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Premature Terminal Exhaustion of Friend Virus-Specific Effector CD8+ T Cells by Rapid Induction of Multiple Inhibitory Receptors

Shiki Takamura,* Sachio Tsuji-Kawahara,* Hideo Yagita,† Hisaya Akiba,‡
Mayumi Sakamoto,* Tomomi Chikaishi,*,† Maiko Kato,*‡ and Masaaki Miyazawa*

During chronic viral infection, persistent exposure to viral Ags leads to the overexpression of multiple inhibitory cell-surface receptors that cause CD8+ T cell exhaustion. The severity of exhaustion correlates directly with the level of infection and the number and intensity of inhibitory receptors expressed, and it correlates inversely with the ability to respond to the blockade of inhibitory pathways. Friend virus (FV) is a murine retrovirus complex that induces acute high-level viremia, followed by persistent infection and leukemia development, when inoculated into immunocompetent adult mice. In this article, we provide conclusive evidence that FV infection results in the generation of virus-specific effector CD8+ T cells that are terminally exhausted. Acute FV-induced disease is characterized by a rapid increase in the number of virus-infected erythroblasts, leading to massive splenomegaly. Most of the expanded erythroblasts strongly express programmed death ligand-1 and MHC class I, thereby creating a highly tolerogenic environment. Consequently, FV-specific effector CD8+ T cells uniformly express multiple inhibitory receptors, such as programmed cell death 1 (PD-1), T cell Ig domain and mucin domain 3 (Tim-3), lymphocyte activation gene-3, and CTLA-4, rapidly become nonresponsive to restimulation and are no longer reinvigorated by combined in vivo blockade of PD-1 and Tim-3 during the memory phase. However, combined blockade of PD-1 and Tim-3 during the priming/differentiation phase rescued FV-specific CD8+ T cells from becoming terminally exhausted, resulting in improved CD8+ T cell functionality and virus control. These results highlight FV’s unique ability to evade virus-specific CD8+ T cell responses and the importance of an early prophylactic approach for preventing terminal exhaustion of CD8+ T cells. The Journal of Immunology, 2010, 184: 4696–4707.
postinfection (15, 20). This rapid and vigorous expansion of erythroid progenitor cells accelerates the viral replication that leads to the high-level viremia and provides a large pool of additional target cells for repeated integration of F-MuLV and SFFV proviruses. Ultimately, FV-infected mice develop erythrolymphocytosis through promoter insertion or the silencing of a tumor suppressor gene (21) or, if able to recover from acute FV-induced disease, develop a life-long chronic infection (22). Although it is well known that the control of Moloney murine leukemia virus infection largely depends on CD8+ T cells (23), the actual contribution of virus-specific CD8+ T cell responses to the control of FV infection is still controversial and inconclusive. Previous studies reported that although virus-specific CD8+ T cells acquired effector functions at an early stage of infection, these cells displayed a profound functional impairment during the chronic phase (24–26). This virus-specific memory CD8+ T cell dysfunction in the chronic phase has been attributed to suppression by virus-induced regulatory T (Treg) cells, because FV infection results in more Treg cells during the chronic phase (24). Support for these ideas has mainly come through an adoptive transfer approach by which TCR-transgenic CD8+ T cells transferred into acutely infected mice developed proper effector function, whereas cells transferred into chronically infected animals showed weaker cytokine production (24). Another piece of evidence for Treg cell-mediated CD8+ T cell dysfunction in chronically infected mice was that endogenous CD8+ T cells in acutely infected mice produced significant amounts of granymes, whereas the production of granymes by CD8+ T cells during the chronic phase of FV infection was much weaker (25, 26). Because these studies did not take into consideration the potentially differential priming conditions (e.g., Ag doses and inflammatory stimuli between the acute and memory phases of FV infection and the stages of CD8+ T cell differentiation from effector to memory), it is unclear whether the dysfunction of virus-specific CD8+ T cells is due solely to the presence of virus-induced Treg cells, as is when and how FV-specific CD8+ T cells suffer functional defects throughout the entire course of FV infection. Furthermore, it has been confirmed that the FV stocks used in the above-reported studies were contaminated with lactate dehydrogenase-elevating virus (LDV), which has been shown to rapidly induce polyclonal T and NK cell activation and enhance Treg cell activities, resulting in delayed and suppressed virus-specific CD8+ T cell responses (27). Taken together, understanding the precise mechanisms and the time course of virus-specific CD8+ T cell dysfunction using LDV-free FV complex is required to fully elucidate the immune evasion strategy of FV.

In this study, we provide evidence that FV infection leads to the global unresponsiveness of virus-specific CD8+ T cells, even during the early stages of infection. Owing to the massive erythroid cell proliferation caused by SFFV, the dysfunctional virus-specific effector CD8+ T cells uniformly express multiple inhibitory molecules and are poorly responsive to the combined blockade of PD-1 and T cell Ig domain and mucin domain 3 (Tim-3), indicating that combined blockade of PD-1 and Tim-3 reduces the frequency of FV-specific CD8+ T cells targeted by FV gp70+ or CD107a+ cells among CD8+ T cells by ∼68% for FV-infected mice and 95% for FBL3-infected mice.

Materials and Methods

Viruses, mice, cells, and infection

A stock of LDV-free B-tropic FV complex was kindly provided by Kim Hasenkrug, National Institutes of Health, National Institute of Allergy and Infectious Diseases, Rocky Mountain Laboratories (Hamilton, MT). FV was expanded, stored, and titrated, as previously described (28). An infectious molecular clone of F-MuLV, FB29, was prepared from culture supernatant of chronically infected Mus dunni cells, as previously described (28). FBL3 is an F-MuLV-induced leukemia of C57BL/6 (B6) origin that expresses FV-related Ags and is known to be highly immunogenic; injection of 107 cells in syngeneic B6 mice results in rejection within 4 wk (29). C57BL/6CrSlc mice were purchased from Japan SLC (Hamamatsu, Japan). A/WySn mice were purchased from The Jackson Laboratory (Bar Harbor, ME). Animals were housed and bred in the experimental animal facilities at Kinki University School of Medicine under specific pathogen-free conditions. Male and female (B6 × A/WySn)F1 mice, 6 to 10 wk old, were infected i.v. with 1000 spleen focus-forming units of FV. Mice were infected s.c. with 5 × 106 FBL3. All animal studies were approved by Kinki University.

