Multi-Layered Polyamide/Collagen Scaffolds with Topical Sustained Release of N-Acetylcysteine for Promoting Wound Healing

Background: Impaired wound healing might be associated with many issues, especially over-active of reactive oxygen species (ROS), deficiency of blood vessels and immature of epidermis. N-acetylcysteine (NAC), as an antioxidant, could solve these problems by inhibiting overreactive of ROS, promoting revascularization and accelerating re-epithelialization. How to deliver NAC in situ with a controllable releasing speed still remain a challenge.

Materials and Methods: In this study, we combined collagen (Col) with N-acetylcysteine to perform the characteristics of sustained release and chemically crosslinked Col/NAC composite with polyamide (PA) nanofibers to enhance the mechanical property of collagen and fabricated this multi-layered scaffold (PA-Col/NAC scaffold). The physical properties of the scaffolds such as surface characteristics, water absorption and tensile modulus were tested. Meanwhile, the ability to promote wound healing in vitro and in vivo were investigated.

Results: These scaffolds were porous and performed great water absorption. The PA-Col/NAC scaffold could sustainably release NAC for at least 14 days. After cell implantation, PA-Col/NAC scaffold showed better cell proliferation and cell migration than the other groups. In vivo, PA-Col/NAC scaffolds could promote wound healing best among all the groups.

Conclusion: The multi-layered scaffolds could obviously accelerate the process of wound healing and exert better and prolonged effects.

Keywords: polyamide, sustained release, N-acetylcysteine, wound healing, multi-layered structure

Introduction

Impaired skin regeneration is a common outcome in patients with diabetic ulcers, burns and pressure ulcers. Among the clinical classical wound treatments, topical therapies such as covering the wound with dressings has been widely used. Over the past decades, many composite synthetic polymers and biological materials have been used to fabricate wound dressing agents.

The wound healing process is extremely complex and is regulated by interactions among several factors, all of which promote the reconstruction of a skin barrier. The reconstruction process is impaired by events such as hematoma, overreactive inflammation (reactive oxygen species) and limited angiogenesis. N-acetylcysteine (NAC) is an antioxidant approved by the Food and Drug Administration for clinical use in the treatment of oxidative stress-related diseases such as diabetic ulcers. Previous studies have reported that NAC promotes wound healing by alleviating the adverse effects of ROS. NAC accelerates this process by...
improving perfusion and angiogenesis at ischemic areas.\textsuperscript{8} NAC has also been reported to facilitate maturation of epidermis through the promotion of MMP-1 expression via the PKC/Stat3 signaling pathway.\textsuperscript{9} NAC has been widely used to treat lung diseases such as chronic obstructive pulmonary disease and hepatic damage, however, rarely is it applied for wound healing.\textsuperscript{10,11} Thus, whether topical administration of NAC promotes the wound healing process has not been sufficiently studied. For this reason, further studies are required to explore its safety and mechanisms.\textsuperscript{12} Besides, chronic wounds that take long to heal may require repeated administration of NAC. This repeated administration might be feasible, but not always the best treatment strategy.\textsuperscript{13} So, the best strategy is to develop a new kind of wound dressing with the ability to achieve the long-term drug release. This strategy could decrease the frequency of changing the wound dressing and relieve the pain of patients.\textsuperscript{14}

To study the topical effects of NAC and its sustained release characteristics, collagen, which has been widely used for tissue engineering and soft tissue repair, was used as the drug carrier.\textsuperscript{15} Collagen has been associated with low antigenicity and excellent biocompatibility with most endogenous tissues. Therefore, collagen-based wound dressings are used for chronic wound and ulcers treatment.\textsuperscript{16–18} Collagen can be crosslinked with drugs to create a sustained release formulation. This ensures that there is sufficient concentration of the drug on the wound during the long healing process.\textsuperscript{19} Unfortunately, collagen is mechanically weak and prone to tear, hence, cannot be applied to wounds in some parts of the body such as joints. Although collagen can be cross-linked to enhance its mechanical properties, it is still sub-optimal to be used as an abrasion-resistant wound dressing.\textsuperscript{20}

