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Absence of Vasoactive Intestinal Peptide Expression in Hematopoietic Cells Enhances Th1 Polarization and Antiviral Immunity in Mice

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Vasoactive intestinal peptide (VIP) induces regulatory dendritic cells (DC) in vitro that inhibit cellular immune responses. We tested the role of physiological levels of VIP on immune responses to murine CMV (mCMV) using VIP-knockout (VIP-KO) mice and radiation chimeras engrafted with syngeneic VIP-KO hematopoietic cells. VIP-KO mice had less weight loss and better survival following mCMV infection compared with wild-type (WT) littermates. mCMV-infected VIP-KO mice had lower viral loads, faster clearance of virus, with increased numbers of IFN-γ+ NK and NKT cells, and enhanced cytolytic activity of NK cells. Adaptive antiviral cellular immunity was increased in mCMV-infected VIP-KO mice compared with WT mice, with more Th1/Tc1-polarized T cells, fewer IL-10+ T cells, and more mCMV-M45 epitope peptide MHC class I tetramer+ CD8+ T cells (tetramer+ CD8 T cells). mCMV-immune VIP-KO mice had enhanced ability to clear mCMV peptide-pulsed target cells in vivo. Enhanced antiviral immunity was also seen in WT transplant recipients engrafted with VIP-KO hematopoietic cells, indicating that VIP synthesized by neuronal cells did not suppress immune responses. Following mCMV infection there was a marked upregulation of MHC-II and CD80 costimulatory molecule expression on DC from VIP-KO mice compared with DC from WT mice, whereas programmed death-1 and programmed death ligand-1 expression were upregulated in activated CD8+ T cells and DC, respectively, in WT mice, but not in VIP-KO mice. Because the absence of VIP in immune cells increases innate and adaptive antiviral immunity by altering costimulatory and coinhibitory pathways, selective targeting of VIP signaling represents an attractive therapeutic target to enhance antiviral immunity. The Journal of Immunology, 2011, 187: 000–000.

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Abbreviations used in this article: BM, bone marrow; DC, dendritic cell; hCMV, human CMV; HSC, hematopoietic stem cells; KO, knockout; Lm-mCMV, Listeria monocytogenes-murine CMV; mCMV, murine CMV; PACAP, pituitary adenyl cyclase-activating polypeptide; PD-1, programmed death-1; PD-L1, programmed death ligand-1; VIP, vasoactive intestinal peptide; WT, wild-type.

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patients, and the pathogenesis of murine CMV (mCMV) infection in mice is similar to that in human CMV (hCMV) infection (18, 19). mCMV and hCMV exhibit 70% sequence similarity, comparable to the global level of DNA sequence homology between their natural hosts (20), and are predicted to contain ~170 and 165 open reading frames, respectively (21, 22). The large number of homologous open reading frames indicates that the two viruses are related, although immune evasive strategies of mCMV infection are quite different from those seen following hCMV infection (20), suggesting specific adaptation of a common ancestor virus to the immune environments of mice and humans (23). Furthermore, mice and humans have similar specific immune responses to their respective CMV (21, 24–26), with coordinated activities of innate and adaptive immune cells, including DC, macrophages, NK cells, T cells, and B cells (27–32). Whereas cellular and humoral immune response to mCMV is robust and effective in clearing the virus, mCMV infection also leads to immunosuppressive effects, including expression of m144, a MHC-I decoy that binds to NK cells and inhibits antiviral cytotoxicity (33, 34), and induction of a paralyzed DC phenotype, characterized by downregulation of MHC-I and MHC-II, costimulatory molecules, and proinflammatory cytokines (32). Hence, we were interested in whether interference with VIP signaling could enhance immune responses to mCMV infection. Previous studies have explored the effect of VIP on inflammation and alloimmune immunity using supraphysiological, pharmacological administration of purified VIP peptide agonist (3, 9). To study the immunomodulatory effects and antiviral immunity of physiological levels of VIP, we used VIP-KO mice (35) and VIP-KO hematopoietic chimeras (36). We hypothesized that mice lacking VIP expression would show an increased response to viral infection due to a lack of immunosuppressive counter-regulatory activity from DCs. We challenged VIP-KO mice and radiation chimeras engrafted with VIP-KO hematopoietic cells with two sources of mCMV Ag: a *Listeria monocytogenes* vaccine that expresses an immunodominant mCMV peptide (*Listeria monocytogenes*-mCMV [Lm-mCMV] vaccine) (37, 38), and an infectious strain of mCMV (37, 39). Our results demonstrate that VIP-KO mice and recipients engrafted with VIP-KO hematopoietic cells have augmented cellular immune responses to mCMV Ag, and improved survival after viral infection. The kinetics of Ag-specific primary and secondary immune responses were accelerated in VIP-KO mice and in mice reconstituted with VIP-KO hematopoietic cells, supporting the role of VIP in immune counter-regulatory pathways.

