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*J Immunol* published online 24 October 2011
http://www.jimmunol.org/content/early/2011/10/23/jimmunol.1102202

Supplementary Material
http://www.jimmunol.org/content/suppl/2011/10/24/jimmunol.1102202.2.DC1

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Altered Differentiation, Diminished Pathogenicity, and Regulatory Activity of Myelin-Specific T Cells Expressing an Enhanced Affinity TCR

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Whereas increased affinity enhances T cell competitiveness after immunization, the role of affinity in modulating the pathogenicity of self-reactive T cells is less established. To assess this, we generated two myelin-specific, class II MHC-restricted TCR that differ only in a buried hydroxymethyl that forms a common TCR β-chain V region variant. The variation, predicted to increase TCR stability, resulted in a ∼3 log10 difference in TCR sensitivity with preserved fine specificity. The high-affinity TCR markedly diminished T cell pathogenicity. T cells were not deleted, did not upregulate Foxp3, and barring disease induction were predominantly naive. However, high-affinity CD4+ T cells showed an altered cytokine profile characterized by the production of protective cytokines prior to experimental allergic encephalomyelitis induction and decreased effector cytokines after. Further, the high-affinity TCR promoted the development of CD4+CD8− and CD8+ T cells that possessed low intrinsic pathogenicity, were protective even in small numbers when transferred into wild-type mice and in mixed chimeras, and outcompeted CD4+ T cells during disease development. Therefore, TCR affinities exceeding an upper affinity threshold may impede the development of autoimmunity through altered development and functional maturation of T cells, including diminished intrinsic CD4+ T cell pathogenicity and the development of CD4−Foxp3− regulatory populations. The Journal of Immunology, 2011, 187: 000–000.
pathogenicity, we generated a mutant self-specific TCR with minimal structural differences compared with its parental receptor, but dramatically increased sensitivity for cognate autoantigen. In a prior report, we identified a variation in the use of G and S residues at position 107 in the TCR β-chain V region (TRBV); TRBV13-2 possesses a G107, whereas most other TRBV have an S. This amino acid lies within the buried N-terminal core of the CDR3, with the side chain intercalating into the loop structure of the CDR3. We predicted based on structural modeling and molecular dynamics that substitution of the TRBV13-2 G with a S residue more typical of other TCR would position the S hydroxymethyl side chain to fill an internal gap within the CDR3 loop and through increased van der Waals interactions and H-bonding stabilize it without substantially altering its structure (20). Indeed, a G107S substitution in the TRBV13-2*1-A\textsuperscript{b}-MOG35–55–specific 1MOG9 TCR yielded a \textasciitilde 3log\textsubscript{10} increase in cognate Ag sensitivity while preserving fine specificity.

We compare in this study development and pathogenicity of T cells expressing these TCR. Either the high- or low-affinity MOG-specific T cells were well selected and predominantly naïve in the absence of immunization. However, the high-affinity TCR markedly diminished disease susceptibility. This resulted from a diminished pathogenicity of autoreactive CD4\textsuperscript{+} T cells. More prominently, the high-affinity TCR led to the development of coreceptor independent, CD4\textsuperscript{+}CD8\textsuperscript{−} (double-negative [DN]), and CD8\textsuperscript{+} T cells that showed low intrinsic pathogenicity, outcompete CD4\textsuperscript{+} T cells, and inhibited disease after transfer to wild-type (wt) mice. Therefore, high-affinity autoantigen-specific T cells can suppress autoimmune responses through both altered lineage differentiation and functional maturation. Implicitly, the pathologic potential of an autoantigen will depend on the affinity distribution of specific TCR; higher affinity clones may be protective.

Materials and Methods

Animals

C57BL/6 (B6), B6.129S7-Rag1\textsuperscript{tm1Mom/J} (Rag1\textsuperscript{−/−}), and B6.SJL-Ptprc\textsuperscript{a} Pep3\textsuperscript{−/−}Boy1 (CD45.1) mice were obtained from The Jackson Laboratory (Bar Harbor, ME). Mice were bred and maintained under specific pathogen-free conditions, including detectable strains of helicobacter. Experiments were performed under a protocol approved by and in accordance with Institutional Animal Care and Use Committee guidelines.

Peptides, Abs, and flow cytometry

MOG\textsubscript{35–55} peptide (MEVGWYRSPFSRVVHLYRNGK) was synthesized and HPLC purified by the St. Jude Hartwell Center for Biotechnology. mAbs specific for CD4 (clone H129.19), CD8 (clone 53-6.7), CD25 (clone 7D4), CD45R (clone H12.2F5), CD45RB (clone 16A), CD44 (clone IM7), and CD45 (clone IM4) were from eBioscience (San Diego, CA). Flow cytometry was performed on an eBioscience FACSCalibur (BD Biosciences), and flow cytometric sorting was performed on a MoFlo high-speed cell sorter (DakoCytomation, Fort Collins, CO).

