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Up-regulated cytotrophoblast DOCK4 contributes to over-invasion in placenta accreta spectrum

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In humans, a subset of placental cells—cytotrophoblasts (CTBs)—invades the uterus and its vasculature, anchoring the pregnancy and ensuring adequate blood flow to the fetus. Appropriate depth is critical. Shallow invasion increases the risk of pregnancy complications, e.g., severe preeclampsia. Overly deep invasion, the hallmark of placenta accreta spectrum (PAS), increases the risk of preterm delivery, hemorrhage, and death. Previously a rare condition, the incidence of PAS has increased to 1:731 pregnancies, likely due to the rise in uterine surgeries (e.g., Cesarean sections). CTBs track along scars deep into the myometrium and beyond. Here we compared the global gene expression patterns of CTBs from PAS cases to gestational age-matched control cells that invaded to the normal depth from preterm birth (PTB) deliveries. The messenger RNA (mRNA) encoding the guanine nucleotide exchange factor, DOCK4, mutations of which promote cancer cell invasion and angiogenesis, was the most highly up-regulated molecule in PAS samples. Overexpression of DOCK4 increased CTB invasiveness, consistent with the PAS phenotype. Also, this analysis identified other genes with significantly altered expression in this disorder, potential biomarkers. These data suggest that CTBs from PAS cases up-regulate a cancer-like proinvasion mechanism, suggesting molecular as well as phenotypic similarities in the two pathologies.

During pregnancy, a subset of placental cells—the villous cytotrophoblasts (CTBs)—differentiate, acquiring the ability to invade the uterus and its vasculature (1–3). Normally, this CTB subpopulation migrates through the uterine lining (decidua), stopping forward progress at the inner third of the organ’s muscular layer (myometrium). Along the way, they invade and remodel the spiral arteries. Completion of this process produces vessels with a unique architecture; placental CTBs of embryonic/fetal origin replace the maternal endothelium and intercalate within their muscular walls. Consequently, these arteries undergo the very significant expansion that is required to carry the ever-increasing amounts of maternal blood that the fetus requires to sustain growth (4). In contrast, CTBs open the termini of uterine veins, which passively expand to accommodate larger and larger amounts of blood flowing back to the maternal circulation. Thus, CTB invasion achieves the physical and physiologic integration of the maternal-fetal unit that is required for normal pregnancy.

More is known about the mechanisms that CTBs use to invade the uterus than why they stop at a circumscribed location. In the majority of early gestation chorionic villi, CTBs form a polarized monolayer with basal attachments to the trophoblast basement membrane and apical contacts with the fused syncytiotrophoblasts (STBs) that are in direct contact with maternal blood. However, at numerous sites, often near the tips of chorionic villi, repulsive mechanisms—including tenasin-integrin and Eph-ephrin interactions (3)—propel CTBs away from the placenta into columns of cells that attach to the inner surface of the uterine cavity. The columns conduct the passage of the CTB subpopulation that is destined to invade the decidua, the myometrium, and the vascular tissue that traverses these regions.

Consistent with this dramatic change in fate, invasive CTBs undergo a unique molecular switch, involving mechanisms that have been implicated in the evolution of tumor cells from benign to malignant (5, 6). They execute an epithelial to vascular transformation, involving numerous adhesion, angiogenic/vasculogenic, and signaling molecules, which has also been described in aggressive forms of melanoma and breast cancer (7–9). Simultaneously they up-regulate matrix-degrading proteinases and many immune regulators, including human leukocyte antigen G (HLA-G), which together affect maternal tolerance of the semiallogeneic placental cells (7).

Unlike cancer cells, CTB penetration of the uterus is tightly regulated to prevent over-invasion. Placenta accreta spectrum (PAS) involves an apparent breakdown in the mechanisms that

Significance

The syndrome of cytotrophoblast invasion beyond the normal boundary (in the superficial myometrium) is collectively termed placenta accreta spectrum. The incidence of this condition is rising. However, little is known about the underlying molecular changes. Global transcriptomic profiling of cytotrophoblasts isolated from these cases, as compared to gestational age-matched controls, revealed numerous changes in gene expression involving diverse pathways, including cell signaling, migration, and immune functions. DOCK4 was the most highly up-regulated mRNA in the cases. Mutations in this gene are mechanistically linked to cancer progression. Overexpression of DOCK4 in primary cytotrophoblasts increased their invasiveness. This study provides molecular insights into the pathways driving placenta accreta spectrum and suggests numerous future directions.

