Heterologous arenavirus vector prime-boost overrules self-tolerance for efficient tumor-specific CD8 T cell attack

Graphical Abstract

Highlights

- Engineered arenaviruses induce potent tumor self-specific CD8 T cell (CTL) response
- Combinations of distantly but not closely related arenavirus vectors eliminate tumors
- Vector backbone-targeted CTL responses compete against tumor self-reactive CTLs
- Optimized vector combinations reshuffle immunodominance to break self-tolerance

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In brief
Therapeutic tumor vaccination should break self-tolerance. Assessing combinations of engineered arenavirus vectors, Bonilla et al. find that distantly related vector combinations reshuffle T cell immunodominance hierarchies to break self-tolerance and eliminate established solid tumors, whereas closely related vectors interfere because of immunodominance of anti-vector CD8 T cells rather than antibodies.

Bonilla et al., 2021, Cell Reports Medicine 2, 100209
March 16, 2021 © 2021 The Author(s).
https://doi.org/10.1016/j.xcrm.2021.100209
Heterologous arenavirus vector prime-boost overrules self-tolerance for efficient tumor-specific CD8 T cell attack

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https://doi.org/10.1016/j.xcrm.2021.100209

SUMMARY

Therapeutic vaccination regimens inducing clinically effective tumor-specific CD8+ T lymphocyte (CTL) responses are an unmet medical need. We engineer two distantly related arenaviruses, Pichinde virus and lymphocytic choriomeningitis virus, for therapeutic cancer vaccination. In mice, life-replicating vector formats of these two viruses delivering a self-antigen in a heterologous prime-boost regimen induce tumor-specific CTL responses up to 50% of the circulating CD8 T cell pool. This CTL attack eliminates established solid tumors in a significant proportion of animals, accompanied by protection against tumor rechallenge. The magnitude of CTL responses is alarmin driven and requires combining two genealogically distantly related arenaviruses. Vector-neutralizing antibodies do not inhibit booster immunizations by the same vector or by closely related vectors. Rather, CTL immunodominance hierarchies favor vector backbone-targeted responses at the expense of self-reactive CTLs. These findings establish an arenavirus-based immunotherapy regimen that allows reshuffling of immunodominance hierarchies and breaking self-directed tolerance for efficient tumor control.

INTRODUCTION

Cytotoxic CD8+ T lymphocytes (CTLs) are central mediators of adaptive immunity. Tumor-infiltrating CTLs in several tumor types are associated with clinical outcome,1–5 and pre-existing CTL infiltration may predict responsiveness to immune checkpoint inhibition.4 Analogously, CTLs are key players in HIV elite control and hepatitis B virus clearance.5–7 Therapeutic vaccination for CTL induction holds great promise for cancer therapy,5,9 but has delivered inconsistent therapeutic benefits, including failure of large clinical trials.10–13 Despite induction of sizeable tumor antigen-specific CD8+ T cell frequencies by modalities such as adjuvanted peptides,14 inefficient tumor infiltration has curtailed the clinical efficacy of these cells.15,16 Delivering tumor-associated antigens (TAAs) in the context of virus-induced inflammation17 has significant potential to overcome these hurdles. Accordingly, several viral vector platforms have been developed for therapeutic use against solid tumors.13,18–25 The immunostimulatory properties of the viral particles themselves, exhibiting pathogen-associated molecular patterns, activate antigen-presenting cells (APCs) to augment and differentiate immune responses.26,27 In addition, certain replicating viral delivery systems trigger release of damage-associated molecular patterns or alarmins, such as interleukin-33 (IL-33).28,29 These signals critically augment activated T cell expansion, effector differentiation, and anti-tumor efficacy.30,31 Anti-vector immunity can inhibit viral delivery systems, impeding their re-administration to augment immune responses. A single immunization with vaccinia virus or modified vaccinia virus Ankara (MVA) elicits neutralizing antibody (nAb) responses in the majority of individuals35 and inhibits vaccinia-vectorized immunization.33 The seroprevalence of common adenovirus (Ad) serotypes such as Ad5 can regionally exceed 90%. Ad5-nAbs
have enabled the tailored design of this virus family and its use to infect the animals. Titers after 72 h are shown. Symbols represent individual virus cultures from one mouse each. Mean (N) BHK-21 cells were infected with viruses re-isolated from individual AGRAG mice on day 224 of the experiment shown in (H)–(J) or with the viral stocks originally depicted schematically in (M). IGR, S segment intergenic region; UTR, untranslated region.

For several decades, members of the Arenaviridae family have found widespread use in basic immunological research because of their capacity to induce CTL responses of exceptional magnitude, functionality, and longevity. Reverse genetic techniques have enabled the tailored design of this virus family and its exploitation for vaccination. Replication-deficient vectors (rLCMV) based on the prototypic arenavirus lymphocytic choriomeningitis virus (LCMV) have demonstrated excellent CTL and nAb induction against vectorized transgenes in mice, non-human primates, and, recently, also in humans. As an important differentiation from other viral vector technologies, rLCMV-based immunization only rarely induces vector-nAbs, facilitating the vector’s repeated administration in homologous prime-boost vaccination. This peculiarity of LCMV-based vectors is due to an N-linked “glycan shield” on the outer globular domain of the viral envelope glycoprotein domain that impairs antibody accessibility to critical neutralizing epitopes. LCMV vectors are currently in clinical phase 2 testing for prevention of cytomegalovirus-associated disease in transplant recipients.

We and others have developed replication-attenuated, tri-segmented, arenavirus-based vector formats (artARENA, r3ARENA; Figures 1B and 1C) that induce even more potent effector CTL responses than replication-deficient rLCMV or commonly used poxvirus and Ad vector systems. An LCMV-based artARENA vector (artLCMV) has recently entered early-stage clinical testing. When used to deliver tumor self-antigens to tumor-bearing mice, artLCMV induced potent anti-tumor CTL responses and extended the animals’ survival. Tumor-specific CTL responses were, however, lower in magnitude than those induced against vectorized non-self-antigens, and complete tumor remission was not achieved. This suggested that self-tolerance limited the therapeutic efficacy of artLCMV-based immunization.

Here we systematically explore a range of additional arenavirus vector backbones for vaccine delivery. We report that, contrary to commonly held concepts, anti-vector CTL responses rather than nAbs curtail the immunogenicity of homologous arenavirus vector prime-boost vaccination. Accordingly, backbones of distant genealogical relationship offered the most potent heterologous prime-boost combinations, resulting in considerable rates of complete tumor rejection.

RESULTS

Arenavirus vector backbone candidates and their genetic and phenotypic stability

Arenaviruses form enveloped particles and contain two segments of negative-stranded RNA. The large (L) segment encodes for the viral polymerase L and the matrix protein Z, whereas the short (S) segment carries the envelope glycoprotein (GP) and nucleoprotein (NP) genes, separated by intergenic regions, respectively (Figure 1A). We and others have incorporated transgenic sequences such as TAAAs into replicating arenavirus vectors by segregating the NP and GP genes onto artificially duplicated S segments (S\text{UP} and S\text{GP}; Figures 1B and S1C). This can be achieved by the artARENA (e.g., artLCMV) or r3ARENA (e.g., r3LCMV) design strategy (Figures 1B and 1C). Here we vectorized additional mammalian arenaviruses (mammarenaviruses) to exploit their immunotherapeutic potential when combined in heterologous prime-boost combinations. Based on phylogentic relationship, mammarenaviruses can be subdivided into the Old World group of viruses and four clades (A–D) of New World viruses (Figure 1D). For vector generation (STAR methods), we selected three mutually very distantly related viruses representing the main branches of the phylogenetic tree: the prototypic Old World virus LCMV, the widely studied New World clade A virus Pichinde virus (PICV), and the Junin virus vaccine strain Candidâ’ (CAND). CAND is in clinical use as a prophylactic vaccine against Junin virus, the causative agent of Argentine hemorrhagic fever, and PICV has no human disease correlate but can infect humans, as documented in accidentally exposed laboratory workers. LCMV infection is mostly asymptomatic or manifests as a flu-like infection. Rare cases of choriomeningitis are generally self-limiting and, despite documented cases of protracted central nervous system manifestations, commonly heal without persisting sequelae.

Figure 1. Arenavirus vector backbone candidates and their genetic and phenotypic stability

(A–C) Schematic of the genome organization of WT arenaviruses (A), artARENA vectors (B), and r3ARENA vectors (C). TAA, tumor-associated antigen.

(D) Genealogy tree of the mammarenavirus family with its clades. Red circles indicate viruses used in this study. The scale bar describes the expected number of mutations per site.

(E–G) Growth curves of the indicated viruses and vectors in BHK-21 cells (E and F) and 293T cells (G) infected at a multiplicity of infection (MOI) of 0.01. Symbols show the mean ± SD of three cell culture wells (error bars mostly within symbol size). **p < 0.01 by unpaired two-tailed Student’s t test.

