TEMPO-Oxidized Sacchachitin Nanofibers (TOSCNFs) Combined with Platelet-Rich Plasma (PRP) for Management of Dry Eye Syndrome

Hong-Liang Lin¹
Ting-Huan Wu²
Hsiu-O Ho²
Fang-Ching Chao²
Meng-Huang Wu³,⁴
Der-Zen Liu⁵
Ling-Chun Chen⁶
Ming-Thau Sheu²

¹School of Pharmacy, College of Pharmacy, Kaohsiung Medical University, Kaohsiung, Taiwan, Republic of China; ²School of Pharmacy, College of Pharmacy, Taipei Medical University, Taipei, Taiwan, Republic of China; ³Department of Orthopedics, Taipei Medical University Hospital, Taipei, Taiwan, Republic of China; ⁴Department of Orthopedics, College of Medicine, Taipei Medical University, Taipei, Taiwan, Republic of China; ⁵Graduate Institute of Biomedical Materials and Tissue Engineering, College of Biomedical Engineering, Taipei Medical University, Taipei, Taiwan, Republic of China; ⁶Department of Biotechnology and Pharmaceutical Technology, Yuanpei University of Medical Technology, Hsinchu, Taiwan, Republic of China

Introduction: In this study, the combination of TEMPO-oxidized sacchachitin nanofibers (TOSCNFs) with chitosan-activated platelet-rich plasma (cPRP) was evaluated for remedying dry eye syndrome (DES).

Methods: TOSCNFs, designated T050SC, were generated. T050SC combined with chitosan-activated (cPRP) was formulated as eye drops for application for severe DES. To evaluate the effects of cPRP and TOSCNFs on the repair of corneal injury, in vitro studies were conducted using Statens Seruminstitut rabbit corneal (SIRC) epithelial cells for cell proliferation and cell migration assays, and a severe DES animal model using rabbits was established with benzalkonium chloride (BAC) treatment for the evaluation.

Results: Results showed that the optimal eye formulation contained PRP activated by 350 μg/mL of the low-molecular-weight chitosan group (L3) combined with 300 μg/mL T050SC (L3+T050SC). In the WST-1 cell-proliferation assay, L3 and L3+T050SC significantly increased Statens SIRC cell proliferation after 24 hrs of incubation. In the SIRC cell migration assay, the L3+T050SC group showed a wound-healing efficiency of 89% after 24-hr treatment. After 5 days of treatment, Schirmer’s test results did not simulate the dry eye animal model. Typical cornea appearance and eye fluorescein staining results showed that the L3 group had the best effect on improving cornea haze and epithelial damage.

Conclusion: This study has determined that TOSCNFs effectively promoted the healing effect on severe cases of corneal damage, and also might enhance the clinical application and medical potential of PRP in ophthalmology.

Keywords: TEMPO-oxidation, sacchachitin, PRP, dry eye syndrome, nanofibers

Introduction: Sacchachitin (SC) is a water-insoluble dry residue from Ganoderma tsugae produced by digesting with KOH and then bleaching with NaClO.¹ SC is composed of approximately 40% chitin and 60% β-1,3-glucan.¹ Several potential applications of these two polysaccharides have been discussed previously. Wound healing¹–³ and eye cornal regeneration⁴ for SC have been studied and reported. A promising wound healing has been demonstrated for an SC film composed of 10–50-μm microfibers formed by filtration followed by dry heat. For eye corneal regeneration, enhancement of corneal epithelial wound healing was observed with application of nanogel formed from micronized sacchachitin (mSC). However, the mechanical micronization of ball milling was only capable to disintegrate neighboring sacchachitin into microfibrils since SC
was toughly organized together via H-bonding among hydroxyl groups or tangibly intertwined by single-chain SC.

