Differential Expression of Granzyme B and C in Murine Cytotoxic Lymphocytes

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Cytotoxic lymphocytes use the granule exocytosis pathway to kill pathogen-infected cells and tumor cells. Although many genes in this pathway have been extensively characterized (e.g., perforin, granzymes A and B), the role of granzyme C is less clear. We therefore developed a granzyme C-specific mAb and used flow cytometry to examine the expression of granzyme B and C in the lymphocyte compartments of wild-type and mutant GzmB<sup>−/−</sup> cre mice, which have a small deletion in the granzyme B gene. We detected granzyme B and C expression in CD4<sup>+</sup> and CD8<sup>+</sup> T cells activated with CD3/CD28 beads or MLRs. Stimulation of NK cells in vitro with IL-15 also induced expression of both granzymes. Granzyme C up-regulation was delayed relative to granzyme B in wild-type lymphocytes, whereas GzmB<sup>−/−</sup> cre cells expressed granzyme C earlier and more abundantly on a per-cell basis, suggesting that the deleted 350-bp region in the granzyme B gene is important for the regulation of both granzymes B and C. Quantitative RT-PCR revealed that granzyme C protein levels were regulated by mRNA abundance. In vivo, a population of wild-type CD8<sup>+</sup>cre intraepithelial lymphocytes constitutively expressed granzyme B and GzmB<sup>−/−</sup> cre intraepithelial lymphocytes likewise expressed granzyme C. Using a model of a persistent murine CMV infection, we detected delayed expression of granzyme C in NK cells from infected hosts. Taken together, these findings suggest that granzyme C is activated with persistent antigenic stimulation, providing nonredundant backup protection for the host when granzyme B fails. The Journal of Immunology, 2009, 182: 6287–6297.

Cytotoxic T lymphocytes, such as activated CD8<sup>+</sup> T cells and NK cells, use a variety of mechanisms to induce the death of virus-infected and tumor cells. These include secretion of proapoptotic cytokines (e.g., TNF-α and IFN-γ), engagement of cell death receptors (e.g., Fas), and granule exocytosis of perforin and granzymes (1–4). When a fully primed CTL recognizes a target cell that is to be killed, an immunological synapse is formed between the two cells. Cytotoxic granules contained within the lymphocytes polarize toward the synapse, fuse with the CTL plasma membrane, and then release their contents into the synaptic cleft (4). Perforin then facilitates the delivery of granzymes into the cytosol of the target cell, where they cleave a variety of substrates to initiate cell death (3).

The granzyme genes are organized into clusters located on three chromosomes (5). Granzymes A and K are clustered together on human chromosome 5 and mouse chromosome 13, respectively. Granzyme M is tightly linked to a cluster of highly related myeloid serine protease genes on human chromosome 19 and mouse chromosome 10. The human granzyme B gene cluster, found on chromosome 14, contains granzyme B and is followed downstream by granzyme H and cathepsin G (which is exclusively expressed in early myeloid cells). In the murine granzyme B gene cluster on chromosome 14, granzyme C, the closest murine homolog of human granzyme H, is also found directly downstream from granzyme B. Six additional granzymes (5’ F, L, N, G, D, and E) are present between granzyme B and cathepsin G in the murine gene cluster, but the precise specificities and functions of these “orphan” granzymes are not yet known.

Granzyme B, the most thoroughly characterized of the granzymes, cleaves a variety of procaspases, BID, inhibitor of caspase-activated DNase, and other important intracellular substrates to initiate classical apoptotic pathways (3, 6–10). Granzyme C-induced cell death is accompanied by phosphatidylserine externalization, nuclear condensation, ssDNA nicking, and mitochondrial depolarization, but virtually nothing is known about the cellular substrates of this enzyme (11). Interestingly, it was recently shown that cell death induced by human granzyme H, an enzyme with chymotrypsin-like activity, similarly resulted in nuclear condensation and loss of mitochondrial membrane potential (12, 13). Given their high amino acid sequence similarity (61%), locations directly downstream from granzyme B, and similar functional characteristics, these data suggest that granzyme C is the functional murine counterpart to granzyme H. Finally, we have previously shown that the human granzyme H 5’ flanking region is regulated similarly to that of granzyme C in transgenic animals, again suggesting that these genes share regulatory homology (14).

Analysis of granzyme-deficient mouse strains has shown distinct roles for granzymes in antiviral and antitumor immune responses. Granzyme A-deficient mice have an increased susceptibility to ectromelia virus infections (15). Our group recently reported that granzyme B-deficient mice are particularly sensitive...
to challenge with murine CMV (MCMV) (16). We have also recently demonstrated that granzyme B-deficient mice are resistant to challenge with syngeneic and allogeneic tumors (17) due to granzyme B-dependent suppression of antitumor responses mediated by CD4⁺Foxp3⁺ regulatory T cells. Alternative, noncytotoxic activities have been proposed for granzyme A in mediating immune responses, but have not been described for granzymes B or C (18).

