Basement Membrane Dissolution and Reassembly by Limbal Corneal Epithelial Cells Expanded on Amniotic Membrane

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**Purpose.** To investigate basement membrane (BM) formation during ex vivo expansion of limbal corneal epithelial cells on intact amniotic membrane (iAM) and epithelially denuded (d)AM.

**Methods.** Human limbal explants were cultured on iAM and dAM. Expression of BM components, including laminin-5, type IV collagen, type VII collagen, perlecan, integrin α6, and epithelial cell differentiation markers such as p63, cytokeratin 3 (K3), and cytokeratin 12 (K12), were investigated by immunostaining. Levels of matrix metalloproteinase (MMP)-2 and MMP-9 and tissue inhibitor of matrix metalloproteinase (TIMP)-1 in the conditioned media were determined by ELISA.

**Results.** All four BM components were preserved in both iAM and dAM before culturing, but dissolved 1 week afterward when MMP-2 was increased. Epithelial outgrowth correlated with increased expression of MMP-2 and -9 for both cultures. Resynthesis of BM began with laminin-5 followed by other components. This process took place at 1 week on iAM but at 2 weeks on dAM after culturing. At 4 weeks, BM was more maturely deposited as a linear band from the explant toward the leading edge on iAM and temporally correlated with a sharp decline of MMP-9 levels. In contrast, such BM deposition began at the leading edge on dAM only when TIMP-1 levels were increased. Epithelial cell outgrowth on iAM expressed more p63 but less K3 and K12 than did that on dAM.

**Conclusions.** After dissolution of original amniotic BM, new BM formed by ex vivo expanded human limbal corneal epithelial cells on iAM deposits much faster and is more mature, resulting in regeneration of a limbal epithelial phenotype. In contrast, BM deposition is delayed and remains immature on dAM, resembling wound healing by a corneal epithelial phenotype. Thus, BM resynthesis may be used as another objective readout resembling wound healing by a corneal epithelial phenotype.

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their discussed in the context of improving the future success of ex vivo expansion of limbal epithelial progenitor cells.

**MATERIALS AND METHODS**

Dulbecco’s modified Eagle’s medium (DMEM), Ham’s/F12 medium, HEPES buffer, amphotericin B, gentamicin, fetal bovine serum (FBS), and mouse epidermal growth factor (EGF) were purchased from Invitrogen (Carlsbad, CA). Dispase II was obtained from Roche (Indianapolis, IN). Hydorcotisone, dimethyl sulfoxide, cholera toxin, insulin-transferrin-sodium selenite medium supplement, propidium iodide, Triton X-100, bovine serum albumin (BSA), mouse anti-type IV collagen and type VII collagen monoclonal antibodies, FITC conjugated anti-mouse, goat, and rat IgGs were from Sigma (St. Louis, MO). Mouse anti-laminin-5 monoclonal antibody, rat anti-perlecan antibody, and mouse anti-cytokeratin 3 (K3) monoclonal antibody were from Chemicon (Temecula, CA). Mouse anti-human integrin α6 and α6 monoclonal antibodies were from DakoCytomation (Carpinteria, CA), goat anti-cytokeratin 12 (K12) polyclonal antibody was from Santa Cruz Biotechnology (Santa Cruz, CA), and ABC-AP kit (mouse IgG) was from Vector Laboratories (Burlingame, CA). A cell viability assay (Live/Dead) was obtained from Invitrogen (Eugene, OR). MMP-2 and -9 and TIMP-1 ELISA kits were from R&D Systems (Minneapolis, MN). Zymogram development buffer was from Bio-Rad Laboratories (Hercules, CA).

**Human Tissue Preparation**

Human tissue was handled according to the Declaration of Helsinki. Corneoscleral tissues from human donor eyes were obtained from the Florida Lions Eye Bank (Miami, FL) immediately after the central corneal button had been used for corneal transplantation. The tissue was rinsed three times with DMEM containing 50 μg/mL gentamicin and 1.25 μg/mL amphotericin B. After careful removal of excessive sclera, conjunctiva, iris, and corneal endothelium, the remaining tissue was placed in a culture dish and exposed to dispase II (1.2 U/mL in complete culture medium) at 37°C in humidified 5% CO2 for 10 minutes. After three rinses with complete culture medium for 10 minutes each, the scleral rim was trimmed to obtain limbal tissue cubes of 1 clock hour of ~1 × 1.5 × 2.5 mm size.

