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Multifunctional silver nanoparticle embedded eri silk cocoon scaffolds against burn wounds-associated infection

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Antimicrobial wound dressings offer enhanced efficacy compared to conventional dressing platforms by limiting bacterial infections, expediting the healing process, and creating a barrier against additional wound contamination. The use of silk derived from silkworm cocoons in wound healing applications is attributed to its exceptional characteristics. Compared to mulberry silk, sericin from non-mulberry cocoons has higher water exchange mobility and moisture retention. Eri, a non-mulberry silkworm, is an unexplored source of silk with an eco-friendly nature of production where the natural life cycle of silkworms is not disrupted, and no moths are sacrificed. This work reports on an eri silk cocoon-based scaffold decorated with silver nanoparticles as a wound dressing material effective against burn-woundassociated multiple-drug-resistant bacteria. The UV-vis spectroscopy showed maximum absorbance at 448 nm due to the surface plasmon resonance of silver nanoparticles. FT-IR spectra exhibited the functional groups in the eri silk proteins accountable for the reduction of Ag⁺ to Ag⁰ in the scaffold. SEM-EDX analysis revealed the presence of elemental silver, and XRD analysis confirmed their particle size of 5.66-8.82 nm. The wound dressing platform showed excellent thermal stability and hydrophobicity, fulfilling the criteria of a standard waterproof dressing material, and anticipating the prevention of bacterial biofilm formation in chronic wounds. The scaffold was found to be effective against both Staphylococcus aureus (MTCC 87) and Pseudomonas aeruginosa (MTCC 1688) multipledrug-resistant pathogens. Electron microscopy revealed the bacterial cell damage, suggesting its bactericidal property. The results further revealed that the scaffold was both hemocompatible and cytocompatible, suggesting its potential application in chronic wounds such as burns. As an outcome, this study presents a straightforward, cost-effective, and sustainable way of developing a multifunctional wound dressing platform, suggesting its significant therapeutic potential in clinical and biomedical sectors and facile commercialization.

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1. Introduction

The skin serves as a vital barrier in the human body, safeguarding organs from environmental variations and microbial invasion.¹ When the skin is injured, numerous complications can occur, such as microbial invasion, increased tissue wound healing, and the promotion of scar formation.^{2,3} Skin tissue has self-repair abilities; however, big cutaneous wounds are prone to bacterial infection and often do not heal properly. Skin wounds compromise the body's physical defense, making patients vulnerable to life-threatening infections caused by

necrosis, hindered epidermal tissue regeneration, delayed

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microbes.⁴ Burns are the most common and dreadful type of trauma. Based on the World Health Organization 2023 report, burns are responsible for an estimated 180 000 deaths per year, primarily in countries with low and moderate incomes.5 From 1950 to 2012, the mean total healthcare cost per burn patient was reported to be \$88218 (ranging \$704-\$717,306).6 Patients with burn injuries tend to survive, yet according to the 2015 United States National Burn Repository, invasive infection and sepsis remain among the top four leading causes of burn morbidity and mortality, accounting for 51% of the deaths.7 Multiple multidrug-resistant (MDR) bacteria now account for the bulk of these deaths.8 According to a five-year clinical report, Pseudomonas aeruginosa was the leading cause of infection (59%), followed by Staphylococcus aureus (17.9%), Acinetobacter spp. (7.2%), and *Klebsiella* spp. (3.9%).⁹ Post-burn, a competition between proliferating skin cells, e.g., keratinocytes, and colonizing microbes exists for wound dominance.¹⁰ Bacteria have the capability to produce extracellular polymeric substances that protect them against antibiotics, which diminishes their effectiveness.11 Therefore, it is crucial to develop highly effective antibacterial materials against MDR strains, in addition to screening for new antimicrobials, to check morbidity and mortality rates.12

The primary and immediate treatment of burn wounds involves topical antiseptic therapy. Antiseptics, being nonselectively toxic to all biological materials, are detrimental for systemic administration, and microbes can develop resistance against them.13 Silver agents work well against both Gram-Gram-negative bacteria, positive and despite their drawbacks.14-17 Silver nitrate or silver sulfadiazine are reported to show cytotoxicity on fibroblasts or keratinocytes, impairing re-epithelialization in burn wounds and often driving bloodrelated disorders.¹⁸ Hence, alternative strategies for microbial control are needed, where the potential of silver as an antibacterial agent remains imperishable. In this connection, silver nanoparticles (Ag NPs) offer broad-spectrum antimicrobial activity against MDR pathogens, such as Pseudomonas aeruginosa.19 The Center for Disease Control and Prevention has classified the antibiotic-resistant strains linked to it as priority infections, requiring an urgent need to discover new therapeutic strategies. Ag NPs can attack the microbes on multiple cellular targets, thus reducing their chance of becoming resistant.^{20,21} The synthesis of Ag NPs involves two approaches, *i.e.*, *in* situ and ex situ. In situ approach of formation of Ag NPs is more convenient than ex situ approaches involving specific reducing agents such as amino acid groups (e.g., carboxylate) residing in silk sericin proteins.22-24

Selecting the appropriate dressing material for a wound is crucial. The material should be able to maintain a moist environment, promote epidermal migration and angiogenesis, allow gaseous exchange, maintain tissue temperature, protect against bacterial infections, and be non-adherent, sterile, nontoxic, and non-allergic.²⁵⁻²⁷ Traditional clinical dressings like gauze, sterilized cotton, and bandages are cost-effective but provide limited wound healing and infection prevention benefits.^{28,29} Thus, to tackle these disadvantages, researchers are exploring silkworm cocoons to address the shortcomings of traditional passive dressings.³⁰⁻³²

