Oncolytic vesicular stomatitis virus quantitatively and qualitatively improves primary CD8⁺ T-cell responses to anticancer vaccines

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Introduction

An outstanding challenge to successful immunotherapy is the development of antigen-specific vaccination regimens that can promote the generation of sufficient CD8⁺ T cell numbers for the treatment of infectious diseases and cancer. One promising approach to circumvent this problem involves prime-boost regimens using heterologous viral vectors that express the vaccination target. Indeed, numerous prior studies have shown that heterologous prime-boosting approaches generate larger amounts of antigen-specific CD8⁺ T cells than one-component vaccines and homologous boosting strategies.¹⁻³ However, there is a limited availability of viral vectors that can be effectively used to boost viral vaccines to achieve an optimal expansion of CD8⁺ T cells. Vectors that can rapidly elicit strong primary and secondary immunological responses to allow for optimal control of progressing diseases are thus highly desirable for clinical application, yet remain poorly characterized.

Replication-incompetent, E1- and E3-deleted recombinant human type 5 adenovirus-based vectors are promising vaccines currently undergoing clinical evaluation. Recombinant adenoviral vectors are indeed highly effective in priming naïve T cells against transgenes, resulting in robust and antigen-specific protection against tumor or viral challenges.¹⁻³ More importantly, adenoviral vectors have been shown to elicit effective immune responses even in individuals with pre-existing adenovirus-specific immunity, making them promising immunizing agents for the development of vaccines that are currently unavailable or unsatisfactory.⁷⁻¹⁰

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The ability of heterologous prime-boost vaccination to elicit robust CD8⁺ T cell responses has been well documented. In contrast, relatively little is known about how this immunotherapeutic strategy impacts the functional qualities of expanded T cells in the course of effector and memory responses. Using vesicular stomatitis virus (VSV) as a boosting vector in mice, we demonstrate that a massive secondary expansion of CD8⁺ T cells can be achieved shortly after priming with recombinant adenoviral vectors. Importantly, VSV-boosted CD8⁺ T cells were more potent than those primed by adenoviruses only, as measured by cytokine production, granzyme B expression, and functional avidity. Upon adoptive transfer, equivalent numbers of VSV-expanded CD8⁺ T cells were more effective (on a per-cell basis) in mediating antitumor and antiviral immunity than T cells only primed with adenoviruses. Furthermore, VSV boosting accelerated the progression of expanded CD8⁺ T lymphocytes to a central memory phenotype, thereby altering the effector memory profile typically associated with adenoviral vaccination. Finally, the functional superiority of VSV-expanded T cells remained evident 100 d after boosting, suggesting that VSV-driven immunological responses are of sufficient duration for therapeutic applications. Our data strongly support the choice of VSV as a boosting vector in prime-boost vaccination strategies, enabling a rapid amplification of CD8⁺ T cells and improving the quality of expanded T cells during both early and late immunological responses.
A significant limitation of immunotherapeutic adenoviral vectors is that they elicit prolonged effector T (T_{eff}) cell and effector memory T (T_{em}) cell responses\textsuperscript{11–13} that require long intervals for an efficient boost. This is mainly due to the killing of migratory antigen-presenting dendritic cells (DCs) by cytolytic T_{eff} cells in the periphery, which prevents their engagement with central memory T (T_{cm}) cells.\textsuperscript{14–16} Seeking to enhance the duration and potency of adenovirus-elicited immune responses, we have recently demonstrated that the intravenous administration of recombinant vesicular stomatitis virus (VSV) can provoke massive secondary expansion at the height of the adenovirus-primed CD8\textsuperscript{+} T-cell response.\textsuperscript{17} Furthermore, in an aggressive melanoma tumor model (B16), we showed that a strong primary immune response to adenovirus-based vaccines was essential for delaying tumor growth, creating a sufficient window to allow VSV boosting to further reduce the growth of—or even eradicate—neoplastic lesions.\textsuperscript{17} Since rapid boosting can be achieved without the need to compromise the T_{eff} population induced by primary immunization, our data suggest that recombinant adenoviruses and VSV can be combined to create a powerful therapeutic prime-boost vaccine.

Although VSV is an effective immunizing agent, it is also oncolytic agent. Thus, the relative contribution of these distinct biological properties towards the overall activity of VSV-based vaccines is difficult to discern, especially in the context of therapeutic (as opposed to prophylactic) settings.\textsuperscript{18} A better understanding of how VSV boosting impacts the functionality of expanded T cells may therefore be gained by investigating this approach in models of infectious disease, which may also broaden its clinical applicability. In the present study, we performed a detailed analysis of T-cell phenotype, cytokine secretion and immunological memory responses in tumor-free animals immunized with adenovirus-based vectors alone or in combination with VSV boosting. Furthermore, we compared the ability of these T cells (on a per cell basis) to protect animals against tumor or viral challenges. We report here that CD8\textsuperscript{+} T-cell responses from VSV-boosted mice are more potent than in mice that received adenovirus alone. More importantly, the superiority of VSV-expanded T cells was evident not only at the peak of secondary responses but was also 70 d after boosting, suggesting that the majority of antigen-specific CD8\textsuperscript{+} T cells elicited by VSV-boosting display a memory phenotype. Our data strongly support that the therapeutic use of recombinant VSV as a boosting vector in prime-boost vaccination strategies enables rapid and massive amplification of CD8\textsuperscript{+} T cells while improving their functional qualities at both early and late time points.

