Analysis of T Cell Responses during Active Varicella-Zoster Virus Reactivation in Human Ganglia

Megan Steain, Jeremy P. Sutherland, Michael Rodriguez, Anthony L. Cunningham, Barry Slobedman, Allison Abendroth

ABSTRACT

Varicella-zoster virus (VZV) is responsible for both varicella (chickenpox) and herpes zoster (shingles). During varicella, the virus establishes latency within the sensory ganglia and can reactivate to cause herpes zoster, but the immune responses that occur in ganglia during herpes zoster have not previously been defined. We examined ganglia obtained from individuals who, at the time of death, had active herpes zoster. Ganglia innervating the site of the cutaneous herpes zoster rash showed evidence of VZV-induced neuronal cell apoptosis, it is possible that immune cell-mediated injury to neurons and/or their protective satellite glial cells (SGCs) play an important role in controlling VZV replication in ganglia during active herpes zoster.

IMPORTANCE

VZV is responsible for both varicella (chickenpox) and herpes zoster (shingles). During varicella, the virus establishes a life-long dormant infection within the sensory ganglia and can reawaken to cause herpes zoster, but the immune responses that occur in ganglia during herpes zoster have not previously been defined. We examined ganglia obtained from individuals who, at the time of death, had active herpes zoster. We found that specific T cell subsets are likely to play an important role in controlling VZV replication in ganglia during active herpes zoster.

Varicella zoster virus (VZV) is a ubiquitous human pathogen responsible for both varicella (chickenpox) and herpes zoster (shingles). Herpes zoster results from reactivation of the virus in sensory ganglia, generally in the setting of reduced VZV-specific cell-mediated immunity (1, 2). Herpes zoster may be followed by postherpetic neuralgia (PHN), which is generally defined as pain persisting for greater than 90 days following the resolution of the herpes zoster rash (3). This pain can persist for years and can have a negative impact on patients’ quality of life (4). The cause of the pain is unknown; however, it has been suggested that damage to neuronal cells plays a role (5). We have previously demonstrated that VZV-infected neurons are resistant to apoptosis (6) due to the expression of the immediate-early gene (IE) product of open reading frame 63 (ORF63) (7). Thus, rather than being a consequence of VZV-induced neuronal cell apoptosis, it is possible that immune cell-mediated injury to neurons and/or their protective satellite glial cells (SGCs) is responsible for the pain associated with herpes zoster and particularly PHN.

Sensory neurons are completely enveloped by SGCs, which provide protection and nutritional support, and they may even play a role in neuronal signaling (8, 9). In most cases, each adult neuron and its surrounding SGCs form discrete individual units; however, the number of SGCs decreases with age (9). SGCs also form an important barrier around peripheral neurons in place of a vascular barrier (9) and share many properties of antigen-presenting cells (10). Thus, SGCs are likely to play an important role in the immune response to viral infections in sensory ganglia.

The immune response has been shown to play an important role in the ganglia during infection with herpes simplex virus (HSV), an alphaherpesvirus closely related to VZV. HSV-1 latency in sensory ganglia is characterized by a persistent CD8+ T cell infiltrate (11–13), and it has been shown that the cytolytic protein granzyme B cleaves the immediate-early HSV protein, ICP4 (homolog of VZV IE62), and that this plays an important role in preventing or limiting viral reactivation (14). Furthermore, it has been demonstrated that during latency the HSV latency-associated transcript (LAT) can prevent granzyme B from cleaving caspase-3 and prevent activation of this apoptotic pathway, ensuring both neuronal and HSV survival (15).

There has been no LAT equivalent described for VZV; however, mRNA transcripts from a number of ORFs, including...
TABLE 1 Antibodies used for immunohistochemistry and immunofluorescence staining

<table>
<thead>
<tr>
<th>Antibody</th>
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<th>Clonea</th>
<th>Dilution</th>
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<td>1:1,000</td>
<td>A gift from Ravi Mahalingam, University of Colorado, Denver, CO, USA</td>
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<tr>
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<td>SG1</td>
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<tr>
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<td>Anti-S100B</td>
<td>Goat polyclonal</td>
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<td>Rabbit polyclonal</td>
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<td>1:200</td>
<td>Novocastra, Melbourne, Australia</td>
</tr>
</tbody>
</table>

a NA, not applicable.

