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Gene expression profile of human endometrial receptivity: comparison between natural and stimulated cycles for the same patients

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Abstract
Background
The adjunction of exogenous hormones for controlled ovarian stimulation (COS) may alter endometrial receptiveness. In order to identify the genes mis-regulated under COS, we compared the endometrium gene expression profiles, from the same patients, in a natural cycle and in a subsequent COS cycle.

Methods
For the same normal-responder patients (n=21), endometrial biopsies (n=84) were collected during the pre-receptive (LH+2) and receptive stages (LH+7) of a natural cycle and, subsequently, on oocyte retrieval day (hCG+2) and on transfer day (hCG+5) of a stimulated cycle. Samples were analyzed using DNA microarrays. Gene expression profiles and biological pathways involved in the endometrial receptivity were analyzed.

Results
Although endometrium transition profiles from pre-receptive to receptive phases are similar between patients, COS regimens alter endometrial receptivity in comparison with natural cycle. Under COS conditions, two endometrial profiles were identified and were associated either with a moderately altered receptivity profile for the majority of the patients or a strongly altered profile for a subcategory of patients. The receptive endometrium transcription profile under COS was defective for biological functions such as TGFβ signaling, leukocyte transendothelial migration and the cell cycle. In addition, several biological functions specifically activated under COS regimens were identified.

Conclusion
Gonadotropin treatments in COS cycles led to disruptions of the transcriptional activation of genes involved in normal endometrial receptivity. We propose that when the receptiveness of the endometrium is seriously compromised by the COS protocol, fresh embryo replacement should be cancelled, the embryo frozen, and thawed embryo replacement should be performed under natural cycles.

MESH Keywords
Adult ; Biological Markers ; Biopsy ; Down-Regulation ; Endometrium ; cytology ; physiology ; Female ; Gene Expression Profiling ; Humans ; Menstrual Cycle ; genetics ; Oligonucleotide Array Sequence Analysis ; Ovulation Induction ; Sperm Injections, Intracytoplasmic

Author Keywords
human endometrium ; microarray ; endometrial receptivity ; natural cycle ; COS

Introduction
In assisted reproductive technology (ART), pregnancy and birth rates following IVF attempts remain low. Indeed, 2 out of 3 IVF cycles fail to result in pregnancy and more than 8 out of 10 transferred embryos fail to implant (Kovalevsky and Patrizio, 2005). Indeed, successful implantation requires a competent embryo, a receptive endometrium and a synchronized dialogue between maternal and embryonic tissues. Although the selection of competent oocyte, embryo and endometrial status assessment has experienced important technological progress, implantation rates are still low under COS for IVF (Donaghay and Lessey, 2007). It is assumed that two-thirds of implantation failures are associated with inadequate endometrial receptivity or with defects in the embryo-endometrium dialogue (Paulson et al., 1990; Simon et al., 1998).

Several parameters have been suggested for assessing endometrial receptivity, including endometrial thickness which is a traditional criterion, endometrial morphological aspect, and endometrial and sub-endometrial blood flow. However, their positive predictive value is still limited (Friedler et al., 1996; Pierson, 2003; Alcázar, 2006). More recently, transcriptomic approaches have been driven to identify bio-markers of the endometrial receptivity. Using microarray technology in human endometrial biopsy samples, several studies have
reported modifications related to the gene expression profile associated with the transition of the human endometrium from a pre-receptive (early-secretory phase) to a receptive (mid-secretory phase) state during a natural cycle (Carson et al., 2002; Riesewijk et al., 2003; Mirkin et al., 2005; Taibi et al., 2006; Haozsi et al., 2009). However, recent data suggested that COS regimens may have a negative impact on endometrial receptivity compared with natural cycles (Paulson et al., 1990; Horcajadas et al., 2005; Simon et al., 2005; Horcajadas et al., 2008; Liu et al., 2008). It has been suggested that the impairment of endometrial receptivity is due to high concentration of steroids resulting from COS (Krikun et al., 2005; Liu et al., 2008).