(H–J) AGAG mice were infected intravenously (i.v.) with the indicated viruses, and viremia was monitored by immunofocus assays detecting PICV-NP (H) or GFP (I) to calculate the NP:GFP infectivity ratio (J). Symbols represent means ± SD of 3–5 mice.

(K–M) RT-PCR strategy (K) to amplify recombinated WT-like PICV S segment RNA species re-uniting NP and GP sequences. Also shown is a gel electrophoresis image of RT-PCR products (L) obtained from serum samples collected at the indicated time points. Each lane represents an individual mouse. Sera from uninfected mice (“none”) and water were included as negative controls. Sequence analysis of the bands numbered in (L) suggested recombination products, as depicted schematically in (M). IGR, S segment intergenic region; UTR, untranslated region.

(N) BHK-21 cells were infected with viruses re-isolated from individual AGAG mice on day 224 of the experiment shown in (H)–(J) or with the viral stocks originally used to infect the animals. Titers after 72 h are shown. Symbols represent individual virus cultures from one mouse each. Mean ± SD is indicated. Number of independent datasets (N) for (H)–(N) = 2. See also Figure S1.
perspective, all three viruses are valid vector backbone candidates for immunotherapy use in humans. We generated artARENA (artLCMV, artPICV, and artCAND) as well as r3ARENA (r3LCMV, r3PICV, and r3CAND) vectors expressing fluorescent reporter proteins. In cell culture, artARENA vectors as well as their respective r3ARENA counterparts reached lower titers than their respective parental wild-type (WT) viruses (Figures 1E–1G). These observations extended and generalized earlier findings of attenuated artARENA and r3ARENA vector growth.29,47,48

Genetic and phenotypic stability are key criteria for the clinical utility of replicating viral vector systems. We have reported previously that artLCMV stably retained its genome organization and transgene expression over extended periods of in vivo replication, whereas r3LCMV underwent inter-segmental recombination, reverting to a non-transgenic bisegmented WT-like virus.29 Here we used artPICV-GFP, r3PICV-GFP, and WT PICV (PICV-WT) to infect highly immunodeficient mice, which lack type I and type II interferon receptors and are devoid of T and B cells because of RAG deficiency (AGRAM mice). In the first 30 days of persistent infection, total viral loads (determined as PICV NP-expressing infectivity) in the blood of artPICV-GFP- and r3PICV-GFP-infected animals were similar and below those of PICV-WT-infected controls (Figure 1H). By day 70, however, r3PICV-GFP viremia exceeded the levels in artPICV-GFP-infected animals and increased continuously thereafter, eventually reaching levels equivalent to PICV-WT. Conversely, artPICV viremia remained consistently below PICV-WT controls. GFP transgene-expressing viral infectivity in r3PICV-GFP- and artPICV-GFP-infected mice was at comparable levels up to around day 70 of infection (Figure 1I). Thereafter it declined continuously in r3PICV-GFP-infected animals but remained stable in artPICV-infected animals. The resulting ratio of total PICV infectivity to GFP-transgenic infectivity documented that r3PICV-GFP progressively lost its transgene, whereas transgene expression by artPICV-GFP remained stable throughout the observation period of more than 200 days (Figure 1J). We performed RT-PCR to detect supposedly recombined WT-like S segments containing PICV NP and GP sequences (Figure 1K). Such RNA species were absent from artPICV-GFP-infected mice but detected consistently in the blood of r3PICV-infected animals (Figure 1L). Sequence analysis of amplicons revealed that some of them contained one or two GFP remnants flanked by partially or completely duplicated viral intergenic regions, identifying them as inter-segmental recombination products of the S\textsubscript{NP} and S\textsubscript{GP} segments of r3PICV-GFP (Figure 1M). On day 224 after infection, RT-PCR assays detected the S\textsubscript{GP} segments of artPICV-GFP but not the corresponding GFP-containing segments of r3PICV-GFP (Figures S1A–S1D), further supporting the notion of r3PICV-GFP transgene loss. When re-isolated after more than 200 days of persistent infection and propagated in cell culture, r3PICV-GFP reached titers equivalent to PICV-WT, whereas re-isolated artPICV-GFP growth was attenuated. This contrasted with the cell culture behavior of the inoculum of r3PICV-GFP and artPICV-GFP, both of which grew to lower titers than the corresponding PICV-WT (Figure 1N). These findings indicated that the genetic instability of r3PICV-GFP was accompanied by phenotypic reversion to PICV-WT-like growth, whereas artPICV-GFP was genetically and phenotypically stable. Analogous findings were made with CAND-based vectors. S\textsubscript{NP}-S\textsubscript{GP} recombination products and loss of GFP-containing segments was also observed in r3CAND-infected AGRAG mice but not in artCAND-infected animals (Figures S1E–S1K). Re-isolated r3CAND-GFP exhibited CAND-WT-like cell culture growth behavior (Figure S1L), whereas artCAND replication in AGRAG mice was at levels too low to allow virus re-isolation. These studies generalized the finding29 that artARENA vectors are genetically and phenotypically stable, whereas r3ARENA vectors are prone to transgene loss and phenotypic reversion to a WT-like virus.

artARENA vectors are attenuated in guinea pig and mouse pathogenesis models

Next we tested whether artPICV and artLCMV were attenuated in animal models. Guinea pigs were infected with titrated doses of PICV-WT, known to cause lethal disease in these animals, or with artPICV-E7E6 expressing a non-oncogenic fusion construct consisting of the HPV16 E7 and E6 open reading frames (ORFs)55 (see chart in Figure 2A). Animals receiving diluent were included as a further control. At PICV-WT doses of 3 × 10\textsuperscript{2} or 3 plaque-forming units (PFUs), three of eight animals reached humane endpoints (Figure 2A), a disease that was always accompanied by high-level viremia (>10\textsuperscript{3} PFUs; Figures 2B and 2C). At the highest PICV-WT dose (3 × 10\textsuperscript{4} PFUs), seven of eight guinea pigs developed high-level viremia and terminal disease (Figures 2A and 2D). In contrast, artPICV-E7E6 infection at 3 × 10\textsuperscript{2} PFUs was aviremic (Figure 2E), and doses of 3 × 10\textsuperscript{4} or even 3 × 10\textsuperscript{3} PFUs (100-fold higher than the highest PICV-WT dose tested) did not result in high-level viremia (Figures 2F and 2G). Transient low-level viremia was detected in only 1 of 8 animals in both of these latter cohorts (Figures 2F and 2G), and none of these animals developed terminal disease (Figure 2A). When administered 3 × 10\textsuperscript{2} PFUs of artPICV-E7E6, the lowest dose tested, seven of eight animals were free of disease throughout the 27-day observation period, and none of the animals had detectable viremia (Figure 2E). On day 24 after vector inoculation, one of these aviremic animals suddenly exhibited signs of disease corresponding to humane study endpoints, necessitating its euthanasia on day 25. The absence of detectable viral loads in this animal suggested, however, that its disease was unrelated to artPICV-E7E6 vector administration. Measurements of body weight loss, a commonly used parameter of PICV-induced disease in guinea pigs, provided additional independent support for the conclusion that artPICV-E7E6 was substantially attenuated (Figure S2).

Intracranial inoculation of mice is the standard model to assess the neurovirulence of LCVM-based vectors.29,41,56 In agreement with earlier data demonstrating attenuation of artLCMV vectors,29 we found that 100–1,000 PFUs of artLCMV-E7E6 had to be administered intracranially to elicit terminal choriomeningitis in about half of the animals, whereas 1 PFU of LCVM-WT resulted in terminal disease in all mice (Figure 2I). LCVM-induced choriomeningitis is mediated by CD8 T cells, which attack virus-infected meningeal cells and
astrocytes, resulting in blood-brain barrier (BBB) breakdown and brain edema. To investigate the mechanisms underlying reduced artLCMV neurovirulence, we infected WT mice intracranially with 10 PFUs of artLCMV-E7E6 or LCMV-WT or with diluent. Seven days later, at the peak of disease in LCMV-WT-infected animals, we determined immunoglobulin deposits in the brain parenchyma as a surrogate of BBB integrity (Figures 2J and 2K). The brain area affected by BBB breakdown was significantly larger in LCMV-WT- than in artLCMV-E7E6-infected mice (Figure 2L). Regions of dense immunoglobulin deposits were evident in LCMV-WT-infected brains, notably around the longitudinal fissure, with substantial extension into deeper cortical layers, whereas in artLCMV-E7E6-infected brains, only small immunoglobulin deposits were detected in proximity to the longitudinal fissure. This morphological correlate of reduced immunopathological damage furthers our understanding of attenuated artLCMV neurovirulence.