**Results**

Quantitative assessment of adenovirus-primed and VSV-boosted CD8\textsuperscript{+} T-cell responses. To compare adenovirus-primed and VSV-boosted CD8\textsuperscript{+} T cells, we reasoned that it would be best to coordinate the vaccinations so that peak expansion of primary and secondary epitope-specific CD8\textsuperscript{+} T-cell responses would coincide. To determine when these peaks occurred, mice were either primed with adenoviral vectors coding for human dopachrome tautomerase (Ad-hDCT) only, or primed with Ad-hDCT and then boosted (14 d later) with hDCT-coding VSV (VSV-hDCT). The dose and injection route of these vectors (i.e., 10\textsuperscript{6} plaque-forming units (PFUs) i.m. for Ad-hDCT, and 10\textsuperscript{9} PFUs i.v. for VSV-hDCT) had been previously optimized.\textsuperscript{17} As measured by the percentage of interferon-γ (IFN-γ)-expressing CD8\textsuperscript{+} T cells, kinetic analyses revealed that the peaks of the primary and secondary responses against the DCT\textsubscript{180–188} epitope occurred about 11 d upon administration of Ad-hDCT (Fig. 1A) and 5 d upon that of VSV-hDCT (Fig. 1B), respectively. Of note, VSV-boosting significantly increased the magnitude of the response (~4.2% vs. ~25% IFN-γ-expressing CD8\textsuperscript{+} T cells, Figure 1A and B).

To evaluate the next phase of T cell-mediated immunity, we extended our observations for 2 weeks after the peak response (designated as "day 0"), confirming our previous finding that adenoviruses typically produce a CD8\textsuperscript{+} T-cell response with a protracted contraction phase (Fig. 1C). Although the frequency of VSV-expanded CD8\textsuperscript{+} T cells remained significantly higher than that of T cells developing in mice receiving Ad-hDCT only with during this 2-week time frame, VSV boosting did not appear to alter the occurrence or duration of the contraction phase (Fig. 1C). To determine if this phenomenon was unique to self-antigens, like hDCT, we repeated the experiment by substituting the transgene in both viral vectors with an immunodominant epitope derived from chicken ovalbumin (OVA), SIINFEKL (OVA\textsubscript{257–264}). Notably, SIINFEKL-specific T cells accounted for more than 22% of total circulating CD8\textsuperscript{+} T cells at the peak of adenoviral priming, a primary response that was increased to more than 80% by VSV boosting (Fig. 1C). Irrespective of both the enhanced magnitude of the response and the nature of the antigen, the protracted nature of the contraction phase remained unaltered (Fig. 1D).

To determine whether VSV could amplify CD8\textsuperscript{+} T-cell responses even at early time points, we boosted mice over shorter intervals with VSV-hDCT from 4 to 14 d after Ad-hDCT priming. In this setting, the frequency of transgene-specific CD8\textsuperscript{+} T cells was remarkably enhanced by boosting with VSV within 4 d of adenoviral priming (Fig. 2A). Also, boosting 7 d after priming was as effective as doing so 14 d thereafter, demonstrating the unique capacity of VSV to accelerate the kinetics of CD8\textsuperscript{+} T-cell responses. We also observed that VSV boosting was effective in the same time frame (i.e., 14 d after primary immunization) in mice primed with hDCT\textsubscript{180–188}-pulsed DCs (DC/SVY) or vaccinia viruses (VV) expressing SIINFEKL (VV/SIIN) (Fig. 2B and C), indicating that the remarkable boosting potency of VSV is not limited to specific antigens or priming methods. Interestingly, however, the magnitude of boosting correlated with the intensity of primary responses (please compare Figure 2A with 2B and Figure 1D with 2C), indicating that the combination of efficient adenoviral vectors with VSV represents a robust prime-boost strategy to achieve therapeutically relevant immune responses.

