Harnessing immunosurveillance: current developments and future directions in cancer immunotherapy

Maureen L Drakes
Patrick J Stiff
Department of Medicine, Division of Hematology and Oncology, Stritch School of Medicine, Loyola University Chicago, Maywood, IL, USA

Abstract: Despite improved methods of cancer detection and disease management over the last few decades, cancer remains a major public health problem in many societies. Conventional therapies, such as chemotherapy, radiation, and surgery, are not usually sufficient to prevent disease recurrence. Therefore, efforts have been focused on developing novel therapies to manage metastatic disease and to prolong disease-free and overall survival, by modulating the immune system to alleviate immunosuppression, and to enhance antitumor immunity. This review discusses protumor mechanisms in patients that circumvent host immunosurveillance, and addresses current immunotherapy modalities designed to target these mechanisms. Given the complexity of cancer immunosuppressive mechanisms, we propose that identification of novel disease biomarkers will drive the development of more targeted immunotherapy. Finally, administration of different classes of immunotherapy in combination regimens, will be the ultimate route to impact low survival rates in advanced cancer patients.

Keywords: cancer, immunotherapy, immunosurveillance, immunosuppression, dendritic cells, T-cells

Introduction
Cancer is a global health problem that affects all socioeconomic groups and people of all ages. Each year, there are approximately 7.6 million cancer deaths worldwide.1,2 The precise causes of cancer have not been defined. It is known that gene mutations induced by host internal or by environmental factors can lead to the growth of cancer cells. Additionally, cancer cells have unique biological properties, including replicative immortality, metastatic capability, and escaping immunosurveillance.3–5 Investigators have also proposed the idea of cancer stem cells as one of the possible causes of cancer.6–9 Many early cancers do not exhibit defined symptoms, and are only detected in the later stages of disease. By this time, there is usually significant spread to other organs and to lymph nodes. If detected early, conventional therapies, such as chemotherapy, radiation, and surgery are generally successful, resulting in excellent 5-year survival rates. However, cancers such as digestive, ovarian, and pancreatic cancer, which lack easily recognizable symptoms for early stage disease and have inefficient screening procedures, are usually discovered at the stage of advanced metastatic disease. At this time, conventional therapy is not sufficiently effective for management, and typically there is a poor response to therapy, a high rate of disease recurrence, and a poor 5-year survival rate.10 To advance the care of cancer patients, over the past few decades, efforts have focused on developing successful immunotherapeutic agents.
Immunotherapeutic agents used to treat cancer began as early as 1891, when streptococcal organisms were injected into a patient with inoperable cancer, resulting in reduced size of the tumor.\textsuperscript{11–14} Today, much more is known about immune escape mechanisms in cancer, and immunotherapy is more targeted and increasingly more successful. Strategies for cancer immunotherapy vary with the affected organ, and include the use of antitumor cytokine administration, monoclonal antibodies to inhibit tumor-cell growth by inducing apoptosis or other mechanisms, targeting of tumor-associated antigens, blocking of immune-checkpoint molecules, dendritic cell (DC) vaccines to boost antitumor immunity, and adoptive transfer of genetically engineered T-cells.

Recently, the era of modern immunotherapy has been marked by significant triumphs, including breakthroughs with the antigen-presenting cell vaccine sipuleucel-T for castration-resistant prostate cancer,\textsuperscript{13} antibody-blocking checkpoint inhibitor cytotoxic T lymphocyte-associated antigen (CTLA)-4 (Yervoy® [ipilimumab]) for metastatic melanoma,\textsuperscript{16,17} and genetically engineered T-cells for lymphomas and leukemias. These immune approaches significantly enhance the management of cancer therapy, resulting in improved survival, and pave the way for expanded use of these or similar immunotherapy regimens for other cancers.

This review provides an up-to-date account of immuno-surveillance in cancer, promising immune-based clinical trials against various cancers, US Food and Drug Administration (FDA)-approved immunotherapies, and future directions that may facilitate the development of novel immunotherapeutic agents, which we hope will be sufficiently cost-effective and available to all patients who require these treatments.

Cancer immuno-surveillance
One in four deaths in the US is due to cancer. It is projected that 1,665,540 new cancer cases of cancer and 585,720 cancer deaths will occur in 2014.\textsuperscript{10} Even so, there has been a decline in cancer deaths over the last two decades, and we anticipate that as scientists unravel more clues to the causes of cancer, combined with the development of novel immunotherapy for management of advanced and recurrent disease, there will be further improvement in survival rates.

The precise causes of cancer are unknown. In 1909, Paul Ehrlich predicted that the immune system repressed the growth of carcinomas that would otherwise occur at high frequency.\textsuperscript{4–18} This initiated the debate in the field of cancer immunosurveillance. With further understanding in tumor immunology, transplantation, and immunogenetics, a broader concept of “cancer immunoediting” emerged. Cancer immunoediting encompasses a complex network of immunosuppressive factors within the host, tumor-evasion mechanisms, and host-elicited antitumor mechanisms.

Cancer immunoediting consists of three dynamic phases. Briefly, normal cells are transformed by oncogenic stimuli, express distinct tumor specific molecules, and generate proinflammatory danger signals, which initiate phase I: the elimination or immuno-surveillance phase.\textsuperscript{4,4} According to Burnet, the thymus-dependent cells of the body constantly survey host tissues for nascently transformed cells.\textsuperscript{19} Components of the innate and adaptive immune system detect and eradicate transformed cells before they are clinically apparent. In healthy individuals, the immuno-surveillance network consists of immune parameters, such as CD4\(^{+}\), CD8\(^{+}\), and γ\(\delta\) T-cells, natural killer (NK) cells, macrophages, IFN\(\gamma\), perforin, and TRAIL (TNF-related apoptosis inducing ligand), which function effectively and prevent the growth of tumors, and hence in theory they should never become clinically apparent.\textsuperscript{5,20–23}

However, if these transformed cells are not completely eliminated by a combination of existing immunity, they enter phase 2 – the equilibrium phase – in which the immune system controls net tumor-cell outgrowth, with immune responses primarily mediated by CD8\(^{+}\) T-cells and IL-12p70 produced by DCs. If at this stage there is lack of sufficiently strong host antitumor immunity, or if tumor cells become less immunogenic via tumor-antigen loss, or there is major histocompatibility complex (MHC)-molecule downregulation,\textsuperscript{1,24} these cells can avoid T-cell attack and enter phase 3, the escape phase. In the escape phase, tumors begin to grow progressively in an unrestrained manner, ie, become malignant, establish an immunosuppressive tumor micro-environment, and eventually become clinically apparent tumors.\textsuperscript{21,22} Understanding immunosuppressive networks in the tumor will guide the development of novel and effective cancer immunotherapy.

Immune components of the cancer micro-environment
Immunosuppressive events occurring in the tumor micro-environment are critical to the clinical onset of cancer and to survival. Firstly, as shown by recent investigations, not all cellular events in the tumor are detrimental to tumor outcome. In this respect, the cell type, abundance, and location of lymphocytes in tumor beds have been identified as useful factors in predicting disease outcome.\textsuperscript{25–28} In colorectal cancer, for example, a diagnosis of stage I cancer with patients having
few tumor-infiltrating lymphocytes (TILs) had a similar outcome to patients with stage IV and metastatic disease.26 Therefore, a high frequency of CD3+ or of CD8+ T-cells at the tumor site is beneficial to outcome. Similar findings were reported in ovarian and other cancers, where the T-cell infiltration in tumor beds is favorable to disease outcome.29,30

On the contrary, elevated numbers of CD4+ T-regulatory cells (Tregs) expressing the transcription molecule FoxP3 in cancer immune infiltrates and in peripheral blood generally have detrimental consequences on cancer outcome.31-38 It should be noted, however, that there are exceptions to this protumor effect of Tregs, as in the case of colon cancer, in which they may have a positive correlation with survival.39 Cytokines and chemokines produced by cancer cells and by infiltrating immune cells may confer a protumor cytokine polarization in the tumor beds (Figure 1).