Cocoons' shielding function is similar to how skin protects our bodies. As a result, the retention of the entire cocoon structure, which includes both fibroin and sericin, may be beneficial for the therapeutic purposes of burns.³³ Silk-fibroinbased protein materials are among the best biomaterials in tissue engineering because of their excellent biocompatibility, bioactivities, and minimal immunogenicity. For instance, dressings made from silk-fibroin can not only assist rapid hemostasis but also enhance the process of healing wounds.³⁴⁻³⁶ The cocoon structure has an antimicrobial function, which gives an added advantage for using entire cocoons.37,38 The cocoon provides a strong protective barrier for the pupa, but it does not prevent the transfer of oxygen or water vapor. Air gaps within cocoons facilitate separate multistage moisture transport processes, resulting in an effective moisture buffer effect.^{39,40} The presence of calcium oxalate hydrate crystals on the outer surface of the cocoon obstructs the flow of CO₂ from outside to inside. This indicates that these crystals play a role in capturing a significant amount of CO₂ in the form of hydrogenbonded bicarbonate on the surface. Entwined silk inside the cocoon structure serves as a second barrier, effectively blocking the flow of residual CO₂. In two extreme natural temperature regimes of 5 °C and 50 °C, the cocoons sustain temperatures from 25 °C to 34 °C, which closely resemble the human body temperature.⁴¹ Therefore, the cocoon mimics the human skin and produces the optimal environment for wound repair and regeneration by allowing preferential CO₂ gating from the internal to the exterior environment while not impeding O₂ and H₂O diffusion, maintaining the correct level of oxygen and moisture content, and promoting speedy wound recovery.42 Therefore, the microstructural and physiological characteristics of silkworm cocoon inspired the team to merge nanoscience with silk biotechnology to design advanced silver nanocomposites silk cocoon scaffolds (ASN-SCSs) using nonmulberry eri silkworm cocoons.

Eri is a domesticated non-mulberry silkworm (Philosamia ricini) under the order Lepidoptera and family Saturnüdae. Eri silkworms, unlike Bombyx mori, live in a complex external environment and require higher protection.43 The use of silk fibroin and sericin obtained from wild silkworms has significant potential in the fields of medicine and healthcare. Compared to Bombyx mori, Saturnüdae silks develop better structural transitions that are well-defined.44 Compared to mulberry silk, sericin from non-mulberry cocoons has higher water exchange mobility and a high moisture retention capability because of its high hydrophilic to hydrophobic amino acid ratio of 9.06-9.8. Additionally, non-mulberry silk fibroin has distinct arg-gly-asp (RGD) motifs with a favorable cell adhesion effect and better mechanical strength than mulberry silk fibroin.45,46 Better cell adhesion stimulates tissue growth, cell migration, and signal transmission. As a result, non-mulberry silk would be a suitable choice for preparing silk cocoon scaffolds. Eri cocoons demonstrate distinct multi-stage moisture transmission processes, providing an excellent moisture buffer effect. They also have a slower temperature change inside than

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outside, indicating temperature damping capability. The high sericin content of the cocoon coat, along with its exceptional ultraviolet absorption and antibacterial properties, effectively inhibits the infiltration of ultraviolet rays and microbes into the pupae.⁴⁷ Understanding the interplay between Eri cocoons' structural properties and performance index can serve as a structural model of bioinspiration in designing or optimizing multifunctional composites that aim to aid the biomedical and public health sectors. Moreover, eri silk, referred to as ahimsā (Sanskrit; non-violence) silk or peace silk, was selected due to the eco-friendly nature of production, as the natural life cycle of silkworms was not disrupted and no moth was sacrificed during the experiment.⁴⁸

This study, for the first time, reports the synthesis of the biomimetic advanced silver nanocomposite silk cocoon scaffold (ASN-SCS) bandage, designed using eri silk cocoons, and includes a systematic investigation of its structure, characterization, antibacterial activity, and biocompatibility. An attempt has been made to pave the way for developing an antibacterial dressing material that could combat burn wounds associated with MDR bacterial infection, which, in turn, could stimulate chronic wound healing.

2. Experimental section

2.1 Sample collection

The Eri (*Philosamia ricini*) silkworm cocoons were procured from the office of the Directorate of Textile Sericulture (under the Government of West Bengal), Bogram Sericulture Complex, Raiganj, Uttar Dinajpur, West Bengal.

2.2 Chemical and biological materials

All the chemicals used in this study were of analytical grade. The silver nitrate (AgNO₃) was procured from Merck (cat. no. 101512), USA. HiMedia Laboratories, Mumbai, India, provided the bacterial culture media. All the working solutions were made using autoclaved, deionized Milli-Q water. To remove the exogenous DNase or RNase, the glassware was autoclaved for 15 minutes at 121 °C. 70% ethanol was applied to the counter area and all the pipetting devices. Lyophilized *S. aureus* (MTCC 87) and *P. aeruginosa* (MTCC 1688) were obtained from the MTCC, IMTECH Chandigarh, India, and were used following the established safety laboratory protocols.

2.3 Preparation of SCS

SCS was synthesized utilizing non-mulberry Eri cocoons following a slight modification of the method described earlier⁴⁹ at a constant temperature of 25 ± 2 °C with a relative humidity of $50 \pm 2\%$ in the laboratory setup. Before selecting the cocoon layers for SCS preparation, the terminal portions of silk cocoons and the internal pupal residues were discarded. An Ajisawa solution comprising ethanol, calcium chloride, and H₂O with a molar ratio of 2 : 1 : 8 was prepared. Successively, the prepared cocoon layers were incubated in a water bath at a temperature of 60 °C for 2–3 hours to obtain SCS. After the Ajisawa solution changed color, the prepared cocoons were pulled out and rinsed three times with MilliQ water to acquire SCS samples (Fig. 1).

2.4 Preparation of ASN-SCS

20 mL of $AgNO_3$ aqueous solution with gradients of concentration (Table 1) was added to sterile borosilicate glass Petri plates (diameter 100 mm and height 15 mm), and to reach equilibrium, the SCS was incubated at room temperature and left overnight. The subsequent product was renamed ASN-SCS. Prepared ASN-SCS was taken out and washed three times with MilliQ water for further study.

2.5 Biophysical characterizations

2.5.1 UV-vis and laser light scattering studies. The aqueous Ajisawa solution-mediated synthesis of SCS and AgNO₃ aqueous solution-mediated synthesis of ASN-SCS were monitored *via* UV-vis spectrophotometer (Varian Inc., USA) within the wavelength range of 200–800 nm using a quartz cuvette of 1 cm optical path length. The laser diffraction analysis was also conducted for preliminary confirmation of NP formation.

