Herpes simplex virus 1 (HSV-1) is a neurotropic human pathogen causing a wide range of disease states from mild orolabial lesions, the most common manifestation, to deadly replicating HSV-1 is required for HSK,5–7 a pathologic immune larization, and potentially loss of vision.2–4 Although actively recurrent syndrome in which chronic inflammation initiated by outcome of HSV-1 infection is herpes stromal keratitis (HSK), a response driven primarily by neutrophils and CD4
tissue.8,9 Herpes virus entry mediator (HVEM, Tnfrsf14) in the murine cornea during the course of herpes simplex virus 1 (HSV-1) infection, the impact of this expression on pathogenesis, and whether alterations in HVEM or downstream HVEM-mediated effects ameliorate corneal disease.

METHODS. Corneal HVEM levels were assessed in C57BL/6 mice after infection with HSV-1(17). Leukocytic infiltrates and corneal sensitivity loss were measured in the presence, global absence (HVEM knockout [KO] mice; Tnfrsf14−/−), or partial absence of HVEM (HVEM conditional KO). Effects of immune-modifying nanoparticles (IMPs) on viral replication, corneal sensitivity, and corneal infiltrates were measured.

RESULTS. Corneal HVEM+ populations, particularly monocytes/macrophages during acute infection (3 days post infection [dpi]) and polymorphonuclear neutrophils (PMN) during the chronic inflammatory phase (14 dpi), increased after HSV-1 infection. Herpes virus entry mediator increased leukocytes in the cornea and corneal sensitivity loss. Ablation of HVEM from CD45+ cells, or intravenous IMP therapy, reduced infiltrates in the chronic phase and maintained corneal sensitivity.

CONCLUSIONS. Herpes virus entry mediator was expressed on two key populations: corneal monocytes/macrophages and PMNs. Herpes virus entry mediator promoted the recruitment of myeloid cells to the cornea in the chronic phase. Herpes virus entry mediator-associated corneal sensitivity loss preceded leukocytic infiltration, suggesting it may play an active role in recruitment. We propose that HVEM on resident corneal macrophages increases nerve damage and immune cell invasion, and we showed that prevention of late-phase infiltration of PMN and CD4+ T cells by IMP therapy improved clinical symptoms and mortality and reduced corneal sensitivity loss caused by HSV-1.

Keywords: herpes simplex keratitis, immunopathology, nanoparticle
not been assessed.\textsuperscript{23,24} We recently have found that HVEM-dependent disease after HSV-1 infection at the corneal surface is mediated by a radiation-resistant cell type or types,\textsuperscript{18} leading us to hypothesize that HVEM on corneal epithelial cells, stromal fibroblasts, or resident stromal macrophages, which incompletely turn over after irradiation,\textsuperscript{25} may promote pathogenesis in our adoptive transfer model. We investigated the expression of HVEM on a variety of cell lineages before, during, and after acute HSV-1 infection and found that corneal monocytes/macrophages are the first population to express HVEM, followed by polymorphonuclear neutrophils (PMN), double-negative (DN) T cells, and CD4\textsuperscript{+} T cells. Increases in corneal HVEM expression were associated with loss of corneal sensitivity and leukocytic infiltrates, while loss of HVEM\textsuperscript{+} cells from the cornea, either by genetic ablation or by administration of immune-modifying particles (IMPs), improved disease. We propose that HVEM on corneal resident monocytes/macrophages is integral for the recruitment of PMNs and other inflammatory cells to the cornea as well as nerve damage and loss of corneal sensitivity. Treatment with IMPs prevents the mobilization of immune cells to the cornea without hindering viral clearance and could show promise as a new therapeutic for treatment of chronic inflammation in HSK.

**Materials and Methods**

**Ethics Statement**

This study was performed according to recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health and adhered to the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research. The Committee on the Ethics of Animal Experiments of the Northwestern University approved the protocol (protocol Nos. 2012-1758 and IS00001532). Ketamine/xylazine anesthesia was used during procedures to minimize suffering.

