Research Article

Topical therapeutic corneal and scleral tissue cross-linking solutions: \textit{in vitro} formaldehyde release studies using cosmetic preservatives

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Our recent tissue cross-linking studies using formaldehyde releasers (FARs) suggest that corneal and scleral tissue strengthening may be possible without using ultraviolet irradiation or epithelial removal, two requirements for the photochemical method in widespread clinical use. Thus, the present study was carried out in order to better understand these potential therapeutic solutions by studying the effects of concentration, pH, buffer, time, and tissue reactivity on formaldehyde release of these FARs. Three FARs, sodium hydroxymethyl glycinate (SMG), DMDM, and diazolidinyl urea (DAU) were studied using a chromotropic acid colorimetric FA assay. The effects of concentration, pH, and buffer were studied as well as the addition of corneal and scleral tissues. The main determinant of release was found to be dilution factor (concentration) in which maximal release was noted at the lowest concentrations studied (submillimolar). In time dependent studies, after 60 min, FA levels decreased by 38\% for SMG, 30\% for DMDM, and 19\% for DAU with corneal tissue added; and by 40\% for SMG, 40\% for DMDM, and 15\% for DAU with scleral tissue added. We conclude that concentration (dilution factor) was found to be the most important parameter governing the percent of FA released.

Introduction

Ultraviolet-A riboflavin photochemical corneal cross-linking (known as 'CXL') has created a new option in the field of corneal therapeutics. In the short period since its development, CXL has been proven both effective and safe in stabilizing patients with keratoconus (KC) [1–4] and post-LASIK keratectasia [5,6], and is becoming standard of care throughout the world and has recently received FDA approval for use in the United States. The cross-linking procedure effectively halts the progression of KC and can be accompanied by an improvement in both corneal curvature (i.e., flattening or ‘normalization’ of astigmatism) and/or visual acuity [5–9]. Furthermore, the indications are ever-expanding to include investigations into treating bullous keratopathy, corneal infections and melts, pellucid marginal degenration, and in combination with refractive surgery (either pre- or postprocedure) [10–13]. As good as it is, CXL has limitations and thus developing a topically delivered adjunct (or alternative) method of inducing corneal tissue cross-linking could improve patient care. Potential benefits include the following: (1) eliminate the need for ultraviolet irradiation, (2) self-administration, (3) more homogeneous cross-linking throughout the extent of the corneal stroma, and (4) the possibility of modulation of degree of cross-linking effect.

Ever since their discovery, cross-linking agents have sparked much interest in their applications, resulting in their use for a variety of purposes in industry and the biomedical sciences. We recently reported on the \textit{in vitro} testing of a group of potential cross-linking compounds in widespread use as chemical preservatives in cosmetics and other personal care products (PCPs) known as formaldehyde releasing agents (or FARs) [14]. FARs are a versatile class of compounds that could potentially act as an adjunct or even an
alternative to CXL treatment. These compounds have received much attention and are found in a wide variety of man-made products; mostly those from the textile industry as well as personal care products, such as toilet deodorizers, shampoos, conditioners, and plasticizers [15]. They also serve as popular and attractive starting compounds or chemical intermediates in the synthetic route of organic compounds used in industry [16]. This includes polymer chemistry, where they act as formaldehyde donors and are used in the production of resins, plastics, polyesters, and polyurethane products to cross-link compounds such as urea, melamine, phenols, and resorcinol [17]. Such widespread use of FARs (which include nitroalcohols) has naturally led to extensive study into the acute toxicity, teratogenicity, and mutagenicity/carcinogenicity profiles of these compounds, which are all favorable [15].