Tissue harvest and flow cytometry

Mice were sacrificed at the indicated time points, and single-cell suspensions of splenocytes were obtained by releasing through nylon mesh, depleted of erythrocytes in buffered ammonium chloride, and panned on goat anti-mouse IgG (H+L) (Kirkegaard & Perry Laboratories, Gaithersburg, MD) for pentamer staining. As a control, mice were injected s.c. with lactate dehydrogenase-elevating virus (LDV), which has been shown to be recognized by CD8+ T cells specific for gag75–83/Db (25). Abs to the following molecules were purchased from BD Biosciences (CD3, CD4, CD8, Ter119, and CD11b), BioLegend (CD43 [1B11], CD25, PD-1, CD69, CD26L, CD122, CD127, Tim-3, Lag-3, CTLA-4, Flu-L, TrAIL, PD-L1, PD-L2, CD11c, and FoxP3), or Invitrogen (granzyme B). Purified F-MuLV gp70 mAb, clone 720 (19), was labeled with biotin by NHS-L2, CD11c, and FoxP3), or Invitrogen (CTLA-4, FL-2, and CD11b), BioLegend (CD43 [1B11], CD25, PD-1, CD69, CD26L, CD122, CD127, Tim-3, Lag-3, CTLA-4, Flu-L, TRAIL, PD-L1, PD-L2, CD11c, and FoxP3), or Invitrogen (Carlsbad, CA; granzyme B). Purified anti–F-MuLV gp70 mAb, clone 720 (19), was labeled with biotin by NHS-PeG2-Biotin (Thermo Scientific, Waltham, MA), as described (31). Labeled pentamer was purchased from ProImmune (Oxford, UK). All three cys. The pentamer used was purchased from ProImmune (Oxford, UK). All three cysteine residues in the epitope peptide were replaced with amino- butyric acid to prevent interpeptide disulfide bonding (25). This variant peptide was shown to be recognized by CD8+ T cells specific for gag75–83/Db (25).

In vitro restimulation and intracellular cytokine staining

Splenocytes isolated from infected mice, as described above, were incubated in a 96-well plate at a concentration of 1 × 106 cells per well. Cells were incubated in the presence of Alexa488-conjugated anti-CD107a, monensin A (BioLegend), and either gag75–83 peptide (5 μM), FBL3 (1 × 106 cells/well), or anti-CD3 (4 μg/ml) (eBioscience, San Diego, CA), anti-CD25 (4 μg/ml) (eBioscience, San Diego, CA), or anti-PD-1 (4 μg/ml) (eBioscience, San Diego, CA) for 15 min on ice to prevent test Abs from binding to FcRs and then stained with an allopolycoykinin-labeled pentamer specific for an immunodominant F-MuLV gag-encoded MHC class I epitope CLCLCLTVFL (gag75–83/Db) for 1 h at room temperature. Pentamer-labeled cells were then washed and stained with FITC-, PE-, PerCP/Cy5.5-, or APC-conjugated Abs for 30 min on ice. In some instances, cells were then fixed and permeabilized with the Cytofix Cytoperm kit (BD Biosciences) or FoxP3 Fix/Perm Buffer Set (BioLegend, San Diego, CA) to detect intracellular granzyme B (without in vitro restimulation) or FoxP3, respectively. The pentamer used was purchased from ProImmune (Oxford, UK), as described (31). Labeled staining was purchased from BioLegend. Samples were run on a FACSCalibur flow cytometer (BD Biosciences). All data were analyzed with FlowJo software (Tree Star, Ashland, OR).

In vitro suppression assay

In vitro suppression assay

In vitro suppression assay was performed as described in a previous report, with minor alterations (32). CD4+CD25+ cells were sorted from the splenocytes of FV-infected or FBL3-infected mice at day 14. The frequency of FoxP3+ cells among sorted cells in each group was ∼68% for FV-infected mice and 95% for FBL3-infected mice. As a target, CD44hiCD8+ splenocytes from naive mice were sorted and then labeled with 5 μM MCFSE (In Vitrogen). Cells were then combined
such that the numbers of FoxP3+ and CD8+ cells were equal and stimulated with anti-CD3 Ab (8 μg/ml). This ratio of Treg cells to CD8+ target cells (1:1) is similar to that of total Treg cells to CD8+ T cells specific for the single dominant FV epitope (gag75–83) at day 14 in FV-infected mice (data not shown). For IFN-γ staining, brefeldin A (10 μg/ml) was added for the last 4 h of culture. Six hours after stimulation, cells were stained for CD8, fixed, permeabilized, and stained for intracellular IFN-γ, as described above. Separate cultures of stimulated cells in the presence or absence of Treg cells were rested for 48 h to assess the CFSE dilution. Similar suppression between each group was also observed when the suppressor/target ratio was adjusted based on the number of CD4+CD25+ instead of FoxP3+ cells (data not shown).

In vivo Ab blockade

Anti–PD-1 (250 μg; RMPI-14) (33), anti–Tim-3 (250 μg; RMT3-23) (34), anti–IL-10 (500 μg; JES5-2A5) (35), or control rat IgG (500 μg; Sigma-Aldrich, St Louis, MO) was administered i.p. every 3 d for 2 wk. The above doses of the blocking Abs used are sufficient to inhibit the relevant receptor–ligand interaction in vivo (36–38). RPMI-14 and RMT3-23 do not deplete PD-1+ or Tim-3–expressing cells in vivo, and they have no agonistic activity (33, 34). For dual blockade of PD-1 and Tim-3, 250 μg each Ab was used.