TCR constructs and generation of retrogenic and chimeric mice

1MOG9 and G107S TCR α- and β-chains were isolated by PCR and cloned into the murine stem cell virus (MSCV)-I–GFP MSCV-based retroviral vector and retroviral mice generated as described (19, 20). Briefly, bone marrow (BM) cells were harvested from the femurs of Rag1\textsuperscript{−/−} mice 48 h after the administration of 0.15 mg 5-fluorouracil/g body weight. The pooled cells were cultured in complete Click’s medium (Invitrogen, Carlsbad, CA) containing 20% FCS, IL-3 (20 ng ml\textsuperscript{−1}), IL-6 (50 ng ml\textsuperscript{−1}), and stem cell factor (50 ng ml\textsuperscript{−1}) for 48 h at 37°C 5% CO\textsubscript{2}. The cells were then cocultured for an additional 48 h with 1200 rad-irradiated retroviral producer cells. The hematopoietic progenitor cells (HPC) were harvested, washed with PBS, transduction confirmed by flow cytometry for GFP, and

FIGURE 1. Immunophenotype of G107S and 1MOG9 retrogenic T cells. Thymocytes, LN cells, and splenocytes were isolated from retrogenic mice \textasciitilde 8 wk following stem cell transfer or B6 controls. A, CD4 and CD8 labeling of thymocytes is shown. B, CD4 and CD8 expression on TCRB-gated LN cells or splenocytes. C, CD44, CD45RB, and CD25 labeling was used to indicate the memory or naive status of CD4\textsuperscript{+} T cells. CD25 and Foxp3 were used as regulatory T cell markers. D, Labeling of CD8\textsuperscript{+} TCRB\textsuperscript{+} T cells to evaluate the distribution of naive and memory populations as in C. E, Naïve and memory cells among G107S DN TCRB\textsuperscript{+} T cells. Data are from representative animals.
injected i.v. into sublethally irradiated (450 rad) Rag1−/− mice at a ratio of two recipient mice per BM donor. For the generation of chimeric mice, transduced HPC were admixed with B6 BM cells and transferred i.v. into 900 rad-irradiated B6 recipients. Mice were assessed at the indicated times posttransplant.

**EAE induction and clinical evaluation**

EAE was induced by s.c. immunization with 100 μg MOG35–55 peptide emulsified in CFA containing 0.4 mg (4 mg/ml) Mycobacterium tuberculosis H37RA (Difco, Franklin Lakes, NJ). Two hundred nanograms pertussis toxin (List Biological Laboratories, Campbell, CA) was administered i.v. on days 0 and 2. In some experiments, CD4+, CD8+, and DN TCRβ+ cells were flow cytometrically isolated and transferred into recipient mice 1 d prior to immunization. Clinical scoring was as follows: 1, limp tail; 2, hind limb paresis or partial paralysis; 3, total hind limb paralysis; 4, hind limb paralysis and body/front limb paresis/paralysis; and 5, moribund.

**T cell proliferation**

CD4+, CD8+, and DN TCRβ+ T cells were isolated by flow cytometry or Ab-based magnetic bead selection (MACS; Miltenyi Biotec, Auburn, CA) per the manufacturer’s instructions. These were from mice either unimmunized or 7 d postimmunization with MOG35–55 peptide. Cells were cultured at 5 × 10⁴/well in 96-well plates with 3 × 10⁵ irradiated B6 APCs and the indicated stimulus, pulsed with 1 μCi [3H]thymidine after 72 h of culture, and then harvested for scintillation counting. Samples were analyzed in triplicate.

**Cytokine analysis**

CD4+, CD8+, and DN TCRβ+ T cells were isolated and cultured as described above. Culture supernatants were collected at 48 h and analyzed for IL-2, IL-4, IL-10, IL-17, and IFN-γ using the Milliplex Map kit assay (Millipore, Billerica, MA).

**Cytotoxicity assay and quantitative flow cytometry**

Flow cytometrically purified CD4+, CD8+, and DN TCRβ+ cells from CD45.1+CD45.2+ G107S mice were stimulated for 4 d with MOG35–55, then incubated at a 1:1 ratio with freshly purified congenic CD45.1+CD45.2+ B cells or BM-derived dendritic cell (DC) targets for 6 or 18 h with or without 100 μg/ml MOG35–55. BM-derived DCs were matured with GM-CSF and IL-4 (R&D Systems, Minneapolis, MN) as described (21). Samples were stained for the indicated target B cell or DC population and then analyzed by quantitative flow cytometry as described (22). Viable cell counts are plotted for individual samples assayed in triplicate.

