Nacre-mimic Reinforced Ag@reduced Graphene Oxide-Sodium Alginate Composite Film for Wound Healing

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With the emerging of drug-resistant bacterial and fungal pathogens, there raise the interest of utilizing versatile antimicrobial biomaterials to treat the acute wound. Herein, we report the spraying mediated assembly of a bio-inspired Ag@reduced graphene-sodium alginate (AGSA) composite film for effective wound healing. The obtained film displayed lamellar microstructures similar to the typical “brick-and-mortar” structure in nacre. In this nacre-mimic structure, there are abundant interfacial interactions between nanosheets and polymeric matrix, leading to remarkable reinforcement. As a result, the tensile strength, toughness and Young’s modulus have been improved 2.8, 2.3 and 2.7 times compared with pure sodium alginate film, respectively. In the wound healing study, the AGSA film showed effective antimicrobial activities towards Pseudomonas aeruginosa, Escherichia coli and Candida albicans, demonstrating the ability of protecting wound from pathogenic microbial infections. Furthermore, in vivo experiments on rats suggested the effect of AGSA film in promoting the recovery of wound sites. According to MTT assays, hemolysis evaluation and in vivo toxicity assessment, the composite film could be applied as a bio-compatible material in vitro and in vivo. Results from this work indicated such AGSA film has promising performance for wound healing and suggested great potential for nacre-mimic biomaterials in tissue engineering applications.

Wounds on human body, as a ruptured skin and tissue, requires appropriate treatments to prevent infections and promote tissue regeneration1. In the past decades, a variety of dressing material have been developed for wound healing2-4. Among them, sodium alginate (SA) has received intensive interests as it was a natural, costless, and easily-obtained polysaccharide, showing good biocompatibility and the ability to facilitate wound healing by maintaining moist microenvironment5,6. However, the poor mechanical property and lack of function to withstand bacterial infection restrict the direct application of sodium alginate in wound healing7.

Inspired by the nature, introducing the inorganic nano-filler and fabricating “bricks-and-mortar” nacre-like microstructures became the most popular method for enhancing the mechanical properties of “soft” polymers in recent decade8-12. Graphene, one of the most popular 2D materials, has been increasingly studied as an ideal inorganic reinforcing constitutes due to their advantages over other materials in light weight, extraordinary mechanical properties and electrical conductivity, high flexibility and good ductility13-16. In addition, with high aspect ratio and superior processibility, graphene and its derivatives are also suitable candidates for growing and assembling functional components on their surface17,18. This unique characteristic could direct a feasible approach for fabricating functional graphene reinforced sodium alginate wound dressing materials.

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Traditionally, organic antibiotics were encapsulated in wound dressing materials to combat with various microbial infections. However, nowadays, the severe antimicrobial drug resistance has gradually weakened the actual anti-bacterial effect of organic medicines and threatens the public health. As effective alternatives, metal or metal oxides inorganic nanoparticles (NPs) have been increasingly reported to kill microbial pathogens or inhibit their growth effectively, thus providing a novel antimicrobial option, which could benefit the decreased usage of organic antibiotics, and in due reduce the emergence of resistance. Moreover, according to our previous work, Ag NPs decorated reduced graphene oxides (rGO) nanocomposites showed enhanced antibacterial effect and good bio-compatibility in skin treatment. These positive results prompt us to further construct Ag NPs enriched rGO/sodium alginate nacre-mimic films with enhanced mechanical properties and anti-infection effect for wound healing.

In this paper, we constructed Ag@reduced graphene oxides-sodium alginate (AGSA) composites film with nacre-mimic microstructures which displayed enhanced mechanical performance comparing with pure SA film and Ag NPs@sodium alginate film. The AGSA film has effective antimicrobial activities against Pseudomonas aeruginosa (P. aeruginosa), Escherichia coli (E. coli) and Candida albicans (C. albicans). Further in vivo investigations showed such composite film prompted wound closure on rats. Moreover, the AGSA film was biocompatible to human endothelia cells and showed low haemolytic potential according to the MTT assay and heamolysis evaluation. We envisage that this AGSA film could find potential clinic translation in antimicrobial wound healing as a promising dressing materials.

