Immunohistochemical Evidence of HLA-G Expression in Extravillous Trophoblast Invading Decidual Tissues

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Abstract

HLA-G is a non classical HLA I gene product involved in the regulation of implantation and in the immune tolerance during pregnancy, by modulating maternal immune responses at the fetal-maternal interface. In pregnancies lacking immune tolerance and ending in miscarriage, the expression of HLA-G is very probably defective or altered. In order to contribute to further investigations about HLA-G expression in autoimmune miscarriages, we have tried to describe HLA-G immunohistochemical patterns of expression in a homogeneous cohort of normal first trimester choriodedicial specimens, in the three different extravillous trophoblast (EVT) populations ("cell islands", "cell columns", and intravascular EVT cells). We expected to demonstrate HLA-G reactivity to the tested antibody (MEMG01) in all three populations. All choriodedicial specimens were histologically and clinically reviewed to exclude any pathologic finding. Immunohistochemical identification of trophoblast cells in the selected specimens was performed via wellknown immunostains such as Cytokeratin 8/18 (CAM5.2) and NCL-PLp; anti-Ki67 was also used to point out proliferating EVT cells. Then, MEM-G01 was tested at various dilutions, with or without pretreatment, to find the optimal protocol. As expected, HLA-G specifically stained all three EVT populations, with decreasing reactivity from EVT cell islands to EVT cell columns or intravascular EVTs. The next step will be the investigation of HLA-G pattern of expression in autoimmune aborters.

Introduction

The Major Histocompatibility Complex (MHC) class I includes two groups of genes. Group la genes encode the classical, highly polymorphic HLA-A, HLA-B and HLA-C antigens, which are expressed on the majority of nucleated cells, while group lb encodes the less polymorphic, nonclassical, HLA-E, HLA-F and HLA-G antigens. HLA-G, first described in 1987 [1] is a nonclassical HLA class I (class lb) gene product with limited polymorphism [2] and restricted tissue distribution [3], known to be involved in the regulation of implantation [4] and in the immune tolerance in human pregnancy [5-6]. At present, seven different isoforms of HLA-G are known: four of them HLA-G are known: four of them (HLA-G1, G2, G3 and G4) are membranebound, while the last three (HLA-G5, G6 and G7) exist as soluble isoforms. Placental cells migrating into the maternal decidua synthesize both membrane and soluble HLA-G isoforms [7], which interact with inhibitory receptors on leukocytes such as ILT2 and ILT4 and, by inhibiting maternal immune responses to foreign (paternal) antigens, facilitate semiallogeneic pregnancy. Such HLA-G proteins act in different ways on immune cells: cytotoxic, CD8 positive, lymphocytes (CTL) are induced by HLA-G either to die or to decrease CD8 production; natural killer cells are immobilized, while mononuclear phagocytes increase production of antiinflammatory cytokines. At the fetalmaternal interface, all these effects of HLA-G modulate the maternal local immune responses in almost every pregnancy [8]. Conversely, in pregnancies lacking immune tolerance and ending in miscarriage, HLA-G expression very probably is defective or altered. The present study was conceived in the setting of a larger interdisciplinary research group which is investigating HLA-G expression in both human fluids and tissues from Systemic Lupus Erythematosus (SLE) patients. Histological assessment of HLA-G in choriodedicial specimens obtained from spontaneous miscarriages in women affected by autoimmune diseases, particularly SLE, could help in the management of future pregnancies in these women. Despite many studies have demonstrated HLA-G in amniotic fluid and maternal blood, only a few of them have investigated HLA-G expression by immunohistochemistry. Moreover, while HLA-G expression on extravillous trophoblast (EVT) is well established [9-10], more controversial results have demonstrated its expression in Hofbauer mesenchimal cells [11-12], in some thymic medullary cells, in syncytiotrophoblast [13], in staminal cytotrophoblast, and in some endothelial or amniotic epithelial cells [14]. In order to provide a contribution to the
study of HLA-G expression in autoimmune miscarriages, we
have tried to describe HLA-G expression patterns in a
homogeneous cohort of normal first trimester
choriodecidual specimens. Three different extravillous
trophoblast populations are known: (1) the so-called "cell
islands" covering the villi; (2) the scattered trophoblast cells
migrating through the decidua ("cell columns") toward the
decidual vessels, and (3) the intravascular trophoblast cells
floating in the lumen of converted decidual arteries. We
expected to demonstrate HLA-G reactivity in all three
populations.

Materials and methods

Clinical and histological criteria of enrollment. We
selected from the database of our Pathology Institute ten
cases of abortion, in order to assess HLA-G reactivity in
normal choriodecidual and, when present, embryofetal
tissues. The four major criteria of enrollment (Tab. 1) were
 gestational age under 13 weeks, voluntary abortion,
presence of basal and parietal decidua as well as chorionic
plate, and the lack of pathological findings in the
histopathological report. Hence, all spontaneous
miscarriages, incomplete samples and specimens showing
morphological signs of infection, abnormal karyotype or
other potential causes of miscarriage were excluded. We
collected clinical data of all women and histologically
reviewed each slide, recording patient's history and
histologic findings in Microsoft® Excel database. The main
purpose of this review was to determine the
composition of any paraffin block: particularly, only those
with basal decidua, including EVT cells, converted vessels
and Nitabuch's stria, would have been submitted to
immunohistochemistry. In addition, given the results
obtained by some investigators (see above), HLA-G was
tested in both fetal and neonatal paraffinembedded thymic
tissue, lacking any clinical and histological evidence of
thymic or immune disease.