ORF21, ORF29, ORF62, ORF63, and ORF66, have been identified by DNA sequencing in latently infected ganglia (16–18). Previous studies examining VZV latently infected ganglia have also reported the detection of a number of VZV proteins by immunohistochemistry (11, 12, 17, 19–21), although examination of ganglia with latent VZV did not find an association between the expression of IE62 during latency and an inflammatory infiltrate, which was in contrast to HSV latency (11–13). It is important to note, however, that VZV protein expression during latency has only been demonstrated by immunohistochemistry (IHC) (3, 22), and that the expressed proteins have been detected within the cytoplasm of neurons, despite many of them typically displaying a nuclear localization within productively infected cells (22). Recently, concerns over the specificity of viral antigen immunostaining in adult human neurons have been raised. These include the presence of the neuronal pigments neuromelanin and lipofuscin and the presence of contaminating antibodies in rabbit and rat preparations that cross-react with blood group A antigens, except in the presence of the neuronal pigments neuromelanin and lipofuscin on specific antigen staining (24). For antibody information, see Table 1.

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RESULTS

Histological analyses of human ganglia during herpes zoster. A major obstacle to studying the interactions between VZV, neurons, and the immune response during herpes zoster is the difficulty in obtaining naturally infected human ganglia. We obtained multiple ganglia at postmortem from 2 patients who died with...
VZV ORF9a was performed. The results are summarized in Table 2. All ganglia used within this study were negative for HSV DNA as determined by PCR (data not shown). DNA extraction was performed on all ganglia, and PCR for VZV ORF9a was performed. The results are summarized in Table 2. All ganglia used within this study were negative for HSV DNA as determined by PCR (data not shown).

Hematoxylin and eosin staining was performed on multiple 5-μm sections from each ganglion. In the reactivated ganglion from HZ1 (LL2), a prominent, widespread infiltrate of inflammatory cells, predominantly small lymphocytes and macrophages, was evident (Fig. 1A). Focally there was perivascular and intramural inflammation with isolated thoracic lesions. The L2 dorsal root ganglion (DRG) was removed (reactivated ganglion, LL2). In addition, single left lumbar (LL1), right lumbar (RL), left thoracic (LT), and right cervical (RC) DRGs from uninvolved dermatomes were available for study.

The second patient (HZ2) was a 62-year-old man with acute myeloid leukemia (diagnosed postmortem) who developed herpes zoster ophthalmicus approximately 17 days prior to death. At autopsy a hemorrhagic vesicular and ulcerated rash was confined to the left L2 dermatome. The L2 dorsal root ganglion (DRG) was removed (reactivated ganglion, LL2). In addition, single left lumbar (LL1), right lumbar (RL), left thoracic (LT), and right cervical (RC) DRGs from uninvolved dermatomes were available for study.

Aspiration pneumonia, dementia Acute myeloid leukemia

Patient 1 (HZ1) was a 93-year-old man with frontotemporal dementia who developed herpes zoster approximately 17 days before death from aspiration pneumonia. At autopsy a hemorrhagic vesicular and ulcerated rash was confined to the left L2 dermatome. The L2 dorsal root ganglion (DRG) was removed (reactivated ganglion, LL2). In addition, single left lumbar (LL1), right lumbar (RL), left thoracic (LT), and right cervical (RC) DRGs from uninvolved dermatomes were available for study.

The second patient (HZ2) was a 62-year-old man with acute myeloid leukemia (diagnosed postmortem) who developed herpes zoster ophthalmicus approximately 17 days prior to death. At autopsy a hemorrhagic vesicular and ulcerated rash was confined to the left L2 dermatome. The L2 dorsal root ganglion (DRG) was removed (reactivated ganglion, LL2). In addition, single left lumbar (LL1), right lumbar (RL), left thoracic (LT), and right cervical (RC) DRGs from uninvolved dermatomes were available for study.