**Immunogenicity and epitope dominance in heterologous artARENA prime-boost vaccination**

Next we tested the utility of artPICV-artLCMV as a heterologous prime-boost regimen. We primed C57BL/6 mice intracranially with titrated doses of LCMV-WT or artLCMV-E7E6 as indicated and monitored them for signs of choriomeningitis ("survival").

To analyze the effect of viral infection on BBB permeability, animals were inoculated intracranially with 10 PFUs of LCMV-WT, 10 PFUs of artLCMV-E7E6, or with diluent. Seven days later, immunoglobulin G (IgG) in the brain parenchyma, indicating leakage across the BBB, was detected by peroxidase-based immunohistochemistry on histological sections. Representative images of coronary brain sections are shown (J), with an enlargement of computer-assisted detection of the IgG-positive surface (K). Quantitation of the detected area is shown in (L). Error bars in (L) show the mean ± SEM, and dots represent individual mice. Scale bars show 1,000 μm (J) and 500 μm (K).

See also Figure S2.
Figure 3. Immunogenicity and epitope dominance in heterologous artARENA prime-boost vaccination

(A–E) C57BL/6 mice were given artPICV-E7E6 and artLCMV-E7E6 homologous or heterologous prime-boost vaccination i.v. on day 0 and day 13. E7-tetramer-binding CD8+ T cell frequencies in the blood were determined at the indicated time points (A). Also shown are representative fluorescence-activated cell sorting (FACS) plots from blood gated on B220− lymphocytes analyzed on day 20 (B). Splenic frequencies of E7-specific (C) or NP396-specific (D) CD8+ T cells on day 51 served to calculate the epitope dominance ratio displayed in (E).

(F–J) BALB/c mice were given artPICV-P1A and artLCMV-P1A homologous or heterologous prime-boost vaccination i.v. on day 0 and day 39. P1A-tetramer-binding CD8+ T cell frequencies in the blood were determined at the indicated time points (F). Also shown are representative FACS plots from blood gated on B220− lymphocytes on day 49 (G). Frequencies of P1A-specific (H) or NP118-specific (I) CD8+ T cells in the blood on day 49 served to calculate epitope dominance ratios as displayed in (J).

Symbols in (A) and (F) represent the mean +/- SEM of five mice, except the “no vaccine” groups: 4 mice in (A) and 2 mice in (F). Symbols in (C), (D), (H), and (I) represent individual mice, and bars indicate the mean ± SD. Boxes in (E) and (J) display the minimal and maximal values. Numbers in (B) and (G) indicate the percentage of tetramer-binding cells among CD8+B220− T cells. See also Figures S3 and S4.
Figure 4. Phenotype of artARENA-induced CTLs and their dependence on IL-33-ST2 alarmin signaling

(A–F) We immunized BALB/c mice with artPICV-P1A and artLCMV-P1A in homologous or heterologous prime-boost vaccination i.v. on day 0 and day 27. On day 34, we analyzed P1A-Tet-binding and CD62L expression by splenic CD8+ T cells (A; gated on CD8+B220– lymphocytes). Unimmunized control mice are shown for comparison in (A) only. Numbers in (A) indicate the percentage of cells in the respective quadrant. Total P1A-Tet+ CTLs (B), P1A-specific effector/effector memory CTLs (CD62Llo; C), and P1A-specific central memory CTLs (CD62Lhi; D) were enumerated in the spleen on day 34. In both subsets of P1A-specific CTLs, CD62Lhi and CD62Llo, we determined the surface expression of KLRG1, CX3CR1, CD27, CD43, and CD127 as well as the master transcription factors Tcf-1, Tbet, and Eomes (E). Total numbers of marker-expressing P1A-specific CTLs were enumerated in (F). (A) shows representative FACS plots from individual mice. Symbols in (B)–(D) and (F) represent individual mice, and bars in (B)–(D) and (F) indicate the mean ± SD. Numbers in (A) and (E) indicate the percentage of gated cells (mean ±

(legend continued on next page)
upon peptide stimulation and comprised E7- as well as E6-specific CTLs (Figure S3). The majority of tumors, however, do not exhibit viral determinants such as HPV E7 and E6, and therefore active immunization has to rely on other classes of TAAAs for which immune responses can be affected by self-tolerance. Studies of the P815 mouse tumor cell-derived cancer-testis antigen P1A, for example, have shown that post-natal expression is restricted to spermatogonia, placenta, and thymic medullary epithelial cells, with the latter being key for central tolerance induction.65,66 Accordingly, P1A knockout mice spontaneously rejected P815 tumors, mounting P1A-specific CD8+ T cell responses of significantly higher magnitude and functional avidity than WT animals.67 To test whether heterologous artARENA immunization facilitates breaking self-tolerance, we immunized BALB/c mice with P1A-expressing artLCMV-P1A or artPICV-P1A. By day 38 after prime, the P1A-specific responses induced by either vector had leveled off in the range of 2%–3% of circulating CD8+ T cells (Figure 3F). Upon heterologous artLCMV-P1A boost, artPICV-P1A–primed mice mounted P1A-specific CTL responses exceeding 50% of the circulating CD8+ T cell pool, substantially higher than upon artLCMV-P1A homologous boost (Figures 3F–3H). These heterologous prime-boost-induced CTL responses contracted slowly, with frequencies of more than 20% persisting in peripheral blood for more than 3 months (Figure 3F). Besides vector transgene-specific responses (E7E6 and P1A), we also determined dominant vector backbone-specific responses directed against the NP-derived epitopes NP396 in C57BL/6 mice (H-2Db restricted) and NP118 in BALB/c mice (H-2Ld restricted). Interestingly, the responses of C57BL/6 and BALB/c mice, respectively, were substantially lower in animals receiving heterologous artPICV-artLCMV immunization compared with homologous artLCMV prime-boost (Figures 3D and 3I). Based on these measurements we calculated the transgene:backbone epitope dominance ratio (E7:NP396 [Figure 3E]; P1A:NP118 [Figure 3J]). It was substantially higher in heterologous compared with homologous prime-boost immunization (~50-fold in Figure 3E; >200-fold in Figure 3J), indicating that heterologous prime-boost immunization biased vaccine-induced CTL responses toward transgene-derived epitopes. In the context of active immunization for immunotherapy, it is important to induce high-frequency CTL responses within a short time window. artPICV-P1A-artLCMV-P1A heterologous prime-boost, administered at an interval of only 4–10 days, induced P1A-specific CTL responses of higher frequencies than obtained with homologous prime-boost immunization given at the same interval (Figures S4A–S4D). A trend of higher P1A-specific CTL frequencies at longer prime-boost intervals (7–10 days compared with 4 days) was accompanied by an inverse trend in NP118 backbone-targeting responses. artPICV-P1A prime followed by artLCMV-P1A boost clearly outperformed the inverse sequence of administration when administered at intervals of 4–10 days (Figures S4A–S4D). The prime-boost regimen of artPICV followed by artLCMV rather than the inverse sequence of administration was therefore used for subsequent studies. Only when the interval between the two vaccinations was substantially longer was the artPICV-P1A boost of artLCMV-P1A–primed responses effective (Figures S4E and S4F).

**Phenotype of artARENA-induced CTLs and their dependence on IL-33-ST2 alarmin signaling**

Next, we determined how heterologous artPICV-artLCMV prime-boost influenced the magnitude and phenotype of P1A-specific CTL populations. Seven days after boost, CD62Llo effector/effectort memory cells dominated the responses of BALB/c mice to homologous artLCMV-P1A prime-boost and heterologous artPICV-P1A-artLCMV-P1A immunization (Figures 4A–4D). Importantly, heterologous prime-boost elicited not only higher P1A-specific CD62Llo effector/effectort memory CTL populations than homologous prime-boost but the CD62Lhi central memory population was also more abundant (Figures 4C and 4D). We studied the phenotype of both CTL subsets by determining the cells’ expression of the surface markers KLRG1, CX3CR1, CD27, CD43, and CD127 and of the transcription factors Tcf-1, Tbet, and Eomes (Figures 4E and S5G). Within the CD62Lhi and CD62Llo subsets of CTLs, heterologous prime-boost immunization promoted expression of the effector differentiation markers KLRG1, CX3CR1, and CD43. 63–67 with a concomitant reduction in the proportion of cells expressing the memory markers CD27 and CD127.65,66 We also observed that CTLs emerging from heterologous prime-boost expressed higher average Tbet levels and lower levels of Tcf-1, further supporting the conclusion that heterologous prime-boost augmented the effector differentiation of CTLs. Irrespective of this relative effector differentiation bias, heterologous prime-boost augmented not only the total number of CTLs expressing the effector differentiation markers KLRG1, CX3CR1, and CD43 but also the population of cells expressing memory precursor markers (CD27 and CD127) and the stemness-defining transcription factor Tcf-1 (Figure 4F). 65 When analyzed 4 weeks after boost (Figures S5A–S5F), most of the differences between CTLs emerging from homologous and heterologous prime-boost persisted but, overall, were less pronounced than on day 7 after boost. Prominent populations of effector-like memory populations, characterized by the marker combinations CD43+, CX3CR1+, and KLRG1+CD27+, were particularly abundant upon heterologous prime-boost immunization (Figure S5G). These observations indicated that CTL responses induced by heterologous prime-boost differed from those emerging from homologous prime-boost primarily by their higher cellularity paired with more pronounced effector differentiation.