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normally restrict advancing CTBs to the inner third of the myometrium (10, 11). As a result, they invade more deeply into the muscle layer of the uterus (placenta accreta), traverse it entirely (placenta increta), or reach the uterine serosa and beyond (placenta percreta) (12). PAS occurs when the decidua is thinner than normal or largely absent. Consequently, CTB invasion to the normal depth produces a deeply adherent placenta. Along the way, CTBs can remodel the spiral arteries to the level of the serosa. As a result, the normal process of placental separation from the uterine lining after delivery of the infant is not possible and attempts to do so can result in significant maternal hemorrhage and the risk of maternal mortality. These cases are often referred to tertiary care centers where surgical removal of the uterus is frequently necessary (13–15).

Placenta accreta was first described in 1937, when Cesarean sections (C-sections) were becoming more common (16). Epidemiological studies have identified prior uterine surgery, with C-sections being the most common type, as the greatest risk factor for the development of PAS (17, 18). In this case, invasive CTBs track along the scar, gaining access to the deeper portions of the uterus from which they are normally excluded. With C-section rates rising, the number of PAS cases is increasing in parallel, from an estimated incidence of 1:30,000 pregnancies in 1950 to more recent estimates of 1:731 (10).

Despite the rapidly increasing incidence, the disease process driving PAS remains poorly understood. For example, it is not known whether CTBs use their normal invasion mechanisms to penetrate the uterus to a greater depth or up-regulate other

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**Fig. 1.** CTB over-invasion and faulty decidualization at the maternal-fetal interface in PAS cases. Tissue sections from each case were: 1) stained with H&E (A, D, G, and J); 2) reacted with anti-CK to identify CTBs (B, E, H, and K); or reacted with anti-vimentin to identify decidual cells (C, F, I, and L). Gestational age-matched samples from nPTB cases served as controls. In nPTB cases, CK-positive CTBs were intercalated among vimentin-expressing cells of the decidua basalis (A–C). In equivalent PAS samples, CK-positive CTBs were evident in the near absence of vimentin-positive cells (D–F). The results from analyses of the basal plate region of additional case and control samples is shown in SI Appendix, Fig. S1. In nPTB, CTBs of the smooth chorion (scCTB) were juxtaposed with vimentin-positive cells of the decidua parietalis (G–I). In comparable specimens from PAS cases, the CK-positive CTB layer had a normal appearance with no signs of invasion (J–L). The results from analyses of the fetal membranes and adjacent decidua of additional case and control samples is shown in SI Appendix, Fig. S2. However, the adjacent maternal cells had low-to-no vimentin signals. PV, placental villi; BP, basal plate; FV, floating villi; FM, fetal membranes; SC, smooth chorion; MYO, myometrium. (Scale bar, 100 μm.)
cancer-associated molecules in the setting of an abnormal uterine environment. To begin addressing this question, we isolated CTBs from PAS cases and performed global transcriptional profiling. The results identified genes and pathways that were dysregulated in samples from the cases versus gestational age-matched controls. They included the guanidine nuclear exchange factor, DOCK4, mutations of which promote cancer cell migration and invasion (19, 20). When overexpressed in CTBs, DOCK4 increased invasion, evidence that PAS may involve a gain in the function of a molecule that promotes cancer progression as well as a loss of the normal architecture that limits placental penetration of the uterus.

**Results**

**Global Alterations in Decidualization Were Observed in the PAS Cases.** The photomicrographs in Fig. 1 illustrate the relative lack of decidua in biopsies of the maternal-fetal interface from PAS cases (n = 7). As comparators, we analyzed the distribution of decidual cells in biopsies of the maternal-fetal interface from pregnancies complicated by preterm birth with no signs of infection (nPTB, n = 5). Previously we showed that the distribution of CTBs and decidual cells is normal in this subset (21). Tissue sections from each case were: 1) stained with hematoxylin and eosin (H&E) (Fig. 1 D–F); 2) reacted with anti-vimentin to identify CTBs (Fig. 1 B, E, H, and K); or reacted with anti-vimentin to identify decidual cells (Fig. 1 C, F, I, and L).