Consequently, in order to overcome the limitations associated with collagen, polyamide (PA) was chosen in this study to strengthen the mechanical properties of drug-loaded collagen. Polyamide was chosen because of the following reasons: i) Polyamide, is a biocompatible material, which is widely used in the clinical treatment of wounds. ii) Polyamide is the most common biomaterial used as a suture in cutaneous surgery.\textsuperscript{21} Previous studies have reported the use of polyamides in the preparation of membranes by electrospinning technique for wound healing application.\textsuperscript{22,23}

In this study, collagen was crosslinked with NAC to achieve sustained release of NAC and then fabricated into triple-layered PA-Col/NAC scaffolds, which consisted of PA nanofibers at the core and the collagen on both sides. This multi-layered structure made use of the mechanical properties of PA nanofibers and improved the stability of the entire scaffolds. Electron microscopy, FTIR spectroscopy and Raman spectroscopy were used to observe the surface structure and characteristics of the scaffolds. The physical properties such as tensile modulus and water absorption were also measured. The migration and proliferation of NIH 3T3 cells were tested after the cells were implanted on the scaffolds to examine the effects of the scaffolds on the activity of fibroblasts. The scaffolds were implanted in Sprague Dawley (SD) rats with a dorsal wound to evaluate their properties in the promotion of wound healing. The full thickness of the skin at the wound area was excised and processed with HE, Masson and CD31 staining to investigate wound closure after 14 days of implantation.

**Materials and Methods**

**Preparation of the PA-Col/NAC Scaffolds**

Figure 1 shows the general preparation procedure of the PA-Col/NAC scaffold. Firstly, polyamide 6,6 (Sigma-Aldrich, America) were dissolved in a mixture of dichloromethane (DCM, Fisher Chemical, America) and N, N-dimethylformamide (DMF, Fisher Chemical, America) in a 4:1 volume ratio and used to prepare a 15% PA solution. Using the PA solution, the PA nanofibers were electrospun (voltage: 18 kV, needle: 20 gauge, speed: 1 mL/h). This was followed by repeated washing (5–6 times) of the PA nanofibers in 75% alcohol (Fisher Chemical, America), deionized water and phosphate buffer saline (PBS, Fisher Chemical, America) to remove all solvent residues. Collagen was dissolved in 0.05 mmol/l acetic acid to prepare a 14% collagen solution. The collagen solution was added into a petri dish with a diameter of 60 mm and a height of 2–3 mm. The processed PA nanofibers (3 cm * 3 cm square) were put on this solution and the petri dish was filled with a similar height of the collagen solution. After lyophilization (40kPa, –20°C), the PA-Col scaffolds were obtained and placed in NAC-EDC/NHS solution, which indicates a mixture of 50 mmol/L 1-(3-Dimethylaminoethyl)-3-ethylcarbodiimide (EDC, Sigma-Aldrich, America) and 13.5 mmol/L N-Hydroxysuccinimide (NHS, Sigma-Aldrich, America) in 75% alcohol solution containing NAC at a concentration of 0.5 g/L. Crosslinking was then performed at room temperature for 24 hrs. Finally, the materials were rinsed and sterilized to obtain PA-Col/NAC scaffolds.
Morphology of the PA-Col/NAC Scaffold
Scanning electron microscopy (JSM-IT300, JEOL, Japan) set at an accelerating voltage of 10 kV was used to observe the microstructure of the scaffolds. The surface of the scaffolds was sprayed with platinum using a sputter coater prior to examination. ATR-FTIR spectroscopy and Raman spectroscopy in VERTEX 70 (Bruker, German) and LabRAM HR800 (HORIBA JobinYvon, French) were used to record the surface characteristics of the scaffolds. FTIR measurements were carried out at a resolution of 4 cm⁻¹ in the frequency range of 4000–500 cm⁻¹ and the Raman spectrum was measured with a scanning range of 50–4000 cm⁻¹. The excitation source was a diode laser with a wavelength of 532 nm adjusted to a power of 30 mW. All the data were analyzed by using the software Origin.

