Follicular helper T cell signature in type 1 diabetes


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The strong genetic association between particular HLA alleles and type 1 diabetes (T1D) indicates a key role for CD4+ T cells in disease; however, the differentiation state of the responsible T cells is unclear. T cell differentiation originally was considered a dichotomy between Th1 and Th2 cells, with Th1 cells deemed culpable for autoimmune islet destruction. Now, multiple additional T cell differentiation fates are recognized with distinct roles. Here, we used a transgenic mouse model of diabetes to probe the gene expression profile of islet-specific T cells by microarray and identified a clear follicular helper T (Tfh) cell differentiation signature. Introduction of T cells with a Tfh cell phenotype from diabetic animals efficiently transferred diabetes to recipient animals. Furthermore, memory T cells from patients with T1D expressed elevated levels of Tfh cell markers, including CXCR5, ICOS, PDCD1, BCL6, and IL21. Defects in the IL-2 pathway are associated with T1D, and IL-2 inhibits Tfh cell differentiation in mice. Consistent with these previous observations, we found that IL-2 inhibited human Tfh cell differentiation and identified a relationship between IL-2 sensitivity in T cells from patients with T1D and acquisition of a Tfh cell phenotype. Together, these findings identify a Tfh cell signature in autoimmune diabetes and suggest that this population could be used as a biomarker and potentially targeted for T1D interventions.

Introduction

Identifying the type of immune response that underlies autoimmune tissue destruction is a critical step in designing appropriate immunomodulatory strategies. In the case of type 1 diabetes (T1D), current dogma holds that Th1 cells cause pathology, and deviating the Th1 response has been a cornerstone of immunotherapeutic efforts. This view stemmed from the seminal observation that islet-specific T cells differentiated to a Th1 phenotype caused aggressive diabetes in neonatal nonobese diabetic mice, while those differentiated under Th2 conditions did not (1). While the latter study suggested Th1 cells to be the most likely drivers of T1D, data emerging over the following 2 decades were not always consistent with this conclusion. Although IFN-γ expression correlated with diabetes in NOD mice (2), other data suggested that diabetes in this model represented a Th2 rather than a Th1 phenomenon (3). Analysis of T cell differentiation in patients with T1D was similarly confusing: some work suggested increased IFNG mRNA (4) or stimulation-induced IFN-γ protein in individuals newly diagnosed with T1D (4, 5); however, others found IFN-γ production to be significantly lower in patients with recent-onset T1D than in healthy control subjects (6, 7). In one study, T cell reactivity to preproinsulin was shown to be Th2 dominant in autoantibody-positive subjects (8), again challenging the Th1 paradigm.

The identification of Th17 cells heralded a shift in our appreciation of autoimmune tissue damage and prompted the first move away from a strict dichotomy between Th1 and Th2 (9). Some data hinted at involvement of Th17 cells in T1D (10, 11), although other studies suggested that IL-17 was dispensable (12) or even protective (13, 14) in this setting. The incorporation of Th17 cells into the Th1/Th2 paradigm focused attention on additional cytokines, outside of those associated with Th1 or Th2 differentiation (IFN-γ and IL-4, respectively). One example, IL-21, was shown to be capable of promoting the Th17 response (15, 16). IL-21 is a member of the common γ chain signaling cytokine family and acts on a broad range of target cell populations, including B cells, CD8 T cells, NK cells, and dendritic cells. Interestingly, abrogation of IL-21 signaling was shown to be protective in mouse models of diabetes (17, 18), while transgenic expression of IL-21 in the pancreatic islets was sufficient to induce diabetes in nonautoimmune (C57BL/6) mice (18). The cellular source of IL-21 in the setting of diabetes is currently unclear, although Th17 cells and follicular helper T (Tfh) cells represent likely candidates. Here, we used an unbiased microarray approach to reassess T cell differentiation in a mouse model of spontaneous autoimmune diabetes. The data indicate that islet-specific T cells responding to pancreatic antigen show the characteristic features of Tfh cell differentiation. Furthermore, analysis of memory CD4 T cells from patients with T1D reveals a striking upregulation of Tfh-associated genes, including CXCR5 and IL21, which are tightly correlated. Collectively, these data implicate IL-21–producing Tfh cells in the immune response associated with T1D.