In a previous study, two kinds of SC nanofibers (SCNFs) designated as MDSCNFs and TOSCNFs, respectively, were produced by either mechanical disintegration or oxidation with various concentrations (mmole/g SC) of NaClO. In a diabetic wound-healing study, both SCN5 (mechanically fragmented SCNFs with NanoLyzer® at 20,000 psi for five cycles) and T050SC (TEMPO-oxidized SCNFs produced with 5.0 mmole NaClO/g SC) embedded in 2-acrylamide-2-methylpropane sulfonate (AMPS)-based wound dressings were showed to accelerate diabetic wound healing to effectively the same as normal skin tissues. T050SC/H furthermore fostered the growth of sweat glands and hair follicles in the healed wound enlightening that the wound had healed as functional tissue. It was concluded that TEMPO-oxidized SCNFs showed appreciably well-improved healing of epithelial tissues. Most crucially being considered to use as a scaffold, TOSCNFs can be suspended in water resulting in the formation of hydrogels that offered an optimal 3D construct for cell growth and differentiation. These might be potentially applicable as a gel form for treating corneal disorders of the ocular surface caused by dry eye syndrome (DES).

DES is a syndrome of the tear film triggered by disturbances in the composition and quantity of tears. It might be attributed to 1) inadequacy of the aqueous phase of the tear, as in Sjögren’s syndrome or lacrimal gland disease; 2) an imperfection of the mucin layer instigated by a vitamin A deficiency, trachoma, diptheric keratoconjunctivitis, mucocutaneous disorders, and certain topical medications; or 3) abnormalities of the lipid tear layer caused by blepharitis and rosacea. Recently, Avila et al reported that platelet-rich plasma (PRP) was recommended to be a beneficial approach for healing several disorders of the ocular surface, including corneal ulcers and persistent ocular epithelial defects. It was also exhibited that PRP has an antiapoptotic effect on corneal stromal cells, and autologous platelets were instilled to manage macular holes. Autologous platelets in the form of eye-drops or as a plasma gel have additionally been used to manage several eye ill conditions depending on the desired effect and the purpose of the clinical intervention. In this way, PRP has shown great usefulness in the management of several ocular diseases.

Thus, it was thought that because they possess better epithelial healing and hydrogel formation ability, TEMPO-oxidized sacchachitin nanofibers (TOSCNFs) combined with PRP would be synergistically beneficial for enhancing tissue-regenerating effects. Therefore, the purpose of this study was to evaluate the use of eye drops composed of TOSCNFs and autologous PRP for treating DES. To evaluate the effects of PRP and TOSCNFs on the repair of corneal injuries, in vitro studies were conducted using Statens Seruminstitut rabbit corneal (SIRC) epithelial cells for cell proliferation and cell migration assays and a severe DES animal model using rabbits was established with benzalkonium chloride (BAC) treatment for the evaluation.

Materials and Methods

Materials

Ganoderma tsugae dry residue (G) (Kang Jian, Nantou, Taiwan), sodium hydroxide pellets (Macron, Radnor, PA, USA), potassium hydroxide (Fluka, Mexico City, Mexico), calcium chloride (Nacalai Tesque, Kyoto, Japan), sodium chloride (80%, Alfa Aesar, Ward Hill, MA, USA), sodium hypochlorite solution (5% available chlorine, J.T. Baker, Pittsburgh, PA, USA), ethanol (Sigma-Aldrich, St. Louis, MO, USA), hydrochloride acid (Fluka), 2,2,6,6-tetramethyl-1-piperidinyloxy (TEMPO) (98%, Alfa Aesar), potassium bromide for the IR spectrum (Fluka), citric acid anhydrous (Riedel-de Haen, Seelze, Germany), acetic acid (Sigma-Aldrich), glycerol (Merek, Billerica, MA, USA), sodium bromide crystals (J.T. Baker), low-molecular-weight chitosan (Sigma Aldrich), PEG1000 (Sigma-Aldrich), N-acetyl-L-cysteine bioXtra (Sigma-Aldrich), L-ascorbic acid 2-phosphate sesquimagnesium salt hydrate (Sigma-Aldrich), minimum essential media (MEM) (1x, Gibco, Waltham, MA, USA), minimum essential medium-alpha (MEM-α) (1x, Gibco), fetal bovine serum (FBS) (Gibco, InvivoGen, Waltham, MA, USA), phosphate-buffered saline (PBS) (10x, Corning, Corning, NY, USA), WST-1 reagent (Roche, Mannheim, Germany), antibiotic antifungal solution (100x) (Corning), plasmocin (Gibco, InvivoGen), DMSO (Corning), and thiazolyl blue tetrazolium bromide (MTT reagent) (Alfa Aesar) were used as received without further purification. All water used was purified with the Milli-Q plus water purification system (Millipore, Billerica, MA, USA).