The divergence between the reports analyzing the effects of stimulation protocols on endometrial receptivity may have several explanations: differences in the day of the endometrial biopsies, different patient profiles, different COS protocols, and inadequate numbers of endometrial samples studied. In addition, no study has compared the early and the mid-secretory phases during a natural and stimulated cycle for the same patients, which seems an essential condition to minimize the impact of inter-patient variability.

The aim of this study was to evaluate the global gene expression profile of human endometrial biopsies of normal responder patients during the early-secretory phase and the mid-secretory phase of natural and stimulated cycles in the same patients.

**Materials and methods**

**Patient characteristics and endometrial biopsies**

The study population included 21 normal responders patients (age 30.9 years ± 3.3) referred for ICSI for male infertility factor recruited after written informed consent. All patients had normal serum FSH, LH and estradiol on day 3 and were normal-responders during a previous first ICSI attempt. This project has received institutional review board (IRB) approval. During the same natural cycle that preceded a second ICSI attempt, two endometrial biopsies were obtained in all women at day 2 (LH2) and day 7 (LH7) after the LH peak and two others were obtained at day 2 (hCG2) and day 5 (hCG5) after hCG administration during COS cycles. The LH surge was estimated by the patient herself according to the first day of her menstruation. Histologic analysis was not performed to verify that the LH timing was accurate. Therefore, the possibility for a delay of one day from the first day of the menstruation cannot be excluded. Each biopsy sample was frozen at ~80°C in RLT RNA extraction buffer (RNeasy kit, Qiagen, Valencia, CA, USA).

**Complementary RNA (cRNA) preparation and microarray hybridization**

Endometrial samples RNA were extracted using the micro RNeasy Kit (Qiagen). The total RNA quantity was measured with a Nanodrop ND-1000 spectrophotometer (Nanodrop Technologies Inc., DE, USA) and RNA integrity was assessed with an Agilent 2100 Bioanalyzer (Agilent, Palo Alto, CA, USA). cRNA was prepared with two rounds of amplification according to the manufacturer's protocol “double amplification” (Two-Cycle cDNA Synthesis Kit, Invitrogen) starting from total RNA (100 ng). Labeled fragmented cRNA (12 μg) was hybridized to oligonucleotide probes on Affymetrix HG-U133 Plus 2.0 arrays containing 54,675 sets of oligonucleotide probes (“probe set”) which correspond to ≈ 30,000 unique human genes or predicted genes. Each endometrial sample was put individually on a microarray chip.

**Data processing**

Scanned GeneChip images were processed using Affymetrix GCOS 1.4 software to obtain an intensity value signal and a detection call (present, marginal or absent) for each probe set, using the default analysis settings and global scaling as the first normalization method, with a trimmed mean target intensity value (TGT) of each array arbitrarily set to 100. Probe intensities were derived using the MAS5.0 algorithm. This algorithm also determines whether a gene is expressed with a defined confidence level or not (“detection call”). This “call” can either be “present” (when the perfect match probes are significantly more hybridized than the mismatch probes, FDR < 0.04), “marginal” (for FDR > 0.04 and <0.06) or “absent” (FDR > 0.06). The microarray data were obtained in our laboratory in agreement with the Minimal Information about Microarray Experiment MIAME recommendations.

**Microarray data analysis**

The Significant Analysis of microarrays (SAM, Stanford University, USA, Tusher et al., 2001) was used to identify genes whose expression varied significantly between the LH+2 (n=21) and LH+7 (n=21) samples and between the hCG+2 (n=21) and hCG+5 (n=21) samples. SAM provides mean or median fold change values (FC) and a false discovery rate (FDR) confidence percentage based on data permutation. To perform the comparison of gene expression profiles between endometrial sample groups, a probe set selection using the Absent/ Present “detection call” and a coefficient of variation (CV) (≥ 40%) between samples was first performed before the SAM.

