Brief Definitive Report

Thrombospondin 1 Is an Autocrine Negative Regulator of Human Dendritic Cell Activation

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Abstract

Thrombospondin 1 (TSP) elicits potent antiinflammatory activities in vivo, as evidenced by persistent, multiorgan inflammation in TSP null mice. Herein, we report that DCs represent an abundant source of TSP at steady state and during activation. Human monocyte-derived immature dendritic cells (iDCs) spontaneously produce TSP, which is strongly enhanced by PGE₂ and to a lesser extent by transforming growth factor (TGF)-β, two soluble mediators secreted by macrophages after engulfment of damaged tissues. Shortly after activation via danger signals, DCs transiently produce interleukin (IL) 12 and tumor necrosis factor (TNF)-α, thereby eliciting protective and inflammatory immune responses. Microbial stimuli increase TSP production, which is further enhanced by IL-10 or TGF-β. The endogenous TSP produced during early DC activation negatively regulates IL-12, TNF-α, and IL-10 release through its interactions with CD47 and CD36. After prolonged activation, DCs extinguish their cytokine synthesis and become refractory to subsequent stimulation, thereby favoring the return to steady state. Such “exhausted” DCs continue to release TSP but not IL-10. Disrupting TSP-CD47 interactions during their restimulation restores their cytokine production. We conclude that DC-derived TSP serves as a previously unappreciated negative regulator contributing to arrest of cytokine production, further supporting its fundamental role in vivo in the active resolution of inflammation and maintenance of steady state.

Key words: PGE₂ • TGF-β • IL-12 • CD47 • CD36

Introduction

DCs play a central role in the induction of immunity and tolerance (1). In the absence of inflammation, immature DCs (iDCs) located in peripheral tissues are specialized in uptake of innocuous and cell-associated self Ag. They continuously capture Ag and migrate to the draining lymph node, where they can induce tolerance (1). Under the influence of danger signals (i.e., pathogens and necrotic cells), DCs undergo a process called maturation manifest by up-regulation of costimulatory molecules, secretion of pro- and antiinflammatory cytokines, and the ability to stimulate the differentiation of naive T cells into effector cells. Cytokine production by activated DCs (IL-12, TNF-α, and IL-10) is transient; after prolonged stimulation, DCs become refractory to subsequent activation signals (2). Although delayed production of IL-10 partially contributes to the arrest of cytokine production by activated DCs and the return to steady state, the immune system relies on additional negative feedback mechanisms to down-regulate proinflammatory cytokine release and/or terminate the response to Ag. These include soluble mediators such as PGE₂, cyclopentenone PGs, and lipoxin A4 (3–5), or ligation of certain cell surface receptors (i.e., CD36, CD47, and CD51/CD61; references 6, 7). The latter receptors bind thrombospondin 1 (TSP), an extracellular matrix (ECM) protein predominantly secreted by platelets, monocytes, macrophages, and several nonhematopoietic cell types (8, 9). Also, TSP binds additional receptors as follows: α3β1, αIIbβ3, and cellular glycosaminoglycans. As a consequence of its binding to several cell surface re-
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Trifugation of heparinized blood from healthy volunteers with blood mononuclear cells were isolated by density gradient centrifugation, suggesting an active role for TSP in the resolution of refractoriness of fully mature DCs to additional activation. Interestingly, only TSP–CD47 interactions appear to contribute to activation, and show that TSP production is regulated by soluble mediators and maturation signals. Our results also support the negative regulatory effects of DC-derived TSP on cytokine release (i.e., IL-12, TNF-α, and IL-10) through interactions with CD36 and CD47 during early DC activation. Interestingly, only TSP–CD47 interactions appear to contribute to refractoriness of fully mature DCs to additional activation signals, suggesting an active role for TSP in the resolution of the inflammatory response after an environmental insult.