Results

Islet-specific T cells in the pancreatic LN show a Tfh gene signature.

To dissect the gene expression changes associated with the T cell response to tissue-specific self antigen, we performed microarray analysis on DO11 T cells responding to pancreas-expressed ovalbumin in the DO11 RIP-mOVA mouse model of diabetes (19).
Gene expression profiles of CD4+ T cells sorted from pancreatic LNs (PanLNs) of DO11 RIP-mOVA mice were compared with those of cells sorted from non-antigen-draining (inguinal) LNs. Conventional (CD4+CD25+) and regulatory (CD4+CD25+) populations were analyzed separately, and cells from the PanLNs were also gated on CD69 expression, since it has been shown that T cells responding to pancreas-derived self antigen upregulate this activation marker (20). Strikingly, 4 of the 20 most significantly upregulated genes in conventional T cells responding to tissue-derived self antigen were archetypal Tfh cell genes (Table 1 and ref. 21). When we hand-curated genes previously associated with Tfh cell differentiation (22), it was clear that these were strongly upregulated in T cells responding to islet antigen. This included Bcl6, the key transcription factor associated with Tfh cell differentiation (refs. 23–25 and Figure 1A). The application of canonical correspondence analysis (CCA) to microarray data has recently been reported (26). This method permits the interrogation of transcriptomic data, in the context of a specific biological problem, using another transcriptome as explanatory data. We adopted this approach to analyze our microarray data in conjunction with a published Tfh data set (25) (see Methods). CCA successfully identified the features of T cells in the PanLNs (magenta circles, Figure 1B), which showed a strong correlation with Tfh cells (blue arrow, Figure 1B). Axis 1 described the major variance (76% of the canonical analysis space; precisely, inertia) and represented the difference in 14 features of T cells in the PanLNs (magenta circles, Figure 1B), which showed a strong correlation with Tfh cells (blue arrow, Figure 1B). Axis 1 described the major variance (76% of the canonical analysis space; precisely, inertia) and represented the difference in 14 features of T cells in the PanLNs (magenta circles, Figure 1B), which showed a strong correlation with Tfh cells (blue arrow, Figure 1B).

**Table 1. The top 20 most significantly upregulated genes in PanLN T cells compared with inguinal LN T cells**

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<th>Rank</th>
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<tr>
<td>1</td>
<td>Cxcr5</td>
<td>11</td>
<td>Bcl6</td>
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<td>2</td>
<td>PtgF2</td>
<td>12</td>
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<td>IIE6</td>
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<td>Godd45b</td>
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<td>Cxcl10</td>
<td>16</td>
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<td>10</td>
<td>Cfr1</td>
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*Four signature Tfh cell genes. The Pdcd1 gene encodes PD-1.*

Gene expression profiles of CD4+ T cells sorted from pancreatic LNs (PanLNs) of DO11 RIP-mOVA mice were compared with those of cells sorted from non-antigen-draining (inguinal) LNs. Conventional (CD4+CD25+) and regulatory (CD4+CD25+) populations were analyzed separately, and cells from the PanLNs were also gated on CD69 expression, since it has been shown that T cells responding to pancreas-derived self antigen upregulate this activation marker (20). Strikingly, 4 of the 20 most significantly upregulated genes in conventional T cells responding to tissue-derived self antigen were archetypal Tfh cell genes (Table 1 and ref. 21). When we hand-curated genes previously associated with Tfh cell differentiation (22), it was clear that these were strongly upregulated in T cells responding to islet antigen. This included Bcl6, the key transcription factor associated with Tfh cell differentiation (refs. 23–25 and Figure 1A). The application of canonical correspondence analysis (CCA) to microarray data has recently been reported (26). This method permits the interrogation of transcriptomic data, in the context of a specific biological problem, using another transcriptome as explanatory data. We adopted this approach to analyze our microarray data in conjunction with a published Tfh data set (25) (see Methods). CCA successfully identified the features of T cells in the PanLNs (magenta circles, Figure 1B), which showed a strong correlation with Tfh cells (blue arrow, Figure 1B). Axis 1 described the major variance (76% of the canonical analysis space; precisely, inertia) and represented the difference in 14 features of T cells in the PanLNs (magenta circles, Figure 1B), which showed a strong correlation with Tfh cells (blue arrow, Figure 1B).
mRNA for \textit{BCL6}, the master transcription factor for Tfh cell differentiation, was also upregulated in the TID samples relative to the control samples; however, levels of other transcription factors associated with T cell differentiation (including \textit{Tbet} and \textit{GATA3}) were not altered between the 2 cohorts. Cytokine mRNAs were generally present at low abundance in T cells examined immediately \textit{ex vivo}. However, we observed elevated \textit{TNFA} mRNA in the patients with TID, along with trends toward higher \textit{IL21} and \textit{IFNG}, although these did not reach significance (Figure 5).

