Endometrial Gap Junction Expression - Early Indicators of Endometriosis and Integral to Invasiveness

Chen-Wei Chen #,1, Jeffery Chavez #,1, Li-Ling Lin2, Chiou-Miin Wang2, Ya-Ting Hsu2, Matthew J. Hart1, Jianhua Ruan3, Laurel Gillette 4, Richard O. Burney4, Robert S. Schenken5, Randal D. Robinson5, Maria Gaczynska2, Pawel Osmulski2, Nameer B. Kirma2*, Bruce J. Nicholson1*

1 Department of Biochemistry and Structural Biology, UT Health San Antonio, Texas, USA
2 Department of Molecular Medicine, UT Health San Antonio, Texas, USA
3 Department of Computer Science, UTSA, San Antonio, Texas, USA
4 Madigan Army Medical Center, Tacoma, WA.
5 Department of Obstetrics and Gynecology, UT Health San Antonio, Texas, USA

# Equal contributions as first authors
*Co-corresponding authors

kirma@uthscsa.edu (NBK)
nicholsonb@uthscsa.edu (BJN)

Abbreviated Title: Gap junctions and invasiveness in endometriosis

Abstract

Endometriosis is an invasive disease, and a leading cause of pain, infertility and disability among women, with an incidence 10 fold that of cancer. A more complete understanding of disease pathogenesis is essential for the development of non-surgical diagnostic assays and non-hormonal therapeutics. Avoidance of immune clearance and implantation of endometrial tissue on peritoneal surfaces are features of endometriosis lesion formation that overlap with cancer metastasis. Connexins, and the gap junctions they form, have been implicated in cancer progression, and may be associated endometriosis pathophysiology. Single cell transcriptomic profiling of endometrial epithelial and stromal cells from women with endometriosis reveals a striking and progressive shift in expression of connexins and related regulatory and junctional genes. We demonstrate that gap junction coupling between endometrial cells and the peritoneal mesothelium is dramatically induced, specifically in endometriosis patients, and is required for invasion by inducing breakdown of the mesothelial barrier function.
INTRODUCTION

Endometriosis is a chronic inflammatory disease affecting 6-10% of reproductive age women (Eskenazi and Warner, 1997). Characterized by the presence of endometrial tissue in extraterine locations including the pelvic peritoneum, ovary and bowel surface, endometriosis is diagnosed in 35-50% of women with pelvic pain and up to 50% of women with unexplained infertility (Rogers et al., 2009). At an estimated annual cost of $12,000 per patient in terms of diagnosis and treatment, and adding in the significant loss of productivity, endometriosis care entails significant socioeconomic burden for both individual patients and healthcare systems. Estimated to cost $80 billion per year for the US alone (Soliman et al., 2016). In the absence of a biomarker, laparoscopic surgery remains the gold standard for diagnosis. The requirement for invasive surgery, which fails to confirm endometriosis almost half the time (Mettler et al., 2003) contributes to an average latency of 6.7 years from onset of symptoms to definitive diagnosis (Bontempo and Mikesell, 2020), and results in 68% of women suffering from endometriosis being incorrectly diagnosed (Hudelist et al., 2012). Diagnostic delay allows time for disease progression, and potentially worsens sequelae and prognosis. An improved understanding disease etiology is critical to developing new diagnostics and therapies.

The original, and still most widely accepted, model for the pathogenesis of endometriosis is retrograde menstruation, in which sloughed endometrial tissue during menses traverses the fallopian tubes and forms invasive lesions within the peritoneal cavity (Sampson, 1927). Peritoneal origins have also been proposed (Mismer et al., 2004; Sasson and Taylor, 2008), but the preponderance of evidence still favors endometrial origins (reviewed in Burney and Giudice, 2012). Abundant evidence supports molecular differences in the eutopic endometrium of women with and without endometriosis (Hastings and Fazleabas, 2006, Tamaresis et al., 2014). Since retrograde menstruation is estimated to occur in 76-90% of women, but only 6-10% develop endometriosis, the interaction of displaced endometrium and the peritoneum is a key differentiating step in disease pathogenesis. Peritoneal factors also influence disease progression, including the hormonal environment (Parente Barbosa et al., 2011), oxidative stress and inflammation (Augoulea et al., 2012) or decreased immune clearance (Oosterlynck et al., 1991), although this could be due to changes in the endometrial cells themselves (Somigliana et al., 1996). This has fueled the debate over whether endometriosis originates from changes in the uterus that predispose the cells to lesion formation (the “seed” model) or if it is a property of a receptive peritoneal environment (the “soil” hypothesis).

To understand the molecular underpinnings of endometrial-peritoneal interaction in lesion formation, we have focused on a class of proteins that has been implicated in tissue invasion, infertility and inflammation in other contexts, but incompletely explored in the pathophysiology of endometriosis. Gap junctions, composed of connexin (Cx) proteins encoded by a family of 21 GJ (A-D) genes, mediate direct contact and communication between most cells of the body via exchange of ions as well as metabolites and signaling molecules <1kD (Goldberg et al., 1999; Weber et al., 2004). Gap junctions have been shown to be essential to many invasive processes, both normal (e.g. blastocyst implantation – Grummer et al., 1996) and pathogenic (e.g. metastasis). Increased Cx43 and Cx26 expression and gap junction intercellular communication (GJIC) have been associated with metastasis and poor prognosis in breast (Kanczuga-Koda et al., 2006; Naoi et al. 2007; Stoletov et al., 2013) and prostate cancers (Zhang et al., 2014; Lamiche et al., 2012), and elevated Cx26 with melanoma metastases (Ito et al., 2000). Connexins...
exert their effects both during intravasation and extravasation (as cells enter and leave the circulation) [el Sabban and Pauli, 1991, 1994; Ito et.al, 2000, Naoi et.al, 2007], as well as establishment of distant lesions by forming heterotypic GJIC (Lamiche et.al., 2012; Stoletov, et.al., 2013; Hong et.al., 2015; Chen et.al., 2016). In a global screen of cervical squamous carcinoma, Cx43 emerged as one of three genes (along with PDGFRA2 and CAV-1) central to cancer invasion and metastasis (Cheng et.al., 2015).

Gap junctions are also critical for a number of steps in human fertility (reviewed in Winterhager and Kidder, 2015). Endometrial gap junctions, comprised of Cx43, are essential for decidualization [Kaushik et.al., 2020], blastocyst implantation [Grummer et.al., 1996; Diao et.al., 2013] and vascularization and endometrial development during pregnancy [Laws et.al., 2008]. Connexins have also been linked to induction of inflammatory processes that either inhibit tissue repair (Willebrords et.al., 2016), such as wound healing of the skin (Montgomery et.al., 2018) and cornea (Ormonde et.al., 2012) processes thought to be driven by ATP release through connexin hemichannels (Mugisho et.al., 2018).

With regard to endometriosis, prior immunohistochemistry studies demonstrated a shift in Cx expression of endometrial epithelial cells (EECs) from primarily Cx26, with some Cx32 (GJB1), in the uterus to Cx43 in peritoneal (ectopic) endometriotic lesions (Regidor et.al., 1997). By contrast, endometrial stromal cells (ESCs) show predominantly Cx43 expression in both eutopic and ectopic locations, although at reduced levels in endometriosis patients (Yu et.al., 2014). This modified expression profile of connexins seen in ectopic lesions of women with endometriosis was recapitulated in eutopic endometrial tissue in an endometriosis model in baboons (Winterhager et.al., 2009), suggesting that early changes in the endometrium might predispose refluxed menstrual endometrial tissue within the pelvic cavity for invasiveness (Guo et.al., 2004; Lucidi et.al., 2005; ).

Our objective in this study was to characterize the contribution of Cxs within the endometrial compartment to its invasive potential in the peritoneum, an essential process for endometriosis lesion formation. We have examined the connexin gene expression profiles in both endometrial stromal (ESC) and epithelial (EEC) cell compartments, using single cell analysis of tissue obtained by endometrial pipelle biopsy from women with and without endometriosis. . We also performed functional studies by assessing GJIC in both homotypic ESC and EEC cultures, as well as heterotypic co-cultures with peritoneal mesothelial cells (PMCs). We finally assessed the role of the primary gap junction (GJ) gene product, Cx43, in the trans-mesothelial invasive process by ESCs and EECs that is essential for endometriosis lesion formation. These studies provide unique insights into the endometrial-peritoneal interaction in lesion formation, and represent the first single cell transcriptomic analysis of connexins and gap junctions in endometriosis.

RESULTS

Expression profiles of Intercellular Interaction Genes in Endometrial Stromal and Epithelial Cells from Women with and without Endometriosis

While studies have addressed different aspects of endometriosis, from changes in hormonal responses and other contributors to infertility to inflammatory responses caused by peritoneal lesions, and some even on lesion formation, none have focused on what common themes may tie all these symptoms together. Given the links, described above, of connexins and gap junctions to each of these processes, we examined their expression patterns, and those of related genes involved in intercellular interactions and their regulation in the endometrial lining of normal and endometriosis patients by analysis of freshly isolated primary cells (i.e. P0). Specifically, the gene panel included: 13 of the total of 21 Cx genes, including the major expressers in the endometrium,
Cx43 (GJA1), 26 (GJB2) and 32 (GJB1), and at least one from each Cx class (A, B and C); adhesion proteins [N-cadherin (CDH2) and Ep-CAM], cytoplasmic proteins associated with junctional complexes [Drebin (DBN1), β-catenin (CTNNB1), caveolin (CAV1), TJAP and ZO1 and 2 (TJP1 and 2)]; regulatory components of junctions, such as transcription factors [Snail 1 (SNAI1)] and kinases (e.g. PKA, PKC, and MAPK isoforms, CK1, SRC, CDK1); components of the extracellular matrix (MME, NOV); housekeeping (UBB, GAPDH, ACTIN, and GUSB) and cell marker (Vimentin (VIM) and cytokeratin (KRT18)) genes. The genes, in the array, in order of presentation in the heat map, are provided in Table S1. Those shaded in grey were control genes for data normalization and are not presented in the arrays.

