Stable inhibition of interleukin 1 receptor type II in Ishikawa cells augments secretion of matrix metalloproteinases: possible role in endometriosis pathophysiology

S Guay1,2 and A Akoum1,2

1Unité d’Endocrinologie de la Reproduction, Centre de Recherche, Hôpital Saint-François d’Assise, Centre Hospitalier Universitaire de Québec, 10 rue de l’Espinay, Local D0-711, Québec, Canada, G1L 3L5 and 2Département d’Obstétrique et Gynécologie, Faculté de Médecine, Université Laval, Québec, Canada, G1V 4G2

Correspondence should be addressed to A Akoum; Email: ali.akoum@crsfa.ulaval.ca

Abstract

Our previous studies showed a marked deficiency in interleukin 1 receptor type II (IL1R2) in the endometrial tissue of women with endometriosis, particularly in epithelial cells. We believe that such a deficiency in IL1R2, a potent and specific IL1 inhibitor, makes endometrial cells more sensitive to IL1 and less capable of buffering the cytokine’s effects, which may lead to functional changes that favor endometriosis development. The main objective of our study was to stably inhibit IL1R2 expression in endometrial cells in order to evaluate the role of IL1R2 deficiency in endometriosis pathophysiology. Stable clones of Ishikawa adenocarcinoma endometrial cells transfected with IL1R2 antisense and showing downregulation of IL1R2 protein expression, or with the empty expression vector alone and showing no noticeable difference in IL1R2 expression, were selected. The downregulation of IL1R2 expression in IL1R2 antisense transfecteds when compared with control cells was confirmed by ELISA, Western blot and immunofluorescence. In these cells, IL1R2 expression was markedly reduced, compared with non-transfected cells or cells transfected with the empty vector, and there was a significant increase in the basal and the IL1β (IL1B)-induced levels of matrix metalloproteinase (MMP)-2 and MMP-9 secretion. Furthermore, a significant decrease in IL1B-induced secretion of tissue inhibitor of MMPs-1, a known MMP-9 inhibitor, was observed. These in vitro data make plausible a role for IL1R2 deficiency in the capability of endometrial cells to invade the host tissue and develop in ectopic locations.

Introduction

Pro-inflammatory cytokines, particularly interleukin 1 (IL1), are involved in the modulation of a variety of endometrial functions (Tabibzadeh 1991, Simon et al. 1998). A tight control of IL1 effects is therefore necessary to maintain homeostasis and normal endometrial functions. IL1 receptor type I (IL1R1), the functional signaling receptor which mediates cell activation by IL1, is expressed in the human endometrium (Simon et al. 1993) and follows a gradually increasing expression pattern throughout the menstrual cycle (Bigonnesse et al. 2001). IL1R2, which has no signaling properties and rather acts as decoy receptor for IL1, is also expressed in the human endometrium and follows a more complex cycle-phase dependent expression pattern during the menstrual cycle (Boucher et al. 2001, Kharfi et al. 2002). Several studies suggest that IL1R2 plays an important physiological role in the regulation of IL1 action in the inflammatory sites by capturing IL1 and preventing its interaction with IL1R1 (Colotta et al. 1993, 1994, Bossu et al. 1995, Orlando et al. 1997, Coulter et al. 1999). Our previous data showed a marked deficiency in IL1R2 expression in the eutopic endometrial tissue of women with endometriosis, occurring at the protein (Akoum et al. 2001a) and the mRNA (Kharfi et al. 2002) levels, particularly in epithelial cells. Therefore, imbalance between the activating IL1R1 and the inhibitory IL1R2 in endometrial cells of women with endometriosis may result in increased cell reactivity to IL1 in the eutopic endometrial tissue, but likely in the ectopic locations as well and in response to local stimuli such as IL1, which may favor endometriosis development. This is all the more possible since either eutopic or ectopic endometrial cells of women with endometriosis showed increased sensitivity to IL1B (Akoum et al. 1995a, 1995b, 2001b, Sillem et al. 1999, 2001,
Lebovic et al. 2000). Furthermore, elevated concentrations of IL1B were found in the peritoneal fluid (Mori et al. 1992, Taketani et al. 1992) and the ectopic endometrial tissue of women with endometriosis (Bergqvist et al. 2001). This emphasizes the role of this cytokine in the pathophysiology of the disease and further supports the relevance of our findings.