Although a granzyme C knockout mouse has not yet been generated, there is evidence that suggests that granzyme C plays a role in CTL-mediated cytotoxicity. We previously compared the cytotoxic function of two granzyme B-deficient strains, one in which expression of granzyme C and F were decreased due to a retained PGK-neo cassette and one in which expression of these downstream genes was restored after removal of this cassette (19). CTLs from mice with diminished granzyme C and F expression had a corresponding reduction in cytotoxicity. More recently, Getachew et al. (20) used small interfering RNA-mediated knockdown of granzyme C in CTLs derived from MLRs to demonstrate a role for this gene during prolonged T cell responses.

The kinetics and patterns of granzyme expression in cytotoxic lymphocytes are dependent on the conditions under which these cells are activated. Kelso et al. (21) previously reported the differential expression of granzyme transcripts in polyclonally activated T cells. By performing nested RT-PCR on single CD8⁺ T cells that were sorted from in vitro-activated cultures at various time points, they found that up-regulation of granzyme B transcripts preceded the induction of granzyme A and C mRNAs. Furthermore, individual T cells had different patterns of expression of these genes. However, until now, there has been no thorough characterization of granzyme C protein expression at the single-cell level.

In this study, we report flow cytometric analyses of granzyme B and C protein expression in activated T and NK cells using a novel granzyme C-specific mAb. Our findings demonstrate the differential expression of granzymes in cytotoxic lymphocytes and shed insight into regulatory mechanisms that control the expression of these genes.

**Materials and Methods**

**Mice**

Wild-type (WT) 129/SvJ, C57BL/6, BALB/c, and Rag1⁻/⁻ (B6) mice were obtained from The Jackson Laboratory. GzmB⁻/⁻ cluster and GzmB⁺ cre mice have been previously described and were derived in the 129/SvJ background (19, 22). All mice were maintained in specific pathogen-free housing and all experiments were conducted in accordance with institutional animal care and use guidelines.

**Hamster mAb production**

Armenian hamsters were immunized with purified recombinant granzyme C in CFA and boosted in IFA. Hamsters showing ELISA seropositivity for granzyme C were boosted and hybridomas were generated as previously described (23). Hybridoma supernatants were screened by flow cytometric analysis of fixed and permeabilized lymphokine-activated killer (LAK) cells. A PE-conjugated goat anti-Armenian hamster IgG secondary Ab (Jackson ImmunoResearch Laboratories) was used for detection. Multiple positive cell lines were identified. One of these, SFC1D8, was selected for further characterization and repeatedly cloned by limiting dilution. Hamster IgG was purified from SFC1D8 hybridoma supernatants by protein A affinity chromatography and subsequently labeled with Alexa Fluor 488 (Invitrogen Protein Labeling Kit). With the exception of the original screening of hybridoma clones, all subsequent staining was performed using this directly conjugated form of SFC1D8.

**Abs and reagents**

Abs used include anti-mouse NK1.1 (PK136), CD3 (145-2C11), CD4 (RM4-5), CD8 (53-6.7), CD16/32 (2.4G2; BD Biosciences), Nkp46 (29A1.4; eBioscience), and granzyme B (GB12; Caltag Laboratories). Mouse CD3/CD28 Dynabeads were obtained from Invitrogen. Cytokines were obtained from R&D Systems (recombinant murine IL-15) or Chiron (recombinant human IL-2). All cytokines were endotoxin free and stored at −80°C after reconstitution in PBS plus 0.1% BSA.

**Cell isolation and stimulation**

LAK cell and MLR preparations were performed as previously described (22, 24). All cells were cultured in K10 medium (RPMI 1640, 10% FCS, 10 mM HEPES, 1% nonessential amino acids, 1% sodium pyruvate, 1% L-glutamine, 1% penicillin/streptomycin, and 0.57 μM 2-ME). For CD3/CD28 activations, bulk splenocytes (2 × 10⁶ cells/well) from WT or GzmB⁻/⁻ cre mice were processed into single-cell suspensions and cultured in 6-well plates with Dynabeads at a 1:1 ratio. At indicated time points, cells were harvested for analysis. IL-15 stimulation of splenocytes was performed as previously described (16). CD8⁺ T cells were purified from the resting spleens of WT and GzmB⁻/⁻ cre mice with a CD8⁺ T cell isolation kit, followed by cell separation on the AutoMACS according to the manufacturer’s instructions (Miltenyi Biotec). For purification of NK cells, splenocytes were surface stained with anti-NKp46 and anti-CD3, and Nkp46⁺ CD3⁻ cells were isolated on a Reflection (iCyt) cell sorter (routinely ≥95% pure).

**Preparation of intraepithelial cells (IELs)**

For isolation of intestinal IELs, small and large intestines were harvested, Peyer’s patches were removed, and tissue was cut open longitudinally. The washed tissue was cut into 1-cm segments. Intraepithelial lymphocytes were released from associated epithelium with two consecutive washes of HBSS with shaking at 220 rpm for 20 min at 37°C. Lymphocytes were then passed through sterile 70-μm nylon mesh and analyzed.

**Intracellular staining and flow cytometry**

One × 10⁶ cells were washed and resuspended in staining buffer (PBS, 0.5% BSA, and 0.5 mM EDTA). Samples were labeled with primary-conjugated Abs against cell surface markers, fixed, permeabilized (Foxp3 staining kit; eBioscience), and stained with primary-conjugated anti-granzyme B Ab and anti-granzyme C Ab. Sample data were acquired on a Cytex-modified FACScan (BD Biosciences) flow cytometer and analyzed with FlowJo (Tree Star) software.