Characteristically, the tumor microenvironment produces high levels of cytokines IL-10 and TGFβ, which are associated with worsening of cancer.38-40 Many cell types also contribute to this immunosuppression. For example Tregs are attracted into the tumor beds by the chemokine CCL22.41-43 Additionally, chemokine receptor CCR4 is highly expressed on some Tregs. Tumors may also promote the release of CCL17 and CCL21, which also causes these Tregs to be recruited to tumor sites. Similarly, Tregs expressing VEGF-A (CXCR4) are recruited into tumors expressing the cognate ligand CXCL12.44,45 In diseases where Tregs are detrimental to tumor outcome, these cells may be enriched by the conversion of immunocompetent T-cells into Tregs or by preferential expansion of resident Tregs over other cell types, as a consequence of the existing protumor cytokine environment. Other immunosuppressive contributing cells in tumors include plasmacytoid DCs, suppressive macrophages, immature myeloid-derived DCs, and myeloid-derived suppressor cells (MDSCs).46-52

The maturation, function, and ability of DCs to migrate may be suppressed by MDSCs, M2 (alternatively activated, protumor) macrophages, Tregs, and other factors in the tumor microenvironment.53,54 Therefore, agents that can reprogram M2 macrophages to M1 (classically activated, antitumor) macrophages may be useful in alleviating immunosuppression in the tumor.55 M1 macrophages produce antitumor cytokines and are associated with prolonged survival, whereas M2 macrophages secrete protumor IL-10 and TGFβ, and are associated with poor outcome in cancer. The chemotherapeutic agent gemcitabine was shown to selectively reduce myeloid-suppressor cells (Gr1+CD11b+) and enhance antitumor activity of CD8+ and NK cells.56

Immature myeloid-derived DCs, which express low levels of costimulatory molecules (CD40, CD80 [B7-1] and CD86 [B7-2]), and high levels of B7-H1 inhibit T-cell proliferation...
and induce FoxP3 Tregs. Also, other investigations showed that patients with a low expression of B7-H4 on tumor-associated macrophages had a higher 5-year survival than those with high expression of B7-H4 on these cells. Furthermore, myeloid DCs and macrophages can induce Th17 cells, which secrete IL-17, a cytokine associated with protumor or antitumor immune responses depending on the nature of the cancer. Plasmacytoid DCs express the enzyme indoleamine 2,3-dioxxygenase, which is associated with poor outcomes in cancer.

In addition to immunosuppression by host immune mechanisms, there are direct immunosuppressive elements imposed by tumor cells in an effort to induce immune evasion. Therefore, tumor cells are capable of downregulating MHC class I, thereby making cross-presentation of tumor-derived antigens (eg, from dead tumor cells) less efficient.

Another immunosuppressive mechanism in the tumor and in the periphery is imposed at immune-checkpoint junctions. Immune checkpoints are necessary to prevent the overstimulation of the immune system; however, coinhibitory molecules that mediate this function can have detrimental consequences in the cancer immunosuppressive environment, inducing protumor immune responses. There are several coinhibitory molecules at immune checkpoints in humans, but notable examples that have recently come full circle as cancer immunotherapy targets are two members of the immunoglobulin gene superfamily, CTLA-4 and programmed death (PD)-1 (CD279) (Figure 2).

Antigen-specific T-cell activation requires at least two signals. Firstly, an antigen is presented to a T-cell receptor (TCR) by an MHC molecule on antigen-presenting cells (APCs). Secondly, B7-1 and B7-2 on APCs bind to CD28 on T-cells and enhance T-cell stimulation. Coinhibitory molecules, such as CTLA-4, on T-cells are useful to prevent inappropriate stimulation of T-cells and possibly autoimmune disease; however, upregulation of CTLA-4 can promote the preferential binding of this molecule (CTLA-4) to B7-1 or B7-2, inducing suppression of T-cell activity. Similarly, binding of PD-1, primarily expressed on tumor-infiltrating T-cells, to PD ligand (PD-L)-1, (B7-H1, CD274) or to PD-L2, represents another coinhibitory and immunosuppressive T-cell mechanism in cancer.

In the clinic, a recent strategy in use to overcome immunosuppression is by antibody blockade of immune-checkpoint inhibitory molecules, such as CTLA-4 and PD-1,66,73,74 and several companies are currently designing molecules to optimize the recently FDA-approved targeted-therapy mechanism of immune-checkpoint antibody blocking.

**Scope of cancer immunotherapy**

Immunotherapy aims to diminish existing immunosuppressive mechanisms in cancer by blocking these mechanisms and/or to potentiate specific memory T-cell or antibody responses by vaccine administration or cellular immunotherapy, with the ultimate goal of disease management or cure, and improved survival. Unfortunately, there are few identified biomarkers in cancer that can be universally targeted in the treatment of a specific disease. Therefore, current immunotherapy approaches primarily target the immunosuppressive mechanisms discussed in the foregoing section. Immunotherapeutic strategies range from the use of tumor cell-based vaccines, virus-based immunotherapy, antibody immunotherapy, and cellular immunotherapy. We now cover these areas and discuss related preclinical studies and clinical trials.

**Tumor cell-based vaccines**

Whole tumor cells have been employed as a useful tool against various forms of cancer. For example, the most clinically advanced trial for pancreatic cancer is the use of algenpantuzumabostin-L immunotherapy. This is an irradiated live combination of two human allogeneic pancreatic cancer cell lines modified to express the murine enzyme α-1,3 galactosyltransferase, which is needed for the synthesis of α-galactosyl epitopes on surface proteins and glycolipids of such cell lines. The mouse gene causes the cells to be recognized as foreign to patients’ immune system, and thus the immune system attacks these cancer cells and destroys many of them. This is classified as HyperAcute™ immunotherapy (designed by NewLink Genetics, Ames, IA, USA), and similar products
are currently being developed with specificity for other types of cancer (non-small-cell lung cancer [NSCLC], advanced melanoma, metastatic castrate-resistant prostate cancer, and renal cancer). HyperAcute immunotherapies do not require tissue from individual patients, but use intact whole cells rather than cell fragments.

For pancreatic cancer, Phase II clinical trials with this technology were very encouraging, where data on 69 patients showed that overall survival at 3 years was 39%. This prompted two Phase III trials for pancreatic cancer using algenpantucel-L immunotherapy (HyperAcute pancreas). IMPRESS (IMmunotherapy for Pancreatic REsectable cancer Survival Study) involves up to 722 patients with surgically resected pancreatic cancer (ClinicalTrials.gov identifier: NCT01072981). PILLAR (Pancreatic Immunotherapy with algenpantucel-L for Locally Advanced non-Resectable; given with or without chemotherapy and radiation) (ClinicalTrials.gov identifier: NCT01836432), is currently enrolling patients with locally advanced pancreatic cancer. Therapeutic advances in pancreatic cancer are desperately needed, as with existing therapy this disease has a 5-year survival rate of below 10%.

Another tumor-cell therapy regimen consisting of autologous tumor cells conjugated to dinitrophenyl, called M-Vax, was used for the treatment of malignant melanoma (ClinicalTrials.gov identifier: NCT00257465).

OncoVAX, a tumor-cell product, is in Phase III clinical trials for colon cancer patients. The OncoVAX regimen consists of intradermally injecting two doses of autologous irradiated (200,000 rads) tumor cells, mixed with fresh-frozen mycobacteria of the Tice strain of bacillus Calmette–Guérin, followed by two injections of irradiated tumor cells alone. Bacillus Calmette–Guérin is a live but weakly pathogenic bacterium, and induces a strong immune response, and hence it is incorporated in the first two vaccines to boost the immune response.

BiovaxID (ClinicalTrials.gov identifier: NCT00091676) is in the regulatory approval process in Canada and Europe for late-stage indolent follicular lymphoma.

Other tumor-cell products that may be used as immunotherapeutic agents are the heat-shock protein (HSP) family molecules. HSPs act as molecular chaperones. They can be induced or released during cellular stress and necrosis. They bind potential antigens on cell death, and deliver them to APCs through several mechanisms. Tumor cells can secrete HSP70-containing exosomes, which recruit MDSCs, thereby contributing to immunosuppression. On the contrary, HSP70 peptide complexes are also secreted from necrotic tumor cells and can trigger anticancer CTL after entering APCs and cross-presenting to CD4 T-cells in afferent lymph nodes, leading to inhibition of tumor growth. HSPs can also bind tumor-associated antigens and deliver them to APCs through MHC I and MHC II molecules, inducing the activation of anti-tumor CD4+ and CD8+ T-cells. The antitumor effects of HSP have been investigated in several preclinical studies, and due to the positive results it has advanced to clinical trials. To mention a few examples, clinical trials with HSP96 are currently in progress for the immunotherapy of diseases, such as glioblastoma (ClinicalTrials.gov identifiers: NCT01814813 and NCT02122822).

**Virus-based immunotherapy**

Some virus infections can lead to cancer, but on the contrary, virus technology is often of significant benefit in cancer immunotherapy. Engineering of viruses to express novel cancer-specific proteins and other molecules is currently being employed in several clinical trials. ProstVac-VF, a prostate-specific antigen (PSA)-targeted therapeutic vaccine, is a combination of recombinant bacterium, and induces a strong immune response, and<br>PsA-Tricom showed clinical benefit and development of specific immune responses in patients. A detailed summary of the scope of viral vector immunotherapy strategies is outlined elsewhere. Oncolytic viruses infect and kill cancer cells and associated endothelial cells, preferentially to normal cells. The death of cancer cells occurs by immunogenic apoptosis, autophagic cell death, necrosis, and pyroptosis, and cell proteins are processed and presented by DCs stimulating antitumor and antiviral immune responses. Current technology allows the manipulation of the viral genome to improve safety, and the insertion of transgenes to augment antitumor activity. Ongoing clinical trials with modified adenovirus, herpesvirus, reovirus, measles, and other viruses will provide critical information regarding the safety and efficacy of oncolytic virus immunotherapy. In general, toxicity with virus clinical trials has been minor.

