The TNFR family members OX40 and CD27 link viral virulence to protective T cell vaccines in mice

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Induction of CD8+ T cell immunity is a key characteristic of an effective vaccine. For safety reasons, human vaccination strategies largely use attenuated nonreplicating or weakly replicating poxvirus-based vectors, but these often elicit poor CD8+ T cell immunity and might not result in optimal protection. Recent studies have suggested that virulence is directly linked to immunogenicity, but the molecular mechanisms underlying optimal CD8+ T cell responses remain to be defined. Here, using natural and recombinant vaccinia virus (VACV) strains, we have shown in mice that VACV strains of differing virulence induce distinct levels of T cell memory because of the differential use of TNF receptor (TNFR) family costimulatory receptors. With strongly replicating (i.e., virulent) VACV, the TNFR family costimulatory receptors OX40 (also known as CD134) and CD27 were engaged and promoted the generation of high numbers of memory CD8+ T cells, which protected against a lethal virus challenge in the absence of other mechanisms, including antibody and help from CD4+ T cells. In contrast, weakly replicating (i.e., low-virulence) VACV strains were poor at eliciting protective CD8+ T cell memory, as only the Ig family costimulatory receptor CD28 was engaged, and not OX40 or CD27. Our results suggest that the virulence of a virus dictates costimulatory receptor usage to determine the level of protective CD8+ T cell immunity.

Introduction
A desirable vaccine should elicit strong, long-lasting humoral and cellular immunity. While all human vaccines provide protection against disease via the generation of antibody responses, increasing attention is being focused on the idea that populations of memory CD8+ T cells are essential for protection from pathogens for which vaccine development has been unsuccessful thus far, such as HIV, malaria, tuberculosis, and herpes simplex virus. For safety reasons, many vaccination strategies use replication-incompetent, or replication-competent highly attenuated, poxvirus-based vectors. However, there is considerable literature acknowledging that highly replicative viruses are better at inducing CD8+ T cell responses. Animal and nonhuman primate studies have suggested that multiple dosing, or higher dosing, with replication-incompetent highly attenuated poxviruses is required to achieve T cell responses, and sometimes antibody responses, comparable to those elicited by replication-competent or highly virulent poxviruses (1–10). Although this is logically related to antigen load or the persistence of antigen to stimulate CD8+ T cells, the effect of reduced virulence on immunogenicity becomes a major issue when attempting to derive a truly effective vaccine that incorporates attenuated vectors.

Vaccinia virus (VACV) represents a relevant example of the importance of understanding how attenuated pathogens might behave as vaccines. Since the successful eradication of smallpox 3 decades ago, a large effort has been made to develop second- and third-generation highly attenuated poxvirus-based vectors as vaccine vehicles for other infectious diseases, including herpes simplex virus, SARS, influenza, HIV, tuberculosis, and malaria, as well as for cancer immunotherapy (11–13). These include modified VACV Ankara (MVA) (14), a variant of Lister strain LC16m8; ACAM1000/2000, derived from New York City Board of Health strain NYCBOH; the highly deleted Copenhagen VACV strain derivative NVVAC; and attenuated canarypox virus (ALVAC). Although some vectors given by certain routes might result in strong CD8+ T cell responses, the literature also suggests there is variability in the levels of CD8+ T cell immunity, raising the question of what determines induction of optimal CD8+ T cell responses (2, 7–10, 15–17). Using several clinically relevant natural and recombinant VACV variants, we show here a quantitative difference in CD8+ T cell immunity elicited depending on the virus and immunization route, with only the most virulent VACVs promoting protective populations of memory CD8+ T cells. This enhanced CD8+ T cell memory was dependent on OX40 (also known as CD134) and CD27, 2 of the many stimulatory receptors in the TNF receptor (TNFR) superfamily that can be expressed on T cells (18, 19). Importantly, OX40 and CD27 were only operative with a VACV strain that could replicate strongly; when higher doses of attenuated VACVs were used for inoculation; or when attenuated VACV was given via an immunization route that allowed for strong replication. In contrast, only CD28/B7 interactions, but not OX40/OX40L or CD27/CD70 interactions, were used in generating responses to VACVs that were cleared quickly; consequently, CD8+ T cell memory was substantially weaker and did not afford protection in the absence of CD4+ T cell help and antibody. These data provide an explanation for why poxvirus-based vaccines can result in good CD8+ T cell immunity and
provide a framework for understanding which molecular interactions to target when using attenuated vaccines to provide the best methodologies to vaccinate individuals against pathogen attack.

