CD300c is an Activating Receptor Expressed on Human Monocytes

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Key Words
CD300 · CD300a · CD300c · Cell surface receptors · Paired receptors

Abstract
Human CD300 molecules comprise a family of receptors that regulate many immune cell processes. They are mostly expressed on myeloid cells, although expression of two members, CD300a and CD300c, has also been described on lymphocytes. However, due to the lack of specific antibodies that distinguish between these two receptors, it has been difficult to determine the expression pattern and function of CD300a and CD300c in primary cells. Here, we have identified a specific monoclonal antibody, clone TX45, that recognizes only CD300c and show that within freshly isolated blood leukocytes, monocytes are the only cells that express CD300c on the cell surface. In vitro differentiation experiments revealed that CD300c is differentially expressed on different monocyte-derived cells, including macrophages and dendritic cells. Furthermore, TLR ligands LPS and flagellin dynamically regulate the expression of CD300c. Cross-linking of this receptor with clone TX45 monoclonal antibody induced calcium mobilization, upregulation of the co-stimulatory molecule CD86 and the production of inflammatory cytokines. Importantly, LPS-mediated production of inflammatory cytokines by monocytes was further enhanced if CD300c was simultaneously engaged by the agonist antibody. Altogether, our results show that human CD300c is an activating receptor expressed on monocytes and that it has a potential role in inflammatory responses.

Introduction

Myeloid cells are main sentinels of the immune system that have a prominent role in the host defense by sensing and monitoring foreign antigens [1, 2]. Many of the myeloid functions are regulated through a set of cell surface molecules belonging to multigenic families that include both activating and inhibitory receptors, which in the end establish and maintain a delicate balance aimed to eliminate the foreign antigen while preserving self [3, 4]. In general, the activating receptors of these multigenic families have charged residues in the transmembrane region and associate with adaptor molecules containing ITAM (immunoreceptor tyrosine-based activating motif) or phosphatidylinositol 3-kinases binding motif YxxM. After receptor engagement, ITAM and YxxM motifs become tyrosine phosphorylated and recruit protein kinases, initiating a downstream activation signaling cascade that regulates many functions such as cytokine secretion, calcium mobilization, differentiation, proliferation, phagocytosis, cytotoxicity and cell survival [5, 6]. On the contrary, the inhibitory receptors have one or more ITIM...
The inhibitory receptors include the surface expression of CD300a and CD300c monoclonal antibodies that could differentiate between difficult to discriminate. This is due to the lack of specific the cell surface expression of these receptors has been difficult. Both lymphoid and myeloid cell lineages encoding both CD300a and CD300c are widely found in human cells. Its function on primary cells has not been elucidated so far. Although human CD300a has a long cytoplasmic tail with three classical ITIMs and one nonclassical ITIM that, after phosphorylation, associates with SHP-1, SHP-2 and SHIP phosphatases and elicits a signal that is capable of inhibiting the functions of natural killer (NK) cells, mast cells, B and T lymphocytes, macrophages, eosinophils, basophils, neutrophils and dendritic cells (DC) [9, 14, 22–28]. In contrast, CD300c has a short cytoplasmic tail, a negatively charged amino acid residue in the transmembrane region and the potential to associate with the adaptor molecules FcεRIγ and DAP12 [17, 18]. Although human CD300c was the first molecule to be identified in this family of receptors [29], its function on primary cells has not been elucidated so far. Ligation of ectopically expressed human CD300c induces activating signals in the mast cell line RBL-2H3, partially through its direct association with FcεRIγ [18]. Nonetheless, its role in human cells that endogenously express the receptor is not known. Although transcripts encoding both CD300a and CD300c are widely found in both lymphoid and myeloid cell lineages [22, 26, 29, 30], the cell surface expression of these receptors has been difficult to discriminate. This is due to the lack of specific monoclonal antibodies that could differentiate between the surface expression of CD300a and CD300c [22, 26, 28, 30]. In this study, using an anti-CD300c-specific antibody, we describe the cell surface expression of CD300c in peripheral blood monocytes and its ability to induce activating signals. Contrary to CD300a, we show that CD300c is not expressed on the surface of the vast majority of freshly isolated lymphocytes or granulocytes. In addition, we also show that CD300c is regulated on monocytes after treatment with the TLR4 and TLR5 ligands LPS and flagellin, respectively, and that it is differentially expressed on monocyte-derived macrophages and monocyte-derived DC (mDC). Furthermore, our data also suggest that CD300c has a costimulatory effect in cytokine secretion on LPS-stimulated monocytes. Altogether, our results support the notion that CD300c is an activating receptor in human monocytes and that it may have a significant role in regulating inflammatory responses.

