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Truncated Form of TGF-βRII, But Not Its Absence, Induces Memory CD8+ T Cell Expansion and Lymphoproliferative Disorder in Mice

Harumichi Ishigame,* Munir M. Mosaheb,*-1 Shomyseh Sanjabi,‡,‡ and Richard A. Flavell*,-8

Inflammatory and anti-inflammatory cytokines play an important role in the generation of effector and memory CD8+ T cells. We used two different models, transgenic expression of truncated (dominant negative) form of TGF-βRII (dnTGFβRII) and Cre-mediated deletion of the floxed TGF-βRII to examine the role of TGF-β signaling in the formation, function, and homeostatic proliferation of memory CD8+ T cells. Blocking TGF-β signaling in effector CD8+ T cells using both of these models demonstrated a role for TGF-β in regulating the number of short-lived effector cells but did not alter memory CD8+ T cell formation and their function upon Listeria monocytogenes infection in mice. Interestingly, however, a massive lymphoproliferative disorder and cellular transformation were observed in Ag-experienced and homeostatically generated memory CD8+ T cells only in cells that express the dnTGFβRII and not in cells with a complete deletion of TGF-βRII. Furthermore, the development of transformed memory CD8+ T cells expressing dnTGFβRII was IL-7- and IL-15-independent, and MHC class I was not required for their proliferation. We show that transgenic expression of the dnTGFβRII, rather than the absence of TGF-βRII-mediated signaling, is responsible for dysregulated expansion of memory CD8+ T cells. This study uncovers a previously unrecognized dominant function of the dnTGFβRII in CD8+ T cell proliferation and cellular transformation, which is caused by a mechanism that is different from the absence of TGF-β signaling. These results should be considered during both basic and translational studies where there is a desire to block TGF-β signaling in CD8+ T cells.

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level of TGF-β signaling. Because of the early-onset lethal autoimmune disease development in CD4Cre-Tgβbr2+/− mice, the dnTGFβRII mice have been used to study the importance of TGF-β signaling in the regulation of effector CD8+ T cell function during autoimmunity and tumor development (20, 21). In fact, these promising results have prompted the idea of overexpressing the dnTGFβRII in CD8+ T cells as a means to enhance the activity of tumor-specific CD8+ T cells toward a feasible strategy for T cell-mediated immunotherapy (22, 23). Furthermore, TGF-β controls the number of CD8+ effector cells that are generated in response to bacterial and viral infections (24, 25). Blockade of TGF-β signaling in this model increased the number of SLECgs by regulating Bcl2 expression induced by IL-15 (24). In addition, homeostatic proliferation of CD8+ memory T cells is also thought to be regulated by TGF-β under steady-state conditions (26, 27). Mice expressing dnTGFβRII under the control of the Cd4 promoter do not develop autoimmunity, but instead develop polyclonal memory-like CD8+ T cell hyperproliferation that eventually leads to cellular transformation (26, 28, 29). However, Cd4 promoter-driven dnTGFβRII mice on a polyclonal background do not exhibit dysregulated expansion of the CD8+ T cells before development of autoimmunity (20). Thus, it is still not clear how TGF-β signaling regulates CD8+ T cell homeostasis as well as peripheral CD8+ T cell lymphoma development. Furthermore, because dnTGFβRII cells can still respond to TGF-β signaling, the exact role of TGF-β signaling in CD8+ T cell function, especially its contribution to memory CD8+ T cell formation and homeostatic maintenance, remains unknown.

In this study, we used two different mouse models, Tg mice expressing dnTGFβRII under the control of Cd4 promoter and Tgβbr2 conditional KO mice crossed to either granzyme B (GzmB)− or Cd4-Cre mice, to address the role of TGF-β signaling in either model. Interestingly, however, we observed a massive lymphoproliferative disorder in both Ag-experienced and homeostatically produced memory phenotypic CD8+ T cells only in the dnTGFβRII model, suggesting that this previously observed phenotype may be caused by the expression of the truncated form of TGF-βRII, rather than the absence of TGF-βRII-mediated signaling, in memory CD8+ T cells.

Materials and Methods

Mice

CD4-dnTGFβRII, Cd11c-dnTGFβRII, and Cd4Cre mice on the C57BL/6 background were previously described (20, 30, 31). Rag1−/−, b2m−/−, OtI−, OtII, and P14 mice were obtained from The Jackson Laboratory, C57BL/6 and CD4Cre-Ly5.2/Cr (CD45.1) congenic mice were obtained from the National Cancer Institute. R15−/− mice were purchased from Taconic. Tgβbr2+/-, Il7−/−, and GzmB-Cre mice were kindly provided by H. Shen (University of Pennsylvania). Memory OTI-WT-Rag1−/− or OTI-dnTGFβRII-Rag1−/− cells were isolated from LNs and spleen of LM-OVA-infected mice based on staining with CD8α and the corresponding congenic markers. For recall experiments, mice were infected i.v. with 5 × 106 of the LM-OVA.

Flow cytometry

Fluorescent dye-labeled Abs against CD8α, TCRβ, CD45.1, CD45.2, CD43 (1B11), CD62L, CD69, and CD25 were purchased from BioLegend; CD127, KLRG1, and CD122 were from eBioscience; IFN-γ, IL-2, and TNF were from BD Biosciences; and GzmB was from Caltag Laboratories. For intracellular cytokine staining, splenocytes from infected mice were stimulated with SIINFEKL peptide in the presence of Golgi-stop (BD Biosciences) for 4 h, followed by fixation and permeabilization according to the manufacturer’s instructions (BD Biosciences). For cell division assay, purified memory OTI cells were stained with CFSE (Molecular Probes) and cultured in the presence of anti-CD3, PMA/ionomycin, IL-2, or IL-15 for 2–4 d. The cells were analyzed on an LSRII or FACS-Calibur (Becton Dickinson), and data were analyzed with FlowJo software (Tree Star).