**Vaccines against pathogens that cause cancer**

Prophylactic vaccination therapy is effective against some viruses that cause cancer. In 1981, the FDA approved the
hepatitis B virus vaccine, now given to infants. This vaccine reduces hepatitis B virus infections and incidences of hepatocellular carcinoma.\textsuperscript{90}

Human papillomavirus (HPV) causes almost all cases of cervical cancer, as well as some anal, vulval, vaginal, penile, and oropharyngeal cancers.\textsuperscript{91,92} Courses of the FDA-approved quadrivalent vaccine Gardasil (Merck Pharmaceuticals, 2006) against HPV types 6, 11, 16, and 18, and the bivalent vaccine Cervarix (GlaxoSmithKline, 2009) against HPV types 6 and 11 are efficient in preventing cervical cancer and genital warts in individuals at risk. Gardasil is approved in many countries for the prevention of cervical cancer. In these vaccines, immune memory is primarily mediated by B cells.\textsuperscript{91–93} The shortcoming with these current prophylactic HPV vaccines, however, is that they are not effective in individuals who already have HPV infections. Therefore, vaccines such as VGX-3100 are being tested in women with cervical dysplasia. This vaccine induces antibodies to HPV serotypes, and specific CD8\textsuperscript{+} T-cells with granzyme B, perforin, and cytolytic potential.\textsuperscript{94,95}

**Cytokine immunotherapy**

The use of cytokine treatment to repolarize the immune system was one of the early forms of immunotherapy. IFN\(\alpha\) was one of the first cytokines approved for use in leukemia\textsuperscript{96,97} and melanoma\textsuperscript{98,99} patients. IL-2, FDA-approved in 1998, is effective in metastatic melanoma and in renal cell carcinoma. IL-2 in combination with granulocyte macrophage colony-stimulating factor (GM-CSF) or with chemotherapy may have added benefits.\textsuperscript{5,100,101} Cytokine GM-CSF stimulates the differentiation of stem cells to granulocytes and monocytes, and this treatment has been used in combination with primed lymphocytes in hematopoietic diseases, such as acute myeloid leukemia.\textsuperscript{102} One of the drawbacks in cytokine immunotherapy is toxic side effects. For example, TNF\(\alpha\) used in the therapy of melanoma patients may induce a septic shock-like condition if not administered at the right doses.\textsuperscript{103} Recent preclinical studies showed that cytokine IL-12 therapy combined with blocking anti-CTLA-4 therapy causes a decrease in FoxP3\textsuperscript{+} T regs and an increase in effector CD4\textsuperscript{+} T-cells.\textsuperscript{104}

**Antibody immunotherapy to reduce immunosuppression**

There are several mechanisms whereby antibody immunotherapy can be used to overcome immunosuppression in cancer or to potentiate antitumor immunity. Some examples follow.

**Anticytokine antibodies**

IL-6 promotes early colitis-associated cancer, and thus anti-IL-6 antibody immunotherapy may be useful in the treatment of such diseases as colorectal cancer.\textsuperscript{105}

**Depleting CD47 on tumor cells**

CD47 on tumor cells binds to its receptor – signal regulatory protein-\(\alpha\) – on phagocytic cells, and prevents phagocytic cells from ingesting tumor cells. This process is mediated by the selective expression of calreticulin on tumor cells. Antibody blocking of CD47 in mouse models of different cancers results in dramatic improvements or failure to induce disease.\textsuperscript{106–109} Engulfment of tumor cells by macrophages after anti-CD47 antibody blockade results in improved antigenspecific CD8\textsuperscript{+} T-effector cell immunity.\textsuperscript{110}

**Abrogation of T regs**

Daclizumab, an anti-CD25 antibody was effective in inducing a prolonged decrease in T regs in breast cancer patients.\textsuperscript{111,112} Denileukin difitox, an IL-2 diphtheria-toxin fusion protein, was used to treat cutaneous T-cell lymphoma and melanoma\textsuperscript{113} and for clinical trials in several other cancers, including one with ovarian cancer by the authors (ClinicalTrials.gov identifier: NCT00703105). Cyclophosphamide is frequently used in clinical trials as a mechanism of targeting T regs.\textsuperscript{114–118}

**Enhancement of immune responses**

CD40 expressed in high levels on the surface of mature DCs binds to CD40 ligand (CD40L) on T-cells, and increases costimulation and effective T-cell immune responses. Ligation of CD40 to CD40L mediates IL-12 production in DCs. Anti-CD40 monoclonal antibodies and IL-2 or IL-15, an NK- and T-cell activator, was effective in the treatment of a murine model of colon cancer.\textsuperscript{119,120} CD137 is expressed on activated T-cells, DCs, and NK cells. It increases T-cell proliferation and IL-2 secretion, and these properties make it a good candidate for antitumor immunotherapy. Clinical trials with anti-CD137 (BMS-663513) have been conducted in melanoma, ovarian cancer, and NSCLC.\textsuperscript{121}

**Immune-checkpoint antibody-blocking immunotherapy**

CTLA-4 and PD-1 prevent immune mediated damage to normal tissues in healthy individuals. In cancer, interaction of these molecules with their ligands B7-1 and B7-2 (for CTLA-4) and PD-L1 or PD-L2 (for PD-1) have been identified as significant immunosuppressive...
pathways.\textsuperscript{71,122–125} A blocking anti-CTLA-4 antibody, ipilimumab, is FDA-approved (2011) for treating metastatic malignant melanoma.\textsuperscript{16,17,72} Ipilimumab is a monoclonal antibody designed to block CTLA-4, thereby preventing the development of tolerance, and augmenting immune responses. Clinical trials are ongoing with this treatment for prostate cancer, NSCLC, renal cancer, and pancreatic cancer, as well as hematologic malignancies.\textsuperscript{73,126–129} In preclinical studies of a murine tumor model, CTLA-4 blockade in synergy with a GM-CSF expressing tumor-derived vaccine enhanced T-cell activation and memory and elicited antitumor T-cell responses in the early stages of tumor growth.\textsuperscript{130}

Phase I, II, or III clinical trials are in progress with ipilimumab, or with another anti-CTLA-4 antibody blocker, tremelimumab, in combination with cellular vaccines, chemotherapy, radiation, cytokine treatment, or adoptive T-cell-transfer therapy, and are reviewed in Gelao et al.\textsuperscript{73} Both agents recognize CTLA-4 and prevent ligation of CTLA-4 with B7 molecules. Even though these treatments show significant promise, ipilimumab therapy is not without side effects, many of which are gastrointestinal.\textsuperscript{72,131,132}

In September 2014, Keytruda (pembrolizumab), to be used for the treatment of unresectable or metastatic melanoma, was the first anti-PD-1 receptor therapy to be FDA-approved. An anti-PD-1 monoclonal antibody, nivolumab, has been investigated in Phase I and II trials in 296 patients with pretreated NSCLC, prostate cancer, renal cell carcinoma, colorectal cancer, and melanoma.\textsuperscript{133,134} Like anti-CTLA-4 antibody-blocking therapy, anti-PD-1 and anti-PD-L1 antibodies are used in single or combination therapy, resulting in some encouraging findings, but with fewer adverse gastrointestinal, skin, liver, and endocrine system side effects than ipilimumab. Combination therapy with these checkpoint inhibitors and other therapy shows more promise than either therapy alone, without additional side effects.\textsuperscript{73,74,130,133,134}

Glioblastoma is a deadly cancer in which patients are expected to survive 1 year after diagnosis.\textsuperscript{135} This disease is characterized by a highly suppressive T$_{reg}$ environment.\textsuperscript{136,137} In a recent mouse-model study of this disease, IL-12 fusion protein (IL-12 Fc) was delivered directly into cerebral tumors via osmotic minipumps over a 28-day period.\textsuperscript{104} This was combined with systemic administration of anti-CTLA-4 antibody treatment. This combined treatment led to full remission in most mice and greatest improved histological response, unlike the outcome for either monotherapy. Additionally, IL-12 Fc treatment led to a dramatic increase in CD4 T-cells, whereas both treatments resulted in increases in CD8 T-cells. Fluorescence-activated cell-sorting analysis showed that between days 21 and 35 after injection, there was an increase in total CD4+ T-cells, and a strong increase in the frequency of these cells that produced IFN-$\gamma$, but a profound reduction in FoxP3+ cells. There was a long-lasting improved effect even after 91 days of therapy. Interestingly, IL-12 Fc increased the expression of perforin in CD8+ T-cells and in NK cells, but not in CD4+ T-cells, indicating that CTLs are primarily responsible for tumor control.\textsuperscript{104} Other studies indicate that selective opsonization of T$_{reg}$ with anti-CTLA-4 antibodies can induce potent antibody-dependent cell-mediated cytotoxicity within the tumor site.\textsuperscript{138}

### Cellular immunotherapy in cancer

Since the FDA approval of the DC immunotherapy sipuleucel-T for castration-resistant prostate cancer, investigators have been prompted to initiate clinical trials and preclinical studies with novel ex vivo manipulations of DCs, and to determine appropriate combination therapies that would enhance and optimize DC-vaccine regimens. Additionally, the field of T-cell immunotherapy for cancer has been rapidly expanding.\textsuperscript{139} Hematopoietic stem cell transplant (HSCT) from a healthy donor is a potent form of cellular immunotherapy to treat hematologic malignancies, and is sometimes the last option in children who have relapsed malignancy.\textsuperscript{140} In some trials, TILs are isolated and used for adoptive cell transfer.\textsuperscript{141} Recent advances in the field of T-cell immunotherapy have been marked by genetic manipulations of infused T-cells, either by TCR or chimeric antigen receptor (CAR) gene transfer.\textsuperscript{142} The following sections will discuss advances in DC-based and T-cell immunotherapy.