**Cells and Viruses**

Plaque assays were performed and virus was propagated on African green monkey kidney cells (Vero) as previously described\textsuperscript{18}; HSV-1 strain 17 was obtained from David Leib (Dartmouth Medical School, Hanover, NH, USA).

**Animal Procedures**

The Animal Care and Use Committee at Northwestern University approved all procedures, in strict adherence to institutional and National Institutes of Health guidelines. Mice were maintained in a specific pathogen-free environment and were transferred to a containment facility after infection. Male 9- to 15-week-old C57BL/6, BALB/c, and Tnfrsf14\textsuperscript{+/-} (HVEM KO) were used in the study. Conditional HVEM KO mice were generated by crossing B6.Tg(Vav1-icre)A2Kio mice (Jackson, KO) were used in the study. Conditional HVEM KOs were for treatment of chronic inflammation in HSK.

**Flow Cytometry**

Corneal pairs and spleens from individual mice were collected in cold PBS. Corneas were digested in 0.7 mg/mL Liberase (Roche, Indianapolis, IN, USA) in RMPI media for 1 hour in a 37°C, 5% CO\textsubscript{2} incubator. Using a 1-mL syringe plunger, corneas were homogenized on top of a 100-μm mesh, washed with cold PBS, strained through a 40-μm mesh, and collected into a small volume. Spleens were prepared similarly to the corneas, but with a red blood cell lysis step between straining steps. After obtaining live cell counts, all of each cornea sample and a portion of each spleen sample were incubated with a 1:1000 dilution of Live/Dead Fixable Aqua Dead Cell Stain Kit (Thermo Fisher Scientific) in PBS in the dark at RT for 30 minutes. Samples were washed with PBS and incubated with Fc block (0.5–1.0 μg/sample anti-mouse CD16/CD32 [eBioscience, San Diego, CA, USA] in PBS + 1% fetal bovine serum + 0.1% sodium azide [FACS buffer]) for 5 minutes at 100°C followed by citrate buffer (pH 6.0). Secondary antibodies labeled with horseradish peroxidase (HRP) were visualized after treatment with chromogen diaminobenzidine (Vector Labs, Burlingame, CA, USA). After washing, slides were stained with Gill’s Hematoxylin and imaged on the EVOS XL Core Imaging System (Thermo Fisher Scientific, Carlsbad, CA, USA).

**Eye Swabs and Viral Plaque Assay**

Eye swabs were collected as previously described into 1 mL DMER media (DMEM containing 5% (vol/vol) fetal bovine serum (FBS), 1% gentamicin, 1% ciprofloxacin, and 1% amphotericin B) and stored at −80°C.\textsuperscript{18} Samples were thawed and vigorously vortexed for 30 seconds, and titers were determined with a standard plaque assay on Vero cells.

**Immunohistochemistry**

Whole eyes were collected 1 day post infection (dpi), rinsed with phosphate-buffered saline (PBS), floated in 10% formalin + neutral buffered PBS for 24 hours, transferred to 70% ethanol, and stored at 4°C until paraffin embedding. Serial 4-μm-thick sections were mounted on glass slides. The Northwestern University Mouse Histology and Phenotyping Laboratory provided naive murine spleen controls. The following antibodies and concentrations were used for immunohistochemical (IHC) staining: rabbit polyclonal antibody (Patricia Spear, Northwestern University) or mouse monoclonal antibody (clone HMMHV-1B18; BioLegend, San Diego, CA, USA) anti-HVEM antibodies diluted 1:200. Antigen retrieval was performed manually with Biocare (Birmingham, UK) decloaker for 5 minutes at 100°C followed by citrate buffer (pH 6.0). Secondary antibodies labeled with horseradish peroxidase (HRP) were visualized after treatment with chromogen diaminobenzidine (Vector Labs, Burlingame, CA, USA). After washing, slides were stained with Gill’s Hematoxylin and imaged on the EVOS XL Core Imaging System (Thermo Fisher Scientific, Carlsbad, CA, USA).
was stopped at 100,000 live cells, and data analysis was performed with FlowJo 10.1 software (Ashland, OR, USA).