Materials and Methods

Study Population

Whole-blood anduffy coats were collected under an institutional review board-approved protocol at the National Institutes of Health Blood Bank from healthy donors. All study subjects provided written informed consent.

Constructs and 293T and YTS Cell Transfectants

RNA was isolated from freshly isolated human NK cells using the RNAqueous-4PCR kit (Ambion-Life Technologies) and transcribed into cDNA using qScript cDNA synthesis kit (Quanta Biosciences). The full-length cDNAs corresponding to CD300a and CD300c were amplified using specific primers and the products were cloned into pcDNA3.1 (+) expression vector (Invitrogen) using standard molecular biology techniques. The constructs were sequenced to confirm their identity. Transient transfection of the constructs into 293T cells, a human fibroblast kidney cell line, was performed using Lipofectamine 2000 according to the manufacturer’s protocol (Invitrogen). YTS stable transfectants were obtained initially by electroporation, then sorted for the positive cells and finally selected in culture medium containing 1 μg/ml of gentamicin (InvivoGen).

Isolation of Peripheral Blood Nucleated Cells, Monocytes and Generation of Monocyte-Derived Cells

The human nucleated cell fraction was isolated from whole blood with HetaSep (Stem Cell Technologies) using either the gravity sedimentation or the centrifugation method according to the manufacturer’s protocol. Enriched human monocytes were obtained from buffy coats of healthy donors using RosetteSep (Stem Cell Technologies), a negative selection method that depletes the unwanted cells with tetrameric antibody complexes recognizing CD2, CD3, CD8, CD19, CD56, CD66b, CD123 and glycophorin A. The purity of monocytes was more than 70%. Macrophages and immature mDC were differentiated from enriched monocytes in differentiation media in the presence of recombinant human GM-CSF (50 ng/ml) and recombinant human GM-CSF (50 ng/ml) plus recombinant human IL-4 (20 ng/ml), respectively, for 6 days. The composition of differentiation media was Iscove’s modified Dulbecco’s medium (plus GlutaMax) sup-

DOI: 10.1159/000350523

J Inattne Immun 2013:5:389–400

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plemented with 10% human AB serum, 2 mM glutamine, 1% sodium pyruvate and 1% non-essential amino acids (Lonza). The medium was replaced 2–3 times during the differentiation period. After 6 days of culture, the immature mDC were further cultured in the presence of 1 ng/ml LPS (InvivoGen) for 48 h to generate mature mDC.

Flow-Cytometric Analysis

The following mouse anti-human antibodies were used for flow-cytometric analysis: purified and PE anti-CD300a/c IRp60 (clone E59.126) from Beckman Coulter; purified anti-CD300a/c (clone TX49) was a generous gift from Dr. Akira Shibuya; eFluor 660 anti-CD300c (clone TX45), PE-Cy7 anti-CD19 (clone H1B19), AF700 anti-CD3 (UCHT1) and PE-Cy7 anti-CD14 (clone 61D3) from eBioscience; FITC anti-CD66b (clone G10F5), purified and PE anti-CD300c (clone TX45) from Biologend; PE anti-CD80 (clone L307.4), FITC anti-CD86 (clone 2331 FUN-1), PE anti-CD83 (clone HB15e), PE anti-CD206 (clone 19.2), APC anti-CD209 (clone DCN46) and APC anti-CD56 (clone NCAM16.2) from BD Biosciences. Purified and fluorescein conjugated MOPC-21 (mouse IgG1; from eBioscience) was used as an isotype-matched control. Secondary antibodies were purchased from Jackson ImmunoResearch. YTS cells stably expressing CD300a or CD300c were washed with staining buffer (1% bovine serum albumin at 5 × 10⁶ cells/ml). Next, cells were labeled with 8.5 μg goat anti-mouse IgG F(ab′)₃, followed by the addition of 1 μg fluorochrome-conjugated anti-CD300a/c (clone E59.126) or anti-CD300c (clone TX45) monclonal antibody for 20 min on ice. Then, cells were washed thoroughly and acquired in a FACSCalibur flow cytometer (BD Biosciences). Regarding 293T cells transiently transfected with CD300a or CD300c encoding constructs, the cells were harvested after 16–20 h of transfection, washed with staining buffer and incubated with fluorescein conjugated anti-CD300a/c (clone E59.126) or anti-CD300c (clone TX45) monoclonal antibody for 30 min on ice. The 293T cells were also stained with purified anti-CD300a/c (clone TX49) for 40 min, washed and incubated with secondary PE conjugated anti-mouse Fc antibody for another 20 min on ice. Then, cells were washed thoroughly and acquired in a FACSCalibur flow cytometer. For the flow-cytometric experiments with human primary cells, the staining buffer contains 1% human AB serum to block the Fc receptors. Before harvest from the plates, monocytes and monocyte-derived macrophages were incubated with Accutase (Innovative Cell Technologies) for 15–20 min at room temperature to detach the cells. After harvesting, cells were washed in staining buffer and incubated with the respective fluorescein conjugated antibodies for 30 min on ice. After extensive washing, the cells were then acquired using either FACSCalibur or LSRII (BD Biosciences) flow cytometers. Flow-cytometric data were analyzed using FlowJo software (Tree Star).