**Materials and Methods**

**Mice**

B6 strain (H-2Kb, CD45.2, CD90.2) VIP/peptide histidine isoleucine–KO mice have been previously described (35). Both male and female VIP-KO mice have been previously described (35). Both male and female VIP-KO mice were used in experiments, using syngenic siblings as wild-type (WT) controls. Congenic strains of B6 mice were purchased from The Jackson Laboratory (Bar Harbor, ME) (H-2Kb, CD45.1, CD90.2) or were bred at the Emory University Animal Care Facility (Atlanta, GA) (H-2Kb, CD45.1/CD45.2). All mice were 8–10 wk old. Procedures conforms to the Guide for the Care and Use of Laboratory Animals, and were approved by the Emory University Institutional Animal Care and Use Committee. According to Institutional Animal Care and Use Committee guidelines, any mouse that lost ≥25% body weight was euthanized and recorded as dying on the following day for statistical analysis.

**Donor cell preparation for transplantation**

BM transplantation was performed to create chimeric mice with hematopoietic cells from VIP-KO donors or WT donors (control). Femora, tibia, and spleens were obtained from VIP-KO or WT mice. BM cells were harvested by flushing the specimens with sterile RPMI 1640 containing 1% heat-inactivated FCS. T cells were purified from splenocytes by negative selection using a mixture of biotinylated non-T cell Abs (anti-CD11b, B220, DX5, and Ter119), streptavidin microbeads, and immunomagnetic separation (MACS; Miltenyi Biotec, Auburn, CA). The average purity of CD3+ T cells was 95%. Lineage (CD3, CD4, CD8, Gr-1, CD11b, I-A^b^, DX5, B220, TER119, and CD19) c-kit+ sca-1+ hematopoietic stem cells (HSC) and lineage- (CD3, DX5, IgM, TER119, and CD19) CD11c+ DC from donor BM were purified using a BD Biosciences FACS Aria cell sorter (36). Purity of FACS-purified HSC and DC averaged 93 and 97%, respectively.

**Radiation chimeras and stem cell transplantation**

On day −1, 8- to 10-wk-old male B6 CD45.1 congenic mice were irradiated with two fractions of 5.5 Gy for a total of 11 Gy (40). On day 0, irradiated mice received 5 × 10^6 T cell–depleted BM cells plus 3 × 10^5 MACS-purified splenic T cells via tail vein injection. Some experiments used an alternate approach, transplanting a combination of 5 × 10^5 HSC, 5 × 10^5 DC, plus 3 × 10^5 T Cells. Mice were monitored for signs of severe infection, including fur texture, posture, activity, skin integrity, and weight loss. Each transplant group was followed for at least 100 dpi (41). Donor cell chimerism in peripheral blood was determined 2 mo after transplantation, and was typically ≥95%. Chimeric mice were then used in vaccination and mCMV infection studies.

**Virus and immunization**

The Smith strain of mCMV passaged in vivo in salivary glands was frozen in aliquots in liquid nitrogen (37, 39), WT and VIP-KO mice, as well as chimeric mice with hematopoietic cells from WT and VIP-KO donors, were given either 5 × 10^6 (10% lethal dose; low-dose) or 1 × 10^7 (LD90; high-dose) PFU mCMV by i.p. injection and then monitored for signs of illness, including hunched posture, decreased activity, and weight loss. Mice were vaccinated i.p. with 1 × 10^5 CFU Lm-mCMV, a *L. monocytogenes* that has been rendered nonpathogenic by knockout of bacterial genes associated with virulence (42) and engineered to express the mCMV H-2^D^ restricted immunodominant peptide M45 aa 985–993 HGRNASFI (43). The vaccine was prepared and supplied by Cerus (Concord, CA, 37, 38).