One of the most common uses for FARs is in cosmetics as preservatives. In the United States and some countries in Europe, DAU, and DMDM are two FARs present in the highest frequencies followed by SMG [18,19]. In addition to use as a cosmetic preservative, DMDM is known to possess antimicrobial properties against fungi, yeast, and Gram-positive and Gram-negative bacteria [20]. SMG has additional application in cleaning/washing agents, rinsing agents, and as a neutralizing agent for acids/acyclics polymers [21]. Although there are many compounds that are suitable for industrial and commercial cross-linking purposes, there are only a limited number of compounds that can be considered for use in vivo. This is due to additional concerns that would normally be irrelevant in the context of in vitro studies and include efficacy under physiologic pH and temperature, permeability, coloration, and effects on light transmission, and cell toxicity.

Using FARs commonly found in personal care products and the textile industry as cross-linking agents is justified by several reasons. First, their applications in commonly used products suggest that these compounds can be used for therapies. Second, many FARs possess antimicrobial activity, which would be useful when considering their application in live subjects, such as patients with keratoconus and post-LASIK keratectasia. Third, these compounds are commercially available and can be readily obtained. Although FARs have been studied in the past for their preservative capabilities (i.e., shelf-life) and industrial cross-linking capabilities, the specific cross-linking capability of these FARs with the intended use for in vivo tissue cross-linking, has not been explored previously.

The current study is an extension of our previous studies [14,22,23] in which we now study the factors governing the release of formaldehyde (FA), the key compound responsible for generating cross-links between collagen molecules. Using a simple, traditional colorimetric assay adapted to a 96-well plate format, we compared three commercially available FARs for their respective FA-releasing ability and reactivity with corneal tissue substrate, including sodium hydroxymethylglycinate (SMG), DMDM, and diazolidinyl urea (DAU), three potentially clinically useful compounds for in vivo use. The data presented were able to show that the release of FA is primarily dependent upon concentration and not pH dependent.

Materials and methods

Chemicals

Formaldehyde solution (36.8–38%), Diazolidinyl urea (N-hydroxymethyl-N-(1,3-di(hydroxymethyl)-2,5-dioxoimidazolidin-4-yl methyl) urea [DAU]), chromotropic acid disodium salt dehydrate (technical grade), sulfuric acid (99.99%), 1N hydrochloric acid, and sodium bicarbonate were purchased from Sigma–Aldrich Corp. (St. Louis, MO, U.S.A.). SMG was obtained from Tyger Scientific Inc. (Ewing, NJ, U.S.A.). DMDM hydantoin (DMDM) was obtained from Oakwood Products Inc. (West Columbia, SC, U.S.A.). All chemical solutions and buffers were prepared fresh using Millipore water (double distilled, deionized water, ρ = 18.2 MΩcm at 25°C) on the day of the experiment.

pH of FAR solutions

SMG, DMDM hydantoin (DMDM), and DAU solutions were dissolved using Millipore water, 0.1N HCl or 0.1M sodium bicarbonate solution to appropriate concentrations at each day of the experiment. pH of solutions was measured using a pH meter (Orion Star™ A221, Thermo Fisher Scientific Inc. Waltham, MA, U.S.A.).

Formaldehyde release assay

The standard formaldehyde solutions and the aqueous chromotropic acid solutions were freshly prepared on each day that calibration lines were determined. Formaldehyde solution (36.8–38%) was diluted to make a stock formaldehyde solution of 10 μg/ml. Stock solution was then further diluted to prepare 5, 2.5, 1.25, and 0.625 μg/ml standard formaldehyde solutions. For the solvent comparison experiment, standard solution was prepared using Millipore water, 0.1N HCl or 0.1M sodium bicarbonate solution. The basic National Institute for Occupational Safety and Health (NIOSH) procedure was followed for the determination of calibration lines for standard formaldehyde solutions and the measurement of formaldehyde release in FAR solutions with slight modifications [24]. Briefly, 200 μl of each
Table 1 Formaldehyde releasers evaluated in the present study and their known characteristics

