Development of a tissue-engineered skin substitute on a base of human amniotic membrane

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Abstract
Allogenic graft material and tissue engineering have recently shown promising results for the improvement of both esthetic and functional outcomes in the treatment of large skin defects. We chose human amniotic membrane as a cellular scaffold in order to develop a skin substitute for later in vivo uses. Various methods of de-epithelialization of the human amniotic membrane were evaluated by histological analysis including hematoxylin–eosin and laminin staining, optic coherence tomography, and scanning electron microscopy with 0.25/0.02% trypsin/ethylenediaminetetraacetic acid treatment and mechanical cell removal showing an almost complete loss of the epithelium and a mainly intact basement membrane. Novel examination of human amniotic membrane by optic coherence tomography was feasible, but difficulties were experienced in handling and interpretation of the tissue as no comparable data exist. Subsequently, we developed an air–liquid interface cell culture to cultivate keratinocytes and fibroblasts on the de-epithelialized human amniotic membrane. We achieved a mostly keratinized surface on the epidermal side with a confluent fibroblast network on the chorion side.

Keywords
Tissue engineering, skin graft, basement membrane, human amniotic membrane, de-epithelialization, air–liquid cell culture, optical coherence tomography, electron microscopy, immunohistochemistry

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Introduction
The skin acts as an important barrier against noxious agents and helps to maintain a stable water balance.1 Various pathologies, such as burn injuries, tumor resections, and chronic wounds, are often responsible for large skin defects that need to be covered properly and in a timely manner.2–5

Diverse autologous and allogenous grafts are currently in use, although limitations include restricted availability, secondary defects, rejection of the graft, and functional and esthetic problems of the resulting scar.6–9 A promising scaffold for tissue-engineered skin without the limitations of the other types of graft is the human amniotic membrane (hAM). It provides a stable basement membrane for cell culture, expresses anti-immunogenic and anti-inflammatory agents, and shows good results in the treatment of wound defects as a wound coverage.10–12

As for all biological scaffolds, gentle but complete decellularization is a critical step in removing allogenous cells, and various methods have been described.13 In view of the importance of the basement membrane in the process of re-epithelialization and the organization of the dermis, our aim has been primarily to maintain the basement membrane through the decellularization process.14–16 As laminin is an essential
component of the basement membrane, it was used to verify the integrity of the membrane.

As typical skin shows an orthokeratinized surface, air–liquid interface cultures have been suggested in order to allow air to contact the surface as the typical stimulus for keratinocytes to differentiate into corneocytes, while simultaneously nutrients are provided from the dermal site of the graft. Choice of the right culture medium is critical to secure the growth of both the fibroblasts on the dermal side and the keratinocytes on the epidermal side. Although serum-containing media are known to be crucial for the cultivation of fibroblasts, it is also described to inhibit the growth of keratinocytes. Therefore, we have compared various combinations of serum and keratinocyte medium in a simple trypan blue viability test and by phase contrast microscopy in order to find a suitable air–liquid medium.

Overall, the aim of this study was to generate a skin substitute on base of hAM for further in vivo comparisons. We suggested that detergents would decellularize the hAM better than enzymes with regard to an intact extracellular matrix and a sufficient cell removal. We also compared the resulting hAM for its re-epithelialization properties. In addition, a feasibility part was conducted, where images of the hAM by optical coherence tomography should be achieved.

Materials and methods

Several methods of de-epithelialization of the hAM were compared, and the tissue was subsequently examined (Figure 1). In addition, various cell-culture media were evaluated to ensure fibroblast and keratinocyte viability on the skin graft. De-epithelialized hAM was then cultured with keratinocytes and fibroblasts and shifted to an air–liquid interface cell culture followed by tissue studies.

General preparation of hAM and quantitative evaluation

Cryoconserved hAMs (Austrian Cluster for Tissue Regeneration, Linz) were gently thawed at room temperature and rinsed thoroughly in DPBS (Gibco, Waltham). Once the tissue had been cut into smaller pieces of approximately $5 \times 5$ cm, they were transferred to the various de-epithelialization methods (Figure 2(a)). After decellularization, the tissue was further rinsed in DPBS+ (Gibco, Waltham) and aprotinin (10 KIU/mL) (Sigma-Aldrich, St. Louis) on a shaker (IKA, Staufen) at 4°C for 48h, followed by a regular medium change to remove loose epithelial cells and enzyme residues.