Therefore, the objective of the present study was to specifically inhibit IL1R2 expression in endometrial cells, and create a stably transected endometrial cell line that will allow the assessment of the role of IL1R2 deficiency in IL1-mediated functional changes that may favor ectopic endometrial cell growth and endometriosis development. As IL1R2 deficiency was more obvious in endometrial epithelial cells (Boucher et al. 2001), a well-differentiated endometrial epithelial adenocarcinoma cell line (Ishikawa cells; Nishida et al. 1985) which responds to IL1B (Makrigiannakis et al. 1999) was used as model and had undergone a stable transfection to block IL1R2 expression and to recreate the imbalance that we found in endometrial cells of patients with endometriosis.

Material and Methods

Cell transfection

For these studies, we used a well-differentiated endometrial adenocarcinoma cell line (Ishikawa cells; Nishida et al. 1985) which was stably transected with the pcDNA expression vector alone or containing IL1R2 cDNA in the antisense direction. This line maintains functional estrogen and progesterone receptors and numerous endometrial epithelial cell functions (Croxtall et al. 1990, Hata & Kuramoto 1992, Lessey et al. 1996, Castelbaum et al. 1997). Cell transfection was performed using Lipofectamine Plus reagent according to manufacturer’s instructions (Invitrogen). Cells were seeded in 24-well culture plates (Costar, Cambridge, MA, USA), cultured in Dulbecco’s modified Eagle medium–F12 medium (DMEM–F12) containing 10% fetal bovine serum (FBS) and 1% antibiotics (Invitrogen) until 70% confluence, and incubated with DNA/Lipofectamine complexes in FBS-free DMEM. After 3 h of incubation at 37 °C, medium containing 10% FBS was added and culture continued for 24 h. Cells were then passed (1/20 dilution) in a fresh culture medium containing 150 µg/ml geneticin (Sigma–Aldrich). After 1–2 weeks, individual growing colonies were selected and separately expanded in culture, in the presence of 150 µg/ml geneticin. A total of 32 clones were isolated from cultures transfected with the plasmid vector harboring or not IL1R2 cDNA in the antisense direction. Clones were assessed as described below and all experiments were repeated at least thrice.

Culture stimulation

Cells were cultured at 37 °C in humidified 5% CO2. Cultures grown to confluence were trypsinized, harvested by centrifugation, and distributed in 12-well culture plates (Costar) in DMEM–F12 containing 150 µg/ml geneticin (only for cells transfected with pcDNA plasmid) with 10% FBS. Medium was changed every 2 days until confluence. Before cell stimulation, the culture medium was replaced by a serum-free medium for 24 h. Cells (three wells/treatment) were exposed or not to IL1B (0–10 ng/ml; Invitrogen) diluted in a fresh FBS-free medium for 24 h. The culture supernatant was collected and kept in small aliquots at −80 °C until used for ELISA and Western blotting. Cells were recovered in a lysis buffer solution containing 0.5% Triton X-100, 10 mM HEPES (pH 7.4), 150 mM NaCl, 2 mM EGTA, 2 mM EDTA and 0.02% NaN3, and a mixture of anti-proteases composed of 5 µM aprotinin, 63 µM leupeptin, and 3 mM phenylmethylsulfonylfluoride and kept at −80 °C.

Protein extraction

Cells recovered in the lysis buffer were disrupted by multiple passages through a tuberculin syringe with a 25-gauge needle and incubated at 4 °C for 45 min under gentle shaking. After centrifugation at 11 000 g for 30 min, soluble protein extracts were collected and total protein concentration was determined using the Bio-Rad DC Protein Assay (Bio-Rad Laboratories Ltd).

Western blotting

The similar procedure was applied for Western blot analysis of IL1R1, IL1R2, and IL1R accessory protein (IL1RAcP). Ten micrograms of protein for IL1R1 and 20 µg for IL1R2 and IL1RAcP were heated in a boiling bath for 5 min in 5 X SDS sample buffer (1.25 M Tris–HCl (pH 6.8), 50% glycerol, 25% β-mercaptoethanol, 10% SDS, and 0.01% bromophenol blue), separated by SDS-PAGE in 10% acrylamide linear gradient slab gels (IL1R1 and IL1R2) or in gradient slab gels ranging from 5 to 18% acrylamide (IL1RAcP), and transferred onto 0.2 µm nitrocellulose membranes (BioTraceNT; Pall Corporation, Ville St-Laurent, Québec, Canada) using electrophoretic transfer cell (Bio-Rad Laboratories Ltd). Recombinant soluble human (RSH) IL1R1, IL1R2, and IL1RAcP (R&D Systems Inc., Minneapolis, MN, USA) were used as positive controls.

