Enhancing Efficacy of Anticancer Vaccines by Targeted Delivery to Tumor-Draining Lymph Nodes

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Abstract

The sentinel or tumor-draining lymph node (tdLN) serves as a metastatic niche for many solid tumors and is altered via tumor-derived factors that support tumor progression and metastasis. tdLNs are often removed surgically, and therapeutic vaccines against tumor antigens are typically administered systemically or in non-tumor-associated sites. Although the tdLN is immune-suppressed, it is also antigen experienced through drainage of tumor-associated antigens (TAA), so we asked whether therapeutic vaccines targeting the tdLN would be more or less effective than those targeting the non-tdLN. Using LN-targeting nanoparticle (NP)-conjugate vaccines consisting of TAA-NP and CpG-NP, we compared delivery to the tdLN versus non-tdLN in two different cancer models, EG7-OVA lymphoma (expressing the nonendogenous TAA ovalbumin) and B16-F10 melanoma. Surprisingly, despite the immune-suppressed state of the tdLN, tdLN-targeting vaccination induced substantially stronger cytotoxic CD8+ T-cell responses, both locally and systemically, than non–tdLN-targeting vaccination, leading to enhanced tumor regression and host survival. This improved tumor regression correlated with a shift in the tumor-infiltrating leukocyte repertoire toward a less suppressive and more immunogenic balance. Nanoparticle coupling of adjuvant and antigen was required for effective tdLN targeting, as nanoparticle coupling dramatically increased the delivery of antigen and adjuvant to LN-resident antigen-presenting cells, thereby increasing therapeutic efficacy. This work highlights the tdLN as a target for cancer immunotherapy and shows how its antigen-experienced but immune-suppressed state can be reprogrammed with a targeted vaccine yielding antitumor immunity. Cancer Immunol Res; 2(5); 436–47. © 2014 AACR.

Introduction

As cancer remains a leading cause of death worldwide, active research is ongoing to develop new treatments that could replace or supplement classical cancer therapies, which include surgery, radiotherapy, and chemotherapy. Immunotherapies, which aim to activate the immune system to kill cancer cells, include strategies to increase the frequency or potency of antitumor T cells, to overcome suppressive factors in the tumor microenvironment, and to reduce T-cell suppression systemically (1–3). Despite promising results from clinical trials (4, 5), such immunotherapies can induce deleterious side effects in healthy nontargeted tissues (2, 3, 6). Furthermore, immunotherapy efficacy is often dampened by tumor-induced immune suppression, which occurs at the primary tumor site and extends to the tumor-draining lymph node (tdLN), thus allowing tumors to evade immune surveillance (6–8). Prophylactic vaccines exist against viruses that can drive certain types of cancer, such as human papillomavirus and hepatitis B virus (9). Therapeutic cancer vaccines, however, remain scarce because of the many challenges involved in breaking tolerance, permeating immunosuppressive barriers in the microenvironment, and attacking a heterogeneous tumor population with antigenic drift, just to name a few (2, 6, 7).

Targeting vaccines to dendritic cells (DC) with appropriate costimulation is crucial for the induction of antigen-specific cytotoxic CD8+ T cells and Th1 cells, which are required to mount an effective antitumor response and to kill tumor cells (7, 10–13). Strategies based on delivering adjuvants or antigens carried by nanoparticles or liposomes have led to improved targeting of LNs, activation of LN-resident DCs, and enhanced induction of antigen-specific CD8+ T cells, hence better protection in tumor challenge studies (14–19). Ultrasmall, functionalizable nanoparticles (NP) developed in our laboratory effectively target DCs in skin-draining LNs upon intradermal delivery (20, 21). These nanoparticles lead to antigen cross-presentation by DCs and to enhanced cytotoxic antigen-specific CD8+ T-cell immunity when coupled with an antigen or an adjuvant (22–24).

LNs could be a strategic target for vaccine delivery because of their role in initiating adaptive immunity. Tumor-derived...
factors and tumor-associated antigens (TAA), along with tumor-educated regulatory T cells (Treg) and myeloid-derived suppressor cells (MDSC), infiltrate the tdLN where they tolerate naïve T cells and prevent DC maturation, while minimally, or not at all, affecting non-tdLNs (8, 25, 26). In this fashion, the tdLN is an immune-privileged site that is remotely controlled and modified by the upstream tumor (25, 27).

Although surgical removal of the tdLN along with the primary tumor is common practice in patients with breast cancer and melanoma (28, 29), targeting the tdLN with a cancer vaccine could take advantage of TAA-primed leukocytes in the tdLN and might enhance therapeutic efficacy and patient survival.

We asked whether specifically targeting the tdLN, which is already TAA primed but also drains immunosuppressive tumor-derived factors, with nanoparticle vaccines would be advantageous or disadvantageous for its efficacy as compared with targeting a naïve non-tdLN. To address this question, we conjugated either the model TAA ovalbumin (OVA) or the endogenous melanoma TAA tyrosinase-related protein-2 (TRP-2) and the TLR9 ligand CpG DNA to nanoparticles separately, and we therapeutically treated E.G7-OVA or B16-F10 tumor-bearing mice with this cancer vaccine. We first determined the therapeutic superiority of nanoparticle-conjugated TAA and CpG over their soluble form. Then, by therapeutic intradermal delivery of our nanoparticle vaccine, we found that targeting the tdLN, as opposed to the non-tdLN, elicited the strongest antitumor response in terms of tumor growth, survival, and effector CD8+ T-cell response in both tumor models, and led to modulation of tumor-infiltrating immunosuppressive cells. Our findings demonstrate that although the tdLN is more immune suppressed than a non-tdLN, it is also TAA primed and can mount a more potent antitumor immune response than the non-tdLN. Thus, anticancer vaccines that target specifically the tdLN may be more effective, and at lower doses, than vaccines delivered at other sites.