2.5.2 Fourier-transform infrared spectroscopy (FT-IR) analysis. The reactive surface sites on SCS responsible for surface reactivity and the formation of ASN-SCS were detected using Fourier transform infrared (FT-IR) spectroscopy. The infrared spectrum was recorded on the PerkinElmer GX FT-IR system (PerkinElmer, USA) in the 4000–800 cm⁻¹ region.

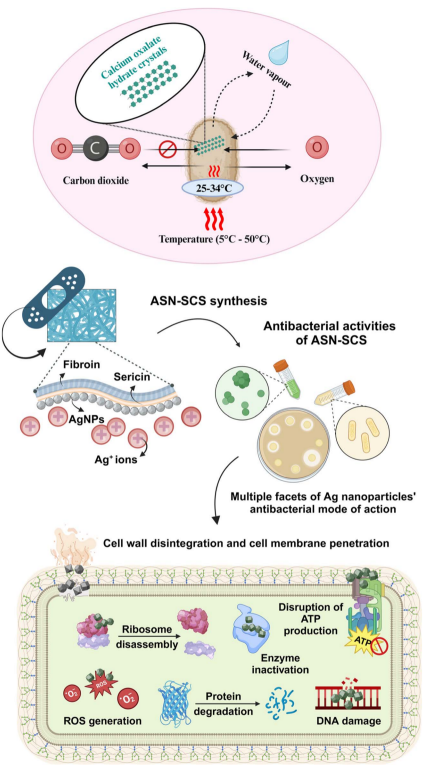
2.5.3 X-ray diffraction (XRD) analysis. The crystallinity characterization of ASN-SCS was obtained by using a Rigaku Miniflex II X-ray diffractometer in the range of 10° – 90° at 40 keV operating at 9 kW and CuK α radiation ($\lambda = 1.54056$ Å). The X'pert Highscore Plus software was used for processing and analyzing the data. The particle size (*d*) of the sample was calculated using Scherrer's equation.

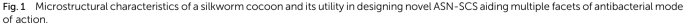
2.5.4 Microscopic analysis. The surface morphology of SCS and ASN-SCS was analyzed through a scanning electron microscope (SEM-EDX; S4160 Hitachi, Japan). Before testing, the samples underwent a direct current sputter technique (Emitechk450X, England) to coat them with a thin layer of gold.

2.5.5 Thermogravimetric–differential thermal analysis (TGA/DTA). The thermal properties of ASN-SCS were measured using a thermal gravimetric analyzer (TA instruments, simultaneous TGA–DTA, Waters LLC, New Castle, DE, USA). The tests were conducted under an inert nitrogen atmosphere in an open bin with a heating rate of $10 \,^{\circ}$ C min⁻¹ in a temperature range of $25-800 \,^{\circ}$ C. Simultaneously, weight loss and temperature difference between the sample and an inert reference were recorded and plotted against temperature for thermogravimetric analysis (TGA) and differential thermal analysis (DTA), respectively.

2.5.6 Hydrophilicity analysis. The contact angle of a liquid on a fabric's surface is a quantitative measure of the substrate's surface energy. A drop of water was placed on SCS and ASN-SCS to measure the contact angle using the drop deposition method. Images were captured, and the water contact angle measurements were performed using the ImageJ software.

Microstructural characteristics of silkworm cocoon





2.6 In vitro application of ASN-SCS

2.6.1 Kirby-Bauer test or zone of inhibition test. ASN-SCS bandages were tested for their antibacterial efficacy against *P*.

aeruginosa (MTCC 1688) and *S. aureus* (MTCC 87) employing an amended Kirby Bauer technique. The cultures of all the test organisms were transferred to the nutrient broth (NB) at 37 $^{\circ}$ C

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 $\label{eq:table_solution} Table 1 \quad \mbox{Concentration of $AgNO_3$ solution used for the preparation of different $ASN-SCS$ samples}$

AgNO ₃ concentration (in mM)	Corresponding sample abbreviations		
0.125	ASN-SCS _{0.125}		
0.25	ASN-SCS _{0.25}		
0.5	ASN-SCS _{0.5}		
1	ASN-SCS ₁		
2	ASN-SCS ₂		
4	$ASN-SCS_4$		
8	ASN-SCS ₈		
10	ASN-SCS10		
20	ASN-SCS ₂₀		
40	ASN-SCS ₄₀		
80	ASN-SCS ₈₀		
160	ASN-SCS ₁₆₀		
320	ASN-SCS ₃₂₀		

in a shaker-incubator operating at 150 rpm and left overnight. Thereafter, the cultures were diluted to 1×10^{6} CFU mL⁻¹, and 200 µL of the diluted cells were spread on sterile Mueller–Hinton (MH) agar plates (100 × 6 mm) and amended with sterile ASN-SCS bandages around the edges ($d = 10.50 \pm 0.50$ mm) and kept overnight at 37 °C to record the inhibitory zones. The MH agar plates with only SCSs were also kept as a control set for comparison.

2.6.2 Bacterial growth inhibition kinetics. The bacterial growth inhibition kinetics were performed using P. aeruginosa (MTCC 1688). Overnight-grown P. aeruginosa (MTCC 1688) suspension (1%) was taken and inoculated into fresh MH broth containing different concentrations of ASN-SCSs, which were 0.125 mM, 0.25 mM, 0.5 mM, 1 mM, 8 mM, 80 mM, and 320 mM, respectively. The inoculated broths were then incubated at 37 °C in a shaker-incubator at 150 rpm. The bacterial growth was assessed at various time intervals by quantifying the O.D. at λ_{600} nm. The minimum inhibitory concentration (MIC), which refers to the lowest concentration of Ag NPs embedded in ASN-SCSs that inhibits visible bacterial growth, was also determined. Each experiment was conducted three times, and the mean values of the results were reported. MH broth containing bacterial inoculums without any ASN-SCS was used as the negative control, and MH broth containing ASN-SCS without any bacterial inoculum was used as the positive control.

