Developmental control of CD8+ T cell–avidity maturation in autoimmune diabetes

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The progression of immune responses is generally associated with an increase in the overall avidity of antigen-specific T cell populations for peptide-MHC. This is thought to result from preferential expansion of high-avidity clonotypes at the expense of their low-avidity counterparts. Since T cell antigen-receptor genes do not mutate, it is puzzling that high-avidity clonotypes do not predominate from the outset. Here we provide a developmental basis for this phenomenon in the context of autoimmunity. We have carried out comprehensive studies of the diabetogenic CD8+ T cell population that targets residues 206–214 of the β cell antigen islet-specific glucose-6-phosphatase catalytic subunit–related protein (IGRP206–214) and undergoes avidity maturation as disease progresses. We find that the succession of IGRP206–214–specific clonotypes with increasing avidities during the progression of islet inflammation to overt diabetes in nonobese diabetic mice is fueled by autoimmune inflammation but opposed by systemic tolerance. As expected, naive high-avidity IGRP206–214–specific T cells respond more efficiently to antigen and are significantly more diabeticogenic than their intermediate- or low-avidity counterparts. However, central and peripheral tolerance selectively limit the contribution of these high-avidity T cells to the earliest stages of disease without abrogating their ability to progressively accumulate in inflamed islets and kill β cells. These results illustrate the way in which incomplete deletion of autoreactive T cell populations of relatively high avidity can contribute to the development of pathogenic autoimmunity in the periphery.

Introduction

The diversity of the preimmune T and B cell repertoires is accomplished by random rearrangement of large arrays of V, D, and J genes during lymphocyte development. The encoded molecules fold into structures in which 6 hypervariable complementarity-determining regions (CDRs) are brought together to form the receptors’ antigen-binding sites (1, 2). A key common feature of the antigen-binding site of these receptors is its conformational flexibility, which affords ligand promiscuity (1–11). This degeneracy in antigen recognition is typically associated with low-affinity interactions and is a hallmark of the preimmune B lymphocyte repertoire. It is via the processes of somatic mutation of B cell receptors and selection of antigen-reactivated B cells within germinal centers that B cell responses improve in both their affinity and fidelity for antigen recognition (12–16). This transition is paralleled by dramatic increases in both the association rate of the antigen-binding reaction and the structural rigidity of the receptors’ antigen-binding site (9, 17).

T cell responses usually undergo an analogous process of avidity maturation. Although there is some controversy as to the underlying kinetic mechanisms (i.e., avidity maturation proceeding through or only up to certain avidity thresholds), most studies concur that this process results from competitive survival of clonotypes bearing high-affinity TCRs for antigen-MHC at the expense of clonotypes bearing low-affinity TCRs for the same ligand (18–22). Unlike B cells, however, T cells do not normally mutate their antigen receptors. Furthermore, whereas a handful of somatic mutations in a B cell receptor are traceable and amenable to detailed developmental and biophysical studies, T cell–avidity maturation usually involves competition among clonotypes expressing highly diverse TCR structures that are difficult to compare. As a result, the molecular basis and physical determinants of T cell–avidity maturation remain poorly understood.

NOD mice develop a form of autoimmune diabetes that closely resembles human type 1 diabetes. The onset of overt diabetes in these mice is preceded by prolonged periods of pancreatic-islet inflammation by β cell–reactive CD4+ and CD8+ T cells (23). A significant fraction of all the CD8+ T cells that are recruited to islets during the earliest stages of diabetogenesis express a restricted set of TCR-α chains (Vβ17 and Jβ42 elements joined by the amino acid sequence MRD) (24–26). We have shown that this T cell population recognizes the mimotopes NRP-A7 and NRP-V7 (KYNKAN(A/V)/FL) in the context of the murine MHC class I molecule Kd (27, 28) and that, over time, this population undergoes a process of avidity maturation (i.e., binds peptide-MHC with progressively lower Kd) that contributes to the progression of islet inflammation to overt diabetes (20). The natural antigenic target of this T cell population has been recently shown to correspond to residues 206–214 (VYLKTNVFL) of islet-specific glucose-6-phosphatase catalytic subunit–related protein (IGRP) (29).

Here we investigate mechanisms underlying the avidity maturation of this T cell population. Unlike other examples of T cell–avidity maturation, which involve repeated exposure of mice to relatively high levels of exogenous antigen and occur within days of antigen exposure (18–22), avidity maturation of IGRP206–214–reactive CD8+ T cells is likely driven by much smaller amounts of self anti-
gen and occurs over a much more prolonged period of time (20). Here we find that this process results, in part, from competition among Vα17-MRD-Jα42+ clonotypes using different Vα17 elements and that small, germline-encoded molecular changes distal to the antigen-binding sites of TCRs can have profound effects on the functional avidity and developmental biology of individual clonotypes. Experiments in TCR-Tg animals further show that, whereas autoimmune inflammation preferentially fuels the local expansion of the highest avidity and most pathogenic T cell pool, systemic tolerance selectively decreases the size of this pool, which explains why there is prevalent recruitment of low-avidity clonotypes in the primary autoimmune response.