**Increased T cell CXCR5 expression in TID.** To extend the above mRNA analysis, peripheral blood T cells from patients with TID and controls were also assessed by flow cytometry. We first ascertained the distribution of CXCR5+ cells within the CD3+CD4+ compartment and found that they resided predominantly within the central memory (CD4+CD45RA–CD62L+) fraction in both patients with TID and controls (Figure 6A and B). However, patients with TID had a significantly higher percentage of CXCR5+ cells in both the central memory and effector memory compartments when compared with healthy controls (Figure 6C). After normalization for differences in the size of the memory compartment between individuals, there was no association between the percentage of CXCR5+ cells and patient characteristics (Supplemental Figure 4, A and B), suggesting that elevations in CXCR5+ cells may be present at all disease stages. It has been shown that peripheral blood...
CXCR5+ cells can be subdivided on the basis of CXCR3 and CCR6 expression and that CXCR3 CCR6+ and CXCR3 CCR6- cells are selectively expanded in juvenile dermatomyositis (28). In our study, the proportion of CXCR5+ cells expressing CXCR3 and/or CCR6 did not differ between patients with T1D and controls (Supplemental Figure 4C). When we analyzed cells coexpressing CXCR5 and ICOS, these were significantly elevated in patients with T1D compared with control individuals (Figure 6D). We did not observe a relationship between autoantibody status and percentage of Tfh cells, although it should be noted that a separate study focusing on patients with new-onset T1D showed a small increase in the percentage of Tfh cells in individuals with IA-2 and ZnT8 autoantibodies but not with GAD autoantibodies (29).

Increased IL-21 production in T1D. No differences in the levels of cytokine expression between patients and controls could be discerned by intracellular staining of isolated cells directly ex vivo (data not shown). However, following a brief period of in vitro activation, T cells from patients with T1D showed a significantly increased propensity to express IL-21 (Figure 7A). Since the IL-21+ cells in the mouse model of diabetes coexpressed IFN-γ and TNF-α, we also tested for polyfunctionality in the T cells from patients with T1D. Significantly increased numbers of IL-21+TNF-α+ and IL-21+TNF-α+IFN-γ+ cells were noted in the T1D samples compared with control samples (Figure 7B). Single TNF-α producers, but not single IFN-γ producers, were also increased in the patients compared with controls (Figure 7B). The IL-21–producing population in patients with T1D largely comprised cells coproducing TNF-α or coproducing both TNF-α and IFN-γ (Figure 7C).

Mouse studies have suggested that IL-21 derives predominantly from cells bearing a Tfh cell phenotype, although the relationship is a complex one, with indications that only a fixed proportion of CXCR5+ cells can produce IL-21 at any one time (30). Strikingly, correlation of ex vivo CXCR5 staining and stimulated...
IL-21 production showed that individuals with higher proportions of CXCR5+ cells also produced the highest level of IL-21 upon activation (Figure 7D). Furthermore, sorting CXCR5+ cells prior to stimulation revealed that IL-21 derived predominantly from CXCR5+ memory cells, while, in contrast, IFN-γ, IL-4, and IL-17 derived mainly from CXCR5− memory cells (Figure 7E).

