Multiple Endpoint Analysis of the 3D-Reconstituted Corneal Epithelium after Treatment with Benzalkonium Chloride: Early Detection of Toxic Damage

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PURPOSE. To investigate the effects of benzalkonium chloride (BAK) on the human reconstituted corneal epithelial model (HCE) and to optimize the operating potential of this model in the field of ophthalmic toxicology.

METHODS. The HCEs were treated with 0.001% to 0.5% BAK for 24 hours followed or not by a 24-hour postincubation period. To complete the histologic analysis, the authors designed a new MTT procedure to assess cellular viability. Frozen sections were analyzed by using fluorescence confocal microscopy for the presence of TUNEL, activated caspase-3, Ki67, ICAM-1, HLA-DR, E-cadherin, and occludin. Occludin gene expression was also investigated by using quantitative RT-PCR.

RESULTS. The MTT test revealed a dose-dependent response of BAK with significant toxic effects for concentrations as low as 0.005%. Increasing BAK concentrations induced an increased number of apoptotic cells, found from the superficial to the deeper layers, with the activation of caspase-3 at 0.01% and 0.02% concentrations. The number of Ki67- and ICAM-1–positive cells increased with 0.01% BAK and with 0.001% to 0.01% BAK, respectively. BAK induced the dose-dependent disappearance of occludin in the superficial layers while increasing its gene expression up to the 0.02% BAK concentration.

CONCLUSIONS. Fluorescence techniques conjugated with confocal microscopy on 3D-reconstructed corneal epithelia were well suited for the investigation of toxicological markers such as cell junction alteration, apoptosis, cell activation, and proliferation and gave relevant results compared with the known unknowns.

CONCLUSIONS. The use of the chicken eye (ICE) assay was found to adequately predict the corneal epithelium of the human eye in morphology, histomorphologic findings in the superficial layers. This procedure may therefore underestimate any mild toxic effect occurring at the apical surface of the constructs. In the literature, systematic histologic analysis and multiple endpoint analysis (MEA) were proposed as practical responses to prevent the occurrence of false-negative MTT results. Although MEA is a valuable approach to determine the mechanisms of eye irritation, the MTT procedure should be optimized separately so as not to neglect the cellular events occurring in the superficial layers. We believe that a small change in the MTT procedure could lead to increased sensitivity, not only making this procedure better suited for risk prediction in case of accidental exposure to the eye, but also adapting it to the prediction of low to very low irritant potential, especially when products are used repeatedly over long periods. The implementation of these very sensitive tools to predict eye irritation is critical for ophthalmologists treating patients who may be exposed to low irritants.
is known to induce oxidative effects and caspase-dependent and -independent apoptosis counteracted by an autophagic process. Chronic use of preservative is responsible for apoptosis of conjunctival cells and conjunctival inflammation, which has demonstrated negative effects (e.g., on the efficacy of glaucoma surgery). Consequently, the use of a highly differentiated, three-dimensional epithelial system of human ocular origin is desirable for prescreening or investigating the effects of ophthalmic drugs. It frees the experimenter from interspecies differences and provides a better approach to ocular epithelial physiology than monolayer models and cells from other organs. It is also a useful alternative to animal testing, which is time consuming and often invasive and which may lack suitable sensitive tools for detecting subclinical reactions. The 3D system models not only the different effects of toxicity on specific cell types, but also the interaction between the cells and the spatial effects induced by the toxic source. Moreover, epithelial cultures at the air-liquid interface are easy to handle and facilitate in vivo-like product exposures. Some of them can inhibit the flow of ionic material such as Na-fluorescein across their surface, suggesting the presence of a functional epithelial barrier, and the 3D-HCE cells were shown to express cytokeratin-3 and to include hemidesmosomes in the basal layers. Different types of intercellular junctions have been identified in the corneal epithelium ex vivo. Among them, adherens junctions, comprising the E-cadherin protein, serve to anchor cells together. Also, the tight junctions, originally defined as zonula occludens (ZO) and comprising occludin, ZO-1, and other proteins, are thought to provide the hydrophobic barrier preventing the free passage of molecules between adjacent epithelial cells.