**Human Limbal Explant Cultures on AM**

Preserved human AM was kindly provided by Bio-Tissue (Miami, FL). Immediately before use, AM was thawed, washed three times with sterile PBS, and cut into pieces approximately 2.5 cm² in size. For preparation of dAM, membranes were deprived of their devitalized amniotic epithelial cells by incubation with 0.02% EDTA at 37°C for 1 hour, to loosen the cellular adhesion, followed by gentle scraping with a cell scraper. After that, AM with the epithelium or the BM side facing up was fastened to a culture insert, as previously reported. On the center of the AM insert, a human limbal explant was placed and cultured in supplemented hormonal epithelial medium (HEM) made of an equal volume of HEPES-buffered DMEM containing bicarbonate and Ham’s/F12. The medium was supplemented with 5% FBS, 0.5% dimethyl sulfoxide, 2 mg/mL mouse EGF, 5 μg/mL insulin, 5 μg/mL transferrin, 5 mg/mL selenium, 0.5 μg/mL hydrocortisone, 1 mM cholora toxin, 50 μg/mL gentamicin, and 1.25 μg/mL amphotericin B. Cultures were incubated at 37°C under 5% CO2 and 95% air, and the medium was changed every 48 hours, conditioned medium was centrifuged (3000g for 15 minutes), and the supernatant was stored at ~8°C. All explants were kept submerged in the culture medium (2 mL) throughout the culture duration. Freeze-thawed, devitalized explants and IAM and dAM without explants were used as the control. Parallel experiments were killed at the end of each week until 4 weeks, and AM together with the explant was removed and embedded in optimal cutting temperature (OCT) compound for cryosectioning.

**Histology and Immunostaining**

Cryostat sections (~4-μm) of the limbal explants as well as outgrowth on AM were fixed in acetone for 10 minutes at ~20°C. The sections used for immunostaining were rehydrated in PBS and then incubated in 0.2% Triton X-100 for 10 minutes. After three rinses with PBS for 5 minutes each and preincubation with 2% BSA to block nonspecific staining, the sections were incubated with primary antibodies (anti-type IV collagen at 1:400, others at 1:100) for 1 hour. After three washes with PBS for 15 minutes, they were incubated with an FITC conjugated secondary antibody (goat anti-rabbit or anti-mouse IgG at 1:100) for 45 minutes. After three additional PBS washes for 15 minutes, they were counterstained with PI (1:2000) for 1 minute, mounted in mounting medium, and analyzed with a fluorescence microscope. For immunohistochemical staining of p63, endogenous peroxidase activity was blocked by 0.6% hydrogen peroxide for 10 minutes. Nonspecific staining was blocked by 1% normal goat serum for 30 minutes. The sections were then incubated with p63 antibody (1:50) for 1 hour. After three washes with PBS for 15 minutes, sections were incubated with biotinylated rabbit anti-mouse IgG (1:100) for 30 minutes, followed by incubation with ABC reagent for 30 minutes. The reaction product was developed with 3,3’-diaminobenzidine (DAB) for 5 minutes, followed by dehydration, and examined under a light microscope.

**ELISA of MMP-2, MMP-9, and TIMP-1**

Quantitative ELISA assays for human MMP-2, MMP-9, and TIMP-1 were performed according to the manufacturer’s protocol. In brief, 100 μL of assay buffer and 50 μL of standard dilutions of recombinant human MMP-2, MMP-9, TIMP-1, and supernatants of the conditioned media were dispensed into a 96-well polystyrene microplate coated with a mouse monoclonal antibody against MMP-2, MMP-9, or TIMP-1. The plate was sealed, incubated for 2 hours at room temperature (RT). After three washes, 200 μL horseradish peroxidase–conjugated secondary antibody was added to each well and incubated for 1 hour at RT. After three washes, 200 μL of the substrate solution tetramethylbenzidine (TMB) was applied for 30 minutes, to develop a blue color, and the reaction was stopped by adding 50 μL of 2 N sulfuric acid. Absorbance was read at 450 nm by an automatic plate reader with a reference wavelength of 570 nm.