Stimulation of Monocytes with LPS and Flagellin

Enriched monocytes (10⁶/ml) from healthy donors were either stimulated with LPS (1 ng/ml) or flagellin (100 ng/ml) in a 24-well plate for different time periods as mentioned in the text and figure legends. Then, cells were harvested and analyzed for CD300c surface expression by flow-cytometric analysis. In addition, cell samples for RNA isolation were stored at 4°C in RNAlater (Ambion-Life Technologies). Real-time quantitative PCR assays were performed on a CFX-96 real-time PCR detection system (Bio-Rad) using 480 SYBR Green I master supermix (Quanta Biosciences) according to the manufacturer’s protocol. Primers for real-time PCR measurement of human CD300c (PPH07153A) and β-actin (PH00073G) were purchased from SA Biosciences. All reactions were made in triplicate, and averages were used to calculate the relative levels of mRNA. Relative quantification of CD300c mRNA was determined using the second derivative maximum using the CFX-Manager software (Bio-Rad), and data were normalized to the housekeeping gene β-actin.

Cross-Linking of CD300c on Monocytes and Measurement of Cytokine Production

Tissue culture plates (24 wells) were coated with 10 μg/ml of either purified anti-human CD300c monoclonal antibody (clone TX45) or the isotype control (MOPC-21) for 2–3 h at 37°C and then extensively washed with PBS to remove the unbound antibody. Enriched monocytes (10⁶/ml) obtained from healthy donors were then added to the antibody-coated plates in RPMI 1640 medium supplemented with 10% FBS, 100 U/ml penicillin-streptomycin, 2 mM glutamine and 10 mM HEPES (Lonza). In the co-stimulation experiments, along with the plate-bound antibody activation, monocytes were also stimulated with the addition of LPS (1 ng/ml) to the culture. After 24 h of stimulation, the cell culture supernatants were harvested and stored at −80°C, and the cells were harvested and analyzed by flow cytometry for the expression of costimulatory molecules.

The cell culture supernatants were thawed on ice and analyzed for cytokine production using the human inflammatory cytokine kit by BD Biosciences cytometric bead arrays according to the manufacturer’s protocol. Samples were acquired on a BD LSRII flow cytometer (BD Biosciences) and analyzed using BD FCAR array software.

Calcium Mobilization Assay

Freshly isolated monocytes were washed and resuspended in Dulbecco’s PBS (GIBCO catalog No. 14287) containing 1% bovine serum albumin at 5 × 10⁶ cells/ml. Next, cells were labeled with Fura Red (5 μg/ml) and Fluo-4 (2 μg/ml), from Invitrogen, for 30 min at 37°C, then extensively washed with PBS to remove the unbound antibodies. Enriched monocytes (10⁶/ml) to the culture. After 24 h of stimulation, the cell culture supernatants were harvested and stored at −80°C, and the cells were harvested and analyzed by flow cytometry for the expression of costimulatory molecules.

Statistical Analysis

Data were analyzed using GraphPad Prism software. The data were plotted as bar graphs, and pairwise comparisons were examined by a paired Student’s t test. * p < 0.05, ** p < 0.01.