### Dendritic cell-based vaccines

DCs are the most potent APCs in the immune system. They were first reported in the skin in 1948,\textsuperscript{143} and later characterized in spleens of mice in 1973.\textsuperscript{144} DCs capture, process, and present antigens, and induce the generation of specific effector and memory T-cells. DCs express costimulatory molecules that regulate immune activation and anergy, T-cell differentiation, and cytokine polarization. DCs prime both T-cell and humoral immune responses.\textsuperscript{145–147}

DCs recognize tumor-associated antigens, and have been used in several clinical trials over the past two decades exploiting their inherent properties.\textsuperscript{148–150} The most successful DC immunotherapy to date is the FDA-approved sipuleucel-T (Provenge; Dendreon) for castration-resistant prostate cancer.\textsuperscript{15} Prostatic acid phosphatase is restricted to prostate tissue, and is thus a specific target in metastatic prostate cancer.\textsuperscript{151} In this personalized vaccine, sipuleucel-T
is prepared with autologous APCs treated with recombinant
fusion protein PA2024. PA2024 is a fusion protein consisting
of full-length human prostatic acid phosphatase and
full-length human GM-CSF. A complete course of
sipuleucel-T therapy, resulting in improved survival in these
prostate cancer patients, consists of three doses of the vaccine
at about 2-week intervals.

Recently, a novel approach was used to treat malignant
glioma in a cohort of 22 patients. Mature DC classified
as alpha-DC-1 were prepared in cytokines GM-CSF, IL-4,
and a potent cytokine-maturation cocktail, consisting of
IL-1β, TNFα, IFNγ, IFNα, and poly-I-C, as previously
described. Two hours before harvesting DCs, cells were
loaded with Pan-DR epitope peptide (PADRE) optimized for
Th-cell response, and with glomerular-associated peptides.
Alpha-DC-1 were administered intranodally in the right or
left inguinal and axillary lymph nodes for alternate injections
at 2-week intervals (one to four vaccines), and polynosinic–
polyuridylic acid stabilized by lysine and carboxymethyl
cellulose (poly-ICLC; 20 μg/kg) was given intramuscu-
larly twice weekly to all patients for 8 weeks. Poly-ICLC
enhances the efficacy of glioma-associated antigen-targeting
vaccinations. Vaccine regimens were subject to later
repeat courses in patients who did not have adverse responses.
IL-12p70 secreted by patient DCs correlated positively
with time to progression, ie, high DC IL-12 secretion was
associated with improved outcome. Nine patients achieved
progression-free status lasting at least 12 months, while
58% of patients showed improved Th1 (eg, IFNγ) responses
in peripheral blood mononuclear cells to at least one of the
vaccination-targeted glioma-associated antigens.

Survival benefits of DC vaccines have been reported in
several clinical trials, but like most immunotherapy there
are still several hurdles to overcome in order to optimize this
treatment for cancer and other diseases. Today, most investi-
gators, including the authors who have ongoing ovarian can-
cer clinical trials with α-DC-1 (ClinicalTrials.gov identifier:
NCT00703105), believe that a mature DC vaccine, such as
α-DC-1, is superior to an immature DC for immunotherapy,
but there are still several unanswered questions concerning
methods of DC optimization for improved therapy. These
questions relate to the cytokine culture environment of
DCs, ex vivo manipulation and selection of antigens for
loading, method of delivery of antigens, the route of vac-
cine administration, the frequency of administration, and
dose of cells. The antigens used for loading DCs varies
with the disease. Proteins, deoxyribonucleic acid (DNA),
ribonucleic acid, or autologous tumor lysate can be used to
load DCs. Some groups have used tumor-associated
antigens or overexpressed tumor antigens, such as the cancer
testis antigens NY-ESO-1, MAGE-family antigens, antigens
derived from melanocyte-differentiation factors (eg, gp100 or
MART), antigens encoded by oncopenes, MUC-1, HER-2/
neu, and other antigens. DC incubation with the anti-
gens or delivery of tumor-associated antigens with bacterial
or viral vectors are strategies for loading DCs. As expected, a consideration of all of these parameters deter-
mines whether CD4+ and/or CD8+ T-cells are activated, and
consequently the benefit of DC therapy to patients.

Another class of cancer antigens currently under investiga-
tion for immune targeting is neoantigens. These antigens
are derived from mutated proteins present in tumors, but not
found in normal individuals. Neoantigens may prove
in future trials to be one of the most successful antigens for
delivery to DCs for use in patient vaccines, since patients
can mount a sustained antitumor immune response against
these unique tumor antigens.

Hematopoietic stem cell transplants
Bone marrow transplants, peripheral blood stem cell trans-
plants, or cord-blood transplants are grouped under the
term HSCTs. Allogeneic HSCTs are the most widely used
form of adoptive T-cell immunotherapy. This method was
developed over 50 years ago to treat patients with anemias
and immune deficiencies, and is now frequently used to
treat hematologic malignancies and some solid cancers.
HSCT has become standard immunotherapy for leukemia
and lymphoma at several large centers, including that of
the authors.

HSCT allows the delivery of myeloablative (high)
doses of radiation or chemotherapy for increased killing
of tumor cells, in comparison with conventional doses of
these standard therapies. The patient’s bone marrow, which
can no longer function, is rescued with intravenous infusion
of HSCs. Peripheral blood SCTs account for 95% of adult
autologous transplants, as well as 70% of adult allogeneic
transplants. In children, bone marrow is commonly used
for these allogeneic transplants, since the outcome is better.
Several milestones have been accomplished in HSCT treat-
ment since the start of this therapy in 1951. Notably, over
the last 20 years, it has been found that cord blood harvested
shortly after birth is very rich in HSCs. Today, there are many
stem cell banks that provide stem cell products for allogeneic
HSCT.

The ultimate goal in HSCT cancer immunotherapy is
to improve engraftment, decrease graft-versus-host disease,
augment graft-versus-tumor effects, and increase survival. To this end, investigators are studying mechanisms to culture and expand HSCs, without further differentiation, by such techniques as using bone marrow stromal cell lines and various cytokine cocktails to optimize this expansion either on a small scale or commercially, or by regulation of molecular pathways in these cells. Since 2006, approximately 40% of allogeneic transplants have used nonmyeloablative conditioning (lower doses of chemotherapy and radiation) of patients. This strategy allows older patients and those with other diseases to benefit from HSCT. Improved techniques and knowledge concerning human leukocyte-antigen matching and posttransplant therapy, such as high-dose cyclophosphamide to deplete T-cells and suppress graft-versus-host disease, allows better identification of suitable donors, and consequently more patients to benefit from HSCT therapy. Furthermore, molecular characterization of the donor and recipient cells can give clues to better matching and to patient outcome after transplant. Additional measures that may improve the success of HSCT include the use of drugs to increase homing of CD34+ cells to the bone marrow. For example, CXCR4 and its ligand CXCL12 (SDF1) direct CD34+ stem cells to migrate to the bone marrow. CD34+ stem cells of bone marrow have the ability to engraft and give rise to stem cells of diverse hematopoietic origins. Plerixafor, a CXCR4 antagonist, given in combination with G-CSF, increases mobilization of stem cells. This combination therapy is FDA-approved for patients with non-Hodgkin’s lymphoma and multiple myeloma.

In a Phase II immunotherapy-combination trial, multiple myeloma patients were given DC-tumor infusions before and after autologous SCT. Twenty-four patients received serial vaccinations with DC-myeloma fusion cells following posttransplant hematopoietic cell recovery. A second group received pretransplant vaccine as well as posttransplant vaccine administration. Seventy-eight percent of patients achieved a best response of complete response + very good partial response. Immune monitoring of patients revealed that vaccination resulted in a marked expansion of myeloma-reactive CD4+ and CD8+ T-cells expressing IFNγ in response to autologous tumor lysate.

