**Pathogenesis of acute and delayed corneal lesions after ocular exposure to sulfur mustard vapor**

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Pathogenesis of Acute and Delayed Corneal Lesions After Ocular Exposure to Sulfur Mustard Vapor

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Purpose: Sulfur mustard (SM) exposure results in dose-dependent morbidities caused by cytotoxicity and vesication. Although lesions resulting from ocular exposure often resolve clinically, an idiopathic delayed mustard gas keratopathy (MGK) can develop after a moderate or severe exposure. Sequelae include persistent keratitis, recurring epithelial lesions, corneal neovascularization, and corneal degeneration, which can lead to impaired vision or loss of sight. The purpose of this effort is to correlate structural changes with injury progression during the development of MGK.

Methods: New Zealand White rabbit corneas were exposed to SM using a vapor cup delivery system. The transition from acute to delayed injury was characterized by clinical, histological, and ultrastructural metrics over 8 weeks.

Results: Exposure dose was correlated to the likelihood of developing MGK but not to its severity. In a 56-animal cohort, a 2.5-minute exposure generated a corneal lesion, with 89% of corneas developing MGK within 5 weeks. A significant decrease in corneal edema at 2 weeks was predictive of the 11% of corneas that underwent asymptomatic recovery. Ultrastructural comparison of asymptomatic and MGK corneas at 8 weeks indicates that MGK is characterized by persistent edema and profound disorganization of the basement membrane zone.

Conclusions: Ultrastructural changes associated with the delayed pathophysiology of corneal SM vapor exposure involve severe degeneration of the basement membrane zone and persistent edema. The mechanisms underlying MGK pathogenesis seem to alter injury progression as soon as 2 weeks after exposure. These data suggest that the vapor cup model system is suitable for therapeutic evaluation.

Key Words: mustard gas keratopathy, ocular toxicity, vapor exposure, sulfur mustard

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Sulfur mustard (2,2′-dichloroethylsulfide; SM) is a highly reactive bifunctional chemical that alkylates proteins and nucleic acids. Battlefield deployment of SM as a chemical weapon in World War I and the Iran–Iraq war resulted in more than 210,000 British and Iranian casualties, 90% of which presented with ocular lesions. In humans, the acute stage of ocular SM toxicity involves dose-dependent morbidities caused by vesication of the corneal epithelium (CE) and keratocytosis in the epithelium and stroma. Eyes are the most sensitive organ to SM injury, and although mild exposures typically resolve uneventfully, those that receive moderate or severe exposures (exceeding 100 μg·min⁻¹·m⁻³) exhibit 3 distinct clinical trajectories: (1) injury resolution, after which the victim remains asymptomatic; (2) persistent keratitis that ultimately results in corneal degeneration (chronic injury); or (iii) an asymptomatic period followed by reemergence of lesions (delayed-onset injury). The latter 2 trajectories comprise the phenomenon known as mustard gas keratopathy (MGK), which has been diagnosed in 16% of casualties receiving a moderate or worse exposure.

MGK is characterized by a persistent inflammatory condition in the cornea (keratitis) followed by development of secondary keratopathies, such as recurring corneal epithelial erosions (RCEs) and neovascularization (NV). In the chronic form of SM toxicity, MGK symptoms develop from the acute injury; alternatively, the delayed-onset form appears after an asymptomatic latent period lasting 0.5 to 40 years. Histology of corneas excised from MGK patients undergoing penetrating keratoplasty displays sequelae of chronic inflammation, such as stromal degeneration and necrosis, suggesting a persistent injury that is beyond the healing capacity of the cornea. Similar findings were reported after in vivo confocal microscopy of SM survivors with moderate symptoms of MGK.
The standard of care for acute SM lesions is similar to that of other corneal abrasions (eg, irritation, topical mydriatics, antibiotics, and steroids); however, in animal models, these do not seem to attenuate late ocular complications.\textsuperscript{1,9} Because the etiologies of the MGK sequelae are unknown, therapeutic strategies have predominantly been supportive and applied succeedent to the onset of symptoms.\textsuperscript{10,11} In the more severe cases, penetrating keratoplasty has been attempted with mixed outcomes.\textsuperscript{2,6} Given the threat of permanent visual impairment and chronic discomfort among victims of ocular SM injury, there is a critical need for an effective therapeutic that mitigates long-term toxicity.