The objective of this study was to optimize the operating potential of this new epithelial model in the field of ophthalmic toxicology. Using a 0.001% to 0.1% range of BAK concentrations, we investigated in vitro the various parameters likely to be modified by the toxic substance. Since the 0.01% BAK concentration failed to induce any significant decrease in cellular viability using the classic MTT procedure and considering that this procedure could be optimized, we made modifications so as to assess, with better sensitivity, the effects of BAK on cellular viability. Then, using en face confocal microscopy on frozen sections and entire epithelia, we investigated the hydrophobic barrier properties of BAK on the cellular surface. We used confocal microscopic analysis to detect cellular changes in the epithelium and to compare the effects of BAK on different cell types. The results were expressed as a percentage of cell viability compared with the negative control. Experiments were conducted three times. The results are expressed as mean ± SD.

Histologic Analysis

The tissue constructs were fixed in 10% formalin and then embedded in paraffin. Micromotive sections (Histolide 2000; Reichert-Jung, Vienna, Austria) were subjected to H&E staining for histomorphologic analysis. Confocal Immunofluorescence Analyses of Frozen Sections and Entire Epithelia

The construct was transferred to a Petri dish containing 500 μL of PBS and cut into two pieces under the microscope. One piece of tissue was fixed in parafformaldehyde 4% for 15 minutes before immunofluorescence labeling of the tight junction protein occludin. The other piece was embedded in OCT embedding medium and frozen at −80°C. Vertical sections (10 μm thick) were cut with a cryotome (CM 3050s; Leica Microsystems AG, Wetzlar, Germany) and stored at −20°C until staining.

The sections were subjected to immunofluorescence with antibodies against E-cadherin, occludin, ICAM-1, HLA-DR, Ki67, and activated caspase-3 as follows. The samples were fixed with 4% paraformaldehyde (PFA) for 5 minutes and then permeabilized with 0.01%-diluted Triton X-100 (Sigma-Aldrich, St. Louis, MO) for 5 minutes, except for occludin, E-cadherin, and CD54 labeling of frozen sections. After they were rinsed with 1% BSA-PBS, different sets of primary antibodies were added to 1% BSA-PBS: the rabbit anti-occludin IgG1 (1:50; Santa Cruz Biotechnology, Santa Cruz, CA), the rabbit anti-activated caspase-3 (1:50; BD-PharMingen, San Diego, CA), and the mouse anti-human IgG1 against E-cadherin (1:50; DakoCytomation, Glostrup, Denmark).
After 1 hour of incubation, the sections were rinsed twice in 1% BSA-PBS and incubated again for 1 hour in the dark with Alexa488-conjugated goat anti-mouse IgG at a 1:500 dilution or Alexa488-conjugated goat anti-rabbit IgG at a 1:500 dilution (Invitrogen-Molecular Probes, Eugene, OR). A terminal deoxynucleotidyl transferase-mediated dUTP nick-end labeling (TUNEL) assay (Roche Diagnostics, Meylan, France) was used to detect apoptosis in the tissue layers. After three washes in PBS, the nuclei were labeled with propidium iodide or DAPI and mounted in an anti-fade medium (Vectorshield; Vector Laboratories, Burlingame, CA). During all the en face immunofluorescence analysis steps, the epithelial side was kept upward to avoid damage. The samples were analyzed under a laser confocal microscope (E800, PCM 2000; Nikon, Champigny-sur-Marne, France).

Quantification
CD-54-, Ki67-, activated caspase-3-, and TUNEL-positive cells were quantified manually, using a microscopic grid on images under ×400 magnification. The results were expressed as the mean cell number per millimeter of epithelial length (mm EL). Standard deviations are indicated.

Occludin mRNA Expression Study
BAK is known as a penetration enhancer. Because occludin is known as one of the most specific tight junction proteins, we focused on its gene expression in response to BAK treatment. The 0.05% BAK concentration was added in this set of experiments so as to better characterize the change in the occludin expression pattern found between the 0.02% and the 0.1% BAK concentrations. Also, the time of 6 hours was added for more comprehensive results.