First, we analyzed the basal plate, which underlies the site of placental attachment. Consistent with our published analyses of this region in nPTB cases, CK-positive CTBs were intercalated among vimentin-expressing cells of the decidua basalis (Fig. 1 A–C). In equivalent PAS samples, CK-positive CTBs were evident in the near absence of vimentin-positive cells (Fig. 1 D–F). Comparable data from additional cases are included in SI Appendix, Fig. S1.

Second, we examined the maternal-fetal interface at the junction of the fetal membranes and the decidua parietalis. In nPTB, CTBs of the smooth chorion were juxtaposed with vimentin-positive cells of the decidua parietalis (Fig. 1 G–J). In comparable specimens from PAS cases, the CK-positive CTB layer had a normal appearance with no signs of invasion (Fig. 1 J–L). However, the adjacent maternal cells had low-to-no vimentin signals. Thus, the decidual defect associated with PAS may be more widespread than previously thought, occurring well beyond the site of placental attachment. Comparable data from additional cases are included in SI Appendix, Fig. S2.

The deeply invasive CTBs of PAS still expressed typical markers. To begin molecular analyses of CTB invasion in PAS cases, we examined the cells’ expression of HLA-G, a stage-specific antigen that is up-regulated in the basal plate and the adjacent myometrium (n = 7). As was evident from H&E staining (Fig. 2A), many of the samples contained the entire uterine wall as they were from hysterectomies done at the time of delivery. Adjacent sections were double immunostained with anti-CK and anti-alpha smooth muscle actin (α-SMA), which enabled visualization of CTBs and uterine muscle cells, respectively (Fig. 2B). As illustrated by this case, placental cells were found throughout the muscle layer of the uterus. Despite the abnormal depth to which they invaded, the cells appropriately expressed HLA-G, which is up-regulated by CTBs within the uterine wall (Fig. 2C). Comparable data from additional cases are included in SI Appendix, Fig. S3.

**CTBs Isolated from PAS Cases Had a Unique Transcriptomic Signature.** Next, we asked whether PAS impacted CTB gene expression. The maternal characteristics of PAS cases used in these experiments are summarized in SI Appendix, Table S1. Consistent with epidemiologic data, all patients had had at least one prior Cesarean section (17). In these experiments, we isolated CTBs from the placentas of gestational age-matched PAS and nPTB cases, and compared their transcriptomes at a global level. In total, 118 genes were differentially expressed (DE) between the sample types; red dots denote up-regulated and blue dots denote down-regulated in PAS (Fig. 3A). Hierarchical clustering of the DE genes separated the two groups (Fig. 3B). The expression of 69 genes was higher in CTBs from PAS vs. nPTB cases; the expression of 49 genes was lower. The 25 most highly up-regulated known genes included a protocadherin cell adhesion molecule, PCDHGA10, a cell cycle regulator, CACUL1 (22), and cancer-associated molecules: DOCK4 (19), PTEN (23, 24), LGF (25), MITF (26), ATPSF1 (27), and ABL1 (28) (Fig. 3A and C). The 25 most highly down-regulated genes included RPL, an endogenous retroviral element (ERV2), a transcriptional activator that mediates fibroblast growth factor (FGF) signaling, CHURC1 (29), an insulin-like growth factor 1 (IGF) binding protein that inhibits growth, IGFBP6 (30), and cancer-associated molecules, PICK1 and LRR1C5 (31) (Fig. 3 A and C). Thus, we identified differences in gene expression that correlated with overly aggressive CTB invasion, including several cancer-associated molecules.

**Numerous Biologic Processes Were Dysregulated in PAS.** Gene ontology (GO) analysis of DE genes identified 23 enriched biological processes (P < 0.05; ≥3 DE genes/term; Dataset S1). Enriched terms included viral defense response, female pregnancy, negative regulation of cell proliferation, cell surface receptor signaling, neuron development, negative regulation of cell migration as well as metabolic processes (e.g., organophosphatase and carbohydrate derivative biosynthetic processes; Fig. 4A). The results suggested PAS was associated with alterations in a diverse array of disease-relevant biological processes.

In keeping with our previous work that showed molecules with important roles in neurogenesis pattern CTB invasion (3), the process of neuron development was populated by DE genes, including PTKEN (1), ABL1 (1), WNK1 (1), CNR1 (1), OMD (1) and OGN (1) (Fig. 4B). The category defense response to virus contained several genes with immune functions—B2M (1), RNF216 (1), MICA (1), IFIT1 (1), and BST2 (1) (Fig. 4C). Cell surface receptor signaling was also well represented among the DE genes: IGFBP5 (1) and IGFBP6 (1), ANGPT2 (1), CHDRL1 (1), DKK1 (1), and PRL (1) (Fig. 4D).

