Morphogenesis of Biologically Active Interfollicular Epidermis from Human Embryonic Stem Cell-derived Keratinocytes

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Abstract

For patients suffering from extensive burns or chronic wounds, allogeneic cell-based therapies provide a means to restore viability and function to severely damaged cutaneous tissues. Pluripotent stem cells have been proposed as an allogeneic source for regenerative medicine applications, such as those required for burn and wound management. Ultimately the clinical utility of pluripotent stem cells relies on the ability to direct differentiation into the desired cell lineage, facilitate appropriate tissue formation, and confirm tissue-specific biological activity. We examined the capacity of human embryonic stem cell-derived keratinocytes (hES-DK) to undergo morphogenesis and form biologically active interfollicular epidermis using methods routinely employed to engineer skin substitutes for clinical applications. Throughout directed differentiation, the orderly sequence of epidermal gene expression mimicked the progression of fetal skin development. When introduced into three-dimensional organotypic culture, hES-DK cells formed a pluristratified tissue with architecture similar to that of the interfollicular epidermis. In hES-DK tissue the expression and localization of cell-cell adhesion proteins, markers of both early- and late-stage keratinocyte terminal differentiation, and host defense peptides were comparable to the patterns observed in keratinized stratified squamous epithelia generated from epidermal keratinocytes. Despite similar tissue morphology, functional analysis revealed that hES-DK tissues did not display robust cutaneous barrier function. However, hES-DK tissues were shown to possess antimicrobial activity, which represents the first demonstration of biological activity in a keratinized stratified squamous epithelium generated from a pluripotent stem cell-derived source. The successful generation of biologically active hES-derived stratified squamous epithelia represents a significant advance in the development of a hES-derived bioengineered human organ for clinical use.

Keywords: Regenerative medicine; Allogeneic cell-based therapy; Human Embryonic Stem cell-derived keratinocyte; hES-derived; Biologically active interfollicular epidermis

Introduction

The utility of pluripotent stem cells, such as human embryonic stem (hES) and human induced pluripotent stem (hiPS) cells, for regenerative medicine applications relies on the cells’ ability to differentiate into clinically relevant cell populations and form constructs that recapitulate the structure and function of the intended organ system. Directed differentiation protocols have been employed to generate multiple cell types, including those of neural, hematopoietic, cardiac, and pancreatic lineages [1,2], as well as epithelial lineages, including keratinocytes [3-9]. For patients suffering from extensive burns or chronic wounds, allogeneic cell-based therapies provide a means to restore the viability and/or function of affected tissues [10]. Fundamentally, stem cell therapy in burn and wound management requires a consistent source of keratinocytes with the capacity to fully restore functional epidermis. To date, a thorough characterization of the structural properties and, more importantly, the biological activities of keratinized stratified squamous epithelia generated from a pluripotent stem cell source has not been reported.

Initial studies describing the derivation of keratinocyte precursors from hES cells relied on spontaneous differentiation events such as teratoma or embryoid body formation [3-5]. The resulting heterogeneous populations displayed sporadic differentiation into a keratinocyte lineage as evidenced by expression of markers such as keratin 14 (K14) or yielded keratinocyte-like cells with limited expansion potential. The efficiency of selecting K14+ keratinocyte-like cells was subsequently enhanced by treating with retinoic acid (RA) and bone morphogenetic protein-4 (BMP4) [6]. Serial cultivation yielded a relatively pure population of keratinocyte-like cells which expressed several proteins associated with early-stage keratinocyte terminal differentiation in monolayer culture. In 2011, a differentiation protocol employing RA and BMP4 was also reported to facilitate the generation of K14-expressing keratinocytes from hiPS cells [7]. Differentiation methods using RA and Activin [8], as well as ascorbic acid and BMP4 [9], have also been shown to generate K14+ keratinocytes from hES cells provided the specified cultivation steps were adhered to.