To optimally understand how changes in patterns of expression might contribute to disease etiology, we separated the two major endometrial cell types [epithelial or glandular (EECs) and stromal (ESCs)], the purity of which was demonstrated to be ~95% by immunocytochemistry (Fig. S1). To further maximize detection of micro-heterogeneities in expression patterns, we used the Fluidigm C1 single-cell capture system followed by the Biomark microfluidic PCR to examine expression at the single cell level within each population. Two normal subjects and 6 endometriosis patients (one in early stage (I/II), and the others later stage (III/IV) endometriosis) were examined, with samples taken from proliferative, early and late secretory menstrual phases, plus one non-cycling patient on birth control (Table 1). Normalized expression results for each cell type from each patient are displayed as a heat map, with each row representing a gene, and each column a cell (Fig. 1A, B). Distinct patterns of expression in stromal (ESCs) and epithelial cells (EECs) were evident, which in this patient sampling clearly distinguished normal from endometriosis samples, and even showed a gradual transition as disease progressed from early to late stages (Fig. 1A, B).

(a) Gap Junction genes: With respect to genes encoding connexins (GJ genes – located at the top of the maps), a subset of ESCs (~25% of total analyzed cell population) showed high relative expression of all GJ genes in normal subjects, which diminished somewhat in early stage (I/II) endometriosis samples, and drops to less than 10% in late stage (III/IV) endometriosis, regardless of the menstrual phase from which the sample is collected (Fig. 1A). Even though single EEC cell preparations could not be made from all patients, it was clear that the pattern was completely reversed compared to ESCs, with low expression of GJ genes in normal subjects (5% of cells show higher expression), which progressively increases in early (I/II) and late stage (III/IV) endometriosis where ~25% of cells show high expression (Fig. 1B). It is striking that this is true for all GJ genes, and not just the most abundantly expressed, like GJA1 (Cx43). The combined expression levels of all GJ genes is illustrated in Violin plots for ESCs (Fig. 1C) and EECs (Fig. 1D) from each patient, with similar plots for specific GJ genes shown in Fig. S2 [top two rows of A (ESCs) and B (EECs)]. A Duncan analysis of the single-cell distributions shows that ESCs from all late stage endometriosis patients had significantly lower expression than ESCs from normal subjects and early stage endometriosis patients. The notable exceptions to this pattern in ESCs were those taken at the mid-secretory menstrual phase when progesterone levels, known to regulate connexin expression in the endometrium (Grummer et.al., 1994), are high [subjects 171 (normal) and 169 (late endometriosis)]. EECs show the inverse pattern that was even detectable in early stage endometriosis (patient 164), In EECs, the pattern was not influenced by the stage of the menstrual cycle when samples were collected, although did seem to be affected in the one non-cycling patient on birth control (003).

Comparison of individual GJ gene expression (Fig. S1) showed that GJA1 (Cx43) was expressed most highly in both ESCs and EECs, almost 10 fold that of other connexins. For ESCs, all GJ genes measured, except GJC2, showed decreases in late stage endometriosis compared to normal subjects and early stage patients (Fig. S1A). In EECs, most of the GJ genes show the inverse behavior seen in ESCs, increasing expression in endometriosis compared to that seen in...
normal subjects. The exceptions were GJA1, the most highly expressed connexin, GJA8, GJB7 and GJC2 (Fig. S1B).

(b) Other genes: Several other genes involved in intercellular or cell-matrix interactions were also found to change in similar pattern to that of the GJ genes. The adhesion related genes EpCAM, β-catenin (CTNNB1), as well as NOV1 show decreased expression in ESCs in endometriosis (Fig. 1E) but increase in EECs (Fig. 1F), while the master transcriptional regulator of epithelial to mesenchymal transitions (EMT), Snail 1 (SNAI1), which negatively regulates several adhesion molecules (Hugo et al., 2011) and Cx43 (deBoer et al., 2007), shows the inverse pattern (increasing with endometriosis in ESCs and decreasing in EECs). Several other genes change in one or other cells type. In ESCs, the metallo-endoproteinase, CD10 (Fig. S2A), NOV1 and the catalytic subunit of protein kinase A gamma (PRKACG) decrease, and ZO2 (TJP2), an accessory protein of tight and gap junctions, increases with disease (Fig. 1E). In EECs, only MAPK1 showed consistent increase in endometriosis patients (Fig. 1F). Some other genes showed increases in the two endometriosis patients in the proliferative or early secretory stages of their menstrual cycle [N-cadherin (CDH2), (Fig. 1F), and PRKACG (Fig. S2B)], but markedly different patterns in the non-cycling patient (Fig. S2B). Independent of these overall patterns, it is notable in the heat maps that the same subset of ESCs from the normal and early endometriosis samples (left three maps in Fig. 1A), and EECs from endometriosis patients (right two maps in Fig. 1B), that show high GJ genes expression, exhibit low expression of several of the tight junction associated genes, but higher expression of adhesion related genes (upper and lower rows, respectively in the group of genes labeled Adh/TJs) (Fig. 1A and B). Similar to what was observed for the tight junction associated genes, several of the regulatory genes (mostly kinases) we tested show the lowest expression in the cells that exhibit the highest expression of GJ genes (left three maps of ESCs (Fig. 1A) and right two maps if EECs (Fig. 1B).

Cell cluster analysis

As an independent means to assess changes in gene expression in endometriosis, intra-sample cellular heterogeneity was analyzed by a graph-based cluster discovery algorithm (Ruan, 2009). By design, the algorithm is able to identify topologically distinct clusters and, importantly, can automatically determine the most appropriate number of clusters for each dataset. Utilizing this algorithm, ESCs and EECs were divided into 6 and 5 sub-populations, respectively (Fig. 2A and C). This analysis confirmed the general impression from the Heat maps, that the cell sub-groups characterized by high GJ gene expression (sc-6 for ESCs, and ec-5 for EECs) consistently decreased as a percentage of the ESC population (Fig. 2B), but increased as a percentage of the EEC population (Fig. 2D) from normal through early stage to late stage endometriosis. This cell cluster most reliably tracked the progression of endometriosis. Stated differently, the abundance of cells showing combined GJ expression levels at the 80th percentile or above are highly predictive of disease state. For example, in ESCs all control subjects show >20%, early endometriosis 5-8% and late stage endometriosis <4%. This pattern was reversed in EECs, although we will need more patient samples to establish specific ranges.

Functional assessment of GJIC in ESCs and EECs, with endometriosis progression

We next assessed the functional consequences of these changes in GJ gene expression by measuring GJIC using an automated variant of the “parachute” technique where dye (calcein) loaded donor cells (D) are dropped onto a monolayer of acceptor cells (A) and the degree of dye spread to the monolayer (A/D ratio) is measured over 10-15 fields in an automated confocal microscope over time (Fig. 3 A and B). Given the changes in GJ gene expression between normal and endometriosis patients in Fig. 1, it was surprising that GJIC did not change significantly in either cell type, although a small decrease was seen in ESCs and increase in EECs (Fig. 3C).
However, as endometriosis is characterized by invasive behaviors following interactions with the peritoneal mesothelium, we also assessed heterotypic GJIC between endometrial cells and a peritoneal mesothelial cell (PMC) line (LP9). This revealed a dramatic induction of GJIC that was most notable in ESCs from endometriosis patients (4 fold increase compared to 2 fold in normal subjects). While some induction was also observed in EECs, this was not significant in either normal or endometriosis samples (Fig. 3C). This induction occurred relatively rapidly, within the context of the assay (~ 2 hours).

Given this time course, which seemed inconsistent with a transcriptional event, we examined the distribution of Cx43 protein in ESCs alone, and after contact with PMCs. Alone, both stromal (Fig. 3D) and mesothelial cells (not shown) show primarily intracellular Cx43 staining. By contrast, in heterotypic cultures, very little intracellular staining is seen within ESCs (fluorescent green labelled cell in Fig. 3E), but plaques are readily observed at sites of contact with PMCs (yellow arrows, Fig. 3E), although significant intracellular labeling within PMCs was still evident. This suggests that the heterotypic contact may trigger enhanced trafficking of Cx43 to the cell surface.

Since the major functional difference we observed related to gap junctions in samples from endometriosis patients were associated with heterotypic contacts with mesothelial cells, especially by ESCs, we predicted that they may influence differential adhesion between the cells. To measure the degree to which heterotypic adhesion properties with mesothelial cells differ between EECs and ESCs, we used atomic force microscopy (AFM) that enables quantitative assessment of the extent of intercellular interactions (Sancho et al., 2017; Roca-Cusachs et al., 2017). Adhesiveness between PMCs (LP9 cells), and EECs and ESCs from endometriosis patients was measured by attaching a single cell to a cantilever of an AFM probe (Fig. 4A). The attached cell was placed in contact with a monolayer of either the same cell, or a different cell type, growing on a culture dish, and the force needed to separate the probe-attached cell from the plate-attached cell was measured (Fig. 4B). Mesothelial cells show low levels of adhesion to one another that was similar to their adhesion to EECs. In contrast, ESCs showed significantly higher adhesion to PMCs (Fig. 4C). This data is consistent with ESCs being the cell type that is the major invasive front into the mesothelium (Nair et al., 2008; Burney and Giudice, 2012).

**Invasive Potential of Endometrial Cells from Normal Subjects and Endometriosis Patients, and dependence on GJIC**

We then directly assessed the invasiveness of primary endometrial cells from normal and endometriosis patients using an established 3D-invasion assay that assessed the efficiency with which ESCs or EECs can pass through a confluent monolayer of PMCs grown on a growth-hormone depleted Matrigel-coated membrane (Fig. 5A). Initial studies using unseparated endometrial cells showed significantly greater invasiveness in samples from endometriosis patients than control subjects (n=3; Fig. 5B). For ESCs (n=3 and 6 for control and endometriosis, respectively) and EECs (n=3 and 6 for normal and endometriosis, respectively) the differences were not significant. In all cases tested the level of invasion was not seen when endometrial cells were dropped on a membrane alone, so that the invasion was dependent on contact with a mesothelial monolayer (data not shown).