Water Absorption of the PA-Col/NAC Scaffold
The water absorption property in vitro was examined. To mimic the areas around the wound and for convenient observations, 0.5 mg/L alkannin PBS solution was prepared and added to the scaffolds. To study the water absorption capacity of the scaffolds, the following formula was used:

$$\text{Water absorption (\%)} = \frac{M_s - M_i}{M_i}$$

where, “Mi” represents the initial weight of the samples. “Ms” represents the weight of samples after being immersed in the PBS.²⁴

Mechanical Properties of the PA-Col/NAC Scaffold
All-electric dynamic test instrument (Instron, British) was used to measure the mechanical properties of the scaffolds. Tensile tests were performed according to the specifications of ASTM D638. The load cell and extension speeds were 100 N and 5 mm/min, respectively. The OriginPro Software (OriginLab, America) was used to calculate Young’s modulus and plot the stress-strain curve.

NIH 3T3 in vitro Culture
NIH 3T3 fibroblasts (ATCC, USA) were used to perform the cytological experiments. Briefly, the cells were cultured at 37°C in an atmosphere with 95% air/5% CO₂ using Dulbecco’s modified Eagle’s medium (DMEM, Thermo Fisher Scientific, America) containing 10% fetal bovine serum (FBS, Thermo Fisher Scientific, America)
and 1% gentamycin/streptomycin (Thermo Fisher Scientific, America).

**In vitro Cell Viability of the PA-Col/NAC Scaffolds**

NIH 3T3 cells were cultured on different samples to test the in vitro cell viability. The scaffolds were sterilized, cut into 5 mm * 5 mm slides and placed in 24-well plates. The NIH 3T3 cells were prepared at a density of $1.0 \times 10^4$ cells per mL and 100 µL of the cell suspension was added to each well. The fluid in each well was removed at 1, 3, 5 and 7 days. Then a mixture of 100 µL complete medium and 10µL CCK-8 was added to each well and incubated for 4 hrs at 37 °C. The supernatant was then transferred into each well of a 96-well plate and absorbance was measured at 450 nm using an absorbance microplate reader (BioTek ELx800, USA).

**Cell Adhesion and Compatibility of the PA-Col/NAC Scaffolds**

NIH 3T3 cells were implanted on the slices made from the scaffolds and cultured for 1, 3 and 7 days to examine the cell adhesion and compatibility of the scaffolds. The NIH 3T3 fibroblasts were stained with fluorescein diacetate/propidium iodide (FDA/PI) (Sigma-Aldrich, America) and then observed using a confocal laser microscope (Leica Microsystems, Germany).

**Cell Migration of the PA-Col/NAC Scaffolds**

NIH 3T3 fibroblasts were seeded in 24-well plates at a density of $0.5 \times 10^6$ cells/well and cultured until they reached a 90% confluence. Next, the migration of cells on the scaffold was evaluated. The fluid in each well was removed and replaced with a different extract liquor (a complete medium in which the scaffolds were immersed for 24 hrs). Scratches were made on the cell surface layers using a pipette tip. The cell images in each well were captured using an optical microscope (Olympus CKX41, Japan) at time 0, 8, 16 and 24 hrs during the cell migration experiment.

**Drug Delivery Release of the PA-Col/NAC Scaffold**

HPLC was used to test the drug release profile of the scaffolds and the absorbance of the solution at different time-points (1 hr, 2 hrs, 4 hrs, 8 hrs, 24 hrs, 48 hrs, 7 days and 14 days) was measured. At each time-point, 2 mL of the solution was taken from each well and added into a bottle filled with phosphate buffered solution (PBS) and scaffolds. Each well was replaced with 2 mL of a fresh PBS solution. The concentration of drug was detected via absolute assessment with HPLC. Briefly, we measured the standard concentration and confirm the characteristic peak around 5 mins. (Figure below) Then we calculated the areas of characteristic peak and established simple unary linear regression between the standard concentration and peak area. According to this unary function, we calculated the concentration of NAC at different time points. The Origin software was used to plot the drug release curve for each scaffold.