Rabbits have been a model system for ocular SM injury for more than 60 years.\textsuperscript{12,13} They exhibit several anatomical and physiological features that facilitate ocular toxicity research, such as a large corneal to sclera ratio, a relative insensitivity to corneal drying, and a low frequency of spontaneous epithelial lesions.\textsuperscript{14} Rabbit corneas have greater structural similarity to human eyes than do mice, rat, or guinea pig corneas, and although rabbits are 4-fold less sensitive to ocular SM injury than humans, at normalized doses, they display similar sequelae and injury progression.\textsuperscript{12,15} Recently, rabbit exposure models have been used to evaluate candidate treatments, test novel ocular delivery systems, and evaluate long-term SM toxicity in the limbus.\textsuperscript{9,16–18}

The traditional model of corneal exposure uses liquid drops of SM applied directly to the cornea.\textsuperscript{12} This generates a localized ulcerative injury as opposed to the milder, more evenly distributed injury incurred by vapor exposure.\textsuperscript{13} Nonetheless, researchers have historically used liquid SM out of concerns that mechanical injury incurred by the cornea during vapor cup delivery might compromise ultrastructural and toxicological evaluation. Although goggle-based delivery of SM vapor to the entire eye avoids these concerns and more closely reflects a battlefield injury, the specific progression of corneal injury and contribution of corneal injury to MGK is complicated by injury to the ocular adnexa, which may influence the extent or degree of corneal pathogenesis.\textsuperscript{19} Thus, although corneal complications are the signature of moderate and severe long-term ocular SM injury, the keratopathogenesis of long-term toxicities after a diffuse exposure has not been well described.\textsuperscript{11} We previously reported on a vapor cup system to deliver SM vapor to the cornea of New Zealand White rabbits.\textsuperscript{20} In preliminary work, we demonstrated that exposure of rabbit corneas to SM vapor resulted in vesicating injuries and at higher doses caused recurring lesions.\textsuperscript{20} This exposure model was not observed to cause direct SM injury to noncorneal tissues, allowing an emphasis on mechanisms of corneal pathogenesis. In this report, we use the vapor cup model to characterize clinical and histological characteristics of the transition from the acute injury to the delayed injury and for the first time present ultrastructural changes associated with the pathophysiology of corneal SM vapor exposure.

**MATERIALS AND METHODS**

**Animal Use**

Rabbit experiments involved 3 cohorts. In the first cohort, 30 corneas were exposed to SM vapor for 1 to 3.5 minutes (n = 5 per dose), followed by weekly evaluations and histopathological analysis at 6 weeks. The second involved exposure of 28 animals for 2.5 minutes followed by histopathological characterization at 1 to 7 days. The third involved exposure of 56 animals for 2.5 minutes followed by regular clinical evaluations and histology/transmission electron microscopy (TEM) at 8 weeks.

Rabbits were ordered from Charles River Laboratories at 2 to 2.5 kg and caged individually on an enriched diet ad libitum. Before exposure, animals underwent a baseline corneal evaluation. On the day of exposure, a surgical plane was induced by intramuscular injection of ketamine HCl (15 mg/kg) and xylazine (7 mg/kg), and a loading dose of buprenorphine HCl (0.05 mg/mL) was administered subcutaneously. Osmotic pumps (Alzet, Cupertino, CA) primed to deliver 10 \textmu L/h of buprenorphine HCl (0.3 mg/mL) for 7 days were implanted subdermally between the scapulae. After exposure, animals were returned to individual cages and monitored daily. Pumps were removed 7 days after implantation. At the termination of each experiment, animals were euthanized by cardiac delivery of 2 mL of pentobarbital sodium (390 mg/mL).