Materials and Methods

Reagents

Chemicals were reagent grade and purchased from Sigma-Aldrich. Cpg-B 1826 oligonucleotide (5’-TCCATGACGTTCCT-GACGT-3’), 5’ SPO3-CpG, and 5’ SPO3-CpG-NH2 3’ were purchased from Microsynth. Low endotoxin grade OVA (<0.01 EU/μg protein) was from Hyglos; OVA grade V and OVA257–264 peptide (SIINFEKL) were purchased from Sigma-Aldrich and GenScript, respectively. OVA, Alexa Fluor 488 conjugate, was purchased from Life Technologies. Endotoxins were removed with the EndoTrap red 10 kit from Hyglos and used according to the manufacturer’s instructions. TRP-2 180-186 Peptide (SYVDFFVVL) was purchased from AnaSpec, and C-TRP-2 180–186 (CSVYDFVVL) was purchased from GenScript. DY633 NHS ester was purchased from Dyomics, and Alexa Fluor 647 C2 maleimide was from Life Technologies.

Mice and cell lines

C57BL/6 female mice, ages 8 to 12 weeks, were obtained from Harlan. All experiments were performed with approval from the Veterinary Authority of the Canton de Vaud, Switzerland, according to Swiss law. E.G7-OVA (CRL-2113), EL-4 (TIB-39), and B16-F10 (CRL-6475) cells were obtained from the American Type Culture Collection and cultured according to the instructions. All cell lines were checked for Mycoplasma; no additional authentication was performed.

Tumor inoculation and immunizations

Mice were anesthetized with isoflurane (5% for induction and 2% for maintenance) and their backs were shaved. A total of 106 E.G7-OVA or EL-4 cells were injected in 30 μL of 0.9% saline solution intradermally on the left side of the back of each mouse. After 4, 7, or 11 days, mice were immunized with 10 μg of OVA and 1 μg of CpG (vaccine) in a dose of 30 μL intradermally in the front footpad ipsilaterally (ipsi) or contralaterally (contra) to the tumor. Mice received a boost 7 days after the immunization. Blood was sampled from the submandibular vein of the cheek pouch with a 4-mm lancet on the day of boost and then every 7 days thereafter.

For TRP-2 studies, 2.5 × 107 B16-F10 melanoma cells were injected intradermally on one side of the back such that lymphatic drainage from the tumor only flowed to the brachial LN (i.e., the tdLN) on one side. Mice were immunized on days 4, 7, 11, and 15 with either 40 μg TRP-2 and 4 μg CpG in all four footpads, or with 10 μg NP-TRP-2 plus 1 μg NP-CpG or 10 μg free CpG in the front footpad either ipsi or contra to the tumor. Blood was sampled on day 10.

Tumors were measured with a digital caliper starting 4 days after inoculation, and volumes (V) were calculated as ellipsoids (V = π/6 × length × width × height). Mice were sacrificed when tumor volumes reached 1 cm3, as required by Swiss law.

Tissue and cell preparation

Spleens, brachial LNs, and tumors were harvested at the time of sacrifice. LNs and tumors were digested 30 and 90 minutes, respectively, in Dulbecco’s Modified Eagle Medium (DMEM) supplemented with 1 mg/mL collagenase D (Roche). Single-cell suspensions were obtained by gently disrupting the organs through a 70-μm cell strainer. Red blood cells were lysed with NH4Cl (ammonium chloride). Cells were counted and resuspended in Iscove’s Modified Dulbecco’s Medium (IMDM) supplemented with 10% FBS and 1% penicillin/streptomycin (full medium; all from Life Technologies). Serum was collected and stored at −20°C. Whole LN cell suspensions were plated in a black 96-well plate, and fluorescence was read on a plate reader.

For real-time PCR (RT-PCR), tumors were snap-frozen and homogenized in Lysis Matrix D tubes (MP Biomedicals). RNA was extracted with RNAqueous kit (Life Technologies) according to the manufacturer’s instructions. OVA primers (forward, TCAACCAAATCACCAAACC; reverse, CCAAGCCTGCTGCTTACAG) were purchased from Microsynth.

Ex vivo restimulation

A total of 2 × 106 spleen or 0.5 × 106 LN cells were plated in 96-well plates and cultured in full medium for 6 hours at 37°C in the presence of 1 μg/mL SIINFEKL peptide or 100 μg/mL OVA protein. Of note, 5 μg/mL brevetoxin A (Sigma-Aldrich) was added to the culture for the last 3 hours of culture before analyzing cells by flow cytometry.
Other methods

Nanoparticle synthesis, conjugation, and labeling, flow cytometry, ELISA, and immunostaining are described in Supplementary Materials and Methods.

Statistical analysis

Statistically significant differences between experimental groups were determined by one-way ANOVA followed by Bonferroni posttest correction with Prism software (v5, GraphPad). Of note, * and ** indicate P values less than 0.05 and 0.01, respectively; n.s., not significant.