In vivo CTL assay

OTI cells of each genotype were FACS sorted based on their congenic markers and mixed with CFSE-labeled target cells at 1:5 ratio. The mixture of effector and target cells was injected i.v. into naïve C57BL/6 hosts. Spleens from the recipient mice were harvested 14–16 h later, and the ratio of CFSE− and CFSE+ cells was measured by FACS analysis, and in vivo CTL activity was determined as described previously (35).

Isolation of lymphocytes from nonlymphoid organs

PBLs were isolated from tail or retro-orbital vein of infected mice. Liver and lung homogenates were digested for 1 h at 37°C with digestion buffer (RPMI + 5% FBS + 100 U/ml DNase I [Sigma] + 0.2 mg/ml Collagenase IV [Sigma]). Liver homogenates were centrifuged at 300 rpm for 3 min to remove hepatocytes, and lung homogenates were run through a 70-μm filter mesh. Nonhepatic supernatants and lung lymphocytes were centrifuged at 1500 rpm for 10 min. The cell pellet was resuspended in 1 ml RPMI + 5% FBS and mixed with 4 ml of 27.5% OptiPrep (Axis-Shield). To make a gradient, we layered 1 ml RPMI on top of the cells and centrifuged it at 2000 rpm for 20 min. Lymphocytes were removed from the interface of the gradient.

Real-time RT-PCR analysis

Total RNA was extracted with Qiagen mRNAEasy kit (Qiagen) according to the manufacturer’s instructions. RNA was reverse transcribed with the High Capacity cDNA Reverse Transcription Kit (Applied Biosystems). The cDNA was subsequently used for quantitative PCR on an ABI Prism instrument using commercial primer-probe sets.

Karyotype analysis

Memory OTI-dnTGFβRII cells were isolated from LNs and spleen of LM-OVA-infected mice, and stimulated with PMA and ionomycin for 2 d. Twenty mitoses were subjected to G-banding analysis (Laboratory of Genome Biology and Cytogenetics, Van Andel Institute).

Statistical analysis

Unpaired Student t test was used to calculate the statistical significance of data comparison. A p value <0.05 was considered to be significant.

Results

Conditional deletion of Tgβbr2 in OTI cells results in increased number of SLECs but does not lead to increased accumulation of total effector cells during acute Listeria infection

Using OTI cells expressing dnTGFβRII driven by the Cd4 promoter, we have shown that TGF-β signaling in CD8+ T cells controls the number of SLECs during acute LM-OVA infection (24). In this model, OTI cells isolated from dnTGFβRII-Rag1−/− mice cannot be adoptively transferred into wild-type (WT) C57BL/6 recipients because of immunological rejection (24, 36). This problem is overcome by using CD11c-dnTGFβRII mice as hosts, as they also express human dnTGFβRII under the control of Cd11c promoter (30). As reported previously (24), effector OTI-dnTGFβRII cells expanded 2- to 3-fold more than OTI-WT cells by days 7 and 14 postinfection (p.i.), and contracted normally by day 21 p.i.
(Fig. 1A, 1B). An increased percentage of SLEC (KLRC3/CD127lo) in OTI-dnTGFβRII cells was also observed during the effector phase of LM-OVA infection (Fig. 1C).

Because of partial blockade of TGF-β signaling and the problem of immunological rejection in dnTGF-βRII cells (18, 20), we used the TGF-βRII conditional KO system as an alternate method to block TGF-β signaling in CD8+ T cells to study the effect of TGF-β on formation and function of memory CD8+ T cells. To delete TGF-βRII in CD8+ T cells, we crossed Tgfr2f/f mice to OTI-Rag1−/−, as well as to mice expressing Cre recombinase driven by the Cd4 promoter (Cd4Cre-Tgfr2f/f), and we used their littersmates (Cd4Cre-Tgfr2f/f or Cd4Cre-negative Tgfr2f/f) as controls. We adoptively cotransferred OTI-control or OTI-Cd4Cre-Tgfr2f/f cells into naive C57BL/6 mice and infected them with LM-OVA. The percentage of transferred cells was measured over the course of infection. We expected that the phenotype of Cd4Cre-Tgfr2f/f cells would be much stronger than that of dnTGFβRII cells. However, in contrast with OTI-dnTGFβRII cells (Fig. 1A, 1B), the expansion of effector OTI-Cd4Cre-Tgfr2f/f cells was comparable with control OTI cells during effector phase of acute LM-OVA infection (Fig. 1D, 1E). By contrast, similar to what was observed in OTI-dnTGFβRII cells (Fig. 1C), the OTI-Cd4Cre-Tgfr2f/f cells had a higher percentage of SLECs and lower percentage of MPECs (Fig. 1F). These data suggest that dnTGFβRII cells and conditional TGF-βRII KO cells have a different phenotype with regard to total effector T cell expansion but show similarly increased percentage of SLECs.