**Results**

**Retrogenic mouse production and characterization**

The parental MOG35–55-specific, I-Aβ–restricted 1MOG9, and G107S TCR are identical except for the added hydroxymethyl group in the TRBV 107 side chain in the G107S receptor. This G/S variant residue is at the immediate junction of the TRBV and TCR β-chain J region, does not directly engage peptide–MHC, and is predicted to fill a buried gap in the 1MOG9 TRBV. The G107S TCR was previously observed in transduced hybridoma cells to possess an ∼3log₈₀ increase in Ag sensitivity (20).

The TCRA and B chains of the 1MOG9 and G107S TCR were subcloned into the MSCV-I–GFP retroviral vector separated by a viral 2A sequence to facilitate their stoichiometric expression. Retrovirus was transduced into Rag1−/− HPC, which were transplanted into lethally irradiated B6 Rag1−/− mice at a ratio of two recipient mice per BM donor. For the generation of chimeric mice, transduced HPC were admixed with B6 BM cells and transferred i.v. into 900 rad-irradiated B6 recipients. Mice were assessed at the indicated times posttransplant.

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planted into Rag1−/− recipients, thereby ensuring that only the retrogenic TCR was expressed.

**G107S lineage infidelity**

Skewed thymic development of CD4+ single-positive T cells was seen in mice receiving HPC transduced either with the 1MOG9 or G107S TCR (Fig. 1A). Both 1MOG9 and G107S retrogenic mice also showed peripheral engraftment with the self-specific TCR (Fig. 1B). In each case, the CD4+ T cells were predominantly naive, bearing a CD44lo, CD45Rbhi, CD62Lhi phenotype (Fig. 1C). Little expression of the activation and regulatory marker CD25 was apparent on either cell type, and despite the enhanced affinity for a self-antigen in the G107S mice, Foxp3+ regulatory T cells were seen. In contrast with the virtually exclusively CD4+ T cell engraftment in mice expressing the wt 1MOG9 TCR, substantial numbers of peripheral CD8+ T cells were observed in G107S mice (Fig. 1B). These CD8+ cells, as the CD4+, were predominantly naive (Fig. 1D). A smaller population of DN T cells was also seen in the G107S mice, whereas few were present in the 1MOG9 animals. These also displayed a largely naive phenotype (Fig. 1E). Therefore, the high-affinity G107S TCR promotes lineage infidelity and T cell subset heterogeneity.

Increased affinity for self is associated with the deletion of autoantigen-specific T cells (23). However, this was not apparent with the G107S TCR. To the contrary, 7–12 wk after stem cell transfer, the G107S mice showed significantly more splenic CD4+ TCR+ T cells than matched wt 1MOG9 TCR retrogenic mice (p < 0.0001; Fig. 2). Therefore, the G107S substitution does not adversely affect the development and engraftment of self-reactive T cells. Rather, it supports increased engraftment as well as production of CD8+ and smaller numbers of DN T cells. Notably, in a prior study of retrogenic mice expressing different MOG-specific TCR (19), we were unable to associate affinity with engraftment level, and the recognition parameters that are promoting this increased engraftment remain to be determined.

**Heightened responsiveness of primary G107S T cells**

Developmental tuning can dampen the responsiveness of high-affinity T cells (24, 25). To determine if the increased Ag sensitivity previously observed in G107S-TCR–transduced cell lines was retained in primary T lymphocytes, we compared the proliferative response of isolated CD4+ T cells from 1MOG9 and G107S retrogenics. Similar to transduced cell lines, a ∼3-log10 increase in MOG sensitivity was seen in primary CD4+ T cells expressing the G107S TCR (Fig. 3A). Isolated CD8+ and DN T cells present in the G107S mice also proliferated strongly in response to cognate Ag with response profiles similar to that of CD4+ T cells (Fig. 4A), verifying our prior observation in hybridomas that the G107S TCR is coreceptor independent. Therefore, a G107S substitution in a TRBV13-2 TCR is fully compatible with the development of functional T lymphocytes bearing markedly enhanced sensitivity for cognate Ag.

