Modified Vaccinia Virus Ankara Can Induce Optimal CD8+ T Cell Responses to Directly Primed Antigens Depending on Vaccine Design

Yik Chun Wong, a Sarah Croft, a Stewart A. Smith, a Leon C. W. Lin, a Tania Cukalac, b Nicole L. La Gruta, b,c Ingo Drexler, d David C. Tscharke a

a John Curtin School of Medical Research, The Australian National University, Canberra, Australian Capital Territory, Australia
b Department of Microbiology and Immunology, University of Melbourne, The Peter Doherty Institute for Infection and Immunity, Melbourne, Victoria, Australia
c Monash Biomedicine Discovery Institute and Department of Biochemistry and Molecular Biology, Monash University, Clayton, Victoria, Australia
d Institute for Virology, Düsseldorf University Hospital, Heinrich Heine University, Düsseldorf, Germany

ABSTRACT A variety of strains of vaccinia virus (VACV) have been used as recombinant vaccine vectors with the aim of inducing robust CD8+ T cell immunity. While much of the pioneering work was done with virulent strains, such as Western Reserve (WR), attenuated strains such as modified vaccinia virus Ankara (MVA) are more realistic vectors for clinical use. To unify this literature, side-by-side comparisons of virus strains are required. Here, we compare the form of antigen that supports optimal CD8+ T cell responses for VACV strains WR and MVA using equivalent constructs. We found that for multiple antigens, minimal antigenic constructs (epitope minigenes) that prime CD8+ T cells via the direct presentation pathway elicited optimal responses from both vectors, which was surprising because this finding contradicts the prevailing view in the literature for MVA. We then went on to explore the discrepancy between current and published data for MVA, finding evidence that the expression locus and in some cases the presence of the viral thymidine kinase may influence the ability of this strain to prime optimal responses from antigens that require direct presentation. This extends our knowledge of the design parameters for VACV vectored vaccines, especially those based on MVA.

IMPORTANCE Recombinant vaccines based on vaccinia virus and particularly attenuated strains such as MVA are in human clinical trials, but due to the complexity of these large vectors much remains to be understood about the design parameters that alter their immunogenicity. Previous work had found that MVA vectors should be designed to express stable protein in order to induce robust immunity by CD8+ (cytotoxic) T cells. Here, we found that the primacy of stable antigen is not generalizable to all designs of MVA and may depend where a foreign antigen is inserted into the MVA genome. This unexpected finding suggests that there is an interaction between genome location and the best form of antigen for optimal T cell priming in MVA and thus possibly other vaccine vectors. It also highlights that our understanding of antigen presentation by even the best studied of vaccine vectors remains incomplete.

KEYWORDS CD8+ T cells, CTL, cytotoxic T cells, MVA, modified vaccinia virus Ankara, antigen presentation, antigen processing, live vector vaccines, vaccinia virus

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Address correspondence to David C. Tscharke, david.tscharke@anu.edu.au.
* Present address: Yik Chun Wong, AIDS Institute and Department of Microbiology, State Key Laboratory of Emerging Infectious Diseases, Li Ka Shing Faculty of Medicine, University of Hong Kong, Hong Kong SAR, People’s Republic of China; Leon C. W. Lin, Institute of Biomedical Sciences, Academia Sinica, Nangang District, Taipei, Taiwan.
Y.C.W. and S.C. contributed equally to this article.

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Vaccinia virus (VACV) was one of the first vectors for recombinant vaccines and candidates have now progressed to clinical trials. The generation of strong CD8+ T cell immunity to foreign antigens encoded in a VACV vector is well agreed upon, in mouse (1–3), nonhuman primate (4–6), and human (7, 8) models. However, while the
bulk of historic studies aimed to understand antigen presentation from VACV were done with virulent strains such as Western Reserve (WR), these viruses have an unacceptable risk profile and are not suitable for use as human vaccines. Safer alternatives have emerged and among these is modified vaccinia virus Ankara (MVA); however, there is a lack of studies that examine MVA and WR in parallel, and such studies are required to unify a broad literature.

MVA is a hyperattenuated VACV strain that lacks immune evasion proteins and virulence factors and does not productively replicate in human tissue (9–11). Clinical trials with MVA as a smallpox vaccine have demonstrated safety, even in HIV- positive individuals (12, 13). These characteristics, along with the capacity to insert up to 25 kbp of foreign genes (10, 14), makes MVA an ideal vector for recombinant vaccines, a potential that is being pursued through the development pipeline into clinical trials (15–24). Further work is progressing to improve immunogenicity and assess the risks of broad release of MVA as a recombinant vaccine (25, 26), but these are large vectors, leaving many aspects of their biology incompletely studied.

A key design parameter for vaccines that induce CD8+ T cells is the form of antigen expressed, be that full-length protein and other stable polypeptides or, conversely, rapidly degraded antigen forms, such as ubiquitin-antigen fusion proteins, artificial polypeptides, and minimal epitope constructs (the latter are referred to here as minigenes). This is an important consideration because of the manner in which each of these is processed and presented on major histocompatibility complex (MHC) class I to activate or prime CD8+ T cells (27). Epitope minigenes bypass the requirement for proteasomal processing from larger polypeptides and so enter the antigen presentation pathway very efficiently in infected cells and are therefore present in great abundance on MHC. However, these short peptides are rapidly degraded, being shown to have a half-life of under 10 s (28). For this reason, although minigenes, and indeed all rapidly degraded polypeptides, prime well by the direct presentation pathway, i.e., via vaccine-infected dendritic cells (DCs), they do not survive long enough to be picked up and cross presented by uninfected DCs (29–31). Epitopes from stable antigens are directly presented on virus-infected DCs at various levels, but in many cases they can also be cross primed (27). Direct presentation is linked closely with translation, and so the structures or functions of full-length proteins are not associated with the levels of presentation by this pathway, unlike factors such as the efficiency of processing and the rate of translation (27, 32–35). Cross presentation is more likely to be influenced by the features of proteins, but with the exception of stability this has not been systematically explored. Further, the relative roles of these two pathways remain difficult to dissect for stable proteins, so it is safest to assume that both pathways may be used; but if neither is efficient, the responses elicited may be modest (36, 37). Finally, we note that while epitope minigenes are themselves not realistic vaccine constructs, because they prime responses to a single epitope presented by a single MHC they are a good model for all rapidly degraded antigen forms, as noted above.