**IL-2 signaling antagonizes CXCR5 upregulation.** The above data indicate that Tfh cell signature genes were upregulated at the mRNA level in T cells from patients with TID and elevations in CXCR5, ICOS, and IL-21 could be confirmed at the protein level. Experiments using murine T cells have shown that IL-2 can inhibit Tfh cell differentiation. Accordingly, signals through the IL-2R can dictate the balance between Tfh and T-effector differentiation (31, 32) via STAT5-dependent skewing of the BCL6/BLIMP1 ratio (33), and endogenous provision of IL-2 directly inhibits Tfh cell differentiation in response to influenza virus (34).

Intriguingly, substantial evidence links genetic alterations in the IL-2 pathway with TID (35–38), and the functional response of T cells to IL-2 has been reported to be defective in TID (39). This may reflect higher expression of PTPN2 (39), a phosphatase that inhibits JAK/STAT signaling and which is genetically associated with TID, or polymorphisms at the IL2RA locus (40), although it is likely that additional mechanistic explanations remain undiscovered. It is therefore possible that suboptimal IL-2 signaling might favor Tfh cell differentiation in the setting of TID. To explore this idea, we used T cells from patients with TID to examine the relationship between IL-2 sensitivity and the propensity to upregulate CXCR5 (Figure 8C and Supplemental Figure 6). Given the multiple defects in the IL-2 pathway documented in TID, this provides...
a putative mechanism for augmented Tfh cell differentiation in this disease setting.

Discussion

The perception of T cell differentiation as a Th1/Th2 dichotomy, with Th1 cells responsible for tissue-specific autoimmunity, has led numerous researchers to hunt for a Th1 signature in T1D, with widely variable results. Data from our mouse model of diabetes prompted us to seek evidence of a Tfh gene signature instead. This resulted in the identification of elevated expression of archepromoted us to seek evidence of a Tfh gene signature instead. This resulted in the identification of elevated expression of arche}

Figure 4. Enrichment for Tfh cells leads to preferential transfer of disease. (A) RIP-mOVA Cd28−/− mice were adoptively transferred with CXCR5-depleted or CXCR5-enriched DO11 T cells sorted from the PanLNs of DO11 RIP-mOVA mice. Blood glucose readings 4 weeks after transfer. **P < 0.01. Central horizontal bars depict the mean and are spanned by bars showing the SEM. (B) Representative pancreas sections stained for T cells (blue) and insulin (brown) are shown (n = 5). Original magnification, ×20.
Tfh cell numbers also have been reported to be subject to control by Qa-1–restricted CD8 T cells (67), and CD8 cells specific for the human equivalent of Qa-1 (HLA-E) have been reported to be functionally defective in T1D (68). More recently, the identification of Tregs expressing CXCR5 (69–71) has reinforced the notion that specialized regulatory mechanisms operate to curb Tfh cell differentiation, highlighting an additional checkpoint that might potentially be dysregulated in T1D.

Frequently, the traits that confer increased susceptibility to autoimmune disease afford an advantage in infectious settings. In this regard, Tfh cell numbers have recently been shown to correlate with protective antiviral immunity (72–74), while loss-of-function mutations in the IL-21 receptor result in immunodeficiency (75). The data presented herein add to a growing body of data linking Tfh cell differentiation with autoimmunity (71, 76, 77). Thus, polymorphisms that augment Tfh development might enhance protective immunity yet confer an increased risk of autoimmunity.

**Methods**

**Mice.** DO11.10 TCR transgenic mice were obtained from The Jackson Laboratory. BALB/c RIP-mOVA mice (expressing the ovalbumin transgene under control of the rat insulin promoter, from line 296-1B) were a gift from W. Heath (The Walter and Eliza Hall Institute, Parkville, Melbourne, Australia). DO11.10 mice were crossed with RIP-mOVA mice to generate DO11 RIP-mOVA mice. Mice were housed in individually vented cages at the University of Birmingham Biomedical Services Unit or within the Comparative Biology Unit at the University College London.