**Analysis of peripheral blood and spleen samples**

Blood and spleen samples were obtained on 3, 7, 10, 14, 17, and 21 d after vaccination or following mCMV infection. Leukocytes, RBCs, and platelets were counted using a Beckman Coulter automated counter. Blood and spleen samples were depleted of RBCs by ammonium chloride lysis and washed twice. NK, NK-T, and T cell subsets were enumerated using CD3 PE/PE-Cy7/TETC, CD4 PE-Alexa610/PE-Alexa700, CD8 PE-Cy7/PerCP, CD62L, FITC/allophycocyanin, CD25 allophycocyanin-Cy7, CD4 PE-Cy5, CD60 PE-Cy7, programmed death-1 (PD-1) NE, and NK1.1 PE (BD Pharmingen, San Jose, CA). Cells were stained with mAbs specific for congenic markers CD3, CD4, CD8, CD90.1, and CD90.2 to determine donor chimerism. Allophycocyanin-labeled mCMV M45 aa 985–993 peptide-HGRNASFI-H-2^D^ tetramer was obtained from the Emory Tetramer Core Facility. All samples were analyzed on a FACS Canto (BD Biosciences, San Jose, CA), and flowcytometry analyses were performed using FlowJo software (Tree Star 2007). Samples for flow cytometric analysis of mCMV-M45 peptide epitope–MHC-I tetramer^+^ CD8^+^ T cells (tetramer^+^ CD8^+^ T cells) were gated for lymphocytes in the area of forward and side light scatter, setting a gate for tetramer^+^ T cells such that 0.1% of control (nonimmune) CD8^+^ T cells were positive (37, 39). Flow cytometric analyses of the T regulatory-associated molecule PD-1 (44), the costimulatory molecule ICOS, the cell-surface molecule CD62L (45), activation markers CD54 and CD69 (46), intracellular cytokines (IFN-γ, TNF-α, IL-4, and IL-10), and DC markers (I-A^b^, CD80, and programmed death ligand-1 [PD-L1]) were analyzed, as previously described (36).

**In vivo killing assay**

Naïve splenocytes were harvested from CD45.1^+^CD45.2^−^ heterozygous C57BL/6 mice and pulsed with 5 μM mCMV M45 aa 985–993 HGRNASFI peptide in RPMI 1640 containing 3% FBS for 90 min at 37°C, and were used three times with ice-cold media. mCMV^+^ peptide-pulsed target splenocytes and nonpulsed splenocytes from CD45.1^+^ B6 congenic mice were mixed together in equal parts, and 40 × 10^6 total target cells per mouse were injected i.v. into CD45.2^+^ VIP-KO or WT C57BL/6 mice that had been infected 9 d earlier with low-dose (10% lethal dose) mCMV, or injected into noninfected WT control mice. Sixteen hours following injection of target cells, recipients were sacrificed, splenocytes harvested, and the numbers of mCMV peptide-pulsed CD45.1^+^CD45.2^−^ and non-pulsed CD45.1^+^ target cells were quantified by FACS analysis. Immune-mediated killing of mCMV peptide-pulsed targets was calculated by first dividing the percentage of peptide-pulsed or nonpulsed targets recovered...
from the spleen of mCMV-immune mice with the mean percentage of the corresponding population of peptide-pulsed or nonpulsed targets from nonimmune mice (ratio of immune killing). The specific antiviral in vivo lytic activity for individual mice was calculated by the following formula:

\[
(1 - \frac{\text{cpm spontaneous release}}{\text{cpm maximum release} - \text{cpm spontaneous release}}) \times 100
\]

**In vitro measurements of immune responses to mCMV peptide**

WT mice, VIP-KO mice, and mice engrafted with either WT or VIP-KO donor cells were infected with low-dose mCMV, and splenocytes were harvested 3 or 7 d later. Splenic DC and T cells were purified by FACS and MACS, respectively (36). DC were plated at 2 \times 10^5 cells/ml in 12-well plates and centrifuged (300 \times g for 30 min) with 3 \mu M mCMV peptide (37). After centrifugation, DC were washed three times with PBS, resuspended in complete medium, and incubated with 2 \times 10^5 T cells at 37°C for 3 or 7 d (47). Cells were treated with Golgi Stop (BD Pharmingen) during the last 6 h of culture. Cells were then harvested from culture plates and stained with fluorescently labeled Abs against DC and T cell lineage markers (36), permeabilized, stained with Abs against IL-10 and IFN-γ, and analyzed by flow cytometry, as previously described, using isotype-matched control Abs to set the gates for distinguishing positive intracellular staining (36). Harvested culture media was stored at −20°C until use for cytokine analysis by ELISA (OpEIA ELISA sets for IL-10 and IFN-γ; BD Biosciences). ELISA plates were read using a SpectraMax 340PC spectrophotometer (Molecular Devices, Sunnyvale, CA) (36).