Results

The tdLN is enlarged and immune suppressed, but contains tumor antigen–primed T cells

Tumor-draining and nondraining brachial LNs were isolated from E.G7-OVA tumor-bearing mice at different time points of growth and analyzed by immunofluorescence and flow cytometry. Although we found that the tdLN was generally inflamed, as seen by size enlargement when compared with a non-tdLN (Fig. 1A), the overall distribution of different lymphocytes and the frequencies of CD11b+ MHCI+ macrophages (Mφ), CD11c+ MHCI+ DCs, and CD11b– MHCI– immature myeloid cells were not affected by the upstream tumor 11 days after tumor inoculation (Fig. 1B). Proportionally, there were more CD200– B cells and fewer CD3+ T cells present in the tdLN than in the non-tdLN (Fig. 1B). Overall, the tdLN contained fewer CD8– CD11b– cross-presenting DCs both as percentage of total live cells (Fig. 1C) and of all DCs (Fig. 1D) as compared with the non-tdLN. DCs in the tdLN expressed more PD-L1 than in the non-tdLN (Fig. 1E). Consistent with this observation, CD8+ T cells expressed higher levels of PD-1 in the tdLN (Fig. 1F). As tumors progressed, more TAA-specific CD8+ T cells infiltrated the tdLN, while the levels in the non-tdLN remained unchanged (Fig. 1G), as determined by SIINFEKL–MHCI pentamer staining. Upon ex vivo restimulation, CD8+ T cells in the tdLN secreted more IFN-γ, TNF-α, and interleukin (IL)-2 than CD8+ T cells isolated from the non-tdLN (Fig. 1H). Taken together, these results show that while being inflamed and immune suppressed, the tdLN is tumor antigen primed and responds to tumor antigen, whereas the non-tdLN is not immune suppressed and not tumor antigen primed.

Nanoparticles target the tdLN

Nanoparticles developed by our group target skin-draining LNs and resident antigen-presenting cells (APC) upon intradermal administration (20, 21). To determine whether nanoparticles could be targeted to the tdLN harnessing lymphatic drainage, we polarized one side of the mouse as being tumor-draining by inoculating 10^6 E.G7-OVA tumor cells intradermally on one side of the back of mice, thus defining an ipsi tumor-draining side and the contra non–tumor-draining side (Supplementary Fig. S1A). After intratumoral injection of 30 μL of propidium iodide (PI) solution 7 days after inoculation, PI accumulated in the ipsi brachial LN, thus identifying it as the tdLN in our model (Supplementary Fig. S1B, left).

To assess LN targeting by nanoparticles, we injected fluorescently labeled nanoparticles (NP), where * is Alexa Fluor 647; Supplementary Materials and Methods) intradermally in the front footpads on the ipsi or contra side 7 days after tumor inoculation. Nanoparticles injected on the contra side accumulated significantly more in the contra brachial LN, whereas ipsi-injected NP* accumulated significantly more in the ipsi brachial LN (Supplementary Fig. S1B). Moreover, we found that contra-injected NP* targeted DCs, defined as CD11c+ MHCI+ in both the contra and ipsi brachial LNs, whereas ipsi-injected NP* targeted more the ipsi brachial and axillary LNs, being taken up by approximately 20% of the DCs resident in those LNs (Supplementary Fig. S1C). These data indicate that nanoparticles administered ipsi to the tumor accumulate in the tdLN and target resident DCs there, thereby providing an opportunity to modulate their behavior and potentially the tumor-associated immune responses.

Nanoparticle conjugation enhances cell targeting and therapeutic efficacy of OVA + CpG cancer vaccines

To explore the concept of antitumor vaccination targeted to the tdLN, we first engineered nanoparticle-conjugated antigen and adjuvant vaccines and explored their therapeutic efficacy in two different tumor models, namely the orthotopic and nonimmunogenic B16-F10 melanoma model and the E.G7-OVA model, a mouse lymphoma cell line (EL-4) that has been engineered to express OVA as model TAA.

We conjugated the TRP-2180–188 peptide, an endogenous and thus centrally tolerized TAA, to nanoparticles by adding a cysteine at the N-terminus of TRP-2180–188 (CSVYDFFWYL) to allow conjugation via a disulfide bond (Supplementary Materials and Methods). We used the TL99-ligand CpG as adjuvant for its Th1-skewing properties (12) and ability to boost antitumor immunity (30–32). B16-F10–bearing mice were immunized intradermally in all four footpads 4, 7, 11, and 15 days after inoculation with 40 μg TRP-2180–188 and 4 μg CpG conjugated onto nanoparticles (NP–TRP-2 + NP–CpG) or in soluble form (TRP-2 + CpG; Fig. 2A). The control group included nonimmunized mice. We found that nanoparticle-conjugated TRP-2 + CpG was therapeutically very efficient and superior to soluble TRP-2 + CpG, as seen by significantly smaller tumors and delayed tumor growth, as well as improved survival compared with mice immunized with soluble TRP-2 + CpG, which showed no improvement over control mice (Fig. 2B and C). Moreover, nanoparticle-immunized mice had approximately 5% circulating TRP-2180–188–specific CD8+ T cells 7 days after immunization, compared with 1% for mice immunized with TRP-2 and CpG in free form (Fig. 2D).

