Improving the clinical impact of biomaterials in cancer immunotherapy

Joshua M. Gammon¹, Neil M. Dold¹, Christopher M. Jewell¹,²,³

¹Fischell Department of Bioengineering, University of Maryland, College Park, MD, USA
²Department of Microbiology and Immunology, University of Maryland Medical School, Baltimore, MD, USA
³Marlene and Stewart Greenebaum Cancer Center, Baltimore, MD, USA

Correspondence to: Christopher M. Jewell, e-mail: cmjewell@umd.edu
Keywords: cancer immunotherapy, nanotechnology, biomaterials, cancer vaccine, translational medicine, Immunology and Microbiology Section, Immune response, Immunity
Received: November 23, 2015  Accepted: January 29, 2016  Published: February 10, 2016

ABSTRACT

Immunotherapies for cancer have progressed enormously over the past few decades, and hold great promise for the future. The successes of these therapies, with some patients showing durable and complete remission, demonstrate the power of harnessing the immune system to eradicate tumors. However, the effectiveness of current immunotherapies is limited by hurdles ranging from immunosuppressive strategies employed by tumors, to inadequate specificity of existing therapies, to heterogeneity of disease. Further, the vast majority of approved immunotherapies employ systemic delivery of immunomodulators or cells that make addressing some of these challenges more difficult. Natural and synthetic biomaterials – such as biocompatible polymers, self-assembled lipid particles, and implantable biodegradable devices – offer unique potential to address these hurdles by harnessing the benefits of therapeutic targeting, tissue engineering, co-delivery, controlled release, and sensing. However, despite the enormous investment in new materials and nanotechnology, translation of these ideas to the clinic is still an uncommon outcome. Here we review the major challenges facing immunotherapies and discuss how the newest biomaterials and nanotechnologies could help overcome these challenges to create new clinical options for patients.

INTRODUCTION

Over the past 50 years cancer diagnosis, prognosis, and treatment have progressed immensely, improving the outcomes and quality of life for millions of patients each year. Many persistent challenges, however, continue to limit our ability to stop cancer upon detection and to prevent relapse following successful treatment. The barriers undermining these broad goals are diverse, spanning drug toxicity and poor selectivity [1], the immunosuppressive nature of the tumor microenvironment [2], and the heterogeneity of disease across cancers and patient populations [3]. One of the most promising means to address these problems is immunotherapy. This field offers the possibility of treatments that are potent, specific, systemic, and durable – major challenges for invasive surgical resection, and non-specific chemotherapeutics or radiation that frequently result in relapse. In the past decade, exciting progress in immunotherapy – for example, with adoptive therapy [4] and monoclonal antibodies [5] – has stimulated new ideas for harnessing the immune system to combat cancer, while rejuvenating interest in other strategies such as cancer vaccination [3, 6, 7]. However, despite increasing investment in biomedical research, the efficiency with which new treatments are delivered is stagnant [8]. For example, since 1965 the rate of new molecular entities (NME) and new drug approvals (NDAs) by the FDA has remained essentially constant, while the publishing of biomedical manuscripts has increased more than 500% [8]. In cancer immunotherapy, some of the challenges facing new progress include generating immune cells with the correct specificity and function to both recognize and attack cancer cells, maintaining the function of these populations over time, and designing combination therapies that are synergistic without compromising...
safety [3]. Multidisciplinary approaches focused on clinical challenges in many biomedical research fields, including cancer immunotherapy, could be improved in terms of clinical impact and translatability by integrating capabilities provided by new technology and engineering. In this review we use recent clinical successes and failures to highlight opportunities to leverage nanotechnology and biomaterials to push cancer immunotherapy forward (Table 1).

Overview of biomaterials: Classes and attractive properties

Biomaterials are ubiquitous in biomedical research, and have had some notable impacts in cancer therapy over the past few decades. Thus far, most of these advances – at least clinically – have involved improving the solubility, reducing the toxicity, or increasing the half-life of small molecule chemotherapeutics such as doxorubicin. These improvements illustrate a few of the properties that make biomaterials of great interest for cancer immunotherapy. Speaking generally, “biomaterial” is a term that spans natural or synthetic polymers, lipids, metal contrast agents, engineered cells, quantum dots, and a multitude of self-assembled structures. These materials are often used to build implantable scaffolds or devices [9, 10], as sensitive biosensors on functionalized surfaces within microfluidic devices [11, 12], or to formulate nanoparticles (NPs) or microparticles (MPs) that can be delivered or conjugated to cells ex vivo or in vivo [13]. One classic advantage of biomaterials is the co-delivery of cargo by encapsulating two or more cargos (e.g., small molecule drugs) in a biodegradable polymeric particle. This approach is frequently used to ensure cells or tissues receive each cargo type to work in synergy, or – by synthesizing polymers with an appropriate degradation rate – to achieve a desired sustained release profile. In addition to co-delivery and controlled release, many particle-based strategies are aimed at improved targeting by surface conjugation of antibodies or ligands for receptors expressed on target cells or tissues.

Another important focus of biomaterials has been in protecting biologic cargo from degradation in the presence of enzymes or extreme pH, and to reduce systemic toxicity by allowing drug to be slowly released over time or upon reaching target tissues such as tumors. This has been particularly important in cancer, where increasing the circulation time of drugs through modification with polyethylene glycol or other molecules has led to better tumor targeting; targeting occurs because of the leaky tumor vascular that causes preferential accumulation at tumors through the enhanced-permeability and retention (EPR) effect [14]. In this last area, liposomes, multi-lamellar vesicles, exosomes, and other lipid-based nanostructures have been particularly useful owing to the highly biocompatible nature of this class of biomaterials [15]. A more recent area of interest is also arising: the intrinsic immunogenic properties of some biomaterials. Many studies demonstrate that common polymers such as poly(lactide-co-glycolide) (PLGA) and poly(styrene) activate pro-inflammatory pathways (e.g., inflammasome) [16–18]. These characteristics, with better understanding, could be exploited to design polymers that serve not only as carriers, but also as agents that help polarize immunity. From another perspective, these materials can complicate rationale design of vaccines and immunotherapies because the carrier itself can alter the response to other vaccine components. Several new strategies are exploring self-assembly of proteins or immune signals to co-deliver vaccine and immunotherapy components [19–21]. In one of these approaches, electrostatic assembly is used to assemble antigens and adjuvants without synthetic polymers or other carrier components [21]. These immune polyelectrolyte multilayer (“iPEMs”) structures thus mimic favorable properties of biomaterial carriers (e.g., tunable sizes, co-delivery) while creating a well-controlled platform for assembling multiple immune signals at high densities without the complicating intrinsic immune effects of many polymers [20, 21].

Disparity between clinical impact and the investment in nanotechnology and biomaterials

Despite the exciting potential of biomaterials, the impact on the clinic has been modest relative to the pre-clinical investment that has been made. For example, by one recent estimate, 3% of the roughly three million paper published in the cancer field are associated with clinical trials [22]. In contrast, cancer papers also involving polymeric materials only connect to clinical trials of some form in 1% of studies. On the other hand, categories such as liposomes and monoclonal antibodies – both much more clinically-established relative to polymeric materials – are ultimately associated with clinical trials in 4% and 8% of cancer papers, respectively [22]. Though this is a single measure, one interpretation is that as more biomaterial-based strategies advance to the clinic, the potential of other material strategies will receive increasing attention. At present, however, there is a significant gap between clinical impact and the investment in basic and pre-clinical nanotechnology research [22–25]. Below we discuss seven areas where challenges or recent progress in the clinic suggests specific opportunities for biomaterials and nanotechnology to make a significant impact. These include i) toxicities associated with current immunotherapies, ii) reprogramming or polarizing anti-tumor T cells in vivo or after adoptive cell therapy, iii) targeting more specific immune pathways, iv) cancer vaccination, v) combination immunotherapies, vi) enhanced homing to and activity of immune cells in the tumor microenvironment, and vii) heterogeneity of disease.
CURRENT CLINICAL CANCER CHALLENGES CREATE OPPORTUNITIES TO HARNESS BIOMATERIALS

Reduce toxicity and enhance pharmacokinetics (PK) or pharmacodynamics (PD)

Similar to chemotherapy and radiation, the potency of cancer immunotherapy can be accompanied by toxicity severe enough to warrant discontinuation. These effects are challenging to address because most of the existing immunotherapies involve large doses (e.g., IL-2 and interferon alpha) delivered systemically to overcome poor PKs or PDs [1]. Thus, many immunotherapies continue to be limited by poor therapeutic targeting, bolus concentrations, and non-specific side effects. The clinical toxicities unique to immunotherapies – including vaccines, cytokines, checkpoint blockades, and cell therapies – have been recently reviewed by Weber et al. [1]. One of the hallmarks of immunotherapy-related adverse events is the induction of unrestrained autoimmune responses toward healthy tissue due to overactive effector cells or due to targeting of tumor-associated antigens (TAAs) that are expressed in both tumors and healthy tissues [1, 26].

Biomaterials can be used to encapsulate and protect therapeutic moieties from biodegradation, to passively

| Clinical challenge/opportunity | Biomaterial strategies | Key references |
|-------------------------------|------------------------|---------------|
| Toxicity                      | Localizing and extending release of immunostimulants in materials at safer doses | Kwong 201329, de Titta 201332 |
| Polarizing adoptively transferred T cells | Ex vivo T cell expansion with artificial antigen presenting cells, Nanoparticle conjugation to T cells, Implantaing T cell seeded scaffolds | Perica 201543, Stephan 201045, 201246, Stephan 201551 |
| Polarizing T cells: cell free therapies | Pathogen mimicking microparticles | Pradhan 201452, Singh 201153 |
| Targeting specific pathways | Delivering siRNA with polymer carriers, Using physiological phenomena to localize therapy in lymph nodes | Alshamsan 201065, 201164; Wang 201331, Liu 201439, Hanson 201534 |
| Revisiting cancer vaccination | Active targeting of nanoparticle vaccines to DCs, Localized delivery of immune signals with Injectable scaffolds, Combining Nanoparticle vaccines with siRNA knockdown of immunosuppressive cytokines | Rosalia 2015102, Ali 2009109, Bencherif 2015110 |
| Combination immunotherapies | Controlled combinatorial delivery of adjuvants | Goldinger 2012120, Thomas 2014121, Roy 2013131, Marrache 2012135 |
| Increasing homing and activity of immune cells in tumor microenvironment | Nanoparticle conjugation to T cells, Sequestering of immunostimulants in tumors, Nanogel co-delivery of IL-2 and TGF-β inhibitor | Huang 2015138, Liu 2011140, Intra 201138, Park 2012143 |
| Addressing tumor heterogeneity | Microparticle or nanoparticle tumor lysate vaccines, Capture of circulating tumor cells for identification of neoantigens or tumor cell phenotype | Prassad 2010154, Gross 2014155, Halo 2014162, Azarin 2015137 |
or actively improve targeting, and to sustain the release of encapsulated cargo. Each of these features can be harnessed to improve PKs or PDs of immunotherapies and to reduce the associated toxicities. Consequently, it is not surprising that the type of biomaterial that has been most successful in clinical cancer therapy is the use of nanoparticles – liposomes, in particular – for therapeutic delivery of toxic chemotherapeutic drugs [27].

In immunotherapy specifically, clinical testing of liposomal or nanoparticle vehicles for altering toxicity, PK, and PD is under-explored, but a number of pre-clinical investigations demonstrate the exciting potential of these ideas [28–31]. Kwong et al. formulated two liposomal therapies that overcome typical toxicity-limited dosing. The first uses liposomes to co-deliver anti-CD40 antibody and CpG oligonucleotide – an immune-stimulatory toll-like receptor agonist (TLRα) – to tumors in a subcutaneous melanoma model (B16-F10) [30]. Compared to soluble delivery, the liposome-coupled therapy resulted in preferential retention of the liposomes and their encapsulated cargos in tumors and the draining lymph nodes (LNs). Notably, the circulating levels of both CpG and anti-CD40 antibody in serum were reduced, leading to significant reduction of pro-inflammatory cytokines such as IL-6 and TNF-α. Alanine aminotransferase (ALT) enzyme – a marker for hepatic damage – was also much lower during liposomal therapies compared with soluble therapies. In a second study, two stimulatory agonists – IL-2/Fc fusion protein and anti-CD137 antibody – were anchored on liposomes, and this therapy was tested against soluble anti-CD137 antibody and IL-2/Fc doses during intraperitoneal (i.p.) and intratumoral (i.t.) injections [29].

Although these liposomal formulations elicited similar proliferation of T cells in vitro in comparison to soluble treatments, the liposomal forms abrogated toxic levels of cytokines in serum in vivo. Specifically, mice receiving liposome-conjugated therapies i.t. exhibited serum cytokine levels (IFN-γ, IL-6, CCL2, and TNF-α) similar to untreated mice without weight loss, whereas mice treated with matched soluble doses had profound increases in serum cytokine levels that caused shivering, >5% weight loss, and some treatment-related fatalities. Both soluble and liposomal formulations were able to induce complete tumor regression of 60-70% of B16-F10 tumors treated i.t. in mice; however, only the liposomal formulations allowed for delivery of larger doses that would have elicited systemic toxicities in soluble form because the liposomes forced sequestration of the therapies in tumors and tumor-draining LNs.