**NK cell lytic activity**

Lytic activity of NK cells was analyzed, as previously described (48). Briefly, YAC-1 cells, a sensitive target for NK cells, were labeled with 37 MBq Na\(^{38}\)CrO\(_4\) at 37°C for 90 min and washed twice with warm RPMI 1640 medium. The labeled target cells (1 \times 10^5) were cocultured with effector splenocytes (containing NK cells) at various ratios of effectors: targets (100:1, 50:1, and 25:1) in a final volume of 0.2 ml fresh medium in 96-well round-bottom microplates. The plates were incubated for 4 h at 37°C with 5% CO\(_2\). The amount of 51Cr released in 0.1 ml supernatant was measured by a well-type gamma counter (beta liquid scintillation counter; EG&G Wallac, PerkinElmer, Ontario, Canada). Specific cytotoxicity was calculated as follows: % \(51\text{Cr} \text{release} = 100 \times \frac{\text{cpm experimental} - \text{cpm spontaneous release}}{\text{cpm maximum release} - \text{cpm spontaneous release}}\).

**Determination of liver viral load**

Viral load was analyzed, as previously described (39). Briefly, livers were collected from CMV-infected recipients, homogenized, and centrifuged. Serially diluted supernatants were added to 3T3 confluent monolayers in 24-well tissue culture plates and incubated for 90 min at 37°C and 5% CO\(_2\), then overlayed with 1 ml 2.5% methylcellulose in DMEM, and returned to the incubator. After 4 d, the methylcellulose was removed and the 3T3 confluent monolayers were stained with methylene blue. mCMV plaques were directly counted under a light microscope (Nikon, Melville, NY). PFUs were calculated.

**Statistical analyses**

The data were analyzed using SPSS version 18 for MAC. In this study, each treatment group (or time point) had four to five mice, and every experiment was repeated at least twice. Data are presented as mean ± SD of all evaluable samples if not specified. Survival differences among groups were calculated with the Kaplan-Meier log-rank test in a pairwise fashion. Differences in tetramer response, cytokine levels, and T cell numbers were compared using a two-tailed Student t test. A p value <0.05 was considered significant.

**Results**

**VIP-KO mice were resistant to mCMV infection**

We first compared the hematological and immunological phenomenotypes of VIP-KO mice. We found no significant differences comparing blood from naive WT and VIP-KO mice in the numbers of total leukocytes, CD4, CD8, αβ TCR T cells, γδ T cells, B cells, myeloid leukocytes, and DCs in blood (Supplemental Fig. 1). VIP-KO and WT mice were infected with a nonlethal dose of mCMV (5 \times 10^5 PFU) and sacrificed 3, 10, and 17 d later. VIP-KO mice had significantly less virus in their liver, a target for mCMV infection (37, 49), with more rapid clearance of virus than mCMV-infected WT mice (\(p < 0.001\); Fig. 1). To test whether VIP-KO mice had better survival following mCMV infection, VIP-KO and WT mice were infected i.p. with either 1 \times 10^7 PFU/mouse (high-dose) or 5 \times 10^6 PFU/mouse (low-dose) mCMV. All WT mice given high-dose mCMV died by day 10 postinfection compared with 65% survival of the VIP-KO mice (\(p < 0.001\); Fig. 2A). Following low-dose mCMV infection, both WT and VIP-KO mice had transient lethargy and weight loss, with recovery to baseline values by day 20 postinfection, with 100% of WT mice and 92% of VIP-KO mice surviving to day 100 postinfection (Fig. 2A, 2B). In a parallel experiment, serial measurements of CD4 and CD8 T cells following mCMV infection showed that VIP-KO mice had more CD4\(^+\) and CD8\(^+\) T cells in their blood and spleen compared with WT mice (Fig. 2C–F).