Results
Avidity maturation of the NRP-A7–reactive CD8+ T cell population is associated with changes in Vα17 usage. (A) Changes in Vα17 usage by islet-derived, NRP-A7–reactive CD8+ cells with age (data from Supplemental Figure 1). (B) Predicted amino acid sequences of the 3 Vα17 elements expressed by NOD mice. Gray boxes highlight sequence differences compared with Vα17.6, the least used Vα17 element by 20 weeks. (C) Average mfi for Vβ8 and CD8 on TCR-αβ transfectants expressing the 8.3 TCR-β chain and the 8.3–TCR-α (Vα17.4) or 8.3-like TCR-α chains (using Vα17.6 or Vα17.5) (n = 3 clones each). (D) Tetramer binding to transfectants. (E) Flow cytometric analysis of 17.6α/8.3β and 17.5α/8.3β transfectants stained with FPLC-purified, Cy5-labeled NRP-V7/Kd tetramers. Numbers on the upper right corner correspond to average mfi values. (F) Production of IL-2 by Vα17.6+, Vα17.5+, and Vα17.4+ transfectants in response to anti-CD3–coated microbeads or IGRP206–214–pulsed NOD splenocytes (average ± SEM for 1, 4, and 4 clones, respectively). The peptide concentrations that give the half-maximum response (SD50) for IL-2 secretion corresponding to Vα17.6+, Vα17.5+, and Vα17.4+ cells were as follows: 173 ± 54 (Vα17.6+), 13 ± 8 (Vα17.5+), and greater than 1000 (Vα17.4+) µg/ml (for NRP-V7), and greater than 1000 (Vα17.6+), 1 ± 1 (Vα17.5+), and 135 ± 17 (Vα17.4+) µg/ml (for IGRP206–214), respectively (P < 0.001) (data are from a separate experiment employing 3 clones of each transfectant type). (G) Flow cytometric analysis of Vα17.5 and Vα17.6 transfectants coexpressing another Vβ8.1+ TCR-β rearrangement cloned from tetramer+ cells of 9-week-old NOD mice (cDNA clone NC28, with the CDR3 sequence CASAPENTL). Numbers on the graphs correspond to average mfi values of 2–6 clones per group.
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Figure 2

Developmental biology of 17.6α/8.3β versus 17.4α/8.3β TCRs in Tg mice. Flow cytometry of thymocytes (A–D) and splenocytes (E–H). (A and E) Representative CD4 versus CD8 dot plots. P values for thymic and splenic CD4:CD8 ratios were P < 0.0001 (for 17.4α/8.3β TCR-Tg vs. non-Tg mice) and P < 0.03 (for 17.6α/8.3β TCR-Tg vs. non-Tg mice). (B–D and F–H) Comparison of the intensity with which the different T cell subsets stain with anti–TCR-β or anti–TCR-α mAbs or with IGRP296–214 tetramers. Values on the panels correspond to percentage of cells within each gate that stained (B–D and F–H); only values for the 2 Tg strains are shown in these panels for ease of comparison) (mean ± SE). **P < 0.05 vs. 17.4α/8.3β Tg mice. Representative histograms are shown. A–D and E–H represent 3–6 mice per strain.

In 9-week-old mice were replaced in 20-week-old mice by Vα17.5 rearrangements (Figure 1A). It is important to emphasize here that the vast majority of these TCR-α rearrangements employing different Vα17 elements shared identical V-J junctional sequences, which suggests that Vα17 element usage per se plays a role in the avidity maturation of this T cell population (Supplemental Figure 1). Although these 3 novel Vα17 elements differ at only 5 residues (Figure 1B), they did not arise by somatic mutation, because all 3 coding sequences were found in NOD genomic DNA (data not shown). Computer modeling of the Vα17.4 TCR derived from the NRP-A7-reactive clone NY8.3 (30) suggests that, of the 4 substitutions which distinguish Vα17.5 from Vα17.6, 3 are located in CDR1 (residues 27–32; Figure 1B) and 1 lies on the lateral side of the Vα interface (data not shown). Given the exquisite heterogeneity of the TCR-β repertoires of both high- and low-avidity NRP-A7–reactive CD8+ T cells, these data suggest that avidity maturation of the NRP-A7–reactive T cell population is driven, at least in part, by competition among clonotypes expressing different Vα17 elements.