Many studies over 20 years have shown that minigenes (or rapidly degraded polypeptides) encoded by VACV WR induce a CD8+ T cell immune response that is always as strong and often significantly stronger than a corresponding full-length stable antigen (37–41). Importantly, this finding holds irrespective of the nature or function of the full-length protein and implies that for recombinant VACV strain WR (rWR), optimizing antigens for effective direct presentation is always adequate and often ideal. In contrast to this, for MVA the prevailing view in the field is that minigenes and other rapidly degraded polypeptides prime poorly. This is based on a key study that directly examined priming requisites for this vector and concluded that stable antigens are best because cross priming is dominant (42). Notably, that report also examined multiple antigens, always finding the same result, suggesting that the finding would generalize broadly. This has been more recently supported by evidence that a second wave of presentation of antigens, which for MVA occurs exclusively by cross presentation, is required for CD4+ T cell help and full development of CD8+ T cell responses (43). Further, mouse knockout models have found that molecules required for cross presen-
tation impact the immunogenicity of MVA more than WR (44). However, many studies have found that MVA can infect DCs, and initial activation of CD8<sup>+</sup> T cells by direct priming has been visualized (42, 43, 45–48). In addition, effective priming of a minigene by MVA has been noted as an incidental finding elsewhere (37).

The purpose of this study was to reconcile these opposing findings for WR and MVA, as well as the possible discrepancy across findings with MVA. To do this, we revisited the preferred form of antigen for CD8<sup>+</sup> T cell priming by recombinant VACV (rVACV) based on strains WR and MVA. Our approach at the outset was to examine a broad range of antigens, which included: herpes simplex virus (HSV) glycoprotein B (gB), a highly immunodominant viral glycoprotein; influenza virus PB1F2, a very weakly immunogenic intracellular viral protein; VACV B8, a dominant native antigen of VACV; and ovalbumin, a classic model antigen for which published data already exist for WR and MVA. By choosing this broad array of antigen/epitope pairs, we sought to identify unifying patterns.

**RESULTS**

A minimal epitope construct of HSV gB<sub>498</sub> is more immunogenic than the full-length protein when expressed both from VACV WR and from MVA. The first foreign antigen/epitope we examined was gB of HSV, from which the highly immunodominant epitope gB<sub>498</sub> is derived (49). Recombinant VACV strain WR (rWR) viruses expressing full-length gB and an endoplasmic reticulum (ER)-targeted minigene (minigene-gB<sub>498</sub>) from the J2R gene under the p7.5 promoter have been published previously (50, 51). ER-targeted epitope minigenes deliver minimal epitope sequences directly into the ER, and so their presentation in infected cells is generally independent of the transporters associated with antigen presentation, but they behave similarly to cytosolic epitope minigenes and other rapidly degraded antigen forms in terms of priming pathway preference (30, 52). J2R encodes the thymidine kinase (TK), and this function is lost in viruses made in this way due to insertional inactivation. This insertion site is referred to here as the TK locus. For this study, we generated recombinant MVA (rMVA) viruses that were matched to the rWRs above in the forms of antigen, site of insertion, and promoter. As controls, we used viruses that had insertions in the TK locus but that expressed no foreign viral protein.

One of the limitations of using epitope minigenes is that their expression cannot be detected by conventional methods, such as Western blotting. For this reason, we needed a way to detect the epitopes presented in association with MHC class I (MHC-I) on cells infected with these viruses to ensure that all were being expressed. Further, by detecting the level of presentation, we have some indication of how well each of the viruses might perform in direct priming, assuming that the vectors can infect the relevant DCs in vivo. To do this for HSV gB<sub>498</sub>, we used an assay based on the ability of cells infected with our rVACV in vitro to restimulate CD8<sup>+</sup> T cells from mice acutely infected with HSV (Fig. 1A). We used the C57BL/6-derived cell line DC2.4, and after 2, 4, or 6 h of infection with the rVACVs, cocultured these cells with splenocytes taken from a mouse 7 days after infection with HSV. The coculture was done in the presence of brefeldin A and restimulation of CD8<sup>+</sup> T cells was determined by the detection of intracellular gamma interferon (IFN-γ) by flow cytometry (Fig. 1A depicted in blue). We cultured another aliquot of the same splenocytes with 1 × 10<sup>-7</sup> M gB<sub>498</sub> peptide and again measured the intracellular IFN-γ to establish a maximum possible response (Fig. 1A, depicted in black). This allowed the response from rVACV-infected cells to be plotted as a percentage of the maximum possible response. This was necessary for standardization across experiments. As expected, for both rWR and rMVA, cells infected with viruses expressing minigene-gB<sub>498</sub> were the better at restimulating CD8<sup>+</sup> T cells from HSV-infected mice than those infected with the viruses expressing full-length gB (Fig. 1B). At the same time, viruses with no form of gB failed to restimulate the HSV-immune splenocytes, showing there was no cross-reactivity between HSV- and VACV-specific CD8<sup>+</sup> T cells. Finally, to ensure that these results were not due to differing efficiencies of the infections or other factors that might impact antigen
presentation in general across the various viruses, portions of the same batches of infected cells used above were tested for their ability to stimulate CD8\(^+\) T cells from VACV WR-infected mice (Fig. 1A, depicted in orange). Restimulation of WR-immune CD8\(^+\) T cells was similar among the cultures infected with the three rWRs and also across those infected with the three rMVA (Fig. 1C). These controls suggest that the infections were all similarly efficient, at least within a VACV strain. Taken all together, we interpret the results of these experiments as showing that (i) full-length gB and minigene-gB\(_{498}\) were expressed and presented on MHC-I as anticipated from the rWR and rMVAs but (ii) that minigene-gB\(_{498}\) was a more efficient form of antigen in terms of direct presentation of the gB\(_{498}\) epitope on infected cells.