The decellularization results were quantified by simple cell counting using Image J of randomly chosen decellularized areas ($n=20$ for each method) of H/E stained light microscopy images (500 µm of straight membrane each). Cells were counted above and below the basement membrane and separately documented; results were presented with mean value ± standard deviation.

![Figure 1. Experimental overview: comparison of different de-epithelialization methods and development of an air–liquid interface culture with subsequent tissue studies.](image-url)
De-epithelialization with trypsin and cell scraper

One method that we evaluated was the use of trypsin and a cell scraper.\textsuperscript{21} The thawed hAM was transferred into 0.25/0.02% trypsin/ethylenediaminetetraacetic acid (EDTA) (Sigma-Aldrich) at 37°C and incubated for 25 min. Cold DBPS\textsuperscript{+} was used to stop the enzyme activity. Under regular phase contrast microscopy control, the epithelium was removed by slow and gentle scraping with a cell scraper (Sarsted, Nümbrecht) (Figure 2(b2)). Following this step, the hAM was placed on an insert (Figure 2(c)) in a cell-culture well (Greiner Bio-One, Kremsmünster), fixed with a PTFE ring (Alwin Höfert, Ammersbek) with the epithelial side pointing up, and rinsed on a shaker as described above (Figure 2(e)).

Decellularization with sodium dodecylsulfate/sodium deoxycholate

Following an experiment in which heart valves were decellularized,\textsuperscript{22} the thawed hAM was placed on an insert as mentioned above. It was then washed with a 0.5% sodium dodecylsulfate (SDS)/sodium deoxycholate (SDC) solution (Sigma-Aldrich) on a shaker (200/min) at room temperature for 48 h (Figure 2(d) and (e)). The solution was changed every 12 h. Subsequently, the tissue was rinsed on a shaker as mentioned above.

De-epithelialization with NaOH

One research group postulated an easy de-epithelialization method involving the use of NaOH and a cotton wool tip.\textsuperscript{23} A cotton stick soaked in 0.5 M NaOH solution (Roth, Karlsruhe) was gently rubbed over a thawed hAM for over a minute to remove the epithelium (Figure 2(b1)). Subsequently, the hAM was rinsed thoroughly in DPBS\textsuperscript{+}, placed on an insert, and rinsed on a shaker as described above.

Decellularization with Triton/SDC

Although similar to the decellularization of the heart valves with SDS/SDC, another composition of the washing solution has been suggested, namely the use of 0.25% Triton/SDC (Roth, Karlsruhe).\textsuperscript{24} The thawed hAM was fixed onto an insert and washed for approximately 48 h on
a shaker (200/min) at room temperature with a medium change every 12 h. The hAM was then rinsed on the shaker as mentioned above.

Cell isolation and cultures
Excess rat skin was used to isolate keratinocytes and fibroblasts, which subsequently were amplified and cultured on the de-epithelialized hAM until a confluent cell population was reached. The culture was then changed to an air-liquid interface culture until final keratinization.

Isolation of keratinocytes. The isolation of fibroblasts and keratinocytes was performed following the protocol of the isolation of various cell lines from corneal tissue.25 Rat skin was stored in keratinocyte medium (Cellsystems, Troisdorf) for up to 4 h after extraction. After removal of the subcutis, the tissue was then cut into pieces of 5 mm which were then incubated in Thermolysin (50 IU/mL) (Sigma-Aldrich) for 15 h at 4°C. The enzyme activity was subsequently stopped with keratinocyte medium. Using sterile forceps (Aeskulap, Tuttlingen), the epidermal layer was then removed and cut into smaller pieces (Figure 3(a)). These samples were transferred into centrifuge tubes (Greiner Bio-One) filled with TrypLE (Gibco, Waltham) and dissociated by occasional gentle shaking for 30 min at 37°C. Subsequently, the dissociation was stopped with cold keratinocyte medium, and the cell suspension was filtered through a cell strainer (Corning, Corning) and centrifuged. The remaining cell pellet was solved in keratinocyte medium and transferred to cell-culture flasks (Greiner Bio-One) (20 × 10⁴/cm²).

