Tolerance to Proinsulin-1 Reduces Autoimmune Diabetes in NOD Mice

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T-cell responses to insulin and its precursor proinsulin are central to islet autoimmunity in humans and non-obese diabetic (NOD) mice that spontaneously develop autoimmune diabetes. Mice have two proinsulin genes proinsulin -1 and 2 that are differentially expressed, with predominant proinsulin-2 expression in the thymus and proinsulin-1 in islet beta-cells. In contrast to proinsulin-2, proinsulin-1 knockout NOD mice are protected from autoimmune diabetes. This indicates that proinsulin-1 epitopes in beta-cells maybe preferentially targeted by autoreactive T cells. To study the contribution of proinsulin-1 reactive T cells in autoimmune diabetes, we generated transgenic NOD mice with tetracycline-regulated expression of proinsulin-1 in antigen presenting cells (TIP-1 mice) with an aim to induce immune tolerance. TIP-1 mice displayed a significantly reduced incidence of spontaneous diabetes, which was associated with reduced severity of insulitis and insulin autoantibody development. Antigen experienced proinsulin specific T cells were significantly reduced in in TIP-1 mice indicating immune tolerance. Moreover, T cells from TIP-1 mice expressing proinsulin-1 transferred diabetes at a significantly reduced frequency. However, proinsulin-1 expression in APCs had minimal impact on the immune responses to the downstream antigen islet-specific glucose-6-phosphatase catalytic subunit-related protein (IGRP) and did not prevent diabetes in NOD 8.3 mice with a pre-existing repertoire of IGRP reactive T cells. Thus, boosting immune tolerance to proinsulin-1 partially prevents islet-autoimmunity. This study further extends the previously established role of proinsulin-1 epitopes in autoimmune diabetes in NOD mice.

Keywords: type 1 diabetes, proinsulin-1, CD4+ T cells, immune tolerance, NOD mice

INTRODUCTION

Recognition of proinsulin by the immune system is a major determinant in the pathogenesis of autoimmune diabetes in both humans and non-obese diabetic (NOD) mice (1, 2). A polymorphic variable number of tandem repeats (VNTR) located in the promoter region of the insulin locus controls the transcription level of the Ins gene and is strongly associated with susceptibility to type 1 diabetes (T1D) in humans (3–5). Mice do not have a VNTR upstream of the insulin locus; however, they have two Insulin genes, Ins1 and Ins2 encoding proteins that are highly homologous with 92% identity at the
amino acid level. Proinsulin 1 and 2 proteins have identical A chains but differ by two amino acids in the B chain, three amino acids in the connecting peptide (C-peptide) and six amino acids in the leader peptide (6). The two proinsulin isoforms are differentially expressed with proinsulin 1 (PIns1) predominantly expressed in the pancreatic beta-cells and proinsulin 2 (PIns2) being the predominant isoform detected in the thymus (7–9).

Immune responses to native insulin peptides, in particular the B chain amino acids 9-23 (Ins B:9-23), are essential for autoimmune diabetes in NOD mice (10, 11). The two proinsulin isoforms differ by a single amino acid in the B: 9-23 region (PIns1: B9 proline, PIns2: B9 serine) and strong cross-reactivity of T cells for the Ins B: 9-23 epitope in both proinsulin molecules has also been reported (12). Despite the high degree of homology in the B:9-23 epitope and cross-reactivity of T cells for the Ins B: 9-23 epitope, a divergent immune response was observed when NOD mice were immunized with either Ins1 B:9-23 or Ins2 B:9-23 peptides, with Ins2 peptide conferring protection from diabetes onset, whereas Ins1 peptide did not prevent disease (13, 14). Further differences in cellular and humoral immune responses to both proinsulin isoforms have been highlighted by individual gene knockouts. NOD mice lacking Ins2 gene develop accelerated diabetes, ascribed to loss of central tolerance to insulin peptides; however, development of insulin autoantibodies (IAA) in Ins2 -/- mice suggests that immune responses against PIns1 epitopes are intact (15). In contrast, genetic deletion of Ins1 or replacement of murine Ins1 with human insulin gene (INS) in NOD mice provides significant protection from diabetes (16, 17). Protection from diabetes in NOD mice lacking Ins1 is likely due to the absence of cognate antigen in the target tissue, indicating that PIns1 peptides may be primarily targeted by insulin reactive T cells. Immunogenic epitopes in the PIns1 molecule have been reported (18), and T cells recognizing PIns1 amino acids 47-64 in the C-peptide region induce diabetes in NOD.SCID recipients (19). Thus, epitopes in PIns1 molecule may contribute to islet autoimmunity.

