Topical Herpes Simplex Virus 2 (HSV-2) Vaccination with Human Papillomavirus Vectors Expressing gB/gD Ectodomains Induces Genital-Tissue-Resident Memory CD8+ T Cells and Reduces Genital Disease and Viral Shedding after HSV-2 Challenge

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ABSTRACT

No herpes simplex virus 2 (HSV-2) vaccine has been licensed for use in humans. HSV-2 glycoproteins B (gB) and D (gD) are targets of neutralizing antibodies and T cells, but clinical trials involving intramuscular (i.m.) injection of HSV-2 gB and gD in adjuvants have not been effective. Here we evaluated intravaginal (ivag) genetic immunization of C57BL/6 mice with a replication-defective human papillomavirus pseudovirus (HPV PsV) expressing HSV-2 gB (HPV-gB) or gD (HPV-gD) constructs to target different subcellular compartments. HPV PsV expressing a secreted ectodomain of gB (gBsec) or gD (gDsec), but not PsV expressing a cytoplasmic or membrane-bound form, induced circulating and intravaginal-tissue-resident memory CD8+ T cells that were able to secrete gamma interferon (IFN-γ) and tumor necrosis factor alpha (TNF-α) as well as moderate levels of serum HSV neutralizing antibodies. Combined immunization with HPV-gBsec and HPV-gDsec (HPV-gBsec/gDsec) vaccines conferred longer survival after vaginal challenge with HSV-2 than immunization with HPV-gBsec or HPV-gDsec alone. HPV-gBsec/gDsec ivag vaccination was associated with a reduced severity of genital lesions and lower levels of viral shedding in the genital tract after HSV-2 challenge. In contrast, intramuscular vaccination with a soluble truncated gD protein (gD2t) in alum and monophosphoryl lipid A (MPL) elicited high neutralizing antibody titers and improved survival but did not reduce genital lesions and viral shedding. Vaccination combining ivag HPV-gBsec/gDsec and i.m. gD2t-alum-MPL improved survival and reduced genital lesions and viral shedding. Finally, high levels of circulating HSV-2-specific CD8+ T cells, but not serum antibodies, correlated with reduced viral shedding. Taken together, our data underscore the potential of HPV PsV as a platform for a topical mucosal vaccine to control local manifestations of primary HSV-2 infection.

IMPORTANCE

Genital herpes is a highly prevalent chronic disease caused by HSV infection. To date, there is no licensed vaccine against HSV infection. This study describes intravaginal vaccination with a nonreplicating HPV-based vector expressing HSV glycoprotein antigens. The data presented in this study underscore the potential of HPV-based vectors as a platform for the induction of genital-tissue-resident memory T cell responses and the control of local manifestations of primary HSV infection.

Genital herpes is a common sexually transmitted disease caused by herpes simplex virus 2 (HSV-2). Worldwide, more than 500 million individuals are chronically infected by HSV-2, and the prevalence of HSV-2 infection is twice as high in women as in men (1). In the United States, the seroprevalence of HSV-2 in 14- to 49-year-olds during the 2005–2010 period was 15.7% (2). During primary infection, HSV-2 infects and replicates in epithelial cells of the genital mucosa and spreads to the regional ganglia, where it establishes a lifelong latent infection. HSV-2 can undergo reactivation and shedding from the genital mucosa, where it can cause recurrent genital lesions, which are associated with an increased risk of HIV-1 acquisition (3, 4). Shedding of HSV-2 may also be subclinical, and HSV-2 transmission can occur in the absence of lesions (5, 6). Immunosuppression is associated with an increased risk of severe disseminated disease. In addition, transmission of HSV-2 from the genital mucosa of acutely infected pregnant women to neonates can cause severe infection.

Several therapeutic and preventive interventions based on antiviral drugs, the use of condoms, abstinence, or circumcision can reduce the burden of HSV-2 infection at the individual level. However, these interventions have not controlled the HSV-2 epidemic (7). Therefore, a vaccine that could prevent primary acquisition of HSV-2 or reduce HSV-2 shedding and/or recurrent lesions in chronically infected individuals might have a substantial impact at both the individual and public health levels.
A variety of HSV-2 vaccine approaches have shown protective efficacy in animal models, including live attenuated, nonreplicating viral vector, subunit, or DNA vaccines (8–20). Recombinant soluble HSV-2 glycoprotein D (gD) combined with an aluminum salt and monophosphoryl lipid A adjuvant (alum–MPL) has been the most promising recent vaccine to undergo extensive clinical evaluation. Although it induced HSV-2 neutralizing antibodies in the sera of vaccinated subjects, this vaccine failed to confer significant protection in a phase III clinical trial (21, 22). It is therefore speculated that a successful HSV-2 vaccine should also induce a robust T cell response (23).

Infection of mice with HSV-2 has provided evidence that CD4+ or CD8+ T cells and gamma interferon (IFN-γ) can contribute to reducing the severity of primary infection, clearing virus from the nervous system, and protecting against reactivation ex vivo (24–28). More recently, it has been shown that, in contrast to circulating memory T cells, a subset of tissue-resident memory (Trm) T cells can confer immediate and enhanced protection against HSV-1 and HSV-2 infections (29–31). In humans, a subset of CD8αααα T cells is induced in the genital epithelium at sites of clinical HSV-2 reactivation, and these cells persist after the lesions have healed (32, 33). The presence of these local T cells is associated with reductions in lesion severity and viral shedding (34). In mouse models, genital Trm T cells can be induced by genital immunization with live attenuated HSV-2 or by systemic immunization followed by topical application to the genital tract of immunomodulatory molecules, which can directly recently activated circulating T cells to the genital tract (29–31, 35, 36).

We previously reported an effective method for transiently transducing the cervicovaginal mucosa with a nonreplicating human papillomavirus (HPV) vector (37). Intravaginal (ivag) immunization with these HPV pseudovirus vectors (HPV PsV) expressing a model antigen induces higher numbers of antigen-specific Trm CD8+ T cells in the cervicovaginal mucosa in mice and nonhuman primates and confers greater control of local viral replication after ivag vaccinia virus challenge than intramuscular (i.m.) immunization at a remote site (38–40).

Here we generated and characterized HPV16 and HPV45 PsV expressing membrane-associated, secreted, and cytosolic forms of HSV-2 glycoprotein B (gB) and gD. We then evaluated the ivag immunogenicities of the HPV PsV constructs, including their abilities to induce cervicovaginal intraepithelial Trm CD8+ T cells and antibodies, and compared the protection conferred by this local vaccination, in a genital tract HSV-2 challenge model, with that conferred by i.m. immunization with a truncated gD protein formulated with alum and MPL.