**Innate and adaptive antiviral responses were enhanced in the absence of VIP**

VIP-KO mice had significantly higher percentages (Fig. 3A) and absolute numbers of Ag-specific tetramer\(^*\) CD8 T cells in the blood (Fig. 3B) and spleen (Fig. 3C) following low-dose mCMV infection than WT mice. The highest frequency of tetramer\(^*\) CD8 T cells in the blood was seen on day +10 postinfection with 9.1 ± 0.8% of blood CD8\(^+\) T cells in VIP-KO mice versus 4.8 ± 0.7% of blood CD8\(^+\) T cells in WT mice (\(p < 0.001\); Fig. 3A). Because lethality was 100% in WT mice receiving high-dose mCMV compared with 35% mortality among VIP-KO mice (\(p < 0.001\)), a longitudinal comparison of the numbers of Ag-specific T cells in WT versus VIP-KO mice could not be performed, but analysis at day 3 showed that VIP-KO mice had greater numbers of tetramer\(^*\) CD8 T cells (295/ml ± 40/ml) compared with WT mice (124/ml ± 38/ml, \(p < 0.001\)). Enhanced innate antiviral immunity among VIP-KO mice was evidenced by higher levels of NK-mediated cytotoxicity against YAC1 targets in VIP-KO splenocytes harvested 3 d postinfection (Fig. 3D). Using mCMV peptide-pulsed and nonpulsed congenic splenocytes as targets in an in vivo cytotoxicity assay in immune mice (previously infected with low-dose mCMV), the specific lysis of mCMV peptide-pulsed targets was significantly enhanced in VIP-KO mice compared with WT mice (Fig. 4). Significantly, VIP-KO mice had similar baseline numbers, but more IFN-γ–expressing NK, NKT cells, and Th1/...
Tc1-polarized (IFN-γ⁺ and TNF-α⁺) T cells on days 3–17 post-infection compared with WT mice (Supplemental Fig. 2A–H).

The absence of VIP expression in donor hematopoietic cells enhanced antiviral immunity in radiation hematopoietic chimeras

Because VIP is expressed in multiple cell lineages (2–6), we tested whether mice lacking VIP expression only in their hematopoietic cells had the same level of enhanced antiviral immunity as we observed in VIP-KO mice. We used VIP-KO mice as donors of hematopoietic cells and created radiation chimeras with syngenic BM transplantation in which recipients had 95% donor cell engraftment (36). The day 59 survival of mice transplanted with VIP-KO 3 × 10⁶ FACS-purified HSC, 5 × 10⁵ FACS-purified DC, and 3 × 10⁶ MACS-purified T cells (75 ± 10%) was similar to the survival seen among mice transplanted with WT HSC, DC, and T cells (80 ± 9%). To explore the effect of VIP expression in hematopoietic cells on primary and secondary immune responses, VIP-KO→WT and WT→WT syngeneic transplant recipients were primed with PBS or the Lm-mCMV vaccine (containing mCMV immunodominant M45 epitope peptide aa 985–993), followed by infection 21 d later with low-dose mCMV (Fig. 5). Peripheral blood samples obtained prior to Lm-mCMV vaccination (day 59 posttransplant), after vaccination, and following mCMV infection (day 80 posttransplant) were analyzed for the numbers of tetramer⁺ CD8 T cells. Nonimmunized WT and VIP-KO chimeric mice had minimal numbers of mCMV-peptide tetramer⁺ CD8 T cells in their blood at baseline (Fig. 5A). Following primary mCMV infection, recipients engrafted with VIP-KO hematopoietic cells had significantly more mCMV peptide tetramer⁺ CD8 T cells in their blood compared with WT mice (Fig. 5A). Vaccination with Lm-mCMV led to a larger increase in blood mCMV tetramer⁺ T cells in the VIP-KO→WT chimeras compared with WT→WT chimeras (Fig. 5B), indicating that mCMV peptide pre-

**FIGURE 2.** Mice lacking VIP had better survival and greater expansion of blood T cells following mCMV infection. VIP-KO and WT mice were infected (day 0) with low-dose 5 × 10⁴ PFU or high-dose 1 × 10⁵ PFU mCMV. Survival was recorded every day, and body weight was recorded twice weekly. Peripheral blood and spleen were collected baseline, prior to mCMV infection, and 3, 7, 10, 14, and 17 d postinfection. Blood cells and splenocytes were stained with fluorescently conjugated mAbs to CD45.2, CD3, CD4, and CD8, and analyzed by flow cytometry, and absolute numbers of cells/ml blood or per spleen were calculated. Data from 12–15 mice per group were pooled from 3 replicate experiments. A and B, Survival and body weight change of WT and VIP-KO mice that received graded doses of 5 × 10⁴, or 1 × 10⁵ PFU mCMV. C and D, Total numbers of CD4⁺ and CD8⁺ T cells in blood following low-dose mCMV infection. E and F, Total numbers of CD4⁺ and CD8⁺ T cells in the spleen following low-dose mCMV infection. *p < 0.05, **p < 0.01, ***p < 0.001, denoting a significant difference between VIP-KO and WT mice.
sentation alone in VIP-KO mice (in the absence of viral infection) was sufficient to result in enhanced expansion of Ag-specific T cells. Subsequent infection of the Lm-mCMV–vaccinated mice with low-dose mCMV led to an accelerated anamnestic response in VIP-KO → WT chimeras compared with mice engrafted with WT BM (Fig. 5B). Because both T cells and accessory cells can secrete VIP (4–6), we further explored the role of VIP synthesis by different immune cell subsets by creating radiation chimeras engrafted with the combination of donor DC and HSC from VIP-KO mice and donor T cells from WT mice. Mice transplanted with the heterogeneous combination of VIP-KO HSC and DC and WT T cells did not show the enhanced immune responses seen in mice engrafted with the homogeneous combination of VIP-KO HSC, DC, and T cells (Fig. 5B), indicating that VIP production by donor T cells was sufficient to attenuate antiviral cellular immunity.