In contrast to NOD mice, non-autoimmune strains lacking Ins2 globally (20), or in medullary thymic epithelial cells (mTECs) did not develop pathological islet destruction however, when C57Bl/6 mice lacking Ins2 in mTECs were crossed to Ins1 knockout mice, the progeny developed spontaneous autoimmune diabetes within 3 weeks after birth (21). These studies suggest that thymic expression of PIns1 may add to the effect of PIns2 in eliminating insulin-specific autoreactive T cells. Constitutive or temporal expression of PIns2 in APCs induces excessive tolerance to PIns2 as it provides lasting protection from autoimmune diabetes in NOD mice (22). These mice were also thought to be tolerant to PIns1 epitopes because of cross-reactivity of the T cells to the conserved Ins B: 9-23 epitope. However, the role of PIns1 specific immune responses in pathogenesis of islet autoimmunity in NOD mice remains unclear, given the differential immune response observed upon immunization with Ins1 B:9-23 or Ins2 B:9-23 peptide. To resolve this, we investigated the impact of induced PIns1 expression in APCs on the development of antigen-specific T cells as well as insulitis and diabetes in NOD mice.

MATERIALS AND METHODS

TetO-Ins1 Mice

To generate the TetO-Ins1 construct, a 411 bp cDNA fragment spanning the coding region of murine PIns1 was amplified by PCR using NOD pancreatic islet cDNA as a template and cloned into HindIII and EcoRV sites of the pTRE-tight plasmid (Clontech). A 1100 bp transgene cassette comprising of the TetO-minimal CMV promoter, followed by the PIns1 gene and a polyA signal was excised between Xho I sites and purified for injection into fertilized NOD/Lt ova using standard techniques. Founders and transgene positive offspring were screened by PCR, using primers spanning the PIns1 gene (5'-TAAAGATATCTCTTTATTAGAGACCTC-3') and the tetO-CMV promoter (5'-TCAGTGTAGAACGTATGTCG-3').

Other Mice

NOD/Lt mice were bred and housed at the bioresources center St. Vincent’s Hospital, Fitzroy. The NOD-IE μT-IE μTα mice that drive the expression of tetracycline transactivator (tTA) under the control of MHC class II IEμ promoter have been previously described (23) and were obtained from Prof. C. Benoist and Prof. D. Mathis (Dept of Pathology, Harvard Medical School, Boston, Massachusetts, USA). Generation of NOD8.3 mice, which express the TCRαβ rearrangements of the H-2Kd-restricted, β cell-reactive, CD8+ T cell clone NY8.3, was previously described in detail (24). TIP-1/8.3 mice were generated by crossing NOD-IE μT-IE μTα-TetO-Ins1double transgenic TIP-1 mice with TCR transgenic NOD8.3 mice. All mice were bred, maintained and used under specific pathogen free conditions at St Vincent’s Institute (Melbourne, Australia). All experimental procedures followed the guidelines approved by the institutional animal ethics committee.

Doxycycline Treatment

Untreated TIP-1 mice constitutively express proinsulin-1 in antigen presenting cells (APCs). To turn-off proinsulin-1 expression, doxycycline hyclate (Dox) (Sigma-Aldrich) was administered via drinking water at concentration of 2mg/ml. Water bottles were changed thrice weekly.

RT-PCR

For total RNA extraction, whole spleen and thymus were harvested in cold Phosphate Buffered Saline (PBS). Tissue homogenates were prepared in RNA lysis buffer RA1 (Macherey-Nagel) from a 15mg slice of tissue using a tissue homogenizer. RNA was isolated using standard techniques. Founders and transgene positive offspring were screened by PCR, using primers spanning the PIns1 gene (5'-TAAAGATATCTCTTTATTAGAGACCTC-3') and the tetO-CMV promoter (5'-TCAGTGTAGAACGTATGTCG-3').

Glyceraldehyde 3-phosphate dehydrogenase (Gapdh; Mm9999915_1) were purchased from Applied Biosystems. To
determine relative expression, Ct values of Insulin were subtracted from Ct values of reference genes for each sample and the difference (dCt) was plotted to determine the abundance of the gene of interest.