**Distinct cytokine profiles of 1MOG9 and G107S T cells**

To assess the differentiation potentials of 1MOG9 and G107S T cells, CD4+ T cells were bead isolated from 8 wk retrogenic mice and equal numbers stimulated with MOG35–55. IL-2, IL-4, IL-10, IFN-γ, and IL-17 were measured as representative Th0, Th2/Tr1, Th1, and Th17 cytokines. Either 1MOG9 or G107S CD4+ T cells showed good IL-2 production, though levels were greater with the G107S T cells (Fig. 3A). Both cell types also produced IFN-γ, associated with differentiation into Th1 cells that are pathologic in EAE (26, 27), though in this study more was produced by the lower affinity 1MOG9 T cells. G107S T cells produced modest amounts of IL-17, also associated with EAE pathogenicity (28), whereas none was seen with the 1MOG9 T cells. IL-4 is associated with disease protection (29). Little or no IL-4 was produced by the 1MOG9 T cells, whereas substantial quantities were produced by the G107S cells. The G107S T cells also exclusively demonstrated significant production of IL-10, a cytokine with pronounced regulatory activity in EAE (30–34). Therefore, CD4+ T cells from 1MOG9 mice showed a restricted cytokine production pattern characterized by IL-2 and IFN-γ production and little IL-4 and IL-10 synthesis. G107S T cells show a more degenerate and pronounced cytokine profile that included cytokines characteristic of Th1, Th2, and Th17 cells. IL-10 and IL-4 production, protective in EAE, was limited to the G107S cells.

CD8+ and DN T cells were similarly isolated, though from older 16-wk retrogenic mice, necessary to obtain adequate DN
numbers by cell sorting. These demonstrated less IL-2 and more IFN-γ than similarly isolated CD4+ G107S T cells (Fig. 4B). Neither DN nor CD8+ T cells produced substantial quantities of IL-10. However, production of other cytokines diverged between the cell types. DN cells produced more IL-17, whereas CD8+ T cells produced little. CD8+ T cells also produced little IL-4 when compared with DN and CD4+ T cells. Therefore, G107S T cell subsets possess distinct functional profiles when stimulated with Ag, with variability in the production of both inflammatory and protective cytokines.

Cytolytic activity of DN and CD8+ T cells
To determine if the T cells were able to kill MOG-expressing APCs, primary B cells or BM-derived DCs were pulsed with MOG35–55 and cocultured for either 6 or 18 h with G107S T cells at a 1:1 E:T ratio (Fig. 5). The DN and CD8+ T cells lysed the B cell populations at either time point. G107S (Fig. 5A) or IMOG9 (not shown) CD4+ T cells were unable to kill these cells. At the 1:1 E:T ratio, DCs proved insensitive to CTL killing (Fig. 5B); such DC resistance to lysis has been previously reported (35). However, at a 4:1 ratio, modest DC killing by the DN and CD8+ populations was evident (data not shown). Therefore, the CD8+ and DN but not CD4+ effector cells are capable of killing cells presenting Ag, though the extent of this is dependent on lineage sensitivity to lysis.

Diminished EAE severity in G107S mice
We would predict that the altered differentiation profile of the higher affinity G107S T cells would influence EAE susceptibility. To test this, IMOG9 and G107S mice were immunized with MOG35–55. The IMOG9 mice developed fulminant disease, which proved nearly uniformly fatal by 2 wk after immunization (Fig. 6A, Table I). In contrast, despite the increased T cell affinity for Ag, the G107S mice manifested milder disease, which, though not

**FIGURE 6.** Diminished EAE severity and altered subset expansion in G107S retrogenic mice. A, EAE was induced by immunization with MOG35–55. Clinical disease score was monitored longitudinally. Mean ± 1 SEM is plotted. Representative of two experiments. Additional data parameters are provided in Table I. B–E, T lymphocytes were isolated from the spleen and CNS of mice 6 d after induction of EAE or spleen of control mice in which disease was not induced. B, Sample plots showing CD4 and CD8 labeling profiles of splenocytes (immunized and unimmunized) or CNS (immunized only) TCRβ+ T cells. Absolute numbers of splenic T cell populations in IMOG9 (C) or G107S (D) mice are plotted. Circles indicate individual animals and lines represent means. E, Absolute numbers of isolated TCRβ+ CNS T cell subsets are similarly plotted.
CD4+ population expanded less than in the 1MOG9 mice. CD8+ mice, a majority of T cells were CD4+ in the absence of immunization of CD4+ T cells in 1MOG9 mice, but CD8+ and DN significantly increased. Analysis of the CNS lymphoid infiltrate showed a predominance of CD4+ T cells, with only small numbers of CD8+ or DN T cells detected. After EAE immunization, all T cell populations expanded, though not equally (Fig. 6). The TCR affinity and pathogenicity of isolated high-affinity subsets, CD4+, CD8+, or DN T cells were transferred into Rag1−/− recipients 1 d prior to immunization. For experiments 1 and 2, retrogenic mice were directly immunized. For experiments 3 and 4, 5 × 10⁵ of the indicated flow cytometrically purified T cells were transferred into Rag1−/− recipients 1 d prior to immunization. Onset time is from MOG immunization. Longitudinal course of experiments 1 and 3 are plotted in Figs. 6 and 9, respectively.

