TRANSLATIONAL SCIENCE

Targeting human plasmacytoid dendritic cells through BDCA2 prevents skin inflammation and fibrosis in a novel xenotransplantation mouse model of scleroderma

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ABSTRACT

Objectives Plasmacytoid dendritic cells (pDC) have been implicated in the pathogenesis of autoimmune diseases, such as scleroderma (SSc). However, this has been derived from indirect evidence using ex vivo human samples or mouse pDC in vivo. We have developed human-specific pDC models to directly identify their role in inflammation and fibrosis, as well as attenuation of pDC function with BDCA2-targeting to determine its therapeutic application.

Methods RNaseq of human pDC with TLR9 agonist ODN2216 and humanised monoclonal BDCA2 antibody, CBS004. Organotypic skin rafts consisting of fibroblasts and keratinocytes were treated with supernatant from TLR9-stimulated pDC and with CBS004. Human pDC were xenotransplanted into NODsebe diabetic/severe combined immunodeficiency (NOD SCID) mice treated with Aldara (inflammatory model), or bleomycin (fibrotic model) with CBS004 or human IgG control. Skin punch biopsies were used to assess gene and protein expression.

Results RNAseq shows TLR9-induced activation of human pDC goes beyond type I interferon (IFN) secretion, which is functionally inactivated by BDCA2-targeting. Consistent with these findings, we show that BDCA2-targeting of pDC can completely suppress in vitro skin IFN-induced response. Most importantly, xenotransplantation of human pDC significantly increased in vivo skin IFN-induced response to TLR agonist and strongly enhanced fibrotic and immune response to bleomycin compared with controls. In these contexts, BDCA2-targeting suppressed human pDC-specific pathological responses.

Conclusions Our data indicate that human pDC play a key role in inflammation and immune-driven skin fibrosis, which can be effectively blocked by BDCA2-targeting, providing direct evidence supporting the development of attenuation of pDC function as a therapeutic application for SSc.

INTRODUCTION

Plasmacytoid dendritic cells (pDC), specialised in the secretion of type I interferon (IFN), activate inflammatory responses through TLR-mediated sensing of nucleic acids released from pathogens during infection or following cell death in autoimmune disease. Self-derived nucleic acids released from damaged tissues, apoptotic/necrotic cells or bound to autoantibodies, can be recognised by TLR7/8/9 and have been shown to induce pDC activation and IFN secretion. The role of CXCL4 has been elucidated as an amplifier of TLR9-mediated pDC hyperactivation and IFN production by organising self-DNA into liquid crystalline immune complexes.

TLR-induced activation of pDC triggers stable cell differentiation into three subtypes, with...
PD-L1(CD274)⁺CD80⁺ (P1) and PD-L1⁺CD80⁻ (P2) subpopulations specialised in type I IFN production both in healthy volunteers (HV) and patients with autoimmune conditions. If pDC have been further implicated in the pathogenesis of autoimmune diseases, such as scleroderma (SSc), systemic lupus erythematosus (SLE) and psoriasis, through their ability to infiltrate the skin and secrete IFNs and proinflammatory chemokines. 

Specifically, SSc is an immune-mediated inflammatory disease (IMID) characterised by vascular and tissue fibrosis, leading to diverse life-altering and life-threatening clinical manifestations. pDC have been observed in affected skin of patients with SSc, and purified peripheral SSc pDC have been shown to spontaneously produce higher levels of type I IFN compared with HV. Indeed, an elevated IFN gene signature in affected organs and in the blood is a common feature of severe disease in SSc, which is present before the onset of clinical fibrosis. These observations collectively support the notion that pDC activation and type I IFN play an important role in SSc pathogenesis.

BDCA2 is a type II transmembrane glycoprotein that belongs to the C-type lectin superfamily receptor that can signal to inhibit pDC type I IFN secretion. BDCA2 signals through an associated transmembrane adaptor, the FcRγ, which recruits the protein tyrosine kinase Syk, inducing protein tyrosine phosphorylation and calcium mobilisation, which reduces TLR-induced activation of pDC, inhibiting type I IFN secretion and other inflammatory mediators. In SLE clinical trials, BDCA2-targeting antibodies induced a significant but partial decrease in IFN response within the blood, and reduced type I IFN-induced response and immune infiltrates in skin lesions.