Similarly, several recent investigations have used non-liposomal materials to minimize toxicity by targeting or retaining immunotherapeutic agents inside tissues vital to disease reduction such as tumors and LNs. These approaches have included NPs and MPs [32–35], self-assembling injectable scaffolds [36, 37], and other materials designed to locally deliver, sequester, and/or preferentially trap cargo in tissues of interest [38–40]. In one study, Hubbell, Swartz, and colleagues reported mixed populations of poly(propylene sulfide) NPs conjugated individually to either a model tumor antigen (ovalbumin, OVA) or CpG-based adjuvants. NP conjugation facilitated co-localization of antigen and adjuvant via trafficking of both NP-OVA and NP-CpGs to lymph nodes following intradermal (i.d.) injection in mice through size-dependent lymphatic draining patterns [32]. Analysis of systemic cytokine levels showed that the dose of CpG used in both NP and free form was not capable of inducing high systemic cytokine concentrations, indicating a low risk of toxicity (Figure 1A, 1B). In a prophylactic model, low dose NP-CpGs enhanced antigen-specific memory recall of cytotoxic T lymphocytes (CTLs) more than low-dose soluble treatments, resulting in greater CTL expansion and cytokine secretion compared with soluble therapies. Additionally, NP-CpGs delivered alongside NP-OVA conferred complete protection against EG.7-OVA lymphoma and significant protection against B16-F10-OVA melanoma tumors, whereas tumor protection was significantly diminished for B16-F10-OVA tumors when the same low-dose CpG formulations were given as free molecules (Figure 1C). Although this approach was tested in tumor models expressing OVA, these data demonstrate the value of localizing multiple therapeutic components in lymphatic tissues and the power of using a biomaterial delivery system to passively target lymphatics through reduced adjuvant doses that limit toxicity while maintaining efficacy. These preclinical examples also suggest new potential for cancer immunotherapies that have been previously considered infeasible strategies. For example, treatments that clinicians might typically avoid or have previously discontinued due to toxicity might be reexamined and enhanced with biomaterials through encapsulation, controlled release, and targeting. Similarly, as discussed later, therapies like cancer vaccination that have not previously been potent enough for efficacy, may be revitalized through better materials and new nanotechnologies.

Reprogramming and polarizing T cells

Adaptive cell therapy

Although recent years have brought significant clinical successes using adoptive cell therapy (ACT), – an approach in which patient cells are isolated, expanded, and re-infused – some of the continuing challenges include rapid death of transferred cells, poor migration to tumors, and deactivation of cells entering the tumor microenvironment [4, 41]. Current protocols effectively expand and activate T cells ex vivo. However, this approach introduces practical challenges that reduce accessibility of treatment owing to the cost and complexity of the protocols. Another challenge facing the field is the limited control over maintaining T cell expansion and polarization after adoptive transfer while minimizing
toxicity and off-target effects of modulatory cues (e.g., drugs, cytokines) administered systemically during these treatments.

There are several routes by which arising nanotechnology might help address these challenges: 1) enhancing and simplifying T cell enrichment and expansion procedures, or 2) providing the ability to locally or selectively modulate T cells after transfer. In one recent example, Fadel et al. utilized high surface area carbon nanotube-polymer composites to improve ex vivo expansion of OT-1 CD8+ T cells. Adoptive transfer of these T cells delayed tumor growth in mice bearing B16-OVA melanoma tumors [42]. Another strategy, recently reported by the Schneck group, simplified ex vivo T cell isolation and expansion to support ACT using artificial antigen presenting cells (aAPCs) [43]. These aAPCs consist of magnetic iron-dextran NPs conjugated to co-stimulatory CD28-activating antibodies and MHC I molecules bound to TAAs. TAA-specific CD8+ T cells are enriched by magnetic isolation after incubation of CD8+ peripheral blood mononuclear cells (PBMCs) with aAPCs. The TAA-specific T cells bound to the aAPCs can then be collected and further cultured, while the co-stimulation and antigen presentation provided by the aAPCs drives expansion of the cells prior to adoptive transfer to mice. In current methods T cells isolated from PBMCs are cultured with DCs pulsed with peptide tumor antigen after isolation from peripheral blood of patients [44]. Thus the aAPC method may offer a simpler alternative for T cell expansion, eliminating the need for patient specific DCs while also increasing the number of TAA-specific T cells. An additional benefit of the aAPC platform is a reduction in the collection time, where tumor-specific CD8+ T cells can be expanded over 1000-fold in as little as one week. The modularity of this design was demonstrated through efficient expansion of T cells specific to a variety of well characterized tumor antigens, including mouse TAAs (Trp2, gp100), human TAAs (NY-ESO1, MART1), and mutated neoantigens from mouse melanoma (B16) or colon carcinoma (CT26). The anti-tumor efficacy of ACT with T cells expanded using this method was tested in mouse models of melanoma. CD8+ T cells specific for Trp2 and gp100 were isolated and expanded with aAPCs. Adoptive transfer of these T cells 8 days after inoculation with B16-F10 tumors significantly enhanced survival compared to mice injected with equivalent numbers of aAPC expanded T cells specific for the model antigen, SIINFEKL. Transfer of aAPC-expanded Trp2 and gp100 T cells caused complete tumor rejection in 25% of mice at 30 days after implantation, whereas all mice treated with SIINFEKL T cells reached terminal endpoints by 22 days post challenge. Building on these and other recent approaches, an important next step is to expand cells from patient samples.

Recent work by the Irvine group has demonstrated a powerful pre-clinical strategy to modulate T cell function in mice by synthesizing NPs containing immunomodulatory signals and conjugating these NPs to the membranes of T cells prior to transfer [45–47]. After adoptive transfer, conjugated NPs provide sustained, local interactions with CD8+ T cells, allowing autocrine delivery of the drugs from the NPs. The particles in these studies are multi-lamellar vesicles conjugated to T cells through maleimide chemistry. This approach leverages disulfide bonds which form between maleimide functionalized NPs and the naturally occurring excess thiols displayed on T cell membranes. In one report, this platform was used to modify CD8+ PMEL T cells with NPs co-encapsulating IL-15 superagonist and IL-21 [45]. T cells from these mice exhibit a transgenic T cell receptor specific for the gp100 tumor epitope. Mice receiving the modified cells exhibited dramatic polarization of anti-tumor T cells toward central memory phenotypes, greatly increasing T cell expansion and persistence compared to transfer coupled with soluble, systemic therapies or unconjugated NPs [45]. The functional impact of this therapy was complete clearance of tumors in all mice bearing metastatic B16-F10 tumors after infusion of NP-conjugated T cells, whereas mice receiving T cells augmented with an identical dose of systemically delivered cytokines all succumbed to the

Figure 1: CpG variants formulated in NPs prevent tumor growth better than free CpG without increasing systemic inflammation. Low doses of NP-CpG-B do not significantly increase peak levels of inflammation as measured by serum levels of A. TNF-α and B. IL-6. C. NP-CpG-B and NP-CpG-C protects mice from B16-F10-OVA tumor establishment better than unconjugated counterparts. Adapted with permission from [32].
tumors. A subsequent study demonstrated that T-cell-conjugated NPs are sequestered to the immune synapse, providing a direct route to control T cell function or target tumor cells during binding by CD8+ T cells [46]. This potential was demonstrated by conjugating cells with NPs containing inhibitors of Shp1 and Shp2 – phosphatases in the immune synapse that downregulate T cell receptor activation [48] – to improve accumulation of T cells in murine prostate tumors and prolong survival of transferred cells. In these studies the NP-conjugated T cells exhibited a 5.7-fold increase in tumor accumulation four days after transfer. Zheng et al. recently reported an alternative approach using liposomes targeted to adoptively-transferred T cells in vivo through antibody fragments specific for Thy1.1 or IL-2 receptor ligand [47]. This strategy expands on ex vivo NP conjugation techniques by allowing NPs to be conjugated to proliferating T cells in vivo after transfer.

The studies above highlight some important opportunities to advance ACT by specifically targeting T cells with continued, localized delivery of immunomodulators at low doses. This ability could reduce toxicities and potential autoimmune reactions associated with repeated soluble administration of high doses of immunostimulatory cues (e.g., cytokines, monoclonal antibodies). These properties are increasingly attractive considering the recent successes of a variety of checkpoint blockade therapies targeting CTLA-4 or PD-1 signaling in the immune synapse [49, 50]. Since one current limitation of checkpoint blockade is the effect on bystander T cells [49], local T cell targeting strategies might also offer further opportunities to enhance the specificity of these combination therapies. An alternative strategy to modifying T cells ex vivo is implantation of biomaterial scaffolds seeded with T cells to provide a reservoir that sustains proliferation and activation after implantation. In a recent demonstration of this idea by Stephan et al., T cell-harboring scaffolds encapsulating immune signals were implanted locally near murine breast or ovarian tumors to overcome T cell death, exhaustion, and poor migration to tumors after adoptive transfer – each significant challenges encountered in the clinic [51]. These devices are prepared from alginate scaffolds coated with collagen receptor ligands to support T cell seeding. Additionally, porous silica MPs incorporating IL-15 superagonist and agonistic antibodies for CD28, CD137 and CD3 (Figure 2A) are embedded in the scaffold. These MPs improve T cell proliferation and survival after migration from the scaffold, as demonstrated in an in vitro assay measuring the ability of T cells to migrate from the scaffold into 3D tissue mimetic collagen gel loaded with an inflammatory cytokine (IP-10). Implantation of T cell-loaded scaffolds at murine 4T1 breast tumor resection sites greatly enhanced survival (Figure 2B), with all mice treated with the T cell harboring scaffolds surviving and remaining relapse-free throughout the study. In contrast, T cells not loaded in a scaffold but pre-stimulated with IL-15, anti-CD3, anti-CD28 and anti-CD137 only increased the median survival by 5 days relative to untreated mice (Figure 2B). Following local injection at resection sites, these cells persisted transiently (Figure 2C) and became functionally exhausted. T cells injected within scaffolds, in contrast, showed drastically increased proliferation at tumor sites and retained functional, non-exhausted phenotypes characterized by low levels of Tim-3, and 2B4 – receptors which inhibit CTL activation and proliferation.

**Cell free therapies**

Immunotherapies aimed at in vivo generation of anti-tumor response offer the potential for simpler, more accessible therapies compared with adoptive transfer. However, the efficacy of these approaches has been limited by the need for not just robust anti-tumor response, but control over the phenotypes of the expanded immune cell populations. Biomaterials are being exploited to achieve this goal by offering improved control over the targeting, release, and co-delivery of immunomodulatory molecules compared with soluble systemic dosing. The Roy group recently used polymeric MPs to polarize T cell differentiation in vivo by directing the function of DCs [52, 53]. These studies sought to alter the balance between CTL-inducing T_h1 responses and suppressive T_reg or T_h2 responses by altering the cytokines secreted by DCs during antigen presentation to CD4+ T cells. CpG has been largely investigated along these lines during clinical trials due to its ability to induce secretion of IL-12 and other T_h1-polarizing cytokines. However, CpG also induces IL-10 which can suppress subsequent T_h1 generations, instead promoting T_h2 and T_reg phenotypes. Pradhan et al. thus co-delivered CpG and IL-10 siRNA with plasmid DNA encoding a B cell lymphoma antigen [52]. These signals adsorbed to cationic pathogen mimicking particles (PMPs) built from PLGA and polyethylenimine (PEI) cores. Treatment of DCs with PMPs resulted in an increased ratio of IL-12 to IL-10, a higher frequency of T_h1 cells, and inhibition of IL-4 – a cytokine associated with T_h2 induction. Implantation of PMPs in a dextran-PEG hydrogel containing MIP-3α – a chemokine which recruits DCs – resulted in enhanced survival in a murine model of B cell lymphoma [53]. Another recent strategy in this area described preferential promotion of a T_h1 response using chitosan as a carrier for intravesical delivery of IL-12 to promote T_h1 cytokine profiles, increased survival, and reduced relapse in the MB49 murine bladder cancer model [54, 55]. These results demonstrate the power of using biomaterials to co-deliver immunomodulators for polarizing T cell phenotype to enhance anti-tumor immunity.

As demonstrated by many studies, the CD8+ T cell compartment plays a crucial role in eradicating tumors, and in generating long-lived anti-tumor memory
populations that can help prevent relapse. CTLs with minimally differentiated phenotypes, including central memory \( (T_{CM}) \) or T memory stem cells \( (T_{SCM}) \) demonstrate enhanced proliferative capacity that supports rapid generation of high numbers of anti-tumor effector T cells [56]. However as T cells proliferate, they lose plasticity and differentiate into terminal phenotypes that eventually become functionally exhausted and exhibit reduced anti-tumor activity [57]. Nanotechnology offers the potential to not only generate large populations of anti-tumor CTLs, but to modulate their phenotype to maintain \( T_{CM} \) or \( T_{SCM} \) to maintain plasticity. For example, recent studies demonstrate sustained release of antigens from particles promotes memory phenotypes compared with the transient antigen presentation characteristic of more traditional vaccines [58]. New strategies that target or sustain the delivery of immunomodulators to regulate differentiation pathways (e.g., mTOR [59]) or induce cytokines that polarize T cells toward \( T_{CM} \) (e.g., IL-15, IL-21 [60]) may be a promising strategy to improve current cancer immunotherapies.