Late memory OTI-dnTGFβRII cells show enhanced accumulation after acute Listeria infection

Using the dnTGFβRII model, we next wanted to study the role of TGF-β signaling in the formation and function of memory CD8+ T cells. OTI-WT cells were either transferred alone into C57BL/6 mice or cotransferred together with OTI-dnTGFβRII cells into CD11c-dnTGFβRII hosts. At early time points, days 25–101 p.i., the frequency of transferred OTI-WT or OTI-dnTGFβRII cells among PBLs ranged from 0.01 to 8.73% in C57BL/6 or CD11c-dnTGFβRII hosts (Fig. 2A). Interestingly, by about day 300 p.i., CD11c-dnTGFβRII hosts that still retained OTI cells showed a dramatically elevated frequency of OTI-dnTGFβRII cells (Fig. 2A, 2B). The time at which this expansion could first be detected varied among the animals and for each experiment, and ranged from day 180 to 250 p.i. (data not shown). OTI-WT cells transferred into the CD11c-dnTGFβRII hosts behaved similarly to those transferred into C57BL/6 hosts, and the percentage of OTI-WT cells in PBLs was only slightly increased in either host (Fig. 2A). However, we observed that the engraftment efficiency of transferred OTI-WT cells in CD11c-dnTGFβRII hosts was significantly lower when compared with their engraftment into C57BL/6 hosts at 28–40 d p.i. (Supplemental Fig. 1A). Although the exact reason for this lower engraftment of the OTI-WT cells in the CD11c-dnTGFβRII hosts is currently unclear, we hypothesize that this may be because of enhanced innate immunity in CD11c-dnTGFβRII hosts (30, 37) and a potential NK-mediated rejection of the OTI-WT cells over time. This long-term rejection of engrafted OTI-WT memory cells in CD11c-dnTGFβRII hosts is different from the early rejection of the OTI-dnTGFβRII cells in C57BL/6 hosts, which occurs within days as previously reported (24, 36) (Supplemental Fig. 1B, 1C). Because the long-term engraftment of OTI-WT cells was considerably reduced in CD11c-dnTGFβRII hosts, we had to use C57BL/6 hosts that received OTI-WT cells as our age-matched controls in most of the subsequent long-term experiments.

Recipient mice were sacrificed at days 374–448 p.i. to measure the number of late memory OTI cells in different tissues. The CD11c-dnTGFβRII hosts that had received OTI-dnTGFβRII cells showed splenomegaly and lymphadenopathy (Fig. 2C). The percentage of OTI-dnTGFβRII cells was significantly higher than that of OTI-WT cells in both lymphoid and nonlymphoid tissues (Fig. 2D), and >20-fold more OTI-dnTGFβRII cells were recovered from each organ than OTI-WT cells (Fig. 2E). Unlike the transferred OTI-dnTGFβRII cells, the frequency of host CD8+ T cells from the spleen of CD11c-dnTGFβRII host did not increase over time (days 374–448 p.i.; Fig. 2F), even though they had the CD44hi CD62Llo activated phenotype (Fig. 2G), suggesting that the OTI-dnTGFβRII phenotype in the CD11c-dnTGFβRII host is unique to the donor OTI cells that express the dnTGFβRII transgene. Analyses of cell-surface markers revealed that OTI-dnTGFβRII cells were phenotypically identical to memory OTI-WT cells before

![FIGURE 1](http://www.jimmunol.org/ Downloaded from by guest on April 12, 2017)

**FIGURE 1.** Conditional deletion of Tgfr2 in OTI cells results in increased number of SLECbs but does not lead to increased accumulation of total effector cells during acute *Listeria* infection (A) Naive OTI-WT and OTI-dnTGFβRII cells on different CD45 congenic backgrounds were adoptively cotransferred into naive CD11c-dnTGFβRII hosts, followed by infection with LM-OVA 1 d later. The percentage of OTI-WT and OTI-dnTGFβRII cells was monitored among the PBLs of infected mice over time. (B) The ratio of OTI-dnTGFβRII over OTI-WT cells in PBLs. (C) The percentage of OTI-WT and OTI-dnTGFβRII SLECs in the blood after LM-OVA infection at the indicated time point (left panel). Representative plots of the expression of CD127 and KLRG1 on blood OTI-WT and OTI-dnTGFβRII cells at day 7 p.i. (right panel). (D) The percentage of OTI-control (CD45.1.1) and OTI-Cd4Cre-Tgfr2f/f cells among PBLs was monitored over time. (E) The percentage of OTI-Cd4Cre-Tgfr2f/f over OTI-control cells in PBLs. (F) The percentage of SLECs in the blood of OTI-control and OTI-Cd4Cre-Tgfr2f/f cells after LM-OVA infection at the indicated time point. Representative plots of the expression of CD127 and KLRG1 in blood OTI-WT or OTI-Cd4Cre-Tgfr2f/f cells at day 5 p.i. (right panel). Each dot represents one mouse. Data are representative of at least three independent experiments. *p < 0.05, **p < 0.01, ***p < 0.001 versus control.
they exhibited lymphoproliferation (data not shown). However, although the expression of CD25, CD122 (IL-2Rβ/IL-15Rβ) and CD127, KLRG1, CD27, CD69, and CD44 was not different, the expanded memory OTI-dnTGFβRII cells showed reduced CD43, CD8α, and CD62L expression when compared with OTI-WT cells at ∼300 d.p.i. (Fig. 2H). As previously reported (29), these results demonstrate that overexpression of dnTGFβRII results in accumulation of dysregulated CD8+ T cells that phenotypically resemble memory CD8+ cells.