AUC, area under the curve analysis using trapezoidal estimation.

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<th>AUC</th>
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In experiments 1 and 2, retrogenic mice were directly immunized. For experiments 3 and 4, 5 × 10⁵ of the indicated flow cytometrically purified T cells were transferred into Rag1−/− recipients 1 d prior to immunization. Mortality was absent among the G107S mice. Therefore, a high-affinity TCR in G107S retrogenic mice, despite promoting significantly enhanced T cell engraftment, led to markedly diminished disease severity.

**Distinct T lineage responses in G107S and 1MOG9 mice**

To better characterize the responding cells in the different mice, we examined splenic and CNS cell numbers prior to and after immunization. As described above, CD4+ T cells were the predominant lymphoid constituent in preimmune 1MOG9 mice, with only small numbers of CD8+ or DN T cells detected. After EAE induction, a dramatic expansion of CD4+ T cells was seen. These remained the dominant cell type in the spleen, and little expansion of CD8+ or DN T cells was apparent (Fig. 6B, 6C). In G107S mice, a majority of T cells were CD4+ in the absence of immunization, though a sizable population of CD8+ T cells and smaller numbers of DN cells were also present. After immunization, all cell populations expanded, though not equally (Fig. 6B, 6D). The CD4+ population expanded less than in the 1MOG9 mice. CD8+ and DN T cells expanded to a much greater degree than CD4+ T cells, reversing the CD4+/CD8+ and CD4+/DN T cell ratios. Analysis of the CNS lymphoid infiltrate showed a predominance of CD4+ T cells in 1MOG9 mice, but CD8+ and DN significantly outnumbered CD4+ cells in G107S mice (Fig. 6E). Therefore, the autoimmune response in 1MOG9 and G107S mice is characterized by the differential expansion and CNS infiltration by CD4+, CD8+, and DN T cell subsets.

**Functional potential of immune T cells**

To functionally assess the responding cells, we analyzed the proliferative potential of purified T cell populations 7 d after immunization. In contrast to the distinct responses of preimmune T99, comparison of the T cell populations from mice with EAE demonstrated nearly equivalent in vitro proliferation of the 1MOG9 CD4+ T cells and G107S T cell subsets (Fig. 7A). This differed from results with preimmune populations (Fig. 3). Indeed, direct comparison of the proliferative responses of purified CD4+ T cells isolated from immunized and unimmunized mice showed a corresponding enhanced responsiveness to Ag among immunized 1MOG9 but not G107S T cells (data not shown) compared with unimmunized controls. Therefore, prior activation differentially influences subsequent Ag sensitivity by G107S and 1MOG9 T cells, with only the lower affinity cells developing increased reactivity.

Cytokine profiles of the 1MOG9 and G107S T cells were further assessed at day 7, which, though early after disease induction, only shortly preceded the demise of most of the 1MOG9 mice (Fig. 7B–F). G107S CD4+, CD8+, and DN T cells showed significantly decreased IL-2, IFN-γ, and IL-17 compared with 1MOG9 cells. This indicates an overall diminished production of Th1 and Th17 cytokines relevant to EAE. Only modest IL-4 and IL-10 was produced by immune cells from either mouse type.

Therefore, an increased TCR affinity in G107S mice is associated with both attenuated disease course and diminished effector cytokine production. Notably, similar to the preimmune mice (Fig. 1), few Foxp3+ regulatory T cells were detectable in either G107S or 1MOG9 mice after induction of EAE (data not shown).

**Pathogenicity and protection by isolated G107S cellular subsets**

Our results indicated that multiple T cell subsets responded in G107S EAE; the dominant T cell response was CD4+, and G107S T cells had diminished effector cytokine production. To examine the pathogenicity of isolated high-affinity subsets, CD4+, CD8+, or DN T cells were purified from G107S and 1MOG9 mice and CD4+ T cells from 1MOG9 mice. These were transferred into Rag1−/− recipients and disease induced. CD4+ T cells from G107S mice manifested diminished disease relative to lower affinity 1MOG9 CD4+ T cells, indicating that increased TCR affinity decreases T cell pathogenicity in this population (Fig. 8, Table I). In contrast to the G107S CD4+ T cells, mild or no disease was seen in recipients of CD8+ or DN G107S T cells. Therefore, pathogenicity is a function of both TCR affinity and cell lineage. DN and CD8+ T cells, which show the greatest expansion in immunized G107S mice, have limited capacity to independently promote disease.

