The regulatory role of DR4 in a spontaneous diabetes DQ8 transgenic model

Li Wen,1 Ning-Yuan Chen,1 Jie Tang,1 Robert Sherwin,1 and F. Susan Wong2

1Section of Endocrinology, Department of Internal Medicine, Yale University School of Medicine, New Haven, Connecticut
2Department of Pathology and Microbiology, University of Bristol, Bristol, United Kingdom

Address correspondence to: Li Wen, Section of Endocrinology, Department of Internal Medicine, Yale University School of Medicine, Box 208020, 330 Cedar Street, New Haven, Connecticut 06520, USA.
Phone: (203) 785-7186; Fax: (203) 737-5558; E-mail: li.wen@yale.edu.

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MHC class II molecules are critical determinants of genetic susceptibility to human type 1 diabetes. In patients, the most common haplotype contains the DRA1*0101-DRB1*0401 (DR4) and DQA1*0301-DQB1*0302 (DQ8) loci. To assess directly the relative roles of HLA-DQ8 and DR4 for diabetes development in vivo, we generated C57BL/6 transgenic mice that lack endogenous mouse MHC class II molecules but express HLA-DQ8 and/or DR4. Neither HLA-DQ nor HLA-DR transgenic mice developed insulitis or spontaneous diabetes. However, when they were crossed to transgenic mice (C57BL/6) expressing the B7.1 costimulatory molecules on pancreatic β cells that do not normally develop diabetes, T cells from these double transgenic mice were no longer tolerant to islet autoantigens. The majority of DQ8/RIp-B7 mice developed spontaneous diabetes, whereas only 25% of DR4/RIp-B7 mice did so. Interestingly, when DQ8 and DR4 were coexpressed (DQ8DR4/RIp-B7), only 23% of these mice developed diabetes, an incidence indistinguishable from the DR4/RIp-B7 mice. T cells from both DR4/RIp-B7 and DQ8DR4/RIp-B7 mice, unlike those from DQ8/RIp-B7 mice, exhibited a Th2-like phenotype. Thus, the expression of DR4 appeared to downregulate DQ8-restricted autoreactive T cells in DQ8DR4/RIp-B7 mice. Our data suggest that although both DQ8 and DR4 can promote spontaneous diabetes in mice with a non-autoimmune-prone genetic background, the diabetogenic effect of the DQ8 allele is much greater, whereas DR4 expression downregulates the diabetogenic effect of DQ8, perhaps by enhancing Th2-like immune responses.


Introduction

Type 1 diabetes mellitus (T1DM), also known as insulin-dependent diabetes mellitus (IDDM), is a polygenic disease, and studies have confirmed that the main locus defining genetic susceptibility is found encoded within the MHC region on human chromosome 6 (1–3). A key function of MHC molecules is to present antigens (Ag’s) to T cells. Specifically, MHC class I molecules present Ag’s to CD8+ T cells, whereas MHC class II molecules present Ag’s to CD4+ T cells. Evidence derived from the nonobese diabetic (NOD) mouse model of T1DM demonstrating both CD4+ and CD8+ T cells are normally required to induce disease (4–7) suggests that the MHC is an important region of genetic susceptibility because of its Ag-presenting function.

CD4+ T cells play an important role in regulating the functions of the immune system, and numerous studies in both human and animals have shown the contribution of Th1 or Th2 to the pathogenesis of a wide variety of diseases. The dichotomy of Th1 and Th2 cells in the development of T1DM has also been well documented (8–15). Most of the studies demonstrate that Th1 cells contribute to the immunopathogenesis of islet β-cell destruction, whereas Th2 cells mediate disease protection. Based on these studies, new strategies in oral or nasal immunotherapy (which promote Th2 immune responses) for diabetes prevention have been tested in experimental animals (16–19), and some of the regimens are currently in clinical trials. On the other hand, other studies have also shown that Th2 cells have the potential to be diabetogenic, as well (20–22). Thus, like most autoimmune disorders, the pathogenesis of T1DM is very complex. In addition to the role of CD4+ T cells as pathogenic cells, a small subset of CD4+ T cells that are positive for CD25 has been reported recently as regulatory cells that prevent or downregulate the development of autoimmunity (23, 24).

