Unique Appearance of Proliferating Antigen-Presenting Cells Expressing DC-SIGN (CD209) in the Decidua of Early Human Pregnancy

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Intact human pregnancy can be regarded as an immunological paradox in that the maternal immune system accepts the allogeneic embryo without general immunosuppression. Because dendritic cell (DC) subsets could be involved in peripheral tolerance, the uterine mucosa (decidua) was investigated for DC populations. Here we describe the detailed immunohistochemical and functional characterization of HLA-DR-positive antigen-presenting cells (APCs) in early pregnancy decidua. In contrast to classical macrophages and CD83+ DCs, which were found in comparable numbers in decidua and nonpregnant endometrium, only decidua harbored a significant population of HLA-DR+/DC-SIGN+ APCs further phenotyped as CD14+/CD4+/CD68+/CD83+/CD25+. These cells exhibited a remarkable proliferation rate (9.2 to 9.8% of all CD209+ cells) by double staining with Ki67 and proliferating cell nuclear antigen. Unique within the DC-family, the majority of DC-SIGN+ decidual APCs were observed in situ to have intimate contact with CD56+/CD16+/ICAM-3+ decidual natural killer cells, another pregnancy-restricted cell population. In vitro, freshly isolated CD14+/DC-SIGN+ decidual cells efficiently took up antigen, but could not stimulate naive allogeneic T cells at all. Treatment with an inflammatory cytokine cocktail resulted in down-regulation of antigen uptake capacity and evolving capacity to effectively stimulate resting T cells. Fluorescence-activated cell sorting analysis confirmed the maturation of CD14+/DC-SIGN+ decidual cells into CD25+/CD83+ mature DCs. In summary, this is the first identification of a uterine immature DC population expressing DC-SIGN, that appears only in pregnancy-associated tissue, has a high proliferation rate, and a conspicuous association with a natural killer subset. (Am J Pathol 2003, 162:887–896)
mucosal surfaces. Thus, such a population could be responsible for maternal tolerance against fetal antigens in the mucosal decidua.

Recently, a novel DC-specific adhesion receptor DC-SIGN (dendritic cell-specific ICAM-grabbing nonintegrin, classified as CD209) has been identified that binds with high affinity to ICAM-2 and ICAM-3. In vivo, DC-SIGN is expressed by immature DCs in peripheral tissue as well as mature DCs in lymphoid tissues. Originally this type II mannose-binding C-type lectin DC-SIGN was cloned from a placenta library through its capacity to bind to the HIV-1 envelope glycoprotein gp120. Herein, we report that in contrast to normal endometrium a huge subpopulation of APCs in the decidua express DC-SIGN. These cells are found in situ in close contact to decidual large granular lymphocytes (LGLs) and exhibit a high proliferative activity. Directly after isolation, DC-SIGN+ APCs show functional features of immature DCs, but can be efficiently matured to potent immunostimulatory CD83+ DCs.

**Materials and Methods**

**Tissue Specimens**

All investigations were approved by the Ethics Committee of the Medical Faculty of the University of Würzburg, Germany, all patients gave informed consent for tissue collection. Decidual tissue (decidua basalis and parietalis) was obtained from 15 healthy women undergoing legal therapeutic abortion of an intact, normally progressing pregnancy with documented fetal heart activity at weeks 7 to 8 of gestation after the last menstrual period. All specimens contained embryonic components as verified by macroscopic and histological examination. Decidual tissue was taken from each specimen to be snap-frozen in liquid nitrogen for histological examination and immunohistochemical staining. The remainder was kept for no more than 30 minutes in phosphate-buffered saline (PBS) before subsequent cell isolation. Endometrial tissues from 17 women at fertile age undergoing hysterectomy because of uterus myomatosus and placental bed biopsies from 10 women during caesarian section at term were snap-frozen in liquid nitrogen for immunohistochemistry.