**MATERIALS AND METHODS**

**Design, construction, and characterization of plasmids expressing HSV-2 gB, HSV-2 gD, or reporter genes.** Plasmids encoding tomato fluorescent protein and firefly luciferase as reporter genes or as controls were generated previously and are described on the Laboratory of Cellular Oncology website (http://home.ccr.cancer.gov/Lco/plasmids.asp).

Plasmids pgB2-333 and pgD2-333 consisted of vector pcDNA3 and the gB or gD DNA fragments that were PCR amplified from HSV-2 strain 333. The entire open reading frames (ORF) were confirmed by sequencing. These HSV-2 DNA fragments were further subcloned into an HPV PsV expression plasmid. Briefly, the full-length open reading frames of gB and gD (gBfl and gDfl) were excised from the pcDNA3 constructs using restriction enzyme HindIII, the end was blunted, and the DNA was further digested using XhoI, the ends were blunt, and the DNA was further digested using NotI. Purified fragments encoding gB and gD were cloned into plasmid pCI to generate plasmids pCgBfl and pCgDfl. To produce the secreted ectodomains of gB and gD (gBsec and gDsec), we generated constructs with deletions of the glycoprotein transmembrane and cytosolic domains from plasmids pCgBfl and pCgDfl by PCR using the primer pair 5’-CATAGATTTCAAAAACTGCGGCGGAGGCGTGC-3’ and 5’-ATACCTTCGCGCGGTTAGGGTCGCGCGGCGG-3’ for gBsec and the primer pair 5’-ACGCTTACAGTAAAGCGGGCCGCGGCTTTGACC-3’ and 5’-ATACCTTCGCGCGGCTAGTGGTGGGCGGCGGCGG-3’ for gDsec. To produce the cytosolic forms of gB and gD (gBcyt and gDcyt), the transmembrane and cytosolic domains were deleted from the glycoprotein constructs, and the signal peptide sequence was replaced by a Kozak sequence and an ATG codon in frame with the rest of the ORF. We generated the cytosolic forms of the glycoproteins by PCR from pcgBfl and pcgDfl constructs using primers 5’-CATAGATTTCAAAAACTGCGGCGGAGGCGTGC-3’ and 5’-ACGCTTACAGTAAAGCGGGCCGCGGCTTTGACC-3’ and 5’-ATACCTTCGCGCGGCTAGTGGTGGGCGGCGGCGG-3’ for gBcyt and primers 5’-ACGCTTACAGTAAAGCGGGCCGCGGCTTTGACC-3’ and 5’-ATACCTTCGCGCGGCTAGTGGTGGGCGGCGGCGG-3’ for gDcyt. All PCR products were cloned into a pCLuc vector backbone after digestion with restriction enzymes XbaI and NotI.

The expression of gB and gD by each plasmid was assessed by Western blotting. Briefly, 293TT cells were transfected with each pcgBfl and pcgDfl construct, or with the pCLuc plasmid as a control, using Lipofectamine 2000 (Invitrogen). Cell lysates were prepared using radioimmunoprecipitation assay (RIPA) buffer 48 h after transfection, and supernatants were obtained for the last 14 h of incubation in serum-free medium (Opti-MEM; Invitrogen) to minimize background. The expression of gB was confirmed by Western blotting under denaturing conditions with rabbit polyclonal antibody R69 (41), a gift from Gary Cohen and Roselyn Eisenberg, and gD expression was confirmed with mouse monoclonal antibody 2C10 (Abcam). The apparent molecular weights were confirmed with a molecular weight marker (MagicMark XP).

**HPV vectors and HSV-2 production.** HPV vectors were produced and purified as described previously (42). Briefly, 293TT cells were cotransfected with a plasmid encoding both the L1 and L2 capsid proteins of HPV16 or HPV45 and either a plasmid encoding a reporter gene (tomato fluorescent protein or firefly luciferase) or a plasmid encoding one of the glycoprotein constructs: gBfl, gBsec, gBcyt, gDfl, gDsec, or gDcyt. After overnight incubation of cell lysates, HPV particles were purified on an OptiPrep gradient. The infectious titer of each pseudovirus preparation was determined on 293TT cells by flow cytometry and was expressed as infectious units (IU) per ml.

HSV-2 (strain R519, plaque purified from strain 333) was propagated in Vero cells as described previously (43), and the infectious titer was determined by a plaque assay on Vero cells and was expressed as PFU per ml.

**Animals.** Six- to 8-week-old female C57BL/6 mice were purchased from the National Cancer Institute (NCI) and were maintained under specific-pathogen-free conditions in the animal care facilities at the NCI and the National Institute of Allergy and Infectious Diseases (NIAID). All procedures were approved by the NCI and NIAID Animal Care and Use Committees.

**Immunizations and HSV-2 challenge.** For primary or single vaccination, HPV16 PsV were used; for booster vaccination, HPV45 PsV were used. HPV PsV ivag immunization was performed on anesthetized mice as described previously (40). Five days prior to immunization, C57BL/6 female mice were treated subcutaneously (s.c.) with medroxyprogesterone acetate (Depo-Provera; 3 mg; Pfizer). On the day of immunization, mice received 4% nonoxynol-9 (N9; Spectrum Chemical and Laboratory) ivag, and 5 h later, they were infected ivag with 1 × 10⁷ or 2 × 10⁷ IU (as indicated for each experiment) of the HPV vectors in carboxymethylcellulose (CMC; Sigma-Aldrich).

For intranasal (i.n.) immunization, anesthetized mice received 1 µg
cholera toxin (CT; List Biological Laboratories) with 20 μg of gB\textsubscript{am-503} (NeoSystems) peptide in 10 μl phosphate-buffered saline (PBS) (5 μl per nostril).

For i.m. immunization, 3 μg of a soluble truncated recombinant gD protein (gD2C; Chiron) was adsorbed onto 50 μg Imject Alum (Thermo Scientific), mixed with 7.5 μg synthetic monophosphoryl lipid A (MPL; Invivogen), diluted in 50 μl with PBS, and injected in the quadriceps muscles of anesthetized mice.

For HSV-2 vaginal challenge, medroxyprogesterone acetate-treated mice were instilled ivag with 10\textsuperscript{5} PFU of HSV-2 strain 333 in 10 μl PBS.