To compare profile expressions of endometrial samples (n=84) from the LH+2, LH+7, hCG+2 and hCG+5 sample groups, we performed an unsupervised classification with both principal component analysis (PCA) and hierarchical clustering (Eisen et al., 1998; de Hoon et al., 2004). The PCA involved original scripts based on the R statistics software through the RAGE web interface (http://rage.montp.inserm.fr) (Rême et al., 2008). Hierarchical clustering analysis based on the expression levels of varying probes were performed with the CLUSTER and TREEVIEW software packages. Genetic expression profiles were analyzed with the RAGE supervised
analysis module, using a non parametric Mann Whitney U test with multiple testing corrections and were confirmed with the SAM software. Selected gene lists (mean fold change $>2$ and FDR $<5\%$) were submitted to Ingenuity (http://www.ingenuity.com) and Fatigo+ (http://babelomics2.bioinfo.cipf.es) software to identify the biological mechanisms altered by these gene expression variations. Fatigo+ is a part of the Babelomics suite and allows any differential distribution of biological terms (Gene Ontology, KEGG pathways) to be found between two groups of genes (Al-Shahrour et al., 2006).

**Ingenuity Pathway Analysis**

Selected gene lists were analyzed using Ingenuity Pathway Analysis (IPA) software (Ingenuity Systems, Redwood City, CA). Those genes with known gene symbols and their corresponding expression values were uploaded into the software. Each gene symbol was mapped to its corresponding gene object in the Ingenuity Pathways Knowledge Base. Networks of these genes were algorithmically generated based on their connectivity and assigned a score. The score is a numerical value used to rank networks according to how relevant they are to the genes in the input dataset but may not be an indication of the quality or significance of the network. The score takes into account the number of focus genes in the network and the size of the network to approximate how relevant this network is to the original list of focus genes. The network identified is then presented as a graph indicating the molecular relationships between genes/gene products. Genes are represented as nodes, and the biological relationship between two nodes is represented as an edge (line).

**Results**

**Gene expression profile of endometrial receptivity during natural and stimulated cycles for the same patients**

We selected 18,389 probe sets based on a CV $\geq40\%$ and a “detection call” present in at least 10 samples per group, and then subjected the 84 samples to an unsupervised clustering (Figure 1A). A first branch allowed the separation of the pre-receptive (LH+2 and hCG+2) from the receptive sample groups (LH+7 and hCG+5), corresponding to 62, 57, 81 and 100$\%$ of the samples for LH+2, hCG+2, LH+7 and hCG+5 respectively. A sub-branch allowed to partition the receptive samples between the natural (LH+7) and stimulated (hCG+5) cycles. A three dimensional PCA was performed (4,237 probe sets), representing 43$\%$ of the data information, and confirmed the unsupervised cluster analysis (Figure 1B).

A SAM analysis between the LH+2 and the LH+7 and between the hCG+2 and the hCG+5 sample groups (LH+2 versus LH+7; hCG+2 versus hCG+5; paired sample analysis) were performed. Among 1071 genes significantly modulated between the pre-receptive and receptive samples during natural cycle, 93$\%$ and 7$\%$ were respectively up- (2 $\leq$ fold change $\leq465$) and down-regulated ($-3.8 \leq$ fold change $\leq -2$) in the LH+7 sample group (fold change $\geq2$ and a FDR $<5\%$) as previously described (Haouzi et al., 2009) (Supplementary Table I and Figure 4). Analysis of samples during stimulated cycle revealed that 998 genes were significantly modulated between the pre-receptive and receptive stages, including 78$\%$ up-regulated (2 $\leq$ fold change $\leq 59$) and 22$\%$ down-regulated ($-5 \leq$ fold change $\leq -2$) genes in the hCG+5 sample group (Figure 2A, Figure 4 and Supplementary Table I). A hierarchical clustering with this significant list of genes allowed the separation of the two endometrial sample groups of the stimulated cycle (Figure 2B). Interestingly, more down-regulated genes were observed during endometrial receptivity of the stimulated cycle in comparison with the natural cycle.

A SAM analysis between the LH+2 and hCG+2 and between the LH+7 and hCG+5 sample groups (LH+2 versus hCG+2, LH+7 versus hCG+5, paired sample analysis) was performed. The number of significantly modulated genes is reported in Table I. In contrast with other studies, the comparison of gene expression profile between natural and stimulated cycles was performed by paired samples (hCG+2 versus their corresponding LH+2), which is an important point to minimize inter-patient variability. For example, the number of genes significantly modulated between LH+2 and hCG+2 samples analysed by paired or unpaired was dramatically different, highlighting the need to perform paired analyses and explaining differences with other studies.