Organotypic culturing techniques, which promote full stratification and differentiation of epidermal and oral keratinocytes under in vivo-like conditions, have fostered the development of the first engineered human tissues for clinical use [11-13]. These specialized techniques have enabled the formation of three-dimensional epithelial tissue constructs from hES-derived cell populations [9,14-16]. Dabelstein et al. [14] first demonstrated epithelial tissue morphogenesis using hES-derived, keratinocyte-like cells transduced with the oncogenic E6 and E7 genes of HPV16 to enable serial cultivation [14]. Three-dimensional tissue has also been generated from hiPS-derived keratinocytes and, although characterization was limited, was shown to express the...
terminal differentiation markers keratin 1 (K1) and loricrin [7]. Metallo et al. [6] utilized hES-derived cells to generate a tissue that, despite several atypical characteristics, displayed many morphological hallmarks of keratinized stratified squamous epithelia [15]. Recently, Leydon and coworkers demonstrated production of a non-keratinized stratified squamous epithelium from hES-derived simple epithelial cells created through employing the initial stages of Metallo’s differentiation method [16]. This tissue was shown to differentially express a subset of proteins in a pattern consistent with vocal fold epithelium. In 2009, Guenou et al. [9] formed three-dimensional tissue from a hES-derived cell source that displayed both the characteristic morphology of keratinized stratified squamous epithelium and appropriate localization of several keratinocyte terminal differentiation markers [9]. A thorough characterization of the distinctive structural and functional proteins typical of a keratinized stratified squamous epithelium generated from both a hES or hiPS cell-derived source and the associated biological activities, has not yet been reported.

Here we describe the morphogenesis of biologically active keratinized stratified squamous epithelium from hES-derived keratinocytes (hES-DK). Treatment of hES cells with RA alone did not induce expression of K14 however, serial cultivation on gelatin-coated plates selected for cells expressing key keratinocyte markers. This hES-DK population was capable of forming a pluristratified epithelium when cultivated under organotypic conditions. Epidermal morphogenesis was confirmed by comprehensively evaluating the expression and appropriate localization of proteins associated with the structure and function of keratinized stratified squamous epithelia, including cell-cell adhesion proteins, markers of both early- and late-stage keratinocyte terminal differentiation, and host defense peptides (HDPs). The epidermis serves as a barrier to transepidermal water loss and functions to prevent bacterial invasion through secretion of HDPS. Although hES-DK tissue did not display cutaneous barrier function, we found that hES-derived epithelial tissue possessed antimicrobial activity, which has not previously been reported in a hES cell-derived tissue. These findings suggest that a well-defined hES-derived cell population, as confirmed by the expression of key epidermal markers, is required for the morphogenesis of biologically active interfollicular epidermis.

Materials and Methods

Monolayer and organotypic cell culture methods

hES cells (H9) [17] were obtained from the WiCell Research Institute (Madison, WI). Undifferentiated hES cells were cultivated on mitomycin C-inactivated murine fibroblasts in hES growth medium [4]. For passaging, cells of the feeder layer were dislodged by pipetting and rinsed free, facilitating hES cell harvest by manual colony dissociation. For directed differentiation, feeder cells were removed and adherent hES colonies were cultivated for 6 days in hES growth medium or medium supplemented with 0.1% dimethyl sulfoxide (DMSO) or 1 μM RA. Cells were dislodged using dispase and transferred into suspension culture in hES medium. The following day, aggregates of differentiated cells were transferred to plates coated with 0.1% gelatin (Sigma-Aldrich, St. Louis, MO) and maintained in defined keratinocyte serum-free medium (dKSFM) (Invitrogen, Carlsbad, CA). For serial cultivation, subconfluent cells were harvested by trypsinization and re-plated onto gelatin-coated plates in dKSFM. Cell and colony morphology were documented using an Olympus IX-70 inverted microscope equipped with a DEI-750 camera (Optronics Engineering, Goleta, CA) and Image-Pro Plus software (Media Cybernetics, Silver Spring, MD). Human epidermal keratinocytes were cultivated on murine feeder layers in standard keratinocyte culture medium as previously described [18,19].

To generate three-dimensional tissues, cells were harvested by trypsinization and seeded at a density of 5 × 10^5 cells/cm^2 onto a cellulized dermal matrix (Stratatech Corporation, Madison, WI) consisting of human dermal fibroblasts embedded in type I collagen in StrataLife media (Stratatech Corporation, Madison, WI) as previously described [19]. Cultures were kept submerged for 5 days, then raised to the air-interface and maintained for an additional 24 days to facilitate terminal differentiation. Tissue samples were prepared for histological evaluation as previously described.