**Immunohistochemistry**

The antibodies applied in this study are listed in Table 1. Serial frozen sections of decidua, endometrium, and placental bed biopsies were cut at 5 μm and placed onto APES (3-aminopropyltriethoxy-silane; Roth, Karlsruhe, Germany)-coated slides, air-dried overnight, fixed in acetone for 10 minutes, and rehydrated in Tris-buffered saline (TBS; 25 mmol/L Tris/HCl, pH 7.4, 137 mmol/L NaCl, 2.7 mmol/L KCl). In proliferating cell nuclear antigen (PCNA) staining, sections were immediately (without drying) transferred into a methanol solution, fixed in 4% buffered formalin, and rehydrated in Tris-buffered saline (5 minutes each) before double stainings. For double/triple-immunohistochemical staining procedures of different cells or proliferating cells, respectively, sections were incubated with two to three cycles of: first, the monoclonal antibody at appropriate dilutions; second, the horse-radish-peroxidase-labeled rabbit anti-mouse-specific secondary antibody (dilution 1:100; DAKO, Hamburg, Germany); and third, the detection reaction followed by 10 minutes of air-drying. The first detection reaction was developed with 3,3′-diaminobenzidine (Sigma, Deisenhofen, Germany), the second with the Vector VIP peroxidase substrate kit (Vector Laboratories, Burlingame, CA), and the third with Vector SG substrate kit (Vector Laboratories) or HistoGreen substrate kit (Linaris, Wertheim, Germany), respectively. Sections were counterstained with hematoxylin (Sigma) or not.

For blue-red double-immunohistochemical staining of co-localized antigens, sections were first incubated with the DC-SIGN-specific monoclonal antibody at appropriate dilution followed by biotin-labeled goat anti-mouse antibody (dilution 1:100, DAKO) and then alkaline phosphatase-labeled streptavidin (dilution 1:300, Sigma) for 30 minutes each. Second the sections were incubated with the fluorescein isothiocyanate (FITC)-labeled specific antibody against the antigen of interest, followed by incubation of the peroxidase-labeled rabbit anti-FITC antibody (DAKO). Before substrate application, endogenous alkaline phosphatase activity was blocked with

**Table 1. Antibodies Used for Immunohistochemistry**

<table>
<thead>
<tr>
<th>Antigen</th>
<th>Clone</th>
<th>Isotype</th>
<th>Major specificity</th>
<th>Supplier</th>
</tr>
</thead>
<tbody>
<tr>
<td>CD4</td>
<td>RPA-T4</td>
<td>IgG1</td>
<td>Helper/inducer T cells, monocyte subsets</td>
<td>Pharmingen</td>
</tr>
<tr>
<td>CD14</td>
<td>UCHM1</td>
<td>IgG2a</td>
<td>Monocytes, macrophages, granulocytes</td>
<td>Serotec</td>
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<tr>
<td>CD34</td>
<td>QBEnd10</td>
<td>IgG1</td>
<td>Capillaries, endothelial cells</td>
<td>Cymbus Biotech.</td>
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<tr>
<td>CD56</td>
<td>NCAM16.2</td>
<td>IgG2b</td>
<td>Natural killer-cells, T-cell subsets</td>
<td>Becton Dickinson</td>
</tr>
<tr>
<td>CD68</td>
<td>KP 1</td>
<td>IgG1</td>
<td>Macrophages, activated platelets</td>
<td>DAKO</td>
</tr>
<tr>
<td>CD83</td>
<td>HB15A</td>
<td>IgG2b</td>
<td>Mature DCs</td>
<td>Immunotech</td>
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<td>DC-SIGN</td>
<td>AZ1-D1</td>
<td>IgG1</td>
<td>DC-SIGN, DC-specific ICAM-3 receptor, does not detect L-SIGN/DC-SIGN-R</td>
<td>Geijtenbeek et al.</td>
</tr>
<tr>
<td>HLA-DR</td>
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<td>IgG1</td>
<td>Monocytes, macrophages, DCs, B lymphocytes</td>
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<td>PC10</td>
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<td>Proliferating nuclear antigen</td>
<td>DAKO</td>
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0.1% levamisole (Sigma) in TBS, pH 8.2. As substrates for the enzymes, first the alkaline phosphatase-detecting APIII-Kit (blue, Vector Laboratories) and then the horseradish peroxidase-specific AEC (red, DAKO) were applied. Sections were not counterstained and were embedded in aqueous mounting media (Aquatex, Sigma).