Deletion of Tgfb2 does not lead to dysregulated expansion of late memory OTI cells upon Listeria infection

To eliminate any potential effect of TGF-β during T cell development and priming, and to further focus on the role of TGF-β signaling on activated and memory CD8+ T cells, we crossed OTI-dnTGFβRII and OTI-CD4Cre-Tgfb2δ cells into naive C57BL/6 hosts, followed by infection with LM-OVA 1 d later. C57BL/6 mice receiving naive OTI-WT were also used as controls. The percentage of OTI-WT and OTI-dnTGFβRII after contraction was monitored in PBLs over time. (B) Representative dot plots of transferred OTI-WT (top panel) and OTI-dnTGFβRII (bottom panel) cells in the PBLs at day 307 p.i. (C) Gross morphology of spleen and LNs from infected hosts at 11 mo p.i. (D and E) The percentage (D) and absolute number (E) of donor OTI cells in the indicated tissues at days 374–448 p.i. Results are combined from four independent experiments. (F) The percentage of host CD8+ T cells in the spleen of the same mice as in (D). (G) Representative dot plots of donor OTI (CD45.1+) and host CD8+ T cells (CD45.1+) in the spleen at day 374 p.i. (top panel). Expression of CD44 and CD62L in the spleen of host CD8+ T cells (bottom panel). (H) Expression pattern of indicated surface cell markers was analyzed on OTI-WT and OTI-dnTGFβRII cells isolated from spleen of LM-OVA-infected mice at day 340 p.i. (I) Naive OTI-control and OTI-GzmB-Cre-Tgfb2δ cells on different congenic backgrounds were adoptively cotransferred into naive C57BL/6 hosts, followed by infection with LM-OVA 1 d later. The percentage of OTI-control and OTI-GzmB-Cre-Tgfb2δ cells in PBLs was monitored over time at indicated time points. (J) The percentage of SLECs, MPECs, and KLRG1+CD127+ (double positive [dp]) cells among the OTI-control or OTI-GzmB-Cre-Tgfb2δ cells in the PBLs at 280 d.p.i. Representative plots of the expression of CD127 and KLRG1 at day 280 p.i. (right panel). (K) The percentage of OTI-control and OTI-GzmB-Cre-Tgfb2δ cells in the spleen at day 438 p.i. (A, B, G–K) Data are representative of three independent experiments. Each dot represents one mouse. *p < 0.05, **p < 0.01, ***p < 0.001 versus OTI-WT. NS, not significant versus OTI-WT.
were bled at indicated time points and CFSE dilution was measured. Data are representative of two independent experiments with similar results. 

Development and proliferation of late memory OTI-dnTGFβRII cells is not dependent on IL-7 or IL-15 

Given that IL-7 and IL-15 are important cytokines for memory CD8+ T cell homeostasis, we next examined whether dysregulated expansion of OTI-dnTGFβRII cells is dependent on IL-7 or IL-15 signaling. We previously reported that effector OTI-dnTGFβRII cells are largely dependent on IL-15 signaling (24). However, we found that a small fraction of effector OTI-dnTGFβRII cells could differentiate into memory cells in Il7−/− or Il7−/−/Il15+/+ CD11c-dnTGFβRII hosts, and they eventually massively expanded at 225 d.p.i. (Fig. 3A, 3B). OTI-WT cells were hardly detected in the CD11c-dnTGFβRII cotransfer setting (Fig. 3A, 3B), possibly as a result of out-competition of OTI-dnTGFβRII cells and reduced engraftment of OTI-WT cells (Supplemental Fig. 1D). To further examine the role of IL-15 in OTI-dnTGFβRII cell expansion, we isolated these dysregulated memory OTI-dnTGFβRII cells 370 d after LM-OVA immunization, labeled them with CFSE, and retransferred them into Il7+/+ or Il7−/−/Il15+/+ CD11c-dnTGFβRII hosts. The proliferation of donor cells was assessed by cell division–induced loss of CFSE fluorescence. Memory OTI-dnTGFβRII cells proliferated similarly in IL-15–deficient and –sufficient hosts over a 2-wk period (Fig. 3C). These results suggest that in contrast with previous reports, IL-15 is not required for the development, survival, and dysregulated proliferation of late memory OTI-dnTGFβRII cells (29).

To determine whether IL-7 plays a critical role in memory OTI-dnTGFβRII cell dysregulated expansion, we cotransferred OTI-WT and OTI-dnTGFβRII cells into IL7−/− CD11c-dnTGFβRII or IL7−/−/CD11c-dnTGFβRII hosts. Interestingly, effector OTI-dnTGFβRII cells expanded at a much higher magnitude compared with OTI-WT cells in the IL-7–deficient environment than in the IL-7–sufficient environment (Fig. 4A). An increased percentage of SLECs in OTI-dnTGFβRII cells was also observed in IL-7–deficient hosts at days 7 and 12 p.i. (Fig. 4B). Furthermore, similar to the results with Il7−/−/CD11c-dnTGFβRII hosts (Fig. 3A), effector OTI-dnTGFβRII cells differentiated into memory T cells and eventually exhibited dysregulated expansion in Il7−/− CD11c-dnTGFβRII mice, and this expansion occurred sooner than in Il7+/+ hosts (Fig. 4C, 4D; compare day 146 p.i.). In addition, when CFSE-labeled dysregulated memory OTI-dnTGFβRII cells were transferred into Il7−/− CD11c-dnTGFβRII or Il7−/−/CD11c-dnTGFβRII host as performed in Fig. 3C, the proliferation of donor cells was accelerated in IL-7–deficient hosts, most likely because of the lymphopenic environment of the Il7−/− hosts (Fig. 4E) (32). Overall, these results suggest that IL-7 is also not essential for the development and dysregulated proliferation of late memory OTI-dnTGFβRII cells.

Dysregulated memory OTI-dnTGFβRII cells can proliferate independent of MHC class I 

Memory CD8+ T cells can persist in the absence of Ag (1, 3, 4). To examine whether proliferation of dysregulated memory OTI-dnTGFβRII cells is dependent on MHC class I, we isolated massively accumulated memory OTI-dnTGFβRII cells from infected CD11c-dnTGFβRII hosts, as performed in Fig. 3C, and transferred them (2 × 10^6) cells into sublethally irradiated C57BL/6 or b2m−/− mice, which are deficient in the expression of functional MHC class I. At day 30 after transfer, the frequency of transferred OTI-dnTGFβRII cells in both spleen and LN was greater in b2m−/− mice most likely because of the lymphopenia caused by the absence of host CD8+ T cells (Fig. 5A, 5B). In addition, dysregulated memory OTI-dnTGFβRII cells proliferated extensively in both irradiated C57BL/6 and b2m−/− mice (Fig. 5C), indicating that dysregulated memory OTI-dnTGFβRII do not need MHC class I to survive and proliferate.