**Corneal Sensitivity**

A Luneau Cochet-Bonnet esthesiometer (No. WO-7760; Western Ophthalmics, Lynnwood, WA, USA) was used to determine the blink threshold of the central cornea. Animals were scruffed, and the length of the monofilament was varied from 6.0 to 0.5 cm and touched perpendicularly to the surface of the central cornea until the first inflection point. A positive response was recorded when two blinks or more were obtained out of three attempts. Absence of a blink response at 0.5 cm was scored as a 0. The same examiner performed all measurements.

**Immune-Modifying Nanoparticle Treatment**

Negatively charged IMPs derived from poly(lactic-co-glycolic acid) (PLGA) were produced by Phosphorex. Immune-modifying nanoparticles were diluted to a previously established maximally effective concentration of 4.7 mg/mL in filtered PBS (0.94 mg total/mouse/injection); 200 μL IMPs or PBS vehicle control were delivered intravenously for 5 days (starting 3 dpi) via tail-vein injection. A control cohort of animals was killed 24 hours after the final dose for analysis of spleens. Associations between corneal blink threshold and viral load or CD45+, PMN, or macrophage cell number were assessed via linear regression. Kaplan-Meier mortality curves were compared by using the log-rank test. All statistics were calculated by using the GraphPad Prism 6.0f software (La Jolla, CA, USA).

**RESULTS**

**Herpes Simplex Virus 1 Infection Expands HVEM+ Populations in the Cornea**

From bone marrow chimera experiments, we localized HVEM-mediated pathogenesis to a radiation-resistant cell type or types. We reinvestigated the expression of HVEM in the cornea by immunohistology and flow cytometry, as a previous report indicates that HVEM is widely expressed in the naive murine corneal epithelium in vivo, and that its expression increases in the epithelium and stroma as early as 1 day after infection in BALB/c mice. The same rabbit polyclonal anti-HVEM antibody, R11874 (Patricia Spear, Northwestern University), produced nonspecific, background staining of the corneal epithelium of both wild-type (WT, C57BL/6) and HVEM KO (Tnfrsf14−/−) 9- to 12-week-old male mice 1 dpi with 2.0 × 10⁶ PFU/5 μL/eye HSV-1 strain 17 or post mock-infection (Vero cell lysate) after scarification (Figs. 1A–D). In contrast, a commercially available mouse monoclonal antibody to HVEM, HMHV-1B18 (BioLegend), produced little positive staining in the cornea 1 dpi (Figs. 1E–H). As a positive control for the monoclonal antibody, we tested HMHV-1B18 in naive murine spleens, which were floridly positive for HVEM after staining (Figs. 1I, 1J), consistent with previously described expression on B cells, T cells, myeloid cells, dendritic cells (DCs), and...
other leukocytes. These findings suggest HVEM expression in the corneas of C57BL/6 mice is limited immediately after scarification or infection.

We used flow cytometry to quantify HVEM expression in pairs of corneas from naive, mock-infected, or infected adult WT or HVEM KO control mice 3 or 14 dpi (Fig. 1K; Supplementary Fig. S1). Herpes virus entry mediator KO samples, an isotype control antibody, and fluorescence minus one control were used to define the threshold for HVEM positivity. Wild-type corneas contained a significantly higher proportion (Fig. 1K) and greater absolute number (Fig. 1L) of HVEM+ cells 3 and 14 dpi than naive corneas or mock-infected corneas, and this increase was specific to the cornea, as no such expansion occurred in the spleen (Fig. 1M). Most of the expanded HVEM+ population was attributable to an increase in HVEM+/CD45+ cells both at 3 dpi and 14 dpi (compare Fig. 1K to 1N; Supplementary Fig. S1). These findings indicate that HVEM, rather than being highly expressed by the murine corneal epithelium, is limited to CD45+ populations and is induced after infection with HSV-1 in a time-dependent manner.

**Most HVEM+ Cells in the Acutely Infected Murine Cornea Derive From the Monocyte/Macrophage Lineage**

To further characterize corneal HVEM expression, we analyzed HVEM+ corneal isolates from adult WT mice 3 dpi for the pan-leukocyte marker CD45, the endothelial marker CD31, and the epithelial markers E-cadherin and ICAM-1 by flow cytometry.

