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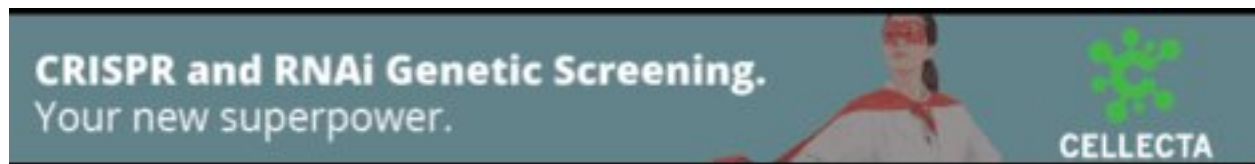
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## Research

# Single-cell transcriptomics of the human placenta: inferring the cell communication network of the maternal-fetal interface

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Organismal function is, to a great extent, determined by interactions among their fundamental building blocks, the cells. In this work, we studied the cell-cell interactome of fetal placental trophoblast cells and maternal endometrial stromal cells, using single-cell transcriptomics. The placental interface mediates the interaction between two semiallogenic individuals, the mother and the fetus, and is thus the epitome of cell interactions. To study these, we inferred the cell-cell interactome by assessing the gene expression of receptor-ligand pairs across cell types. We find a highly cell-type-specific expression of G-protein-coupled receptors, implying that ligand-receptor profiles could be a reliable tool for cell type identification. Furthermore, we find that uterine decidual cells represent a cell-cell interaction hub with a large number of potential incoming and outgoing signals. Decidual cells differentiate from their precursors, the endometrial stromal fibroblasts, during uterine preparation for pregnancy. We show that decidualization (even *in vitro*) enhances the ability to communicate with the fetus, as most of the receptors and ligands up-regulated during decidualization have their counterpart expressed in trophoblast cells. Among the signals transmitted, growth factors and immune signals dominate, and suggest a delicate balance of enhancing and suppressive signals. Finally, this study provides a rich resource of gene expression profiles of term intravillous and extravillous trophoblasts, including the transcriptome of the multinucleated syncytiotrophoblast.

[Supplemental material is available for this article.]

The long duration of eutherian fetal development requires a considerable level of negotiation between fetal and maternal needs and capacities. The key locus of this negotiation is between trophoblast cells of the placenta and the endometrium of the maternal uterus, in addition to hormonal signals. Eutherian trophoblasts are ancestrally invasive (Wildman et al. 2006), and it is likely that even secondarily noninvasive placentation, such as that of hoofed animals or lemurs, is due to the evolution of a less permissive uterus, rather than the evolution of less invasive trophoblasts (D'Souza and Wagner 2014). Differentiation of endometrial stromal fibroblasts to form the decidua, which accepts the implanting conceptus, is an additional evolutionary novelty to accommodate trophoblast invasiveness also found in humans (Wagner et al. 2014).

Maternal-fetal interactions are also central in human pregnancy. In addition to standard communication between adjacent cells within an individual, the maternal-fetal interface also inte-

grates two semiallogenic individuals, the fetus and the mother. The manifold functions in negotiating maternal and fetal interests (e.g., nutrient and gas exchange, anchoring, immunity) are reflected in heterogeneous placental structure, encompassing many distinct cell types. Placental cell fate is determined in early eutherian development, as the outer layer of the blastocyst, the trophoctoderm, is the precursor of placental tissue. Enveloped by the trophoctoderm is the inner cell mass, which gives rise to the embryo proper and further extraembryonic tissues, like the yolk sac, the amnion, and the allantois. Following the implantation of the blastocyst into decidualized endometrium, a specialized population of placental trophoblasts, extravillous trophoblasts (EVTs), invade the maternal decidua and vessels and thereby generate lacunas filled with maternal blood in which the developing surface-enlarging fetal villi become bathed. EVT migrate out of the

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anchoring villi into maternal endometrium and partially into myometrium. A subset of EVT is involved in remodeling maternal spiral arteries, thereby acquiring endothelial character, while another subset fuse to form the placental bed, and yet another set is involved in the uterine gland remodeling (Ji et al. 2013; Maltepe and Fisher 2015). The placental chorionic villi, which are bathed in maternal blood, contain fetal blood vessels and are covered by a continuous multinucleated layer of syncytiotrophoblast. This layer arises and is maintained through pregnancy by the fusion of the underlying cytotrophoblasts and represents an alternative differentiation fate to EVT. The syncytiotrophoblast is in direct contact with maternal blood and is the main interface between maternal and fetal circulation.

Implantation outside of the decidualized uterine region is deeper, often reaching the myometrium (i.e., placenta accreta) and can be fatal to the mother at birth (Hannon et al. 2012). In addition to forming a maternal barrier against invasive trophoblast, decidualization has been suggested to actively control trophoblasts. For example, the invasiveness and growth of cultured trophoblast cells is decreased in conditioned medium from decidual cells (Lewis et al. 1993; Zhu et al. 2009; Godbole et al. 2011). Decidual effects are not uniformly suppressive; rather they involve fine-tuned interactions via numerous pathways (Knofler 2010). For example, a number of decidual growth factors secreted starting at implantation likely support trophoblast invasion. Similarly, the maternal immune system is not simply suppressed at the introduction of the allograft but rather modulated to add active support (Knofler and Pollheimer 2012; Erlebacher 2013; Fock et al. 2013). Given the pervasive evidence for mutual interaction between maternal and fetal tissues, the resulting utero-placental phenotype is best conceptualized as a unit resulting from a rich network of developmental interactions.

Identification of cell-cell crosstalk requires examining maternal and fetal cells jointly. This cannot be done in healthy human pregnancy, and therefore most questions are addressed either on the term placenta alone, or in maternal or fetal cell culture. The utility of animal models for this purpose is often questioned, as the evolutionary relationships, and therefore homology between placental cell lineages across species, are not well understood (Mess 2014; Maltepe and Fisher 2015). Here, we approach this question by inferring the cell interactome of trophoblast cells and maternal decidua from single-cell transcriptomes, complemented with laser microdissected syncytiotrophoblast and cell culture of endometrial stromal cells. While the inferred cell communication network remains hypothetical, the coexpression of ligands and receptors in adjacent cells is stronger evidence for functional importance than the gene expression of a single cell type. In combination with downstream gene expression effects of the signaling, this information promotes rich functional insights, because cell interactions are part of the core biological function of these tissues (Semrau and van Oudenaarden 2015). In addition, the high level of resolution in the single-cell transcriptome allows identification of potentially important rare cell types, whose signal is too diluted to be detected in tissue-level transcriptomes (Buettner et al. 2015; Crosetto et al. 2015; Stegle et al. 2015).

## Results

Sampling the villous tissue of two human term placentas resulted in 87 single-cell transcriptomes, which grouped into five distinct clusters (Fig. 1; Supplemental Fig. S6). Of these, three large clusters consisted of 23, 28, and 28 cells, respectively, while sets of three

and five cells represent the remaining two small clusters. We identified the three large and one small cluster as belonging to different types of placental trophoblasts and the remaining small cluster as maternal immune cells. These data were complemented by two transcriptomes of syncytiotrophoblast, collected from a single placenta by laser microdissection, the transcriptomes of primary undifferentiated endometrial stromal fibroblast (ESF), and the transcriptome of in vitro differentiated primary decidual cells from two patients. The cells included in this study thus represent adjacent maternal (decidua) and fetal (syncytiotrophoblast and extravillous trophoblast) cells of utero-placental interface, as well as the precursors of decidualized cells and syncytiotrophoblast (ESF and cytotrophoblast). This approach enables identification of putative interactions and their development. Finally, we generated two tissue-level transcriptomes of human term placenta for comparison. All transcriptomes were normalized as TPM (transcripts per million) to the same set of human coding genes prior to comparison.