**Targeting more specific immune pathways**

While clinical immunology is targeting an array of signaling pathways to enhance anti-tumor immunity and suppress tumor-evasion mechanisms, the bulk of biomaterial-based strategies have focused primarily on materials as carriers of established cargos to, for example, sustain delivery. Further, many recent breakthroughs in checkpoint blockade and other areas demonstrate that pathways beyond traditional lymphocyte expansion are important in the efficacy of immunotherapy. Thus, there is substantial opportunity for biomaterials to be harnessed in studying and targeting specific immune pathways. As discussed further in the conclusion, this is a challenging undertaking that requires cross-disciplinary collaboration across engineering and immunology, but these approaches could be transformative.

The potential to harness biomaterials in targeting more sophisticated immune pathways is highlighted by several recent research directions. One pathway of interest is signal transducer and activator of transcription

![Figure 2: Implantable scaffolds enhance ACT. A. BMPs encapsulating T cell stimulants are embedded in the implantable scaffold. B. Scaffold delivered T cells prevent relapse after tumor resection and greatly enhance survival. C. Locally administered T cells within scaffolds persist and proliferate at the tumor site compared to prestimulated T cells administered at the resection site in the absence of a scaffold. Adapted with permission from Macmillan Publishers Ltd: Nature Biotechnology [51] copyright (2015).]
3 (STAT3), a transcription factor which in addition to regulating tumorigenic and metastatic properties [61], is involved in regulating pro-immune function of tumor cells [62]. In particular for cancer immunotherapy, STAT3 is overexpressed in many tumors and is associated with regulatory pathways that include inhibition of DC maturation and inflammatory cytokine production [63]. Thus, siRNA knockdown to reduce STAT3 expression and the associated tolerance of DCs toward tumor antigens has recently been investigated using PEI modified with stearic acid (StA) lipids to form electrostatic siRNA/PEI-StA complexes [64, 65]. These complexes protect the siRNA from degradation and are readily internalized by tumor cells. When soluble or complexed treatments were administered i.t. following s.c. inoculation of B16-F10 tumors in mice, the PEI-StA complexes mediated greater anti-tumor effects. Specifically, these particles reduced both the total concentration of STAT3 and the active, phosphorylated form, while also upregulating inflammatory cytokines and classical DC activation markers (CD40, CD86). These effects were also observed with increased tumor infiltration of DCs and CD3+ lymphocytes and significantly reduced tumor growth [64, 65]. The non-specific cytotoxicity of siRNA complexes was mitigated in additional experiments by encapsulating the PEI-StA/siRNA complexes in PLGA NPs [65]. Similar efficacy was also achievable using small-molecule STAT3 inhibitors encapsulated in PLGA NPs, demonstrating the robustness of biomaterials to deliver different classes of cargos to target specific pathways [66].

In the examples above, STAT3-inhibitors were administered via i.t. injection, which is only feasible in patients with solid tumors accessible for injection. Wang et al. recently reported a systemic knockdown therapy using siRNA loaded in liposome-protamine-hyaluronic acid NPs to target CD47, a protein found on the surface of cancer cells that prevents phagocytic destruction [31]. Intravenous administration of these particles in mice every two days beginning on day 8 after B16-F10 tumor inoculation resulted in efficient delivery and retention of siRNA in tumors due to the EPR effect. In these studies, treatment reduced CD47 mRNA five-fold and tumor growth by ~93% [31].

Beyond targeting cellular pathways co-opted by tumors, pathways that control physiological structures and immune cell trafficking to lymphoid tissues can also be harnessed in new strategies that emerge from immunology and materials science [16]. The benefits of targeting lymph nodes are apparent because lymphatic tissues are key sites where antigen presenting cells and lymphocytes interact to mount anti-tumor responses. One fascinating new strategy to shuttle vaccine cargos to lymph nodes exploits the natural trafficking of albumin from blood, across lymphatics, and into lymph nodes. In these studies by Liu et al., CpG and antigenic peptides were covalently modified with albumin-binding domains to allow “hitchhiking” of these immune signals to lymph nodes [39]. This vaccine method was highly potent and generated efficient immune responses in mice against a variety of antigens, including tumor antigens that caused regression of TC-1 tumors and slowed growth of B16-F10 tumors. Another recent strategy for targeting lymph nodes uses intra-lymph node injection to introduce degradable polymer depot vaccines to lymph nodes for local sustained release of adjuvants and other immune signals in these tissues [67, 68]. This approach generates potent antigen-specific responses in mice and could offer a direct route for locally engineering the lymph node microenvironment to control specific immune pathways [67].

Lastly, some very recent strategies simultaneously make use of physiological organization of immune cells and intracellular signaling pathways. Activation of the stimulator of interferon genes (STING) complex, for example, is now known as a major mechanism the innate immune system uses to sense tumor cells [69]. STING agonists like cyclic di-GMP (cdGMP, a bacterial cyclic dinucleotide) are internalized by immune cells after binding STING and are able to activate inflammatory transcription factors such as those that drive type I interferon production. Like many adjuvants, STING agonists may exhibit poor activity in soluble form, partly due to the fact that these molecules do not accumulate in a free form near immune cells in tissues like lymph nodes [67, 70]. In a recent study, subcutaneous administration of soluble cdGMP showed very low accumulation in lymph nodes, whereas nanoparticle formulations (NP-cdGMP) increased cdGMP accumulation ~15-fold and sustained its presence in lymph nodes using the same injection route (Figure 3A) [34]. To test functionality, mice were inoculated with EG.7-OVA tumors then vaccinated s.c. with either OVA or OVA combined with soluble or NP-formulated cdGMP. Vaccination with NP-cdGMP and OVA stimulated the highest frequencies of antigen-specific CTLs (Figure 3B), along with robust antibody production equivalent to the levels achieved with a 30-fold higher soluble dose. Moreover, NP-cdGMP therapeutic vaccination on days 6, 13, and 20 following EG-OVA tumor inoculation attenuated tumor growth (Figure 3C) and prolonged survival (Figure 3D) [34]. Each of the above examples demonstrates the utility of combining detailed immunological knowledge with the features of biomaterials, underscoring the need to cultivate effective cross-disciplinary collaborations and strategies in future studies.

Renewed potential of cancer vaccination

The success of several new immunotherapy strategies in the clinic have also revitalized the potential of cancer vaccination. One important example is targeting of the immune checkpoints, CTLA-4 and PD-1, which have been highly effective in recent trials [49, 50]. Ipilimumab,
an antibody targeting CTLA-4, was the first approved checkpoint blockade therapy, approved in 2010 for treatment of melanoma. In phase III trials conducted on patients with advanced melanoma, Ipilimumab conferred durable and long-term response in a subset of patients, with greater than 20% of patients surviving for over 4 years [50, 71]. Another example is the recent approval of sipuleucel-T (Provenge®). Sipuleucel-T is a vaccine consisting of autologous DCs pulsed with a fusion protein of human GM-CSF linked to a prostate cancer antigen (PAP) expressed in more than 90% of prostate tumors [72]. Phase I and II clinical trials demonstrated that sipuleucel-T is safe and can generate PAP-specific immune response [72]. In phase III clinical trials, patients treated with sipuleucel-T had enhanced survival compared to patients treated with a vaccine in which DCs were reinfused without being pulsed with the PAP fusion protein. [72, 73]. These benefits secured FDA approval of sipuleucel-T for treatment of minimally symptomatic or asymptomatic castrate-resistant prostate cancer in 2010, and a variety of clinical trials investigating sipuleucel-T are ongoing [74]. With these recent demonstrations of the renewed potential for therapeutic cancer vaccines, biomaterials are positioned to help overcome the persisting hurdles.

One of the fundamental challenges facing cancer vaccines in the clinic is poor immunogenicity of TAAs. Biomaterials can help combat this challenge by improving the likelihood that vaccine components reach DCs or other immune cells or tissues together. TLRas, for example, are being used to enhance TAA-specific immune response; however, systemic co-administration of soluble TLRas with TAAs does not ensure that both components reach DCs together. Additionally, soluble vaccine adjuvants are rapidly cleared in vivo, and may not persist long enough to strongly activate DCs-effects that could limit efficacy, or even promote tolerance against TAAs if DCs present the antigens after adjuvants are cleared [75]. Biomaterials have been utilized to address these limitations by co-delivering and sustaining the release of multiple classes of vaccine cargos (e.g., antigens, adjuvants). Recent approaches along these lines have exploited biomaterials to adsorb, self-assemble, or encapsulate whole tumor cells [76], tumor lysates [77–80], proteins [10, 81–83], minimal epitope peptides [84–89], DNA [90] and mRNA [91].

In addition to increasing immunogenicity of TAAs by providing co-delivery with immunostimulants, biomaterials have also been utilized to facilitate cross-presentation of antigens for the generation of CTL responses [92]. Antigens delivered with particulate carriers can enhance processing of antigen onto the MHC-I pathway compared to soluble delivery of exogenous antigen [93–95]. Additionally pH responsive carriers have been used to enhance this effect by efficiently delivering antigens into the cytosol of APCs. A variety of biomaterial carriers have been developed to release antigens within the acidic pH of endosomes/lysosomes [96–98], and some have been utilized to the promote endosomal escape of released antigens by causing lysis of endosomes [99–101].

Figure 3: STING agonists exhibit enhanced adjuvant properties and anti-tumor efficacy when administered in NPs rather than soluble form. A. Only NP-formulated cdGMP accumulates in lymph nodes. B. Following s.c. inoculation of EG.7-OVA tumors, NP-cdGMP elicits higher antigen-specific T cell frequencies (SIINFEKL) among CD8+ cells compared with soluble cdGMP. C. Tumor growth is attenuated by NP-cdGMP, but not soluble cdGMP, when administered therapeutically (arrows) after tumor inoculation. D. Mice receiving NP-cdGMP exhibit improved survival. Adapted from Figures 1 and 2 of [34] with permission.
Another key advantage of biomaterial vaccine delivery systems, in particular for MPs or NPs, is the ability to preferentially target DCs, which specialize in phagocytosing particulate matter such as cells, antigens, or vaccine particles. The targeting of NPs to DCs can actively be enhanced through the conjugation of antibodies or other specific ligands. Rosalia et al. recently demonstrated that NP cancer vaccines can be enhanced by targeting DCs using PLGA NPs conjugated to CD40 antibody [102]. These particles were used to co-encapsulate and deliver the model antigen OVA with Pam3CSK4 (TLR1/2) and Poly IC (TLR3). CD40-NPs effectively targeted and accumulated inside DCs in lymph nodes, driving increased antigen internalization and enhanced expression of DC activation markers. Therapeutic vaccination regimens with CD40-NPs also drove enhanced survival of mice implanted with B16-OVA tumors [102].

Beyond the use of biomaterials as passive vaccine delivery vehicles, a developing area in the biomaterials and vaccine fields is the intrinsic and self-adjuv Canting properties of polymers and other materials [16]. Some of the materials now being studied along these lines for cancer vaccination include cationic lipids [82, 83], or polymer micelles [103], dendrimers [87, 88], polyanhydrides [81], self-assembling peptides [84], and fibronectin [86]. These intrinsic immunostimulatory effects – activation of the inflammasome [17], for example – seem to result from the features that many biomaterials share with pathogens: a particular nature/form, repetitive polymeric structures that loosely resemble the repetitive nature of bacterial polysaccharides or other pathogen-associated polymers, and hydrophobic moieties or regions that activate pathogen/danger-associated molecular patterns (PAMPs/DAMPs) [104]. Intriguingly, recent studies by Andorko et al. have also revealed that the intrinsic immunogenicity of degradable polymeric carriers can change during degradation [18]. Understanding how the properties of materials exhibit or drive immunostimulatory pathways could improve DC targeting or activation to create a new generation of vehicles that serve not just as carriers but that also help actively direct immune response during cancer vaccination.

Several of the NP-based strategies discussed in previous sections are also relevant to improved cancer vaccination. The aAPCs discussed earlier for ex vivo expansion of anti-tumor T cells, for example, have also recently been investigated as vaccines to prime effective anti-tumor CTLs in vivo [105, 106]. Additionally, particulate vaccine delivery systems generally accumulate in lymphoid organs through passive drainage, which serves as a passive targeting system for uptake and processing of vaccine particles by lymph node-resident APCs [32]. This process also helps ensure vaccine components encapsulated in particles reach APCs together to maximize the enhancement to antigen processing and presentation, and the subsequent expansion of tumor-specific lymphocytes. As discussed above, there are also some other exciting new strategies for active targeting, including the work of Liu et al., in which albumin shuttling was exploited in mouse models of melanoma and cervical cancer [39].