Isotype controls were used to set thresholds for each. Mock-infected corneas contained few HVEM+ cells, and few of those were CD45+, while the HVEM+ cells from infected corneas 3 dpi were mostly CD45+ leukocytes (Figs. 2A, 2B; Supplementary Fig. S1). Epithelial cells, defined in this study as CD45+/E-cadherin+ICAM1 epithelial cells, or CD45+/CD31+ endothelial cells. Corneal isolates were also evaluated for leukocytic lineage markers, including CD3, CD11b, Ly6C, Ly6G, and CD11c. (C) Percentage of HVEM+/CD45+ cells expressing the lymphoid marker CD3 or the myeloid marker CD11b 3 dpi. (D) Percentage of HVEM+/CD45+/CD3+/CD11b+ myeloid cells categorized as inflammatory monocyte/macrophage lineage (IMs; CD11c+/Ly6C+Ly6G–), PMNs (CD11c+/Ly6C+Ly6G+), or dendritic cells (CD11c+/Ly6G–). (E) Representative dot plot of CD11c versus Ly6C expression of HVEM+/CD45+/CD3+/CD11b+ myeloid cells from a WT cornea pair 3 dpi. Values for (B–E) are means ± SEM (n = 8 cornea pairs, two replicates).

**Herpes Virus Entry Mediator Promotes Loss of Corneal Sensitivity and Corneal Leukocytic Infiltration**

To determine whether the increase in HVEM+ cells over the course of HSV-1 infection impacts corneal physiology and function, we determined the corneal touch threshold in WT and HVEM KO mice corneally infected with HSV-1(17) with a Luneau Cochet-Bonnet esthesiometer as previously described. Briefly, the length of the filament was varied from 6 to 0.5 cm in increments of 0.5 cm and touched perpendicularly to the central cornea until the first inflection point. The blink threshold was counted as the length at which the animal blinked two times or more out of three; if no
response occurred at 0.5 cm, the eye was scored as 0. After infection, WT mice rapidly and dramatically lost sensitivity in the central cornea, compared to HVEM KO mice, in which corneal reflexes were largely maintained (Fig. 3A). Along with experiencing a loss of sensitivity, WT corneas also became more heavily infiltrated by CD45$^+$ leukocytes, as determined by flow cytometry on day 14 (Fig. 3B).

To determine if a correlation between these two factors existed, we performed a linear regression on the average blink threshold length at the end of the experiment, day 14, compared to the number of CD45$^+$ cells in the cornea on that same day (Fig. 3C), and found a significant negative association (R$^2$ = 0.338, P = 0.0005). Characterized by cell type, similar negative correlations also existed between number of PMNs (Fig. 3D) and number of monocytes/macrophages (data not shown) and the blink threshold length (R$^2$ = 0.374, P = 0.0002 and R$^2$ = 0.2992, P = 0.0012, respectively). We collected eye swabs from infected mice 1, 3, and 5 dpi to determine viral loads in the tear film, and found higher day-1 titers from HVEM KO eye swabs than from WT (data not shown), similar to what we have previously reported. Comparison of day 1 (but not day 3 or day 5) eye swab titers to the blink threshold (day 14) for each eye individually revealed a significant but small negative correlation (Fig. 3E; R$^2$ = 0.145, P = 0.0155).

We next investigated the specific identities of the CD45$^+$ populations in WT and HVEM KO corneas 14 dpi (see Fig. 2C; see Supplementary Figs. S2, S3 for gating strategy). Herpes simplex virus 1–infected WT corneas tended to contain higher proportions of CD4$^+$ and CD8$^+$ T cells on day 14 than HVEM KO corneas, although these differences did not reach statistical significance (Fig. 3F). Levels of mDCs (CD45$^+$/CD3$^-$/CD11b$^-$/Ly6G$^-$/CD11c$^-$), monocytes/macrophages (Ms: CD45$^+$/CD3$^-$/CD11b$^-$/Ly6G$^-$/CD11c$^-$), and specifically, inflammatory monocytes/macrophages (Ly6C$^+$ monocytes/macrophages) in WT corneas were higher than in HVEM KO corneas to the 0.05 significance level, but after correction for multiple comparisons, only the level of PMNs remained statistically significantly higher in WT corneas than in HVEM KO corneas (Fig. 3G).