<table>
<thead>
<tr>
<th>Major criteria of enrollment</th>
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<tbody>
<tr>
<td>- gestational age &lt; 13 weeks (90 days)</td>
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<td>- voluntary abortion</td>
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<tr>
<td>- presence of basal and parietal decidua and chorionic plate</td>
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<tr>
<td>- lack of pathological findings in the histopathological report</td>
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</tbody>
</table>

Table 1 - Clinical and histological criteria of enrollment

Immunohistochemistry. Curettage specimens from
voluntary abortions were investigated by immunostaining
for different antigens (Tab. 2), performed on serial sections
(the first stained with hematoxylin and eosin (H&E), to
compare their morphological expressions. Trophoblast cells
were identified by using antiCyto keratin 8 and 18 (Clone
SD3, DiaPath, USA) and NCL-PLp (Novocastra, Newcastle
upon Tyne, UK) antibodies. HLA-G reactivity was assessed
using the antibody MEM-G/01 (Exbio, Prague, Czech
Republic), which is known to recognize all HLA-G
isoforms. Finally, proliferating cells were marked with
Confirm™ anti-Ki67 antibody (Ventana Medical Systems,
Tucson, Arizona, USA).

<table>
<thead>
<tr>
<th>Antibody</th>
<th>Source</th>
<th>Dilution</th>
<th>Target</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anti-Cytokeratin 8/18 (clone SD3)</td>
<td>DiaPath, USA</td>
<td>1:100</td>
<td>Trophoblast cells</td>
</tr>
<tr>
<td>NCL-PLp</td>
<td>Novocastra UK</td>
<td>1:200</td>
<td>Trophoblast cells</td>
</tr>
<tr>
<td>MEM-G/01</td>
<td>Exbio CZ</td>
<td>1:100</td>
<td>EVT cells</td>
</tr>
<tr>
<td>Confirm™ anti-Ki67</td>
<td>Ventana Medical Systems USA</td>
<td>RTU</td>
<td>Proliferating cells</td>
</tr>
</tbody>
</table>

Table 2 - Employed antibodies and their tissue targets (RTU=ready-to-use).

Cytokeratin 8/18 Ab1 (Clone SD3), also known as
CAM5.2, recognizes all simple epithelia including thyroid,
breast, gastrointestinal and respiratory glandular epithelia,
as well as urogenital structures, including transitional
epithelium. Keratin 8 belongs to the type B and Keratin 18
to the type A family of HMW cytokeratins. Clone SD3 was
obtained by using cytokeratins purified from breast cancer
MFC7 cells as an immunogen, and its pattern of expression
is cytoplasmic. Tissue was pretreated by digestion with
trypsin (1 mg/ml PBS, pH 7.4) for 15 minutes at RT, and
antibody was incubated at 1:100 dilution for 30 minutes at
RT. [15-16] MEM-G/01 reacts with denatured HLA-G
heavy chain of all isoforms. It is obtained using as an
immunogen recombinant human HLA-G denatured heavy
chain; the purified antibody is conjugated with BiotinLC-
NHS and the reagent is free of unconjugated biotin. It is
stored (1 mg/ml) in phosphate buffered saline (PBS) with
15 mM sodium azide (approx. pH 7.4). As a starting point,
we tried to find the appropriate working dilution and
antigen retrieval technique. To find the optimal working
dilution, MEM-G/01 was tested at the suggested dilution
(1:50) and at different decreasing concentrations (1:100,
1:500, 1:1000, 1:5000) on: A) chorio decidual specimen from
elective cessation of pregnancy at 8 gestational weeks in a
36 year old woman; B) fetal thymus from stillbirth at 41
weeks, following intrauterine asphyxia; and C) neonatal
thymus from a newborn deceased immediately after
delivery due to gastrointestinal complications. Such dilutions
of MEM-G/01 were tested on each sample: 1) without
antigen retrieval; 2) with EDTA and 98.1°C bath; 3) with
EDTA and heating in microwave oven for 3 minutes at
900W and for 13’ at 360W. Results were compared and
allowed us to choose the best pretreatment for further
experimental investigations. Finally, MEM-G/01 was tested
at the reduced dilution of 1:100 on chorio decidual tissue
taken from all ten cases. NCL-PLp is a rabbit polyclonal
antibody specific for human placental lactogen (hPL),
obtained from placental lactogen isolated from human
placenta. It stains membrane and cytoplasm of
syncytiotrophoblast cells and it is useful for the detection of
HPL in the placenta and in all cells showing trophoblastic differentiation. It was incubated for 60 minutes at 25°C at 1:200 dilution, performing standard ABC technique in a Benchmark automatic immunostainer. NCL-PPlp is the trophoblaststaining antibody routinely used in our Institution in the setting of the investigation of early spontaneous abortions [17]. Confirm™ anti-Ki67 (clone K2) is a mouse monoclonal antibody which identifies the Ki67 nuclear antigen in proliferating cells and it is obtained by an immunogen recombinant protein corresponding to its C-terminal. As for cytokeratins 8/18 and for NCL-PPlp, a Benchmark automatic immunostainer was used.