**Quantitative real-time RT-PCR**

Total RNA was isolated from resting and activated CD8⁺ T cells and NK cell samples (2.5 × 10⁶) with the RNeasy Micro Kit (Qiagen). Quantitative RT-PCR (qRT-PCR) was performed as described for granzyme B and C (19). Comparisons were made by the comparative threshold cycle (Ct) method, with β-actin serving as the comparator. Data are presented as fold change relative to resting NK or T cell samples, which are set to a value of 1.

**MCMV infection**

A salivary gland stock of Smith strain MCMV was prepared from BALB/c mice that had been i.p. injected with tissue culture-propagated MCMV, and the titer was determined via standard plaque assay using permissive NIH3T12 fibroblasts (American Type Culture Collection) (25). Rag1⁻/⁻ mice were injected i.p. with 2 × 10⁶ PFU/mouse of a salivary gland MCMV stock. Twenty-one days after infection, splenocytes were harvested and analyzed.

**Results**

Characterization of granzyme expression in CD3/CD28 bead-activated CD4⁺ and CD8⁺ T cells

To define the expression of granzymes at the single-cell level in activated lymphocytes, we generated a granzyme C-specific mAb. Hybridoma clones were obtained after immunization of Armenian hamsters with purified recombinant granzyme C and CFA. Supernatants from these hybridoma clones were screened using flow cytometric analysis of LAK cells derived from 129/SvJ WT mice and two types of strain-matched granzyme B-deficient mice (granzyme B⁻/⁻ cluster and granzyme B⁻/⁺ cre mice). GzmB⁻/⁻ cluster mice have a PGK-neo cassette retained within the granzyme B
gene, which produces a neighborhood effect that reduces the expression of downstream granzymes, including granzyme C (19, 26). This PGK-neo cassette was removed by cre-mediated recombination in targeted embryonic stem cells, which were then used to create GzmB−/− cre mice. As a result, no neighborhood effect is observed in these mice. In fact, greater amounts of granzyme C mRNA and protein were detected in activated CTLs from GzmB−/− cre mice compared with their WT counterparts (19). Since no knockout mice for granzyme C currently exist, we used the differential expression of granzyme C in these mice to define the specificity of candidate hybridoma clones in a flow-based assay. Several reactive clones were identified. One of the clones

FIGURE 1. Characterization of granzyme protein expression in activated lymphocytes using a novel granzyme C-specific mAb. LAK cells were generated by culturing splenocytes from WT, granzyme B−/− cre, or granzyme B−/− cluster-deficient mice in K10 medium supplemented with high-dose IL-2 (1000 U/ml). After 10 days of culture, LAK cells were harvested, fixed, permeabilized, and stained for intracellular granzyme C followed by staining with a PE-conjugated anti-hamster IgG secondary Ab. A secondary Ab alone condition (gray) was included as a negative control. Representative histograms are shown in A. A summary graph plotting MFI of granzyme C expression (mean ± SD) from three independent experiments is shown in B. * p < 0.0001. WT and granzyme B−/− cre splenocytes were cultured in K10 medium with CD3/CD28 beads and harvested for flow cytometric analysis at various times during activation. T cell expression of granzymes B and C are shown. Results from an individual experiment are shown, gating on CD4+ T cells (C) and CD8+ T cells (D).
stained GzmB<sup>−/−</sup> cre LAK cells with a higher mean fluorescence intensity (MFI) than WT cells and also stained GzmB<sup>−/−</sup> cluster LAK cells with a significantly lower MFI than WT cells (Fig. 1, A and B). This pattern of staining is consistent with that of a granzyme C-specific Ab. However, this Ab did not recognize either native or recombinant murine granzyme C with Western blotting of proteins separated using SDS-PAGE, suggesting that it recognizes a folded epitope (data not shown). This was confirmed by direct spotting of serially diluted, non-denatured recombinant murine granzymes A, B, and C followed by detection with the anti-granzyme C mAb (19, 27, 28). Under conditions where the Ab could detect 12.5 ng of granzyme C, 500 ng of recombinant granzyme B and 1000 ng of granzyme A yielded no signal with this Ab. These data show that this Ab has minimal cross-reactivity with granzymes A and B, the most closely related granzyme family member by amino acid sequence similarity (66% identical) (supplemental Fig. 1A). Granzyme F also has a similar expression pattern to granzyme C in LAK cells, which raises the possibility that this Ab may potentially cross-react with granzyme F in the flow-based assay (19). Because granzymes C and F share less sequence similarity (59.9% amino acid identity) than granzymes C and B (66.1% amino acid identity) and because there was minimal cross-reactivity between granzymes C and B in both the flow-based and dot blot assays, it is not likely that this Ab cross-reacts with granzyme F. In the absence of a granzyme C-deficient mouse; the flow cytometric analysis of LAK cells derived from mice with genetically defined amounts of granzyme C expression, coupled with dot blot analysis of available granzymes in recombinant form, strongly suggests that this reagent is granzyme C specific. Using this Ab, along with a commercially available granzyme B-specific Ab (GB12) (16), we characterized the expression of granzymes B and C in polyclonally activated T cells over a 6-day time course.