Materials and Methods

Dendritic Cell Preparation and Culture Conditions. Peripheral blood mononuclear cells were isolated by density gradient centrifugation of heparinized blood from healthy volunteers with lymphoprep (Nycomed). Monocytes were enriched by cold aggregation, followed by T and NK depletion as reported previously (14). Monocyte purity was shown to be >95% CD14+ cells by flow cytometry. Human monocyte-derived iDCs were prepared exactly as described previously (16). Every other day, two thirds of culture medium was replaced by fresh medium (RPMI 1640 10% FCS supplemented with 2 mM glutamine, 1 mM sodium pyruvate, 10 mM Hepes, 100 IU penicillin, and 100 µg/ml streptomycin, containing GM-CSF and IL-4); nonadherent cells were harvested at day 5 to obtain iDCs. Mature DCs (mDCs) were generated after stimulation of iDCs (0.5 × 10^6/ml) in complete HB101 medium (Irvine Scientific) or complete medium (RPMI 1640 10% FCS containing low TSP levels [<30 ng/ml]) for the indicated periods of time. DCs were activated by Staphylococcus aureus Cowan I strain (SAC; 0.0025%) (wt/vol; Pansorbin; Calbiochem), 1 µg/ml LPS or CD40-L L-transfectants (or 1 µg/ml sCD40-L and 500 U/ml IFN-γ). IL-10 and TGF-β were used at 10 ng/ml and purchased from R&D Systems, and PGE2 was used at 100 nM (Sigma–Aldrich). Neutralizing anti-TSP mAbs III (clone 6.7) and I (clone A4.1) obtained from Lab Vision (Clontech) were used at 10 µg/ml. To examine DC exhaustion, mDCs were washed, counted, and restimulated overnight with L-transfectants CD40-L or sCD40-L and IFN-γ in RPMI 1640 10% FCS.

GeneChip Expression Analysis. Human genome–wide gene expression was examined with the Human Genome U133A probe array (GeneChip; Affymetrix, Inc.), which contains the oligonucleotide probe set for ~22,000 full-length genes, according to the manufacturer’s protocol. Total RNA was extracted from ~5 × 10^6 cells. Double-stranded cDNA was synthesized by means of Superscript Choice system (Life Technologies) and a T7-(dt)24 primer (Amersham Biosciences). The cDNA was subjected to the in vitro transcription in the presence of biotinylated nucleoside [Q]triophosphate by means of a BioArray High Yield RNA Transcript labeling kit (Enzo Diagnostics). The biotinylated cRNA was hybridized with a probe. After washing, the hybridized biotinylated cRNA was stained with PE-streptavidin (Molecular Probes) and scanned with an HP Gene Array Scanner (Affymetrix, Inc.). The fluorescence intensity of each probe was quantified with a computer program (Suite 4.0; Affymetrix, Inc.).

Figure 1. Immature and mature DCs express TSP. 0.5 × 10^6/ml monocyte-derived iDCs were cultured for 48 h (A) or the indicated time (B) in the absence or presence of 1 µg/ml LPS or 0.0025% SAC. Cells were permeabilized, fixed, and stained for intracellular TSP as described in Materials and Methods. (A) TSP staining is depicted as follows: iDCs (left, thick line); iDCs + LPS (thick line); and iDCs (right, thin line) versus iDCs + SAC (thin line). (dotted lines) Isotype control mAbs. Data are from three representative experiments out of five (A) and out of five (B).
Flow Cytometric Analysis. The phenotypes of iDCs and mDCs were determined by direct staining with PE-CD40 (clone BU63) and FITC-CD83 (clone HB15e; Becton Dickinson). A three-step procedure was used for intracytoplasmic detection of TSP. In brief, brefeldin A was added for the last 5 h of the culture; DCs were permeabilized with 0.5% saponin and fixed for 5 min with 0.1% paraformaldehyde. Cells were incubated with 2 μg/ml anti-TSP mAb (clone P10; Lab Vision) for 30 min at 4°C, followed by biotinylated goat anti–mouse IgG + IgM (1/500; Biosource International) for 30 min and stained with streptavidin-PE (Becton Dickinson). Stained cells were analyzed with a FACScalibur™ (Becton Dickinson).

Cytokine Measurement. IL-12p70, TNF-α, and IL-10 release were assessed by two-site sandwich ELISA as described previously (7). The sensitivity of the assays was 6 pg/ml for IL-12 and 50 pg/ml for the other cytokines. For TSP measurement, plates were coated overnight with 0.5 μg/ml anti-TSP mAb (clone P10; Chemicon) and blocked with PBS containing 2% BSA and 0.1% Tween 20 for 2 h. After washing, 100-μl samples diluted in HB101 medium were added overnight. TSP purified from platelets (Calbiochem) was used as a standard. Plates were washed and incubated with detection polyclonal rabbit anti–TSP Ab (Calbiochem) and developed by HRP-labeled mouse anti–rabbit IgG and substrate. The sensitivity of the assay was 10 ng/ml.