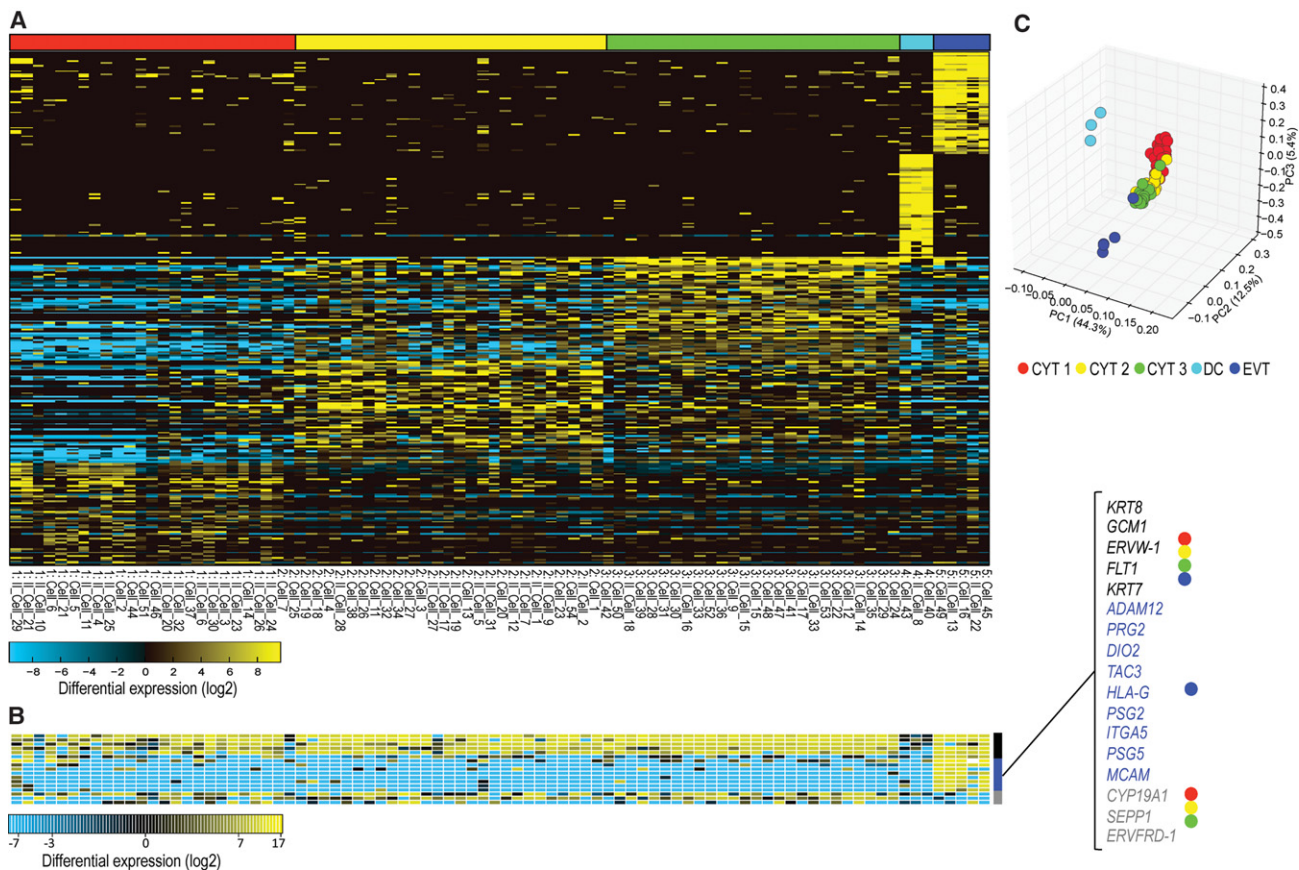
In the following, we first present the main expression patterns starting with comparison of tissue-level transcriptomes to the sum total of single-cell transcriptomes and then characterize the clusters by their diagnostic signatures relative to the particular assembly of cells. The description is limited to diagnostic gene expression, with the noteworthy observations of potential interest for experimental follow-up described in greater depth in Supplemental Information. The 30 most highly expressed genes in each cluster relative to other clusters are listed in Table 1 (averaged values) (full normalized expression levels in Supplemental Table S1). The second part of the study focuses on the cell communication network, its structure, and maternal and fetal contributions to pathways known to play major roles at the maternal-fetal interface. To our knowledge, this is the first characterization of the cell communication network at the human maternal-fetal interface.

## Expression profiles

### *Tissue-level villous placenta RNA-seq*

Various factors may result in cells acquired by the pipeline not being represented in proportion to their frequency in the starting tissue. For example, the cells' adhesive properties or size may bias their passage through the Fluidigm cell, as is most apparent in the large multinucleated syncytiotrophoblast, which we targeted separately. In addition, rare cell types will evade limited sampling. Our tissue preparation enriched for trophoblasts, which is the likely reason that the endothelial cells of placental vasculature or the villous mesenchymal cells are not represented. To estimate what portion of placental gene expression is covered by the single-cell transcriptomes, we compared the single-cell data with the tissue-level transcriptomes, generated from different individuals but the same type of sample (term Cesarean section, not in labor).

The top 25% (2108) most highly expressed genes in the whole placental transcriptome comprise 80% of the total aligned placental mRNA. Only 20 of these genes were not detected at all in the combined single-cell transcriptomes, and a further 34 genes were detected in less than three cells (syncytiotrophoblast-specific genes excluded) (Supplemental Table S2). The list includes several highly expressed RNA-genes with placental function (e.g., *H19* and *MIRLET7*-family); however, it is not clear what missing cell population may be implicated. These genes do not show high expression in the decidual cells, which is the cell type commonly included in



**Figure 1.** The gene expression in the single-cell experiment. (A) Expression of the top 300 genes with greater than twofold differential expression between clusters that were previously established by hierarchical clustering. *x*-axis: The cells with roman II denote the second batch of cells. Note that the cells from both batches are represented in all clusters. (B) Known trophoblast markers are shown separately, with general trophoblast markers in black, extravillous trophoblasts in blue, and cytotrophoblast-specific expression in gray. (C) First three principal components based on 300 marker genes (full PCA in Supplemental Fig. S8). Clusters correspond to the clustering in the heat map in A.

preparations of whole placenta. Given this high level of correspondence between tissue-level transcriptomes and the single-cell data, we conclude that the trophoblast cells at term are by far the most abundant cells, whereas the missing cells are rare and/or overlap greatly with respect to their expression with the cells captured in our samples. Consistent with the view that single-cell analysis may reveal expression profiles of rare cells, our analysis recovers expression of over 4000 genes, which are expressed at levels  $>2$  TPM in five or more cells but are not detected in the tissue-level transcriptome at levels  $\geq 1$  TPM. In this comparison, we chose a lower threshold for the tissue-level transcriptome, as this transcriptome has been sequenced at greater depth than the single-cell transcriptomes. We report the top 1000 of these genes in the Supplemental Table S3, ranked by the number of expressing cells.

#### Placental trophoblast cells

We clustered cells by their single-cell gene expression profiles into five clusters. A combination of known trophoblast markers (*KRT7*, *KRT8*, *GCM1*, *CYP19A1*) distinguishes clusters 1, 2, 3, and 5 from cluster 4 and from endometrial stromal cells (Fig. 1B). Further diagnostic genes are listed in Table 2, showing genes with  $>10$ -fold higher expression in trophoblast than in uterine and immune cells. Among the trophoblast clusters, clusters 1 and 5 show lower

expression of ribosomal proteins, a signature typical of later stages of differentiation in cell lineages leading toward extravillous trophoblast (Ji et al. 2013) and the syncytiotrophoblast (Handwerker and Aronow 2003). The large trophoblast clusters 1–3 were identified as intravillous cytotrophoblast (see Supplemental Text) and the smaller cluster 5 as extravillous trophoblast.

#### Cytotrophoblast (CYT; Clusters 1–3)

These clusters likely represent a continuum in the differentiation of cytotrophoblasts. Genes specifically expressed in cytotrophoblasts are summarized in Table 2. Intravillous cytotrophoblasts are a very heterogeneous group, a finding often encountered at single-cell-level transcriptomes (Brunskill et al. 2014; Olsson et al. 2016). The expression profiles of these cells suggest that differentiation proceeds by individual cells turning on—or off—sets of genes in a different order rather than a uniform progression toward a terminal expression profile. Detailed characterization of intercellular heterogeneity and cell differentiation is beyond our present scope and will be addressed separately.

#### Extravillous trophoblast (EVT; Cluster 5)

This cluster of cells is unique in its expression of nonclassical human leukocyte antigen *HLA-G* and integrin 5 alpha *ITGA5*—both