Unfractionated splenocytes from WT and GzmB<sup>−/−</sup> cre mice were cultured with CD3/CD28 beads and harvested at various times for flow cytometric analysis. Representative flow plots of granzyme B and C expression in CD4<sup>+</sup> T cells are shown in Fig. 1C and a summary graph is shown in Fig. 2A. Although the percentage of granzyme B-expressing WT CD4<sup>+</sup> T cells peaked on day 3 of activation at ~25%, the proportion of granzyme C-expressing CD4<sup>+</sup> T cells continued to increase throughout the 6-day time course. Approximately 75% of all WT CD4<sup>+</sup> T cells in culture had detectable amounts of granzyme C protein on day 6 of activation. Notably, the rise in granzyme C expression lagged behind the induction of granzyme B protein by ~2 days. As expected, no granzyme B was detectable at any time using GzmB<sup>−/−</sup> cre-derived splenocytes. However, a greater proportion of CD4<sup>+</sup> T cells were granzyme C<sup>+</sup> on days 5 and 6. By day 6, 100% of GzmB<sup>−/−</sup> cre CD4<sup>+</sup> T cells expressed granzyme C. Analysis of Foxp3<sup>+</sup> regulatory and Foxp3<sup>−</sup> effector CD4<sup>+</sup> T cells showed that this activation protocol preferentially expanded effector T cells; almost all of the CD4<sup>+</sup> T cells were Foxp3<sup>−</sup> at the end of the time course (data not shown). In addition, based on the fluorescence intensity of granzyme C staining, the amount of granzyme C protein in GzmB<sup>−/−</sup> cre CD4<sup>+</sup> T cells on a per-cell basis was ~5-fold greater than in WT cells (Fig. 1C).

Representative flow plots of granzyme B and C expression in CD3/CD28 bead-activated CD8<sup>+</sup> T cells is shown in Fig. 1D and a summary graph is shown in Fig. 2B. Like WT CD4<sup>+</sup> T cells, the percentage of granzyme B-expressing WT CD8<sup>+</sup> T cells peaked on day 3 of activation at ~80%, whereas the proportion of granzyme C-expressing CD8<sup>+</sup> T cells began rising above baseline on day 4, reaching ~80% on day 6. For CD8<sup>+</sup> T cells from GzmB<sup>−/−</sup> cre cultures, the proportion of granzyme C-expressing CD8<sup>+</sup> T cells began rising on day 2 and continued to increase until all CD8<sup>+</sup> T cells in culture were granzyme C<sup>+</sup>. Again, as was seen for CD4<sup>+</sup> T cells, a higher granzyme C MFI (~5-fold on day 6) was observed in GzmB<sup>−/−</sup> cre CD8<sup>+</sup> T cells.

Characterization of granzyme expression in CD4<sup>+</sup> and CD8<sup>+</sup> T cells activated during MLRs

In these experiments, we used allogeneic mismatch as another stimulus for T cell activation. Our group had previously reported the expression of granzyme C by Western blot analysis of bulk populations harvested from MLRs (19). A flow-based assay using our mAb allows for a more detailed analysis of discrete T cell populations as a function of time. Splenocytes from 129/SvJ WT and GzmB<sup>−/−</sup> cre mice (H-2<sup>b</sup>) were cultured with irradiated splenocytes from BALB/c mice (H-2<sup>k</sup>) and harvested at various times for flow cytometric analysis. Representative flow plots gated on CD4<sup>+</sup> T cells are shown in Fig. 3A and a summary graph is shown in Fig. 3C. No granzyme B or C was detectable during the first 3 days of culture in either WT or GzmB<sup>−/−</sup> cre MLRs. The proportion of granzyme B- and C-expressing CD4<sup>+</sup> T cells began rising on day 4 and peaked on day 5 with ~10% granzyme B<sup>+</sup> and ~30% granzyme C<sup>+</sup> in WT cultures. All CD4<sup>+</sup> T cells from GzmB<sup>−/−</sup> cre MLRs were granzyme C<sup>+</sup> by day 5. Further analysis of CD4<sup>+</sup> effector and regulatory subsets showed that granzyme C expression was restricted to the Foxp3<sup>+</sup> compartment (data not shown).

Flow plots and summary graphs for MLR-activated CD8<sup>+</sup> T cells are shown in Fig. 3, B and D, respectively. Similar to MLR-activated CD4<sup>+</sup> T cells, neither granzyme B nor C was detected in CD8<sup>+</sup> T cells during the first 3 days of culture, and there was only a slight increase in the percentage of WT CD8<sup>+</sup> T cells that were granzyme B<sup>+</sup> on day 4. After 5 days of culture, however, there was

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4 The online version of this article contains supplemental material.
a substantial increase in granzyme B and C double-positive cells, and there were few granzyme single-positive T cells. Analysis of MLR cultures in an extended time course showed that there was little change in the proportion of granzyme B- and granzyme C-expressing CD4$^+$ and CD8$^+$ T cells from day 5 through day 8 (supplemental Fig. 2). Interestingly, $GzmB^{-/-}$ cre CD4$^+$ T cells had distinct patterns of granzyme C positivity. Although CD4$^+$ T cells are homogeneously granzyme C positive, the population of granzyme C-positive CD8$^+$ T cells is distributed over a 1-log range in fluorescence intensity, suggesting that there may be greater heterogeneity in CD8$^+$ T cell subsets activated in culture relative to CD4$^+$ T cells. Unlike CD3/CD28-activated T cells, no lag in granzyme C expression was observed relative to granzyme B, although expression of granzyme B and C protein during MLR was delayed and less robust. We attribute these differences to the strength of the activation signal, since allogeneic mismatch stimulates only a small proportion of the total T cells in culture, whereas CD3/CD28 activation stimulates virtually the...
entire T cell population (29). Granzyme C was maximally
expressed in \textit{GzmB}^+/H11002/+/H11002/cre CD8^+/H11001 T cells on day 5 and little to no
granzyme C was observed at earlier time points.