Next, we used an established vaccination method to determine whether the gB-containing antigens expressed by our viruses in infected cells might be able to cross prime CD8\(^+\) T cells in vivo (30, 37, 52, 53). As a source of antigen, human 293A cells were infected with rWR and rMVA viruses and then heat killed to destroy any infectivity, which includes residual viral inoculum as well as any replicated virus. These cells were used to vaccinate mice by intradermal (i.d.) injection of ear pinnae and, after 7 days, epitope-specific responses were determined by a short ex vivo culture with synthetic peptides and staining for CD8 and IFN-γ (Fig. 2A). We measured responses to gB\(_{498}\) and,
to ensure similar immunization across viruses, selected VACV epitopes, including the dominant B820 and a set of less-dominant epitopes comprising A47138, A47171, L253, and A3270, for which responses were summed and are referred to as P4. All of these epitopes are shared between WR and MVA (54, 55). Both for rWR and for rMVA, only cells infected with viruses that expressed the full-length gB were able to prime a gB498-specific CD8^+ T cell response above background in mice (Fig. 2B). The sizes of these responses were low, at around 0.2% of CD8^+ T cells, but this is in the range published for similar types of experiments (30, 37). At the same time, rWR-infected cells primed responses to the VACV strain as shown above the graphs expressing the form of gB498 as shown in the key. The epitopes are shown on the x axis; P4 is the sum of responses to A47138, A47171, L253, and A3270. Means and standard errors of data from six mice combined from two independent experiments are shown (*, P < 0.05).

Finally, we wanted to determine for rWR and rMVA the form of HSV gB that was most immunogenic after a standard infection of mice, which is the key experiment with these viruses. To do this, mice were infected by the i.d. route with the gB498-presenting rWR and rMVA viruses, and CD8^+ T cell responses to gB498, P4, and B820 in the spleen were measured 7 days later (Fig. 3A). As published previously for rWR (51), minigene-gB498 primed significantly more CD8^+ T cells than full-length gB, but more surprisingly, this result was recapitulated by the rMVA viruses (Fig. 3B). Indeed, the strength of the minigene-gB498 priming from both vectors appeared to compete with native VACV epitopes, including the usually dominant B820 epitope, such that responses were reduced in mice infected with the minigene, compared with full-length gB-expressing viruses. However, we cannot be absolutely sure that reduction of response to the VACV epitopes does not reflect some disadvantage in infection or general antigen presentation for viruses expressing minigene-gB498. For this reason, we also plotted gB498-specific responses normalized against the total VACV-specific response (sum of P4 and B8), which highlighted further the difference between minigene and full-length HSV gB-expressing viruses (Fig. 3B).
From these data, we concluded that for the highly dominant HSV gB498 epitope, the priming requisites for rWR and rMVA are the same when similar constructs are compared, the form of gB optimized for direct presentation (Fig. 1) and not able to cross prime (Fig. 2) being the most immunogenic (Fig. 3).

Minigene-IAV PB1-F262 is more immunogenic than full-length PB1-F2 when expressed from VACV strains WR and MVA. We were concerned that the gB498 epitope might not be representative, perhaps because of its immunodominant nature or its structure as a surface glycoprotein, so we wanted to determine whether the results above would be consistent for a very weak, cytoplasmic antigen. We chose a subdominant influenza A virus (IAV) antigen/epitope, namely, the nonstructural PB1-F2 protein and its epitope PB1-F262 (57). The viruses used were matched with those expressing HSV gB above in promoter and insertion site. The rWRs were made by others (53, 58), but the rMVAs were made for this study. The same TK– control viruses were used.

Following the experimental pattern established in the previous section, we first tested the ability of these rVACVs to express their antigens in infected cells, as reflected by the presentation of PB1-F262 on MHC-I. The experimental scheme was as shown in Fig. 1A, but this time we used splenocytes from IAV-primed mice to detect the presentation of PB1-F262. All viruses with PB1-F2, irrespective of the form, were detected by the IAV-immune CD8+ T cells, but the control WR and MVA were not detected. As expected, both for rWR and for rMVA, cells infected with viruses expressing minigene-PB1-F262 were the best at restimulating CD8+ T cells from IAV-infected mice (Fig. 4A, left). The control restimulations of WR-immune splenocytes showed that the VACV-derived epitopes were presented equally across the set of rWRs and across the set of rMVAs, so that within each strain the infection was equally efficient (Fig. 4A, right). We interpret these results, taken together, to show that these sets of viruses...
expressed their antigens and that there was more efficient presentation of PB1-F2 on infected cells expressing the minigene than the full-length PB1-F2.