Effects of Vα17 element usage on T cell avidity for peptide-MHC. In order to characterize TCR-β heterodimers binding NRP-A7/K4 complexes with a range of avidities, we attempted to generate clones from islet-derived, NRP-A7/K4–reactive CD8+ T cell lines
that bound tetramers with either low or high avidity. Unexpectedly, we found that these clones, which were derived from single, originally tetramer-positive cells, reexpressed recombination-activating genes, transcribed new TCR rearrangements, and progressively lost their ability to bind tetramers (31). We therefore resorted to cotransfecting cDNAs encoding V\(\alpha_{17.6}\)-, V\(\alpha_{17.5}\)-, or V\(\alpha_{17.4}\)-MRD-J\(\times\)c42 \^ chains with a TCR-\(\beta\) cDNA derived from the diabeticogenic V\(\alpha_{17.4}\)-MRD-J\(\times\)c42 NY8.3 clone into a TCR-\(\alpha\)-\(\beta\), CD8^+ T cell hybridoma. Note that these 3 TCR-\(\alpha\) cDNAs were identical in sequence to the 3 corresponding TCR-\(\alpha\) rearrangements expressed by islet-associated CD8^+ T cells of NOD mice. Stable transfectants expressing comparable levels of TCRs and CD8 (Figure 1C) were tested for their ability to recognize IGRP\(_{206-214}\) (29) and NRP-V7 (a higher-avidity ligand of IGRP\(_{206-214}\) compared with IGRP\(_{206-214}\) and NRP-A7) (28). Whereas V\(\alpha_{17.6}\) TCR transfectants bound rather poorly to IGRP\(_{206-214}\) tetramers and only somewhat better to NRP-V7 tetramers (Figure 1D), V\(\alpha_{17.5}\) and V\(\alpha_{17.4}\) transfectants bound well to both tetramers (Figure 1D). Furthermore, Scatter plots of fluorescence units (median of CD8^+ population tetramer staining)/nM of tetramer versus fluorescence units of tetramer staining revealed that the V\(\alpha_{17.5}\) cells bound NRP-V7 tetramers with significantly lower \(K_e\) values (higher avidity) than did those expressing V\(\alpha_{17.4}\) or V\(\alpha_{17.6}\) TCRs, in that order (7 ± 1 vs. 13 ± 1 vs. 16 ± 2 nM, respectively; \(P < 0.05\)). Experiments employing tetramers labeled with the low-molecular weight fluorochrome Cy5 and purified by fast-performance liquid chromatography (FPLC) from small amounts of contaminating octamers, trimers, and dimers were consistent with the concept that V\(\alpha_{17.5}\) cells bind ligand with higher avidity than do V\(\alpha_{17.4}\) cells. Although the low-mean fluorescence intensity (mfi) of Cy5 labeling precluded measurements of avidity, these tetramers stained V\(\alpha_{17.5}\) cells with higher mfi than V\(\alpha_{17.4}\) cells (44 ± 7 vs. 20 ± 2; \(P < 0.05\)) (Figure 1E). In keeping with these results, V\(\alpha_{17.5}\) cells secreted more IL-2 in response to NRP-V7 and IGRP\(_{206-214}\), but not anti-CD3 mAb, than did V\(\alpha_{17.4}\) or V\(\alpha_{17.6}\) cells, indicating that V\(\alpha_{17.5}\) and V\(\alpha_{17.6}\) cells have the highest and lowest functional avidity for peptide-MHC, respectively (Figure 1F and data not shown).

Comparison of the TCR-\(\beta\) repertoires of the tetramer^+ and tetramer^− cells from 9- and 20-week-old NOD mice (Supplemental Figure 1) suggested that, notwithstanding its heterogeneity, TCR-\(\beta\) usage also contributes to the specificity of NRP-A7–reactive CD8^+ T cells. V\(\beta_{8.1}\) usage was significantly increased in the tetramer^− compared with the tetramer^+ subsets of both age groups (33% vs. 7%, respectively; \(P < 0.001\)). To determine whether the avidity hierarchy described above (highest for V\(\alpha_{17.5}\) and lowest for V\(\alpha_{17.6}\)) was a peculiarity of cells expressing the 8.3–TCR-\(\beta\) chain, we cotransfected V\(\alpha_{17.5}\)-J\(\times\)c42 or V\(\alpha_{17.6}\)-J\(\times\)c42 cDNAs with cDNAs encoding 2 additional V\(\beta\) genes, expressing 2 additional V\(\alpha\)-V\(\beta\) TCR transfectants (cloned from the tetramer-positive cells of 9-week-old NOD mice). We then compared the ability of stable transfectants expressing similar levels of CD3 and CD8 (data not shown) to bind tetramers. As expected, these additional V\(\alpha_{17.5}\) TCRs bound NRP-V7 tetramers better than did V\(\alpha_{17.6}\) TCRs in the context of identical TCR-\(\beta\) chains (Figure 1G shows representative results for 1 of these TCR-\(\beta\) chains). Thus, although we cannot exclude a contribution of TCR-\(\beta\) usage to avidity maturation of the NRP-A7–reactive CD8^+ T cell subset, our data are consistent with the concept that 1 mechanism for the increased avidity of the islet-associated T cells from 20-week-old mice is the replacement of low-avidity, V\(\alpha_{17.6}\) clonotypes with high-avidity, V\(\alpha_{17.5}\) ones. The data also show that the single amino acid substitution that distinguishes V\(\alpha_{17.5}\) from V\(\alpha_{17.4}\) (Figure 1B), probably distal to the antigen-binding site, has significant effects on the avidity of T cells for peptide-MHC.