**Results**

*Virulence correlates with the generation of persistent VACV-specific CD8+ T cell populations.* Numerous VACV strains have been isolated, including the prototypical strain used in the laboratory, Western Reserve (WR), as well as others such as Lister (used as a smallpox vaccine in Europe) and the New York City Board of Health strain (NYCBOH) used as a smallpox vaccine in the United States (Dryvax, produced by Wyeth). All viruses are known to differ in expression of several virulence factors, although the exact extent of variation is not fully documented. For example, the VACV WR encodes B18R, which produces an IFN-I–binding protein that neutralizes this cytokine. Lister does not possess this gene, and NYCBOH produces a truncated B18R gene product that is inactive (20). Another difference is the activity targeting the IL-1β–converting enzyme encoded by the B13R gene that is found in WR, but is produced as a truncated protein in NYCBOH and not present in Lister (21). As a direct example of an engineered attenuated vaccine vector, we also used a mutant WR strain in which only the B18R gene was deleted (22), referred to herein as WR-B18R.

When infection was i.p., WR was slowly cleared, with a strong reduction in viral load only obvious between days 14 and 15, whereas NYCBOH and Lister were cleared by day 3 in the spleen and day 7 in the ovaries (Figure 1A). WR-B18R exhibited intermediate characteristics, with reduced titers in the spleen between days 3 and 4, and moderately lower titers compared with WR in the ovaries at day 7. To further demonstrate altered virulence between these strains, mice were exposed to i.n. virus at varying doses (Figure 1B). Whereas even a low dose of 10^4 PFU WR was not controlled well, resulting in severe weight loss, 10^6 PFU Lister, NYCBOH, or WR-B18R was not pathogenic, and little to no weight loss was observed. This was not dependent on the background of the host, as similar results were obtained after i.p. or i.n. infection of BALB/c mice (Supplemental Figure 1; supplemental material available online with this article; doi:10.1172/JCI42056DS1). Thus, in i.p. and i.n. infection models, the VACV’s WR-B18R, Lister, and NYCBOH were highly attenuated compared with the native VACV WR.

To assess induction of CD8+ T cell memory, the immunodominant VACV-reactive CD8+ T cell population was tracked with a tetramer of a peptide of B8R (23–25) or through the production of IFN-γ following stimulation with vaccinia epitopes. At 40 days after i.p. infection, the frequency of memory B8R tetramer–reactive (i.e., B8R+) CD8+ T cells (and CD8+IFN-γ+ T cells; data not shown) in both secondary lymphoid organs and peripheral organs such as the lung was 2- to 4-fold greater in mice infected with WR than in those infected with WR-B18R, Lister, or NYCBOH (Figure 2A). T cell avidity was also assessed by intracellular IFN-γ staining of freshly isolated splenocytes from WR- and Lister-infected mice. At 40 days after infection, memory CD8+ T cells were stimulated for 6 hours with graded concentrations of B8R, and cytokine-producing cells were analyzed by intracellular cytokine staining. As shown in Supplemental Figure 2, the amount of IFN-γ production (as measured by mean fluorescent intensity) on a per-cell basis was almost identical between WR- and Lister-infected groups over a wide range of peptide concentrations, similar to other assessments of IFN-γ (Figure 2, C and D). This suggests that exposure to varying levels of viral antigen primarily affects the frequency of CD8+ T cells elicited rather than the functional activity of individual cells.

Furthermore, mice exposed to virus via the i.n. route (Figure 1B) also generated 2- to 5-fold fewer lung-resident memory CD8+ T cells in response to WR-B18R, Lister, or NYCBOH, even when 30-fold more virus was inoculated (Figure 2B). This did not simply apply to the B8R+ population, but was observed when CD8+ T cells were examined reactive with peptides of A3L, A8R, and B2R. Notably, 540 days after i.p. infection, the frequency of persisting memory VACV-reactive CD8+ T cells in both spleen and lung was 3- to 10-fold greater in mice infected with WR compared with those infected with WR-B18R, Lister, or NYCBOH (Figure 2, C and D).

Analysis of the initial effector CD8+ T cell response showed that this difference was likely to be explained by altered molecular regulation during the primary infection, as the size of these populations of VACV-specific T cells correlated to an extent with the size of the memory T cell pool and, again, opposite to the rate of virus clearance (Figure 2, E and F). Therefore, the immune system generates greater numbers of VACV-reactive memory CD8+ T cells with virus that replicates most extensively and is not rapidly cleared, consequently exposing the immune system to viral antigens for extended periods.