The MHC class II loci have been investigated extensively as determinants of genetic susceptibility both in human T1DM and the NOD mouse model. In NOD mice, the predisposing allele is I-Ag7, characterized by a substitution of serine for the charged aspartic acid residue at position 57 of the β chain. However, the picture is likely to be more complex. For example, many studies suggest that a single residue does not determine susceptibility and that the combination of DQA1 and DQB1 determinants is important (25–28). Whereas a
number of DQA1-DQB1 pairs are found in Caucasian populations, the most common haplotype seen in patients with T1DM is the DRA1*0101-DRB1*0401 (DR4)-DQA1*0301-DQB1*0302 (DQ8) haplotype. The predisposing DQ8 allele, the homologue of I-A in the mouse, has a structure similar to I-Aβ.

Transgenic mice that carry only human MHC class II DQ8 molecules and are deficient in murine MHC class II molecules (DQ8+/mII−) have been generated recently (29–32). However, in studies of diabetes they have been used mainly to identify T-cell determinants of pancreatic β-cell autoantigens (29, 31, 32). In no case did these transgenic mice (HLA-DQ8 and/or HLA-DR4) develop diabetes spontaneously, even if they were expressed on the NOD genetic background (32). To investigate whether we could break the tolerance exhibited by the HLA-DQ8 and HLA-DR4 transgenic mice that were not on the NOD genetic background, by providing locally the costimulatory signal B7-1, we introduced this costimulatory molecule using the rat insulin promoter (RIP). The RIP-B7.1 transgene alone normally does not cause diabetes on the C57BL/6 genetic background (33). However, we have reported that the expression of DQ8 in which murine MHC class II was knocked out provoked autoimmune diabetes in the majority of non–diabetes-prone RIP-B7.1 C57BL/6 mice (34). In this study we compared the effect of DQ8 with that of DR4, alone or in combination, on disease development. Our goal was to use this “humanized” spontaneous diabetes model system to study the relative immunopathological effect of individual diabetes susceptibility genes in strong linkage disequilibrium. The novel information regarding the role of DR4 in disease development is important for the clinical development of effective immunotherapy, especially because of the fact that DQ8/DR4 HLA genes are commonly found in diabetic patients.

Methods
Generation of DR4+/mII−/RIP-B7.1 and DQ8+/DR4+/mII−/RIP-B7.1 transgenic mice. HLA-DQA1*0301/ DQB1*0302 (DQ8) transgenic, murine MHC class II molecule–deficient (mII−), and RIP-B7.1 C57BL/6 mice were generated as described previously (34). HLA-DRB1*0401 (DR4) transgenic mice (B10) were kindly provided by Wicker and Zaller (Merck & Co. Inc., Rahway, New Jersey, USA) (35). After back-crossing these mice to C57BL/6 genetic background (N4 to N5), we introduced disrupted murine MHC class II molecule (mII−) and B7.1 transgene on pancreatic β cells (DR4+/mII−/RIP-B7.1), both on the C57BL/6 background. We then intercrossed DQ8+/mII−/RIP-B7.1 mice with DR4+/mII−/RIP-B7.1 mice in order to obtain DQ8+/DR4+/mII−/RIP-B7.1 mice and all different types of control mice. The expression of the DQ8 and/or DR4 transgene and murine class II I-Aβ molecules were screened by flow-cytometric analysis of peripheral blood lymphocytes (PBL), and the RIP-B7.1 transgene was screened using PCR (34). All the mice used in this study were littermates. As mentioned above, all the different types of control mice, such as DQ8+/DR4+/mII−/RIP-B7.1, DQ8+/mII−/RIP-B7.1, DR4+/mII−/RIP-B7.1, and mII−/RIP-B7.1, were derived from the same pairs for breeding. The possibility that transgenic mice expressed hybrid I-Aα/DQ8β and DR4α/Eβ was excluded by negative staining of PBL with mAb’s specific for I-Aα (AF6-120.1; PharMingen, San Diego, California, USA) and I-E (Y17; Charlie Janeway), respectively.

To establish the C57BL/6 genetic background of the mice used in this study, we also screened the genomic markers in these mice using microsatellite analysis. Seventeen markers were used for the screening: idd2 to idd15 (36) and three additional markers (D1Mit232, D1Mit22, and D7Nds6) that were reported to be associated with diabetes in the C57BL/6 genetic background (37).

The mice were housed in specific pathogen-free facilities, and all experiments were undertaken in accordance with approved Yale Animal Care and Use Committee protocols.