Naive CD8^+ T cells expressing low-affinity V\(\alpha_{17.6}\) TCRs recognize peptide-MHC with low avidity and are not diabetogenic. To investigate the in vivo significance of the above differences in avidity, we compared the developmental biology of the 3 TCRs in TCR-Tg NOD mice. Figure 2 compares the thymic– and splenic–flow cytometry profiles of non-Tg NOD mice and NOD mice expressing an intermediate-avidity TCR (referred to as 8.3–17.4\(\alpha\)/8.3\(\beta\))...
or a low-avidity TCR (8.3–17.6α/8.3β). Note that high, intermediate, and low are used to rank the 3 different TCRs in relative rather than absolute terms. Results shown here are representative of at least 2 different Tg lines per strain (data not shown). 17.6α/8.3β mice displayed significantly higher thymic and splenic CD8α/CD4 ratios than did non-Tg NOD mice (Figure 2, A and E). This phenotype required expression of both chains of the Tg TCRs because it was not seen in single chain 8.3β−TCR-Tg mice (25), suggesting that 17.6α/8.3β-TCR fosters positive selection of CD8+ T cells, as expected. These differences between Tg and non-Tg mice were much more pronounced with 17.4α/8.3β mice (Figure 2A), suggesting that the 17.4α/8.3β TCR is more efficient at promoting positive selection of CD8+ cells than is the 17.6α/8.3β TCR. In support of this concept, upregulation of Tg TCR on double-positive thymocytes, which usually accompanies thymocyte-positive selection, was more pronounced in 17.4α/8.3β mice than in 17.6α/8.3β mice (Figure 2, B and C). The absolute number of CD8+ splenocytes expressing the Tg-TCR-β chain was slightly higher in 17.4α/8.3β mice, but not by much ([22 ± 4] × 10^6 vs. [16 ± 8] × 10^6). Strikingly, 17.6α/8.3β mice contained significantly fewer NRP-V7 tetramer-reactive CD8+ thymocytes and splenocytes than did 17.4α/8.3β mice, and the 17.6α/8.3β CD8+ cells that bound tetramers did so with much lower mfi than the tetramer-positive cells of 17.4α/8.3β mice (Figure 2, D and H). This apparent lower avidity of naive 17.6α/8.3β vs. 17.4α/8.3β CD8+ T cells was confirmed by comparing the tetramer reactivity and cytolytic activity of NRP-V7–specific CTL lines from both types of mice. Whereas virtually all 17.4α/8.3β CD8+ CTLs bound NRP-V7 tetramers, most 17.6α/8.3β CD8+ CTLs did not or did so weakly (Figure 3A). In addition, 17.6α/8.3β CD8+ CTLs were much less efficient at killing NRP-V7 peptide-pulsed, K4-transfected RAM-S (RAM-S-K4) cells targets than were 17.4α/8.3β CD8+ CTLs (Figure 3B). Furthermore, naive tetramer+ CD8+ splenocytes of 17.6α/8.3β mice secreted significantly lower levels of IFN-γ than did those derived from 17.4α/8.3β mice (on a cell by cell basis) (Figure 3C). In addition, 17.6α/8.3β mice developed diabetes with much lower incidence and with significantly delayed kinetics compared to 17.4α/8.3β mice (Figure 3D). We thus conclude that naive CD8+ T cells expressing the low-avidity 17.6α/8.3β TCR have low avidity for ligand and are not (or are only very poorly) diabetogenic.