**Table 1.** Top 30 expressed genes per cluster, ranked according to the average level of expression on a TPM scale

IVT <sup>F</sup> cluster 1	IVT <sup>F</sup> cluster 2	IVT <sup>F</sup> cluster 3	EVT <sup>F</sup> cluster 5	SYN <sup>F</sup>	DC <sup>M</sup> cluster 4	ESF <sup>M</sup>	DEC <sup>M</sup>
<i>RPS28</i>	<i>MIF</i>	<i>RPS11</i>	<i>PRG2</i>	<i>RN7SK</i>	<i>APOC1</i>	<i>TMSB4X</i>	<i>TMSB4X</i>
<i>RPLP1</i>	<i>RPL36</i>	<i>RPLP1</i>	<i>TMSB10</i>	<i>RPS15</i>	<i>FTL</i>	<i>TMSB10</i>	<i>TMSB10</i>
<i>RPS27</i>	<i>RPLP1</i>	<i>RPL36</i>	<i>FTL</i>	<i>CSH1</i>	<i>TMSB4X</i>	<i>ACTB</i>	<i>FTL</i>
<i>MT2A</i>	<i>RPS27</i>	<i>MIF</i>	<i>RPLP1</i>	<i>FTL</i>	<i>TMSB10</i>	<i>IGFBP5</i>	<i>SPARC</i>
<i>TMSB10</i>	<i>RPS11</i>	<i>RPS28</i>	<i>RPS27</i>	<i>RPS13</i>	<i>RPS14</i>	<i>RPL41</i>	<i>B2M</i>
<i>SAT1</i>	<i>RPS14</i>	<i>GSTP1</i>	<i>PHKA2</i>	<i>ATP6V0C</i>	<i>RPS27</i>	<i>THBS1</i>	<i>FTH1</i>
<i>KRT18</i>	<i>RPS28</i>	<i>RPS14</i>	<i>SAT1</i>	<i>RPL37A</i>	<i>CD74</i>	<i>B2M</i>	<i>FSTL1</i>
<i>RPS10</i>	<i>TMSB10</i>	<i>RPS27</i>	<i>FN1</i>	<i>RPL35</i>	<i>RPLP1</i>	<i>SPARC</i>	<i>THBS1</i>
<i>RPL24</i>	<i>RPS18</i>	<i>RPS18</i>	<i>RPL36</i>	<i>RPL41</i>	<i>RPL35</i>	<i>COL1A1</i>	<i>RPL41</i>
<i>RPS18</i>	<i>GSTP1</i>	<i>RPL18A</i>	<i>CDKN1C</i>	<i>RPL12</i>	<i>IFI30</i>	<i>FSTL1</i>	<i>ACTB</i>
<i>RPS11</i>	<i>RPL35</i>	<i>RPL35</i>	<i>AOC1</i>	<i>CSH2</i>	<i>AIF1</i>	<i>VIM</i>	<i>COL1A1</i>
<i>CHMP2A</i>	<i>RPL18A</i>	<i>RPS15</i>	<i>MIF</i>	<i>MIF</i>	<i>RPS19</i>	<i>FTL</i>	<i>IGFBP4</i>
<i>RPLP2</i>	<i>RPS15</i>	<i>NDUFA13</i>	<i>S100A11</i>	<i>RPS16</i>	<i>RPL41</i>	<i>LGALS1</i>	<i>IGFBP3</i>
<i>RPL35</i>	<i>RPLP2</i>	<i>TMSB10</i>	<i>RPS14</i>	<i>RPL31</i>	<i>RPS11</i>	<i>CCND1</i>	<i>IGFBP1</i>
<i>S100P</i>	<i>RPL41</i>	<i>RPL23</i>	<i>EBI3</i>	<i>RPL23</i>	<i>HLA-DRB1</i>	<i>TPM4</i>	<i>WNT5A</i>
<i>RPL8</i>	<i>RPS19</i>	<i>RPL41</i>	<i>RPS28</i>	<i>RPLP2</i>	<i>RPL23</i>	<i>RND3</i>	<i>CALR</i>
<i>CARS</i>	<i>NDUFA13</i>	<i>RPS19</i>	<i>RN7SK</i>	<i>CGB8</i>	<i>HLA-DRA</i>	<i>TAGLN</i>	<i>RND3</i>
<i>NDUFB2</i>	<i>RPL23</i>	<i>RPLP2</i>	<i>S100P</i>	<i>NPIP5</i>	<i>RPL21</i>	<i>FTH1</i>	<i>TXN</i>
<i>CDKN1C</i>	<i>S100P</i>	<i>HIST1H4C</i>	<i>SSR4</i>	<i>RPL36</i>	<i>RPS15</i>	<i>S100A6</i>	<i>LEPROT</i>
<i>RPL23</i>	<i>PAGE4</i>	<i>PAGE4</i>	<i>RPS11</i>	<i>RPS11</i>	<i>RPS28</i>	<i>PTMA</i>	<i>IGFBP5</i>
<i>EEF1D</i>	<i>KRT18</i>	<i>S100P</i>	<i>RPL9</i>	<i>HIST2H2AC</i>	<i>ISG15</i>	<i>MT2A</i>	<i>TPT1</i>
<i>LDHB</i>	<i>RPS29</i>	<i>RPL8</i>	<i>RPL18A</i>	<i>ATP9B</i>	<i>TYROBP</i>	<i>C12orf75</i>	<i>GABARAPL1</i>
<i>RPL41</i>	<i>HIST1H4C</i>	<i>RPS29</i>	<i>RPLP0</i>	<i>RPS10</i>	<i>SAT1</i>	<i>ITGB1</i>	<i>IFITM3</i>
<i>RPL12</i>	<i>FTL</i>	<i>S100A11</i>	<i>RPL41</i>	<i>GNAS</i>	<i>RPS25</i>	<i>TPM1</i>	<i>COL1A2</i>
<i>MIF</i>	<i>XAGE3</i>	<i>RPS16</i>	<i>RPS18</i>	<i>HBB</i>	<i>MT2A</i>	<i>RPL39</i>	<i>SPON2</i>
<i>DDX50</i>	<i>RPL21</i>	<i>XAGE3</i>	<i>EEF1A1</i>	<i>RPL26</i>	<i>RPL36</i>	<i>CALM2</i>	<i>CTSB</i>
<i>RPL21</i>	<i>RPS25</i>	<i>ALDOA</i>	<i>RPL21</i>	<i>HIST2H2AB</i>	<i>ACTB</i>	<i>ACTG1</i>	<i>LGALS1</i>
<i>TMEM258</i>	<i>RPL8</i>	<i>UBB</i>	<i>KRT18</i>	<i>HBA2</i>	<i>RPS29</i>	<i>ANXA2</i>	<i>TPM4</i>
<i>S100A11</i>	<i>S100A11</i>	<i>FTL</i>	<i>ACTB</i>	<i>RPL18</i>	<i>FTH1</i>	<i>CD59</i>	<i>OST4</i>
<i>ATPS1</i>	<i>RPLP0</i>	<i>RPLP0</i>	<i>RPS19</i>	<i>RPS15A</i>	<i>RPS18</i>	<i>CFL1</i>	<i>RPL39</i>

Top 30 expressed genes in each of the cell types (average across cluster or across repeated measurements), ranked according to their expression level (all highly expressed >200 TPM). (IVT) Intravillous cytotrophoblast, (EVT) extravillous trophoblast, (SYN) syncytiotrophoblast, (ESF) endometrial stromal fibroblast, (DEC) decidual cell, (DC) dendritic cell. Superscripts "M" and "F" refer to maternal and fetal source, respectively.

considered markers of extravillous trophoblast, *KRT7* (Telugu et al. 2013; Borbely et al. 2014) and *ADAM12* (Biadasiewicz et al. 2014). Further EVT-specific genes and other highly expressed genes are listed in Tables 1 and 2 and discussed in the Supplemental Material. These cells differ from cytotrophoblasts by expression of some endothelial genes, such as adhesion molecule *MCAM* (Jouve et al. 2013; Kaspi et al. 2013), lowered E-cadherin (*CDH1*) (Zhou et al. 1997), and the receptor for vascular endothelial growth factor *FLT1*. Note that *MCAM* may be a receptor for *WNT5A*, affecting cell motility (Ye et al. 2013). While EVTs are involved in invasion in early pregnancy (Bischof et al. 2000), some of the invasion-associated signature may be lacking at term (Ji et al. 2013). At term, genes up-regulated in EVT relative to cytotrophoblast are involved in modulation of extracellular matrix, vascularization, and immune pathways (Table 2; Supplemental Fig. S1B).

#### Syncytiotrophoblast (SYN; laser microdissection)

This cell type was collected in two samples from a single term placenta by laser microdissection and analyzed by the conventional RNA-seq pipeline. The two samples were analyzed separately and show a high correlation of expression ( $r=0.99$ , in square root-transformed TPM), indicating a technically robust transcriptome. Among the most highly expressed syncytiotrophoblast-specific genes are hormones (*CSH2*, *CSHL1*, *GH2*), *CGA*, and pregnancy-specific glycoproteins (*PSG*-family) (detail in Supplemental Fig. S2). Several highly expressed genes are concomitantly restricted to syncytiotrophoblast and extravillous trophoblast: *PSG2* and

*-5*, chorionic somatotropin (*CSH1* and *CSH2*), and the key prostaglandin catabolic enzyme, *HPGD*. An intriguing observation is the expression of human MHC type II antigen in syncytiotrophoblast; uniquely among trophoblasts, this cell layer expresses a profile of human leukocyte antigens similar to the profile of the uterine dendritic cell cluster (*HLA-DRA*, *HLA-DRB1* and *-5*, *HLA-DMA*, *HLA-DQB1*, *HLA-DPA1*), albeit at levels of expression 20- to 30-fold lower than those in dendritic cells.