After each time point of treatment, RNA was extracted from the 3D-HCE cells (RNAqueous; Applied Biosystems, [ABI] Monza, Italy), according to the manufacturer’s protocol. Briefly, the tissue was lysed with a high-concentration guanidinium salt-containing buffer to inactivate endogenous RNase and then homogenized. Total RNA was isolated on a spin column and then eluted in a low-ionic-strength solution.

The cDNAs were then synthesized from 2 μg RNA template in a 20-μL reaction (High-Capacity cDNA Reverse Transcription Kit; ABI). The master mix used for the reverse transcription contained 2 μL 10× RT buffer, 0.8 μL 25× dNTP mix, 2 μL 10× random primers, 1 μL RT (Multiscribe; ABI), 1 μL RNase inhibitor, and 3.2 μL nuclelease-free water. Ten microliters of RNA was added to the master mix and then subjected to reverse transcription in a thermal cycler (Prism 7500 Real Time PCR System; ABI). The cDNA was amplified with Taq polymerase (TagMan Universal PCR Master Mix; Invitrogen) and the following gene expression assays (TagMan; Invitrogen): the human GAPDH probe GAPDH Hs09999990_m1 (calibrator gene) and the human occludin probe OCLN Hs00170162_m1 (occludin gene; Taqman probe; Invitrogen). PCR amplifications were performed with 25 ng of cDNA in 25 μL of total volume. Briefly, we used 12.5 μL of 2× PCR master mix (TagMan Universal PCR Master Mix; Invitrogen), 1.25 μL of 20× gene expression assay, 6.25 μL of water, and 5 μL of cDNA. The PCR conditions were 95°C for 10 minutes (AmpliTaq Gold DNA Polymerase Activation; ABI) followed by 40 amplification cycles on a real-time PCR system (95°C for 15 seconds and 60°C for 1 minute; Prism 7500; ABI). For each sample, analyses were performed in triplicate. Expression rates were expressed as relative quantification data (RQ), with changes considered significant when RQ was decreased or increased twofold compared with the control.

Statistical Analysis
Statistical comparisons of control and treatment groups for MTT data and quantification of Ki67-, activated caspase-3-, and TUNEL-positive cells were performed by two-way analysis of variance (ANOVA), followed, if necessary, by multiple pair-wise comparisons with the Bonferroni adjustment (Statview V for Windows; SAS Institute, Cary, NC).

RESULTS
Histologic Analysis
The morphology of the control consisted of a layer of nonkeratinized, more or less flattened superficial cells, an intermediate cell layer, where cells displayed lateral cytoplasmic extension similar to the wing cells, and a basal layer presenting regular column cuboidal cells. After 72 hours of treatment with any concentration of BAK, the histologic analysis as well as the MTT test revealed drastic cell death in all cellular layers associated with low viability levels (data not shown). The histologic analysis of the 3D-HCE revealed a dose- and time-dependent effect of BAK, with a reduced epithelial thickness and the presence of an increasing number of necrotic cells from the apical to the basal layers with increasing BAK concentrations, compared to the control (Fig. 1). BAK at 0.01% induced modifications of the superficial layer continuity, nucleus condensation, and vacuole formation in the superficial epithelial layer after 24 hours of treatment. These signs were more pronounced after the 24-hour postincubation period. With 0.1%...
BAK, all cellular layers were affected as early as 24 hours, with most of the cells detaching from each other and undergoing cell death after the 24-hour postincubation period.

**Modified MTT Test**

BAK induced a dose-dependent significant decrease in cellular viability as assessed by the MTT test after 24 hours and 24/24 hours (Fig. 2). The 0.001% BAK concentration tended to reduce cell viability at 86.6% and 82.6% at 24 hours and 24/24 hours, respectively. The 0.005% to 0.1% range of concentrations showed significant cytotoxic effects on 3D-HCE cells (P < 0.0001 compared to control), with 0.005% BAK, 0.01% BAK, and 0.02% BAK inducing, respectively, 72.8%, 61.7%, and 27.3% viability at 24 hours and 59.8%, 60.9%, and 15.3% viability at 24/24 hours. The damage tended to be greater after the 24-hour postincubation period, and all 3D-HCE cells were found to be dead with 0.1% BAK. When only two values were used for each treatment group, the MTT test discriminated between the toxicity levels of 0.001% BAK versus 0.01% BAK, 0.005% BAK versus 0.01% BAK, and 0.01% BAK versus 0.02% BAK.