Differential granzyme B and C protein expression in WT and
\textit{GzmB}^+/H11002/+/H11002/cre CD8^+/H11001 T cells is regulated by mRNA abundance

Our flow cytometric analysis of polyclonally activated T cells re-
vealed that granzyme C lags behind the induction of granzyme B
in WT T cells and that the induction of granzyme C in \textit{GzmB}^+/H11002/+/H11002/cre T cells occurs earlier and is more abundant on a per-cell basis
when compared with WT T cells. These data suggest that gran-
zymes B and C may be regulated by distinct mechanisms during T
cell activation and subsequent differentiation into a cytotoxic lym-
phocyte. To determine whether this process is regulated at the level
of mRNA abundance, we performed quantitative real-time RT-
PCR on CD3/CD28-activated CD8^+/H11001 T cells obtained from WT or
\textit{GzmB}^+/H11002/+/H11002/cre mice (Fig. 4A). The abundance of granzyme B
mRNA between days 0 and 2 in WT CD8^+/H11001 T cells increased by at
least a factor of 10, whereas the fold change for granzyme C was
much less on day 2. By day 4, the fold changes for granzyme B and
granzyme C mRNAs were comparable. In contrast, mRNA levels
of granzyme C in \textit{GzmB}^+/H11002/+/H11002/cre CD8^+/H11001 T cells increased on day 2
and the fold changes from baseline on days 4 and 6 were higher
than that of WT cells. Consistent with the flow data, there is a
reproducible lag in the induction of granzyme C mRNA abundance
compared with granzyme B in WT CD8^+/H11001 T cells, and the fold
increase in granzyme C mRNA abundance for \textit{GzmB}^+/H11002/+/H11002/cre
CD8^+/H11001 T cells was significantly higher than that observed in WT T
cells at all time points after activation. These data suggest that the
differential expression of granzymes in CD3/CD28-activated T
cells is regulated by mRNA abundance.

To demonstrate that the difference in fold change observed on
day 4 in granzyme C mRNA between WT and \textit{GzmB}^+/H11002/+/H11002/cre
CD8^+/H11001 T cells is due to higher granzyme C mRNA abundance in
\textit{GzmB}^+/H11002/+/H11002/cre T cells (and not due to altered actin mRNA abun-
dance, which was used to normalize expression levels), we plotted
the raw threshold cycles detected in each biological replicate for

\begin{figure}[h]
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\includegraphics[width=\textwidth]{figure4}
\caption{Quantitative real-time RT-PCR comparing WT and gran-
zyme B^+/H11002/+/H11002/cre T cell expression of granzyme C mRNA. CD8^+/H11001 T cells
were cultured in K10 medium with CD3/CD28 beads and RNA was har-
vested for qRT-PCR at various times during activation. The mean \pm SEM
expression from three independent experiments is shown as fold change of
activated compared with naive T cells (A). The comparative Ct method
was used and granzyme C expression was normalized to \textit{\beta}-actin. Raw threshold
cycles for actin and granzyme C mRNA on day 4 are plotted on an inverted
scale in B.}
\end{figure}

FIGURE 4.
per panels is shown in A. A summary of the percentage of granzyme Band C-positive NK cells from three independent experiments (mean ± SD) is shown in B, demonstrating that granzyme C protein is expressed after granzyme B following IL-15 activation.

FIGURE 6. Granzyme C protein is expressed later than granzyme B protein in IL-15-activated NK cells from WT C57BL/6 mice. Splenocytes were isolated from WT C57BL/6 mice, stained, and analyzed by flow cytometry for the expression of intracellular granzyme B and C at rest (day 0) or after 1, 2, 3, or 4 days of activation with recombinant murine IL-15 (100 ng/ml). Representative flow plots (gated on NK1.1 T cells) showing the majority of the cells express granzyme B and C at rest (day 0) or after 1, 2, 3, or 4 days of culture in the presence of IL-15. The majority of NK cells were granzyme B+ after 1 day of culture, the majority did not become granzyme C+ until day 3. By day 4, almost all NK cells expressed both granzymes B and C. Similar kinetics were also observed in NKp46+/CD3− NK cells from WT 129/SvJ mice (Fig. 6A). A summary graph is shown in Fig. 6B. Consistent with our previously published observations, there was little to no granzyme B protein expression in resting NK cells, but after 1 day of culture in the presence of IL-15, the majority of NK cells were granzyme B+. The proportion of granzyme B+ NK cells continued to rise, peaking on days 3 and 4 at ~95%. Similar to T cells, NK cell expression of granzyme C lagged behind granzyme B. Whereas most NK cells were granzyme B+ after 1 day of culture, the majority did not become granzyme C+ until day 3. By day 4, almost all NK cells expressed both granzymes B and C. Similar kinetics were also observed in NKp46+/CD3− NK cells from WT 129/SvJ mice (Fig. 7A). However, granzyme C up-regulation was not delayed in GzmB+/− cre NK cells (Fig. 7B). Notably, the kinetics of granzyme C expression in GzmB+/− cre NK cells was nearly identical to that of granzyme B expression in WT NK cells (Fig. 7C).