**Immunofluorescence and confocal microscopy.** For analysis of the expression of tomato fluorescent protein in the cervicovaginal mucosa, tissues were collected 72 h after immunization and were fixed for 1 h in 4% paraformaldehyde (EMS), followed by an incubation of 24 h in 1% sucrose solution and then 24 h in 30% sucrose. Tissues were snap-frozen in tissue freezing medium (Tissue-Tek OCT; Sakura). Six-micrometer tissue sections were cut, transferred to glass slides (Superfrost Plus; VWR), and mounted with an antifade reagent (Prolong Gold with 4\textsuperscript{′},6-diamidino-2-phenylindole [DAPI]; Molecular Probes).

For analysis of CD8 and CD4 cell infiltrates, cervicovaginal tissues were collected 3 weeks after the final immunization, snap-frozen directly in Tissue-Tek compound, and processed as described previously (38). Briefly, 6-μm ethanol-fixed tissue sections were incubated with an antibody against CD16 and CD32 to block FcR (clone 24G2; BioXcell) and with normal donkey serum and were stained using Alexa Fluor 488-conjugated anti-CD8 and Alexa Fluor 594-conjugated anti-CD4 antibodies (BioLegend). Tissue sections were mounted with Prolong Gold with DAPI.

**Preparation of cell suspensions for flow cytometry analysis.** Cervicovaginal cell suspensions were obtained after mincing of the cervicovaginal tissue into small pieces using dissecting scissors. The minced tissue was incubated for 1 h at 37°C in a shaker at 250 rpm in RPMI 1640 medium with 2% fetal bovine serum (FBS; Sigma-Aldrich), 0.5 mg/ml collagenase A (Roche), and 0.1 mg/ml DNase I (Roche). Spleen cell suspensions were obtained by following the protocol described above with a 20-min collagenase A–DNase I incubation. All cell suspensions were filtered through a 70-μm cell strainer. To remove red blood cells, cell suspensions and EDTA-treated blood were incubated for 5 min at room temperature (RT) in an ammonium chloride solution, washed, and kept on ice before further analysis.

**Fluorescence-activated cell sorter (FACS) analysis of CD8^+ T cell responses against the HSV-2 gB immunodominant epitope gB\textsubscript{am-503}.** After FcR blocking, cell suspensions were stained for 30 min at 4°C with an allophycocyanin (APC)-conjugated H-2K\textsuperscript{b}gB\textsubscript{am-503} tetramer (NIH Tetrramer Facility) and the following antibodies: Pacific Blue-conjugated anti-CD3 (clone 17A2), APC- and Cy7-conjugated anti-CD4 (clone RM4-5), phycoerythrin (PE)- and Cy7-conjugated anti-CD8 (clone IM7), PE-conjugated anti-CD49a (clone HM41), PE- and Cy7-conjugated anti-CD69 (clone H1.2F3), fluorescein isothiocyanate (FITC)-conjugated anti-CD62L (clone ME-L14), peridinin chlorophyll protein (PerCP)- and Cy5.5-conjugated anti-CD103 (clone 2E7), PE-conjugated anti-CD127 (clone SB/199), PE- and Cy7-conjugated anti-CXCR3 (clone CXCR3-173) (BioLegend), FITC-conjugated anti-KLRG-1 (clone 2F1; SouthernBiotech), and Pacific Orange-conjugated anti-CD8 (clone 53.6.7; Invitrogen).

To measure intracellular cytokine content, cells were incubated for 5 h at 37°C under 5% CO\textsubscript{2} in RPMI 1640 medium containing 10% FBS, sodium pyruvate, L-glutamine, B-mercaptoethanol, and GolgiPlug (final concentration, 1 μg/ml brefeldin A; BD Biosciences), with 1 μg/ml gB\textsubscript{am-503} peptide. After incubation, cells were washed, FcR blocked, and stained with antibodies against CD8, CD3, and CD4 as described above; dead cells were labeled with Live/Dead yellow dye (Invitrogen); the cells were fixed and permeabilized with Cytofix/Cytoperm (BD Biosciences); and intracellular staining was performed for 30 min at 4°C with FITC-conjugated anti-IFN-γ (clone XMG1.2), PerCP- and Cy5.5-conjugated anti-interleukin 2 (anti-IL-2) (clone JES65H4), and APC-conjugated anti-tumor necrosis factor alpha (anti-TNF-α) (clone MP6-XT22) antibodies (BioLegend).

**In vitro neutralization assays.** The *in vitro* neutralization of HPV pseudovirions has been described previously (44). Serial dilutions of serum samples from individual mice were incubated with pseudovirus expressing secreted alkaline phosphatase (SEAP) at 4°C for 1 h. The pseudovirus–serum mixtures were added to 293TT cells, and 3 days later, SEAP activity was measured in the supernatants. The 50% effective concentration (EC\textsubscript{50}) was defined using Prism software as the dilution of serum corresponding to a 50% reduction in SEAP activity.

To determine the HSV-2 neutralizing titer in serum, 2-fold serial dilutions of serum samples were incubated with HSV-2 strain 333 for 1 h at RT and were added to Vero cell monolayers for 1 h at 37°C. The virus–antibody inocula were replaced with a plaque assay overlay consisting of Vero cell growth medium containing 0.3% (vol/vol) G418 (Ganumex-C; Talecris Biotherapeutics), and cells were incubated for 48 h at 37°C. Cells were then stained with a plaque staining solution containing 10% (vol/vol) acetic acid, 60% (vol/vol) methanol, and 1% (wt/vol) crystal violet and were washed, and plaques were counted.

**ELISA for measurement of gD antibodies.** Titers of IgG against HSV-2 gD in serum were determined by enzyme-linked immunosorbent assays (ELISA). Briefly, a high-binding-capacity microplate (2HB; Immunon) was coated with purified recombinant gD21 protein (50 ng/well; Chiron). Plates were blocked with 1% (wt/vol) PBS, and serial dilutions of serum samples were incubated in each well. Immunoglobulin binding to the plate was measured using a horseradish peroxidase (HRP)-conjugated affinity-purified goat antibody to mouse IgG (Southern Biotechnology Associates). Specific titers were defined as the reciprocals of the highest sample dilutions giving signals equal to or at least 3-fold the background signal.

**Titration of virus shedding.** Vaginal swabs were collected daily from day 1 to day 7 after HSV-2 genital challenge, placed in culture medium containing penicillin, streptomycin, and amphotericin B, and stored at ~80°C until titration by plaque assay on Vero cells.

**Statistics.** Statistical significance (P ≤ 0.05) was determined using the nonparametric Mann-Whitney U test in experiments with two independent groups. One-way analysis of variance (ANOVA) and multiple comparisons were performed for experiments with more than two groups. Spearman’s coefficient (r) was calculated to determine the correlation between variables.