Aspiration pneumonia, dementia Acute myeloid leukemia

**TABLE 2 Patient ganglion samples used in this study**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value(s) for case no.(^a)</th>
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<tr>
<td>Age (yr)</td>
<td>CI C2 HZ1 HZ2</td>
</tr>
<tr>
<td>Gender</td>
<td>Female Male Male Male</td>
</tr>
<tr>
<td>Cause of death and underlying conditions</td>
<td>Metastatic adenocarcinoma Blunt-force head injury</td>
</tr>
<tr>
<td>Notes on herpes zoster</td>
<td>NA NA Rash at L2 dermatome, 17 days prior to death, rash still present at autopsy Rash at L2 dermatome, 17 days prior to death, rash still present at autopsy</td>
</tr>
<tr>
<td>Ganglia examined</td>
<td>1 DRG 2 DRG LL2 DRG (reactivated, ganglionitis) RTG (reactivated, ganglionitis)</td>
</tr>
<tr>
<td>Blood group</td>
<td>Non-A Non-A Non-A A</td>
</tr>
<tr>
<td>PCR for VZV DNA</td>
<td>ND ND LL2-Pos RL-ND LL1-ND RC-ND</td>
</tr>
</tbody>
</table>

**Abbreviations:** DRG, dorsal root ganglia; TG, trigeminal ganglia; IHC, immunohistochemistry; LL, left lumbar; RL, right lumbar; RT, right thoracic; LT, left thoracic; RC, right cervical; LC, left cervical; RTG, right trigeminal ganglia; NA, not applicable; ND, not detected; Pos, positive.

In HZ2, there was a significant infiltrate of inflammatory cells, including small lymphocytes in the RTG (reactivated ganglionitis) as well as the right cervical DRG (RC1). In one region of the RTG there was a localized hemorrhage (Fig. 1E and F). In all 7 ganglia examined from this patient, neurons and SGCs appeared to be intact. There was no inflammation in the three control ganglia, and the neurons and SGCs were intact (Fig. 1H).

**Detection of VZV antigens in human ganglia during herpes zoster.** To determine whether VZV antigen, indicative of viral replication, was present in ganglia innervating or distal to the site of the herpes zoster rash, we performed immunohistochemical staining for the VZV immediate-early protein IE63 using a rabbit polyclonal antibody (19), the late viral gene product glycoprotein E (gE; Millipore, Billerica, MA, USA), and the precursor and mature forms of gE and the gI complex (gEgI; Meridian Life Sciences, Saco, ME).
In the reactivated ganglion from patient HZ1 (LL2), intense cytoplasmic staining for VZV IE63 (Fig. 2A) was observed in neuron-SGC units within the area of necrosis. However, precise neuronal or SGC localization was not possible due to the degenerated state of the neurons and SGCs within this region (Fig. 2A). Antibody specificity was confirmed by staining consecutive sections with the appropriate nonimmune isotype antibody (Fig. 2B). In addition, IE63 staining was identified within axons in the necrotic area (data not shown). No IE63 staining was observed in the non-reactivated ganglia from HZ1 or in the control ganglia (Fig. 2C and data not shown).