**Atypical gene expression profile in endometrium under stimulated cycles**

The hierarchical clustering based on the stimulated cycle endometrium signature (Figure 2B) revealed that 14$\%$ (3/21 patients) of the hCG+5 samples were misclassified. Interestingly, the same samples also displayed an altered expression profile at day hCG+2 (Figure 3 and 2B). Hence, gene expression analysis of the endometrium transcriptome after COS can distinguish a “standard” profile shared by most patients, and an “atypical” profile found in 14$\%$ of patients. A SAM analysis performed between the hCG+2 and hCG+5 samples reveals that only one gene, TSPAN3 (x39.3, FDR<0.0001) was significantly modulated between these two sample groups in these three “atypical” patients compared to 1296 genes which were modulated in the 18 “standard” patients. This gene was not modulated during the transition from the pre-receptive to the receptive stages in natural nor in the 18 “standard” patients of the stimulated cycles. We also performed a SAM analysis between the hCG+2 of the 18 “standard” and the 3 “atypical” samples. 197 genes were significantly modulated, including 188 up- and 9 down-regulated genes, suggesting a strong difference in gene expression at hCG+2 between “standard” and “atypical” samples. Among these genes, 24 up- and 2 down-regulated genes were specifically restricted to the hCG+2 “atypical” profile. In addition, the SAM analysis between the “standard” hCG+5 sample group and the “atypical” hCG+5 sample group revealed that 14 genes were significantly modulated, among which, 4 up- and 1 down-regulated genes were restricted to the “atypical” samples.
Differential expression of biological functions between natural and stimulated cycles

The two gene lists identified with SAM analyses between the pre-receptive and receptive samples from the natural and stimulated cycles were then intersected to determine their overlap. While 361 up- and 32 down-regulated genes were in common to the natural and stimulated cycles, each category displayed a specific gene expression profile (Figure 4). These specific profiles included 632 up- and 46 down-regulated genes during the natural cycle, 416 up- and 189 down-regulated genes during the stimulated cycle. Interestingly, during the endometrial receptivity of the stimulated cycles, numerous genes which were down-regulated compared to natural cycles, were articulated around the cell cycle function (Figure 5). Among this function, the most representative genes were those coding for Cyclins, Cell Division Cycle (CDC) members, Cyclin Dependent Kinases (CDK), and members of the E2F family of transcription factors. They included: CCNA2 (x-2.7, FDR<0.0001), CCNB1 (x-2.3, FDR<0.0001), and CCNE2 (x-2.1, FDR<0.0001), CDC25C (x-3.3, FDR<0.0001), CDC20 (x-2.8, FDR<0.0001), CDC7 (x-2, FDR<0.0001), CDKN3 (x-2.9, FDR<0.0001), E2F1 (x-2.1, FDR<0.0001), E2F7 (x-2.9, FDR<0.0001), E2F8 (x-3.2, FDR<0.0001), as well as FOXM1 (x-2.1, FDR<0.0001), MYBL2 (x-2.8, FDR<0.0001), TK1 (x-2.2, FDR<0.0001), PAX5 (x-3.2, FDR<0.0001), CHAF1B (x-2.2, FDR<0.0001), and CDKN2A (x-2.9, FDR<0.0001).

