Isolation and culture of human endometrial cells in a three-dimensional culture system

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A cell culture system was established in which endometrial stromal cells were embedded in a collagen matrix and separated from the endometrial epithelium by basement membrane material (Matrigel). The epithelium, seeded on top of the collagen matrix, grew in a monolayer. The cultures were evaluated by light microscopy and transmission and scanning electron microscopy. Light and transmission electron microscopy indicated a polarized columnar epithelium in monolayer with basally positioned nuclei. Scanning electron microscopy revealed a confluent epithelium with an abundance of microvilli and cilia, as well as pinopodes on the apical surface. Immunohistochemical staining for cytokeratin confirmed the epithelial origin of the surface cells, and staining for human collagen IV demonstrated its presence underneath the epithelial cells. This culture system represents a three-dimensional system that imitates the normal endometrium.

Introduction

Several culture systems have been used to study various aspects of mammalian endometrial metabolism (for review, see Findlay et al., 1990) and embryo–endometrial interactions (Lindenberg et al., 1985). However, none of these culture systems in vitro fully imitates the normal structure and function of the endometrium in vivo (Hearn, 1986).

Previous studies on endometrial epithelial monolayers in contact with the human embryo (Lindenberg et al., 1985) have given information only about the attachment of the embryo. As normal implantation is considered to be an active process involving both trophoblastic cells and cells of the secretory endometrium, it is questionable whether observations on ‘in vitro implantation’ reflect the true situation in vivo.

Endometrial stroma cells influence the development of epithelial cells (Mahfoudi et al., 1992) and may, alone or together with the extracellular matrix, play an important role in implantation. A possible significance of the extracellular matrix in implantation comes from a preliminary study in which human blastocysts were co-cultured on human endometrial epithelial cells supported by an extracellular matrix in vitro (B. Pedersen, A. Philip and S. Lindenberg, unpublished).

The object of this study was to establish a model in vitro that imitates the normal human secretory endometrium in vivo.

The model consisted of polarized epithelial cells resting on an artificial basement membrane (Matrigel), with underlying stromal cells embedded in a collagen matrix.

Materials and Methods

Isolation of tissues

Endometrial tissues were obtained by curettage (Cunell biopsy) from 12 normal ovulating women admitted to our department for tubal sterilization. The biopsies, 6–8 per patient, were taken at 3–14 days after the occurrence of the endogenous LH peak. All biopsies used were histologically classified as normal endometrium corresponding to the measured LH peak (Johannisson et al., 1982, 1987; Li et al., 1988).

Biopsy material from each patient was processed for light and electron microscopy and for immunohistochemistry (see below). One biopsy was frozen for further studies, and the remaining material was used for cell culture.

The study was approved by the National Ethical Committee (Denmark) and the women had signed informed consent forms.

Preparation of endometrial cultures

Tissue samples (approximately 30 mg wet mass) were transferred to flasks containing Ham’s F-10 medium (Nutrient
mixture F-10: GIBCO, Life Technologies, Roskilde), supplemented with 0.2% penicillin–streptomycin (10 000 IU ml\(^{-1}\); GIBCO). After washing, the tissue was minced into small pieces with a scalpel and incubated in 5 ml trypsin–EDTA (1 \(\times\) trypsin–EDTA solution; GIBCO), and supplemented with 0.05 g pancreatin ml\(^{-1}\) (Sigma Chemical Co, St Louis, MO) for 30 min at 4°C. This incubation was continued at room temperature in a Petri dish and the detachment of the glands and surface epithelium was carefully followed through a stereo microscope. The large epithelial glands and surface epithelium were sampled with a pipette into Ham’s F-10 medium, sedimented at 60 \(\times\) g for 10 min and resuspended in Ham’s F-10 medium. After this sampling the remaining ‘stromal cell fraction’ was supplemented with 5 ml fetal calf serum (GIBCO) to stop the enzyme digestion. After sedimentation, the supernatant was removed with a pipette and sedimented at 60 \(\times\) g for 10 min. The resulting pellet was treated with 150 IU collagenase ml\(^{-1}\) (Sigma) for 45 min at 37°C to obtain a single cell suspension. After sedimentation at 60 \(\times\) g for 10 min, the cells were washed in Ham’s F-10 and the two cell fractions could be seeded. The ‘stromal cell fraction’ was sedimented at 60 \(\times\) g for 10 min and the pellet was mixed with Vitrogen 100 (Celtix Laboratories, Palo Alto, CA), which consisted of bovine dermal collagen type I, adjusted to pH 7.4, and 2.3 mg collagen ml\(^{-1}\). The final cell concentration was 2–10 \(\times\) 10\(^6\) cells ml\(^{-1}\), and 300 \(\mu\)l of the mixture was placed in a Falcon Cell Culture insert 9 mm in diameter (Becton Dickinson, NJ).