Methods

Fabrication of TEMPO-Oxidized SCNFs

TEMPO-oxidized SCNFs were fabricated following a report by Chao et al. In brief, ganoderma dry residues (50 g) were processed at 85°C for 24 hrs with 400 mL 1
N NaOH solution. After treatment, the fibrous residue was gathered and rinsed with deionized water to remove any remaining NaOH. H₂O₂ at an one-third weight ratio to the residue was then used for bleaching. After decolorizing, the fibrous pulp was cleaned thoroughly with double-distilled (dd)H₂O. The fibrous pulp, so-called SC, was assembled by filtration then lyophilized. For oxidation by TEMPO, 5 g of SC was suspended in 375 mL ddH₂O comprising 0.0625 g TEMPO and 0.625 g sodium bromide. The TEMPO-mediated oxidation of the SC slurry was started by adding 5% NaClO (aq.) at an NaClO amount of 5.00 mmol/g SC to the slurry mixture with continuous stirring at room temperature. During reaction, pH 10 was preserved by adding 5 N NaOH and 1 N HCl for 2 hrs or until no alkali solution was consumed. Thereafter, oxidation was stopped by adding 5 mL ethanol followed by adding 200 mL acetone to precipitate the TEMPO-oxidized SC. The TEMPO-oxidized SC fibrous slurry was then dialyzed with OrDial® D80 (MWCO: 6000–8000) for 7 days in ddH₂O at room temperature. The dialyzed slurry was further homogenized with high-pressure homogenizer (NanoLyzer® N-2) under 20,000 psi for one cycle, and then lyophilized. SC oxidized with TEMPO followed the conditions as described above was designated as T050SC.

Physical and Chemical Characterizations of TEMPO-Oxidized SCNFs

Physical characteristics including fourier transform infrared (FT-IR) spectra, 13C cross-polarized magic-angle spinning (CP-MAS) NMR spectra, X-ray diffraction (XRD) spectra, determination of the size and zeta Potential (ζ), evaluation of gel-forming properties, transmission electronic microscopy (TEM), scanning electronic microscopy (SEM), water-retention ability, were measured followed the same procedure as that reported by Chao et al. The carboxylate and aldehyde contents of T050SC were determined according to the same method as that reported by Chao et al.

Preparation of PRP from Rabbit Blood

Blood (9 mL) was collected from the marginal ear vein of New Zealand white dutch rabbits. Whole blood was loaded into a blood-collecting tube coated with anticoagulant citrate dextrose solution A (ACD-A), then centrifuged at 2500 rpm (720 g) for 15 mins. After centrifugation, the blood was separated into three layers. We collected the middle layer which contained the platelet-poor plasma for a second centrifugation. The second centrifugation was carried at 4800 rpm (2650 g) for 15 min, and then the PRP was at the bottom of the tube. We diluted the PRP to a final concentration of 4.75 x 10⁶ platelets/µL for further application. Low-molecular-weight chitosan (135 µg) was added to each milliliter of PRP for activation just before its usage.

Evaluation of the Effects of PRP Combined with TEMPO-Oxidized SCNFs on Cell Proliferation and Cell Migration of SIRC Fibroblasts

SIRC fibroblasts were used in the same culture conditions as described above. Six different treatments were used to evaluate proliferation including PRP, low-molecular-weight chitosan-activated PRP (LPRP), 300 µg/mL T050SC combined with PRP, and 300 µg/mL T050SC combined with LPRP. Cell proliferation was determined at 72 hrs with the WST-1 reagent. The WST-1 assay was the same as described above.

SIRC fibroblasts (purchased from BCRC, Bioresource Collection and Research Center) were used in the same culture condition as described above. Before adding treatment medium, cells were scraped out of a straight line with a 200-µL pipette tip at the bottom of each well. Then, four different treatments were used to evaluate cell migration including PRP, 300 µg/mL T050SC, 5% PRP, 5% LPRP, and 300 µg/mL T050SC combined with 5% PRP. Photos were taken of cells under a microscope at 0, 6, 18, and 24 hrs to determine cell migration.