**Exposure Procedures**

Before exposure, 10 \textmu L of neat SM was distributed on Whatman filter paper firmly lodged in the bottom of a 14 \times 5.2-mm deep screw cap modified with a 12-mm rubber O-ring, designed to minimize mechanical injury to the cornea and to form a capillary seal with the tear film. Loaded caps were inverted for 1 minute and placed directly over the center of the rabbit eye for 1 to 4 minutes. After exposure, animals were monitored for 2 minutes, and remaining agent was gently flushed from the eye with 10 mL of sterile saline. The contralateral eye served as an exposure control. No adverse outcomes were observed in control eyes with saline-soaked vapor cups. Exposure hoods were maintained at 22 to 22.5°C and at 35% to 45% humidity.

**Clinical Evaluations**

Rabbits were evaluated at regular intervals starting 24 hours after exposure. The presence of corneal epithelial lesions was characterized by blue light photography of eyes 1 minute after treatment with 40 \textmu L of fluorescein; retained fluorescein is indicated of a disruption in epithelial integrity. Planimetric lesion size was calculated on photographs using NIH Image after image registration. Pachymetry and slit-lamp examination were used to quantify corneal edema, and white light evaluation was used to identify opacity and NV. Corneal epithelial erosions and NV were graded as present or absent. Changes in mean corneal thickness and planimetric lesion size were compared using a t test.

**Histopathology and TEM**

After enucleation, eyes were fixed by injection of buffered 1.6% paraformaldehyde and 2.5% glutaraldehyde into the posterior chamber, followed by immersion in the same at 4°C for 24 to 28 hours. Corneas were excised and
representative areas of tissue to include any gross lesions were isolated by scalpel and processed for routine histopathology and TEM. Light microscopy was performed using an Olympus BX51 microscope. Categorical scoring for histopathology used the following system: “0” = no evidence of injury, “1” = injury is present over <10% of the area, “2” = injury over 11% to 25% of area, “3” = injury over 26% to 50% of area, and “4” = injury over 51% to 100% of area. The degree and extent of epithelial and stromal sequelae were graded similar to Milhorn et al20 to include ulceration, edema, necrosis, deformation, NV, and inflammation, and scores were summed, then compared by the Mann–Whitney rank sum test with Bonferroni adjustment. The mean number of heterophils in 2 hematoxylin and eosin–stained sagittal sections per cornea were averaged across 4 cornea per time point and compared using a t test.

Tissues selected for TEM were postfixed in buffered 1% osmium tetroxide, dehydrated in graded ethanol, and embedded in Poly/Bed 812 resin. Semithin sections (1.5 μm thick) were stained in 1% methylene blue and 1% azure II. Ultrathin sections (90 nm thick) were mounted on copper mesh grids and counterstained using uranyl acetate and lead citrate. The observation of sequelae in micrographs from at least 3 corneas were considered to be genuine. TEM was performed using a JEOL JEM-1230 transmission electron microscope.

Gas Chromatography–Mass Spectrometry

Rabbit plasma was collected and clarified by centrifugation at 2000g for 30 minutes at 5°C. Protein from plasma samples obtained from exposed animals or spiked standards was isolated and prepared as described with the following modifications.21 Plasma protein was extracted from 1 mL of plasma by 3 washes with acetone, and precipitated protein was dried. The protein pellet was weighed and digested at 125 mg/mL in 1M NaOH at 70°C. Digested protein was doped with 0.6 ng of the internal standard octadeuterothiodiglycol, pH adjusted with 3M HCl, and extracted with neat ethyl acetate. Extracts were derivatized, passed over preconditioned Bond Elute silica 100-mg Si SPE cartridges and evaluated by gas chromatography–mass spectrometry. Calibration curves were prepared using rabbit plasma treated with 0.78 to 50 nM SM for 2 hours at 37°C. Protein precipitated from the calibration curve samples was processed as described above. Calibration curves established a limit of detection of 0.07 ng/mL and a limit of quantitation of 0.24 ng/mL and were used with each set of rabbit samples to calculate plasma SM concentrations. Sample means were compared across dose and time by t test.