Results and Discussion
The spraying assisted assembly of AGSA composite film. Biomimetic synthesis is a fast growing and the most promising method for developing advanced materials. The “brick-and-mortar” ordered microstructures, which endow nacre superior strength, stiffness, and toughness, have encourage us to reinforce SA film mechanical properties by filling inorganic building blocks and mimicking nacre-like microstructures. As illustrated in Fig. 1, the AGSA film was assembled via a simple process including the absorption of Ag@rGO sheets with SA molecules and spraying-assist assembly, and the content of Ag@rGO in this composite was 20% (w/w). Initially, Ag@rGO sheets were coated with SA molecules which could enable the formation of basic assembly building blocks. When the suspension of SA coated Ag@rGO sheets were sprayed onto a hot substrate, the ordered lamellar microstructures were formed after water-evaporation. According to previous studies of alginate/graphene and Sodium alginate/montmorillonite bionanocomposites, in the evaporation process of nano-platelets and alginate aqueous mixtures, the wet nanocomposites form a gel state, which push the nano-platelets to arrange layer-by-layer. Moreover, in our manuscript, small drops of rGO-Ag and alginate mixture were sprayed continuously onto a hot plate, which accelerated the evaporation of water and the highly ordered arrangement of Ag@reduced graphene oxide-sodium alginate composite.

Ag@rGO composites sheets were prepared according to a facile approach reported previously by our group. Herein, as shown in Fig. 2a, mass production of this high quality composite nanosheets aqueous solution was successfully obtained at the concentration of 2 mg/mL, which could meet the demand of further fabrication of macroscale assemblies. Transmission electron microscopy (TEM) images confirmed that small Ag NPs were decorated uniformly on rGO sheets, with the size about 10 nm (Fig. 2b–d). The cross-section scanning electron microscopy (SEM) images of as-prepared films are presented in Fig. 3, and the thicknesses of these four samples (SA, ASA, GSA and AGSA films) were approximately 30 μm. Among them, SA and Ag-sodium alginate (ASA) films showed typical fracture morphology of polymers without lamellar microstructures (Fig. 3a,b,e,f). In contrast, ordered arrangement of rGO sheets can be clearly seen in graphene-sodium alginate (GSA) and AGSA films, demonstrating successful aligning and assembling of SA-coated rGO or Ag@rGO sheets during spraying and water-evaporation (Fig. 3c,d,g,h). Similar ordered microstructure have been reported in nacre-like film prepared from GO and various polymers. EDS images and X-ray diffraction (XRD) analysis also confirmed the presence and well-distribution of Ag@rGO sheets in SA matrix (Fig. 3i–l). AGSA films with 10, 20 and 40 μm thick were also prepared respectively (Fig. 4a–c), indicating the thickness of as-prepared films increased with the increasing of spraying solution amount (Fig. 4d). From the digital pictures, as-prepared AGSA film was...
Figure 2. Preparation of Ag@rGO. (a) Mass production and (b–d) TEM images of the Ag@rGO sheets for assembling AGSA film.

Figure 3. Cross-sectional SEM images of different samples: (a) SA film, (b) ASA film, (c) GSA film and (d) AGSA film; (e–h) High resolution images for (a–d), respectively; EDS analysis of cross-section of the AGSA film: (i–k) The corresponding full EDS mapping images of C, Na and Ag elements; (l) Spectrum of AGSA film.

Figure 4. The thickness and dimension of AGSA film. (a–c) Cross-sectional SEM images for 10, 20, and 40 μm AGSA films, respectively; (d) Linear correlation between thickness of AGSA films and volume of solutions for spraying (error bars are standard deviation, n = 5); (e,f) Pictures of the AGSA film; (g) XRD and (h) UV-Vis spectrums of the samples.
6.5 cm in size, displaying a black appearance with slight metal luster (Fig. 4e), and the films were flexible to withstand bending (Fig. 4f). As shown in XRD patterns of ASA and AGSA films (Fig. 4g), the typical broadened diffraction peaks at 38.2° and 44.3° were indexed as the Ag (111) and (200) planes, demonstrating the presence of small sized Ag NPs in such films. The absorption peaks around 400 nm arisen from surface plasmon resonance of colloidal silver further confirmed the presence of silver content in AGSA film (Fig. 4h). The typical transmission peaks of SA can be found in Fourier transform infrared spectroscopy (FTIR) patterns of SA, ASA, GSA and AGSA films (Supplementary Fig. 1), indicating SA serves as polymer matrix in all composite films. These results demonstrated the well-aligned AGSA composite film could be facilely fabricated by spraying method, and the thickness together with the dimension could be well controlled.