**Targeting of tumors with modified T-cells**

TIL therapy has been used in preclinical models and in clinical trials with some success. However, the production of these cells is a complex process, generally limited to specialized centers. Isolated TILs (T-cells) are harvested, activated, and expanded in culture and then infused back into the patient, usually with IL-2 administration. In this form of adoptive cell transfer, these cells can traffic to the tumor and lead to prolonged tumor eradication.

There are two main types of genetically engineered T-cells used for adoptive cell therapy for cancers: TCR gene-modified and CAR gene-modified T-cells. These cells have high avidity and high reactivity to tumor antigens.

T-cells may be genetically modified to enhance the expression of selected high-affinity TCR before infusion into patients (ClinicalTrials.gov identifier: NCT01586403). T-cells are isolated from patients and cultured and expanded in cytokine cocktails. Cells are genetically altered using vectors containing nucleic acids encoding molecules that enable T-cells to recognize and mount immune responses to cancer cells. These vectors, often replication-deficient retroviruses or lentiviruses, or DNA plasmid-based vectors, deliver the selected genes to T-cells; these genes are incorporated into the T-cell genome, and hence passed down to dividing cells. Genes inserted into T-cells can enhance tumor-cell recognition, tumor-cell killing, cause T-cells to migrate into tumors, increase T-cell proliferation, or overcome factors in the tumor-suppressive microenvironment.

CARs combine antigen specificity with T-cell activation in a single fusion molecule. The structure consists of an antigen-binding domain, an extracellular domain spacer/hinge region, a transmembrane domain, and an intracellular signaling domain, leading to T-cell activation after antigen binding. For CAR T-cell therapy, a patient’s T-cells are collected and genetically altered to produce special receptors on their surface. These CAR proteins allow the T-cells to recognize a specific antigen on tumor cells. The altered cell is expanded in culture several-fold and then administered to the patient. These T-cells multiply in vivo and recognize and kill cancer cells bearing the target antigen independently of MHC I. This therapy has proved very effective, especially in children. There have been a limited number of successful cases so far, as this form of therapy is still developing. However, recently the CAR T-cell therapy CTL019 gained FDA breakthrough designation for the treatment of acute lymphoblastic leukemia. In this trial, the T-cells were released into the patient’s blood, where they proliferated and bound to the targeted CD19+ cells and destroyed them. CD19 is commonly expressed on B cells, and hence this is a critical tool against B-cell leukemia. There are several other ongoing clinical trials in this area for hematologic malignancies and solid cancer, but most
genetically modified T-cell therapy is in the experimental stages.\textsuperscript{142,188,194,195}

**Conclusion**

Since the original debate on cancer immunosurveillance about a century ago, there has been significant knowledge gained concerning events in the tumor microenvironment and immunosuppression in the host. Scientists have been able to use this information to design immunotherapy strategies to exploit the patient’s immune system to improve patient survival. Therefore, the use of anticytokine antibody therapy, immune-checkpoint inhibition, autologous DC immunotherapy, and T-cell immunotherapy are all designed to overcome immunosuppression, improve patient care, and ultimately prolong survival.

Today, immunotherapy has advanced from improvements in a few cases to FDA approval of several immunotherapeutic agents. However, even with achievements in DC immunotherapy, such as sipuleucel-T, immune-checkpoint blocking antibodies, such as ipilimumab and Keytruda (pembrolizumab) and the recent FDA breakthrough designation for CAR T-cell therapy CTL019, the greatest barrier to new immunotherapy developments is the lack of relevant disease biomarkers for each disease so that the immune system can be more effectively targeted.

Recently, investigators reported a detailed genomic analysis of adenocarcinoma, a cancer found in the tissues near the outer parts of the lungs. They identified 18 statistically significant gene mutations in a study of 230 adenocarcinomas,\textsuperscript{186} a cancer with very low 5-year survival. Similar genomic analysis has been done for ovarian, colon, and breast cancer and acute myeloid leukemia, findings that we hope will facilitate the identification of additional biomarkers for diagnosis and/or therapy.

Many immunotherapeutic agents are still in the experimental or clinical trial phase, but we are encouraged, because many of these trials have shown sufficient success to prompt Phase II and III clinical trials. Additionally, several large and mid-size biotechnology companies are providing experimental drugs for preclinical and clinical trial testing. Combined targeting of the immune system with different classes of immunotherapeutic agents, DC, and/or adoptive T-cell therapy, will deliver the optimum therapeutic force against cancer.

In the case of DC immunotherapy, it is likely that improved nanoparticle delivery of relevant molecules, such as those of chemokine receptor CCR7, which guide DCs to lymph nodes, will enable DCs to traffic, persist in vivo, and maintain function to stimulate potent and sustained antigen-specific T-cell responses. The use of target molecules, such as neoantigens in DC immunotherapy, is also a promising strategy.

CAR T-cell immunotherapy is also an attractive immunotherapeutic mechanism, because benefit is derived through an MHC I-independent mechanism, and thus the effect of tumors downregulating MHC I to evade detection is not a concern. There are several ongoing clinical trials with genetically redirected T-cells in solid and hematologic cancers.\textsuperscript{142} However, different delivery vectors need to be tested, as do the best conditions under which to expand T-cells, as well as incorporation of specific/multiple target genes required to polarize the immune system from immunosuppressive to antitumor, and to avoid immune escape mechanisms of the tumor cells.

In the US, about a quarter of deaths each year is due to cancer. This is a very exciting and productive phase in the era of cancer-immunotherapy development, but given the complexity of immunosuppression in cancer, researchers must continue to elucidate new biomarkers for the disease and test effective immunotherapy combinations to treat this disease. Additionally, efforts need to be directed toward minimizing and managing adverse reactions to immunotherapy, and precautions need to be taken to reduce infection risks of potent immunotherapy. Finally, there is also the need for industry and health care systems to design creative ways to overcome the high cost of immunotherapy, so that this novel approach to cancer treatment can reach all who need it.

**Disclosure**

The authors report no conflicts of interest in this work.

**References**

1. Dimberu PM, Leonhardt RM. Cancer immunotherapy takes a multi-faceted approach to kick the immune system into gear. *Yale J Biol Med.* 2011;84(4):371–380.

2. Jemal A, Bray F, Center MM, Ferlay J, Ward E, Forman D. Global cancer statistics. *CA Cancer J Clin.* 2011;61(2):69–90.

3. Dunn GP, Bruce AT, Ikeda H, Old LJ, Schreiber RD. Cancer immunoeediting: from immunosurveillance to tumor escape. *Nat Immunol.* 2002;3(11):991–998.

4. Dunn GP, Old LJ, Schreiber RD. The immunobiology of cancer immunosurveillance and immunoeediting. *Immunity.* 2004;21(2):137–148.

5. Zhou J. Advances and prospects in cancer immunotherapy. *N J Sci.* 2014;2014:745808.

6. Das B, Tsuchida R, Malkin D, Koren G, Baruchel S, Yeger H. Hypoxia enhances tumor stemness by increasing the invasive and tumorigenic side population fraction. *Stem Cells.* 2008;26(7):1818–1830.

7. Maitland NJ, Collins A. A tumour stem cell hypothesis for the origins of prostate cancer. *BJU Int.* 2005;96(9):1219–1223.