To determine whether G107S T cells could also suppress disease development and progression in a competitive environment, we analyzed their impact in B6 mice. Small numbers (1 × 10⁵) of flow cytometrically purified CD4+, CD8+, or DN T cells were transferred into Rag1−/− recipients.
ferred into B6 recipients, and 1 d later, EAE was induced (Fig. 9 A).
Transfer of G107S CD4+ T cells diminished disease severity when
compared with control treated mice. However, CD8+ T cells
blocked G107S disease development and DN T cells markedly
delayed time to disease onset as well as disease severity, indicating
that these subsets are protective in wt mice.

To determine if the MOG-specific CD4+ T cells would develop
not only in Rag2−/− mice but in the context of a wt thymus, we
diluted G107S, 1MOG9, or retroviral vector-transduced Rag2−/−
HPC with an excess of untransduced B6 BM cells and trans-
planted these into B6 recipients. Only small numbers of retrogenic
T cells, identifiable through their coexpression of GFP, were
identified. They averaged fewer than 2% of TCRAB+ T cells in
mice receiving 1MOG9-transduced progenitors and were com-
prised virtually exclusively of CD4+ T cells (Fig. 9 B). Approxi-
mately 7% of TCRAB+ cells in the G107S chimeras expressed the
retrogenic TCR. Quantities of CD4+ retrogenic T cells were not
significantly different from those in the 1MOG9 mice. However,
CD8+ and DN G107S T cells were also observed, making up for
the balance of cells. Therefore, the high-affinity MOG-specific
TCR promotes the cross-differentiation of T cells into corecep-
tor independent CD8 and DN T cell subsets when developing as
a minority population in wt mice.

To determine if the MOG-specific CD4+ T cells would protect against EAE
development, disease was induced in these chimeric mice (Fig.
9 C). Whereas mice receiving control MSCV vector- or 1MOG9
TCR-transduced HPCs demonstrated similarly high levels of dis-
ease, mice receiving G107S-transduced cells demonstrated a milder disease course. Therefore, as with T cell transfer into wt
mice, G107S T cells developing in mixed stem cell chimeras di-
minish EAE severity.
cytochrome C is dominated by high-affinity cells that following in vitro priming generates a Th1-biased response. Purging the high-affinity cells leads to lower affinity repertoire that develop a Th2 bias (38). In a consistent manner, low Ag dose or weak APL signaling leads to a Th2 responses in TCR Tg cells, whereas high dose promotes Th1 maturation (10, 39). In contrast, in matured T cell lines, it has also been observed that Th2 cells require greater signal strength on rechallenge than Th1 cells for cytokine production (40).

Affinity likewise is essential in determining cell survival and proliferation. In the Listeria-specific response, early stimulation activates both low- and higher-affinity cells. The lower affinity cells, however, exit the lymph nodes (LNs) earlier and have a more abbreviated expansion phase, potentially explaining the affinity maturation that is observed (41). In other conventional responses, affinity maturation is also present, and higher affinity cells dominate the secondary response. Despite these findings, in autoimmune systems, strong agonists may be protective. Thus, a myelin APL superagonist did not lead to EAE, but promoted the desensitization and activation-induced cell death of autoantigen-specific T cells in a Fas-dependent manner (11, 12).

Our results are consistent with a model in which T cell immunopathologic potential is bracketed by both low- and high-affinity thresholds and describe new mechanisms for this. Importantly, the G107S TCR differs only in the presence of a buried hydroxymethyl group in the TRBV (20). The S107 residue supplying this moiety is present in the majority of expressed TCR; 20 out of 23 mouse and 49 out of 54 human TRBV possess an S, whereas only 1 and 2, respectively, incorporate a G, and therefore, the substitution represents a natural variation at this site (20). The G leaves a gap at the base of the CDR3β, which is filled by the S side chain. This minor structural perturbation influences affinity for MOG35-55 without substantially altering fine specificity. Yet it has several functionally significant effects. Most prominently, whereas the wt 1MOG9 TCR drives T cells into the CD4+ T cell lineage, the G107S TCR leads to lineage infidelity, with CD4+, CD8+, and DN T cells identifiable. These CD4+ populations also develop in mixed HPC chimeras in wt mice, indicating that they reflect the natural differentiation potential of this TCR.

Whether increased G107S TCR affinity for a self-ligand is itself responsible for the lineage infidelity is uncertain from this study. In one other model, a high-affinity self-specific TCR also guided T cell development into CD4+CD8- T cells (42), suggesting a role. Assessments of additional TCR series of varying affinity may help determine the recognition parameters controlling the fidelity of lineage assignments. Increased affinity is, however, essential for coreceptor-independent Ag recognition. Studies of MHC class I-restricted Tg1 TCR identified an affinity threshold in the micromolar range above which coreceptor is no longer required (43). The 1MOG9 TCR is class II MHC restricted, and threshold affinities for coreceptor independent class II MHC recognition are, to our knowledge, unavailable. Nevertheless, we previously demonstrated that CD4+ hybridomas expressing the high-affinity G107S TCR are coreceptor independent and remain class II MHC restricted, class II-/- APCs failing to stimulate them (19). This finding was replicated with the primary retrogenic T cells studied in this paper (data not shown).