Mechanical properties. The curing of wound sites may locate at motional parts of human body, and this requires wound healing materials having adequate mechanical stability. Dramatic enhancement in tensile strength was observed in typical stress–strain curves of the prepared films with inorganic fillings (Fig. 5a). As shown in XRD patterns of ASA and AGSA films (Fig. 4g), the typical broadened diffraction peaks at 38.2° and 44.3° were indexed as the Ag (111) and (200) planes, demonstrating the presence of small sized Ag NPs in such films. The absorption peaks around 400 nm arisen from surface plasmon resonance of colloidal silver further confirmed the presence of silver content in AGSA film (Fig. 4h). The typical transmission peaks of SA can be found in Fourier transform infrared spectroscopy (FTIR) patterns of SA, ASA, GSA and AGSA films (Supplementary Fig. 1), indicating SA serves as polymer matrix in all composite films. These results demonstrated the well-aligned AGSA composite film could be facilely fabricated by spraying method, and the thickness together with the dimension could be well controlled.

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expected to have better mechanical stability during wound healing treatment. With reinforced strength, toughness and Young's modulus, this AGSA film was prompted. In Fig. 5d, the fracture morphology of AGSA film displays a typical nacre-mimic structure, which was in agreement with previous reports. Moreover, as has been reported before, the presence of nanograins on aragonite platelets in nacre can play an important role in enhancing the mechanical properties. In this case, the Ag NPs decorated on rGO sheets may act similarly to those nanograins. Through retarding the sliding of rGO sheets under tensile stress and introducing mechanically interlocking, Ag NPs contributed to the higher strength, which was in agreement with previous reports. The role of Ag NPs can also be confirmed as AGSA film has higher tensile stress and modulus than GSA film (Fig. 5). With reinforced strength, toughness and Young's modulus, this AGSA film was expected to have better mechanical stability during wound healing treatment.

**Antimicrobial activity.** The antimicrobial effects of as-prepared films were evaluated by investigating the inhibition zones. As shown in Fig. 6, inhibition zones around AGSA films were 1.30, 0.65 and 2.20 cm in diameter P. aeruginosa, E. coli and C. albicans respectively. These findings indicated that AGSA film exhibited effective protection capability against bacterial and fungal infection, especially towards P. aeruginosa which was the main concerned pathogen in wound. Pure SA, ASA and GSA films were examined to illustrate the origin of antimicrobial effect. Because the inhibition zones were absent in SA and GSA treatments but can be seen in ASA treatment (Fig. 6a), Ag NPs contributed majorly to the antimicrobial effect of AGSA. As previously reported, Ag NPs exhibited excellent inhibiting capability to microorganism. Very recently, ultrafine Ag/AgCl nanoparticles coated on graphene was reported to inhibit microbial growth by generating oxidative species under visible-light irradiation, which can further promote healing in burn wound. Herein, to illustrate the antimicrobial mechanism of Ag NPs in AGSA film, ICP analysis was conducted to investigate the leaching of Ag\(^{+}\) ions. According to Supplementary Fig. 3, the content of released silver increased with the monitoring time. After 7 days, 16.5% (w/w) of the total silver in AGSA film were released. This result was in consistent with previous conclusion regarding the role of released silver ions in antimicrobial effect of Ag NPs and Ag-based materials. In our study, the antimicrobial effect of AGSA film could be attributed to the released silver ions, which could destroy cell membrane and disturb DNA replication.