Rabbit Model of Severe DES

Captive of all animals was conducted in keeping with a protocol (no. LAC-2017-0152, 2017-06-07) approved by the Laboratory Animal Center of Taipei Medical University, and all experiments were implemented in accordance with the Guidelines for the Care and Use of Laboratory Animals. Animals were accommodated under a 12-hr light-dark cycle with free admittance to water and food. The rabbit model of severe DES was modified from that reported by Tseng et al. DES was induced in New Zealand white Dutch rabbits aged 15 weeks by 20 µL 0.1% benzalkonium chloride three times a day for 3 weeks, and then benzalkonium chloride was increased to 0.2% for a week. After the DES model was successfully induced, four different treatments including 1% PBS, LPRP, T050SC, and 300 µg/mL T050SC combined with LPRP were applied twice a day for 5 days.

To evaluate the healing outcomes, we used corneal fluorescein staining. Rabbits were anesthetized with...
1~1.5 mL Zoletil+Xylazine (Roumpun) (5 mg/kg+5 mg/kg), and laid flat on an operating table. Corneal fluorescein staining was conducted using fluorescein sodium ophthalmic strips (Fluoro Touch, New Delhi, India). Strips were first wetted with PBS and placed on a rabbit’s cornea. After staining, the corneal topography was evaluated by cobalt-blue light ophthalmoscopy (Topcon Medical Systems, Oakland, NJ, USA).

Histological Analysis
On day 5, treated rabbit corneas were sampled and fixed in a 10% (w/v) formaldehyde in aqueous phosphate buffer solution. Collected samples were then dehydrated with a gradient series of ethanol and implanted within paraffin. Embedded tissue blocks were sectioned and stained with hematoxylin and eosin (H&E). Histological images of all stained samples were filmed underneath of optical microscopy (BX41, Olympus, Tokyo, Japan).

Statistical Analysis
All physical and chemical measurements were presented as the mean ± standard deviation (SD). All in vitro and in vivo studies are presented as the mean ± standard error of the mean (SEM). Data were compared among treatment groups using one-way Analysis of Variance (ANOVA), and \( p<0.05 \) was regarded as statistical significance.

Results
Physical and Chemical Characterization of TEMPO-Oxidized SCNFs
SCNFs obtained by TEMPO oxidation with 5.0 mmole/g of SC were designated as T050SC. Several physical characteristics including FTIR, CP-MASS \(^{13}\)C ssNMR, and X-ray powder diffraction were measured and were reported in a previous publication.\(^5\) Those results are summarized as follows. Both FTIR spectra of T050SC and SCNF (Figure 1A) showed representative cellulose or chitin peaks, ie, O–H, C–H, and C–O stretching vibrations at 3400, 2900, and 1060 cm\(^{-1}\), respectively. After oxidation, T050SC appeared a sharp peak at 1720 cm\(^{-1}\) which was C=O stretching, while the moderate C-H stretching peak at 2920 cm\(^{-1}\) had decreased. Both peak changes were attributed to the transformation of -CH\(_2\)OH to -COOH at C6 in the T050SC structure. Figure 1B shows \(^{13}\)C cross-polarized magic-angle spinning (CP-MAS) NMR spectra for T050SC. Since T050SC was mainly composed of \(\beta\)-1,3-glucan and chitin, it showed carbon signal for C1, C2, C3, C4, C5, and C6 located at 103, 62, 74, 87, 77, and 68 ppm, respectively. Basically, the carbonyl group of acetyl amide was presented in chitin structure showing a carbon peak for C8 centered at 23 ppm. Further, it was expectable that the signal intensity of the C6 peak should decrease accordingly with oxidation of the hydroxyl group.