**Partial but constitutive deletion of high-avidity Va17.5+ thymocytes in TCR-Tg mice.** The developmental fate of the high-avidity TCR (8.3–17.5α/8.3β) was strikingly different from that of its intermediate-avidity counterpart (8.3–17.4α/8.3β) (Figure 4). Note that these differences in developmental biology were seen in 2 different founder lines per strain (data not shown) and were not associated with differences in transgene copy number (~29 and ~31 vs. ~33 and ~20, respectively). Although both TCRs skewed T cell development toward the CD8− subset, the 17.4α/8.3β TCR did so much more efficiently (Figure 4A). Interestingly, 17.5α/8.3β

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**Figure 4**

Striking differences in the developmental biology of 17.4α/8.3β and 17.5α/8.3β T cells in Tg mice. Flow cytometry of thymocytes (A–D) and splenocytes (E–H). (A and E) Representative CD4 versus CD8 dot plots. P values for percentage of thymic and splenic CD8−CD4+ cells were as follows: P < 0.017 (for 17.4α/8.3β Tg vs. 17.5α/8.3β Tg, 5- to 9-week-old mice). (B–D and F–H) Comparison of mfi with which the different T cell subsets stain with anti–TCR-Vβ8 or anti–TCR-β mAbs or with IGRP206–214 tetramers. Values on the panels correspond to percentage of cells contained within each quadrant (A and E) or to percentage of cells within each gate that stained (B–D and F–H) (mean ± SE). **P < 0.05 versus 17.4α/8.3β Tg mice. Representative histograms are shown. A–D and E–H are representative of 4–5 mice per strain per age group. Note that the representative dot plots and histograms corresponding to 17.4α/8.3β Tg mice are the same as those shown in Figure 2. The data are rendered twice to facilitate comparisons with results obtained with Tg mice bearing lower− (Figure 2) or higher-avidity T cells (this figure).
CD8+CD4+ thymocytes expressed lower levels of Tg-TCRβ chains and total TCRs than did 17.4α/8.3β CD8+CD4+ thymocytes (Figure 4, B and C) (P < 0.007 for mfi values corresponding to total TCRs; data not shown). Such downregulation of the Tg TCR was already seen on double-positive thymocytes (Figure 4B), was inherited by peripheral CD8+ (but not CD4+) cells (Figure 4, F and G; P < 0.004 for mfi values, data not shown), and was confirmed by staining thymocytes with IGRP206–214 tetramers (Figure 4D; P = 0.05 for mfi values of tetramer staining; not shown). This TCRαβ phenotype was not age dependent, as it was already seen in 0.5-week-old animals (Figure 4D). We conclude that thymocytes expressing high levels of 8.3–17.5α/8.3β TCR, which is also the TCR that affords persistent splenic flow cytometry profiles (CD4, CD8, and Vβ8) of RAG-2−/− mice (17.4α/8.3β-Tg) and 8 (17.5α/8.3β-Tg) RAG-2−/− mice. (E) Representative splenic flow cytometry profiles (CD4, CD8, and Vβ8) of RAG-2−/− mice.

Depletion of the peripheral Vβ17.5+ T cell pool. Analyses of splenic T cells suggested that the 17.5α/8.3β CD8+ cells that were selectively tolerant also undergo peripheral deletion. The CD8+CD4+ cell ratios in the spleens of 5- to 9-week-old 17.4α/8.3β-Tg mice were significantly higher than those seen in age-matched 17.5α/8.3β-Tg mice (−2 vs. −0.33, P = 0.017; Figure 4E). In addition, whereas most CD8+ splenocytes from 17.4α/8.3β-Tg mice bound IGRP206–214 tetramers, most CD8+ splenocytes from 17.5α/8.3β-Tg mice did not (P < 0.05; Figure 4H). This phenotype was not caused by thymic tolerance, as the CD8+CD4+ ratios in 17.5α/8.3β-Tg spleens were significantly lower than those seen in 17.5α/8.3β-Tg thymi (−0.33 vs. −1; P < 0.04; Figure 4A) and 17.5α/8.3β CD8+ thymocytes bound tetramers (Figure 4D). Notably, this phenotype was absent in the spleens of 0.5-week-old 17.5α/8.3β-Tg animals; these young mice were shown to have more CD8+ cells than CD4+ cells (Figure 4E), and their CD8+ cells bound tetramers (Figure 4H). In addition, the number of CD8+ splenocytes increased significantly with age only in 17.4α/8.3β-Tg mice (17.4α/8.3β-Tg mice: from 1.6 × 106 cells at 1.5 weeks to [23 ± 3] × 106 cells at 9 weeks; 17.5α/8.3β-Tg mice: from [0.3 ± 0.3] × 106 cells at 0.5 weeks to [2.3 ± 0.2] × 106 cells at 9 weeks; P < 0.05). These phenomena of severely reduced numbers of splenic CD8+ cells and decreasing CD8+CD4+ cell ratios in spleen compared with thymus in older 17.5α/8.3β-Tg animals are all consistent with mechanisms of peripheral tolerance emerging to suppress the transgene-expressing high-avidity T cells.