RESULTS

Acute and Delayed SM Toxicity Is Dose Dependent

To correlate exposure dose with long-term outcomes, we used the vapor cup model to deliver SM vapor to rabbit corneas for 1 to 3.5 minutes, evaluated the clinical status at weekly intervals, then scored the corneal histopathology at 6 weeks. All animals underwent an acute injury characterized by corneal edema, corneal opacity, and epithelial lesions (Fig. 1A, top panels). After partial healing of the acute injury, a second set of pathologies developed between 3 and 5 weeks, in which corneas exhibited increased corneal edema, immune cell infiltrates, RCEs, and NV (Fig. 1A, bottom panels). Clinical evaluations indicated a strong relationship between dose and the

FIGURE 1. Severity of acute injury and frequency of delayed symptoms correlate well with dose, whereas severity of delayed injury does not. A, Corneas demonstrated escalating severity of acute sequelae at 1 week but similar severity in delayed-onset injury at 8 weeks. B, Frequency of development of late-onset symptoms at different durations of SM vapors exposure. C, Cumulative histopathology scores at different doses.

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likelihood of developing delayed keratopathies (Fig. 1B), but histopathology suggested that the dose was not related to the severity of delayed keratopathies (Fig. 1C). Based on these data, we selected a 2.5-minute exposure for future work because this produced corneal injuries similar to those resulting from moderate-to-severe SM exposures in humans, with a high prevalence of recurring lesions for therapeutic testing.\textsuperscript{11}

The Acute Lesion Involves Vesication of the CE and Corneal Cytotoxicity

All exposed corneas developed an epithelial lesion within 1 day (Fig. 2A). Although lesion morphology changed between 1 and 2 days, the total lesion area did not, suggesting competing processes of in-migration and lesion expansion (Fig. 2B). The CE regenerated and became refractory to the fluorescein assay in 96% of corneas by 4 days (a photographic panel of reepithelialization is presented in Supplemental Digital Content 1, http://links.lww.com/ICO/A32). Stromal edema was present immediately under the denuded cornea as soon as 1 day after exposure (Figs. 2C, D).\textsuperscript{20}

Ultrastructural changes associated with acute toxicity were evaluated by TEM (Fig. 3). As is typical with vesicating injuries, separation between the epithelium and stroma occurred within the lamina lucida (Fig. 3B).\textsuperscript{22} Migrating epithelial cells could be observed as soon as 2 days. Desmosomal attachments between epithelial cells suggest that migration occurred as an epithelial sheet (Fig. 3C), and the basal membranes were highly involuted, indicative of a mobile population forming transient adhesions to the basement membrane (Fig. 3D). By 4 days, nascent hemidesmosomal densities in the basal membranes of CE cells and associated anchoring plaques in the Bowman-like layer (BLL) became apparent (Figs. 3E, F). By 7 days, the CE was stratified with maturing basement membrane zone (BMZ) architecture (Fig. 3F). Conversely, the stroma continued to show signs of edema and immune cell infiltration, with necrotic fibrocytes distributed through the stromal volume underlying the exposure site appearing as soon as 1 day after SM treatment (Figs. 3H, I).

Delayed Injury Frequencies Within Experimental Cohorts Suggest That Multiple Clinical Outcomes Are Possible

To evaluate the development of MGK from the acute injury, 56 corneas were longitudinally evaluated using clinical metrics over 8 weeks. Despite regeneration of an intact epithelium within 5 days (which also is the point of maximum edema during the acute injury), 89% continued to exhibit a persistent keratitis characterized by elevated edema (Fig. 4A) and immune cell infiltrates (Fig. 4B). Of these, 80% subsequently developed secondary keratopathies between 3 and 5 weeks with RCEs (Figs. 4C, D) and NV (Fig. 4C). Interestingly, although recurring lesions were only about one third to one half the size of the acute lesions, the morphology varied on a weekly basis, suggesting cyclic attempts to reepithelialize followed by the development of new lesions (Fig. 5, bottom panels, days 21–56). The remaining 11% of animals progressively became asymptomatic, with healthy appearing corneas and minimal opacity by 2 weeks, a decline in corneal thickness to baseline values by 5 weeks, and failure to develop RCEs or NV. Notably, corneas undergoing asymptomatic recovery could be distinguished as soon as 2 weeks after exposure by a significant decrease in corneal thickness (Fig. 4E, dashed line). Despite persistent markers of inflammation in all corneas that would develop MGK, there was only mild evidence of corneal injury until secondary keratopathies developed (Fig. 5, days 7–21). No cornea that transitioned to a persistent keratitis subsequently exhibited clinical improvement or resolution within 8 weeks. A summary of clinical outcomes and characteristic sequelae is presented in Table 1.

