Aerosol vehicle for delivery of epidermal cells – An in vitro study

Andre Bahoric DVM MSc, A Robertson Harrop MD MSc FRCSC, Howard M Clarke MD PhD FRCSC FACS, Ronald M Zuker MD FRCSC FACS
Division of Plastic Surgery and Cryopreservation Laboratory, The Hospital for Sick Children, Toronto, Ontario

Early coverage of large burn wounds is often limited by the availability of donor sites for skin grafts. One solution to this problem is the use of cultured epithelial autografts, whereby a small sample of uninjured skin is obtained from the patient and its keratinocytes are grown in a cell culture environment. Over a three to four week period the number of cells can be expanded more than 5000-fold, yielding enough cells to cover virtually the entire body surface area of an adult patient (1-3). Most commonly, the cultured cells are then attached to a backing of fine mesh gauze and placed onto the debrided wound (4-7). The process of applying the cells onto the gauze, and then onto the patient, is time-consuming and labour intensive. In addition, a large container is required to house the sheets of gauze while the cells are transferred from the laboratory to the clinical facility where the cells will be applied to the patient. It is speculated that a simpler process would involve suspending keratinocytes in a solution that could be sprayed as an aerosol onto the wound.

This in vitro study investigated the possibility of delivering epidermal cells to a cell culture plate via an aerosol vehicle. A simple and inexpensive aerosolization apparatus was used to deliver a suspension of viable epidermal cells to a culture plate. The aerosolization process was successful in consistently delivering a uniform distribution of suspended epidermal cells. By day 4 there was evidence of cell proliferation, and by days 7 to 9 a confluent layer of cells was achieved on the plates. The monolayer consisted primarily of keratinocytes interspersed with a few fibroblasts. The aerosol method was shown to be effective at delivering a suspension of viable epidermal cells to a culture plate.

Key Words: Aerosol, Delivery, Epidermal cells, Keratinocytes
MATERIALS AND METHODS

Epidermal cells used in this experiment were obtained during setback otoplasty procedures, from the full-thickness skin on the medial surface of the ear – skin that is normally discarded. Full-thickness skin samples measuring approximately 1x2 cm were obtained in this way from seven children (ages 5 to 14 years) undergoing surgery for prominent ears at The Hospital for Sick Children, Toronto, Ontario.

Full-thickness skin samples from each patient were placed in chilled RPMI 1640 transport medium (JRH Biosciences, Kansas), supplemented with polymixin B (100 µg/mL), vancomycin (50 µg/mL), lincomycin (120 µg/mL), cefoxitin (240 µg/mL) and amphotericin B (10 µg/mL). The samples were stored at 4°C for up to 6 h. The samples were then treated with trypsin (0.25% in ethylenediamine-tetraacetic acid) and incubated for 90 mins at 37°C with periodic, gentle agitation. This process separates epidermal cells from one another in preparation for aerosolization. After trypsinization was complete, the cells were suspended in aerosolization medium (RPMI 1640 medium supplemented with 10% fetal calf serum [Cansera], penicillin [50 U/mL] and streptomycin [50 µg/mL]). A hemocytometer was used to determine the concentration of cells in the aerosol. Fifteen millilitres of suspended epidermal cells (1x10^6 cells/mL) were placed in 50 mL sterile vials (Corning, New York), which were then attached to a sterilized pump-action aerosol nozzle (commercially available, modified in our laboratory) (Figure 1). The suspension was agitated just before aerosolization, then sprayed (using the ‘coarse mist’ mode on the nozzle) onto 10 cm cell culture plates (Becton-Dickinson, New Jersey). The plates were then incubated at 37°C, 5% Carbon dioxide and 75% humidity. The medium was replaced after 24 h with complete keratinocyte medium (Gibco, New York), supplemented with penicillin (50 U/mL) and streptomycin (50 µg/mL) and then exchanged with fresh medium every two days. The plates were examined by phase contrast microscopy on a daily basis, and photographs were taken to document progress.

RESULTS

The aerosolization process was successful at delivering a uniform distribution of suspended epidermal cells from each of the seven patients to the culture plates (Figure 2). In each case, most of the suspension consisted of individual cells; however, some aggregates of cells were noted. After incubation for 24 h (day 1), some cells were attached to the surface of the culture plates (Figure 3). It was apparent that many of these attached cells were keratinocytes; however, fibroblasts were also present. By day 3 (Figure 4) there was evidence of considerable cell proliferation with increased density of keratinocytes and, to a lesser extent, fibroblasts. A further increase in cell density was observed on day 5 (Figure 5). A confluent layer of cells was achieved on the plates by days 7 to 9 (Figure 6): the monolayer consisted primarily of kera-
tinocytes interspersed with a few fibroblasts and the occasional dendritic cell (presumed to be Langerhan’s cells).