SD or the mean fluorescence intensity (MFI ± SD). Means were calculated from six mice per immunization group (A–F) or from three unimmunized controls (A).

N = 2. **p < 0.01 by unpaired two-tailed Student’s t test.

(G and H) We immunized ST2−/− and WT mice with artLCMV-E7E6, artPICV-E7E6 or arCAND-E7E6 i.v. Controls were left unimmunized (“no vaccine” in G). E7-tetramer-binding cells in blood were determined on day 7. Representative FACS plots are shown in (G); values indicate E7-Tet+CD8+ T cells as a percentage of lymphocytes. Symbols in (H) represent individual mice (n = 5 per group) with mean ± SD. N = 2. **p < 0.01 by two-way ANOVA with Sidak’s post-test.

See also Figures S5 and S6.
IL-33 signals through its receptor ST2 are key for protective CTL responses to replicating viruses, and IL-33 signals critically augment CTL responses to artLCMV-vectorized immunization and the resulting tumor control. Here, we compared the ST2 dependence of artARENA-induced CTL responses by immunizing ST2-deficient and WT control mice with artLCMV-E7E6, artPICV-E7E6, or artCAND-E7E6. artLCMV- and artPICV-induced CTL responses to E7 were significantly lower in the blood of ST2-deficient mice than in WT controls (Figures 4G and 4H). Conversely, ST2 deficiency had no clear effect on artCAND-induced CTL responses. ST2-dependent differences in artLCMV- and artPICV-induced CTL responses were also evident when enumerating IFN-γ and TNF-α co-producing CTLs in the spleen (Figure S6). These findings indicated that, besides artLCMV, artPICV- but not artCAND-induced CTL responses also benefited from IL-33 signaling.

Genealogic artARENA vector backbone relatedness dictates interference by pre-existing immunity and potency in heterologous prime-boost immunization

Our observations shown in Figures 3E and 3J suggested that heterologous artARENA prime-boost immunization overruled the immunodominance of vector backbone-directed CTL responses to focus immune responses on vaccine targets. To systematically investigate this hypothesis, we preimmunized mice with LCMV-WT, PICV-WT, CAND-WT, or WT Mopeia virus (MOPV-WT) and studied the animals’ ability of responding to artLCMV-E7E6 vaccination 1 month later. MOPV is an Old World mammarenavirus and, thus, has a close genealogic relationship with LCMV, whereas the New World viruses PICV and CAND are only distantly related to LCMV (Figure 1D). Because of epitope sequence homology between LCMV and MOPV (Figure 5A), both viruses elicited clearly detectable CTL responses to NP396 whereas the more distantly related PICV-WT and CAND-WT did not (Figure 5B). Pre-existing immunity to LCMV or MOPV almost completely abrogated E7-specific CTL induction by artLCMV-E7E6, whereas the responses of PICV-WT- or CAND-WT-immune mice were only modestly below those of control mice without prior arenavirus infection (Figure 5C). This interference by LCMV-WT and MOPV-WT immunity was accompanied by high-frequency NP396-directed responses and a biased E7:NP396 immunodominance hierarchy upon artLCMV-E7E6 vaccination (Figures 5D and 5E). We relied on the immunodominant NP396 epitope as an indicator of backbone cross-reactivity, which likely comprised additional epitopes in the four viral backbone proteins NP, GP, L, and Z.

Next, we tested whether these interference and immunodominance hierarchies correlated with the immunogenicity of heterologous artARENA vector prime-boost combinations. To further expand the quiver of arthARENA vectors, we generated a reverse genetic system for MOPV and, based thereupon, developed an artMOPV-P1A vector. Cell culture experiments demonstrated attenuated growth of artMOPV-P1A compared with its parental virus (Figure S7), analogous to the artARENA vectors described in Figures 1E–1G. We compared the ability of artLCMV-P1A to boost P1A-specific CTL responses induced by artPICV-P1A, artCAND-P1A, or artMOPV-P1A in BALB/c mice. Animals undergoing homologous artLCMV-P1A, artPICV-P1A, or artCAND-P1A prime-boost immunizations served as comparators and, after boost, remained in the 10% range or below (Figure 5F). Among the heterologous artARENA vector combinations, artPICV-P1A- and artCAND-P1A-primed animals were boosted efficiently by artLCMV-P1A, reaching frequencies in the 20%–30% range. Conversely, P1A-specific CTL responses of artMOPV-P1A-primed and artLCMV-P1A-boostered animals remained below 3%, failing to exceed the responses upon homologous artLCMV-P1A prime-boost. This indicated that the immunogenic benefit of heterologous prime-boost over homologous prime-boost was abolished when the closely related MOPV and LCMV vector backbones were combined. The above hierarchy of heterologous prime-boost combinations correlated inversely with the LCMV NP118-specific CTL responses to prime and boost (Figure 5G). artPICV-P1A and artCAND-P1A prime did not induce NP118-specific CTLs above technical background and repressed NP118-directed responses upon artLCMV-P1A boost. Conversely, artMOPV-P1A-primed and artLCMV-P1A-boostered animals had more than 50% NP118-specific CD8+ T cells in peripheral blood with a corresponding P1A:NP118 epitope dominance ratio of less than 0.1 (Figure 5H). Thus, genealogic vector backbone relatedness and vector backbone-biased CTL responses correlated with inefficient induction of P1A-specific responses.

Interference by vector backbone-specific CTLs rather than by nAbs

Next, we investigated the induction and cross-reactive neutralizing activity of artARENA vector-induced antibody responses. Homologous prime-boost immunization with artCAND induced sizeable vector-nAb titers (Figure 6A). We failed to detect artLCMV- or artPICV-nAb after homologous prime-boost immunization (Figure 6A), and neither artLCMV nor artPICV or artCAND immune sera cross-neutralized any of the other viruses. nAb induction by artCAND, but not artLCMV or artPICV, was in line with differentially dense glycan shields on the respective viruses’ envelope proteins.

These findings argued against vector-nAb as a limiting factor in artLCMV- or artPICV-based homologous or heterologous prime-boost immunization. To formally rule out antibody-mediated inhibition, we performed homologous and heterologous prime-boost immunizations in B cell-deficient mice (JHT mice) and WT control mice. E7-specific responses to artPICV-E7E6 and artLCMV-E7E6 prime were indistinguishable in the two strains of mice (Figure 6B). More importantly, however, JHT and WT mice had a virtually identical response pattern to homologous artLCMV-E7E6 or heterologous artPICV-E7E6 boost, respectively. The superior immunogenicity of heterologous compared with homologous prime-boost despite a lack of anti-vector antibodies in JHT mice excluded anti-vector antibody responses as a major limitation in homologous artLCMV prime-boost immunization (Figure 6B).

To address whether anti-vector CTL responses, independently of other components of anti-vector immunity, can interfere with artLCMV-based vaccination, we “preimmunized” mice with a recombinant vaccinia virus expressing the NP396 epitope as a minigene (VACC-NP396). Control animals were either given vaccinia virus expressing an irrelevant transgene...
Figure 5. Genealogic artAREN vector backbone relatedness dictates interference by pre-existing immunity and potency in heterologous prime-boost immunization

(A) Alignment of the LCMV NP118-126 and NP396-404 epitope sequences with the respective homologous sequences in the Old World arenavirus MOPV and the New World arenaviruses PICV and CAND. Epitope scores were predicted by the SYFPEITHI algorithm. Major histocompatibility complex (MHC) anchor positions are boxed, and amino acids corresponding to consensus anchor residues for H-2L^d and H-2D^d, respectively, are shown in bold. 

(B–E) We preimmunized C57BL/6 mice on day 0 with 10^5 PFUs of MOPV-WT, PICV-WT, CAND-WT, or LCMV-WT i.v. or left them uninfected. On day 9, we determined NP396-specific CD8^+ T cell frequencies in peripheral blood (B). On day 30, all mice were vaccinated with artLCMV-E7E6. Frequencies of E7-specific (C) and NP396-specific (D) CD8^+ T cells in the blood on day 38 served to calculate the epitope dominance ratio as displayed in (E). Symbols in (B)–(D) represent individual mice, and bars in (B)–(D) show their mean ± SD.