Ligand-binding assay

Single-cell suspensions of splenocytes were obtained from Ab-treated mice, as described above. Blockade of PD-L1 interaction with PD-1 on the surface of CD8+ T cells was measured by the ex vivo binding of the chimeric mouse B7-H1/Fc protein (R&D Systems, Minneapolis, MN). In brief, after FcR blocking, cells were incubated with the chimeric protein (1 μg/10^6 cells) for 1 h on ice and stained with anti-human IgG (Valeant Pharmaceuticals, Costa Mesa, CA), anti-CD8, and anti–PD-1 (RMP1-30), which is noncompetitive with RMPI-14. Because the Tim-3 ligand, galecitin-9, can bind CD44, blockade of cell-surface Tim-3 was determined by the binding of fluorescent-conjugated anti–Tim-3 mAb (RMT3-23). Tim-3 expression on RMT3-23–treated cells was detected by using a different clone of anti–Tim-3 mAb (RMT3-8), which is noncompetitive with RMT3-23.

Infectious-center assays

Splenic cell suspensions were serially diluted and plated in duplicate at concentrations between 1 × 10^5 and 1 × 10^6 cells onto monolayers of M. dunni cells. After being washed and fixed with methanol on the second day of coculturing, cells were stained with biotinylated 720 mAb, and F-MuLV–infected foci were visualized using an Elite ABC Kit (Vector Laboratories, Burlingame, CA), as described (31).

Statistical analysis

One- and two-way ANOVA with Bonferroni or Dunnett posttests, as indicated, were used to determine statistically significant differences between groups of repeated measurements. For the comparison of groups or repeated measurements with different sample numbers, the Student or Welch t test was used, depending on whether variances were regarded as equal or not, and Bonferroni correction was used to determine α values based on the number of independent hypotheses. Prism software (GraphPad, San Diego, CA) was used to calculate p values.

**FIGURE 1.** FV infection results in the generation of excessively activated CD8+ T cells. Mice were infected with FV (1000 spleen focus-forming units) or injected with FBL3 (5 × 10^5 cells). Splenocytes were isolated at day 14 (B, E) or the indicated time points (A, C, D) and stained with Abs and/or FV gag75–83/Dβ pentamer. A. Actual numbers of CD8+pentamer+ cells in the spleen. B. Intracellular expression of granzyme B determined without in vitro restimulation. Shown are the representative staining patterns for CD43 and granzyme B of CD8+pentamer+ or CD8+pentamer− cells at day 14. The numbers indicate the percentage of cells gated in each quadrant. C. Percentages of granzyme B+ cells among splenic CD8+ T cells. D. FV-producing infectious centers were enumerated by coculturing the splenocytes with M. dunni cells. Data are from the same experiment as in C. E. A representative staining pattern for gp70 and granzyme B of CD8+ T cells. Numbers indicate the percentage in each quadrant. All data are representative of at least three independent experiments with essentially the same results.

**Results**

**FV infection results in the generation of excessively activated CD8+ T cells**

Previous studies demonstrated that although FV infection elicits vigorous Ag-specific CD8+ T cell responses, they are insufficient to totally eradicate FV-producing cells, and the virus persists after regression of acute splenomegaly even in the resistant strains, suggesting that FV-specific CD8+ T cell responses are compromised during an early stage of infection (25). It has also become clear that FV stocks used in the previous studies were contaminated with LDV, which compromises CD8+ T cell responses (27). To thoroughly investigate the potential factors that influence the onset of virus-specific CD8+ T cell responses following FV infection in the absence of LDV, we compared the Ag-specific CD8+ T cell responses elicited by FV infection or injection of FBL3 tumor cells, a highly immunogenic murine leukemia cell line induced in a B6 mouse by F-MuLV (39). As shown in Fig. 1A, FV infection elicited a robust virus-specific CD8+ T cell response in the spleen at 14 d postinfection, with >5% of the CD8+ T cell population stained brightly with the gag75–83/Dβ pentamer. As a control, an s.c. injection of FBL3 elicited comparable numbers of gag75–83/Dβ-specific CD8+ T cells at day 14 (Fig. 1A). However, despite the elicitation of comparable levels of acute CD8+ T cell responses, there was a notable difference in the effector phenotypes between Ag-specific CD8+ T cell responses elicited by FV infection and those elicited by FBL3 tumor injection. Although only less than one quarter of Ag-specific CD8+ T cells generated by FBL3 injection displayed a highly activated effector phenotype, as determined by CD43 expression and intracellularly stored granzyme B, Ag-specific CD8+ T cells generated by FV infection were markedly skewed toward the CD43+/granzyme B+ population; detection of this without in vitro restimulation (Fig. 1B) indicates that this was excessive. Substantial accumulation of granzyme B was also observed in the CD43+ population of gag75–83/Dβ-negative CD8+ T cells in FV-infected mice, indicating that excessive activation is not a unique feature of gag75–83/Dβ-specific CD8+ T cells. The frequency of CD8+ T cells that contained granzyme B reached up to 48% of the total CD8+ T cell population at 2 wk postinfection, and it declined after 3 wk postinfection (Fig. 1C). Importantly, this decrease in granzyme B expression was apparently associated with the course of virus elimination from the spleen (Fig. 1D). Because the expression of cytolytic molecules detected intracellularly without in vitro restimulation depends mainly on the recent Ag exposure in vivo (40), the reduction in granzyme B+ cells through the late stage of infection is likely due to the combined consequences of the contraction of Ag-experienced effector
CD8+ T cells, the transition of CD8+ T cells from effector to memory stages, and the resolution of FV infection in vivo. A relatively low number of CD8+ T cells was positive for the virally encoded envelope protein (gp70), even at the peak of FV infection (Fig. 1E), indicating that direct infection of CD8+ T cells with FV does not have much impact on the generation of excessively activated effector CD8+ T cells.