Reverse transcription-polymerase chain reaction (RT-PCR) analysis

Total RNA was isolated using TRIzol Reagent according to the manufacturer’s instructions (Invitrogen, Carlsbad, CA). Following DNase I treatment of 2 μg total RNA using the DNA free kit (Ambion, Austin, TX), reverse transcription was performed using oligo dT primers and M-MLV reverse transcriptase (Invitrogen, Carlsbad, CA) per manufacturer’s instructions. cDNA was analyzed by PCR amplification using the primer pairs listed in Supplementary Table 1. Amplified products were visualized by agarose gel electrophoresis and gene expression was confirmed by identification of the appropriate bands. GAPDH served as a loading control.

Indirect immunofluorescence

Cryopreserved sections were incubated for 1 hour at 37°C with primary antibody (Supplementary Table 2) diluted in blocking agent followed by incubation for 30 minutes at room temperature with goat anti-mouse or anti-rabbit IgG Alexa-594 secondary antibody (Invitrogen, Carlsbad, CA). Sections incubated with blocking agent served as controls for non-specific staining. Nuclei were counterstained with 5 μg/ml Hoechst 33258 in PBS. Digital image capture of individual fluorescence signals was performed using an Olympus IX-70 inverted fluorescent microscope equipped with Texas Red and Hoechst band pass filters, an Olympus DP70 digital camera, and DP Controller software (Olympus, Center Valley, PA). Identical image manipulations were performed for each set and dual-color images were created by overlay of single color captures of the same field.

Barrier function analysis

Electrical impedance, as measured by a Nova Derma Phase Meter (DPM) 9003 equipped with a DPM 9107 sensor probe (NOVA Technology Corporation, Gloucester, MA), was used to evaluate barrier function. The magnitude of impedance change during a defined measurement period correlates with the rate of transepidermal water loss [20], thus a large change in impedance as measured by DPM units is indicative of impaired barrier function. The probe was placed in direct contact with the tissue surface and impedance measurements were recorded at 0.5 second intervals for 10 seconds. hES-DK tissues resulting from three independent differentiation series were evaluated. For comparison studies, triplicate stratified squamous epithelial tissues generated from epidermal keratinocytes were analyzed. Three measurements were also taken from native skin. All values are reported as the mean ± SD. Contiguous epithelialization of hES-DK tissues was confirmed by the presence of a uniformly-stained culture surface after exposure to 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl tetrazolium.
findings show that although RA induced expression of the epithelial collection of cells with no histological features characteristic of simple organotypic culture of RA-treated hES cells resulted in a disorganized organized pseudostratified columnar epithelia (Figure 1D). In contrast, cells cultivated under organotypic conditions primarily formed well-organotypic culture to investigate the ability of these cells to form early marker of keratinocyte differentiation, was not induced by 6 days epithelia [23], increased with RA treatment. Expression of K14, an intermediate filament primarily associated with simple in the proliferative compartment of epithelial tissues, and keratin 18 mRNA expression of both ΔNp63, a transcription factor expressed was observed for the neural ectoderm differentiation marker Sox 1. mRNA decreased. RA has previously been reported to attenuate neural differentiation [22]. Although mRNA expression of the intermediate filament nestin remained unchanged in RA-treated cells, a reduction of expression and differentiation of hES cells were examined. After 6 days, both untreated (data not shown) and vehicle-treated control monolayer cultures (Figure 1A) contained fusiform, fibroblastic cells at the periphery of hES colonies. In contrast, a cobblestone, epithelial morphology was observed on the outer third of hES colonies treated with RA (Figure 1B). RA-treated hES cells were analyzed for gene expression of neural ectoderm and epithelial differentiation markers by RT-PCR (Figure 1C). Following RA treatment, expression of Oct4 mRNA decreased. RA has previously been reported to attenuate neural differentiation [22]. Although mRNA expression of the intermediate filament nestin remained unchanged in RA-treated cells, a reduction was observed for the neural ectoderm differentiation marker Sox 1. mRNA expression of both ΔNp63, a transcription factor expressed in the proliferative compartment of epithelial tissues, and keratin 18 (K18), an intermediate filament primarily associated with simple epithelia [23], increased with RA treatment. Expression of K14, an early marker of keratinocyte differentiation, was not induced by 6 days of RA treatment alone.