#### Maternal cells

##### Uterine dendritic cells (DCs; Cluster 4)

Cluster four consists of three cells lacking the trophoblast markers but expressing MHC II subunits (*HLA-DRA* in particular) (Table 2), characteristic of antigen-presenting cells. These cells lack expression of *CD19*, *CD209*, and *CD163* but express the combination of markers *ITGAX*<sup>+</sup> *CD14*<sup>+</sup>/*CD4*<sup>+</sup>/*CD83*<sup>+</sup> *CD86*<sup>+</sup>. The genes *CLEC4C*, *THBD*, *CD1C*, *CD80*, *IL10*, and *IL12B* that characterize some of the decidual DCs (Kammerer et al. 2008; Talayev et al. 2010) were not found. *CD14* is often considered a marker of macrophages, with which dendritic cells are closely related. It has, however, been shown that *CD14* expression is retained in dendritic cells when they develop in the presence of trophoblasts (Talayev et al. 2010). These facts, together with the source of cells in this experiment, suggest that these are uterine dendritic cells and thus of maternal origin (Kammerer et al. 2000; Tagliani and Erlebacher 2011). Even if we cannot exclude that some of the

**Table 2.** Local markers—genes greater than fivefold up-regulated in the cluster relative to other clusters present in the current data set; in addition, genes discussed in the text due to their status as markers are added

Fetal					Maternal		
Intravillous cytotrophoblast			EVT	SYN	ESF	Decidua	Dendritic cells
Cluster 1	Cluster 2	Cluster 3	Cluster 5	(microdiss.)	(primary cells)		Cluster 4
		XAGE3				CITED2	AIF1
		XAGE2				COL1A2	C1QB
		SERINC5				NBL1	CD11c
		S100P				CXCL12	CD14
		RASA1				DKK1	CD4
		MFAP5				DKK3	CD68
		LIN28B				GREM1	CD83
		KRT8				SULF1	CD86
		KRT7				IGFBP4	CTSS
		INS-IGF2				IGFBP5	CXCL10
		GCM1				LOXL2	CXCL11
		GATA3				MMP14	CXCL16
		ERVW-1				MMP2	CXCL9
		ERVFRD-1				MYL9	LGALS2
		EGR1				LOX	FGL2
		EFNA1				CDH2	GPR183
		CYP19A1				SPOCK1	HLA-A
		PHLDA2				OXTR	HLA-B
		ACKR2				SPON2	HLA-DMB
	ADR81			PSG5		MSRB3	HLA-DPA1
	PUF60			CST6		PXDN	HLA-DPB1
	SNORD3A			PSG11		WNT5A	HLA-DQA1
	PRMT7			PSG2	TRIP13	ABI3BP	HLA-DQB1
	EIF1AY			HPGD	TOP2A	ADRA2C	HLA-DRA
	FXYD3			CSH2	TACC3	C1QTNF1	HLA-DRB1
	GRAMD2			CSH1	SSTR1	PRL	HLA-DRB4
	INSL4		TAC3		SPDL1	SST	IRF4
	ITGB8		AOC1	SEMA3B	SPC24	CDKN2B	LAPTM5
	PAGE4		B3GNT7	CSH2	SMC4	CEMIP	MS4A6A
	SLC13A4		COL17A1	CSHL1	SHCBP1	CRELD1	PTPN7
	SLC22A11		CORO6	FCGR2A	RACGAP1	CSF1	RGS1
			DAPP1	GH2	PRR11	FGF9	VMO1
			DIO2	HBA1	PRPS1	FOXO1	CCRL2
			EGLN3	HBA2	PODXL	GALNT15	MMP9
			FAM174B	HBB	PLK1	GAS1	LGALS17A
			FN1	HBG1	MMP1	GPR137B	BCL2L14
			HLA-G	HBG2	MCM4	HSD17B2	IRF4
			TUBB2A	HLA-DMA	MAD2L1	HSPA2	UBD
			HTRA1	HLA-DPA1	KNSTRN	IGFBP1	CD38
			IFNGR2	HLA-DQB1	HMGA2	IL15	CST7
			IL1R2	HLA-DRA	FOXM1	KCND2	ACHE
			IL1RAP	HLA-DRB1	CEP55	LAYN	CD37
			ITGA5	HPGDS	CDC45	LRIG1	MED140S
			TNFSF10	CGB8	ANLN	MCC	ZNF497
			MCAM	LGALS14	ACTC1	NID2	GAL
			SELM	LYVE1	AURKB	NPTX1	DHRS4L1
			PAPPA2	NPIP3	NTM	RNF182	AZU1
			PLAC8	CGB5	CDC20	WNT2	NTM
			PRG2	PSG1	RRM2		AURKB
			PROCR	PSG3			CCL7
			PTPRF	PSG6			PINX1
				PSG9			ARMC6
				ZNF117			FAM26F
				ZNF91			MT1X
				HIST2H2AB			ASF1B
				RAMP2			SLCO4A1
				MUC20			

markers are missing in our data as a consequence of methodological or biological variability, particularly due to low number of cells of this type in our sample ( $N = 3$ ), the expression profile is nonetheless sufficiently consistent for the identification of these cells as uterine dendritic cells.

#### Endometrial stromal fibroblasts and decidual cells (ESFs and DEC; cell culture)

These cells stem from primary cultures from two patients at days 3 and 9 of the menstrual cycle and were cultured following standard procedures. Decidual cells in this study have been differentiated in vitro (see Methods; for associated gene expression changes, see

**Table 3.** Genes up- and down-regulated during decidualization

Up-regulated				
IGFBP1	IL1R1	MCC	EDNRB	MAOA
SST	GAS1	CEBPD	BDKRB2	ARSG
CGA	SLA	FGF9	PDGFD	FAM19A2
EPAS1	MUM1L1	KLF9	MMP11	GDF15
RGCC	WNT4	C1QTNF7	CPM	FIBIN
SLC40A1	ACKR3	LRIG1	PTPRE	TM4SF1
ADRA2C	PRL	CLU	MAOB	CEND1
DHRS3	FOXO1	ABCA8	HSPA2	FAXDC2
CEMP1	SNCA	SIK1	RAMP1	OLFML2B
HSD17B2	ALDH1A1	TSKU	GPX3	MAPK10
PDK4	HSD11B1	CDKN1C	PTGES	NR4A2
KCND2	RASL11B	PAPPA	PPARGC1A	GALNT15
EFEMP1	MASP1	RDH10	FAM167A	
NPTX1	WISP1	BMP2	CYP26B1	
RGS2	DCN	CRISPLD2	RBKS	
Down-regulated				
ANLN	NCAPG	CDCA3	PRSS3	MCM2
NTM	ZWINT	CDCA2	GIN52	RAD51
AURKB	E2F7	SERPINB7	OIP5	APCDD1L
CDC20	FOXM1	DTL	RECQL4	E2F1
RRM2	DLGAP5	MYBL2	AURKA	EPHB1
BIRC5	CDCA8	PLK4	MND1	TMPO
SKA3	PRR11	CCNA2	SGOL1	CHAF1B
MKI67	KIF2C	TPX2	CIT	KIF18A
UBE2C	NUF2	RNASEH2A	SPAG5	SPDL1
PLK1	CENPA	MELK	FEN1	TNFSF4
CEP55	CCNB2	EXO1	CENPK	PTTG1
KLF2	GTSE1	CDKN3	DES	MCM7
ASF1B	TTK	MYBL1	KCNMA1	AKS
CKAP2L	FAM64A	ACTC1	RAB27B	FOSL1
PBK	TK1	CENPW	FANCI	FAM196B
MALL	KIF20A	CENPE	MAD2L1	FANCA
CDCA5	NRGN	ESCO2	RACGAP1	FBXO5
SHCBP1	CDT1	RAD54L	CKS1B	TRAIP
MMP1	BUB1	BRIP1	OPCML	BARD1
CENPU	CENPM	NCAPH	NDC80	ITGA2
CALB2	PODXL	HMGGA2	S100A10	PSMC3IP
SPC24	CASC5	NEK2	FGF5	SSTR1
PKMYT1	MCMS5	CCNE2	MMP3	IQGAP3
KIAA0101	TYMS	KIF23	ARHGAP11A	HMGB2
BUB1B	RAD51AP1	HMMR	GIN53	HJURP
TOP2A	CENPF	DEPDC1	ORC6	GIN51
CDK1	CCNB1	GIN54	KIF18B	
PRC1	SPC25	RGS4	FANCD2	
TRIP13	TACC3	FAM83D	MCM4	
NUSAP1	PLEK2	TROAP	CHTF18	

Brar et al. 2001). For the most significantly up- and down-regulated genes, see Table 3; a portion of these will be discussed in the context of interactions below. Differentiation marks a transition from the proliferative into decidual state. Correspondingly, the bulk of down-regulated genes in decidua relative to the endometrial stromal fibroblasts are involved in cell cycle, replication, and DNA repair (Supplemental Fig. S3). In contrast, multiple metabolic, immune, and hormonal pathways are up-regulated in decidualization.

### Maternal-fetal cell-cell communication

We used public databases of ligand-receptor relationships (Methods) to infer cell interactions. While based on mRNA expression rather than function, many putative interactions are supported by the expression of downstream target genes in the particular cells or have empirical support in prior studies and have been tested here (below).