**FV-specific CD8+ T cells are not responsive to restimulation**

Because there is considerable evidence showing that gag75–83/Db-specific CD8+ T cells generated by FBL3 injection are fully functional (39), it is of interest to compare the actual functionalities of gag75–83/Db-specific CD8+ T cells generated by FV infection with those generated by FBL3 injection, especially during the early stages postinfection/injection. To do so, splenocytes from mice infected with FV or injected with FBL3 were cultured in vitro with gag75–83 peptide for 6 h and evaluated for intracellular expression of IFN-γ and cell-surface expression of CD107a in conjunction with granzyme B expression. Upon peptide restimulation, CD8+ T cells from FBL3-injected mice showed a significant increase in the intracellular expression of IFN-γ and the surface expression of CD107a at 2 and 3 wk after inoculation compared with those prior to the tumor injection (Fig. 2A, 2B). In contrast, CD8+ T cells from FV-infected mice were nonresponsive to the restimulation; neither upregulation of cell-surface CD107a nor an increase in intracellular IFN-γ was observed (Fig. 2A). Surprisingly, we could not find any responses to the peptide restimulation during the entire course of FV infection (Fig. 2B), despite the presence of significant numbers of epitope-specific CD8+ T cells (Fig. 1A). This contrasts remarkably with other models of chronic infection, such as infection with LCMV clone 13, in which virus-specific CD8+ T cells still exhibit an ability to produce IFN-γ upon restimulation during the early stage of infection (10, 41). Furthermore, the excessively activated (granzyme B+) CD8+ T cell populations in FV-infected mice displayed severely impaired IFN-γ production even following anti-CD3 stimulation (Fig. 2A), suggesting that a vast majority of Ag-experienced CD8+ T cells in FV-infected mice have a possible defect in TCR signaling. A strong staining for CD107a was observed intracellularly, supporting the notion that there was a defect in CD107a relocalization from intracellular lytic granules to the plasma membrane in response to TCR signaling in FV-infected mice (data not shown). The impaired responsiveness was not due to a dysfunction of APCs in the spleens of FV-infected mice, because similar results were observed even when restimulated with FBL3 (Fig. 2A). Thus, we hypothesized that although Ag-specific CD8+ T cells from FV-infected mice expressed significant amounts of effector molecules, they were incapable of responding to the target, thereby failing to exert cytolysis. To directly test this, we performed ex vivo [3H]-release assays. CD8+-enriched splenocytes from FV-infected or FBL3-injected mice at day 14 were cocultured with gag75–83 peptide-loaded EL-4 cells at an equivalent E:T ratio, based on the number of gag75–83/Db pentamer+ cells. As shown in Fig. 2C, virus-specific CD8+ T cells from FV-infected mice exerted only marginal levels of cytotoxicity, whereas an equivalent number of virus-specific CD8+ T cells from FBL3-injected mice exhibited significantly greater cytolytic activities. Collectively, these data provide convincing evidence that FV infection results in the generation of virus-specific effector CD8+ T cells that are functionally impaired.

**Minor contribution of Treg cells to the early loss of FV-specific CD8+ T cell function**

Given that infection of mice with FV plus LDV induced CD4+ Treg cells that suppressed virus-specific CD8+ T cells during the chronic phase of infection (24), we first considered the possibility that FV-induced Treg cells might contribute to the dysfunction of Ag-specific effector CD8+ T cells. Because previous studies evaluated the presence of virus-induced CD4+ Treg cells by the expression of CD25, we first compared the frequencies and the actual numbers of CD4+CD25+ T cells in the spleen between FV-infected and CD8+ T cells without stimulation). Data are from the same experiment as in A. Each symbol represents an individual mouse. *p < 0.01; †p < 0.05.

**FIGURE 2.** FV-specific CD8+ T cells are not responsive to restimulation. Mice were infected with FV or injected with FBL3, as described for Fig. 1. **A.** Splenocytes were isolated at day 14 and stimulated with gag75–83 peptide, FBL3, or anti-CD3 mAb for 6 h. The intracellular expression of IFN-γ and granzyme B and the surface expression of CD107a were measured by flow cytometry. Shown are representative staining patterns for granzyme B, IFN-γ, and CD107a of CD8+ T cells. The numbers indicate the percentage of cells gated in each quadrant. As CD8+ T cells were stimulated for 6 h in the presence of anti-CD107a Ab, granzyme B+/CD107a+ cells were observed, indicating degranulation. Data are representative of five independent experiments with essentially the same results. **B.** Ag-specific increase in the percentages of IFN-γ+ and CD107a+ cells among CD8+ T cells calculated by the percentage of IFN-γ+ [CD107a+] cells following peptide stimulation — (the percentage of IFN-γ+ [CD107a+] cells without stimulation). Data are from the same experiment as in A. Each symbol represents an individual mouse. *p < 0.01; †p < 0.05 by two-way ANOVA, with Bonferroni posttests between the groups (FV versus FBL3). One-way ANOVA with Dunnett posttests indicated that the percentages of IFN-γ+ and CD107a+ cells at 2 and 3 wk after FBL3 injection were significantly higher than prior to injection (p < 0.001), and the percentage of CD107a+ cells at 4 wk also was significantly increased (p < 0.05). C, Ex vivo cytotoxic activity of gag75–83/Db-specific effector CD8+ T cells analyzed by [3H]-release assay. Erythroblast (Ter119−)–depleted splenocytes were further enriched for CD8+ cells by negative selection. Cells were incubated with [3H]-labeled EL-4 target cells that had been loaded with gag75–83 peptide for 5 h. Data are representative of two independent experiments with essentially the same results. Bars indicate mean ± SEM.
and FBL3-injected mice. No significant groupwise differences were observed in the frequency or actual numbers of CD4+CD25+ cells between FV-infected and FBL3-injected mice, and the frequency and absolute numbers of CD4+CD25+ cells increased significantly from those before infection only at days 14 and 7 after FV infection, respectively (Fig. 3A, 3B). An additional phenotypic analysis showed that ∼30% of the CD4+CD25+ cells in FV-infected mice were negative for FoxP3, whereas most of the CD4+CD25+ cells in FBL3-injected mice were FoxP3+ (Fig. 3C, 3D). Moreover, a large majority of the FoxP3+ CD4+CD25+ cells in FV-infected mice displayed highly activated phenotypes, as determined by the expression of PD-1 and CD69 (Fig. 3C, 3D), suggesting that the increased numbers of activated cells, but not an increase in Treg cells, were responsible for the slightly elevated levels of CD4+CD25+ T cells after FV infection. In fact, the actual numbers of FoxP3+ Treg cells in FV-infected mice at day 14 were not different from those in FBL3-injected mice. Frequencies of CD4+CD25+ cells in FV-infected and FBL3-injected mice. Frequencies of CD25+ cells among CD4+ T cells and the actual numbers of CD4+CD25+ cells in the spleen at the indicated time points after FV infection or FBL3 injection. Bars indicate SEM. Because the number of samples at each time point differs, t tests with Bonferroni correction for the number of hypotheses were performed. Significant increase from day 0 by p < α (n = 5) = 0.0102. C, Relative expression of activation markers (PD-1 and CD69) and the Treg cell marker (FoxP3) on CD4+CD25+ cells in the spleen of FV-infected or FBL3-injected mice. Bars indicate mean ± SEM. Because the number of samples at each time point differs, t tests with Bonferroni correction for the number of hypotheses were performed. Significant increase from day 0 by p < α (n = 5) = 0.0012. D, Phenotypic differences of CD4+CD25+ cells in FV-infected and FBL3-injected mice. Frequencies of the cells positive for the indicated markers among CD4+CD25+ cells in the spleen at day 14 are shown. p < 0.001; p < 0.05 by two-way ANOVA with Bonferroni posttests. E, Actual numbers of CD4+CD25+FoxP3+ cells in the spleen of FV-infected or FBL3-injected mice at day 14. F, CD8+ T cells were stimulated in vitro by anti-CD3 in the presence or absence (w/o) of Treg cells separated from FV-infected or FBL3-injected mice at day 14 and evaluated for intracellular IFN-γ expression (6 h) and proliferation (48 h). Data are representative of two independent experiments with essentially the same results. p < 0.01; p < 0.05 by one-way ANOVA with Bonferroni posttests.