Wild-type corneas tended to contain more CD4$^+$ and DN T cells, mDCs, and monocyte/macrophages than HVEM KOs during acute infection (5 dpi), although these differences were not statistically significant (data not shown). Polymorphonuclear neutrophils were notably absent from the corneas of both genotypes at this time point, consistent with previously published reports that neutrophils invade the cornea in two temporally distinct waves. At 14 dpi, most HVEM$^+$/CD45$^+$ cells in WT corneas remained CD3$^-$/CD11b$^+$ myeloid cells, although approximately 20% of HVEM$^-$/CD45$^+$ cells were CD3$^+$ (data not shown). Most HVEM$^+$/CD3$^-$/CD11b$^+$ cells were PMNs, signaling a switch
from the acute phase infection, when corneal monocytes/macrophages represented the greatest HVEM⁺ population (comparing Fig. 2D to Fig. 3H). Of HVEM⁺/CD3⁺/CD11b⁺ cells, most were CD8⁻/CD4⁻ (as well as NK1.1⁻) cells, here called DN T cells, although this population could potentially represent γδ T cells. A small population of HVEM⁺/CD4⁺ also occurred in the cornea 14 dpi (Fig. 3I). These data indicate that HVEM promotes an increase in viral replication, as previously reported, as well as increased corneal infiltration, both of which correlated with a loss of central corneal sensitivity.

**Ablation of HVEM From CD45⁺ Lineages Reduces Myeloid Infiltrates Late in Infection**

We specifically ablated HVEM from CD45⁺ cells by using a Cre/lox system. HVEM₀/₀ mice on the C57BL/6 background contain loxP sites flanking exons 3 and 6 of the HVEM gene. These animals were bred to B6.Tg(Vav1-icre)A2Kio mice (Jackson), which express Cre recombinase under control of the Vav promoter, resulting in expression in 98% to 100% of hematopoietic cells with no littermate mosaicism. Genotyping was performed by PCR analysis of floxed HVEM and Vav-Cre genes. In addition, we confirmed that homozygous floxed animals expressing Vav-Cre (Vav⁺ HVEM₀/₀) had successful ablation of HVEM from CD45⁺ cells in comparison to homozygous HVEM₀/₀ controls by analysis of peripheral blood (data not shown) and infected corneas via flow cytometry for HVEM expression (Fig. 4A). Although CD11b⁺ myeloid cells had lost HVEM expression, this did not impact the presence of CD11b⁺ cells in the cornea 3 dpi (Fig. 4B). Consistent with HVEM KOs, Vav⁺ HVEM₀/₀ corneas had significantly lower titers than controls early after infection (Fig. 4C). We also assessed late-phase populations of corneal leukocytes to determine if HVEM-dependent infiltration required HVEM expression on CD45⁺ cell types. Infiltration of CD4⁻ and CD8⁻ T cells was diminished in Vav⁺ HVEM₀/₀ mice compared to controls, although this did not reach statistical significance (Fig. 4D). However, myeloid cell infiltration into the cornea 14 dpi was markedly blunted when HVEM was ablated from CD45⁺ cells (Fig. 4E). Corneal sensitivity in Vav⁺ HVEM₀/₀ mice was also maintained, similar to that of HVEM KOs (data not shown). These findings suggest that the establishment of CD11b⁺ myeloid cells in the cornea in the acute phase of HSV infection is independent, but that the recruitment or maintenance of myeloid lineages in the chronic phase of infection cannot occur effectively without HVEM.