**qRT-PCR Confirmed a Subset of the Microarray Results.** We used qRT-PCR to validate dysregulated expression of specific genes identified in the microarray analysis (Fig. 5A). To examine the generalizability of our initial findings, we assayed two of the original PAS samples and five that were subsequently collected (n = 7). An equal number of CTBs isolated from nPTB cases served as controls. The results confirmed higher expression (P < 0.05) of DOCK4, B2M, PAPD4, and ANKR34D in PAS vs. nPTB; PRL was down-regulated. In general, the microarray and the qRT-PCR data were highly correlated (R² = 0.78; SI Appendix, Fig. S4).

**B2M Overexpression in PAS Was Confirmed at the Protein Level.** In two cases, we took the extra step of confirming that differences in mRNA expression were also observed at the protein level, which would increase the probability of the changes having functional relevance. Thus, we immunolocalized B2M in biopsies from PAS (n = 7) and nPTB (n = 5) cases. Representative results are shown in Fig. 5B with data from additional cases included in SI Appendix, Fig. S5. Higher immunoreactivity with anti-B2M was observed in association with CK-positive CTBs from PAS cases as compared to pregnancies that resulted in nPTB. Immunoblot analyses of CTB lysates corroborated these observations (n = 3 technical replicates; Fig. 5C; uncropped gels are shown in SI Appendix, Fig. S6A). On average, B2M was expressed...
DOCK4 Overexpression in PAS Was Also Confirmed at the Protein Level. Since the microarray analysis identified DOCK4 as the most highly PAS-associated DE gene in CTBs, we also confirmed up-regulated expression of this molecule at the protein level. Immunolocalization failed to detect binding of anti-DOCK4 to STBs or CTBs in floating villi from either cases or controls (Fig. 6A). Analysis of basal plate biopsies from pregnancies complicated by nPTB (n = 5) revealed a relatively faint but consistent signal that was associated with invasive CTBs, which was significantly stronger in equivalent PAS samples (n = 7). Data from additional cases are included in SI Appendix, Figs. S7 and S8. Immunoblotting confirmed and enabled quantification of this result (Fig. 6B and C; uncropped gels are shown in SI Appendix, Fig. S6B). CTB lysates from the increta (ICT) and percreta (PCT) cases had an immunoreactive band that corresponded to the molecular weight of DOCK4 (225 kDa); the additional bands are likely splice variants (32). CTB lysates from the accreta (ACT) and nPTB cases were immunonegative. Overall, DOCK4 expression was ~threefold higher in the setting of PAS as compared to nPTB (P < 0.05; Fig. 6C).

Overexpression of DOCK4 Increased CTB Invasion. Given the positive correlation between DOCK4 expression/activity and the progression of some cancers (33, 34), we overexpressed this molecule (~7.5-fold, P < 0.01) in primary CTBs and analyzed its function in terms of CTB invasion (Fig. 7A, quantified in Fig. 7B; uncropped gels are shown in SI Appendix, Fig. S6C). Up-regulation of DOCK4 increased CTB invasion in all three independent experiments (P < 0.01) and on average by ~3.7-fold as compared to the level observed in the control cultures (P < 0.001; Fig. 7C).

Discussion
Here we report the results of using a global transcriptional profiling approach to identify the potential mechanisms driving overly aggressive CTB invasion in PAS. In total, we identified 118 DE genes as compared to gestational age-matched control nPTB cases in which CTB invasion and uterine decidualization were ostensibly normal. GO analyses suggested that they participated in numerous biological processes that could contribute to the cancer-like properties of the cells, which traverse the myometrium and sometimes encroach on neighboring organs such as the bladder. In this context, it was logical that the enriched terms were related to processes that are integral to cell migration/invasion, cell signaling, and maternal-fetal immune tolerance.