**Gelatin Zymography**

To determine the relative concentration and activity of gelatinases in the conditioned medium from limbal explant cultures on IAM or dAM, SDS-PAGE gelatin zymography was performed by using a previously reported method. Briefly, 10 μL of each conditioned medium was treated with SDS-PAGE sample buffer without boiling and reduction. Samples were fractionated in a 10% polyacrylamide gel by electrophoresis at 100 V for 90 minutes at 4°C. The gels were soaked in 0.25% Triton X-100 for 30 minutes at RT, to remove the SDS, and incubated in a development buffer (50 mM Tris-HCl [pH 7.5] 200 mM NaCl, 5 mM CaCl₂, and 0.02% Brij-35) containing 5 mM phenylmethylsulfonyl fluoride, a serine protease inhibitor, at 37°C overnight, to allow proteinase digestion of the substrate, and then stained with 5% Coomassie blue R-250 in 40% isopropanol for 2 hours and destained with 7% acetic acid. Gelatinolytic activities appeared as clear bands of digested gelatin against a dark blue background of stained gelatin.

**Statistical Analysis**

All experiments were repeated three times, each in triplicate. For MMP-2, MMP-9, and TIMP-1 detection, each group had six samples, and each sample was assayed in duplicate. Group means were compared using the appropriate version of Student’s unpaired t-test. Test results were reported as two-tailed, where P < 0.05 was considered statistically significant. Summary data are reported as the mean ± SD.
RESULTS

Epithelial Morphology

Limbal corneal epithelial cells started to migrate from the explant to iAM and dAM 4 to 5 days after culturing (Fig. 1A, 1E), which is consistent with our previous report. A prominent leading edge of outgrowth was visible on iAM cultures at 1 week (Fig. 1B), and spread out gradually, but did not reach confluence at 4 weeks of culturing (Fig. 1D). In contrast, the leading edge of outgrowth on dAM was not discernible by the naked eye, but could only be traced under the microscope (Fig. 1F, 1G). The outgrowth on dAM was subconfluent at 2 weeks and reached confluence at 4 weeks, indicating that the outgrowth rate on dAM was higher than that on iAM, which is also in agreement with our previous report. The epithelial morphology visualized by the cell viability assay showed that epithelial cells on iAM were uniformly small and had a cuboidal shape (Fig. 1I), whereas cells on dAM were large and had a migratory spindle shape (Fig. 1J). Hematoxylin and eosin (H&E) staining of the 4-week culture showed that the leading edge of the epithelial outgrowth on iAM was stratified whereas the remaining area had one to two cell layers (Fig. 2A). In contrast, the leading edge of the epithelial outgrowth on dAM was not stratified but was uniformly one to two cell layers throughout (Fig. 2B). Because the cell viability assay staining did not reveal any dead cells underneath the epithelial outgrowth on iAM (Fig. 1I), we concluded that there were no devitalized AM epithelial cells residing underneath the epithelial outgrowth. This interpretation was also supported by the finding that the devitalized amniotic epithelial cells were pushed away on iAM by expanding epithelial outgrowth (not shown).

BM Components in iAM or dAM before Culturing

H&E staining showed that a monolayer of amniotic epithelial cells was well kept on iAM before culturing, whereas these cells were completely removed on dAM after EDTA treatment and mechanical scraping (Fig. 3). The BM components of laminin-5, type IV collagen, type VII collagen, and perlecan were strongly expressed as a continuous band underneath the epithelial outgrowth, but decreased and became negative farther out from the outgrowth’s leading edge (arrow). In contrast, when an explant was cultured on dAM for 4 weeks (B), Ln-5, Col IV, Col VII, and perlecan started to deposit in a linear-segmented band close to the leading edge of the outgrowth (arrow) and showed a spotted and cytoplasmic staining in the major part of the outgrowth. Bar, 200 μm.
Degradation of Amniotic BM Components and Reassembly of New BM Components during Epithelial Outgrowth