**Figure 3.** Minor contribution of Treg cells to the early loss of FV-specific CD8+ T cell function. A, Representative staining patterns of CD4 and CD25 in FV-infected or FBL3-injected mice at day 14. B, Frequencies of CD25+ cells among CD4+ T cells and the actual numbers of CD4+CD25+ cells in the spleen at the indicated time points after FV infection or FBL3 injection. Bars indicate mean ± SEM. Because the number of samples at each time point differs, t tests with Bonferroni correction for the number of hypotheses were performed. Significant increase from day 0 by p < α (n = 5) = 0.0102. C, Relative expression of activation markers (PD-1 and CD69) and the Treg cell marker (FoxP3) on CD4+CD25+ cells in the spleen of FV-infected or FBL3-injected mice. Bars indicate mean ± SEM. Because the number of samples at each time point differs, t tests with Bonferroni correction for the number of hypotheses were performed. Significant increase from day 0 by p < α (n = 5) = 0.0012. D, Phenotypic differences of CD4+CD25+ cells in FV-infected and FBL3-injected mice. Frequencies of the cells positive for the indicated markers among CD4+CD25+ cells in the spleen at day 14 are shown. p < 0.001; p < 0.05 by two-way ANOVA with Bonferroni posttests. E, Actual numbers of CD4+CD25+FoxP3+ cells in the spleen of FV-infected or FBL3-injected mice at day 14. F, CD8+ T cells were stimulated in vitro by anti-CD3 in the presence or absence (w/o) of Treg cells separated from FV-infected or FBL3-injected mice at day 14 and evaluated for intracellular IFN-γ expression (6 h) and proliferation (48 h). Data are representative of two independent experiments with essentially the same results. p < 0.01; p < 0.05 by one-way ANOVA with Bonferroni posttests.

**Figure 4.** FV-specific CD8+ T cells show highly exhausted phenotypes. A, Relative levels of PD-1 expression and the pentamer binding on CD8+ T cells and their granzyme B contents in the spleens of FV-infected or FBL3-injected mice at day 14. Profiles shown are of gated CD8+ T cells. B, Representative FACS plots showing the coexpression of inhibitory molecules with PD-1 on the gag75–83/D3-specific CD8+ T cells. Each symbol represents an individual mouse. p < α (n = 3) = 0.017 by two-tailed t test with Bonferroni correction. C, Expression of the indicated surface molecules was assessed on gag75–83/D3-specific CD8+ T cells at the indicated time points after FV infection or FBL3 injection. Each symbol represents an individual mouse. p < α (n = 3) = 0.017 by two-tailed t test with Bonferroni correction. D, Representative FACS plots showing the coexpression of inhibitory molecules with PD-1 on the gag75–83/D3-specific CD8+ T cells in FV-infected and FBL3-injected mice at day 14. Profiles shown are of gated gag75–83/D3+ CD8+ T cells. All data are representative of at least three independent experiments with essentially the same results.