Membranous and cytoplasmic gE staining in neuron-SGC
units was identified in the reactivated DRG from HZ1 (LL2). As with IE63 staining, positive neuron-SGC units were restricted to the area of necrosis (Fig. 2D). Isotype antibody staining of consecutive sections confirmed the specificity of the gE staining (Fig. 2E). Staining of nonreactivated ganglia from HZ1 or ganglia from the control cases did not yield any detectable gE staining (Fig. 2F and data not shown). In addition, a small number of gE-positive infiltrating cells were observed scattered throughout this ganglion (data not shown). Staining with the gE:gI antibody confirmed the presence of VZV glycoprotein-positive cells within the region of necrosis in the reactivated ganglion (LL2) from HZ1 (Fig. 2G). This antibody also stained membrane and cytoplasmic regions within neuron-SGC units. Staining was not observed when consecutive sections were incubated with isotype antibody (Fig. 2H) or when nonreactivated ganglia from HZ1 or ganglia from the control cases were stained for gE:gI (Fig. 2I and data not shown). No IE63-positive cells were identified in any of the 7 ganglia, including the reactivated RTG from the second patient with herpes zoster (HZ2). Staining for gE:gI was not observed in any ganglia from HZ2, with the exception of a few small infiltrating cells which were gE:gI positive within the reactivated RTG (data not shown). The gE antibody could not be used to assess ganglia from HZ2. This is because, unlike HZ1 and all of the control cases, HZ2 was blood group A, as determined by staining of ganglionic sections from all cases with an anti-blood group A antibody (Table 2), and it has been reported that this gE antibody (which is produced from ascites fluid) can induce a mouse ascites Golgi (MAG) reaction typified by punctate cytoplasmic/trans-Golgi network localized staining within neurons in ganglia from blood group A individuals (23). It was concluded that VZV antigens were readily detectable in the reactivated ganglion LL2 from HZ1 but only rarely detectable in the reactivated ganglion RTG from HZ2.

Characterization of the T cell infiltrate in human ganglia during herpes zoster. To characterize the T cell infiltrate within ganglia during herpes zoster, serial sections were immunofluorescently stained for the T cell markers CD3, CD4, and CD8, as well as granzyme B, to identify cytotoxic T cells and NK cells. Sections were costained with S100B to identify SGCs. Sections of control DRGs (negative control) and tonsil (positive control) were stained in parallel, and antibody specificity was confirmed by staining with the appropriate nonimmune isotype antibody (Fig. 3E and F and data not shown).

In the reactivated ganglia of HZ1 and HZ2 (LL2 and RTG, respectively), a widespread infiltrate of CD3+ T cells was apparent (Fig. 3A). Infiltrating T cells consisted of both CD4+ and CD8+ subsets (Fig. 3B and C). There were also numerous infiltrating cells that stained positive for the cytolytic marker granzyme B (Fig. 3D). The numbers of cells expressing CD3, CD4, CD8, or granzyme B were determined by counting multiple sections and averaging the results from 2 independent stains (Fig. 3G). The density of CD3+ T cells averaged across whole ganglia was highest in the reactivated ganglia, ranging from 525 cells per mm² (RTG) to >1,200 cells per mm² in LL2. Compared to control ganglia, CD4+ T cell densities were also slightly elevated in several ganglia in proximity to LL2 (i.e., LL1, RL, and LT) and RTG (i.e., RC) but not in more distant ganglia. Both CD4+ and CD8+ T cells were well represented in both reactivated ganglia, although there was a trend toward CD4+ T cell predominance, which was most evident in LL2. There was also an abundance of granzyme B-positive cells in the reactivated ganglia (501 per mm² [LL2] and 307 per mm² [RTG]). Compared to control DRGs, gran-
zyme B-positive cells were also increased in ganglia in proximity to LL2 (LL1, RL, and LT) and RTG (RC) but not in more distant ganglia. To further characterize the granzyme B-positive cells, dual staining for granzyme B and CD8 or CD4 was performed on sections from reactivated ganglia. Many CD8*/granzyme B-positive cells were identified (Fig. 4A to C), whereas there were no CD4*/granzyme B-positive cells identified (Fig. 4D to F). The number of granzyme B-positive cells within both reactivated ganglia exceeded the number of CD8-positive T cells (Fig. 3G), and given that CD4 T cells were shown to be noncytolytic (Fig. 4D to F), this difference may be attributed to infiltrating cytolytic natural killer (NK) cells. Staining for CD16, which is present on NK cells, showed that a significant number

FIG 3 Characterization of T cell subsets in ganglia during herpes zoster. (A to D) Immunofluorescent staining was performed with CD3, CD4, CD8, granzyme B, and S100B. Bound antibodies were visualized using species-specific Alexa Fluor antibodies, and immune cell markers are shown in red and S100B in green. Representative images are shown for HZ2. (E and F) No specific staining was seen using isotype control antibodies. Sections were counterstained with DAPI (blue). Scale bars are indicated. (G) The number of positive cells per mm² was determined from 2 independent stains and averaged (with the exception of HZ1 RL granzyme B counts, for which only 1 stain could be counted due to a lack of sample material).
of these cells were present within reactivated ganglia (data not shown).