Recently, pDC’s role in fibrosis was elucidated as elimination of mouse pDC reduced bleomycin-induced skin fibrosis, further highlighting the therapeutic potential of BDCA2-targeting for SSc. However, exploring the efficacy of BDCA2-targeting during fibrosis is difficult, as BDCA2 is only expressed in primates, highlighting the need for a human-specific pDC in vivo model. Furthermore, there are key differences between mouse and human pDC; thus, functions determined in mouse models may not be fully transferable to human pDC.

We developed human-specific models to uncover the role of pDC biology in inflammation and fibrosis, as well as attenuation of pDC function with BDCA2-targeting to determine its therapeutic application for SSc.

**RESULTS**

**TLR9-induced activation of human pDC goes beyond type I IFN secretion and is hindered by BDCA2-targeting**

Using RNAseq, we set out to discover the transcriptome of human pDC when stimulated with TLR9 agonist, A-class oligodeoxynucleotides containing CpG motifs (ODN2216/ODN), to understand the pathways that could contribute to the pathogenesis of chronic inflammation and immune-driven tissue damage seen in IMID. We performed RNAseq analysis of human pDC purified from peripheral blood mononuclear cells (PBMC) from four HV (online supplemental figure S1), as previously described. Transcriptome analysis revealed 328 differentially expressed genes (DEGs ≥2 fold change, FDR≤0.05) between unstimulated (control/CTR) and ODN-stimulated pDC (ODN), with donor heterogeneity observed with ODN stimulation, suggesting pDC response donor variability (figure 1A and online supplemental table 1). Pathway analysis identified genes involved in immune response against viruses and other organisms as key enriched biological processes (figure 1B). These innate immune processes match to those previously identified in characterised inflammatory SSc skin subsets, suggesting involvement of pDC activation in this specific subset. Consistent with this notion, we observed upregulation of many type I IFN-dependent pathways and IFN-related genes, such as IFN-A2, IFN-A21, IFN-B1 and CCL5, a common feature seen in SSc. Pathway analysis also showed JAK/STAT, nuclear factor kappa B subunit 1 (NF-κB) and angiogenesis pathways to be major biological processes upregulated by ODN stimulation (figure 1B), which have been shown to be dysregulated in SSc, but not shown in pDC before. These data suggest that TLR stimulation of pDC can induce a multitude of genes beyond IFN, which could contribute to the pathogenesis of inflammation in SSc and other IMID. Interestingly, a recent publication showed that the majority of SSc skin samples with higher fibroblast scores had significantly increased macrophage and/or dendritic cell scores, suggesting a link between the two cell types that are important for inducing the fibroinflammatory signature.

A monoclonal BDCA2 antibody (clone AC144) has previously been shown to suppress human pDC TLR-induced IFN type I secretion by interfering with the FcRγSyk signalling. To aid our understanding of human pDC in IMID, we generated mouse monoclonal antibodies (mAb) against human BDCA2 and fully humanised the lead mAb, CBS004, which had a greater affinity for BDCA2 compared with AC144 control (online supplemental figure S2A). *Ex vivo* direct competition assays showed that CBS004 and AC144 bind alternative pDC epitopes, as indicated by double staining of the pDC population gated within PBMC (LIN⁻ HLA⁺ CD123⁺ CD304⁺) (online supplemental figure S2B,C).

BDCA2-targeting using CBS004 reduced the expression of 60% of ODN-inducible DEGs ≥1.5-fold (figure 1C and online supplemental table 2). It has been recently shown that TLR-induced pDC triggers stable cell differentiation into three stable subtypes, with PD-L1(CD274)⁺CD80⁻ (P1) and PD-L1(CD274)⁺CD80⁺ (P2) subpopulations specialised in type I IFN production both in HV and patients with autoimmune conditions. Consistent with these observations, ODN induced CD274 expression, which was also suppressed by BDCA2-targeting (figure 1D). To validate the RNAseq findings at protein level, we measured the PD-L1 and CD80 positive subpopulations by fluorescence-activated cell sorting (FACS) (figure 1D,E). Similarly to what has been observed following viral stimulation, ODN induced 62% of pDC differentiation into P1 and P2, with no differences observed between healthy and SSc samples (figure 1D,E).

In this context, BDCA2-targeting caused an increase of the P3 subpopulation, which has been previously shown to produce less IFN. These data were further validated by performing FACS analysis of IFN-positive pDC gated within human PBMC (online supplemental figure S2B). Functionally, ODN led to a dramatic induction of IFN positive pDC, which was reduced by 76% with BDCA2-targeting (figure 1F,G).