The inserts were placed in six-well culture dishes (Nunclon multidish 6 wells Delta SI: NUNC A/S, Roskilde). The collagen gel was allowed to polymerize for 45 min at 37°C, after which the upper surface was coated with a thin layer of murine basement membrane material (Matrigel: Collaborate Research Inc, Bedford, MA). Matrigel was diluted 1:1 (v:v) with Alpha Modification of Eagle’s medium (Flow Laboratories, Irvine) and 500 \(\mu\)l was applied to the surface for 30 s. The suspension was then removed with a pipette and 3 ml culture medium was added to the bottom of the six-well dishes. After an additional 30 min at 37°C the ‘epithelial gland fraction’ was seeded on top of the artificial basement membrane and 400 \(\mu\)l culture medium was added into the cell culture insert. The optimal seeding density was achieved when the glands covered approximately 20% of the surface, allowing them to explant in a monolayer. Usually the cultures were confluent after 3 days. On average, six cultures per patient were established from the 30 mg tissue extracted; the number of stromal cells was the limiting factor.

Culture conditions

The culture medium used was Alpha Modification of Eagle’s medium (Flow Laboratories); 100 ml were supplemented with 0.2 ml penicillin–streptomycin (10 000 IU ml\(^{-1}\); GIBCO), 0.5 ml L-glutamine (200 mmol l\(^{-1}\); GIBCO), 2 ml Ultra-roser G (IBF-Biotechnics, Villeneuve-la-Garenne), 5 ml fetal calf serum (GIBCO), 0.5 g BSA (Behring Institut, Marburg), 10 \(\mu\)g retinoic acid (Sigma) and 1.75 \(\mu\)g progesterone. Since oestradiol concentration in the medium was already 35 nmol l\(^{-1}\), it was not supplemented with oestrogen.

All cultures were maintained in a humidified atmosphere with 5% CO\(_2\) in air and the medium was changed the day after seeding and then every other day. The cultures were examined under a Zeiss inverted phase contrast microscope.

Preparation for electron microscopy

For transmission electron microscopy and scanning electron microscopy, the cultures and fresh biopsied material were fixed in Karnovsky’s fixative (Karnovsky, 1965) in a sodium cacodylate buffer (0.1 mol l\(^{-1}\), pH 7.2; Merck, Darmstadt) for 60 min at 20°C.

For transmission electron microscopy, the cultures and fresh biopsied material were washed in 0.1 mol phosphate buffer l\(^{-1}\), pH 7.4, and fixed in 1% osmium tetroxide in 0.1 mol sodium cacodylate buffer l\(^{-1}\) for 2 h at 5°C. Subsequently, they were dehydrated and embedded in Epon (Merck). The embedded cultures and biopsies were sectioned into semi-thin and ultra-thin sections. Semi-thin sections were stained with toluidine blue, while ultra-thin sections were contrasted with uranyl acetate and lead citrate and examined under a Jeol JEM 100B electron microscope.

For scanning electron microscopy, the cultures and fresh biopsied material were washed in phosphate buffer, fixed in osmium tetroxide and dehydrated in graded ethanol. After drying to the critical point, the specimens were sputter-coated with gold and examined under a Jeol 25 SII microscope at 25 kV.

Preparation for light microscopy and immunohistochemistry

The cultures and fresh biopsied material were fixed in neutral-buffered 4% formaldehyde for 30 min at 4°C and dehydrated. The material was embedded in paraffin wax and cut to 5 \(\mu\)m sections. For light microscopy the sections were stained with haematoxylin–eosin. For immunohistochemistry (cultures only) a three-stage immunoperoxidase-staining procedure was used to demonstrate the presence of cytokeratin and human collagen IV. Endogenous peroxidase was inhibited by incubating the slides with 1% hydrogen peroxide in distilled water for 5 min. They were then washed in 0.05 mol Tris–HCl l\(^{-1}\) buffer, pH 7.6. Sections were pretreated with 0.2 mg trypsin ml\(^{-1}\) (Sigma) for 10 min at 37°C. After incubation in normal swine serum (Dako, Glostrup) that had been diluted 1:5 (v:v) in Tris buffer for 20 min at room temperature, the sections were incubated with monoclonal antibodies against collagen IV (mouse anti-human collagen IV, Dako) at a concentration of 1:50 (v:v) or cytokeratin (mouse anti-human cytokeratin, Dako) at a concentration of 1:10 (v:v) overnight at 4°C. Both antibodies were diluted in Tris buffer. Sections were then overlaided with rabbit anti-mouse immunoglobulin/ horseradish peroxidase (Dako) diluted 1:100 (v:v) in Tris buffer for 30 min at room temperature, followed by washing twice for 5 min each in Tris buffer. The last step was an incubation with swine anti-rabbit immunoglobulin/horseshad peroxidase (Dako) diluted 1:100 (v:v) in Tris buffer for 30 min followed by washing twice in Tris buffer for 5 min.