(F–H) We immunized BALB/c mice with 10^5 PFUs of artARENA-P1A vectors i.v. in various homologous and heterologous prime (day 0) to boost (day 35) combinations as indicated. On day 32 after prime and on day 55 (20 days after boost), we determined the frequencies of P1A-specific (F) and NP118-specific (G) CD8^+ T cells in the blood and calculated the P1A:NP118 epitope dominance ratio on day 55 (H). The lack of NP118-specific responses over technical background in artPICV-artPICV and artCAND-artCAND immunized mice precluded this latter assessment ("n.a."). Bars in (F) and (G) represent the mean ± SEM of 3 (artPICV-artPICV) to 4 mice (other groups). Boxes in (E) and (H) display the minimal and maximal values. N = 2.

See also Figure S7.
(Vacc-lacZ) as preimmunization or were left unimmunized. As expected, VACC-NP396 preimmunization, but not VACC-lacZ, induced a CTL response to NP396 (Figure 6C), whereas both viruses triggered CTLs to the immunodominant vaccinia virus backbone epitope B8R20–27 (Figure 6D). When subsequently vaccinated with artLCMV-E7E6, the E7-specific CTL responses of VACC-NP396-preimmune animals were 7- to 8-fold lower than those of VACC-lacZ-preimmune mice or controls not previously exposed to vaccinia virus (“none”). On day 15, we determined LCMV NP396-specific (C) and vaccinia BBR-specific (D) CTLs in the blood by MHC class I tetramer staining. On day 18, i.v. artLCMV-E7E6 vaccination was performed. Eight days later (day 26), we determined E7-specific (E) as well as NP396-specific (F) CD8+ T cell frequencies in the blood and calculated the E7:NP396 epitope dominance ratio (G). Bars represent the mean ± SEM of 5 mice per group. Boxes display the minimal and maximal values. N = 2.

Heterologous artARENA vector immunotherapy increases TIL numbers and tumor cure rates, resulting in long-term anti-tumor immunity

To assess whether the augmented immunogenicity of heterologous artARENA vector prime-boost translated into superior therapeutic efficacy, we exploited two transplantable syngeneic mouse tumor models. The P815 mastocytoma cell line is derived from a DBA/2 mouse and expresses the P1A cancer-testis antigen, whereas the C57BL/6-derived TC-1 cell lines serves as a model of HPV16 E7E6-expressing cancer. We implanted P815 and TC-1 tumors subcutaneously into the flank of mice and initiated artPICV or artLCMV vector therapy when tumors were palpable (P815, day 9) or had reached an average critical volume (C24100 mm3; TC-1, day 8). artPICV-artLCMV heterologous therapy delivering the respective tumor antigen, P1A or E7E6, was compared with homologous artPICV-artPICV or artLCMV-artLCMV prime-boost, all administered at an interval of 7 days (P815 model) or 10 days (TC-1 model). Homologous prime-boost with TAA-expressing artLCMV or artPICV extended...
Figure 7. Heterologous artARENA vector immunotherapy increases TIL numbers and tumor cure rates, resulting in long-term anti-tumor immunity

(A–L) We implanted tumor cells subcutaneously in the flanks of mice. (A), (C), (E), and (G–L): DBA/2 mice, P815 tumor cells. (B), (D), and (F): C57BL/6 mice, TC-1 tumor cells. When animals exhibited palpable tumor masses on day 9 (P815) or when tumors had reached an average volume of ~100 mm³ on day 8 (TC-1), mice were assigned randomly to the indicated prime-boost regimens administered i.v. on day 9 and day 16 (P815) or day 8 and day 18 (TC-1).

(A and B) Symbols show tumor volumes (mean ± SEM) from two (A) and one (B) independent experiments. The curves end when more than 50% of animals in a group have reached humane endpoints. Entire tumor volume curves were statistically compared as described in STAR methods.

(C and D) Kaplan-Meier survival curves of the animals, with median survival and percent long-term survivors indicated.

(A and C) n = 21 (untreated), 21 (artLCMV-P1A-artLCMV-P1A), 8 (artPICV-P1A-artPICV-P1A), 22 (artPICV-P1A-artLCMV-P1A), and 8 (artPICV-GFP-artLCMV-GFP).

(E and F) Animals, which had rejected their tumors ("cured") and tumor-naive controls were re-challenged subcutaneously on day 160 and day 140 (E) or day 118 (F) after primary tumor implantation. Re-challenged mice did not form palpable tumors (data not shown).

(E) Combined data from tumor-free mice (n = 5) and tumor-naive mice (n = 10) in two independent experiments.

(F) Tumor-free mice (n = 3) and tumor-naive mice (n = 5) from one experiment.

(legend continued on next page)
the survival of tumor-bearing mice, albeit to a lesser extent than heterologous prime-boost, and all mice receiving homologous prime-boost immunization eventually reached humane endpoints (Figures 7C and 7D). In contrast, 18 percent of mice with P815 tumors and 37.5 percent of animals with TC-1 tumors rejected their respective tumors when undergoing heterologous TAA-vectorizing artPICV-artLCMV immunotherapy (P1A and E7E6 but not GFP), resulting in the animals’ survival for more than 125 days (Figures 7C and 7D). When these long-term survivors were re-challenged with the same tumor cells, no tumor growth was recorded, whereas tumor- and therapy-naïve control animals progressed rapidly to humane endpoints (Figures 7E and 7F). This observation indicated that elimination of established tumors by artPICV-artLCMV immunotherapy resulted in long-term anti-tumor immunity.

We enumerated and characterized tumor-infiltrating CTLs on day 5 after homologous or heterologous artARENA vector immunotherapy of P815 tumors (Figures 7G–7L). Heterologous artPICV-P1A-artLCMV-P1A immunotherapy did not substantially augment total tumor-infiltrating CTL numbers compared with homologous immunotherapy with either one of these vectors (Figure 7G), but P1A-specific CTLs in tumors were significantly more numerous upon heterologous compared with homologous artARENA immunotherapy (Figure 7H). In line with this observation, we found an increase in proliferating (Ki67+) and granzyme B-expressing (cytotoxic) P1A-specific, tumor-infiltrating CTLs in mice with heterologous artARENA immunotherapy (Figures 7I and 7J). Heterologous artPICV-P1A-artLCMV-P1A therapy triggered significantly less NP118-specific TILs than homologous artLCMV-P1A prime-boost vaccination (Figure 7K), with a corresponding shift in P1A:NP118 epitope dominance in the tumor (Figure 7L). These data indicated that heterologous artARENA vector immunization shifted the TAA:vector backbone immunodominance patterns of tumor-infiltrating CTLs.

**DISCUSSION**

Elicitation of clinically effective tumor-specific CTL responses remains an important unmet medical need. Here we identify epitope dominance and interference by vector backbone-directed T cell responses as an important hurdle in vectored vaccine delivery. Based on this concept in conjunction with arenaviral genealogy analyses, we developed potent heterologous artARENA-based immunization regimens inducing effective anti-tumor immunity.

Therapeutic cancer vaccination is a particularly demanding field of active immunization. Induction of high-frequency CTL responses against TAA is often encumbered by central as well as peripheral tolerance mechanisms, limiting the available T cell repertoire and its responsiveness. Additional challenges arise from chronic antigenic exposure, which can lead to functional adaptation or impairment of specific T cell responses. In this context, potent viral vector systems and their optimal combination are of particular importance. Although poxvirus delivery systems have been evaluated clinically as cancer vaccines for more than 2 decades, a wide variety of viral vector systems has entered clinical testing in recent years. These approaches comprise alphavirus vectors, human and sAd-based vectors, lentivirus delivery systems, as well as combinations thereof, attesting to the promise of and broad interest in virally vectored cancer vaccination.

Animal studies have shown that arenavirus vectors are immunogenic when administered intravenously, intradermally, subcutaneously, or intramuscularly, and the latter route has also been validated clinically in a human trial. Earlier studies from our lab revealed, however, that the IL-33-ST2 alarmin pathway contributes essentially to the effectiveness of artARENA-vectorized cancer immunotherapy, and the vector’s ability to trigger this pathway correlates with its spread into IL-33-expressing splenic stromal cells. In line with other investigators’ work, these findings highlight the importance of vaccine delivery to specialized compartments of secondary lymphoid organs. The intravenous route provides optimal access to these tissues and, therefore, has been used here; it is also exploited in an ongoing clinical trial for artLCMV-based therapy of HPV16-positive head and neck cancer.

A discriminating feature of artLCMV and artPICV vector technology is the lack of vector-nAb induction because of the viral glycan shield and globally low arenavirus seroprevalence. The reason for the latter is the natural host range of mammarenaviruses being restricted to rodents, combined with only rare transmission from rodents to humans. Although they do not elicit nAbs against their own glycan-shielded envelope protein, replication-deficient rLCMV vectors elicit potent antibody immunity against vectorized cargo. While future work should investigate the utility of artARENA vector technology for prophylactic antibody induction against infectious diseases, potent CD8 T cell responses position heterologous artARENA vector prime-boost immunization as a promising strategy for therapeutic vaccination in persistent microbial diseases, notably HIV and hepatitis B.