**Immunofluorescence Analysis and Quantification on Frozen Sections**

Rare, homogenously disseminated, TUNEL-, and caspase-3–positive cells were found in the control 3D-HCE cultures (Figs. 3A1, 3A2). BAK concentrations ranging from 0.01% to 0.1% significantly increased the number of TUNEL-positive cells in a dose-dependent manner with 83, 85, and 310 cells/mm EL after 0.01%, 0.02%, and 0.1% BAK treatments, respectively (Fig. 4A). At these three concentrations, the cells did not recover after the postincubation period, and more cells tended to undergo apoptosis with 140, 261, and 390 cells/mm EL, respectively. Of

**Figure 2.** Cellular viability of PBS, 0.001%, 0.005%, 0.01%, 0.02%, and 0.1% BAK-treated 3D-HCE after 24 hours of treatment with and without the 24-hour postincubation period. BAK induced a dose-dependent decrease in cellular viability as assessed by the MTT test. This decrease was considered significant for concentrations as low as 0.005% compared to the control and according to multiple pair-wise comparisons using the Bonferroni adjustment (n = 3).

**Figure 3.** Immunolocalization of TUNEL (row 1), activated caspase-3 (row 2), Ki67 (row 3), and ICAM-1 (row 4) green positive cells. (A1) TUNEL immunostaining of 3D-HCE cells showing an epithelium disclosing no or very rare apoptotic cells after PBS treatment; (B1) apoptotic cells in the most apical cell layer; and (C1) a great number of apoptotic cells in all cellular layers. (A2) Activated caspase-3 immunostaining showing an epithelium with no or rare positive cells after PBS treatment; (B2) with positive cells distributed in all cellular layers after 0.02% BAK treatment for 24 hours; and (C2) with few positive cells throughout the epithelium after 0.1% BAK treatment for 24 hours. (A3) Ki67 immunostaining showing an epithelium with proliferating cells not restricted to the basal layer after PBS treatment for 24 hours; (B3) numerous proliferating cells, with a great number located in the basal layer after 0.01% BAK treatment for 24 hours; and (C3) disclosing no proliferating cells after 24 hours of treatment with 0.1% BAK. ICAM-1 immunostaining showing an epithelium (A4) with ICAM-1–positive cells after PBS treatment; (B4) with an increased number of ICAM-1–positive cells after 0.01% BAK treatment; and (C4) with few remaining ICAM-1–positive cells after 0.1% BAK treatment. The nuclei were stained with DAPI (blue, rows 1 and 2) or propidium iodide (PI, red, rows 3 and 4).
** statistically significant compared to all the other groups at 24h+24h with p<0.003

* statistically significant compared to concentrations ≥ 0.01% at 24h and at 24h+24h with p<0.003

† statistically significant compared to control at the same time with p<0.003

pin 24h and 24h+24h significantly different with p<0.0001

**The number of cells is given per 1 mm of epithelial length (mm.E.L.**

**Figure 4.** Quantification of TUNEL, activated caspase-3–, Ki67–, and ICAM-1–positive cells after 24 hours of treatment with PBS or BAK followed or not by a 24-hour postincubation period. (A) The quantification of TUNEL-positive cells shows BAK-induced apoptosis with a dose-dependent effect reaching significance for 0.01% BAK and higher concentrations. (B) BAK at 0.01% and 0.02% significantly increased the number of activated caspase-3–positive cells at 24 hours, and at 24+24 hours compared with control cells, respectively. (C) A pool of Ki67-expressing cells was found in the control group. The number of proliferating cells was significantly increased by 24-hour 0.01% BAK treatment. This number, although decreasing slightly after the 24-hour postincubation period, remained significantly higher than in the control. After 0.02% and 0.1% BAK treatments, no proliferating cells could be counted because most of the cells underwent cell death. (D) ICAM-1–expressing cells were found in the control group. After 24 hours of treatment with BAK concentrations ranging from 0.001% to 0.01%, the number of these positive cells significantly increased with and without the 24-hour postincubation period. The number of ICAM-1–expressing cells significantly decreased during the 24-hour postincubation period after 24 hours of treatment with 0.02% BAK. After 0.1% BAK treatment, only a few cells still expressed ICAM-1. Mean ± SD are represented.