**FV-specific CD8+ T cells show highly exhausted phenotypes.** It is well established that virus-specific CD8+ T cells eventually become dysfunctional during chronic diseases as the result of increased expression of inhibitory receptors, including PD-1. To test whether this could be the case for the observed dysfunction of FV-specific effector CD8+ T cells in FV-infected mice, we analyzed the surface expression of PD-1 on the gag75–83/D3-specific CD8+ T cells. Strikingly, as early as 14 d after FV infection, nearly all gag75–83/D3-specific CD8+ T cells expressed high levels of PD-1 (Fig. 4A). Such a high expression of PD-1 on the gag75–83/D3-specific CD8+ T cells was maintained until ≥10 wk postinfection (Fig. 4B). In contrast to other models of chronic diseases, FV-specific CD8+ T cells show highly exhausted phenotypes.
infection in which the expression levels of PD-1 on exhausted CD8$^+$ T cells were heterogeneous, Ag-specific CD8$^+$ T cells in FV-infected mice were predominantly PD-1$^{hi}$ regardless of the stage of infection (Fig. 4B), revealing a unique severity of exhaustion caused by FV infection. Highly elevated expressions of PD-1 were also observed in the entire population of CD8$^+$ T cells that contained granzyme B in FV-infected mice (Fig. 4A), suggesting that a vast majority of Ag-experienced CD8$^+$ T cells, not just those specific for the gag$^{75–83}$ epitope, shared a propensity for severe exhaustion. Additional phenotypic profiles of Ag-specific CD8$^+$ T cells in FV-infected mice included higher expression levels of CD69 and lower expression of all three of the other markers (CD62L, CD122, and CD127) in comparison with those in FBL3-injected mice, revealing typical characteristics of exhaustion (Fig. 4C) (41). To more thoroughly pursue the severity of exhaustion, Ag-specific CD8$^+$ T cells were examined for the expression of additional inhibitory receptors. Concomitant with the elevated expression of PD-1, gag$^{75–83}$$^D^P$-specific effector CD8$^+$ T cells from FV-infected mice expressed three other cell-surface inhibitory receptors at high levels: Tim-3, LAG-3, and CTLA-4; this expression was obviously distinct from CD8$^+$ T cells with the same epitope specificity in FBL3-injected mice (Fig. 4D). A lack of the elevated expression of FasL and TRAIL suggests that these cells are not yet programmed to undergo activation-induced cell death (Fig. 4D). Taken together, these data suggest that the severe exhaustion of Ag-specific CD8$^+$ T cells occurs even in the early stage of FV infection.

Elevated expression of PD-L1 and MHC class I on FV-infected erythroblasts

The pronounced expression of PD-1 on FV-specific effector CD8$^+$ T cells prompted us to investigate the impact of FV infection on the expression of PD-1 ligands: PD-L1 and PD-L2. In contrast to a modest influence of FV infection on PD-L2 expression, significant upregulation of PD-L1 was observed in subpopulations of CD11b$^+$ dendritic cells and erythroblasts at 14 d after FV infection (Fig. 5A). The elevated expression of PD-L1 on erythroblasts was of particular interest because of the rapid expansion of these cells in the early phase of FV infection. In fact, the percentage of Ter119$^+$ erythroblasts among nucleated splenocytes increased to 75%, with a >45-fold increase in their number at its peak, which was accompanied by the expansive increase in FV-infected cells (Fig. 5B). Because an engagement of PD-L1 and concomitant TCR stimulation, but not the engagement of PD-1 alone, is necessary for providing the negative signal to T cells (42), we further examined the possible expression of MHC class I molecules on the erythroblasts. In agreement with the previous findings that inflammatory stimuli activate hematopoiesis, resulting in increased numbers of nucleated erythroblasts that express MHC class I molecules (43), elevated expression of H-2Db was even observed on the surfaces of erythroblasts from FBL3-injected mice (Fig. 5C). However, in the case of FV infection, nearly all expanded erythroblasts were positive for H-2Db (Fig. 5C). Visualization of the erythroid cell expression of PD-L1 and viral protein revealed that ~50% of virus-infected cells were PD-L1$^{hi}$ (Fig. 5D). Together, these results suggest that the FV-infected erythroblasts can be a potent provider of PD-1–mediated negative signals to CD8$^+$ T cells.

Exhaustion of virus-specific CD8$^+$ T cells is irreversible in FV-infected mice

Thus far, the data have demonstrated that virus-specific CD8$^+$ T cells are exposed to abundant cell-associated viral Ags and immunosuppressive signals; therefore, they are highly exhausted at the early stage of FV infection. To determine whether the functionally exhausted CD8$^+$ T cells could be revived by a blockade of immunosuppressive signals, we attempted to interrupt two pathways by administering blocking Abs to PD-1 and Tim-3, alone or in combination. In the initial experiments, in vivo blockade was started at day 14 (Fig. 6A), when most Ag-specific CD8$^+$ T cells had already expressed these inhibitory receptors at high levels (Fig. 4). Despite the effective blockade of these inhibitory receptors on cell surfaces (Fig. 6B), no reversal of exhaustion was observed, even when in vivo blockade was performed by a combination of anti–PD-1 and anti–Tim-3 (Fig. 6C, 6D). Furthermore, the combined blockade of PD-1 and Tim-3 had no impact on the pathogen control, because the numbers of virus-producing spleen cells detected in the Ab-treated mice were similar to those in the control mice (data not shown). The failure of reinvigoration was probably not caused by the presence of abundant Ags during the blockade, because FV-specific memory CD8$^+$ T cells were still in the nonresponsive state, even when the blockade was operated between days 28–42 postinfection, when initial splenomegaly had regressed (data not shown). Given that CD8$^+$ T cell production of IFN-γ is the last function to be ablated in the stepwise hierarchy of T cell exhaustion (1), it is tempting to hypothesize that high-level viral replication in FV-infected mice results in the generation of terminally exhausted FV-specific effector CD8$^+$ T cells at a very early stage.