2.6.3 Hemocompatibility analysis. The hemocompatibility of the ASN-SCS samples was evaluated through a hemolytic assay, as described previously.⁵⁰ A 0.4 mg mL⁻¹ leaching aqueous solution of three samples of different ASN-SCSs (synthesized using 8 mM, 80 mM, and 320 mM AgNO₃ concentrations, respectively) was prepared. The aqueous dispersion was prepared through incubation of 0.4 mg mL⁻¹ ASN-SCS in PBS solution at 37 °C for 24 h, 48 h, and 72 h, respectively. For the hemocompatibility assay, an EDTA-stabilized human blood sample was freshly collected. The diluted suspension of human red blood cells (RBCs, 0.4 mL) was then mixed with varied concentrations of ASN-SCSs (1.6 mL). Diluted suspensions of RBCs mixed with 1.6 mL of PBS and 1.6 mL of double-distilled water were used as negative and positive controls, respectively. The mixtures were incubated at room temperature for 2 h. The % hemolysis was calculated by measuring the absorbance of the supernatant at 541 nm in an UV-vis spectrophotometer (Varian Inc., USA) and using the following equation:

% hemolysis =
$$[(A_{\rm S} - A_{\rm N})/(A_{\rm P} - A_{\rm N})] \times 100$$

here, A_S , A_N , and A_P are the absorbances of the sample, negative control, and positive control, respectively. Moreover, RBCs present in the pellet are used to prepare smears and were examined under a microscope at a magnification of 400× to document any changes in the morphology of RBCs by ASN-SCSs.

2.6.4 MTT assay. Healthy human adult primary epidermal keratinocytes (HEKa) cells were seeded in 96-well microtiter plates with DMEM/F-12 medium and incubated at 37 °C in 5% CO_2 for 24 h. On the next day, a 0.4 mg mL⁻¹ leaching aqueous solution of ASN-SCS8, ASN-SCS80, and ASN-SCS320 was prepared for three different time periods (24 h, 48 h, and 72 h) and added to each well, except the control, and kept further for incubation for another 24 hours at 37 °C in 5% CO₂. Subsequently, the plate was removed from the incubator, and the culture media were aspirated. Next, 10 µL of freshly prepared MTT reagent was added to each well and kept for 3 h under the same conditions. Afterward, 50 µL of isopropanol, a formazan solubilizer, was added to each well, and the absorbance was measured using an ELISA reader at 570 nm using a SPECTROstar Nano BMG LAB-TECH microtiter plate reader. All the factors such as dilution, mixing, trypsinization, etc., were synchronized before calculating the cell viability. The cell viability percentage was determined using the formula:

> [(O.D. treated well – blank)/ (mean O.D. control well – blank)] × 100.

2.6.5 Statistical analysis. All the triplicated data were analyzed using GraphPad Prism v. 8.4.2 and the values are presented as the mean \pm standard deviation (SD).

3. Results and discussion

3.1 Preparation of SCS and ASN-SCS

During the preparation of SCS, the Eri cocoons were separated from exterior to interior into five segments, *i.e.*, cocoon coat (*i.e.*, floss), cocoon layers (*i.e.*, upper, middle, and lower layers), and cocoon lining (*i.e.*, pelade).⁴⁴ The cocoon coats, containing loose silk fibers along with discontinuous static air gaps, were discarded. Furthermore, non-mulberry cocoons possess better water exchange mobility than mulberry cocoons and, hence, are a potential candidate in biomedical applications,^{51,52} especially in designing scaffold-based bandages. In Eri cocoons, the sericin content tends to decrease gradually from the outer to the inner layers.⁵³ Thus, the cocoon lining (pelade) with lower sericin content (5.95 \pm 0.30%) and lower moisture regain capacity (11.94 \pm 0.22%) was excluded from this study. Cocoon layers possessing the potential to qualify as a bandage material with higher sericin content (9.64 \pm 0.47%) and higher moisture regain capacity (12.34

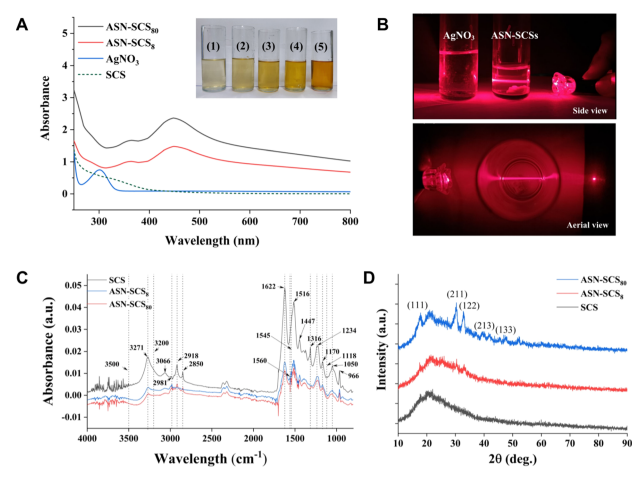


Fig. 2 (A) UV-vis absorption spectra of leached aqueous dispersions of ASN-SCSs in deionized water, $AgNO_3$ solution and Ajisawa solution during SCS synthesis. (B) Laser beam passing through aqueous solution of $AgNO_3$ and ASN-SCS solutions; upper image is side view, whereas the lower image is the aerial view. (C) FT-IR spectra of SCS and ASN-SCSs. (D) XRD analyses of SCS and ASN-SCSs.

 \pm 0.31%) were screened for further processing. Ajisawa solution was utilized for partially dissolving fibroin molecules to obtain SCS, which was used for ASN-SCS synthesis and characterization.