interest, increasing BAK concentrations induced apoptosis in an increasing number of cellular layers, from the most superficial to the deepest basal layers (Figs. 3A1–C1). The number of activated caspase-3–positive cells increased significantly at 24 hours after 0.02% BAK treatment (87 cells/mm EL) and at 24+24 hours after 0.01% (33 cells/mm EL) and 0.02% BAK (39 cells/mm EL) treatments (Figs. 3B2, 4B). These cells were mainly located in the superficial layer (Figs. 3A2–C2). Approximately, 21 cells/mm EL were found to be positive for Ki67 in 3D-HCE cells treated with PBS (Fig. 4C). These cells were not restricted to the basal layer, but few of them were found in the suprabasal layers (Fig. 3A3). The 24-hour treatments with 0.001% and 0.005% BAK did not significantly change the number of proliferating cells. However, the 0.01% BAK concentration significantly increased this number at levels as high as 81 and 53 cells/mm EL at 24 hours and 24+24 hours, respectively. Most of them were found in the basal layer (Fig. 3B3, 4C), and no proliferating cells were observed after 0.02% and 0.1% BAK treatments (Fig. 3C3). Control 3D-HCE cultures were found to express CD54 (ICAM-1) at levels of approximately 12 cells/mm EL (Figs. 3A4, 4D). After 24 hours of treatment with concentrations ranging from 0.001% to 0.01%, BAK significantly increased the number of CD54-positive cells (P < 0.003 compared with control) at levels greater than 137 cells/mm EL, with positive cells mainly located in the basal layers (Figs. 3B4). The number decreased to 98 cells/mm EL after 0.02% BAK treatment. The decrease reached significance with 0.1% BAK (P < 0.003 compared to control) at 24 hours and 24+24 hours. Of note, the postincubation period appeared to be critical for the 0.02% BAK concentration, as it showed a significant decrease in CD54– and activated caspase-3–positive cells compared with the treatment without a postincubation period. Neither HLA-DR nor E-cadherin was detected in the 3D-HCE cultures treated with PBS or BAK (data not shown).

**En Face Confocal Microscopic Analysis of Occludin Protein Expression**

En face confocal microscopic analysis of 3D-HCE cultures treated with PBS revealed a fine membrane immunostaining of
occludin in the large superficial cells, forming a ring around the cells (Fig. 5). A slight occludin staining was also founded in the deeper cell layers (data not shown). After treatment with 0.001% and 0.005% BAK for 24 hours, a persistent membrane staining in the superficial layers was observed, but it disappeared at 0.01% and more concentrated BAK solutions. After the 24-hour postincubation period, no significant change in occludin protein distribution was found compared with treatment without the postincubation period (data not shown).

Occludin mRNA Expression Assessed by RT-PCR

BAK showed dose- and time-dependent effects on occludin gene expression, with inverse kinetic expression patterns for the 0.001% to 0.02% and the 0.05% to 0.1% ranges of concentrations (Fig. 6). When increasing the duration of treatment (6–24 hours) or adding a postincubation period (24–48 hours), occludin gene expression tended to increase at the low range of concentrations, whereas it decreased at the high range of concentrations. The very low 0.001% and 0.005% BAK concentrations were unable to increase occludin gene expression at 24 hours but tended to increase this expression at 24 hours of treatment followed by the 24-hour postincubation period. At 24 hours, the 0.01% and 0.02% BAK concentrations upregulated the occludin gene expression at levels reaching 1.9- and 2.6-fold the control level, respectively. This expression increased to a greater extent at 24+24 hours with an RQ of 3.3 and 6, respectively. The highest 0.05% and 0.1% BAK concentrations induced increased expression levels at the early time of 6 hours with a RQ of 3.2 and 2.9, respectively. Then, for both concentrations, occludin expression was decreased at levels still greater than the control at 24 hours and less than the control at 24+24 hours. During the 6-hour treatment, occludin mRNA expression increased with increasing BAK concentrations up to 0.05%. Then, the expression decreased at the highest 0.1% concentration.