Results

Immature and Mature DCs Synthesize TSP. TSP produced by macrophages facilitates the phagocytosis of apoptotic cells by establishing a molecular bridge between the phagocyte and the dying cell (17, 18). The iDC is another Ag-presenting cell specialized in the uptake of apoptotic cells, and is the only cell type capable of cell-associated Ag cross-presentation to naive T cells. Therefore, we postulated that iDCs could represent an additional source of TSP. Our initial microarray analysis performed on iDCs isolated from five donors revealed the constitutive presence of TSP transcript and its up-regulation by SAC (11-fold) and LPS (∼8-fold) stimulation. By using two complementary experimental approaches (intracellular staining and ELISA), we demonstrated that TSP is secreted spontaneously by iDCs and that its production is significantly increased during DC maturation. First, we confirmed that DCs express TSP by intracellular staining (Fig. 1). Indeed, the staining reveals that TSP is readily detected in the whole population of unstimulated iDCs. Engagement of toll-like receptors (TLR) 4 or 2, by LPS and SAC respectively, significantly increases TSP expression (Fig. 1 A) with maximum levels observed after 48 h of stimulation (Fig. 1 B, P < 0.05, n = 5). TSP is hardly detectable in nonpermeabilized cell preparations, indicating that the vast majority of TSP is located intracellularly and not on the cell surface (unpublished data). Second, the measurement of TSP in the culture supernatant of unstimulated and maturing DCs confirmed that TSP is indeed secreted by DCs (Fig. 2). TSP production is strongly increased during DC maturation in response to T cell–dependent or TLR-mediated activation signals (Fig. 2 A, P < 0.05, n = 9). The levels of spontaneous TSP production plateau between 48 h and 72 h (Fig. 2 B). Kinetic studies indicate that TSP production by maturing DCs is delayed when compared with that of IL-12 and TNF-α, but simultaneous with that of IL-10 (Fig. 2, B and C). These data establish that DCs are an abundant source of TSP in the steady state and that production can be enhanced by inflammatory signals.

Soluble Mediators Regulate TSP Production by Immature and Activated DCs. TSP potentiates the phagocytosis of apoptotic cells by macrophages, leading to the up-regulation of TGF-β and PGE2 production by these cells (17, 18). Therefore, we examined whether PGE2, TGF-β, or IL-10, each of which exhibits well-established antiinflammatory properties, would in turn modulate TSP production by immature and maturing DCs. As shown in Fig. 3, TGF-β but not IL-10 significantly increases TSP production by iDCs (Fig. 3 A). In contrast, both cytokines both significantly potentiate TSP production after 24 h of stimulation by SAC (Fig. 3 B). This enhancing effect is no longer observed after 48 h, when TSP levels reach a plateau (unpublished data).
Strikingly, PGE_2 increases the release of TSP by iDCs by >50-fold (Fig. 3 C) and 1-nM physiological concentrations of PGE_2 are sufficient to increase TSP production (Fig. 3 E); this enhancing effect is confirmed by intracytoplasmic staining (Fig. 3 F). PGE_2 further potentiates TSP secretion by activated DCs (Fig. 3 C), whereas it significantly down-regulates IL-12 and TNF-α release (Fig. 3 D).

We conclude that antiinflammatory molecules, including PGE_2, IL-10, and TGF-β, provide a positive signal for TSP secretion by both iDCs and activated DCs. Because exogenous TSP reportedly down-modulates DC maturation (7), next we examined the role of endogenous TSP in the regulation of cytokine production.