A small contingent of peripheral high-avidity Vβ17.5+ T cells evades tolerance and triggers diabetes. Despite expressing lower levels of total TCRs (Figure 4G), the few tetramer-reactive CD8+ splenocytes of 17.5α/8.3β-Tg mice bound IGRP206–214 tetramers with higher avidity than did 17.4α/8.3βCD8+ splenocytes (K0.5 of 5.9 ± 0.5 vs. 14.9 ± 2 nM). In addition, when equal numbers of tetramer-reactive cells were stimulated with IGRP206–214, 17.5α/8.3β CD8+ splenocytes proliferated more efficiently than did 17.4α/8.3β CD8+ splenocytes (Figure 5A), particularly at low concentrations of antigen. Strikingly, despite having approximately 28 times fewer IGRP206–214-reactive CD8+ cells and lower insulitis scores than 17.4α/8.3β-Tg mice (in prediabetic animals; Figure 5B), 17.5α/8.3β- and 17.4α/8.3β-Tg mice developed diabetes with similar penetrance (Figure 5C).
Furthermore, whereas abrogation of recombination activating gene (RAG) expression decreased the diabetogenic potential of the 17.4α/8.3β-TCR (owing to partial dependence of 17.4α/8.3β-CD8+ cells on CD4+ T cell help) (30) (Figure 5D), it did not impair the pathogenicity of the severely depleted CD8+ T cell pool of 17.5α/8.3β-Tg mice (Figure 5, D and E). Because Vx17.5+ cells progressively replace Vx17.6+ cells in the islets of non-Tg NOD mice, we propose that in non-Tg mice the islet microenvironment selects the few high-avidity CD8+ cells that survive peripheral tolerance from further effects and fosters the manifestation of their high avidity (outgrowth of the fittest and most pathogenic). Alternatively, some of these high-avidity clonotypes escape tolerance by (or upon) undergoing productive antigen-induced activation in the regional lymph nodes and/or pancreatic islets. Whatever the mechanisms, these results explain why, in the islets of non-Tg NOD mice, high-avidity cells only come to predominate with age.

**Discussion**

The natural evolution of any lymphocyte response results from competition among cells expressing antigen receptors of varying affinities. B cell affinity maturation is driven by somatic hypermutation of the preimmune low-affinity antigen-receptor repertoire (9, 17, 32). Since T lymphocytes cannot somatically mutate their TCRs, T cell avidity maturation relies exclusively on competition among clonotypes with highly heterogeneous TCR structures. As a result, the mechanisms responsible for the recruitment of predominantly low-avidity T cells in primary immune responses and their progressive replacement by high-avidity clonotypes upon antigen reencounters are poorly understood (18–20). Here we have provided a developmental basis for this phenomenon by studying the diabetogenic CD8+ T cell population that targets the β cell antigen IGRP (30) and undergoes avidity maturation as disease progresses (20). Importantly, our observations shed new light on our understanding of how pathogenic autoimmunity develops in a genetically predisposed background.