Beyond type I IFN signature genes, ODN-induced inflammatory interleukin (IL)-6 expression was also modulated by BDCA2-targeting. Interestingly, IL-6 and IFN secretion can synergistically activate B cells. Among other genes of interest, serglycin (SRGN) also showed a TLR-induced and BDCA2-dependent pattern. SRGN has been shown to be secreted into the extracellular matrix and linked to promoting lymphoid cells adhesion and activation, storage of chemokines and cytokines, as well as being able to induce epithelial–mesenchymal transition. These data demonstrate that TLR stimulation of human pDC goes beyond IFN secretion induction and predicts a greater biological relevance of pDC activation in IMID. Our analyses show that TLR-induced pDC activation can be drastically suppressed by BDCA2-targeting.
**Systemic sclerosis**

Figure 1  TLR9-induced activation of human pDC goes beyond type I IFN secretion and is hindered by BDCA2-targeting *in vitro*. Transcriptome analysis of human pDC cultured in media alone (CTR), with 1 µM ODN2216 (ODN) or with ODN and CBS004 (10 µg/mL) (added 15 min prior to stimulation) for 16 hours (n=4). (A) Heatmap of reduced, centred normalised read counts for DE transcripts among CTR and ODN populations, <5% FDR, calculated using Benjamini-Hochberg multiple testing correction. DE transcripts≥or ≤2-fold (FDR<0.05) shown in online supplemental table 1. (B) Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway analysis showing top biological processes enriched in the set of DE transcripts between CTR and ODN pDC, and their associated p values. (C) Average fold change of DEGs for ODN and ODN+CBS004 relative to CTR (n=4). For repeat transcripts, the highest fold change was used for comparison. Red bars represent the 87 genes that were increased ≥2-fold (FDR<0.05) between CTR and ODN that are dependent on BDCA2 treatment (reduced ≥1.5-fold by CBS004) (full transcript data, online supplemental table 2). (D,E) Validation of CD274 (PD-L1) TLR-induced expression. (D) Example of subtyping FACS analysis of pDC (CD123+CD304+) in the three culture conditions (CTR, ODN and ODN+CBS004). Gating illustrates P1, P2 and P3 subtypes, with the former two previously shown to be IFN-secreting cells.17 PBMC were cultured for 16 hours and pDC sorted as Lin HLA-DR+CD45+CD123+CD304+ (online supplemental figure S2B) and gated for PD-L1 and CD80 expression to determine P1, P2 and P3 sub-types. P1, PD-L1+CD80−; P2, PD-L1+CD80+; P3, PD-L1 CD80+. (E) Quantification of subtypes based on FACS analysis of PBMC samples (healthy and SSc n=3, 4) between ODN and ODN+CBS004 culture conditions. (F,G) Validation of IFN-related gene expression. (F) Representative histogram of intracellular IFN alpha staining of pDC gated within PBMC when cultured in RPMI alone, with ODN (1 µM) and with CBS004 (10 µg/mL). (G) Percentage of IFN-positive pDC from FACS analysis from F (n=6). (E,F) Error bars represent mean±SEM, and statistical significance was evaluated using paired two-tailed t-test. *P<0.05, **P<0.01. CTR, control; DE, differentially expressed; DEG, differentially expressed gene; FACS, fluorescence-activated cell sorting; FDR, false discovery rate; IFN, interferon; PBMC, peripheral blood mononuclear cell; pDC, plasmacytoid dendritic cell; SEM, standard error of mean; SSc, scleroderma.
Human pDC induced IFN response in organotypic skin rafts (OSR), which is inhibited by BDCA2-targeting

Growing evidence shows pDC skin infiltration and induced IFN signature within the skin of patients with IMID, such as SSc. To determine whether TLR9-stimulated pDC could also induce this response, and whether treatment with CBS004 may have a functional effect on target tissue cell activation, we set out to measure IFN-induced genes in an OSR model of keratinocytes and fibroblasts co-culture following exposure to pDC supernatants (figure 2A).