In the E.G7-OVA tumor model, conjugating the model antigen OVA to nanoparticles and codelivering it with NP-CpG was also therapeutically more beneficial than their soluble counterparts. E.G7-OVA tumor-bearing mice were ipsi immunized 4 days after inoculation and boosted 7 days later with 10 μg OVA and 1 μg CpG conjugated onto nanoparticles (NP–OVA + NP–CpG) or in soluble form (OVA + CpG; Fig. 2E). Control groups included nonimmunized tumor-bearing mice and NP-CpG–immunized mice (i.e., without antigen) to assess the contribution of innate immunity in the immune response to the vaccine. NP-OVA + NP-CpG significantly enhanced tumor regression compared with soluble OVA + CpG, as seen by
Figure 1. The tdLN is enlarged and immune suppressed, but contains tumor antigen-primed T cells. A total of 10^6 E.G7-OVA cells were inoculated intradermally on one side of the back such that tumor-draining and non-tumor-draining lymph nodes (tdLN and non-tdLN, respectively) could be compared at various time points. A, representative LN sections of tumor-bearing mice demonstrate enlargement of the tdLN compared with non-tdLN 14 days after tumor inoculation (white, lyve-1; red, CD3; green, B220); scale bar, 500 μm. B–H, characterization of LN-resident leukocytes 11 days after tumor inoculation. B, relative cell distributions show an increase in B-(B220^+T-cell (CD3^+)) ratios in the tdLN versus non-tdLN, while percentages of total leukocytes (CD45^+), as well as macrophages (CD11b^+ MHCII^+), DCs (CD11c^+ MHCII^+), and immature myeloid cells (CD11b^+ MHCII^-) were unchanged 11 days after tumor inoculation. C and D, cross-presenting DCs (CD8^+CD11b^+) as percentages of CD11c^+ MHCII^+ DCs (G) and total live cells (H). E, PD-L1 expression by mature DCs (CD11c^+ MHCII^-). F, PD-1 expression by CD8^+ T cells. G, relative numbers of LN-resident TAA-specific CD8^+ T cells as determined by SIINFEKL-MHCII pentamer staining on days 4, 7, and 11 after tumor inoculation. H, functionality of CD8^+ T cells after 6 hours restimulation as reflected by intracellular cytokine staining. Data reflect two independent experiments with 8 mice per group. *, P < 0.05; **, P < 0.01.

smaller tumor sizes and earlier onset of tumor regression (Fig. 2F and G). Moreover, NP-OVA + NP-CpG improved survival (Fig. 2H) and elicited 13% circulating OVA 257–264-specific CD8^+ T cells 7 days after immunization, whereas immunization with soluble OVA + CpG induced only 0.6% circulating OVA257–264-specific CD8^+ T cells (Fig. 2I). NP-CpG, lacking antigen, did not affect therapeutic outcomes versus nonimmunized mice.

By fluorescent labeling of OVA (with Alexa Fluor 488) and of CpG (with Dy633; Supplementary Materials and Methods), we found that therapeutic efficacy of nanoparticle-conjugated antigen and adjuvant relied on enhanced APC targeting in the LNs. Because we had determined that the ipsi brachial LN was the tdLN in our tumor model (Supplementary Fig. S1B), we chose the symmetric contra brachial LN as non-tdLN. Nanoparticle-conjugated or soluble labeled OVA and labeled CpG were injected ipsi or contra to the tumor 7 days after inoculation, and mice were sacrificed 24 hours later. Noninjected tumor-bearing mice were used as control for gating purposes. Nanoparticle conjugation led to significantly more co-uptake of OVA and CpG together by Mφ, while it also significantly enhanced the uptake by DCs in the tdLN after ipsi injection (Fig. 2J). Similarly, delivering labeled NP-OVA + NP-CpG contra significantly enhanced the uptake of OVA and CpG by Mφ in the non-tdLN (Fig. 2K) compared with OVA + CpG. These results show a clear advantage of nanoparticle conjugation for OVA and CpG targeting to APCs in the tdLN.

The tdLN is a more effective vaccine target site than the non-tdLN for a tumor antigen, but a less effective site for a nontumor antigen

Although we found that the tdLN was tumor antigen primed but immune suppressed, we asked whether a tdLN or a non-tdLN would induce a stronger immune response to an
Figure 2. Nanoparticle conjugation enhances cell targeting and therapeutic efficacy of OVA + CpG cancer vaccines. Responses of B16-F10 (A–D) and E.G7-OVA (E–K) tumor-bearing mice to therapeutic vaccination are shown. Vaccination schedule (A), B16-F10 tumor volumes (B), overall survival rates (C), and circulating TRP-2180–188-specific CD8+ T cells (D) 11 days after tumor inoculation (as determined by TRP-2180–188-MHCI pentamer staining). Vaccination schedule (E), E.G7-OVA tumor volumes (F), percentage of mice with tumor shrinkage after vaccination (G), overall survival rates (H), and circulating OVA257–264-specific CD8+ T cells (I) on days 11 and 18. J and K, on day 7 after tumor inoculation, E.G7-OVA tumor-bearing mice were injected intradermally with Alexa Fluor 488-labeled OVA and DY633-labeled CpG, either free or nanoparticle conjugated, in the front footpad either ipsi or contra to the tumor, and brachial LNs were analyzed 24 hours later. Nanoparticle conjugation leads to better targeting of OVA and CpG by mature macrophages (Mφ, MHCIiCD11b+) and DCs (MHCIiCD11c+) in the tdLN after ipsi injection (J) and in the non-tdLN after contra injection (K). Data reflect two independent experiments with 8 mice per group. *, P < 0.05; **, P < 0.01; ***, P < 0.001.
irrelevant antigen. We immunized EL-4- and E.G7-OVA tumor-bearing mice with NP-OVA + NP-CpG ipsi or contra to the tumor and compared the immune response in both tumor models (Fig. 3A). Although EG7-OVA tumors of immunized mice regressed as compared with control mice, EL-4 tumors were not affected by the vaccine 11 days after tumor inoculation (Fig. 3B). In the EG7-OVA model, ipsi immunization with a TAA was more beneficial than contra immunization, as seen by tumor volumes and circulating OVA257–264-specific CD8+ T cells 11 days after tumor inoculation (Fig. 3B and C). However, contra immunization with NP-OVA + NP-CpG in the EL-4 tumor model led to more OVA257–264-specific CD8+ T cells in the blood than ipsi immunization (Fig. 3C). Taken together, these data indicate that targeting a tumor-irrelevant vaccine to a non-tdLN, which is not immune suppressed, is more beneficial than targeting a tdLN.