DMSO- and RA-treated hES cells were next introduced into organotypic culture to investigate the ability of these cells to form stratified tissue. Histological analysis revealed that DMSO-treated cells cultivated under organotypic conditions primarily formed well-organized pseudostratified columnar epithelia (Figure 1D). In contrast, organotypic culture of RA-treated hES cells resulted in a disorganized collection of cells with no histological features characteristic of simple or stratified squamous epithelia (Figure 1E). Taken together these findings show that although RA induced expression of the epithelial markers ΔNp63 and K18 in hES cells, RA treatment alone did not promote K14 expression or induce differentiation of hES cells into an epithelial cell type capable of forming stratified squamous epithelia.

**hES-DK cells form keratinized stratified squamous epithelia displaying characteristics of the interfollicular epidermis**

Enrichment of RA-treated hES cultures for cells differentiating into the keratinocyte lineage was next explored. We used serial cultivation on gelatin-coated plates to select for cells exhibiting epithelial morphology. hES cells treated with DMSO or RA were transferred onto gelatin-coated culture plates and maintained in monolayer culture for two passages (Figure 2A). Typical of spontaneous differentiation, the cell population resulting from DMSO treatment was morphologically heterogeneous (data not shown). Conversely, after one passage onto gelatin-coated culture plates, RA-treated hES cells yielded a cell population, hES-derived epithelial (hES-DEp) cells, that exhibited a more flattened, epithelial morphology at the periphery of the expanding colony (Figure 2B). Harvest and subsequent passage of these hES-DEp cells onto gelatin-coated culture plates resulted in a population where the predominant cell type (hES-DK) exhibited a cobblestone morphology (Figure 2C) typical of epidermal keratinocytes (Figure 2D). These results strongly suggest that serial cultivation onto gelatin-coated plates served as a selection for cells with a keratinocyte phenotype.

**Results**

**Retinoic acid-treated hES cells do not form stratified squamous epithelia in organotypic culture**

To determine if retinoic acid alone was sufficient to induce keratinocyte differentiation, the effects of RA treatment on gene expression and differentiation of hES cells were examined. After 6 days, untreated (data not shown) and vehicle-treated control monolayer cultures (Figure 1A) contained fusiform, fibroblastic cells at the periphery of hES colonies. In contrast, a cobblestone, epithelial morphology was observed on the outer third of hES colonies treated with RA (Figure 1B). RA-treated hES cells were analyzed for gene expression of neural ectoderm and epithelial differentiation markers by RT-PCR (Figure 1C). Following RA treatment, expression of Oct4 mRNA decreased. RA has previously been reported to attenuate neural differentiation [22]. Although mRNA expression of the intermediate filament nestin remained unchanged in RA-treated cells, a reduction was observed for the neural ectoderm differentiation marker Sox 1. mRNA expression of both ΔNp63, a transcription factor expressed in the proliferative compartment of epithelial tissues, and keratin 18 (K18), an intermediate filament primarily associated with simple epithelia [23], increased with RA treatment. Expression of K14, an early marker of keratinocyte differentiation, was not induced by 6 days of RA treatment alone.

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hES-DEp cells, hES-DK cells, and human epidermal keratinocytes were then cultivated using organotypic conditions to assess the capacity of these cells to form stratified squamous epithelial tissue. The tissue architecture resulting from hES-DEp cells did not resemble stratified squamous epithelia and was disorganized with numerous, atypical cells (Figure 2E). Surface coverage was incomplete and although limited regions displayed an epithelial appearance, highly vacuolized cells were observed throughout the tissue. In contrast, hES-DK cells formed a multi-layered stratified squamous epithelium (Figure 2F). Epithelial KO