*Selective use of OX40 to drive VACV-specific CD8+ T cells related to virulence.* We then assessed whether the molecular control of CD8+ effector and memory T cells elicited by the attenuated VACV strains paralleled that in response to WR. We recently demonstrated a major role for OX40/OX40L interactions in generating CD8+ T cell memory in response to WR (25) and therefore initially focused on these molecules. OX40 is not constitutively present on T cells, but has to be induced by antigen recognition and inflammatory stimuli; similarly, OX40L is only expressed on APCs such as dendritic cells and B cells after their activation by a variety of stimuli and cytokines (18, 26). Importantly, the absence of OX40 only impaired memory in response to WR, but did not affect the number of memory CD8+ T cells induced by WR-B18R, Lister, or NYCBOH (Figure 3, A and B). In fact, the magnitude of the CD8 response to WR in OX40−/− mice highly correlated with the response to WR-B18R, Lister, and NYCBOH in WT mice. Therefore, a large cytokine-competent, VACV-reactive memory CD8+ T cell pool only formed when OX40 became active in response to strongly replicating virus. Virus-specific memory T cells did form when OX40 was not brought into play, but their frequency was markedly reduced.

To again determine whether this related to initial priming events during the periods of active virus replication, the CD8+ T cell response at day 7 was assessed. There were approximately 75% fewer primary CD8+ T cells induced in OX40−/− mice after WR infection. In contrast, little difference in priming of CD8+ T cells reactive with B8R (or another peptide, A23R; data not shown) was observed in OX40−/− mice infected with WR-B18R, Lister, or NYCBOH (Figure 3C). To further show that the overall response and use of OX40 was related at least in part to the extent of attenuation and to viral load, mice were infected with 10 times as much virus (Figure 3D). The CD8+ T cell response to WR-B18R was strongly increased, to levels similar to WR, when given at the lower dose; OX40 became relevant and accounted for the enhanced response. The requirement for OX40 observed with a higher inoculum of WR-B18R directly correlated with increased viral titers recovered from the spleen (Figure 3E). A similar trend was also seen with NYCBOH: a 10-fold higher infection dose resulted in slightly elevated CD8+ T cell responses, albeit not to the same extent as WR or WR-B18R, and much of the enhanced response was attributable to OX40.
To extend these findings, we injected poly:IC at the time of WR infection. Treatment with poly:IC is known to lead to elevated levels of cytokines like TNF and IFN-I that can suppress viral replication. Strikingly, poly:IC resulted in only low numbers of VACV-reactive CD8$^+$ T cells being generated, to levels approximating those induced in response to Lister and NYCBOH (Figure 3, F and G). Most importantly, the weaker CD8 expansion seen with WR in the context of poly:IC injection directly correlated with reduced viral titers in the spleen (Figure 3H), and this T cell response was independent of OX40. These data further show that the use of OX40 to drive enhanced CD8$^+$ T cell priming and memory is related to the extent of viral load.

CD28 and CD27 are differentially active, depending on the stage of CD8 response and viral virulence. To determine whether this differential use of a stimulatory receptor applies to other, similar molecules, CD28 and CD27 were examined. CD28 is a costimulatory receptor in the Ig superfamily known to regulate initial activation of T cells in many immune responses in concert with antigen signals (27), and it has variably been reported to be required for CD8$^+$ T cell responses against viruses (28). CD28 is constitutively expressed on T cells and binds B7.1 and B7.2 expressed on many APCs, and CD28/B7.2 interactions play a role in generating pools of VACV-reactive CD8$^+$ T cells after infection with WR (28, 29). CD27 is another TNFR family member that acts independent of CD28 to promote T cell responses. CD27 is also constitutive on all T cells, but CD70, its ligand, is similar to OX40L and only induced once APCs such as dendritic cells or B cells receive certain inflammatory signals (18, 19). CD27/CD70 interactions were previously shown to be active in driving the CD8$^+$ T cell response to WR (30). Furthermore, CD27 and OX40 have previously been shown to cooperate together to regulate T cell memory to influenza virus (31, 32).