Purification of CD4+ T cells. After removing erythrocytes, total splenocytes were incubated with anti-CD8 mAb (TIB-105; rat IgG2a) on ice for 30 minutes and washed once with cold Click’s medium. The cells were then incubated with magnetic beads conjugated with goat anti-mouse IgG/IgM and goat anti-rat IgG (PerSeptive Biosystems, Framingham, Massachusetts, USA) on ice with gentle agitation for 45 minutes. B cells and CD8+ cells were removed by using a magnetic plate (PerSeptive Biosystems). The purity of the CD4+ population using this method is routinely over 90%.

Proliferation and cytokine assays. Splenic T cells (105/well) from diabetic or nondiabetic mice were assayed for antigenic response against three main putative β-cell autoantigens—GAD (30), insulin (Eli Lilly and Co., Indianapolis, Indiana, USA), and IA-2 (kindly provided by Peter van Endert, Hôpital Necker, Paris, France), as described previously. Secreted cytokine proteins (IL-4, IL-6, IL-10, and IFN-γ) from those responses were measured by ELISA using mAb’s and recommended protocols (PharMingen).

Adoptive-transfer experiments. Mice used for these experiments were irradiated (6 Gy) one day before the adoptive transfer. Diabetic splenocytes were injected intravenously into the recipients (107/recipient). All the mice were monitored for glycosuria, and the experiments were terminated 8 weeks after adoptive transfer unless the mice developed diabetes, which was confirmed by blood glucose (>250 mg/dl), and they were sacrificed immediately.

Generation of bone marrow chimera. MII−/RIP-B7.1 mice were used as recipients in the chimeric experiment, and they were irradiated (8.5 Gy) the day before the bone marrow (BM) reconstitution. Our preliminary BM chimera study showed a high mortality rate in recipients that received over 9 Gy irradiation. This may be due to the fact that the recipients are deficient for
MHC class II and therefore they are immunocompromised. BM cells harvested from nondiabetic DQ8+/mII–/RIP-B7.1, DR4+/mII–/RIP-B7.1, and DQ8+/DR4+/mII–/RIP-B7.1 donor mice (8–12 weeks of age) were injected intravenously into the recipients (10^7/recipient). All the mice were monitored for glycosuria, and the experiments were terminated 8 weeks after adoptive transfer unless the mice developed diabetes, which was confirmed by blood glucose, and they were sacrificed immediately.

RT-PCR for the detection of expression of chemokine receptors. Purified CD4+ T cells (2×10^6) from DQ8+/DR4+/mII–/RIP-B7.1 and DQ8+/mII–/RIP-B7.1 mice were activated by anti-CD3 cross-linking (24 hours), and total cellular RNA was isolated using TriZol (Life Technologies Inc., Grand Island, New York, USA). The cDNA was synthesized using oligo-dT primer (Amersham Pharmacia Biotech Inc., Piscataway, New Jersey, USA), which recognizes the poly(A) tail of mRNA. PCR was performed using specific primers for mouse CCR5 and CCR2 and same amount of cDNA templates. Standard conditions were used for the PCR, and an internal control, CD3ε, was used for normalization of RNA levels.

Ig isotype assay. Blood samples were taken, and sera were separated when experiments were terminated. The levels of Ig isotypes were measured using ELISA, as described previously (38). All the reagents for this assay were purchased from Southern Biotechnology Associates (Birmingham, Alabama, USA).

Immunohistology. Pancreata, kidneys, livers, and salivary glands from all the mice used in this study were examined by immunohistochemistry as described previously (39).

Results

Expression of DR4-single and DQ8/DR4-double transgenes in mice lacking endogenous mouse MHC class II molecules. We have shown recently that DQ8+/mII–/RIP-B7.1 C57BL/6 mice develop a high incidence of spontaneous diabetes (34). In the present study we generated DR4+/mII– mice, since the most common haplotype seen in patients with T1DM contains DR4 and DQ8. As shown in Figure 1, the expression of the HLA-DR4 transgene in mice deficient in murine MHC class II molecules partially restored CD4+ T cells to a level similar to that in the DQ8+/mII– transgenic mice. The TCR variable β chain (Vβ) repertoire of the DR4+/mII– mice showed no significant differences as compared with wild-type (MHC class II–sufficient) mice (C57BL/6), except that the total number of CD4+ T cells was lower (approximately 25% of total T cells). This was also the case in the DQ8+/mII– mice (30). More CD4+ T cells (about twofold) were selected in mice expressing both DQ8 and DR4 (Figure 1), however, and the coexpression of DQ8 and DR4 did not appear to skew the TCR Vβ repertoire (data not shown). The expression level of DQ8 and DR4 in DQ8/DR4+/mII– mice was indistinguishable, and neither DR4+/mII– nor DQ8/DR4+/mII–
mice (like their DQ8+/mII− counterparts) developed spontaneous insulitis or diabetes over a period of 10 months’ observation (data not shown).