OTI-dnTGFβRII cells, but not OTI-CD4Cre-Tgfbr2fl/fl cells, show dysregulated expansion under steady-state condition 

C57BL/6 mice expressing dnTGFβRII driven by the Cd2 promoter developed polyclonal memory-like CD8+ T cell hyperproliferation under steady-state condition without developing autoimmunity (29). However, our dnTGFβRII mice on the Rag1−/− background did not show this phenotype as they develop colitis so rapidly it was not possible to assess this (20). However, we did observe the hyperproliferation phenotype in pathogen-induced memory OTI-dnTGFβRII-Rag1−/− cells (Fig. 2). To eliminate the autoimmune phenotype in dnTGFβRII mice, as well as the complications associated with the CD11c-dnTGFβRII hosts, we analyzed the

![FIGURE 3.](http://www.jimmunol.org/Downloadedfrom)
and CFSE dilution was measured. Each dot represents one mouse. Data are representative of two independent experiments. **p < 0.01, ***p < 0.001.

The expression of CD44 on OTI cells was analyzed in OTI-dnTGFβRII-Rag1<sup>−/−</sup> and OTI-CD4Cre-Tgfb2<sup>−/−</sup>-Rag1<sup>−/−</sup> mice under steady-state conditions, in which endogenous TCRs are excluded and autoimmunity was eliminated. Despite the fact that OTI-dnTGFβRII-Rag1<sup>−/−</sup> mice do not express the cognate OVA Ag, a large proportion (35–95%) of OTI-dnTGFβRII cells isolated from LNs and spleen were CD44<sup>hi</sup> memory phenotype even at 32 wk of age (Fig. 6C). In contrast with OTI-dnTGFβRII cells, more than half of OTI-CD4Cre-Tgfb2<sup>−/−</sup> cells, similar to control cells, still retained CD44<sup>lo</sup> naive phenotype at 11–25 wk of age (Fig. 6B). Interestingly, OTI-dnTGFβRII-Rag1<sup>−/−</sup> mice displayed a considerable increase in the absolute number of OTI cells in both LNs and spleen by 12–32 wk of age (Fig. 6C). Most of the expanded OTI cells were CD44<sup>hi</sup> memory-like cells, but CD44<sup>lo</sup> naive phenotype cells were also expanded in some of the animals (Fig. 6A, 6C). Furthermore, the dysregulated dnTGFβRII cell expansion is not OTI-Tg line specific, as it was also seen in lymphocytic choriomeningitis virus–specific CD8<sup>+</sup> P14-dnTGFβRII cells (Supplemental Fig. 2A–D). As expected, consistent with the late memory OTI cell phenotype, this dysregulated expansion of OTI cells was detected only when TGF-β signaling was blocked by overexpression of dnTGFβRII, and not when the whole receptor was deleted (Fig. 6D). Indeed, no significant difference in the absolute number of OTI cells in the LNs and spleen was observed between control and OTI-CD4Cre-Tgfb2<sup>−/−</sup>-Rag1<sup>−/−</sup> mice (Fig. 6D), further demonstrating that the deletion of the TGF-βRII itself, at the same time as when the dnTGFβRII would be expressed (thymic double-positive stage), does not lead to dysregulated expansion of memory-like OTI cells, even though a fraction of naive OTI-CD4Cre-Tgfb2<sup>−/−</sup> cells do become CD44<sup>hi</sup> over time (Fig. 6B).
Abrogation of TGF-βRII Signaling in CD8+ T Cells

Rapid differentiation of naive OTI-dnTGFβRII cells into CD44hi memory-like cells occurs even in the absence of endogenous Tgfr2 gene

Because the effects of TGF-β signaling in T cell differentiation are context dependent (38), it is possible that a low level of leaked TGF-β signaling in OTI-dnTGFβRII cells promotes CD44hi memory CD8+ T cell differentiation, which then somehow causes progressive expansion. To examine this possibility, we crossed OTI-dnTGFβRII-Rag1+/− mice with CD4Cre-Tgfr22f−/− mice to completely delete TGF-β signaling in naive OTI cells. We still found an increased percentage of CD44hiCD122hi memory-like cells in the LNs of OTI-dnTGFβRII-Rag1+/− × CD4Cre-Tgfr22f−/− mice when compared with OTI-CD4Cre-Tgfr22f−/− cells (Fig. 6E), suggesting a dominant effect of truncated form of TGF-βRII on CD8+ T cell expansion that is independent of blocking TGF-β signaling in these cells.