VSV-boosted CD8\textsuperscript{+} T cells produce more cytokines and granzyme B than those primed with adenoviruses. Although the ability of VSV to rapidly and quantitatively amplify T_{eff}
responses elicited by adenoviral vaccines was highly attractive and unique, its impact on the qualitative function and memory development of adenovirus-primed T cells remained to be determined. As an initial approach to these incognita, we assessed the cytokine secretion profile of hDCT180–188- and SIINFEKL-specific cells developing in mice that had been primed with adenovirus as a standalone immunotherapeutic intervention or in combination with VSV boosting at various time points. Compared with primary T cells (i.e., cells developing in response to adenoviral priming), a higher frequency of secondary T cells (i.e., cells developing upon adenoviral priming and VSV boosting) could simultaneously produce IFN\(\gamma\) and tumor necrosis factor \(\alpha\) (TNF\(\alpha\)) (Fig. 3A), regardless of the nature of the immunizing epitope. VSV-boosted cells also produced more IFN\(\gamma\) (Fig. 3B) and TNF\(\alpha\) (Fig. 3C) than adenovirus-primed cells on a per-cell basis. The dynamics of the response specifically showed that the amount of IFN\(\gamma\) produced by VSV-boosted T cells generally plateaued starting from 1-week post-peak (Fig. 3B), but the proportion of cells producing TNF\(\alpha\) (Fig. 3A) and the amount of TNF\(\alpha\) per cell (Fig. 3C) were still increasing at the final 2-week time point. In addition to enhanced cytokine production, CD8+ T cells from VSV-boosted mice also exhibited substantially increased expression levels of granzyme B than those from mice subjected to adenoviral priming only, particularly at the peak of response (Fig. 3D).

VSV boosting enhances the functional avidity of CD8+ T cells. In light of our observations that secondary T cells had a superior cytokine secretion profile, we hypothesized that VSV boosting selectively induces the expansion of T-cell subsets that exhibit the highest avidity within the adenovirus-primed cell pool. To test this hypothesis, we assayed the functional avidity of CD8+ cells isolated from mice that were primed with either

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**Figure 1.** Quantitative assessment of adenovirus-primed and VSV-boosted CD8+ T cell responses. (A) C57BL/6 mice were primed by intramuscular injection of 1 x 10^8 plaque-forming units (PFUs) of Ad-hDCT. Blood samples were drawn on various days post-priming to quantify CD8+ T-cell response to the immunodominant epitope DCT180–188 by cytofluorometric detection of intracellular interferon \(\gamma\) (IFN\(\gamma\)) upon in vitro stimulation with the cognate peptide in the presence of brefeldin A. Data were pooled to plot the kinetics of the response. (B) Mice primed with Ad-hDCT were boosted after a 14-d interval by intravenous injection of 1 x 10^9 PFUs of VSV-hDCT. Transgene-specific CD8+ T-cell responses were measured in the blood at various days post-boosting to establish the kinetics of secondary responses. (C and D) Mice were primed by intramuscular injection of 1 x 10^8 PFUs of Ad-hDCT or Ad-SIINFEKL. Half of these mice were then boosted by intravenous injection of 1 x 10^9 PFUs of VSV-hDCT or VSV-SIINFEKL. The vaccinations were offset such that the peak of primary transgene-specific CD8+ T-cell responses in mice subjected to priming only (11 d post-priming) coincided with the peak of secondary responses in VSV-boosted animals (5 d post-boosting). The frequency of circulating CD8+ T cells specific for DCT180–188 (C) and SIINFEKL (D) was quantified by flow cytometry in terms of IFN\(\gamma\)-expressing cells upon in vitro antigenic stimulation. In all cases, \(n = 5\) animals/group; data are reported as means ± SEM and are representative of two experiments. \(\triangleright\) adenovirus-induced primary response; \(\triangledown\) adenovirus-primed, VSV-boosted secondary response. Statistical significance was determined by two-way ANOVA: **\(p < 0.01\), ***\(p < 0.001\).

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Ad-hDCT or Ad-SIINFEKL alone or in combination with a heterologous boost VSV-hDCT or VSV-SIINFEKL, respectively. At multiple time points, circulating lymphocytes were harvested and stimulated with serial dilutions of cognate peptides in the presence of brefeldin A (to trap cytokines in the cytoplasm of responding T cells). T-cell responses to each concentration of peptide were monitored by flow cytometry in terms of the percentage of IFN-γ+ CD8+ T cells relative to that induced by the highest peptide concentration for each immunogen. In this assay, high T-cell avidities are defined by a large proportion of T cells that can respond to target peptides at low concentrations. Both at the peak of the immune response (Figure 4A) and 1-week later (Fig. 4B), VSV-boosted T cells were of higher avidity than adenovirus-primed cells upon exposure to both hDCT180-188 (Fig. 4, left panels) and SIINFEKL (Fig. 4, right panels). Interestingly, however, by 2-weeks post-peak, the average avidity of primary and secondary T cells was equivalent (Fig. 4C).