Given that we had demonstrated that endometrial cell contact with mesothelial cells specifically induced GJIC, we tested the degree to which invasive behavior was dependent on GJIC by targeting Cx43, the dominantly expressed Cx in both cell types (Fig. S2). Four approaches were used to test this. Firstly, we pre-treated both PMCs and endometrial cells (not separated into ESCs and EECs) with GAP27, a peptidomimetic to the extracellular domain of Cx43 that we, and many others (Evans and Leybaert, 2007), showed blocked the formation of gap junctions between newly contacting cells, in this case by ~85%. This reduced
invasiveness of unseparated endometrial cells from control subjects by 57% (although not significant at the 0.05 level), and from endometriosis patients by 70% (p<0.01) (Fig. 6A).

Secondly, we tested purified ESCs (as only they showed increased adhesiveness to PMCs (Fig. 4)), using a combination of two siRNAs targeted to Cx43, where the PMC monolayer was transfacted immediately prior to the invasion assay. This caused an average of 65 ± 9% reduction in Cx43 protein (Fig. 6B) and ~70% reduction in GJIC between endometrial and mesothelial cells (p<0.001 in control and <0.0005 in endometriosis) (Fig. 6C). Invasiveness was reduced by 30-40% (p<0.05 in control, and <0.02 in endometrial samples) (Fig. 6D). In all cases, Cx43 siRNA effects were compared to effects by either a scrambled siRNA, or siRNA directed to GAPDH.

Thirdly, in order to avoid any negative effects of the transient transfection process for siRNA on the integrity of the PMC monolayer, or invasive potential of ESCs, we prepared stably expressing LP9 cells and ESCs from a normal and endometriosis patient (172 and 163, respectively), by infection with lentivirus that either expressed shRNAs targeted to Cx43 that were inducible by doxycycline. In the presence of doxycycline, shRNA suppressed Cx43 protein levels by 35 ± 6% (Fig. 6E), and GJIC was by 88 ± 4% in the cell types tested (Fig. 6F). Invasive behavior was also inhibited by 86 ± 2% when Cx43 was suppressed in the ESCs from either control or endometriosis patients, or in the PMCs (Fig. 6G).

Finally, we used the same Lentivirus system to express a dominant negative Cx43 construct, Cx43 T154A (DN Cx43), which we have previously shown preserves normal gap junctional plaque structures, but fails to open functional channels, and prevents the opening of co-expressed wt Cx43 (Beahm et.al, 2006). Expression of DN Cx43 increased total Cx43 levels by 2 fold in ESCs and 1.4 fold in PMCs (Fig. 6E), decreased GJIC by 90% (Fig. 6F) and invasive behavior by a remarkable 98-99%, whether expressed in ESCs or PMCs (Fig. 6G).

This analysis clearly shows that both EECs and ESCs can be invasive across a mesothelium, and that this is even true when they are derived from control subjects, although the levels are higher in cells from endometriosis patients. Three independent treatments that specifically block either functional GJIC between ESCs and PMCs (GAP27 or DN Cx43) or expression of Cx43 in either of the cell types, greatly reduce the invasive behavior in both control and endometrial samples. Tests on EEC invasiveness (data not shown) showed similar dependence of invasiveness on Cx43, but the higher variability between samples from different patients led to the effect not being significant at the 0.05 level.

AFM analysis of ESC effects on the PMC monolayer

Given the clear dependence of invasiveness of ESCs through the mesothelium on Cx43 mediated GJIC, we turned to AFM to provide insights into how this occurs. We specifically probed the influence of ESCs, in the presence or absence of Cx43, on the “barrier function” of the mesothelium (i.e. the “tightness” of contact between PMCs that prevent transmigration of cells). ESCs from a control subject (172) or endometriosis patients (169 and 170) were first labeled with the membrane dye DiO, and dropped onto a PMC monolayer at a ratio (ESC:PMC) of 1:20. After ~3 hrs, the monolayer was imaged with AFM by pressing it with a ‘sharp’ conical probe at a constant pressure of 1 nN to obtain a 3-D contour map of the monolayer (Figs. 7A-B). This readily allowed the interfaces between cells to be identified (Fig. 7C), and to measure the depth of penetrance between cells. ESCs from several patients all induced an increase in penetrance, measured ~10um away (~2 cell diameters) from site of the dropped cells, identified based on prior DiO labeling (Fig. 7D). This was also evident as a widening in the gap between cells (compare Figs 7A and B). This effect on penetrance between cells was most pronounced in ESCs from an endometriosis patient (169) compared to a control patient (171) (Fig. 7D), and also showed a
dependence on the density of ESCs, as penetrance was less when a 1:50 ratio of ESCs:PMCs was used instead of 1:20 (data not shown). We have also conducted a similar test using an AFM probe to which a 3 μM diameter glass bead is attached to mimic the shape of a cell, and similarly shown that ESCs induce greater ability for larger objects to penetrate the monolayer (data not shown).

We then used the shRNA, DN Cx43 and wt Cx43 infected LP9 cells, characterized in Fig. 6E, to test the dependence of these changes on Cx43 GJIC. First, we observed that the “barrier function” of mesothelial cells alone (i.e. in the absence of ESCs) was dependent on Cx43 expression, as the degree of penetrance was reduced when Cx43 was overexpressed, and increased when Cx43 was inhibited by shRNA (Fig. 7E). Strikingly, this effect was exactly inverted when we introduced ESCs, as now Cx43 overexpression significantly increased penetrance, while Cx43 inhibition by shRNA completely eliminated the effect of ESCs, so that penetrance was the same as in their absence. This effect was due to the channel forming role of Cx43, as the same reduction in penetrance was observed in LP9 cells infected with a DN Cx43, which would preserve, or even enhance, any adhesive functions of Cx43 between ESCs and PMCs.

DISCUSSION

Invasive processes of any cell into a “foreign” environment occurs in many instances, both normal (during development, extravasation, blastocyst implantation) and pathological (metastasis) and involve close intercellular interactions. While the need for initial adhesion events has long been recognized, it has become increasingly evident that the formation of heterotypic gap junctions between the invading and target tissue is an early event in implantation (Grummer et.al., 1996), extravasation (el Sabban and Pauli, 1991, 1994; Ito et.al, 2000, Naoi et.al, 2007) and metastasis (Lamiche et.al., 2012; Stoletov, et.al., 2013; Hong et.al, 2015; Cheng et .al., 2015). However, despite endometriosis being the most common form of pathological invasiveness (affecting 10% of women worldwide), there are few studies on the molecular characterization of gap junctions, (or other intercellular junction components) in this disease. Herein, we sought to determine if dysregulated expression of these endometrial genes associated with disease status.

Expression of GJ and other genes: In a test sample of 6 endometriosis patients and 2 healthy controls, we found highly significant shifts in the expression of several junctional genes and proteins that regulate junctional contacts transcriptionally and post-transcriptionally. The most predictive change associated with endometriosis was in the gap junction (GJ) gene family encoding connexins, specifically in terms of the subpopulation of cells showing high GJ expression (Fig. 1A-D). This became even more evident in a cell clustering analysis, which demonstrated that the only group of cells where fraction of the cell population reliably tracked disease progression were those characterized by high GJ expression with (Fig. 2). This pattern has likely escaped prior screens of endometrial tissue using bulk PCR approaches, as the changes in GJ gene expression in the two major cell types of the endometrium were in opposite directions, with stromal cells showing decreased expression (Fig 1C), and epithelial cells evidencing upregulation (Fig 1D) as disease stage increased.

Notably, these changes were not restricted to the connexins expressed most abundantly in the endometrium (Cx43, 26 and 32), but applied to most of the GJ gene members screened. It was also notable that Cx43 was the dominantly expressed Cx in both ESCs and EECs. For the former, this was consistent with what has been reported in the literature, but EECs have been reported to display mostly Cx26 and Cx32 expression in eutopic endometrium (Jahn et.al., 1995). This has been shown to switch to Cx43 in ectopic lesions (Regidor et.al., 1997), and even eutopically in a baboon model of endometriosis (Winterhager et.al., 2009). While it is possible that the short time in culture modified this expression pattern of EECs in our study, it is also possible
that the switch to Cx43 occurs very early in release of cells from the endometrial lining, even in normal subjects. In either event it is clear that the expression of Cx43 is greatly enhanced in EECs from endometriosis patients compared to control, even while they are still resident in the eutopic endometrium.

In terms of other genes involved in intercellular interactions, many do not show consistent changes, at least in both cell types, but two genes associated with adhesion (EpCAM and β-catenin) show the same expression shifts as the GJ genes, while Snail 1, a master transcriptional repressor that promotes endothelial to mesothelial transition (EMT), shows the opposite pattern. This is consistent with the demonstrated suppressive effect Snail 1 has been shown to have on Cx43 (deBoer et.al., 2007), and the promotion of a migratory phenotype in ESCs, and its repression in EECs. However, it is not clear if in this instance Snail1 is activating a typical EMT response, as the expression of E-cadherin, which is a primary target for suppression by Snail1, was not detected in the endometrium, and N-cadherin, the expression of which is normally increased by Snail1, increases in endometriosis EECs coincident with a Snail 1 decrease. Furthermore, the increase in Snail1 in ESCs is associated with a reduction in β-catenin levels, which may limit EMT effects as they typically depend on activation of β-catenin signaling. This is consistent with the view that EMT transitions in endometriosis are only partial (Konrad et.al., 2020).

With regard to kinases that regulate many intercellular junctional components, it is notable that the cells with the highest GJ expression (control ESCs and endometriosis EECs) in general show the lowest expression of most kinase genes (see heat maps in Fig. 1 A-B), suggesting an inhibitory relationship. However, in terms of global kinase levels, the only kinases to show significant shifts with endometriosis were PKA gamma in both ESCs and EECs, and MAPK1 in EECs (Fig. 1E-F, and Fig. S2B), all of which changed in the same direction as the GJ genes. This is a striking example of how overall expression levels can be misleading when compared to the details of expression patterns at the single cell level.