Equal loading in each lane was confirmed by staining the blots with Ponceau S. (2%). Nitrocellulose membranes were then immersed in PBS containing 3% skimmed milk and 0.1% Tween 20 (blocking solution) for 1 h at room temperature and cut into strips. Membranes were incubated for 3 h at room temperature with a polyclonal goat anti-human IL1R1 (R&D Systems;
2 μg/ml in blocking solution), a polyclonal goat anti-human IL1R2 (R&D Systems; 2 μg/ml in blocking solution), or a polyclonal goat anti-human IL1RAcP (R&D Systems; 2 μg/ml in blocking solution). Equivalent concentrations of normal goat IgGs were used as controls. After six washes (5 min each in PBS/0.1% Tween 20), the strips were incubated for 1 h at room temperature with Fc-specific peroxidase-labeled rabbit anti-goat antibody (1:10,000 dilution in blocking solution; Jackson ImmunoResearch Laboratories Inc., West Grove, PA, USA). Membranes were finally washed and incubated for 1 min with an ECL Western blotting detection reagent (GE Healthcare Bio-Sciences Inc., Baie d’Urfé, Quebec, Canada) and exposed to Fujifilm for 2–15 min for optimal detection (all bands visible but not overexposed). The α-tubulin was used as a second control to ensure equal protein loading in each lane. Membranes were incubated for 1 h at room temperature with mouse anti-α-tubulin antibody (Sigma–Aldrich; 1:50,000 dilution in washing solution). After three washes (5 min each), the strips were incubated for 45 min at room temperature in Fc-specific peroxidase-labeled goat anti-mouse antibody (1:10,000 dilution in washing solution; Jackson ImmunoResearch Laboratories), washed six times (5 min each), incubated for 1 min with an ECL Western blotting detection reagent (GE Healthcare Bio-Sciences Inc.), and exposed to Fujifilm for 2–15 s for optimal detection (all bands visible but not overexposed).

**Immunofluorescence**

The same immunofluorescence procedure was applied for IL1R1, IL1R2, and IL1RAcP. Cells were seeded on eight-well sterile culture slides (5 × 10⁴ cells/well; BD Biosciences, Mississauga, Ontario, Canada), cultured overnight without stimulation, washed once with PBS, and fixed for 15 min at room temperature with PBS/3.7% formaldehyde. After washing with PBS, cells were incubated with PBS containing 1% Triton X-100 for 15 min at room temperature, washed in PBS, and incubated with a monoclonal mouse anti-human IL1R1 (R&D Systems; 10 μg/ml in PBS containing 0.2% bovine serum albumin (BSA) and 0.01% Tween 20), a monoclonal mouse anti-human IL1R2 (R&D Systems; 10 μg/ml in PBS/BSA/Tween 20), or a polyclonal goat anti-human IL1RAcP (R&D Systems; 10 μg/ml in PBS/BSA/Tween 20) for 1 h at room temperature in a humid chamber. For controls, cells were incubated with PBS/BSA/Tween 20 only. After washing with PBS containing 0.1% Tween 20 and subsequently with PBS alone, cover slips were incubated for 1 h at room temperature with a biotin-conjugated horse anti-mouse antibody (Vector Laboratories, Burlingame, CA, USA; 1:100 dilution in PBS/BSA/Tween 20) for IL1R1 and IL1R2 or with a biotin-conjugated rabbit anti-goat (H + L; Jackson Immuno Research Laboratories; 1:100 dilution in PBS/BSA/Tween 20) for IL1RAcP. Subsequently, culture slides were washed with PBS and incubated with 1% streptavidin–fluorescein isothiocyanate in PBS/BSA/Tween 20 for 1 h at room temperature in a humid chamber. After a final wash in PBS, samples were mounted in Mowiol containing 10% para-phenylenediamine (Sigma–Aldrich), an anti-fading agent, and observed under a Leica microscope (Leica Mikroskopie und Systeme, Wetzlar, Germany) equipped for fluorescence with a 100 W u.v. lamp and connected to an image analysis system (ISIS; Metasystems, Altlussheim, Germany).

