurrent Melanoma

Interventions: Drug: dendritic cell-gp100-MART-1 antigen vaccine; Drug: sargramostim