Isolation of fibroblasts. After removal of the epidermis, the remaining pieces of dermis were transferred to fibroblast medium (Gibco, Waltham) and cut into smaller pieces, which were then placed on a cell-culture dish.
(Greiner Bio-One) (Figure 3(b)). After 4 h, the dish was filled with fibroblast medium with the pieces staying adherent to the dish. After several days with regular medium changes, the pieces were removed when fibroblast colonies were visible.

Medium changes and cell passages. To amplify the cells, regular medium changes and cell passages had to be performed. The medium was changed every second or third day for keratinocytes or fibroblasts, respectively. For a medium change, the remaining medium was removed, the flask/dish was rinsed with DPBS, and new medium was administered. To passage the cells, a small amount of TrypLE was used for 15–40 min at 37°C. Dissociation was observed under phase contrast microscopy. When almost all the cells had dissociated, the enzyme was blocked with cold keratinocyte or fibroblast medium. The solution was then centrifuged, and the cell pellet was dispersed in keratinocyte or fibroblast medium and recultured in cell-culture flasks (keratinocytes 10,000/cm²; fibroblasts 4000/cm²).

Comparison of various air–liquid media

In the air–liquid model, only one medium was present on the basolateral side of the insert for both keratinocytes and fibroblasts. Cell cultures of newly passaged fibroblasts and keratinocytes were incubated for several days in fibroblast medium with 10% fetal calf serum (FCS) (Gibco, Waltham) (n = 5), in keratinocyte medium (n = 5), or in keratinocyte medium with 5% FCS (n = 5) or 10% FCS (n = 5). Morphology and growth characteristics were regularly controlled by phase contrast microscopy, and a Trypan Blue exclusion test was performed and the viability documented.

Cell culturing on hAM and by air–liquid interface culture

We developed an air–liquid interface culture based on another published model. De-epithelialized hAM was placed upside down with the chorion side facing up. Then, 1 mL fibroblast medium with approximately 100,000 fibroblasts were pipetted on this side using the surface tension to avoid run-off (Figure 3(c)). After an incubation time of 6 h, the insert was transferred to a cell-culture well with the epithelial side facing up, and 1.5 mL fibroblast medium was pipetted onto the basolateral side (Figure 3(d)). Subsequently, 0.5 mL keratinocyte medium with 500,000 keratinocytes was placed on the epithelial side of the hAM. Medium changes were performed daily. When confluence of both cell lines was achieved, the cultured tissue was lifted to enable air–liquid interface culture with only keratinocyte medium with 5% FCS being present on the basolateral side of the insert. Medium was changed regularly until final keratinization was accomplished.

Similar to protocols used before, using phase contrast microscopy, the re-cellularization for each decellularized hAM (n = 20) of both sides was then semi-quantitatively scored as 0 = absent, 1 = focal (<50%), 2 = non-confluent (>50%), 3 = confluent layer of cells. Regarding the keratinization, the following additional score was obtained: A = no keratinization, B = keratinization < 50% of re-cellularized areas, C = keratinization > 50% of re-cellularized areas, D = complete keratinization.

Tissue examination

Hematoxylin and eosin staining, immunohistochemistry. The tissue was fixed in formalin, embedded in paraffin (over 48 h), and subsequently cut into 5-µm-thick sections. Hematoxylin–eosin staining was performed using an established protocol. For immunohistochemical analysis, 5-µm-thick sections mounted on slides were deparaffinized using xylene and ethanol. After treatment of the slides with a peroxide and a protein block, they were incubated with primary antibody (polyclonal rabbit antibody against laminin) for 30 min and subsequently with secondary antibody (anti-rabbit antibody) for 45 min. The sections were then treated with streptavidin and counter-stained with hematoxylin.

Scanning electron microscopy. The tissue was rinsed in Sörensen buffer, fixed in 2.5% glutaraldehyde (Sigma-Aldrich), rinsed in water, and dried with ethanol and hexamethyldisilazane. It was then sputter-coated and transferred to a scanning electron microscope (Zeiss, Oberkochen).

Optical coherence tomography. As a pilot project, hAM was evaluated using optical coherence tomography (OCT), a non-invasive method that allows images to be taken at the cellular level with the depth of the tissue (millimeters). The hAM, which was fixed in the insert, was simply positioned vertically in front of the OCT apparatus (Heidelberg Engineering, Heidelberg).