FIGURE 2. Clinical and histopathological metrics of the acute injury. A, Planimetric area of fluorescein staining in exposed and sham-exposed corneas (n = 25). B, Percent of corneas exhibiting fluorescein retention over the first week after exposure (n = 75). C, Thin sections demonstrating histological changes during the acute injury (×4). Scale bar = 200 μm. D, Full corneal strips (×10) from same slides in (C). At 4 days, it was not possible to image the entire stroma due to the extent of edema. Arrowheads represent margin of acute lesion.
Resolved Corneas Do Not Exhibit the Severe Ultrastructural Disorganization Displayed in Corneas Experiencing Late SM Lesions

In resolved corneas 8 weeks after exposure (Fig. 6A; also see Supplemental Digital Content 2, http://links.lww.com/ICO/A33, for clinical metrics and gross images), the CE was fully stratified and of normal thickness, with extensive desmosomes and without evidence of intercellular edema (Fig. 6B). Although the BMZ was not yet fully mature, there were numerous hemidesmosomal densities and associated anchoring plaques in the BLL, and the lamina lucida and lamina densa were well developed without disruption (Fig. 6C). Although necrotic remnants of basal epithelial cell intrusion into the stroma were occasionally observed, they were...
structurally isolated from the epithelium by a regenerated intact BMZ. The stroma exhibited a low-grade edema, with mild disruption of the lamellar structure and occasional infiltration of inflammatory cells (Fig. 6D).

Conversely, MGK corneas at 8 weeks exhibited severe ultrastructural degeneration consistent with the clinical and histological sequelae (Fig. 6E; also see Supplemental Digital Content 2, http://links.lww.com/ICO/A33, for clinical metrics and gross images), including extensive disruption of the BMZ, necrotic and apoptotic CE cells, incomplete and disorganized stratification of the CE, and significant stromal edema (Figs. 6E–H). In general, the BMZ was highly disorganized. The presence of redundant basal lamina was common, and numerous examples of basal cell processes extending through interruptions in the basement membrane into the BLL were observed (Figs. 6F, G). Basal epithelial cells exhibited an irregular distribution of hemidesmosomal densities ranging from rudimentary to complex in nature (Fig. 6G). Moderate to severe edema was present within the basal and suprabasal epithelial cell layers. Apoptotic bodies and cellular debris were also observed within extracellular gaps. Necrotic basal cells overlay the fragmented basement membrane, whereas desmosome-rich wing and superficial cell layers remained intact. The stroma was characterized by full-thickness edema, disorganized collagen fibrils, distorted lamella, heterophil infiltrates, and necrotic fibrocytes (Fig. 6H).

**Differences in Clinical Outcome Cannot be Attributed to Variable Dosing**

The observation that similar exposures resulted in significantly different outcomes raised the concern that corneas were receiving variable doses of SM vapor. Because SM-alkylated proteins can be identified in the plasma after cutaneous and inhalational exposures in a dose-dependent manner, we evaluated whether plasma adducts could be detected after ocular exposures. Plasma adduct concentrations were tightly correlated with duration of exposure (dose; Fig. 7A; \( R^2 = 0.98 \)) and cleared with a half-life of 130 ± 21 hours (Fig. 7B; \( R^2 = 0.98 \)), similar to the half-life of rabbit albumin. Applying this analysis to plasma isolated from the 56-animal cohort revealed a coefficient of variation of 31%, indicating a consistent exposure (Fig. 7C). Retrospective correlation of plasma adduct concentrations with clinical outcomes demonstrated no significant relationships with corneal thickness at 1 week (Fig. 7D; \( R^2 < 0.01 \)), incidence of RCEs at 4 weeks, incidence of NV at 6 weeks (Fig. 7E), or animal weight at time of exposure (Fig. 7F; \( R^2 < 0.12 \)). The
slight correlation between adduct levels and animal weight is consistent with the change in plasma volume with respect to animal weight and therefore could reflect dilution in a larger plasma volume.\textsuperscript{26} In some cases, animals receiving relatively large doses of SM exhibited no symptoms of MGK and vice versa. These data support the observation that factors other than dose can influence whether late SM lesions will develop and are likely to underlay the observation of diverse clinical outcomes within exposure cohorts.