Whole cancer cell based vaccines have been widely investigated in the clinic and have demonstrated the potential of localized co-delivery of tumor antigens and immunomodulators. These strategies aim to overcome the generally poor immunogenicity of cancer cells by modifying the cells with a variety of immune signals to amplify the response against cancer antigens [107]. A common example of this strategy is whole cancer cells engineered to express the immunostimulant, GM-CSF, which have been studied as a vaccine for a variety of cancers [108]. In these approaches, allogenic or autologous tumor cells are transduced to express high levels of GM-CSF in vitro and are then administered to patients. However these strategies have resulted in limited efficacy, likely due to cell death. This hurdle leads to poor persistence and co-localization of the signals and tumor antigens from the engineered cancer cells. To address these limitations, injectable or in situ forming scaffolds have been widely studied as next generation cancer vaccines [10, 76–80, 90, 109]. In addition to vaccine delivery vehicles, scaffolds provide sustained delivery of multiple immune signals and can function as sites for robust recruitment, activation and priming of DCs against TAAs in vivo. Scaffolds delivering a variety of immune cues have been investigated, including chemokines (CCL20) or cytokines (GM-CSF, Flt3L) for DC recruitment, TLRs (CpG, PolyIC, MPLA) for activation of DCs, and antigens (tumor lysates, model antigens) for the generation of strong and specific anti-tumor response [10, 76–80, 90, 109].

The Mooney Lab is one of the pioneers in this area, developing PLGA or other polymer-based scaffolds to serve as a platform for local juxtaposition of multiple immune signals to promote effective anti-tumor response [10, 76–80, 109]. Work by Ali et al. revealed these scaffolds function to enhance anti-tumor immunity by recruiting and priming phenotypically distinct populations of DCs towards TAAs [109]. Scaffolds encapsulating GM-CSF, for example, promote infiltration of conventional DCs (cDCs) that secrete high levels of T\textsubscript{H}2, and incorporation of CpG in the scaffold results in infiltration by plasmacytoid DCs (pDCs) that secrete high levels of type-1 IFNs. The recruitment of heterogeneous populations of DCs corresponded with high amounts of CTLs specific for Trp2 – a melanoma antigen – infiltrating tumors, and increased ratios of CD8\textsuperscript{+} effector T cells to T\textsubscript{REG} in tumors. These effects provided therapeutic anti-tumor immunity in the B16-F10 mouse model, where implantation of vaccine scaffolds 9 days after the establishment of tumors resulted in survival of 47% of mice at the end of the 100 day study, while
all mice vaccinated with GM-CSF-secreting irradiated B16-F10 cells succumbed by 36. Bencherif et al. recently demonstrated this strategy can be expanded to scaffolds consisting of implantable cryogels – porous hydrogels formulated at subzero temperatures to mimic tissue properties – for co-delivering irradiated whole B16-F10 cells, CpG and GM-CSF. These cryogel-based scaffolds can be injected with a syringe [110], demonstrating a minimally invasive alternative to surgically implantable scaffolds while still conferring similar therapeutic anti-tumor immunity. Vaccination with the cryogel scaffold vaccine 3 days after tumor challenge resulted in the survival of over 40% of mice during the 100 day experiment, compared to mice vaccinated with bolus vaccine which all reached terminal end points by day 38 post-challenge [76].

The clinical success of immunotherapies targeting immune checkpoints has sparked interest in combining vaccination with these ground breaking therapies. Through this combination, pathways that restrain pro-immune function might be released to create opportunities for vaccines to generate efficacious responses. For example, GVAX, a whole cell pancreatic cancer vaccine utilizing GM-CSF-expressing tumor cells showed initial promise but was ultimately halted in phase III clinical trials due to lack of efficacy [111]. GVAX is now being investigated in combination with PD-1 blockade (Nivolumab) during an ongoing phase II clinical trial [112, 113]. Additionally, sipuleucel-T is recruiting for phase II clinical trials investigating combination with CTLA-4 blockade [114].

The idea that GVAX vaccine may be enhanced by the addition of checkpoint blockade therapies highlights one important area related to this idea of combination therapy – how to overcome the immunosuppressive microenvironment of tumors. Although vaccines may generate anti-tumor immune responses, these cells are often inactivated after infiltrating tumors. In addition to the expression of checkpoint blockade ligands by tumor cells, the anti-tumor efficacy of infiltrating immune cells may also be inhibited by suppressive cytokines such as TGF-β [115] or enzymes such as indolamine 2,3-dioxygenase (IDO) [116] secreted by tumor cells or tumor-resident suppressive immune cells (e.g T<sub>REG</sub>, myeloid derived suppressor cells (MDSC)). These immunosuppressive signals can be targeted using small molecule inhibitors or siRNA as a strategy to overcome tumor immunosuppression. However, the effectiveness of systemic delivery of these cargos is limited by poor targeting to tumors and rapid degradation.

Biomaterials have recently been used to improve the delivery of immunomodulators targeting a variety of immunosuppressive pathways to actively enhance the cancer vaccination. Xu et al. recently demonstrated the potential to combine biomaterial based immunotherapies and vaccines to alter the tumor microenvironment for enhanced immunity in the B16-F10 mouse melanoma model [117]. In this study, lipid-calcium-phosphate NP (LCP NPs) based vaccines co-encapsulating Trp2 and CpG were first administered subcutaneously to generate a robust anti-tumor CTL response. This NP vaccine was augmented by intravenous injection of TGF-β siRNA delivered in liposome-protamine-hyaluronic acid NPs (LHA NPs) – a formulation the investigators optimized for cargo protection and efficient delivery to the tumor site [117]. LHA NPs efficiently knocked down up to 50% of TGF-β expression in tumors, while leaving TGF-β expression in the spleen and lymph nodes unaffected. The effect on tumor growth was investigated using both early (4 days post tumor challenge) and late stage (13 days post tumor challenge) therapeutic vaccination regimens. The early vaccine regimen was sufficient to clear small established tumors, while late stage vaccination, representing advanced disease with highly suppressive tumors, was ineffective. Systemic CTL responses generated by both vaccines was similar. This finding was demonstrated using an in vivo cytotoxicity assay which tested the ability of vaccinated mice to lyse splenocytes from naïve mice, after the splenocytes were pulsed with Trp2 peptide and adoptively transferred. However, the late stage vaccine was able to match the anti-tumor efficacy of early vaccination only with the combination of LCP NP-mediated TGF-β knock down. Importantly, mechanistic investigations revealed TGF-β knockdown depleted T<sub>REGs</sub> in tumors and caused higher levels of CD8<sup>+</sup> T cells infiltrating tumors in late-stage vaccination. These effects were minimal following immunization with the monotherapy vaccine. Together, the ideas and examples discussed in this section demonstrate the ability of biomaterials to enhance pre-clinical cancer vaccination, but these combination approaches still remain uncommon and are largely untapped in the clinic, highlighting an important investment area that could “rescue” or improve previously ineffective strategies such as cancer vaccination.

Combination therapies

With recent developments in immunotherapies, new attention is warranted to strategies combining immunotherapy with not just checkpoint blockade, but also with chemotherapy, radiation, ablation, or resection. This year the FDA granted accelerated approval for the first combination of cancer immunotherapies – checkpoint inhibitors nivolumab and ipilimumab – following phase II studies in patients with unresectable stage III or IV BRAF V600 wild-type melanoma [118]. However, the potential toxicity and difficulty in delivering multiple cargos to tumors or other target tissues is amplified in combination therapies [118]. Material delivery systems may help address this challenge by improving control over existing combination therapy schemes and make...
new multi-pronged therapies possible through controlled release, dose-sparing, or better targeting during radiation and ablation.

One promising area already being explored as a combination therapy using biomaterials is the stimulation of multiple TLRs through co-delivery of multiple TLR agonists (TLRs) \[119–121\]. Signaling through specific TLRs occurs because TLRs are recognized by receptors for their pathogen associated molecular patterns uncommon in mammals, such as double stranded RNA (e.g., polyIC, TLR3a) or endotoxins characteristic of certain pathogen categories. These TLRs can be used to drive inflammation and can amplify antigen-specific immune responses. Interestingly, recent work has shown that mixtures of TLRs can be used to elicit synergistic adjuvant responses that are distinct from the responses attributed to the administration of individual TLRs \[70, 122–126\]. Further, many biomaterial approaches aim to create pathogen-mimicking particles that incorporate multiple stimulatory signals as an alternative to the inactivated or attenuated pathogens often found in vaccines \[92, 127, 128\]. Virus-like particles (VLPs) are one good example, as these particles mimic viruses through intrinsic immunogenicity resulting from self-assembly of isolated viral proteins without use of viral genomic material.

VLPs known to accumulate in draining lymph nodes were recently studied alongside TLRs in a phase IIA human trial with stage III and IV melanoma patients \[120, 129\]. The VLPs were derived from a bacteriophage then loaded with TLR9 agonist and a peptide derived from the Melan-A/MART-1 melanoma tumor antigen. The VLPs were administered in a multi-injection regimen using one of four formulations: i) s.c. with Incomplete Freund's Adjuvant (IFA), ii) s.c. with IFA and a TLR-7 agonist (Imiquimod) applied topically at the injection site, iii) i.d. with topical Imiquimod application, and iv) intra-lymph node injection using ultrasound guidance \[120\]. Doses were matched for each immunization, except group IV which received lower doses. All patients experienced increased antibody production, with the highest levels in patients receiving IFA. These patients all experienced at least two-fold increases of Melan-A/HLA-A2 tetramer positive T cells in peripheral blood. Vaccines containing IFA also increased frequencies of effector memory T cells most efficiently (Figure 4A), while Imiquimod best expanded antigen-specific central memory T cells (Figure 4B). PET/CT scans from a patient that received s.c. immunization revealed swelling in draining lymph nodes, as well as local inflammation at vaccine sites, confirming drainage to lymphatics still plays a role in this vaccine’s efficacy (Figure 4C). This particular patient was studied for an extended period and showed increased metabolic response activity in draining lymph node hundreds of days after the last vaccine dose (Figure 4C) \[120\]. Moreover, because particle size influences particle trafficking and retention, preferential drainage of these VLPs to lymph nodes near vaccine sites may explain some of the inflammation and metabolic activity observed in patients’ lymph nodes using the peripheral injection routes. These outcomes indicate a role for the addition of multiple TLRs administered via distinct routes.

Another new combination therapy being tested in the B16-F10 melanoma model exploits the size of nanoparticles to preferentially deliver CpG and paclitaxel to tumor draining lymph nodes \[121\]. Here, both CpG and paclitaxel serve as TLR agonists (TLR9 and TLR4, respectively), rather than the more traditional role of paclitaxel as a chemotherapeutic. Particles loaded with both agonists activated DCs in vitro with no cytotoxicity. Four to nine days after implating B16-F10 tumors, the NPs were administered i.d. to limbs of mice on the same or opposite side of the tumors as an in situ vaccine. This experiment studied immune responses toward tumors directed only by TLR stimulation and the natural levels of tumor antigen expressed in the mice, instead of administration of additional tumor antigen. Particles administered on the same side and loaded with both CpG and paclitaxel accumulated in the tumor draining axillary and brachial lymph nodes, causing the greatest reduction in tumor growth \[121\]. These studies revealed that both the combination of different TLRs and the accumulation of particles in the tumor draining lymph node each influenced the therapeutic outcomes without inclusion of a tumor antigen. Thus new approaches should leverage biomaterials to break tumor tolerance and promote infiltration of immune cells that discover new tumor antigens for improved adaptive response.

Although in the clinic chemotherapeutics are regularly combined with immunotherapy, application of biomaterials to this area has been relatively sparse \[130, 131\]. This clinical promise however, along with the ease of co-encapsulating small molecule drugs (e.g. paclitaxel, doxorubicin) and biological molecules (e.g. TLRas), warrants more investment. In one recent example, a derivative of the TLR4 agonist LPS and paclitaxel were encapsulated in PLGA nanoparticles \[131\]. Peritumoral delivery of the particles elicited stronger anti-tumor effects, increasing the frequencies of tumor-infiltrating DCs (CD11c+), macrophages (CD14+), and CD4+ and CD8+ T cells while driving a 40% reduction in tumor burden relative to paclitaxel delivered alone \[131\]. Further study of these types of combinations may reveal opportunities to build on and optimize material platforms to enhance the individual effects of each therapeutic.