**GJIC and invasiveness:** These expression profiles initially pose an interesting conundrum, as stromal cells are thought to be the primary compartment involved in endometriotic lesion formation (Lucidi et.al., 2005; Burney and Giudice, 2012), and show the greatest EMT-like response in endometriosis (see above), yet they also show repressed levels of Cx43, which is needed for invasive behavior in metastasis and extravasation. However, functional studies reveal that ESCs, while poorly coupled homotypically, show a dramatic induction of heterotypic coupling when contacting mesothelial cells that is greatly enhanced in endometriosis samples (Fig. 3C). This induction, linked to increased trafficking of Cx43 to the cell surface (Fig. 3D and E), is very analogous to what has been observed in breast cancer, where internalized Cx43 and 26 in the primary tumor, traffics to the surface in lymph node metastases (Kanczuga-Koda et.al., 2006). Enhanced heterotypic coupling with capillary endothelia (el Sabban and Pauli, 1991, 1994; Ito et.al., 2000) or target tissues (Lamiche et.al., 2012; Hong et.al., 2015) have been shown to be critical to invasive behavior during early metastasis.

The importance of the enhanced heterotypic coupling between ESCs and PMCs, unique to endometriosis, to their invasiveness across the mesothelium was demonstrated using four different modes of inhibition of Cx43. Specifically, invasion was not only inhibited by suppression of Cx43 expression (Fig. 6D and G), but also by a peptide blocker GAP27 (Fig. 6A), which inhibits all channel function, but leaves protein expression unaffected, and by DN Cx43 expression (Fig. 6G), which actually increases protein levels (Fig. 6E) and gap junction structures (Beahm et.al., 2006), but blocks all channel activity. Notably, inhibition of invasion is the same whether Cx43 function is ablated in ESCs or PMCs (cf. shRNA and DN Cx43 treatments for 163S and 169S in Fig. 6G). This suggests that heterotypic GJIC likely triggers the invasive behavior, which, as we demonstrate with AFM (Fig. 7), is associated with increased separation between PMCs, likely...
resulting from a disruption of the adhesion and tight junctions between the cells (i.e. breakdown of the “barrier function”), akin to what happens during extravasation (Ito et al., 2000; Reymond et al., 2013).

In Fig. 8 we present a summary of how interactions between the mesothelium and endometrial cells arriving in the peritoneum by retrograde menstruation differ in normal and endometriosis patients, resulting in lesion formation. The enhanced GJIC seen in endometriosis ESCs (Fig. 8B) would allow signals, indicated by green triangles, to pass effectively from endometrial cells to the mesothelium, where they are further propagated through GJIC, to promote disruption of “barrier function” between PMCs, allowing invasion. Interestingly, in the absence of signals from ESCs, GJIC was shown to be important to maintenance of the barrier function (Fig. 7E). This may also be why DN Cx43 was more effective in preventing invasion than Cx43 knock-down by sh- or si-RNA (Fig. 6G), as it may reinforce the barrier functions by enhancing adhesiveness between PMCs. The model emphasizes how increased receptiveness of the ESCs to form heterocellular gap junctions in endometriosis can lead to invasive lesion formation. However, it is also possible that they endometriotic ESCs may make more of the hypothetical signals that induce disruption of the barrier function, emphasizing the importance in identifying such factors. It should be noted that we cannot definitively exclude involvement of factors released through Cx43 hemichannels in the invasive process, but there are no obvious signals that would induce their opening in these experiments, and effects on invasion are the same independent of the cell in which Cx43 is inhibited.

Much remains to be done to identify the intercellular signals that mediate barrier function disruption, and also to understand reciprocal effects of the mesothelium on the ESCs, aside from the demonstrated induction of heterotypic gap junction formation. However, the current study clearly implicates changes in gap junction expression in the earliest phases of the development of endometriosis, and their critical role in initiating lesion formation. They also demonstrate that changes within the uterine endometrium prime these cells to be invasive once they reach the peritoneal cavity, in much the same way that the metastatic potential of cancer cells is determined as they leave the primary tumor, suggesting that understanding each of these processes will inform the other.

**MATERIALS AND METHODS**

**Primary endometrial epithelial cell isolation from endometrial biopsies**

Primary ESCs and EECs were isolated from endometrial biopsies obtained from women with and without endometriosis. All women provided informed consent prior to participating in this Institutional Review Board approved protocol. Study subjects were premenopausal women between 20 and 45 years of age with regular menstrual cycles undergoing laparoscopic surgery for gynecologic indications (Table 1). Women with pelvic inflammatory disease/hydrosalpinx, endometrial polyps, or submucosal fibroids were excluded. Endometriosis was staged according to the revised ASRM criteria and confirmed by histopathologic review of peritoneal or cyst wall biopsy in all cases. Fertile women undergoing tubal sterilization and without endometriosis at surgery were considered healthy controls. Menstrual cycle phase (proliferative or secretory) was determined by cycle history and confirmed by serum estradiol and progesterone levels. Endometrial tissue was obtained by pipelle biopsy at the time of laparoscopic surgery.

The biopsy material was dissociated by shaking in 5mg/ml collagenase and 2.5mg/ml DNase in Hanks Balanced Salt Solution at 37°C for 1 hour. Isolation of primary ESCs and EECs from the biopsies was performed using a combination of straining (45uM nylon filter) and differential sedimentation (EECs cluster and sediment faster), followed by differential attachment.
(EECs adhere less well to culture plates), in a modification of the method developed by Kirk and Irwin (1980) used in prior studies De La Garza, et.al., 2012; Chen et.al., 2016). In some experiments the differential attachment step was replaced by using an Ep-CAM affinity column to enrich EECs. Both methods achieve about 97% purity for EECs and ESCs, as illustrated in Fig. S1 by immunostaining for epithelial [EpCAM (ab71916 from Abcam) and CK 7 (ab902 and 1598 from Abcam)] and stromal [Vimentin (MA1-10459 from Thermo Fisher; NBP1-92687 from NovusBio)] markers. Gene analysis was performed exclusively on P0 or P1 cells, while functional tests of coupling and invasion used cells between P0 and P5 (ESCs) or P0 to P3 (EECs).

Cell Culture

Primary ESCs were cultured in Dulbecco's Modified Eagle Medium (DMEM)/F12 (1:1) (Gibco, Buffalo, NY) containing antibiotic/antimycotic mix (Gibco, Buffalo, NY), 10 µg/ml insulin (Sigma, St. Louis, MO) and 10% heat inactivated fetal bovine serum (FBS - Gibco, Buffalo, NY) as described previously (Ferreira et al., 2008). EECs were cultured in MCDB/Medium 199/MEMα (1:1:0.6) containing antibiotic/antimycotic mix, 10μg/ml insulin, D-Glucose (0.45%) (Sigma, St. Louis, MO), GlutaMax and 10% FBS (Gibco, NY). Prolonged culture was in defined KSFM with supplement, 1% FCS, and antibiotics/antimycotics (Gibco, NY) to preserve differentiated state of the EECs (Chen et.al., 2016) although this generally was only possible to 3 - 4 passages. All experiments were performed using low passages (≤5 for ESCs ≤3) to avoid loss of differentiated characteristics. Established LP9 cells (Corriell Cell Repositories, Camden, NJ) were used as a model for peritoneal mesothelial cells and cultured as described previously (De La Garza, 2012, Liu et al., 2009) and grown in MCDB 131. Medium 199 (1:1 - Gibco, NY) with 15% FBS, sodium pyruvate, GlutaMax, antibiotic/antimycotic mix (Gibco, NY), hEGF (20ng/ml) and hydrocortisone (0.4ng/ml) from Sigma, St. Louis, MO. Previous studies, including our work, have validated and used LP9 cells as a model for peritoneal invasion by endometrial cells (Nair et al., 2008).

Single-cell RNA expression of a connexin gene panel by microfluidic PCR

Automated single cell isolation and processing for cDNA amplification was performed in the C1 (Fluidigm Inc) system, using the C1 integrated fluidic circuit (IFC) chips. Captured single cells were visually confirmed by viewing the microfluidic wells of the C1 IFCs on an inverted light microscope. Empty wells or those containing doublets were excluded. Single-cell cDNA from the IFC chips is then subjected to microfluidic PCR amplification in the Biomark platform (Fluidigm, Inc), using Biomark PCR IFC chips with1936 (48x48) or 9216 (96x96) microfluidic wells (Polusani et al., 2019). The gap junction gene panel and corresponding PCR primer sequences for expression profiling selected from the PrimerBank database (http://pga.mgh.harvard.edu/primerbank) are outlined in Supplemental Table 1. In each Biomark chip assay, universal RNA (200 pg) from human normal tissues (cat #4234565, BioChain, Newark, CA) and no template control (NTC) served as positive and negative controls, respectively. Valid PCR products were determined by amplicon melting temperature curves of each gene. Expression values were determined based on cycle threshold (Ct) normalized to the housekeeping gene Actin. For cell quality control, UBB and Actin expression with Ct threshold of >20 were excluded (generally 1-3% of captured cells), as this was indicative of RNA degradation.