**In vitro and in vivo biological test.** The as-prepared AGSA film was composed of Ag NPs, reduced graphene oxide nanosheet and sodium alginate, and all components were biocompatible. In our previous report, no skin irritation on the epidermal laceration of rat was induced by Ag@rGO composite nanosheet. In addition, water-soluble sodium alginate is a commercially used anionic natural polymer with broad applications in biomedical area including drug delivery, tissue engineering and wound dressing. First of all, the cytotoxicity of pure SA film and obtained ASA, GSA and AGSA composite films was studied using Human Umbilical Vein Endothelial Cells (HUVEC). After 2 days exposure to these films, no inhibition of cellular viability was detected in comparison with control group (untreated cells), confirming the good cyto-compatibility of these composite films including the AGSA film (Fig. 7a). In addition, the haemolysis evaluation of as-prepared AGSA film was performed using rat whole blood. In comparison with negative control (normal saline) and positive control (distilled water), the haemolytic potential was measured to be only 0.52% (Fig. 7b). According to previous report,

![Figure 6. Results of antimicrobial investigations. (a) Pictures of the inhibition zones of different films; (b) Diameters of inhibition zones (Error bars are standard deviation; n = 3 for P. aeruginosa, n = 4 for E. coli and C. albicans; * p < 0.05, ** p < 0.01).](image-url)
the haemolytic limit was considered to be 5%, which was 10 times higher than that of our AGSA film. As a result, our AGSA film with non-cytotoxicity and such a low haemolytic potential was compatible.

In order to assess whether the release of silver ions will cause adverse effects on hepatic and renal functions, the serum biochemistry tests were carried out. The blood of five rats was collected before the wound healing experiment to serve as normal samples. 3 and 7 days after wound-cutting and treatment with AGSA dressing on the same five rats, serum samples were further collected and measured respectively. The major serum markers of liver function including aspartate transaminase (ALT), alanine transaminase (AST), albumin (ALB) and alkaline phosphatase (ALP) were evaluated, and major indexes of kidney function including creatinine (CREA) and blood urea nitrogen (BUN) were measured in the meantime. All of these measured serum parameters stayed within normal ranges and revealed no obvious liver or kidney injury compared to the conditions without skin damage (Fig. 7c–h). As a result, our AGSA film was confirmed to be cyto- and haemo-compatible, and no disturbance of hepatic or renal function was induced after exposure to this film.

**In vivo Wound healing.** Alginate dressing has been widely used for the treatment of acute surgical wounds, for example, alginate matrix could load with stromal cell-derived factor-1 and release them continuously to accelerate wound closure rates and reduce scar formation. Nowadays, various alginate dressings have been developed commercially, including Tegagen™ (3 M Healthcare), Comfeel Plus™ (Coloplast) and Kaltostat™ (ConvaTec). Herein, five rats were employed to study the wound healing efficiency of our nacre-mimic reinforced Ag@rGO-sodium alginate composite film. After disinfection of skin using 75% ethyl alcohol and iodophor disinfectant, two round skin wounds with the diameter of about 0.5 cm were cut at each side of depilated back skin of hip. AGSA film was cut into proper size to cover the wound of right side in each rat. The wounds of left side were disinfected by rubbing alcohol and iodophor, and then covered with a sterile cloth as standard treatment control. In comparison with standard treatment control, significant wound shrunk and small scar could be observed in AGSA film treated group, without induction of inflammation (Fig. 8a–e). The size of wound area was measured by vernier caliper at 3, 5 and 7 days after the wound-cutting. As shown in Fig. 8f, the wound in standard treatment control group cured for 13.8%, 39.3% and 51.5% in area after 3, 5 and 7 days, respectively. Compared with the standard treatment control group, the wound area of AGSA film treatment group decreased quickly by 19.4%, 79.5% to 92% after 3, 5 and 7 days respectively, and new epidermis was regenerated prominently. In addition, as shown in Supplementary Fig. 4, when the wounds were remained without disinfection and dressing treatment, the control group showed much slow wound healing, and swelling and inflammation in wound appeared obviously.

The wound healing progress was further evaluated by histology analysis and Hematoxylin & Eosin and Masson’s Trichrome staining. After 7 days, a large number of inflammatory cells could be observed from the wound of standard treatment control (Fig. 8g), while some collagen fibers and glandular cavity appeared on the wound treated with AGSA dressing (Fig. 8h). Moreover, Masson’s Trichrome staining was used to examined collagen fibers formation in the wounds. As shown in the Fig. 8i,j, wound in standard treatment control group had a loose reticular arrangement of collagen fibers, and the space between the collagen was relatively large. In comparison, collagen bundles were relatively compact and showed a better ordered arrangement in the AGSA dressing treated wounds. These results indicated as-prepared AGSA film obviously accelerated effect on wound healing over a short period of time, and this composite film showed a great potential for further clinical translation.