Radiation and ablation also provide unique opportunities for combination with immunotherapy because recruitment of immune cells to the local tissues in the treatment area can allow identification of neoantigens by immune cells infiltrating these sites. This mechanism has been hypothesized as a potential explanation for tumor regression observed in areas of the body not directly
treated via radiation or ablation [132–134]. In one effort to exploit this effect, researchers combined a polymeric NP platform with gold NPs, CpG, and photosensitive zinc phthalocyanine [135]. These complex NPs enable photothermal ablation while simultaneously delivering a TLR agonist. This approach might create an *in situ* vaccine by delivering an adjuvant to sites where ablation kills tumor cells and exposes additional tumor antigens. Thus far, however, the platform has been investigated *in vitro*. For these studies, 4T1 tumor cells were ablated with the NPs and exposed to DCs. The DCs were activated by the TLRa-loaded NPs and phagocytosed the remnants of the ablated tumor cells [135]. This strategy represents a rational approach toward combining multiple therapies: ablation and immunotherapy. Building on this broad concept, researchers should explore new approaches for using immunomodulatory agents to create *in situ* vaccines by combining them with treatments like radiation, ablation and surgeries that expose new tumor antigens. Moreover, as we learn more about the unique connections between each arm of combination therapies, this synergy may also be able to help reduce relapse, for example, by promoting memory T cell populations specific for neoantigens.

**Increased homing of immune cells to the tumor microenvironment and improved activity at these sites**

Despite the clinical success of some immunotherapies and vaccines in generating tumor specific CTLs, the anti-tumor efficacy of these treatments is limited by poor migration to tumors, as well as inactivation of cells that do reach the tumor microenvironment [2]. Biomaterials offer the potential to specifically target the tumor microenvironment through the EPR effect, as well as exciting new strategies to alter

---

**Figure 4:** A VLP-peptide vaccine containing CpG delivered *s.c.* or *i.d.* with adjuvants, or intra-nodally without adjuvants modulates antigen-specific T cell immunity. VLPs with adjuvants promote A. higher overall frequencies of effector memory T cells (CCR7^−/CD45RA^−), and B. higher frequencies of central memory T cells (CCR7^+/CD45RA^−) among antigen-specific T cells. C. PET/CT imaging of a draining lymph node in a patient receiving subcutaneous treatment (IFA only). Shown is enlargement (arrows), inflammation visible both arms and one thigh (vaccination sites), and sustained increases in glucose metabolism after vaccination indicated by standard uptake value (SUV) measurements. Adapted by permission from Macmillan Publishers Ltd: European Journal of Immunology [120] Copyright Wiley-VCH Verlag GmbH & Co. KGaA. (2012).
the trafficking of immune cells and cancer cells. This section discusses these advances.

In addition to targeting tumors at the tissue level, NP size can be tuned to passively target specific immune cells within tumors. Kourtis et al. demonstrated 30nm poly(propylene sulfide) NPs administered i.d. accumulate at high levels in the spleen and draining LNs, and are efficiently taken up by cellular populations of the myeloid lineage. In particular, NPs are efficiently internalized by myeloid derived suppressor cells (MDSC) in tumors, tumor-draining LNs and spleens. This represents a potential clinical strategy to deliver immunomodulatory cues to suppressive cell subsets [136]. Another approach integrating passive targeting of tumor cells – discussed below in the heterogeneity of disease section – was recently developed by the Shea lab to recruit tumor cells to a polymeric scaffold implanted at a distant site from the tumor [137].

NPs may also be actively targeted to lymphatic tumors for the delivery of therapeutics by taking advantage of the ability of lymphocytes to home to lymphoid organs. Huang et al. demonstrated a strategy to deliver NPs encapsulating a chemotherapeutic (SN-38) to lymphoma tumors through the conjugation of NPs to polyclonal T cells from C57BL/6J mice [138]. T cells were expanded ex vivo in the presence of rapamycin, which polarized T cells toward phenotypes that retain high levels of lymphoid homing receptors (e.g., CD62L, and CCR7) after expansion. NP-conjugated T cells effectively trafficked to lymphoid tumors after adoptive transfer, causing increased concentrations of SN-38 in these lymph nodes compared to administration of unconjugated NPs. This active targeting of tumors resulted in increased survival of tumor bearing mice at greatly reduced doses.

The above examples demonstrate the benefits of passively targeting tumors after administration at a distant site from the tumor. In some cases, i.t. injection of non-resectable tumors offers a simple route to directly deliver immunostimulants that locally reactivate exhausted immune cells in the tumor microenvironment. By causing these reactivated APCs to be primed against TAAs within tumors, localized i.t. therapy may thus offer exciting potential to generate systemic anti-tumor immunity to combat distal tumors, or memory that helps address relapse [139]. Using biomaterials with this strategy capitalizes on the ability of particulate material to be retained in tumors, while soluble immunostimulants are rapidly drained from tumors. Liu et al. demonstrated that CpG can be retained locally in tumors by conjugating CpG to lipophilic moieties, which readily insert into tumor cell membranes after i.t. injection [140]. A strategy reported by Intra et al. involves activation of the immune system to clear tumor cells after resection to prevent relapse. In this study, PLGA sutures closing the incision from resection were used to encapsulate and sustain the release of CpG locally at this site [38]. Using these CpG-loaded sutures during resection of N2a neuroblastoma – an extracranial pediatric cancer – protected mice from relapse and prolonged survival compared to mice sutured with unloaded PLGA and locally injected with CpG.

In addition to targeting the tumor microenvironment with immunostimulants, strategies aimed at neutralizing immunosuppressive cytokines may also allow anti-tumor immunity to overcome the suppressive mechanisms of the tumor microenvironment. IL-2 therapy, for example, shows potential by augmenting the activation and expansion of anti-tumor CTLs. Clinical trials testing IL-2 during metastatic melanoma demonstrate 4-6% of patients achieve complete remission [141, 142]. One major barrier limiting the robustness of this idea is the restraint of IL-2 induced expansion of CTLs by TGF-β and other immunosuppressive cytokines present in tumors [115]. Park et al. has enhanced IL-2 based immunotherapy in mice using nanoparticles co-loaded with IL-2 and a small molecule inhibitor of TGF-β1 (SB505124) [143]. In this strategy nanoparticle liposomal polymeric gels (nLGs) were designed. The nLGs were formed from a biodegradable polymer core loaded with drug solubilized in cyclodextrans and then enveloped in a PEGylated lipid shell. This design allowed efficient encapsulation and co-delivery of both hydrophilic IL-2 and hydrophobic SB505124. Passive targeting due to the EPR effect led to nLG accumulation in both subcutaneous tumors (Figure 5A, 5B) and metastatic pulmonary tumors after nLGs were administered intravenously to mice bearing B16-F10 tumors. Weekly administration of nLGs encapsulating IL-2 and SB505124 protected mice from tumors and enhanced survival compared to soluble systemic treatment, which conferred no benefit compared with untreated mice (Figure 5C). Mechanistically, these therapeutic effects correlated with increased NK cell frequencies in tumors and improved CTL recruitment. These examples demonstrate new ways in which biomaterial strategies might help overcome inactivation of CTLs in the tumor microenvironment, a challenge that connects with the hurdle discussed next, heterogeneity of disease.

**Addressing heterogeneity of disease**

A major barrier to cancer immunotherapy is the highly heterogeneous nature of tumors within individual patients, in different types of cancer, and across patient populations [3, 144, 145]. Tumor cells are dynamic, exhibiting phenotypic and antigenic heterogeneity that creates variable susceptibilities to different vaccines and immunotherapies, as well as evolution of these responses as treatment continues or during relapse [144].

The effectiveness of a cancer vaccine depend in part on the individual MHC alleles expressed by the patient and the level of expression of TAAs targeted by the vaccine [146]. Frequent mutations or immunoselection in response to vaccination may drive the loss of expression
of targeted antigens [147]. One potential strategy to overcome this challenges is development of personalized vaccines for individual patients. For example, antigens present in a patient’s tumor that exhibit a high affinity for their individual MHC alleles could be identified and utilized in antigen selection [148]. Additionally, vaccines might utilize multiple patient specific TAAs for a broad immune response less susceptible to loss of antigens or antigenic editing by tumors [149]. However, while vaccines in the clinic targeting well characterized TAAs, such as in melanoma, have been effective at causing partial tumor regression, complete regression is rare, likely due to antigenic shift of tumors in response to immune attack of tumors induced by vaccination [147, 150]. Importantly, recent preclinical studies reveal neoantigens driven by mutations are seen as non-self and preferentially targeted by CTLs after checkpoint blockade [151]. This outcome suggests neoantigens – if identifiable during treatment – could be used effectively for vaccination by pairing with checkpoint blockade [152, 153]. A majority of neoantigens driven by mutation are patient-specific, and one current challenge is that the genome of a patient’s own tumor cells must be sequenced if prepared de novo.

Resected tumors theoretically contain all antigens present in an individual’s tumor, including neoantigens resulting from mutations. Thus these tissues may be used as effective antigen sources in personalized cancer vaccines. Using irradiated tumor cells or tumor lysates as antigen sources in vaccines may likewise offer the potential to better combat tumor immunoselection [149]. Biomaterials can further enhance this approach by targeting to immune cells, sustained delivery of antigens, and co-localization with adjuvants – all discussed previously for cancer vaccines based on well-defined antigens. The previously-discussed scaffold vaccines from the Mooney lab employed irradiated whole tumor cells or tumor lysates as a source of antigens. PLGA MP and NP loaded with tumor lysates have also been developed [154] and have recently been shown to enhance immunogenicity in preclinical breast cancer models [154, 155]. Another interesting idea recently reported is the use of exosomes secreted from DCs after DCs were stimulated with PolyIC and loaded with tumor lysates to create an effective vaccine in murine models of melanoma [156].

Advances in sequencing technologies have enabled efficient exosome mapping of tumor cells [157]. In this area, biomaterials and nanotechnology are rapidly developing as powerful tools to enable capture of rare circulating tumor cells (CTCs) [11, 158–162], capabilities that may enhance personalized cancer immunotherapy by allowing identification of neoantigens in individual patients [148]. This approach might open new potential even for patients with haematological tumors or unresectable solid tumors. Additionally, capture of CTCs may allow for early detection of cancer and patient-specific drug or immunotherapy screening to determine the best treatment options [65].

Another new strategy aimed at CTCs is based on altering or exploiting the homing of these cells. Azarin et al. recently demonstrated a biomaterial strategy that enables efficient capture of CTCs using implantable scaffolds, which function to preferentially redirect metastatic tumor cells from other solid organs [137]. In this study PLGA scaffolds were designed to mimic the pre-metastatic niche – an environment consisting of matrix proteins, cytokines and chemokines as well as immune cells – to create a conducive location for new tumors from circulating metastatic cells. Tumor cells accumulated in scaffolds after subcutaneous or intra-peritoneal implantation of scaffolds in mice bearing breast cancer tumors in the mammary fat pads (Figure 6A) [137]. In contrast, tumor cells were not detected in the peritoneal fat pads of mice not implanted with scaffolds (Figure 6B). Additionally, the accumulation of metastatic tumor cells in scaffolds correlated with decreased incidence of metastases in the lungs and...
liver, as well as decreased metastatic tumor burdens in the lungs (Figure 6C). Mechanistically, the recruitment of CTCs was mediated by the infiltration of leukocytes (Figure 6D). These infiltrating leukocytes consisted of populations of immune cells at similar relative levels to those seen in the lungs of tumor bearing mice, which are sites of frequent metastasis. In particular Gr1<sup>hi</sup>CD11b<sup>+</sup> MDSCs accumulated in scaffolds to a high degree, consisting of 66% of total leukocytes in the scaffold, compared to 89% of total leukocytes in the lungs 28 days after tumor inoculation [137]. By utilizing scaffolds to locally deliver lentiviral vectors encoding the chemokine CCL22, Gr1<sup>hi</sup>CD11b<sup>+</sup> cell infiltration in the scaffold was increased significantly by day 14 after implantation, compared to scaffolds delivering a control vector (Figure 6D). Additionally, this chemokine increased infiltrating Gr1<sup>hi</sup>CD11b<sup>+</sup> cells to enhance tumor cell capture in the scaffold. This study also demonstrated a potential strategy to address heterogeneity of disease with respect to metastatic potential, where early metastases can be detected by label-free in situ imaging of scaffolds based on optical properties of cancer cells. This capability may allow for patients to be treated based on expected aggressiveness of their disease. Additionally, metastatic cancer cells may be removed from scaffolds or other “sensor” devices for analysis of biomarkers or antigen expression. An especially exciting aspect is the multi-functionality that these types of devices could play as both a diagnostic tool to address heterogeneity, and as a strategy to delay or inhibit metastases to allow time for parallel treatments to work.

CONCLUSIONS

It is clear that immunotherapies for cancer treatment have progressed greatly in recent years, revealing new possibilities through which the immune system can be used to treat cancer. The successes and failures of immunotherapies in the clinic have given researchers newfound understanding of the significant challenges presented by cancers that limit current immunotherapies. Nanotechnologies have been investigated extensively for cancer immunotherapies and vaccination pre-clinically, but have had comparatively little impact for patients thus far. One common occurrence is clinicians and engineers coupling cutting-edge ideas from their respective field with basic – or sometimes outdated – ideas from the other field. For example, much nanoparticle research is being performed on small molecules drugs delivered as a monotherapy that are never used as a primary therapy in patients. Similarly, most clinical therapies involve systemic delivery, while many pre-clinical engineering strategies exist for targeting specific cells or tissues. Further, as can be observed by range of papers reviewed here, a great deal of pre-clinical work involving biomaterials is conducted in fairly basic models – for example, B16-F10. Although these are valuable starting points, utilization of more sophisticated animal models that recapitulate cardinal features – such as pathology and metastatic tendency – could improve the translatability of new material strategies. This gap highlights the potential for impact that materials can have as the focus on cross-disciplinary understanding and collaboration grows; there is outstanding synergy at the interface of nanotechnology, bioengineering, and immunotherapy. This nexus creates enormous opportunities for teams of researchers that understand both the clinical and technological needs of new cancer therapies.