Characterization of granzyme C expression in IL-15-stimulated NK cells

Our group previously demonstrated that resting murine NK cells express abundant amounts of granzyme A protein but not granzyme B or perforin, although transcripts for granzyme A, granzyme B, and perforin were all detected in resting NK cells (16). Granzyme B and perforin protein were induced upon cytokine stimulation, with IL-15 having the most potent effect. To define the kinetics of granzyme C expression in NK cells under these conditions, bulk spleen cells were stimulated with IL-15 over a 4-day time course. Splenocytes from WT B6 mice were cultured and harvested daily for flow cytometric analysis of granzyme B and C expression gated on NK1.1− CD3+ NK cells (Fig. 6A). A summary graph is shown in Fig. 6B. Consistent with our previously published observations, there was little to no granzyme B protein expression in resting NK cells, but after 1 day of culture in the presence of IL-15, the majority of NK cells were granzyme B+. The proportion of granzyme B+ NK cells continued to rise, peaking on days 3 and 4 at ~95%. Similar to T cells, NK cell expression of granzyme C lagged behind granzyme B. Whereas most NK cells were granzyme B+ after 1 day of culture, the majority did not become granzyme C+ until day 3. By day 4, almost all NK cells expressed both granzymes B and C. Similar kinetics were also observed in NKp46+/CD3− NK cells from WT 129/SvJ mice (Fig. 7A). However, granzyme C up-regulation was not delayed in GzmB+/− cre NK cells (Fig. 7B). Notably, the kinetics of granzyme C expression in GzmB+/− cre NK cells was nearly identical to that of granzyme B expression in WT NK cells (Fig. 7C).

Granzyme C is constitutively expressed in CD8αα+ IELs of GzmB−/− cre mice

To further define the conditions under which granzymes B and C are expressed in vivo, we isolated IELs from WT and GzmB−/− cre mice. Within this lymphoid compartment, there are well-defined populations of thymus-dependent and thymus-independent T cells expressing the α/α homodimeric form of CD8 along with either αβ or γδ TCRs. Other groups have reported that these cells constitutively express perforin, granzyme A, and granzyme B mRNA (30, 31). These cells are able to kill target cells in redirected cytotoxicity assays as well as lymph node-derived CD4+ T cell blasts (32). We therefore analyzed these T cell subsets by flow cytometry to determine whether they express granzyme B and C protein. Immunophenotyping analysis gated on CD8α− IELs revealed that the majority of the cells express γδ TCRs, while the rest express αβ TCRs. Almost all of the gated cells are CD8β−, thereby confirming that these cells express the CD8αα homodimer (Fig. 5A). Within each of these two subsets, there are discrete populations of granzyme B+ and granzyme B− T cells, with the majority of the gated population expressing granzyme B (Fig. 5, B and C). Costaining experiments further showed that no granzyme C was detected in WT CD8αα+ T cells. However, ~30% of CD8αα+ IELs harvested from GzmB−/− cre mice were granzyme C+.

Characterization of granzyme C expression in IL-15-stimulated NK cells

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Granzyme C protein expression in WT and GzmB−/− cre NK cells is regulated by mRNA abundance

Next, we used quantitative real-time RT-PCR to examine the expression of granzyme B and C mRNAs in flow-sorted NK cells during a time course of IL-15 activation (Fig. 8). Consistent with our published findings, there is a modest increase in granzyme B
mRNA after 1 day of IL-15 stimulation (16). WT NK cells had a similar increase in granzyme C mRNA abundance on day 1. Although granzyme C mRNA levels continued to rise throughout the 3-day time course, granzyme B mRNA abundance stabilized after 1 day. For GzmB−/− cre NK cells, there was a greater fold increase in granzyme C mRNA at each time point after IL-15 activation. Microarray analyses revealed that no granzyme C transcripts were detectable in resting NK cells, which stands in distinct contrast to that of granzyme B and perforin, which have abundant mRNA in resting NK cells, but no protein expression (16). After NK cell activation with IL-15, granzyme B and perforin mRNAs are rapidly translated, arming the cells to kill their targets. Taken together, these data suggest that the regulation of granzyme C in activated NK cells is fundamentally different from that of granzyme B or perforin.