**In vivo Implantation of the PA-Col/NAC Scaffold**

All procedures were performed in accordance with the guidelines approved by the Animal Ethical Committee of Huazhong University of Science and Technology. Eight adult SD rats (male, age: 8–10 weeks; weight: 180–220g) were used for this in vivo implantation tests. All operations were performed by the same surgeon in a laminar clean bench under anesthesia (2% isoflurane inhalation). The sterilization of the scaffolds was done according to the ISO 11137 standard requirements. The hair was shaved, and the skin was disinfected and then three equal ovals (major axis = 2 cm, minor axis = 1 cm) full-thickness skin excisions were made on the dorsal skin via the wound model producer fabricated by 3D printing. Subsequently, the wound area was treated with different scaffolds and gross images were recorded at different time-points.

**Histological Analysis**

The animals were sacrificed 14 days after implantation by CO₂ asphyxiation. The wound together with the surrounding skin was excised and fixed in 10% formaldehyde. The tissues were stained with Hematoxylin, Eosin and Masson’s Trichrome (Sigma-Aldrich, America) to examine the epidermis, collagen and formation of new tissues. The new vessels formed during the wound healing were evaluated by CD31 staining (Sigma-Aldrich, America). The samples on the rats performed experiment were sectioned along the longest axis. Because the shapes of samples were not regular, two or three sections per specimen were evaluated. Ten sections were randomly selected and calculated for further histological analysis.
Statistical Analyses
Statistical analysis was performed using GraphPad Software (USA). Data are shown as the mean ± standard deviation. Parametric Student’s t-test and Analysis of Variance (ANOVA) with a post hoc test by Tukey’s method were used to determine the significance levels. The *** and **** refer to the P-value of <0.05, <0.01 and <0.001, respectively.

Results
Physical Characterization of the Scaffolds
Figure 2A and B shows the porous structure of the PA-Col/NAC scaffold. The multi-layered structure, “collagen-PA nanofibers-collagen” composite, is shown in Figure 2C and D. So, it could be inferred that the surface structure of PA-Col/NAC scaffold was porous, and the spatial structure of collagen and PA nanofibers accumulated layer by layer. The ATR-FTIR spectrum and Raman spectrum analysis results of the PA-Col/NAC scaffolds are shown in Figure 2E and F. It can be observed that PA had a characteristic C-H peak at 960 cm⁻¹ and 930 cm⁻¹.²⁸ PA-Col and PA-Col/NAC exhibited the >NH and -OH peaks at 3300 cm⁻¹, amide I, amide II and amide III peaks at 1657 cm⁻¹, 1528 cm⁻¹ and 1300 cm⁻¹, which were ascribed to the presence of collagen in the scaffolds.²⁹,³⁰ In the PA, the amide peaks were obtained at the wavenumber mentioned previously. The FTIR spectrum revealed the NAC peaks as shown in Figure 2F. The PA-Col/NAC scaffold exhibited characteristic V(CH₃) (symmetric) and the V(CH₃) (asymmetric) peak at 2940 cm⁻¹.³¹

Figure 2G shows the tensile stress-strain curve of different groups under the tensile loading. The three groups reached the ultimate tensile stress at the same ultimate strain of about 16.42%±0.41%. Before the ultimate tensile stress, the three curves were linear. The Young’s modulus of PA, PA-Col and PA-Col/NAC group was calculated as follows: 20.34±2.22MPa, 8.44±1.20MPa and 7.88±1.90MPa, respectively. The Young’s modulus of PA-Col and PA-Col/NAC

Figure 2
(A and C) Macro photograph of PA-Col/NAC scaffolds from vertical view and from the side view (scale bar 1 cm). (B and D) Magnified view of A and C (scale bar 100 nm). (E) ATR-FTIR spectroscopy, (F) Raman spectroscopy and (G) tensile stress-tensile modulus curve of the scaffolds was displayed. (H–J) Water absorption of three scaffolds indicated by stained normal saline (scale bar 1 cm). (K) Water absorption histogram of three scaffolds.
Abbreviation: N.S., not significant.
scaffold was lower than that of PA scaffold, indicating that PA-Col/NAC and PA-Col scaffolds were much softer than the PA scaffold and hence more suitable for wound dressing.