**Patients with T1D.** Peripheral venous blood was obtained from patients attending the clinical T1D service at the University Hospital Birmingham National Health Service Foundation Trust, United Kingdom. A total of 102 patients (mean age, 37 years; mean duration of T1D, 19 years; 60 men and 42 women) were recruited to this study, although not all assays were performed on every patient (see Supplemental Figure 7). Patients were selected on the basis of a clin-
shown that T cells responding to pancreas-derived self antigen upregulate CD69 (20). Antigen-specific T cells were gated on CD69+ in the PanLNs (antigen-specific T cells in the inguinal LNs were CD69−). To obtain sufficient cells of each type, cells from 14 mice were pooled to generate each sample. Mice were randomly assigned into groups. Three to six replicates were collected per experimental group (each replicate deriving from 14 mice). Sort purities were 97.3%–99.7%. Sorted cells were snap frozen immediately in liquid nitrogen, and RNA was subsequently extracted using the RNeasy Micro Kit (Qiagen). Microarray analysis was performed using the GeneChip Mouse Genome 430A 2.0 Array, and data were acquired using the Affymetrix Model 450 Fluidics Station and Affymetrix Model 3000 scanner. Affymetrix GeneChip data were normalized using the robust multichip averaging (RMA) algorithm of the Bioconductor package, affy. Quality control was done by the Bioconductor package, affyQCReport. CCA was performed as previously described (26). Briefly, logged expression values of the diabetes T cell data set were analyzed by CCAM using fold change of Tfh and naive Tfh data set (Gene Expression Omnibus [GEO] accession no. GSE40068) (25) as an explanatory variable. For the Tfh-ness anal-

Figure 6. Increased frequency of CXCR5+ T cells in patients with T1D. (A) Representative staining of PBMCs with CXCR5 and canonical T cell memory markers CD45RA, CD62L, and CCR7. Plots are gated on CD3+CD4+ and central memory (C. memory) and effector memory (E. memory) subsets as shown. The percentage of events within each quadrant (top left) is shown on the graph, and the frequency of CXCR5+ events within the naive, central memory, and effector memory fraction is shown as boxed on each subsequent plot. (B) The proportions of CD4+CXCR5+ T cells that fall into naive (CD4+CD45RA+CD62L−), central memory (CD4+CD45RA−CD62L−), and effector memory (CD4+CD45RA−CD62L+) subsets in patients with T1D (n = 24) and healthy controls (n = 15). Central horizontal bars depict the mean and are spanned by bars showing the SEM. (C) Frequencies of CXCR5+ cells within naive, central memory, and effector memory CD4+ T cell subsets in patients with T1D (n = 24) and healthy controls (n = 15). (D) A subset of the above individuals was examined for coexpression of CXCR5 and ICOS. Graph shows the percentage of CXCR5+ICOS+ cells within CD3+CD4+ T cells of patients with T1D (n = 11) and healthy controls (n = 9). Box and whisker plots show the median, interquartile range, and 10th to 90th percentile.