**Zymography**

Gelatinase activity in samples of culture supernatants (40 μl) was analyzed by zymography on 7.5% SDS-polyacrylamide gels containing 0.5 mg/ml gelatin under non-reducing conditions as described previously (Collette et al. 2004, Bellehumeur et al. 2005). An equal volume of culture supernatant from human fibrosarcoma HT1080 cell line known for releasing elevated proteolytic activity was used as control in each gel (a gift from Dr Eric Petitclerc, Quebec City, PQ, Canada). Quantification of detectable gelatinases was achieved by computer assisted densitometry (Bioline, Visage 110s, Genomic Solutions Inc., Ann Arbor, Michigan, USA). Data were expressed as percentage of control (HT1080 gelatinases). Assays were performed thrice in duplicates.

**Statistical analysis**

Data followed a parametric distribution and were shown as means ± S.E.M. Comparison of two groups was performed using the unpaired t-test, whereas one-way ANOVA and the Bonferroni’s test post hoc on selected pairs were used for multiple comparisons, i.e. controls (NT) versus each cell line for each dose of IL1B. Statistical analyses were performed using GraphPad Software, Prism 4.0 (GraphPad Software, San Diego, CA, USA). Differences were considered as statistically significant whenever a P value <0.05 occurred.

**Results**

**Selection of cell clones**

In total, 15 clones with the empty plasmid vector (E clones) and 17 clones with the IL1R2 antisense
plasmid (A clones) were generated. IL1R2 concentration in culture supernatants and cell lysates from each of these clones were measured by ELISA, so as to select E clones showing IL1R2 expression similar to that of NT Ishikawas and A clones having weak or absent IL1R2 expression. Two E clones (E03 and E11) and two A clones (A08 and A17) were selected. As shown in Fig. 1A and B, IL1R2 concentrations in cell lysates and supernatants from E03 and E11 clones were equivalent to those of NT cells, whereas lower concentrations of IL1R2 in the cell lysates from A08 and A17 clones were noted (P<0.01). Only A17 showed lower levels of IL1R2 in the culture supernatant (P<0.05).

**Immunofluorescence analysis of IL1R1, IL1R2, and IL1RAcP expression**

IL1 receptors’ expression in E and A clones was also compared with that of NT Ishikawas by immunofluorescence. Representative photomicrographs exhibited in Fig. 2 show IL1R1, IL1R2, and IL1RAcP expression. Only IL1R2 immunostaining was reduced in A08 and A17 clones compared with NT cells, whereas no noticeable change in IL1R1 and IL1RAcP immunostaining was noted.

![Effect of IL1B on MMP-2 and MMP-9 secretion](image)

**Western blot analysis of IL1R1, IL1R2, and IL1RAcP expression**

To further evaluate IL1R1, IL1R2, and IL1RAcP expression, equivalent amounts of total cell protein extracts were analyzed by Western blotting. Western blot analysis of IL1R1 showed a 90 kDa band whose apparent molecular weight (MW) corresponds to the membrane-bound form of IL1R1 (mbIL1R1) as we previously reported (Bigonnesse et al. 2001), a generally faint 55 kDa band corresponding to the soluble form of the receptor (sIL1R1) and three other bands of 79, 73, and 69 kDa, which may correspond to degraded IL1R1 protein (Fig. 3A). Western blot analysis of IL1R2 showed a major band of 68 kDa, which sometimes appeared as doublet of 68/70 kDa, corresponding to the reported MW of mbIL1R2, and a doublet of 44/45 kDa, which may correspond to two forms of sIL1R2 as we reported previously (Akoum et al. 2001a; Fig. 3B). Western blot analysis of IL1RAcP (Fig. 3C) showed one band at ~61 kDa which corresponds to mbIL1RAcP. Pre-absorption of IL1R1, IL1R2, and IL1RAcP antibodies with an excess of rshIL1R1, rshIL1R2, and rshIL1RAcP respectively before incubation with blotted proteins markedly reduced the intensity of the detected above-described bands, thereby demonstrating specific binding (data not shown). For the same amount of total proteins, the intensity of both mbIL1R2 and sIL1R2 was clearly lower in the A clones when compared with NT cells, whereas no noticeable change in the levels of IL1R1 and IL1RAcP expression was noted.