3.2 Biophysical characterization

3.2.1 UV-vis spectroscopy and laser light scattering analvses. Once the Ajisawa solution altered its color (Fig. 2A inset), the solution was analyzed using a UV-vis spectrophotometer. It revealed no significant peak of sericin, suggesting that the majority of sericin content is retained in SCS and can potentially embed Ag NPs on their surface. The color variation in the Ajisawa solution was visualized due to the naturally occurring pigments present in the Eri cocoon structure as well as due to NP formation. UV-vis spectroscopy analyses of ASN-SCSs were used to inspect the reduction of Ag⁺ ions to Ag⁰. The light exposure causes the polarization of free electrons that conduct in Ag NPs, leading to electron dipolar oscillation and the development of a surface plasmon resonance (SPR) spectrum at 448 nm.54 There was an absence of an absorption peak in the identical wavelength spectrum for the aqueous AgNO₃ solution that was used as a control. Fig. 2A presents the UV-vis absorption spectra of aqueous dispersions of ASN-SCSs containing Ag NPs. The laser beams passing through colloidal solution after the formation of ASN-SCSs exhibit the Tyndall effect of light scattering (Fig. 2B). The light scattering pattern due to the Tyndall effect preliminarily detects the formation of Ag NPs in the solution. The Tyndall effect was demonstrated with Ag NPs, which were microscopical colloidal particles yet sufficiently sized to scatter a light beam. The density of particles with colloidal structure and the frequency of the incident light both influence the magnitude of scattered light.

3.2.2 Fourier-transform infrared spectroscopy (FT-IR) analysis. The presence of a phenolic hydroxyl group in a molecule like tyrosine can typically exhibit characteristic peaks in the FTIR spectrum in the region between 3200 cm⁻¹ and 3500 cm⁻¹ due to the stretching vibration of the O–H bond.⁵⁵ The peaks of phenolic hydroxyl disappeared and were altered in the treated sample, indicating the interaction between the molecules and the NPs. Aromatic CH stretching vibrations usually appear as bands around 2800–3000 cm⁻¹. Bands at 2918 and 2850 cm⁻¹ regions rising from stretching C–H of aromatic compounds were detected, which might participate in the synthesis process as capping agents or stabilizers. Hence, the weakening or disappearance in the intensity of these bands could reflect

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alterations in the interactions between aromatic compounds and silver ions.⁵⁶ Moreover, in ASN-SCSs, the presence of a peak at 2981 cm⁻¹ indicated the same.⁵⁷ The infrared spectral range across 1700 and 1500 cm⁻¹ is frequently utilized for assessing diverse secondary features of silk fibroin. The peaks at 1610-1630 cm⁻¹ and 1510–1520 cm⁻¹ indicate the β -sheet secondary structure of silk fibroin, whereas the absorptions at 1648-1654 cm^{-1} and 1535–1542 cm^{-1} indicate the α -form conformation. The peaks at 1622 cm^{-1} , 1514 cm^{-1} , and 1236 cm^{-1} exhibit sericin amide-I (C=O stretching), amide-II (C-N stretching, N-H deformation), and amide-III (C-N stretching), suggesting a random coil/silk-I structure (Fig. 2C). The peak values at 1620–1625 cm⁻¹ show C=O stretching of carboxylate ions,⁵⁸ whereas those at 1560 cm^{-1} , 1550–1545 cm^{-1} , and 1540 cm⁻¹ denote association with silver-carboxylate complexes (-COO- + Ag). The development of large new absorption bands is due to NH deformation.59 The peak at 1118 cm⁻¹ reveals the stretching of the C–O bond of a phenolic hydroxyl group in the tyrosine of sericin. Following the reaction, the absorbance intensity at 1118 cm⁻¹ weakens, indicating that the phenolic hydroxyl groups were cleaved and oxidized to create quinone structures. The strong vibration of 1316 cm⁻¹ is assigned to the symmetric C=O stretching vibration of oxalate ions.⁶⁰ Here, the development of Ag NPs might be facilitated by the oxalate ions, which reduce Ag⁺ ions to Ag⁰ atoms. The peak at 1170 cm⁻¹ corresponds to C-N stretching of tyrosine, and the intensity declined in the ASN-SCSs.⁶¹ The band at 1050 cm⁻¹ corresponds to sericin C-O stretching.62

3.2.3 X-ray diffraction (XRD) analysis. The analysis revealed the presence of silver peaks in the ASN-SCS samples. When the XRD graph was examined, there were characteristic peaks at positions $2\theta^{\circ} = 17.668^{\circ}$, 29.757° , 32.778° , 39.312° , and 47.835° assigned to the 111, 211, 122, 213 and 133 planes of crystalline Ag (JCPDS card no: 00-001-0856) (Fig. 2D). Ag NPs were crystallized, as the measured values closely correlated with the standard reference of the Joint Committee on Powder Diffraction Standards for Ag NPs. The strong and apparent peaks of the XRD spectrum suggested the crystalline structure of ASN-SCSs.

The particle size was determined using the line width of the XRD peak, *i.e.*, *hkl* data, using Debye–Scherrer's equation (Table 2). In the equation, wavelength of X-ray (λ), peak width, *i.e.*, full width at half of its maximum intensity in radians (β), angle between the incoming and outgoing beam directions (2 θ), Bragg's diffraction angle (θ), Scherrer constant (K), and nanoparticle crystalline size or diameter ($D = K\lambda/\beta \cos \theta$) in nanometers.

Table 2 XRD data for the Debye–Scherrer equation (where $\lambda = 0.1541$ and K = 0.9) of ASN-SCSs

hkl	FWHM, deg	B, rad	2θ , deg	θ , deg	$\cos \theta$	<i>D_{hkl}</i> , nm
111	1.19455	0.02084	17.6680	8.8340	0.9881	6.65
211	1.08026	0.01885	29.7570	14.8785	0.9664	7.36
122	0.90165	0.01573	32.7780	16.3890	0.9593	8.82
213	1.40425	0.02450	39.3120	19.6560	0.9417	5.66
133	1.04384	0.01821	47.8350	29.9175	0.8667	7.62

The smallest NPs appeared to be primarily in their monocrystalline form. These characteristic peak values are consistent with previous reports.^{16,63} An earlier study demonstrated that the bactericidal properties of Ag NPs within a range of 5–100 nm were dramatically increased as the NP size was reduced to <10 nm. This may be attributable to an elevation in the surface area to volume ratio (SA/V) with a reduction in particle size and an increase in relative particle concentration.⁶⁴