CONFLICTS OF INTEREST

The Authors do not have any conflicts of interest.
GRANT SUPPORT

This work was supported in part by NSF CAREER Award # 1351688, Alex’s Lemonade Stand “A” Award # 27120, and support to C.M.J. through the Damon Runyon-Rachleff Innovator Award (# DRR3415), the Alliance for Cancer Gene Therapy Young Investigator Award (# 15051543), and the Melanoma Research Alliance Young Investigator Award (# 348963). J.M.G. is a grantee of the Pediatric Oncology Student Training award from Alex’s Lemonade Stand Foundation # 15082537.

REFERENCES

1. Weber JS, Yang JC, Atkins MB and Disis ML. Toxicities of Immunotherapy for the Practitioner. J Clin Oncol. 2015; 33:2092-U2154.
2. Motz GT and Coukos G. Deciphering and Reversing Tumor Immune Suppression. Immunity. 2013; 39:61-73.
3. Doroshow JH and Kurnar S. Translational research in oncology-10 years of progress and future prospects. Nat Rev Clin Oncol. 2014; 11:649-662.
4. Rosenberg SA and Restifo NP. Adoptive cell transfer as personalized immunotherapy for human cancer. Science. 2015; 348:62-68.
5. Galluzzi L, Vacchelli E, Bravo-San Pedro JM, Buque A, Senovilla L, Baracco EE, Bloy N, Castoldi F, Abastado JP, Agostinis P, Apte RN, Aranda F, Ayyoub M, et al. Classification of current anticancer immunotherapies. Oncotarget. 2014; 5:12472-12508. doi: 10.18632/oncotarget.2998.
6. Kim J and Mooney DJ. In vivo modulation of dendritic cells by engineered materials: Towards new cancer vaccines. Nano Today. 2011; 6:466-477.
7. Sahdev P, Ochyl LJ and Moon JJ. Biomaterials for Nanoparticle Vaccine Delivery Systems. Pharm Res-Dordr. 2014; 31:2563-2582.
8. Bowen A and Casadevall A. Increasing disparities between resource inputs and outputs, as measured by certain health deliverables, in biomedical research. P Natl Acad Sci USA. 2015; 112:11335-11340.
9. Singh A and Peppas NA. Hydrogels and Scaffolds for Immunomodulation. Adv Mater. 2014; 26:6530-6541.
10. Kim J, Li WA, Choi Y, Lewin SA, Verbeke CS, Dranoff G and Mooney DJ. Injectable, spontaneously assembling, inorganic scaffolds modulate immune cells in vivo and increase vaccine efficacy. Nat Biotechnol. 2015; 33:64-U241.
11. Reategui E, Acco N, Lim EJ, Sullivan JP, Jensen AE, Zeinali M, Martel JM, Aranyosi AJ, Li W, Castleberry S, Bardia A, Sequist LV, Haber DA, Maheswaran S, Hammond PT, Toner M, et al. Tunable Nanostructured Coating for the Capture and Selective Release of Viable Circulating Tumor Cells. Adv Mater. 2015; 27:1593-1599.
12. Domachuk P, Tsioris K, Omenetto FG and Kaplan DL. Biomechanics: Biomaterials and Biomimetic Designs. Adv Mater. 2010; 22:249-260.
13. Shao K, Singha S, Clemente-Casares X, Tsai S, Yang Y and Santamaria P. Nanoparticle-Based Immunotherapy for Cancer. Acs Nano. 2015; 9:16-30.
14. Fang J, Nakamura H and Maeda H. The EPR effect: Unique features of tumor blood vessels for drug delivery, factors involved, and limitations and augmentation of the effect. Adv Drug Deliver Rev. 2011; 63:136-151.
15. Maruyama K. Intracellular targeting delivery of lipidosomal drugs to solid tumors based on EPR effects. Adv Drug Deliver Rev. 2011; 63:161-169.
16. Andorko JI, Hess KL and Jewell CM. Harnessing Biomaterials to Engineer the Lymph Node Microenvironment for Immunity or Tolerance. Aaps J. 2015; 17:323-338.
17. Sharp FA, Ruane D, Claass B, Creagh E, Harris J, Malyala P, Singh M, O’Hagan DT, Petrilli V, Tschopp J, O’Neill LAJ and Lavelle EC. Uptake of particulate vaccine adjuvants by dendritic cells activates the NALP3 inflammasome. P Natl Acad Sci USA. 2009; 106:870-875.
18. Andorko JI, Hess KL, Pineault KG and Jewell CM. Intrinsic immunogenicity of rapidly-degradable polymers evolves during degradation. Acta Biomater. 2015; 32:24-34.
19. Hudalla GA, Sun T, Gasiorowski JZ, Han H, Tian YF, Chong AS and Collier JH. Graded assembly of multiple proteins into supramolecular nanomaterials. Nature materials. 2014; 13:829-836.
20. Zhang P, Chiu YC, Tostanoski LH and Jewell CM. Polyelectrolyte Multilayers Assembled Entirely from Immune Signals on Gold Nanoparticle Templates Promote Antigen-Specific T Cell Response. Acs Nano. 2015; 9:6465-6477.
21. Y. C. Chiu JMG, J. I. Andorko, L. H. Tostanoski, and C. M. Jewell. Modular Vaccine Design Using Carrier-Free Nanoparticles. Biomater Sci Eng. 2015; 1:1200-1205.
22. Venditto VJ and Szoka FC. Cancer nanomedicines: So many papers and so few drugs! Adv Drug Deliver Rev. 2013; 65:80-88.
23. Cheng ZL, Al Zaki A, Hui JZ, Muzykantov VR and Tsourkas A. Multifunctional Nanoparticles: Cost Versus Benefit of Adding Targeting and Imaging Capabilities. Science. 2012; 338:903-910.
24. Juliano R. Nanomedicine: is the wave cresting? Adv Drug Deliver Rev. 2013; 65:80-88.
25. Domachuk P, Tsioris K, Omenetto FG and Kaplan DL. Biomechanics: Biomaterials and Biomimetic Designs. Adv Mater. 2010; 22:249-260.
26. Caspi RR. Immunotherapy of autoimmunity and cancer: the penalty for success. Nat Rev Immunol. 2008; 8:970-976.
27. Svenson S. What nanomedicine in the clinic right now really forms nanoparticles? Wiley Interdisciplinary Reviews: Nanomedicine and Nanobiotechnology. 2014; 6:125-135.
28. Carstens MG, Camps MGM, Henriksen-Lacey M, Franken K, Ottenhoff TTH, Perrie Y, Bouwstra JA, Osendorf F and Jiskoot W. Effect of vesicle size on tissue localization and immunogenicity of liposomal DNA vaccines. Vaccine. 2011; 29:4761-4770.
29. Kwong B, Gai SA, Elkhader J, Wittrup KD and Irvine DJ. Localized Immunotherapy via Liposome-Anchor Anti-CD137 + IL-2 Prevents Lethal Toxicity and Elicits Local and Systemic Antitumor Immunity. Cancer Research. 2013; 73:1547-1558.
30. Kwong B, Liu H and Irvine DJ. Induction of potent anti-tumor responses while eliminating systemic side effects via liposome-anchored combinatorial immunotherapy. Biomaterials. 2011; 32:5134-5147.
31. Wang Y, Xu Z, Guo S, Zhang L, Sharma A, Robertson GP and Huang L. Intravenous delivery of siRNA targeting CD47 effectively inhibits melanoma tumor growth and lung metastasis. Molecular therapy. 2013; 21:1919-1929.
32. de Titta A, Ballester M, Juliet Z, Nembrini C, Jeanbart L, van der Vlies AJ, Swartz MA and Hubbell JA. Nanoparticle conjugation of CpG enhances adjuvancy for cellular immunity and memory recall at low dose. Proc Natl Acad Sci U S A. 2013; 110:19902-19907.
33. Lemke CD, Geary SM, Joshi VB and Salem AK. Antigen-coated poly α-hydroxy acid based microparticles for heterologous prime-boost adenovirus based vaccinations. Biomaterials. 2013; 34:2524-2529.
34. Hanson MC, Crespo MP, Abraham W, Moynihan KD, Szeto GL, Chen SH, Melo MB, Mueller S and Irvine DJ. Nanoparticulate STING agonists are potent lymph node–targeted vaccine adjuvants. Journal of Clinical Investigation. 2015; 125:2523-2546.
35. Radovic-Moreno AF, Cherryak N, Mader CC, Nallagatla S, Kang RS, Hao LL, Walker DA, Halo TL, Merkel TJ, Rische CH, Anantatmula S, Burkhart M, Mirkin CA and Gryaznov SM. Immunomodulatory spherical nucleic acids. P Natl Acad Sci USA. 2015; 112:3892-3897.
36. Hori Y, Stern PJ, Hynes RO and Irvine DJ. Engulfing tumors with synthetic extracellular matrices for cancer immunotherapy. Biomaterials. 2009; 30:6757-6767.
37. Wen Y, Kolonich HR, Kruszewski KM, Giannoukakis N, Gawalt ES and Meng WS. Retaining Antibodies in Tumors with a Self-Assembling Injectable System. Molecular Pharmaceutics. 2013; 10:1035-1044.
38. Intra J, Zhang XQ, Williams RL, Zhu X, Sandler AD and Salem AK. Innate immunostimulatory suture that treat local disease recurrence following primary tumor resection. Biomedical materials. 2011; 6:011001.
39. Liu H, Moynihan KD, Zheng Y, Szeto GL, Li AV, Huang B, Van Egeren DS, Park C and Irvine DJ. Structure-based programming of lymph-node targeting in molecular vaccines. Nature. 2014; 507:519-522.
40. Vo JL, Yang L, Kurtz SL, Smith SG, Koppolu BP, Ravindranathan S and Zaharoff DA. Neoadjuvant immunotherapy with chitosan and interleukin-12 to control breast cancer metastasis. Oncoimmunology. 2014; 3:e968001-e968001.
41. June CH. Adoptive T cell therapy: Grand challenges and opportunities. J Immunother. 2007; 30:858-858.
42. Fadel TR, Sharp FA, Vudattu N, Ragheb R, Garyu J, Kim D, Hong E, Li N, Haller GL, Pfefferle LD, Justesen S, Herold KC and Fahmy TM. A carbon nanotube-polymer composite for T-cell therapy. Nature nanotechnology. 2014; 9:639-647.
43. Perica K, Bieler JG, Schutz C, Varela JC, Douglass J, Skora A, Chiu YL, Oelke M, Kinzler K, Zhou SB, Vogelstein B and Schneck JP. Enrichment and Expansion with Nanoscale Artificial Antigen Presenting Cells for Adoptive Immunotherapy. Acs Nano. 2015; 9:6861-6871.
44. Wolff M and Greenberg PD. Antigen-specific activation and cytokine-facilitated expansion of naive, human CD8(+) T cells. Nat Procot. 2014; 9:950-966.
45. Stephan MT, Moon JJ, Um SH, Bershteyn A and Irvine DJ. Therapeutic cell engineering with surface-conjugated synthetic nanoparticles. Nature medicine. 2010; 16:1035-1041.
46. Stephan MT, Stephan SB, Bak P, Chen J and Irvine DJ. Synapse-directed delivery of immunomodulators using T-cell-conjugated nanoparticles. Biomaterials. 2012; 33:5776-5787.
47. Zheng Y, Stephan MT, Gai SA, Abraham W, Shearer A and Irvine DJ. In vivo targeting of adoptively transferred T-cells with antibody- and cytokine-conjugated liposomes. Journal of controlled release. 2013; 172:426-435.
48. Lorenz U. SHP-1 and SHP-2 in T cells. Nat Protoc. 2014; 9:639-647.
49. Pardoll DM. The blockade of immune checkpoints in cancer immunotherapy. Nat Rev Cancer. 2012; 12:252-264.
50. Sharma P and Allison JP. Immune Checkpoint Targeting in Cancer Therapy: Toward Combination Strategies with Curative Potential. Cell. 2015; 161:205-214.
51. Pradhan P, Qin H, Leleux JA, Gwak D, Sakamaki I, Kwak LW and Roy K. The effect of combined IL10 siRNA and CpG ODN as pathogen-mimicking microparticles on Th1/Th2 cytokine balance in dendritic cells and protective immunity against B cell lymphoma. Biomaterials. 2014; 35:5491-5504.
52. Singh A, Qin H, Fernandez I, Wei J, Lin J, Kwak LW and Roy K. An injectable synthetic immune-priming center mediates efficient T-cell class switching and T-helper 1
response against B cell lymphoma. Journal of controlled release. 2011; 155:184-192.