These data indicate that during VZV reactivation there is extensive infiltration of CD4⁺ and CD8⁺ cytotoxic T cells into the affected ganglia. Infiltrating cells also appeared in neighboring ganglia, such as LL1 and RL (HZ1) and RC (HZ2), at numbers that were greater than what was seen in ganglia further from the site of reactivation and in control ganglia (Fig. 3G).

Proximity of T cells to neurons in ganglia during herpes zoster. Further examination of the localization of T cells within reactivated ganglia revealed that many T cells were in close proximity to neuron-SGC units. Counts were performed on sections stained with antibodies to CD3⁺, CD8⁺, or CD4⁺ T cells as well as S100B (marker of SGCs) to determine the proportion of neurons with T cells in very close proximity. In reactivated ganglia from HZ1 and HZ2, 23% and 12% of neuron-SGC units, respectively, had one or more CD3⁺ T cells in very close proximity. In contrast, in unaffected ganglia from these patients, only 2 to 4% of neuron-SGC units had CD3⁺ T cells in close proximity. A breakdown of the T cell subsets involved in this association revealed that despite there being a greater number of CD4⁺ than CD8⁺ T cells in both reactivated ganglia, CD8⁺ T cells appeared more likely to be associated with neuron-SGC units, as 9.9% (73/740) of neurons had a CD8⁺ T cell in close proximity, whereas 4.3% (28/651) of neurons had a CD4⁺ T cell in close proximity in HZ2 RTG.

Confocal microscopy was used to examine the relationship between neurons and T cells in closer detail. In some neuron-SGC units there was a focal absence of S100B staining, and, as a result, SGC cytoplasm, separating the CD3⁺ T cells from the neuron (Fig. 5A to C). This suggests that the protective SGC layer was breached, which could allow interactions between neurons and T cells to occur in reactivated ganglia during herpes zoster.

This observation was confirmed with dual immunofluorescence staining for NCAM (expressed on the surface of the neurons and satellite cells) and CD8. In both reactivated ganglia, for some neuron-SGC units the NCAM membrane was discontinuous and gaps could be observed when an infiltrating CD8⁺ cell appeared adjacent to a neuron (Fig. 5D to F). These data suggest that there is potential for direct interaction between neurons and CD8⁺ T cells in reactivated ganglia during herpes zoster. Furthermore, staining for granzyme B and S100B also showed that cells with cytolytic potential could be found in close proximity to some neurons (Fig. 5G to I).

Cleaved caspase-3 detection in ganglia during herpes zoster. In order to determine if the infiltrating cytotoxic CD8⁺ T cells contribute to increased neuron apoptosis, in herpes zoster, dual immunofluorescent staining for cleaved caspase-3 and granzyme B was performed. Interestingly, within all ganglia, including both reactivated ganglia, the majority of neurons and satellite glial cells did not stain for cleaved caspase-3. However, some of the nonneuronal cells, particularly those in close proximity to neurons, were dual positive for cleaved caspase-3 and granzyme B (Fig. 6). Counting of 30 nonoverlapping images taken (representing 0.486 mm²) from a single stain from the reactivated ganglia from HZ2 showed 11 small cells which were positive for cleaved caspase-3, 7 of which (64%) were also granzyme B positive (Fig. 6). Thus, the cleavage of caspase-3 does not appear to be a feature in neurons in affected ganglia during VZV reactivation, despite the abundance of cells expressing cytolytic markers.

