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## Activation of T cell checkpoint pathways during $\beta$ -cell antigen presentation by engineered dendritic cells promotes protection from type 1 diabetes

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### Abstract

Defective immune regulation has been recognized in type 1 diabetes (T1D). Immune regulatory T cell check-point receptors, which are generally upregulated on activated T cells, have been the molecules of attention as therapeutic targets for enhancing immune response in tumor therapy. Here, we show that pancreatic  $\beta$ -cell antigen (BcAg) presentation by engineered tolerogenic dendritic cells (tDCs) that express CTLA4 selective ligand (B7.1wa) or a combination of CTLA4, PD1 and BTLA selective ligands (B7.1wa, PD-L1, and HVEM-CRD1 respectively; multiligand-DCs) causes an increase in regulatory cytokine and T cell (Treg) responses and suppression of the effector T cell function as compared to engineered control-DCs. Non-obese diabetic (NOD) mice treated with BcAg-pulsed CTLA4-ligand-DCs and multiligand-DCs at pre-diabetic and early-hyperglycemic stages showed significantly lower degree of insulinitis, higher frequencies of insulin-positive islets, profound delay in, and reversal of, hyperglycemia for a significant duration. Immune cells from the tDC treated mice not only produced lower amounts of IFN $\gamma$  and higher amounts of IL10 and TGF $\beta$ 1 upon BcAg challenge, but also failed to induce hyperglycemia upon adoptive transfer. While both CTLA4-ligand-DCs and multiligand-DCs were effective in inducing tolerance, multiligand-DC treatment produced an overall higher suppressive effect on effector T cell function and disease outcome. These studies show that enhanced engagement of

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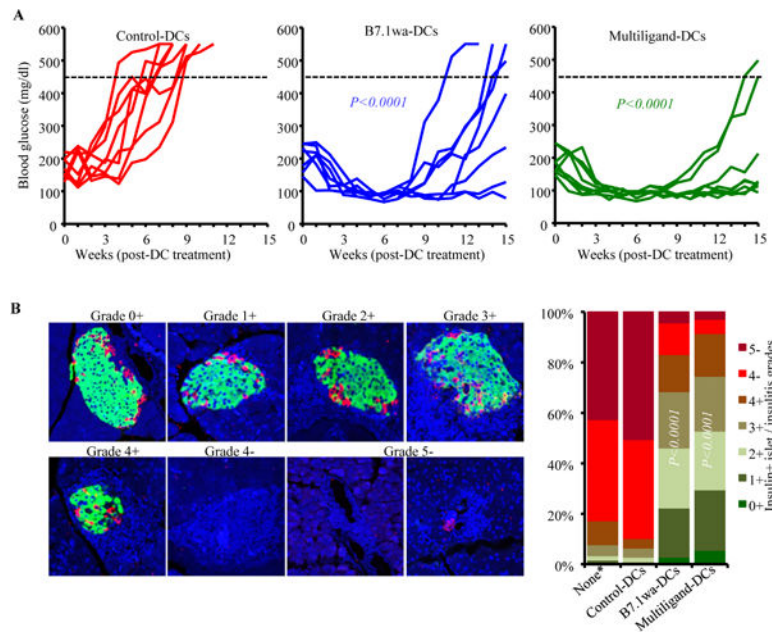
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T cell checkpoint receptors during BcAg presentation can modulate T cell function and suppress autoimmunity and progression of the disease in T1D.

## Graphical Abstract



## Keywords

Type 1 diabetes; autoimmunity; dendritic cells; T cell repressor/inhibitory receptors; beta-cell antigen; T cell tolerance

## Introduction

Type 1 diabetes (T1D) is an autoimmune disorder resulting from the progressive destruction of insulin-producing pancreatic  $\beta$  cells by autoreactive lymphocytes, leading to insulin deficiency and hyperglycemia. Pancreatic islet  $\beta$ -cell antigen (BcAg)-specific T cells play a major role in the disease process, and they arise and expand under genetic susceptibility due to defective immune regulation(1). Immune regulatory defects and disease susceptibility in both humans and experimental animals have been linked to many genetic factors including the loci encoding for T cell inhibitor/repressor receptors such as Cytotoxic T-lymphocyte-associated antigen 4 (CTLA4)(2, 3). Hence, approaches targeting these defects as well as enhancing the T cell repressor function could be valuable for T1D therapy.

The inhibitory/repressor receptors, also referred to as checkpoint receptors, expressed on T cells are critical for preventing and regulating self-antigen specific immune responses(4–6). CTLA4, programmed cell death-1 (PD1) and B- and T-lymphocyte attenuator (BTLA) are the major checkpoint receptors of CD28 family expressed on T cells(7, 8). Surface expression levels of these receptors are upregulated upon activation of T cells as a homeostatic mechanism. CTLA4 is a master regulator of peripheral T cell tolerance, T cell

homeostasis and regulatory T cell (Treg) function<sup>(9–19)</sup>. CTLA4 deficiency causes massive lymphoproliferative condition in mice, leading to their death at as early as 4 weeks of age<sup>(19)</sup>. While CTLA4 shares ligands CD80 and CD86 with CD28, a single amino acid mutation eliminates CD28 binding ability of CD80 producing a CTLA4 selective ligand, B7.1wa<sup>(20)</sup>. PD1 is induced on T and B cells upon activation and its interaction with PD-L1 and PD-L2 activates negative regulatory function<sup>(21, 22)</sup>. PD1 deficiency leads to increased susceptibility to autoimmunity<sup>(23)</sup>. BTLA is expressed on activated T and B cells, and its signals can also suppress the T-cell response<sup>(8)</sup>. BTLA interacts with the TNFR family member, HVEM (a herpes virus entry mediator)<sup>(24, 25)</sup>. While HVEM also interacts with T cell activation receptor LIGHT, its interaction with BTLA and CD160 on activated T cells through its CRD1 region induces an inhibitory response in T cells<sup>(25, 26)</sup>. Considering their role in immune tolerance, T cell checkpoint pathways, initiated by CTLA4 and PD1 particularly, are being targeted to treat cancers<sup>(27–29)</sup>. However, these therapies also produced autoimmune etiology in patients<sup>(30–33)</sup>, further substantiating their role in maintaining peripheral self-antigen tolerance.

Previously, we have shown that the dominant engagement of CTLA-4 on T cells from target tissues and antigen presenting dendritic cells (DCs) can induce Treg responses, antigen specific T cell tolerance, and prevent and suppress experimental autoimmune thyroiditis (EAT) and spontaneous type 1 diabetes (T1D) in mouse models<sup>(34–38)</sup>. Recently, we tested if exogenous expression of checkpoint-receptor selective ligands in DCs for enhancing T cell inhibitory signals can generate tolerogenic antigen presenting cells (tDCs/tAPCs)<sup>(39)</sup>. DCs that exogenously express selective ligands for CTLA4, PD1 and BTLA individually (monoligand-DCs) or in combination (multiligand-DCs) induced an antigen specific immune tolerance. These engineered DCs, CTLA4-ligand-DCs and multiligand-DCs particularly, can induce profound inhibition of T cell proliferation, modulation of cytokine response, generation of T cells with a regulatory phenotype and suppress autoimmunity in a thyroglobulin immunization induced EAT model. However, whether this engineered DC approach can be employed to suppress spontaneous autoimmunity is not known. Here, we show that engineered CTLA4-ligand-DCs and multiligand-DCs that are pulsed with pancreatic  $\beta$ -cell antigen peptide cocktail (BcAg) induced immune regulatory cytokine and Treg responses and prevented T1D in non-obese diabetic (NOD) mice for a profound duration, when treated at pre-diabetic stage. Furthermore, early-hyperglycemic mice that received these tDCs showed reversal of hyperglycemia for a significant duration. Overall, this study demonstrates that enhancing checkpoint receptor signaling during self-antigen presentation by DCs can effectively suppress autoimmunity and progression of the disease in T1D.