Homotypic and Heterotypic GJIC Assays

GJIC was measured using a novel automated parachute assay. Recipient cells are grown to confluence in a 96-cell flat bottomed plate, and the media changed to (Phenol Red-free DMEM, sodium pyruvate and 5%FBS – Assay Media) immediately before the assay. Donor cells in separate wells are incubated for 20 mins with 10uM calcein AM (Invitrogen), a membrane permeable dye that on cleavage by intracellular esterases becomes membrane impermeable, but
permeable to gap junctions. After washing, trypsinization and addition of assay media, ~2500
calcine-labeled donor cells per well are dropped (‘parachuted’) onto the recipient cell layer, and
calcine transfer between donor and recipient cells observed by fluorescent microscopic imaging
(Fig. 3A). For homotypic interactions, ESCs, EECs or LP9 donor cells were parachuted onto
recipient cells of the same type. For heterotypic GJIC assays, ESCs or EECs were parachuted
onto LP9 recipient cells. Fluorescent, bright field and digital phase contrast images of 10-15 fields
per well were captured on an Operetta automated microscope (Perkin Elmer) at 30 min intervals
for approximately 2 hours. A program (developed in consultation with Perkin Elmer) allowed
identification of all cells on the plate, (from phase contrast image), original donors (5-15 per field),
and dye-filled recipients (based on calcine intensity – see Fig. 3B). Data are expressed as # of
fluorescent recipient cells/# of donor cells for each condition (R/D ratio), plotted over time, and a
linear regression line drawn through the data, with the slope used as a measure of coupling (Fig.
3C) and regression coefficient (mostly >0.8) used as a measure of assay reliability.

Trans-mesothelial Invasion Assay

The 3-D invasion assay modeling trans-mesothelial invasion (Fig. 5A) has been described
previously (De La Garza et. al., 2012, Ferreira et. al., 2008, Nair et. al., 2008). Briefly, LP9 peritoneal
mesothelial cells (PMCs) were grown to confluence in 24-well invasion chamber inserts containing
growth-factor-reduced Matrigel™, coated on 8-µm pore membranes (Corning, NY). ESCs or
EECs were then labeled with CellTracker Green® or DiO (Invitrogen), trypsinized and counted,
before dropping onto the confluent layer of LP9 PMCs in the prepared inserts (~20,000 cells per
insert), at which time the media below the insert was changed to the appropriate media for the
invading cell (ESC or EEC). After 24 hr. incubation, non-invading cells on the upper surface of the
insert were mechanically removed. Invading cells on the bottom of the membrane insert, were
stained with DAPI, and 10 fields counted using an Inverted Nikon 2000 fluorescence microscope
with 20x objective. Invasion assays for each cell type were performed in triplicates.

To test the role of gap junctions in the invasive process, we initially pretreated both the
monolayer and dropped cells for 24 hours with 300uM GAP27 (Zealand Pharma, Copenhagen,
Denmark), a peptidomimetic of the extracellular loops of Cx43 which competitively blocks Cx43
gap junction formation. Due to difficulties in obtaining consistently active peptide preparations, we
subsequently shifted to 24 hour pre-treatment of the LP9 monolayer with a combination of two
siRNAs to Cx43 (10 pmoles/well or 5nM final concentration) - Ambion Silencer™ Select) in Opti
MEM (Gibco, NY) with RNAi MAX (1/100 dilution, Invitrogen), diluted 1:1 with assay media, per
manufacturer’s instructions. As the siRNA transfections done immediately prior to invasion could
impact cell behavior, we also knocked-down Cx43 expression using 3 different shRNAs
(Dharmacon, UK) introduced into either Endometrial or LP9 Mesothelial cells via Lentivirus
(carrying an RFP reported driven by a separate promoter) infection. Average infection efficiency
was 58% (range 30-80%). In assessing invasion, cells expressing shRNA were identifiable as
RFP positive, while unlabelled cells in the same experiment served as internal controls.

AFM measurements of cell-cell adhesion and mesothelial integrity.

We applied a Nanoscope Catalyst AFM (Bruker) interfaced with an epifluorescent inverted
microscope Eclipse Ti (Nikon). AFM images were acquired with the Peak Force Quantitative
Nanomechanical Mapping (QNM) mode with cells immersed in appropriate culture media.
ScanAsyst probes (Bruker) with the nominal spring constant 0.4 N/m were used for imaging. The
exact spring constant for each probe was determined with the thermal noise method (Butt and
Jaschke, 1995). For each cell culture dish, at least 5 fields 100 by 100 µm were collected with the
Peak Force set point of 2nN, and electronic resolution of 256 by 256 pixels. Nanomechanical data
were processed with Nanoscope Analysis software v.1.7 (Bruker) using retrace images. Then,
the Sneddon model (Sneddon, 1965) and the rules established by Sokolov (Sokolov and Dokukin, 2014) were applied for calculations of mechanical parameters.

**Cell to cell adhesion:** We attached a tester cell to a tipless probe MLCT-O10 (Bruker, cantilever A, spring constant 0.07N/m) using polyethyleneimine (PEI) as a glue (Friedrichs et. al., 2013) (Fig. 4). Briefly, the probes were immersed in 0.01% PEI in water for 30 min. Tester cells attachment to a culture dish was weaken by replacement of the culture medium with a non-enzymatic cell dissociation solution (Millipore) for 15-30 min in a cell culture incubator (37°C, 5%CO). Next, a single tester cell loosely attached to a culture dish was attached to a PEI covered cantilever by pressing it at 1 nN for 5-10 min. After visual inspection of successful cell attachment, the tester cell was lifted and transferred to a dish containing single tested cells. Then the tester cell was positioned over a tested cell and the cantilever slowly lowered till detection of cell-cell interactions with a force plot. The cells were left interacting for 30 to 180 sec at forces 0.5 to 5 nN and then the tester cell was lifted. During this step a force plot was recorded and the collected data applied to calculate cell – cell adhesion parameters. The force plots were baseline corrected and a maximum of adhesion between cells during their detachment was calculated (units of force, Newton) (Taubenberger, Hutmacher, and Muller 2014; Dufrêne et al. 2017).

**Integrity of LP9 layer:** LP9 cells were grown to confluence in a 60mm culture dish (Fig. 7). ESC cells grown in separate wells were stained with DiO, suspended, and dropped on to the LP9 monolayer at either a 1:50 or 1:20 ratio to the LP9 cells. Three hours later the cells were imaged by AFM. To calculate a tip penetration depth, cell boundaries were identified using images collected by the peak force error (PFE) channel. To exclude gap areas between cells or areas of cells growing in multilayers, PFE images were overlaid with height channel images after processing them with the flatten function of 1st order. Tip penetration was calculated based on a height histogram of all data points using a difference between the prevalent maximum of cell monolayer height and the prevalent maximum depth between cells accessible for the tip.

**Data and Statistical Analysis**

For microfluidic PCR analysis, cycle thresholds (Ct) were used to calculate gene expression relative to the housekeeping (HK) gene ACTIN. Relative expression was calculated by the formula $2^{-\Delta Ct}$, where $\Delta Ct = Ct_{\text{gene}} - Ct_{\text{HK}}$. Heat maps were generated for data visualization by MeV (http://mev.tm4.org/). In addition, violin plots were used to visualize single-cell expression distribution. To statistically assess this distribution between samples, Duncan’s multiple range test by R package was used to compare expression in different patient samples. For statistical analysis of cell assays, the Shapiro-Wilk normality test was first performed. t-test and ANOVA (with Holm-Sidak post-hoc analysis) were used as parametric assays after normality of data was ensured. In cases where normality test failed, Mann-Whitney rank sum test was used as a nonparametric analysis method. P values $\leq 0.05$ were considered statistically significant. As data on GJIC and invasion of ESC and EEC populations from patients was normally distributed, comparisons utilized two tailed student t-tests, with a cut-off off of p $< 0.05$ (degrees of freedom ranged from 8 – 20). Statistical tests of all AFM data were performed and corresponding graphs prepared with OriginPro 2000b (Origin Lab). All data showed normal distribution by Shapiro-Wilk, except the 170S-LP9 (Cx43) data set in Fig. 7F, which did pass the Kolgomorov test of normality. P values were corrected using the more stringent Shapiro method.

**Cell Cluster Analysis**

Single cells were clustered based on the normalized expression values using an in-house graph-based community discovery algorithm. Briefly, the algorithm starts by computing the Euclidean distances between pairs of cells, and then constructs k nearest neighbor graph, where each cell is connected to k cells that are closest to it. The best k is chosen by the algorithm with a
Finally, a community discovery algorithm optimizing the well-known modularity function is applied to find dense subgraphs as cell clusters (Ruan and Zhang, 2008). Note that the number of clusters is determined automatically during the optimization process.

ACKNOWLEDGEMENTS: We would like to express gratitude to the members of the members of the High-Throughput Screening Facility who helped with the assessment of GJIC, particularly Srikanth Polusani, the members of the clinical teams who helped in collection of patient samples, particularly Jessica Perry at UTHealth SA, and Peter Binkley curator of the UTHealth SA Ob/Gyn tissue bank. We would also like to thank Taryn Olivas and Taylor Williams who made critical contributions to the early characterization of patient samples. We would like to also acknowledge the CPRIT-funded Bioanalytics and Single-Cell Core (BASiC) facility at the University of Texas Health San Antonio for C1 cell capture and microfluidic PCR analysis and the High-throughput Screening Facility (HTSF) in the CPRIT-funded Center for Innovative Drug Discovery (CIDD). Thanks to Mr. Nicholas Lucio at the BASiC facility for his assistance with running the C1 system, and Dr. Srikanth Polusani for running the Perkin Elmer Operetta system in the HTSF.

FUNDING: This work was supported by an award from the Women’s Health Initiative through the Joe R. and Theresa Long School of Medicine (BJN and NBK), a Presidential Entrepreneurial Fund Award (NBK) and a pilot award from the Mays Cancer Center in conjunction with philanthropic donations form the Circle of Hope (BJN), all affiliated with UT Health San Antonio, as well as a pilot award from the Endometriosis Foundation of America (NBK). The BASiC facility at the University of Texas Health San Antonio is funded by the Cancer Prevention and Research Institute of Texas (CPRIT, grant # RP150600), and the HTSF is within the CIDD funded by CPRIT, (# RP160844) and the Institute for Integration of Medicine and Science (NIH-NCATS grant #UL1 TR 002645).