VSV-boosted CD8+ T cells are functionally superior in vivo to those primed by adenoviruses only. Having determined that VSV-boosted CD8+ T cells display higher avidity and produce more cytokines than adenovirus-primed cells, we sought to determine if these characteristics translated into enhanced immunotherapeutic activity in vivo, in appropriate tumor and viral challenge models. We selected SIINFEKL as the target epitope for this experiment because an almost 14-fold increase in transgene-specific T cells can be achieved with SIINFEKL as compared with DCT180-188 (Fig. 1), significantly reducing the number of donor animals required. Also, we excluded quantitative differences as a confounding factor by using flow cytometry to sort transgene-specific CD8+ T cells from mice primed with either Ad-SIINFEKL alone or subjected to a prime-boost regimen involving Ad-SIINFEKL and VSV-SIINFEKL. CD8+ T cells purified from each group of mice were then adoptively transferred into naïve recipients in equal amounts (4.5 × 105 cells/mouse). Negative control animals received the same number of CD8+ T cells obtained from mice that had been sequentially treated with empty adenoviral (Ad-BHG) and VSV (VSV-MT) vectors. Transferred SIINFEKL-specific CD8+ T cells were allowed to home to their target tissues for 24 h prior to the administration of the immunogen.

In order to test the utility of heterologous VSV boosting against both malignant and infectious diseases, we chose to test two different challenge models. Mice first received 5 × 10^5 OVA-expressing B16 (B16-OVA) murine melanoma cells. Twenty-one days after challenge, mice were euthanized and lung metastases were enumerated. Transferring SIINFEKL-targeting CD8+ T cells from Ad-SIIN-primed donor mice effectively decreased the frequency of metastases (relative to CD8+ T cells derived from Ad-BHG-primed donor mice), confirming that adenoviruses efficiently prime protective CD8+ T-cell responses. As expected, VSV-boosted T cells reduced the proportion of metastases even further (Fig. 5A; p < 0.0001). These observations provide compelling evidence that VSV boosting indeed improves the cytotoxic function of adenovirus-primed CD8+ lymphocytes.

**Figure 2.** VSV enables ultra-rapid boosting that is not limited to a single antigen or priming method. (A-C) C57BL/6 mice were primed by intramuscular injection of 1 × 10⁸ plaque-forming units (PFUs) of Ad-hDCT (A) subcutaneous injection of 1 × 10⁶ dendritic cells pulsed with the DCT180-188 Peptide (DC/SVY) (B), or subcutaneous injection of 1 × 10⁹ PFUs of VV-SIINFEKL (VV-SIIN). Four to 14 (A) or 14 (B and C) days later, mice were boosted by intravenous injection of 1 × 10⁹ PFUs of VSV-hDCT (A and B) or VSV-SIINFEKL (VV-SIIN) (C). Notes: In all cases, n = 5 animals/group; data are reported as means ± SEM and are representative of two experiments.
at least in the context of antitumor immunity. To extend the relevancy of our findings to the treatment of infectious diseases, we assayed the relative efficacy of adenovirus-primed vs. adenovirus-primed and VSV-boosted CD8+ T cells in a model of viral challenge. Following adoptive T-cell transfer, mice were challenged i.p. with 1 × 10^6 PRUs of a recombinant VV engineered to express SIINFEKL (VV-SIINFEKL). Immunizations were scheduled so that the peaks of primary or secondary transgene-specific CD8+ T-cell responses coincided. Circulating antigen-specific T cells were identified by flow cytometry upon staining of intracellular interferon γ (IFNγ) and tumor necrosis factor α (TNFα) after in vitro stimulation with cognate peptides in the presence of brefeldin A. (A) Multi-cytokine producing transgene-specific CD8+ T cells were defined as those capable of simultaneously producing IFNγ and TNFα. The amount of IFNγ (B) and TNFα (C) produced on a per cell basis was also evaluated. (D) Primary and secondary SIINFEKL-specific CD8+ T cells identified by tetramer staining were fixed, permeabilized and then assessed for granzyme B expression by intracellular immunostaining followed by cytofluorometric analysis. In all cases, n = 5 animals/group; data are reported as means ± SEM and are representative of two experiments. *p < 0.05, **p < 0.01, ***p < 0.001.

Figure 3. VSV-boosted CD8+ T cells produce more cytokines and granzyme B than those primed with adenoviruses only. (A–D) C57BL/6 mice were primed by intramuscular injection of 1 × 10^8 plaque-forming units (PFUs) of Ad-hDCT or Ad-SIINFEKL, and part of them boosted 14 d later by intravenous injection of 1 × 10^9 PFUs of VSV-hDCT or VSV-SIINFEKL. Immunizations were scheduled so that the peaks of primary or secondary transgene-specific CD8+ T-cell responses coincided. Circulating antigen-specific T cells were identified by flow cytometry upon staining of intracellular interferon γ (IFNγ) and tumor necrosis factor α (TNFα) after in vitro stimulation with cognate peptides in the presence of brefeldin A. (A) Multi-cytokine producing transgene-specific CD8+ T cells were defined as those capable of simultaneously producing IFNγ and TNFα. The amount of IFNγ (B) and TNFα (C) produced on a per cell basis was also evaluated. (D) Primary and secondary SIINFEKL-specific CD8+ T cells identified by tetramer staining were fixed, permeabilized and then assessed for granzyme B expression by intracellular immunostaining followed by cytofluorometric analysis. In all cases, n = 5 animals/group; data are reported as means ± SEM and are representative of two experiments. *p < 0.05, **p < 0.01, ***p < 0.001.
We observed a dominance shift from effector (T\(_{\text{EFF}}\)) to memory (comprising T\(_{\text{EM}}\) and T\(_{\text{CM}}\)) cells 70 d after the peak of responses (Fig. 6B). Of particular interest, VSV boosting exaggerated the predominance of CD127+ memory T cells as compared with adenovirus priming only (Fig. 6B). We further analyzed DCT-specific CD8+ T cells 113 d post-peak to elucidate the relative contribution of T\(_{\text{EM}}\) and T\(_{\text{CM}}\) cells towards the increased abundance of memory T lymphocytes. Consistent with previous observations,\(^6,11\) adenoviral immunization mainly resulted in the generation of T\(_{\text{EM}}\) (tetramer+CD127+CD62L−) cells (Fig. 6C). By contrast, the vast majority of the memory compartment in VSV-boosted animals (at this time point) displayed a central memory phenotype (tetramer+CD127+CD62L+), suggesting that VSV boosting does not only increase the abundance and promote the early cytotoxic functions of adenovirus-primed T\(_{\text{EM}}\) cells, but also influences the frequency and phenotype of memory responses to priming.