| Name             | MW    | Structure | Mole of FA released per 1 mole of FAR | Predicted octanol/water partition coefficient, LogP | Max allowed concentration (mM) | Mutagenicity | Toxicity |
|------------------|-------|-----------|---------------------------------------|-----------------------------------------------|-------------------------------|--------------|----------|
| SMG [CAS # 70161-44-3] | 127.07 | ![Structure of SMG](image) | 1 | −1.197 [41] | 39.06 [42] | N [43] | LD50 oral, rat, 2100 mg/kg; LD50 dermal, rabbit, >2000 mg/kg [45] |
| DMDM hydantoin (DMDM) [CAS # 6440-58-0] | 188.18 | ![Structure of DMDM](image) | 2 | −2.3 [44] | 31.88 [42] | N [45] | LD50 oral, rat, 3720 mg/kg; LD50 oral, rat, >2000 mg/kg [45] |
| DAU [CAS # 78491-02-8] | 278.22 | ![Structure of DAU](image) | 4 | −5.398 ± 0.866 [46] | 17.97 [42] | N [47] | LD50 oral, rat, 2600 mg/kg; LD50 dermal, rabbit, >2000 mg/kg [48] |

standard formaldehyde solution and FAR solutions at appropriate concentration was aliquoted into clean Eppendorf tubes. To each of these solutions, 30 μl of 5% chromotropic acid solution was added and thoroughly mixed by vortexing. Then 300 μl of concentrated sulfuric acid (99.99%) was added. The mixtures were thoroughly vortexed. Tubes were then placed in a water bath (80–90°C) for 1 h. The tubes were cooled down to room temperature and vortexed before spectrophotometric determinations. Absorbance was measured at 570 nm by using a spectrophotometer (ELx800; Bio-Tek Instruments, Inc., Winooski, VT, U.S.A.).

**Time course experiments using corneal and scleral tissue added to the FAR solution**

At least 4 ml of standard formaldehyde solutions (10, 5, 2.5, 1.25, and 0.625 μg/ml), and stock FAR solutions were prepared freshly at each day of the experiment. Total 80–100 mg of porcine corneal or sclera tissue (previously frozen, purchased from Clements Food Group, Hatfield, PA, U.S.A.) was incubated with 2 ml FAR in water. At every time point (0, 20, 40, and 60 min) after tissue was in contact with the solution, 2 × 200 μl aliquots were taken and subjected to chromotropic formaldehyde release assay (refer to the previous formaldehyde release assay section). The reactions were carried out in sealed tubes and the opening of tubes was minimized in order to limit any potential FA release to the air.

**Results**

Due to their widespread use as cosmetic preservatives, their safety with regard to mutagenicity/carcinogenicity has been well studied. That being said, the safety of FAR intended for the corneal and scleral cross-linking has been limited to our initial and ongoing live rabbit studies [23,25]. Table 1 and Scheme 1 summarize the characteristics of FARs used in the present study, including the theoretical FA release, partition coefficient, max allowed concentration in cosmetics, mutagenicity, and LD50 toxicity. Initial *ex vivo* rabbit cross-linking studies suggested that SMG, DMDM, and DAU were the FARs which demonstrated the highest cross-linking efficacy and also thought to be the most clinically useful based on size and lower toxicity profiles (data not shown) [14]. Therefore, formaldehyde release experiments in the present study, evaluating pH, buffer dependency, and dilution factor, were limited to these three compounds.

**Determination of FAR pH in various solutions**

Figure 1A–C show the effect of active FAR compounds on pH in three different solutions, namely unbuffered water, 0.1N HCl and 0.1M NaHCO₃. All except for SMG in unbuffered water did not have impact on solution pH, at any concentration. SMG in unbuffered water, however, seem to have caused pH to fluctuate, especially at the lower concentration. At high concentration of SMG, pH of unbuffered water reaches as high as pH of 10. DMDM and DAU
Scheme 1. Formaldehyde release/decomposition of FAR compounds
Known/proposed chemistry of (A) SMG decomposition into formaldehyde and glycine, (B) DMDM into formaldehyde and dimethylhydantoin, and (C) DAU into formaldehyde and allantoin.