**Conclusion**

In summary, we demonstrated the assembly of AGSA composites film with nacre-like lamellar microstructures aiming at improving the performance of SA materials in wound healing. The as-prepared AGSA film showed enhanced mechanical properties compared with pure SA film. Its tensile strength, modulus and toughness were 2.8, 2.3 and 2.7 times of pure SA film, reaching up to 161.2 ± 4.6 MPa, 0.63 ± 0.01 MJ m⁻³ and 9.9 ± 0.5 GPa.
respectively. Meanwhile, AGSA film exhibited effective antimicrobial activities against both bacterial (P. aeruginosa, E. coli) and fungal pathogens (C. albicans). Importantly, MTT assay, heamolysis evaluation and in vivo toxicity assessment results suggested the promising safety for further medical applications. Furthermore, in vivo experiment on rat confirmed the effect of AGSA film in promoting the recovery of skin wound. These achievements, clearly demonstrated that our AGSA film possessed better mechanical stability and anti-infection capability than SA film, meanwhile maintained the safety in vivo. It is expected that AGSA film would have outstanding performance for wound healing and suggested a great potential for nacre-mimic biomaterials in tissue engineering applications.

Methods

Chemicals. Sodium alginate was purchased from Sangon Biotech Co. Ltd (Shanghai, China). Graphite powder, sulfuric acid (H₂SO₄), potassium permanganate (KMnO₄), phosphorus pentoxide (P₂O₅), potassium peroxydisulfate (K₂S₂O₈), hydrogen peroxide (H₂O₂), hydrochloric acid (HCl), and silver nitrate (AgNO₃) were obtained from Sinopharm Chemical Reagent Co. Ltd (Shanghai, China). Poly (sodium 4-styrenesulfonate) (PSS), (Mw = 70000) was purchased from Sigma-Aldrich.

Preparing of Ag NPs, rGO and Ag@rGO nanocomposites. Ag NPs were prepared according to Micro-wave assisted method reported by Hu et al. Graphene oxide nanosheets (GO) were synthesized from a modified Hammer’s method Ag@rGO composites were prepared according to our previous method Briefly, GO were firstly reduced to rGO using hydrazine hydrate as reductant. Then rGO-PSS solution were prepared and kept at 60°C. AgNO₃ solution was added into the rGO-PSS solution slowly with the assistance of double-jet pump.
at a speed of 0.5 ml h⁻¹. Finally, Ag@rGO nanocomposites solution (1.5 mg/mL) were obtained after washing and diluting with deionized water.

Assembly of nacre-like Ag@rGO-sodium alginate composites film. Spray assembly was employed to fabricate AGSA composites film. In a typical process for assembling AGSA film with the Ag@rGO nanocomposites water solution (pH~6.5, 1.5 mg/mL) was mixed with sodium alginate water solution (pH~6.5, 2 mg/mL) under vigorous stirring for 10 min, and then subjected to sonication for 10 min in order to completely disperse Ag@rGO. Then, the mixture was sprayed onto heated glass substrate (140°C). Finally, films were peeled off from substrate after solvent evaporated. The proportion of Ag@rGO in the film was fixed at 20%. To confirm the controllability of AGSA film, 7.5, 15 and 30 mL Ag@rGO-sodium alginate mixture solutions with the same compound proportion were sprayed into films with different thickness respectively. In addition, pure SA film, GSA film and ASA film were also prepared by similar spraying assisted process. The typical thicknesses of these composite films (SA, ASA, GSA and AGSA films) were about 30 μm.