When memory was assessed 40 days after VACV i.p. infection, the requirement for CD27 followed the principles demonstrated for OX40: although CD27 contributed substantially to development of the high frequency of CD8$^+$ T cells elicited to WR, it was not active in the weaker responses to Lister and NYCBOH (Figure 4A). This selective use of CD27 was again reflected in the primary effector response at day 8, in that CD27 was not required for the smaller CD8$^+$ T cell populations elicited in response to WR in the absence of CD8$^+$ T cell populations elicited in response to Lister or NYCBOH, but controlled those induced in response to WR (Figure 4B). There was a small defect in the CD8$^+$ T cell response in Cd27$^{−/−}$ mice infected with the mutant WR-B18R virus, but again, the response in Cd27$^{−/−}$ mice largely followed that in Ox40$^{−/−}$ mice. Cd28$^{−/−}$ mice revealed a divergent pattern of usage. CD28 was required for development of primary CD8$^+$ T cells induced to all virus variants when assessed at day 8. In contrast, at day 40, CD28

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**Figure 1**

Altered virulence of VACV strains. C57BL/6 WT mice were infected i.p. with different strains of VACV ($2 \times 10^5$ PFU). (A) On the indicated days after infection, ovaries and spleens were removed from individual mice, and VACV titers were determined as described in Methods. (B) WT mice were infected i.n. with $10^6$ PFU WR and $10^5$ or $10^6$ PFU WR-B18R, Lister, and NYCBOH as indicated. Animals were weighed daily. Mean percent of initial body weight is shown. Results are mean (n = 4 per group) from 1 of 3 experiments. *P < 0.05 vs. WR.
Figure 2

VACV virulence correlates with magnitude of CD8+ T cell memory. WT mice were infected i.p. (A and C–F) or i.n. (B) with the indicated VACV strains. Splenocytes and lung cells were harvested on day 40 (A), day 60 (B), day 540 (C and D), or days 4, 5, 6, 7, and 8 (E and F) after infection and stained with anti-CD8, -CD44, and -B8R or for intracellular IFN-γ after restimulation with the designated VACV peptides. (A) Representative plots of gated CD8+ T cells staining for CD44 and B8R in spleen and lung. Numbers indicate percent CD44+B8R+ cells after gating on CD8+ T cells. Total numbers of CD8+CD44+B8R+ T cells per organ were determined as described in Methods. *P < 0.05 vs. WR. (B) Number of CD8+IFN-γ+ T cells per lung specific for peptides B8R, A3L, A8R, and B2R. (C and D) Percent CD8+IFN-γ+ T cells per spleen (C) or lung (D) specific for B8R and A8R. (E and F) Total number of CD8+CD44+B8R+ (E) and CD8+IFN-γ+ (F) T cells per spleen at the indicated time points after infection with VACV variants. Results are mean ± SEM (n = 4 per group) from 1 of 2 experiments. *P < 0.05 vs. all other strains or as otherwise denoted.
still contributed significantly to memory CD8\(^+\) T cell populations generated in response to WR-B18R, Lister, and NYCBOH, but memory T cell development in response to native WR was unaffected (Figure 4, A and B). The kinetics of development of VACV-reactive CD8\(^+\) T cells defined 2 obvious phases: a CD28-dependent phase, regulating primary effector cells, and a CD28-independent phase, beginning at day 12, in which the frequency of CD8\(^+\) T cells was equivalent in WT and \(\text{Cd}28\)–/– mice (Figure 5, A and B). Further demonstrating that this CD28-independent memory generation was driven by the TNFR family molecules, blocking CD70 in \(\text{Cd}28\)–/– animals from day 5 after infection largely recapitulated the defect seen in the absence of CD27 (Figure 5, C and D). Thus, the strong T cell memory elicited in response to the virulent virus strain was dependent on 2 TNFR family interactions, whereas the weaker memory in response to Lister, NYCBOH, and WR-B18R strains was driven by CD28 and largely independent of the TNFR family molecules.

Immunization with WR, but not with NYCBOH or Lister, protects mice lacking MHC class II against lethal respiratory virus challenge. Next, this alternate use of costimulatory receptors that drove expanded memory was studied in terms of protection against subsequent viral
encounter. Because antibody can protect against VACV, we chose a model whereby CD8+ T cell activity can be separated from antibody-mediated protection. After i.n. infection with WR, naïve mice exhibit weight loss and die within 6–9 days, but memory CD8+ T cells induced by peptide vaccination can afford protection in this situation (25). MHC class II–deficient (MHCII−/−) mice, which lack CD4+ T cells and cannot mount a VACV-specific neutralizing antibody response, were vaccinated with WR and Lister and challenged 70 days later with a lethal i.n. dose of WR. All mice vaccinated with WR survived the infection and exhibited only 10%–15% weight loss (Figure 6, A and B). Protection was dependent on CD8+ T cells, since their depletion before challenge resulted in 100% mortality. In contrast, no protection was evident in mice vaccinated with Lister or NYCBOH, and all succumbed to the infection with kinetics similar to those of naïve mice (Figure 6, A and B, and data not shown). This response paralleled the titers of virus detected in the lungs, which were strongly reduced (100–1,000-fold) in mice vaccinated with WR, but not in those vaccinated with Lister (Figure 6C). Consistent with the notion that the frequency of memory T cells dictated protection, far fewer B8R-specific memory CD8+ T cells were present in the spleen and lungs of MHCII−/− mice after immunization with Lister (Figure 6, D and E). This shows that the CD8+ T cell memory elicited because of varying levels of replication, and the differential use of OX40 and CD27, correlated with the degree of protection that could subsequently be provided.