This analysis identified DOCK4 as the most highly DE/up-regulated gene in CTBs isolated from PAS placentas. This membrane-associated guanine nucleotide exchange factor enhances the formation of adherens junctions between cells (19). Mutations that interfere with the molecule’s ability to activate RhoGTPases, required for the assembly of these specialized attachment sites, are associated with the progression of many tumor types (19, 20). DOCK4 also augments tumor angiogenesis by enabling the complex process by which the lumen of a blood vessel forms (35). The primary mechanism involves a pathway in which RhoG acting through DOCK4 triggers a signaling cascade involving Rac1, DOCK9, and Cdc42. Here we discovered CTB overexpression of DOCK4 in the cancer-like setting of PAS. Whether this is in response to mutations such as those described in certain cancer types remains to be determined. If this is not the case, it appears that DOCK4 must have unique functions in CTBs as compared to other cells as its overexpression significantly increased invasion rather than cell-cell adhesion, which should limit this process. In this regard, our findings are more in line with recent reports that up-regulation of DOCK4 expression is associated with breast and lung cancer metastases (33, 34).

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Beyond DOCK 4, our analyses suggested PAS-associated dysregulation of other genes with suspected roles in cell signaling and migration/invasion in the context of cancer, including BST2 (36, 37), ANGPT2 (38, 39), and IGFBP5 (40). While the specific roles of these genes in CTBs are not yet understood, these molecules may drive the hyperinvasive state of CTBs in PAS placentas and will be an interesting area of investigation for future studies. If a less restrictive model is used for the transcriptomic analysis, additional pathways of interest are identified (Dataset S2). For example, a subset of the genes involved in microtubule movement are highlighted. These proteins have diverse roles in the primary cilium, including ARL3 [signal transduction and function (41)]; DYNC1L1 and DYNC1I2 [protein transport (42)]; VAMP7 [cilia length (43)]; and spastin [microtubule cleavage (44)]. Although the specifics are not yet understood, the primary cilium has a role in regulating the cell cycle and dysfunction has been observed in multiple cancers (45). These cell surface specializations are reported to facilitate invasion of a transformed trophoblast cell line (46). Given our findings, the functions of primary cilia in CTBs from PAS placentas is another potential area of investigation.

One explanation for the increased CTB invasiveness in PAS is lack of decidualization at the placental attachment site. We confirmed this longstanding observation in the cases that were collected for this study. Thus, the decidual signals that may normally constrain invasion are missing in PAS. Here we provide preliminary evidence that this defect may be more widespread than previously reported as there was also a lack of vimentin-positive decidual cells adjacent to the CTBs that formed the smooth chorion layer of the fetal membranes. We speculate that one reason could be an abnormal wound healing response to uterine surgery—e.g., C-section or myomectomy—a risk factor for PAS. This could explain why only some women who undergo these procedures subsequently have overly invasive placentas. The fact that the smooth chorion CTB subpopulation, unlike those of the placenta proper, fails to invade highlights the cells’ fundamental lack of cancer-like potential despite their extravillous location.

Our data suggested that in addition to a PAS-associated decidual defect, there were also CTB aberrations that manifested as changes in gene expression. Whether this is evidence of autocrine dysfunction or a paracrine response to absent, deficient, or

Fig. 3. Transcriptomic profiling revealed PAS-associated CTB gene signatures. We compared the gene expression profiles of CTBs from PAS (over-invasion; \( n = 6 \)) and gestation-matched nPTB cases (normal invasion; \( n = 3 \)). There was no statistically significant difference in the gestational ages between the two sample sets (\( P = 0.11 \)). (A) The statistical significance versus gene expression fold change is displayed as a volcano plot. In total, 118 genes were DE between the sample types (blue dots down-regulated and red dots up-regulated; \( P < 0.05 \)). (B) Hierarchical clustering of the DE genes separated the two groups. (C) The 25 most highly up-regulated (Top cluster) and down-regulated genes (Bottom cluster) are shown. OE, overexpressed.
deranged decidual signals remains to be determined. Of note, the CTBs we analyzed were primarily (but not exclusively) from the placenta; therefore, largely a villous rather than extravillous/invasive population. The fact that differences in gene expression as compared to the control cells were readily apparent prior to invasion suggested that “precancerous” changes may be taking place before the CTBs leave the placenta and enter the uterine wall. We interpret these data as favoring the theory that CTBs from PAS placentas have cancer-like features that include overexpression of genes that play a role in tumor progression.

In this analysis, we considered CTBs from PAS cases to be one experimental group as we had too few of each subtype to draw additional conclusions. Relative quantitation of DOCK4 immunoblot data showed lower CTB expression in the less invasive subtype (accreta) as compared to the more aggressive forms of PAS (increta and percreta). These preliminary data suggest the value of comparing the different forms of this syndrome in terms of both CTB and decidual effects.