For purple-green immunohistochemical staining of colocalized antigens, the DC-SIGN antibody was detected by incubation with horseradish peroxidase-labeled rabbit anti-mouse-specific secondary antibody (DAKO) and the detection reaction (very short incubation time) with the Vector VIP (Vector Laboratories). After blocking with mouse-IgG, remaining horseradish peroxidase activity was blocked by 10 minutes of air-drying. Second the sections were incubated with the FITC-labeled specific antibody against the antigen of interest, followed by incubation of the peroxidase-labeled rabbit anti-FITC antibody. Second detection reaction was performed with the HistoGreen substrate kit.

To evaluate the average density value of APCs in human first pregnancy decidua and endometrium, DC-SIGN+, CD14+, CD68+, and CD83+ cells were counted at the 100-fold magnification in 30 single fields of 1,008 mm² (=30.24 mm²) for each of two sections per patient per antibody by two independent observers in 11 decidual and 17 endometrial samples. Fields were randomly selected over the entire cryosection and completely filled with nonnecrotic cells.

Single Cell Isolation

For isolation of decidual cells in 10 cases, specimens were dissected free of products of conception and washed twice in PBS. The total decidual tissue (4 to 10 g) was then minced into fragments of ~1 mm³ and digested for 20 minutes at 37°C under slight agitation in PBS with 200 U/ml of hyaluronidase (Sigma), 1 mg/ml collagenase type IV (Seromed, Berlin, Germany), 0.2 mg/ml DNase I (2500 Kunitz U/mg, Sigma), and 1 mg/ml bovine serum albumin/fraction V (Sigma). The cell suspension obtained was filtered through sterile stainless steel 50-µm wire mesh and washed once in PBS. The mononuclear cell population was then separated by centrifugation over a Histopaque-1077 density gradient. After depletion of T cells with neuraminidase treated sheep erythrocytes (rosetting), the remaining non-T cells were resuspended in RPMI 1640 medium supplemented with 10% heat-inactivated fetal calf serum (PBS/IgG, Berglabor; Ceton, Marburg, Germany) and separated in a positive selection column in a magnetic cell separator (VarioMACS, Miltenyi).

For positive selection of CD14⁺ cells the cells were labeled with anti-CD14 Microbeads (clone Leu-M3; Miltenyi, Bergisch Gladbach, Germany) following the manufacturer’s instructions in PBS supplemented with 10% human immunoglobulin (PBS/IgG, Berglabor; Ceton, Marburg, Germany) and separated in a positive selection column in a magnetic cell separator (VarioMACS, Miltenyi).

The positive CD14-enriched fraction containing decidual DCs was analyzed by flow cytometry and used for subsequent analysis in a primary mixed leukocyte reaction or antigen uptake experiments or maturation experiments, respectively.

Cell Labeling and FACS

Flow cytometry of CD14-enriched decidual cells was performed with the monoclonal DC-SIGN (AZN-D1) antibody, with isotype control (mixture of mouse IgG1, IgG2a, IgG2b; all from Pharmingen, Heidelberg, Germany) and with the following primary antibodies directly labeled with FITC: HLA-DR (TU36; Caltag, Hamburg, Germany), CD14 (UHCM1; Cymbus/Chemicon, Hofheim, Germany), CD25 (M-A251, Pharmingen), CD68 (Ki-M7, Caltag), and CD83 (HB15a, Caltag). All incubation steps were followed by a wash in cold PBS/0.1% fetal calf serum. In brief, aliquots of 2 to 5 x 10⁶ isolated cells were resuspended in 20 µl of PBS/IgG. First, the DC-SIGN antibody (1:100) was added and incubated for 30 minutes on ice. After one wash, phycoerythrin (PE)-labeled goat anti-mouse antibody (Southern Biotechnology, Birmingham, AL) was incubated for another 15 minutes. After washing, unspecific Fc binding was then blocked with mouse IgG followed by a wash step. FITC-labeled antibodies (10 µl each) were added to the cell suspension and the preparation was then incubated for 30 minutes at 4°C. CD68 was stained intracellularly using the Fix and Perm kit (Dianova, Hamburg, Germany) according to the manufacturer’s instructions. ICAM-3 staining on LGLs was performed following the above procedure but with CD14-depleted decidual cells stained for ICAM-3 (clone 186-269, T. Geijtenbeek) followed by PE-anti-mouse and CD56/FITC (NCAM16.2, Becton Dickinson, Heidelberg, Germany). Cell suspensions were washed once, resuspended in 200 µl of PBS, and then analyzed in a FACScan flow cytometer (Becton-Dickinson). A total of 20,000 cells per sample were evaluated for specific staining. Results were analyzed using the WinMDI-software (Version 2.8, Joseph Trotter).