Negative controls were processed simultaneously by omitting the primary antibody; positive controls were skin.
Fig. 1. Light micrograph of histochemically stained cultured cells. (a) Positive cytokeratin staining (arrowheads) concentrated at the cell borders of the epithelium confirms the epithelial origin of these cells. (b) Staining for human collagen IV indicates its presence underneath the epithelial cells (arrowheads). (c) Negative control cultured cells were not stained. (Scale bar represents 30 µm.)

Fig. 2. Light micrograph showing (a) cultured polarized epithelial cells on the surface and stromal cells embedded in the collagen gel after culture for 4 days. The density of the stromal cells is lower than in (b) the biopsy material. The stain is toluidine blue. (Scale bars represent 15 µm.)

B 006 (Monocarb A/B, Lund) was used to detect nonspecific binding. Antibody binding was visualized using 3-amino-9-ethylcarbazole (Sigma) dissolved in acetone and 0.05 mol acetate buffer 1⁻¹, pH 5.0, for 9 min. After rinsing in distilled water the nuclei were counterstained with Mayers haematoxylin (Merck) for 5 min. The sections were mounted in Glycergel (Dako).

Fig. 3. (a) Scanning electron micrograph of cultured epithelial cells. Note the abundance of microvilli and cilia formation. Arrows indicate a pinopode. (b) Scanning electron micrograph of epithelial cells in a biopsy. (Scale bars represent 10 µm.)

Results

All biopsy specimens were evaluated by light microscopy and by transmission and scanning electron microscopy. The number of culture wells established from the biopsies of each of 12 patients was between 2 and 12. All cultures were examined using an inverted microscope with phase contrast, and confluent cultures from each patient were examined by transmission and scanning electron microscopy or immunohistochemistry. Altogether, 27 cultures were prepared for transmission electron microscopy, at least one per patient, and four cultures from four different patients for scanning electron microscopy. Twenty-four cultures, at least one per patient,
were embedded in paraffin wax for immunohistochemistry. Fifteen non-confluent or very shrunken cultures were omitted from the study.

**Immunohistochemistry**

Cytokeratin was detected in epithelial cells (Fig. 1a) concentrated at the cell borders, whereas cells beneath the basement membrane did not stain for cytokeratin. Staining for human collagen type IV demonstrated its presence underneath the epithelium (Fig. 1b). Matrigel is of murine origin and did not stain with this antibody. Negative controls were unstained (Fig. 1c).

**Morphology**

Light microscopy. The epithelial compartment of the cell cultures comprised polarized columnar cells with basally located nuclei (Fig. 2a). The stromal cells were distributed just beneath the epithelium, whereas the deeper part of the Vitrogen contained very few cells. Occasional gland formation was detected. Morphological resemblance between cultured and biopsied material (Fig. 2b) was demonstrated, but the cell cultures contained fewer stromal cells and showed no sign of capillary formation.

**Scanning electron microscopy.** The cultured epithelium was confluent and revealed an abundance of both cilia and microvilli. Pinopodes were seen (Fig. 3a). There were similarities between the cultured (Fig. 3a) and the biopsied material (Fig. 3b).

Transmission electron microscopy. The polarization of the cultured epithelium was confirmed, and the nuclei of the epithelial cells were located basally, similar to the biopsied material (Fig. 4a,b). The surface of both cultured and biopsied epithelial cells was covered with numerous slender microvilli (Fig. 4a,b). The cells were attached to each other by desmosomes (Fig. 4a, insert). The cytoplasm of both cultured and biopsied cells contained various amounts of glycogen and a well-developed organelle apparatus. The cells contained long, slender mitochondria in close association with rough endoplasmic reticulum, polyribosomes and glycogen deposits in the basal compartment (Fig. 5).

At the interface between the epithelium and stroma, a discontinuous basal lamina was observed (Fig. 6), and collagen was detected underneath the basement membrane, as well as in stromal protrusions in close association with the basal surface of the epithelial cells (Fig. 5).