Persistent viral infection induces NK cell expression of granzyme C in vivo

To determine whether NK cells express granzyme C in vivo during a viral infection, we challenged WT B6 mice with a sublethal dose (5 × 10⁶ PFU) of Smith strain MCMV and analyzed the spleens of infected mice by flow cytometry. In this well-characterized mouse model of an acute infection, there is an early NK cell-dependent phase of viral clearance. We previously reported that granzyme B and perforin are rapidly up-regulated in NK cells during the first 8 days following infection (16). However, we were unable to detect granzyme C during this time frame (data not shown). Because granzyme C expression is relatively delayed and has been
postulated to serve as a fail-safe mechanism in killing target cells, we hypothesized that a sublethal dose of MCMV that is rapidly cleared may not provide sufficient time and/or activation signals for the induction of granzyme C in NK cells. To address this hypothesis, we infected immunodeficient B6 Rag1−/− mice with a lower dose (2 × 10⁷ PFU) of MCMV. In this model, NK cells were able to mount a response to the viral challenge initially, but weeks later, the infected mice died due to consequences of uncontrolled MCMV replication (33, 34). The selection pressure exerted by Ly49H⁺ NK cells in the absence of adaptive immunity generated escape mutants that ultimately resulted in the death of these immunodeficient hosts. We hypothesized that a viral infection that persists beyond the initial NK cell-dependent phase of clearance may induce the expression of granzyme C in NK cells. Splenocytes from B6 Rag1−/− mice were harvested on day 21 after infection and analyzed for granzyme B and C expression in NK1.1⁺ CD3⁺ NK cells. Representative flow plots are shown in Fig. 9A and summary graphs are depicted in Fig. 9B. As expected, NK cells from uninfected mice had little or no detectable granzyme B or C. Approximately 70% of NK cells from infected hosts were granzyme B⁺ and a smaller proportion (~15%) expressed granzyme C. Thus, although immunocompetent hosts challenged with a sublethal dose of MCMV failed to induce NK cell expression of granzyme C, we did detect granzyme C expression in NK cells from immunodeficient Rag1−/− mice that rely on innate NK cells for transient control of a persistent and ultimately lethal MCMV infection.

Discussion
In this report, we developed a novel mAb to characterize the expression of granzyme C in cytotoxic lymphocytes at single-cell resolution. Allogeneic mismatch and coculture with CD3/CD28 beads induces granzyme C expression in both CD4⁺ and CD8⁺ T cells. Activation of NK cells with IL-15, a cytokine previously shown to induce potent perforin-dependent cytotoxicity in NK cells, induces granzyme C mRNA and protein expression. Granzyme C activation was delayed relative to granzyme B in WT T and NK cells, while granzyme C was expressed earlier and was more abundant on a per-cell basis in GzmB−/− cre CTLs. The expression of granzyme C protein was regulated at the level of mRNA abundance in CD3/CD28-activated T cells and in IL-15-activated NK cells. In addition to in vitro-activated lymphocytes, we also detected granzyme C expression in vivo in CD8αα⁺ intestinal IELs harvested from GzmB−/− cre mice as well as in NK cells from immunodeficient mice that had been challenged with a persistent and lethal viral infection.

The delayed expression of granzyme C protein relative to granzyme B is in concordance with previous studies that measured mRNA abundance of various components of the perforin/granzyme pathway during T cell activation. Kelso et al. (21) activated purified naïve CD8⁺ T cells with immobilized Abs to CD3, CD8, and CD11a and used single-cell PCR to demonstrate that up-regulation of granzyme B mRNA preceded granzyme C by 1 day. The fact that this delay in granzyme C expression was observed across multiple lymphocyte subsets and activation methods suggests that the granzyme genes are subject to distinct regulatory mechanisms. Furthermore, these data also support the notion that orphan granzymes serve as fail-safe mechanisms for the induction of target cell death.

Recently, Thiele and colleagues (20) provided functional data in support of this hypothesis. By comparing CTLs from WT mice and mice deficient for dipeptidyl peptidase I (DPPI, a protease that is required for the functional activation of granzymes A and B, but not granzyme C), they showed that DPPI−/− CTLs that had been primed over a period of 5 days exhibited reduced cytotoxicity. However, after restimulating the cells for an additional 3 days, cytotoxic function in DPPI−/− CTLs had returned to WT levels. The restoration of cytotoxicity in DPPI−/− CTLs correlated with late granzyme C up-regulation. Furthermore, knockdown of granzyme C during restimulation decreased the cytotoxic activity of DPPI−/− (but not WT) CTLs. Together, these data suggest that granzyme C can maintain cytotoxic activity during late T cell responses in the absence of granzymes A and B.

These observations complement our expression studies of NK cells activated during MCMV infection. Under conditions where granzyme B and perforin are rapidly induced following challenge of WT mice with a sublethal dose of MCMV, no granzyme C was detected. However, when mice that lack an adaptive immune system were challenged with a dose that allowed the virus to persist and eventually kill the host, a population of granzyme C-expressing NK cells arose relatively late during the course of the infection. These findings suggest that the persistence of the virus in the host may provide sufficient time for the up-regulation of granzyme C in NK cells. Alternatively, additional signals generated during the prolonged course of the infection may be required for granzyme C to be expressed.
FIGURE 10. Model of granzyme B and C regulation in WT, GzmB−/− cre, and GzmB−/− cluster-derived cytotoxic lymphocytes. In WT cytotoxic lymphocytes (A), the LCR first interacts with regulatory elements within or near the AvrII fragment of granzyme B to induce transcription. Granzyme C is induced only after prolonged activation. GzmB−/− cluster-derived cytotoxic lymphocytes (B) have the normal AvrII fragment replaced with a PGK-neo cassette and fail to express granzyme B due to disruption of the gene. In addition, granzyme C expression is diminished due to the neighborhood effect created by the retained PGK-neo cassette. In GzmB−/− cre CTLs (C), granzyme C is expressed earlier and more abundantly (similarly to granzyme B in WT cells), because missing regulatory elements in granzyme B cause the LCR to “scan” downstream and activate the next gene in the cluster. Removal of the PGK-neo cassette eliminates the neighborhood effect.