**RESULTS**

**Generation and characterization of HPV PsV that target expression of HSV–2 gB and gD to different subcellular compartments.**

We hypothesized that targeting the accumulation of HSV–2 gB and gD to different subcellular compartments after *in vivo* transduction of cervicovaginal epithelial cells of mice with HPV PsV (shown in Fig. 1A as a 3-dimensional reconstruction (45)) might affect their immunogenicity. *In vivo* transduction of cervicovaginal epithelial cells by HPV16 and HPV45 PsV was confirmed by confocal microscopy analysis of genital tract tissue sections after intravaginal instillation of PsV expressing tomato fluorescent protein (Fig. 1B), as reported previously by Roberts et al. (37).

To convert wild-type membrane-associated gB and gD (gBfl and gDfl) into secreted (gBsec and gDsec) or cytosolic (gBcty and gDcty) forms, we deleted the transmembrane and cytoplasmic sequences or the transmembrane, cytoplasmic, and signal peptide
sequences of gB and gD, respectively. The remaining DNA sequences were introduced into a modified expression plasmid (pCI), as depicted in Fig. 1C.

The expression of gB and gD was assessed by Western blotting of cell lysates and supernatants of 293TT cells transfected with the plasmids expressing full-length, secreted, and cytosolic forms of HSV-2 gB and gD. Full-length, secreted, and cytosolic gB and gD migrated with apparent molecular sizes of 120, 100, and 80 kDa (gB) or 50, 45, and 35 kDa (gD), respectively (Fig. 1D and E). The apparent molecular masses of the two cytosolic constructs of gB and gD in the cell lysates corresponded to their predicted molecular masses without posttranslational modification of 79 kDa and 33 kDa, respectively, strongly suggesting restricted expression of the gBcyt and gDcyt constructs in the cytosolic compartment (Fig. 1D and E). While gB and gD were detected in the supernatants of cells transfected with the secreted constructs, they were not seen in the supernatants of cells transfected with the cytosolic constructs or the full-length construct for gB (Fig. 1D and E). A faint band of the same apparent molecular mass as the gD secreted form was detected in the supernatant of cells transfected with the full-length construct (Fig. 1E), in agreement with a previous report that a fraction of full-length gD is subject to proteolytic cleavage and release of the ectodomain into the supernatant of HSV-2-infected cells (46).

CD8+ T cell responses after intravaginal immunization with HPV PsV that target the expression of HSV-2 gB to different cellular compartments. We used C57BL/6 mice for immunogenicity studies because this haplotype permits the measurement of CD8+ T cell responses against the immunodominant epitope gB496–503 restricted to H-2Kb (47, 48). C57BL/6 mice were immunized ivag with HPV-gBsec elicited detectable Kb/gB496 tetramer-positive CD8+ T cells in cervical mucosa cell suspensions (Fig. 2A and B). While the percentage of Kb/gB496 tetramer-positive CD8+ T cells in the cervical mucosa cell suspensions was slightly higher after a booster immunization than after HPV-gBsec priming immunization alone (means, 63.6% versus 41.3%, respectively), the total number of Kb/gB496 tetramer-positive CD8+ T cells in the cervi-
covaginal mucosa was 4 times higher after the booster immunization. Importantly, prime-boost ivag immunization with HPV-gBsec resulted in 10 times more Kb/gB496 tetramer-positive CD8+ T cells in the cervicovaginal mucosa than prime-boost immunization with i.n. CT plus gB496–503 peptide. In contrast, prime-boost i.n. immunization with CT plus gB496–503 peptide induced higher frequencies of Kb/gB496 tetramer-positive CD8+ T cells in blood (mean, 6.8%) and spleen (mean, 2.7%) than prime-boost ivag immunization with HPV-gBsec (mean frequencies, 4.0% in blood and 1.5% in spleen) (Fig. 2C and D). As with the CD8+ T cell
responses observed in the cervicovaginal mucosa, HPV-gBsec ivag prime-boost immunization induced higher frequencies of K\(^6\)/gB\(_{496}\) tetramer-positive CD8\(^+\) T cells in blood and in spleen than HPV-gBsec ivag priming immunization only. Interestingly, a strong correlation was observed between blood and genital CD8\(^+\) T cell responses, whether the mice received only the priming dose or the prime-boost regimen (\(r = 0.93; P = 0.0007\)) (Fig. 2F). In the HPV-gBsec group, the total number of CD8\(^+\) T cells was increased by 20-fold after ivag prime-boost immunization, whereas we observed only a 2-fold increase at most in CD4\(^+\) T cells (data not shown). In agreement with these results, immunofluorescence staining with antibodies against CD8 and CD4 confirmed the intraepithelial distribution of CD8\(^+\) cells and of some CD4\(^+\) T cells in the cervicovaginal mucosa after ivag immunization with the HPV-gBsec vector, in contrast to observations for animals treated with the HPV control vector (Fig. 2F).

Together, these data indicate that the secreted ectodomain of gB expressed by cervicovaginal epithelial cells after HPV PsV immunization is much more immunogenic for T cells than the cytoplasmic or membrane-bound form of the protein and, in particular, that it can induce robust CD8\(^+\) T cell responses in the cervicovaginal epithelium. Therefore, all subsequent immunization experiments were carried out with the secreted ectodomain constructs, gBsec and gDsec.

**Surface marker expression by HSV-2-specific CD8\(^+\) T cells after ivag immunization with HPV-gBsec.** We performed phenotypic analyses by flow cytometry of HSV-2 gB-specific CD8\(^+\) T cells in order to define which type of memory cells was induced by ivag HPV-gBsec immunization. The expression of various surface molecules involved in T cell migration, activation, and function was measured on K\(^6\)/gB\(_{496}\) tetramer-positive CD8\(^+\) T cells in blood and cervicovaginal cell suspensions 1 month after the last immunization. In all of the tissues analyzed, HSV-specific CD8\(^+\) T cells induced by ivag HPV-gBsec immunization displayed an effector or effector/memory phenotype characterized by low expression of CD62L, but heterogeneous expression of CD127 (Fig. 3). There is considerable evidence that tissue-resident memory CD8\(^+\) T cells and other intraepithelial lymphocytes selectively express the integrin CD103 and that its expression is associated with improved protection in several infectious models, including HSV-1 models (50). CD103 was selectively upregulated by HSV-specific CD8\(^+\) T cells in cervicovaginal cell suspensions but not in blood. CXCR3, which is the receptor for chemokines CXCL9 and CXCL10, is involved in the migration of memory CD8\(^+\) T cells into inflamed tissue and was recently associated with the differentiation of tissue-resident memory CD8\(^+\) T cells (51). Expression of CXCR3 was detected on the majority (70%) of cervicovaginal HSV-specific CD8\(^+\) T cells and on half of HSV-specific CD8\(^+\) T cells in blood induced by ivag HPV-gBsec vaccination. Although expression of CD69 is usually associated with recent activation of antigen-specific CD8\(^+\) T cells, all genital HSV-specific CD8\(^+\) T cells still expressed CD69 1 month after the final ivag HPV-gBsec immunization. However, CD69 was not expressed by HSV-specific CD8\(^+\) T cells in blood (Fig. 3). Interestingly, KLRG-1 was expressed by most HSV-specific CD8\(^+\) T cells in blood but was not expressed by HSV-specific cervicovaginal CD8\(^+\) T cells. As expected, CD44 was expressed by both cervicovaginal and blood HSV-specific CD8\(^+\) T cells. Finally, CD49a (\(\alpha1\) integrin) was expressed by HSV-specific cervicovaginal CD8\(^+\) T cells and, to a lower extent, by HSV-specific CD8\(^+\) T cells in blood (Fig. 3).