**Negatively Charged IMP Treatment Ameliorates Disease After Ocular HSV-1(17) Infection**

Negatively charged, 500-nm-diameter IMPs derived from PLGA limit tissue infiltration of inflammatory monocytes and other circulating phagocytic cells by rerouting them to the spleen for degradation. Immune-modifying nanoparticle therapy is effective in a wide range of inflammatory diseases, including West Nile infection, experimental autoimmune encephalitis, cardiac and kidney reperfusion injury, and others. Herpes virus entry mediator, which we found localized mostly on corneal inflammatory monocytes/macrophages during acute infection, and on PMNs during the chronic phase, promotes infiltration and is associated with sensory loss at the corneal surface. We hypothesized that IMP treatment could prevent the migration of circulating immune cells into the cornea, even in the presence of HVEM, and thus ameliorate corneal disease.

We performed a pilot study in adult male BALB/c mice, which are highly susceptible to neurologic morbidity after corneal HSV-1 infection, to determine if a previously established dose of 0.94 mg IMPs/mouse via tail-vein injection delivered daily for 5 days could improve survival when compared to mice receiving the vehicle, PBS. To target the later, pathologic influx of immune cells without hindering viral clearance, we began treatment 3 dpi, before the onset of clinical symptoms such as ruffled fur, periorbital swelling, and lesion development, which occur around day 5, but after an...
 innate response had been mounted. Treated mice survived at significantly higher rates (40% vs. 0%) than mice that received the vehicle (data not shown). We next tested the therapeutic potential of IMPs in C57BL/6 mice to prevent loss of corneal sensitivity, reduce corneal infiltration, and improve symptoms after HSV-1 infection. Immune-modifying nanoparticle–treated mice maintained corneal blink thresholds similar to preinfection levels, while corneal sensitivity in vehicle-treated animals significantly declined by 6 dpi (Fig. 5A). Viral titers in eye swabs collected 1, 3, and 5 dpi did not differ between the treatment groups (Fig. 5B).

Flow cytometry of corneas collected on day 14 revealed that IMP treatment significantly reduced CD45\(^+\) populations (Fig. 5C), particularly in all myeloid populations we investigated (Fig. 5D), and tended to decrease CD4\(^+\) and DN T cells (Fig. 5E). Populations of leukocytes present in the spleens of treated animals significantly declined by 6 dpi (Fig. 5A). Viral titers in eye swabs collected 1, 3, and 5 dpi did not differ between the treatment groups (Fig. 5B).

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indicates that HVEM-mediated pathogenesis is not strain-specific.\textsuperscript{21} It would be informative to repeat our assessment of leukocytic infiltrates in WT corneas instead infected with HSV-1(McKrae), as this strain does not require scarification before infection (to cause pathology\textsuperscript{26}), prior report\textsuperscript{27} we were unable to detect significant levels of HVEM on corneal epithelial cells by IHC 1 dpi; flow cytometry 3 and 14 dpi indicated most (roughly 75%) HVEM\textsuperscript{+} cells in the cornea were CD45\textsuperscript{+} leukocytes. In a previous study using bone marrow chimeras with WT and HVEM KO mice, we have found that HVEM on radiation-resistant lineage(s) is sufficient to mediate disease after HSV infection.\textsuperscript{18} The naive cornea, once thought to be devoid of immune cells, contains numerous CD11b\textsuperscript{+} cells with variable major histocompatibility complex (MHC) class II expression; most lack CD11c and Ly6G and therefore are of the monocyte/macrophage lineage.\textsuperscript{25,56} Because a sizable portion (25\%) of in vivo corneal resident macrophages do not turn over post irradiation even after 8 weeks of recovery,\textsuperscript{25} we hypothesize that this cell type, essentially the only lineage to express HVEM early after infection, is the radiation-resistant population sufficient for HVEM-mediated HSV-1 pathogenesis.