**Effect of IL1B on MMP-2 and MMP-9 secretion**

MMPs play a major role in endometriosis pathophysiology (Osteen et al. 2003, 2004), and according to our data there is an increased production and secretion of MMP-9 in the eutopic endometrial tissue of women with endometriosis, and, interestingly, a relationship with decreased sIL1R2 levels (Collette et al. 2004, 2006, Bellehumeur et al. 2005). The secretion of MMP-2 (Fig. 4A) and MMP-9 (Fig. 4B) in E and A clones stimulated with varying concentrations of IL1B was therefore analyzed. Data shown in Fig. 4A showed that the basal level of MMP-2 secretion was significantly higher in A17 when compared with NT cells (P<0.01). Furthermore, a significant increase in MMP-2 in response to 1 ng/ml IL1B was observed in the A08 clone when compared with NT cells (P<0.05), whereas in the A17 clone a significant increase in MMP-2 secretion was observed at 0.1 and 1 ng/ml IL1B (P<0.05). MMP-9 secretion in the A08 clone was significantly increased at 0.1 and 10 ng/ml IL1B (P<0.05). However, in the A017, a statistically significant increase was observed at 10 ng/ml IL1B (P<0.05). No significant difference between E03 or E11 clone and NT cells was noted.

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Culture supernatants from NT Ishikawa cells and from E and A clones stimulated with varying concentrations of IL1B were then analyzed by gelatinolytic zymography. As shown in Fig. 5 (assays run under the same conditions), one distinct band of gelatinase activity at 72 kDa and corresponding to the latent form of MMP-2 (pro-MMP-2) was detected in the culture supernatants of NT and E clones. In the culture supernatants of A clones, the pro-MMP-2 band together with two additional 170 and 92 kDa bands were detected. The 170 kDa faint band might correspond to the putative dimeric MMP-9 pro-form (put-MMP-9) described by Watari et al. (1999) and identified by Western blotting in our previous studies (Collette et al. 2006), whereas the 92 kDa band is consistent with the latent form of MMP-9. Densitometric analysis of MMPs’ lysis bands showed that pro-MMP-2 band was significantly more intense in the A08 and A17 clones than in NT Ishikawa cells incubated with the culture medium alone ($P<0.05$ and $P<0.01$ respectively) or supplemented with 0.1 ($P<0.01$) and 1 ($P<0.01$) ng/ml IL1B. At 10 ng/ml IL1B, only A08 showed a statistically significant increase in pro-MMP-2 gelatinolytic activity when compared with NT cells ($P<0.05$). No statistically significant change in pro-MMP-2 activity was found in E03 and E11 clones when compared with NT cells was found, either with or without IL1B treatment (Fig. 6A). The gelatinolytic activity corresponding to pro-MMP-9 was detected only in the culture supernatants of A08 and A17 cells (Fig. 6B). However, no statistical analysis in comparison...
Reproduction cDNA in the antisense direction. The purpose of this pcDNA expression vector alone or containing IL1R2 endometrial cell line was stably transfected with the ectopic development, the Ishikawa adenocarcinoma on the functional modifications which can promote their To study the effects of IL1R2 deficiency in endometrial cells and 1 ng/ml IL1B (\(P<0.05\)). No statistically significant change in TIMP-1 secretion in NT cells or the E03, E11, and A08 clones was seen (Fig. 7).

**Effect of IL1B on TIMP-1 secretion**

Measurement of TIMP-1 concentrations in the culture supernatants by ELISA and statistical analysis of the data showed a diminution of TIMP-1 secretion in the A17 clone when compared with NT cells in response to IL1B, with statistically significant differences observed at 0.1 and 1 ng/ml IL1B (\(P<0.05\)). No statistically significant change in TIMP-1 secretion in NT cells or the E03, E11, and A08 clones was seen (Fig. 7).

**Discussion**

To study the effects of IL1R2 deficiency in endometrial cells on the functional modifications which can promote their ectopic development, the Ishikawa adenocarcinoma endometrial cell line was stably transfected with the pcDNA expression vector alone or containing IL1R2 cDNA in the antisense direction. The purpose of this transfection was to recreate the deficiency in IL1R2 that we found in endometrial cells of women with endometriosis and the resulting imbalance in cell responsiveness to IL1. In fact, a decreased expression of IL1R2, a specific downregulator of IL1 action, was shown in endometrial cells, either at the protein or the mRNA level (Boucher et al. 2001, Kharfi et al. 2002). This is in line with other studies showing that IL1A/IL1B exerts more pronounced effects on endometrial cell adhesion (Sillem et al. 1999), and that MMP-3 (stromelysin-1) expression is increased after exposure to IL1A/IL1B and may promote invasion and remodeling of the host tissue (Sillem et al. 2001). Lebovic et al. (2000) showed enhanced production of angiogenic molecules such as vascular endothelial cell growth factor and IL6 in response to IL1B in endometriotic cells when compared with endometrial cells from normal women. Therefore, IL1, a major pro-inflammatory and multifunctional cytokine, may play a central role in the inflammatory cascade associated with endometriosis and in propagating endometriotic implants through a pro-inflammatory stimulus and the synthesis of chemokines, growth, remodeling, and angiogenic factors (Akoum et al.)
In women with endometriosis, peritoneal macrophages, whose number and activation level were shown to increase in the peritoneal cavity of patients (Mori et al. 1992), secrete elevated levels of IL1B. Elevated concentrations of IL1B were found in the peritoneal fluid of women suffering from endometriosis (Mori et al. 1992, Taketani et al. 1992). Interestingly, recent studies showed an increased expression of IL1B in the ectopic endometrium of women with endometriosis (Bergqvist et al. 2001) which highlights the role of this cytokine in the pathophysiology of the disease.