Acute SFFV-induced pathogenesis is responsible for the terminal exhaustion of effector CD8$^+$ T cells in FV-infected mice

We next asked whether SFFV-induced acute pathogenesis causes the terminal exhaustion of effector CD8$^+$ T cells in FV-infected animals. To test this, we examined the severity of CD8$^+$ T cell exhaustion in
mice infected with F-MuLV alone. At the peak of infection, FV infection induced severe splenomegaly due to the massive expansion of virus-infected erythroid progenitor cells, whereas such pathologies were not observed in mice infected with F-MuLV (Fig. 7A). Although ~25% of erythroid cells were infected with F-MuLV, these cells did not show any increase in the expression of PD-L1 (Fig. 7B), indicating that infection with F-MuLV alone results in a less immunosuppressive environment compared with that created by infection with FV complex. In the absence of the SFFV-induced acute pathogenesis, we noted a significantly reduced frequency and intensity of PD-1 expression on gag75–83/Db-specific effector CD8+ T cells in F-MuLV-infected mice (Fig. 7C). Because high-level expression of PD-1 can be observed by the time of first cell division under the tolerogenic environment (44), and the massive expansion of FV-infected erythroid cells begins as early as 1 wk postinfection (45), it is possible that Ag-specific CD8+ T cells rapidly enter the state of full functional exhaustion during the priming/differentiation phase in FV-infected animals. To examine this, we next treated FV-infected mice with Abs to PD-1 and Tim-3 alone that was started at day 14 significantly improved Ag-specific IFN-γ production from and CD107α mobilization on CD8+ T cells, indicating that exhaustion of CD8+ T cells in F-MuLV-infected mice was reversible. In contrast, blockade of Tim-3 alone had little effect on the functions of CD8+ T cells (Fig. 7G, 7H), which is likely due to the relatively low expression of Tim-3 on gag75-83/D3-specific effector CD8+ T cells (Fig. 7D). Taken together, these results suggest that SFFV causes rapid terminal exhaustion of virus-specific CD8+ T cells by creating a highly tolerogenic environment.

Simultaneous blockade of PD-1 and Tim-3 pathways during priming/differentiation phase rescues CD8+ T cells in FV-infected mice from becoming terminally exhausted

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Annexin V on their surfaces at day 14 (Fig. 8H). Finally, in contrast to the model of LCMV clone 13 infection (46–48), blockade of IL-10 during the early stage of infection had no impact on the responsiveness of FV-specific effector CD8⁺ T cells or the viral clearance, indicating that FV-specific effector CD8⁺ T cell exhaustion occurred in an IL-10–independent fashion (Fig. 8B–F). Overall, these data demonstrate that, in contrast to F-MuLV infection, infection with FV complex induces a massive expansion of virus-infected PD-L1Hi and MHC class I+ erythroblasts during the priming/differentiation phase, and virus-specific effector CD8⁺ T cells rapidly succumb to terminal exhaustion through accelerated high-level expression of multiple inhibitory receptors.

**Discussion**

It is well established that chronic Ag exposure leads to the stepwise functional exhaustion of CD8⁺ T cells (1, 41). The cells lose the ability to produce IL-2, exert cytosis, and proliferate in early stages of exhaustion, followed by the loss of TNF-α production. IFN-γ production is eventually compromised in the final stage of exhaustion. Although this hierarchical loss of effector functions takes place during the initial phase of chronic infection and causes the failure in viral clearance, terminal exhaustion of Ag-specific CD8⁺ T cells is observed exclusively in the late stages of infection (41). In this study, we have revealed an exception to the above stepwise exhaustion: following FV infection, gag75–83/Db-specific effector CD8⁺ T cells rapidly lose their ability to produce IFN-γ, a function most resistant to being lost, even during the early stage of infection. CD8⁺ T cells specific for an immunodominant epitope are generally more prone to functional exhaustion than are those specific for other less-immunogenic epitopes, resulting in the inverted immunodominance (1). However, FV-driven terminal exhaustion of virus-specific effector CD8⁺ T cells may occur at the population level, because the failure to produce IFN-γ upon restimulation was observed in the entire population of Ag-experienced CD8⁺ T cells in FV-infected mice, not just in the cells specific for the
immunodominant gag epitope. This global nonresponsiveness of Ag-specific CD8+ T cells is consistent with the previous finding that FV-infected mice were incapable of rejecting tumor cells that express FV-related epitopes, although these mice still have an ability to reject other tumor cells that express FV-unrelated epitopes (49).

The most notable feature of these dysfunctional effector CD8+ T cells is the excessive expression of multiple inhibitory receptors that can negatively regulate the CD8+ T cell function, such as PD-1, Tim-3, LAG-3, and CTLA4. PD-1 has been identified as a major cell-surface inhibitory receptor capable of regulating CD8+ T cell exhaustion in mice, as well as in humans and nonhuman primates (50). Tim-3 and CTLA-4 were recently found to be overexpressed on HIV- and hepatitis C virus-specific CD8+ T cells and to act to suppress effector functions of activated T cells (11–13, 51).

There has been considerable evidence that the initial viral load inversely correlates with the functionality of Ag-specific CD8+ T cells (10, 52, 55). Recently, a combined in situ tetramer staining and in situ hybridization approach showed that relative rates of increase in Ag-specific effector CD8+ T cells and virally infected targets during the early stage predict the outcome of viral infection (56). For example, LCMV clone 13 preferentially infects fibroblastic reticular cells,
which enables this clone to spread and replicate tremendously faster than LCMV-Armstrong, thereby exceeding the capacity of initial immune response to reduce the basic rate of replication. Moreover, fibroblastic reticular cells express high levels of PD-L1 soon after LCMV clone 13 infection, resulting in the debilitated proliferation and function of virus-specific CD8\(^+\) T cells (57). These observations strikingly resemble the early events in FV infection, which is represented by the rapid expansion of virus-infected, PD-L1\(^{+}\) erythroblasts and rapid exhaustion of CD8\(^+\) T cells, although the severity of virus infection and CD8\(^+\) T cell exhaustion between these models differs in degree. Considering that CD8\(^+\) T cells upregulate inhibitory molecules at very early stages under the tolerogenic environment and that the negative signals provided at this stage have a profound impact on the early fate-decision of CD8\(^+\) T cells (44, 58), prevention of the potential negative signals during the priming/differentiation phase is rational for rescuing virus-specific CD8\(^+\) T cells from terminal exhaustion and for the effective elimination of virus-infected cells, as we successfully demonstrated in the current study.