To formally demonstrate that the frequency of memory CD8+ T cells is a primary determinant of the extent of protection afforded in the absence of CD4+ T cells and antibody, mice were infected with WR; 35 days later, varying numbers of naïve (CD8+CD44hi) or VACV-reactive memory (CD8+CD44loB8R+) T cells were isolated and then transferred directly into the lungs of naïve recipients by intratracheal injection. As a positive control, infected mice were bled at the same time, and VACV-immune serum was prepared and transferred i.p. into separate groups of naïve mice. All groups were subsequently infected i.n. with a lethal dose of WR. Mice that received naïve CD8+ T cells did not survive the infection, whereas 100% were protected by direct intratracheal transfer of 1 × 104 B8R+ T cells (Figure 7A). Lower numbers of VACV-specific CD8+ T cells either protected a fraction of mice or failed to protect at all, with the effect essentially proportional to the number of memory cells transferred. Similarly, the extent of disease as measured by weight loss directly correlated with the number of B8R-reactive memory CD8+ T cells transferred (Figure 7B).

Targeting OX40 enhances virus-specific CD8+ T cell responses to attenuated VACVs. We further investigated whether engagement of OX40 would boost the response to an attenuated VACV strain by treating with an agonist antibody after infection. Anti-OX40 enhanced CD8+ T cell priming in response to Lister, with 300%–400% more IFN-γ–producing cells detected regardless of epitope specificity (Figure 8, A and B). To exclude the possibility that the enhanced VACV-specific CD8+ T cell response was dependent on CD4+ T cells, MHCII−/− mice were infected with Lister. Anti-OX40 treatment again strongly enhanced the accumulation of these CD8+ T cells (Figure 8, C and D).

Most importantly, anti-OX40 provided complete protection against death in MHCII−/− mice vaccinated with Lister, which alone was ineffective at preventing lethality (Figure 8E). Notably, the extent of protection when mice were vaccinated with the combination of Lister and anti-OX40 was similar to that when mice were vaccinated with the virulent VACV strain WR. Thus, exogenous OX40 signals given at the time of activation of naïve VACV-specific CD8+ T cells promote a large population of CD8+ T cells that have the ability to fully protect against subsequent lethal virus infection.

**Figure 4**

Virulence of VACV and stage of CD8+ T cell response determine the selective use of CD28, OX40, and CD27. WT, Cd28−/−, OX40−/−, or Cd27−/− mice were infected i.p. with VACV variants (2 × 106 PFU) as indicated. At day 40 (A) or day 7 (B), splenocytes were stained with CD8 plus CD44 and B8R, and the number of VACV-reactive CD8+ T cells was calculated. Results are mean ± SEM (n = 6 per group). *P < 0.05 vs. WT. Similar results were obtained in 2 separate experiments. Inset in A shows the same data for WR-B18R, Lister, and NYCBOH with the scale enlarged.
The recombinant vaccine. Cottontop tamarins vaccinated with a VACV vector also critically influence the protective efficacy of et al. (17) and supported by our present data, the virulence of a CD4
tility (40% response), but the responses induced were exclusively
same vaccines at a higher dose showed enhanced immunogenic
importantly, and correlating with our prior data and hypotheses,
number of primed CD8
memory, because they did not allow molecules
significant implications for understanding mechanisms by which
antiviral immunity is induced, and also for methodologies used
to vaccinate individuals against pathogen attack. Increasing the
number of primed CD8
T cells by agonist engagement of OX40
Lístter resulted in numbers of CD8
major molecules in the TNFR superfamily — to retain the ability to use
including OX40 and CD27 — and perhaps similar costimulatory
which suggests that future vaccine strategies should consider
improved the efficacy of vaccination with attenuated VACV,
importantly, and correlating with our prior data and hypotheses,
OX40 and CD27, as well as CD28, were then involved in driving this
CD8
T cell response to Lister (Figure 9, C and D). Immunization of MHCII–/– mice with Lister via scarification resulted in
enhanced priming of CD8
T cells, and use of these TNFR family costimulatory molecules.