Memory OTI-dnTGFβRII cells develop lymphoma

Because memory-like CD8+ T cells that express dnTGFβRII under the control of the Cd2 promoter have been shown to develop lymphoma (28), we analyzed whether overexpression of dnTGFβRII by Cd4 promoter also causes cellular transformation. Dysregulated memory-like OTI-dnTGFβRII cells or Listeria-induced memory OTI-dnTGFβRII cells were isolated from 6-mo-old OTI-dnTGFβRII-Rag1+/− mice or CD11c-dnTGFβRII hosts at day 420 p.i., respectively, and karyotype analysis was performed. These cells showed chromosomal abnormality, including trisomy of chromosome 15 and reciprocal translocation between chromosomes 4 and 15 (Fig. 7A, Table I). Gain of chromosome 15 is frequently present in several mouse models of lymphoma (39–41), and it has been suggested that Myc proto-oncogene located on mouse chromosome 15 contributes to trisomy 15 and leukemic transformation (42). We therefore analyzed Myc expression in memory-like OTI-dnTGFβRII cells from 2- to 3-mo-old OTI-dnTGFβRII-Rag1+/− mice before they exhibited dysregulated expansion. We found that although the expression level of Myc target genes, such as Id2 (inhibitor of DNA binding 2) (43) and Cdkn1a (cyclin-dependent kinase inhibitor 1A; also called p21) (44), were increased in CD44hi memory-like OTI-dnTGFβRII cells, Myc expression did not differ between these groups (Fig. 7B). However, when we analyzed dysregulated memory-like OTI-dnTGFβRII cells isolated from OTI-dnTGFβRII-Rag1+/− mice at 6 mo of age, Myc expression was increased in memory-like OTI-dnTGFβRII cells compared with that of OT-I WT cells (Fig. 7C). These results indicate that dysregulated memory-like OTI-dnTGFβRII cells can progress to overt lymphoma.
suggest that overexpression of dnTGFβRII in OTI cells by Cd4 promoter results in cellular transformation most likely through an Myc-mediated mechanism.

**Transformed memory OTI-dnTGFβRII cells produce effector cytokines and have CTL activity but cannot proliferate after recall response**

To determine whether transformed memory OTI-dnTGFβRII cells retain their functionality as memory cells, we assessed the capacity of these cells to produce effector cytokines and to kill Ag-sensitized target cells in vivo. We isolated memory OTI-WT cells or transformed memory OTI-dnTGFβRII cells from infected C57BL/6 or CD11c-dnTGFβRII mice at day 380 p.i., respectively, and restimulated them with SIINFEKL peptide ex vivo. Interestingly, the transformed memory OTI-dnTGFβRII cells were functional in terms of IFN-γ, TNF, and GzmB production, when compared with memory OTI-WT cells (Fig. 8A). Furthermore, when the same number of each memory OTI population was retransferred into naive hosts along with SIINFEKL-pulsed target cells, no difference was observed in the in vivo ability of OTI-dnTGFβRII cells to kill target cells when compared with memory OTI-WT cells (Fig. 8B), suggesting that they also retain their cytolytic activity. Similar results were also obtained when we used early memory OTI-WT and OTI-dnTGFβRII cells that had not yet exhibited dysregulated expansion (Supplemental Fig. 3A, 3B). Similarly, early memory OTI-Cd4Cre-Tgfbr2<sup>fl/fl</sup> cells exhibited a normal ability to produce IL-2 and IFN-γ after SIINFEKL peptide stimulation ex vivo (Supplemental Fig. 4A). Interestingly, however, transformed memory OTI-dnTGFβRII cells proliferated less than memory OTI-WT cells in response to SIINFEKL peptide or anti-CD3 ex vivo, whereas they responded similarly to OTI-WT cells to PMA/ionomycin stimulation (Fig. 8C). Furthermore, transformed memory OTI-dnTGFβRII cells did not respond to IL-2 or IL-15 stimulation (Fig. 8D), whereas early memory OTI-dnTGFβRII cells isolated from the hosts at day 70 p.i. proliferated similarly to OT-WT cells in response to these cytokines (Supplemental Fig. 3C). To determine the ability of transformed memory OTI-dnTGFβRII cells to respond to a secondary infection in vivo, we isolated transformed memory OTI-dnTGFβRII cells and memory OTI-WT cells at day 461 p.i., and retransferred the same number of cells from each into naive hosts followed by LM-OVA infection. Consistent with results from ex vivo experiments, OTI-dnTGFβRII cells responded poorly to a secondary infection, whereas memory OTI-WT cells mounted a proliferative response (Fig. 8E). In contrast, no difference in proliferative response was observed when we transferred memory OTI-WT and OTI-dnTGFβRII cells isolated 180 d p.i. that had not exhibited dysregulated expansion (Supplemental Fig. 3D). The early memory OTI-control and OTI-GzmB- or CD4-Cre TGFβRII in OTI cells by Cd4 signaling on memory CD8<sup>+</sup> T cells does not impact their function and recall response. In addition, although transformed late memory OTI-dnTGFβRII cells are capable of producing effector cytokines and killing target cells, they have defects in Ag- or cytokine-induced proliferative responses. The consequence of long-term overexpression of dnTGFβRII should be significantly considered for studies where the overexpression of this construct is used for enhancing T cell function during T cell–mediated immunotherapy.

**Discussion**

In this study, we used two different mouse models, Tg mice expressing truncated form of TGF-βRII under the Cd4 promoter and TGF-βRII conditional KO mice crossed to GzmB- or CD4-Cre mice to address the role of TGF-β signaling in memory CD8<sup>+</sup> T cell development, function, and homeostatic proliferation. We demonstrate that Ag-induced expansion and cytokine production of memory CD8<sup>+</sup> T cells is not effected by the blockade of TGF-β signaling in both genetic models. Interestingly, however, we observe a massive lymphoproliferative disorder in both Ag-experienced

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**Table I. Karyotype analysis of OTI-dnTGFβRII cells**

<table>
<thead>
<tr>
<th>ID</th>
<th>Cell Types</th>
<th>Sampling</th>
<th>Karyotype</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Memory-like OTI</td>
<td>6 mo old</td>
<td>41,XY +15 [17/30]&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>40,XY [13/30]</td>
</tr>
<tr>
<td>2</td>
<td>Memory-like OTI</td>
<td>6 mo old</td>
<td>41,XY +15 [8/30]</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>40,XY [22/30]</td>
</tr>
<tr>
<td>3</td>
<td><em>Listeria</em>-induced memory OTI</td>
<td>420 d.p.i.</td>
<td>40,XX [3/20]</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>41,XX +15 [15/20]</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>40,XX Der(4)/T(4E;15A2) [2/20]</td>
</tr>
</tbody>
</table>

<sup>a</sup>Number of chromosomes with aberration/total number of chromosomes studied.
and homeostatically generated memory CD8+ T cells in the Tg model only, suggesting that the truncated form of TGF-βRII has functions that are independent of endogenous TGF-βRII signaling. This study has therefore revealed a previously unrecognized function of truncated TGF-βRII in CD8+ T cell differentiation and proliferation, which should be considered during the potential therapeutic use of the dnTGFβRII as a means to enhance Ag-specific CD8+ T cell function.