Figure 1. pH of FAR (A: SMG, B: DMDM, C: DAU) solutions measured in various solvents (unbuffered water, 0.1N HCl, and 0.1M NaHCO₃)
FAR solution was made by dissolving appropriate amount of solid in unbuffered water, 0.1N HCl, and 0.1M NaHCO₃. The pH was determined with 3 ml aliquot of each solution in a falcon tube.

were able to keep the unbuffered water at around a constant pH range of 6. Unlike the other FARs, SMG is not a cyclic compound and it is this structural difference, namely the greater difficulty of a linear molecule to stabilize the negative charge (concentrated on the nitrogen atom after the liberation of FA) that may contribute to its relatively high basicity (Scheme 1).

Formaldehyde release assay
The amount of FA released in each solution was determined by the chromotropic acid method. Chromotropic acid is one of the reagents which reacts with FA with high selectivity [24,26]. Though, the chemistry of the reaction of chromotropic acid with FA with sulfuric acid has not been completely solved, it is believed that the reaction is carried out in two steps. First, 2 moles of chromotropic acid reacts with 1 mole of FA to form a series of diazotized derivatives of aromatic compounds, in which para, para-quinoidal adduct was the most often quoted structure. Then the sulfuric acid oxidizes p-quinoidal adduct to mono-cationic dibenzoxanthylum, which absorbs light in the range of 570–580 nm [26].
Figure 2. Effect of concentration (dilution factor) and pH on FA release for three FAR compounds

(A) SMG, (B) DMDM, and (C) DAU. A colorimetric FA assay, chromotropic acid assay was used for the determination of free formaldehyde in FAR solutions. For SMG, concentrations ranging from 0.075 to 1 mM were assayed in three different solutions, a very acidic 0.1N HCl (pH = 2), a 100 mM NaHCO₃ solution maintaining a pH close to 8.5, and an unbuffered solution made up in H₂O. For DAU and DMDM, concentrations ranging from 0.025 to 1 mM were assayed in the three different solutions. The amount of formaldehyde released into solution at time = 10 min for each compound was expressed as a percentage of that predicted, based on the theoretical complete release of formaldehyde. (A) SMG releases 1 mole/mole, (B) DMDM releases 2 moles/mole and (C) DAU releases 4 moles/mole.

**Formaldehyde release by SMG in various solutions (Figure 2)**

SMG is an amine based formaldehyde releaser, and 1 mol of SMG releases 1 mol of FA and results in decomposition into glycine. At a concentration lower than 0.3 mM, SMG decomposes into FA and glycine at any pH.

Unlike the study by Emeis et al., our test was done at a lower concentration and we did not observe complete dissociation in 1 mM, the highest concentration we tested. At 1 mM, the equilibrium was reached and only 50% of FA was released. Buffer types/pH did not affect the release percentage. This could be clinically relevant as SMG has known storage and shelf life properties based on monitoring of a very concentrated solution.

**FA release by DMDM and DAU in various solutions (Figure 2)**

DMDM and DAU are also known to reliably decompose and we have found DMDM and DAU to release FA in water without any addition of bicarbonate buffer. According to the structures of DMDM and DAU, DMDM can release 2 moles of FA and DAU can release 4 moles of FA. Our FA assay showed that DMDM at 0.5 mM concentration released 48% of its releasable FA in water, 49% in HCl, and 45% in NaHCO₃. When DMDM was further diluted to 0.1 mM or below, DMDM seemed to reach complete dissociation. When DAU was dissolved to 1 mM concentration in water, it only released less than 20% of releasable FA and reached equilibrium. At 0.25 mM, DAU released 38% (in water), 42% (in HCl) and 43% (in NaHCO₃) of releasable FA. The highest % of free FA (ca. 60%) was detected when DAU was diluted to 0.1 mM or below. There was no significant difference in the release % in different solutions.