MHC-I and -II are upregulated in human ganglia during herpes zoster. Sensory neurons typically lack expression of major histocompatibility complex (MHC) molecules; however, expression is upregulated following HSV-1 infection of mouse sensory neurons (26), which may facilitate neuron-T cell interactions. In order to determine if MHC molecules are expressed in ganglia during active VZV reactivation, immunofluorescent staining was
performed on all ganglia for β2-microglobulin (β2m) and the human leukocyte DR (HLA-DR) antigen as markers of MHC class I (MHC-I) and MHC-II, respectively. No neuronal β2m or HLA-DR expression was detected in ganglia from both HZ1 and HZ2 or in control ganglia. However, β2m expression was observed on all other cell types within the ganglia, and HLA-DR was expressed on SGCs as well as on some infiltrating inflammatory cells. Compared to control ganglia, there was marked upregulation of β2m and HLA-DR expression on SGCs in both reactivated ganglia (RTG and LL2) (Fig. 7 and 8). In HZ1, herpes zoster was confined to a single dermatome without clinical evidence of dissemination, which suggests that ganglia in proximity to the site of reactivation also undergo some degree of immune activation despite the lack of demonstrable viral antigens.

DISCUSSION

VZV reactivation (herpes zoster) is associated with pain and postherpetic neuralgia with severe pain, persisting for months to years following the resolution of the herpes zoster rash. Neither the cellular mechanisms involved in viral reactivation from latency nor the cause of postherpetic neuralgia is known. The strict species specificity of VZV, coupled with the difficulty of obtaining ganglia from patients with herpes zoster or fresh human ganglia for experimental infection, have contributed to the paucity of information about these key aspects of VZV infection. Previously, we reported the first detailed characterization of the immune response in naturally infected ganglia from 3 patients who died 1 to 5 months after contracting herpes zoster (25). However, no detailed studies of ganglia obtained during acute herpes zoster while the rash was still present have been reported; consequently, the constituents of the immune response in these ganglia have not been defined. In this study, we analyzed multiple ganglia from 2 patients (HZ1 and HZ2) with active herpes zoster at the time of death. While direct quantitative comparisons between the current study using ganglia during active herpes zoster and our previous study using ganglia after recovery from herpes zoster (25) are complicated by the different approaches used to stain and enumerate different cell types, there were several apparent differences of note. During active herpes zoster, there was a large infiltrate of cytotoxic T cells, and the predominant infiltrating cells were CD4+ T cells and, to a lesser degree, CD8+ T cells, whereas following resolution of the rash (i.e., 1 to 5 months after contracting...
herpes zoster) the predominant infiltrating cells are CD8+ T cells, yet the majority lack cytotoxic markers (25). Further, the current study demonstrates that despite an abundance of cytotoxic CD8+ T cells, some of which were in close proximity to neurons and may have breached the SGC barrier, cleaved caspase 3 was not identified, indicating that neurons and SGCs were unlikely to be undergoing apoptosis. The strong upregulation of MHC class I and II molecules on SGCs suggests that these cells play an important role in directing the immune response during herpes zoster.

Previous histological studies of ganglia from patients with herpes zoster close to the time of death noted necrosis in part or the whole of the ganglia, with some hemorrhage and lymphocytic infiltration (27–32). Some cases also showed vascular involvement with perivascular lymphocytic cuffing and thrombosis noted (28, 31). Similar features were noted in ganglia from the current study, particularly in the reactivated ganglia of HZ1, where focal necrosis was apparent. Given the case history for this patient as well as the histology, the observed necrosis was most likely an infarct caused by vasculitis rather than by vascular occlusion from an embolus or other cause. A localized hemorrhage was also noted in the reactivated ganglia from HZ2, and both reactivated ganglia contained a dense lymphoid infiltrate.