Generation and Maturation of Monocyte-Derived DCs (MoDCs)

For control purposes, DCs were generated from blood monocytes following standard procedures in four cases. In brief, peripheral blood mononuclear cells derived from buffy coats of healthy volunteer donors were prepared by isolation over a Histopaque-1077 density gradient. After depletion of T cells with neuraminidase treated sheep erythrocytes (rosetting), the remaining non-T cells were resuspended in RPMI 1640 medium (Seromed) containing 10% heat-inactivated fetal calf serum (PAN Biotech GmbH, Aidenbach, Germany) and 50 µg/ml of gentamicin (Seromed) (R10) and incubated at 5 x 10⁶ cells/ml in a tissue-grade Petri dish to allow monocytes to adhere to the bottom. The nonadherent cells were removed after 1 hour. The adherent fraction was then cultured in R10/IL-4/GM that is R10 supplemented with GM-CSF (1000 U/ml; Sandoz, Basel, Switzerland) and IL-4 (800 U/ml; Strathmann Biotech AG, Hamburg, Germany) for 7 days. Half the volume of medium was replaced with fresh R10/GM-CSF/IL-4 at days 3 and 5. For mature MoDC culture, at day 7 in half of the culture dishes, half the volume of medium was replaced by fresh
R10 supplemented with maturation cocktail to a final concentration of: IL-1β (1000 U/ml; Strathmann), tumor necrosis factor-α (1000 μg/ml; Strathmann), IL-6 (1000 μg/ml; Strathmann) and PGE2 (E2, 10⁻⁸ mol/l; Calbiochem-Novabiochem GmbH, Bad Soden, Germany). The remaining half of culture dishes was fed with R10/IL-4/GM without cocktail, resulting in immature MoDCs. At day 10 of culture, nonadherent cells with typical appearance of DCs were collected, and used in the mixed leukocyte reaction (only mature DCs) and antigen uptake experiments. FACS analysis confirmed cocktail-matured MoDCs to express HLA-DR in 85.7 ± 7.8% SD and CD83 in 67.8 ± 9.8% SD.

Analysis of Antigen Uptake

Antigen uptake was determined by incubating 1 × 10⁶ cells with different amounts of FITC-conjugated chicken OVA/FITC (Isomer I and OVA protein, conjugated as previously described; Sigma, Deisenhofen, Germany) for 1 hour at 37°C in R10. Incubated cells were washed extensively with ice-cold PBS, stained with PE-conjugated HLA-DR on ice, and immediately analyzed by flow cytometry. Control incubation was performed at 4°C. Dead cells and cell debris were excluded by propidium iodide labeling and forward and side scatter gating.

Mixed Leukocyte Reaction

Different amounts of decidual APCs (dAPCs) or MoDCs were co-cultured in triplicates for 5 days at 37°C in 5% CO₂ with 9 × 10⁵ allogeneic T lymphocytes (prepared by 0.8% NH₄Cl lysis of neuraminidase-treated sheep red blood cell rosettes of peripheral blood mononuclear cells) in a final volume of 200 μl of R10 in round bottom 96-well plates (Falcon/Becton Dickinson, Franklin Lakes, NJ). During the final 16 hours of co-culture, 1 μCi of [³H]thymidine (Amersham International, Arlington Heights, IL) per well was added and incorporated radioactivity was determined.