Results

Specific Recognition of Human CD300c by the Monoclonal Antibody Clone TX45

Human CD300a and CD300c mRNA transcripts are broadly detected in cells from both the myeloid and lymphoid lineages [22, 26, 29, 30]. However, the cell surface
expression of these two highly homologous receptors has been difficult to determine because of the lack of specific antibodies that are able to discriminate between CD300a and CD300c. The majority of available antibodies show cross-reactivity and recognize both receptors on the cell surface [22, 28, 30]. Thus, it has been a very difficult, if not an impossible task to investigate and understand the specific role of these two molecules in primary cells.

As a first step, we wanted to examine if there is a monoclonal antibody that binds specifically to cell surface CD300c but not to CD300a. Plasmids encoding full-length CD300a and CD300c were generated and transfected into 293T and YTS cells. After testing several antibodies, we found that clone TX45, a commercially available monoclonal antibody, did not bind to 293T cells transfected with either empty vector control or with a construct encoding CD300a. However, clone TX45 showed a strong binding to CD300c transfected cells (fig. 1a). On the contrary, the monoclonal antibodies clones E59.126 and TX49 did recognize transfectants expressing both CD300a and CD300c, indicating that these two antibodies are not specific (fig. 1a). In addition, we wanted to test the specificity of clone TX45 on lymphoid cells expressing CD300a and CD300c. To perform that, we generated CD300a or CD300c stable transfectants of YTS, an NK cell tumor cell line. Similar to the results mentioned with the 293T cells, clone TX45 bound only to CD300c-expressing YTS cells and clone E59.126 bound to both CD300a and CD300c transfected YTS cells. As expected, none of these antibodies showed any binding to the empty vector transfected

**Fig. 1.** Monoclonal antibody clone TX45 binds specifically to CD300c. **a** 293T cells were transiently transfected with empty vector (upper panels), vectors expressing full-length CD300c (middle panels) or CD300a (lower panels). After 16 h of transfection, cells were stained with the following monoclonal antibodies: anti-CD300c, clone TX45 (left lane); anti-CD300a/c, clone E59.126 (middle lane), and anti-CD300a/c, clone TX49 (right lane). Shaded histograms represent unstained cells and the open histograms represent the staining with the specific antibodies. These data are representative of 3 independent experiments. **b** YTS cells stably transfected with empty vector (upper panel), vector expressing CD300c (middle panel) and CD300a (lower panel) were stained with anti-CD300c, clone TX45 (left lane) and anti-CD300a/c, clone E59.126 (middle lane). The shaded histograms represent the isotype-matched controls and the open histograms the staining with the specific monoclonal antibodies. Results are representative of 3 independent experiments.
YTS cells (fig. 1b). Thus, despite the very high homology between the extracellular domains of CD300a and CD300c, we show that the anti-CD300c monoclonal antibody, i.e. clone TX45, binds specifically to human CD300c and does not cross react with CD300a.

**CD300c is Expressed on the Surface of Monocytes and Monocyte-Derived Cells**

The identification of a specific monoclonal antibody against CD300c, clone TX45, was of great interest, and therefore we decided to study the cell surface expression of this receptor on cells of both lymphoid and myeloid lineages. The nucleated cell fraction was isolated from the whole blood of healthy donors and the binding of the monoclonal antibodies anti-CD300a/c, clone E59.126, and anti-CD300c, clone TX45, to different cell subsets was tested. We confirmed previously published results [23, 27, 28, 31, 32] and showed that clone E59.126 binds to all NK cells (CD3− CD56+), monocytes (CD14+), granulocytes (CD66b+) and subsets of T (CD3+) and B cells (CD19+; fig. 2a). Intriguingly, the specific monoclonal antibody against CD300c, clone TX45, binds almost exclusively to the CD14+ population and not to any other cell type (fig. 2a). In some donors, a very small subset of T cells, less than 2%, bound clone TX45 (data not shown). These results indicate that CD300a and CD300c are differentially expressed. In freshly isolated peripheral blood cells, CD300a is expressed on the cell surface of lymphocytes, monocytes and granulocytes, while CD300c is almost exclusively expressed on monocytes.