8. Odoux C, Fohrer H, Hoppe T, et al. A stochastic model for cancer stem cell origin in metastatic colon cancer. *Cancer Res.* 2008;68(17):6932–6941.
9. Selby P, Bizzini J, Buick R. Therapeutic implications of a stem cell model for human breast cancer: a hypothesis. Cancer Treat Rep. 1982; 67(7–8):659–663.
10. Siegel R, Ma J, Zou Z, Jemal A. Cancer statistics, 2014. CA Cancer J Clin. 2014;64(1):9–29.
11. Bickels J, Kollender Y, Merinsky O, Meller I. Coley’s toxin: historical perspective. Isr Med Assoc J. 2002;4(6):471–472.
12. McCarthy EF. The toxins of William B. Coley and the treatment of bone and soft-tissue sarcomas. Iowa Orthop J. 2006;26:154–158.
13. Nauts HC, Swift WE, Coley BL. The treatment of malignant tumors by bacterial toxins as developed by the late William B Coley, MD, reviewed in the light of modern research. Cancer Res. 1946;6: 205–216.
14. Nelson D, Fisher S, Robinson B. The “Turan horse” approach to tumor immunotherapy: targeting the tumor microenvironment. J Immunol Res. 2014;2014:799069.
15. KantoFF PW, Higano CS, Shore ND, et al. Sipuleucel-T immunotherapy for castration-resistant prostate cancer. N Engl J Med. 2010;363(5): 411–422.
16. Lipson EJ, Drake CG. Ipilimumab: an anti-CTLA-4 antibody for metastatic melanoma. Clin Cancer Res. 2011;17(22):6958–6962.
17. Prieto PA, Yang JC, Sherry RM, et al. CTLA-4 blockade with ipilimumab: long-term follow-up of 177 patients with metastatic melanoma. Clin Cancer Res. 2012;18(7):2039–2047.
18. Ehrlich P. Über den jetzigen stand der karzinomforschung. Ned Tidsskr Geneeskd. 1909;5:273–290.
19. Burnet F. The concept of immunological surveillance. Prog Exp Tumor Res. 1969;13:1–27.
20. Doll R, Kinlen L. Immunosurveillance and cancer: epidemiological evidence. Br Med J. 1970;4(5732):420–422.
21. Matsueda S, Graham DY. Immunotherapy in gastric cancer. World J Gastroenterol. 2014;20(7):1657–1666.
22. Staney CY, Rautela J, Parker BS. The emerging role of immunosurveillance in dictating metastatic spread in breast cancer. Cancer Res. 2013;73(19):5852–5857.
23. Romero JM, Jiménez P, Cabrera T, et al. Coordinated downregulation of the antigen presentation machinery and HLA class I/β2-microglobulin complex is responsible for HLA-ABC loss in bladder cancer. Int J Cancer. 2005;115(4):605–610.
24. Pagès F, Galon J, Pagès F, Tartour E, Sauvé-Fridman C, Kroemer G. Prognostic and predictive impact of intra- and peritumoral immune infiltrates. Cancer Res. 2011;71(17):5601–5605.
25. Galon J, Costes A, Sanchez-Cabo F, et al. Type, density, and location of immune cells within human colorectal tumors predict clinical outcome. Science. 2006;313(5795):1960–1964.
26. Pagès F, Galon J, Dieu-Noël Jean MC, Tartour E, Sauvé-Fridman C, Fridman WH. Immune infiltration in human tumors: a prognostic factor that should not be ignored. Oncogene. 2010;29(8):1093–1102.
27. Pagès F, Kirilovsky A, Mlecnik B, et al. In situ cytotoxic and memory T cells predict outcome in patients with early-stage colorectal cancer. J Clin Oncol. 2009;27(35):5944–5951.
28. Stiff PJ, Potkul RK, Venkataraman G, Sojitra P, Drakes ML. Immune surveillance tissue antigen profiling in advanced ovarian cancer. J Immunother. 2012;35(2):78–86.
29. Zhang L, Conejo-Garcia JR, Katsaros D, et al. Intratumoral T cells, recurrence, and survival in epithelial ovarian cancer. N Engl J Med. 2003;348(3):203–213.
30. Zeng YZ, Grote DM, Ziesmer SC. Al. Soluble and membrane-bound TGF-β-mediated regulation of intratumoral T cell differentiation and function in B-cell non-Hodgkin lymphoma. PLoS One. 2013;8(3):e59456.
31. Faget J, Biota C, Bachelot T, et al. Early detection of tumor cells by innate immune cells leads to T(reg) recruitment through CCL22 production by tumor cells. Cancer Res. 2011;71(19):6143–6152.
32. Gobert M, Treilleux I, Bendriss-Vermare N, et al. Regulatory T cells recruited through CCL22/CCR4 are selectively activated in lymphoid infiltrates surrounding primary breast tumors and lead to an adverse clinical outcome. Cancer Res. 2009;69(5):2000–2009.
33. Qin XJ SH, Deng JM, Liang QL, Jiang J, Ye ZJ. CCL22 recruits CD4-positive CD25-positive regulatory T cells into malignant pleural effusion. Clin Cancer Res. 2009;15(7):2231–2237.
34. Jaafar F, Righi E, Lindstrom V. Al. Correlation of CXCL12 expression and FoxP3+ cell infiltration with human papillomavirus infection and clinicopathological progression of cervical cancer. Am J Pathol. 2009;175(4):1525–1535.
35. Wald O, Izhar U, Amir G, et al. CD4+CXCR4+ highCD11b+ regulatory T cells accumulate in lung adenocarcinoma. J Immunol. 2006;177(10):6938–6990.
36. Capece D, Fischetti M, Verzella D, et al. The inflammatory microenvironment in hepatocellular carcinoma: a pivotal role for tumor-associated macrophages. Biomed Res Int. 2013;2013:187204.
37. Ostrand-Rosenberg S, Sinha P, Beury DW, Clements VK. Cross-talk between myeloid-derived suppressor cells (MDSC), macrophages, and dendritic cells enhances tumor-induced immune suppression. Semin Cancer Biol. 2012;22(4):275–281.
38. Ostrand-Rosenberg S, Sinha P, Chornoguz O, Ecker C. Al. Regulating the suppressors: apoptosis and inflammation govern the survival of tumor-induced myeloid-derived suppressor cells (MDSC). Cancer Immunol Immunother. 2012;61(8):1319–1325.
39. Pollard JW. Al. Tumour-educated macrophages promote tumour progression and metastasis. Nat Rev Cancer. 2004;4(1):71–78.
40. Drake CG, Jaffee E, Pardoll DM. Al. Mechanisms of immune evasion by tumors. Adv Immunol. 2006;90:51–81.
41. Tien AH, Xu L, Helgason CD. Al. Manrique SZ, Correa MA, Hoelzinger DB, et al. Foxp3-positive macrophages display immunosuppressive properties and promote tumor growth. J Exp Med. 2011;208(7):1485–1499.
54. Gabrilovich D, Ishida T, Oyama T, et al. Vascular endothelial growth factor inhibits the development of dendritic cells and dramatically affects the differentiation of multiple hematopoietic lineages in vivo. Blood. 1998;92(11):4150–4166.

55. Guiducci C, Vicari AP, Sangaletti S, Trinchieri G, Colomba MP. Redirecting in vivo elicited tumor infiltrating macrophages and dendritic cells towards tumor rejection. Cancer Res. 2005;65(8):3437–3446.

56. Suzuki E, Kapoor V, Jassar AS, Kaiser LR, Albelda SM. Geminibehinselectively eliminates splenic Gr-1+CD11b+ myeloid suppressor cells in tumor-bearing animals and enhances antitumor immunity. Clin Cancer Res. 2005;11(18):6713–6721.

57. Dong H, Chen L. B7-H1 pathway and its role in the evasion of tumor immunity. Mol Med (Brl). 2003;8(5):281–287.

58. Kryczek I, Wei S, Zhu G, et al. Relationship between B7-H4, regulatory T cells, and patient outcome in human ovarian carcinoma. Cancer Res. 2007;67(18):8900–8905.

59. Kryczek I, Zou L, Rodriguez P, et al. B7-H4 expression identifies a novel suppressive macrophage population in human ovarian carcinoma. J Exp Med. 2006;203(4):871–881.

60. Hinrichs CS, Kaiser A, Paulos CM, et al. Type 17 CD8+ T cells display enhanced antitumor immunity. Blood. 2009;114(3):596–599.

61. Llosa NJ, Geis AL, Orberg ET, Housseau F. Interleukin-17 and type 17 helper T cells in cancer management and research. Immunother Ther. 2014;2014(3):39–54.

62. Zou W, Restifo NP. T(H)17 cells in tumour immunity. Int J Biochem Cell Biol. 2014;57:1351–1361.

63. Inaba T, Ino K, Kajiyama H, et al. Role of the immunosuppressive enzyme indoleamine 2,3-dioxigenase in the progression of ovarian carcinoma. Gynecol Oncol. 2009;115(2):185–192.

64. Malachowski WP, Metz R, Prendergast GC, Muller AJ. A new cancer immunosuppression target: indoleamine 2,3-dioxigenase (IDO). A review of the IDO mechanism, inhibition and therapeutic applications. Drugs Future. 2005;30(9):879–813.

65. McKenna K, Beignon AS, Bhardwaj N. Plasmacytoid dendritic cells: linking innate and adaptive immunity. J Virol. 2005;79(1):17–27.

66. Sharma MD, Babin B, Chandler P, et al. Plasmacytoid dendritic cells from mouse tumor-draining lymph nodes directly activate mature Tregs via indoleamine 2,3-dioxigenase. J Clin Invest. 2007;117(9):2570–2582.

67. Wei S, Kryczek I, Zou L, et al. Plasmacytoid dendritic cells induce CD8+ regulatory T cells in human ovarian carcinoma. Cancer Res. 2005;65(12):5020–5026.

68. McDonnell AM, Robinson BW, Currie AJ. Tumor antigen cross-presentation and the dendritic cell: where it all begins? Clin Dev Immunol. 2010;2010:539519.

69. Seliger B. Different regulation of MHC class I antigen processing components in human tumors. J Immunother. 2008;31(4):361–367.

70. Inman BA, Frigola X, Dong H, Kwon ED. Costimulation, coinhibition and cancer. Curr Cancer Drug Targets. 2007;7(1):15–30.