Incidence of spontaneous diabetes in DR4+/mII−/RIP-B7.1 and DQ8+/mII−/RIP-B7.1 mice and in mice coexpressing DR4 and DQ8. The introduction of the RIP-B7.1 transgene into DR4+/mII− mice caused the development of spontaneous diabetes in 25% (5 of 20) of DR4+/mII−/RIP-B7.1 mice. Like the DQ8+/mII−/RIP-B7.1 mice, there was no sex preference, and disease development occurred with similar kinetics in DR4+/mII−/RIP-B7.1 mice (Figure 2). However, the overall incidence of diabetes in DR4+/mII−/RIP-B7.1 mice was much lower than that found in DQ8+/mII−/RIP-B7.1 mice (Figure 2). The incidence of diabetes was threefold higher in the DQ8+/mII−/RIP-B7.1 mice (73%; 11 of 15) (Figure 2), which are values similar to those in our recent report (34). Of particular interest, when DR4 was introduced into DQ8+/mII−/RIP-B7.1 mice, the incidence of diabetes was markedly reduced to a level similar to that found in DR4+/mII−/RIP-B7.1 mice (23%; 5 of 22) (Figure 2).

None of the murine MHC class II–sufficient mice (DR4+/mII+/RIP-B7.1 or DQ8+/DR4+/mII+/RIP-B7.1) developed diabetes (data not shown). As in previous experiments, one of ten (10%) of the mouse MHC class II–deficient RIP-B7.1 mice (mII−/RIP-B7.1) also developed diabetes (Figure 2).

Fewer CD8+ T-cell infiltrates in pancreatic islets of DR4+/mII−/RIP-B7.1 and DQ8+/DR4+/mII−/RIP-B7.1 diabetic mice compared with diabetic DQ8+/mII−/RIP-B7.1 mice. Pancreata from mice that developed diabetes were analyzed for lymphocytic infiltration by immunohistochemistry staining. Although the number of peripheral CD8+ T cells was very similar between DQ8+/mII−/RIP-B7.1 and DR4+/mII−/RIP-B7.1 mice, it appeared that fewer CD8+ T cells were found in diabetic DR4+/mII−/RIP-B7.1 islet infiltrates than were seen in diabetic DQ8+/mII−/RIP-B7.1 islet infiltrates (Figure 3). Fewer CD8+ T-cell infiltrates were also seen in diabetic DQ8+/DR4+/mII−/RIP-B7.1 mice (Figure 3). However, in this case coexpression of DQ8 and DR4 led to better selection and maturation of CD4+ T cells (see above) on the murine MHC class II–deficient background; therefore, the ratio of CD4/CD8 in DQ8+/DR4+/mII−/RIP-B7.1 mice is relatively greater than that in DQ8+/mII−/RIP-B7.1 and DR4+/mII−/RIP-B7.1 mice.

Insulitis and/or diabetes could be transferred by diabetic splenocytes of DR4+/mII−/RIP-B7.1 or DQ8+/DR4+/mII−/RIP-B7.1 mice. Because the expression of the RIP-B7.1 transgene is required for the adoptive transfer model in the RIP-B7.1 transgenic systems (34, 40, 41), adoptive transfer recipients in this study were RIP-B7.1–transgene positive. We have shown that diabetic DQ8+/mII−/RIP-B7.1 splenocytes can adoptively transfer diabetes to irradiated syngeneic recipients (34). In a similar manner, splenocytes (depleted of erythrocytes) from diabetic DR4+/mII−/RIP-B7.1 and DQ8+/DR4+/mII−/RIP-B7.1 mice were adoptively transferred into sublethally irradiated DR4+/mII−/RIP-B7.1 mice and DQ8+/DR4+/mII−/RIP-B7.1 mice, respectively. The experiments were terminated 8 weeks after adoptive transfer. Diabetes developed in 38% of the DR4+/mII−/RIP-B7.1 recipients (three of eight; 4, 6, and 6 weeks after transfer) that received DR4+/mII−/RIP-B7.1 diabetic splenocytes and in four of eight (50%; 4, 5, 6, and 6 weeks after transfer) of DQ8+/DR4+/mII−/RIP-B7.1 recipients given diabetic DQ8+/DR4+/mII−/RIP-B7.1 splenocytes.