Microarray. Antigen-specific (KJ-126+) CD4+CD25− cells (conventional T cells) or CD4+CD25+ cells (Tregs) were sorted from pooled PanLNs or inguinal LNs from 6-week-old DO11 RIP-mOVA mice by FACS (MoFlo, Dakocytomation). Antibodies used for the sort stain were CD4 (LT34; Ebioscience), KJ-126 (Caltag Laboratories), CD25 (PC61; BD Biosciences), and CD69 (HL2F3; BD Biosciences). Cells were sorted on CD69 as a marker of activation, since it has been shown that T cells responding to pancreas-derived self antigen upregulate CD69 (20). Antigen-specific T cells were gated on CD69+ in the PanLNs (antigen-specific T cells in the inguinal LNs were CD69−). To obtain sufficient cells of each type, cells from 14 mice were pooled to generate each sample. Mice were randomly assigned into groups. Three to six replicates were collected per experimental group (each replicate deriving from 14 mice). Sort purities were 97.3%–99.7%. Sorted cells were snap frozen immediately in liquid nitrogen, and RNA was subsequently extracted using the RNeasy Micro Kit (Qiagen). Microarray analysis was performed using the GeneChip Mouse Genome 430A 2.0 Array, and data were acquired using the Affymetrix Model 450 Fluidics Station and Affymetrix Model 3000 scanner. Affymetrix GeneChip data were normalized using the robust multichip averaging (RMA) algorithm of the Bioconductor package, affy. Quality control was done by the Bioconductor package, affyQCReport. CCA was performed as previously described (26). Briefly, logged expression values of the diabetes T cell data set were analyzed by CCAM using fold change of Tfh and naive Tfh data set (Gene Expression Omnibus [GEO] accession no. GSE40068) (25) as an explanatory variable. For the Tfh-ness anal-
Flow cytometry and histology. Mouse cells were stained with mAbs against FOXP3 (FJK-16s), CD4 (LT34), CD3 (17A2), CXCR5 (2G8; BD Biosciences), PD-1 (J43), IL-21 (mhalx21), IL-17 (TC11-18H10; BD Biosciences), TNF-\(\alpha\) (MP6-XT22), and IFN-\(\gamma\) (XMG1.2). All antibodies were purchased from eBioscience unless otherwise indicated. For quantitation of Tfh cells, Tregs were gated out using CD25 stain-
were treated with 100 U IL-2 for 10 minutes and then fixed and stained for phosphorylated STAT5 (pSTAT5). The percentage of pSTAT5+ is plotted against the proportion of CXCR5+ cells induced by culture in the presence of IL-12 (as in enriched CD4+DO11+ T cells sorted by FACS (MoFlo, Dakocytoma-
injected with equivalent numbers of CXCR5-depleted or CXCR5-

Figure 8. Patients who respond poorly to IL-2 show an increased propensity to upregulate CXCR5. (A) Sorted CD4+CD45RA+ naive T cells from patients with T1D were cultured for 5 days with anti-CD3/anti-CD28 beads in the presence or absence of IL-12. Plots show representative CXCR5 staining. The percentage of gated CXCR5+ cells is shown on the graph. (B) Graphs show CXCR5 MFI of 5 independent experiments in which naive T cells were cultured for 5 days as above with IL-12 and or IL-2 (n = 15). Box and whisker plots show the median, interquartile range, and 10th to 90th percentile. *P ≤ 0.05, ***P ≤ 0.001, ****P ≤ 0.0001. U, untreated. (C) Relationship between CXCR5 induction and sensitivity of T cells to IL-2. CD4+ T cells from patients with T1D were treated with 100 U IL-2 for 10 minutes and then fixed and stained for phosphorylated STAT5 (pSTAT5). The percentage of pSTAT5+ is plotted against the proportion of CXCR5+ cells induced by culture in the presence of IL-12 (as in A and B) (n = 13).

ing. For intracellular cytokine staining, cells were stimulated for 5 hours with phorbol myristate acetate and ionomycin in the presence of brefeldin A (Sigma-Aldrich) for the final 4 hours. For cytokine and FOXP3 staining, cells were fixed and permeabilized according to the manufacturer’s instructions (eBioscience). Human cells were stained with CD4 (SK3), CD3 (UCHT1; Life Technologies), CXCR5 (R88B2; BD Pharmingen), CCR7 (G043H7; Biologend), CD45RA (HI100), CD62L (DREG-56), CCR6 (R6H1), CXCR3 (IC6/CXCR3; BD Biosciences), ICOS (ISA-3) biotin (streptavidin APC) and PE-Cy7, IL-21 (eBio3a3-N2), TNF-α (MAb11), IFN-γ (B27; BD Biosciences), IL-4 (BD4-8), IL-17 (CZB-23G1; Miltenyi Biotec), and pSTAT5 (47/STAT5/pY694; BD Biosciences). All antibodies were purchased from eBioscience unless otherwise indicated. For cytokine staining, cells were stimulated for 4 hours with phorbol myristate acetate and ionomycin in the presence of brefeldin A for the final 2 hours. Cells were then fixed and permeabilized according to the manufacturer’s instructions (eBioscience). For the pSTAT5 assay, stimulation was performed using 100 U recombinant human IL-2 (Peprotech) for 10 minutes. Fixation and staining were performed per the manufacturers instructions (BD Phospho Fix/Perm Buffer III). All data were acquired using a Dako CyAn (DakoCytomation) or Fortessa (BD) flow cytometer and analyzed using FlowJo software.