We then enriched monocytes from the peripheral blood using negative selection and in vitro generated mDC and macrophages. Phenotypic analysis confirmed the purity of the different cell types: monocytes (CD14+), macrophages (CD14high CD206++ CD83− CD80− CD86+), immature mDC (CD14low CD209+++ CD80+ CD86+ CD83−) and mature mDC (CD14low CD209++ CD80+++ CD86+++ CD83+; data not shown). Our data showed that monocyte-derived macrophages are positive for CD300c, with similar expression levels to those observed on peripheral blood monocytes. On the contrary, we observed that the cell surface expression of CD300c was reduced on immature and LPS-induced mature mDC relative to monocytes and macrophages (fig. 2b). Additionally, the anti-CD300a/c (clone E59.126) monoclonal antibody binds to all monocyte-derived cells without any change in the intensity of cell surface expression (data not shown). Thus, the surface expression of CD300c is exclusive to monocytes and monocyte-derived cells. Since it has been reported that CD300c transcripts are found in other cell types, further studies are needed to understand the mechanism regulating the cell surface expression of this receptor.

**Expression of CD300c is Regulated on Monocytes upon LPS Treatment**

To further investigate how CD300c expression is regulated, we treated freshly isolated enriched monocytes with the TLR4 ligand LPS. Gene expression analysis showed that after 2-hour treatment with LPS, the levels of CD300c transcripts were significantly downregulated, while after 24 h of treatment the amount of CD300c transcripts increased to the levels expressed by nonstimulated monocytes (fig. 3a). On the other hand, although the cell surface expression of CD300c decreased after LPS treatment for 2 h, in line with the amount of CD300c transcripts, after 24 h of LPS stimulation CD300c expression showed a significant increase when compared with untreated monocytes (fig. 3b). Similar results were observed when monocytes were treated with the TLR5 ligand flagellin (online suppl. fig. 1; for all online suppl. material, see www.karger.com/doi/10.1159/000359523). Therefore, our results suggest that the regulation of CD300c expression on monocytes is a highly dynamic process, probably involving both transcriptional and posttranscriptional mechanisms.

**Triggering of CD300c Induces Calcium Mobilization and Upregulation of CD86**

To characterize the function of CD300c in circulating monocytes, we studied whether the anti-CD300c-specific monoclonal antibody clone TX45 is able to induce activation signals in monocytes. We first investigated its ability to induce intracellular calcium mobilization. Engagement of CD300c with soluble anti-CD300c monoclonal antibody followed by cross-linking with anti-mouse IgG F(ab′)2 induced a transient and rapid increase in intracellular calcium (fig. 4a), which was not observed when cells were stimulated under the same conditions with the isotype-matched control (clone MOPC-21). Engagement of CD300c only with soluble anti-CD300c monoclonal antibodies, in the absence of cross-linking with anti-mouse IgG F(ab′)2, did not induce calcium mobilization (data not shown). To further prove that CD300c is an activating receptor, we stimulated freshly isolated monocytes with plate-bound anti-CD300c monoclonal antibody. After 24 h of culture, monocytes were able to significantly upregulate the cell surface expression of the costimulatory molecule CD86, while monocytes cultured on plates coated with the isotype-matched control did not show an increase in the expression of CD86 (fig. 4b). Interestingly,
in our experimental settings, no change was found in the cell surface expression of CD80 after stimulation of monocytes with plate-bound anti-CD300c monoclonal antibody (data not shown). Thus, our data support the notion that CD300c is a functional activating receptor in monocytes.

**Cross-Linking of CD300c Upregulates Proinflammatory Cytokine Secretion from LPS-Treated Monocytes**

As monocytes are very important players in the control of inflammatory processes, we investigated if CD300c engagement has a role in the production of pro-

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**Fig. 2.** CD300c is expressed on human monocytes and monocyte-derived cells. **a** Cells were isolated from normal healthy donors and stained with specific antibodies to distinguish different cell subsets. In addition, cells were stained with anti-CD300a/c (clone E59.126) and anti-CD300c (clone TX45) monoclonal antibodies. Shaded histograms represent the staining with isotype-matched controls and empty histograms the staining with anti-CD300a/c (clone E59.126; upper panel) and anti-CD300c (clone TX45; lower panel). Data from a representative out of 6 are shown. **b** Enriched monocytes were differentiated into mDC and macrophages according to the protocol described in the Materials and Methods. The expression of CD300c was determined using anti-CD300c (clone TX45) monoclonal antibody. The shaded histograms represent the staining with isotype-matched control and the open histograms correspond to the specific staining with anti-CD300c (clone TX45). Results from a representative out of 4 are shown.
inflammatory cytokines. Freshly isolated monocytes were cultured either with plate-bound anti-CD300c monoclonal antibody (clone TX45) or with isotype-matched control (clone MOPC-21). Cross-linking of CD300c induced significant production of the inflammatory cytokines TNF-α and IL-8, while there was not a significant increase in IL-6 and IL-1β secretion compared with monocytes cultured with plate-bound isotype-matched control (fig. 5). TLRs are crucial in the recognition of pathogens, and their engagement on monocytes activates the secretion of pro- and anti-inflammatory cytokines [33]. We sought to determine if cross-linking of CD300c is able to modulate TLR-mediated cytokine production. Our results showed that CD300c engagement significantly increased LPS-mediated TNF-α, IL-1β, IL-6 and IL-8 secretion (fig. 5). In our experimental conditions, no IL-12 was detected (data not shown). Intriguingly, the LPS-mediated secretion of the anti-inflammatory cytokine IL-10 did not increase when CD300c was cross-linked (fig. 5). Altogether, these data further point out that CD300c is a novel activating receptor expressed on human monocytes and that is able to cooperate with TLR4 in the production of inflammatory cytokines.