Absence of VIP augmented antiviral CD8+ T cell proliferation and Th1/Tc1 polarization in vitro

To study the effect of VIP on antiviral immunity in vitro, we analyzed cultures of T cells and mCMV peptide-pulsed DC for tetramer+ CD8 T cells and for Th1 and Th2 cytokines. DC and T cells were purified from WT or VIP-KO mice (36), and the DC were pulsed with mCMV peptide, and then mixed with T cells. The numbers of tetramer+ CD8 T cells generated over 10 d of culture were measured by flow cytometry. Significantly greater numbers of Ag-specific tetramer+ CD8 T cells were detected after 3 d in cultures of T cells with DC that had been isolated from mCMV-immune VIP-KO mice compared with similar cells isolated from mCMV-immune WT mice (Fig. 6A). To rule out an effect of VIP synthesized by nonhematopoietic cells on in vitro immune responses to mCMV peptides, donor-derived T cells and DC were recovered from syngeneic transplant recipients of VIP-KO → WT or WT → WT radiation chimeras. Homogeneous cultures of DC and T cells recovered from VIP-KO → WT radiation chimeras generated more tetramer+ CD8 T cells than cultures of DC and T cells from WT → WT radiation chimeras. Homogeneous cultures of DC and T cells recovered from VIP-KO → WT radiation chimera generated more tetramer+ CD8 T cells than cultures of DC and T cells from WT → WT radiation chimera generated more tetramer+ CD8 T cells than cultures of DC and T cells from WT → WT radiation chimeras (Fig. 6D), indicating the absence of VIP synthesis by hematopoietic cells in radiation chimeras programmed T cells and DC toward enhanced cellular immune responses. Supernatants from cultures of T cells and mCMV peptide-pulsed DC from WT mice had higher levels of IL-10, and lower levels of IFN-γ compared with supernatants from cultures of T cells and mCMV peptide-pulsed DC from VIP-KO mice (Fig. 6D). To determine whether synthesis of VIP by T cells was sufficient to downregulate immune responses to mCMV, we cultured WT T cells and VIP-KO DC isolated from radiation chimeras originally transplanted with the heterogeneous combination of WT T cells plus VIP-KO DC and VIP-KO HSC. In contrast to the larger numbers of tetramer+ CD8 T cells seen in homogeneous cultures of T cells and DC from VIP-KO mice, heterogeneous cultures of WT

FIGURE 3. Mice lacking VIP had larger increases of Ag-specific T cells following mCMV infection. VIP-KO and WT mice were infected (day 0) with low-dose 5 × 10^4 PFU or high-dose 1 × 10^5 PFU mCMV. Peripheral blood and spleen were collected at baseline, prior to infection and 3, 7, 10, 14, and 17 d post-mCMV infection. Blood cells and splenocytes were stained with fluorescently conjugated mAbs to CD45.2, CD3, CD4, CD8, and mCMV M45-epitope peptide-specific MHC-I tetramer reagents and analyzed by flow cytometry, and absolute numbers of cells/ml blood and per spleen were calculated. NK cell-killing activity was measured by 51Cr releasing assay using YAC-1–pulsed 51Cr. A, Percentages of tetramer+ CD8 T cells in blood following low-dose mCMV infection. Dot graphs showed concatenated list mode files from analysis of four mice per group 10 d following mCMV infection (mean ± SD), and are representative of five replicate experiments. B, Absolute numbers of tetramer+ CD8 T cells/ml in blood following low-dose mCMV infection (data are from 5 replicate experiments involving a total of 12–20 mice per time point). C, Absolute numbers of tetramer+ CD8 T cells in the spleen following low-dose mCMV infection (data are from 5 replicate experiments involving a total of 12–20 mice per time point). D, NK cells mediated cytolytic activity (data are from 3 replicate experiments with 12 mice per time point). **p < 0.01, ***p < 0.001, denoting a significant difference between VIP-KO and WT mice.
In this study, we explored the immunoregulatory effect of VIP in immune responses to mCMV infection, hypothesizing that the absence of VIP would increase innate and adaptive immune responses to viral infection. Our data using VIP-KO mice demonstrate that the absence of physiological levels of VIP in hematopoietic cells led to striking enhancement of innate and adaptive antiviral cellular immune responses. VIP-KO mice had less mortality and faster viral clearance compared with WT mice. The increased expansion of tetramer+ CD8 T cells and increased cytolytic activity of NK cells seen in VIP-KO mice are most likely responsible for their greater resistance to mCMV infection (50). Whereas we used the M45 epitope peptide to measure mCMV-specific T cells, and T cells recognizing this epitope have been shown to be relatively ineffective in clearing virus-infected cells due to m152/gp40-mediated immune interference (51), the enhanced killing of M45 epitope-containing peptide-pulsed target cells supports the contribution of M45-reactive T cells to functional antiviral cytotoxic activity in vivo.