Finally, to directly determine the functionality of adenovirus-primed or VSV-boosted T cells well beyond their peak responses, we purified primary or secondary T cells from SIINFEKL-immunized mice 70 d post-peak and adoptively transferred them in equal amounts (4.5 × 10\(^5\) cells/mouse) into distinct naïve hosts. These mice were intravenously challenged with 5 × 10\(^5\) B16-OVA cells 24 h after T-cell transfer. Similar to what was observed when T cells derived from peak immune responses were transferred (Fig. 5A) and in line with our previous findings,\(^6,11\)
Figure 5. VSV-boosted CD8+ T cells are functionally superior to T cells subjected to adenoviral priming only. (A-B) Donor C57BL/6 mice were primed by intramuscular injection of 1 × 10^8 plaque-forming units (PFUs) of Ad-SIINFEKL (Ad-SIIN), or primed and then boosted 14 d later by intravenous injection of 1 × 10^9 PFUs of Vsv-SIINFEKL (Vsv-SIIN). Immunizations were scheduled so that the peaks of primary or secondary transgene-specific CD8+ T-cell responses coincided. At the peak of these responses, 4.5 × 10^5 of either primary or secondary SIINFEKL-specific CD8+ T cells (4.5 × 10^5) was adoptively transferred into naïve C57BL/6 hosts. Control animals received an equal number of CD8+ T cells obtained from mice that had been treated with 1 × 10^8 PFUs of Ad-BHG i.m. followed by 1 × 10^9 PFUs of Vsv-MT i.v. (both are empty vectors). (A) Twenty-four hours after adoptive cell transfer, mice were challenged with 5 × 10^5 B16-OVa cells i.v.. Twenty-one days later, lung metastases, and representative images are shown. (B) Twenty-four hours after adoptive cell transfer, mice were challenged with 1 × 10^6 PFUs of Vv-SIINFEKL (VV-SIIN) i.p.. Three days later, VV titers were quantified in ovaries. Both these challenge experiments were repeated twice with n = 3–6 mice/group and yielded similar results. In all cases, data are reported as means ± SEM. Statistical significance was determined by one-way ANOVA.

fewer metastases developed in mice receiving adenovirus-primed T cells than in animals treated with control lymphocytes, suggesting that the therapeutic potential of adenoviral priming persists even 70 d after the peak response. In addition, VSV-boosted cells were significantly superior to adenovirus-primed cells in attenuating the metastatic spread of B16-OVA cells to the lung (Fig. 7; p = 0.0001). This provides firm experimental evidence indicating that the functional improvement of T cells achieved by VSV boosting is long lasting, presumably due to enhanced immunological memory.

Discussion

The success of immunotherapeutic vaccines is dependent upon sufficient elicitation of antigen-specific cytotoxic T lymphocyte-mediated cell death and immune responses that persist over time. Here, we report that the magnitude, cytotoxic function and memory phenotype of antigen-specific CD8+ T lymphocytes produced by adenoviral vaccines can be reshaped by VSV boosting, leading to dramatically enhanced CD8+ T-cell immunity in mouse models of cancer and viral infection. Combining adenoviruses and VSV into a prime-boost regimen can rapidly stimulate increased numbers of both effector and memory CD8+ T cells, which are both essential for improved clinical outcomes. These data have broad implications for the treatment of both cancer and infectious diseases, especially in therapeutic (as opposed to prophylactic) settings.