**FIGURE 5.** Immunofluorescence analysis of occludin protein expression by using en face confocal microscopy after treatment with PBS (A), 0.005% BAK (B), and 0.01% BAK (C) for 24 hours. The epithelium showed a fine occludin staining in the most superficial large cells, forming a continuous ring around them (A). After treatment with 0.005% BAK for 24 hours, the epithelia showed a persistent occludin expression in the apical cells (B), which disappeared after treatment with 0.01% BAK for 24 hours (C), leaving a diffuse cytoplasmic staining of occludin visible in the wing cells. Note that the undulation of the epithelium gives artifactual images of some epithelial defects, but the cell borders are visible when focusing.

**FIGURE 6.** Quantitative analysis of occludin mRNA expression in 3D-HCE treated with BAK. BAK concentrations ranging from 0.001% to 0.02% induced a dose- and time-dependent increased expression of occludin mRNA in 3D-HCE cells, whereas for the two next higher BAK concentrations—0.05% and 0.1%—the expression seemed to decrease with higher doses and longer treatment periods. Twofold quantitative ratios (QR) were considered significant.
DISCUSSION

In the rabbit Draize test, the eye irritancy score is strongly influenced by the response of the cornea, which accounts for approximately 75% of the overall score. Because of the problems of interspecies differences and given that its structure is similar to the living human corneal epithelium, the 3D-HCE cell was thought to be a good surrogate for in vivo testing in the field of ocular irritation and ocular safety assessment. A relatively small number of large-scale studies have focused on the evaluation of this model. Doucet et al. tested 65 cosmetic products and 35 chemicals and demonstrated a good correlation between MMAS values generated by the Draize test and in vitro results using the 3D-HCE system. In 2006, a multicenter prevalidation study was conducted to assess the ability of 3D-HCE to assess the eye irritating potential of chemicals using individual rabbit data described in the ECETOC (European Centre for Ecotoxicology & Toxicology of Chemicals) data bank and in 2008. The authors found the model to be a quality-controlled and easy-to-perform test system, with a high level of reproducibility. They emphasized that the selection of quality in vivo data on eye irritation is one of the major issues for the (pre)/validation process of new in vitro methods and provided a proof of principle that the 3D-HCE system can help discriminating between eye irritants and nonirritants. However, Van Goethem et al. recommended considering additional protocol modifications that may increase the discriminative power of the assay and indicated that false-negative results could be prevented by performing histologic analysis when irritants fail to induce a significant reduction in the MTT-determined viability. MEA were therefore proposed for testing mild irritants, since the tissue viability might remain unaffected, resulting in false-negative results. Although we agree that an additional endpoint may help differentiate the mechanisms of irritation and also increase confidence in negative results, we kept in mind that numerous studies confirmed the hypothesis that area and depth of chemical-induced corneal injury correlates with cell death and can predict the extent of subsequent ocular responses.

The present study, we optimized the classic MTT protocol used for irritation assessment to increase its sensitivity and more accurately correlate with histologic findings. With this new procedure, we demonstrated that BAK induced a significant dose-dependent decrease of cell viability, with worsened effects after a 24-hour postincubation period for concentrations greater than 0.005%. These results correlated with the number of TUNEL-positive cells and correlated inversely with the number of Ki67-positive cells. This was consistent with our previously published study showing a dose-dependent increase in epithelial apoptosis in rat conjunctival brushings. Moreover, the MTT test succeeded in detecting decreased cellular viability for concentrations as low as 0.005% BAK, whereas the former procedure failed to detect the effects of a 0.01% BAK concentration at 24 hours. This new, more sensitive MTT procedure is therefore expected to better predict potential adverse effects that may occur with the long-term use of ophthalmic products. It may also help optimize the protocol currently used for acute eye irritation assessment and help future validation studies. Indeed, one prediction model currently proposed to distinguish irritants from nonirritants was defined based on a viability cutoff value of 60%. The use of a more sensitive MTT protocol may thus decrease this cutoff value and enlarge the scale for discriminating mild and moderate irritants. Of note, low doses of BAK induced apoptosis restricted to the superficial cell layers and proliferation mainly located in the basal layers, whereas high doses of BAK induced apoptosis in all cellular layers, including the deepest one, and hindered cell proliferation. BAK-induced cell apoptosis, at least partly mediated by caspases, as demonstrated by increased activated caspase-3-positive cells with 0.01% and 0.2% concentrations. With 0.1% BAK, activated caspase-3 was not detected, probably because most of the cells were already undergoing apoptosis and because the activation of caspase-3 precedes the induction of apoptosis. It is noteworthy that BAK used at the low 0.001% to 0.01% concentrations seemed to elicit a defense response by increasing Ki67 and ICAM-1 expression as well as by increasing occludin mRNA expression; Ki67 distribution patterns were modified by 0.01% BAK only. BAK showed dose- and time-dependent effects on occludin gene expression, with inverse toxicokinetic profiles observed at the 0.001% to 0.02% and the 0.05% to 0.1% ranges of BAK concentrations. This suggests that for concentrations greater than or equal to 0.05%, the cellular defense system was no longer able to respond for treatments exceeding 6 hours.