KEGG pathways between natural and stimulated cycles

We performed a FatiGO+ analysis with the two lists of up-regulated genes during endometrial receptivity of the natural and stimulated cycles. There were numerous differences, between the natural and stimulated cycles, in main systems involved in the implantation process, such as the TGFβ signaling pathway, the complement and coagulation cascades, and leukocyte transendothelial migration (Figure 6). While CHRD (x2.4, FDR<0.0001), BMP6 (x3.5, FDR<0.0001), DCN (x2, FDR=0.02), FST (x8.7, FDR<0.0001), ACVR1B (x2.9, FDR=0.02) and SMAD3 (x2.4, FDR=0.003) were the most representative up-regulated genes around the TGFβ signaling pathway in the natural cycle from the pre-receptive and receptive stages, there were no changes in these genes the stimulated cycle (Figure 6A). Also, MASP1 (x2.2, FDR=0.004), C2 (x3.6, FDR<0.0001), C3 (x4.7, FDR<0.0001), SERPINF2 (x2.2, FDR=0.0006), FGB (x46.4, FDR=0.005), PROS1 (x2.9, FDR=0.0006), and BDKRB2 (x2.1, FDR=0.006) were the major up-regulated genes of the complement and coagulation cascades in the natural cycle, while BDKRB1 (x2.6, FDR=0.008) was the only gene associated with this pathway to be up-regulated during receptivity in the stimulated cycles (Figure 6B). Numerous genes involved in leukocyte transendothelial migration were up-regulated during endometrial receptivity of the natural cycle, such as ITGB2 (x3.1, FDR<0.0001), ITGB3 (x7, FDR=0.002), ITGB4 (x2.6, FDR<0.0001), ITGA3 (x2.2, FDR<0.0001), ITGAL (x2.6, FDR<0.0001), ITGAM (x2, FDR=0.005), CD99 (x3.3, FDR=0.001), VCAM1 (x4.2, FDR=0.003), VIL2 (x2.5, FDR<0.0001) and RAC2 (x2.6, FDR=0.0007). Whereas, only ITGA4 gene (x2.6, FDR=0.003) was up-regulated during endometrial receptivity of the stimulated cycle (Figure 6C).

Biomarkers of the endometrial receptivity during stimulated cycles

We also performed a PCA analysis of the hCG+2 and hCG+5 samples with the predictor list of endometrial receptivity as previously described (Haouzi et al., 2009). The majority of the hCG+2 samples were separated from the hCG+5 samples, except for the three “atypical” samples, in this PCA using the first two dimensions representing 39% of the data information (Figure 7). During the pre-receptive to the receptive transition in stimulated cycles, four of our biomarkers of endometrial receptivity were significantly up-regulated. For natural cycles compared to stimulated cycles, NLF2 was increased by a factor 12.6 compared to 2, PROK1 was increased by a factor of 20.4 compared to 11.1. The upregulation observed in the microarray analyses was validated by QRT-PCR (Haouzi et al., 2009; Supplementary Figure 1).

ICSI outcome and biomarker genes

Seven ongoing pregnancies were obtained in the patients participating in this study. There were no significantly differences in the modulated genes between the pregnant and non pregnant patients either in the hCG+2 or hCG+5 subgroups.

Discussion

This is the first study analyzing by paired samples the endometrial gene expression profiles from the same patients during the pre-receptive to the receptive transition both in a natural and in a subsequent stimulated cycle. Although endometrium transition profiles are similar between patients, COS regimens affect endometrial receptivity in comparison with natural cycles. Two endometrium genes profiles were observed and associated either with a moderately altered receptivity for the majority of the patients or a strongly altered receptivity under COS conditions.

Differential gene expression profiles of endometrial receptivity between natural and stimulated cycles