Results

De-epithelialization

The membranes treated with trypsin showed good results under phase contrast and light microscopy (Figures 4(b) and 5(a1) and (c1)). The epithelium was almost completely removed with just 3.4 ± 3.3 cells remaining (Figure 6), and the basement membrane was consistent and prominent. The mesenchymal connective tissue showed some loosening compared with the native hAM, while decellularization below the basement membrane was not complete with 16.8 ± 6.3 cells remaining (Figure 6). In addition, some shaded lines were apparent by phase contrast microscopy, although the related layer was difficult to allocate. The
lines might represent the effects of the mechanical impact or be a normal aspect of the basement membrane of the hAM. The hAM treated with NaOH presented incomplete de-epithelialization with $9.4 \pm 4.1$ cells above the basement membrane (Figures 4(d), 5(a3) and (c3), and 6). Furthermore, a loosened mesenchymal connective tissue and a largely consistent but blurred basement membrane were observed following this treatment with no removal of cells below the basement membrane (Figure 6). The two surfactant-containing decellularization solutions also gave incomplete de-epithelialization, but better decellularization below the basement membrane compared to the other methods with $10.8 \pm 4.3$ remaining cells for SDS/SDC and $7.2 \pm 3.2$ for Triton/SDC (Figures 4(c) and (e), 5(a2), (a4), (c2) and (c4), and 6). Although the epithelial cell boundaries were no longer visible under phase contrast optics, the epithelium still did not provide a clear view of the mesenchymal connective tissue. By light microscopy, the basement membrane was largely lost and a loosened mesenchymal connective tissue was presented.

**Colonization of hAM—air–liquid culture**

Fibroblast growth was completely inhibited when incubated with keratinocyte medium DermaLife K (Figure 7(a1)). When FCS was added, the fibroblasts showed pronounced...
vital cell growth with 10% FCS showing slightly better results than 5% FCS (Figure 7(a2) and (a3)). As expected, the best growth conditions were provided by Dulbecco’s Modified Eagle’s Medium (DMEM) with 10% FCS (Figure 7(a4)). The keratinocyte growth under keratinocyte medium showed a typical hexagonal vital appearance of the cell culture (Figure 7(b4)). Almost the same aspect was observed when 5% FCS was added (Figure 7(b3)). With 10% FCS and DMEM, the cells showed typical signs of cell damage, such as a change in shape and a numeral decrease (Figure 7(b1) and (b2)). The trypan blue exclusion test confirmed these findings. Whereas the three groups growing in culture

\[ \text{Figure 5. De-epithelialized hAM stained with HE and for laminin. Light microscopy. 20–40×. (a1–a4) HE staining of de-epithelialized hAM. Whereas the hAM treated with trypsin and NaOH shows mostly removed epithelium, the hAM treated with either Triton/SDC or SDS/SDC presents a consistent epithelium. (b1–b2) HE and laminin staining of native hAM, which presents a typical pseudostratified columnar epithelium and a consistent basement membrane. (c1–c4) Laminin staining of de-epithelialized hAM. The hAM treated with trypsin and NaOH present a consistent, strongly stained, basement membrane, whereas the other two groups exhibit no obvious basement membrane.} \]
media with added FCS showed a cell viability $>85\%$, the only group without FCS had a viability of only $0\%$–$5\%$. The opposite effects were presented by the keratinocyte cultures. The viability of the cultures incubated without FCS or with 5% FCS had a viability of 90%–95%. When more FCS was added, the viability decreased to almost 0%. As the keratinocytes and fibroblasts had to be supplied with the same medium for several days while being in the air–liquid culture, the keratinocyte medium with 5% FCS additive, which showed the best results, was therefore used for the air–liquid interface culture.

The colonizing of the hAM started with the fibroblasts. The formation of the differentiated cell networks on the chorion side occurred acceptably, no matter which de-epithelialization method was used (Table 1). By contrast, the growth of the keratinocytes on the epithelial side differed greatly between the methods used (Table 1). The culture was harvested 2 days after relocation and at the end of the air–liquid culture. The hAM treated with trypsin presented a dense and confluent epithelial layer after 2 days (Figure 8(a1)). At the end of the air–liquid culture, some keratinization at the chorion side (Figure 9(a1)) similar to the findings presented by other authors. The trypsinized hAM exhibited the complete removal of the epithelium with only some detritus remaining and a smooth appearance as an indication of an intact basement membrane (Figure 9(b)). On the stromal side of the membrane, some fissures could be found (Figure 9(c2)) but, otherwise, no different morphology compared with the native hAM. The hAM analyzed at the end of the air–liquid culture showed partially single-layered, partially multilayered, or pseudostratified epithelium (Figure 9(d1) and (d2)). Similar to the aspect under phase contrast optics, a large variation could be seen between raised epithelial cell complexes and areas not covered by any epithelial cells. A confluent fibroblast network on the chorion side also confirmed the findings of the analysis carried out by phase contrast microscopy (Figure 9(c3)).