Water absorption capacity is one of the important factors affecting drug release. Besides, it is important that wound dressings possess the ability to absorb blood, necrotic tissues and exudate, which is beneficial to the prevention of infection. Figure 2H–J shows the condition at which the scaffolds absorbed a liquid similar to the body fluid. According to this results, PA-Col and PA-Col/NAC scaffolds performed better and had higher water absorption capacity than the PA nanofibers. To further examine this property of the scaffolds, Figure 2K was plotted to reveal the water absorption capacity of each scaffold. This plot indicated that the water absorption of PA-Col and PA-Col/NAC was superior to that of PA nanofibers, and this was ascribed to the high water absorption capacity of collagen. However, there was no significant difference between PA-Col and PA-Col/NAC scaffold, implying that the NAC or the process of crosslinking did not influence the water absorption property of the scaffolds.

The Cell Biocompatibility and Cell Migration of the PA-Col/NAC Scaffolds in vitro

In this study, NIH-3T3 cell types were selected for testing the effects of the scaffolds on wound healing in vitro, as fibroblasts were the mainly cell types involving in process of wound healing. The biocompatibility of each group was inferred from the results of FDA/PI staining and CCK8 assay. Figure 3A shows that there was no significant difference among the three groups on day 1 and 3. However, on the 7th day, the density of the live cells (marked in green) on the PA-Col and PA-Col/NAC samples was much higher than that of the PA group. This could be due to good cell biocompatibility or water absorption property of collagen. The number of live cells in PA-Col/NAC group was higher than that in PA-Col group whereas the number of dead cells (marked in red) in PA-Col/NAC group was higher than that in PA-Col group, and this was associated with the effects of NAC.

The results of CCK8 assays were also consistent with those indicated in Figure 3A. Figure 3B shows that there was no significant difference among the three groups in the 1st and 3rd day. However, on the 7th day, the absorbance of PA-Col and PA-Col/NAC at 450 nm was higher than that of PA, whereas there was no significant difference between PA-Col and PA-Col/NAC. This indicated that NAC had no detectable effect on cell proliferation, and it did not induce cell toxicity.

The general cell migration trend for the three groups is shown in Figure 4A. The closure area was calculated using the Image-Pro software and results are shown in Figure 4B. The analysis revealed that the percent of wound closure area of the PA-Col/NAC group (65.93±3.91% at the 16th hour and 87.18±2.98% at the 24th hour) was higher than that of PA-Col group (61.47%±11.29% at the 16th hour and 69.27%±8.87% at the

Figure 3 (A) Confocal fluorescent images for FDA/PI staining of NIH 3T3 fibroblasts cultured on the scaffolds for 1, 3 and 7 days (scale bars 200 μm). (B) The CCK8 assay of NIH 3T3 fibroblasts seeded onto the scaffolds (mean ± SD, *P<0.05, **P<0.01).

Abbreviation: N.S., not significant.
24th hour) and PA group (35.53%±13.99% at the 16th hour and 39.86%±15.97% at the 24th hour). To further examine the wound closure rate, the wound width-time curve (Figure 4C) was plotted. Linear regression was used to examine the average rate of wound healing and calculate the rate of closure. The rate of closure of PA-Col/NAC group 26.87 μm/h ($r^2=0.96$) was higher than that of PA group 19.18 μm/h ($r^2=0.84$) and PA-Col group 22.93 μm/h ($r^2=0.87$).

Sustained Release of PA-COL/NAC Scaffolds

The drug release profiles of the scaffolds are as shown in Figure 5 and the process of sustained release was divided into two parts. For the first 24 hrs, we could infer that most of the NAC were released for the two scaffolds and the cumulative release percentage of PA-Col/NAC (54.53%±7.45%) was higher than that of Col-NAC (35.75%±0.84%) in Figure 5B. There was a sustained drug release from day 7 to day 14. Moreover, the cumulative drug released from PA-Col/NAC (58.56%±8.77%) was higher than that of Col-NAC (40.41%±0.49%). This indicated that the PA-Col/NAC scaffolds displayed a good drug sustained release profile.