**Intravaginal immunization with HPV-gBsec elicits the production of HSV-specific IFN-\(\gamma\)- and TNF-\(\alpha\)-secreting systemic and cervicovaginal CD8\(^+\) T cells.** The production of multiple cytokines in memory CD8\(^+\) T cells is often associated with better function and quality of the response (52). In particular, IFN-\(\gamma\) and TNF-\(\alpha\) production by local T cells has been shown to contribute to protection in the context of HSV-2 primary infection or in experimental vaccination settings (26–28). Spleen and genital tract cell suspensions were obtained 4 weeks after the last ivag immunization with HPV-gBsec, HPV-gBCyt, HPV-gBIII, or the HPV control or 4 weeks after the last i.n. immunization with CT plus gB\(_{496-503}\). The cell suspensions were incubated for 5 h with 1 \(\mu\)M gB\(_{496-503}\) peptide, and cytokine production in CD8\(^+\) T cells was measured by intracellular staining with antibodies against IFN-\(\gamma\), TNF-\(\alpha\), and IL-2, followed by flow cytometry analysis. The number of IFN-\(\gamma\)-producing CD8\(^+\) T cells was higher in the cervicovaginal mucosa after ivag prime-boost immunization with HPV-gBsec than after i.n. immunization with CT plus the gB\(_{496-503}\) peptide (Fig. 4A). In contrast, the frequency of IFN-\(\gamma\)-producing CD8\(^+\) T cells in the spleen was higher after i.n. immunization with CT plus the gB\(_{496-503}\) peptide than after ivag immunization with HPV-gBsec (Fig. 4B). In agreement with the data shown in Fig. 2, neither HPV-gBCyt nor HPV-gBIII ivag vaccination induced IFN-\(\gamma\)-producing cells in vivo stimulation with the gB\(_{496-503}\) peptide.

The frequency of CD8\(^+\) T cells producing at least one cytokine in response to *in vitro* stimulation with gB\(_{496-503}\) (responding CD8\(^+\) T cells) was higher in cervicovaginal suspensions (21%) than in the spleen (0.7%) (Fig. 4C and D). Cervicovaginal or spleen CD8\(^+\) T cells from HPV control-immunized mice did not produce any cytokine in response to *in vitro* stimulation with gB\(_{496-503}\) (Fig. 4C and D). In the cervicovaginal mucosa of mice immunized with HPV-gBsec, 47% of the responding CD8\(^+\) T cells produced IFN-\(\gamma\) only, 11% produced TNF-\(\alpha\) only, and 37% produced both IFN-\(\gamma\) and TNF-\(\alpha\), and 5% produced IFN-\(\gamma\), TNF-\(\alpha\), and IL-2 simultaneously (Fig. 4E). Responding CD8\(^+\) T cells in the spleen produced a slightly narrower range of cytokines than those in the cervicovaginal mucosa; the majority (72%) of the responding splenic CD8\(^+\) T cells produced only one cytokine, IFN-\(\gamma\) or TNF-\(\alpha\). These data indicate that gB-specific cervicovaginal CD8\(^+\) T cells and, to a lesser extent, gB-specific CD8\(^+\) T cells in the spleen induced by ivag HPV-gBsec immunization can produce multiple cytokines for antiviral defense.

**Combined ivag immunization with HPV-gBsec/gDsec vaccines confers enhanced survival after HSV-2 vaginal challenge over that obtained by immunization with HPV-gBsec or HPV-gDsec alone.** We evaluated whether ivag immunization with HPV vectors expressing gBsec or gDsec alone or combined could confer protection in the mouse HSV-2 lethal genital challenge model. Medroxyprogesterone acetate-treated mice (n = 10 per group) were immunized ivag with 10\(^6\) IU of HPV16-gBsec or HPV16-gDsec alone or with a mixture of HPV16-gBsec and HPV16-gDsec (10\(^6\) IU each), followed 1 month later by 10\(^8\) IU of HPV45 expressing the respective antigens. As expected, none of the sham-treated (2% CMC) mice survived the genital challenge with 10\(^6\) PFU HSV-2 strain 333, and sham-treated mice started to die by day 9 after HSV-2 genital challenge. Only 20% or 30% of the mice immunized ivag with HPV-gBsec or HPV-gDsec, respectively, survived the HSV-2 genital challenge (P = 0.284 and 0.048 for comparison to sham-
treated mice). Interestingly, combining both HSV-2 glycoproteins in the HPV PsV conferred higher protection (50% survival) than either sham treatment ($P = 0.003$) or immunization with HPV-gBsec ($P = 0.069$) or HPV-gDsec ($P = 0.093$) alone (Fig. 5). These results suggest that HPV vectors can confer partial protection against HSV-2 lethal genital challenge and that delivering more than one antigen can enhance this protection.