## Materials and methods

### Mice, antibodies, antigens, and other reagents

NOD/Ltj and NOD-BDC2.5 TCR-transgenic (BDC2.5 mice) were originally purchased from the Jackson laboratory (Bar Harbor, ME, USA). Foxp3-GFP-knockin (ki)<sup>(40)</sup> mice in the B6 background were kindly provided by Dr. Vijay Kuchroo (Harvard Medical School, MA). NOD-Foxp3-GFP-ki mice and NOD-BDC2.5-Foxp3-GFP-ki mice were generated as

described in our previous reports(41, 42). Breeding colonies of these mice were established and maintained in the specific pathogen free facility of the Medical University of South Carolina (MUSC). Glucose levels in the tail vein blood samples of mice were monitored with the Ascensia Microfill blood glucose test strips and an Ascensia Contour blood glucose meter (Bayer, Pittsburgh, PA). The mice with glucose levels >250 mg/dl for two consecutive bleeds were considered diabetic. Mice with blood glucose levels: 140–250 mg/dl and >450 mg/dl were considered early hyperglycemic and overt-diabetic respectively. All experimental protocols were approved by the Institutional Animal Care and Use Committee (IACUC) of MUSC. All methods in live animals were carried out in accordance with relevant guidelines and regulations of this committee and performed according to National Institutes of Health (NIH) Guidelines on Humane Care and Use of Laboratory Animals.

Immunodominant  $\beta$ -cell-Ag (BcAg) peptides [namely: 1) insulin B(9–23); 2) GAD65(206–220); 3) GAD65(524–543); 4) IA-2 $\beta$ (755–777); 5) IGRP(123–145), 6) GAD65(286–300), 7) Insulin B(15–23), and 8) IGRP(206–214)], and BDC2.5 TCR-reactive peptide (YVRPLWVRME; referred to as BDC2.5-peptide) were used. Peptides 1–8 were pooled at an equal molar ratio and used as BcAg to pulse the DCs for in vitro and in vivo experiments, as done in our earlier studies, using some of these peptides(36, 38, 42, 43).

### Generation of cDNA vectors and lentivirus production

The third-generation replication-incompetent lentiviral cDNA cloning vector and packaging plasmids were purchased from System Bio Inc and modified(39). Details on the generation of vector constructs: pCDH1 vector with cDNA for B7.1wa, PD-L1 or HVEM-CRD1 and GFP reporter under a separate promoter are described in our previous report(39). Most experiments employed DCs engineered using these constructs. pCDH1-vector encoding a “multi-gene/cDNA sequence” (B7.1wa, HVEM-CRD1 and PDL1 cDNAs separated by self-cleaving P2A and T2A peptide sequences: pCDH1-EF1-copGFP-B7.1wa-T2A-HVEM-CRD1-P2A-PDL1; pCDH1-multiligand vector) (shown in supplemental Fig. 1) was also constructed by gene synthesis and custom cloning service of Genscript and used in some experiments where specified. Comparable results were obtained when single-cDNA and multi-cDNA constructs were used in in vitro assays. For lentivirus production, HEK293T cells were transfected with cDNA expression vectors along with packaging vectors by calcium phosphate method. GPRG cells(44), provided by National Gene Vector Biorepository (Indianapolis, IN, USA), were also used as virus packaging cells in some experiments. Preparation and titration of virus particles are described in our recent report(39).

### Generation of engineered BMDCs

Generation of DCs from BM cells (BMDCs) and engineering using lentiviral particles are described in our previous reports(36–39). Briefly, BM cells from prediabetic NOD mice were cultured in complete RPMI 1640 medium containing 10% FBS, in the presence of GM-CSF (20 ng/ml), at 37°C in 5% CO<sub>2</sub> for 3 days and then for further 3 days in fresh complete RPMI 1640 medium, containing GM-CSF (20 ng/ml) and IL-4 (5 ng/ml). For lentiviral transduction, cells from 3-day-old cultures were used. A final concentration of >5×10<sup>8</sup> TU/ml virus (for individual ligand-DCs and control-DCs) or a cocktail of >1.5

$\times 10^9$  TU/ml virus (for multiligand-DCs) was used for transducing cells in the presence of polybrene (8  $\mu\text{g/ml}$ ) and protamine sulfate (10  $\mu\text{g/ml}$ ) for up to 24h, washed and cultured in fresh medium containing GM-CSF and IL4 for an additional 48h. Transduction efficiency was examined by microscopy and/or FACS, and the cells that showed transduction efficiency of  $>90\%$  were washed thoroughly and counted before in vitro or in vivo use. Ligand functionality was determined by assessing the increase in soluble receptor binding by ligand-DCs as compared to control-DCs having endogenous ligand expression.

### **In vitro antigen presentation assay**

BDC2.5-peptide pulsed NOD BMDCs ( $2 \times 10^4$  cells/well) were plated in triplicate in 96-well U-bottom tissue culture plates along with unlabeled or CFSE-labeled purified CD4<sup>+</sup> T cells from NOD-BDC2.5 or NOD-BDC2.5-Foxp3-GFP mice ( $1 \times 10^5$  cells/well) in complete RPMI 1640 medium. After 4 days of culture, cells were subjected to surface and/intracellular staining using fluorochrome-labeled Abs and examined for CFSE dilution or cytokine production by FACS. Cells were subjected to brief activation using PMA and ionomycin (4h) in the presence of Brefeldin A, before staining to detect intracellular cytokines. In some assays, equal number of live cells from primary cultures were seeded along with BDC2.5 peptide and freshly isolated NOD-mouse splenic DCs for 24 h, and the spent media were tested for cytokine levels by ELISA or Luminex multiplex assay. In some assays, BcAg (peptide cocktail) pulsed DCs were incubated with T cells from the pancreatic lymph nodes (PnLNs) of early-hyperglycemic mice for up to 4 days and subjected to various assays.

### **Treatment of NOD mice with engineered DCs**

Female NOD mice of different age groups and disease stages were injected i.v. with  $2 \times 10^6$  BcAg-pulsed control-DCs or ligand-DCs (B7.1wa-DCs or multiligand-DCs) once or twice, at a 15-day interval. Cohorts of animals were examined for blood glucose levels every week. In some experiments, two or four-weeks post-treatment, mice were euthanized to test for T cell response to ex vivo challenge with the BcAg, T cell phenotype, and/or insulinitis.

### **Adoptive transfer of immune cells to NOD-Rag1<sup>-/-</sup> mice**

Total PnLN cells of engineered DC treated mice were cultured overnight in round-bottom 96-well plates in the presence of anti-CD3-Ab (2  $\mu\text{g/ml}$ ), washed, and injected into 6–8-week-old NOD-*Rag1*<sup>-/-</sup> mice. In some experiments, splenic T cells enriched from the spleens of untreated early-hyperglycemic mice were cultured in the presence of BcAg pulsed DCs for 4 days, and live T cells enriched from these primary cultures were injected ( $1 \times 10^6$  cells/mouse) i.v. into female NOD-*Rag1*<sup>-/-</sup> mice. These T cell recipients were tested for blood glucose levels every week. In some experiments, purified CD4<sup>+</sup> T cells from 4-week-old BDC2.5 TCR-Tg mice that were cultured in the presence of BDC2.5 peptide-pulsed various engineered DCs, as described above and cells from these cultures ( $1 \times 10^6$  cells/mouse) were injected i.v. into 4-week-old male WT-NOD mice. Blood glucose levels in these mice were tested every other day. Cohorts of mice were euthanized at different time-points to determine the degree of insulinitis.