AUTHOR CONTRIBUTIONS:

CWC and JC performed all functional experiments on cell coupling and invasion, isolated cells, and prepared them for single cell analysis; CWW, CMW, YTH and LLL conducted single cell analyses, and LLL and YTH performed statistical analyses; MJH designed and supervised the automated system for analysis of GJIC; JR performed all cell clustering analysis; LG isolated patient cells; ROB, RSS and RDR collected patient samples for analysis; MG and PO conducted all AFM experiments and analyses; NBK and BJN conceived and designed all experiments, coordinated patient sample collections and wrote the manuscript.
**TABLE 1: Patient database**

| Patient | Endo stage | ethnicity  | age | BMI | Menst. stage |
|---------|------------|------------|-----|-----|--------------|
| 171     | Norm       | ND         | 35  | 27  | Mid sec      |
| 172     | Norm       | hispanic   | 25  | 28  | Early sec    |
| 164     | I-II       | Pac Is     | 30  | 28  | Early sec    |
| 170     | III-IV     | white      | 30  | 22  | prolifer     |
| 002     | III-IV     | white      | 26  | 26  | Early prolifer |
| 163     | III-IV     | white      | 23  | 25  | Early sec    |
| 169     | III-IV     | white      | 37  | 23  | Mid sec      |
| 003     | III-IV     | hispanic   | 31  | 31  | Non-cycl     |
Figure 1: Single cell gene expression profiles show a consistent, but opposite, shift in Gap Junction (GJ) gene expression in endometrial stromal (ESCs) and epithelial cells (EECs) with endometriosis progression.

(A-B) Heat maps of single cell (columns) expression of a panel of genes, plotted in descending order of Cx gene expression (rows) involved in intercellular interactions and their regulation in ESCs (A) and EECs (B) isolated from 2 control and 6 endometriosis patients, with samples taken at different disease stages and at different phases of the menstrual cycle, as indicated. Due to poor yields or their adhesive nature preventing single cell isolation for analysis, EECs could not be analyzed for all endometriosis patients. Most notable were changes in the abundance of cells with high levels of expression of all GJ genes (top third of each map), which steadily decreased with disease progression in ESCs and increased in EECs. A full list of genes in order (top to bottom) in Table S1.

(C-F) Violin plots showing the distribution of RNA expression levels (log scale) for selected genes, or groups of genes, in stromal (C, E) and epithelial cells (D, F) derived from each of the patients studied (indicated by numbers on the X-axis). C and D show the aggregate of combined GJ gene expression, while E and F show 6 genes with the most consistent changes with endometriosis. Green outlines indicate increased expression and red outlines decreased expression in endometriosis patients compared to control (EECs) or control and early endometriosis, excluding the sample taken during mid-secretory phase of the menstrual cycle (ESCs), indicated by the red horizontal line in each plot. Horizontal bulges in each plot indicate larger numbers of cells with that expression level. The black dot in the middle of the plot indicates mean expression. Significance was determined by Duncan’s multiple range test and statistical differences (P<0.05) between samples, as indicated by letters (e.g. a vs. b; b vs. d). In addition, ‘a’ indicates the highest mean expression and ‘d’ indicates the lowest mean expression value for each factor (i.e. a>b>c>d). Violin plots of the remaining genes are shown in Fig. S2.
Figure 2: Cell clustering analysis comparing control and endometriosis samples show progressive changes in the fraction of cells characterized by high GJ expression. Cells from ESCs (A) and EECs (C) were grouped into 6 and 5 clusters, respectively, based on patterns of gene expression across all patients, with the bottom clusters [sc6 for ESCs, and ec5 for EECs] characterized by high GJ expression. Following the distribution of each cluster in all patients, endometriosis patients have consistently lower percentages of subpopulation sc-6 (B), and higher percentages of subpopulation ec-5 (D) compared to controls (trends highlighted by yellow dotted lines).

Figure 3: Heterotypic coupling between ESCs and PMCs is dramatically induced in endometriosis.

Gap junction intercellular coupling (GJIC) was measured by a modified “parachute assay” where calcein ester loaded donors are dropped onto a monolayer of acceptors, and dye transfer measured over time as a linear increase in fluorescent acceptor/donor ratio (A). This can measure homotypic (donor and acceptor the same), or heterotypic (endometrial donors and mesothelial acceptors) GJIC (B). Although there are slight changes in homotypic GJIC in ESCs and EECs consistent with the changes in expression levels of connexins, they were not significant (C-green bars). By contrast, heterotypic interactions with mesothelial cells (PMCs) caused a significant induction of GJIC, most notably in ESCs from endometriosis (C-black bars). Immunocytochemistry showed this was likely due to a redistribution of Cx43 (red) from primarily cytoplasmic in homotypic ESCs (D), to the interfaces of ESCs (green) and PMCs in heterotypic cultures (E). Samples tested: ESC control (8) and endometriosis (11); EEC control (8) and endometriosis (5), which included 2-4 biological replicates for each patient. Significance level was calculated by two tailed t-test.
Figure 4: Atomic Force Microscopy (AFM) demonstrates enhanced adhesion between ESCs and PMCs. 
Cells attached to an AFM tip are brought into contact with PMCs growing on a dish (A), and the force needed to separate them is measured (B). PMCs (LP9 cells) show similar adhesion to one another as to EECs, but much stronger adhesion to ESCs (C). Significance determined by two-tailed t-test.

Figure 5: Enhanced invasiveness of endometrial cells from endometriosis patients across a mesothelial monolayer. 
Using a 3-D model of endometrial cell invasion across a peritoneal cell monolayer (A), endometrial cells from patients show greater invasion than those from control subjects that was significant with a mix of ESCs and EECs (B). Number of independent patients tested - All Controls (3); Endometriosis – ESC+EEC (3 patients); ESC (5); EEC (4). Significance was determined by two-tailed t-test.
Figure 6: Invasiveness of ESCs across a PMC monolayer is dependent on Cx43 GJIC:

(A) Averaging ESCs from all control (black bars, n=3) and endometriosis patients (grey bars, n=6), invasion was inhibited by a peptide inhibitor of GJ channels, GAP27 (percent GJIC compared to untreated shown below each bar).

(B - D) Cx43 siRNA transfection of either ESCs or PMCs (LP9) reduced protein levels of Cx43 (arrow) compared to Laminin A and B controls (upper bands) (B), inhibited heterotypic GJIC compared to scrambled siRNA [Control n=5 (independent replicates from 2 patients); Endo n=8 (independent replicates from 5 patients)] (C), and also inhibited invasion [Control n=6 (2-3 independent replicates of two patients); Endo n=14 (2-4 independent replicates of 6 patients)] (D).

(E – G) Stable, doxycycline inducible shRNA infections of ESCs from a normal (172) and endometriosis (163) patient, as well as PMCs (LP9), reduced levels of Cx43 (arrow) compared to Laminin controls (E), inhibited GJIC (n=3-7 independent experiments with 3 different shRNAs) (F) and invasiveness by ~90% [10 technical replicates of all samples; 3-4 biological repeats for +/- shRNA using 3 different shRNAs (controls were internal)] (G). The block of invasiveness was observed independent of whether Cx43 expression was inhibited in ESCs or PMCs. DN Cx43 (which blocks channel function, but not protein assembly into GJs) was also expressed in ESCs and PMCs at approximately equal levels to wt Cx43 (total Cx43 levels doubled - E). This caused an even greater inhibition of invasiveness (~98% - G). All statistical significance based on one-tailed t-tests.
Figure 7: AFM demonstrates that ESCs induce disruption of the barrier function of a mesothelial monolayer, which is dependent on Cx43 channels.

The topology of the mesothelial monolayer surface shows a significant increase in the spacing between cells in the presence of ESCs (A-B). Using constant pressure, the tip can measure the depth of penetration between cells (C). This was increased from that measured in an LP9 PMC monolayer alone (n=25) when ESCs were dropped onto the monolayer, particularly from an endometriosis patient (169, n=19) compared to control (172, n=13) (D). Cx43 appears to help preserve the barrier function of the mesothelium alone, as overexpression reduced penetration, while Cx43 shRNA infection increased penetration in the PMC monolayer in the absence of ESCs (E). By contrast, in the presence of ESCs from a patient (170), Cx43 overexpression in PMCs enhances penetration, while blocking Cx43 with either shRNA or DN Cx43 neutralize the effect of ESCs (F). Each dot represents a cell measurement from plates prepared on the same day for comparison. Statistical tests of significance by two-tailed t-test (see methods).
Figure 8: Model of GJIC induction of transmesothelial invasion.

(A) In healthy patients, when endometrial cells (brown) encounter a mesothelium (following arrival in the peritoneum via retrograde menstruation), modest GJIC provides limited exchange of signals from ESCs to PMCs (green triangles) or PMCs to ESCs (purple dots). (B) In endometriosis, endometrial cells have Cx43 mostly in intracellular stores, but encounters with mesothelial cells triggers Cx43 trafficking to the cell surface. The increased GJIC that results mediates transfer of signals to PMCs (green triangles), which propagate through the mesothelium, inducing disruption of the adhesive and tight junctions between PMCs, facilitating invasion of the ESCs.
REFERENCES:

Augoulea, A., Alexandrou, A., Creatsa, M., Vrachnis, N and Lambrinoudaki, I. Pathogenesis of endometriosis: the role of genetics, inflammation and oxidative stress, Arch. Gynecology and Obstetrics, 286: 99-103 (2012). PMID: 22546953

Beahm DL, Oshima A, Gaietta GM, Hand GM, Smock AE, Zucker SN, Toloue MM, Chandrasekhar A, Nicholson BJ, Sosinsky GE. Mutation of a conserved threonine in the third transmembrane helix of alpha- and beta-connexins creates a dominant-negative closed gap junction channel. J Biol Chem. 281: 7994-8009 (2006) PMID: 16407179

Bontempo A.C., Mikesell L., Patient perceptions of misdiagnosis of endometriosis: results from an online national survey, Diagnosis (Berl) 7: 97-106 (2020) PMID: 32007945

Burney RO, Talb S, Hamilton AE, Vo KC, Nyegaard M, Nezhat CR, et al. Gene expression of endometrium reveals progesterone resistance and candidate susceptibility genes in women with endometriosis. Endocrinology 148:3814-26 (2007) PMID 17510236

Burney R.O., Giudice L.C., Pathogenesis and pathophysiology of endometriosis, Fertil. Steril., 98: 511-519 (2012) PMID: 22819144

Butt H.-J., and Jaschke M. Calculation of Thermal Noise in Atomic Force Microscopy. Nanotechnology 6: 1–7. (1995).