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**Figure 1:** RA-treated hES cells express epithelial markers, but do not form organized structures when cultivated under organotypic conditions. Representative phase contrast images depicting the cell and colony morphology of hES cells after treatment with hES growth medium supplemented with DMSO (A) or RA (B). RT-PCR detection of neural or epithelial markers for untreated, DMSO-treated, and RA-treated samples (C). Results are representative of three independent experiments. Histological analysis of tissue generated from DMSO-treated hES cells revealed formation of well-organized pseudostratified columnar epithelia (D). Culture of RA-treated hES cells resulted in disorganized structures with limited areas of stratified epithelia (E). Scale bar represents 500 µm (A and B); 50 µm (D and E).
coverage was contiguous, although variation in the thickness of stratified hES-DK tissue was observed. Basal, spinous, and granular layers were evident throughout the tissue. Flattened, enucleated squames were present at the apical surface of stratified hES-DK tissue, however hES-DK tissue lacked a well-defined stratum corneum. Squames were more frequently observed in thinner regions. Stratified squamous epithelia generated from human epidermal keratinocytes (Figure 2G) exhibited a consistent thickness with well-defined basal, spinous, granular and cornified layers.

To evaluate the progression of hES-DK cell development in monolayer culture, samples from each differentiation step were analyzed for gene expression of markers indicative of neural ectoderm and epithelial differentiation, as well as keratinocyte terminal differentiation (Figure 2H). Oct4 mRNA expression was not detected in hES-DEp and hES-DK monolayer cultures or in tissues generated by hES-DK cells. Sox1 and nestin expression in hES-DEp cells was similar to RA-treated hES cells, but absent in hES-DK cells and hES-DK tissue. Compared to ΔNp63 levels in stratified squamous epithelia generated from human epidermal keratinocytes, low levels of ΔNp63 mRNA were detected at all stages of hES-DK differentiation. Robust K18 expression was observed in RA-treated, hES-DEp, and hES-DK cells, as well as in tissues generated by hES-DK cells or epidermal keratinocytes. K14 mRNA levels were low in the hES-DK population; however K14 mRNA levels increased substantially in hES-DK cells. K14 levels in hES-DK tissue were comparable to tissues generated with human epidermal keratinocytes. Expression of the late-stage keratinocyte differentiation markers involucrin and loricrin were first detected within hES-DK cells. These findings suggest that a defined cell population robustly expressing key markers of epithelial differentiation was required for epidermal morphogenesis.

Proteins essential to the structure and function of keratinized stratified squamous epithelia are expressed and appropriately localized in hES-DK tissue

hES-DK cell morphogenesis into stratified squamous epithelia prompted an investigation of the expression and localization of proteins necessary for the structure and function of stratified squamous epithelia. In these studies hES-DK tissues were compared to epithelial keratinocyte-generated tissues. E-cadherin appropriately localized to the membrane of cells positioned within the basal and immediately suprabasal layers (Figures 3A and 3B), although staining was generally more intense in hES-DK tissue compared to stratified squamous epithelia generated from epidermal keratinocytes. P-cadherin protein expression was localized to the cell membrane and restricted to cells of the basal and immediately suprabasal layers (Figures 3C and 3D), although staining appeared substantially diminished in hES-DK tissues relative to epidermal keratinocyte-generated tissue. The β-catenin expression pattern (Figures 3E and 3F) was similar to that of E-cadherin and no nuclear localization was detected. The expression of keratin intermediate filaments is highly regulated and dependent upon the stage of keratinocyte terminal differentiation [24]. Typical for stratified squamous epithelia, the expression of K14 protein was found in basal and suprabasal layers of tissues generated from hES-DK and epidermal keratinocytes (Figures 3G and 3H). For all tissues examined, K1 was appropriately localized to the suprabasal layers (Figures 3I and 3J) while keratin 2 (K2) was properly restricted to the upper spinous and granular layers (Figures 3K and 3L). Nuclear localization of the transcription factor p63 was detected exclusively in cells of the basal layer in hES-DK and control epidermal keratinocyte tissues (Figures 3M and 3N). Punctuate staining for filagrin, a late-stage keratinocyte terminal differentiation marker, was found in the cytosol of cells located in the granular layer for hES-DK and human epidermal keratinocyte-derived tissues (Figures 3O and 3P). Similarly, keratinocyte type 1 transglutaminase (Tgase1) was properly localized to the cell membrane of the spinous and granular layers (Figures 3Q and 3R). These results demonstrated that key structural proteins of stratified squamous epithelia, and proteins associated with both early- and late-stage keratinocyte terminal differentiation, were appropriately expressed and localized in hES-DK tissues relative to stratified squamous epithelia generated from epidermal keratinocytes.