**Therapeutic vaccines targeting the tdLN are more effective than those targeting the non-tdLN, but only after tumor growth has begun**

After validation of nanoparticle-conjugated OVA and TRP-2 as effective therapeutic cancer vaccines, we used them as a tool to assess whether targeting the tdLN with a TAA is therapeutically different from targeting a non-tdLN. EG7-OVA tumor-bearing mice were immunized ipsi or contra to the tumor with NP-OVA + NP-CpG at different stages of tumor growth, namely on day 4, 7, or 11 after inoculation. Blood was sampled 7, 14, and 21 days after immunization, and cells were stained with a SIINFEKL–MHC-I-specific pentamer.

Both ipsi and contra day 4 vaccines induced regression of early-stage tumors. Ipsi immunization led to significantly smaller tumors on days 9 to 13 (Fig. 4A, top) and slightly enhanced survival (Fig. 4A, middle) compared with contra immunization. As seen by averaged results, ipsi delivery of day 4 vaccine induced significantly more circulating OVA257–264-specific CD8+ T cells than contra delivery 7 days after immunization (Fig. 4A, bottom).

Targeting the tdLN of established tumors (day 7 psi) led to significantly smaller tumors on days 13 to 16 (Fig. 4B, top) compared with targeting the non-tdLN (day 7 contra). Of note, ipsi immunization on day 7 led to 100% survival (Fig. 4B, middle), as no tumors grew back after regressing, and to significantly more circulating OVA257–264-specific CD8+ T cells than contra delivery 7 days after immunization, with a 5-fold increase in OVA257–264-specific CD8+ T cells for ipsi vaccine compared with contra vaccine (Fig. 4B, bottom). At the latest immunization time point (day 11), ipsi injection of NP-OVA + NP-CpG afforded tumor regression of 1-cm3 tumors, whereas contra delivery did not (Fig. 4C, top). Consequently, ipsi-immunized mice had significantly improved survival (Fig. 4C, middle). Although initially offering no advantage over contra vaccination in the blood, day 11 ipsi immunization led to approximately 15% antigen-specific CD8+ T cells in the blood 14 days after immunization (Fig. 4C, bottom). NP-OVA + NP-CpG-induced OVA257–264-specific CD8+ T cells with an activated, functional phenotype, i.e., CD44+, CD62L−, PD-1+/−, CTLA-4−, and LAG-3+ (Supplementary Fig. S2A). Finally, it is interesting to note that ipsi immunization with NP-OVA + NP-CpG also led to enhanced production of anti-OVA IgG antibodies 3 weeks after the day 4 vaccine compared with contra immunization (Supplementary Fig. S2B).

These findings were corroborated in the B16-F10 melanoma model with the peptide TRP-2180–188 as TAA. Tumor-bearing mice were immunized with 10 µg NP-TRP-2 + 1 µg NP-CpG intradermally in the front footpad ipsi or contra to the tumor. Ipsi immunization resulted in significantly smaller tumors than those in control mice and led to improved survival compared with contra-immunized mice (Fig. 4D and E). Furthermore, ipsi immunization led to over 5% circulating TRP-2180–188-specific CD8+ T cells 7 days after immunization, as opposed to 2.5% in contra-immunized mice (Fig. 4F). Even when comparing with immunization of melanoma-bearing mice with a higher dose of free CpG (10 µg, a dose that we cannot easily achieve with nanoparticle-conjugated CpG)
along with NP-TRP-2 (10 μg as usual), the benefit of targeting the tdLN compared with the non-tdLN with a NP-TAA vaccine is evident (Supplementary Fig. S3A and S3B). The observed tumor regression and humoral responses, consistent both in the E.G7-OVA and B16-F10 models, highlight the advantage of delivering cancer vaccines to the tdLN for improved therapeutic antigen-specific tumor outcomes.

**Targeting a nanoparticle vaccine to the tdLN enhances the effector CD8^+ T-cell response locally and systemically**

With a cancer vaccine, induction of an antigen-specific effector immune response locally in the tdLN and especially in the tumor is key to obtaining tumor regression. To address this question and to understand the mechanisms behind the previous findings, we sacrificed mice at the peak of the vaccine-
and tumor regression (Fig. 4). Ipsi immunization led to specific CD8+ T-cell responses 7 days after vaccination in E.G7-OVA tumor-bearing mice immunized with 10 µg NP-OVA + 1 µg NP-CpG ipsi or contra to the tumor. A, vaccination schedule. B, representative flow cytometry plots of SIINFEKL−MHCII pentamer staining of tumor-infiltrating CTLs (CD4+ CD62L− effector CD8+ T cells). C, proportion of OVA327−264−specific CTLs (CD44+ CD62L− effector CD8+ T cells) in the tumor, spleen, tdLN, and non-tdLN as determined by SIINFEKL−MHCII pentamer staining. D, tumor-infiltrating OVA327−264−specific CD8+ T cells with an effector (CD44+ CD62L−) versus an exhausted (PD-1+/C3/C3) phenotype. E–H, cells from spleens and LNIs were restimulated ex vivo with SIINFEKL (1 µg/mL) for 6 hours before staining for intracellular cytokines and analysis by flow cytometry. E, representative flow cytometry plots of spleen CD8+ T cells stained for IFN-γ+ and TNF-α+. Values in the dot plots represent the percentage of CD8+ T cells in each gate. Proportion of IFN-γ+, IFN-γ− TNF-α+, and IFN-γ− TNF-α− IL-2− cytotoxic CD8+ T cells in the spleen (F), tdLN (G), and non-tdLN (H) of ipsi, contra, and control mice after SIINFEKL restimulation. Data reflect two independent experiments with 6 mice per group. *, P < 0.05; **, P < 0.01.