Results

Decidua but Not Endometrium Contains DC-SIGN-Positive Cells

To quantify the cellular expression of monocyte/DC markers within endometrial and decidual tissue, morphometric
Phenotypical Characterization of Decidual APCs in Situ

**DC-SIGN** Decidual Cells Exhibit the in Situ Phenotype HLA-DR/CD14+/CD68+/−/CD4+/CD83−

Immunohistochemistry staining on cryostatic sections of decidual tissue revealed DC-SIGN+ cells to exhibit a veiled shape with long, irregular dendritic protrusions (Figure 2A) and to be regularly scattered through the tissue, with a preference for the areas with clear visible spiral arteries (Figure 2B). DC-SIGN+ cells could be clearly distinguished from mature DCs and were never found to be positive for CD83 (Figure 2, C and H).

To further characterize the phenotype of DC-SIGN+ cells immunohistochemical double stainings were performed. All DC-SIGN+ cells in decidual tissue expressed HLA-DR (Figure 2, D and I), CD14 (Figure 2, E and J), and CD4 (Figure 2, F and K). Most of them expressed CD68+ with a perinuclear pattern (Figure 2, G and L). Next to the double-positive cells (HLA-DR+/DC-SIGN+, CD14+/DC-SIGN+, CD4+/DC-SIGN+, and CD68+/DC-SIGN+) some single-positive cells for HLA-DR, CD14, CD4, and CD68 were detected (Figure 2; D to F), but not vice versa.

In contrast to the smaller CD83+ mature DCs, which were often found in clusters with CD3+ T cells, DC-SIGN+ cells were only occasionally seen in clusters with more than three T cells (Figure 2C). However, the majority of DC-SIGN+ cells was found in close contact to CD68+ LGLs (Figure 3A). Quantification of LGL/DC-SIGN+ cell clusters revealed 52 ± 4.9% SD of DC-SIGN+ cells to be associated with one LGL, 9.6 ± 4.5% SD with two LGLs, and 1.9 ± 1.2% SD with three LGLs. Only very occasionally DC-SIGN+ cells were found in intimate contact with more than three LGLs. Co-staining of DC-SIGN+ with ICAM-3, a ligand for DC-SIGN, revealed, that many of ICAM-3-positive cells resembling LGLs were attached to the DC-SIGN+ cells (Figure 3B). Indeed, double labeling clearly demonstrated all LGLs to be positive for ICAM-3 and only occasionally single cells to be ICAM-3+/CD68− (Figure 3C). FACS analysis confirmed the immunohistochemical findings (Figure 3D). Previously published data19 has shown, that besides proliferating LGLs (7 to 23% Ki-67+) there are numerous Ki-67-positive CD68+ cells found in decidua. Immunohistochemical double staining (n = 10) confirmed that 9.3 ± 0.4% SD of all DC-SIGN+ cells co-expressed Ki-67 (Figure 3, E and G). Additional double-immunohistochemical analysis using DC-SIGN/PCNA demonstrated 9.8 ± 0.3% SD of DC-SIGN+ cells to be PCNA+ (Figure 3, E and F).

Phenotypical Characterization of Decidual APCs in Vitro

**Decidual DC-SIGN** Cells Exhibit the Phenotype of Immature DCs

To further estimate the amount of DC-SIGN+ cells in the decidua FACS analysis was performed on freshly isolated decidual cells. As shown in Figure 4A, decidual cells contain 5 to 8% DC-SIGN+ cells. Double staining confirmed the immunohistological findings and revealed that all DC-SIGN+ cells were positive for CD14 and that the vast majority of the cells was positive for HLA-DR. DC-SIGN+ cells were clearly negative for CD83 and CD25. Intracellular staining performed with the CD14+-enriched decidual cell population revealed about 50% of the DC-SIGN+ cells to be positive for CD68 (Figure 4B).