Next, we used the in vivo assay based on immunization of mice with infected and heat-treated 293A cells to determine whether full-length and minigene-PB1-F2 were able to cross prime CD8⁺ T cells. As above, we measured responses against PB1-F2₆₂ and as controls, VACV epitopes including B₈₂₀, and the P₄ set of epitopes. None of the four viruses was able to prime a PB1-F2₆₂-specific CD8⁺ T cell response that was clearly above background in these experiments, despite the VACV antigens being able to elicit

**FIG 4** Presentation to, and priming of, CD8⁺ T cells by PB1-F2₆₂ expressed in different forms by rWR and rMVA. (A) Results obtained according to the experimental design in Fig. 1A show the extent of restimulation of CD8⁺ T cells from IAV-immune (left) or VACV WR-immune (right) splenocytes by cells infected by the virus strain, as shown above graphs, expressing the forms of PB1-F2₆₂ as shown in the key. For the cocultures with IAV-immune splenocytes (left), data are presented relative to the maximum possible value obtained by stimulation of the same spleen cells with PB1-F2₆₂ peptide. Means and standard errors of triplicates are shown. The experiment was repeated with similar results. (B) Results obtained according to the experimental design shown in Fig. 2A, except viruses expressed versions of PB1-F2₆₂ (as shown). Epitope-specific responses are shown as the percentage of CD8⁺ T cells making IFN-γ. (C) Results obtained according to the experimental design in Fig. 3A. Mice were infected with rWR and rMVA viruses expressing versions of PB1-F2₆₂ (as shown) and, 7 days later, the epitope-specific responses were measured. The graph on the right for each VACV strain shows the PB1-F2₆₂-specific response divided by the total VACV-specific response. For panels B and C, the means and standard errors of data from nine mice from three independent experiments are shown (*, P < 0.05; **, P < 0.01).
responses at the expected levels (Fig. 4B). These data are consistent with the poor immunogenicity of PB1-F2<sub>62</sub> when mice were immunized with IAV-infected MC57G cells lacking TAP (59). Further, it is possible that expression of PB1-F2 protein is not well tolerated by cells in the context of infection with one or both rVACVs, and this is reflected in the lack of a response. Arguing against this is that the responses to the native VACV epitopes (P4 and B8) were similar for the viruses expressing full-length PB1-F2 (presumably functional) and minigene-PB1-F2<sub>62</sub> (not functional). So, for PB1-F2<sub>62</sub> we were not able to confirm whether either form of antigen was able to cross prime CD8<sup>+</sup> T cells, probably due to the poor immunogenicity of this epitope.

Finally, mice were infected with the PB1-F2<sub>62</sub>-presenting viruses to determine for each vector which form of antigen was most immunogenic in a live-virus vaccination. All responses to PB1-F2<sub>62</sub> were exceptionally weak, being just above background; however, all the average responses were above zero by more than the 95% confidence level (not shown). Further, the minigene-PB1-F2<sub>62</sub> construct primed significantly more CD8<sup>+</sup> T cells than full-length PB1-F2, when expressed both from rWR and from rMVA (Fig. 4C). The CD8<sup>+</sup> T cell response to the native VACV epitopes appeared to be similar across the viruses from each strain; however, to take this into account formally, we normalized PB1-F2<sub>62</sub>-specific responses against the total anti-VACV response. Doing this confirmed that the responses primed by minigenes were stronger. Therefore, we conclude that minigenes are the optimum form of PB1-F2<sub>62</sub> for stimulating CD8<sup>+</sup> T cell responses for rWR and rMVA vectors, even though responses were very low and close to the limit of detection.

With the exception of our inability to detect cross priming from infected cells, these data recapitulate what was seen for HSV gB<sub>498</sub>: there was no difference in the preferred form of antigen for priming CD8<sup>+</sup> T cells between rWR and rMVA, and directly presented minigenes were more immunogenic than a full-length antigen.

**Minigene-B8<sub>20</sub> is more immunogenic than full-length B8 when expressed from MVA.** The two antigens examined thus far were expressed from the same promoter and were inserted into the TK gene, so we then extended the study to a VACV antigen expressed by its own promoter, from its native location. We chose the highly immunogenic VACV B8<sub>20</sub> epitope and compared antigen presentation and immunogenicity from wild-type viruses and from recombinants where the BBR open reading frame had been replaced with a short sequence encoding a B8<sub>20</sub> minigene. It should be noted here that B8 from MVA is truncated and predicted to be nonfunctional, so the nonrecombinant WR and MVA viruses that serve to provide full-length B8 protein are not strictly equivalent. However, (i) in well-controlled studies WR B8 fails to function in mice due to an inability to adequately bind mouse IFN-γ, so neither WR or MVA form of this protein should impact immune responses (60), and (ii) our data from Fig. 2 and 4B, in which B8 is used as a control native VACV antigen, confirm that both forms of B8 are stable enough to support very effective cross priming.

Direct presentation in vitro was used to ensure that all viruses expressed their antigens and present B8<sub>20</sub> as described above, but for these experiments the source of primed CD8<sup>+</sup> T cells was mice infected with an IAV that expresses the B8<sub>20</sub> epitope as part of the neuraminidase stalk. Again as expected, all antigens were expressed, and cells infected with the minigene-B8<sub>20</sub>-expressing viruses were better at restimulating CD8<sup>+</sup> T cells from IAV-B8<sub>20</sub>-immune splenocytes (Fig. 5A). At the same time, DC2.4 cells infected with viruses lacking B8 failed to restimulate the B8<sub>20</sub>-immune splenocytes but cells infected with all viruses were able to restimulate CD8<sup>+</sup> T cells from VACV WR-infected mice. Again, this verified that our rWR and rMVA viruses were expressing antigen and presenting B8<sub>20</sub> on infected cells as expected.

Next, we used heat-killed rVACV-infected cells to test the ability of B8 from these viruses to cross prime a CD8<sup>+</sup> T cell response. Cells infected with wild-type WR and MVA viruses, but not the recombinants expressing minigene-B8<sub>20</sub>, were able to provide antigen that cross primed B8<sub>20</sub>-specific CD8<sup>+</sup> T cells (Fig. 5B). This result for the wild-type B8 versions back up the data from Fig. 2 and 4B, again confirming that not only is the truncated B8 from MVA able to be cross primed, it is better in this assay than.
the full-length B8 from WR. It is possible that B8 from WR is largely lost from infected cells because it is secreted and that the truncated version from MVA is either retained or perhaps remains associated with cells to a greater extent. We also noted that the mean response to the P4 set of epitopes was lower for the minigene-B820 viruses than for corresponding wild types, but this did not reach statistical significance for WR or MVA. These data together support the idea that both versions of full-length B8 are able to be cross primed but, as expected, minigene-B820 is unable to cross prime CD8+ T cells.