Results

Clinical records. The 10 women enrolled in the study were 19 to 42 years old (mean age: 31 yrs). Gestational age at the time of cessation of pregnancy ranged from 8 to 11 weeks (mean gestational age: 9 weeks). Seven women (70%) were facing their first pregnancy, while the other three had yet experienced term pregnancies (Fig. 1). Moreover, two of the latter (20% of all cases) had previously undergone elective cessation of pregnancy. None of them referred any spontaneous miscarriage, late fetal loss, stillbirth, threatened abortion, preterm or otherwise pathologic pregnancy.

![Fig. 1 - Obstetric history: main clinical records; 3 out of 10 women had yet experienced term pregnancy, and 2 among them had undergone voluntary abortion as well.](image)

Specimen composition. Basal and parietal decidua and parts of chorionic plate were present in all cases, owing to the enrollment criteria, while only one case featured decidua capsularis as well. Embryo or fetus (depending on gestational age) were present in all cases; anyway, embryonic structures were seen only in 7 of the selected slides. Umbilical cord was found in 4 of 10 cases, and in two specimens amniotic membranes were seen as well. In three specimens decidua, chorionic disc, membranes and umbilical cord were present together (Fig. 2).

Histological features. All decidual specimens (Fig. 3) showed Arias-Stella reaction and hemorrhage (100% each), while aspecific/necrobiotic infiltrate was seen in 7 out of 10 cases (70%), thrombosis in 5 (50%), edema or necrosis in 3 (30%) and a true lymphoplasmocytic infiltrate in only 2 cases (20%).

![Fig. 2 - Histological composition of the specimens.](image)

Immunohistochemical findings. Preliminary tests on thymic and choriodendritic tissue demonstrated that the optimal procedure at 1:50 dilution was antigen retrieval by heating with EDTA and 98.1°C bath. Anyway, at the working dilution of 1:50, EVT showed strong positivity, but a weaker membrane reactivity was seen in some thymic structure as well, such as adipocytes, endothelial cells, and scattered medullary cell both in fetal and neonatal thymus. Once the dilution was increased to 1:100, immunohistochemical staining for HLA-G resulted more specific, since only EVT cells invading decidual tissues were stained, while the "false" reactivity of the thymic cells disappeared. At the higher dilutions tested, both EVT and thymus turned out negative. HLA-G was found to mark all

![Fig. 3 - Histological features in the decidua.](image)

Main findings in the chorionic plate (Fig. 4) were light fibrin deposition, seen in 7 cases (70%), villar edema in 3 cases (30%), waxy degeneration of the villi in 2 cases (20%), and collapsed vessels in one case (10%). All the above mentioned are non specific findings, usually seen in elective pregnancy termination owing to surgical procedures. No actual pathologic finding was seen in the embryo/fetus, umbilical cord or amniotic membranes, when present.

![Fig. 4 - Histological features in the chorionic plate.](image)
EVT populations: (1) HLA-G was expressed by cell islands covering the villi (Fig. 5b) previously identified in H&E (Fig. 5a), (2) by column cells migrating through the decidual tissues (Fig. 6a, 6b) and (3) by intravascular EVT (Fig. 7). To identify EVT elements on H&E slides (Fig. 8a), we marked them with CAM5.2 (Fig. 8b) and hPL (Fig. 8c) and we searched for proliferating cells with Ki67 (Fig. 8d).

**Discussion**

Our study was intended to contribute to a larger research project about the role of HLA-G and its expression in the setting of autoimmune diseases. Previous studies demonstrated HLA-G presence in body fluids, but the few investigating its immunohistochemical expression did not
accurately differentiate between the three populations of EVT. Moreover, expression in such nontrophoblast cells such as Hoffbauer mesenchimal cells, some medullary cells of the thymus and some endothelial or epithelial cells needed further investigation. We found that the latter are nonspecific stainings, while, at the optimal dilution of 1:100 applied to normal first trimester chorionicdecidual specimens, MEM-G/01 specifically stains all three EVT populations, as we expected. We also demonstrated a decreasing HLA-G reactivity from EVT cell island to EVT cell columns or intravascular EVT; anyway, in the latter immunostaining remain easily detectable and specific. In conclusion, we have tried to provide a background for further investigations in the setting of immunohistochemical characterization of curettage specimens from autoimmune patients. The next step will consist in the immunohistochemical investigation of the HLA-G pattern of expression in autoimmune aborters.

References


Fig. 8 - Trophoblast cells invading maternal vessels (serial sections): a) hematoxylin and eosin; b) immunostaining with CAM5.2: cytokeratins are expressed by EVT; c) immunostaining with NCL-Pep identifies EVT; d) immunostaining with Ki67 marks proliferating EVT cells.
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