The molecular differences we identified between the cases and controls are possible evidence of biomarkers that could be used...
to more reliably screen for and identify pregnancies complicated by PAS. Overall, this diagnosis has significant implications for the mother and fetus. Most often, this includes a preterm delivery to decrease the risk of spontaneous labor and hemorrhage as well as a hysterectomy at the time of C-section. Prenatal diagnosis is crucial so that affected women can deliver at a tertiary care center and with proper planning (14). This decision is commonly made based on prenatal imaging, such as ultrasound or magnetic resonance imaging (MRI). A recent meta-analysis showed the sensitivity of ultrasound to be 81 to 93% (47). Thus, there remains room for improvement, a critical issue given the repercussions of a missed diagnosis. As the incidence of PAS rises, the importance of developing a diagnostic biomarker panel that could be assayed using a maternal blood or urine sample increases. This seems feasible given our discovery of numerous CTB genes that are dysregulated in this condition and the fact that excessive invasion of the uterus and its arteries should result in higher circulating levels of placental proteins. Such a test would improve outcomes by allowing appropriate planning in women with PAS and avoiding unnecessary anxiety and procedures in women who do not.

Materials and Methods
Collection of Placentas from Cases and Controls. The University of California San Francisco Institutional Review Board (Committee on Human Research) and the Stanford Institutional Review Board approved this study. Written informed consent was obtained from all donors. Women with preexisting medical conditions such as (but not limited to) thyroid insufficiency, chronic hypertension, or diabetes mellitus were excluded from the case and the control groups. Other exclusion criteria included premature rupture of membranes and/or fetal anomaly. Cases were identified based on prenatal imaging—ultrasound or MRI—consistent with an invasive placenta (47). Intraoperatively, findings were consistent with PAS as the placenta did not separate after delivery. Final confirmation of this diagnosis was made by a gynecologic pathologist. Gestational age-matched control samples were obtained from women who had a preterm birth (PTB), diagnosed according to the criteria recommended by Herron et al., including regular uterine contractions at ≥20 or <37 wk of gestation, which were 5 to 8 min apart and accompanied by one of the following events (1): progressive changes in the cervix (2), cervical dilation ≥2 cm, and/or (3) cervical effacement ≥80% (48). Patients with evidence of inflammation were excluded on the basis of the following criteria: maternal fever >100.4°F, uterine tenderness, fetal tachycardia (fetal heart rate >160 beats per min), and (placental) histologic criteria compatible with inflammation.

CTB Isolation. CTBs were isolated as previously described (1). The basic steps included extensive washing of the chorionic villi, sequential digestion with collagenase/trypsin, and isolation via Percoll density gradient centrifugation. For the first three isolations, CTB purity was confirmed by immunostaining for the CTB marker, cytokeratin 7, which showed ≥90% purity. The morphology of the cells in subsequent preparations was consistent with this result. A portion of each CTB preparation was transferred to TRIzol for RNA extraction or to medium salt lysis buffer plus 1:100 protease inhibitor (Roche) and 200 mM diithiothreitol (DTT) for preparation of protein lysates.