Our findings highlight anti-vector T cell immunity as a mechanism of interference in artARENA vector homologous prime-boost vaccination. For delivery systems such as Ad and poxvirus vectors, which readily elicit potent vector-nAbs, the latter are commonly taken as surrogate and supposed main mechanisms of interference by anti-vector immunity. Several observations suggest, however, that pre-existing anti-vector T cell responses impede responses to Ad and poxvirus vector systems, too. Inefficient responses to MVA-vectorized TAA has been credited to epitope dominance and competition by vector backbone-directed T cell responses. Similarly, the inhibitory activity of Ad vector-nAbs may have obscured the contribution of additional interference mechanisms. Certain pairs of adenoviral vectors, although serologically distinct, failed to effectively boost each other. This indicates that the mere absence of vector-nAbs cannot predict efficient boosting. Conversely, pre-existing Ad-specific T cell immunity is correlated inversely with
HIV-specific CTL induction by rAd5 vectors in human vaccine trials. Pre-existing T cell reactivity to Ad5 is found in more than 80% of healthy adults, irrespective of Ad5-specific serostatus. It originates from prior exposure to Ads of unrelated serotypes, targets epitopes in conserved regions of the viral genome, and expands significantly upon rAd5-vectored vaccination. These considerations highlight advantages of viral vector platforms based on virus families such as Arenaviridae, which circulate almost exclusively in the animal kingdom.

Limitations of study
We relied on the transgene:backbone epitope dominance ratio of CD8 T cell responses as an indicator and correlate of backbone-directed T cell interference. Besides inter-clonal competition of CD8 T cell responses as an indicator and correlate of backbone-directed T cell interference. Besides inter-clonal competition of transgene- and backbone-specific T cells, an alternative bone-directed T cell interference. We relied on the transgene:backbone epitope dominance ratio as an indicator and correlate of backbone-directed T cell interference. Besides inter-clonal competition of transgene- and backbone-specific T cells, an alternative bone-directed T cell interference.

The present data obtained for mice may imperfectly predict efficacy in humans, but they call for immediate clinical evaluation of heterologous artPICV-artLCMV prime-boost immunization regimens to benefit patients. artPICV-E7E6 and artLCMV-E7E6 are currently entering clinical phase 1 testing as a repeated alternating prime-boost immunization regimen for HPV16-positive head and neck cancer. Irrespective of the limited predictive value of transplantable mouse tumor models, our findings regarding potent CTL induction to self-antigens such as P1A suggest that self-tolerance can be broken efficiently and provide an incentive for our plan of applying artARENA vector technology to a broader range of cancers, most of which do not express viral target antigens. The arenavirus vector platform accommodates transgenes up to ~2,000 bp, lending itself for delivery of all types of proteinaceous cancer targets, but it remains unknown which ones provide the best clinical efficacy.

STAR METHODS
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SUPPLEMENTAL INFORMATION
Supplemental Information can be found online at https://doi.org/10.1016/j.xcrm.2021.100209.

ACKNOWLEDGMENTS
We wish to thank Nadège Lagarde, Beatrice Dolder Schlienger, Felix Stemmeseder, and Sophie Schulha for excellent technical assistance and Theresa Kleissner, Goran Bekic, Daniel Oeler, and Sonja Feher for diligent implementation of animal studies. This work was supported by the Swiss National Science Foundation (310030, 185318/1 to D.D.P., CRSII3_160772/1 to D.D.P. and D.M., 310030, 173010 to D.M., and CRSII5_170929 to A.Z.), Hookipa Pharma, Inc. (to D.D.P., A.Z., and D.M.), and by Austrian Research Promotion Agency (Österreichische Forschungsförderungsgesellschaft [FFG]).

AUTHOR CONTRIBUTIONS
W.V.B., N.K., A.-F.M., S.M.K., S.S., J.R., U.B., I.V., S.G., C.M., D.M., K.K.O., A.Z., and D.D.P. contributed to experimental conception and design. W.V.B., N.K., A.-F.M., S.M.K., M.A.K., M.L., S.D., J.R., I.V., M.P., R.K., S.H., and C.M. performed experiments. W.V.B., N.K., A.-F.M., S.M.K., S.S., J.R., U.B., I.V., M.P., C.M., D.M., K.K.O., A.Z., and D.D.P. analyzed and/or interpreted the data. W.V.B., N.K., A.Z., and D.D.P. wrote the manuscript.

DECLARATION OF INTERESTS
S.S., J.R., U.B., and K.K.O. are employees and stock option holders of Hookipa Pharma, Inc. D.D.P. is a founder, consultant, shareholder, and stock option holder of Hookipa Pharma, Inc. W.V.B., S.M.K., S.S., U.B., D.D.P., and D.M., and K.K.O. are listed as inventors on patents describing artARENA vector technology. W.V.B. and D.D.P. are married.

Received: September 15, 2020
Revised: December 16, 2020
Accepted: February 4, 2021
Published: March 3, 2021

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## STAR METHODS

### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| Rat monoclonal anti-| Dr. D.D. Pinschewer Battegay et al. 91 | PMID: 1939506 |
| LCMV-NP: VL4        |        |            |
| Rat monoclonal anti-| Biolegend | Cat# 338002; RRID: AB_1279414 |
| GFP                 |        |            |
| Nakauchi et al. 92  | PMID: 19553554 | |
| mouse monoclonal anti-| Nakauchi et al. 92 | PMID: 19553554 |
| PICV-NP: 17.2.E4-2  |        |            |
| mouse monoclonal anti-| Hufert et al. 93 | PMID: 2476109 |
| CAND-NP: 17.1.C6.0  |        |            |
| mouse monoclonal anti-| Biolegend | Cat# 562611 |
| MOPV-NP: 2B5       |        |            |
| Rat monoclonal anti-| BD Biosciences | Cat# 553088 |
| mouse CD8a; Clone 53-6.7 |        |            |
| Syrian Hamster monoclonal anti-| Biolegend | Cat# 138418 |
| mouse KLRG1; Clone 2F1/KLRG1 |        |            |
| Rat monoclonal anti-| Biolegend | Cat# 135027 |
| mouse CD127; clone A7R34 |        |            |
| mouse monoclonal anti-| Biolegend | Cat# 149031 |
| mouse CX3CR1; clone SA011F11 |        |            |
| Armenian Hamster monoclonal anti-| Biolegend | Cat# 124216 |
| mouse CD27; clone LG3A10 |        |            |
| Rat monoclonal anti-| Biolegend | Cat# 121214 |
| mouse CD43; clone 1B11 |        |            |
| mouse monoclonal anti-| Thermo Fisher | Cat# MHGB04 |
| mouse Granzyme B; clone GB12 |        |            |
| Rat monoclonal anti-| Thermo Fisher | Cat# 56-5698-80 |
| mouse IFNg; clone XMG1.2 |        |            |
| Rat monoclonal anti-| Biolegend | Cat# 505810 |
| mouse TNFa; clone MP6-XT22 |        |            |
| Rat monoclonal anti-| Biolegend | Cat# 506324 |
| mouse IL2; clone JES6-5H4 |        |            |
| Rat monoclonal anti-| BD Biosciences | Cat# 553310 |
| mouse CD11b; clone M1/70 |        |            |
| Armenian Hamster monoclonal anti-| BD Biosciences | Cat# 553801 |
| mouse CD11c; clone HL3 |        |            |
| Rat monoclonal anti-| BD Biosciences | Cat# 553785 |
| mouse CD19; clone 1D3 |        |            |
| Rat monoclonal anti-| Biolegend | Cat# 137605 |
| mouse NKp46; clone 29A1.4 |        |            |
| Rat monoclonal anti-| Biolegend | Cat# 100548 |
| mouse CD4; clone RM4-5 |        |            |
| Rabbit monoclonal anti-| Cell Signaling | Cat# 11/2014 |
| mouse TCF1; clone C63D9 |        |            |
| Donkey polyclonal anti-| Biolegend | Cat# 406410 |
| mouse Poly4046 |        |            |
| Rat monoclonal anti-| eBioscience | Cat# 50-4875-82 |
| mouse Eomes; clone Dan11mag |        |            |
| mouse monoclonal anti-| Thermo Fisher | Cat# 12-5825-82; RRID: AB_925761 |
| mouse T-bet; clone eBio4B10 (4B10) |        |            |

(Continued on next page)
| REAGENT or RESOURCE NAME | SOURCE | IDENTIFIER |
|--------------------------|--------|------------|
| EnVision+ System-HRP Labeled Polymer | Dako | Cat# K4001 |