Figure 1 Physical characterization of mechanically nanosized saccharachitin (SC) and chemically TEMPO-oxidized sacchachitin nanofibers (T050SC). (A) FT-IR spectrum, (B) \(^{13}\)C ssNMR, (C) X-ray powder diffraction, (D) Water retention.
on C6. On the other hand, a new peak at 176 ppm was appeared resulting from transformation of primary alcoholic group (-CH₂OH) into a carbonyl group (-C=O) by TEMPO oxidation. These alterations additionally confirmed the conversion of the C6 primary hydroxyl group to a carboxylate group. There was also a slight change in the C5 peak. C5 exhibited an upfield chemical shift from 77 to 74 ppm, which overlapped with the C3 peak. This upfield shift might have been triggered by the newly created carboxylate group based on observation from the 3D structure.

As illustrated in Figure 1C for the X-ray powder diffraction analysis, T050SC nanofibers possessed an amorphous form with a lower crystallinity as well. There appeared two main peaks at 2θ = 6.2° and 20.8°. The amorphous form might be due to the existence of T050SC fiber size in the nanoscale range, since as small as nanosize of the fiber lengths was expected to offer a more-amorphous structure. This amorphous structure also facilitated the resuspension of lyophilized fibers. Acid–base titration method was employed to determine the carboxylate and aldehyde contents in T050SC to be 1.803 and 0.119 mmol/g, respectively. With the formation extent of aldehyde/g SC less than 1 mmol, it warranted that TEMPO oxidation had minor concerns of potential toxicity. The size and SD for T050SC nanofibers were observed to be 539.2 and 49.3 nm, respectively, while its zeta potential was measured to be −51.1 mV. This more negativity than that for SC nanofibers without TEMPO oxidation also definitely verified that the C6 primary hydroxyl group had been transformed into a carboxylate group. With the contribution by greater hydration abilities with the existence of hydrophilic carboxylate groups on the surface of these nanofibers, T050SC had water-retention abilities of 48.48 ± 2.54 g water retention/g film as shown in Figure 1D.

Figure 2A shows that immediately after dispensing 2% (w/v) T050SC in water, T050SC was viewed like wet fibrous hydrogels. Thirty minutes after stirring, T050SC displayed a gel form monitored by the vial-inversion method. However, the gel-forming character of TS50SC vanished at the 60-min time point, and it never happened again thereafter. The results indicated that the transformation of C6 hydroxyl group into carboxylic group led to negatively charged nanofibril surfaces, which allowed the formation of homogeneous electrostatically stabilized aqueous colloidal suspensions. In TEM as shown in Figure 2B, T050SC had fiber lengths of 825.54 ± 109.18 nm and widths of 27.87 ± 5.04 nm. T050SC had less-tangled fibers due to its higher zeta potential of −51.6 mV. With a higher negative charge on the surfaces of nanofibers, the nanofibers of T050SC displayed better separation and dispersion. In the SEM morphology as shown in Figure 2C, T050SC nanofibers existed as highly spongy structure, not only on the surface of the lyophilized sponge film but also on its cross-section.

In vitro and in vivo Regeneration Studies for DES

The T050SC dressing was used to evaluate chronic wound healing for DES based on previous chemical and physical evaluations. In the WST-1 assay of SIRC fibroblast cells, cell viability in the T050SC groups was first evaluated. T050SC was demonstrated to be highly biocompatible at all concentrations as shown in Figure 3A. However, the higher the concentration, the more viscous the solution was, and we also considered the transparency which are two important properties that should be evaluated in developing eye drop products. We picked 300 µg/mL for further experiments. T050SC was combined with PRP or L3 to assess cell proliferation, and results illustrated in Figure 3B show that the PRP and PRP+T050SC groups did not produce quite as promising cell-proliferation effects. This possibly indicates that T050SC had less effect of activating platelets to release growth factors to promote cell proliferation. Combining L3, which is activated-PRP with chitosan (135 µg/mL), with T050CS showed promising cell-proliferation effects. L3 and L3+T050SC showed over 150% cell viability after 24 hrs. However, L3 had the highest cell proliferative effect at around 180% ± 3.75% cell viability. After 72 hrs of incubation, L3 and L3+T050SC dropped to 134.86% ± 12.87% and 142.16% ± 7.86% cell viability, respectively. In comparison between 24 and 72 hrs, cell viabilities of L3 and L3+T050SC dropped 25% and 13%, respectively. L3 showed a greater decrease indicating that T050SC can provide a more-viscous formulation for the sustained release of growth factors. In other words, TOSCNF formulations can work as plant-based scaffolds in this system and provide a sustained-release effect of growth factors released by activated PRP. Sustained release can prolong the effect of the formulation which can decrease the drug administration time for higher convenience and patient compliance. Since L3+T050SC showed the most
promising effect on cell proliferation, it was selected to perform subsequent experiments.