To clarify the effect of various physiological sources of VIP (hematopoietic versus neuronal), we used C57BL/6 radiation chimeras engrafted with syngeneic VIP-KO or WT hematopoietic cells following myeloablative radiation. Recipients of VIP-KO hematopoietic grafts showed accelerated kinetics of cellular immune responses to primary mCMV infection and Lm-mCMV vaccination, as well as greater amnestic responses following Lm-mCMV vaccination and mCMV infection compared with recipients of WT grafts. These data indicate that VIP produced by hematopoietic cells has a dominant-negative effect on antiviral cellular immune responses, and that VIP synthesis by non-hematopoietic neuronal cells does not significantly affect antiviral immune responses in this system.
Immune cells in VIP-KO mice had more Th1 polarization (52, 53), less Th2 polarization, and higher MHC-II expression (47) than those of WT mice following mCMV infection, consistent with the reports that VIP is a negative regulator of Th1 immune responses (3, 54). A simple in vitro model of T cells cocultured with mCMV peptide-pulsed DC recapitulated the in vivo immunology of VIP-KO mice. Cocultures of DC and T cells from VIP-KO mice had higher levels of IFN-γ+ CD4+ and CD8+ T cells and more Ag-specific antiviral CD8+ T cells compared with cultures of WT DC and WT T cells. Conditioned media from cultures of WT

![Figure 6](http://www.jimmunol.org/)  
**FIGURE 6.** The generation of Ag-specific antiviral T cells and Th1 polarization was increased in cultures of DC and T cells from VIP-KO mice compared with cells from WT mice. DC and T cells were isolated from spleens of VIP-KO and WT mice, and from radiation chimeric mice that received homogeneous grafts from VIP-KO or WT (3 × 10^3) HSC, (5 × 10^4) DC and (1 × 10^6) T cells, and heterogeneous grafts from the combination of VIP-KO HSC and DC and WT T cells 15 d following infection with 5 × 10^4 PFU (low-dose) mCMV (15–18 per group were studied). FACS-purified DC from these mice were incubated with 5 μM mCMV peptide for 30 min, washed, and then triple-well cocultured with T cells from the same groups. On days 3 and 7 of culture, Ag-specific T cells were measured by FACS using mCMV-M45 epitope peptide–MHC-I tetramer reagent. A and B, The absolute numbers of tetramer+ CD8 T cells/ml in cultures of cells from nontransplanted (A) and radiation chimeric mice (B). Day 0 data were obtained using cells from noninfected mice. Culture media from day 3 cultures of cells from radiation chimeric mice were assayed for IL-10 (C) and IFN-γ (D) by ELISA. *p < 0.05, ***p < 0.001, comparing VIP-KO mice and WT groups. Means ± SE from pooled results of three repeat experiments. The experiment was repeated three times.