hES-DK stratified squamous epithelia do not exhibit cutaneous barrier function

Human skin maintains tissue homeostasis by providing a barrier to transepidermal water loss. To evaluate the level of cutaneous barrier function of hES-DK and human epidermal keratinocyte tissues, skin surface electrical impedance was measured. Stratified squamous epithelial tissue generated from epidermal keratinocytes yielded impedance values similar to that of native skin indicating the presence of cutaneous barrier function (Figure 4). In contrast, hES-DK tissue did not form an epidermal permeability barrier. To ensure all tissues possessed contiguous epithelialization, which is critical for this assay, hES-DK tissues were incubated with the metabolic dye 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl tetrazolium bromide [25] which confirmed complete culture surface coverage (data not shown). Since the barrier to transepidermal water loss results primarily from...
lipid deposition in the intercellular spaces between the enucleated squames of the stratum corneum, these findings were consistent with the histological observation noting the lack of a well-defined cornified layer in hES-DK tissue.

**Stratified squamous epithelia generated from hES-DK cells express proteins associated with innate immunity and demonstrate antimicrobial activity in vitro**

HDPs are integral to the innate immunity of epithelial tissues (reviewed in [26]), including oral and epidermal stratified squamous epithelia, and function as a barrier against entry of infectious agents in stratified human tissues [27-29]. In human skin, both human β defensin 3 (hBD3) and human cathelicidin (LL37) are typically stored in lamellar bodies within differentiated keratinocytes [30,31]. To determine if HDPs were expressed in hES-DK stratified squamous epithelia, the expression patterns of hBD3 and LL37 were examined. hBD3 protein was localized to the granular layer in hES-DK tissue (Figure 5A), consistent with the pattern observed in stratified squamous epithelia generated from epidermal keratinocytes (Figure 5B). Staining for LL37 protein was observed in all layers of hES-DK tissue (Figure 5C). Similarly, LL37 was distributed throughout tissues generated from human epidermal keratinocytes, however slightly more intense signal was noted in the basal and immediately suprabasal layers (Figure 5D).

The presence of HDPs in hES-DK tissue implied the potential for innate immune system functionality, therefore an in vitro assay was used to evaluate antimicrobial activity. In this assay bacterial levels were shown to significantly decrease in a dose-dependent manner upon treatment with increasing concentrations of purified LL37 peptide (Figure 5E). Samples treated with 0.4 μg/ml LL37 peptide resulted in a 43% inhibition in bacterial growth when compared to the level of bacterial growth found in untreated control samples, indicative of bacteriostatic activity. Similarly, media conditioned by hES-DK stratified squamous epithelia significantly inhibited bacterial growth by 39% compared to the bacterial growth control. Detection of antimicrobial activity confirmed formation of a biologically functional stratified squamous epithelia and represents the first description of this activity in a hES cell-derived three-dimensional tissue.
Discussion

To successfully restore damaged tissues, allogeneic stem cell therapies for the treatment of severe cutaneous injuries require the derived cell types to generate biologically functional keratinized stratified squamous epithelial tissue. hES- and hiPS-derived cells have been reported to generate stratified epithelia in vitro with varying levels of success [7,9,14-16]. To date, no functional bioactivities for stratified squamous epithelia generated using pluripotent cell-derived sources have been reported. Here we evaluated the capacity of hES-derived epithelial cell populations to form organized structure, thoroughly characterized the expression patterns of proteins essential for keratinized stratified squamous epithelia structure and function, and assessed key bioactivities typical of the interfollicular epidermis.