**Bacterial and virus strains**

| Strain Name | Source | PMID |
|-------------|--------|------|
| LCMV-WT | Sommerstein et al. | 26587982 |
| artLCMV-GFP/RFP | Kallert et al. | 28548102 |
| r3LCMV-GFP/RFP | Kallert et al. | 28548102 |
| artLCMV-P1A | Kallert et al. | 28548102 |
| artLCMV-E7E6 | This paper | N/A |
| PICV-WT | This paper | N/A |
| artPICV-GFP/RFP | This paper | N/A |
| r3PICV-GFP/RFP | This paper | N/A |
| artPICV-P1A | This paper | N/A |
| artPICV-E7E6 | This paper | N/A |
| CAND-WT | This paper | N/A |
| artCAND-GFP/RFP | This paper | N/A |
| r3CAND-GFP/RFP | This paper | N/A |
| artCAND-P1A | This paper | N/A |
| artCAND-E7E6 | This paper | N/A |
| MOPV-WT | This paper | N/A |
| artMOPV-P1A | This paper | N/A |
| VACC-NP396 | Probst et al. | 14607945 |
| VACC-lacZ | Ludewig et al. | 10704461 |

**Chemicals, peptides, and recombinant proteins**

| Name | Source | Identifier |
|------|--------|------------|
| P1A-Tetramer-PE: H2-Ld-LPYLGWLVF | Tetramer Core facility University Lausanne | TA P815 35-43 PE |
| LCMV-NP(396-404)-Tetramer-PE: H2-Db-FQPONGQFI | NIH Tetramer Core Facility at Emory University | N/A |
| LCMV-NP(118-126)-Dextramers: H2-Ld-RPQASVGYYM | Immudex, Denmark | JG2750-APC |
| HPV-E7(49-57)-Dextramers: H2-Db-RAHYNIVTF | Immudex, Denmark | JA219S-APC |
| VACC-B8R(20-27)Tetramer-PE: H-2Kb-TSYKFESV | Tetramer Core facility University Lausanne | MVA-B8R-20-27-PE |
| HPV-E7 peptide set: 22 peptides (peptide scan 15/11) | JPT | HPV16-E7 overlapping peptides |
| HPV-E6 peptide set: 37 peptides (peptide scan 15/11) | JPT | HPV16-E6 overlapping peptides |
| Lipofectamine 2000 Transfection Reagent | Thermo Fisher / Invitrogen | Cat# 11668019 |
| TRI Reagent | Sigma Aldrich / Merck | T9424 |

**Critical commercial assays**

| Name | Source | Cat# |
|------|--------|------|
| QIAamp-Viral RNA-Mini | QIAGEN | 52906 |
| QIAGEN OneStep RT-PCR Kit | QIAGEN | 21020 |
| SuperScript IV First-Strand Synthesis System | Thermo Fisher | 18091050 |
| Zombie UV Fixable Viability kit | Biolegend | 423108 |
| eBioscienceFoxp3/ Transcription Factor Staining Buffer Set | Thermo Fisher | 00-5523-00 |

**Experimental models: cell lines**

| Name | Source | Cat# |
|------|--------|------|
| Hamster: BHK 21 (clone 13) cells | ECACC | 85011433 RRID: CVCL_1915 |
| Hamster: BHK 21-GP cells | Flatz et al. | 20139992 |
| Human: HEK293T cells | ECACC | 12022001 RRID: CVCL_0063 |

(Continued on next page)
**RESOURCE AVAILABILITY**

**Lead contact**
Further information and requests for resources and reagents should be directed to the Lead Contact, Daniel D. Pinschewer (Daniel.Pinschewer@unibas.ch).

**Materials availability**
Material transfer agreements with standard academic terms will be established to document reagent sharing by the lead contact’s institution. artPICV vector materials will be supplied by Hookipa Biotech GmbH under MTA.”

**Data and code availability**
The accession number for the data reported in this study is Zenodo: https://doi.org/10.5281/zenodo.4488070.

**Experimental model and subject details**

**Animals and ethics statement**
AGRAM mice (IFNα/βR−/−, IFNγR−/−, RAG1−/− triple-deficient), B cell-deficient JHT mice and ST2-deficient Il1rl1−/− mice on C57BL/6J background have been described and they were bred at the Laboratory Animal Sciences Center (LASC) of the University of Zurich, Switzerland. C57BL/6J, BALB/c and DBA/2 wild-type mice were either purchased from Charles River and Janvier Labs or were bred at LASC and at the University of Geneva, Switzerland, under specific pathogen-free (SPF) conditions.

TC-1 tumor therapy studies in mice were performed at Hookipa Biotech GmbH using C57BL/6N mice purchased from Charles River, Sulzfeld, Germany. These experiments were approved by the Austrian authorities and were carried out in accordance with the approved guidelines for animal experiments at Hookipa Biotech GmbH.
All other mouse experiments were performed at the Universities of Basel and Geneva in accordance with the Swiss law for animal protection. Permission was granted by the Veterinäramt Basel-Stadt and by the Direction générale de la santé, Domaine de l’expérimentation animale, of the Canton of Geneva, respectively. Animals in experimental groups were sex- and age-matched. In general, adult animals of both genders were used to reduce the number of animals bred for research purposes. P1A-specific immunogenicity assessments and P815 tumor control studies were conducted in female mice. Mice in tumor therapy experiments were assigned to groups in a manner to assure even distribution of tumor volumes between groups at the time of tumor therapy. In accordance with the Swiss law for animal protection mice exhibiting wounds on the tumor or displaying signs of distress (evident namely in lethargy, hunchback, piloerection, emaciation and agonal breathing) were immediately euthanized irrespective of tumor size and diameter. Study sample sizes in animal experiments were chosen based on experience in our labs with respect to group sizes readily revealing biologically significant differences in the experimental models used. The groups were neither randomized nor were experiments conducted in a blinded fashion.

The PICV virulence study in guinea pigs was conducted at Meditox (Czech Republic) and was approved by the Institutional Animal Care and Use Committee (IACUC) and the Committee for Animal Protection of the Ministry of Health of the Czech Republic. Dunkin Hartley guinea pigs were from Charles River, France, and weighed 370 – 520 g at the start of the study.

Cell lines
BHK-21 cells, HEK293T cells were purchased from ECACC (Clone 13, Cat #85011433), P815 mastocytoma cells (TIB-64), NIH 3T3 and BSC40 cells from ATCC. FreeStyle 293-F suspension culture cells were purchased from Invitrogen/ThermoFisher. LCMV-GP-expressing BHK-21 cells (BHK-21-GP) and 293T-GP cells have previously been described. All cell lines were regularly tested for mycoplasma and were negative. Owing to their origin from renowned international repositories and vendors they were not authenticated.

METHOD DETAILS

Viruses, titration and neutralization test
The titration of LCMV, PICV, MOPV and derived vectors by immuno-focus assay has been described and was performed using NIH 3T3 cells as a substrate. CAND and derived vectors were titrated using HEK293T cells by analogous techniques. For detection of GFP-expressing artPICV and r3PICV infectivity by immuno-focus assay we used rat-anti-GFP antibody (Biolegend). To quantify PICV and derived vectors by immuno-focus assays, monoclonal antibody (mAb) 17.2.E4-95 served as primary antibody. mAb 17.1.C6-95 was used for detection of CAND and mAb 2B5 for MOPV. LCMV, PICV and derived vectors batches were produced on BHK-21 cells and 293F cells, MOPV and artMOPV-P1A on BHK-21, CAND and derived vectors on 293T cells.

Recombinant vaccinia viruses expressing the NP396 miniepitope or lacZ, respectively (VACC-NP396, VACC-lacZ) have been described. They were grown and titrated on BSC40 cells.

The neutralizing capacity of immune serum was determined by immuno-focus reduction assays.

Viral virulence testing
Intracranial LCMV infection was administered through the skull and mice developing signs of terminal disease were euthanized in accordance with the Swiss law.

The wellbeing of guinea pigs undergoing PICV or artPICV infection was monitored twice daily during the entire study and clinically scored. Moribund animals were sacrificed. Humane endpoints were hypothermia (body temperature < 35°C, determined at two independent monitoring time points) and/or body weight loss ≥ 20%.

Tumor implantation and tumor measurement
P815 cells (10^6 per mouse) were implanted subcutaneously in the right flank. Tumor growth was assessed three times per week. The longest and the shortest diameters were determined using a caliper. Tumor volumes (mm^3) were calculated as ½ (length*width^2). When tumor volumes exceeded 1500 mm^3 or when the longest median tumor diameter exceeded 20 mm, mice were euthanized in accordance with the Swiss law.

TC-1 cells expressing HPV 16 E6 and E7 were obtained from Johns Hopkins University. For tumor implantation, 10^5 cells were injected subcutaneously into the flank of C57BL/6 mice and, in accordance with the Austrian law, the experiment was terminated when tumor sizes exceeded 20 mm in any dimension.