Isolated DN and CD8+ G107S T cells were only weakly pathogenic. Moreover, these populations outcompete CD4+ T cells during disease development. They expanded to form the majority of cells infiltrating the CNS, potentially explaining the dramatically reduced disease in G107S mice. After transfer into wt mice, very small numbers of the cells (10^7, Fig. 9, and as few as 4 × 10^4 [not shown]) were protective. This indicates a broader capacity

**FIGURE 9.** Regulatory activity of G107S CD8+ and DN T cells in B6 mice and mixed chimeras. A, Total of 10^5 of the indicated flow cytometrically purified T cell subset or saline was transferred i.v. into B6 mice 1 d prior to EAE induction and disease course monitored. Data are pooled from two similar experiments. n = 10 per group. Mean ± 1 SEM is plotted. B, Engraftment of G107S or 1MOG9 T cell subsets in mixed chimeras. HPC transduced with the indicated TCR were admixed with wt B6 BM cells and transplanted into irradiated B6 recipients. The retrogenic T cells were identified in blood cells by coexpressed GFP. Mean + 1 SD is plotted. Representative of two experiments. C, EAE was induced in mice in B and disease clinically monitored. n = 7, G107S; n = 6 other groups. Mean ± 1 SEM is plotted.

**Discussion**

The presence of self-specific T cells is inadequate for autoimmunity. These cells must also be stimulated in a manner that triggers their maturation into destructive effector forms, promotes their expansion, and sustains their pathogenicity over time. To achieve this balance experimentally, specific induction regimens are often needed. In most experimental systems, relatively few epitopes have been identified against which T cell responses can provoke autoimmunity. For example, in B6 mice, MOG35-55 and the cross-reactive NF-M18-30 epitope are the only Ags identified able to induce EAE (36, 37). The available T cell repertoire will determine the quality of the response to Ag and hence susceptibility to autoimmunity. Thymic and peripheral T cell deletion may lead to specificity holes and thereby tolerance. As described in this study, increased affinity can also lead to diminished disease susceptibility by altering differentiation potential.

TCR affinity plays a crucial role in conjunction with environmental signals in the functional maturation of T cells. Th subset maturation is avidity dependent. Thus, the response to pigeon immunopathologic potential is bracketed by both low- and high-affinity thresholds and describe new mechanisms for this. Importantly, the G107S TCR differs only in the presence of a buried hydroxymethyl group in the TRBV (20). The S107 residue supplying this moiety is present in the majority of expressed TCR; 20 out of 23 mouse and 49 out of 54 human TRBV possess an S, whereas only 1 and 2, respectively, incorporate a G, and therefore, the substitution represents a natural variation at this site (20). The G leaves a gap at the base of the CDR3β, which is filled by the S side chain. This minor structural perturbation influences affinity for MOG35-55 without substantially altering fine specificity. Yet it has several functionally significant effects. Most prominently, whereas the wt 1MOG9 TCR drives T cells into the CD4+ T cell lineage, the G107S TCR leads to lineage infidelity, with CD4+, CD8+, and DN T cells identifiable. These CD4+ populations also develop in mixed HPC chimeras in wt mice, indicating that they reflect the natural differentiation potential of this TCR.

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to modulate autoimmune disease. Implicitly, in contrast to the competitive maturation to high-affinity responses typical after immunization or infection, the low or moderate affinities often observed in autoimmunity may be necessary for the generation of pathologic responses.

The mechanism of protection of these CD4+ cells in this model remains uncertain, though some clues are present. DN T cells have been observed more broadly to possess regulatory properties, as evidenced in other models of allograft and autoimmune diseases (44, 45). It would seem likely in this study that the low pathologic potential of the DN cells coupled with their ability to outcompete more pathologic CD4+ T cells provides for their immunomodulatory activity. The source of these cells in natural circumstances has been debated. Small numbers of DN αβ T cells normally circulate, typically <2%, though these are increased in certain conditions such as autoimmune lymphoproliferative syndrome (46, 47). Defects in Fas signaling there suggest that alterations in this death-signaling pathway may allow the accumulation of DN cells. Indeed, DN T cells survive chronic stimulation better than their coreceptor-positive peers (48). In some situations, it is hypothesized that DN cells are generated through coreceptor downregulation with stimulation, which may occur genetically or epigenetically (49–52). However, other sources are possible (53), and the largely naive state of the DN cells in unimmunized G107S mice and their distinct cytokine profile when compared with either CD4+ or CD8+ cells would indicate that they are a primary population. Indeed, diversion of a large proportion of autoreactive T cells to DN cells was observed when a high-affinity PLP-specific TCR was transgenically expressed on T cells in a disease-resistant genetic background. Whereas DN TCRαβ T cells form a small portion of most immune responses, this evidence supports their formation from high-affinity, self-specific cells through a common differentiation pathway (42). Further evaluation of these DN T cells and their role in EAE and other autoimmune diseases in unmanipulated mice is warranted.