**Figure 3**
Immunohistochemistry staining of pancreatic sections of diabetic mice, with the diabetic DQ8+/mII−/RIP-B7 mouse in the upper panel, diabetic DR4+/mII−/RIP-B7 mouse in the middle panel, and diabetic DQ8+/DR4+/mII−/RIP-B7 mouse in the lower panel. Islet infiltrates were stained for CD4+ T cells (left), CD8+ T cells (middle), and B220+ B cells (right).
Reconstitution of insulitis by nondiabetic BM cells from DQ8+/mII–/RIP-B7.1, DR4+/mII–/RIP-B7.1 and DQ8+ DR4+/mII–/RIP-B7.1 mice. To evaluate the role of BM-derived human MHC class II+ cells in disease development, we generated BM chimeras by transferring BM cells (after depletion of erythrocytes) from human MHC class II+/RIP-B7.1 transgenic mice (8 to 12 weeks of age) into lethally irradiated human MHC class II–/RIP-B7.1 transgenic recipients (10 to 12 weeks of age, n = 3–5/group). We aimed to study the prediabetic phase in these experiments and chose 8 weeks as the observation period. All the recipients except one (which received DQ8' DR4+/mII–/RIP-B7.1 BM cells), were diabetes free in the 8-week observation period. The reconstitution was successful and stable as determined by the expression of the appropriate human MHC class II molecules on B220+ cells in human MHC class II–negative recipients (Figure 4a) and by the positively selected mature CD4+ or CD8+ splenocytes (Figure 4a). However, when we analyzed the pancreata of the recipients, a different picture was revealed. Although insulitis was found in all the recipients, the severity of the insulitis was different: insulitis in the DQ8+/mII–/RIP-B7.1 and DQ8+ DR4+/mII–/RIP-B7.1 mice was similar and much greater than in DR4+/mII–/RIP-B7.1 (Figure 4b). It is interesting that the only recipient who developed diabetes at the time when the experiment was terminated exhibited infiltration in both islets and exocrine tissue (data not shown). The same cohort of the experiments was repeated using anti-CD3–depleted BM cells, and very similar results were obtained. This indicates that the T cells infiltrating the islets are more likely derived from BM stem cells rather than the 0.5–1% of CD3+ cells in the untreated BM. BM chimeras were also generated using BM derived from mII–/RIP-B7.1 or DQ6+/mII–/RIP-B7.1 mice as controls, and none of the chimeras in the control groups developed insulitis when the experiment was terminated (8 weeks, data not shown).

Spontaneous GAD and insulin reactivity in DQ8' DR4+/mII–/RIP-B7.1 mice. We have shown in our recent study that diabetic splenocytes from DQ8' mII–/RIP-B7.1 mice responded to islets and the putative autoantigens GAD and/or insulin (34). This reactivity was accompanied by the production of IFN-γ, but not IL-4 (34). To investigate whether the low incidence of spontaneous diabetes seen in DQ8' DR4+/mII–/RIP-B7.1 mice was due to the loss of this autoreactivity restricted to DQ8', we performed in vitro proliferation assays using purified CD4+ T cells from splenocytes of nondiabetic (Figure 5) and diabetic (data not shown) DQ8' DR4+/mII–/RIP-B7.1 mice. CD4+ T cells from DR4+/mII–/RIP-B7.1 mice showed moderate responses to GAD and insulin (1,400 and 1,260 cpm, respectively; over 470 cpm in the absence of the Ag’s), whereas CD4+ T cells from DQ8' mII–/RIP-B7.1 mice showed similar results (albeit a lower level as compared with diabetic individuals; data not shown) to our previous study (34). Introduction of DR4 into DQ8' mII–/RIP-B7.1 mice (namely DQ8+ DR4+/mII–/RIP-B7.1 mice) did not abolish the islet autoantigen reactivity (Figure 5). Moreover, the responses were restricted to both DQ8 and DR4, as shown by Ab-inhibition assay (Figure 5, a and b).
blocking experiments were repeated on two further occasions with similar results (data not shown). However, the profile of cytokines produced by islet autoantigens in the mice expressing DQ8 and DR4 was different from that seen in DQ8+/mII–/RIP.B7-1 mice, namely the suppression of IFN-γ and the production of IL4, albeit at low levels (Figure 6a). We also studied another cohort of control mII– mice that express DQ8 or DR4 or both (DQ8DR4), but without the expression of RIP-B7 transgene. No CD4+ T cells from any of the groups showed notable proliferative responses to either GAD or insulin (data not shown), and cytokine production (IFN-γ and IL-4) was only slightly above detection levels (Figure 6a). However, anti-CD3 stimulation drove the production of both IFN-γ and IL-4 by those CD4+ T cells (Figure 6b), and the profile of either IFN-γ or IL-4 was similar to that seen in mice expressing RIP-B7 transgene (Figure 6b). All the experiments shown above were repeated using total T cells (CD4+ and CD8+), and very similar results were obtained (data not shown). This suggests that the effect of DR4 on a Th2-like immune response is confined to CD4+ T cells in these mice.