Adoptive transfer. Adoptive transfer experiments were carried out as previously reported (8). RIF-mOVA Cd28− mice were injected with equivalent numbers of CXCR5-depleted or CXCR5-enriched CD4+DO11 T cells sorted by FACS (MoFlo, Dakocytomation) from the pooled PanLNs of DO11 RIP-mOVA mice. Blood glucose was monitored by Glucometer Ascensia Elite XL (Bayer). Blood glucose readings above 250 mg/dl are typically considered diabetic. All injections were carried out in the absence of anesthesia and analgesia, and mice were returned immediately to home cages following the procedure. The welfare of experimental animals was monitored daily. No unexpected adverse events were noted during the course of these experiments.

Immunofluorescence staining and confocal microscopy. PanLNs and inguinal LNs from DO11 RIP-mOVA and DO11 mice were embedded in O.C.T. compound (Tissue-Tek 4583) and frozen in liquid nitrogen vapor. Five- to seven-μm tissue sections were prepared, acetone fixed, and stored at −20°C. Prior to staining, slides were rehydrated in PBS and subjected to biotin blocking and preincubation with 10% goat serum (Sigma-Aldrich). Sections were incubated with goat anti–IgM-555 (Jackson), Ki67-FITC (BD 556026) detected with rabbit anti-FITC (DakoCytomation VO403) and then donkey anti-rabbit 488 (Invitrogen A21206), and hamster anti-CD3 (BD 553058) detected with goat anti-hamster Cy5 (Stratech Scientific). Images were captured by confocal microscopy (Zeiss LSM 510 Meta).

Realtime PCR. PBMCs were isolated from the venous blood of patients with TID or healthy controls by density centrifugation. CD4+CD45RA− T cells were selected by magnetic negative selection (EasySep Human Memory CD4+ T cell Enrichment Kit; Stemcell) and stored in RNAlater (Qiagen). RNA extraction was performed using the RNeasy Mini Kit (Qiagen). RNA was reverse transcribed and interrogated using TaqMan probes for BCL6, CXCR5, ICOS, IL21, PDCDI1, FOXP3, GATA3, IL2, IL4, IL10, IL17A, TXB21, and IFNG. qPCR was carried out using the Stratagene Mx3000P. Gene expression was normalized to GAPDH using the 2−ΔΔct calculation.

In vitro T cell differentiation. PBMCs were isolated from venous blood by density centrifugation, and naive CD4 T cells were purified...
by negative magnetic separation (EasySep Human Naive CD4+ T cell Enrichment Kit, Stem Cell Technologies). Cells were plated at a density of 100,000 per well in a 96-well round bottom plate and stimulated with anti-CD3/28 beads (Human T-Activator Dynabeads, Life Technologies) for 5 days. Where indicated, cultures were treated with 10 ng/ml IL-12 and/or 20 ng/ml IL-2 (both from Peprotech).

Statistics. Data were analyzed using Prism statistical software. Statistical significance was assessed using the Mann-Whitney test. Paired data were analyzed using a 2-tailed paired Student’s t test. A P value of less than 0.05 was considered significant.

Study approval. Murine experiments were performed in accordance with the relevant Home Office project and personal licences following institutional ethical approval (from the University of Birmingham and University College London). Informed consent was obtained from all participants, and the study was approved by the National Research Ethics Committee.

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27. Wellcome Trust Case Control Consortium. Genome-wide association study of 14,000 cases