First, we performed IFN secretion ELISA assays on pDC purified from PBMC (online supplemental figure S1) to establish the concentration of IFN within ODN-stimulated pDC supernatants. ODN stimulation was for 16 hours to allow IFN secretion into the supernatant to accumulate, since pDC–IFN maximum production has previously been shown between 12 and 18 hours. To determine whether CBS004 induced a striking increase in IFN-α secretion. The addition of CBS004 15 min prior to ODN stimulation did not induce this secretion, whereas human IgG (HIgG) had no significant effect (figure 2B). To ensure functional inhibition driven by BDCA2-targeting of human pDC was maintained in HV and SSc PBMC, we conducted the same experiment ex vivo. ODN stimulation of PBMC induced a similar but substantial increase in IFN secretion in HV and SSc samples, which was again suppressed by >98% in all samples by BDCA2-targeting (figure 2C). To determine the dose–response of CBS004 and to determine the concentration needed for maximal inhibition of IFN secretion, similar experiments were carried out on ODN-treated HV PBMC, identifying IC50 0.06 nM and IC90 0.86 nM values, with HIgG not effecting IFN production (figure 2D). The inhibitory activity of CBS004 was 17-fold higher than previously tested mAb AC144, which supports the in vitro data showing enhanced BDCA2 affinity and binding (online supplemental figures S3A and S2B). Importantly, BDCA2-targeting did not significantly reduce pDC viability as determined by 7AAD assay gated within HV PBMC (online supplemental figures S2B, S3B).

OSR were treated with supernatants from pDC cultured in cell media alone (CTR), ODN or ODN+CBS004 (figure 2A,B). The volume of supernatant was calculated to produce a final concentration of 6000 pg/ml of IFN in the ODN experiment, as determined via ELISA (figure 2B). Real-time quantitative reverse transcription PCR (qRT-PCR) of 78 key interferon signalling genes (ISGs) was performed on RNA collected from the keratinocyte and fibroblast collagen matrix. ODN supernatants resulted in an increase of 1.8-fold to 32-fold in 35 ISGs relative to CTR (figure 2E). Despite the limited number of models tested (n=3), eight genes showed a statistically significant upregulation, including ISG15, IFITM1, BST2, IFI6, IFIH1, NMI, HLA-B and IFITM3 (induction of 3-fold to 19-fold relative to control and p<0.05). These data suggest the use of OSR to explore the effect of TLR-activated pDC in a preclinical human model. Importantly, ODN+CBS004 supernatants resulted in suppressed upregulation in all of those genes, ranging from 1.8-fold to 11-fold compared with gene expression induced by ODN (figure 2F; ANOVA, p<0.0001). This resulted in a transcription profile similar to CTR (figure 2G). Together these results suggest that BDCA2-targeting of pDC can suppress the IFN signature within skin cells.

Xenotransplant of human pDC in NOD SCID mice increased skin IFN response to TLR stimulation in a BDCA2-dependent manner

To advance our understanding of the role of circulating human pDC in IFN-induced response within the skin, we developed a novel in vivo model. We implemented a xenotransplant transfer of purified normal human primary pDC into nonobese diabetic/severe combined immunodeficiency (NOD SCID) mice (XenoSCID) via intravenous (iv) injection followed by topical application of imiquimod-containing cream (Aldara, TLR7 agonist), with or without anti-BDCA2 (CBS0004) or human IgG (HIgG) (online supplemental figure S4A). Topical imiquimod contains a TLR7 agonist that when applied to resting skin induces expression of type I IFN, primarily in macrophages. Repeated application of topical imiquimod results in induction of ISG in an IFNAR1-dependent manner, the recruitment of leukocytes, skin thickening and the development of an inflammatory lesion. Xenotransplantation of human pDC into this model would allow us to determine pDC role in ISG response to imiquimod and whether it is sensitive to BDCA2-targeting. In vitro imiquimod-stimulation of healthy PBMC induced IFN secretion, which was BDCA2-dependent (online supplemental figure S4B). The purity of pDC isolated from healthy PBMC for in vivo experiments and functional responses to TLR-9 were assessed (online supplemental figures S1, S4C). FACS analysis of CD45+CD123+CD304+ cells indicated pDC skin infiltration within the Aldara+pDC condition (0.3% of total cells, figure 3A), which resulted in a functional increase in mouse skin ISG expression, as mice receiving human pDC induced a 3.2-fold increase in composite ISG score (including Ifi1, Isg15, Ifx1, Cxcl10 and Viperin), compared to CTR (ANOVA P<0.05) (figure 3B). Importantly, this is greater than Aldara treatment alone in absence of pDC, where a 1.7-fold increase in composite ISG score compared to CTR was observed (figure 3B). In agreement, IHC staining showed only a slight increase in MX1 protein expression levels between CTR and Aldara (online supplemental figure S4D). Interestingly, epidermal thickening was observed (online supplemental figure S4D), which has previously been shown to be Aldara-induced response independent of type I IFN and TLR7.