Our finding that VZV antigens were detected only in neuron-SGC units within the area of necrosis in HZ1 and not within the adjacent viable ganglia or in ganglia from HZ2 was unexpected, given that both patients had active herpes zoster with a rash at their time of death, 17 (HZ1) and 21 (HZ2) days after the onset of the rash. It is possible that following a burst of viral replication within the ganglia with transport of new virus particles to the skin resulting in a vesicular rash, VZV replication is rapidly contained within the ganglia, resulting in the loss of viral antigen staining. This is supported by the observation that herpes zoster skin lesions begin to resolve in less than a week (3), implying little reseeding of the skin from the ganglia. Also, antiviral treatment is most effective when administered within 72 h of the onset of the rash (33–36), suggesting that this is when the majority of virus replication occurs. Alternatively, low levels of viral replication may persist but are below the level of detection by immunohistochemistry, or viral replication is ongoing but is not uniform within the ganglia and was not present in the sections of the ganglia we examined. A persistent low level of virus replication may also explain our previous results where a persistent immune infiltrate was evident in ganglia several months after clinical resolution of herpes zoster (25). Ongoing viral replication was also suggested by Schmidbauer et al. (29), who detected viral antigens (gH and nucleoprotein) and VZV DNA in ganglia from 7 patients up to 7 weeks after the onset of the herpes zoster rash. Interestingly, as in our study, they only identified viral antigens in necrotic regions of the ganglia (29). We have previously shown that VZV-infected neurons expressing IE63 are resistant to apoptosis (7). Therefore, the necrosis of neuronal cells may be due to the inflammatory response to viral reactivation, which may also lead to vasculitis with or without an infarction. Alternatively, the viral antigens, which were observed in necrotic cells, may be the result of secondary viral reactivation occurring subsequent to infarction. Studies have shown that a loss of connection between a sensory ganglion and the central nervous system results in HSV reactivation (37–41); thus, it is possible that by disrupting the architecture and function of the ganglion, necrosis triggers viral reactivation.

In the reactivated ganglia in the current study, a dense T cell infiltrate was observed, although there were relatively more CD4+ than CD8+ T cells identified and most of the CD8+ T cells were cytotoxic (granzyme B positive). Given the expression of MHC-II molecules on SGCs and their phagocytic capacity (10), they may play an important role in presenting exogenous VZV antigens to CD4+ T cells (13). However, although cytotoxic CD4+ T cells...
expressing granzyme B play an important role in controlling herpesvirus infections independent of CD8$^+$ T cells and B cells (reviewed in reference 42) and VZV-specific CD4$^+$ T cells are capable of controlling VZV infection in human retinal pigment epithelial cells (43), we did not identify any CD4$^+$/granzyme B$^+$ cells. The temporal profile of the immune response to VZV reactivation is poorly understood, and it is possible that significant numbers of cytotoxic CD4$^+$ cells are present only for a short period during reactivation. We were able to study ganglia taken 17 (HZ1) and 21 (HZ2) days after the onset of the rash, which may be outside this period. The observed infiltrating CD4$^+$ T cells may still play an important role in controlling VZV replication through the secretion of cytokines or via FAS-FAS ligand interactions. Further characterization of the cytokine and chemokine profile within reactivated ganglia and the expression of immune-inhibitory molecules will form a part of future studies.

With the currently available tools it is not possible to determine what proportion of the infiltrating T cells are VZV antigen specific or whether the majority of the infiltrating cells are not VZV specific but are recruited following secretion of chemokines released in response to VZV infection. Previously we have reported CXCL10 production and the presence of CXCR3-positive cells in both human fetal DRG explants experimentally infected with VZV and in one of the reactivated ganglia used in the present study (LL2) (40). It is likely that other chemokines also play an important role in immune cell recruitment into ganglia following VZV reactivation, and their identification remains to be determined.

A number of neurons within both reactivated ganglia from HZ1 and HZ2 were shown to have at least one T cell within close proximity, which in many cases appeared to have breached the protective SGC layer, as evidenced by focal loss of S100B and NCAM staining. Many of these cells were cytotoxic (granzyme B$^+$) (Fig. 5G to I). Although neuronal expression of MHC-I and -II molecules was not detected, expression may have been below the limit of detection by immunohistochemistry. Similarly, β2m expression was not identified on neurons within ganglia several months after herpes zoster reactivation (25).