**Decidual DC-SIGN** Cells Can Be Matured in Cell Culture into CD83+/CD25− DCs

Analysis of APC surface marker expression by decidual CD14+ cells either freshly isolated or cultured for 3 days in the presence of a cocktail of inflammatory cytokines is shown in Figure 5. In comparison to a slight increase in the mean fluorescence intensity (MFI) of HLA-DR-expression (144.9 ± 23.7 SEM on fresh cells versus 177.5 ± 22.8 SEM on cultured cells), the level of CD83 and CD25 strongly increased on culture. Although the MFI of CD83 increased sevenfold (from 10.8 ± 1.1 SEM on fresh cells to 70.9 ± 14.8 SEM on cultured cells), the MFI of CD25 increased dramatically (13.4 ± 0.9 SEM on fresh cells versus 146.9 ± 48.7 SEM on cultured cells). In contrast to the maturation markers, treatment of decidual CD14+ cells with cocktail resulted in a decrease of DC-SIGN and CD14. Whereas the MFI for DC-SIGN on cultured cells was reduced to half (48.7 ± 0.5 SEM to 267 ± 1.9 SEM), expression of the monocyte/macrophage marker CD14 dropped extensively (200.4 ± 60.0 SEM on cultured cells versus 860.0 ± 40.6 SEM on fresh cells).

Functional Analysis of Fresh Isolated and Cocktail Matured Decidual APCs in Comparison to Monocyte-Derived DCs

Because of the fact that the DC-SIGN antibody blocks DC-T-cell interaction13 and thereby inhibits T-cell prolif-
Figure 3. Interaction of proliferating DC-SIGN⁺ cells with CD56⁺ LGLs. A: Interference contrast clearly demonstrated the intimate contact between DC-SIGN⁺ cells (violet, white arrowhead) with CD56⁺ LGLs (brown, black arrow). B: DC-SIGN⁺ cells (green, white arrowhead) together with ICAM-3-positive cells (brown, arrow) that resemble the morphology of LGLs. C: Double labeling of ICAM-3 (blue) with CD56 (red) demonstrated all CD56⁺ cells to be ICAM-3⁺ (purple), one single ICAM-3⁺/CD56⁻ cell was seen (blue, black arrow). White arrowheads indicate examples of ICAM-3⁺/CD56⁻ LGLs. D: FACS analysis of isolated decidual lymphocytes. Left: A high percentage of the cells express ICAM-3. Right: All CD56⁺ LGLs co-express ICAM-3, representing the largest population of all ICAM-3-positive cells. E: Bar chart of proliferating DC-SIGN⁺ cells detected by double staining with Ki-67 (white bar, ~9.2% of all DC-SIGN⁺ cells) and PCNA (black bar, ~9.8%). F: Immunohistochemistry shows clear double staining for DC-SIGN (green) and PCNA (brown, white arrowhead) some PCNA⁻/DC-SIGN⁺ cells were seen (black arrow) next to proliferating non-DC-SIGN⁺ cells (black arrowhead). G: Simultaneous staining for DC-SIGN (green), CD56 (brown), and Ki-67 (violet) revealed a proliferating DC-SIGN⁺ cell (arrow) next to a proliferating LGL (white arrowhead) and closely attached to a nonproliferating LGL (black arrowhead). Original magnifications: ×400 (A, F), ×160 (B, C), ×1000 (G).
eration in allogeneic T-cell proliferation assays, decidual DC-SIGN+ APCs were enriched by CD14 selection, resulting in a population of ~80% HLA-DR+ cells containing 50 to 70% DC-SIGN+ cells.

To further analyze the maturation stage of these DC-SIGN+ APCs antigen uptake and allogeneic T-cell stimulation capacity assays were performed as shown in Figure 6. In a dose-dependent manner, fresh CD14+ enriched dAPCs had a very high FITC-OVA uptake capacity (fourfold higher than immature MoDCs) that decreased on maturation to levels exhibited by immature MoDCs (Figure 6A).
After cocktail maturation, decidual APCs stimulated allogeneic T cells as efficiently as cocktail matured MoDCs (Figure 6B). Interestingly freshly isolated decidual APCs did not have any stimulatory capacity, even at the highest APC:T cell ratio no T-cell proliferation was apparent. Nevertheless, the allostimulatory capacity of cocktail matured dAPCs was in all cases slightly higher than those of the control MoDCs. These data reflect the particular ability of the isolated decidual APCs to efficiently mature into potent stimulators of allogeneic T cells on exposure to DC maturation stimuli such as a cocktail of inflammatory cytokines.