**Immunogenicity of simultaneous ivag immunization with HPV-gBsec/gDsec vectors and i.m. immunization with gD2t protein in alum and MPL.** A soluble truncated gD protein (gD2t) in alum-MPL was the basis of an HSV-2 vaccine tested recently in a phase III clinical trial. This vaccine induced HSV-2 neutralizing antibodies in serum, though with low titers, and provided no protection against genital HSV-2 ($21, 22$). In the guinea pig HSV-2 genital model, immunization with gD2t in alum and MPL induced high titers of neutralizing antibodies in serum and conferred full survival upon challenge but did not fully control viral replication during acute infection ($53$). We speculated that combining a vaccine that induces high HSV-2 neutralizing antibody titers with our HPV PsV to induce high numbers of intraepithelial HSV-specific memory CD8$^+$ T cells might confer enhanced protection against disease and improve control of local replication of the virus. Medroxyprogesterone acetate-treated mice ($n$, 10/group) were immunized ivag with a mixture of HPV16-gBsec and HPV16-gDsec ($2 \times 10^8$ IU each) and were boosted 1 month later with $2 \times 10^8$ IU each of HPV45-gBsec and HPV45-gDsec (HPV-gBsec/gDsec). Another group of mice was immunized i.m. twice, 1 month apart, with 3 $\mu$g gD2t and 7.5 $\mu$g MPL adsorbed on 50 $\mu$g of alum (gD2t-alum-MPL). A third group of mice was prime-
Cytokine production by CD8⁺ T cells induced after ivag immunization with HPV-gBsec, gBcyt, or gBfl. Medroxyprogesterone acetate-treated mice (4 per group) were immunized ivag with $1 \times 10^8$ IU of HPV16 expressing gBsec, gBcyt, or gBfl, followed 1 month later by an ivag booster immunization with $10^8$ IU of HPV45 expressing the same glycoprotein. Another group of mice was immunized i.n. twice, 1 month apart, with $20\mu g$ of the minimal immunodominant peptide gB496–503 admixed with $1\mu g$ CT. As a control, a group of mice was immunized with $10^8$ IU of HPV16 expressing luciferase, followed 1 month later by a second immunization with $10^8$ IU of HPV45 expressing luciferase. (A and B) Levels of IFN-γ-producing cells in the vagina (A) and spleen (B) measured by intracellular cytokine staining after in vitro stimulation with the gB496–503 peptide 4 weeks after the final immunization. (C and D) Representative plots of IFN-γ and TNF-α production, and of IFN-γ and IL-2 production, in cells from the vagina (C) and spleen (D). (E) Pie diagrams of production of multiple cytokines by cervicovaginal and spleen CD8⁺ T cells in mice immunized with HPV-gBsec.
boost immunized ivag with HPV-gBsec/gDsec and i.m. with gD2t-alum-MPL simultaneously. For control purposes, a group of mice was prime-boost immunized ivag with HPV16 and HPV45 PsV expressing red fluorescent protein (RFP). As expected, i.m. immunization with gD2t-alum-MPL induced high HSV-2 gD-specific antibody titers and high neutralizing antibody titers, whereas ivag immunization with HPV-gBsec/gDsec induced modest levels of HSV-2 gD and HSV-2 neutralizing antibodies in the serum (Fig. 6A and B). Interestingly, the HPV-gBsec/gDsec vaccine, which delivers a small amount of HPV capsid protein (2 μg L1) to the vaginal tract in the absence of adjuvant, induced high titers of HPV type-specific neutralizing activity in the serum (Table 1).

The two groups receiving HPV-gBsec/gDsec had similar frequencies of circulating HSV-specific CD8+ T cells after a single immunization (Fig. 6C). However, after the second immunization, the group receiving HPV-gBsec/gDsec ivag alone displayed a pronounced increase in circulating gB-specific CD8+ T cells that was not observed in the group receiving HPV-gBsec/gDsec ivag combined with i.m. gD2t-alum-MPL. The latter data suggest that i.m. immunization with gD2t-alum-MPL might have interfered with the boosting of gB-specific CD8+ T cells by the HPV vectors given ivag (Fig. 6C).

Intravaginal immunization with the HPV-gBsec/gDsec vaccine, but not i.m. immunization with gD2t-alum-MPL, reduces the severity of genital lesions and viral shedding after HSV-2 genital challenge. As expected, none of the mice immunized with the control HPV vector survived the lethal HSV-2 genital challenge with 10^4 PFU of HSV-2 strain 333, whereas 80% of the mice immunized i.m. with gD2t-alum-MPL survived (Fig. 7A). Sixty percent of the mice immunized ivag with HPV-gBsec/gDsec survived, while 90% of mice immunized with a combination of ivag HPV-gBsec/gDsec and i.m. gD2t-alum-MPL survived. Interestingly, there was no difference in HSV-2 titers in genital swabs between HPV control-treated mice and gD2t-alum-MPL i.m.-immunized mice (Fig. 7B), despite high levels of neutralizing activity in the sera of the latter group (Fig. 6B). Mice immunized ivag with

![FIG 5](https://example.com/figure5)

**FIG 5** Combined immunization with HPV vectors expressing gB and gD induces enhanced protection against HSV-2 vaginal challenge. Medroxyprogesterone acetate-treated mice (n, 10/group) were immunized ivag with HPV16 expressing gBsec, gDsec, or a mixture of both (each at 10^9 IU), followed 1 month later by ivag booster immunization with HPV45 expressing the respective antigens at 10^9 IU. As a control, a group of mice was sham treated (2% CMC) twice, 1 month apart. The graph shows the survival of mice challenged ivag 1 month after the last immunization with 10^4 PFU of HSV-2 strain 333. Statistical significance was measured by a log rank (Mantel-Cox) test (*, P ≤ 0.05; **, P ≤ 0.01; ns, nonsignificant).

![FIG 6](https://example.com/figure6)

**FIG 6** Immunogenicity of simultaneous ivag immunization with HPV-gBsec/gDsec and i.m. immunization with gD2t with alum and MPL. Medroxyprogesterone acetate-treated mice (n, 10/group) were immunized twice, 1 month apart, with 3 μg gD2t with 50 μg alum and 7.5 μg MPL (gD2t/A/MPL). A second group of mice was immunized ivag with a mixture of 2 × 10^8 IU of HPV16-gBsec and 2 × 10^8 IU of HPV16-gDsec, followed 1 month later by ivag booster immunization with HPV45 expressing the respective antigens (2 × 10^8 IU each) (HPV-gBsec/gDsec). A third group of mice was immunized simultaneously with HPV-gBsec/gDsec ivag and gD2t-alum-MPL i.m. As a control, a group of mice was prime-boost immunized ivag with HPV16 and HPV45 expressing RFP. (A) gD-specific IgG antibody titers were measured in serum samples. Each bar represents the geometric mean of the endpoint titers of serum samples (error bars, 95% confidence intervals). Symbols represent results for individual serum samples. (B) HSV-2 neutralization titers of serum samples. Data are expressed as means ± standard deviations. (C) Flow cytometry analysis of the K7/gBsec tetramer in peripheral blood CD8+ T cells after priming and booster immunization. Each set of symbols connected by a line represents the results for an individual mouse after primary immunization (open circles) and booster immunization (open squares).
HPV-gBsec/gDsec, whether alone or combined with i.m. gD2t in alum-MPL, had lower levels of viral shedding in genital swabs than HPV control-immunized mice (Fig. 7B). The development of genital disease is a hallmark of primary human HSV-2 infection and can be characterized by redness, swelling, hair loss, and genital lesions (Fig. 7C). All mice in the groups immunized ivag with the HPV control vector or i.m. with gD2t in alum-MPL developed mild to severe genital disease (Fig. 7C). Strikingly, 40% of mice immunized ivag with HPV-gBsec/gDsec (with or without i.m. gD2t-alum-MPL) did not develop genital disease and survived (Fig. 7D).