The density of the stromal cells was less pronounced in the cultures (Fig. 4a,b). The stromal cells were rich in rough endoplasmic reticulum and polyribosomes (Figs 5 and 7), and
Fig. 5. Transmission electron micrograph of cultured cells: a stromal cell (bottom) is in close association with the basal surface of an epithelial cell (top). The cytoplasm of the epithelial cell contains glycogen, long slender mitochondria, rough endoplasmic reticulum and polyribosomes. Note the collagen beneath the epithelial compartment. The cytoplasm of the stromal cell contains polyribosomes and rough endoplasmic reticulum. (Scale bar represents 1 µm.)

Fig. 6. Transmission electron micrograph of basal lamina (arrows) underneath an epithelial cell. (Scale bar represents 1 µm.)

Fig. 7. Transmission electron micrograph of cultured stromal cells. The cells are not polarized and the cytoplasm is rich in slightly dilated rough endoplasmic reticulum (arrows) and polyribosomes. (Scale bar represents 2 µm.)

discernible only small amounts of glycogen and microfilaments in various amounts along cell borders.

Discussion

In mammals, blastocyst implantation is the process by which the fertilized ovum, which has developed to the hatched blastocyst stage, becomes intimately connected to the maternal tissue in the uterus. Since this process requires interactions between the embryo and the maternal tissue, it is useful to separate the events of implantation into several phases (Larsen, 1974; Schlafke and Enders, 1975). The study of these early stages of human implantation in vivo is for practical purposes impossible; several culture systems in vitro have therefore been developed.

In early experiments Gwatkin (1966) permitted a murine blastocyst to attach to plastic in order to determine culture requirements. Glenister (1961) and Grant et al. (1975) examined blastocyst implantation on rabbit endometrial strips and cultured murine uteri, allowing the blastocyst to penetrate the epithelium and basement membrane. However, necrosis meant that these culture systems were of no value. Lindenberg et al. (1984) made similar observations when culturing human endometrial tissue.

Studies of trophoblast interactions with isolated components of the endometrium, such as stromal cells (Salomon and Sherman, 1975), basement membrane material (Arman et al., 1986a,b) or extracellular matrix (Farach et al., 1987), or interactions with cultured murine epithelial monolayers gave only...
limited information (Van Blerkom and Chavez, 1981). A technique by which human endometrial epithelial monolayers were cultured on plastic was developed by Lindenberg et al. (1984). This system was suitable for studying apposition and attachment of human blastocysts to epithelial cells (Lindenberg et al., 1985), but provided no information about further stages of the implantation process.

We modified the model (Lindenberg et al., 1984) to examine trophoblast attachment to, and penetration through, the epithelial layer and interaction with the extracellular matrix and stromal cells. In a preliminary series of experiments (B. Pedersen, A. Philip and S. Lindenberg, unpublished), human endometrial epithelial cells were cultured on a collagen membrane and a hatched blastocyst was attached to the monolayer. This in vitro system was extended into a co-culture system, including stromal cells, that was embedded in a collagen matrix and separated from the epithelial monolayer by a basement membrane material, Matrigel, that was extracted from Engelbreth Holm Swarm mouse tumours (Kleinman et al., 1986). Matrigel has been demonstrated to promote a differentiated morphology of the endometrial epithelial cells in primary culture (Mahfoudi et al., 1992).

To study implantation, it is important to have a polarized epithelium with well-developed intercellular junctions that rests on a basement membrane. Furthermore, it is crucial to have a three-dimensional stromal compartment with both extracellular matrix and stromal cells, since interaction(s) between epithelium and the underlying stromal cells is (are) essential for the function of the endometrium (Inaba et al., 1988). It has been reported that the stroma may mediate oestrogen responsiveness in cultured murine uterine epithelia (Inaba et al., 1988).

In the present culture system the epithelial compartment was derived from glandular and surface epithelial cells. Human collagen IV was observed under the epithelial cells (Fig. 1b) showing cell polarization and indicating that the present culture model is a dynamic system. Although deposition of basement membrane, as revealed by transmission electron microscopy, could not be demonstrated initially (after 3 days) it did appear later, giving further support to the contention that this culture system is a dynamic environment. However, a prolonged culture time resulted in considerable shrinkage of the gel, which could be due to release of proteolytic enzymes by the cells.

The present culture system seems to fulfil the morphological three-dimensional criteria for an implantation system by presenting a polarized epithelium resting on a basement membrane, with an underlying collagen matrix containing stromal cells.

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