In the migration assay, we evaluated formulations which promoted SIRC cell migration for 24 hrs, and results of photo imaging of migration and quantification of gap closure are shown in Figure 3C and D, respectively. PRP, L3, and L3+T050SC all showed a significant difference in gap closure compared to the control group at the same time point. In contrast, T050SC alone did not show a significant difference with the control. In other words, T050SC alone had a limited effect on promoting cell migration. In previous studies, multiple growth factors from PRP were reported to promote cell migration. PRP, L3, and L3+T050SC all showed significant differences with the control group in the first 6 hrs, and these significant differences persisted until 24 hrs. In other words, growth factors in PRP can promote cell migration and showed a significant difference at 6 hrs. However, L3+T050SC showed the most outstanding cell migratory effect among all groups. Comparing L3+T050SC to L3, one can see that L3+T050SC had a smaller cell gap at the late time point of 24 hrs, which may have resulted from T050SC providing sustained release of growth factors. L3+T050SC promoted cell
migration, and 89% of the wound had healed, which was the best result in this experiment.

In an in vivo study, we applied L3+T050SC to severe DES, which was generated in 15-week-old New Zealand white dutch rabbits by applying 0.1% BAC t.i.d. for 3 weeks and 0.2% BAC t.i.d. for 1 week. Figure 4A shows that healthy rabbit eyes had a transparent cornea and no corneal fluorescein staining. After BAC induction, the cornea became cloudy, and one can also see fluorescein staining of the cornea. In severe DES, the cornea becomes cloudy, and the cornea can also be damaged, which will affect the visual ability of the patient. On day 0 of treatment, one can also see the cornea status of all groups with severe DES. After treatments with L3 and L3+T050SC, the cornea had again become transparent, and the corneal ulcer was also relieved. In contrast, in the control and T050SC groups, the cornea remained non-transparent, and the ulcer could still be observed. Corneal fluorescence staining as illustrated in Figure 4B showed no staining in the T050SC, L3, and L3+T050SC groups after 5 days of treatment. In other words, the corneal damage of these three groups had been relieved. Under cobalt-blue light ophthalmoscopy, one can see that the L3 group had the best optical properties compared to the others. However, the L3+T050SC groups showed a better outcome of corneal ulcer relief than L3.

Histological images of corneal severity are shown in Figure 4C, and qualification and quantification evaluations of the corneal severity level after 5 days of treatment are illustrated in Table 1. Results showed that there were about three to five epithelial layers properly aligned and well organized on the surface of the cornea, which showed a more-purple stain under H&E staining in the healthy rabbit eye cornea. The central thickness of the cornea was 49.2 ± 2.2 µm. After BAC induction, epithelial cells of the eye cornea exhibited a poor arrangement, and the central thickness of the cornea dropped to 12.3 ± 1.9 µm. Furthermore, some epithelial cell necrosis and stromal swelling were observed. T050SC produced better recovery than the control, and the central thickness of the cornea
Figure 4 (A) Preoperative and postoperative appearances of the rabbit cornea. All groups presented a severe corneal ulceration appearance on day 0. After 5 days of treatment with T033SC, L3, and L3+T050SC, photos were collected as postoperative appearances with a digital camera. (B) Eye corneas were stained with fluorescein, and photos were collected under cobalt-blue light ophthalmoscopy. (C) Histological analysis of normal rabbit cornea of untreated, and T005SC, L3, and L3+T050SC-treated corneas on day 5. Scale bars, 125 µm.
was 45.1 ± 1.2 µm, which was similar to the thickness of the normal cornea. The arrangement of epithelial cells became more orderly, and no stromal swelling was observed. However, some eosinophil cell infiltration was observed, and Bowmen’s layer became swollen, which may have been linked to the reason that the T050SC group had poor corneal transparency. The L3 and L3+T050SC groups showed promising corneal recovery after 5 days of treatment. Epithelial cells were well arranged, and the central thicknesses of the corneas were 43.7 ± 1.1 and 40.3 ± 2.3 µm, respectively. However, the L3+T050SC group appeared to still be undergoing remodeling, as there were more fibroblast cells present in the upper part of the stroma and epithelial layer. The results indicated that T050SC and the growth factors secreted by PRP can repair severe DES of the cornea. However, L3 showed the best overall outcome. This may have resulted from the rapid release of growth factors, which play important roles in the early stage of healing of severe DES, whereas L3+T050SC instead exhibited sustained release. However, T050SC can act as a water-retention agent that provides a moist environment for DES and may result in better outcomes in the late stage of the healing process.