**Time course experiments using corneal and scleral tissue added to the FAR solution (Figures 3 and 4)**

In the final set of experiments, we evaluated changes in FA levels when corneal tissue substrate is introduced into the mixture, similar to that which would occur in vivo. Porcine corneal and scleral tissues (previously frozen, cut pieces) were utilized as substrates and incubated with FAR solution up to 60 min. At several time points, aliquots were taken from the reaction mixture and subjected to the formaldehyde release assay. The results are shown in Figures 3 and 4. The level of FA at 0 min represents the level of FA in the solution before the substrate was added. At 20 min of SMG 0.2mM incubated with tissue substrates, the level of free FA in the solution was reduced by approximately 20% compared with control (SMG solution without tissue substrate), and the level of free FA in the solution continued to decrease for the time period studied. Similar trends to SMG 0.2 mM were obtained using DMDM at 0.1 mM, which theoretically can release 0.2 mM of FA. When incubated with corneal substrate, the amount of free FA released from DMDM decreased from 86 to 56% within an hour, while the free FA of the control solution without tissue substrate did not change. When incubated with scleral substrate, the amount of free FA released from DMDM decreased from 83 to 43%. When 0.05 mM DAU was incubated with tissue substrates, free FA level decreased from 61 to 41% in a sample incubated with corneal tissue and from 43 to 28% with scleral tissue. Slight differences between corneal versus scleral tissue experiments could be attributed to the differences in reactivity between the tissue samples. It should be
Figure 3. Change in FA level was measured over 0–60 min at various FAR concentrations incubated with porcine corneal tissue (40–50 mg/ml).

FAR concentrations used were as follows: (A) SMG at 0.2 mM, (B) DMDM at 0.1 mM, (C) DAU at 0.05 mM, and (D) FA at 0.2 mM. All solutions were made in unbuffered water. At \( t = 0, 20, 40, 60 \) min, an aliquot was taken from the reaction mixture, and free FA in the solution was measured using the chromotropic acid assay. FA levels decreased in a time dependent manner. The levels at 60 min decreased by (A) 38% for SMG, (B) 30% for DMDM, (C) 19% for DAU, and (D) 32% for FA. The difference in levels between tissue containing solutions and control at 60 min were all statistically significant at \( P < 0.05 \).

Remembered that because DMDM (1:2) and DAU (1:4) produce greater amounts of FA per mole than SMG (1:1), the lower percentage differences noted for DMDM and DAU as compared with SMG (Figures 3 and 4), do not necessarily reflect a lower amount of FA consumed by the tissue substrate. In other words, for example, a 40% decrease in SMG (1:1) would be equivalent to 20% decrease in DMDM (1:2), when considering absolute amounts of FA. Finally, FA levels of FAR solutions without tissue substrate added did not change over the course of 60 min. This suggests that the tissue substrate added provided reactive sites for FA. Such FA protein reactions are well known to occur and are discussed in greater detail below.

Discussion

Our previously reported screening for more effective, yet safe, chemical agents that can induce tissue cross-linking has yielded several potential candidates, including SMG, DMDM, and DAU [14]. Thus, due to their demonstrated efficacy in inducing tissue cross-linking in the cornea and sclera in that previous study, we chose to focus our analysis of FA release on these same compounds, utilizing a simple colorimetric assay for quantitation. There are other, more sophisticated methods for measuring formaldehyde and following the decomposition of FARs such as \(^1\)H and \(^13\)C NMR. However, specialized analytical instrumentation is necessary and may not be available to many labs, particularly in institutions with more limited resources. Also, when the dilution is the crucial factor for the FA release and
Figure 4. Change in FA level was measured over 0–60 min at various FAR concentrations incubated with porcine scleral tissue (40–50 mg/ml).