Endogenous TSP Secretion Renders Fully Mature DCs Refractory to Further Activation. TSP can potentially use two pathways to decrease proinflammatory cytokine release by maturing DCs. Ligation of CD36 and CD47, two TSP receptors on DCs, by their respective mAbs reportedly impairs IL-12 secretion (6, 7). Therefore, we attempted to block TSP binding to CD36 or CD47 at an early stage of DC activation by adding neutralizing anti-TSP mAbs together with a TLR activation signal. We found that anti-TSP mAb III and I, which interfere with the binding of TSP to CD47 and CD36 respectively, enhance TNF-α, IL-12, and IL-10 synthesis in response to SAC or in response to CD40L plus IFN-γ stimulation (Fig. 4 A and not depicted). It is worth noting that these mAbs do not by themselves modulate the expression of costimulatory molecules or affect the ability of DCs to stimulate the proliferation of allogeneic naïve T cells (unpublished data). This indicates that endogenous TSP negatively regulates pro- and antiinflammatory cytokine release during DC maturation.

Next, we examined the possible role of endogenous TSP in DC exhaustion. DCs were cultured for 48 h with SAC (or LPS plus IFN-γ), washed, and restimulated for 24 h with L-CD40L transfectants and IFN-γ (to mimic T cell interactions) in the presence or absence of the two different neutralizing anti-TSP mAbs (Fig. 4 B). As reported previously by Langenkamp et al. (2), DCs have almost extinguished their cytokine release after 48-h maturation in response to secondary stimulation, resulting in the state referred to as exhaustion. Interestingly, we found that exhausted DCs still produced large amounts of TSP (ranging from 1,320 to 3,890 ng/ml; n = 5). Interruption of TSP–CD47 interactions (by anti-TSP mAb III) allows DCs to recover their ability to secrete cytokines. This is not the case with anti-TSP mAb I (i.e., inhibiting binding of TSP to CD36) nor with neutralizing anti-IL-10 mAb (unpublished data). The differential effect of the two anti-TSP mAbs can be partly explained by the loss of CD36 contrast-
The present findings demonstrate that DCs represent a previously unknown and abundant source of TSP, both in the steady state and during activation by danger signals. Additionally, we present evidence that endogenous TSP actively renders DCs refractory to subsequent stimulation and, thus, contributes to the arrest of the inflammatory response. In fact, TSP is rapidly expressed at high levels in inflamed and damaged tissues (20). Resolution of inflammation is now considered to be an active process tightly regulated by stop signals (21), including antiinflammatory molecules such as lipid mediators (lipoxins and prostaglandin derivatives), TGF-β, IL-10, and TSP. Our data are consistent with the known predominant antiinflammatory functions of TSP in vitro and in vivo (10). TSP null mice have persistent inflammation in multiple organs, including the pancreas and the lung, and display a similar phenotype as TGF-β null mice (11, 13). TSP is reported to be a major activator of TGF-β in vivo because a TSP peptide-activating TGF-β reverses the TSP null phenotype toward wild type. Here, we show that TGF-β further increases TSP secretion by DCs. These results are in agreement with recent findings that TGF-β augments TSP expression by macrophages, and TGF-β-treated APCs promote the generation of regulatory T cells (22, 23). This positive feedback loop creates an environment favorable for the induction of anterior chamber-associated immune deviation, a form of toler-
unidentified receptor, possibly CD47, on dying cells (17). In that context, we postulate that iDC-derived TSP is likely to be involved in phagocytosis of damaged tissues by steady-state DCs that continuously tolerate lymph node T cells. Furthermore, TSP inhibits early T cell activation and promotes naive T cell anergy (28, 29), and active TGF-β displays potent immunoregulatory functions, including prevention of DC maturation and activation of regulatory T cells (15, 30). Altogether, it is possible that PGE₂ and TGF-β further amplify a TSP/TGF-β/TSP-positive immunoregulatory loop established to ensure tolerance in the steady state. Consistent with numerous analyses, we observed that exogenous PGE₂ strongly inhibits cytokine release by activated DCs (3). Under these inflammatory conditions, PGE₂, like TGF-β and IL-10, markedly increases TSP.

Herein, we provide evidence that DC-derived TSP mediates DC exhaustion. After prolonged stimulation, DCs secrete TSP and TGF-β and promote the generation of nonpolarized T cells, a subset of central memory T cells (2). We propose that TSP exerts its antiinflammatory activity both in the steady state, contributing to the tolerogenic function of DCs, and at the end of the inflammatory response, allowing the rapid and active resolution of inflammation. The in vivo biological relevance of the present in vitro findings is supported strongly by the inflammatory phenotype of TSP null mice.

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