To investigate whether there was any change in the BM components of AM during expansion of epithelial outgrowth from the limbal explant, we performed immunostaining of each BM component at different times of iAM and dAM culture. We noted unexpectedly that all four BM components that preexist in both iAM and dAM (Fig. 3) disappeared at 1 week of culture, suggesting that they were degraded after 1 week (Fig. 4). To strengthen our interpretation further, we cultured freeze-thawed, devitalized limbal explants for 2 weeks on either iAM or dAM as a control, as well as iAM or dAM without explants as another control, and noted that all four BM components were well preserved (data not shown). These findings further indicate that degradation of BM components was dependent on live cells derived from the limbal explant.

Continuous observation revealed that there was new synthesis of BM components by outgrowth of epithelial cells on iAM, as shown by initial intracellular expression, followed by punctate deposition beneath the cells, and finally by a continuous band formed at the interface between outgrowth of epithelial cells and AM stromal matrix. When explant was cultured on iAM, the earliest BM component expressed by the epithelial outgrowth was laminin-5 at 1 week, which exhibited strong intracellular staining, followed by the same staining pattern of type IV collagen, type VII collagen, and perlecan, stromal matrix. The new synthesis of Ln-5 was first noted in the stromal matrix. The new synthesis of Ln-5 was first noted in the epithelial outgrowth at 1 week (1W), but that of Col IV, Col VII, and perlecan started at 2 weeks (2W) and finished as a linear band on iAM by 4 weeks (4W). In contrast, new synthesis of these four components was delayed 1 week in dAM and there was still no linear staining by 4W. Expression of integrin α6 by epithelial outgrowth was intracellular at 1W on both iAM and dAM, but became linear at 2W on iAM but at 3W on dAM. Bar, 50 μm.

Figure 4. Immunostaining of BM components of laminin-5 (A), type IV collagen (B), type VII collagen (C), perlecan (D) and integrin α6 (E) during ex vivo expansion on iAM and dAM. At 1 week of culture, all four BM components disappeared in both iAM and dAM, indicating the degradation of preexisting BM components of AM. The reexpression of new BM components followed the order of intracellular punctuate staining, extracellular punctate deposition beneath the cells, and a continuous linear staining between the epithelial outgrowth and AM stromal matrix. The new synthesis of Ln-5 was first noted in the stromal matrix. The new synthesis of Ln-5 was first noted in the epithelial outgrowth at 1 week (1W), but that of Col IV, Col VII, and perlecan started at 2 weeks (2W) and finished as a linear band on iAM by 4 weeks (4W). In contrast, new synthesis of these four components was delayed 1 week in dAM and there was still no linear staining by 4W. Expression of integrin α6 by epithelial outgrowth was intracellular at 1W on both iAM and dAM, but became linear at 2W on iAM but at 3W on dAM. Bar, 50 μm.
beginning at 2 weeks. Compared with the iAM culture, there was a 1-week delay in the expression of all BM components in dAM cultures. At the end of 4 weeks of culturing, all four BM components showed a continuous band on iAM, but not on dAM, where the staining was of lower intensity and remained fragmented and an intracellular pattern.

At 4 weeks of culturing, the outgrowth on iAM still did not reach confluence whereas outgrowth on dAM had already reached confluence (Fig. 1). Low-magnification composites showed that laminin-5, type IV collagen, type VII collagen, and perlecan were all strongly stained continuously underneath the outgrowth on iAM. Moreover, expression of type IV collagen and perlecan was also found in the AM stroma with a gradient coming from the explant. Staining of all four BM components was notably stronger and more prominent toward the explant, but became more attenuated toward the leading edge of outgrowth and was negative in iAM outside the leading edge of outgrowth. These results indicate that deposition of BM components was mature after 4 weeks of culturing on iAM and constituted a gradient starting from the explant to the leading edge of the outgrowth. In contrast, when the explant was cultured on dAM, the staining of all four BM components was most prominent at the leading edge of the outgrowth in contact with the plastic wall, became markedly attenuated toward the explant, and showed an intracellular punctuate pattern in nearly all of the outgrowth. These results indicate that the deposition of BM components remained immature after 4 weeks of culturing on dAM and adopted a gradient starting from the leading edge of the outgrowth toward the explant, the opposite the gradient of iAM cultures.