The plot in Figure 2 shows the inferred cell-cell interactions, with the cell types indicated by color, and expressed ligands and

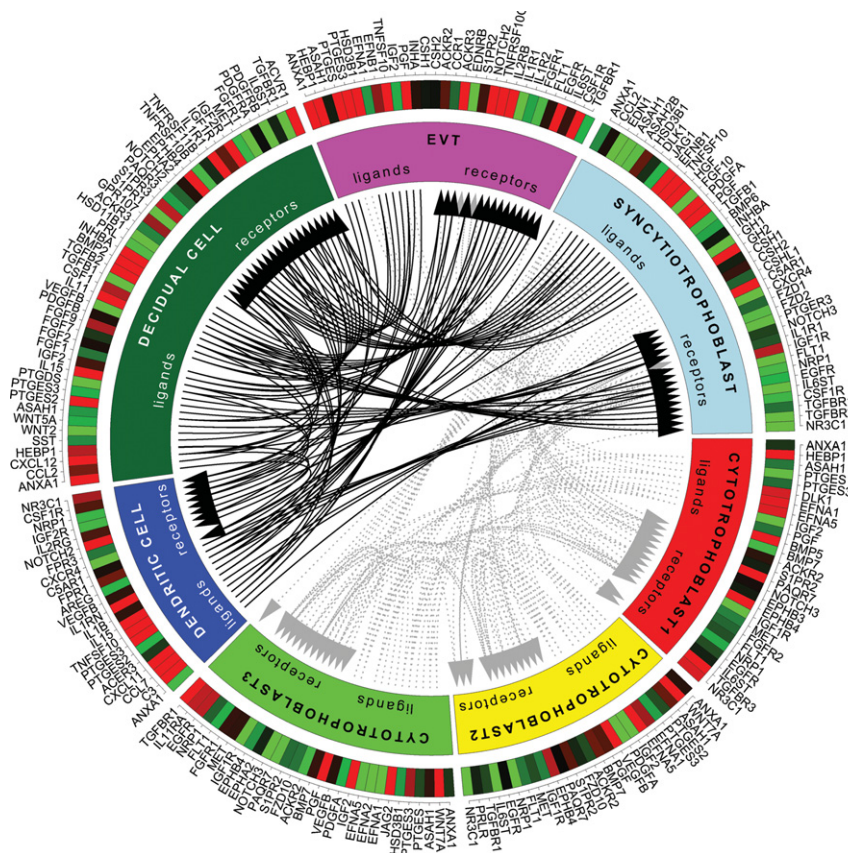
receptors listed along the outer margins. We included ligands and receptors expressed beyond 30 TPM, to allow visualization. The arrows connect expressed ligand and receptor genes, indicating an interaction, with gray arrows representing the full interactome, whereas the black arrows signify a set of interactions between adjacent maternal and fetal cells. Figure 3 illustrates the increase in the potential for interactions between uterine and adjacent trophoblast cells during decidualization as interaction plots (Fig. 3A,B) and quantitatively as the number of edges (Fig. 3C).

### The structure of the cell-cell communication network: DEC's are an interaction hub

The most striking aspect of the interactome in Figure 2 is the large number of interactions between decidual cells and the syncytium, suggesting that the uterine decidua acts as a hub of fetomaternal cell signaling. Figure 3C summarizes the total number of outgoing and incoming edges of each cell type. The interactions with placental and dendritic cells are assessed for decidualized cells (rather than ESF), as this reflects the *in vivo* situation during pregnancy. Uterine cells and the syncytiotrophoblast thus lead in the potential for interactions among the cells of maternal-fetal interface. The dominance of decidual signaling is maintained if the overlap in ligands and receptors across closely related trophoblasts is removed (which could enhance the pattern). We further asked how likely the observed structure is based merely on the observed number of ligands/receptors per cell type, using a permutation test. We find that, given the observed number of ligands and receptors across cells, only 20% of the randomly generated networks have a higher number of connected decidual ligands (Supplemental Fig. S10). As this is at least in part due to the high number of ligands and receptors expressed in maternal endometrial stroma, we next asked how likely it is that decidual cells will express the number of ligands and receptors given the expectations from other cells in the data set. This is highly unlikely for the number of expressed receptors and ligands, as well as the number of those involved in putative interactions (in all cases,  $P < 0.05$  for decidualized, nondecidualized cells, and syncytiotrophoblast), confirming the significance of the characterization as a hub. To specifically assess the contribution of decidualization to the potential for cell communication, we compared the inferred interactions of fetal cells with ESFs to those with DEC's in Figure 3, A and B. The up-regulation of uterine ligands and receptors during decidualization enhances signaling to the syncytiotrophoblast and EVT, as well as from syncytiotrophoblast. In fact, a majority of receptors and ligands up-regulated in decidualization likely communicate with trophoblast cells (Supplemental Fig. S4). The involved pathways will be discussed below.

### GPCR spectrum characterizes cell types

Cells exert their biological function, in large part, by interacting with other cells. This suggests that cell types with different biological functions would show distinct profiles of ligands and receptors, as these enable the cell-type-specific interactions. Therefore, the receptor/ligand profile may serve as a robust biological characterization of the cell types. In our data, this is the case in particular with respect to G-protein-coupled receptors (Fig. 4; Supplemental Table S4). Figure 4A shows the heat map of GPCR expression among cells, showing the specificity of GPCR signaling among clusters. Distinct signatures of different cell types are contrasted in our data set with very similar signatures of the three cytotrophoblast clusters. This supports the inference that these three clusters are closely related groups of cells representing the same cell type.



**Figure 2.** Putative signaling between expressed receptors and their ligands in pregnant uterus at term. Compartments represent cell types, and their expressed ligands and receptors are noted along the outer margin. Average cluster gene expression was used, with expression threshold 30 TPM. Black arrows denote putative interactions between directly adjacent maternal and placental cells (maternal decidua and placental EVT and SYN), and dotted gray lines denote putative interactions with cytotrophoblasts.

Specificity of GPCR profiles suggests an interesting way to characterize cell-type identity; namely, by focusing specifically on the cell's interaction potential. Thereby, it may be possible to distinguish between the cell's biological role within a tissue captured in interactions from the more variable aspects of the cell's gene expression phenotype.

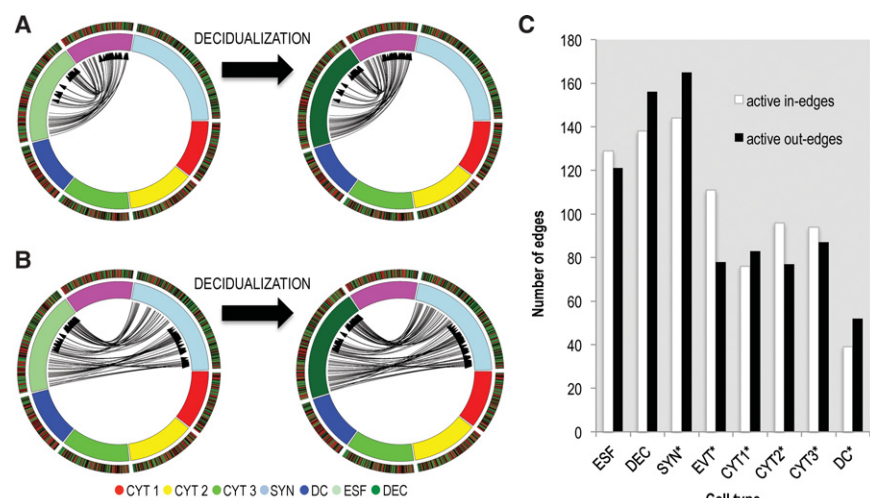
Of over 400 GPCRs in the IUPHAR data set (<http://www.guidetopharmacology.org>), 116 are expressed >2 TPM in at least one of our cells, most in a cell-type-specific manner. For example, GPCRs expressed in cytotrophoblasts include multiple members of the WNT-pathway (mainly Frizzled and LGR families), purinergic and chemokine receptors. Dendritic cells are rich in chemokine receptors, complement system receptors, and prostaglandin E receptors. Decidua differs from endometrial stromal fibroblast in the 260-fold up-regulation of adrenergic receptor 2C and the expres-

sion of endothelin and chemokine receptors, while it shares with endometrial stromal fibroblasts the expression of serotonin and oxytocin receptors (down-regulated in decidualization), thrombin receptor-like 2 (*F2RL2*), and several Frizzled receptors (*FZD4*, -6). The downstream G-protein subunits, in particular the G-protein alpha subunits, can be associated with functional classes via their specific second messenger systems. We used this association to compare cell types with respect to the types of GPCR signaling they are involved in. The plot in Figure 4B shows that the most distinct cell type with respect to GPCR signaling is the dendritic cell, in which most GPCR signaling is associated with inhibition of cAMP signaling. Somewhat distinct is EVT, in which cAMP inhibition represents a greater proportion of GPCR signaling than in other trophoblasts or endometrium.