71. Krempsji K, Karyampudi L, Behrens MD, et al. Tumor-infiltrating programmed death receptor-1 dendritic cells mediate immune suppression in ovarian cancer. J Immunol. 2011;186(12):6905–6913.

72. Wolchok JD, Neys B, Linette G, et al. Ipilimumab monotherapy in patients with pretreated advanced melanoma: a randomised, double-blind, multicentre, phase 2, dose-ranging study. Lancet Oncol. 2010;11(2):155–164.

73. Gelaio L, Criscitelli C, Esposito A, Goldhirsh A, Curigliano G. Immune checkpoint blockade in cancer treatment: a double-edged sword cross-targeting the host as an “innocent bystander”. Toxins. 2014;6(3): 914–933.

74. Lesterhuis WJ, Salmons J, Nowak AK, et al. Synergistic effect of CTLA-4 blockade and cancer chemotheraphy in the induction of antitumor immunity. PloS One. 2013;8(4):e61895.

75. Galili U, Clark MR, Shohet SB, Buchler J, Machet BA. Evolutionary relationship between the natural anti-Gal antibody and the Gal α1→3Gal epitope in primates. Proc Natl Acad Sci U S A. 1987;84(5):1369–1373.

76. Gunteru KS, Rossi GR, SafiMW. Immunotherapy updates in pancreatic cancer: are we there yet? Ther Adv Med Oncol. 2012;5(1):81–89.

77. Rossi GR, Mautino MR, Awwad DZ, et al. Allogeneic melanoma vaccine expressing αGal epitopes induces antitumor immunity to autologous antigens in mice without signs of toxicity. J Immunother. 2008;31(6):545–554.

78. Hanna MG Jr. Immunotherapy with autologous tumor cell vaccines for treatment of occult disease in early stage colon cancer. Hum Vaccin Immunother. 2012;8(8):1156–1160.

79. Murshid A, Gong J, Stevenson MA, Calderwood SK. Heat shock proteins and cancer vaccines: developments in the past decade and chaperoning in the decade to come. Expert Rev Vaccines. 2011;10(11):1553–1568.

80. Calderwood SK, Stevenson MA, Murshid A. Heat shock proteins, autoimmunity, and cancer treatment. Autoimmune Dis. 2012; 2012:486069.

81. Solit DB, Rosen N. Hsp90: a novel target for cancer therapy. Curr Top Med Chem. 2006;6(11):1205–1214.

82. Antonarakis ES. Combining active immunotherapy with immune checkpoint blockade for the treatment of advanced prostate cancer. Asian J Androl. 2012;14(4):520–521.

83. Madan RA, Mohebthash M, Arlen PM, et al. Iplumimub and a poxviral vaccine targeting prostate-specific antigen in metastatic castration-resistant prostate cancer: a phase 1 dose-escalation trial. Lancet Oncol. 2012;13(5):501–508.

84. Ursello V, Koski A, Vähä-Koskela M, Hemminki A. Chapter eight – Oncolytic adenoviruses for cancer immunotherapy: data from mice, hamsters, and humans. Adv Cancer Res. 2012;115:265–318.

85. Guo ZS, Thorne SH, Bartlett DL. Oncolytic virotherapy: molecular targets in tumor-selective replication and carrier cell-mediated delivery of oncolytic viruses. Biochim Biophys Acta. 2008;1782(2):217–231.

86. Patel MR, Kratzke RA. Oncolytic virus therapy for cancer: the first wave of translational clinical trials. Transl Res. 2013;161(4):355–364.

87. Russell SJ, Peng KW, Bell JC. Oncolytic virotherapy. Nat Biotechnol. 2012;30(7):658–670.

88. Chang MH, You SL, Chen CJ, et al. Decreased incidence of hepatocellular carcinoma in hepatitis B vaccinees: a 20-year follow-up study. J Natl Cancer Inst. 2009;101(19):1348–1355.

89. Olsson SE, Villa LL, Costa RL, et al. Induction of immune memory following administration of a prophylactic quadrivalent human papil-lomavirus (HPV) types 6/11/16/18 virus-like particle (VLP) vaccine. Vaccine. 2007;25(26):4931–4939.

90. Van Kriekinge G, Castellsague X, Cibula D, Demarteau N. Estimation of the potential overall impact of human papillomavirus vaccination on cervical cancer cases and deaths. Vaccine. 2014;32(6):733–739.

91. Villa LL, Ault KA, Giuliano AR, et al. Immunologic responses following administration of a vaccine targeting human papillomavirus types 6, 11, 16, and 18. Vaccine. 2006;24(27):5571–5583.

92. Bagarazzi ML, Yan J, Morrow MP, et al. Immunotherapy against HPV16/18 generates potent TH1 and cytotoxic cellular immune responses. Sci Transl Med. 2012;4(155):155ra138.

93. Maldonado L, Teague JE, Morrow MP, et al. Intramuscular therapeutic vaccination targeting HPV16 induces T cell responses that localize in mucosal lesions. Sci Transl Med. 2014;6(22):221ra213.

94. Bogdanov KV, Frolova OI, Marinets OV, Ogordnikova IuS, Afanas'ev BV, Zaritski AIu. [The effectiveness of interferon-alpha therapy in Ph-positive chronic myeloid leukemia]. Vopr Onkol. 2003;49(2):189–192. Russian.

95. Lutz D. [Interferon-alpha therapy in chronic myeloid leukemia]. Wien Med Wochenschr. 1993;143(16–17):416–419. German.

96. Moelling S, Pasquali S, Rossi CR, Nitti D. Interferon alpha adjuvant therapy in patients with high-risk melanoma: a systematic review and meta-analysis. J Natl Cancer Inst. 2010;102(7):493–501.
99. von Wuswass P, Block B, Hartmann F, Deicher H. Intraläsionales interferon-alpha therapy in advanced malignant melanoma. *Cancer*. 1986;61(6):1071–1074.

100. Elias EG, Zapas JL, Beam SL, Brown SD. GM-CSF and IL-2 combination as adjuvant therapy in cutaneous melanoma: early results of a phase II clinical trial. *Oncology*. 2005;19(4 Suppl 2):15–18.

101. Fateh S, Schell TD, Gingrich R, Neves RI, Drabick JJ. Unsuccessful low dose temozolomide can result in rapid durable complete and near-complete remissions in metastatic melanoma. *Cancer Biol Ther*. 2010;10(11):1091–1097.

102. Borrello IM, Levitsky HI, Stock W, et al. Granulocyte-macrophage colony-stimulating factor (GM-CSF)-secreting cellular immunotherapy in combination with autologous stem cell transplantation (ASCT) as postremission therapy for acute myeloid leukemia (AML). *Blood*. 2009;114(9):1736–1745.

103. Sorokin P, Abu-Abid S, Lev D, et al. Systemic leakage and side effects of tumor necrosis factor alpha administered via isolated limb perfusion can be manipulated by flow rate adjustment. *Arch Surg*. 1995;130(10):1079–1084.

104. Von Berg J, Vrolihings M, Hailer S, et al. Intratumoral IL-12 combined with CTLA-4+ blockade elicits T cell-mediated glioma rejection. *J Exp Med*. 2013;210(13):2803–2811.

105. Wang K, Grivennikov SI, Karin M. Implications of anti-cytokine mediators of tumor immunosurveillance. *Trends Immunol*. 2010;31(6):212–219.

106. Sick E, Jeanne A, Schneider C, Dedieu S, Takeda K, Martiny L. CD47 update: a multifaceted actor in the tumour microenvironment of potential therapeutic interest. *Br J Surg*. 2012;167(7):1415–1430.

107. Edris B, Weiskopf K, Volkmer AK, et al. Antibody therapy targeting the CD47 protein is effective in a model of aggressive metastatic leiomyosarcoma. *Proc Natl Acad Sci U S A*. 2012;109(17):6656–6661.

108. Jaiswal S, Chao MP, Majeti R, Weissman IL. Macrophages as mediators of tumor immunosurveillance. *Trends Immunol*. 2010;31(6):98–102.

109. Vermeij R, Leffers H, Hoogeboom BN, et al. Potentiation of a p53-SLP vaccine by cyclophosphamide in ovarian cancer: a single-arm phase II study. *Int J Cancer*. 2012;131(5):E670–E680.

110. Fonsatti E, Maio M, Altomonte M, Hersey P. Biology and clinical applications of CD40 in cancer treatment. *Semin Oncol*. 2010;37(5):517–523.

111. Zhang M, Yao Z, Dubois S, Ju W, Muller JR, Waldmann TA. Interleukin-15 combined with an anti-CD40 antibody provides enhanced therapeutic efficacy for murine models of colon cancer. *Proc Natl Acad Sci U S A*. 2009, 106(18):7513–7518.

112. Szolov M, Hodi F, Margolin K, et al. Phase I study of BMS-663513, a fully human anti-CD137 agonist monoclonal antibody, in patients (pts) with advanced cancer (CA). In: *J Clin Oncol*. 2008; 26(Suppl 15):3007.