**DISCUSSION**

Burns covering more than 60% of the patient’s total body surface present a special challenge because donor skin is only available in small amounts relative to the size of the open wound requiring coverage. The traditional approach to treating these patients is to graft as much of the burned wound as possible, given the amount of donor skin available, and then to wait one to two weeks until the donor sites have reepithelialized, after which time the donor sites are re-harvested. This process is repeated until all wounds are closed.

One solution to the problem of limited donor sites, first described by Rheinwald and Green in 1975 (1), is the multiplication of epidermal cells in a cell culture environment. This process involves obtaining small samples of skin from uninjured donor areas. The dermis is separated from the epidermis with dispase, a protein-digesting enzyme. The remaining epidermis is treated with trypsin, an enzyme that dissociates epidermal cells (mostly keratinocytes) from one another. The keratinocytes are transferred to a cell culture environment, placed onto a bed of lethally irradiated fibroblasts within a culture vessel and grown in the presence of a medium containing epidermal growth factor (2). Starting with a 3 cm² split-thickness skin biopsy, the area of epithelium can be expanded more than 5000-fold within three to four weeks, yielding enough epithelial sheets to cover the body surface of an adult human (approximately 1.7 m²) (3).

In 1981, O’Connor et al (4) successfully performed grafts of cultured epithelial cells in human burn patients. The sheets of keratinocytes, expanded in culture by the Rheinwald and Green technique, were attached to a backing of fine mesh gauze and applied to the patients’ wounds. Since then there have been other reports of successful clinical use of cultured epithelial autografts in patients, adult and child, with burns covering up to 90% of the total body surface area (5-7). Cultured epithelial autografts have also been used in large cutaneous wounds other than burns. Examples include giant congenital pigmented nevi (8), epidermolysis bullosa (9) and even smaller wounds such as chronic leg ulcers (10,11).

Current clinical techniques for application of cultured epithelial autografts involve carefully transferring the confluent epithelial cells from the culture dish to a backing of fine mesh gauze. This technique is time-consuming and labor intensive. Also, a large container is required to house the sheets of gauze while the cells are transferred from the labo-

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**Figure 3** Epidermal cells on the culture plate after 24 h incubation (day 1). Many cells have adhered to the plate surface (10x magnification)

**Figure 4** After incubation for three days there is evidence of cell proliferation (10x magnification)

**Figure 5** After five days there is further increase in cell density. The cells with the cobble-stone pattern are keratinocytes while those with a spindle shape are fibroblasts (10x magnification)

**Figure 6** Confluent layer of cells is seen by day 8 (10x magnification)
ratory to the clinical facility where the cells will be applied to the patient. One proposed method, studied in pigs, to circumvent delivery problems involved the use of water-tight vinyl chambers placed over full-thickness wounds. Autologous keratinocyte suspensions were then placed inside these chambers and were found to adhere to the wound and proliferate on its surface (12-14).

The authors of this paper examined another solution, whereby a suspension of epidermal cells was delivered to the target (in this case a cell culture plate) in aerosol form. This method was successful in delivering a uniform distribution of cells to the plate. The epidermal cells were, in turn, observed to proliferate over the surface of the plate, forming a confluent monolayer within seven to nine days. The monolayer consisted chiefly of keratinocytes interspersed with a small number of fibroblasts. Fibroblasts were present because the epidermal cell suspension was prepared directly from full-thickness skin biopsies. In the culture setting, where the cells are first multiplied in a culture environment, gradual overgrowth of fibroblasts relative to keratinocytes may be a problem. For this reason the Rheinwald and Green technique involves culturing the epidermal cells in the presence of lethally irradiated fibroblasts, which suppress fibroblast multiplication (2).

It is believed that the aerosol vehicle may be useful in other laboratory and clinical settings. The authors are currently in the process of developing an in vivo model, in which a suspension of epidermal cells is delivered to a wound in an animal. This work may eventually lead to the development of a cost- and personnel-efficient way of providing cultured epithelial cell coverage in clinical settings such as extensive burns or other cutaneous injuries. Another potential application of this technique, particularly in the pediatric population, is in scald burns. In these cases, there is often a deep, partial-thickness injury that leaves behind a small portion of the dermis, from which the epidermis can gradually be regenerated from epithelial cells present in the remnants of glands and hair follicles. It was speculated that ‘spraying’ the wound with a layer of autologous epithelial cells may speed up the healing process in this situation.

**SUMMARY**

A method was developed for delivering a suspension of epidermal cells to a cell culture plate using an aerosol vehicle. It was possible to deliver consistently a uniform distribution of cells to the plate. Over a subsequent period of seven to nine days the cells proliferated on the plate to form a confluent monolayer, comprised primarily of keratinocytes. The authors are now in the process of applying the technique to an experimental animal model. It is hoped that the aerosol vehicle may eventually prove clinically useful, for delivery of keratinocytes to human wounds.

**REFERENCES**