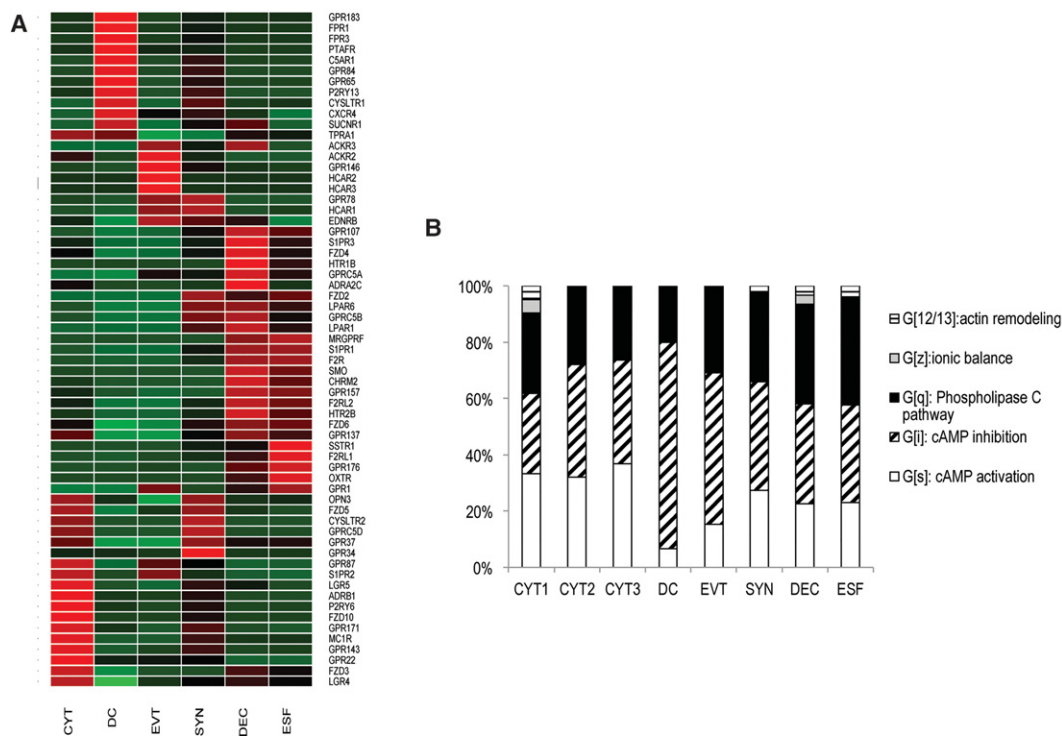
#### Uterine decidualization changes the interaction potential

The cell-cell interactions that are enabled by decidualization of the uterine stroma are of particular interest. These can be found by comparing the interactions of the decidualized with those of undifferentiated stromal fibroblasts.

Decidualization up-regulates growth factors *FGF9*, *BMP2*, and *PDGFB*, for which the receptors are expressed in syncytiotrophoblast and EVT, and the receptors are also up-regulated in decidua itself. In addition, decidual up-regulation of *IL15* and *CSF1* introduces potential to signal to the receptors in EVT



**Figure 3.** Decidualization increases the cell-cell signaling potential with the adjacent EVT (purple, *A*) and syncytiotrophoblast (light blue, *B*). Average cluster gene expression is shown, with genes expressed >5 TPM included. (*C*) Average numbers of putative interactions per cell type, separated into in- and out-edges. Asterisks refer to the number of interactions in the context of decidua, rather than undifferentiated endometrial stromal fibroblasts.



**Figure 4.** Distribution of G-protein-coupled receptors among cell types. (A) The expression heat map of GPCRs across sampled cells. (B) Characterization of cell clusters by class of G-protein alpha subunit used.

and dendritic cells. Decidual cells also likely communicate with dendritic cells via *IL1B*, which is produced by dendritic cell and signals to decidual cell receptor *IL1R1*. Further potential for interaction is the decidual up-regulation of *CCL2*, which binds to atypical chemokine receptor *ACKR2* expressed in trophoblast and dendritic cells and is associated with termination of inflammatory response by chemokine scavenging. Up-regulated autocrine signaling includes somatostatin (*SST*) and likely neurostatin signaling via down-regulated *SSTR1* and up-regulated *GPR107* receptor (Samson et al. 2008; Yosten et al. 2012, 2015). *WNT4* expression is initiated, with its receptors expressed in trophoblast and decidua (below). Decidualization down-regulates uterine neuropilin (*NRP1*), reducing its potential inhibition of *VEGFB* in trophoblast.

#### Signaling changes in trophoblast differentiation

The comparison of receptor/ligand spectra expressed in differentiated extravillous trophoblasts and syncytiotrophoblasts to that of cytotrophoblasts also suggests the signaling differences that arise during trophoblast differentiation in the two lineages. The signaling interactions likely gained in EVT involve interleukin receptors *IL1R1* and *IL1R2* potentially interacting with maternal decidua and dendritic cells, *EFNB1* and *TNFSF10*, as well as growth factor receptors of decidual growth factors. Similarly, the differentiation toward syncytiotrophoblast involves gain of chemokine and growth factor receptors (*IL1R1*, *TGFBR1* and *-2*, *PDGFRA*, *BMPR2*) but in addition adds growth signaling toward decidua by of *IGF1*, *HGF*, and *DLK1*.

#### Maternal–fetal interactions in main signaling pathways

Here, we characterize the fetal and maternal contributions to the main signaling pathways and their change upon decidualiza-

tion (see Supplemental Material for IGF, GH and prolactin pathways).

#### Wnt signaling

Decidualization entails differentiation toward epithelial-like character, reflected in up-regulation of several Wnt ligands, above all, the induction of *WNT4* and up-regulation of *WNT2* and *WNT5A*. Intravillous cytotrophoblasts show moderate expression of Wnt ligands (*WNT3*, *WNT7A*), and *WNT6* is expressed in syncytiotrophoblast. Expression of Wnt receptors (Frizzled family) in fetal and maternal cells suggests autocrine as well as paracrine signaling. Specifically, *FZD2*, *-4*, and *-6* are expressed in decidualized cells, and *FZD3*, *-4*, *-5*, and *-6* are expressed in cyto- and syncytiotrophoblast. *WNT4*, a marker of decidualization, likely binds to the *FZD6* receptor, possibly similar to signaling during mesenchymal-to-epithelial transition in kidney development (Lyons et al. 2004). Canonical and noncanonical pathways have been suggested for *WNT4* signaling in endometrium (Hou et al. 2004; Knofler and Pollheimer 2013). The up-regulation of *WNT2* and putative binding to the *FZD4* receptors (Klein et al. 2008) in cytotrophoblast and decidua may augment canonical *WNT4* signaling. The expression of downstream transcription factors *TCF7L2* in trophoblast and decidua, as well as *LEF1* in syncytiotrophoblast, further supports activation of canonical Wnt signaling in maternal and placental cells via autocrine and paracrine signals.

#### Prostaglandins

Short-distance signaling of prostaglandins makes their expression highly informative. The enzyme-expression of the prostaglandin biosynthesis pathway is summarized in Supplemental Figure S5. Briefly, the phospholipase A (*PLA2G16*) catalyzing the early step

is broadly expressed, and cyclooxygenase 1 (*PTGS1*) is up-regulated in decidualization. The enzymes active in the later stages of the pathway seem complementarily present in maternal and placental term cells: The prostaglandin D synthetase *PTGDS* increased fourfold in decidualization; *HPGDS* is expressed in syncytiotrophoblast; prostaglandin E synthetase *PTGES* in EVT and cytotrophoblast, and moderately in decidua; and the oxidoreductase *FAM213B* catalyzing the biosynthesis of prostaglandin F2alpha (PGF2 $\alpha$ ) is expressed in decidua. Interestingly, while prostaglandin F receptor is weakly expressed in syncytiotrophoblast and decidua, prostaglandin F receptor antagonist, *PTGFRN*, is highly expressed in decidua. In addition, the main prostaglandin catabolic enzyme, 15-hydroxy dehydrogenase (*HPGD*), is highly expressed in all term trophoblast cell types (in EVT >2000 TPM) and absent in maternal cells. Collectively, the data suggest prostaglandin E2 synthesis in decidua and trophoblast and prostaglandin F2alpha predominantly in decidua. Receptor expression is generally weak, and prostaglandin F receptor antagonist and the catabolic enzyme *HPGD* may further inhibit the prostaglandin signaling at the maternal-fetal interface. These patterns suggest a uterine autocrine signaling of PGF2 $\alpha$ , whereas the trophoblast PGE2 signaling may be directed toward decidua. In vivo analysis will reveal whether low levels of decidual PGE-receptors are artifacts of decidualization in the absence of trophoblast.