FIGURE 5. Longitudinal characterization of injury progression in a representative cornea over 8 weeks. White light photographs (×6.3, top panels) and images from fluorescein exclusion assay (×10, bottom panels) visualizing the acute injury and development of MGK in a characteristic rabbit.

DISCUSSION

Human and Rabbit Injury Progression After Corneal SM Exposure

In humans suffering from MGK, chronic inflammation results in corneal degeneration, permanently reducing visual acuity and potentially causing complete loss of vision.\textsuperscript{5} The etiology of late SM lesions has been described as the product of a complicated series of processes, starting with direct SM injury to epithelial, stromal, and possibly endothelial tissues and leading to secondary responses in multiple cell types converging on the damaged cornea.\textsuperscript{22} Multiple mechanisms of MGK have been proposed, including limbal stem cell deficiency, collagenase-resistant stromal–SM adducts inducing proinflammatory responses, basement membrane dystrophy, and barrier dysfunction resulting in persistent edema.\textsuperscript{17,20,27} No therapeutic approach has succeeded in preventing late SM toxicity.

Our data suggest that vapor cup delivery to rabbit corneas produces a milder and more diffuse injury than liquid SM drops, providing increased consistency in ocular response with greater similarity to the characteristic human casualty.\textsuperscript{12,13,28} Sham-exposed eyes exhibited neither clinical nor histopathological morbidities, and clinical outcomes were
similar to those reported for goggle-based vapor delivery models, suggesting that the vapor cup exposure model does not induce a mechanical injury to the cornea. After a 2.5-minute vapor cup exposure, 89% of corneas transition to a persistent keratitis, with 80% subsequently developing symptoms similar to human MGK (RCEs, persistent edema, corneal degeneration, and inflammatory infiltrates) between 2 to 5 weeks after exposure. Although the eye appears grossly quiet during the period between acute injury and the development of secondary keratopathies, the injury is not clinically asymptomatic. There was no significant improvement in clinical or histological metrics of corneal health after the onset of delayed SM lesions, suggesting that this persistent keratitis is the rabbit equivalent of the chronic form of human MGK. It is unknown whether the 11% of animals that exhibited a persistent keratitis without developing secondary keratopathies would do so beyond 8 weeks.

Overview of Pathogenesis and Early Identification of Corneas That Will Not Develop MGK

In rabbit corneas undergoing delayed MGK, injury progression can roughly be divided into 3 stages: (1) the acute injury, which is partially resolved within 1 week; (2) a persistent keratitis characterized by persistent inflammatory sequelae; and (3) development of delayed keratopathies 3 to 5 weeks after exposure. Acute SM toxicity results in cell death and sloughing of the CE through the lamina lucida of the BMZ. Disruption of the epithelial barrier and proinflammatory signaling due to necrosing corneal tissues then lead to corneal edema. Corneas subsequently develop a progressive inflammatory injury between 3 and 5 weeks, with a frequency that is dose dependent. This delayed injury is characterized by RCEs, inflammatory cell infiltrates, bullae, and elevated corneal edema, with associated degradation of the stromal architecture and the BMZ. Notably, significant decreases in corneal edema by 2 weeks after exposure are predictive of injury resolution, suggesting that the underlying pathology of the late SM lesions is present early in the injury process. Kadar et al also reported the early ability to identify resolving corneas based on corneal thickness after goggle-based vapor delivery to the orbital tissues.