113. Blank C, Mackensen A. Contribution of the PD-L1/PD-1 pathway to T-cell exhaustion: an update on implications for chronic infections and tumor evasion. *Cancer Immunol Immunother*. 2007;56(5):739–745.

114. Ghiringhelli F, Larmonier N, Schmitt E, et al. CD40 blockade depletes and selects for CD4+ regulatory T cells suppressing tumor immunity but are sensitive to cyclophosphamide which allows immunotherapy of established tumors to be curative. *Cancer Res*. 2012;72 Suppl 2:ii100–ii103.

115. Rech AJ, Vonderheide RH. Clinical use of anti-CD25 antibody therapy target regulatory T cells. *Nat Rev Clin Oncol*. 2013;10(1):ii10–ii13.

116. Vermeij R, Leffers H, Hoogeboom BN, et al. Potentiation of a p53-SLP vaccine by cyclophosphamide in ovarian cancer: a single-arm phase II study. *Int J Cancer*. 2012;131(5):E670–E680.
138. Selby MJ, Engelhardt JJ, Quigley M, et al. Anti-CTLA-4 antibodies of IgG2a isotype enhance antitumor through reduction of intratumoral regulatory T cells. Cancer Immunol Res. 2013;1(1):32–42.

139. Knutson KL, Wagner W, Disis ML. Adoptive T cell therapy of solid cancers. Cancer Immunol Immunother. 2006;55(6):96–103.

140. Jenq RR, Van den Brink MR. Allogeneic haematopoietic stem cell transplantation: individualized stem cell and immune therapy of cancer. Nat Rev Cancer. 2010;10(3):213–221.

141. Goff SL, Smith FO, Klapper JA, et al. Tumor infiltrating lymphocyte therapy for metastatic melanoma: analysis of tumors resected for TIL. J Immunother. 2010;33(8):840–847.

142. Kershaw MH, Westwood JA, Darcy PK. Gene-engineered T cells for cancer therapy. Nat Rev Cancer. 2013;13(8):525–541.

143. Billingham R. Dendritic cells. J Anat. 1948;82(1 Pt 2):93–109.

144. Steinman RM, Cohn ZA. Identification of a novel cell type in peripheral lymphoid organs of mice. I. Morphology, quantitation, tissue distribution. J Exp Med. 1973;137(5):1142–1162.

145. Banchereau J, Briere F, Caux C, et al. Immunobiology of dendritic cells. Annu Rev Immunol. 2000;18:767–811.

146. Hart DN. Dendritic cells: unique leukocyte populations which control immune responses. J Immunol. 1997;159(7):3133–3137.

147. Liu YJ. Dendritic cell subsets and lineages, and their functions in innate and adaptive immunity. Cell. 2001;106(3):259–262.

148. Banchereau J, Palucka AK. Dendritic cells as therapeutic vaccines against cancer. Nat Rev Immunol. 2005;5(4):296–306.

149. Harada Y, Yonemitsu Y. Recent developments in patented DC-based immunotherapy for various malignancies. Recent Pat Regen Med. 2011;1(1):72–87.

150. Palucka K, Banchereau J. Cancer immunotherapy via dendritic cells. Nat Rev Cancer. 2012;12(4):265–277.

151. Graggis TJ, McMahan CJ, Tamman J, Page KJ, Trager JB. Prostatic acid phosphatase expression in human tissues. Int J Clin Exp Pathol. 2011;4(3):295–306.

152. Pong L, Ruegg CL, Brockstedt D, Engleman EG, Laus R. Induction of tissue-specific autoimmune prostates with prostatic acid phosphatase immunization: implications for immunotherapy of prostate cancer. J Immunol. 1997;159(7):3133–3137.

153. Thakur A, Vaishampayan U, Lumb LG. Immunotherapy and immune evasion in prostate cancer. Cancers (Basel). 2013;5(2):569–590.

154. Okada H, Kalinski P, Ueda R, et al. Induction of CD8+ T-cell responses against novel glialoma-associated antigen peptides and clinical activity by vaccinations with [alpha]-type I polarized dendritic cells and polyinosinic-polycytidylic acid stabilized by lysine and carboxymethylcellulose in patients with recurrent malignant glioma. J Clin Oncol. 2011;29(33):330–336.

155. Lee JJ, Foon KA, Mailliard RB, Muthuswamy R, Kalinski P. Type I-polarized dendritic cells loaded with autologous tumor are a potent immunogen against chronic lymphocytic leukemia. J Leukoc Biol. 2008;84(1):319–325.

156. Mailliard RB, Wankowicz-Kalinska A, Cai Q, et al. [alpha]-Type I polarized dendritic cells: a novel immunization tool with optimized CTL-inducing activity. Cancer Res. 2004;64(17):5934–5937.

157. Zhu X, Falletti-Juneczo BA, Fujita M, et al. Poly-ICLC promotes the infiltration of effector T cells into intracranial gliomas via induction of CXCL10 in IFN-α and IFN-γ-dependent manners. Cancer Immunol Immunother. 2010;59(9):1401–1409.

158. Zhu X, Nishimura F, Sasaki K, et al. Toll like receptor-3 ligand poly-ICLC promotes the efficacy of peripheral vaccinations with tumor antigen-derived peptide epitopes in murine CNS tumor models. J Transl Med. 2007;5(1):10.

159. Eyrich M, Rachor J, Schreiber SC, Wolff M, Schlegel PG. Dendritic cell vaccination in pediatric gliomas: lessons learnt and future perspectives. Front Pediatr. 2013;1:12.

160. Fong L, Brockstedt D, Benike C, Wu L, Engleman EG. Dendritic cells injected via different routes induce immunity in cancer patients. J Immunol. 2001;166(6):4254–4259.
182. Knudtzon S. In vitro growth of granulocytic colonies from circulating cells in human cord blood. Blood. 1974;43(3):357–361.
183. Wright DE, Bowman EP, Wagers AJ, Butcher EC, Weissman IL. Hematopoietic stem cells are uniquely selective in their migratory response to chemokines. J Exp Med. 2002;195(9):1145–1154.
184. Berenson R, Andrews R, Bengsner W, et al. Antigen CD34+ marrow cells engraft lethally irradiated baboons. J Clin Invest. 1998;81(3):951–955.
185. DiPersio JF, Stadmayer EA, Nademanee A, et al. Plerixafor and G-CSF versus placebo and G-CSF to mobilize hematopoietic stem cells for autologous stem cell transplantation in patients with multiple myeloma. Blood. 2009;113(23):5720–5726.
186. Rosenblatt J, Avivi I, Vasir B, et al. Vaccination with dendritic cell/tumor fusions following autologous stem cell transplant induces immunologic and clinical responses in multiple myeloma patients. Clin Cancer Res. 2013;19(13):3640–3648.
187. Dudley ME, Gross CA, Somerville RP, et al. Randomized selection design trial evaluating CD8+–enriched versus unselected tumor-infiltrating lymphocytes for adoptive cell therapy for patients with melanoma. J Clin Oncol. 2013;31(17):2152–2159.
188. Restifo NP, Dudley ME, Rosenberg SA. Adoptive immunotherapy for cancer: harnessing the T cell response. Nat Rev Immunol. 2012;12(4):269–281.
189. Morgan RA, Dudley ME, Wunderlich JR, et al. Cancer regression in patients after transfer of genetically engineered lymphocytes. Science. 2006;314(5796):126–129.
190. Robbins PF, Morgan RA, Feldman SA, et al. Tumor regression in patients with metastatic synovial cell sarcoma and melanoma using genetically engineered lymphocytes reactive with NY-ESO-1. J Clin Oncol. 2011;29(7):917–924.
191. Rosenberg SA, Restifo NP, Yang JC, Morgan RA, Dudley ME. Adoptive cell transfer: a clinical path to effective cancer immunotherapy. Nat Rev Cancer. 2008;8(4):299–308.
192. Curran KJ, Pegram HJ, Brentjens RJ. Chimeric antigen receptors for T cell immunotherapy: current understanding and future directions. J Gene Med. 2012;14(6):405–415.
193. Grupp SA, Kalos M, Barrett D, et al. Chimeric antigen receptor-modified T cells for acute lymphoid leukemia. N Engl J Med. 2013;368(16):1509–1518.
194. Han EQ, Li XL, Wang CR, Li TF, Han SY. Chimeric antigen receptor-engineered T cells for cancer immunotherapy: progress and challenges. J Hematol Oncol. 2013;6:47.
195. Porter DL, Levine BL, Kalos M, Bagd A, June CH. Chimeric antigen receptor-modified T cells in chronic lymphoid leukemia. N Engl J Med. 2011;365(8):725–733.
196. Cancer Genome Atlas Research Network. Comprehensive molecular profiling of lung adenocarcinoma. Nature. 2014;511(7511):543–550.