Chen Q, Boire A, Jin X, et al. Carcinoma-astrocyte gap junctions promote brain metastasis by cGAMP transfer Nature, 533 : 493-498 (2016) PMID: 27225120

Chen JC, Hoffman JR, Arora R, Perrone LA, Gonzalez-Gomez CJ, Vo KC, Laird DJ, Irwin JC, Giudice LC. Cryopreservation and recovery of human endometrial epithelial cells with high viability, purity, and functional fidelity. Fertil Steril. 105: 501-10. (2016) PMID: 26515378

Cheng Y., Ma, D. Zhang Y., Li Z., Geng L. Cervical squamous cancer mRNA profiles reveal the key genes of metastasis and invasion, Eur J Gynaecol Oncol, 36: 309-317 (2015) [PMID: 26189259]

de Boer TP, van Veen TA, Bierhuizen MF, Kok B, Rook MB, Boonen KJ, Vos MA, Doevendans PA, de Bakker JM, van der Heyden MA. Connexin43 repression following epithelium-to-mesenchyme transition in embryonal carcinoma cells requires Snail1 transcription factor. Differentiation. 75:208-18. (2007) PMID: 17359298.

De La Garza EM, Binkley PA, Ganapathy M, Krishnegowda NK, Tekmal RR, Schenken RS, Kirm NB. Raf-1, a potential therapeutic target, mediates early steps in endometriosis lesion development by endometrial epithelial and stromal cells. Endocrinology. 153: 3911-21. (2012) PMID: 22619359

Diao H., Xiao S., Howerth E.W, Zhao F., Li R., Ard M.B., Ye X. Broad gap junction blocker carbenoxolone disrupts uterine preparation for embryo implantation in mice, Biol. Reproduction, 89: 31. (2013) PMID: 23843229

Dufrêne YF, Ando T, Garcia R, Alsteens D, Martinez-Martin D, Engel A, Gerber C, and Müller DJ. 2017. "maging Modes of Atomic Force Microscopy for Application in Molecular and Cell Biology. Nature Nanotechnology 12: 295–307.

el-Sabban M.E., Pauli B.U. Cytoplasmic dye transfer between metastatic tumor cells and vascular endothelium, J Cell Biology, 115: 1375-1382 (1991) PMID: 1955478
el-Sabban M.E., Pauli B.U. Adhesion-mediated gap junctional communication between lung-metastatatic cancer cells and endothelium, Invasion & Metastasis 14: 164-176 (1994) PMID: 7657509

Evans W.H., Leybaert L. Mimetic peptides as blockers of connexin channel-facilitated intercellular communication, Cell Communication & Adhesion, 14 265-273. (2007) PMID: 18392994

Eskcnazi B, Warner ML. Epidemiology of endometriosis. Obstet Gynecol Clin North Am. 24:235-58 (1997) PMID: 9163765

Ferreira MC, Witz CA, Hammes LS, Kirma N, Petraglia F, Schenken RS, Reis FM. Activin A increases invasiveness of endometrial cells in an in vitro model of human peritoneum. Mol Hum Reprod. 14:301-7 (2008) PMID: 18359784

Friedrichs J, Legate KR, Schubert , Bharadwaj M, Werner C, Müller DJ and Benoit M.. A Practical Guide to Quantify Cell Adhesion Using Single-Cell Force Spectroscopy. Methods 60: 169–78 (2013).

Grümmer R, Chwalsiz K, Mulholland J, Traub O, Winterhager E. Regulation of connexin26 and connxin43 expression in rat endometrium by ovarian steroid hormones. Biol Reprod. 51:1109-16. (1994) PMID: 7888490

Grummer R., Reuss B., Winterhager E. Expression pattern of different gap junction connexins is related to embryo implantation, Intl. J. Dev. Biol. 40: 361-367. (1996) PMID: 8735949

Guo SW, Wu Y, Strawn E, Basir Z, Wang Y, Halverson G, et al. Genomic alterations in the endometrium may be a proximate cause for endometriosis. Eur J Obstet Gynecol Reprod Biol.116:89–99. (2004) PMID 15294375

Hastings J.M., Fazleabas A.T., A baboon model for endometriosis: implications for fertility, Reprod Biol Endocrinol, 4 (Suppl 1): S7 (2006). PMID: 17118171

Hong X., Sin W.C., Harris A.L, Naus., C.C. Gap junctions modulate glioma invasion by direct transfer of microRNA, Oncotarget, 6 :15566-15577 (2015) PMID: 25978028

Hudelist G., Fritzer N., Thomas A., Niehues C., Oppelt P., Haas D., Tammaa A., Salzer H. Diagnostic delay for endometriosis in Austria and Germany: causes and possible consequences, Hum Reprod, 27: 3412-3416 (2012). [PMID: 22990516]

Hugo HJ, Kokkinos MI, Blick T, Ackland ML, Thompson EW, Newgreen DF. Defining the E-cadherin repressor interactome in epithelial-mesenchymal transition: the PMC42 model as a case study. Cells Tissues Organs. 193:23-40 (2011) PMID: 21051859

Ito A., Kato H., Kataoka T.R., Okada M., Tsubota N., Asada H., Yoshikawa K., Maeda S., Kitamura Y., Yamasaki H., Nojima H. A role for heterologous gap junctions between melanoma and endothelial cells in metastasis, J. Clin. Invest. 105: 1189-1197 (2000) PMID: 10791993

Jahn E, Classen-Linke I, Kusche M, Beier HM, Traub O, Grümmer R, Winterhager E. Expression of gap junction connexins in the human endometrium throughout the menstrual cycle. Hum Reprod. 10: 2666-70 (1995). PMID: 8567789

Kanczuga-Koda L., Sulkowski S., Lenczewski A., Koda M., Wincewicz A., Baltaziak M., Sulkowska M. Increased expression of connexins 26 and 43 in lymph node metastases of breast cancer, J. Clin. Pathol. 59: 429-433. (2006) PMID: 16567471

Kaushik T., Mishra R., Singh R.K., Bajpai S. Role of connexins in female reproductive system and endometriosis, J Gynecol Obstet Hum Reprod, 49:101705. (2020) PMID: 32018041
Kirk D, Irwin JC. Normal human endometrium in cell culture. Methods Cell Biol. 21B: 51-77 (1980). PMID: 7412575

Konrad L, Dietze R, Riaz MA, Scheiner-Bobis G, Behnke J, Horne F, Hoerscher A, Reising C, Meinhold-Heerlein I. Epithelial-Mesenchymal Transition in Endometriosis-When Does It Happen? J Clin Med. 9:1915. (2020) PMID: 32570986

Lamiche C, Clarhaut J, Strale P.O., Crespin S., Pedretti N., Bernard F.X., Naus C.C., Chen V.C., Foster L.J., Defamie N., Mesnil M., Debiais F., Cronier L. The gap junction protein Cx43 is involved in the bone-targeted metastatic behaviour of human prostate cancer cells. Clin. Exp. Metastasis, 29: 111-122. (2012) PMID: 22080401

Laws M.J., Taylor R.N., Sidell N., DeMayo F.J., Lydon J.P., Gutstein D.E., Bagchi M.K, Bagchi I.C. Gap junction communication between uterine stromal cells plays a critical role in pregnancy-associated neovascularization and embryo survival, Development 135: 2659-2668 (2008) PMID: 18599509

Lucidi RS, Witz CA, Chrisco M, Binkley PA, Shain SA, Schenken RS. A novel in vitro model of the early endometriotic lesion demonstrates that attachment of endometrial cells to mesothelial cells is dependent on the source of endometrial cells. Fertil Steril. 84:16–21 (2005) PMID: 16009148

Mettler L., Schollmeyer T., Lehmann-Willenbrock E., Schuppler U., Schmutzler A., Shukla D., Zavala A., Lewin A. Accuracy of laparoscopic diagnosis of endometriosis, JSLS, 7: 15-18 (2003). PMID: 12722993

Missmer SA, Hankinson SE, Spiegelman D, Barbieri RL, Michels KB, Hunter DJ. In utero exposures and the incidence of endometriosis. Fertil Steril. 2004; 82:1501–8. PMID: 15589850

Montgomery J, Ghatnekar GS, Grek CL, Moyer KE, Gourdie RG. Connexin 43-Based Therapeutics for Dermal Wound Healing. Int J Mol Sci. 19:1778. (2018) PMID: 29914066;

Mugisho O.O., Green C.R., Kho D.T., Zhang J., Graham E.S., Acosta M.L., Rupenthal I.D. The inflammasome pathway is amplified and perpetuated in an autocrine manner through connexin43 hemichannel mediated ATP release, Biochim Biophys Acta 1862: 385-393 (2018) PMID: 29158134

Nair AS, Nair HB, Lucidi RS, Kirchner AJ, Schenken RS, Tekmal RR et al. Modeling the early endometriotic lesion: mesothelium-endometrial cell co-culture increases endometrial invasion and alters mesothelial and endometrial gene transcription. Fertil. and Steril. 90:1487-95 (2008). PMID: 18163995

Nair R.R., Jain M., Singh K. Reduced expression of gap junction gene connexin 43 in recurrent early pregnancy loss patients, Placenta 32: 619-621 (2011) PMID: 21669459

Naoi Y., Miyoshi Y., Taguchi T., Kim S.J., Arai T., Tamaki Y., Noguchi S. Connexin26 expression is associated with lymphatic vessel invasion and poor prognosis in human breast cancer, Breast Cancer Res.Treat. 106: 11-17 (2007) PMID: 17203385

Oosterlynck DJ, Cornillie FJ, Waer M, Vandeputte M, Koninckx PR. Women with endometriosis show a defect in natural killer activity resulting in a decreased cytotoxicity to autologous endometrium. Fertil Steril. 56:45–51 (1991) PMID: 2065804