## Histochemical analysis of pancreatic tissues

Pancreatic tissues were fixed in 10% formaldehyde and 5- $\mu$ m paraffin sections were made and stained with H&E. Stained sections were analyzed in a blinded fashion using a grading system: 0, no evidence of infiltration; 1, peri-islet infiltration (<5% of the islet area); 2, 5–25% infiltration; 3, 25–50% infiltration; and 4, >50% infiltration as described in our earlier studies(36, 38, 42, 43, 45, 46). In some experiments, pancreatic sections were stained using anti-insulin antibody followed by Alexa fluor 488- or 568- linked secondary antibodies and DAPI and scored for insulinitis based on DAPI-positive cells in islet areas and insulin expression. Insulinitis was scored as described for H&E stained sections and insulin positive and negative islets were counted as described in our previous reports(41–43). Briefly, we employed the grading system: 0+, insulin+ with no evidence of infiltration; 1+, insulin+ with peri-insulinitis (<5% of the islet area with infiltration); 2+, insulin+ with 5–25% infiltration; 3+ insulin+ with 25–50% infiltration; and 4+, insulin+ with 50–100% infiltration; 4-, insulin- fully infiltrated islets; and 5-, insulin- with infiltrates in residual islets structures.

## Statistical analysis

Mean, SD, and statistical significance ( $p$ -value) were calculated using Microsoft Excel or GraphPad prism statistical application. In most cases, values of the individual test group (ligand-DC group) were compared with that of control group (control-DC group). For most comparisons, Mann-Whitney or paired  $t$ -test was employed. Statistical analysis of insulinitis severity was done by Fisher's exact test. T1D incidence in different groups was compared by employing log-rank test. Specific methods used for obtaining  $p$ -values were described under figure legends.  $p < 0.05$  was considered significant.

## Results:

### T cell-checkpoint receptor selective ligand expressing DCs modulate T cell response in vitro

In a recent report, we showed that exogenous expression/overexpression of selective ligands of CTLA4, PD1 and BTLA receptors could make the DCs produce, upon antigen presentation, tolerogenic effects on T cells, presumably through enhanced receptor engagement(39). We engineered the BMDCs to express B7.1wa, PD-L1 and HVEM-CRD1 exogenously, individually and in combination and characterized their receptor-binding, antigen presenting and tolerogenic properties extensively(39). WT B7.1 binds to both CTLA4 and CD28, and WT HVEM binds to BTLA and LIGHT. CTLA4- and BTLA-selective binding abilities of B7.1wa and HVEM-CRD1 were described before(20, 47). We found that B7.1wa-DCs and multiligand-DCs have a better ability to induce robust Treg responses and overall immune modulation compared to control-DCs, and other mono-ligand DCs such as PD-L1-DCs and HVEM-CRD1-DCs(39). This report which used a mouse thyroglobulin immunization model showed that B7.1wa-DCs and multiligand-DCs are effective in modulating T cell function and suppressing autoimmune thyroiditis. However, whether such engineered DCs are useful for modulating spontaneous autoimmunity such as T1D is not known. Considering the superior tolerogenic effects of B7.1wa-DCs

and multiligand-DCs, the current study examined the antigen presenting and tolerogenic properties of these engineered-DCs (Fig. 1A & B) in the context of T1D in NOD mice.

To assess the abilities of inhibitory ligand expressing NOD-mouse DCs to activate T cells, control-DCs, B7.1wa-DCs and multiligand-DCs were pulsed with BDC2.5 peptide and cultured in the presence of CD4<sup>+</sup> T cells enriched from NOD-BDC2.5 TCR-Tg mice or NOD-BDC2.5-Foxp3-GFP-ki mice. As observed in Fig. 1C, T cells cultured with B7.1wa-DCs and multiligand-DCs showed lower frequencies of IFN $\gamma$ <sup>+</sup> cells, higher frequencies of IL10<sup>+</sup> and Foxp3<sup>+</sup> cells, and a lower rate of proliferation as compared to those cultured with control-DCs. We also observed that considerable proportion of sorted GFP negative cells (Foxp3<sup>-</sup> cells) from Foxp3-GFP-ki mice expressed GFP upon activation using ligand-DCs (Supplemental Fig. 2A) suggesting that, in addition to potential expansion of existing Tregs, new Tregs are induced by the tDCs. Ligand-DC activated BDC2.5 T cells produced higher amounts of immune regulatory cytokines IL10 and TGF $\beta$ 1 and showed diminished production of the pro-inflammatory cytokine IFN $\gamma$  upon rechallenge with BDC2.5 peptide (Fig. 1D and supplemental Fig. 2B). Of note, as expected, DCs that were not pulsed with BDC2.5 peptide failed to induce significant Treg response or cytokine production in BDC2.5 T cells (Supplemental Fig. 3A & B). Further, T cell viability in primary cultures containing peptide-pulsed, ligand-DCs was considerably higher compared to that contained control-DCs (Supplemental Fig. 4). We then examined the ability of BcAg pulsed engineered DCs to modulate the properties of splenic T cells obtained from early-hyperglycemic WT NOD mice in vitro. T cells from these primary cultures were challenged with BcAg and examined for the degree of proliferation and levels of secreted cytokines. As observed in Fig. 1E, Ag presentation by B7.1wa and multi-ligand DCs resulted in the modulation of T cell phenotype, as indicated by significantly lower levels of proinflammatory IFN- $\gamma$  and IL17 and higher immune regulatory IL-10 and TGF- $\beta$ 1 cytokine responses, compared to T cells activated using control-DCs. These results show that enhanced DC-directed negative signaling, during BcAg presentation, suppresses the pro-inflammatory activation of, and promotes immune regulatory features in, T cells.

### **B7.1wa-DC and multiligand-DC activated T cells fail to induce hyperglycemia**

To assess if the in vitro activation of T cells by ligand-DCs has an impact on their diabetogenic properties, WT-NOD and BDC2.5 T cells were cultured in the presence of control-DCs, B7.1wa-DCs and multiligand-DCs in vitro for 4 days as described for Fig. 1, equal number of enriched live cells from these cultures were injected i.v. into NOD-*Rag1*<sup>-/-</sup> mice or young WT NOD mice and monitored for hyperglycemia. Fig. 2A shows that 100% of NOD-*Rag1*<sup>-/-</sup> mice that received control-DC activated WT T cells developed hyperglycemia during 8–12-week period after cell transfer. However, mice that received B7.1wa-DC and multiligand-DC activated WT T cells failed to develop hyperglycemia, at least for 18 weeks post-cell transfer. Pancreatic tissues of B7.1wa-DC and multiligand-DC activated T cell recipients that were euthanized 4-weeks post cell-transfer, showed significantly lower degree of insulinitis compared to those which received control-DC activated T cells (Fig. 2B). In BDC2.5 T cell recipient WT NOD mice, while 100% of mice that received control-DC activated T cells developed overt-hyperglycemia within 8 days, recipients of B7.1wa-DC and multiligand-DC activated T cells failed to show hyperglycemia

for at least 15 days (Fig. 2C). Further, recipients of ligand-DC activated T cells showed little or no insulinitis on day 5 post-cell transfer as opposed to 100% of the islets of control group showing severe grade 3 or grade 4 insulinitis (Fig. 2D). Overall, these results show that T cell inhibitory ligand expressing-DCs can suppress the diabetogenic/pathogenic activation of T cells upon antigen presentation in vitro.

### **Single injection of B7.1wa-DCs or multiligand DCs causes significant modulation of BcAg specific T cell response in NOD mice**

To determine the effect of treatment with engineered DCs on immune cells, ten-week-old pre-diabetic female NOD mice were injected once with control-DCs, B7.1wa-DCs and multiligand-DCs, euthanized after 15 days, and PnLN cells were examined for their proliferative response upon challenge with BcAg, and cytokine and Foxp3 expressing CD4<sup>+</sup> T cell frequencies. As observed in Fig. 3A, PnLN T cells from B7.1wa-DC and multiligand-DC recipients proliferated at significantly lower rate compared to the cells from control-DC recipient mice. Intracellular staining of fresh PnLN cells showed that while IFN $\gamma$ <sup>+</sup> and IL17<sup>+</sup> T cell frequencies were significantly lower in B7.1wa-DC and multiligand-DC recipients, IL10<sup>+</sup> and Foxp3<sup>+</sup> T cell frequencies were higher in these mice compared to control-DC recipients.