Having again established that the B820-presenting viruses used in this section behaved as expected in the previous experiments, we infected mice with these viruses to test them as live vaccines. When mice were infected with these viruses in the case
of WR, the wild-type virus initially appeared to induce a significantly larger B820-specific CD8+ T cell response than the minigene-B820 recombinant; however, this trend was also noted for the P4 set of other VACV epitopes. To account for a possible difference in infections with these viruses, the data were normalized as a ratio of the B820-specific to the total anti-VACV response. In this analysis, there was no longer any significant difference between the two WR viruses, so it seems likely that the two forms of B820 were equally effective (Fig. 5C, left). In contrast, for MVA, the minigene-B820-expressing virus induced a very slightly larger B820-specific CD8+ T cell response than the wild-type virus (Fig. 5C, right). Furthermore, this difference between rMVAs remained significantly different when normalized against the total anti-VACV responses.

Thus, we concluded that for a VACV antigen expressed in its native condition there was no advantage from expression as a minigene by strain WR. However, this was not true for MVA, where the direct-priming minigene construct elicited a slightly, but significantly higher CD8+ T cell response. Whether this difference between strains is due to the variants of native B8 expressed by these viruses or other factors such as the extent of B8 immunodominance remains to be determined (54, 61).

Replication of a published result for MVA expressing the OVA257 peptide from the delIII locus. All the data above seemed to contradict the key study on antigen presentation from MVA by suggesting that minigenes, which require direct presentation, are an optimal form of antigen to induce a CD8+ T cell response using this VACV strain. So, we next used one of the same sets of rMVAs used by Gasteiger et al. (42) to repeat a published experiment, albeit using our i.d. infection route compared to the original intraperitoneal infections. These rMVAs, expressed either ovalbumin (OVA) or an minigene-OVA257 from the delIII locus, the site of one of the major genomic deletions in MVA. This site does not exist in the WR genome, so for comparison we used rWRs that expressed the same antigens from the TK locus, maintaining consistency with the other WR viruses used here. The presentation of OVA257 can be directly measured using a monoclonal antibody (25D1.16), which detects this peptide when presented by H-2Kb, and flow cytometry (62). This is a much simpler and more strictly quantitative compared to the assays using splenocytes described for the other antigens above, so we used this method to ensure all antigens were expressed and to examine direct presentation. As has been published elsewhere, cells infected with these viruses expressing minigene-OVA257 presented substantially more OVA257 compared to those infected with rVACVs expressing full-length OVA (Fig. 6A) (42, 63).

Next, we used heat-killed virus-infected cells to test the ability of these forms of OVA to cross prime CD8+ T cells in vivo. Irrespective of virus strain, only cells infected with viruses that express full-length OVA were able to induce a CD8+ T cell response (Fig. 5B), which is consistent with published results for rWRs (30).

Finally, mice were infected with the OVA257-expressing rWR and rMVA viruses to test the optimal form of antigen in our i.d. infection model. Consistent with all the data here for other antigens, and as published previously (41), rWR encoded minigene-OVA257 was more immunogenic than full-length OVA (Fig. 6C, left). However, unlike our data here for other antigens and consistent with the published pattern for rMVA (42), minigene-OVA257 was significantly less immunogenic than the full-length protein. Again, to formally take into account possible variations in infections across these viruses, we normalized OVA257-specific response to the total anti-VACV response (Fig. 6C). This additional analysis supported the initial comparison made with the unnormalized values. Thus, we were able to confirm previously published results for MVA, irrespective of the different route of infection, which is consistent with route not being important for determining antigen preference for MVA (42). This result confirms a contradiction between the rMVA results obtained with OVA as published previously (42) and those acquired for all three other antigens examined here.

Genome location and TK gene function affect the relative immunogenicity of OVA and minigene-OVA257 expressed by rMVAs. To address the discrepancy noted above, we considered the differences between the OVA-expressing MVAs and other foreign gene-expressing viruses used here. These were (i) the antigen, (ii) the genomic
location from which antigens were expressed, and (iii) the presence of a functional TK gene, which is inactivated when this locus is used to insert foreign antigens. We examined these differences with six new viruses: an OVA and minigene-OVA257 pair of rMVAs with these antigens inserted into the TK locus; a second pair of rMVAs with these antigens expressed from the intergenic space between A11R and A12L; and finally, TK– variants of the rMVAs with OVA and mini-OVA257 expressed from the delIII region (Fig. 5A). We tested expression and direct presentation on cells infected with these pairs of rMVAs in vitro and the immunogenicity of OVA257 in mice infected with these viruses.

Expression and antigen presentation on infected DC2.4 cells was tested with the 25D1.16 monoclonal antibody and flow cytometry. We found that all of the pairs of rMVAs behaved as expected with all presenting OVA257, and cells expressing the minigene viruses presented this epitope at higher levels than those infected with their paired full-length OVA-expressing counterpart (Fig. 7B to D, left). There were some differences in the data across the various pairs, with the A11R/A12L insertion and TK-
FIG 7 Presentation to, and priming of, CD8+ T cells by OVA257, expressed in different loci of rMVA and without a functional TK. (A) MVA genome maps showing HindIII fragments and the site of insertion of OVA antigenic constructs and the promoter (arrow). The “m” denotes the minigene (OVA257). (B to D, left) Extent of cell surface presentation of MHC-I:OVA257 complexes on DC2.4 cells infected with viruses. (B to D, right) OVA-257 response (normalized). (Continued on next page)
variant of MVA expressing OVA from the delIII locus apparently presenting OVA$_{257}$ less efficiently than the other viruses. However, we are presenting raw mean fluorescence data and that limits direct comparisons.