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In natural as well as stimulated cycles, the majority of the genes modulated during endometrial receptivity were up-regulated (93% in the natural cycle compared to 78 % in the stimulated cycle), suggesting that the receptive endometrium requires transcriptional activation as suggested by Horcajadas et al. (2008). Among the up-regulated genes of endometrial receptivity in the stimulated cycles, only 46 % were in common with those of the natural cycles. These data suggest that either the duration or FSH dose in gonadotropins treatment under COS cycles leads to the transcriptional activation of other genes which are not involved in physiological endometrium receptivity. Major differences in biological functions known to be involved in the implantation process, such as the TGFβ signaling pathway (Jones et al., 2006), the complement and coagulation cascades and the leukocyte transendothelial migration (Berlin et al., 1995; Ivetic et al., 2004; Dominguez et al., 2005; da Costa Martins et al., 2006), were observed between the natural and stimulated cycle. The up-regulation of genes involved in these signaling pathways was found in natural cycles, but was lacking in stimulated cycles. On the other hand, the expression profile of the receptive endometrium in stimulated cycles during COS showed more down-regulated genes than in the natural cycle (7% in natural compared to 22 % in stimulated cycles respectively), suggesting that COS treatment, partly, induced gene repression. The majority of these down-regulated genes was not observed in the natural cycle (2% in common) and were related to the 'cell cycle' function, suggesting a negative impact of gonadotropins treatment during COS on this function. Several cyclins (cyclin A2, B1 and E2), cell-division cycles (CDCA5, A8, A7, 20 and 25C), and members of E2F family (E2F1, 7 and 8), which are key regulators of cell cycle, as well as numerous genes articulated around them, were not upregulated in stimulated cycles. Moreover, genes involved in checkpoint regulation, which are molecular circuits monitoring DNA integrity and cell growth prior to replication and division at the G1/S and G2/M transitions respectively, were also not upregulated in stimulated cycles (Branzei and Foiani, 2008). These data can be considered in conjunction with reports which show that both GnRH antagonist and agonist exert an inhibitory effect on the cell cycle of human endometrial cancer cells in vitro, probably by decreasing the synthesis of growth factors (Emons et al., 1993; Hershkovitz et al., 1993; Kleinman et al., 1993).

In contrast to Horcajadas et al. (2005) who showed that genes up-regulated during the formation of the receptive endometrium in the natural cycle tended to be down-regulated during stimulated cycles, we report similar variations but with a lesser amplitude during COS in comparison with natural cycles (Supplementary Tables I and II). One possible explanation is that, this amplitude difference reflects the sampling time of the receptive samples between natural (LH+7) and stimulated cycles (hCG+5). However, as embryos transfer was generally performed on day five after hCG administration (hCG+5) of a stimulated cycle, endometrial biopsies just before embryo transfer fitted with clinical practice. Recently, it has been reported that there is a two-day delay at hCG+7, with hCG+7 samples having a gene expression profile close to day 5 of stimulated cycle samples (Horcajadas et al., 2008). However, in the current study, the unsupervised cluster and PCA analyses showed that a proportion of LH+7 (67%) and hCG+5 (30%) samples tend to cluster close together. In addition, the study design (LH+7/9 compared to hCG+7/9 in the Horcajadas’s study, LH+2 and LH+7 compared to hCG+2 and hCG+5 in the present study), the size of patient samples (n=5 per group in the Horcajadas’s study, n=21 per group in the present study) as well as the samples, which were obtained from different fertile patients candidates for oocytes donation in the Horcajadas’s study, could contribute to these diverging results (Horcajadas et al., 2008). By contrast, in the present study the comparison of gene expression profiles between natural and stimulated cycles was performed in the same normal-responder patients, which is an important point to minimize inter-patient variability.

We can wonder about the impact of multiple biopsies performed on the same patient. Kalma et al. (2008) reported that the first endometrial biopsy modulated a wide variety of genes in the same cycle as well as in the following cycle, which can consequently mask true gene expression profile differences. However, all patients were subjected similarly to four biopsies. If biopsies affect all samples equally, so this should not prevent the identification of different gene expression profiles. Therefore, gene expression profile differences observed during the pre-receptive to the receptive endometrial transition under COS regimens (hCG+2 compared to hCG+5) between the two described endometrial profiles (endometrial receptivity moderately and strongly altered) suggested, contrariwise, that true differences are not masked. In addition, although Barash et al. (2003) showed that multiple endometrium biopsies during the spontaneous menstrual cycle increased implantation and pregnancy rates at the following cycle of treatment, and that Zhou et al. (2008) demonstrated that local injury of the endometrium executed in IVF patients during their cycle of treatment, before ovum retrieval, gains the same effect of increasing implantation and clinical pregnancy rates, we don’t known the consequences to perform a biopsy on the embryo transfer day. However, in this cohort, pregnancy rate (7/21 patients) was similar to those habitually acquired, suggesting a no detrimental effect on implantation process.