**Histology and Immunohistochemistry**

For immunohistochemistry, pancreata were snap-frozen in optimal cutting temperature compound (OCT Compound; Sakura Finetek, Torrance, CA) and stored at −80°C. For histological analysis 5-µm frozen sections of pancreas were prepared from three levels (200 µm apart), acetone fixed, stained with guinea pig anti-insulin followed by horse radish peroxidase–conjugated anti–guinea pig Ig (Dako Cytomation, Carpenteria, CA) and counterstained with hematoxylin. Insulitis was graded using the following scale: 0 = no infiltrate, 1 = peri-islet infiltrate, 2 = extensive (>50%) peri-islet infiltrate, 3 = intraislet infiltrate, and 4 = extensive intraislet infiltrate (>80%) or total β-cell loss. The percentage of islets with each grade per pancreas was calculated by addition of the grades for the three sections. Individual insulitis scoring for each mouse was performed as previously described (22).

**Incidence of Diabetes and Insulitis**

Diabetes onset was monitored by weekly measurement of urine glucose levels using Diastix (Bayer Diagnostics). Blood glucose levels were measured in mice with glycosuria using Advantage II Glucose strips (Roche). Animals displaying two consecutive blood glucose measurements of ≥15 mmol/L were considered diabetic. For adoptive transfer of diabetes, 2 × 10⁷ splenocytes from 13-17 week old pre-diabetic TIP-1 mice or control NOD mice were transferred (i.v.) into 9-12 week old NOD Rag−/− recipients and diabetes development was monitored as above.

**Flow Cytometry**

Antibodies used were anti-CD4 (RM4-5) conjugated to PerCP-Cy5.5, anti-CD3 (145-2C11) conjugated to FITC or anti-CD3 (500 A2) V500, anti-CD44 (1M7) conjugated to AlexaFlour488 (all BD Biosciences), anti-CD11c (N418), anti-CD3 (145-2C11) conjugated to FITC or anti-CD8 (RM4-5) conjugated to APC-Cy7 (BD Biosciences). Data were collected on an LSR Fortessa flow-cytometer (BD) and analyzed by FlowJo (Treestar) software.

**Tetramer and Magnetic Bead-Based Enrichment**

The tetramer and magnetic bead-based enrichment method was previously described (25). I-Ag7 tetramers were obtained from NIH tetramer core facility (Emory University, Georgia, USA), Kd-tetramers were obtained from ImmunoID (Parkville, Victoria, Australia). To enrich insulin-specific CD4+ T cells single cell suspensions from peripheral lymphoid organs (PLO), (pooled spleen and non-draining lymph nodes), were stained with phycoerythrin (PE)-conjugated I-Ag7-INSB₁₀₋₂₃ (HLVERLYVC GGEG) tetramer for 1 hour at room temperature. The Ins B₁₀₋₂₃ peptide in the I-Ag7-INSB₁₀₋₂₃ tetramer has been mutated (Glutamic acid to Glycine (E-G) at position 20 and Arginine to Glycine (R-G) at position 21) to improve its binding to the I-Ag7 molecule, which allows for better detection of insulin-specific CD4+ T cells (26). Insulin-specific CD8+ T cells were enriched from pooled PLO by staining the cell suspensions with APC-conjugated H-2Kd-INSB₁₅₋₂₃ (LYLVCGGGE) tetramer for 1 hour on ice. Hen Egg Lysozyme I-Ag7-HEL (AMKRHGLDNYRGYSL) tetramer or H-2Kd-TUM (KYQAVTTTL) were used as controls. Cells were then washed and stained with anti-PE or anti-APC microbeads (Miltenyi Biotec) followed by magnetic separation using an AutoMACSpro (Miltenyi Biotec) according to manufacturer’s instructions. IGRP₂₀₆₋₂₁₄ specific CD8+ T cells (H2-Kd, VYLTNNVFL) were stained and enriched as previously described (27). The separated fractions were stained and analyzed by flow cytometry. Gating strategy for tetramer enrichment was as follows: single cells were gated on forward and side scatter, and dead cells excluded using propidium iodide. From the live cell population, CD11c-CD11b-B220-F4/80-CD3+ CD8+ T cells were gated as the T cell population for analysis. Further selection of CD4+ T cells or CD8+ T cells was followed by analysis of the insulin or IGRP tetramer positive population respectively.