To assess if BcAg loaded ligand-DC treatment, compared to control-DC treatment, impacts the global T cell function, spleen cells from the treated mice were examined for proliferative, IL2 and IL10 responses upon stimulation using anti-CD3 antibody. Fig. 3B shows that the degree of anti-CD3 antibody induced proliferation of CD4<sup>+</sup> T cells was not significantly different between control-DC and ligand-DC treated mice. However, modestly lower IL2 responses, albeit not significant statistically, were observed with splenic T cells from ligand-DC treated mice compared to control-DC recipients. On the other hand, immune regulatory cytokine, IL10 was produced at higher levels by cells from ligand-DC treated mice, multiligand-DC recipients particularly, compared to control-DC recipients. Overall, sustained ability of most T cells to proliferate and produce comparable amounts of IL2 upon CD3 stimulation suggests that BcAg-pulsed ligand-DC induced modulation of immune response is primarily BcAg specific and immune regulatory in nature.

### **B7.1wa-DC and multiligand-DC treatments, at pre-diabetic stage, exert profound protection from T1D and immune modulation**

To assess the impact of treatment using engineered DCs on T1D incidence in NOD mice, ten-week-old pre-diabetic female NOD mice were injected i.v., twice at 15-day interval, with BcAg-pulsed control-DCs, B7.1wa-DCs or multiligand-DCs and monitored for hyperglycemia. As shown in Fig 4A, diabetes onset was detected in the control groups of mice within 4–6 weeks of monitoring as compared to 18-weeks post-treatment in multiligand DC treated mice and 12 weeks post-treatment in B7.1wa-DC treated mice. While about 60% and more than 80% respectively of B7.1wa-DC and multiligand-DC treated mice remained euglycemic for at least 30-weeks post treatment initiation, 100% of the control-DC recipient mice turned hyperglycemic during this period. Of note, recipient of ligand-DCs that were not pulsed with BcAg showed only a modest, but not statistically significant, delay in the clinical disease onset (Supplemental Fig. 5).



To assess if treatment using engineered-DCs causes protection of pancreatic islets from destruction, tissues from treated mice were examined for insulinitis, 30 days post treatment. Fig. 4B shows that mice that received B7.1wa-DCs and multiligand-DCs had significantly lower frequencies of islets with severe insulinitis compared to control-DC recipients. While about 70% of pancreatic islets from control-DC recipients showed severe insulinitis grade of 3, only about 30% and 10% of pancreatic islets in mice that received B7.1wa-DCs or multiligand-DCs respectively, had severe insulinitis. Overall, these results show that treatment using BcAg-presenting, T cell checkpoint-receptor ligand DCs, multiligand-DCs particularly, can prevent hyperglycemia for a profound duration.

Fifteen days post-treatment, cohorts of mice were examined for Foxp3+ T cell frequencies and BcAg-challenge induced cytokine secretion. As observed in Fig. 4C, Foxp3+ T cell frequencies were profoundly higher in the spleen and PnLN of B7.1wa-DC and multiligand-DC treated mice compared to control-DC treated mice. PnLN cells from B7.1wa-DC and multiligand-DC treated mice, compared to control-DC recipients, showed significantly lower IFN $\gamma$ , and higher IL10 and TGF $\beta$ 1 secretion upon ex vivo challenge exposure to BcAg (Fig. 4D and supplemental Fig. 6). These results suggest that the B7.1wa-DC and multiligand-DC treatments promote protection of  $\beta$ -cells from destruction, perhaps through enhancing immune regulation, leading to prevention of hyperglycemia.

### **Treatment with B7.1wa-DCs or multiligand-DCs at early-hyperglycemic stage results in transient reversal of hyperglycemia**

To determine if blood glucose levels in mice with early-hyperglycemia can be altered by treating with ligand-DCs, female-NOD mice with blood glucose levels between 140–250 mg/dl (12–25 weeks of age) were injected twice, 15-days apart, with control-DCs, B7.1wa-DCs or multiligand-DCs and examined for blood glucose levels every week. As shown in Fig. 5A, while 100% of the control-DC treated mice progressed to overt hyperglycemia within 8 weeks, all mice that received ligand-DCs reverted to and remained euglycemic for at least 8 weeks post-treatment initiation. Thirty days post-treatment (45 days post-treatment initiation), pancreatic tissues were collected from cohorts of mice to examine for insulinitis and to assess the abundance of insulin positive islets. Tissue sections were stained using anti-insulin and anti-glucagon antibodies and examined for the frequencies of “functional islets”. As shown in Fig. 5B, about 80% and 90% of islet areas, with or without immune cell infiltration, of B7.1wa-DC and multiligand-DC recipient mice respectively showed some degree of islet function, as indicated by both insulin and glucagon positivity, compared to about 5% insulin positive islet remnants in control-DC treated mice. These results show that reversal of hyperglycemia is possible by treatment with T cell-checkpoint receptor ligand expressing engineered DCs.

As observed in prediabetic mice (Fig. 4C and 4D), PnLN cells from these early-hyperglycemic groups of mice showed significantly higher frequencies of Foxp3+ T cells and higher production of IL10 and TGF $\beta$  and lower secretion of IFN $\gamma$  compared to control-DC recipients (not shown). To determine the pathogenic effect of these immune cells, PnLN cells from DC treated individual mice (harvested 30-days post-treatment) were cultured in the presence of anti-CD3 antibody for 24h, injected into NOD-*Rag1*<sup>-/-</sup> mice and monitored

for hyperglycemia. As observed in Fig. 5C, all recipients of PnLN cells from control-DC treated donors turned diabetic within 11 weeks of cell transfer. However, mice that received PnLN cells from B7.1wa-DC and multiligand-DC treated mice showed a significant delay in the onset of hyperglycemia, or failed to develop hyperglycemia, during the 20-week monitoring period. This suggested that the diabetogenic properties of immune cells of early-hyperglycemic mice were profoundly suppressed in ligand-DCs treated mice. Overall, these results show that T cell checkpoint-receptor selective ligand expressing DCs, multiligand-DCs particularly, can be effective tolerogenic DCs and used to induce self-antigen specific T cell immune modulation for preventing and suppressing spontaneous autoimmunity in T1D

## Discussion

BcAg-specific T cells, which are induced/expanded in susceptible subjects and spontaneous disease models, likely because of defective immune regulation(1, 48, 49), play an important role in islet destruction and T1D onset. Immune regulatory defects and T1D disease susceptibility in humans and mouse models have been linked to multiple genetic loci including that encoding CTLA-4(50). Splice variants of CTLA-4 have been identified as potential risk factors contributing to the development of T1D(2). While CTLA4 is considered as the master checkpoint receptor on T cells, other receptors such as PD1 and BTLA also play important roles in maintaining peripheral tolerance(6, 8, 51, 52). Checkpoint inhibitor therapy for cancers show that interfering with their functions, CTLA4 and PD1 functions particularly, can effectively break the T cell tolerance and treat many cancers(53). Recent reports show that, although these therapies are effective against cancers, they produce autoimmune manifestations that include T1D and rheumatic diseases(30–32, 54, 55), substantiating their role in peripheral self-tolerance. Therefore, these checkpoint receptors can, individually or in combinations, be targeted for promoting BcAg-specific T cell tolerance and preventing/suppressing spontaneous autoimmunity in T1D. Importantly, inhibition of T cell response through checkpoint receptors and inducing antigen specific tolerance requires delivering active negative signals through them in conjunction with TCR engagement(51, 56–59). Therefore, induction of BcAg-specific tolerance is possible because targeting CTLA4, PD1 and/or BTLA on T cells from professional antigen presenting cells such as DCs and enhancing the negative signaling strength occurs concurrently with the engagement of specific TCRs on T cells.