#### Interleukin and TNF signaling

The interleukin expression among the collected cells is highest in dendritic cells and is comprised of *IL1B*, *IL15*, and *IL32*. *IL1B* expression is matched by high expression of its receptor *IL1R1* in decidua, up-regulated >40-fold during decidualization. The adjacent fetal EVT and the syncytiotrophoblast express *IL1R1*, and EVT also highly expresses *IL1R2*, a decoy receptor for both *IL1B* and *IL1R1*. Interestingly, maternal dendritic cells also express anti-inflammatory *IL1RN*, the *IL1*-receptor antagonist. Thus, while maternal immune cells produce pro-inflammatory signals and decidua and trophoblasts are capable of perceiving them, maternal immune cells and placental cells concurrently express strong multifaceted anti-inflammatory signals, potentially inducing a sustained anti-inflammatory state (see details in Supplemental Material).

Tumor necrosis factor alpha and beta (*TNF*, *LTA*) are not expressed in our cells; however dendritic cells express genes of the tumor necrosis factor ligand family, *TNFSF*. In addition, EVT and syncytiotrophoblast highly express the apoptosis-inducing *TNFSF10*. This is a gene for the ligand of death receptor *TNFRSF10B*, a gene which is up-regulated in decidualization. Decidualization also increases expression of *TNFRSF11B*, coding for a decoy receptor for *TNFSF11* and *TNFSF10*. These data suggest that active signaling from the trophoblast to decidua via *TNFSF10* and *TNFRSF10B* is modulated by decidual *TNFRSF11B* expression. The downstream targets are described in the Supplemental Material.

#### Steroid hormones progesterone, estrogen, and glucocorticoids

Steroid hormone biosynthesis is well represented at the feto-maternal interface (Supplemental Fig. S6). The upstream enzymes catalyzing steroid and specifically progesterone synthesis (*CYP11A1* and *HSD3B1*) are expressed in all trophoblast cells. On the maternal side, decidualization is accompanied by a strong up-regulation of the enzyme with a primary role in converting estrone to estradi-

ol (*HSD17B2*) but also catabolizing progesterone. Decidua expresses a series of progesterone-catabolizing enzymes: *AKR1C1*, *SRD5A1*, and *SRD5A3*. The conversion of progesterone to estrogen via androgens is inhibited in human pregnancy by the lack of *CYP17A1*, the enzyme catalyzing the first step of the conversion. In contrast, aromatase (*CYP19A1*), the enzyme converting the second step, is highly expressed in term syncytio- and cytotrophoblast. Whereas progesterone- and estrogen nuclear receptors *PGR* and *ESR1* are weakly expressed in decidual or placental cells, the less-understood membrane-associated receptors *PGRMC1* and -2 are highly expressed in all trophoblasts and up-regulated in decidualization (*PGRMC2*: fourfold to reach >800 TPM). Collectively, data suggest high progesterone (P4) production in the fetal compartment, a moderate increase in P4-catabolism in decidualization, and high overall expression of membrane-bound P4 receptors.

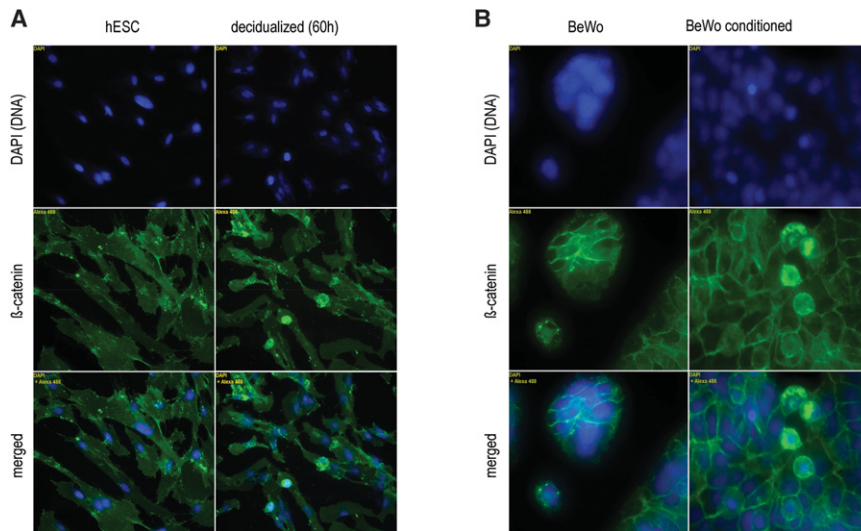
The progesterone signal may also be mediated via the glucocorticoid receptor (GR; gene *NR3C1*) (Karalis et al. 1996), which is generally expressed most highly in dendritic cells and is down-regulated in uterine decidualization. Signaling via GR could explain P4's modulation of maternal immune cells (for review, see Butts et al. 2010). Dendritic cells are a candidate mediator of maternal tolerance for fetal antigens (Tagliani and Erlebacher 2011), with P4 proposed as a signal reducing DCs' capacity to stimulate T-cells (Ivanova et al. 2005; Hughes et al. 2008; Butts et al. 2010). Previous work using PGR antagonist RU486 to reverse the effect of progesterone concluded that the effect is mediated by PGR (Butts et al. 2010). However, the fact that RU486 also antagonizes GR, the absence of *PGR*, and high expression of *GR* in uterine dendritic cells suggest that the observed effect on dendritic cells is mediated by P4 binding to GR. In accordance, P4 signaling via GR has been recently shown to modulate *IL1B*-induced *COX2* expression in myometrium (Lei et al. 2012).

#### Empirical validation

We chose Wnt-signaling to validate some of our results. Transcriptome analysis predicted induction by *WNT4* and/or *WNT2* of canonical Wnt pathway in maternal and fetal cells upon decidualization. The downstream effect of the canonical Wnt signal is nuclear translocation of beta-catenin. Using endometrial stromal cells (hESC) and cytotrophoblast cells (BeWo), we tested the predicted activation by conditioning the growth medium of the trophoblast cells with the supernatant of the maternal cells and determined beta-catenin localization using immunocytochemistry. As shown in Figure 5A, decidualization indeed activates the canonical pathway in endometrial stromal cells. Adding supernatant from decidualized cells to the growth medium of the trophoblast cells also activates the canonical Wnt pathway in trophoblast cells (Fig. 5B), whereas this does not occur upon adding supernatant from nondecidualized cells (or fresh decidualization medium) (Supplemental Fig. S9).

## Discussion

The data presented in this study reveal two main patterns in addition to the detailed characterization of the cells. The first is a model of the cell-cell communication network at the feto-maternal interface, with some interesting aspects that will be discussed below. The second interesting pattern is the cell-type specificity of GPCR profiles, suggesting a novel way of characterizing cell types.



**Figure 5.** Proof of concept: Canonical Wnt signaling is induced in decidualization in maternal cells, as well as in fetal cells conditioned with supernatant of maternal decidualized cells. Anti-beta-catenin antibody is shown in green, and DAPI nuclear staining in blue. The translocation of beta-catenin into the nucleus marks activation of the canonical Wnt signaling pathway. (A) Endometrial stromal fibroblasts (hESC) show beta catenin nuclear localization upon decidualization (Nikon, 20 $\times$ ). (B) BeWo show no nuclear localization in normal growth medium; however, the nuclear localization occurs upon addition of the supernatant from decidualized cells (Nikon, 40 $\times$ ).

When discussing our cell-cell network model, it is important to note that this model is inferred from RNA expression rather than the presence of corresponding proteins, which is dependent upon the degree of post-transcriptional regulation. Nevertheless, the expression of both interacting partners arguably offers strong suggestive evidence of actual cell-cell signaling. Maternal decidual cells and the syncytium are at the center of the cell-cell interaction network. Many ( $\geq 20$ ) decidual signals are also directed toward and received by the extravillous trophoblast cells. Data show that the majority of receptors and particularly ligands up-regulated during decidualization have their complement ligand or receptor expressed in placental cells. Note that the decidual-trophoblast match in this study involves the portion of changes intrinsic to the decidualization program and does not depend on the presence of interaction partners, because the complementary gene expression changes were observed *in vitro*. Decidualization *in vitro* predominantly activates ligands of receptors in the adjacent trophoblast cells. This is consistent with the observations that the presence of factors from the maternal decidua affects the growth of trophoblast cells (Menkhorst et al. 2012). The potential to receive incoming trophoblast signals is similar between endometrial stromal fibroblasts and *in vitro* decidualized cells.

Interestingly, immune-regulatory as well as growth-regulatory interactions between maternal and placental cells show mutual concurrent expression of the enhancing and the inhibitory signals. We interpret this pattern as a signature of the fine-tuned regulation, which, as in any control circuit for maintaining homeostasis, must involve both inhibitory and enhancing effectors (Bernard 1856; Bernard et al. 1927). This principle is currently best appreciated in the immune system (Kotas and Medzhitov 2015) and is reflected in cytokine and prostaglandin pathways in these data. Yet, the principle likely also applies to regulation of another energetically costly process, namely growth during pregnancy. In the case of pregnancy, a dynamic balance must accom-

modate not just different cells or organs within an individual but also transient close interaction with another individual and its growing requirements. It will therefore be particularly useful to focus on the interaction dynamics occurring as new stable states are reached in pregnancy or even just anticipated in the nonpregnant cycle. The fact that the uterine cells decidualized in the absence of trophoblasts show many changes required for maternal-fetal interactions that likely also occur in the sterile menstrual cycle, strongly suggests that information collected in the nonpregnant cycle (i.e., in menstrual shedding) could be predictive of pregnancy outcomes.