Studies of TCR usage by islet-associated, NRP-A7 tetramer-reactive CD8+ T cells of prediabetic NOD mice showed that most of these T cells employ Vx17-Jα42 rearrangements with identical CDR3 sequences. This contrasts with the heterogeneity of TCR-β usage by these T cells, which suggests that Vx17-Jα42 chains play a major (albeit perhaps not exclusive) role in defining the antigen specificity of this T cell population. Detailed analyses of the Vx17-Jα42 sequences used by high- compared with low-avidity subpopulations suggest that, in addition to helping define their antigenic specificity, TCR-β chain usage also contributes to defining the avidity of IGRF206–214-reactive CD8+ T cells for peptide-MHC. Specifically, avidity maturation of this T cell population was associated with changes in the frequency with which 3 Vx17 elements were used. As mice aged, islet-associated T cells expressing Vx17.5 replaced those expressing Vx17.6. Studies with TCR transfectants confirmed that Vx17.5+ cells coexpressing the 8.3-TCR-β chain (or 2 other β chains) recognize peptide-MHC with higher physical and functional avidity than do those expressing Vx17.4+ or Vx17.6+ TCRs. Studies in TCR-Tg NOD mice were also consistent with these observations. The Vx17.6+ TCR fostered positive selection of CD8+ T cells, but less efficiently than did the Vx17.4+ TCR. In addition, whereas naïve Vx17.6+ T cells could readily expand in response to stimulation with NRP-V7 in the presence of rIL-2, the resulting CTLs killed peptide-pulsed targets and bound peptide-MHC tetramers much less efficiently than did CTLs expressing the Vx17.4+ TCR. In contrast, expression of the Vx17.5+ TCR promoted central and peripheral tolerance of CD8+ T cells. In addition, the few Tg Vx17.5+ T cells that survived tolerance recognized peptide-MHC with significantly higher avidity than the nontolerized cells of Vx17.4+ TCR-Tg mice. Furthermore, Vx17.5+ T cells were much more diabetogenic than were Vx17.4+ or Vx17.6+ T cells. Taken together, these observations support the views that Vx17.5+ T cells recognize peptide-MHC with higher avidity than do Vx17.4+ or Vx17.6+ T cells and that avidity, as measured with tetramers, is a determinant of both competitiveness and pathogenic potential. It is worth noting here that, although it has previously been shown that the frequency of spontaneous encephalomyelitis in 2 EAE-susceptible strains of TCR-Tg mice also correlates with the affinity of the Tg TCRs for peptide-MHC (33), this, is, to the best of our knowledge, the first report of a high-avidity, pathogenic autoreactive TCR that contributes to the progression of spontaneous autoimmunity despite undergoing tolerance. Our observations further indicate that small, germ-line-encoded molecular changes distal to the antigen-binding site of the TCR can have profound effects on the developmental biology of individual clonotypes. Since some of the molecular differences that are associated with avidity maturation of the IGRF206–214-reactive T cell population map to the CDR1α, and since some CDR1α residues (i.e., position 27) interact directly with MHC molecules (34–36), it is possible that the above differences in biophysical, functional, and developmental behavior are determined by differences in the strength with which these TCRs engage MHC molecules in the absence of cognate peptide ligand. In support of this hypothesis, we have observed that, unlike Vx17.4+ CTLs, Vx17.5+ CTLs can kill TUM-pulsed RMA-S-K4 cells and can produce substantial amounts of IFN-γ in response to TUM-pulsed dendritic cells (our unpublished observations).

The evolutionary advantage of involving low-avidity T and B lymphocytes in primary immune responses lies in the conformational flexibility of the binding site of their antigen receptors, which allows a finite repertoire of receptors to cope with a limitless array of epitopes (1–10). Whereas B cell affinity maturation can proceed in the absence of high-affinity antigen receptors in the preimmune repertoire (37). Our data provide evidence that both self and foreign antigens predominantly recruit low-avidity T cells (18–20, 22). Our observation that tolerance selectively decreases the size of the high-avidity IGRF206–214-reactive T cell pool provides one explanation for the prevalent recruitment of low-avidity clonotypes in the primary autoimmune response and for the progressive replacement of these low-avidity clonotypes with higher-avidity clonotypes as the animals age. The observation that the Vx17.6+ TCR was not as good a positive selector of CD8+ thymocytes as the Vx17.4+ TCR is compatible with the Vx17.6+ rearrangement accounting for about twice the total number of Vx17.4+ cDNAs derived from the low-avidity, islet-associated CD8+ T cell pool of non-Tg 9-week-old NOD mice.

It has been established that thymocytes expressing TCRs recognizing self-antigen–MHC complexes with high affinity/avidity undergo central deletion (a process also referred to as thymocyte negative selection). In contrast, thymocytes expressing low-affinity TCRs for the same complexes differentiate into mature T cells and populate the peripheral lymphoid organs, where they become available for recognizing foreign antigens (37). Our data provide...
direct evidence that autoreactive T cells with relatively high avidity and pathogenic potential can escape thymocyte negative selection and elicit autoimmunity in the absence of adequate peripheral regulation. Whereas thymocytes expressing high levels of the 8.3–17.6a/8.3β TCR were deleted, those expressing lower levels of the same TCR survived and migrated to the periphery as TCRαβ cells. These T cells were found to recognize self-peptide-MHC with an avidity that was high enough to afford exquisite pathogenic activity but below the threshold required for thymocyte negative selection. Accordingly, these “high-avidity” cells that survive thymocyte negative selection are not truly high avidity but rather cells that can be classified, along with T cells expressing the 8.3–17.4a/8.3β TCR, as cells of “intermediate avidity” with pathogenic potential.

We do not think that central tolerance of V17.5α IGRPαβ reactive CD8+ T cells is related to the fact that IGRP is a self protein for 3 reasons. First, although β cell–specific proteins can be expressed in thymic epithelium (38, 39), we could not detect the presence of either of the 2 isoforms of IGRP mRNA (or other hypothetical splice variants) in unsorted NOD thymic stroma or purified UEA-1/AIRE+ cells by RT-PCR (our unpublished data). Second, the occurrence of central tolerance preceded the initiation of cross-presentation of IGRPAβ
d in the periphery (40). Third, whereas the magnitude of peripheral tolerance was paralleled by an age-dependent increase in the magnitude of cross-presentation of IGRPAβ
d in the pancreatic lymph nodes (40), central tolerance was not.