**Insulin Autoantibody (IAA) Assay**

A non-competitive IAA assay was performed in a 96 well ELISA format as previously described (28, 29). Briefly, an ELISA plate (Costar) was coated with or without human insulin (10 µg/ml, Actrapid, Novo Nordisk) overnight at 4°C. Wells were blocked with PBS containing 2% BSA for 2 hours and room-temperature and then probed with sera from 12-15 weeks old TIP mice, NOD or C57BL/6 mice (1:10 dilution) for 2 hours. Wells were washed 4 times and a biotinylated anti-mouse IgG1 (AbCam, 1:10000 dilution) antibody was added for 30 minutes. After washing, horse-radish-peroxidase conjugated streptavidin (BioLegend) was added for 15 minutes. The plate was washed five times, TMB substrate solution (BioLegend) was added and absorbance was measured at 450 nm using a Polarstar (BMG labtech) microplate reader. Each sample was run in duplicate and absorbance (450 nm) of test sample without plate bound insulin was subtracted from absorbance of test sample with plate bound insulin to calculate the actual absorbance value for each sample.

**Statistical Analysis**

Statistical analysis was performed using GraphPad Prism 8 Software (GraphPad, San Diego, CA, USA). Pooled data are shown as dot-plots with individual mice and the mean ± SEM. Data were tested for normal distribution using D’Agostino-Pearson omnibus normality test or Shapiro-Wilk’s test. Comparisons between two groups were performed using two-tailed unpaired student t-tests. Multiple comparisons were performed using One-way ANOVA with Sidak’s post-hoc test. Survival curves were compared using Log-Rank (Mantel-Cox) test. Statistical significance was defined as P < 0.05.
RESULTS

Conditional Expression of Proinsulin-1 in NOD Mice

To test whether inducing immune tolerance to proinsulin-1 (PIns1) influenced autoimmune diabetes we generated transgenic NOD mice to facilitate conditional expression of PIns1 in the antigen presenting cells (APCs). Reporter NOD mice expressing PIns1 under the control of the tetracycline-responsive CMV promoter (TetO-Ins1 mice) were bred with previously described driver NOD mice expressing TetR-VP16 tetracycline transactivator protein (tTA) under the control of IEα-MHC-II promoter referred to as TA-NOD mice (23). Bi-transgenic progeny referred to as TIP-1 (Tet Inducible PIns1) mice (Figure 1A) express PIns1 in the APCs, which can be turned-off upon doxycycline (Dox) treatment. Analysis of PIns1 expression in TIP-1 mice revealed that PIns1 transgene was expressed in the thymus and spleen as measured by RT-PCR (Figure 1B). After one week of Dox treatment, PIns1 expression dropped to baseline levels (Figure S1). Thus, PIns1 expression in TIP-1 mice was conditional, and tightly regulated.

TIP-1 Mice Have Reduced Insulitis and Insulin Autoantibody Expression

We recently reported that constitutive or temporal expression of PIns2 (PIns2) in the APCs limited to the perinatal period prevented insulitis and diabetes in NOD mice (22). To test whether PIns1 expression in the APCs influenced the progression of islet autoimmunity, we examined the immune infiltrate (insulitis) in the pancreata of TIP-1 mice expressing PIns1 continuously. At 12-14 weeks of age, insulitis was significantly reduced in TIP-1 mice expressing PIns1 compared to age matched NOD mice or TIP-1 mice fed dox to suppress PIns1 expression (Figures 2A–C). Analysis of pancreas histology from TIP-1 mice at 20-25 weeks of age revealed that approximately 50% of the islets examined were free of insulitis, whereas more than 80% of the islets examined from non-transgenic littermates were infiltrated (Figures 2D, E and Table S1), indicating that PIns1 expression in the APCs decreased but did not completely abolish development of insulitis, which progressed over time. Production of insulin autoantibodies (IAA) indicates spontaneous anti-insulin autoimmunity and IAA are frequently detected prior to diabetes onset in both humans and NOD mice (30, 31). We examined whether induced PIns1 expression in TIP-1 mice influenced B cell mediated humoral responses against insulin by measuring IAA in TIP-1 mice. IAA was significantly reduced in 12-15 weeks old TIP-1 mice as compared to age matched non-transgenic NOD mice (Figure 2F). Previously described PIns2 tolerant NOD-PI mice that are protected from diabetes and non-autoimmune prone C57BL/6 mice were used to set the baseline. Collectively, these results indicate that immune tolerance to PIns1 influenced progression of insulitis and reduced the development of IAA.