It is currently not possible to determine if an immunological synapse between the T cells and neurons is occurring within the ganglia. In vitro studies have shown that neurons are susceptible to T cell-mediated contact-dependent cytotoxicity that does not require MHC-I expression (44). Interestingly, although granzyme B-positive CD8$^+$ cells appeared in close proximity to neurons in the reactivated ganglia, no neuronal staining for cleaved caspase-3

![FIG 7 Upregulation of MHC-I and -II in human ganglia during herpes zoster. Representative images stained for β2 microglobulin (A, C, and E) and HLA-DR (B, D, and F) in the reactivated LL2 ganglion, a neighboring lumbar ganglion (LL1), and a cervical ganglion (RC) from HZ1. Bound antibodies were visualized with Alexa Fluor antibodies (green), and sections were counterstained with DAPI (blue). Corresponding images were taken with identical exposure times.](http://jvi.asm.org/)
was identified in either the reactivated or unaffected ganglia. A small number of small infiltrating cells, often granzyme B positive, were cleaved caspase-3 positive. The phenotype of these cells remains to be determined. A similar phenotype has also been observed in rabbit TG experimentally infected with another alphaherpesvirus, bovine herpesvirus type 1. Two days after infection, the infiltrating mononuclear cells surrounding many of the neurons were terminal deoxynucleotidyltransferase-mediated dUTP-biotin nick end labeling (TUNEL) positive, although only rare TUNEL-positive neurons were observed (45). While this may reflect the presence of an innate protective mechanism within sensory ganglia to prevent the irreversible loss of neurons, the presence of replicating or latent VZV may protect neurons from T cell-mediated apoptosis.

It is possible that the neurons we observed with T cells in close proximity harbored latent VZV. Unfortunately, degradation of nucleic acids due to the fixation conditions and the age of our samples did not allow us to perform in situ hybridization to detect VZV nucleic acid. Studies of trigeminal ganglia harboring latent HSV-1 have shown that antigen-specific granzyme B+ CD8+ T cells preferentially surround LAT+ neurons (13). It has also been demonstrated that granzyme B cleaves the HSV-1 immediate-early protein ICP4, which may play a role in limiting viral reactivation (14). Similarly, expression of LAT protects neuroblastoma cells from granzyme B-mediated apoptosis (15). Further studies are required to determine if similar protective mechanisms exist for VZV during productive infection and/or latency.

The expression of MHC-I and particularly MHC-II was increased on SGCs within both reactivated and neighboring ganglia. Kennedy et al. (46) examined ganglia from patients with a history of herpes zoster near the time of death and also found intense staining for MHC-II in areas with infiltrating lymphocytes. These findings suggest a role for SGCs in controlling the immune response, and combined with our results demonstrating that T cell counts were higher in ganglia close to the primary site of reactivation, they suggest that the effects of viral reactivation are not limited to a single ganglion. Further studies into the role of SGCs and bystander neurons during VZV reactivation are warranted.

**FIG 8** Upregulation of MHC-I and -II in human ganglia during herpes zoster. Representative images were stained for β2 microglobulin (A, C, and E) and HLA-DR (B, D, and F) in the reactivated RTG ganglion, a neighboring cervical ganglia (RC), and a lumbar ganglion (LL1) from HZ2. Bound antibodies were visualized with Alexa Fluor antibodies (green), and sections were counterstained with DAPI (blue). Corresponding images were taken with identical exposure times.
In summary, this study has identified VZV antigen staining using 3 independent antibodies in necrotic neurons during VZV reactivation. This is similar to other studies and shows that this may be a common feature of herpes zoster, which has implications regarding the development of PHN. Further, infiltrating CD4+ and cytotoxic CD8+ T cells were also a hallmark of the host immune response within the ganglia during herpes zoster, and some of these T cells appeared in close proximity with neurons but did not appear to be inducing apoptosis. Upregulation of MHC-I and -II molecules on SGCs suggests that these cells play a role in directing the immune response during viral reactivation. Further studies are required to elucidate the mechanisms that may lead to or prevent neuronal necrosis during herpes zoster and the role of SGCs during viral reactivation.

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