In vivo Evaluation of the Therapeutic Effects of PA-Col/NAC on Rat Wound Model

The wound model producer fabricated by 3D printing method was used to create three oval (major axis=2 cm, minor axis = 1 cm) full-thickness wounds on the dorsa of the rat. These were then covered with the three types of scaffolds. After implantation, the dorsal area was covered with an air-permeable film to prevent the scaffolds from falling from the wound. The macro views of the wound at different time-points are shown in Figure 6A. To further analyze the percentage of wound area, the curve shown in Figure 6B was plotted. The percentage wound area for the three scaffolds before day 3 was almost similar. However, the percent of wound closure for PA-Col/NAC (69.3%±11.47%) was higher than that of PA (37.95%±10.36%) and PA-Col (54.18%±12.51%) on day 7. On day 14, all the rats in PA-Col/NAC group presented almost complete wound healing (86.17%±5.46%), while the percentage of

Figure 4 (A) Images from scratch experiment for three groups at different time point (scale bars 200 μm). (B) Closure area (%) of the scaffolds at different time points (*P<0.05, PA-Col vs PA-Col/NAC). (C) A scatter plot was used to display the width of the wound over time and the rate of closure (ROC) was calculated. To generate the $r^2$ value, a linear regression was run on the graph using the GraphPad Prism software.
wound closure for PA (55.15%±12.24%) and PA-Col (73.5%±7.46%) were much lower.

The results of the histological tests are shown in Figure 7. The unhealed areas in the three groups were distinguished from normal areas by examining the subcutaneous tissues. The sections between the black dash lines showed lighter color with no cutaneous appendages such as hair follicles and sebaceous glands, indicating that the process of wound healing was not complete. The plot shown in Figure 7M presents the length of the unhealed wound. The wound length for PA-Col/NAC (1757.3 μm ± 1185.2 μm) was shorter than that of PA-Col (3927.8 μm ± 1849.5 μm) and PA (5223.8 μm ± 2087.7 μm).

Formation of new epidermis reflected the status of the wound healing process. The HE staining images (Figure 7A–F) revealed that the epidermis of the PA group was discontinuous, whereas the other groups presented complete and continuous epidermis. The thickness of the new epidermis is shown in Figure 7B, D and F. To quantify this measurement, the curve shown in Figure 7N was plotted. The thickness of PA-Col/NAC (67.19 μm ±10.60 μm) was much higher than that of PA-Col (46.71 μm ± 13.40 μm) and PA (38.99 μm ± 13.59 μm).

Collagen regeneration plays an important role in the wound healing process. Figure 7G–L shows the Masson staining results of the samples indicating that the new collagen mass in PA and PA-Col at 14 days appears to be irregularly aligned, with high infiltration of inflammatory cells, and high deposition of collagen bundle in PA-Col /NAC, uniformly and regularly arranged. This phenomenon indicated that PA-Col/NAC group exhibited better wound healing than PA and PA-Col groups.

To examine the angiogenesis process in the samples, CD31 staining was performed to calculate the number of blood vessels. Figure 8A–F shows the CD31 staining results of the sample. The results shown in Figure 8G revealed that the number of vessels in PA-Col/NAC group (75.5±26.1) was higher than that of PA (49.3 ±19.7) and PA-Col (48.3±15.4).

Discussion

In this study, multi-layered PA-Col/NAC scaffolds were fabricated from PA nanofibers and collagen loaded with NAC. We found that the PA-Col/NAC scaffolds displayed excellent water absorption capacity, porosity, biocompatibility, and sustained drug release profile. Most importantly, the scaffolds promoted wound healing in SD rats.

Polyamide (PA) is a polymer with broad applications in a variety of fields. It is a commonly used biomaterial in medicine due to its good mechanical strength and excellent biocompatibility. Moreover, compared to poly (lactic acid) and polycaprolactone, PA possess certain advantages such as high mechanical strength, flexibility, rigidity and structural similarity to peptides (due to amide bonds). Although its biodegradability is not adjustable like other polyesters, it is still effective for wound dressing since this property has little influence on this process. One of the essential applications of PA is in membranes used in medicine and life science such as matrices and scaffolds. Using the electrospinning technology, PA nanofibers can be manipulated for various applications. PA nanofibers have been shown to possess three-dimensional interactive porous network structure and high specific surface area. This structure is equivalent to ECM in biological tissues, and therefore provide ideal environment for cell adhesion and proliferation. Besides, the direction and orientation of nanofibers that were feature of circumferentially or axially aligned structures might affect the
vascularization of the scaffolds. Therefore, in this study, PA was used to prepare PA electrospun nanofibers for fabrication of the PA-Col/NAC scaffold.