Figure 5. Targeting a nanoparticle vaccine to the tdLN enhances the effector CD8+ T-cell response locally and systemically. CD8+ T-cell responses 7 days after vaccination in E.G7-OVA tumor-bearing mice immunized with 10 µg NP-OVA + 1 µg NP-CpG ipsi or contra to the tumor. A, vaccination schedule. B, representative flow cytometry plots of SIINFEKL−MHCII pentamer staining of tumor-infiltrating CTLs (CD44+ CD62L− effector CD8+ T cells). C, proportion of OVA327−264−specific CTLs (CD44+ CD62L− effector CD8+ T cells) in the tumor, spleen, tdLN, and non-tdLN as determined by SIINFEKL−MHCII pentamer staining. D, tumor-infiltrating OVA327−264−specific CD8+ T cells with an effector (CD44+ CD62L−) versus an exhausted (PD-1+/C3/C3) phenotype. E–H, cells from spleens and LNIs were restimulated ex vivo with SIINFEKL (1 µg/mL) for 6 hours before staining for intracellular cytokines and analysis by flow cytometry. E, representative flow cytometry plots of spleen CD8+ T cells stained for IFN-γ+ and TNF-α+. Values in the dot plots represent the percentage of CD8+ T cells in each gate. Proportion of IFN-γ+, IFN-γ− TNF-α+, and IFN-γ− TNF-α− IL-2− cytotoxic CD8+ T cells in the spleen (F), tdLN (G), and non-tdLN (H) of ipsi, contra, and control mice after SIINFEKL restimulation. Data reflect two independent experiments with 6 mice per group. *, P < 0.05; **, P < 0.01.

induced T-cell response. Briefly, E.G7-OVA tumor-bearing mice were immunized with NP-OVA + NP-CpG 7 days after tumor inoculation ipsi or contra to the tumor and sacrificed 7 days later (Fig. 5A), which corresponded to the time point of peak number of circulating OVA327−264−specific CD8+ T cells and tumor regression (Fig. 4). Ipsi immunization led to enhanced infiltration of CD8+ T cells (Supplementary Fig. S4A) and OVA327−264−specific CD8+ T cells (Supplementary Fig. S4B) in the tumor. As seen by representative flow cytometry plots (Fig. 5B), ipsi NP-OVA + NP-CpG enhanced tumor-infiltrating OVA327−264−specific effector CD8+ T cells (defined as CD44+ CD62L−), with almost 70% of effector CD8+ T cells in the tumor being antigen-specific, compared with 40% for contra and control mice. Targeting the tdLN (ipsi) also induced significantly more OVA327−264−specific effector CD8+ T cells in the spleen and tdLN compared with contra immunization (Fig. 5C), which induced significantly more OVA327−264−specific effector CD8+ T cells in the non-tdLN than ipsi immunization (Fig. 5C). Similar results were obtained when all CD8+ T cells and effector CD8+ T cells were considered in the tumor, spleen, and tdLN (Supplementary Fig. S4B). Of note, tumor-infiltrating CD8+ and OVA327−264−specific CD8+ T cells expressed more effector markers (CD44+ CD62L−) and less PD-1 in ipsi-immunized mice than in contra-immunized and control mice (Supplementary Fig. S4C and Fig. 5D).

In a cancer vaccine, it is important that T cells be licensed to kill tumor cells and thus secrete cytotoxic factors when encountering cells expressing tumor antigens (7, 33). Cells from the spleens, tdLN, and non-tdLN were restimulated ex vivo with the full OVA protein or its immunodominant peptide
SIINFEKL for 6 hours and then stained for intracellular cytokines. As shown by representative flow cytometry plots and averaged results, targeting the tdLN with NP-OVA + NP-CpG (ipsi) led to 5% IFN-γ*, 2% IFN-γ* TNF-α*, and 1% triple-positive IFN-γ* TNF-α* IL-2* cytotoxic CD8* T cells in the spleen upon SIINFEKL restimulation, which are significantly more when compared with vaccines targeting the non-tdLN (contra; Fig. 5E and F). Similarly, ipsi immunization significantly more polyfunctional CD4+ T cells than contra immunization as indicated by OVA restimulation on cells from the spleens (Supplementary Figs. S5A and S5B). Delivering NP-OVA + NP-CpG to the tdLN (ipsi) induced significantly more IFN-γ* and TNF-α*–secreting cytotoxic CD8* T cells (Fig. 5G) and polyfunctional CD4* T cells (Supplementary Fig. S5C) in the tdLN compared with targeting the non-tdLN and control mice. Contra administration of NP-OVA + NP-CpG also led to a cytotoxic CD8* (Fig. 5H) and polyfunctional CD4* (Supplementary Fig. S5D) T-cell response in the non-tdLN, while ipsi delivery had no effect. These results illustrate the role of the tdLN in the induction of a potent effector T-cell response both systemically and locally in the tdLN and tumor, which is where tumor antigen recognition and tumor cell killing happen, respectively.