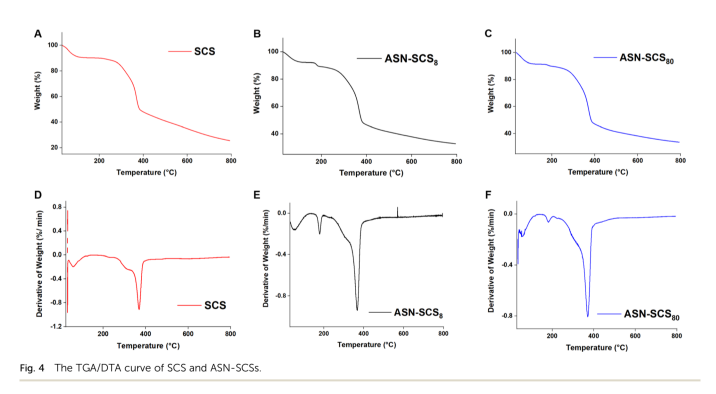
3.2.4 SEM-EDX. SEM is a surface imaging technique that can determine the size, shape, and texture of particles. SEM images revealed the embedding of Ag NPs on the surface of ASN-SCSs (Fig. 3). Interestingly, ASN-SCS₈₀ showed deposited NPs on silk that were absent in the control sample (SCS). The SCS, in turn, exhibits calcium oxalate crystals that got reduced on the ASN-SCS surface, indicating its role as a reducing agent during NP synthesis, which agrees with the FT-IR data. In an earlier report, a similar SEM study was performed to confirm nanosilica deposited on eri silk fabric.⁶⁵ Moreover, the EDX spectra of ASN-SCS revealed a percentage of Ag of 26.85%, which indicates that ASN-SCS was embedded with Ag NPs.

3.2.5 Thermogravimetric-differential thermal analysis (TGA/DTA). The thermal behavior and thermal stability of ASN-SCSs were investigated via TGA/DTA analysis at high temperatures (Fig. 4). The weight loss pattern in TGA curves can be categorized into three different regions depending on mass loss rates. The first region revealed a slight reduction in weight for silk scaffolds up to 110 °C, mostly due to water adsorption. The second zone (200-380 °C) showed a sharp increase in weight reduction, as most of the peptide chains were broken and intermolecular interactions decreased. The third zone, beginning at 380 °C, demonstrated a steady and persistent breakdown at a lower rate up to 800 °C. The final residual weight in control remained at 25.73% after 800 °C whereas, the remaining residual weight in treated samples (ASN-SCS₈ and ASN-SCS₈₀) was found to be 32.77% and 33.69%, respectively, which concludes enhanced stability of ASN-SCS bandages due to NPs embedded on SCS. Moreover, the stability of the ASN-SCS was much greater than previously reported studies, suggesting it as durable bandage material for future biomedical а applications.66

3.2.6 Hydrophilicity analysis. To measure the hydrophobicity of textiles, the water drop contact angle was measured. The surface becomes more hydrophobic when the water drop contact angle is >90° and superhydrophobic when the contact angle is >150°.67 Fig. 5 shows the water contact angles of SCS and ASN-SCS. It depicts an increased contact angle as the percentage of embedded NPs increases in treated ASN-SCS relative to untreated SCS. The SCS exhibited a contact angle of 108.24°, which was raised to 123.63° in the case of ASN-SCS₈₀, indicating that both the samples are hydrophobic. Moreover, hydrophobicity intensified with increased NP deposition on ASN-SCS. An earlier study found that the roughness of the surface and the composition of its chemicals have a significant impact on the contact angle.68 Aggregated NPs on the scaffold surface may enhance surface micro-roughness, potentially increasing fabric hydrophobicity because of the lotus effect.69 Hence, the increased hydrophobicity of ASN-SCS, along with its

A C	°	,	S
\$3400 15 0KV 14 8mm x100 SE 2/22/2024 16 20 500um	0 1 2 3 Full Scale 1231 cts Cursor: 0.000	4 5 6	7 8 9
B			
	Element	Weight %	Atomic %
	СК	40.11	59.06
	ОК	33.05	36.53
	Ag L	26.85	4.40
	Totals	100.00	
			1

Fig. 3 SEM micrograph of (A) SCS and (B) ASN-SCS. Red circles in the inset of (B) indicate ASN-SCS embedded with NPs. (C) EDX spectrum of ASN-SCS.



microstructured surface, is anticipated to inhibit biofilm formation in the case of one of the antibacterial actions of ASN-SCS. Generally, microbes have a lower ability to adhere to structured surfaces compared to flat films. This is because micro-structured surfaces minimize the amount of surface area accessible for microorganisms to adhere to.⁷⁰ Previous studies revealed that hydrophobic dressings could inhibit biofilm formation, leading to an adverse impact on antimicrobial resistance and promoting immune system mobilization.⁷¹ As a result, synthesized ASN-SCS has the potential to be excellent wound dressing materials.

3.3 In vitro tests

3.3.1 Kirby–Bauer test of synthesized ASN-SCSs. The synthesized ASN-SCSs were investigated for antibacterial activity against two major burn wound-inhabiting multi-drug-

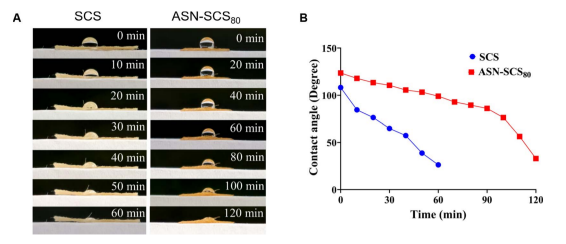


Fig. 5 (A) Time lapse photographs of water droplets on the SCS and ASN-SCS₈₀. (B) Contact angles of water droplets on the SCS and ASN-SCS₈₀ as a function of time.

resistant pathogens: *P. aeruginosa* (MTCC 1688) and *Staphylococcus aureus* (MTCC 87). ASN-SCSs synthesized using three different cocoon layers [upper (1), middle (2), and lower (3)] exhibited strong antibacterial efficacy against these tested MDR pathogens with distinct inhibition zones (Fig. 6).