Hemidesmosome Assembly of Epithelial Cell Outgrowths

Hemidesmosomes are specialized junctional complexes that mediate adhesion of epithelial cells to the underlying BM. Hemidesmosomal components include the α6β4 integrin, HD1 plectin, and the bullous pemphigoid antigens BP180 and BP230 (for review, see Ref. 23). We used integrin α6 as a marker to investigate hemidesmosome formation. The results showed that integrin α6 was present intracellularly along the outgrowth on both iAM and dAM from 1 week (Fig. 4B). The staining of integrin α6 became concentrated along the basal epithelial membrane on both iAM and dAM at 3 weeks and formed a continuous band at 4 weeks. At this time, the staining was much stronger on iAM than dAM, suggesting mature formation of hemidesmosomes and restoration of polarity of epithelial cell outgrowth.

Correlation of Degradation and Reassembly of BM Components with Levels of MMP-2, MMP-9, and TIMP-1 in Conditioned Media

To investigate the mechanism of degradation and reassembly of BM components in our culture system, we measured levels of MMP-2, MMP-9, and TIMP-1 proteins in the conditioned media with ELISA assays. MMP-2 and -9 activity was further analyzed by gelatin zymography. The results showed that MMP-2 concentration increased in the first week in iAM and dAM cultures. After that, MMP-2 decreased at the second week and maintained at a stable plateau at the third and fourth weeks in iAM cultures. However, a high level of MMP-2 was maintained from the second to fourth weeks in dAM cultures, and the concentration was higher in dAM than in iAM from days 10 to 26 (P < 0.05; Fig. 5A). In contrast, MMP-9 was kept at a negligible low level in the first week and increased dramatically in the second week on both iAM and dAM. A high level of MMP-9 was maintained from the second to fourth weeks on dAM, but decreased in the fourth week on iAM. MMP-9 concentration was higher in iAM than in dAM in the third week, but was lower in iAM in the fourth week (P < 0.05 at days 16, 22, 24, and 26; Fig. 5B). TIMP-1 also increased in the first week in both iAM and dAM cultures. After that, TIMP-1 decreased gradually and was kept at a low level on iAM from the second to the fourth weeks. However, TIMP-1 remained at a high level in dAM and increased dramatically in the fourth week (P < 0.01 at day 10, 12, and days 18–26; Fig. 5C). The control without explants showed negligible or undetectable levels of MMP-2, MMP-9, and TIMP-1 throughout the culture duration.

Gelatin zymography assay results showed an active form of MMP-2 only in the explant-conditioned medium (Figs. 6A, 6B), but not in the control culture with iAM (Fig. 6C) or...
dAM (Fig. 6D) alone. MMP-2 expression was relatively stable throughout the culture period and was higher on dAM (Fig. 6B) than on iAM (Fig. 6A), in agreement with ELISA assay results. MMP-9 expression was undetectable at 4 days when the explant was cultured on either iAM or dAM, but it quickly increased in the second week. Expression of MMP-9 remained high in the second and third weeks on both iAM and dAM, but decreased in the fourth week on iAM but not on dAM. This finding was also in agreement with ELISA assay results. MMP-9 activities were undetectable in the control of either iAM or dAM without explants.

Correlation of Epithelial Differentiation and Reassembly of New BM Components

Deposition and assembly of the BM occurs concurrently with the normalization of epithelial growth, morphogenesis, and differentiation. To study the differentiation of corneal limbal epithelial cells expanded on iAM or dAM, we investigated the expression of p63, K3, and K12 in the outgrowth. The results showed that after 4 weeks of culture, there was more nuclear staining of p63 in the outgrowth on iAM. K3 was negative or weakly positive in the basal epithelial cells on iAM, but was strongly positive in all epithelial cell outgrowths on dAM. K12 was weakly positive on iAM, but almost all cells were strongly positive on dAM (Fig. 7). These results indicated that the epithelial cell outgrowths on iAM remained poorly differentiated, resembling limbal basal epithelial cells, whereas cells on dAM were more differentiated, similar to corneal epithelial cells.