Discussion

Human decidual tissue appears to play an important role in the immune surveillance of the implanted embryo. Many features of the local maternal-fetal interaction in the uterus are related to the primitive innate immune systems. Two of the immune cell types appearing early in ontogeny are phagocytes and natural killer cells, which seem to be the most primitive cytolytic cells. Both, phagocytes and natural killer cells persist during ontogeny and can be found in the human decidualized endometrium (decidua), where the embryo implants. Compared with the uterine endometrium, early decidual tissue has been shown to contain a markedly increased number of granulated CD56+ natural killer cells. These LGLs are the predominant cell population during early human pregnancy. CD14+ cells, generally described as decidual macrophages, constitute the second main cell population. Here we report, that nonpregnant endometrium and decidua harbor macrophage/dendritic-like cells expressing CD4/HLA-DR/CD68/CD14, but only in the decidua do these APCs co-express the DC-SIGN antigen. DC-SIGN+ cells, however, were found in nonpregnant uterine samples in outer regions of the myometrium (not shown).

DC-SIGN+ cells were recently described in placenta and decidual tissue. In line with our findings, Soilleux and colleagues found co-expression of CD4 by DC-SIGN+ cells. The reported high number of CD4+ cells contrasts previous characterizations of leukocyte populations in human decidua but can be explained by optimized immunohistochemical staining systems that allow the detection of even weak CD4 expression by DC-SIGN+ cells.

The appearance of DC-SIGN+ cells in the decidua is a pregnancy-associated event and may result from up-regulation of this surface marker on a subpopulation of endometrial macrophage/dendritic-like cells induced by signals related to the exogenous and endogenous tissue damage caused by invading trophoblasts. A candidate signal is IL-13 expressed at high levels in the placenta. This Th2-type cytokine has recently been shown in vitro to significantly up-regulate the DC-SIGN surface expression on macrophages. Alternatively, with onset of pregnancy DC-SIGN+ cells of the myometrium or blood might invade the decidua and then proliferate in situ. This hypothesis is supported by our finding of a remarkable number of proliferating Ki67+/DC-SIGN+ or PCNA+/DC-SIGN+ cells within the decidua (~10%).

Because human endometrium appears to harbor DC-SIGN+ cells only during decidualization, these cells might play a crucial role for the local immune response at the barrier between immunity and tolerance. Of special interest seems to be the close topographical relationship of pregnancy-related DC-SIGN+ cells with ICAM-3-expressing LGLs. Corresponding to the high affinity of DC-SIGN for ICAM-3, this intimate cell contact is likely mediated by DC-SIGN/ICAM-3 interactions. Hypothetically, binding of ICAM-3-positive LGLs to DC-SIGN+ decidual APCs may on one hand prevent the interaction of DC-SIGN+ cells with T cells, and furthermore impede maturation of DC-SIGN+ APCs into potent immunostimulatory DCs. Indeed, LGLs have been shown to produce a wide range of cytokines, especially high local concentrations of GM-CSF and IL-10 known as a potent inhibitor of DC maturation. On the other hand, interaction of DC-SIGN+ APCs with LGLs might result in activation of LGLs and even stimulate their proliferation. Further studies on the interaction of these two pregnancy-related cell populations should help understanding how immune responses in human pregnancy are regulated.

Finally, we show that decidual DC-SIGN+ cells in situ and in vitro not only resemble immature DCs by phenotype and function, but in addition very effectively may be turned into potent immunostimulatory CD83+CD25+ mature DC in vitro if maturation stimuli such as a cocktail of inflammatory cytokines are provided. From these data we conclude, that at least three different macrophage/DC populations do occur in human early pregnancy decidua: classical macrophages, mature CD83+ DCs, and DC-SIGN expressing immature DC-like APCs prone to mature into potent immunostimulatory DCs. Because macrophages and immature MoDCs principally are able to convert into one another until late stages of their respective differentiation/maturation process, our DC-SIGN+ immature DC-like decidual cell may correspond to the transient differentiation cell type (the “veiled immediate DC precursors”) postulated by Palucka and colleagues. It is tempting to speculate that the local cytokine environment and cellular interactions within the decidua may well constitute the final decision-making factors determining whether DC-SIGN+ APCs will acquire characteristics and functions of macrophages or mature DCs.

References

5. Thomas R, Davis LS, Lipsky PE: Comparative accessory cell function...