FAR concentrations used were as follows: (A) SMG at 0.2 mM, (B) DMDM at 0.1 mM, (C) DAU at 0.05 mM, and (D) FA at 0.2 mM. All solutions were made in unbuffered water. At $t=0, 5, 20, 40, 60$ min, an aliquot was taken from the reaction mixture, and free FA in the solution was measured using the chromotropic acid assay. FA levels decreased in a time dependent manner. The levels at 60 min decreased by (A) 40% for SMG, (B) 40% for DMDM, (C) 15% for DAU, and (D) 11% for FA. The difference in levels between tissue containing solutions and control at 60 min were all statistically significant at $P < 0.05$.

when it is required to work with such diluted solutions, NMR measurement is not feasible. Thus, we believe that this is a good starting point for trying to understand how the FARs may function in in vivo systems. Consistent with prior literature, concentration (or dilution) was determined to be the primary factor governing formaldehyde release [27]. The equilibrium of FA for these three compounds has been studied and reported previously using $^{13}$C NMR by Emeis et al (2007). In that study, although the concentration ranges tested were significantly higher than those reported in our study, the authors found that concentration was a major factor which drives the release of FA. The reason for studying this concentration range in our study (Figure 2), which is much lower than those reported by Emeis et al., was in order to evaluate FA release at concentrations that would be expected intrastromally to result from topical application of a ‘max allowed’ concentration. In other words, since it is known that approximately 1% or less of a topically applied eye drop will reach the stroma [28,29], this is the concentration range that we will be producing intrastromally. For example, for SMG, a 40 mM ( = 0.5% max allowed) eye drop will allow for approximately 0.4 mM intrastromal concentrations. For DMDM, the levels would be approximately 32 mM ( =0.6% max allowed) for a 0.32 mM intrastromal concentration and for DAU, the level would be approximately 18 mM ( =0.5% max allowed) for a 0.18 mM intrastromal concentration.
DAU is an allantoin derivative, where allantoin reacts with four equivalents of formaldehyde under basic conditions to form the parent compound [19]. Dilution encourages the decomposition reaction, overcoming possible steric interference, and facilitating the separation of the formaldehyde moiety from the mother compound. Lehmann et al. demonstrated that DAU exists as a mixture of isomers, with 'compound BHU' (1-(3,4-bis-hydroxymethyl-2,5-dioxo-imidazolidin-4-yl)-1,3-bis-hydroxymethyl-urea) as the dominant form (30–40%) [30]. It is hypothesized that the remainder consists of many polymers of allantoin-formaldehyde condensation products. It is conceivable that the impurity of the DAU sample could result in inconsistent reaction thermodynamics and kinetics.

Regarding FA release, DAU was reported for two concentrations by Emeis et al., approximately 18 mM and approximately 7 mM. Higher levels of free FA occurred at the highest pH 9 (compared with pH 6 and pH 4). As well, higher FA was detected at the lower concentration (7 mM), once again consistent with a concentration dependent release effect. Of note, the maximum amount of free FA detected was approximately 2 × of the starting material which is significantly less than the expected amount based on DAU’s chemical structure, which suggest that 4 moles of FA can be theoretically released from each mole of DAU [31]. This lower than expected release was also noted in our study (Figure 2C). One explanation for the low amount of free FA release from DAU is that the experimental environment in the first two sets of experiments was in a 'closed system' in that there is no substrate available for FA modification. Thus, the reaction would be approaching equilibrium during which time it is possible that the liberated FA can react with other reaction products of the FAR decomposition. This includes the reverse reaction to regenerate starting material, resulting in a lower quantity of free formaldehyde. This fact notwithstanding, DAU consistently released the most formaldehyde out of the selected FARs of the present study, since it possesses a much higher molar formaldehyde releasing capacity than the other FARs, DMDM, and SMG.

DMDM is a hydantoin with a theoretical yield of two FA moieties. DMDM has one of the higher maximum allowed concentrations of the FARs used in the present study (0.6%). Regarding FA release, DMDM is a smaller size molecule and has a theoretical yield of two moles of FA per mole of DMDM. For DMDM at concentrations from approximately 3 mM to approximately 1.3 M a more alkaline pH (8.5–9 as compared with pH 6–6.5 and pH 4–4.5) and a lower concentration favored higher levels of free FA, consistent with the release characteristics of DAU, which is also increased at lower concentrations [31]. Once FA is liberated from either of the nitrogen atoms of the five-membered ring, the resulting negative charge on the nitrogen atom is delocalized into the π-system provided by both of the adjacent carbonyl moieties. The formation of intramolecular hydrogen bonds between local DMDM molecules stabilizes any additional negative charge.