All the cocoon layers showed approximately similar inhibition zones when tested against either Gram-negative or Grampositive bacteria. Therefore, utilizing multiple cocoon layers from the same cocoon would be economically beneficial for industrial use in the future. The antibacterial efficacy of the ASN-SCSs was slightly higher in Gram-negative bacteria when compared to the results of Gram-positive bacteria. The plausible explanation for these differential inhibition zones is connected to the fact that the cell walls of Gram-negative bacteria possess a thin layer of peptidoglycans. On the other hand, the likelihood of Ag NPs penetrating Gram-positive bacterial cells is reduced by their thick peptidoglycan coating. The antibacterial performance of ASN-SCSs prepared with varied concentrations of AgNO₃ was evaluated against P. aeruginosa (MTCC 1688), which is the major burn wound-colonizing bacteria (Fig. 6C-L). Following a 24 h incubation period at 37 °C, varying levels of antibacterial activity were seen in each sample, with the ASN-SCS prepared using an 80 mM concentration of AgNO₃ depicted as ASN-SCS₈₀ clearly demonstrating the highest antibacterial zones (5.50 mm) against the tested bacterium. These results point to an improved antimicrobial efficacy at a lower dosage compared to previous reports where the highest zone of inhibition was observed for silkworm cocoon-based wound dressings synthesized using 200 mM of AgNO3.33 The control SCSs displayed no inhibition against P. aeruginosa, which suggests that silk has no antibacterial effect (Fig. 6M). It was previously stated that the chemical residues that remained after isolating or purifying the cocoon components were accountable for the antibacterial characteristics of pure silk cocoons.72 However, in this study, the zone diameters of ASN-SCS samples gradually increased up to ASN-SCS₈₀; thereafter, they decreased, as noted for ASN-SCS₁₆₀ and ASN-SCS₃₂₀ (Fig. 6K and L). Thus, ASN-SCS₈₀

was chosen for further study and large-scale production. The antibacterial effect of ASN-SCSs could be attributed to the release of Ag⁺ from the sample discs, which resulted in increased antibacterial activity. In contrast, ASN-SCSs synthesized with higher AgNO₃ concentrations, *viz.*, ASN-SCS₁₆₀ and ASN-SCS₃₂₀, exhibited a sharp decrease in the zone of inhibition which might be linked to the fact that the higher concentration of Ag NPs result in agglomeration as reported previously.³³ However, further study is needed to validate this.

3.3.2 Microbial growth inhibition study. The results of the Kirby Bauer test analysis revealed that the antibacterial effect is prominent in each sample of ASN-SCSs synthesized using 1, 2, 4, 8, 10, 20, 40, 80, 160, and 320 mM AgNO₃ solution. To explore the efficacy of these ASN-SCSs, the growth kinetics of P. aeruginosa were checked in the presence of ASN-SCSs, and the MIC value was estimated. The bacterial inhibition of the tested bacterium treated with ASN-SCSs (ASN-SCS_{0.125} to ASN-SCS₃₂₀) was almost 100% when recorded after 24 h of incubation amended with 0.2 g of different concentrations of ASN-SCSs in a 20 mL bacterium culture broth with an initial O.D. of 0.06 at λ_{600} (Fig. 3A). Our results further showed that the bacterial inhibition from ASN-SCS_{0.5} to ASN-SCS₃₂₀ against P. aeruginosa was nearly 100% even after 72 h of incubation. Furthermore, MIC was also assessed to precisely determine the lowest concentration of ASN-SCSs, preventing the visible growth of P. aeruginosa. Among all the experimental sets, ASN-SCS_{0.5} was depicted as the MIC value.

The bacterial population began proliferating after 24 h of incubation in broths treated with ASN-SCS_{0.25} and ASN-SCS_{0.125} due to delayed lag-phase (Fig. 7A) and degradation of silk proteins by the test bacterium to utilize the amino acids as a nitrogen source for its growth. Therefore, the released proteins might have contributed to the increased absorbance value in the ASN-SCS_{0.125} sample. A similar interpretation was also described in a separate study where *P. aeruginosa* could potentially carry out silk biodegradation in minimal medium by proteolytic enzymes called elastases.⁷³ While most silk-

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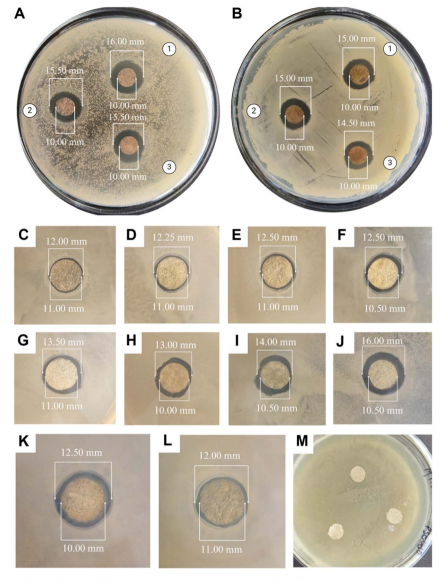


Fig. 6 Antibacterial assay of ASN-SCS (using 80 mM AgNO₃ abbreviated as ASN-SCS₈₀) synthesized using three different cocoon layers against (A) Gram negative bacteria *P. aeruginosa* and (B) Gram positive bacteria *S. aureus*; where the ASN-SCSs prepared utilizing upper, middle, and lower layers of Eri cocoons are indicated with 1, 2 and 3 respectively. Zone of inhibition diameters (in mm) of (C–L) ASN-SCSs synthesized using 1, 2, 4, 8, 10, 20, 40, 80, 160, 320 mM AgNO₃ solution respectively; and (M) SCSs as negative control against *P. aeruginosa*.

degrading gram negatives use sericin to obtain amino acids, another species of the genus *Pseudomonas* named *Burkholderia cepacia* (formerly *Pseudomonas cepacia*) was utilizing fibroin proteins as a nitrogen as well as carbon source. To visualize the morphological changes in ASN-SCS-treated bacteria, the SEM study was carried out, where the membrane stability was altered after ASN-SCS application, indicating the bactericidal effect of ASN-SCSs (Fig. 7A inset).

3.3.3 *In vitro* hemocompatibility analysis of ASN-SCSs. The hemolytic assay is a crucial test for any biomaterials that contact blood, assessing their hemocompatibility to determine their selection for further use. According to the ISO standard, a material is deemed safe if its hemolysis index is <5%.⁷⁴ The hemocompatibility analysis on human red blood cells was accomplished using the leaching aqueous dispersion of ASN-

SCS bandages. Most of the samples exhibited <1% hemolysis, indicating their hemocompatible nature (Table 3, Fig. 7B).