Several native, trypsinized, and recolonized hAM were analyzed within a pilot project to test whether the OCT was suitable for visualizing the hAM. In particular, the positioning of the hAM in front of the OCT was difficult and only possible under non-sterile conditions. We were however able to prepare an overview image and a thin-layered series of images (Figure 10(a)–(c)). Whereas the native (Figure 10(a)) and recolonized hAM (Figure 10(c)) were significantly thicker, mostly by $>1\,$mm, the de-epithelialized hAM (Figure 10(b)) was comparatively thin. Exact measurements were not obtained because of possible artifacts from light refraction. The epithelial layers seemed to be well definable with respect to the underlying matrix. Furthermore, the epithelia of the different hAM showed visible differences in thickness with the recolonized hAM seeming to have the thickest epithelium. At various areas of the membranes, we were able to recognize defects in the membrane that were not detectable by phase contrast microscopy.

Various de-epithelialization methods were compared in this study. Trypsin showed the best results with regard to the removal of the epithelium and the integrity of the basement membrane. We also obtained good results with NaOH, which until now is not as established as

![Bar graph of the decellularization success above and below the basement membrane (BM).](image-url)
**Figure 7.** Comparison of various culture media on keratinocytes and fibroblasts. Phase contrast microscopy, 100×. Left: Top down (a1–a4), fibroblast cultures are displayed after 5 days incubation in different culture media. The viability increases with the use of FCS. Right: Top (b1–b4) down, keratinocyte cultures are visible after 5 days incubation in media of different compositions. The smaller the amount of FCS added, the higher the viability.
trypsinization and should therefore be further evaluated. The SDS/SDC and Triton/SDC protocols were used because of their effective decellularization of heart valves. Contrary to our expectations that the amniotic epithelium, which has to resist mechanical strains in the uterus, would act similarly to the endothelium of the equally mechanically stressed heart valves, the two surfactant-containing decellularization methods seemed to be insufficient to remove the epithelium in contrast to other authors who have presented better results using such methods.

Often, more rigorous decellularization is accompanied with more damage to the extracellular matrix. Trypsin is well known for its strong collagenase effect, which might however lead to mechanical instability of the stroma. Light microscopy (LM) and Scanning electron microscopy (SEM) revealed some tears in membranes treated with trypsin; this indicates that trypsin causes tissue damage. Moreover, the further handling of the de-epithelialized hAM was difficult because some membranes, often when fixed on the insert, showed macroscopic lesions and had to be discarded. Regarding the non-epithelial side, there were differences in the decellularization properties of the different methods with the detergents being more efficient. However, when observing the re-cultivation, there were no differences with complete confluent fibroblast networks regardless of the previous decellularization.

Alternative methods have been published in which only the epithelial side comes into contact with the enzymatic agent. These should be further evaluated with regard to possible tissue damage and to the effect on re-cultivation of the non-epithelial side. In addition, considering the wide variance of the amniotic tissue used, a continuous phase contrast microscopy control of the de-epithelialization should be performed to minimize the exposure time to the enzyme. Overall, despite there being many different de-epithelialization methods for amniotic membranes, no ideal method has been established as yet; similar conclusions have been expressed by other authors.