Atypical gene expression profile of endometrial cells during COS

The comparisons of gene expression from the same patients between natural and stimulated cycles revealed endometrial profiles associated either with a moderately altered receptivity in most cases (86%) or a strongly altered receptivity during the COS protocol in a few cases (14%). In these few cases, the transition from the oocyte collection day (hCG+2) to the embryo transfer day (hCG+5) phase introduced only minor differences in gene expression profiles. Only one gene, TSPAN3 was significantly modulated between their pre-receptive to receptive samples. The protein encoded by this gene is a member of the tetraspan family and plays a role in the regulation of the cell development, activation, growth and motility (Hemler, 2005). Impairment of gene expression of endometrial cells
was already apparent at hCG+2 in these samples as judged by the gene number significantly modulated (197 genes) between these samples and those associated with a moderately altered receptivity. However, on hCG+5, there were fewer differences between the two endometrial profiles.

We previously described a predictor list (comprising 60 probe sets) of endometrial receptivity during natural cycle (Haouzi et al., 2009). These predictors confirmed the presence of the two endometrial profiles: either receptive or not receptive. However, among these predictors, only four were significantly up-regulated during the pre-receptive to the receptive transition in the stimulated cycle. In addition, the fold changes between the pre-receptive to the receptive phases in endometrial profiles associated with a positive receptive status were less pronounced in stimulated than in natural cycles, suggesting incomplete receptivity during COS. Therefore, our predictors further suggest the moderately altered receptivity status for the majority of the samples, as well as the strongly altered receptivity status for a few patients. When the receptiveness of the endometrium is seriously compromised during COS, fresh embryo replacement could be reconsidered. In these cases, embryos freezing enabling the IVF attempt to be saved and the embryo transfer can be done later during a natural cycle. For these patients, the endometrial receptivity gene expression signature is of high value (Haouzi et al., 2009).

In conclusion, the transcriptomic pattern of endometrial cells in natural and stimulated cycles in the same patients reveals either moderate or strong alterations of endometrial receptivity under COS protocols. This information could open new perspectives, particularly in patients with multiple implantation failures. In this case, analysis of the endometrial profile could reveal a strongly altered profile during COS protocols, prompting the clinician to either adapt the IVF stimulation protocol or to perform embryo transfer later during a natural cycle.

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Unsupervised classification with both principal component analysis (PCA) and hierarchical clustering of 84 endometrium samples during the pre-receptive and receptive stages from natural and stimulated cycles.

(A) Unsupervised hierarchical clustering was performed, allowing a separation between the pre-receptive (LH+2 and hCG+2) and receptive samples (LH+7 and hCG+5). (B) PCA using the three dimensions was performed representing 43% of the data information, and confirmed the unsupervised cluster analysis. Pink circle, LH+2 and hCG+2 samples; grey circle, LH+7 samples; green circle, hCG+5 samples.
Figure 2
Supervised hierarchical clustering of hCG+2 and hCG+5 samples from stimulated endometrium cycles
(A) The majority of the genes specifically modulated during endometrial receptivity in stimulated cycles (998 genes) were up-regulated (777 up-regulated genes, 221 down-regulated genes). Red, up-regulated genes; green, down-regulated genes. (B) A hierarchical clustering was performed with this gene list, allowing the separation of the two sample groups. Rhomb black outline: "Atypical" samples.

Figure 3
The two endometrium profiles during the pre-receptive and the receptive stages in stimulated cycles
The majority of patients (18/21 patients) were characterized by a common profile, called "standard profile". Other patients (3/21) presented an "atypical profile". PR, pre-receptive; R, receptive; red, up-regulated genes; green, down-regulated genes.
**Figure 4**
Venn diagram of transcripts up-regulated and down-regulated during endometrial receptivity in the natural cycle compared to the stimulated cycle
Each list of genes was determined with the SAM software with a FDR < 5% and a fold change > 2.