An interesting result of potentially broad significance is the cell-type specificity of G-protein-coupled receptor profiles. GPCRs are used by a variety of ligands in numerous organismal contexts, from immune system to hormonal and neuronal signaling. GPCR signaling is evolutionarily old, and the diversification of receptors has been related to the increase of organismal complexity following the evolution of multicellularity

and the associated diversification of cell types (Bradford et al. 2013; de Mendoza et al. 2014). Indeed, GPCRs are expressed in a cell-specific manner, with less heterogeneity of expression among cells of the same type. We interpret this as a signature of the biological role of a cell type, which is in part realized by the information transferred in cell-cell communication. If this observation can be confirmed in studies of other cell types, GPCR expression may constitute a mechanism for conveying cell-type identity within a focal species.

The novel insights from this work are enabled by the single-cell resolution of transcriptomes. However, the approach has limitations to consider. The cell collection was intentionally not guided by cell sorting using specific markers; nonetheless, the sample is biased due to morphological and chemical properties of the cells, such as cell size or cell adhesion. This is important for the comparison of single-cell data with tissue-level transcriptomes. Tissue transcriptomes are dominated by the transcription profiles of the common cells, whereas single-cell transcriptomes are biased by the physical properties of the cells. Provided that the most common cells are not also the most easily attainable in the single-cell pipeline, the two approaches provide complementary information.

We extended the analysis of sampled placental cells with *in vitro* decidualized primary endometrial stromal fibroblasts as a proxy for uterine lining. The transcriptome of these cells exhibits most of the known markers of decidualization. A potential limitation to these data is that *in vitro* cell cultures could lack some gene expression that arises due to the *in vivo* context or by interaction with adjacent cell types (e.g., Calabria and Shusta 2008; Gellersen and Brosens 2014; LoVerso et al. 2015). Future *in vivo* decidual cell transcriptomes will provide an answer to this question.

Finally, the study required combining multiple approaches to obtain transcriptomic data. We have accounted for this diversity to the extent possible in the computational part of the analysis.

There are also methodological differences prior to and during the sequencing pipeline, such as tissue handling, amplification methods, and facility, which may, to some extent, influence the overall comparability of expression levels. In spite of these limitations, our results are consistent with previous experimental studies. We are thus encouraged to consider the novel insights of this study to be robust. It will be interesting to broaden this approach in the future with additional cell types, in particular myometrial cells, and earlier gestational stages, as well as to use these results as a standard against which to interpret disease-related deviations.

## Methods

### Generating transcriptomes

#### *Single-cell transcriptomes*

Fresh, term placental tissue from normal pregnancies was obtained following elective Cesarean section and minced finely. Tissue underwent serial trypsin and collagenase digests as previously described (Troja et al. 2014), and the resulting cell suspension was layered on a continuous Percoll gradient. Cells were collected from the 20%–60% fractions. Collected cells were filtered through a 40- $\mu$ m cell filter and underwent Trypan blue exclusion to assess viability. Cell suspension at  $3 \times 10^5$  cells/mL was submitted to Auto Prep System and handled according to the manufacturer's instructions. Libraries were subjected to paired-end 75-bp RNA sequencing on an Illumina HiSeq2500, resulting in a median of 3.65 million reads per cell. The reads were aligned to human assembly (hg19) (see Supplemental Fig. S7 for fragment lengths and mapped reads) using Bowtie, and assessed for abundance by eXpress (<http://www.rnaexpress.org>) (see also <http://bio.math.berkeley.edu/eXpress/index.html>; Roberts and Pachter 2013).

#### *Placental tissue-level transcriptome*

Two human placentas (separate from above) were collected at term C-section without labor (IRB: CCHMC IRB 2013–2243), flash-frozen, and kept at  $-80^\circ\text{C}$  until RNA extraction using TRIzol. All samples passed quality tests using Bioanalyzer 2100 (Agilent), assuring RIN  $> 9$ . Libraries were prepared with TruSeq Stranded mRNA HT (Illumina) and subjected to paired-end 50-bp RNA sequencing on an Illumina HiSeq2500, at the depth of 30 million reads per sample.

#### *Syncytiotrophoblast transcriptome*

We used a Leica LMD 7000 Laser Microdissection system to collect syncytiotrophoblast from fresh-frozen sectioned tissue. Microdissected material was collected directly into the lysis buffer of the Qiagen RNeasy Plus Micro kit, and RNA was extracted subsequently. Two samples were collected from the same placenta. Due to low concentration, the Ovation RNA Amplification System (Ribo-SPIA technology, NuGen) was used to amplify cDNA. Libraries were sequenced using Illumina HiSeq2500, to obtain a minimum of 30 million 75-bp paired-end reads per sample. Reads were aligned using Bowtie, counted and normalized using RNA-eXpress.

#### *Primary human endometrial stromal fibroblast culture transcriptome*

Two primary human endometrial stromal fibroblast lines (HsESC\_217S and HsESC\_218S) were obtained from two patients undergoing a polypectomy. The cells were grown in DMEM-based medium and decidualized *in vitro*. Total RNA was extracted from undifferentiated and differentiated cells using the RNeasy Mini

kit (Qiagen), followed by on-column DNase I treatment. RNA quality was assayed with the Bioanalyzer 2100 (Agilent) and samples with RIN  $> 9.5$  sequenced using the Illumina HiSeq2500, to a minimum of 36 million paired-end reads per sample. Reads were aligned with TopHat2 to the human Ensembl cDNA build, counted using HTSeq, and normalized as transcripts per million. Two technical repeats of each of the two primary cell cultures were processed independently and the expression values averaged over four transcriptomes.

For greater detail on tissue/cell preparation as well as sequencing, see Supplemental Methods.

### Immunocytochemistry

We used human trophoblast cell line BeWo (ATCC CCL98) and endometrial stromal fibroblasts (T HESC, Mor lab, Yale University, corresponding to ATCC CRL-4003). Uterine cells were decidualized *in vitro* by addition of cAMP and progesterone. Trophoblast cells were conditioned with decidualized cell-supernatant for 30 min, upon which the cells were fixed. Uterine non-decidualized and decidualized cells were also fixed and treated the same way. Immunocytochemistry was performed on all cells using anti-beta catenin primary antibody (Abcam, ab16051) and secondary fluorescent antibody (Abcam, ab150077).

### Analysis

#### *Single-cell transcriptomes*

The resulting transcriptomes were hierarchically clustered and subsequently analyzed for specific marker expression using AltAnalyze with default settings (<http://www.altanalyze.org>; Emig et al. 2010). Downstream analyses included the characterization of the cluster-specific gene expression with GO-terms (using GO Elite from Gladstone Institutes: [http://www.genmapp.org/go\\_elite/](http://www.genmapp.org/go_elite/)) to determine cell-type identities of the clusters. The cells from both batches are represented in all clusters, enabling us to combine the two batches in common analyses.

#### *Combining transcriptomes to explore cell communication*

For the cell-interaction analyses, the expression levels were recalculated relative to the total reads mapping to the same set of coding genes in all transcriptomes, on a TPM scale (Wagner et al. 2012). The expression values were averaged within each single-cell cluster/cell sample (Supplemental Table S1). The ligand-receptor relationships are based on data from two databases (Supplemental Table S5; <http://www.guidetopharmacology.org>; <http://dip.doe-mbi.ucla.edu/dip/DLRP.cgi>). Ligands and receptors above the cut-off threshold of 10 TPM were used to construct lists of putative interactions discussed, but only those beyond 30 TPM are shown in Figure 2, to allow better visualization. We used the freely available R package Circlize (Gu et al. 2014; <https://cran.r-project.org/web/packages/circlize/>) to script interaction plots with arrows connecting the ligands and corresponding receptors.

### Data access

Raw and processed data from this study have been submitted to the NCBI Gene Expression Omnibus (GEO; <https://www.ncbi.nlm.nih.gov/geo/>) under accession numbers GSE87692 (uterine cells) and GSE87726 (placental tissue-level transcriptome, single-cell transcriptomes, and syncytiotrophoblast transcriptome).

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**Author contributions:** L.M. and H.J. initiated the project; H.J. and K.O. collected and prepared placental tissue for single-cell analysis; S.G.K. collected tissue for bulk analysis; C.D.-F. extracted RNA from tissue samples; G.P.W. and A.R.C. performed the uterine cell culture experiments; A.R.C. and M.P. performed laser microdissection of syncytiotrophoblast; J.M. collected the data from the endometrial stromal fibroblasts and decidual cells; M.P. and G.P.W. analyzed the data and drafted the manuscript; all authors contributed to writing the manuscript.

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