All generated clones were first analyzed by ELISA in cell lysates and we report here a marked diminution in IL1R2 concentrations in two cell clones stably transfected with IL1R2 antisense when compared with NT cells or to cells transfected with the empty expression vector. It is noteworthy that transfection of the A clones was not always successful, and varied largely between the selected clones. Considering that the receptor can be cleaved and shed from the cell surface by the proteolytic action of matrix metalloproteases (Orlando et al. 1997), where sIL1R2 can prevent the interaction of IL1B with the signal-transducing IL1R1 (Colotta et al. 1993), IL1R2 ELISA in the cell-free supernatants was necessary and demonstrated a reduced release of IL1R2. Further analysis of IL1 receptors by immunofluorescence and

![Figure 5](https://example.com/figure5.png)

**Figure 5** Assessment of gelatinase activity by zymography. Non-transfected (NT), E03, E11, A08, and A17 cells were stimulated for 24 h with different concentrations of IL1B (0–10 ng/ml). Samples of cell-free culture supernatants were analyzed by zymography in gels impregnated with gelatin under non-reducing conditions. Representative zymograms showing lysis bands corresponding to the latent and active forms of MMP-2 and MMP-9. Only pro-MMP-2 band was detected in the culture supernatants of NT and E cells, whereas in addition to pro-MMP-2, two bands corresponding to putative (put)-MMP-9 and pro-MMP-9 were detected in the culture supernatants of A cells. Active MMP-2 and MMP-9 forms, which were obvious in the culture supernatant of HT-1080 cell line, were not observed in the culture supernatants of NT, E, or A cells.

![Figure 6](https://example.com/figure6.png)

**Figure 6** Densitometric analysis of MMP-2 (A) and MMP-9 (B) lysis bands following zymography. Data are means ± S.E.M. and expressed as percentage of control (corresponding HT1080 MMP-9 and MMP-2 bands). *, **Significant difference with NT cells for similar IL1B concentration (P<0.05 and P<0.01 respectively).
In the same way, it has been reported that MMP-2 (Wenzl & Heinzl 1998) and MMP-9 expression was increased in rheumatoid arthritis, tumor invasion, and endometriosis. An imbalance between MMP and TIMP expression has been involved in the invasive establishment of ectopic endometriotic lesions since they regulate MMP activity. An especially important in regulating extracellular matrix remodeling of the MMPs 2 and 9 and that of TIMP-1.

To IL1B may contribute to an imbalance in the expression levels and consequently increasing cell responsiveness to IL1B. In view of the pluripotent effects of IL1 and its well-documented role in endometriosis, the role of MMPs in tissue remodeling and the marked upregulation of these factors in ectopic and eutopic endometrial tissues of endometriosis women, our results showed that abnormal IL1R2 expression was associated with a significant increase in MMP-2 and MMP-9 secretion and a concomitant decrease in TIMP-1 secretion in response to IL1B. In view of the pluripotent effects of IL1 and its well-documented role in endometriosis pathophysiology, the role of MMPs in tissue remodeling and the marked upregulation of these factors in ectopic and eutopic endometrial tissues of endometriosis women, our in vitro data make plausible the role of IL1R2 deficiency in the acquisition by endometrial cells of endometriosis women of abnormal functions that may enable them to invade and implant into the host tissue. Delineation of the mechanisms underlying the capacity of endometrial cells to implant ectopically and respond differently to IL1 and identification of the resulting abnormal functions may be of interest for a better understanding of endometriosis pathophysiology and ultimately for the development of new diagnosis tools and targeted therapies.

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