To identify the T-cell epitopes giving the spontaneous GAD response in DQ8’DR4+/mII–/RIP-B7.1 mice, we tested six GAD peptides (GAD 61-80, 201-220, 281-300, 247-266, 509-528, and 524-543). However, none of these peptides generated notable T-cell reactivity either in assays of proliferation or cytokine production. The antigenic peptide(s) recognized by CD4+ T cells remain to be determined. There were no spontaneous responses to IA-2, another well-characterized islet β-cell autoantigen.

Expression of DR4 may cause CD4 T cells to differentiate toward a Th2 phenotype. To confirm the cytokine production in response to autoantigens, we stimulated the CD4+ T cells from DQ8+/mII–/RIP-B7.1, DR4+/mII–/RIP-B7.1, or DQ8+DR4+/mII–/RIP-B7.1 mice with anti-CD3, which induces maximum stimulation of T cells. Supernatants were harvested after 48 hours and assayed for cytokine production (IFN-γ, IL-4, IL-6). As shown in Figure 6b, CD4+ T cells from DQ8+/mII–/RIP-B7.1 mice secreted a high level of IFN-γ and a relatively low level of IL-4, whereas the expression of DR4 (either alone or together with DQ8) reversed this pattern of the cytokine production. The production of IL-6 could be detected, albeit at low level, in the supernatants of stimulated CD4+ T cells from mice expressing DR4 (either alone or together with DQ8), whereas it was undetectable in the supernatant of stimulated CD4+ T cells from DQ8+/mII–/RIP-B7.1 mice (data not shown).

Chemokines are chemotactic cytokines, and the interaction of chemokines and their receptors are important for both primary and secondary (memory) immune responses. It has been shown that certain chemokine receptors, such as CCR5, CXCR3, and CCR1, are preferentially expressed in Th1 cells (42–45). To investigate whether the downregulation of endogenous Th1 immune response by the expression of DR4 also affects chemokine-receptor expression, we examined the mRNA levels of CCR5 and CCR2 on anti–CD3-activated CD4+ T cells from DQ8’DR4+/mII–/RIP-B7.1 and DQ8’/mII–/RIP-B7.1 mice. Although this is not a real-

**Figure 6**

Cytokine production by CD4+ T cells after GAD (a) and anti-CD3 (b) stimulation. Purified CD4+ cells (2 × 10^6/ml) from spleens of nondiabetic mice, as indicated, were cultured with GAD protein or anti-CD3 (2C11 supernatant) in the presence of irradiated splenocytes for 72 and 48 hours, respectively. The culture supernatants were then tested for the production of IFN-γ and IL-4 using ELISA.
The mRNA expression of chemokines CCR5 and CCR2. Expression of chemokines CCR5 and CCR2 mRNA was assessed using RT-PCR (see Methods), as shown in the ethidium-bromide stained gel. CD3ε was used as the internal control for the normalization of RNA levels of tested samples, as indicated. Lane 1: 100-bp ladder; lanes 2–5: CCR5 or CCR2, as shown. RT-PCR was performed on oligo-dT–synthesized cDNA, which was derived from total RNA of anti-CD3–stimulated T cells from spleens of non-diabetic mice as indicated.

The mRNA expression of chemokines CCR5 and CCR2. Expression of chemokines CCR5 and CCR2 was downregulated in CD4+ T cells from DNQ8+/mII–/RIP-B7.1, DR4+/mII–/RIP-B7.1, and DQ8+DR4+/mII–/RIP-B7.1 mice. We are aware that genetic background can influence immune responses, and to examine whether the mice used in the study were on the same genetic background (C57BL/6) we performed microsatellite analysis using 17 genetic markers that span mouse chromosomes 1 to 7, 9, 11, 13, and 14 (see Methods). The mapping results are identical at the loci for the C57BL/6 strain (Massachusetts Institute of Technology database) and do not support the possibility of some genes carried over with the expression of DR4, contributing to the disease modulation described above. However, it is clear that this does not cover the entire mouse genome.