Based on these findings, we defined three subgroups of mice: those that did not survive, those that survived but developed genital disease, and those that survived and were free of genital disease. Within the group immunized i.m. with gD2t-alum-MPL, there was no difference in viral shedding over time between the subgroup of mice that did not survive and the subgroup that survived with genital disease (Fig. 8A and B). Similarly, in the groups immunized ivag with HPV-gBsec/gDsec, with or without i.m. gD2t-alum-MPL, there was no difference in viral shedding over time between the subgroup that did not survive and the subgroup that survived with genital disease. However, for the subgroups of

### TABLE 1 HPV16 and HPV45 neutralization activities in serum after ivag immunizationa with HPV PsV

<table>
<thead>
<tr>
<th>Immunization</th>
<th>Geometric mean neutralization titerb (95% CI) in serum samples</th>
</tr>
</thead>
<tbody>
<tr>
<td>HPV control</td>
<td>862.6 (232–3,202) 348 (699–221)</td>
</tr>
<tr>
<td>gD2t</td>
<td>&lt;40 (N/A)</td>
</tr>
<tr>
<td>HPV-gBsec/gDsec</td>
<td>1,936 (942–3,552) 1,087 (367–3,214)</td>
</tr>
<tr>
<td>HPV-gBsec/gDsec + gD2t</td>
<td>1,879 (994–3,552) 1,301 (509–3,321)</td>
</tr>
</tbody>
</table>

a Mice were primed and boosted 1 month apart, and serum was collected 4 weeks after each immunization.
b Expressed as the 50% inhibitory concentration, determined using GraphPad Prism. 95% CI, 95% confidence interval; N/A, not applicable.

**FIG 7** Protection induced by combined immunization with HPV-gBsec/gDsec ivag and gD2t-alum-MPL i.m. Medroxyprogesterone acetate-treated mice (*n*, 10/group) were immunized twice, 1 month apart, with 3 μg gD2t with 50 μg alum and 7.5 μg MPL (gD2t/A/MPL). A second group of mice was primed ivag with a mixture of 10⁸ IU of HPV16-gBsec and 10⁸ IU of HPV16-gDsec, followed 1 month later by ivag booster immunization with HPV45 expressing the respective antigens (10⁸ IU each). A third group of mice was primed simultaneously with a mixture of 10⁸ IU of HPV16-gBsec and 10⁸ IU of HPV16-gDsec given ivag and gD2t-alum-MPL given i.m., followed 1 month later by booster immunization with HPV45 expressing 10⁸ IU of each of the respective antigens given ivag and with gD2t-alum-MPL given i.m. As a control, a fourth group of mice was prime-boost immunized ivag with HPV16 and HPV45 expressing RFP. Mice were challenged ivag 1 month after the last immunization with 10⁴ PFU of HSV-2 strain 333. (A) Survival of mice after HSV-2 challenge. Statistical significance was measured by a log rank (Mantel-Cox) test. (B) Viral shedding measured at the indicated days in genital swabs by plaque assay titration. Means ± standard deviations are shown. Statistical analysis by two-way ANOVA (*P = 0.02) was followed by multiple-comparison analysis. Symbols indicate significant differences between the HPV control and HPV-gBsec/gDsec (*, *P < 0.05) or HPV-gBsec/gDsec plus gD2t-A/MPL (#, *P < 0.05; ##, *P < 0.01). (C) Representative pictures of local manifestations of pathology in mice infected with HSV-2. (D) Percentages of mice that did not survive (filled bars), that survived but developed genital disease (hatched bars), or that survived and had no genital disease (open bars).
mice that survived free of genital disease, viral titers in the genital swabs remained under 10 PFU per ml over time (Fig. 8C and D). Finally, the frequency of circulating gB-specific CD8⁺ T cells was inversely correlated with viral shedding in the genital swabs of mice immunized ivag with HPV-gBsec/gDsec (r = −0.78; P = 0.005) or immunized ivag with HPV-gBsec/gDsec and i.m. with gD2t-alum-MPL (r = −0.69; P = 0.014) (Fig. 9A and B), whereas serum gD-specific antibody titers did not correlate or showed a weak positive correlation with viral shedding in groups immunized i.m. with gD2t-alum-MPL only (r = 0.57; P = 0.05) or i.m. with gD2t-alum-MPL and ivag with HPV-gBsec/gDsec (r = 0.15; P = 0.34) (Fig. 9C and D). Similarly, HSV-2 neutralizing antibody titers did not correlate with lower viral shedding (data not shown).

Together, these results indicate that ivag immunization with HPV-gBsec/gDsec is more effective than i.m. immunization with gD2t-alum-MPL at reducing genital disease and genital virus shedding after challenge with HSV-2.

**DISCUSSION**

Our study shows that transducing cervicovaginal epithelial cells in mice with HPV vectors expressing the gB and gD glycoprotein ectodomains of HSV-2 can preferentially induce HSV-specific intraepithelial-tissue-resident memory CD8⁺ T cells with a polyfunctional phenotype, together with modest levels of HSV-2 neutralizing antibodies. Secretion of gB was crucial to prime a CD8⁺ T cell response. Since gB expression in cultured cells was similar for the secreted, cytosolic, and membrane constructs, it is unlikely that a putative in vivo difference in expression levels accounted for the observed differences in immunogenicity. We have shown previously that CD4⁺ T cells are crucial for the priming of CD8⁺ T cells upon HPV vector ivag vaccination (38), so we speculate that secretion of gB and gD may facilitate the induction of CD4⁺ T helper cells, which, in turn, allow the priming of naïve CD8⁺ T cells. Our results are consistent with those of studies showing that secreted, but not intracellular, ovalbumin was able to induce CD4⁺ T cells upon i.n. immunization with an adenoviral vector (54) and that a secreted form of gD expressed by a DNA HSV vaccine induced a more potent IgG response than nonsecreted constructs (55).