In this context, we could assess the in vivo efficacy of BDCA2-targeting. CBS004 and HIgG control mAb were injected into XenoSCID 12 hours before pDC intravenous injection (online supplemental figure S4A). Aldara-induced pDC skin infiltration, as detected by human CD45+CD123+CD304+ cells in the mouse treated skin, was not hindered by HIgG (0.3% of total cells) but reduced 3-fold by BDCA2-targeting (figure 3C). Most importantly, BDCA2-targeting suppressed 44% of the ISG expression observed in Aldara+pDC, which was significantly reduced compared with HIgG administration (figure 3D).

Consistent with these findings, immunohistochemistry (IHC) analysis showed a strong induction of MX1 and pSTAT1 (Tyr701) with pDC transplantation compared with NOD SCID mice, which was dramatically reduced by BDCA2-targeting and unaffected by HIgG (figure 3E,F).

Xenotransplant of human pDC in NOD SCID mice increased the skin profibrotic response to bleomycin treatment in a BDCA2-dependent manner

Ah Koon et al have shown that depletion of mouse pDC can ameliorate bleomycin-induced skin fibrosis in a mouse model of SSc. While supporting the role of pDC in fibrosis in mice, the model could not directly demonstrate the role of human pDC in this setting. Thus, we developed our XenoSCID model with bleomycin-induced skin fibrosis. We supplemented every other day subcutaneous injection of bleomycin with weekly tail vein injection of human pDC for 3 weeks (online supplemental figure S5A). As anticipated in an immunocompromised mouse, bleomycin alone induced a blunted fibrotic response at 3 weeks,
Figure 2  Human pDC induced IFN response in OSR, which is suppressed by BDCA2-targeting. (A) Systematic outline of OSR protocol; fibroblasts are embedded into a collagen matrix and keratinocytes seeded above until confluence. OSR is brought to ALI to sustain epithelium differentiation. After 5 days, ALI media spiked with 6000 pg/mL of IFN (generated by TLR-stimulated pDC, ODN (B)) for 48 hours. CTR contains equivalent supernatant from untreated pDC (undetectable IFN) and from pDC treated with ODN+CBS004 (10 µg/mL). (B) Quantification of IFN secretion from purified HV pDC (n=7) after 16 hours of culturing in cell media alone (CTR), with ODN (1 µM) and with CBS004 or human IgG1 (10 µg/mL) (added 15 min prior to stimulation) measured by ELISA to determine volume to add to ALI. CTR contains equivalent supernatant from untreated pDC (undetectable IFN) and from pDC treated with ODN+CBS004 (10 µg/mL). (C) CBS004 suppresses TLR-induced IFN secretion in HV and SSc PBMC. PBMC from donors were cultured in media alone (CTR), with 1 µM ODN, or with ODN and CBS004 [10 µg/mL] for 16 hours (n=15). IFN was quantified in the supernatants by ELISA. (B,C) Error bars represent mean±SEM and statistical significance was evaluated using unpaired two-tailed t-test. (D) Percentage of IFN alpha secretion, measured by ELISA, from PBMC from four donors stimulated with ODN in the presence of CBS004 or HIgG (0–66 nM) relative to ODN-stimulated pDC with no antibody (100%). Dotted lines highlight IC50s and IC90 with mean values±SEM. (E,F) RNAs harvested from 3 mm biopsies from OSR and subjected to type I IFN inducible genes superarray. Volcano plots illustrate the fold change of 79 IFN type I-related genes (black dots) between CTR and ODN (E) and between ODN and ODN+CBS004 (F) (pDC supernatant from three different donors for each condition). Grey lines represent the 1.8-fold change cut-off for statistical significance of p=0.05 calculated using Student’s t-test (two-tailed distribution and equal variances between the two samples) on the triplicate 2−ΔCt values for each gene in each treatment group compared with the CTR group. (F) bar chart illustrates the IFN type I-related genes that were >1.8 fold increased in ODN relative CTR and the effect of CBS004. results are represented as means±SEM. *P<0.05, **P<0.001. ALI, air–liquid interface; CTR, control; HIgG, human IgG; HV, healthy volunteer; IFN, interferon; OSR, organotypic skin raft; PBMC, peripheral blood mononuclear cell; pDC, plasmacytoid dendritic cell; SEM, standard error of mean; SSc, scleroderma.