Herpes virus entry mediator on corneal macrophages may promote immunopathogenesis by recruiting other inflammatory cells to the eye. Herpes virus entry mediator expression increases the levels of macrophage-associated chemotactic factors in the cornea, such as CXCL10 and CCL3, which are known to recruit PMNs, monocytes, and T cells during ocular HSV-1.\textsuperscript{18,57,58} Herpes virus entry mediator upregulates these factors independently of viral entry,\textsuperscript{18} consistent with the observation that, specifically in the cornea, levels of infectious virus do not correlate with chemokine expression.\textsuperscript{59} In line with this hypothesis, we found that genetic ablation of HVEM from CD45\textsuperscript{+} cells through a Cre/lox system prevented infiltration by myeloid cells and tended to limit CD4\textsuperscript{+} populations. We also found that viral titers 1 dpi were higher in the control mice than in the conditional HVEM KO s, suggesting that HVEM on CD45\textsuperscript{+} cells also has an early impact on viral replication. Herpes virus entry mediator increases viral titer independently of entry.\textsuperscript{18} Herpes virus entry mediator on leukocytes may indirectly improve replication in or survival of infected corneal epithelial cells or fibroblasts, perhaps through increased expression of other HSV receptors or replication factors, although this mechanism requires further investigation.

In human patients with HSK, corneal sensitivity to mechanical stimulation is significantly impaired.\textsuperscript{40} In our murine model, HVEM promoted loss of corneal sensitivity, which others have correlated with loss of sensory nerve endings from the cornea.\textsuperscript{44} Immune cells and inflammatory mediators, such as IL-6, promote nerve retraction rather than direct effects of viral replication, and treatment with dexamethasone reduces corneal sensory nerve losses.\textsuperscript{41,42} Some corneal monocytes/macrophages reside in close proximity to corneal nerve endings,\textsuperscript{44} raising the intriguing possibility that resident HVEM\textsuperscript{+} macrophages could aggravate nerve damage during acute infection. Consistent with this theory, WT mice had already begun losing corneal sensitivity, compared to HVEM KO s, as early as 2 to 3 dpi, at which time corneas contained significant HVEM\textsuperscript{+} monocyte/macrophage cells but few other infiltrates, indirectly suggesting nerve damage precedes large-scale leukocyte invasion.

In fact, nerve damage likely contributes to further leukocytic infiltration of the cornea. Neuropeptides released from corneal nerve endings, such as substance P and calcitonin gene-related peptide, can induce IL-8 transcription in corneal epithelial cells, recruiting PMNs,\textsuperscript{45,46} substance P has also been reported to increase the severity of HSK lesions.\textsuperscript{47} Another mechanism proposed by Yun and colleagues\textsuperscript{41} posits that HSV (or, more likely, the initial inflammation associated with viral replication) damages corneal nerves, and subsequent desiccation due to lack of blinking leads to chronic, bilateral leukocytic infiltration. In addition, aberrantly organized sensory nerves have been found to reinnervate the corneal stroma after sensory nerve retraction during HSV-1 infection.\textsuperscript{43} The elimination of these fibers through superior cervical ganglionectomy reduces neovascularization and opacity, suggesting the influx of sympathetic fibers rather than a paucity of sensory fibers may be responsible for HSK symptoms.\textsuperscript{43} In this study, we found that large populations of leukocytes in the cornea, especially PMNs and monocytes/macrophages, correlated with a prolonged lowered blink threshold 14 dpi. When HVEM ablation or treatment with IMPs 3 to 7 dpi prevented late-phase corneal infiltration, corneal sensitivity was preserved, suggesting nerve damage is exacerbated or perpetuated by PMN, CD4\textsuperscript{+}, and other chronic-phase infiltrates.\textsuperscript{41} Coincidently, these populations were also HVEM\textsuperscript{+}, although expression on these radiation-sensitive populations did not contribute significantly to pathogenesis in our earlier bone marrow chimera experiments.\textsuperscript{18} We find it likely that both early and late inflammatory cells contribute to loss of corneal sensitivity during HSV infection: resident macrophages may secrete cytokines or other factors that damage corneal nerves, decreasing blinking and drying the eye, leading to immune cell infiltration, which in turn damages corneal nerves further.