DCs are considered the most effective APCs and the only type of cells that can activate naïve T cells(60, 61). Hence, engineering the DCs to ensure maintenance of their tolerogenic function, even under pro-inflammatory conditions, is critical for establishing antigen specific T cell tolerance to prevent and treat T1D. In this regard, DCs that are selectively deleted of costimulatory molecules or ectopically expressing cytokine and non-cytokine factors have shown the ability to induce T cell tolerance(62–64). We also reported that co-stimulation by CD80, preferential ligand of CTLA4 in terms of binding avidity, results in the generation of IL-10 dependent TGF- $\beta$ 1+ regulatory T cells (34–38). We reported that enhancing CTLA4 agonist strength on mature DC surface could lead to the dominant engagement of this repressor-receptor on T cells upon antigen presentation and induce antigen specific tolerance and prevent/suppress autoimmune thyroiditis and T1D(37, 38). Recently, using antigen immunization models, we demonstrated that lentiviral transduction approach can be

employed to generate tDCs that are exogenously expressing selective ligands for multiple T cell repressor receptors (viz: CTLA4, PD1 and BTLA) and these DCs, particularly B7.1wa-DCs and multiligand-DCs as described in this study, can hyper-activate T cell inhibitory pathways individually or in combination during antigen presentation and induce antigen specific immune tolerance(39). Our current study shows that treatments using B7.1wa-DCs and multiligand-DCs effectively induce BcAg tolerance and protects NOD mice from T1D.

Our current study shows that tDC, B7.1wa-DC and multiligand-DC, treatments promote protection from  $\beta$ -cell destruction and prevent T1D for a significant period when treated at pre-diabetic stage. These treatments also reverse the elevated blood glucose levels and improves the islet function significantly, when the treatment was initiated at an early-hyperglycemic stage. These observations suggest that a well-designed tDC approach such as the one described here has the potential to prevent and treat T1D in human subjects. Importantly, our observation that multiligand-DCs are more effective than B7.1wa-DCs in promoting protection from T1D in NOD mice suggests that evidence-based approaches for targeting multiple pathways is important for achieving effective and long-lasting T cell tolerance. Of note, our previous report also showed that while B7.1wa-DCs are more efficient in inducing Treg response and immune tolerance than the other checkpoint-ligand expressing DCs such PD-L1-DCs and HVEM-CRD1-DCs, and treatment using multiligand-DCs produced the most effective overall immune modulation(39). Further, our previous reports have consistently demonstrated that enhanced engagement of CTLA4 during antigen presentation by DCs or autoimmune target tissue can produce immune tolerance and protection from autoimmunity, in both spontaneous and immunization-induced models(37, 38). However, the current study and our recent report(39) demonstrates that concurrent engagement of other checkpoint receptors along with CTLA4 could enhance the tolerogenic ability of CTLA4 targeting DCs to achieve longer-lasting protection from spontaneous autoimmune manifestations.

Antigen specificity of the T cell tolerance induced by BcAg-pulsed B7.1wa- and multiligand-DCs is evident from different observations of this study. tDCs that were not pulsed with antigens failed induce proliferation or modulate the T cell function. Importantly, not only the recipients of tDC-activated BDC2.5 T cells, but also those that received tDC-activated WT T cells failed to develop hyperglycemia. Moreover, challenge of immune cells from tDC treated mice using BcAg, but not anti-CD3 antibody, showed significantly diminished proliferation. Further, the release of large amounts of immune regulatory cytokines occurred primarily when the cells were challenged with BcAg. Notably, treatment with tDCs that were not pulsed with BcAg could produce only a modest, but not statistically significant, delay in the onset of T1D supporting the notion that the disease protection achieved using tDCs is an antigen specific effect. Although it is possible that inoculated tDCs could capture small amounts of endogenous antigens, particularly in the pancreatic microenvironment, and induce limited tolerogenic response, exogenous antigen loading appears to be necessary for achieving significant protection. Our earlier study using animals that were immunized with ovalbumin or mouse-thyroglobulin(39) has shown a profound increase in Foxp3+ T cells when treated with B7.1wa-DCs and multiligand-DCs. The current study using a spontaneous autoimmune diabetic model also showed significantly

higher frequencies of Foxp3<sup>+</sup> cells in ligand expressing DC treated mice compared to control-DC recipients.

Overall, our observations show that powerful tAPCs for T1D therapy can be generated by engineering the DCs to express the selective ligands of multiple T cell checkpoint receptors simultaneously, and this could be an attractive and effective approach for inducing BcAg specific T cell tolerance. Moreover, our previous report(39) and the current study demonstrating the ability of these engineered DCs, multi-ligand DCs particularly, to affect multiple aspects of immune function, viz: higher Treg frequencies and immune regulatory cytokine responses, and lower pro-inflammatory IFN $\gamma$  and IL17 responses against self-antigens suggest that such tDCs could have therapeutic value not only in T1D, but also can be exploited to treat other autoimmune diseases. Importantly, efficient engineering of the DCs for constitutive overexpression of multiple T cell inhibitory ligands, as described in this study, makes them less susceptible to changes, compared to previously employed immature DCs, in the functional properties upon in vivo therapeutic delivery; hence is desired for clinical translation for treating T1D.

## Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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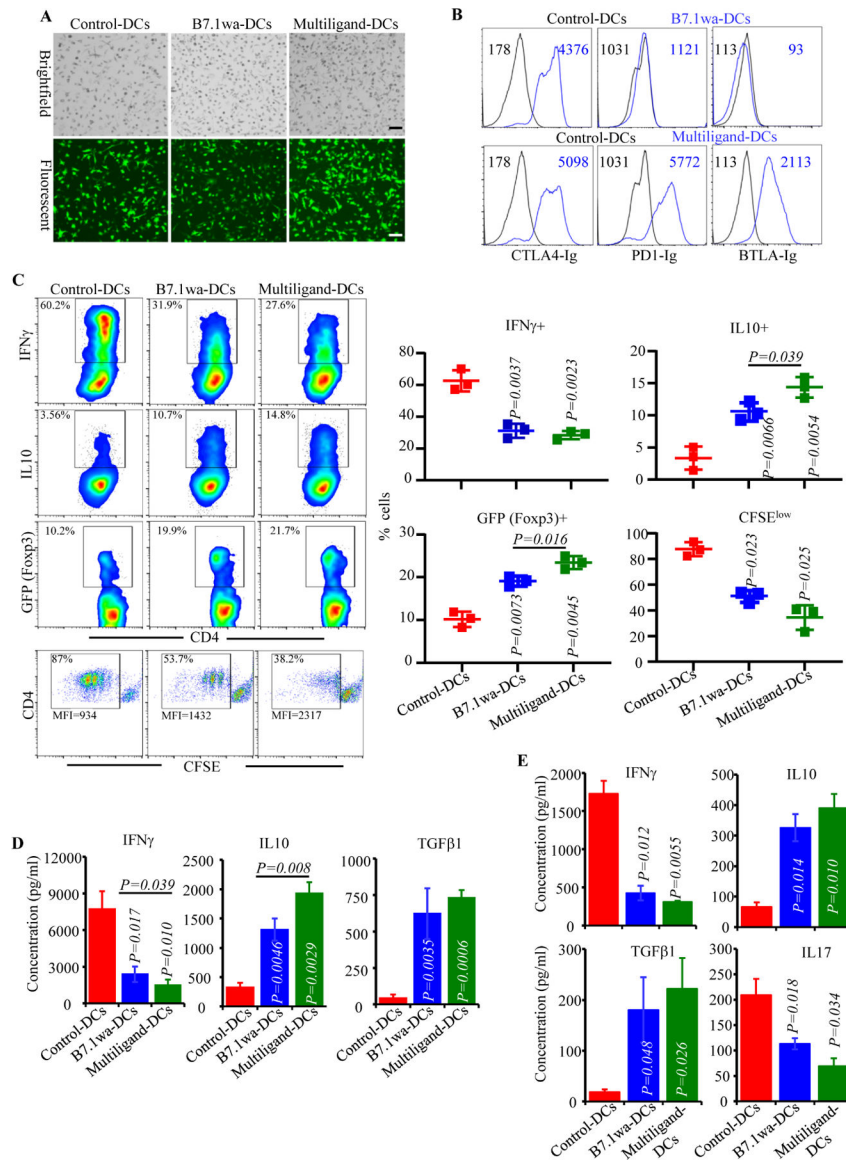
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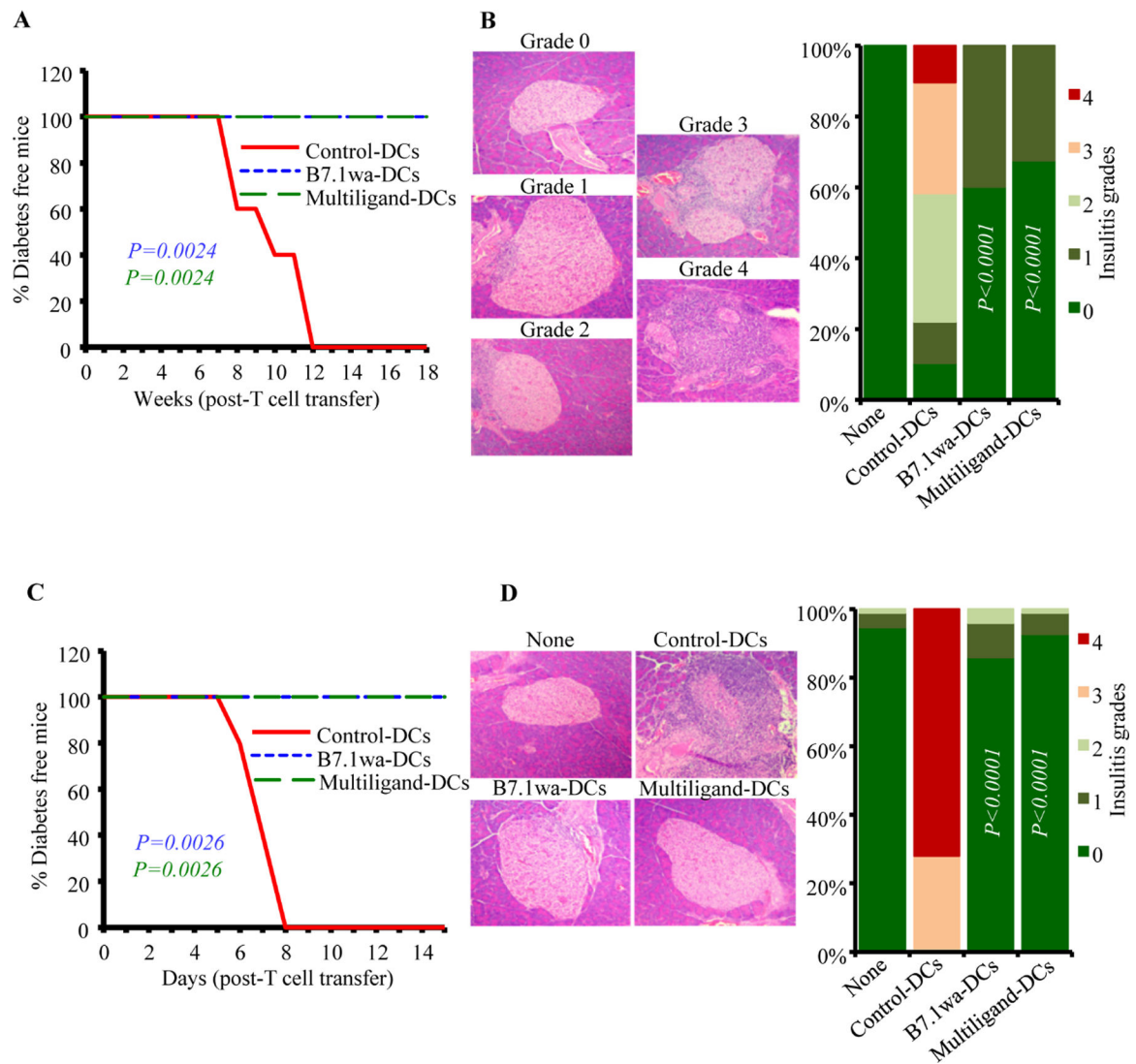


**FIGURE 1: Characterization of the antigen presenting properties of engineered NOD mouse DCs.**

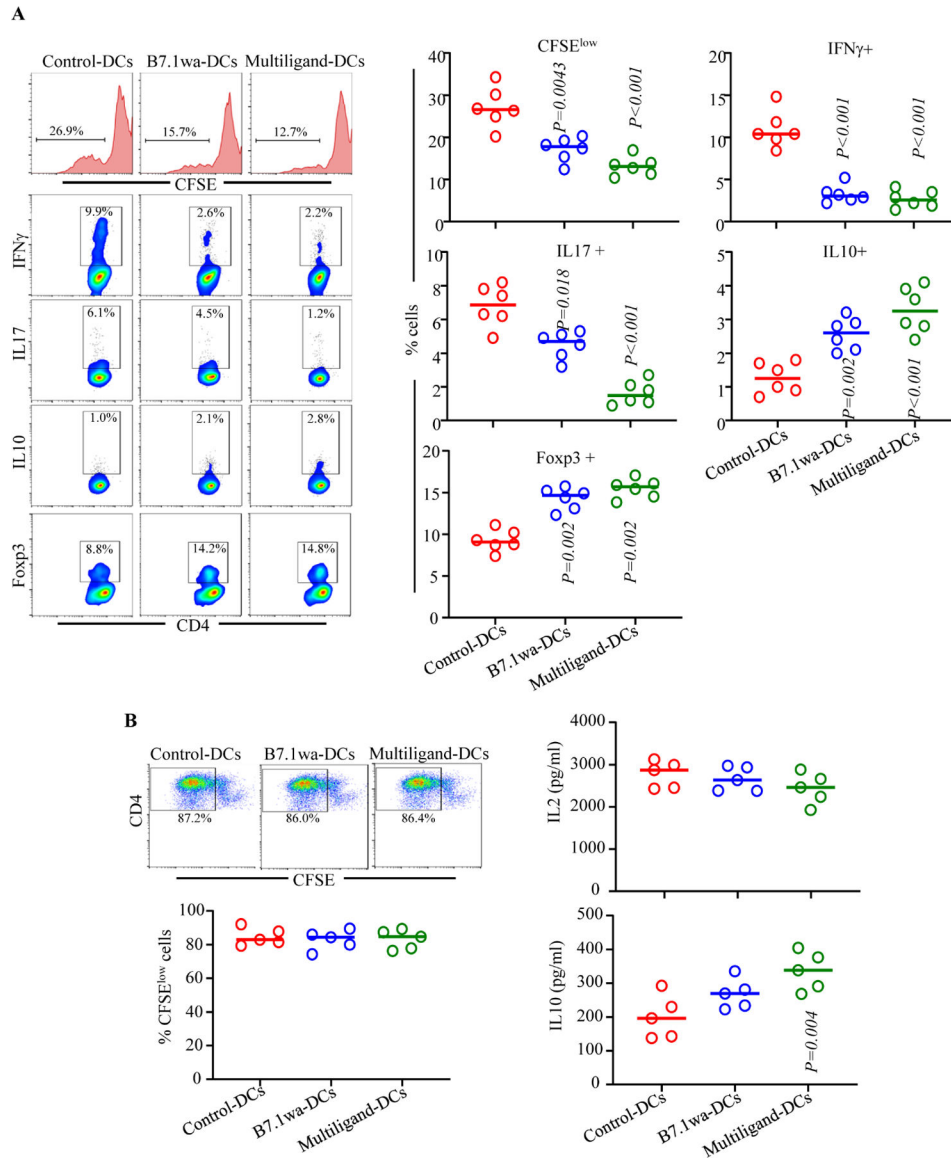
NOD mouse BM DCs were transduced with lentiviral vectors as described in Materials and methods. **A**) Examples of transduction of DCs using various preparations of virus with GFP reporter are shown. **B**) Functional ligand expression levels on DCs were determined after incubating with soluble receptors (CTLA4-Ig, PD1-Ig or BTLA-Ig), followed by PE-labeled Fab(2) fragment of anti-IgG (Fc specific) Ab and testing by FACS. Histogram overlay graphs (representative of 3 independent experiments) showing ligand specific staining of control virus transduced and ligand virus transduced DCs along with mean fluorescence intensity (MFI) values are shown. Both control and ligand DCs were also stained using control Ig reagents to assess the background staining (not shown). **C**) BDC2.5-peptide pulsed engineered DCs were used in antigen presentation assays by culturing them with CD4<sup>+</sup> cells isolated from BDC2.5 mouse spleens. After 4 days, cells from the primary cultures were stimulated using PMA and ionomycin, in the presence of Brefeldin A for