Figure 3  XenoSCID with human pDC increased skin IFN response to TLR stimulation in a BDCA2-dependent manner. Intravenous tail injection of $2.5 \times 10^5$ human purified pDC and intraperitoneal injection of CBS004 mAb (5 mg/kg) or CTR human IgG to NOD SCID mice treated with topical Aldara cream application (online supplemental figure S4A, systematic diagram and timeline), with five different treatment conditions consisting of CTR (n=5), Aldara (n=5), Aldara+pDC (n=5), Aldara+pDC+CBS004 (n=9) and Aldara+pDC+HIgG (n=9). Treated skin was harvested using a 3 mm punch biopsy and processed for FACS analysis of human pDC (CD45+CD123+CD304+) (representative analyses A and C), qRT-PCR analysis for type I IFN inducible genes (B,D), and IHC staining for MX1 and pSTAT1 (Tyr701) (E,F). (B) Composite ISG score within Aldara and Aldara+pDC conditions relative to CTR. Score shows average fold difference between relative expression of Mx1, Isg15, Cxcl10, Ifit1, Isg15 and Viperin between the test condition and CTR. Different symbols represent the different mice litters/human pDC donors. Statistical significance was evaluated using analysis of variance test. (D) Illustration of the composite ISG scores for +CBS004 and +HIgG conditions relative to that of Aldara+pDC. Statistical significance was evaluated using unpaired two-tailed t-test, *P<0.05, **P<0.01. CTR, control; HIgG, human IgG; IHC, immunohistochemistry; ISG, interferon signalling gene; NOD SCID, nonobese diabetic severe combined immunodeficiency; pDC, plasmacytoid dendritic cell; qRT-PCR, real-time quantitative reverse transcription PCR; NOD SCID.
Figure 4  XenoSCID with human pDC increased the pro-fibrotic skin response to bleomycin treatment in a BDCA2-dependent manner. Intravenous tail injection of 2.5×10⁵ human purified pDC and intraperitoneal injection of CBS004 mAb (5 mg/kg) or CTR human IgG into NOD SCID mice treated with Bleo or PBS injections (online supplemental figure S5A); systematic diagram and timeline, with five different treatment conditions consisting of PBS/CTR, Bleo, Bleo+pDC, Bleo+pDC+CBS004 and Bleo+pDC+HiG, each in triplicate. Treated skin was harvested using a 3 mm punch biopsy and processed for H&E (A) and MT staining (B). (C) Epidermis and dermal thickness were measured from 20 areas in each condition. (D) An additional punch biopsy was taken and used to extract protein. Total collagen content measured by Sircol™ assay and shown relative to total protein concentration. Results represented as means±SEM of triplicate experiments. Statistical significance was evaluated using paired two-tailed t-test. (E,F) IHC analysis of MX1 and pSTAT1 (Tyr701) representative images with zoomed in areas, arrows highlight positively stained cells. *P<0.05, **P<0.01, ***P<0.001. Bleo, bleomycin; CTR, control; HiG, human IgG; IHC, immunohistochemistry; MT, Masson trichrome; NS, no significance; NOD SCID, nonobese diabetic severe combined immunodeficiency; PBS, phosphate buffered saline; pDC, plasmacytoid dendritic cell; SEM, standard error of mean.
as shown by partially retained fatty layer, as well as no significant differences observed in skin thickness and collagen content compared with control (figure 4A–D). Furthermore, no significant increase in MX1 protein expression was observed (online supplemental figure S5B,C). Injection of human pDC resulted in bleomycin-induced loss of all subdermal fat, along with increased collagen formation and a 40% increase in overall skin thickness (figure 4A–D). The fibrotic response was associated with type I IFN signalling activation as MX1 and pSTAT1 (Y701) protein expression was increased (figure 4E,F and online supplemental figure S5B,C).

To determine the therapeutic implications of BDCA2-targeting, bleomycin-XenoSCID were treated with intraperitoneal injection of PBS004 or HlgG (online supplemental figure S5A). pDC-induced skin fibrosis was dramatically reduced by BDCA2-targeting as demonstrated by the retention of some fatty layer tissue, similar to bleomycin-only treated mice, a 2-fold reduction in dermal and epidermal thicknesses and 1.5-fold reduction in collagen content compared with HlgG (figure 4A–D). Furthermore, reduction in MX1 and specific pSTAT1 dermal fibroblast (based on morphology) protein expression was observed compared with HlgG treatment (figure 4E,F and online supplemental figure S5B,C).