Fig. 3. Expression of CD300c is regulated on monocytes by LPS. a Enriched monocytes from healthy donors were either left untreated (Alone) or treated with LPS for 2 and 24 h. RNA was extracted and the levels of CD300c mRNA were determined by real-time PCR. Graph bars represent the average ± SEM. Data are from 5 independent experiments. b Enriched monocytes were either left untreated or treated with LPS for 2 and 24 h. The cells were harvested and checked for CD300c cell surface expression. The shaded histograms represent the staining with the isotype control (MOPC-21) and the open histograms represent the staining with anti-CD300c (clone TX45) monoclonal antibody. The numbers indicate the values of median fluorescence intensity. Results are representative of 5 healthy donors.
Discussion

Monocytes have a very important role in both innate and adaptive immune responses, and their tasks are in part controlled and regulated by cell surface receptors that are able to deliver activating or inhibitory signals. The CD300 family of molecules regulates a large range of immune cell processes via their paired activating and inhibitory receptors [9, 20]. Among these molecules, it has been shown that the activating members, CD300b and CD300e, and the inhibitory receptors, CD300a and CD300f, have a prominent role in regulating functions of human primary monocytes and monocytic cell lines [15, 16, 19, 34, 35]. In this article, we show that CD300c, another member of the CD300 family of receptors, is only expressed on monocytes within blood leukocytes. Furthermore, cross-linking of CD300c with an agonist antibody led to the activation of monocytes, as shown by mobilization of intracellular calcium, upregulation of CD86 and production of cytokines. We also show that the expression of CD300c on primary monocytes is regulated upon stimulation with TLR4 and TLR5 ligands, and that it has a costimulatory effect with LPS in the secretion of proinflammatory cytokines. Altogether, our results indicate that CD300c is an activating receptor expressed by human monocytes that may have a very important role during inflammation.

Elucidating the expression and function of human CD300c in primary cells has been hampered by the unavailability of specific monoclonal antibodies. CD300a and CD300c are paired receptors with more than 80% homology in the amino acid sequence of the extracellular domain [22], and the majority of available monoclonal
antibodies recognize both CD300a and CD300c [22, 26, 28, 30]. Here, we have analyzed the specificity of the monoclonal antibody clone TX45 and show that it binds to cell surface CD300c but not CD300a. The availability of the specific anti-CD300c monoclonal antibody clone TX45 clarifies, at least in part, the issue of the cell surface expression of CD300a and CD300c on human immune cells. Although previous reports have described that CD300a and CD300c transcripts are expressed on lymphoid and myeloid cells, the cell surface expression of both receptors was not clarified. Our data demonstrate that CD300c is expressed exclusively on the surface of human monocytes, and also on other monocyte-derived cells like macrophages and DC. On the other hand, the

Fig. 5. Cross-linking of CD300c in monocytes induces the secretion of inflammatory cytokines. Freshly isolated monocytes from healthy donors were either stimulated with plate-bound isotype-matched control antibody, MOPC-21 (empty bars), or with anti-CD300c antibody, TX45 (black bars), in the absence (Untreated) or presence of LPS for 24 h. Culture supernatants were harvested and tested for the secretion of human inflammatory cytokines using flow-cytometric bead analysis. The values on the y-axis correspond to the concentrations of cytokines: TNF-α, IL-1β and IL-10 (pg/ml), and IL-8 and IL-6 (ng/ml). Graph bars represent the average ± SEM. Data are from 5 independent experiments.
binding of monoclonal antibodies that recognize both CD300a and CD300c, including clones E59.126 and TX49, to other cell types, such as T, B and NK cells and granulocytes [22, 23, 27, 28, 30–32], suggest that what those antibodies are recognizing on these cells is probably CD300a but not CD300c. The latter has been suspected in some way because cross-linking experiments with anti-CD300a/c monoclonal antibodies have been shown to deliver inhibitory signals in several cell types [9, 23, 27, 28, 31, 34].