4h, stained for intracellular cytokines IFN $\gamma$  and IL10, and subjected to FACS analysis. CD4<sup>+</sup> T cells from BDC2.5-Foxp3-GFP mice were also used in primary cultures for 4 days and subjected to FACS to determine GFP<sup>+</sup> CD4<sup>+</sup> cell frequencies by FACS. In addition, CFSE labeled CD4<sup>+</sup> T cells from BDC2.5 mice were used in primary culture for 4 days and examined for percentage of cells with CFSE dilution and their MFI by FACS. CD4<sup>+</sup> cells were gated and representative FACS plots (left) and Mean $\pm$ SD values of 3 independent experiments (transduction using three preparations of lentivirus), each done in triplicate (right), are shown. **D**) Equal number of live BDC2.5 T cells isolated from primary cultures, by negative selection magnetic sorting, were cultured with BDC2.5 peptide pulsed fresh splenic DCs for 48h, and supernatants were subjected to Luminex multiplex assay or ELISA. **E**) PnLN cells from early hyperglycemic NOD mice were cultured with BcAg-pulsed engineered-DCs for 4 days, equal number of T cells isolated from these primary cultures were challenged using BcAg-pulsed fresh splenic DCs, and supernatants were tested for cytokine levels as done for BDC2.5 T cells. pCDH1-multiligand vector, instead of individual ligand vectors, was used for generating multiligand-DCs for panel E. Mean $\pm$ SD values of 3 independent experiments (transduction using three preparations of lentivirus), each done in triplicate, are shown for panels D and E. *P*-value by paired *t*-test for panels C-E. All *P*-values are in comparison with control-DC group unless indicated (for B7.1wa group vs multiligand-DC group). All in vitro experiments were repeated multiple times and comparable results were obtained at least thrice. In some experiments, two to three batches of viral preparations were tested in parallel and results from one such representative experiment are shown.



**FIGURE 2: Diabetogenic properties of T cells activated in vitro using engineered-DCs.** PnLN T cells from early-hyperglycemic WT-NOD mice and CD4<sup>+</sup> T cells from BDC2.5 mice were subjected to primary culture in the presence of various DC preparations for 4 days as described for Fig. 1. **A&B**) Live T cells from primary cultures of WT PnLN T cells were enriched and injected i.v. into 6-week-old female NOD-*Rag1*<sup>-/-</sup> mice ( $1 \times 10^6$  cells/mouse) and tested for blood glucose levels every week for up to 18 weeks (A; 5 mice/group) or euthanized 30 days post-cell transfer (B; 4 mice/group). pCDH1-multiligand vector, instead of individual ligand vectors, was used for generating multiligand-DCs for panels A&B. **C&D**) Live T cells from the primary cultures of BDC2.5 cells were injected i.v. into 4-week-old WT male NOD mice and tested for blood glucose levels every week for up to 15 weeks (C; 5 mice/group) or euthanized 5-days post-cell transfer (D; 4 mice/group). At least 120 islets from intermittent sections were examined for each group (at least 30 islet areas/mouse) for panels B and D. *P*-values by log-rank test for panels A and C, and Fisher's exact test for panels B and D. All *P*-values are in comparison with control-DC group. Group "none" did not receive T cells.



**FIGURE 3: Treatments using B7.1wa-DCs and multiligand-DCs result in modulation of immune response to BcAg in vivo.**

BcAg-pulsed engineered DCs were injected i.v. once into 10-week-old pre-diabetic female NOD mice ( $2 \times 10^6$  cells/mouse; single injection). **A**) Cohorts of mice (6 mice/group) were euthanized 15-days post-treatment, PnLN cells were labeled with CFSE, cultured in the presence of BcAg for 72h and examined for CFSE dilution by FACS. Fresh PnLN cells were also cultured in the presence of PMA/Ionomycin and Brefeldin A for 4h and subjected to intracellular staining and FACS to detect cytokine positive cell frequencies. In addition, fresh cells were examined for Foxp3+ cell frequencies by FACS. CD4+ T cells were gated for these analyses. **B**) Splenocytes from additional cohorts of mice (5 mice/group) were labeled using CFSE, stimulated using anti-CD3 antibody ( $2 \mu\text{g/ml}$ ) for 72h, and CFSE dilution was examined by FACS. Supernatants from these cultures were tested for secreted cytokines by multiplex assay. *P*-value by Mann-Whitney test. All *P*-values are in comparison

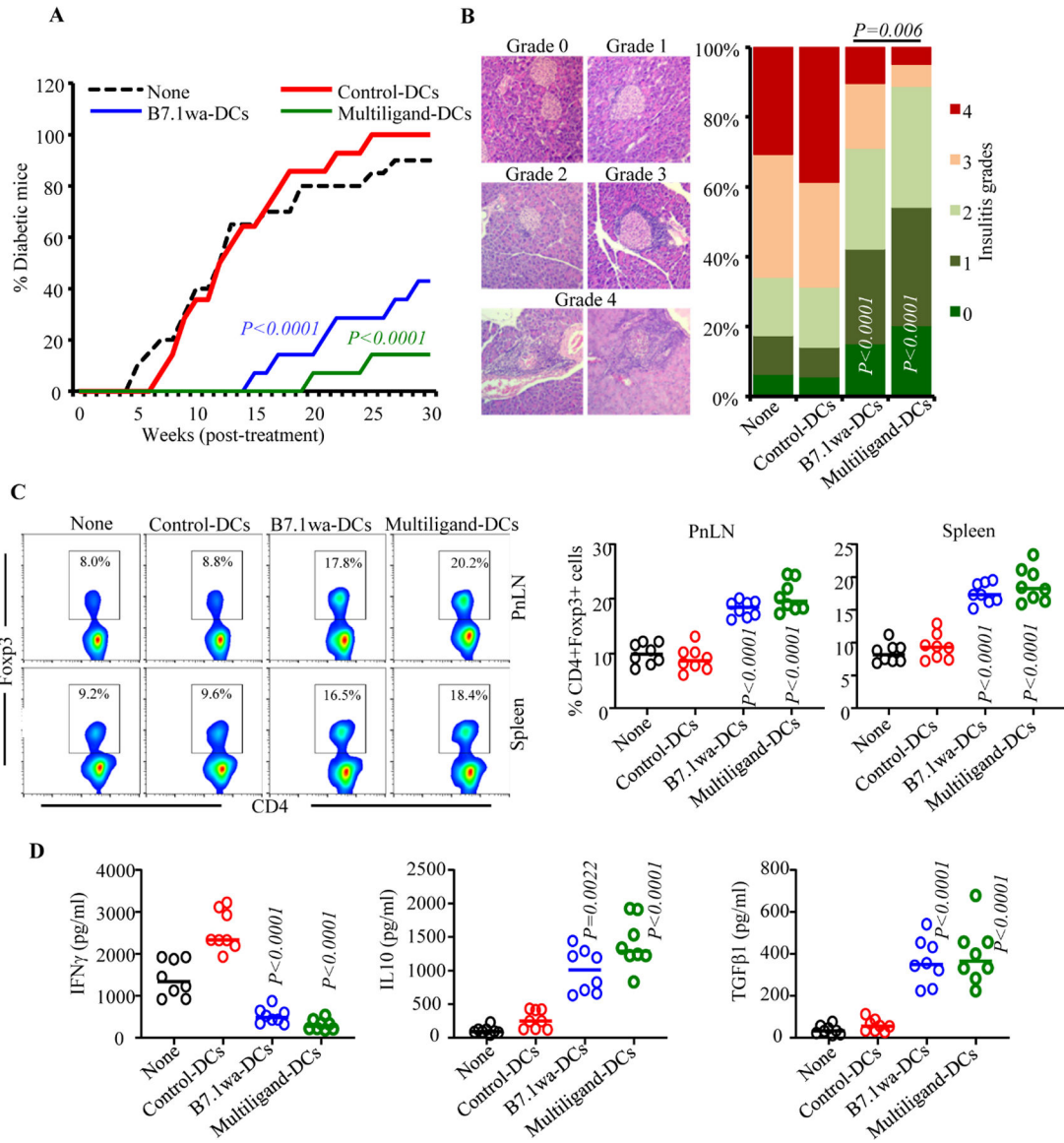
with control-DC group. This experiment was repeated once using an additional 4 mice/group and obtained comparable results.