**Discussion**

Recent animal and clinical studies have questioned whether attenuated poxvirus-based vectors that are currently under
evaluation for vaccination elicit optimal long-lived CD8
T cell immunity. For example, the ALVAC HIV-1 vaccine in humans elicited HIV-1–specific CD8
T cell responses in less than 25%
of normal volunteers (8–10). Similarly, MVA expressing the GAG protein (consensus of HIV-1 clade A) and several immunodominant CD8
T cell epitopes resulted in responses detectable in only 17% of individuals (34). A smaller study using the same vaccines at a higher dose showed enhanced immunogenicity (40% response), but the responses induced were exclusively the result of CD4
T cells (7). As previously shown by Morgan et al. (17) and supported by our present data, the virulence of a VACV vector also critically influences the protective efficacy of the recombinant vaccine. Cottontop tamarins vaccinated with a

**Figure 5**

CD27 signaling is critical for CD28–independent CD8
T cell responses to WR. (A and B) WT or Cd28–/– mice were infected i.p. with WR (2 × 10
PFU). At the indicated days after infection, splenocytes were stained with CD8 plus CD44 and B8R (A), or intracellular IFN-γ or TNF (B), and the number or percentage of VACV-reactive CD8
T cells was calculated. (C and D) WT, Cd28–/–, or Cd27–/– mice were infected i.p. with WR and injected i.p. on days 5, 6, 7, and 8 (C) with 150 μg nondepleting anti-CD70 blocking Ab in PBS. At day 13 after infection, splenocytes were harvested and stained for CD8, CD44, and B8R. (D) Representative plots of B8R staining, gating on CD8
T cells, as well as total number of CD8
CD44–B8R
T cells per spleen. Numbers within dot plots denote percentage of cells in the respective quadrants/gates. Results are mean ± SEM (n = 4 per group). *P < 0.05. Similar results were obtained in 2 separate experiments.
become highly active with certain strains of LCMV (35), and with HIV (36), that are persistent and infect chronically. PD-1 limits CD8$^+$ T cell reactivity over extended periods, in this case in favor of the virus. Our data represent the opposite effect, in favor of the host. OX40 and CD27 did not participate in any substantial way in priming of CD8$^+$ T cells against VACV strains that were rapidly cleared. Thus, OX40 and CD27 were not active when the immune system was very efficient in eliminating the primary infection. Variable results in use, and timing of use, of OX40 and CD27 have been reported before in systems with other viruses, such as LCMV Armstrong, influenza virus, vesicular stomatitis virus (VSV), and MCMV; in some cases, these molecules played limited or no roles in primary T cell responses, strong roles in memory T cell generation, or were found essential for second- ary reactivity of memory populations (30–32, 37–41). Similarly, reports of the dependency for CD28 signaling to generate virus-specific CD8$^+$ T cells also diverge considerably. $\text{Cd28}^{-/-}$ mice infected with LCMV generated a normal CD8 response (42, 43), but CD8$^+$ T cell priming in response to VSV (44) and influenza (41, 45) was severely impaired. As proposed before (19, 31, 32), some of this variability implied that the route of infection, the dose of virus used for infection, and whether the infection was acute or chronic might control the requirement for these individual interactions, perhaps reflecting differences related to the rate of viral replication, antigenic load, and the cytokine milieu induced in response to each virus. We now provide direct data in support of these hypotheses. The use and activity of OX40 and CD27 were characterized by the generation of a high frequency of long-lasting and functional memory CD8$^+$ T cells, but only when replication was for extended periods or if the inoculum of virus used for infection was markedly increased (also translating to a greater viral load over time). This is particularly significant in that the memory CD8$^+$ T cells induced had the ability to protect against future viral challenge in the absence of other mechanisms, including help from CD4$^+$ T cells and neutralizing antibody, and this was directly proportional to their numbers.