Characterization. Scanning electron microscopy (SEM, Zeiss Supra 40) and transmission electron microscopy (TEM, Hitachi HT7700) were used to investigate the morphology and structure of the samples. Energy dispersive spectroscopy (EDS, X-Max, Oxford) was used to analyze the constituent elements of the samples. X-ray diffraction (XRD) analysis was performed with a Philips X’Pert PRO SUPER X-ray diffractometer (Cu Kα radiation). The mechanical properties of the as-prepared films were measured in tensile modern using Instron 5565 A testing machine at a load speed of 0.04 mm sec⁻¹. The as-prepared films mechanical performance in wet condition were measured immediately after spraying water mist (~5μL/s) onto the tested films for 3 s. The UV-visible (UV-Vis) spectrums of SA and AGSA films with the thickness of around 30 μm were measured by Shimadzu UV-2600 spectrometer. Inductive coupled plasma atomic emission spectrometer (ICP-AES, Perkin Elmer Optima 7300 DV) was used to determine the concentration of the released Ag⁺ ions. The mechanical properties of as-prepared films were measured using dual column electromechanical testing systems for tensile (Instron 5565 A equipped with 500 N load cells).

Antimicrobial test. P. aeruginosa (ATCC 15692), E. coli (DH5α) and C. albicans (Sc 5314) were employed to evaluate the antimicrobial effect of the samples. Suspensions of P. aeruginosa and E. coli were obtained after culturing the two strains in Luria–Bertani (LB) medium for 20 h respectively. C. albicans was cultured in Yeast Extract Peptone Dextrose (YPD) medium for 20 h. All the microbial cells suspensions were diluted to 10⁶ CFU/mL, and each 100 μL such suspensions were spread onto LB or YPD agar plates uniformly. Then, film samples in round pieces with a diameter of 10 mm were attached onto such agar plates respectively. After incubation at 37°C, 24 h for P. aeruginosa, E. coli and C. albicans respectively, antimicrobial effects of each sample were evaluated by measuring the diameters of inhibition zones.

Detection of the released silver ions. In order to investigate the release of silver ions from AGSA film, the dialysis of AGSA film was carried out using a dialysis bag (MWCO = 1000 Da, Sangon Biotech, Shanghai, China). The filtrates were collected after 1, 3, and 7 days respectively, and ICP-AES was carried out to determine the amount of silver released from AGSA film.

MTT assay. Human umbilical vein endothelial cell (HUVEC) was purchased from Anhui Medical University, which was employed to evaluate the cellular toxicity of films. Control group is normal cells without the exposure to any films, while cells in the other four groups were exposed to SA, AS, GSA and AGSA films, respectively. After exposure to UV overnight, a series of films were cut into small pieces (2 mm). HUVEC cells cultured in DMEM medium (n = 3). Then these pieces were removed, and cells were washed with PBS and incubated with MTT for 4 h. The formazan crystals in each well were formed by the living cells, which was dissolved by DMSO. Finally, the absorbance at 490 nm was measured with Multiskan FC microplate photometer (Thermo, USA) to quantitatively calculate the cell viability.

Animal experiments. According to previous procedure, haemolytic potential of the final obtained AGSA film was evaluated using whole blood of rat with distilled water and PBS solution served as positive and negative control, respectively. Five SD rats (220 g bw, male) were anesthetized by chloral hydrate (10% in normal saline, 0.3 ml/100 g bw), and the depilated back skin of hip was chosed for the wound healing test. For the safety evaluation of AGSA film on five rats, blood samples were collected through the orbital venous plexus at different time points. These tests were carried out in accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health. All protocols including hemolysis evaluation and wound healing experiment were reviewed and approved by the Institutional Animal Care and Use Committee (IACUC) of Anhui Medical University (LJSC20150134).

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Author Contributions
X.Y., F.L., and K.-D.H. contributed equally to this work. J.X., Y.L., X.Y., K.-D.H., T.H., and W.-P.X. designed the experiments. J.X. and Y.L. supervised the research. X.Y., F.L., J.X., L.D., Y.-H.S. and Y.L. synthesized the sample and performed the characterization. X.Y., J.X., X.-F.P., X.-Y.W. worked on the spray assembly of Ag/reduced graphene oxide–sodium alginate film. K.-D.H. completed the antimicrobial test. I.D. and Y.-D.W. performed the
MTT assays. X.Y. and Y.L. conducted the hemolysis evaluation. X.Y. performed the in vivo experiments on rats. All authors analyzed and discussed the results. X.Y., F.L., K.-D.H., J.X. and Y.L. wrote the paper.

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