The presence of CD300c transcripts in other cell types, in addition to monocytes, raises the question of what mechanisms control the cell surface expression of this receptor. It is important to note that stimulation of cultured NK clones and of cultured primary CD4+ cells with anti-CD300a/c monoclonal antibodies do not always induce a negative signal [22, 28], suggesting the possibility that in these circumstances, the monoclonal antibodies may also be recognizing CD300c. Clearly, further investigations are required in this regard to determine if cell culture and activation induces cell surface expression of CD300c. In our study, we report that both transcripts and cell surface expression of CD300c on monocytes were downregulated with LPS or flagellin treatment after 2 h. On the other hand, the cell surface expression is significantly upregulated after 24 h of treatment, without any apparent increase in mRNA levels compared with nonstimulated monocytes. These results suggest that the regulation of CD300c surface expression on monocytes is regulated both at transcriptional and posttranscriptional level. The expression of other CD300 molecules has been shown to be regulated by posttranscriptional mechanisms. For example, mouse CD300b expression on neutrophils is regulated by the action of matrix metalloproteinases, and TLR4 stimulation increases the release of CD300b from the cell surface [36]. It has also been described that stimulation of human neutrophils with LPS and GM-CSF causes a rapid translocation of an intracellular pool of CD300a to the cell surface [23]. Our preliminary data suggest that matrix metalloproteinases may also have a role in the downregulation of CD300c from the surface of monocytes after 2 h of stimulation with LPS or flagellin (data not shown).

Monocytes are recruited and activated at the sites of inflammation, secrete cytokines and upregulate the expression of costimulatory molecules that are important for optimal T-cell responses [37]. Similar to CD300e [16], we have also shown that cross-linking of CD300c with agonist antibodies induces mobilization of intracellular calcium, cytokine production and upregulation of CD86 expression on human monocytes. On the other hand, we could not see an upregulation of CD80 expression in our experimental settings. This could imply that CD300c is a bona fide activating receptor, since it has been proposed that CD80 and CD86 have different functional identities, namely that CD86 is the main ligand for CD28, a T-cell costimulatory molecule, and CD80 is the major ligand for CD152, a T-cell inhibitory receptor [38]. Clearly, more studies are required to address this issue.

Similar to ligation of CD300e, engagement of CD300c with antibodies triggered the secretion of TNF-α and IL-8 by monocytes. However, as opposed to CD300e [16], the cross-linking of CD300c further enhanced LPS-mediated inflammatory cytokine production, while it did not increase the secretion of IL-10, emphasizing the proinflammatory role of CD300c. Nonetheless, it is very important to note that despite the fact that agonist monoclonal antibodies are helpful tools to functionally characterize cell surface receptors, data should be carefully interpreted for comparative analysis between different molecules, unless they are validated with their natural ligands. Recently, it has been published that certain CD300 molecules have the ability to functionally interact with lipids [10–12, 14]. Specifically, human CD300a has been shown to interact with phosphatidylserine and phosphatidylethanolamine, two lipids that are exposed on the outer leaflet of the plasma membrane of apoptotic and activated cells [14]. Although the natural ligand of human CD300c is unknown, given that CD300a and CD300c show more than 90% homology in their IgV like domain, it would not be unreasonable to expect that CD300c may bind the same or structurally similar lipids to those that are recognized by CD300a. Further studies are required to identify the ligand of human CD300c and to fully understand its physiological role.

In conclusion, here we show that the clone TX45 monoclonal antibody specifically recognizes CD300c. We demonstrate that this activating receptor is only expressed on the surface of human monocytes within peripheral blood leukocytes, and that its engagement results in the production of proinflammatory cytokines and upregulation of ligands for T-cell costimulatory receptors. Altogether, this suggests that CD300c may have a very important role in regulating different immune responses.
Acknowledgments

The authors would like to thank Dr. Akira Shibuya for the generous gift of clone TX49 antibody. This work was funded by the intramural program of the Food and Drug Administration.

Disclosure Statement

The authors declare no competing financial interests.

References