The isolation and cultivation of the keratinocytes and fibroblasts was largely problem-free. Moreover, primary cultivation with trypsin-de-epithelialized hAM showed confluent results. With regard to the composition of the air–liquid medium, a keratinocyte-based medium with the addition of 5%–10% serum provided both good fibroblast and good keratinocyte viability. Noticeably, the confluence of the recultivated hAM was partially lost with a few areas without any epithelium and many areas with significant multilayered epithelium and keratinization. Other research groups cultivating hAM with non-keratinizing epithelium have not described such findings. As we did not perform cell viability tests on the hAM, the reason for these findings remains uncertain: Possible explanations are that the remaining trypsin molecules might have caused local cytotoxicity or heterogeneous membrane conditions that might have inhibited nutrition from the chorion side while the tissue was under air–liquid conditions. To date, no investigations have been published regarding the filter characteristics of de-epithelialized hAM, and therefore, it remains uncertain where the medium might have penetrated the membrane sufficiently to achieve confluent keratinization. Other authors have mentioned such perfusion as the most important aspect for obtaining confluent and differentiated keratinocytes.

Further testing, such as viability essays with the hAM after de-epithelialization should provide further data regarding this issue. Moreover, more air–liquid media should be evaluated to enhance further the keratinocyte viability. Some serum-free synthetic media for fibroblasts are the subject of ongoing research. They might, with the addition of specific growth factors for keratinocytes, be adapted to the requirement of air–liquid culture. Overall, the re-epithelialized hAM can still be improved regarding gap-less epithelialization. However, they should already be tested in vivo as skin substitutes to look for possible advantages regarding wound healing to justify further research efforts.

The OCT examination was considered as a pilot project. Whereas the presentation of the hAM was feasible, we found it difficult to adjust the hAM in the sterile cell-culture well in front of the OCT apparatus. Furthermore, refraction at the various surfaces complicated the location of the membrane structures. In addition, no comparable images have previously been published for any comparisons to be made. Nonetheless, the advantages of the OCT include the non-invasive fast characterization of the hAM with regard to the thickness of the membrane and the identification of lesions not visible by phase contrast microscopy, both important features for a mechanically stable membrane. In the future,
Figure 8. Colonized epithelial side of de-epithelialized hAM. Phase contrast microscopy. 200×. (a1–a4) Re-epithelialized hAM processed by various de-epithelialization methods after 2 days. After 2 days, only the hAM treated with trypsin showed a confluent gap-less epithelium. Membranes prepared with NaOH and Triton/SDC showed no differentiated epithelial cell complex, and the cells remained round. The hAM treated with SDS/SDC showed no difference from the state after de-epithelialization. (b1–b4) The same membranes at the end of air–liquid culture. The hAM treated with trypsin and NaOH showed incomplete multilayered epithelialization with signs of keratinization. In the membranes washed with Triton/SDC, the epithelium was much more incomplete, whereas the hAM treated with SDS/SDC presented almost no adherent keratinocytes.
Figure 9. Various hAM preparations. Scanning electron microscopy. (a1) (1000×) Epithelial side of a native hAM in plain shot. (a2) (5000×) Epithelial side of a native hAM in inclined shot. The hAM showed the typical hexagonal architecture of the epithelium. Marked with red: exemplary cell boundaries. (b) (100×) Epithelial side of a de-epithelialized hAM in plain shot. The
a standardized OCT examination might be employed to exclude unsuitable membranes for further experimental and clinical use in order to improve the results obtained. Such standardization should be approached in future studies.

When using hAM, its variable inter-individual conditions should be considered: Many reasons, such as age, general condition, gestational age, and the sex of the fetus, play significant roles in the expression of cytokines, hyaluronic acid, and other factors. Furthermore, intra-individual factors such as the sampling location are relevant: hAM from the cervical zone is often thinner with less epithelium than hAM sampled from other zones, and the expression of cytokines also varies. In addition, the alignment of the collagen fibers of the hAM is irregular with occasional weak spots. For logistical reasons, we used cryoconserved hAM. Other authors have found the low expression of angiogenin, IL-6, and CP-1 in such processed membranes compared with fresh membranes, whereas still others could find no such differences. As the conservation of the hAM has possible harmful effects, it should be avoided in further studies. These limitations must be considered when using biological membranes, and findings have to be interpreted with great caution.

Conclusion

Treatment with trypsin and a cell scraper showed the best de-epithelialization results in hAM but were accompanied with detectable stromal strains. Nevertheless, with the use of serum-containing keratinocyte-based growth medium, the culturing of keratinocytes and fibroblasts on the de-epithelialized hAM in an air–liquid culture was possible and resulted in a mostly keratinized surface. Whereas tissue-engineered skin grafts on a base of hAM can still be improved with regard to gap-less epithelialization, they should be evaluated in in vivo experiments to justify further research.

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