The observation that corneas developing MGK did not improve within 8 weeks suggests that either MGK is irreversible at this dose or animals were euthanized before improvement could be detected. The former supposition is supported by electron micrographs of 8-week MGK corneas, which exhibit progressive corneal degeneration, including extensive disruption of the basement membrane, severe stromal distortion, and corneal edema. The clinical status of the small percentage of animals that exhibited a persistent keratitis but did not develop MGK within 8 weeks is unclear. Assuming that persistent keratitis is causally linked to development of the late-onset keratopathies, it seems probable that these animals will ultimately develop MGK. Similarly, even though resolved corneas appear significantly improved compared with MGK corneas, we do not know how these corneas would behave beyond 8 weeks. It may be that some ultimately develop MGK, belying the apparent improvement. If so, this pathology might be equivalent to the delayed-onset injury observed in human casualties, in which an asymptomatic period precedes the appearance of MGK sequelae.

Table 1. Summary of Clinical Outcomes Observed Within 8 Weeks of Ocular Vapor SM Exposure

<table>
<thead>
<tr>
<th>Outcome</th>
<th>n (%)</th>
<th>Common Symptoms/Sequelae</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acute injury</td>
<td>56/56 (100)</td>
<td>Edema, epithelial lesion, corneal cystosis</td>
</tr>
<tr>
<td>Long-term outcomes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Injury resolution</td>
<td>6/56 (11)</td>
<td>Return to baseline values</td>
</tr>
<tr>
<td>Chronic injury</td>
<td>50/56 (89)</td>
<td>Persistent edema, immune cell infiltration</td>
</tr>
<tr>
<td>Secondary keratopathies</td>
<td>45/50 (90)</td>
<td>Recurring corneal erosions increased edema and</td>
</tr>
<tr>
<td></td>
<td></td>
<td>increased opacity, NV, epithelial bullae,</td>
</tr>
<tr>
<td></td>
<td></td>
<td>corneal degeneration</td>
</tr>
<tr>
<td>Persistent keratitis</td>
<td>5/50 (10)</td>
<td>No evidence of secondary keratopathies</td>
</tr>
</tbody>
</table>

The delayed appearance and increased severity of the delayed injury in cornea provide clues to the specific etiology of MGK. The observations that (1) persistent edema is present in all unresolved animals, (2) changes in edema at 2 weeks are predictive of injury resolution, and (3) many of the secondary keratopathies are associated with persistent edema suggest that edema is the principal pathology of the late SM injury. Corneal edema results from fluid accumulation in the cornea and derives from corneal inflammation or disruption of the corneal epithelial or endothelial barrier. Complications of persistent edema may be symptomatically evidenced through multiple sequelae. For example, unresolved edema within the BLL is likely to hamper successful remodeling of the basement membrane, resulting in persistent cellular necrosis, inflammation, and further degeneration of the anterior segment. A stable and functional basal epithelial cell layer is unlikely to develop because there is no structural scaffold on which to deposit or assemble a functional basement membrane. This is consistent with ultrastructural findings demonstrating severe BMZ disorganization and rari- fication of the BLL. The inability of basal CE cells to establish permanent adhesions is likely to be destabilizing, leading to the increased deposition of redundant basement membrane components, edema, and CE cell death, resulting in a cyclical phenomenon of sloughing, edema, and failed attempts to reepithelialize.

Whether endothelial cell disruption contributes to persistent corneal edema is currently unknown. Although
damage to the corneal endothelium would contribute to a loss of barrier function and fluid maintenance within the stroma, there has been no evaluation of endothelial cell toxicity during the acute or delayed SM injury in animal models. In vivo confocal microscopy of human survivors suggests that MGK involves profound decreases in keratocyte and corneal endothelial cell density, providing a mechanism for corneal endothelial barrier dysfunction. The reduction of corneal edema after reepithelialization of the acute lesion between 5 and 14 days indicates some active level of osmotic regulation. However, both delayed corneal endothelial toxicity (eg, due to apoptosis) and RCEs that allow the sustained penetration of protein-rich tear fluid into the stroma could overwhelm the capacity of endothelial cells to deturgesce the stroma.