Ormonde S, Chou CY, Goold L, Petsoglou C, Al-Taie R, Sherwin T, McGhee CN, Green CR. Regulation of connexin43 gap junction protein triggers vascular recovery and healing in human ocular persistent epithelial defect wounds. J Membr Biol. 245:381-8. (2012) PMID: 22797940.
Parente Barbosa C., Bentes De Souza A. M., Bianco B., and Christofolini D. M. The effect of hormones on endometriosis development, Minerva Ginecol. 63: 375–386 (2011). PMID: 21747346

Polusani SR, Huang YW, Huang G, Chen CW, Wang CM, Lin LL, Osmulski P, Lucio ND, Liu L, Hsu YT, Zhou Y, Lin CL, Aguilera-Barrantes I, Valente PT, Kost ER, Chen CL, Shim EY, Lee SE, Ruan J, Gaczynska ME, Yan P, Goodfellow PJ, Mutch DG, Jin VX, Nicholson BJ, Huang TH, Kirmia NB. Adipokines Deregulate Cellular Communication via Epigenetic Repression of Gap Junction Loci in Obese Endometrial Cancer. Cancer Res. 79:196-208 (2019) PMID: 30389702

Regidor P.A., Regidor M., Schindler A.E., Winterhager E. Aberrant expression pattern of gap junction connexins in endometriotic tissues, Mol. Human Reprod. 3: 375-381 (1997) PMID: 9239721

Reymond N, d’Água BB, Ridley AJ. Crossing the endothelial barrier during metastasis. Nat Rev Cancer. 13: 858-70. (2013) PMID: 24263189

Roca-Cusachs P, Conte V, and Trepat X. Quantifying Forces in Cell Biology. Nature Cell Biology 19: 742–51 (2017).

Ruan J, Zhang W. Identifying network communities with a high resolution. Phys Rev E Stat Nonlin Soft Matter Phys. 77: 016104 (2008). PMID: 18351912.

Ruan J. A Fully Automated Method for Discovering Community Structures in High Dimensional Data. Proc IEEE Int Conf Data Min. 968-973. (2009). PMID: 25296858

Sampson J.A. Peritoneal endometriosis due to menstrual dissemination of endometrial tissue into the peritoneal cavity. Am J Obstet Gynecol. 14: 422-469 (1927)

Sasson IE, Taylor HS. Stem cells and the pathogenesis of endometriosis. Ann N Y Acad Sci. 1127:106–15. (2008) PMID: 18443337

Sancho A., Vandersmissen I, Craps S, Luttun A, and Groll J. A New Strategy to Measure Intercellular Adhesion Forces in Mature Cell-Cell Contacts. Scientific Reports 7: 46152 (2017).

Sneddon IN. “The Relation between Load and Penetration in the Axisymmetric Boussinesq Problem for a Punch of Arbitrary Profile.” International Journal of Engineering Science 3: 47–57 (1965).

Sokolov, I, and Dokukin M. Mechanics of Biological Cells Studied with Atomic Force Microscopy. Microscopy and Microanalysis 20: 2076–77 (2014).

Soliman A.M., Yang H., Du E.X., Kelley C., Winkel C. The direct and indirect costs associated with endometriosis: a systematic literature review, Hum Reprod, 31: 712-722 (2016) PMID: 26851604

Somigliana E, Vigano P, Gaffuri B, Guarneri D, Busacca M, Vignali M. Human endometrialstromal cells as a source of soluble intercellular adhesion molecule (ICAM)-1 molecules. HumReprod. 11:1190–4 (1996) PMID 8671421

Stoletov K., Strnadel J., Zardouzian E., Momiyama M., Park F.D., Kelber J.A., Pizzo D.P., Hoffman R., VandenBerg S.R., Klemke R.L. Role of connexins in metastatic breast cancer and melanoma brain colonization, J Cell Sci. 126: 904-913. (2013) PMID: 23321642
Tamaresis J.S., Irwin J.C., Goldfien G.A., Rabban J.T., Burney R.O., Nezhat C., DePaolo L.V., Giudice L.C., Molecular classification of endometriosis and disease stage using high-dimensional genomic data, Endocrinology, 155: 4986-4999 (2014) PMID: 25243856

Taubenberger AV., Hutmacher DW, and Muller DJ. Single-Cell Force Spectroscopy, an Emerging Tool to Quantify Cell Adhesion to Biomaterials. Tissue Engineering Part B: Reviews 20: 40–55 (2014)

Ulukus M., Cakmak H., Arici A. The role of endometrium in endometriosis, J Soc Gynecol Investig, 13: 467-476 (2006) PMID: 16990031

Weber PA, Chang HC, Spaeth KE, Nitsche JM, Nicholson BJ. The permeability of gap junction channels to probes of different size is dependent on connexin composition and permeant-pore affinities. Biophys J 87:958-973. (2004) PMID 15298902

Winterhager E, Grümmer R, Mavrogianis PA, Jones CJ, Hastings JM, Fazleabas AT. Connexin expression pattern in the endometrium of baboons is influenced by hormonal changes and the presence of endometriotic lesions. Mol Hum Reprod. 15:645-52. (2009) PMID: 19661121

Winterhager E., Kidder G.M. Gap junction connexins in female reproductive organs: implications for women's reproductive health, Hum Reprod Update, 21: 340-352. (2015) PMID: 25667189

Yu J., Boicea A., Barrett K.L., James C.O., Bagchi I.C., Bagchi M.K, Nezhat C., Sidell N., Taylor R.N., Reduced connexin 43 in eutopic endometrium and cultured endometrial stromal cells from subjects with endometriosis, Mol. Human Reprod. 20: 260-270 (2014) PMID: 24270393

Zhang A., Hitomi M., Bar-Shain N., Dalimov Z., Ellis L., Velpula K.K., Fraizer G.C., Gourdie R.G., Lathia J.D. Connexin 43 expression is associated with increased malignancy in prostate cancer cell lines and functions to promote migration, Oncotarget, 6: 11640-11651 (2015) PMID: 25960544
Endometrial Gap Junction Expression - Early Indicators of Endometriosis and Integral to Invasiveness

Chen-Wei Chen #, 1, Jeffery Chavez #, 1, Li-Ling Lin2, Chiou-Miin Wang2, Ya-Ting Hsu2, Matthew J. Hart1, Jianhua Ruan3, Laurie Gillette 4, Richard O. Burney4, Robert S. Schenken5, Randal D. Robinson5, Maria Gaczynska2, Pawel Osmulski2, Nameer B. Kirma2*, Bruce J. Nicholson1*

SUPPLEMENTARY MATERIAL
**TABLE S1: Genes in gene panel microarray**

| Group          | Gene   | Protein       | Group          | Gene   | Protein       |
|----------------|--------|---------------|----------------|--------|---------------|
| GJs            | GJA1   | Cx43          | Kinases        | MAPK1  | ERK1          |
|                | GJA3   | Cx46          |                | MAPK3  | ERK3          |
|                | GJA5   | Cx40          |                | PRKACA | Prot. Kin.Aα (cat. su) |
|                | GJA8   | Cx50          |                | PRKACB | Prot. Kin.Aβ (cat. su) |
|                | GJA9   | Cx59          |                | PRKACG | Prot. Kin. Aγ (cat. su) |
|                | GJB1   | Cx32          |                | PRKCA  | Prot. Kinase Cα |
|                | GJB2   | Cx26          |                | PRKCB  | Prot.Kinase Cβ |
|                | GJB3   | Cx31          |                | CDK1   | Cyclin Dep.kinase1 |
|                | GJB4   | Cx30.3        |                | CSNK1D | Casein Kinase 1Δ |
|                | GJB5   | Cx31.1        |                | SRC    | c-src         |
|                | GJB6   | Cx30          |                | NOV    | CCN3          |
|                | GJB7   | Cx25          |                | PLCB1  | Phospholipase B1 |
|                | GJC2   | Cx47          |                |        |               |
| Adhesion/TJs.  | TJAP1  | TJ assoc. protein | Markers        | VIM    | Vimentin      |
|                | TJP1   | ZO1           |                | MME/CD10 | Metallo-endopept. |
|                | TJP2   | ZO2           |                | EPCAM  | Ep CAM        |
|                | CAV1   | Caveolin      |                | KRT 18 | Cytokeratin 18 |
|                | CDH2   | N-cadherin    |                |        |               |
|                | SNAIL1 | Snail 1       |                |        |               |
|                | CTNNB1 | Beta-catenin  |                |        |               |
|                | DBN1   | Drebin 1      |                |        |               |

| Stand.         | GAPDH  | GAP dehydrog. |
|----------------|--------|---------------|
|                | ACTIN-B | Actin        |
Figure S1: Immunocytochemical assessment of Epithelial and Stromal cell isolations from two patients.

Endometrial cells are separated by differential sedimentation and adhesion into epithelial (left) and stromal populations (right). Immunohistochemistry [40x (top) and 10x (bottom)] demonstrate the purity of the isolations by dual staining with EpCAM antibodies to mark epithelial cells (red), and Vimentin antibodies to mark stromal cells (green). Nuclei are stained blue with DAPI in both.
Figure S2: Violin plots of average cell expression levels for all genes for which signals were detected and for all patients in stromal (A) and Epithelial cells (B).

Patients are color coded in legend, and are arranged in each plot from left to right as control (N), early stage (I/II) and late stage endometriosis. Phase of the cell cycle when samples were collected are indicated in the legend (P = proliferative; ES = early secretory; MS = mid-secretory; NC = non-cycling). The average value of control and early stage endometriosis in ESCs (A) or just control samples in EECs (B) is shown as a red horizontal line for reference, and the general pattern with endometriosis is indicated by arrows, and color of margin (red = decreased expression with endometriosis, green = increased expression). Note that in ESC samples collected at the MS stage and EECs (and in a few cases ESCs) collected from non-cycling patients, some genes show a divergence from general pattern of gene expression, indicating effects of the hormonal environment on expression.