![Figure 7](http://www.jimmunol.org/)  
**FIGURE 7.** Higher levels of CD80 and MHC-II expression on DC and lower levels of PD-1 and PD-L1 expression on CD8+ T cells and DC, respectively, from VIP-KO mice following mCMV infection. Splenocytes were isolated from VIP-KO and WT mice at baseline and 3, 10, and 17 d postinfection with 5 × 10^4 PFU mCMV. Expression patterns of CD80 (A), MHC-II (B), and PD-L1 (C) on conventional DC (cDC, lineage^−^, CD11c^high, B220^−^) and plasmacytoid DC (pDC, lineage^−^, CD11c^dim, B220^+) and the percentages of CD8+ T cells expressing PD-1 (D) were analyzed by flow cytometry. Histograms depict analysis of concatenated list mode files from four mice per group at each time point, and are representative of three replicate experiments. Dashed lines represent the staining profile using an isotype-matched control Ab; filled lines represent specific staining.
T cells and WT DC had higher levels of IL-10, and lower levels of IFN-γ, compared with culture media from VIP-KO T cells and VIP-KO DC, consistent with other reports (55). Heterogeneous cocultures of VIP-KO DC and WT T cells had the same (lower) numbers of Ag-specific antiviral CD8 + T cells as cultures of WT DC and WT T cells, confirming that T cells making VIP are sufficient to polarize Th2 immunity and suppress Th1 immunity, and that VIP made by T cells is a dominant-negative regulator of antiviral immune responses (56, 57).

The mechanisms for the enhanced antiviral cellular immunity and greater Th1/Th1c1 immune polarization seen in VIP-KO mice following mCMV infection appear to be due to a profound shift in the pattern of costimulatory and co-inhibitory molecule expression on DC and CD8 + T cells. The higher levels of MHC-II and CD80 and greater Th1/Tc1 immune polarization seen in VIP-KO mice and that VIP made by T cells is a dominant-negative regulator of DC and WT T cells, confirming that T cells making VIP are protective Th1 immune responses following vaccination or infection, and are distinct from the role that PD-1 signaling plays in T cell longevity, immunosuppression (62) due to cell cycle–dependent apoptosis (63). We found that DC from mice transplanted with VIP-KO–producing cells had dramatically reduced PD-L1 expression on DC and PD-1 expression on activated memory CD8 + T cells that were associated with increased quantitative and qualitative antiviral T cell responses following mCMV infection. Our studies indicate that physiological levels of VIP contribute to the upregulation of PD-L1/PD-1 interaction seen in WT mice following mCMV infection. This work further suggests that induction of VIP may be part of the active suppression of adaptive immune responses that occurs following mCMV infection.

In summary, these data indicate that VIP synthesis by hematopoietic cells is a key factor in regulating the development of protective Th1 immune responses following vaccination or infection with mCMV. The absence of VIP synthesis by hematopoietic cells leads to lower levels of counter-regulatory co-inhibitory molecules and changes in serum cytokines consistent with global Th1 immune polarization. The increased antiviral immunity seen in the absence of VIP suggests that VIP antagonists may be of clinical benefit for patients with viral infection.

Disclosures
The authors have no financial conflicts of interest.

References


Supplemental Figure 1

Mice lacking VIP had equivalent numbers of immune cell populations in the blood compared with WT mice. Peripheral blood was collected from naïve VIP-KO and WT mice and stained with fluorescently conjugated monoclonal antibodies to CD45.2, CD3, CD4, CD8, CD11b, B220, TCRαβ, TCRγδ, B220, CD11c, and a lineage cocktail of CD3, DX5, IgM, CD19 and TER119, and analyzed by flow cytometry. Numbers of total leukocytes, or the immune cell subsets shown in the legend, were calculated per μL blood. Open bar represents VIP-KO mice (n=18), Filled bar represents WT mice (n=18). Data are pooled results of 3 replicate experiments.
Supplemental Figure 2

Mice lacking VIP had Th1/Tc1 immune polarization following mCMV infection. VIP-KO and WT mice were infected (day 0) with $5 \times 10^4$ PFU mCMV. Splenocytes were collected at baseline prior to infection and 3, 10, and 17 days post-infection, cultured with PMA plus ionomycin and Golgistop for 6 hrs, and then stained with fluorescently conjugated monoclonal antibodies to CD45.2, CD3, CD4, CD8, NK1.1, IFN-$\gamma$ and TNF-$\alpha$, and analyzed by flow cytometry. Numbers (per $\mu$L of blood) of IFN-$\gamma$+ T-cells (A-D), TNF-$\alpha$+ T-cells (E-H), NK cells (A, E), NKT-cells (B, F), CD4+ T-cells (C, G) and CD8+ T-cells (D, H) following low-dose mCMV infection are shown. * Signifies $p<0.05$, ** signifies $p<0.01$, and *** signifies $p<0.001$, denoting a significant difference between VIP-KO and WT mice. Data pooled from 3 replicate experiments; 12-16 mice were studied at each time-point.