FIGURE 6. Histological and ultrastructural differences between resolved corneas versus those undergoing delayed SM toxicity at 8 weeks. A, Thin section of a resolved cornea. B–D, Electron micrographs of resolved cornea with emphasis of the: (B) BMZ, (C) higher magnification view of BMZ, and (D) stroma. E, Thin section of a MGK cornea. F–H, Electron micrographs of MGK cornea with emphasis of the: (F) BMZ, (G) higher magnification view of BMZ, and (H) stroma. Asterisk indicates representative examples of necrotic remnants, and viable projections (cp) of basal CE cells into the stroma are marked. Arrows indicate basement membrane, and arrowheads indicate hemidesmosomal plaques. bc, basal cell; hp, infiltrating heterophils; ld, lamina densa; nf, necrotic fibrocyte; and st, stroma. Scale bars = 2 μm.
Implications of Plasma Adduct Measurements to Toxicological and Therapeutic Evaluations

The ability to quantify SM-adducted proteins in the plasma validated the consistency of ocular exposures. Although the specific routes by which SM and/or adducted protein are absorbed into the vascular system are not clear, radiolabeled SM vapor has been shown to penetrate through the cornea to the lens within 5 minutes. The observed disjunction between plasma adduct levels and metrics of acute and delayed injury suggests that individual rabbits exhibit an inherent variability to SM injury, similar to observations from human clinical reports and animal models. The mechanism underlying this variability is of interest and may provide a novel therapeutic approach. Conversely, the inclusion of clinically resistant animals in therapeutic trials can complicate therapeutic evaluation, particularly for therapies applied during the first 2 weeks after exposure (ie, before resolving corneas can be identified).

Therapeutic Considerations

Treatment of SM keratopathies is predominantly symptomatic, aimed at managing ocular irritation and improving vision. Due to the broad degenerative nature of delayed-onset sequelae and persistent keratitis, it seems a priori that the optimal therapeutic approach would be to intervene early in the injury process. In animal models, topical corticosteroids, nonsteroidal antiinflammatory agents, and a matrix metalloprotease inhibitor have been therapeutically evaluated within hours to weeks of SM exposure. In all cases, these treatments resulted in the transient depression of corneal injury, which subsequently rebounded to levels statistically indistinguishable from exposed controls after cessation of therapy. An alternative delivery mechanism that avoids cyclic dosing may provide a stable therapeutic improvement, but current data suggest that reducing edema or decreasing NV without treating the causative pathology does not prevent long-term corneal keratopathies.

The eye is a complex structure, with multiple tissues interacting to maintain ocular health. Although no direct SM injury was observed in orbital adnexa using the vapor cup model, it is likely that under typical battlefield conditions, both corneal and noncorneal ocular tissues will suffer direct SM toxicity. In such a scenario, the nature and extent of MGK sequelae may be secondary to injury processes in other tissues. For example, it has been reported that goggle-based SM vapor exposure alters limbal stem cell viability, providing a potential contributory mechanism for RCEs that may not be observed in the corneal vapor exposure model. Thus, to expand the range of therapeutic targets, we are currently exploring how the interaction between corneal and noncorneal injury processes influences MGK development.

The temporal gap observed between the acute injury and clinical detection of delayed corneal keratopathies may imply a secondary etiology. Potential mechanisms include injuries to supportive tissues or destructive modifications to extracellular material. For example, cells responsible for corneal remodeling may encounter chemically adducted proteins and subsequently initiate a proinflammatory cascade. Alternatively, chemical modification may interfere with the ability of CE cells to stably attach. It is possible that intoxication of limbal or corneal cell populations may have a delayed effect on cell viability, such as inducing apoptosis through disruption of DNA synthesis or RNA transcription. Although alklylation of protein and nuclear material can influence cell fate, the direct consequences of adduct formation on acute and long-term corneal sequelae have not yet been addressed.

These data demonstrate a system-based approach combining ultrastructural analysis, histochemistry, and molecular evaluation that links architectural changes in
corneal structure with clinical outcomes and posits potential mechanisms for the pathogenesis of SM corneal lesions. The identification of such mechanisms is critical to designing effective therapeutic modalities that prevent the development of acute and delayed lesions and may be applicable to persistent keratopathies unrelated to SM alkylation.

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