CD8+ T cells recognizing epitopes internal to MOG35–55 can modify MOG-EAE, and both pathologic and protective properties have been attributed to them (54–56). EAE in β2-microglobulin (β2m)-/- mice shows increased disease severity, suggesting that, overall, the class I-restricted response is protective (57). In our analyses, the G107S CD8+ T cells recognized class II MHC-restricted Ag. Such lineage crossover by coreceptor-independent cells has been previously observed in immune responses (58), though the extent to which T cells are capable of crossing MHC restriction is unclear. Nevertheless, coreceptor independent signaling is adequate to activate T cells with high affinities (43). Further, the sheer magnitude of the degeneracy inherent in TCR recognition, in which it is anticipated that a single TCR may recognize >10^6 unique Ags, would suggest that some high-affinity, coreceptor-independent responses will exist, and these may cross MHC class barriers (59).

To our knowledge, it has not been previously shown that high-affinity cells specific for a class II-restricted Ag may actually develop in situ into CD8+ T cells. To test the MHC requirements for this development, we generated G107S retrogenic mice using transduced B2m-/- Rag2-/- HPCs transferred into β2m-/- Rag1-/- recipients. Although the relative numbers of CD8+ T cells was diminished compared with Rag1-/- retrogenic mice, a significant population was nevertheless present, and DN cells were increased (Supplemental Fig. 1). Therefore, both CD8+ and DN G107S T cells can develop in a class I MHC-independent manner.

It is of interest, though not surprising, that the CD8+ and DN G107S T cells can kill Ag-pulsed APCs, particularly B cells. Cytotoxicity is a characteristic of these populations, and it has been hypothesized that this can restrict autoreactive responses by reducing APC availability (54). The role of APC killing in the diminished pathogenicity and protection in this study will have to be established. However, DCs are a critical APC type in the EAE model system (60), and these possess mechanisms to resist lysis, including the expression of granzyme-inhibitory serpins (35). Indeed, at E:T ratios capable of easily killing B cells, DCs proved impervious to lytic attack.

The functional profile of the G107S and 1MOG9 T cells, defined in this study by their cytokine production patterns, differed significantly between the cells, and further explains the reduced pathogenicity of the higher affinity cells. Primary CD4+ G107S T cells produced increased IL-10 and IL-4, which are regulatory in EAE, compared with wt 1MOG9 cells. In mice with active disease, substantive IL-10 and IL-4 responses were not detected. Nevertheless, the G107S T cells produced less IFN-γ and IL-17, indicating a disruption in Th1/Th17-mediated immunopathogenesis. Further, isolated CD4+ G107S T cells were less potent than CD4+ 1MOG9 T cells in transfer models. Therefore, consistent with data on the use of superagonistic APL to stimulate autoreactive T cells, and the observed diminished pathogenicity of high-affinity TCR transgensics, G107S CD4+ T cells possess an intrinsically reduced pathogenicity compared with paired 1MOG9 T cells. Further, lower affinity 1MOG9 but not G107S T cells demonstrated enhanced in vitro response to Ag after immunization, suggesting that prior stimulation differentially upregulates responsiveness and potentially supports expansion of the lower affinity cells upon Ag restimulation.

In summary, we demonstrate highly distinct T cell immunopathologic potential in two cohorts of mice expressing retrogenic TCR with minimal structural differences though more dramatically altered affinity for MOG35,55. Different sources are responsible, including cell-intrinsic effects on cytokine patterns by pathologic CD4+ T cells as well as altered T cell differentiation and the induction of protective DN and CD8+ T cell responses. The presence of high-affinity self-antigen–specific T cells may therefore provide a barrier, protecting the host against pathologic autoimmune responses. Further analyses of the effect of TCR affinity on autoantigen-specific T cell differentiation profiles are warranted to better define how T cell repertoire properties influence autoimmune susceptibility.

Acknowledgments
We thank Richard Cross, Greig Lennon, Stephanie Morgan, and the Immunology Core Flow Cytometry facility for assistance with flow cytometric sorting.

Disclosures
The authors have no financial conflicts of interest.

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