It has been previously reported that recombinant adenoviruses are effective vectors that can consistently foster robust CD8+ T cell immunity in multiple infection models as well as in patients affected by various diseases. CD8+ T cells induced by adenovirus-based vaccines exhibit unique characteristics amenable
to therapeutic use, including the preferential accumulation of T\textsubscript{EM} cells and their persistence at high numbers\textsuperscript{6,11,12}. These clinically relevant responses may potentiate immune protection against peripheral (e.g., skin, mucosal) infections and, by similar mechanisms, against cancer\textsuperscript{23–26}. Our results have confirmed this adenovirus-induced CD8\textsuperscript{+} T cell phenotype. Moreover, by adoptive cell transfer experiments, we have further evinced the robust immunotherapeutic functions of adenovirus-primed cells at both the early and late stages of the immune response.

Remarkably, VSV boosting can further magnify the high number of effector T cells that develop in response to adenoviral priming. Strikingly, this immunologically advantageous boosting effect can be achieved over a short interval (within 4–14 d upon priming) and offers an effective vaccination strategy for circumstances in which immediate protection is essential (e.g., malignancy or potentially pandemic infections).

Interestingly, VSV does not only increase the magnitude of adenovirus-primed CD8\textsuperscript{+} T-cell responses but also enhances the cytotoxic activity of these cells, as measured by increased cytokine production, granzyme B expression, and avidity for their targets.

**Figure 6.** VSV-boosted CD8\textsuperscript{+} T cells exhibit enhanced functional memory. (A-C) C57BL/6 mice were primed by intramuscular injection of 1 × 10\textsuperscript{8} plaque-forming units (PFUs) of Ad-hDCT or Ad-SIINFEKL, or primed and then boosted 14 d later by intravenous injection 1 × 10\textsuperscript{9} PFUs of VSV-hDCT or VSV-SIINFEKL. Immunizations were scheduled so that the peaks of primary or secondary transgene-specific CD8\textsuperscript{+} T-cell responses coincided. Antigen-specific T cells were identified in the blood by tetramer staining or cytofluorometric detection of intracellular interferon \(\gamma\) (IFN\(\gamma\)) upon in vitro stimulation with cognate peptides (DCT\textsubscript{180-188} or SIINFEKL) in the presence of brefeldin A. (A) Frequencies of transgene-specific cells on day 70 post-peak as measured by tetramer and intracellular cytokine staining (ICS). (B) Kinetics of CD127 expression on transgene-specific cells as measured by tetramer staining. (C) Proportion of DCT\textsubscript{180-188} tetramer-positive CD8\textsuperscript{+} T cells with effector (T\textsubscript{eff}), effector memory (T\textsubscript{EM}), and central memory (T\textsubscript{CM}) phenotypes at day 113 post-peak. Data are representative of two experiments. O adenovirus-primed, + VSV-boosted secondary response. Statistical significance in (A) and (C) was determined by one-way ANOVA, \(p\) values are reported. Statistical significance in (B) was determined by two-way ANOVA: *\(p < 0.05\), **\(p < 0.01\), ***\(p < 0.001\).
We observed that VSV-boosted T cells are functionally superior—on a per-cell basis (i.e., in equal amounts)—to adenovirus-primed T cells in mediating immune responses against either viral or tumor challenges. Improving the quality of T cells has been a problem in particular for the treatment of cancer or chronic infections, two pathological conditions in which T cells either intrinsically exhibit a low avidity for cognate antigens or, alternatively, become functionally exhausted upon prolonged antigenic stimulation in the context of an inflammatory milieu, such as it occurs in the tumor environment. Combining adenoviruses with VSV may be able to potentiate the magnitude as well as the quality of immune responses, both of which are essential for the successful outcome of vaccination in patients. The precise molecular mechanism by which VSV boosting enhances the quality of expanded CD8+ T lymphocytes remains unclear; however, we postulate that it may relate to antigen cross-presentation by DCs, which has been linked to the immunological profile of CD8+ T cells. In our experience, VSV appears to primarily infect follicular B cells while antigen presentation in this setting relies on DCs (Bridle and Wan, manuscript in preparation), suggesting that cross-presentation may constitute the predominant mechanism that mediates the increased immunological efficacy of VSV-boosted CD8+ T cells that we report here. VSV vectors possess self-replicating, cytopathic and oncogenic properties that may facilitate the cross-presentation of antigens released in the follicles upon cell death, leading to superior CD8+ T-cell responses. Furthermore, like other viral vectors, VSV generates a variety of “danger signals” that engage host Toll-like receptors (TLRs) and other pattern recognition receptors to activate the innate immune response. Specifically, it has been reported that ligation of TLR4 and/or TLR7/TLR8 on APCs can influence the phenotype of T cells, enhancing the generation of polyfunctional CD8+ T lymphocytes. We and others have previously shown that VSV infection stimulates DCs to produce large amounts of IFNα/β, IL-12 and IL-15, APC responses that may contribute to the functional improvement of expanding CD8+ T cells. It has been previously shown that adenoviral immunization favors the commitment of antigen-specific CD8+ T cells to the TEM lineage, mainly due to prolonged antigenic stimulation. In our hands, prolonged antigen presentation upon adenoviral vaccination is exclusively mediated by non-hematopoietic APCs that persistently stimulate existing memory cells. We surmise that the rapid VSV boosting of effector CD8+ T cells with a high functional avidity may clear such non-hematopoietic APCs, resulting in shortened antigen presentation, a situation that resembles the outcome of extinction of transgene expression. On the contrary, boosting with VSV over a short interval may prolong antigen presentation by conventional hematopoietic APCs, effectively accelerating CD8+ T-cell responses and the acquisition of a TCM phenotype.