One of the key findings from the current study is that severe FV-specific effector CD8\(^+\) T cell dysfunction seemingly occurred, irrespectively of increased Treg cell numbers or functions. In fact, the suppressive ability of Treg cells did not differ between FV infection and FBL3 injection, despite the fact that the latter did not induce CD8\(^+\) T cell exhaustion. Several studies have shown that FV infection results in a slight increase in the number and suppressive ability of Treg cells during the chronic phase (24, 25, 59); however, the precise role of FV-induced Treg cells in inducing the dysfunction of FV-specific CD8\(^+\) T cells in vivo has not been directly elucidated. In fact, previous evidence for the involvement of Treg cells in the CD8\(^+\) T cell dysfunction in vivo is based entirely on the adoptive transfer approach, in which the abilities to secrete effector cytokines were determined by using TCR-transgenic CD8\(^+\) T cells that had been transferred into FV-infected animals at different stages [e.g., acute phase versus memory phase (24)]. However, these studies did not take into consideration the possible impact of different Ag/ inflammation levels at the time of transfer on the priming of TCR transgenic CD8\(^+\) T cells and the subsequent acquisition of effector function. The contribution of Treg cell-mediated suppression to the dysfunction of FV-specific CD8\(^+\) T cells in vivo was also demonstrated by depletion of total Treg cells (24, 60). However, all animals have naturally occurring Treg (nTreg) cells that confer basal suppression of effector cell functions; the depletion of nTreg cells results in the induction of autoimmune and/or inflammatory diseases, which are associated with enhanced autoreactive cellular immune responses (61). Therefore, it is common that total Treg cell depletion in mice results in stronger cellular immune responses than do those in Treg cell–competent mice, and this could be observed with any kind of infection (62, 63). A strategy to deplete virus-induced Treg cells alone, without affecting the numbers and functions of nTreg cells, would be necessary to precisely determine the role of induced Treg cells in vivo, which has never been demonstrated. Thus, it is unknown whether FV-induced Treg cells have any roles in inducing FV-specific CD8\(^+\) T cell dysfunction. In addition, because FV-infected animals are capable of mounting a CD8\(^+\) T cell response against FV-unrelated tumor cells (49), the severe CD8\(^+\) T cell dysfunction seems to occur in an Ag-dependent manner. Because Treg cells seem to suppress a broad spectrum of effector cells, it is unlikely that virus-induced Treg cells provide a selective contribution to the FV-specific CD8\(^+\) T cell dysfunction. Moreover, direct cell-to-cell interaction is required for Treg cells from FV-infected mice to exert suppressive activity on CD8\(^+\) T cells (59), and Treg cell-mediated suppression of CD8\(^+\) T cell function in vivo is known to be reversible (64). These facts strongly support our conclusions on the rather minor role of Treg cells in inducing FV-specific CD8\(^+\) T cell dysfunction, because the opportunity for Treg cells to interact with virus-specific CD8\(^+\) T cells in FV-infected mice must be lower than that in FBL3-injected mice because of the presence of expanded erythroblasts as potential interceptors. Considering the current observations that Ag-specific CD8\(^+\) T cells rapidly succumbed to terminal exhaustion in the early phase of FV infection and could not be reinvigorated, memory CD8\(^+\) T cell dysfunction that has been observed in FV-infected animals may be better explained as a consequence of the currently observed premature terminal exhaustion of effector CD8\(^+\) T cells rather than by Treg cell–mediated suppression.

Another issue to be considered is the possible requirement of virus-specific effector CD8\(^+\) T cells for the control of FV infection. It has been believed that FV-specific CD8\(^+\) T cells initially acquire effector functions during the early phase of infection, based mainly on the observation of their significant expression of granzyme B (25). However, our results draw attention to the importance of determining the responsiveness to restimulation as an indicator of CD8\(^+\) T cell functionality, which is severely compromised in FV-infected mice. Another piece of evidence for the functionality of FV-specific effector CD8\(^+\) T cells also relies on the approach of using supraphysiological numbers of TCR-transgenic CD8\(^+\) T cells specific for the gag epitope (24). However, we clarified the nonresponsiveness of effector CD8\(^+\) T cells by focusing on the endogenous FV-specific CD8\(^+\) T cells induced in FV-infected mice. Because the optimal induction of functionally competent FV-specific CD8\(^+\) T cell responses by early blockade of inhibitory receptors significantly improved the virus control and led to diminished FV-induced splenomegaly, it is reasonable to conclude that FV infection results in the generation of virus-specific effector CD8\(^+\) T cells that are functionally impaired. One question is whether virus-specific CD8\(^+\) T cells contribute to the early control of FV in vivo. Although previous studies demonstrated that the depletion of CD8\(^+\) T cells significantly accelerated the acute splenomegaly (65–67), the FV stock used in these studies had been contaminated with LDV, a potent suppressor of CD8\(^+\) T cell responses (27). Using the LDV-free FV stock, we confirmed an exacerbated acute splenomegaly in CD8\(^+\) T cell–depleted animals upon FV infection (S. Takamura, S. Tsuji-Kawahara, and M. Miyazawa, unpublished observations), which seems to conflict with our current observations regarding the early loss of FV-specific CD8\(^+\) T cell function. One potential explanation is that a short lag exists between the functional maturation and the subsequent exhaustion of CD8\(^+\) T cells in the early stage of FV infection, during which FV-infected cells are at least partially eliminated by effector CD8\(^+\) T cells. Alternatively, because weak, but spontaneous, degranulation was observed on granzyme B–CD8\(^+\) T cells in FV-infected, but not FBL3-injected, mice, the expanded FV-infected erythroblasts that dominated in the spleen can be eliminated in an Ag-independent manner. We are currently investigating these possibilities.

Overall, the data presented in this article indicate that FV, but not F-MuLV, silences antiviral CD8\(^+\) T cell responses during the early phase of infection by inducing the rapid and high expression of multiple inhibitory receptors, leading to irreversible terminal exhaustion. These findings have to be considered a newly recognized strategy for FV to escape from acute CD8\(^+\) T cell responses. Finally, manipulation of CD8\(^+\) T cell exhaustion by early blockade of multiple inhibitory receptors would be necessary to control the pathogens, such as HIV, that exhibit widespread terminal exhaustion of CD8\(^+\) T cells.

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