54. Smith SG, Koppolu BP, Ravindranathan S, Kurtz SL, Yang L, Katz MD and Zaharoff DA. Intravesical chitosan/interleukin-12 immunotherapy induces tumor-specific systemic immunity against murine bladder cancer. Cancer immunology, immunotherapy. 2015; 64:689-696.

55. Zaharoff DA, Hoffman BS, Hooper HB, Benjamin CJ, Jr., Khurana KK, Hance KW, Rogers CJ, Pinto PA, Schlom J and Greiner JW. Intravesical immunotherapy of superficial bladder cancer with chitosan/interleukin-12. Cancer Res. 2009; 69:6192-6199.

56. Crompton JG, Sukumar M and Restifo NP. Uncoupling T-cell expansion from effector differentiation in cell-based immunotherapy. Immunol Rev. 2014; 257:264-276.

57.Gattinoni L, Klebanoff CA, Palmer DC, Wrzesinski C, Kerstann K, Yu Z, Finkelstein SE, Theoret MR, Rosenberg SA and Restifo NP. Acquisition of full effector function in vitro paradoxically impairs the in vivo antitumor efficacy of adoptively transferred CD8+ T cells. The Journal of clinical investigation. 2005; 115:1616-1626.

58. Dementol SL, Cui W, Criscione JM, Stern E, Tulipan J, Kaech SM and Fahmy TM. Role of sustained antigen release from nanoparticle vaccines in shaping the T cell memory phenotype. Biomaterials. 2012; 33:4957-4964.

59. Araki K, Turner AP, Shaffer VO, Gangappa S, Keller SA, Bachmann MF, Larsen CP and Ahmed R. mTOR regulates memory CD8 T-cell differentiation. Nature. 2009; 460:108-U124.

60. Zeng R, Spolksi R, Finkelstein SE, Oh SK, Kovanen PE, Hinrichs CS, Pise-Masison CA, Radonovich MF, Brady JN, Restifo NP, Berzofsky JA and Leonard WJ. Synergy of IL-21 and IL-15 in regulating CD8(+) T cell expansion and function. J Exp Med. 2005; 201:139-148.

61. Fofaria NM and Srivastava SK. Critical role of STAT3 in melanoma metastasis through anoikis resistance. Oncotarget. 2014; 5:7051-7064. doi: 10.18632/oncotarget.2251.

62. Mulders PF, De Santis M, Powles T and Fizazi K. Targeted therapy of metastatic, asymptomatic hormone refractory prostate cancer with sipuleucel-T immunotherapy. Cancer Immunol. 2015; 64:655-663. doi: 10.18632/oncotarget.15439.
77. Ali OA, Doherty E, Mooney DJ and Emerich D. Relationship of vaccine efficacy to the kinetics of DC and T-cell responses induced by PLG-based cancer vaccines. Biomatter. 2011; 1:66-75.
78. Ali OA, Huebsch N, Cao L, Dranoff G and Mooney DJ. Infection-mimicking materials to program dendritic cells in situ. Nature materials. 2009; 8:151-158.
79. Ali OA, Tayalia P, Shvartsman D, Lewin S and Mooney DJ. Inflammatory cytokines presented from polymer matrices differentially generate and activate DCs. Advanced functional materials. 2013; 23:4621-4628.
80. Ali OA, Verbeke C, Johnson C, Sands RW, Lewin SA, White D, Doherty E, Dranoff G and Mooney DJ. Identification of immune factors regulating antitumor immunity using polymeric vaccines with multiple adjuvants. Cancer Res. 2014; 74:1670-1681.
81. Joshi VB, Geary SM, Carrillo-Conde BR, Narasimhan B and Salem AK. Characterizing the antitumor response in mice treated with antigen-loaded polyanhydride microparticles. Acta biomaterialia. 2013; 9:5583-5589.
82. Vasiievich EA, Chen W and Huang L. Enantiospecific adjuvant activity of cationic lipid DOTAP in cancer vaccine. Cancer immunology, immunotherapy. 2011; 60:629-638.
83. Vasiievich EA, Ramishetti S, Zhang Y and Huang L. Trp2 peptide vaccine adjuvanted with (R)-DOTAP inhibits tumor growth in an advanced melanoma model. Mol Pharm. 2012; 9:261-268.
84. Black M, Trent A, Kostenko Y, Lee JS, Olive C and Tirrell M. Self-assembled peptide amphiphile micelles containing a cytotoxic T-cell epitope promote a protective immune response in vivo. Adv Mater. 2012; 24:3845-3849.
85. Campbell DF, Saenz R, Bharati IS, Seible D, Zhang L, Esener S, Messmer B, Larsson M and Messmer D. Enhanced anti-tumor immune responses and delay of tumor development in human epidermal growth factor receptor 2 mice immunized with an immunostimulatory peptide in poly(D,L-lactic-co-glycolic) acid nanoparticles. Breast cancer research. 2015; 17:48.
86. Julier Z, Martino MM, de Titta A, Jeambart L and Hubbell JA. The TLR4 agonist fibronectin extra domain A is cryptic, exposed by elastase-2; use in a fibrin matrix cancer vaccine. Scientific reports. 2015; 5:8569.
87. Liu TY, Hussein WM, Giddam AK, Jia Z, Reiman JM, Zaman M, McMillan NA, Good MF, Monteiro MJ, Toth I and Skwarczynski M. Polyacrylate-based delivery system for self-adjuvanting anticancer peptide vaccine. Journal of medicinal chemistry. 2015; 58:888-896.
88. Liu TY, Hussein WM, Jia Z, Ziora ZM, McMillan NA, Monteiro MJ, Toth I and Skwarczynski M. Self-adjuvanting polymer-peptide conjugates as therapeutic vaccine candidates against cervical cancer. Biomacromolecules. 2013; 14:2798-2806.
89. Phua KK, Boczkowski D, Dannull J, Pruitt S, Leong KW and Nair SK. Whole blood cells loaded with messenger RNA as an anti-tumor vaccine. Advanced healthcare materials. 2014; 3:837-842.
90. Liu Y, Xiao L, Joo Ki, Hu B, Fang J and Wang P. In situ modulation of dendritic cells by injectable thermosensitive hydrogels for cancer vaccines in mice. Biomacromolecules. 2014; 15:3836-3845.
91. Phua KKL, Staats HF, Nair SK and Leong KW. Intranasal mRNA nanoparticle vaccination induces prophylactic and therapeutic anti-tumor immunity. Journal of Controlled Release. 2015; 213:E66-E67.
92. Irvine DJ, Swartz MA and Szeto GL. Engineering synthetic vaccines using cues from natural immunity. Nature materials. 2013; 12:978-990.
93. Moon JJ, Suh H, Bershteyn A, Stephan MT, Liu HP, Huang B, Sohail M, Luo S, Um SH, Khant H, Goodwin JT, Ramos J, Chiu W and Irvine DJ. Interbilayer-crosslinked multilamellar vesicles as synthetic vaccines for potent humoral and cellular immune responses. Nature materials. 2011; 10:243-251.
94. Kovacsovicsbankowski M and Rock KL. A Phagosome-to-Cytosol Pathway for Exogenous Antigens Presented on Mhc Class-I Molecules. Science. 1995; 267:243-246.
95. Shen H, Ackerman AL, Cody V, Giodini A, Hinson ER, Cresswell P, Edelson RL, Saltzman WM and Hanlon DJ. Enhanced and prolonged cross-presentation following endosomal escape of exogenous antigens encapsulated in biodegradable nanoparticles. Immunology. 2006; 117:78-88.
96. Hirose S, Kourtis IC, van der Vlies AJ, Hubbell JA and Swartz MA. Antigen delivery to dendritic cells by poly(propylene sulfide) nanoparticles with disulfide conjugated peptides: Cross-presentation and T cell activation. Vaccine. 2010; 28:7897-7906.
97. Scott EA, Stano A, Gillard M, Maio-Liu AC, Swartz MA and Hubbell JA. Dendritic cell activation and T cell priming with adjuvant- and antigen-loaded oxidation-sensitive polymersomes. Biomaterials. 2012; 33:6211-6219.
98. Keller S, Wilson JT, Patilea GI, Kern HB, Convertine AJ and Stayton PS. Neutral polymer micelle carriers with pH-responsive, endosome-releasing activity modulate antigen trafficking to enhance CDS(+) T cell responses. Journal of Controlled Release. 2014; 191:24-33.
99. Wilson JT, Postma A, Keller S, Convertine AJ, Moad G, Rizzardo E, Meagher L, Chieffari J and Stayton PS. Enhancement of MHC-I Antigen Presentation via Architectural Control of pH-Responsive, Endosomolytic Polymer Nanoparticles. Aaps J. 2015; 17:358-369.
100. Foster S, Duvall CL, Crownover EF, Hoffman AS and Stayton PS. Intracellular Delivery of a Protein Antigen with an Endosomal-Releasing Polymer Enhances CD8 T-Cell Production and Prophylactic Vaccine Efficacy. Bioconjugate Chem. 2010; 21:2205-2212.
101. Wilson JT, Keller S, Manganelli MJ, Cheng C, Lee CC, Opara C, Convertine A and Stayton PS. pH-Responsive Nanoparticle Vaccines for Dual-Delivery of Antigens and
102. Rosalia RA, Cruz LJ, van Duikeren S, Tromp AT, Silva AL, Jiskoot W, de Grujil T, Lowik C, Oostendorp J, van der Burg SH and Ossendrop F. CD40-targeted dendritic cell delivery of PLGA-nanoparticle vaccines induce potent antitumor responses. Biomaterials. 2015; 40:88-97.

103. Zeng Q, Jiang H, Wang T, Zhang Z, Gong T and Sun X. Cationic micelle delivery of Trp2 peptide for efficient lymphatic draining and enhanced cytotoxic T-lymphocyte responses. Journal of controlled release. 2015; 200:1-12.

104. Seong SY and Matzinger P. Hydrophobicity: an ancient damage-associated molecular pattern that initiates innate immune responses. Nat Rev Immunol. 2004; 4:469-478.

105. Perica K, De Leon Medero A, Durai M, Chiu YL, Bieler JG, Siblener L, Niemoller M, Assenmacher M, Richter A, Eddin M, Oelke M and Schneck J. Nanoscale artificial antigen presenting cells for T cell immunotherapy. Nanomedicine. 2014; 10:119-129.

106. Ugel S, Zoso A, De Santo C, Li Y, Marigo I, Zanovello P, Scarselli E, Cipriani B, Oelke M, Schneck JP and Bronte V. In vivo administration of artificial antigen-presenting cells activates low-avidity T cells for treatment of cancer. Cancer Res. 2009; 69:9376-9384.

107. de Grujil TD, van den Eertwegh AJM, Pinedo HM and Scheper RJ. Whole-cell cancer vaccination: from autologous to allogeneic tumor- and dendritic cell-based vaccines. Cancer Immunol Immun. 2008; 57:1569-1577.

108. Dranoff G, Jaffe E, Lazenby A, Golubek P, Levitsky H, Brose K, Jackson V, Hamada H, Pardoll D and Mulligan RC. Vaccination with Irradiated Tumor-Cells Engineered to Secrete Murine Granulocyte-Macrophage Colony-Stimulating Factor Stimulates Potent, Specific, and Long-Lasting Antitumor Immunity. P Natl Acad Sci USA. 1993; 90:3539-3543.

109. Ali OA, Emerich D, Dranoff G and Mooney DJ. In Situ Regulation of DC Subsets and T Cells Mediates Tumor Regression in Mice. Sci Transl Med. 2009; 1.

110. Bencherif SA, Sands RW, Bhatta D, Arany P, Verbeke CS, Edwards DA and Mooney DJ. Injectable preformed scaffolds with shape-memory properties. Proc Natl Acad Sci U S A. 2012; 109:19590-19595.

111. Arlen PM, Mohebtash M, Madan RA and Gulley JL. Promising novel immunotherapies and combinations for prostate cancer. Future Oncol. 2009; 5:187-196.

112. Bartholomew RM. GVAX Pancreas Vaccine (With CY) and CRS-207 With or Without Nivolubam. ClinicalTrials.gov Bethesda (MD): National Library of Medicine (US). 2015 (Accessed 12/14/2015):Available using NLM Identifier NCT02243371.

113. Zheng L. Neoadjuvant/Adjuvant GVAX Pancreas Vaccine (With CY) With or Without Nivolubam Trial for Surgically Resectable Pancreatic Cancer. ClinicalTrials.gov Bethesda (MD): National Library of Medicine (US). 2015 (Accessed 12/14/2015):Available using NLM Identifier NCT02451982.

114. Fong S. A Randomized Phase 2 Trial of Combining Sipuleucel-T With Immediate vs. Delayed CTLA-4 Blockade for Prostate Cancer. ClinicalTrials.gov Bethesda (MD): National Library of Medicine (US). 2015 (Accessed 12/14/2015 ):Available using NLM Identifier NCT01804465.