DISCUSSION

BM formation is an important criterion for structural and functional regeneration of the epidermis when skin equivalents are engineered and transplanted. Only tissues with well-structured BM show optimal epithelial tissue architecture after transplantation. This study is the first attempt at determining whether BM formation can be used as another readout to assess the success of ex vivo expansion of limbal epithelial progenitor cells before transplantation.

Our study demonstrates dynamic changes of BM degradation and reassembly during ex vivo expansion of limbal corneal epithelial cells on AM, and provides four important findings. The first surprising finding was the degradation of preexisting BM components such as type IV collagen, type VII collagen, laminin-5, and perlecan on iAM and dAM before migration of epithelial outgrowth (Figs. 3, 4). In mouse corneas, epithelial BM is found partially disassembled after extensive epithelial debridement before complete epithelialization. If what we observed herein also had taken place in vivo, it would seriously challenge our prior belief that preexisting amniotic BM components serve as a scaffold to promote epithelialization after AM transplantation (for review, see Ref. 29). Because these BM components were still well preserved on both iAM and dAM without limbal explants or with devitalized limbal explants after 2 weeks of culture in the same medium (data not shown), we conclude that the degradation of preexisting BM is due to the release of gelatinases by cells in live limbal explants. Because MMP-2 levels were readily detectable and dramatically increased in both iAM and dAM cultures, whereas MMP-9 levels were negligible during the first week when epithelial migration just started, we strongly speculate that MMP-2, not MMP-9, is responsible for BM degradation. This speculation is supported by the finding that MMP-2 can degrade type IV collagen, type VII collagen, and laminin-5. Because selective action of MMP-2 and -9 in explant culture conditions, but not in the control without explant cultures (C, D).

FIGURE 6. Gelatin zymography of MMP-2 and -9 in explant culture conditioned medium. The active form of MMP-2 was only present in explant culture on iAM (A) and dAM (B) conditioned medium, but not in the control with iAM (C) or dAM (D) alone without explants. MMP-2 activities were relatively stable throughout the culture period, but the activity was higher on dAM than on iAM. MMP-9 activities were undetectable at 4 days when the explant was cultured on both iAM (A) and dAM (B), but quickly increased in the second week. A high expression of MMP-9 was maintained in the second and third weeks on both iAM and dAM, but it decreased in the fourth week on iAM, but remained at a high level on dAM. MMP-9 activities were undetectable in the control without explants (C, D).

FIGURE 7. Differentiation markers expressed by outgrowth epithelial cells expanded on iAM or dAM for 4 weeks. There was more nuclear staining of p63 on iAM than dAM. K3 was negative or weakly positive in basal epithelial cells on iAM, but was strongly positive in all outgrowth epithelial cells on dAM. K12 was weakly positive on iAM, whereas almost all cells were strongly positive on dAM. Bar, 50 μm.
MMP-2 on laminin-5 results in stimulation of mammary epithelial cell migration, we speculate that the removal of preexisting laminin-5 is an important stimulus to promote epithelial outgrowth migration onto either iAM and dAM. Because BM also acts as a reservoir for latent growth factors and cytokines, such as transforming growth factor-β and basic fibroblast growth factor, proteolysis of BM components by MMP-2 may help release and activate these factors-cytokines that may play an important role in triggering or supporting subsequent epithelial migration, proliferation, and differentiation. Because MMP-2 may be secreted by both epithelial cells and fibroblasts in the anterior segment of the eye (for review, see Ref 37), future studies are needed to determine whether limbal epithelial cells, stromal mesenchymal cells, or both are responsible for the early release of MMP-2 into the conditioned medium and how such an early release is regulated.