When mice were infected with these viruses, two different patterns of immunogenicity were observed between full-length and minigene-OVA$_{257}$ rMVAs. For the first pair of viruses, where antigen was expressed from the TK locus, minigene-OVA$_{257}$ induced higher CD8$^+$ T cell responses than full-length OVA (Fig. 7B, middle). This result was then tested further by normalization against the total VACV-specific responses and the difference remained in favor of minigene-OVA$_{257}$ and was statistically significant (Fig. 7B, right). This was the opposite of the results obtained in the previous figure using rMVAs expressing OVA$_{257}$ constructs from the delIII-based recombinants. It did, however, match the results for all the other antigen pairs that we have tested when expressed from the TK region and also for B8$_{20}$. This suggests that the locus of expression, rather than the antigen examined (OVA), is responsible for the apparent contradiction in previous data. For the remaining pairs of rMVAs, the full-length and minigene constructs elicited an equivalent OVA$_{257}$-specific CD8$^+$ T cell response; again, this was supported by formal normalization (Fig. 7C and D, middle and right). These data suggest that genome location and TK function can contribute to the relative immunogenicity of unstable and stable polypeptides that present OVA$_{257}$ when expressed from rMVA.

**DISCUSSION**

This study started with the aim of carrying out a comprehensive side-by-side comparison of the antigen requisites and therefore priming pathway preferences for virulent VACV, strain WR, and for MVA. The results for rWR viruses were entirely in line with what was first shown more than 2 decades ago, specifically that short-lived constructs were the optimum antigen form to elicit CD8$^+$ T cell responses (41). We have simply extended the data set to more antigens and expression sites in the virus and used more quantitative tools to quantify T cell responses (64). In contrast, the results for rMVAs were unexpected and so merit further discussion.

The first study of the optimum antigen forms for priming CD8$^+$ T cells by rMVA examined multiple antigens and antigen forms, but all being expressed from delIII (42). These included full-length antigens, as well as an ubiquitylated version of tyrosinase, an epitope of tyrosinase that is derived from the leader sequence of that protein and the minigene-OVA$_{257}$. There are further unpublished constructs with antigens inserted into the delIII region that all behave in the same way as shown elsewhere (I. Drexler, data not shown). Thus, there is strong evidence that whenever antigens are expressed from the delIII region of MVA, unstable antigens that can support only direct priming will induce poor CD8$^+$ T cell responses compared to stable antigens. This was the current state of understanding in the field when we began our experiments. However, these published results contrast with the data for all of the pairs of rMVAs that we made for this study. We present data for several antigens and, in our case, three different genomic loci and come to the opposite conclusion. The differences across these two studies suggest strongly that the antigen form that primes optimal CD8$^+$ T cell responses when expressed from rMVA differs according to the expression locus. Further support for this conclusion comes from our direct comparison of rMVAs that present the OVA$_{257}$ epitope. We used a set of OVA-expressing rMVAs examined in the original study noted above and confirmed the published result with our model (Fig. 6C). This was followed by our own recombinants that differed only in that expression was from rMVA.

**FIG 7 Legend (Continued)**

D, middle) Mice were infected with rMVA viruses expressing versions of OVA$_{257}$ (as shown) and, 7 days later, the responses to peptides were measured. (B to D, right) Using the data shown in the middle graphs, the OVA$_{257}$-specific response was normalized by dividing by the total VACV-specific response. Means and standard errors of data from at least nine mice combined from three independent experiments are shown (*, $P < 0.05$).
the TK locus and not delIII and found the opposite result (Fig. 7A). These results were supported by a rigorous normalization process to take into account any difference in infections (e.g., that might be caused by inaccurate virus titers used to infect the mice). In looking at these findings, it is pertinent to ask whether it was the immunogenicity of the full-length gene or the minigene (or both) that varied because this might suggest the priming pathway that is working with differing efficiency across the pairs. While a direct statistical comparison is not possible, across these two pairs of viruses (Fig. 6C to 7A) it can be seen that the immunogenicity of the minigene varies substantially, but the full-length construct elicits a similar response irrespective of expression locus. This is seen most clearly in the normalized data (Fig. 6C, far right, and Fig. 7A, right), where for full-length constructs the OVA_257/total VACV-specific responses are almost identical (0.129 and 0.123), but for the minigenes they vary almost 3-fold (from 0.071 to 0.211). Taken together, the best explanation these findings, ours in the present study and those of Gasteiger et al. is that the genome location of a foreign antigen influences whether antigens designed for direct or cross priming are likely to be most immunogenic when expressed from an rMVA.

MVAs that have a transgene inserted in the delIII region differ in two ways from the majority of our viruses in which the viral TK gene was used: the genomic site of insertion and the presence of the viral TK. In teasing these two factors apart, we found that when OVA_257 antigens were inserted in the A11R/A12L intergenic space, the full-length gene and the minigene were equally immunogenic (Fig. 7C). This result was halfway between those for the corresponding rMVA with delIII and TK locus insertions. This suggested that inactivation of TK was played some role in improved immunogenicity of minigenes over full-length protein expressed from the TK locus. Further support for this was provided by our results when we inactivated TK in the original delIII-inserted MVAs (Fig. 7D). However, we also note that when B820 was expressed as a minigene from its native location in MVA, it outperformed the full protein in terms of immunogenicity (Fig. 5C). This suggests that there is a role for other factors, perhaps the antigen/epitope, e.g., if the antigen is very poorly cross primed, or the precursor frequency of T cells that recognize a particular epitope, may also be important. Taken together, however, our data establish roles both for insertion site of transgene and for TK function in determining the immunogenicity of epitope minigenes from rMVA at least for one model antigen.