115. Wrzesinski SH, Wan YSY and Flavell RA. Transforming growth factor-beta and the immune response: Implications for anticancer therapy. Clin Cancer Res. 2007; 13:5262-5270.

116. Holmgard RB, Zamarin D, Li YY, Gasmì B, Munn DH, Allison JP, Merghoub T and Wolchok JD. Tumor-Expressed IDO Recruits and Activates MDSCs in a Treg-Dependent Manner. Cell Rep. 2015; 13:412-424.

117. Xu ZH, Wang YH, Zhang L and Huang L. Nanoparticle-Delivered Transforming Growth Factor-beta siRNA Enhances Vaccination against Advanced Melanoma by Modifying Tumor Microenvironment. Acs Nano. 2014; 8:3636-3645.

118. Larkin J, Chiarion-Sileni V, Gonzalez R, Grob JJ, Cowey CL, Lao CD, Schadendorf D, Dummer R, Smylie M, Rutkowski P, Ferrucci PF, Hill A, Wagstaff J, Carlino MS, Haanen JB, Maio M, et al. Combined Nivolumab and Ipilimumab or Monotherapy in Untreated Melanoma. The New England journal of medicine. 2015; 373:23-34.

119. Fox CB, Sivananthan SJ, Duthie MS, Vergara J, Guderian JA, Moon E, Coblentz D, Reed SG and Carter D. A nanoliposome delivery system to synergistically trigger TLR4 AND TLR7. Journal of nanobiotechnology. 2014; 12:17-17.

120. Goldinger SM, Dummer R, Baumgaertner P, Mihic-Probst D, Schwarz K, Hammann-Haenni A, Willers J, Geldhof C, Prior JO, Kündig TM, Michielin O, Bachmann MF and Speiser DE. Nano-particle vaccination combined with TLR-7 and -9 ligands triggers memory and effector CD8+ T-cell responses in melanoma patients. European journal of immunology. 2012; 42:3049-3061.

121. Thomas SN, Vokali E, Lund AW, Hubbell JA and Swartz MA. Targeting the tumor-draining lymph node with adjuvanted nanoparticles reshapes the anti-tumor immune response. Biomaterials. 2014; 35:814-824.

122. Bagchi A, Herrup EA, Warren HS, Trigilio J, Shin HS, Valentine C and Hellman J. MyD88-dependent and MyD88-independent pathways in synergy, priming, and tolerance between TLR agonists. J Immunol. 2007; 178:1164-1171.

123. Zhu Q, Egelston C, Gagnon S, Sui Y, Belyakov IM, Fox CB, Sivananthan SJ, Duthie MS, Haanen JB, Maio M, et al. Combined Nivolumab and Ipilimumab or Monotherapy in Untreated Melanoma. The New England journal of medicine. 2015; 373:23-34.
125. Tom JK, Mancini RJ and Esser-Kahn AP. Covalent modification of cell surfaces with TLR agonists improves & directs immune stimulation. Chem Commun. 2013; 49:9618-9620.

126. Delamarre L, Mellman I and Yadav M. Cancer immunotherapy. Neo approaches to cancer vaccines. Science. 2015; 348:760-761.

127. McCarthy DP, Hunter ZN, Chackerian B, Shea LD and Miller SD. Targeted immunomodulation using antigen-conjugated nanoparticles. Wires Nanomed Nanobi. 2014; 6:298-315.

128. Rosenthal JA, Chen LX, Baker JL, Putnam D and DeLisa MP. Pathogen-like particles: biomimetic vaccine carriers engineered at the nanoscale. Curr Opin Biotech. 2014; 28:51-58.

129. Speiser DE, Schwarz K, Baumgaertner P, Manolova V, Devevre E, Sterry W, Walden P, Zippelius A, Conzett KB, Senti G, Voelter V, Cerottini JP, Guggisberg D, Willers J, Geldhof C, Romero P, et al. Memory and Effector CD8 T-cell Responses After Nanoparticle Vaccination of Melanoma Patients. J Immunother. 2010; 33:848-858.

130. Makkouk A, Joshi VB, Wongrakpanich A, Lemke CD, Gross BP, Salem AK and Weiner GJ. Biodegradable Microparticles Loaded with Doxorubicin and CpG ODN for In Situ Immunization Against Cancer. The AAPS Journal. 2014; 17:184-193.

131. Roy A, Singh MS, Upadhyay P and Bhashkar S. Nanoparticle mediated co-delivery of paclitaxel and a TLR-4 agonist results in tumor regression and enhanced immune response in the tumor microenvironment of a mouse model. International Journal of Pharmaceutics. 2013; 445:171-180.

132. Chu KF and Dupuy DE. Thermal ablation of tumours: biological mechanisms and advances in therapy. Nat Rev Cancer. 2014; 14:199-208.

133. Lumnieczyk K and Safrany G. The impact of radiation therapy on the antitumor immunity: Local effects and systemic consequences. Cancer Lett. 2015; 356:114-125.

134. Formenti SC and Demaria S. Systemic effects of local radiotherapy. Lancet Oncol. 2009; 10:718-726.

135. Marrache S, Choi JH, Kundup S, Zaver D, Ham D and Dhar S. Immune stimulating photoactive hybrid nanoparticles for metastatic breast cancer. Integrative Biology. 2012; 215-223.

136. Kourtis IC, Hirose S, de Titta A, Kontos S, Stegmann T, Hubbell JA and Swartz MA. Peripherally administered nanoparticles target monocytic myeloid cells, secondary lymphoid organs and tumors in mice. PloS one. 2013; 8:e61646.

137. Azarin SM, Yi J, Gower RM, Aguado BA, Sullivan ME, Goodman AG, Jiang EJ, Rao SS, Ren Y, Tucker SL, Backman V, Jeruss JS and Shea LD. In vivo capture and label-free detection of early metastatic cells. Nature communications. 2015; 6:8094.

138. Huang BN, Abraham WD, Zheng YR, Lopez SCB, Luo SS and Irvine DJ. Active targeting of chemotherapy to disseminated tumors using nanoparticle-carrying T cells. Sci Transl Med. 2015; 7.

139. Sagiv-Barfi I, Kohlt HE, Burckhardt L, Czerwinska DK and Levy R. Ibrutinib enhances the antitumor immune response induced by intratumoral injection of a TLR9 ligand in mouse lymphoma. Blood. 2015; 125:2079-2086.

140. Liu H, Kwong B and Irvine DJ. Membrane anchored immunostimulatory oligonucleotides for in vivo cell modification and localized immunotherapy. Angewandte Chemie. 2011; 50:7052-7055.

141. Atkins MB, Lotze MT, Dutcher JP, Fisher RI, Weiss G, Margolin K, Abrams J, Szno M, Parkinson D, Hawkins M, Paradise C, Kunkel L and Rosenberg SA. High-dose recombinant interleukin 2 therapy for patients with metastatic melanoma: Analysis of 270 patients treated between 1985 and 1993. J Clin Oncol. 1999; 17:2105-2116.

142. Smith FO, Downey SG, Klapper JA, Yang JC, Sherry RM, Royal RE, Kammula US, Hughes MS, Restifo NP, Levy CL, White DE, Steinberg SM and Rosenberg SA. Treatment of metastatic melanoma using interleukin-2 alone or in conjunction with vaccines. Clin Cancer Res. 2008; 14:5610-5618.

143. Park J, Wrzesinski SH, Stern E, Look M, Criscione J, Ragheb R, Jay SM, Demento SL, Agawu A, Licona Limon P, Ferrandino AF, Gonzalez D, Habermann A, Flavell RA and Fahmy TM. Combination delivery of TGF-beta inhibitor and IL-2 by nanoscale liposomal polymeric gels enhances tumour immunotherapy. Nature materials. 2012; 11:895-905.

144. Burrell RA, McGranahan N, Bartek J and Swanton C. The causes and consequences of genetic heterogeneity in cancer evolution. Nature. 2013; 501:338-345.

145. Meacham CE and Morrison SJ. Tumour heterogeneity and cancer cell plasticity. Nature. 2013; 501:328-337.

146. Stevanovic S. Identification of tumour-associated T-CELL epitopes for vaccine development. Nat Rev Cancer. 2002; 2:514-520.

147. Yamshchikov GV, Mullins DW, Chang CC, Ogino T, Thompson L, Presley J, Galavotti H, Aquila W, Deacon D, Ross W, Patterson JW, Engelhard VH, Ferrone S and Stingluff CL. Sequential Immune Escape and Shifting of T Cell Responses in a Long-Term Survivor of Melanoma. The Journal of Immunology. 2005; 174:6863-6871.

148. Schumacher TN and Schreiber RD. Neoantigens in cancer immunotherapy. Science. 2015; 348:69-74.

149. Neller MA, Lopez JA and Schmidt CW. Antigens for cancer immunotherapy. Semin Immunol. 2008; 20:286-295.

150. Schreiber RD, Old LJ and Smyth MJ. Cancer Immunoeediting: Integrating Immunity’s Roles in Cancer Suppression and Promotion. Science. 2011; 331:1565-1570.
J, Weinschenk T, Modrusan Z, Mellman I, Lill JR and Delamarre L. Predicting immunogenic tumour mutations by combining mass spectrometry and exome sequencing. Nature. 2014; 515:572-576.

152. Gubin MM, Zhang XL, Schuster H, Caron E, Ward JP, Noguchi T, Ivanova Y, Hindal J, Arthur CD, Krebber WJ, Mulder GE, Toebes M, Vesely MD, Lam SSK, Korman AJ, Allison JP, et al. Checkpoint blockade cancer immunotherapy targets tumour-specific mutant antigens. Nature. 2014; 515:577-581.

153. Castle JC, Kreiter S, Diekmann J, Lower M, Van de Roemer N, de Graaf J, Selmi A, Diken M, Boegel S, Paret C, Koslowski M, Kuhn AN, Britten CM, Huber C, Tuerecu O and Sahin U. Exploiting the Mutanome for Tumor Vaccination. Cancer Research. 2012; 72:1081-1091.

154. Prasad S, Cody V, Saucier-Sawyer JK, Saltzman WM, Sasaki CT, Edelson RL, Birchall MA and Hanlon DJ. Polymer nanoparticles containing tumor lysates as antigen delivery vehicles for dendritic cell-based antitumor immunotherapy. Nanomedicine. 2011; 7:1-10.

155. Gross BP, Wongrakpanich A, Francis MB, Salem AK and Norian LA. A Therapeutic Microparticle-Based Tumor Lysate Vaccine Reduces Spontaneous Metastases in Murine Breast Cancer. Aaps J. 2014; 16:1194-1203.

156. Damo M, Wilson DS, Simeoni E and Hubbell JA. TLR-3 stimulation improves anti-tumor immunity elicited by dendritic cell exosome-based vaccines in a murine model of melanoma. Scientific reports. 2015; 5:17622.

157. Lohr JG, Adalsteinsson VA, Cibulsikis K, Choudhury AD, Rosenberg M, Cruz-Gordillo P, Francis JM, Zhang CZ, Shalek AK, Satija R, Trombetta JJ, Lu D, Tallapragada N, Tahirova N, Kim S, Blumenstiel B, et al. Whole-exome sequencing of circulating tumor cells provides a window into metastatic prostate cancer. Nat Biotechnol. 2014; 32:479-U202.

158. Sarioglu AF, Aceto N, Kojic N, Donaldson MC, Zeinali M, Hamza B, Engstrom A, Zhu H, Sundaresan TK, Miyamoto DT, Luo X, Bardia A, Wittner BS, Ramaswamy S, Shioda T, Ting DT, et al. A microfluidic device for label-free, physical capture of circulating tumor cell clusters. Nat Methods. 2015; 12:685-691.

159. Park GS, Kwon H, Kwak DW, Park SY, Kim M, Lee JH, Han H, Hoo S, Li XS, Lee JH, Kim YH, Lee JG, Yang W, Cho HY, Kim SK and Kim K. Full Surface Embedding of Gold Clusters on Silicon Nanowires for Efficient Capture and Photothermal Therapy of Circulating Tumor Cells (vol 12, pg 1638, 2012). Nano Lett. 2012; 12:2176-2176.

160. Fierer JO, Veggiani G and Howarth M. SpyLigase peptide-peptide ligation polymerizes affibodies to enhance magnetic cancer cell capture. P Natl Acad Sci USA. 2014; 111:E1176-E1181.

161. Li P, Mao ZM, Peng ZL, Zhou LL, Chen YC, Huang PH, Truica CI, Drabick JJ, El-Deiry WS, Dao M, Suresh S and Huang TJ. Acoustic separation of circulating tumor cells. P Natl Acad Sci USA. 2015; 112:4970-4975.

162. Halo TL, McMahon KM, Angeloni NL, Xu YL, Wang W, Chinen AB, Malin D, Strekalova E, Cryns VL, Cheng CH, Mirkin CA and Thaxton CS. NanoFlares for the detection, isolation, and culture of live tumor cells from human blood. P Natl Acad Sci USA. 2014; 111:17104-17109.