Previous studies have already described the distribution of T-junction proteins occludin in rat, rabbit, and human corneal epithelia as well as in human corneal cell cultures. Our results agreed closely with those in these studies, since we found a strong membrane occludin staining in the most apical superficial cells of the control 3D-HCE cultures. BAK induced the dose-dependent disappearance of occludin, suggesting the disruption of the epithelial tight junctions in the superficial cells. These results are consistent with the known enhancer properties of BAK used to increase corneal penetration of active compounds and they confirmed the results of other studies showing that the occludin distribution in monolayer cultures could be altered by a chemical treatment. However, in two very recent studies, the expression of functional tight junctions, which is usually characterized by electrophysiological parameters such as transepithelial electric resistance (TEER), has been investigated in various corneal models including the 3D-HCE model (SkinEthic Laboratories). The studies reported that only one of these models, the HCE-T system, which is also based on transformed human corneal epithelial cells, exhibited epithelial tightness and functional integrity, whereas the TEER values of the other models were far below the TEER values reported for isolated cornea. These results were confirmed in permeation studies, with permeation coefficients that were increased by about 10-fold relative to the isolated cornea. In eye irritation testing, because intrinsic cytotoxicity potential of chemicals is a decisive factor affecting cornea after contact, the evaluation of cell viability remains the principal evaluation endpoint and epithelial barrier properties and compound permeability are typically not an issue. The permeation studies suggested, however, that the differentiation in most of the today available 3D ocular models does not occur as completely as in the human corneal epithelium.

ICAM-1 is not expressed on normal corneal epithelium. However, we found the expression of ICAM-1 in our control 3D-HCE cultures. This result was consistent with that in previous work on cultured HCE cells that demonstrated ICAM-1 to be involved in corneal epithelial cell movement. The BAK-induced overexpression of ICAM-1 on 3D-HCE suggests that it can contribute to corneal epithelial cell injury by aiding the attachment of inflammatory cells such as eosinophils, which express the receptor for ICAM-1 and the β2 integrins (CD11a, -b, -c/CD18). Ki67 protein expression was observed in the nuclei of a few basal cells within the central and peripheral corneal epithelium of human donors, suggesting that individual cells within this layer undergo asynchronous cell division to replace cells lost from the epithelial surface. Organotypic epidermal and oral cultures demonstrated an increased proliferation pattern (Ki67 staining) in basal and suprabasal layers after exposure to acetone and low concentrations of sodium lauryl sulfate, respectively. Our results are in agreement
with these studies and suggest that mild to moderate stress can trigger proliferation as a mechanism of tissue defense. Finally, neither HLA-DR nor E-cadherin was detected in control and treated 3D-HCE cells.

Taken together, these results suggest that the MEA and the use of our modified, more sensitive MTT protocol will be useful for the preclinical irritation screening of new ophthalmic products in this type of tissue model. In particular, this approach may be a better predictor of the subclinical signs of irritation and toxicity of substances applied topically to the skin and mucous membranes. *J Pharmacol Exp Ther.* 1994;82:377–390.