**Targeting the tdLN with nanoparticle-conjugated OVA and CpG reduces the frequencies of tumor-infiltrating MDSCs and Tregs**

Immunosuppressive cells, such as MDSCs and Foxp3+ CD4+ Tregs, contribute to tumor-induced tolerance (34, 35) and classically hinder cancer immunotherapies, making them an important target when designing cancer vaccines (7). Here, the goal was to assess whether targeting a nanoparticle-based vaccine to a tdLN could have an impact on the aforementioned cells locally in the tumor. To investigate this, we analyzed tumor-infiltrating cells from NP-OVA + NP-CpG–immunized mice (experiment setup as in Fig. 5A). Targeting the tdLN (ipsi)
led to a decrease in the proportion of tumor-infiltrating Gr1\textsuperscript{int}-expressing MDSCs (defined as Gr1\textsuperscript{+} CD11b\textsuperscript{+} MHCII\textsuperscript{−}), as shown by representative flow cytometry plots and averaged results (Fig. 6A and B). Gr1\textsuperscript{int} MDSCs represented 28% of tumor-infiltrating CD11b\textsuperscript{+} MHCII\textsuperscript{−} cells in ipsi-immunized mice, compared with 35% and 38% in contra-immunized and control mice, respectively. Moreover, not only was the proportion of Gr1\textsuperscript{int} cells among MDSCs reduced by ipsi immunization with NP-OVA + NP-CpG, the total number of Gr1\textsuperscript{int} MDSCs also decreased in the tumor (Fig. 6C), while leaving the Gr1\textsuperscript{hi}, Gr1\textsuperscript{lo}, and Gr1\textsuperscript{−} cells unaffected compared with contra-immunized and control mice. Interestingly, Gr1\textsuperscript{int} MDSCs were recently shown to be the most immunosuppressive MDSCs in tumor-bearing mice (36). Furthermore, we observed that significantly more MHCII\textsuperscript{−}-expressing CD11b\textsuperscript{+} Gr1\textsuperscript{+} MDSCs infiltrated the tumors of tdLN-targeted (ipsi) mice compared with control and non-tdLN-targeted (contra) mice, suggesting maturation of MDSCs (Fig. 6C, left). Indeed, one promising approach to inhibit the immunosuppressive ability of MDSCs has been the induction of their maturation into APCs (37). Targeting the tdLN (ipsi) also led to a decrease in tumor-infiltrating Foxp3\textsuperscript{+} CD4\textsuperscript{+} Tregs (Fig. 6D), and to a more immunogenic, less suppressive Tregs to effector T-cell ratio (Fig. 6E). The proportions of MDCs and Tregs decreased locally in the tumor microenvironment but were left unaffected by the nanoparticle vaccine systemically as seen in the spleen (Fig. 6F and G). Taken together, these data suggest that targeting the tdLN with a therapeutic cancer vaccine can also affect the tumor microenvironment locally by decreasing the number of immunosuppressive cells, both MDSCs and Tregs.

**Discussion**

In this study, we explored the use of a therapeutic nanoparticle-particle cancer vaccine platform to target the antigen-experienced, yet immune-suppressed tdLN of tumor-bearing mice. We found that this strategy was substantially more beneficial in terms of inducing a potent adaptive cellular immune response, both enhancing the numbers of antigen-specific CTLs and reducing the numbers of suppressive MDSCs and Tregs in the tumor, which correlated with reduction in tumor volume and prolongation of survival. Results from this study indicate that antigen-experience outweighs immune suppression in the tdLN, an immune organ whose role in clinical oncology should be reconsidered as it may play a useful role in facilitating cancer vaccination.

The tdLN is usually regarded as a site with prevailing immune suppression and tolerance to the upstream tumor (25, 27). We found that with tumor progression, the tdLN contained more OVA\textsubscript{257-264}-specific CD8\textsuperscript{+} T cells and that tdLN-resident CD8\textsuperscript{+} T cells were more responsive to antigen stimulus than those in a non-tdLN as seen by cytokine secretion, but that the tdLN was also enlarged and immune suppressed as seen by enhanced PD-1 and PD-L1 expression by T cells and DCs, respectively (Fig. 1; ref. 2). We hypothesized that targeting a nanoparticle-based vaccine to the tdLN could break immune suppression and boost already TAA-primed immune cells in the tdLN.

Upon intradermal injection, sufficiently small nanoparticles take advantage of interstitial flow into the dermal lymphatics to efficiently drain through lymphatic vessels and target skin-draining LNs (21). Although intratumor, intra-LN, or subcutaneous administrations have been explored to target LNs with cancer immunotherapies, our experimental model permitted passive but direct and efficient targeting of nanoparticles from the skin to the tdLN (Supplementary Fig. S1; refs. 17, 31, 38). Our nanoparticles were in the range of 30 nm, thus smaller than particles used in many vaccination studies, hence affording enhanced lymphatic drainage and targeting of the skin-draining LNs (14–19).