As first described by Metallo et al. [6], treatment of hES cells with RA to initiate epithelial differentiation and BMP to inhibit neural differentiation enhanced differentiation efficiency of cells displaying an epithelial-like morphology. Our studies confirm and expand upon the role of RA in the generation of hES-DK cells. We found that mRNA expression of epithelial differentiation markers (ΔNp63, K18) in RA-treated cells was upregulated in comparison to DMSO-treated or untreated hES cell controls, while neural ectoderm differentiation marker expression (nestin, Sox 1) remained unaffected or was attenuated. Expression of K14 mRNA was not detected in RA-treated cells. Our findings replicate those recently reported by Leydon et al. [16] where p63+, K18+, K14- simple epithelial cells were generated from hES cells using nearly identical methods. Although RA supported differentiation of a simple epithelial cell type, it appears insufficient to induce keratinocyte differentiation. We further found that RA-treated hES cells cultured under organotypic conditions formed tissue that lacked organized structure. This is in contrast to Leydon’s results where p63+, K18+, K14- RA-treated hES cells developed a uniform multilayered, non-keratinized squamous epithelium when cultivated on a vocal cord fibroblast-containing dermal matrix. Leydon and coworkers also demonstrated that these same cells failed to form three-dimensional structure when a non-cellularized collagen matrix was used, implying that stromal-derived paracrine signaling was inadequate to promote the formation of organized tissue from RA-treated hES cells.

RA-treated hES cells aggregates attached to the culture surface and supported the outgrowth of hES-DP cells exhibiting a cuboidal, epithelial morphology. Selection by subsequent passage of hES-DP cells onto gelatin-coated culture plates resulted in clonal growth of hES-DK cells which exhibited epidermal keratinocyte morphology and, unlike hES-DP cells, formed keratinized stratified squamous epithelia in organotypic culture. Gene expression analysis was completed at each step of differentiation and hES-DK tissue formation to assess changes in epidermal marker expression. A low level of ΔNp63 mRNA was detected at all stages of differentiation, consistent with its function in the maintenance of epithelial proliferation [32]. Similarly K18, commonly associated with simple epithelia, was expressed throughout hES-DK differentiation. Expression of K14, first evident in the hES-DP population, became more pronounced after serial cultivation and subsequent organotypic culturing. Although previous studies required prolonged exposure to differentiation agents [8], clonal isolation and expansion [9], or extended serial cultivation [6,15] to enrich for a K14+ keratinocyte-like population, we found that a single serial cultivation step onto gelatin-coated plates was sufficient to substantially enrich for cells expressing markers of a keratinocyte lineage. Integrin-mediated adhesion signaling plays critical roles in stem-cell-niche interactions [33], as well as keratinocyte proliferation and differentiation (reviewed in [34]). Since adhesion to gelatin during cultivation enhanced selection for cells with keratinocyte properties, it is possible that integrin-mediated signaling may also be critical for keratinocyte lineage differentiation. In addition to K14, expression of involucrin and loricrin, components of the keratinocyte cornified envelope, were detected in both hES-DK cells and stratified tissue. It is interesting to note that the succession of gene expression acquired as hES-derived populations adopted a keratinocyte phenotype paralleled the progression of epidermal marker expression found during fetal epidermal morphogenesis [35,36]. Importantly, we found that expression of key keratinocyte markers, most notably K14, was predictive of the cells’ ability to form a keratinized stratified squamous epithelium.