Collagen is one of the most common biomaterials used in wound healing. In this study, collagen was chosen due to the following: i) collagen can achieve sustained drug release via a simple chemical crosslinking method (using the NHS-EDC mixture); ii) given that PA is highly resistant to solvents and body fluids, it might negatively influence the wound healing process. Therefore, incorporating PA to biomaterials such as collagen which have high hygroscopicity remarkably transforms the hydrophobicity of PA. In this study, the water absorption capacity of PA-Col and PA-Col/NAC scaffolds were similar and higher than that of PA. iii) as the material to directly interact with the wound surface, the collagen mimics the extracellular matrix and provides favorable conditions for cell adhesion and proliferation.

NAC was used to promote wound healing in this study. As a ROS scavenger, NAC regulates excessive inflammatory response and prevents oxidative stress triggered by ROS. Feng et al leveraged on the ROS scavenging properties of NAC and anti-apoptosis effects to promote bone generation. Deniz et al demonstrated that NAC is a common potent antioxidant which prevents necrosis of stasis when applied topically. Tsai et al demonstrated that NAC promotes cell proliferation, cell migration, wound healing and collagen expression via the MMP-1 via PKC/Stat3 signaling pathway. NAC has been shown to accelerate the wound healing process via promotion of the epidermal maturity and angiogenesis. Gill et al illustrated that MMP-1 is important for the reconstruction of the basement membrane and the proliferation of epidermal cells and actin cytoskeletal reorganization all of which participate in for the regeneration of the epidermis.

In this study, we could observed that PA-Col/NAC group achieved the best repairing effects of the epidermis among the three groups. Albasanz-Puig et al reported that NAC activates Stat3 to regulate vascular endothelial growth factor (VEGF) and facilitate the growth of new vessels. This is in accordance with the results of our study in which the number of CD31 positive vessels in PA-Col/NAC group was higher than that in PA and PA-Col group.

We successfully fabricated scaffolds with multi-layered structure. This method has been widely used in the
fabrication of composites. Zhang et al prepared sandwich-like fibers/sponge composites using a similar method to prevent the recurrence of tumor and metastasis and combined it with chemotherapy and hemostasis. The scaffold carrying the gelation loaded with cisplatin produced good hemostasis and chemotherapy effects. In this study, the multi-layered structure also played an important role in the fabricated PA-Col/NAC scaffolds. The PA nanofibers enhanced the stability of the scaffolds at the wound area. The collagen loaded with NAC on each side had direct contact with the wound. Collagen provided a favorable environment for wound healing and controlled the sustained release of the NAC to the wound. NAC promoted the wound healing process by resisting ROS, accelerating the maturation of the epidermis and facilitating angiogenesis.

**Conclusion**

In conclusion, this study successfully fabricated a multi-layered scaffold for wound dressing. The scaffold consisted of PA nanofibers and collagen loaded with NAC, which promoted the wound healing process. This novel PA-Col/NAC scaffold displayed excellent water absorption capacity, mechanical properties, biocompatibility, and ability to steadily release NAC. These properties...
promoted cell proliferation and migration in vitro. The scaffold also enhanced the repair of an oval rat wound indicating a promising application in wound healing.

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**Disclosure**

The authors report no conflicts of interest in this work.

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**Figure 8** Immunohistochemical staining for CD31 showed significant increase of number of vessels in PA-Col/NAC group compared with the other groups. 

Notes: (A–F) Vascular endothelial cells that showed CD31 antibody positive were marked by black arrows. Scale bars: 200 μm. (A and B) PA group. (C and D) PA-Col group. (E and F) PA-Col/NAC group. (G) Analysis of number of CD31 positive vessels on skin tissue sections. *P<0.05.

Abbreviation: N.S., not significant.


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