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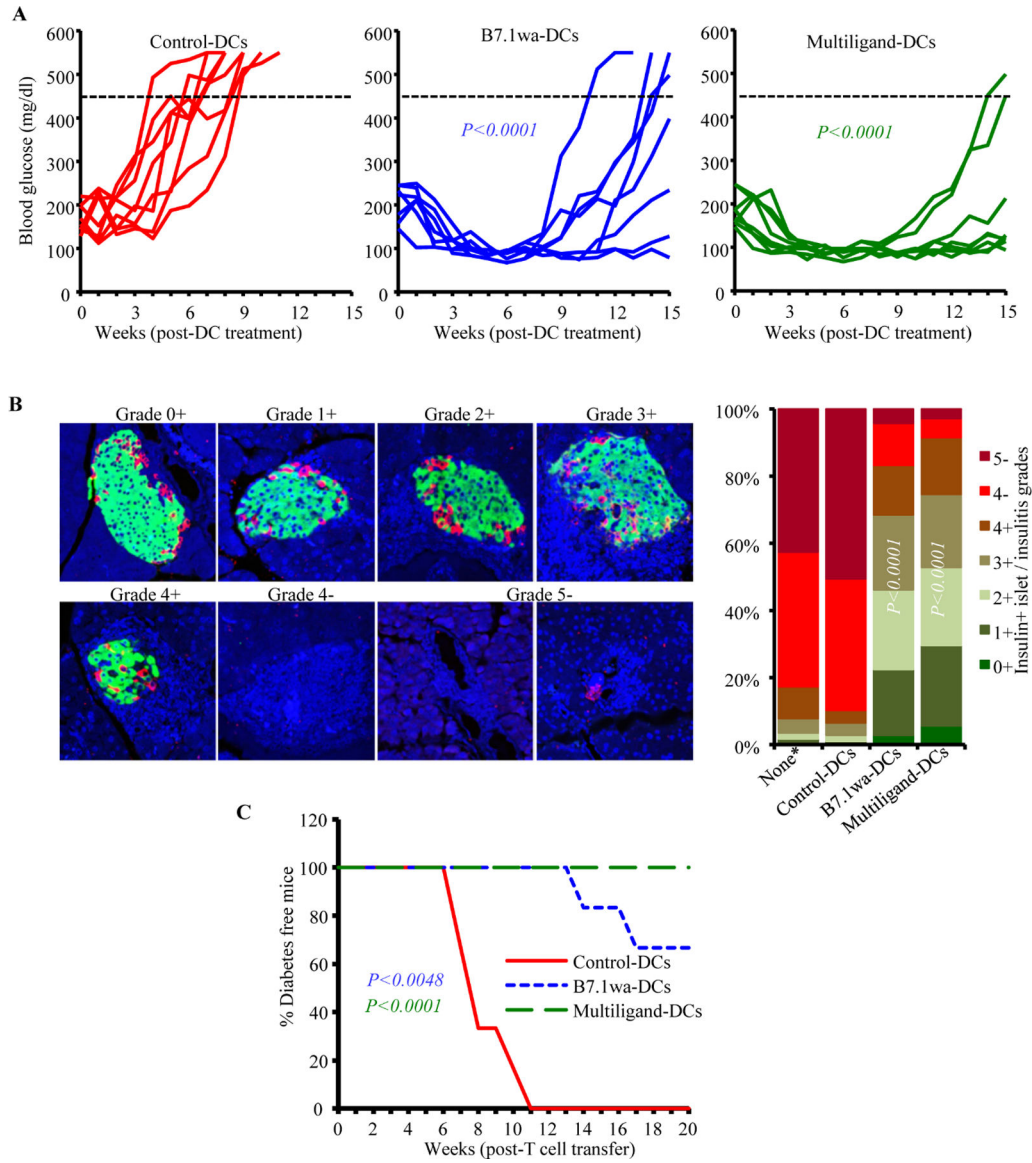
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**Figure 4: B7.1wa-DC and multiligand-DC treated pre-diabetic NOD mice show significant delay in disease onset.**

Ten-week-old pre-diabetic female NOD mice were left untreated (none) or injected i.v. with BcAg-pulsed, engineered-DCs twice at a 15-day interval ( $2 \times 10^6$  cells/mouse/injection). **A)** Cohorts of mice (14 mice/group for treated groups and 20 mice for “none” group) were monitored for hyperglycemia by testing for blood glucose levels every week. This experiment was conducted in 3 small batches of 4–5 DC-treated mice/group and the cumulative results are shown. At least five untreated mice were tested in every batch. **B)** Cohorts of mice (5 mice/group) were euthanized 30 days post-treatment, pancreatic sections were subjected to H&E staining and grading for insulinitis severity. At least 150 islet areas from intermittent sections were examined for each group (at least 30 islet areas/mouse). **C)** Cohorts of mice (8 mice/group) were euthanized 15-days post-treatment, PnLN and spleen cells were stained for Foxp3 and analyzed by FACS. **D)** PnLN cells were also cultured in the presence of BcAg for 72h and supernatants were tested for secreted cytokines by multiplex

assay or ELISA. *P*-value by log-rank test for panel A, Fisher's exact test for panel B, and Mann-Whitney test for panels C and D. All *P*-values are in comparison with control-DC group unless indicated for B7.1wa group vs multiligand-DC group.



**Figure 5: B7.1wa-DC and multiligand-DC treatments at early-hyperglycemic stage results in reversal of hyperglycemia for significant durations.**

Early-hyperglycemic female NOD mice (blood glucose: 140–250 mg/dl) were treated i.v. with control-DCs and B7.1wa-DCs and multiligand-DCs twice as described for Fig. 4. pCDH1-multiligand vector, instead of individual ligand vectors, was used for generating multiligand-DCs in this experiment. **A**) Blood glucose levels were examined every week and glucose values of individual mice are shown. Dotted line represents overt-hyperglycemic (blood glucose: >450 mg/dl). This experiment was carried out in 2 batches of 4 mice/group and the cumulative results are shown. **B**) Cohorts of mice were euthanized 30-days post treatment, and pancreatic sections were subjected to insulin (green), glucagon (red) and DAPI (blue) staining and insulin positive islets and immune cell infiltration, based on DAPI staining, were scored as detailed under materials and method section. \*Pancreatic tissues from untreated mice that were at early-hyperglycemic stage at the time of euthanasia were included as “none” control group to assess the pre-treatment frequency of insulin

positive islets. Pancreatic tissues from at least 5 mice were used for each group. C) PnLN cells from individual mice (6 mice/group), cultured for 24h with BcAg, and injected (i.v.) into NOD-*Rag1*<sup>-/-</sup> mice (about  $1 \times 10^6$  cells/mouse; one recipient/donor) and examined for blood glucose levels every week, for up to 20 weeks. *P*-value by log-rank test with overt hyperglycemia for two consecutive weeks as the end-point for panel A, Fisher's exact test for comparing grades 3+ and 4+ among groups for panel B, and log-rank test for panel C. All *P*-values are in comparison with control-DC group.

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