Virus engineering, infection and immunization
The reverse genetic engineering of LCMV-WT, r3LCMV and artLCMV vectors using a polymerase I- / polymerase II-based plasmid system has been described. PICV-based, CAND-based and MOPV-based vectors and the corresponding cDNA-derived WT viruses were generated using analogous expression cassettes and transfection procedures using BHK-21-GP cells as a cell substrate. In brief, we transfected 5x10^6 cells, seeded the day before into an M6 cell culture well, with 0.8 μg of each pol-I-driven S segment expression plasmid, 1 μg of pol-I-driven L segment, 1.4 μg of pol-II-driven L ORF expression plasmid and 0.8 μg of pol-II-driven NP expression plasmid using 12 μl Lipofectamine 2000. Six hours after transfection for the rescue of CAND and derived vectors,
$10^6$ 293T-GP cells were added to each well. 72 hours after transfection, the cells were trypsinized and transferred to a T75 tissue culture flask. Virus- and vector-containing supernatants, respectively, were harvested 6-10 days after transfection. Wild-type CAND virus serving as a template for vector generation was generously provided by R. Charrel, Marseille, France. Its sequence of vectors was synthesized by Genscript, the Netherlands, and were ligated into polymerase-II- and polymerase-I-driven expression cassettes, respectively. MOPV cDNAs for virus and vector rescue (GenBank accession numbers JN561685.1 and JN561684.1) were generated by RT-PCR cloning (L and WT S segment, NP and L ORFs) and by gene synthesis (transgenic S segments of vectors). cDNAs of the full-length cancer-testis antigen P1A (comprising the immunodominant LPYGWLVF epitope), a non-oncogenic fusion protein consisting of the complete HPV16 E7 and E6 sequences (comprising the immunodominant epitope RAHYNIVTF), GFP or Tomato (TOM), were used for insertion into the respective vectors and viruses using a seamless cloning strategy previously described in detail.

Infections and immunizations of mice with arenaviruses and arenavirus-based vectors were performed at a dose of 10e5 PFU.i.v. unless specified otherwise. Vaccinia virus vectors were given at an intravenous dose of 2x10e6 PFU.

**Assessment of blood-brain-barrier integrity**

Blood-brain-barrier leakage was assessed by detecting IgG deposits in mouse brain parenchyma.

Cryosections of 10μm were fixed with 4% PFA for 15 min, endogenous peroxidases were inactivated and tissue sections were incubated with HRP-labeled-anti-mouse-IgG (Dako, K4001). Bound peroxidase polymers were visualized using polymerized 3,3'-diaminobenzidine (DAB, Dako, K5001). The stained sections were scanned using a Panoramic Digital Slide Scanner 250 Flash II at 200x magnification. Quantifications were performed on scanned slides applying a custom-programmed script to detect the DAB+ area in Cognition Network Language (Definiens Developer D software). For representative images, white balance was adjusted and contrast was linearly enhanced using the tools "levels," "curves," "brightness" and "contrast" in Photoshop CS6 (Adobe).

**Virus sequencing and genealogy tree building**

Viral RNA was extracted from cell culture supernatant and from serum of infected mice using the QIAamp Viral RNA Mini Kit (QIAGEN, Cat No. 52906). RNA from spleens of mice was extracted using TRI Reagent (Sigma Aldrich). Reverse-transcription PCR was performed with the One Step RT-PCR kit (QIAGEN) and gene-specific primers. Amplified products were gel-purified for Sanger sequencing.

A mammarenavirus genealogy tree was built based on S segment sequences of the following viruses and GenBank accession numbers: Allpahuayo virus (AY081210.1), Amapari virus (AF485256.1), Junin virus (AY358023), Bear Canyon virus (AY924391), Sabinia virus (U41071), Pichinde virus (K02734), Chapare virus (EU260463), Cupixi virus (AF512832), Flexal virus (AF512831), Gaibo virus (KJ855308), Guanarito virus (AY129247), Ippy virus (DQ328877), Lassa virus (AF181854.1), Latino virus (AF512830), Loei River virus (KC669698), Lujo virus (FJ952384), Luna virus (AB586864), Lunk virus (AB693150), Machupo virus (AY129248), Mariental virus (KM272987), Merino Walk virus (AY342390), Mopeia virus (AY772170), Okahandja virus (KM272988), Olliver virus (U34248), Parana virus (AF485261), Pirital virus (AF485262), Ryukyu virus (KM020191), Solwezi virus (AB972428), Souris virus (KP052277), Taborine virus (M20304), Tamiami virus (AF485263), Wenzhou virus (KJ909794), Whitewater Arroyo virus (AF228063). To build the phylogenetic tree, we used the software package BEAST2 with a TN93 site model (the corresponding .xml file will be deposited for further reference). The MCMC chain ran for 10^000’000 steps. All ESS were well above the critical threshold. The maximum credibility tree was constructed with TreeAnnotator and the phylogeny displayed with FigTree v1.4.4.

**Flow cytometry**

Antibodies against CD8 (53-6.7 or Ly-3.2), CD45R/B220 (RA3-6B2), Klrg1 (2F1), CD11a (A7R34), CX3CR1 (SA011F11), CD27 (LG3A10), CD43 (1B11), GrzB (GB12), Ki67 (solA15), IFN-γ (KM1G.2), TNF (MP6-XT22), IL-2 (JES6-5H4) CD11b (M1/70) CD11c (HL3) CD19(1D3) Nkp46 (29A1.4) and CD4 (RM-5) were from Biolegend, BD Biosciences/PharMingen and eBioscience/Thermo-Fisher. To assess intracellular levels of the transcription factor Tcf1, primary antibody binding (C63D8, Cell Signaling) was detected using donkey anti-rabbit IgG PE (Poly4064-eBioscience). Eomes (Dan11mag) and T-bet (4B10) were detected using the eBioscience™ FOXP3 transcription factor staining kit (Invitrogen).

Dead cells were excluded with Zombie UV Fixable Viability Kit (Biolegend, Cat. #423108). P1A epitope- (LPYLGLVLF), BBR epitope- (TYSKFESV) and NP396 epitope- (FOPQNGOFI) specific CTLs were identified by peptide-MHC class I tetramers after gating on CD8+2B20+ lymphocytes. The H-2Dβ tetramer loaded with the NP396 epitope and conjugated to PE was obtained through the NIH Tetramer Core Facility, the H-2Lα tetramer loaded with the P1A epitope and conjugated to PE as well as the H-2Kb tetramer loaded with the BBR epitope and conjugated to PE were purchased from the University of Lausanne Tetramer core facility. For tumor-infiltrating CTL analyses, cells expressing CD11b, CD11c, CD19 or Nkp46 were excluded. For detection of E7- (RAHYNIVTF epitope) and NP118- (RPQASGVYM epitope) specific CTLs the corresponding dextramers (Immundex) were used analogously (for simplicity referred to as "tetramer" in the text). Splenic single-cell suspensions were prepared by mechanical disruption and were counted...
using the respective single-use chambers in a Immunospot S6 device (C.T.L.). Total numbers of peptide-MHC tetramer-binding CTLs were back calculated. Cytokine profiles after restimulation with overlapping peptide sets spanning the E7 and E6 proteins of HPV16, respectively (JPT) were determined in intracellular cytokine assays as previously described. Samples were measured on BD LSRFortessa flow cytometers and were analyzed using FlowJo software (Becton Dickinson).

For TIL analysis, tumors were dissected and digested with accutase (Sigma Aldrich), Collagenase IV (Worthington), Hyaluronidase (Sigma Aldrich) and DNaseI (Sigma Aldrich) for 60 min at 37°C, followed by red blood cell lysis. Single cell suspensions were filtered using a cell strainer (70 µM).

QUANTIFICATION AND STATISTICAL ANALYSIS

Statistical testing
For statistical analysis, GraphPad Prism software (Version 9.0, GraphPad Software) was used unless stated otherwise. Differences between two groups were generally assessed using unpaired two-tailed Student’s t tests. To compare one group against multiple other groups we used one-way Analysis Of Variance (ANOVA), followed by Dunnett’s post hoc test (Figures 7G–7J). Two-way ANOVA with Sidak’s post-test was used to compare multiple cytokine-producing T cell subpopulations of two groups.

Tumor volume curves were compared as described. In brief, the area under the curve (AUC) for each individual animal was calculated using GraphPad Prism software and groups were compared pairwise by two-tailed Mann-Whitney tests. To calculate AUCs for all groups and animals throughout the 41d and 60d periods in Figures 7A and 7B, respectively, animals reaching humane endpoints of tumor volume were assigned either the maximally permitted tumor volume (1500 mm³, termination criterion for Figure 7A) or the maximally reached tumor (2815 mm³ for Figure 7B) for time points after sacrifice. For comparison of long-term survival rates chi-square tests were performed using the GraphPad QuickCalcs online tool.

P values of p < 0.05 were considered significant (*), p < 0.01 (***) as highly significant.