Blockade of TGF-β signaling by overexpressing dnTGFβRII using a tissue-specific promoter has been widely used to study the role of TGF-β signaling in vivo. We expected that conditional deletion of Tgfb2 in CD8+ T cells would result in a much stronger phenotype compared with CD8+ dnTGFβRII cells, because the latter still express a functional TGF-βRII that can induce some level of TGF-β signaling (18, 20). However, we found that memory OTI-dnTGFβRII cells expanded massively and developed a lymphoproliferative disorder both in Listeria-infected hosts and in immunized OTI-dnTGFβRII-Rag1−/− mice, whereas the OTI-conditional TGF-βRII KO cells lacked this phenotype. These results indicate that dnTGFβRII itself has functions independent of the endogenous TGF-βRII in CD8+ T cells. In support of this, increased differentiation of naive OTI cells into CD4+ memory-like T cells was still observed in OTI-dnTGFβRII × CD4Cre-Tgfb2f/f mice on the Rag1−/− background. Previous reports showed that immunocompetent mice expressing dnTGFβRII driven by Cd2 promoter exhibited dysregulated memory CD8+ T cell homeostasis and cellular transformation without any sign of autoimmune disease (26), whereas the dnTGFβRII-Rag1+/− strain driven by Cd4 promoter developed autoimmune disease before any obvious signs of memory CD8+ T cell expansion (20). By contrast, a recent study demonstrated that lymphoproliferative disorder can also be seen in the dnTGFβRII-Rag1+/− strain driven by Cd4 promoter, in which dnTGFβRII are homozygously expressed, and IL-6 or IL-12p40 was also deleted to prevent autoimmune disease development (45). The reason for the differences observed between the two dnTGFβRII strains is unknown, but it is suggested that differential expression of the dnTGFβRII transgene may affect the efficiency of TGF-β signaling blockade. However, we showed in this study that dnTGFβRII phenotype is not recapitulated in the conditional KO system, suggesting that lymphoproliferative disorder seen by increasing copy number of dnTGFβRII is not simply due to increased suppression of TGF-β signaling. In fact, dysregulated memory OTI-dnTGFβRII cell homeostasis was observed despite the use of heterozygous OTI-dnTGFβRII-Rag1−/− mice in this study. In this regard, overexpression of dnTGFβRII in a tumor cell line has been shown to impair both TGF-β1 and BMP2 signaling (46). Thus, it is possible that dnTGFβRII interacts with receptors other than TGF-β and also blocks TGF-β-independent signaling pathway such as BMP signaling, which may have synergistic effects on proliferation and transformation of memory CD8+ T cells. Further studies, outside the scope of this study, are required to identify such a dnTGFβRII-binding ligand, the composition of the receptor complex, as well as the downstream signaling pathway(s) in CD8+ T cells.

Because the human origin of the dnTGFβRII rendered this molecule immunogenic in mice, we had to adoptively transfer the OTI-dnTGFβRII cells into the CD11c-dnTGFβRII hosts, which also express the same transgene in the NK and dendritic cell compartments, and thus are tolerant to the transgene (24, 36). The naive OTI-dnTGFβRII and OTI-WT cells highly expanded after immunization, and differentiated memory cells successfully engrafted in the CD11c-dnTGFβRII hosts; however, the OT-WT cell engraftment of memory cells diminished over time in these hosts. On the rare occasions that we were able to find engrafted OTI-WT late memory cells in the CD11c-dnTGFβRII hosts, they did not display the same expansion phenotype as the OTI-dnTGFβRII cells, and the host CD8+ T cells did not expand in this setting either. Furthermore, we also found that OTI-dnTGFβRII-Rag1−/− or P14-dnTGFβRII-Rag1−/− mice, where the dnTGFβRII transgene is expressed only in the CD8+ T cells, also showed accumulation...
of memory-like OTI or P14 cells under steady-state condition. Therefore, we believe that the expansion and transformation of late memory OTI-dnTGFβRII cells is cell intrinsic, although we cannot completely exclude the contributions from the CD11c-dnTGFβRII transgene on our findings.

We found that the kinetics of proliferation between OTI-dnTGFβRII cells isolated from Listeria-infected CD11c-dnTGFβRII host and those from uninfected OTI-dnTGFβRII-Rag1−/− mice is clearly different; in the former case, cells start expanding around 180–220 d (25–32 wk) p.i., whereas homeostatic proliferation begins around 10–12 wk of age in the latter hosts. After Listeria infection in the CD11c-dnTGFβRII hosts, the sorted naive OTI-dnTGFβRII cells clonally expand, contract, form proper memory cells, and remain quiescent until ~180 d p.i., after which they form lymphomas and begin proliferating again. Several studies have shown that CD8+ T cell clonal expansion, which is non-malignant expansion of monoclonal populations of CD8+ T cells, can appear with increasing frequency as individuals and mice age (47, 48). In fact, we did observe this phenomenon in our aged mice as the frequency of OTI-WT cells in the PBLs of C57BL/6 hosts was slightly increased over time. However, unlike the CD11c-dnTGFβRII host that received OTI-dnTGFβRII cells, mice having TEC do not show increased number of total CD8+ T cells, but rather an expansion of a subclone of the CD8+ T cells (47, 48). It has been reported that transfer of oncogene-expressing TCR-Tg T cell (OTI or P14) or Rag1−/− hosts induces lymphoma, and transformation of these cells is suppressed by cotransferring polyclonal T cells (49). This clonal competition model may explain the difference between these two experimental models; presence of polyclonal T cells in CD11c-dnTGFβRII hosts may delay proliferation and transformation of memory OTI-dnTGFβRII cells after Listeria infection.