PIns1 Overexpression Partially Suppresses Spontaneous Diabetes in NOD Mice

Reduced insulitis and IAA suggest that diabetes development may be altered in TIP-1 mice. A cohort of female TIP-1 mice expressing PIns1 continuously and control NOD mice were observed for incidence of spontaneous diabetes. TIP-1 mice developed diabetes but at a significantly reduced incidence compared to non-transgenic control NOD mice. By 300 days of age 40% of TIP-1 mice and 65% of the control mice developed diabetes (Figure 3A). In addition, we investigated whether PIns1 expression in TIP-1 mice influenced the pathogenic potential of effector T cells. Splenocytes from 15-18 weeks old TIP-1 mice with ongoing expression of PIns1 and age matched control NOD mice were transferred into NOD.Rag1 -/- recipients. All recipient mice receiving control splenocytes developed diabetes between 50-70 days post-transfer, whereas only 2 out of 6 (33%) animals that received splenocytes from TIP-1 mice developed diabetes 70-90 days post-transfer (Figure 3B). Taken together these results suggest that overexpression of PIns1 in APCs is able to partially dampen immune responses against insulin and reduce diabetes incidence in NOD mice.
Proinsulin-Specific Tolerance in TIP-1 Mice

The partial protection from insulitis and diabetes in TIP-1 mice expressing PIns1 in the APCs could be due to immune tolerance to PIns1 epitopes. To demonstrate tolerance to PIns1, we enumerated the frequency of Insulin B9-23 reactive CD4+ T cells and Insulin B15-23 reactive CD8+ T cells in the peripheral lymphoid organs (PLO) (pooled spleen and non-draining lymph nodes) of 20-25 weeks old non-diabetic TIP-1 mice and age matched control mice using respective I-A (g7) and Kd tetramers. There was a significant reduction in the absolute number of insulin B9,23 specific CD4+ T cells binding to insulin B9,23/I-A (g7) tetramer (26) and the antigen-experienced CD44hi subset of insulin B9,23 specific CD4+ T cells in TIP-1 mice (Figures 4A–D). The absolute number of CD8+ T cells recognizing insulin B15,23 epitope (32) as well as the number of antigen-experienced CD44hi subset of insulin B15,23 specific CD8+ T cells were comparable in both TIP-1 mice and controls (Figures 4E–H). While the significant reduction of insulin-specific CD4+ and CD8+ T cells in TIP-1 is suggestive of deletional tolerance, it is possible that transgenic antigen expression in APCs may induce regulatory T cells (Tregs) that confer dominant tolerance and prevent diabetes in TIP-1 mice. We examined the expression of Foxp3 on insulin B9,23 specific CD4+ and CD8+ T cells in TIP-1 is suggestive of deletional tolerance, it is possible that transgenic antigen expression in APCs may induce regulatory T cells (Tregs) that confer dominant tolerance and prevent diabetes in TIP-1 mice. We examined the expression of Foxp3 on insulin B9,23 specific CD4+ and CD8+ T cells in TIP-1 mice (Figures S2A, B). In addition we examined the frequency of Foxp3+ CD4+ Tregs in the thymus and pancreatic lymph node (PLN). The proportion of Tregs was similar in both TIP-1 and control mice (Figures S2C, D). Taken together our data indicate that ectopic PIns1 expression induces deletion of cognate CD4+
FIGURE 4 | Immune tolerance to insulin specific T cells in TIP-1 mice. Insulin B:10-23-specific CD4+ T cells or Insulin B:15-23-specific CD8+ T cells were stained with respective tetramers and enriched from pooled peripheral lymphoid organs (PLO) of 20-25 weeks old TIP-1 mice and NOD mice using magnetic beads and enumerated by flow-cytometry. Representative FACS plots (A, C, E, G) and enumeration of insulin tetramer+ CD4+ T cells (B), insulin tetramer+ CD8+ T cells (F), CD44hi Insulin tetramer+ CD4+ T cells (D) and CD44hi Insulin tetramer+ CD8+ T cells (H) in TIP-1 and NOD mice. Values in the FACS plots indicate absolute number of tetramer binding cells. Each symbol in the scatter plots (Mean ± SEM) represents data from an individual mouse. **P < 0.01, *P < 0.05, ns= not significant. Data compared using 2-tailed unpaired t-test.
T cells, but does not induce antigen specific Tregs. The few remaining insulin reactive CD4+T cells could not be activated by the expressed antigen, whereas the low-affinity insulin B15-23 reactive CD8+ T cells (33) are not influenced by transgenic PIns1 expression.