Figure 6
Vaccination of $\text{MHCII}^{-/-}$ mice with WR, but not NYCBOH or Lister, protects against lethal respiratory virus challenge. $\text{MHCII}^{-/-}$ mice were immunized i.p. with VACV variants ($2 \times 10^5$ PFU). Naïve $\text{MHCII}^{-/-}$ mice were used as control. 10 weeks after vaccination, mice were infected i.n. with a lethal dose of WR ($4.5 \times 10^6$ PFU; i.e., 400 $\times$ LD$_{50}$). Some $\text{MHCII}^{-/-}$ groups were depleted of CD8$^+$ (αCD8) T cells before i.n. challenge with VACV. Animals were weighed daily and euthanized if weight loss was greater than 30% body weight. Mean percent survival (A) and percent of initial body weight (B) are shown. Mean weight data in some cases were not plotted beyond the point at which mice died and beyond day 7 reflected only mice that survived infection. (C) On day 7 after challenge, tissues from individual mice that survived the infection were collected, and virus titers were determined by plaque assay as described in Methods. Results are mean ($n = 4$ per group) from 1 experiment. (D and E) Percent and total number of CD8$^+$CD44$^+$B8R$^+$ cells and B8R-reactive, IFN-$\gamma$–producing CD8$^+$CD62L$^-$ cells in the spleen (D) and lungs (E) of $\text{MHCII}^{-/-}$ mice 90 days after immunization with WR or Lister. Results are mean ± SEM ($n = 4$ per group) from 1 experiment. *$P < 0.05$ vs. WR.
research article

Although antiviral antibody can protect mice in the absence of CD8+ T cells (4, 46, 47), mice deficient in B cells or MHC class II are dependent on CD8+ T cells in protecting against VACV (4, 48). Thus, this argues that the development of such high frequencies of protective memory CD8+ T cells will offset any deficiency in the antibody arm of the antiviral response.

Our data further highlight the importance of TNFR/TNF superfamily molecules in eliciting protective T cell memory. A group of these receptor-ligand pairs are inductive molecules on T cells and/or APCs, often appearing late in the response, typified by — although not exclusive to — the interactions of OX40 with OX40L, CD27 with CD70, TNFR with TNF, GITR with GITRL, CD30 with CD30L, 4-1BB with 4-1BBL, and HVEM with LIGHT (18, 19, 32, 49, 50). The genes for these molecules are tightly clustered in groups on several human chromosomes and are thought to have arisen through duplication and genome expansion, with their products arguably possessing similar intrinsic functional activity. Over the years, we and others have proposed models whereby such costimulatory molecules act together at any one time to allow for both quantitative and qualitative control of T cells, as well as acting at different times in a temporal sequence (18, 19, 31, 49, 50). It has been hypothesized that the immune system might need this type of cooperative action because of natural or induced suppressive activities that limit immune reactivity. Certainly, there are many suppressive or coinhibitory receptor-ligand pairs, including PD-1 and its ligands, that directly oppose viral replication, which can lead to differential use of costimulatory receptors for T cells. We demonstrate that molecular plasticity can occur during antiviral responses, in which certain immune mechanisms thought nonessential could become highly relevant, and provide insights into factors that may allow for better vaccine design.

Methods

Mice. The studies reported here conform to the animal Welfare Act and the NIH guidelines for the care and use of animals in biomedical research. All experiments were done in compliance with the regulations of the La Jolla Institute Animal care committee in accordance with the guidelines by the Association for Assessment and Accreditation of Laboratory Animal Care. Female C57BL/6 mice and C57BL/6 background (8–12 weeks old) were purchased from the Jackson Laboratory. C57BL/6 mice were bred in house.

Peptides and tetramers. VACV peptide epitopes used in this study were predictably and synthesized as described previously (23, 24, 28). The aa sequences for the peptides used were as follows: B8R (aa 20–27; TSXKFESV), A3L (aa 270–277; KSINYMLL), A8R (aa 189–196; ITYRFYLI), B2R (aa 54–62; YSQVNRLY), A23R (aa 297–305; JGMNPLTFI), MHC/peptide tetramers for the WR epitope B8R (20–27; TSXKFESV)/H-2 Kb, which were conjugated to allophycocyanin, were obtained from the NIH Tetramer Core facility at Emory University.

Viruses. The WR and Lister VACV strains were purchased from the American Type Culture Collection, grown in HeLa cells, and titered on VeroE6 cells. NYCBOH was generated as low-passage stocks from commercial Dryvax (Wyeth) and grown using the same conditions described for WR.