HPV PsV have several characteristics that make them attractive vehicles for generating tissue-resident T cells after ivag delivery. First, they have a natural and highly restricted tropism for the genital epithelium, and the transduced gene(s) is only transiently expressed in the epithelium. Second, no HPV genes are expressed from the pseudogenomes, so there is no potential reduction in immunity to the target antigen because of immunodominance of an HPV viral antigen. Third, the recent cessation of an HIV trial (HVTN 502) highlighted the risk of potentiating HIV infection through the induction of HIV-susceptible CD4⁺ T cells (56). Because ivag genetic vaccination with HPV vectors results in highly skewed CD8⁺ T cell responses, it should minimize the risk of...
inducing HIV-susceptible target CD4+ T cells in the cervicovaginal mucosa. Occasionally, we observed a <2-fold increase in the level of CD4+ T cells in HPV vector ivag-immunized groups, which is much less pronounced than the 20-fold increase in the level of CD8+ T cells. Also, HPV vector ivag immunization did not induce expression of HIV coreceptor α4β7 by cervicovaginal CD4+ T cells (data not shown) (57, 58), in contrast to marked induction of this coreceptor after vaccination with an adenovirus 5 vector (56). Finally, there are many human and animal papillomavirus types that are distinct serotypes. Therefore, the induction of neutralizing antibodies by the priming dose of vaccine can easily be overcome by boosting with a heterologous type, as demonstrated here (38).

The mouse model of genital herpes disease has been useful for studying the roles played by innate and adaptive immunity in protection during primary infection or the mechanisms of protection conferred by vaccination; however, it has limitations (59). Vaginal HSV-2 infection in the mouse does not lead to recurrent genital lesions. Consequently, the primary endpoint of many mouse challenge studies has been protection from lethal HSV infection. Protection from death appears to be primarily antibody mediated (60, 61), in agreement with our finding that i.m. injection of HSV-2 gD in alum-MPL induced high titers of HSV-2 neutralizing antibodies and strong protection from lethal challenge, whereas ivag immunization with HPV-gDsec/gBsec induced lower neutralizing antibody titers and less protection from lethal challenge. However, death is an extremely rare consequence of human HSV-2 infection; thus, this endpoint would not be appropriate for clinical trials of HSV-2 vaccines.

Genital lesions and viral shedding are frequent and important manifestations of HSV-2 infection in humans, and therefore, these parameters, if they are predictive of what happens in people, may be more relevant endpoints for preclinical animal studies of prophylactic HSV vaccines. It is striking that i.m. gD2t in alum-MPL provided no protection against these endpoints, despite inducing high levels of serum neutralizing antibodies. If these interpretations are correct, our findings are consistent with the negative results of the recent clinical trial of an HSV-2 gD subunit vaccine, which did not prevent HSV-2 infection or disease (22). Interestingly, the HSV-2 vaccine results contrast with those observed in preclinical and clinical trials of the HPV virus-like particle (VLP) vaccine, where serum neutralizing antibodies are believed to be the principal mediator of sterilizing immunity to genital HPV infection. It is currently unclear why serum neutralizing antibodies can protect against death due to HSV infection but not against genital lesions or shedding, whereas serum antibodies can induce strong protection against genital HPV infection. Possible explanations include the facts that HSV has a large number of glycoproteins and capsid proteins relative to HPV, that HSV infects the external epithelial cells while HPV infects the cells at the basement membrane, and that HSV establishes latency and reactivates from neurons while HPV does not.

In contrast to i.m. injection of gD2t protein, ivag instillation of HPV PsV expressing secreted gB and gD induced strong protection against genital disease and virus shedding in a substantial percentage of mice. The direct correlation between the induction of gB-specific CD8+ T cell responses after genital immunization, but not systemic antibody responses after i.m. immunization, and shedding of virus from the vagina supports the hypothesis that these T cells are the primary effectors of local protection. In agreement with this conjecture, intravital microscopy studies have shown that intraepithelial CD8+ T cells translocate extensively through the lower layers of stratified squamous epithelium (as found in the mouse vaginal tract) and arrest when they contact keratinocytes that present their cognate peptide/major histocompatibility class I (MHC-1) complex (62).

Several strategies might be employed to increase the level of protection with ivag PsV vaccination. Our finding that the combination of gB and gD ectodomains conferred better protection than each antigen given separately suggests that including additional HSV antigens may improve the breadth of the immune response induced by vaccination. In addition, a recent study showed that adding the immediate early gene ICP27 to a DNA vaccine expressing gB and gD dramatically increased the efficacy of the vaccine (63, 64). Since the induction of T cells that can interrupt initial genital epithelial infection, rather than a response that is limited to neutralizing antibody induction, is the main goal of our vaccine strategy, it will be important to evaluate HSV-2 genes that are expressed very early after infection so as to favor a rapid response to virus-infected cells prior to the expression of viral immune evasion genes.

Other strategies designed to preferentially generate local T cell responses to HSV-2 have been reported. A live attenuated HSV-2 vaccine with the thymidine kinase gene deleted (HSV-TK), delivered topically, is considered among the most effective experimental HSV vaccines. This vaccine induced sterilizing immunity to genital infection that was mediated by CD4+ and CD8+ T cells producing IFN-γ (28, 36, 65). Recent studies showed that topical immunization with live attenuated HSV-1-TK induced a pool of long-lived tissue-resident memory CD8+ T cells close to the site of initial vaccination that was able to respond rapidly and to confer enhanced protection against subsequent HSV-1 skin infection (29, 51). However, this attenuated vaccine has generated potential safety concerns because it can establish latent infection in the regional ganglia, and the vaccine strain is resistant to most antivirals (66). Other studies have shown that, compared with systemic vaccination alone, the combination of systemic vaccination followed by either local inflammation or topically administered chemotactic molecules can redirect recently activated circulating HSV-2-specific CD8+ T cells to the genital tract and confer better control of viral replication in HSV-2 genital infection models (30, 31). However, although this “prime-pull” strategy can recruit T cells into the genital tract, it does not lead to their local amplification. In contrast, intravaginal boosting with HPV PsV can induce substantial local proliferation of antigen-specific tissue-resident CD8+ T cells in response to local antigen production in the transduced keratinocytes (38).

Based on the findings reported in this study, we believe that intravaginal delivery of HPV PsV expressing secreted versions of multiple HSV-2 genes is an attractive strategy to pursue in developing an HSV-2 vaccine for women. While this study has concentrated on a prophylactic vaccine, the ability to preferentially induce intraepithelial CD8+ T cells might also be useful for a therapeutic HSV-2 vaccine. This possibility could be evaluated in the guinea pig model of genital HSV-2 infection, which, unlike the mouse model, results in spontaneous recurrent vesicular genital lesions.
ACKNOWLEDGMENTS

This work was supported by the intramural research programs of the National Institute of Allergy and Infectious Diseases and the National Cancer Institute, Center for Cancer Research, NIH.

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