Discussion

Studies examining MHC types associated with T1DM have identified individual loci associated with the disease, although haplotypes (an association of several genes together) are likely to be much more important in defining diabetes susceptibility or resistance. DR4 and/or DR3 have been shown to be strongly associated with T1DM in various types of studies (populations, families, and twins) (49–54). Among DR4 subtypes, DRB1*0401 has been reported to be most strongly associated with the disease (55). However, the strong linkage disequilibrium within the MHC involving DQ8 (DQA1*0301/DQB1*0302) and DR4 (DRB1*0401) makes it difficult to ascribe inde-

![Figure 7](https://example.com/figure7.png)

**Figure 7**
The mRNA expression of chemokines CCR5 and CCR2. Expression of chemokines CCR5 and CCR2 mRNA was assessed using RT-PCR (see Methods), as shown in the ethidium-bromide stained gel. CD3ε was used as the internal control for the normalization of RNA levels of tested samples, as indicated. Lane 1: 100-bp ladder; lanes 2–5: CCR5 or CCR2, as shown. RT-PCR was performed on oligo-dT–synthesized cDNA, which was derived from total RNA of anti-CD3–stimulated T cells from spleens of non-diabetic mice as indicated.

![Figure 8](https://example.com/figure8.png)

**Figure 8**
Expression of Ig isotypes in DQ8+/mII–/RIP-B7, DR4+/mII–/RIP-B7, and DQ8+DR4+/mII–/RIP-B7 mice. Serum levels of IgG1 and IgA were measured using ELISA (see Methods) in the three groups of mice as indicated. The number of mice in each group was 15, 20, and 22 for DQ8+/mII–/RIP-B7, DR4+/mII–/RIP-B7, and DQ8+DR4+/mII–/RIP-B7 mice, respectively. The results are presented as mean ± SD. Student’s t test was used for the statistical analysis.
pendent associations with diabetes. It is noteworthy that T1DM patients more commonly express DR4 in combination with DR3 haplotypes than DR4 homozygotes. The findings of this study may suggest a reason why this is so in that DR4 may have an inhibitory effect on the susceptibility given by DQ8 and that in a homozygous state, the relative inhibitory effect of DR4 may override DQ8. In this study we have dissected the role of DR4 (DRBI*0401) independently or in association with DQ8 on the development of spontaneous diabetes by using HLA-transgenic mice. To our surprise, DR4, to a large extent, downregulated the strong diabetogenic effect of DQ8 when they were coexpressed, and the diabetogenic effect of DR4 alone was much milder than that of DQ8 in our spontaneous diabetes mouse model and only slightly greater than in mice not expressing MHC class II (25% vs. 10%). The data suggest that the inhibitory effect of DR4 may be mediated by selection of islet β-cell–specific Th2 regulatory CD4 T cells or by preferential presentation of islet β-cell–associated self-proteins to the immune system in a manner that activates a protective Th2 type of CD4+ T cell response.

It has been reported by several groups that expression of an I-E transgene in NOD mice protects them from diabetes development (56–58). Although the mechanism of how the protection occurs has not been fully elucidated, deletion of self-reactive T cells (clonal deletion) due to the expression of I-E molecules did not appear to be the cause. Hanson et al. (59) had shown that expression of Eβ4 transgene in NOD mice blocked diabetes development by altering the balance of cytokines produced by immunization with islet β-cell–associated autoantigen, directing autoreactive T cells toward a Th2-type immune response. Our data presented here are consistent with and extended those findings. In our model, expression of DR4 molecules altered the cytokine profile of spontaneous GAD and insulin autoreactivity. The change of serum Ig isotypes by the expression of DR4 provides further support for the Th2-like phenotype seen in these mice. It is conceivable that the expression of DR4 may have altered general immune responses (not just limited to islet β-cell–associated autoreactive T cells) to Th2 type of responses because it is unlikely that the increased levels of IgG1 and IgA seen in DR4+ or DQ8×DR4 mice are all islet β-cell–associated autoantibodies. In fact, the results from the anti-CD3 cross-linking experiments suggested a general Th2 immune response confined to CD4+ T cells in these mice.