Discussion

SCNFs were prepared by chemical nanonization with TEMPO-oxidation in this study. The T050SC produced by chemical nanonization had a size of 539.2 nm with an SD of 49.35 nm and a zeta potential of −51.6 mV. The vial-inversion method showed that 2% of T050SC nanofibers could form hydrogel after initial hydration and so formed hydrogel liquefied in water only after 30 min stirring. Probably, due to the gel-forming character with a viscous 3D network after hydration, T050SC nanofibers offered sustain release of growth factors from activated PRP.

In the in vitro study examining chronic epithelial wound-healing regeneration to relieve DES, nanofibers of T050SC acted as plant-based scaffolds to provide a sustained-release effect of growth factors from activated PRP (L3). The sustained release of such factors can prolong the effect of the formulation, which can decrease the frequency of drug administration, thereby promoting higher convenience and patient compliance. The combination of T050SC and L3 showed the greatest enhancement in cell proliferation and promotion of cell migration leading to 89% healing of the wound. In the in vivo study of applying L3+T050SC to severe DES, L3+T050SC showed a better outcome of corneal ulcer relief than L3 alone. In the histological evaluation of rabbit eye corneas, results indicated that T050SC and growth factors secreted by PRP could repair corneas with severe DES. However, L3 produced the best overall outcomes. This may have resulted from the rapid release of growth factors that play important roles in the early stage of healing of severe DES, while L3+T050SC exhibited sustained release instead. However, T050SC can act as a water-retention agent that provides a moist environment for DES and may result in better outcomes in the later stages of the healing process.

Conclusions

TEMPO-oxidized SCNFs possessing a gel-forming character with a 3D network were developed as a functional scaffold with ideal characteristics. PRP combined with TEMPO-oxidized SCNFs designated as T050SC has pronounced potential applications in severe dry eye syndrome. These SCNFs combined with activated PRP also possesses conceivable medical practices for any epithelial cell disorders similar to severe dry eye syndrome. Furthermore, the multifunctional biomaterial based on SCNFs combined with activated PRP also might have additional clinical applications such as hair regeneration in the future. However, the most challenge issue in practical applications of SCNFs combined with activated PRP is the stability of released various growth factors in the hydrogel.

Table 1 H&E Staining of Corneal Severity Levels After 5 Days of Treatment

<table>
<thead>
<tr>
<th>Center Corneal Severity Level</th>
<th>Normal</th>
<th>Control</th>
<th>T050SC</th>
<th>L3</th>
<th>L3+T050SC</th>
</tr>
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<tbody>
<tr>
<td>Corneal staining</td>
<td>None</td>
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<td>None</td>
<td>None</td>
<td>None</td>
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<tr>
<td>Center epithelial thickness (n=2–4)</td>
<td>49.2±2.2 µm</td>
<td>12.3±1.9 µm</td>
<td>45.1±1.2 µm</td>
<td>43.7±1.1 µm</td>
<td>40.3±2.3 µm</td>
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<tr>
<td>Epithelial damage</td>
<td>None</td>
<td>Severe</td>
<td>Moderate</td>
<td>None</td>
<td>Moderate</td>
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<tr>
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Disclosure
The authors report no conflicts of interest in this work.

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