The reasons why these two factors might alter priming preferences for CD8^+ T cells are not clear. Differences in immunogenicity across sites and with variation due to TK function have been noted previously for virulent VACV, but in that case these differences were linked to transgene expression level (65). As noted above, in every experiment we found enhanced presentation of epitopes from minigenes over those from full-length proteins on infected DC2.4 cells. In addition, cells infected with MVAs that express full-length OVA generate enough of this protein to cross prime CD8^+ T cell responses in vivo as shown in Fig. 6B for the delIII insertion and as published previously for a TK insertion (37). Having said this, we are unable to know how much of each full-length antigen is expressed and more importantly its capacity to cross prime in vivo. However, differences in immunogenicity not linked to expression have been seen previously for MVA (66). We speculate that differences in genomic insertion sites can change the interaction of rMVA with the DCs responsible for priming CD8^+ T cells. In the case of rMVAs that used the delIII region, while no functional genes are disrupted, we wonder whether the transcription that is driven into the neighboring viral genes alters their expression, leading to changes in infected DCs. In support of this general concept, we have noticed that insertions that include an expressed gene into the A11R/A12L intergenic space can alter the immunogenicity of multiple native VACV epitopes, whereas a promoterless insertion did not have this effect (L. C. W. Lin and D. C. Tscharke, unpublished data). In contrast to these loci, use of the TK region leads to a loss of TK function. Indeed, this locus was originally chosen because this phenotype provided a selectable marker for recombinant viruses (67–69). VACV TK functions to phosphorylate thymidine, producing dTMP, which after a series of phosphorylation
events is utilized in de novo DNA synthesis (70). In vitro, the function of this gene is not required, but the loss of TK is substantially attenuating for virulent VACVs in vivo, reducing viral loads (54, 71). However, MVA fails to replicate in vivo, and so we did not expect to find any impact of TK deletion. Further, TK deletion from WR does not substantially change the specificity of CD8+ T cell responses, despite the attenuation noted above (54). Finally, although MVA does replicate its genome in some cells, and so inhibition of this DNA synthesis might have an impact, in the case of DCs MVA aborts infection at an earlier stage, suggesting that this is not relevant for direct priming (72).

It remains possible that there is some effect of TK function on DCs, perhaps prolonging their lifespan during infection. We have explored this in vitro and found no impact of TK expression from MVA on DC viability (Y. C. Wong and D. C. Tscharke, unpublished data), but our in vitro DC cultures are unlikely to model the situation in vivo very faithfully, all the more so given that a second wave of presentation has been shown to be required for full immunogenicity of VACVs (43). The role of TK and of transcription that runs into neighboring genes in MVA vaccines requires a more thorough investigation.

Our data may also have implications for our understanding of the ability of MVA to support direct priming in general. While there is one well-characterized exception, namely, minigene-IAV PA224, minigenes (either ER targeted or cytosolic) and other rapidly degraded constructs have never been found to be cross presented. This is consistent over multiple studies, and the expression of these antigens can therefore be used to establish the effectiveness of direct presentation (30, 38, 42, 51, 52, 56, 73, 74). Thus, the finding by Gasteiger et al. that multiple rapidly degraded proteins were poorly immunogenic suggested that direct priming is inefficient for MVA in general. We now find here that several rMVA minigene viruses are able to elicit CD8+ T cell responses at least as effectively as full-length proteins and often significantly better. This includes four antigens expressed from four loci in the virus, including under a natural promoter in the native location. The only exception is minigene-OVA257 expressed from delIII. The weight of evidence then suggests that MVA in general supports efficient direct priming of CD8+ T cells. There is likely to be a contribution by cross priming as well, but we are unable to determine the relative importance of these pathways for nonrecombinant MVA.

From a practical perspective, our data show that MVA can be an efficient vector for the delivery of antigens for direct presentation. Having noted this general rule, it is clear that the use of the delIII region, and perhaps others yet to be determined, to make rMVAs creates exceptions. This remains important because delIII was the original and remains a commonly used site for the introduction of genes encoding foreign antigens in rMVAs (14, 75). It is also necessary for the interpretation of any experiments where rMVA are used to dissect mechanisms of CD8+ T cell priming. Indeed, the finding that the XCR1+ DCs required for a fully functional CD8+ T cell response used exclusively cross priming was made using H-2Kbm1 mice infected with an rMVA that expressed H-2Kb and OVA from the delIII region. Finally, we have only examined systemic responses and, for the priming of resident memory populations (Trm), cross priming has been shown to be important using mouse knockout models (76).

In conclusion, we show that directly priming minigenes are optimal for CD8+ T cell priming by virulent (WR) and attenuated (MVA) rVACV, with the notable exception of rMVAs that express antigens from the DelIII region. Minigenes, while they are not likely to be used as vaccines themselves, are a model for all forms of rapidly degraded antigens. Other forms, such as ubiquitin fusions or polyepitope constructs, are more practical antigens for vaccines because they can induce responses to multiple epitopes in the context of multiple MHC allomorphs. MVAs with insertions into delIII are common, but it is not the only site used. Indeed, we are not the only group to use rMVAs with antigens expressed from the TK locus, and some of these vaccines have advanced to clinical trials, so our findings have practical implications (22, 77–83). The findings presented here in general highlight that there remain many wrinkles to iron
out in our understanding of antigen presentation, even for well-studied viral vectors that have been used in human clinical trials.

MATERIALS AND METHODS

Mice. Specific-pathogen-free, female, 7- to 14-week-old C57BL/6 mice were obtained from the Australian Phenomics Facility (Canberra, Australia) or ARC (Perth, Australia). All experiments were conducted according to relevant ethical requirements that were approved by the Australian National University Animal Ethics and Experimentation Committee (protocols F.BMB.38.08, A2011.001, A2013.037, and A2016.045).