Another low molecular weight FAR, SMG, demonstrates similar behavior to DMDM with regards to near total theoretical formaldehyde release. SMG has a theoretical yield of one mole of FA per mole of SMG. In spite of this lower ratio when compared with the other FARs of the present study, SMG appears to release FA more readily than DAU and behaved differently from the other FARs in certain regards. Solutions of SMG in water tend to be highly alkaline (Figure 1) but can be modulated downward with the addition of bicarbonate buffer (approximately pH 9.2) which buffers maximally in the pH 9.2–10.8 range. The solution pH can be lowered significantly further into the acidic range using 0.1N HCl without having an effect on the formaldehyde yield (Figure 2) which is in some contrast with the findings from Emeis et al. that indicate SMG releases more formaldehyde under acidic conditions. The degradation of SMG seems to be favored compared with other compounds that reaches equilibrium. Further studies are required to elucidate the details of this trend and can include 1H NMR, 13C NMR, and HPLC applications.

The reactions of free FA and related preparations (i.e., paraformaldehyde) have been studied in depth owing to their importance as tissue cross-linking agents for histologic purposes. Known reactivity of formaldehyde with tissue for histology are primarily with proteins and avoid carbohydrates, lipids, and nucleic acids, which are trapped in cross-linked matrix and are not chemically altered unless fixation is prolonged [32]. Reaction with DNA nucleotide bases such as adenine and cytosine have been reported to occur. However, the reactions require high concentration of free FA and relatively long incubation times, making them less likely to be relevant in our types of reactions [33].

Several reactions have been shown to occur between formaldehyde and free amino acids, and include reaction with lysine, asparagine, threonine, cysteine, histidine, and tryptophan [34]. In addition, stable intermolecular cross-links can form between the sulfydryl group of cysteines and side chains of lysine, arginine, asparagine, and glutamine. In reactions with proteins, formaldehyde reactions with several different proteins have been studied and include bovine serum albumin (BSA) [35], insulin [36], and cottonseed protein [37], for example. Of these, BSA has been studied the most extensively. Both acid-labile and acid-stable products can form and include hydroxymethyl adducts and methylene bridges between combinations of lysine, arginine, asparagine, glutamine, and tyrosine. Reactions with insulin have been studied by Metz et al. (2006) and indicated reactivity with lysine, arginine, and tyrosine. Similarly, reactions with cottonseed protein involve primarily lysine and tyrosine [37]. The reactions with collagen and...
formaldehyde have been reported to take place at ε-amino groups of lysine residues [38], and further form cross-links with the nitrogen atom of a peptide linkage [34,39,40]. These studies, while helpful in understanding the chemical moieties that may be involved in the FAR TXL reactions, do not necessarily predict the types of cross-links and protein modification that may actually be induced by FARs since the formaldehyde is likely to be released locally and at significantly lower concentration than using preparations of free formaldehyde and its polymers.

Although no attempt was made in the present study to identify specific reaction sites or products of the reactions (either cross-link or non-cross-link adducts), lysines and hydroxylysines, are likely sites since these residues are widely involved in collagen cross-linking and non-cross-linking reactions, both enzymatic (lysyl oxidase mediated) and non-enzymatic (i.e., non-enzymatic glycation, non-enzymatic nitration, and oxidations).

Competing Interests
The authors declare that there are no competing interests associated with the manuscript. Patent pending through Columbia University.

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Author Contribution
A.T. designed, performed the experiments, analyzed the data, and wrote the manuscript. K.C. performed the experiments, analyzed the data, and wrote the manuscript. E.M.O. performed the experiment and analyzed the data. T.N. contributed to the analysis of the result and proofreading of the manuscript. D.C.P. conceived the study, interpreted the results, and wrote the manuscript.