Within the interfollicular epidermis, proteins associated with cell-cell adhesion and keratinocyte terminal differentiation are expressed and localized to specific layers. Although examination of several of these proteins in hES cell-derived tissue constructs has previously been described [9,14,15], a comprehensive examination of proteins essential for the structure and function of keratinized stratified squamous epithelia has not yet been reported. Our study provides the first examination of the cell-cell adhesion proteins E-cadherin, P-cadherin, and β-catenin in three-dimensional hES-derived epithelia. E-cadherin and β-catenin staining was cell membrane-associated, localized to the basal and immediately suprabasal layers, and more pronounced in hES-DK tissue relative to epidermal keratinocyte-derived tissue. Upregulation of these proteins could inhibit keratinocyte terminal differentiation [37], contributing to the morphological differences noted between hES-DK and human epidermal keratinocyte-generated tissues. Although substantially reduced relative to tissues generated using epidermal keratinocytes, P-cadherin staining in hES-DK tissue was cell membrane-associated and localized to the basal and immediately suprabasal layers. This pattern is characteristic of the epidermis, but not of non-keratinized stratified squamous mucosal epithelia where P-cadherin expression is found throughout the suprabasal layers [38,39]. Keratin expression patterns not only change as terminal differentiation proceeds, but reflect tissue specificity. Although K14 is found in many epithelial tissues, K1 is differentiation-specific and primarily associated with keratinized stratified squamous epithelia [40]. Both K14 and K1 expression was observed suggesting morphogenesis of a keratinized tissue. In Dabelsteen et al. [14] and Metallo et al. [15], expression of keratin 10 (K10), the partner of K1, was absent or displayed a localization pattern that differed considerably from that of epidermal keratinocyte-generated tissues. In our study, K1 expression was appropriately restricted to the immediately suprabasal layers, similar to the K10 expression pattern observed by Guenou and coworkers [9]. We found that the epidermal-specific keratin K2, which has not previously been examined in hES-derived stratified tissues, K2, typically restricted to the upper spinous and granular layers of the epidermis and not expressed in other keratinized stratified squamous epithelial tissues, is expressed to varying levels in different regions of the body during fetal skin development [41]. The level and variability of K2 protein expression observed in hES-DK tissue are consistent with this pattern. p63 expression in hES-DK tissue was restricted to the basal layer, unlike the findings of Dabelsteen et al. or Metallo et al. [14,15] where p63 was dispersed throughout the tissue as is commonly
found in neoplastic epithelia [42,43]. Expression of transglutaminase 1, which functions to crosslink proteins into the cornified envelope during keratinocyte terminal differentiation, was also found to be appropriately localized. Filaggrin, which is integral to the epidermal permeability barrier [44], was properly restricted to the granular layer in hES-DK tissue. This supports the earlier finding of Guenou and coworkers [9] but stands in contrast to Metallo et al. [15] where filaggrin in hES-derived epithelial tissues was virtually absent. The presence and appropriate localization of proteins essential for the structure and function of keratinized stratified squamous epithelia, and specifically the epidermis, confirms morphogenesis of interfollicular epidermal tissue from hES-derived keratinocytes.

In vivo the epidermis serves as a barrier to transepidermal water loss, maintaining fluid homeostasis via cutaneous barrier function, and as a barrier to infectious agents, producing HDPs integral to innate immunity. Evaluation of these biological activities in epithelial tissue generated from a pluripotent stem cell source has not previously been reported. Cutaneous barrier function is generated by deposition of lipids in the intercellular spaces between cells of the cornified envelope in the stratum corneum [45-47]. Unlike native human skin or tissue formed from human epidermal keratinocytes, we found that hES-DK tissues did not possess a functional permeability barrier most likely resulting from the lack of a well-defined stratum corneum. It is possible that extending the period of air-interface cultivation would promote further accumulation of enucleated squames, establish a more substantial stratum corneum, and enhance the development of cutaneous barrier function. hBD3 protein, a HDP associated with the more terminally-differentiated layers of the epidermis [28], was found to localize to the granular layer in both hES-DK and human epidermal keratinocyte-generated tissue. Protein expression of the cathelicidin LL37 was primarily restricted to the basal and immediately supra-basal layers in tissues generated from human epidermal keratinocytes. In contrast, intense staining for LL37 was detected throughout hES-DK tissue. The reason for this difference is not currently known. The presence of HDPs implied the potential for innate immune system functionality in hES-DK stratified squamous epithelia, therefore antimicrobial activity was evaluated. Bacteriostatic activity was confirmed for hES-DK tissue, which inhibited bacterial growth by 39%. This level of antimicrobial activity. Such allogeneic cell-based therapies offer a means to restore appropriately localized. Filaggrin, which is integral to the epidermal permeability barrier [44], was properly restricted to the granular layer in hES-DK tissue. This supports the earlier finding of Guenou and coworkers [9] but stands in contrast to Metallo et al. [15] where filaggrin in hES-derived epithelial tissues was virtually absent. The presence and appropriate localization of proteins essential for the structure and function of keratinized stratified squamous epithelia, and specifically the epidermis, confirms morphogenesis of interfollicular epidermal tissue from hES-derived keratinocytes.

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