**Downstream Responses to IGRP Are Delayed in TIP-1 Mice**

Previous work from our group has demonstrated that autoreactive responses to islet-specific glucose-6-phosphatase catalytic subunit-related protein (IGRP) are dependent upon immune response to PIns2 (1). To investigate if tolerance to PIns1 influenced the immune response to IGRP we examined the frequency of pathogenic IGRP 206-214 reactive CD8+ T cells in TIP-1 mice. The number of IGRP 206-214 specific CD8+ T cells was significantly reduced in 12-14 weeks old TIP-1 mice expressing PIns1 as compared to age matched controls. However, the frequency of IGRP 206-214 specific CD8+ T cells in TIP-1 mice expressing PIns1 did not differ from age-matched controls at 20-25 weeks of age (Figures 5A, B). This indicates that tolerance to PIns1 delays but does not prevent the spreading of immune responses to downstream antigen IGRP.

**Immune Response to Proinsulin-1 Is Not Required for Diabetes in NOD 8.3 Mice**

Autoreactivity to PIns2 is required for diabetes development in NOD 8.3 mice that have a pre-existing repertoire of IGRP specific T cells (34). Since we observed reduced frequency of IGRP reactive CD8+ T cells in 12-14 weeks old TIP-1 mice, we wished to know if immune responses to PIns1 were necessary for diabetes development in NOD 8.3 mice. TIP-1 mice were crossed with NOD 8.3 mice to generate offspring that were TIP-1/NOD8.3 double transgenic or NOD 8.3 transgenic alone. TIP-1/NOD8.3 mice developed diabetes with significantly delayed kinetics (median survival 97 days) compared to NOD8.3 mice (median survival 70 days) but all mice eventually developed disease (Figure 5C). The frequency of insulin specific T cells is very low even in NOD mice and with a skewed T cell repertoire in NOD 8.3 transgenic mice it is not possible to detect any insulin specific T cells. We were unable to detect insulin-specific T cells

![Figure 5](image-url)
in TIP-1/8.3 mice (data not shown). Therefore, tolerance to Plns1 significantly delays but does not prevent diabetes development in NOD 8.3 mice.

**DISCUSSION**

In this study we generated transgenic NOD mice to induce Plns1 expression in the APCs and examined the impact of antigen specific tolerance on autoimmune diabetes. The main findings of this study are 1) TIP-1 mice expressing Plns1 in the APCs show significantly reduced incidence of diabetes, which is associated with reduced insulitis and insulin autoantibody (IAA) expression. 2) Proinsulin specific CD4+ T cells are detectable in TIP-1 mice at a reduced frequency and are not activated. 3) Immune responses to downstream antigen IGRP are delayed but not absent in TIP-1 mice.

Given the high degree of homology between proinsulin 1 and 2 proteins, especially in the immunodominant insulin B chain epitope Ins B_9-23_ we expected to achieve robust protection from diabetes onset in TIP-1 mice, similar to previously described proinsulin-2 tolerant NOD mice (35); however, the partial protection from insulitis and diabetes observed in TIP-1 mice points to the existence of distinct pathogenic peptide epitopes in the Plns2 protein that can precipitate autoimmunity in NOD mice. A previous study characterizing immunogenic epitopes in NOD mice reported existence of multiple epitopes on both Plns1 and Plns2 molecules recognized by CD4+T cells (18). Importantly, epitopes outside of the highly homologous Ins B_9-23 peptide were identified in the leader and A chain sequences of Plns2 molecule. Thus, it is likely that Plns2 reactive T-cells recognizing these unique epitopes may induce islet destruction and subsequent diabetes onset in TIP-1 mice.

Our data complement the previous observations that reported detection of Plns1 reactive T cells (18, 19) in NOD mice. While the previous studies did not directly demonstrate the role of Plns1-reactive T cells in spontaneous disease, the significant reduction in diabetes incidence in TIP-1 mice suggests that Plns1 specific T cells participate in autoimmune destruction of beta cells. On the other hand, development of IAA and diabetes in TIP-1 mice may be related to ongoing immune responses to Plns2 epitopes.