Regardless of the precise recruitment mechanism of immune cells in the chronic phase, it is well established that their presence in the cornea promotes neoangiogenesis and neolympangiiogenesis,\textsuperscript{48,49} opacification,\textsuperscript{50} and scarring.\textsuperscript{51,52} CD4\textsuperscript{+} T cells also prevent reinnervation of the cornea, sustaining loss of the blink response and desiccation-related inflammation.\textsuperscript{53} However, loss of immune cells, particularly macrophages, can be equally devastating, as unchecked viral replication leads to central nervous system (CNS) invasion and mortality; depletion of Gr-1\textsuperscript{+} cells with a monoclonal antibody before and during infection increases replication, spread to the skin and brain, and mortality in BALB/c mice.\textsuperscript{53} Cladonidine liposomes, which nonspecifically deplete phagocytic cells including macrophages, injected subconjunctivally before infection severely heighten viral replication, blepharitis, and epithelial keratitis; however, stromal keratitis is mildly improved after depletion.\textsuperscript{54–56} In contrast, delaying depletion to 2 and 4 dpi does not enhance viral replication.\textsuperscript{54} These effects are likely due to macrophages, as targeted depletion of neutrophils with a Ly6G antibody does not alter viral replication or pathogenesis.\textsuperscript{57}

Knowing this, we hypothesized that therapy in the prechronic phase (3–7 dpi) with IMPs, which are efficacious in the treatment of a host of inflammatory disorders,\textsuperscript{26,58} could limit HSK symptoms without enhancing viral replication. Immune-modifying nanoparticles are absorbed by circulating engulfing cells; in inflammatory monocytes/macrophages, negatively charged particles are taken up in an opsonin-independent manner by the MARCO (macrophage receptor with collagenous structure) receptor.\textsuperscript{26} This process redirects them to the spleen, where they undergo apoptosis, preventing tissue damage caused by these cells at extrasplenic sites.\textsuperscript{26} In this study, the corneas of IMP-treated mice contained significantly fewer CD45\textsuperscript{+} cells in general; both lymphoid (CD4\textsuperscript{+}, natural killer T [NKT], and DN) and myeloid cell types (PMNs, monocytes, macrophages, and mDCs) were excluded from the cornea, with a concomitant preservation of corneal blink response. Initiation of treatment 3 dpi had no impact on viral loads, and actually improved mortality due to CNS involvement, indicating immune control of viral spread occurs within the first 72 hours after infection; after this, immune...
populations in the cornea exacerbate rather than ameliorate pathology.\textsuperscript{4}

Although previous work has focused on the role of IMPs in preventing inflammatory monocyte-driven pathology, in our model IMP therapy was also effective in limiting PMN, CD4\textsuperscript{T} and DN T cells from the cornea.\textsuperscript{29} There is some evidence that IMPs can act directly on these cell types,\textsuperscript{38} although it is also possible that without inflammatory monocytes/macrophages in the cornea, T cell and neutrophil populations are not mobilized to the cornea. The characteristics and role of CD3\textsuperscript{\beta}/CD4\textsuperscript{T}/CD8\textsuperscript{\beta} DN cells during HSK requires a more thorough investigation, as these cells could represent γδ T cells rather than true DN cells. Regardless, it is clear that IMP therapy limits their presence in the cornea. While acutely induced populations control viral spread, PMNs and macrophages in the chronic phase orchestrate much of HSK-related tissue damage.\textsuperscript{4,8,59} Importantly, CD4\textsuperscript{T} T cells are considered the primary pathologic cell type in HSK, and recent findings suggest depletion of this cell type allows for reversal of nerve loss and corneal damage.\textsuperscript{41} Because IMP treatment significantly limited PMN and CD4\textsuperscript{T} T cells in the cornea, this therapy would seem to be beneficial to the treatment of HSK inflammation.

In conclusion, we propose that HVEM on resident corneal monocytes/macrophages promotes early nerve damage, leading to increased infiltration of the cornea by a variety of leukocytes and furthering loss of corneal sensitivity. Interruption of this process, either by HVEM ablation from CD4\textsuperscript{\beta} cells or by treatment with IMPs, prevents the influx of circulating immune cells, maintaining corneal health. Given these promising results, we are hopeful that IMP therapy could be adapted to treat recurrent HSK in human disease.

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