Whether the quality or the magnitude of the immune response is a better predictor of the clinical efficacy of vaccination is an important and hitherto unresolved question for the field. It is likely that an optimal protection against individual pathogens relies on T cells with a distinct functional profile, while both the magnitude and quality of CD8+ T cells are required to effectively
control neoplastic diseases.69 Accumulating data highlight the potential efficacy of recombinant adenoviruses as highly versatile vectors that are capable of generating high numbers of CD8+ effector-like T cells in peripheral tissues to provide immediate immunological protection.20,48 Heterologous boosting with VSV offers a new strategy not only to improve the already desirable properties of memory T cells elicited by adenoviruses (notably their persistence at high levels and their Th1 phenotype) but also to increase the amounts of Tcm cells, a subset of CD8+ T cells that may be preferable in the context of certain pathogens. Thus, the administration of recombinant adenoviral vectors (priming) followed by that of VSV expressing the same antigenic epitope (boosting) represents a novel combinatorial approach to vaccination that elicits CD8+ T cells with optimal magnitude and phenotype at both effector and memory stages.

**Materials and Methods**

**Mice and cell cultures.** Age-matched (8–10 weeks old at initiation of each experiment) female C57BL/6 (haplotype H-2b) mice (Charles River Laboratories) were housed in a dedicated pathogen-free facility. Animal studies complied with Canadian Council on Animal Care guidelines and were approved by McMaster University’s Animal Research Ethics Board. B16-OVA cells, a murine melanoma line stably transfectected with an expression vector encoding full-length OVA, were cultured in MEM-F11 medium supplemented with 10% fetal bovine serum (FBS), 2 mM l-glutamine, 55 µM 2-mercaptoethanol, 1× vitamin solution and antibiotics. G418 (at the dose of 800 µg/mL) was used to maintain OVA expression. Vero cells were cultured in Eagle’s minimum essential medium containing Earle’s salts supplemented with 10% FBS, 2 mM l-glutamine and antibiotics. All cell culture reagents were from Life Technologies.

**Viruses.** The E1/E3-deleted, replication-deficient, recombinant human serotype 5 adenovirus and the replication-competent, recombinant vesicular stomatitis virus (VSV) with the methionine residue deleted at position 51 of the matrix protein (to abrogate its ability to inhibit Type I IFN responses) have been described previously.6,41 Ad-hDCT and VSV-hDCT vectors expressing the human melanoma-associated antigen dopachrome tautomerase (hDCT); Ad-SIINFEKL and VSV-SIINFEKL express the H-2d-restricted dominant CD8+ T-cell epitope OVA257-264 (SIINFEKL). Recombinant VVs expressing SIINFEKL (VV-SIINFEKL) has previously been described.6,50 Ad-BHG and VSV-MT were empty control vectors.

**Peptides.** The immunodominant peptide from DCT that binds to H-2Kb (DCT180–188, SVYDFFVWL; shared by human and murine DCT) was synthesized by PepScan Systems (Lelystad). The H-2Kb-restricted OVA-derived SIINFEKL peptide was synthesized by Biomer Technologies.

**Dendritic cell-based vaccine.** Murine bone marrow-derived DCs were generated in the presence of 40 ng/mL recombinant murine granulocyte macrophage colony-stimulating factor (GM-CSF; from PeproTech) for 7 d, as previously described,42 and then loaded with 1 µg/mL DCT180–188 for 4 h in the presence of 2 µg/mL lipopolysaccharide LPS (Sigma-Aldrich). To vaccinate mice, 5 × 10⁶ peptide-pulsed DCs were injected s.c. into each hind footpad (total dose = 1 × 10⁶ cells).

**Administration of viral vaccines.** Anesthetized mice were immunized by injection of 1 × 10⁸ PFUs of adenoviral vectors in 100 µL PBS (50 µL/hamstring) i.m., or 1 × 10⁸ PFUs of VV vectors in 200 µL PBS i.p. When appropriate, boosting was performed by injection of 1 × 10⁸ PFUs of VSV in 200 µL PBS i.v., into the tail vein.

**Antibodies and tetramers.** The following monoclonal antibodies were used in flow cytometry assays. Anti-CD16/CD32 (clone 2.4G2) antibodies were employed to block Fc receptors; anti-CD8 (clone 53–6.7), anti-CD62L (clone MEL-14) and anti-CD127 (clone SB/199) antibodies were used for cell-surface staining; and anti-IFNγ (clone XMG1.2), anti-TNFα (clone MP6-XT22) and anti-granzyme B (clone GB11) antibodies were employed for intracellular staining. All antibodies were from BD Biosciences. For the quantification of antigen-specific T cells, the following allophycocyanin (APC)-conjugated tetramers were used: Kb-DCT180–188-APC and Kb-OVA257–264-APC (MHC Tetramer Lab, Baylor College of Medicine).