A drawback of our study is that we have analyzed a single transgenic founder line expressing Plns1 in the APCs. Varying levels of transgenic insulin expression in the thymus may influence the diabetes progression in NOD mice. Plns2 levels were 7-fold higher in the spleen (~140 pmol/L) as compared to thymus (~20pmol/L) in the partially protected Plns2 tolerant mice previously described by Jaeckel et al. (12), whereas in the recently described TIP mice with robust protection from autoimmune diabetes upon conditional Plns2 expression in APCs, the level of thymic Plns2 expression (100pmol/L) was 5-fold more compared to peripheral tissues(20pmol/L) (22). TIP-1 mice may have relatively reduced transgenic expression of Plns1 in the thymic APCs as compared to transgenic Plns2 expression in the previously described TIP mice, thus imparting incomplete protection from autoimmune diabetes. Chentoufi and Polychronakos previously reported that Ins2 is expressed at a level more than 3-fold higher than Ins1 in the thymus of NOD mice (9). In TIP-1 mice analyzed here, induction of Plns1 results in approximately 5-fold higher expression as compared to non-transgenic NOD mice or uninduced TIP-1 mice. Moreover, protection from insulitis in TIP-1 mice is associated with the expression of Plns1 transgene, as TIP-1 mice fed doxycycline to suppress Plns1 expression develop islet infiltration comparable to non-transgenic controls indicating that ectopic Plns1 expression in APCs influences anti-islet immunity.

Does the reduction in the incidence of spontaneous diabetes in TIP-1 mice correlate with deletion of Plns1 specific T cells? Insulin B_{10-23} and Insulin B_{15-23} specific tetramers used in our study are likely to detect both Plns1 and 2 reactive CD4+ and CD8 +T cells, due to the invariant nature of the Insulin B_{9-23} peptide between the two isoforms. Immune responses to Insulin B_{9-23} epitope are required for both priming and effector phase of islet autoimmunity in NOD mice. Moreover, Insulin B_{9-23} primed CD4+ T cells are able to induce islet autoimmunity evidenced by IAA production (36). The significant reduction in absolute number of Insulin B_{9-23} tetramer binding CD4+T cells, and the antigen-experienced subset of tetramer binding CD4+ T cells, coupled with reduced IAA production in TIP-1 mice is suggestive of antigen-specific tolerance. While Tregs are an important tolerance mechanism, we did not find any evidence to suggest that the partial protection from diabetes in TIP-1 mice is due to antigen-specific Tregs. We are currently unable to conclude whether central or peripheral tolerance mechanisms regulate the insulin specific T cells in TIP-1 mice; however, future studies with ectopic antigen expression induced after the exit of antigen-specific T cells from the thymus may resolve this question.

Autoimmunity to insulin determines immune responses to other downstream antigens such as IGRP (1). IGRP_206-214 reactive CD8+ T cells were reduced in TIP-1 mice at 12-14 weeks; but ongoing tolerance to Plns1 did not prevent development of diabetes onset in TIP-1/8.3 mice. The precursor frequency of IGRP reactive CD8+ T cells is low in NOD mice (27), and the residual immune response to Plns2 in TIP-1 mice may be reduced as compared to control mice. The reduced CD4+ T cell help possibly accounts for the delayed expansion of IGRP specific T cells seen in TIP-1 mice. However, the residual immune response to Plns2 in TIP-1/8.3 mice with a pre-existing repertoire of IGRP specific T cells may be sufficient to help IGRP specific CD8+ T cells to mediate beta-cell destruction.

In summary, we find that immune tolerance to Plns1, whilst partly protective, is not sufficient to prevent spontaneous diabetes in NOD mice. Our data clarifies the role of Plns1 in the pathogenesis of autoimmune diabetes in NOD mice and extends the previously established role of Plns1 in autoimmune diabetes. The experimental model we have presented here, with its conditional gene-expression system, has the potential to delineate whether antigen-specific interventions can induce immune tolerance after islet autoimmunity is well established. Understanding this is important for development of strategies to induce antigen-specific tolerance clinically in people with stage 1 or 2 type 1 diabetes (37).
DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Material. Further inquiries can be directed to the corresponding author.

ETHICS STATEMENT

The animal study was reviewed and approved by Animal Ethics Committee, St Vincents Hospital, Melbourne.

AUTHOR CONTRIBUTIONS

GJ performed experiments, analyzed data, and wrote the manuscript. CS, JC, C-TK, and EP performed experiments and analyzed data. HT, BK, and TK designed the study, analyzed data, and wrote the manuscript. BK and TK supervised the study. All authors contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fimmu.2021.645817/full?supplementary-material
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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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