Given the essential role of IL-15 and IL-7 in regulating memory CD8+ T cell proliferation and survival, and the critical role of these cytokine signaling pathways in lymphoma development (10, 12), we examined the involvement of these cytokines in the expansion and transformation of memory OTI-dnTGFβRII cells. However, we found that neither IL-15 nor IL-7 was required for the development and expansion of memory OTI-dnTGFβRII cells, as the OTI-dnTGFβRII cells showed enhanced accumulation even in the Il7−/− and II15−/− mice. Furthermore, we also observed that dysregulated memory-like OTI-dnTGFβRII cells transferred into Il7−/−II15−/− mice massively proliferated (data not shown). Although transformed memory OTI-dnTGFβRII cells displayed normal CD122 and CD127 expression, cytokine production, and cytotically activity, they had a poor proliferative response to IL-2 or IL-15, and exhibited a reduced capacity to participate in recall proliferative response both in vivo and in vitro, whereas early memory OTI-dnTGFβRII cells that had not yet exhibited dysregulated expansion did proliferate and responded to cytokines. These results indicate that the normal homeostatic regulation of memory CD8+ T cells becomes lost once they become lymphomas. However, naive OTI-CD2-dnTGFβRII cells proliferated extensively in Il7−/−II15−/− hosts (27), suggesting that additional cytokines or mediators may regulate the proliferation and maintenance of CD8+ T cells expressing dnTGFβRII.

TCR signaling has been shown to play an important role in lymphoma development in several murine lymphoma models (50, 51). Furthermore, it has been reported that the TCR–MHC interaction is required for the development of memory-like CD8+ CD2- dnTGFβRII T cells (27, 29). Indeed, male, but not female, HY- TCR-Tg CD2- dnTGFβRII mice on the Rag1−/− background showed an increased accumulation of CD44hi CD8+ T cells (29), and naive OTI-CD2-dnTGFβRII cells failed to expand in MHC class I−deficient host (27). However, we found that transformed memory OTI-dnTGFβRII cells could expand in an MHC class I−deficient environment, although the molecular mechanisms by which these cells can proliferate so extensively are still under investigation. It is likely that presence of dnTGFβRII in CD8+ T cells may induce cell-intrinsic alterations, such as acquisition of mutations or epigenetic alteration that affect a common signaling pathway that leads to cell proliferation. Indeed, we found chromosomal abnormalities in infection-induced late memory OTI- or memory-like OTI-dnTGFβRII cells. Upregulation of proto-oncogenes, such as Myc, and epigenetic changes may provide a mechanism by which the growth properties of memory CD8+ dnTGFβRII T cells could be altered, ultimately resulting in a CD8+ T cell expansion and transformation (45). Thus, we suggest that TCR signaling is most likely required for initial proliferation and transformation of CD8+ dnTGFβRII cells; however, it does not drive the proliferation of the lymphoma after transformation.

We previously reported that TGF-β selectively controls the number of SLECs during the effector phase of Listeria infection (24). However, in contrast with OTI-dnTGFβRII cells, this study and a recent published article showed that the total number of effector OTI-1 cells was not affected by depletion of TGF-β signaling using a conditional KO system (52). We do, however, show in this article that TGF-β signaling regulates the ratio between SLECs and MPECs; nonetheless, TGF-β signaling−deficient early memory CD8+ T cells produce a normal level of effector cytokines and mount recall responses, suggesting that TGF-β signaling is not essential for memory CD8+ T cell function during Listeria infection. The role of TGF-β signaling in CD8+ T cells in infectious diseases is still unclear. Previous studies showed that dnTGFβRII mice showed rapid viral eradication, and more functional effector and memory CD8+ T cells after chronic viral infection (25, 53), whereas therapeutic blockade of TGF-β signaling by a neutralizing Ab or small inhibitor did not recapitulate the phenotype of dnTGFβRII mice (53, 54). This discrepancy might be because of the effects of spontaneous accumulation of memory-like CD8+ T cells in dnTGFβRII mice (53). Because blockade of TGF-β signaling by dnTGFβRII results in unexpected effects in CD8+ T cells, future studies need to address the mechanism of regulation of SLEC formation by TGF-β signaling and its contribution to host protection against pathogens using conditional KO system.

In summary, we have provided evidence showing that TGF-β signaling in CD8+ T cells controls the formation of SLECs during the effector phase of Listeria infection. However, TGF-β signaling does not regulate the homeostatic proliferation of memory CD8+ T cells, and transgenic overexpression of dnTGFβRII leads to dysregulated expansion and ultimately transformation of memory CD8+ T cells. Thus, investigators should be cautious concerning the use of these different models to study the role of TGF-β signaling in CD8+ T cells and the potential use of the dnTGFβRII for therapeutic purposes. Nonetheless, identification of the dnTGFβRII-dependent alternative signaling pathway might provide a new mechanism for the regulation of memory CD8+ T cell expansion and lymphoma development.

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Disclosures
The authors have no financial conflicts of interest.
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