Cells and nonrecombinant viruses. For cross presentation assays, 293A cells (ATCC, CRL-1573) were used as antigen donor cells. The C57BL/6 mouse-derived, dendritic cell-like cell line DC2.4 was used for in vitro presentation assays (84). Unmodified Western Reserve vaccinia virus (VACV WR, ATCC VR1354) and MVA were originally a gift from B. Moss (National Institutes of Health, Bethesda, MD). Influenza A virus (IAV) strain PR8 was provided by C. Goodnow (ANU, Canberra Australia). HSV strain KOS was provided by F. Carbone (University of Melbourne, Melbourne, Australia). All strains were grown and titrated according to standard methods.

Recombinant viruses and virus construction. The VACVs used here are listed in Table 1 with descriptions and origins, if not made for this study. With the exception of the minigene-B8, WR and MVA, which used the native B8 promoter, all antigens were expressed from the VACV p7.5 promoter. Homologous recombination between appropriately designed transfer plasmids supplied by transfection and VACV genomes provided by infection was used to make the new viruses required. Transfer plasmids were based on pSC11 and p7.5GB-ins for insertions into the TK and A11R/A12L regions, respectively (69, 85). Briefly BHK-21 or 293A cells were infected with MVA or WR, respectively, at a multiplicity of infection (MOI) of 0.05 in Dulbecco modified Eagle medium (DMEM) supplemented with 2% fetal bovine serum (FBS) and incubated at 37°C and 5% CO2 for 1 h. The inoculum was then removed, and the transfer plasmid was added in a preincubated transfection mix of plasmid, Lipofectamine 2000 (Life Technologies), and DMEM. Infected cell cultures were incubated

### Table 1: Viruses used in this study

| Virus | Recombinant details | Source or reference |
|-------|---------------------|---------------------|
| MVA   | Wild-type VACV strain MVA | 89 |
| WR    | Wild-type VACV strain WR | 90 |
| WR-Full-gB | Full-length HSV gB in TK | 51 |
| WR-ESmini-gB | ER-targeted minigene-gB_{498} in TK | 50 |
| MVA-Full-gB | Full-length HSV gB in TK | This study |
| MVA-ESmini-gB | ER-targeted minigene-gB_{498} in TK | This study |
| WR-TK | Insertional inactivation of TK | 69 |
| MVA-TK | Insertional inactivation of TK | This study |
| WR-Full-PB1-F2 | Full-length IAV PB1F2 in TK | 58 |
| WR-ESmini-PB1-F2 | Minigene-IAV PB1F2_{452} in TK | 53 |
| MVA-Full-PB1-F2 | Full-length IAV PB1F2 in TK | This study |
| MVA-ESmini-PB1-F2 | Minigene-IAV PB1F2_{452} in TK | This study |
| IAV-miniB8 | IAV PR8 with sequence encoding B8_{20} inserted into the neuraminidase stalk | This study |

| Virus | Recombinant details | Source or reference |
|-------|---------------------|---------------------|
| WR-delB8 | B8R deleted | 50 |
| WR-delB8-miniB8 | B8R replaced with minigene-B8_{20} | 85 |
| MVA-delB8 | B8R deleted | 91 |
| MVA-delB8-miniB8 | B8R replaced with minigene-B8_{20} | This study |
| WR-TK-OVA | Full-length OVA in TK | 41 |
| WR-TK-miniOVA | Minigene-OVA_{257} in TK | 92 |
| MVA-dellIII-OVA | Full length OVA in dellIII | 42 |
| MVA-dellIII-SIINFEKL | Minigene-OVA_{257} in dellIII | 42 |
| MVA-TK-OVA | Full-length OVA in TK | This study |
| MVA-TK-miniOVA | Minigene-OVA_{257} in TK | 37 |
| MVA-A11R/A12L-OVA | Full-length OVA in A11R/A12L | This study |
| MVA-A11/A12-SIINFEKL | Minigene-OVA_{257} in A11R/A12L | This study |
| MVA-dellIII-OVA-delTK | Full-length OVA in dellIII with insertional inactivation of TK | This study |
| MVA-dellIII-SIINFEKL-delTK | Minigene-OVA_{257} in dellIII with insertional inactivation of TK | This study |

*The original nonrecombinant parent virus is indicated by the first letters of each name: WR, MVA, or IAV. The order of the viruses is as they appear in the main text, and they are separated into groups according to the figures and legends in which they are described.*

November 2019 Volume 93 Issue 21 e01154-19
TABLE 2 Synthetic peptides used in this study

| Peptidea | Originb | Sequence | MHC | Reference(s) |
|----------|---------|----------|-----|--------------|
| A3270    | VACV, A3270–277 | KSYNYMILL | H-2Kb | 33 |
| A47138   | VACV, A47138–146 | AAEFINSL | H-2Kb | 61 |
| A47171   | VACV, A47171–180 | YAHINALEYI | H-2Kb | 55 |
| L253     | VACV, L253–61 | VIYIFTVL | H-2Kb | 33 |
| B820     | VACV, B820–27 | TSYKFEVL | H-2Kb | 61 |
| gB608    | HSV-1 gB608–505 | SSIEFARL | H-2Kb | 93 |
| OVA257   | Chicken, OVA257–264 | SIINEFKL | H-2Kb | 94, 95 |
| P81F22   | IAV strain PR8, P81F22–70 | LSLRPILV | H-2Dd | 58 |

Data from the first four peptides (A3270, A47138, A47171, and L253) were pooled and are referred to as P4.

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Author/s:
Wong, YC; Croft, S; Smith, SA; Lin, LCW; Cukalac, T; La Gruta, NL; Drexler, I; Tscharke, DC

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