Histology. Placental and placental bed biopsies from cases and controls were fixed in paraformaldehyde, embedded in optimal cutting temperature compound (Fisher Scientific), and stored at −80 °C. Sections (6 μm) were stained with H&E. Images were obtained using a Leica inverted microscope and processed in Photoshop (Adobe).
Global RNA Expression Profiling. The microarray analysis used methods that we published (49). Total RNA was isolated using the TRIzol reagent according to the manufacturer’s directions. The RNA concentration was determined via a Nanodrop spectrophotometer (Thermo Scientific), and quality was assessed using an Agilent 2100 Bioanalyzer. Only samples with a RNA integrity number (RIN) greater than 7 were used in subsequent analyses. We used Affymetrix Human Gene 2.0 ST arrays to profile the global gene expression patterns of CTBs isolated from gestational age-matched PAS cases ($n = 6$) vs. placentas from pregnancies that ended in nPTB ($n = 3$). Sample processing and hybridization was performed by the University of California San Francisco (UCSF) Gladstone Institute as previously described (49). Affymetrix CEL files were processed and annotated using the Oligo software package (50). Raw values were normalized via the Robust Multi-Array Average (RMA) algorithm. Raw and normalized data were deposited in the Gene Expression Omnibus (accession no. GSE136048). We applied linear models for microarray data (LIMMA) to identify DE genes. If there were multiple probes per gene, we used the probe with the lowest P value (differences between cases and controls) for downstream analyses. We selected probes with average log2 intensities > 3 to remove nonspecific probes/low abundant transcripts (lower ~15th percentile), and to limit the detection of false positives in our downstream analyses. In addition, only probes linked to an Official Gene Symbol [HUGO Gene Nomenclature Committee (HGNC)] were included. In total, we evaluated the expression of 20,289 unique genes. We calculated the ratio of average log2 intensities between PAS and nPTB to determine the average fold change (FC) difference between groups. DE genes were defined by applying a cutoff (unadjusted $P < 0.05$) and an absolute fold change > 0.5 (log2 scale), which distinguished the two groups. Clustering of relative expression values was performed by Pearson correlation. We utilized the Database for Annotation, Visualization and Integrated Discovery (DAVID) to perform GO enrichment analysis of DE genes. Functional enrichment analysis of Gene Ontology (GO) Biological Processes was performed for all DE genes via DAVID (uncorrected $P < 0.05$; number of DE genes associated with enriched term $\geq 3$) (51). DAVID enables evaluation of enrichment of GO terms at five levels (1 to 5) of ontology based on size and specificity (51). Enrichment analysis was performed at all levels, although we only reported Level 4 Biological Processes—a category of moderate to high specificity within the GO organizational structure—to reduce redundancy of categories with similar functions.

Validation of DE Genes Using TaqMan qRT-PCR. cDNA libraries were prepared with 500 ng RNA using the iScript Kit (Bio-Rad) and diluted 20-fold in water. For qRT-PCR, we employed TaqMan Universal Master Mix II, no AmpErase Uracil N-Glycosylase (UNG) (Life Technologies), and TaqMan primers for DOCK4, PRL, ANKR44, B2M, PAPD4, GAPDH, and ACTB (SI Appendix, Table 2A).
Reactions were carried out for 40 cycles. Seven biological and three technical replicates were compared for all targets. Differences among target expression levels were estimated by the ΔΔCT method with normalization to the geometric mean of two housekeeping genes, GAPDH and ACTB. Differences between means were assessed using a 2-tailed Student’s t test (P = 0.05), assuming unequal variance.

**Immunolocalization and Immunoblotting.** The experiments were performed as previously described (52, 53). Sections of seven case and five control samples were examined for the expression of each antigen. Antibody reactivity enabled identification of the following cell types using our published methods: cytokeratin antibody (7D3), which enabled the counting of CTBs that were transduced with the DOCK4-containing vector as compared to control cells that received the empty vector (n = 3 biological replicates). Asterisk and double dagger symbols indicate significant differences between DOCK4+ and control within each experiment (P < 0.01).

**DOCK4 Overexpression and CTB Invasion.** Invasion assays were performed as previously described (53). Briefly, 250,000 CTBs isolated from second trimester placenta as described above were plated in Transwell inserts (6.5 mm; Costar Corp.) on Matrigel-coated polycarbonate filters (8-µm pores). To up-regulate DOCK4 expression, the open-reading frame of DOCK4 (constitute from ORI-GENE) was inserted into an adenovirus vector (Vector Biolabs). At the time of plating, the CTBs were transduced with the DOCK4 adenovirus vector or an empty adenovirus vector, which served as a negative control. An aliquot (2.5 µL of Ad-hDOCK4 or the empty vector (106 plaque forming units/mL) was added in medium (final volume = 0.25 mL) to each well (multiplicity of infection = 100). After 36 h, the Transwell filters were stained with an anti-cytokeratin antibody (7D3), which enabled the counting of CTBs that reached the filter’s underside.

To evaluate the data, we applied Student t tests to examine pairwise differences in cell counts between DOCK4+ and control cultures within each experiment. We also assessed differences across experiments utilizing a generalized linear regression model to control for differences in baseline activity (n = 3). Relative % invasion was expressed as the ratio of the average number of cell projections per filter in control vs. DOCK4-transduced wells. All experiments included at least three technical replicates. DOCK4 overexpression was confirmed by immunoblotting as described above.

**Data Availability.** Raw and normalized data were deposited in the Gene Expression Omnibus (accession no. GSE136048) as described in the Methods section Global RNA Expression Profiling.

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