Abbreviations
BSA, bovine serum albumin; CXL, corneal cross-linking; DAU, diazolidinyl urea; DMDM, DMDM hydantoin; FA, formaldehyde; FAR, FA releasing agent; KC, keratoconus; PCP, personal care product; SMG, sodium hydroxymethylglycinate; TXL, tissue cross-linking.

References
1 Wollensak, G., Spoerl, E. and Seiler, T. (2003) Riboflavin/ultraviolet-a-induced collagen crosslinking for the treatment of keratoconus. Am. J. Ophthalmol. 135, 620–627, https://doi.org/10.1016/S0002-9394(02)02220-1
2 Phillips, J.R., Khalaj, M. and McBrien, N.A. (2000) Induced myopia associated with increased scleral creep in chick and tree shrew eyes. Invest. Ophthalmol. Vis. Sci. 41, 2028–2034
3 Caporossi, A., Baiocchi, S., Mazzotta, C., Traversi, C. and Caporossi, T. (2006) Parasurgical therapy for keratoconus by riboflavin-ultraviolet type A rays induced cross-linking of corneal collagen: preliminary refractive results in an Italian study. J. Cataract Refract. Surg. 32, 837–845, https://doi.org/10.1016/j.jcrs.2006.01.091
4 Wollensak, G. (2006) Crosslinking treatment of progressive keratoconus: new hope. Curr. Opin. Ophthalmol. 17, 356–360, https://doi.org/10.1097/01.ioc.0000233954.86723.25
5 Hersh, P.S., Greenstein, S.A. and Fry, K.L. (2011) Corneal collagen crosslinking for keratoconus and corneal ectasia: One-year results. J. Cataract Refract. Surg. 37, 149–160, https://doi.org/10.1016/j.jcrs.2010.07.030
6 Hafezi, F., Kanellopoulos, J., Wiltfang, R. and Seiler, T. (2007) Corneal collagen crosslinking with riboflavin and ultraviolet A to treat induced keratectasia after laser in situ keratomileusis. J. Cataract Refract. Surg. 33, 2035–2040, https://doi.org/10.1016/j.jcrs.2007.07.028
7 Caporossi, A., Baiocchi, S., Mazzotta, C., Traversi, C. and Caporossi, T. (2006) Parasurgical therapy for keratoconus by riboflavin-ultraviolet type A rays induced cross-linking of corneal collagen: preliminary refractive results in an Italian study. J. Cataract Refract. Surg. 32, 837–845, https://doi.org/10.1016/j.jcrs.2006.01.091
8 Wittig-Silva, C., Hoyer, A., Espinosa, E. and Pillunat, L.E. (2008) Collagen crosslinking with riboflavin and ultraviolet-A light in keratoconus: long-term results. J. Cataract Refract. Surg. 34, 796–801, https://doi.org/10.1016/j.jcrs.2007.12.039
9 Menge, K.M. and Hayes, S. (2013) Corneal cross-linking - a review. Ophthalmic Physiol. Opt. 33, 78–93, https://doi.org/10.1111/opo.12032
10 Al-Sabai, N., Koppen, C. and Tassignon, M.J. (2010) UVA/riboflavin crosslinking as treatment for corneal melting. Bull. Soc. Belge Ophtalmol. 13–17
11 Vinciguerra, P., Albe, E., Tranza, S. et al. (2009) Refractive, topographic, tomographic, and aberrometric analysis of keratoconic eyes undergoing corneal cross-linking. Ophthalmology 116, 369–378, https://doi.org/10.1016/j.ophtha.2008.09.048
12 Makdouni, K., Mortensen, J. and Crafoord, S. (2010) Infectious keratitis treated with corneal crosslinking. Cornea 29, 1353–1358, https://doi.org/10.1097/ICO.0b013e3181d2de91
13 Babar, N., Kim, M., Cao, K. et al. (2015) Cosmetic preservatives as therapeutic corneal and scleral tissue cross-linking agents. Invest. Ophthalmol. Vis. Sci. 56, 1274–1282, https://doi.org/10.1167/iovs.14-16035