To address our hypothesis, we first engineered nanoparticle cancer vaccines by conjugating the endogenous peptide TRP-2\textsubscript{190-198} and the nonendogenous model antigen OVA to the surface of nanoparticles and codevolved them with NP-CpG. Nanoparticle conjugation led to the induction of more antigen-specific CD8\textsuperscript{+} T cells and stronger therapeutic outcomes, benefits that relied on enhanced dual targeting of APCs in the LNs (Fig. 2). Compared with current published studies, our TRP-2-based therapeutic nanoparticle vaccine is very potent at the low doses used for antigen and adjuvant (18, 39, 40). Similarly, other studies have shown improved therapeutic outcomes by either conjugating the adjuvant or the antigen onto or into delivery systems, such as poly(lactide-co-glycolide) particles and liposomes, but, to our knowledge, this is the first study to show enhanced co-uptake of both adjuvant and antigen by LN-resident APCs after nanoparticle conjugation (14–19). Furthermore, it has been shown recently that encapsulating antigens or TLR agonists in nanoparticles enhanced humoral responses compared with immunization with soluble antigen or adjuvant (41, 42).

By comparing delivery of our model cancer vaccine with the tdLN versus a non-tdLN, we found that targeting the tdLN afforded significantly improved tumor reduction and survival both in the E.G7-OVA and the aggressive B16-F10 melanoma mouse models (Fig. 4 and Supplementary Fig. S3). This was associated with a stronger antigen-specific effector immune response, in terms of cytokotoxic CD8\textsuperscript{+} T cells and polyfunctional CD4\textsuperscript{+} Th1 cells, both systemically and locally in the tumor and the tdLN (Fig. 5 and Supplementary Fig. S5). Furthermore, efficient LN targeting by nanoparticles and conjugation of CpG onto nanoparticles allowed a very low dose of CpG (1 \mu g/mouse) to be effective, while typically much higher doses of free CpG (8–100 \mu g) are needed for effective adjuvant function (14–16, 18). Our results showed that the vaccine-induced local immune response in the tdLN translated to greater tumor reduction than the one elicited in a non-tdLN. This complements our previous study using adjuvant targeting alone; specifically, we demonstrated that NP-CpG or NP-paclitaxel targeted to the tdLN could partially reverse the immunosuppressed milieu there to slow tumor growth (43). In that study, however, the therapeutic benefit of adjuvant alone was very moderate compared with the present study with TAA-targeting vaccines (e.g., as shown in Fig. 4).

The very high number of circulating OVA\textsubscript{257-264}-specific CD8\textsuperscript{+} T cells (up to 30%; Fig. 4B) induced by targeting the vaccine to the tdLN led to regression of early-stage tumors and
more impressively of tumors having reached the Swiss legal limit of 1 cm³. In the literature, cancer vaccines are usually administered when the tumors are 5 to 7 mm in diameter or even before they are detectable, and also do not usually induce this high number of TAA-specific CD8⁺ T cells in the blood and the LNs (17, 19, 30, 38, 44), thus emphasizing the benefits of nanoparticle-mediated targeting of antigen and adjuvant to DCs in the tdLN (18, 38, 45). We noted, however, that after regressing some tumors grew back. OVA-transfected tumor cells can lose OVA expression over time (46, 47), a trend that we also observed by RT-PCR analysis of OVA expression at days 4, 7, and 11 after tumor inoculation (Supplementary Fig. S6); the loss of TAA expression may explain why some tumors grew back afterregressing.

Finally, targeting the tdLN with nanoparticle-conjugated OVA and CpG affected the tumor microenvironment by reducing the frequency of immunosuppressive cells, including MDSCs and Tregs, while leaving those cells unaffected systematically (Fig. 6). While we previously showed that tumor-infiltrating monocytic MDSCs are NP⁺, delivering our model cancer vaccine to the tdLN decreased the number of tumor-resident Gr1⁺MDSCs, which are described as monocytic and the most immunosuppressive subset, and led to MDSC maturation, thus making them less suppressive (21, 36, 48). CpG has been shown to affect the phenotype of MDSCs, especially monocytic MDSCs, in tumor-bearing mice, but at much higher CpG doses than we used here (32, 49). We believe that our findings may be due to an effect of CpG on MDSCs and to a change in the cellular and cytokine environment in the tumor following immunization. We noted a decrease in the frequency of Tregs and a subsequent reduction in the ratio of Tregs to effector CD8⁺ T cells, suggestive of a benefit in antitumor response (50, 51). These findings indicate that the model cancer vaccine is shifting the balance from a tolerogenic to an immunogenic microenvironment in the tumor. Similarly, other studies have found that Tregs infiltrated the tdLN less when targeted with a cellular therapy in tumor-bearing mice (44).

Here, we show that the tdLN may be capable of playing a special role for the induction of a strong TAA-specific immune response using TAA vaccines, its antigen experience outweighing its immune suppressed state. Our results also show the advantage of using nanocarriers that are sufficiently small to drain to the LNs and to deliver adjuvants and antigens to resident APCs for effective induction of immune responses; this drainage enabled tdLN targeting. Finally, on the basis of our results in tumor models with both exogenous and endogenous antigens, we believe that delivering cancer immunotherapies to the tdLN is more effective than targeting them to a non-tdLN. In conclusion, this study shows the important role played by the tdLN in the induction of a potent antigen-specific immune response to a nanoparticle-based cancer vaccine; our results show that the benefits of TAA experience in the tdLN outweigh the detriments of exposure to immunosuppressive cytokines produced by the tumor.

Disclosure of Potential Conflicts of Interest
J.A. Hubbell and M.A. Swartz have ownership interest (including patents) in Locat Bio Sarl. No potential conflicts of interest were disclosed by the other authors.

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