Immunization protocols. For most experiments, mice were infected i.p. with 2 × 10^6 PFU VACV. Efferor responses were analyzed between days 7 and 8 after infection, while memory responses were analyzed 30 or more days after infection, after restimulating in vitro with VACV peptides as described previously (52). For dermal scarification, virus (10^6 PFU) was deposited at the base of the tail, and the skin at the site of the droplet was scarified 25–30 times with a 25-gauge needle. After 3–4 days, pustules or scabs were observed at the scarification site, indicating a localized VACV infection. Results are mean (n = 4 per group) from 1 experiment.
VACY i.n. challenge. Mice were anesthetized by inhalation of isoflurane and inoculated i.n. with $1 \times 10^6$ or $4.5 \times 10^5$ PFU WR as indicated. Mice were weighed daily for 2 weeks following challenge and were euthanized when they lost 25%–30% of their initial body weight. For protection experiments, mice were immunized i.p. or via dermal scarification at the base of the tail with $2 \times 10^6$ PFU WR, Lister, or NYCBOH. Body weight was calculated as percentage of the mean weight for each group on the day of challenge.

Flow cytometry. After lysing rbcs, splenocytes from infected mice were resuspended in RPMI-1640 medium (Invitrogen) supplemented with 10% FCS (Omega Scientific), 1% t-glutamine (Invitrogen), 100 μg/ml streptomycin, 100 U/ml penicillin, and 50 μM 2-mercaptoethanol (Sigma-Aldrich). 1–2 × 10^6 cells were plated in round-bottomed 96-well microtiter plates in 200 μl with medium or the indicated VACV peptides at 1 μg/ml for 1 hour at 37°C. GolgiPlug (BD Biosciences) was then added to the cultures according to the manufacturer’s instructions, and the incubation continued for 7 hours. Cells were stained with anti-CD8 (PerCP) and CD62L (PE), followed by fixation with cytofix/cytoperm (BD Biosciences) for 20 minutes at 4°C. Fixed cells were stained with anti-CD8 (PerCP) and CD62L (PE), followed by fixation and with anti-OX40 (FITC) and IFN-γ staining after stimulation with the indicated VACV peptides (B and D). Data are either representative plots of tetramer staining in gated CD8+ T cells, with percent positive indicated, or total number (mean ± SEM) of CD8+IFN-γ+ T cells per spleen from 4 individual mice. *P < 0.05. Similar results were obtained in 2 separate experiments. (E) MHCII−/− mice were immunized i.p. with WR or Lister (2 × 10^6 PFU). 1 day later, cohorts of Lister-immunized mice were treated with 150 μg anti-OX40. Naive MHCII−/− mice were used as control. 10 weeks after vaccination, mice were infected i.n. with a lethal dose of WR (4.5 × 10^6 PFU; i.e., 400 × LD50). Animals were weighed daily and euthanized if weight loss was greater than 25% body weight. Mean percent of initial body weight is shown, with percent survival indicated.
Serial dilutions were made, and virus titers were determined by plaque assay on confluent VeroE6 cells. For dermal scarification, the tail was cut at both sides of the scarification site. The excised tail was then homogenized and assessed for viral titers as described for spleens and ovaries.

**In vivo CD8**^+^ **T cell depletion.** Hybridomas were cultured in protein-free hybridoma medium-II (Invitrogen), and mAbs were isolated by dialysis of supernatant. Groups of WR-immunized MHCII^−/−^ mice (deficient in MHC class II) were depleted of CD8^+^ T cells with anti-CD8 (clone 2.43; 200 μg/mouse) given in 1 i.v. injection 3 days before, and 2 i.p. injections on days 1 and 4 after, i.n. challenge with WR. CD8 depletion was confirmed by flow cytometry of spleens and lungs of treated mice.

**Avidity titration.** Groups of WT mice were infected i.p. with 2 × 10^5 PFU WR or Lister. 40 days after infection with VACVs, the avidities of CD8^+^B8R^+^ T cells were titrated by IFN-γ intracellular staining after stimulation of splenocytes with graded concentrations of B8R peptide. Results are mean ± SEM (n = 4 per group) from 1 of 3 experiments. *P < 0.05. (E) MHCII^−/−^ mice were immunized by dermal scarification with WR and Lister (2 × 10^5 PFU). Naive MHCII^−/−^ mice were used as control. 10 weeks after vaccination, mice were infected i.n. with a lethal dose of WR (4.5 × 10^6 PFU; i.e., 400 × LD50). Animals were weighed daily and euthanized if weight loss was greater than 30% body weight. Mean weight data in some cases were not plotted beyond the point at which mice died and beyond day 7 reflected only mice that survived infection.

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