It is interesting that fewer CD8+ T cells were found in diabetic DR4+/mII /RIP-B7.1 islet infiltrates than were seen in diabetic DQ8+/mII /RIP-B7.1 islet infiltrates, although the number of peripheral CD8+ T cells was very similar between these two types of mice. It is possible that the reduction of IFN-γ-producing Th1 immune responses due to the expression of DR4 may also diminish the recruitment and/or homing of diabetogenic T cells, including CD8+ T cells.

CD4+ T cells play an important role in regulating the functions of the immune system, and numerous studies in both human and animals have shown the contribution of Th1 or Th2 to the pathogenesis of a wide variety of diseases (60–68). The dichotomy of Th1 and Th2 cells in the development of T1DM has also been well documented (8-15, 69). Most of the studies demonstrate that Th1 cells contribute to the immunopathogenesis of islet β-cell destruction, whereas Th2 cells mediate disease protection. Based on these studies, new strategies in oral or nasal immunotherapy (which promote Th2 immune responses) for diabetes prevention have been tested in experimental animals (16–19), and some of the regimens are currently in clinical trials. More recently, Lyons et al. (70) demonstrated the influence of the idd 9 genetic interval on the pathogenesis of insulitis and diabetes in NOD mice. Using genomic-mapping analysis, the authors showed that the accumulation of IL-4–producing cells is associated with disease protection in the idd 9 loci. On the other hand, other studies have also shown that Th2 cells have the potential to be diabetogenic, as well (20–22). Thus, like most autoimmune disorders, the pathogenesis of T1DM cannot be explained simply by the dichotomy of Th1 and Th2, even though many studies, including our own, support this notion. Although control of autoimmune disease, such as diabetes, may be achieved by therapeutic interventions designed to enhance Th2 responses, the cure or prevention of the disease possibly will require the combination of different strategies.

To identify the T-cell epitopes giving the spontaneous GAD response in DQ8′DR4+/mII /RIP-B7.1 mice, we tested six GAD peptides (GAD 61-80, 201-220, 281-300, 247-266, 509-528, and 524-543). However, none of these peptides generated notable T-cell reactivity either in assays of proliferation or cytokine production. Using B10M/DR4-transgenic mice, the founder of our DR4+/mII/C57BL/6 mice, Wicker et al. have identified several strongly binding GAD peptides restricted to DR4 using binding assays and an immunization approach (35). Peptides 201-220 and 524-543 that we have tested were either the same as Wicker et al. reported (201-220) or in the vicinity of a strongly binding GAD peptide in Wicker et al.’s study (531-550). Peptide 509-528 used in this study is in the vicinity of two relatively weak binding peptides (501-520 and 511-530) in Wicker et al.’s study (35), but a similar peptide (511-525) showed potent immunogenicity in another study using a similar system (71). There are several possibilities that could explain the nonresponsiveness to those GAD peptides, especially the strong binding peptides. First, the development of tolerance to those strong binding self-peptides may contribute to the nonresponsiveness; second, it is possible that the lack of peptide response was due to the low frequency of T cells reactive to any given GAD peptide tested in our study; and last, it is likely that different GAD epitopes may be generated by Ag-presenting cells in an in vivo and immunized setting such as ours. Thus, DR4 molecules may present preferentially these GAD peptides leading to a Th2 type of
immune response. It is also possible that subdominant antigenic peptides or a determinant(s) generated during Ag spreading, which has been well documented in TIDM, may also induce protective, rather than pathogenic, immune responses (10,72,73).

The present study provides both in vivo and in vitro evidence for a possible inhibitory role of DR4 in the development of diabetes. Our data as a whole are most consistent with the view that expression of DR4 enhances more endogenous Th2 type of immune responses. The precise mechanism of how DR4 molecules promote Th2 regulatory immune responses is currently under investigation. However, we believe that the idea that expression of DR4 promotes more endogenous Th2 type of immune responses may not be the only mechanism for disease modulation seen in our study. Regardless of what mechanisms are involved, our “humanized” spontaneous diabetes model provides us not only a novel means to study the immunopathological effect of individual genes in strong linkage disequilibrium that is impossible to dissect in humans, but will also provide novel information regarding the role of DR4 in disease development. This information is important for the development of effective immunotherapy, especially as we are studying the effect of DQ8/DR4 HLA genes – those commonly found in diabetic patients.

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