**Figure 5**
Down-regulated genes related to the cell cycle function in the receptive endometrium of stimulated cycles
Genes involved in (A) the cell cycle, (B) G1/S checkpoint regulation, and (C) the G2/M DNA damage checkpoint. All of these genes were down-regulated in stimulated cycles in comparison with natural cycles. Green: down-regulated genes.
Figure 6
Major differences in main systems involved in implantation between natural and stimulated cycles
To identify differential distributions of KEGG pathways, we performed a FatiGO+ analysis with the two lists of genes up-regulated gene during endometrial receptivity of the natural and stimulated cycles. There were numerous differences, between the natural and stimulated cycles, in main systems involved in the implantation process, such as the TGFβ signaling pathway, the complement and coagulation cascades, and leukocyte transendothelial migration. Red, genes up-regulated in natural cycles; pink, genes up-regulated in stimulated cycles.

Figure 7
Unsupervised classification with principal component analysis (PCA) of the hCG+2 and hCG+5 samples with the predictor list
Our predictor list of the endometrial receptivity allows the separation of 90% of hCG+5 samples from the hCG+2 samples in this PCA using the first two dimensions, representing 39% of the data information.
### Table I
Number of genes significantly modulated in 6 microarray analyses comparing the pre-receptive or receptive stages between natural and stimulated cycles.

<table>
<thead>
<tr>
<th>Study</th>
<th>Number of samples</th>
<th>Natural cycle (number of samples)</th>
<th>Stimulated cycle (number of samples)</th>
<th>Paired analysis</th>
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<tr>
<td>Mirkin et al. (2004)</td>
<td>13</td>
<td>LH+8 (n=5)</td>
<td>hCG+9 Atg (n=5)</td>
<td>Partially</td>
<td>≥ 1.19</td>
<td>6</td>
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<tr>
<td></td>
<td></td>
<td>LH +8 (n=5)</td>
<td>hCG+9 Ag (n=3)</td>
<td></td>
<td>≥ 1.2</td>
<td>5 1</td>
</tr>
<tr>
<td>Horcajadas et al. (2005)</td>
<td>19</td>
<td>LH +7 (n=14)</td>
<td>hCG +7 (n=5)</td>
<td>No</td>
<td>≥ 3</td>
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<td></td>
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<td>277</td>
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<td>Simon et al. (2005)</td>
<td>28</td>
<td>LH +7 (n=14)</td>
<td>hCG +7 Atg standard dose (n=4)</td>
<td>No</td>
<td>≥ 2</td>
<td>22 69</td>
</tr>
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<td></td>
<td></td>
<td>LH +7 (n=14)</td>
<td>hCG +7 Atg high dose (n=5)</td>
<td></td>
<td>88 24</td>
<td>22 100</td>
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<td></td>
<td>LH +7 (n=14)</td>
<td>hCG +7 Ag (n=5)</td>
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<tr>
<td>Horcajadas et al. (2008)</td>
<td>49</td>
<td>LH+1 (n=5)</td>
<td>hCG+1 (n=5)</td>
<td>No</td>
<td>-</td>
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<tr>
<td></td>
<td></td>
<td>LH+2 (n=5)</td>
<td>hCG+2 (n=5)</td>
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<tr>
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<td></td>
<td>LH+3 (n=5)</td>
<td>hCG+3 (n=5)</td>
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<tr>
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<td></td>
<td>LH+5 (n=5)</td>
<td>hCG+5 (n=4)</td>
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<td></td>
<td></td>
<td>LH+7 (n=5)</td>
<td>hCG+7 (n=5)</td>
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<td>69 73</td>
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<tr>
<td></td>
<td></td>
<td>LH+9 (n=5)</td>
<td>hCG+9 (n=5)</td>
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<tr>
<td>Liu et al. (2008)</td>
<td>13</td>
<td>LH+7 (n=5)</td>
<td>hCG+7 high serum E2 levels (n=4)</td>
<td>No</td>
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<td>LH+7 (n=5)</td>
<td>hCG+7 low serum E2 levels (n=4)</td>
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<td>Current study</td>
<td>84</td>
<td>LH+2 (n=21)</td>
<td>hCG+2 (n=21)</td>
<td>Yes</td>
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<td>LH+7 (n=21)</td>
<td>hCG+5 (n=21)</td>
<td></td>
<td>657 0</td>
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</table>

This table compares our gene lists significantly modulated between LH+2 and hCG+2, and between LH+7 and hCG+5 with five other studies using the same approach. Atg, antagonist; Ag, agonist.