Overall, our xenotransplant models clearly show that human pDC have a crucial and direct role in skin inflammation and fibrosis and highlight pDC as a viable therapeutic target for SSc.

**DISCUSSION**

In the past decade, substantial evidence has pointed to the involvement of pDC in the pathogenesis of many IMID, including SSc.24–28 Nevertheless, research on pDC has focused only on indirect evidence using *ex vivo* human samples or mouse models. A recent key study supports that mouse pDC have a pathogenic indirect evidence using human samples or mouse models.

Injection of human pDC resulted in bleomycin-induced loss of all subdermal fat, along with increased collagen formation and a 40% increase in overall skin thickness (figure 4A–D). The fibrotic response was associated with type I IFN signalling activation as MX1 and pSTAT1 protein expression was increased (figure 4E,F and online supplemental figure S5B,C).

Our study shows that TLR stimulation of pDC activates a gene expression profile mapping to activation of inflammation, JAK/STAT, NF-κB and angiogenesis pathways, predicting a greater biological relevance of pDC activation in IMID. A time course of ODN stimulation would be beneficial in the future to determine the transcriptome over time and to ensure key gene expression has not been overshadowed, as well as determining the effect of restimulation of previously stimulated pDC.39 Crucially, we have shown that BDCA2-targeting is effective at blocking pDC IFN production, as well as the ODN-induced transcriptome. Furthermore, we show that BDCA2-targeting strongly suppresses the differentiation of IFN-secreting CD274+ pDC17 with a prevalent differentiation towards CD274+CD80+ pDC. Further functional studies will shed light on the effects of BDCA2-targeting on T-cell costimulation, which has been suggested to be affected by CD274+CD80+ pDC.17

Our preclinical organotypic model of human skin allowed us to show target cell activation by TLR-activated pDC and further supports the biological relevance of BDCA2 inhibition. Our XenoSCID model using human pDC greatly expanded this observation. By inducing local TLR-activated skin, we have shown that human pDC can migrate efficiently into Aldara-treated skin and enhance mouse IFN response, as seen in patients with SSc.42–48 Aldara treatment alone increased a small increase in mouse IFN skin response. pDC can be found in mouse blood and lymphoid tissue of NOD SCID mice but are undetectable in skin biopsies either at rest or following imiquimod application for 24 hours,49 indicating our observations are unlikely to be caused by host pDC. Furthermore, we clearly see that human pDC are capable of inducing IFN and fibrotic skin response when introduced into our bleomycin mouse model. While these data directly support a pro-fibrotic effect of pDC in response to bleomycin, they do not directly show that IFN is driving this effect. As shown in our *in vitro* data, pDC produce other cytokines that could plausibly drive the tissue fibrosis in this model. Nevertheless, BDAC2-targeting of pDC in situ prevented the pathogenic responses to proinflammatory and pro-fibrotic stimuli, identifying specific pDC targeting to be a viable therapeutic application for SSc. Our data are supported by similar observations seen when mouse pDC were depleted in a bleomycin-induced fibrotic model of SSc and when specific BDCA2-targeting of resident human pDC in a xenograft Psoriasis model prevented progression into psoriatic skin.50–52

The development of our XenoSCID model is a novel tool that can be used to study the biology of human pDC in mice and can be applied in the research of other IMID affecting the skin, such as psoriasis or SLE. A limitation of this approach is the lack of adaptive immune response in these animals. Therefore, the consequence of pDC inhibition in a competent immune system remains unknown. Nevertheless, the studies from Rowland *et al* showed that in a mouse model of SLE, elimination of pDC strongly impaired expansion and activation of T and B cells.21 In this context, xenotransplant models of human PBMC with and without pDC depletion would be extremely informative although falling beyond the scope of this study.

Together, our data indicate that human pDC, and their cytokine production, are a key cell type in the pathogenesis of SSc. As shown in our *in vitro* and *in vivo* models, BDCA2-targeting of human pDC can reduce ISG response and inflammation, as well as prevent fibrosis. For effective therapeutic application, stratification of patients for those with pDC infiltration and higher IFN score should aid responsiveness to BDCA2 suppression of fibrosis.

**MATERIAL AND METHODS**

Detailed description of experimental methods is available in online supplemental file 1.

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