In sum, regardless of the role of IGRPAβ
d in tolerance of high-avidity clonotypes, our work demonstrates that a small contingent of self reactive T cells with relatively high avidity can escape thymic negative selection. In the absence of peripheral regulation, these cells gain access to the target organ and, upon undergoing local expansion and/or accumulation, contribute to the progression of benign autoimmune inflammation to overt clinical disease. These observations provide an explanation of how potentially pathogenic autoimmunity develops in the periphery.

Methods

Mice. 17.4a/8.3β (8.3-NOD) mice have been described (30). 17.6a/8.3β- and 17.5a/8.3β-Tg mice were produced by microinjection of TCR transgenes (as described in ref. 30) into NOD zygotes by or by intercrossing 17.6α and 8.3β-Tg NOD mice. Expression of the 17.6α chain was confirmed by assessing the ability of the transgene to foster normal thymocyte development in the presence of this anti-CD8 mAb does not change the percentage or absolute number of cells that bind the tetramers. After washing, cells were resuspended in 100 μl of wash media and analyzed with a flow cytometer. To calculate the apparent Kd values, we plotted the negative reciprocal of the slope of the line fit to Scatchard plots of fluorescence units (median of CD8+ population tetramer staining)/nM versus fluorescence units. As expected, the Scatchard plots of tetramer binding obtained at the range of concentrations tested was curvilinear. For low-avidity transfectants (i.e., those expressing the 17.6 TCR), stains performed with the 4 lowest concentrations of tetramers were not informative, as significant numbers of staining events fell outside the tetramer-positive gate (defined with the negative-control tetramer TUM/Kd). Accordingly, Kd values were calculated from the slope of the line fit to Scatchard plots obtained with the 4 highest concentrations of tetramers. It should be noted that, although the absolute Kd values for specific groups of transfectants varied considerably depending on the slope that was chosen for calculation (higher vs. lower 4 concentrations), the relative differences were unaffected by the choice of slope. For example, the Kd values for 17.4 and 17.5 TCR–transfectants obtained with the lowest 4 concentrations of tetramer were: 3.4± 0.7 vs. 1.77 ± 0.2, respectively. This corresponds to a Kd 17.4:17.5 ratio of 1.94, which is statistically similar to the ratio calculated using Kd values obtained with the 4 highest concentrations of tetramers (1.76).

TCR repertoires. Pancreatic islets pooled from 15–20 NOD mice per age group were cultured in the presence of 0.5 U/ml of Takeda rIL-2 for 6–7 days, stained with NRP-A7 tetramer and anti-CD8 mAb, and sorted into tetramer+ and tetramer− CD8+ cells by flow cytometry. TCR-specific cDNA libraries were generated by anchor-PCR and analyzed by sequencing (25). Vβ7 gene element usage by NOD splenic CD8+ cells was determined by sequencing Vβ7 TCR−α cDNAs (generated by RT-PCR). 3D models of 8.3-like TCRs were obtained with a Swiss-pdb viewer using the 2C TCR crystal coordinates.

TCR transfectants. Full-length TCR cDNAs were expressed in a TCR−TCRζ−CD8αβ T cell hybridoma. TCR expression was verified by flow cytometry using anti-Vβ (MR5-2) and anti-CD3ε (145-2C11) mAbs. Peptide recognition by transfectants was measured by IL-2 production using irradiated NOD splenocytes as APCs.

Flow cytometry. Thymi and spleens were analyzed by 3-color flow cytometry using anti-CD8-PerCP (53-6.7), anti-CD4-FTTC/PE (IM7), and tetramer-PE, anti-Vβ1/8.2-FTTC (MR5-2), or anti-TCR-β–FTTC (H57-597) analysis. All mAbs were from Pharmingen.

Proliferation, cytokine, and cytotoxicity assays. Proliferation and cytokine secretion assays were done using peptide-pulsed, irradiated NOD splenocytes (27) or anti-CD3/anti-CD28 mAb–coated microbeads (Dynal Biotech) following the manufacturer’s instructions. Values obtained with TUM were subtracted. Cytotoxicity assays were performed using peptide-pulsed (1 μg/ml) [51Cr]-sodium chromate-labeled RMA-S-K2 cells as targets (27).

Pathology. Frozen or formalin-fixed, paraffin-embedded pancreas sections were stained with H&E and scored for islets (30).

Diabetes. Diabetes was monitored by measuring urine glucose levels with Diastix (Miles Laboratories Inc.) twice weekly. Animals were considered diabetic after 2 consecutive readings greater than or equal to +3.

Statistical analyses. Data were compared using linear regression, variance analysis, the Mann-Whitney U test, or the χ² test.

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