**Intracellular cytokine staining.** To assess antigen-specific T-cell responses, blood was collected from the peri-orbital sinus into heparinized tubes and red blood cells were lysed. Cells were kept on ice during handling and the time from sample collection to the end of processing was less than 2 h. Only fresh cells were used in cytfluorometric assays. To this aim, cells were counted on an improved Neubauer hemocytometer and cell viability was ensured consistently greater than 90% (as assessed based on the exclusion of trypan blue). Mononuclear cells were stimulated with 1 µg/mL peptides (controls were exposed to irrelevant peptides at the same concentration) in RPMI medium supplemented with 10% FBS, 2 mM l-glutamine, antibiotics and 1 µg/mL brefeldin A (GolgiPlug, BD Biosciences, added after 1 h of incubation). During the 5 h total incubation time, Fc receptors were blocked with anti-CD16/CD32 antibodies and then cells were stained with fluorescent anti-CD8 antibody in PBS supplemented with 5% bovine serum albumin (BSA). Cells were then fixed/permeabilized with Cytofix/Cytoperm (BD Biosciences) and stained for the detection of intracellular cytokines. Data were acquired using a FACSCanto flow cytometer with the FACSDivadiva v.5.0.2 software (BD Biosciences) and analyzed with the FlowJo software (Tree Star).

**Functional avidity assays.** The functional avidity of antigen-specific CD8+ T cells was determined by intracellular cytokine staining, as described above, following stimulation with serial log-dilutions of peptides in vitro. Peptide concentration varied from 1,000 to 0.1 ng/mL. Data are expressed as percentages of the response to the maximal peptide concentration, calculated as follows: (% of CD8+ cells responding to a given concentration of peptides / % of CD8+ cells responding to the highest concentration of peptides) × 100.

**Tetramer staining.** The flow cytometry-assisted phenotyping of antigen-specific T cells for the expression of memory markers was accomplished using circulating lymphocytes stained with fluorochrome-conjugated tetramers and anti-CD8, anti-CD62L and anti-CD127 antibodies. The cytolytic potential...
of antigen-specific T cells was assessed by surface-staining with anti-CD8 antibodies and tetramers followed by fixation, permeabilization and intracellular staining for granzyme B.

Gating strategy for analyzing cytofluorometric data. Using forward vs. side scatter-width parameters (FSC vs. SSC), doublets were excluded from analyses and single lymphocytes were gated on. Single CD8+ cells were subsequently gated on using a histogram so that their staining with tetramers or intracellular cytokine-specific antibodies could be queried.

Adaptive T-cell transfer. Equal numbers of SIINFEKL-specific T cells were transferred into naïve recipient mice. To accomplish this, spleens from donor mice that has been previously immunized with adenoviral vectors alone or coupled to VSV boosting were harvested after 12 days post-priming (i.e., day 12 post-priming) or secondary (i.e., day 5 post-boosting) responses or 70 d post-pull. Aliquots from single splenocyte suspensions were labeled with tetramers and anti-CD8 antibodies so that the frequency of antigen-specific T cells could be quantified by flow cytometry. Equal doses of SIINFEKL-specific adenovirus-primed and VSV-boosted T cells were then prepared. The total amounts of splenocytes within each dose was equalized equal by taking up VSV-boosted T cells with splenocytes derived from naive mice. Preparations containing equal numbers of both antigen-specific T cells and total splenocytes were then injected i.v. into recipients. Controls received splenocytes from mice receiving Ad-BHG followed by VSV-MT.

Tumor and virus challenge models. Mice into which SIINFEKL-specific T cells had been adoptively transferred (see above) were assessed for their ability to clear OVA-expressing tumor cells and viruses. In both cases, adoptively transferred T cells were given 24 h to achieve homeostatic trafficking patterns. In the lung metastasis model, mice received 5 × 10^6 B16-OVA cells i.v. After 21 d, mice were euthanized and tumor nodules growing on the surface of the lungs were counted. When too many tumor nodules were present to obtain a reliable count, a value of 800 was assigned based on a semi-quantitative estimation. In the infection model, 1 × 10^6 PFU of VV-SIINFEKL were injected i.v.. After 3 d, mice were euthanized and VV titers in homogenized ovaries were determined by a plaque assay on Vero cells.

Statistical analyses. GraphPad Prism version 4.00 for Windows (GraphPad Software) was used for all graphing and statistical analyses. Differences in T-cell responses were analyzed for statistical significance by two-way ANOVA. Data from tumor and viral challenge experiments were analyzed for statistical significance by one-way ANOVA. If required, data were normalized by log transformation prior to statistical analyses. Differences between means were considered statistically significant when p < 0.05. Data are reported as means ± SEM.

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Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

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