The GzmB−/− cre mice were made after our group discovered that activated CTLs from GzmB−/− cluster mice also had diminished expression of granzymes C and F. The null mutation in GzmB−/− cluster mice was originally generated by replacing a 350-bp AvrII fragment (containing most of exon 1, including the start codon, and 283 bp of intron 1) with a PGK promoter-driven neomycin phosphotransferase cassette. Pham et al. (26) reported that activation of GzmB−/− cluster CTLs resulted in robust induction of PGK-neo mRNA at the expense of the expression of downstream granzymes, suggesting that PGK-neo had been captured by regulatory elements in the granzyme B gene cluster (26). We postulated that the retained PGK-neo cassette exerted a “neighborhood effect” on the downstream granzyme genes in the cluster.

Consistent with that hypothesis, we found that deletion of a loxp-flanked PGK-neo cassette targeted to the same location resulted in the earlier, more abundant expression of granzyme C mRNA and protein relative to WT CTLs. Additionally, although CD8αα− IELs from WT mice expressed granzyme B but not granzyme C, GzmB−/− cre IELs had constitutive expression of granzyme C in vivo. In the case of CD3/CD28 activation, we found that up-regulation of granzyme C in GzmB−/− cre CD8+ T cells occurred 2 days earlier than WT CD8+ T cells. Interestingly, the kinetics of granzyme C expression in GzmB−/− cre CTLs was identical to that of granzyme B expression in WT CTLs (true for both CD3/CD28-mediated T cell activation and IL-15-stimulated NK cell activation). Since the only difference between WT and GzmB−/− cre mice is the presence or absence of the 350-bp AvrII fragment (and the retained loxp site), we hypothesize that deletion of this fragment causes a recently defined locus control region (LCR) to “skip” granzyme B and proceed to activate the next gene downstream, granzyme C; this gene must contain regulatory elements that permit the LCR to activate it (35). Although we have demonstrated that levels of granzyme C protein expression are regulated by mRNA abundance, it remains unclear whether regulation of transcriptional activity or mRNA half-life account for the differential expression of granzyme C mRNA in WT and GzmB−/− cre lymphocytes. Alternatively, GzmB−/− cre mice may generate signals that activate additional granzymes as a fail-safe mechanism when these animals fail to clear viruses or tumors, or granzyme B mRNA or protein may repress granzyme C expression, a process that could be overcome during persistent stimulation.

Bleackley and colleagues (35) have recently developed strong evidence for a LCR that regulates gene expression in the granzyme B gene cluster. These investigators identified a DNA element located upstream from granzyme B that is involved in the control of its transcription (35). In their studies, a DNase I-hypersensitive site (HS2) was detected 3.9 kb upstream from the transcriptional start site of granzyme B in activated T cells. When this DNA element was included in transgenes that were transferred to a CTL line, HS2 conferred position-independent expression of granzyme B. Transgenic mice expressing a tagged version of granzyme B (to distinguish transgenic and endogenous granzyme B) with or without HS2 were also made. Splenocytes from transgenic mice containing HS2 resulted in a 10-fold greater amount of transgene mRNA when compared with transgenic mice lacking this element. Since this element confers position independence of granzyme B expression in vivo, these features are most consistent with that of a LCR.

The interpretation of the findings described here is informed by many studies of the β-globin cluster LCR elements. Similar to the β-globin genes, the granzyme genes are structurally organized in clusters and expression of genes within these clusters are temporally regulated (36). In WT CTLs, we have observed the delayed
expression of granzyme C relative to granzyme B in both T and NK cells (Fig. 10). The two granzyme B-deficient mouse strains that were developed in our laboratory have provided some clues into how this differential expression occurs. The retained PGK-neo cassette in GzmB+/−/− cluster mice is thought to function as a transcriptional “sink” within that locus, thereby diminishing the transcriptional activity of downstream granzymes C and F. When the PGK-neo cassette is deleted in GzmB+/−/− mice, the interaction between the 5′ LCR and the PGK promoter is presumably lost, causing the LCR to scan downstream to the next gene, granzyme C. Knockout mouse models and analysis of regulatory components upstream of these genes will be required to confirm these hypotheses.

Granzyme C and its human ortholog, granzyme H, can potently induce cell death via mechanisms that are distinct from that of granzyme B (11, 12). The diversity of death pathways targeted by granzymes allows the host to respond to a variety of immunological challenges. In the setting of granzyme B deficiency or inhibition, as has been demonstrated during certain viral infections, persistent antigenic stimulation may cause the delayed activation of downstream granzymes that act via novel mechanisms to remove the threat (37). The gene regulatory mechanisms controlling this late switch in granzyme gene expression were serendipitously uncovered by deleting the AvrII fragment in the granzyme B gene. These findings provide an important clue for the location of the critical elements and will be the subject of future studies.

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