The second finding was that active epithelial outgrowth from the explant correlated with a dramatic increase of MMP-9 in the conditioned medium of both iAM and dAM cultures (Fig. 5B). Recently, Sun et al. also reported elevation of MMP-9 in the conditioned medium during the outgrowth of limbal explants on AM. MMP-9 has been found expressed mostly in migrating epithelial cells in branchial epithelial wounds and in migrating basal corneal epithelial cells in scrape or excimer laser-ablated wounds. The mechanism by which MMP-9 is upregulated in our system remains unclear. In epidermal keratinocytes, absence of laminin-5 triggers upregulation of phosphorylation of p38 MAPK, which has been linked to corneal epithelial migration and to upregulation of MMP-9 in other cell systems. Therefore, it is possible that early dissolution of BM components by MMP-2 triggers phosphorylation of p38 and as a result induces MMP-9 expression. At this moment, we do not know why MMP-9 decreased dramatically in iAM culture, but stayed at a high level in dAM during the fourth week.

The third finding revealed different patterns of BM formation on iAM and dAM. In several models of epidermal wound healing, BM deposition is a late event and starts at the leading edge after migrating epithelial cells fill in the wounded area, presumably due to “contact inhibition.” A similar phenomenon was observed in iAM cultures where linear BM deposition started at the leading edge when epithelial outgrowth reached the plastic wall at the end of 4 weeks and became markedly attenuated toward the explant (Fig. 4B). In contrast, BM linear deposition in iAM cultures started much earlier, before the leading edge reached the plastic wall, and formed a gradient starting from the explant toward the leading edge, a pattern exactly opposite to that of dAM cultures (Fig. 2A). This surprising finding prompted us to speculate that epithelial cell growth and migration on dAM resemble epithelial wound healing, whereas cells expanding on iAM resemble epithelial regeneration. The major difference between iAM and dAM is the presence of devitalized amniotic epithelial cells in the former. Besides the possibility that such devitalized amniotic epithelial cells serve as a physical barrier to slow down the outgrowth, which may trigger BM deposition, the other possibility is that they serve as a feeder layer to promote BM formation. This concept was suggested by a recent report showing the success of using devitalized amniotic epithelial cells as a feeder layer to support growth and maintain the undifferentiated status of primate embryonic stem cells. In epidermal organotypic cultures, BM assembly is facilitated when keratinocytes are cocultured with dermal fibroblasts. Future studies are needed to determine whether the protocol of coculturing limbal epithelial outgrowth with T3T fibroblast feeder layers may also promote BM assembly in dAM cultures, and if so whether such a mechanism lends support of this method in expanding limbal epithelial progenitor cells.

Temporal and spatial correlation further indicated that deposition of BM components started from laminin-5 followed by type IV, VII collagen and perlecan, and that the pattern started from subepithelial punctuate staining to a linear band in both iAM and dAM cultures (Fig. 3). These results further support the pivotal role of laminin-5 in paving the way of BM deposition. Although at this moment, we do not know what factor(s) triggers laminin-5 deposition.

In epidermal organotypic cultures, addition of MMP inhibitors promotes BM formation, indicating that a decrease in MMP leads to BM deposition. If this interpretation is correct, we noted herein the fourth intriguing finding that a sharp decline of MMP-9 levels without a concomitant increase of TIMP-1 was temporally correlated with mature deposition of BM in iAM cultures. In contrast, an increase of TIMP-1 was correlated with BM deposition on dAM during the fourth week. These findings further suggest that different mechanisms are used for BM formation during limbal epithelial outgrowth on iAM and dAM. Future studies are needed to investigate how MMPs and TIMP-1 are differentially modulated in iAM and dAM culture.

We noted that early BM reassembly by iAM correlated with higher expression of p63 and lower expression of K3 and K12, compared with dAM (Fig. 7). Although studies from different groups showed discrepancies of p63 expression patterns in corneal and limbal epithelium, it was generally believed that limbal basal p63-positive cells include corneal epithelial stem cells, and that p63 positivity also represents cells in a proliferative state, such as transit amplifying cells (for review, see Ref. 54). Together with expression of K3 and K12, which had been used as differentiated corneal epithelial cell marker, we believe early BM formation correlates well with the preservation of limbal epithelial phenotype. These results were consistent with our previous report showing that ex vivo expansion on iAM preserves the limbal epithelial phenotype, whereas that on dAM promotes a corneal epithelial phenotype. We then propose that BM formation may be used as another objective sign for assessing the experimental variables used in different protocols for successful ex vivo expansion of limbal epithelial progenitor cells in the future.

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References