The positive control-lysed RBCs exhibited two significant peaks in their absorption spectra (Fig. 7B). According to the literature, the two high absorption peaks in hemoglobin's (Hb) spectrum, nearly at 540 and 576 nm, along with an absorption valley at 560 nm, indicate that oxygen partial pressure (PO₂) is higher than 100 mm Hg and an oxy-Hb structure was formed.⁷⁵ Various studies have shown that silk bandages are hemocompatible. As well, dose-dependent Ag NPs do not have high hemolysis potential.⁷⁶ In our previous study, 13% hemolysis was observed in the presence of 40 µg mL⁻¹ of bioconjugated Ag NPs.⁷⁷ Elsewhere, it was suggested that Ag NPs affect platelet aggregation, and blood coagulation over a concentration of 40



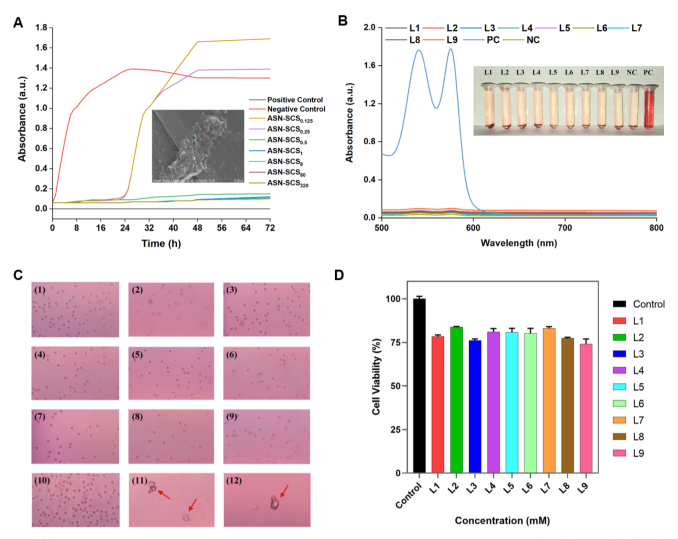


Fig. 7 (A) Growth kinetics of *P. aeruginosa* in the presence of different concentrations of Ag NPs embedded in ASN-SCSs for 72 h (ASN-ASN-SCS_{0.125}, ASN-SCS_{0.25}, ASN-SCS_{0.5}, ASN-SCS₁, ASN-SCS₈, ASN-SCS₈, and ASN-SCS₃₂₀ denote ASN-SCSs prepared in 0.125 mM, 0.25 mM, 0.5 mM, 1 mM, 8 mM, 80 mM, and 320 mM AgNO₃ solution). Figure inset denotes the SEM micrograph of *P. aeruginosa* treated with ASN-SCSs; red arrows indicate changes in bacterial morphology. (B) Hemolysis study of human RBCs treated with varied concentrations of ASN-SCSs. RBCs suspended in PBS and distilled water were used as negative and positive controls, respectively. The inset represents the photographs of centrifuged samples, ensuring the supernatant contains no hemoglobin (Hb). (C) Microscopic images of RBCs present in blood smears prepared from samples L1–L9 (1–9), NC (10), and PC (11–12), containing ghost RBCs (red arrow). (D) Cell viability analysis using the MTT assay on HEKa cells. The results shown are the mean \pm SD of three separate experiments performed in triplicate.

Table 3 Hemolysis assay of leaching liquors from different ASN-SCSs

ASN-SCSs	Leaching duration	Sample	% Hemolysis	
ASN-SCS ₈	24h	L1	0.14	
ASN-SCS ₈₀	24h	L2	0.45	
ASN-SCS320	24h	L3	0.63	
ASN-SCS ₈	48h	L4	0.15	
ASN-SCS80	48h	L5	0.57	
ASN-SCS ₃₂₀	48h	L6	0.81	
ASN-SCS ₈	72h	L7	0.61	
ASN-SCS ₈₀	72h	L8	0.94	
ASN-SCS320	72h	L9	1.88	

 μ g mL⁻¹.⁷⁸ Despite synthesis with greater concentrations of AgNO₃, ASN-SCSs did not exhibit significant hemolysis (Fig. 7B,

Table 3), indicating their hemocompatible nature, which supports their potential use in tissue engineering and wound healing. Moreover, blood smears prepared from the RBCs present in the sample pellet after final centrifugation during hemocompatibility analysis procedures revealed the presence of intact RBCs in all the samples. No significant changes in the shape and morphology of RBCs or in absorbance spectra (Fig. 7C) were recorded in the negative control. In turn, the positive control samples were composed of ghost RBCs.

3.3.4 MTT assay. To estimate the effect of the synthesized ASN-SCSs on humans, healthy human adult primary epidermal keratinocytes (HEKa) cells were used. The MTT assay relies on the respiratory system of the mitochondria and secondarily helps in the evaluation of the cellular energy capacity of a viable cell. MTT is a yellow dye that is reduced by oxido-reductase, where NADPH

serves as an electron donor, resulting in the blue-colored product, formazan. Such transformation is feasible only in live cells; thus, the amount of formazan is directly correlated to the number of viable cells. No cytotoxicity was detected, regardless of the leaching time of ASN-SCSs. The ASN-SCS₈₀ exhibited high biocompatibility with a cell viability of approximately 80% (Fig. 7D), which agrees well with a previous study where silkworm cocoon wound film exhibited no significant cytotoxity against murine L929 fibroblast cells and the cell viability was > 80%.³³

4. Conclusion

In conclusion, ASN-SCSs synthesized using eri silkworm cocoon were biomimetic, preserving both the benefits of fibroin and sericin, and qualified as an advanced dressing compared to conventional dressings. In situ assembly of Ag NPs on SCS was carried out using natural functional groups present in sericin and fibroin molecules as reducing and stabilizing agents. Moreover, the ASN-SCS bandages showed excellent thermal stability and water retention properties. The embedded Ag NPs exhibited a particle size ranging from 5.66 to 8.82 nm. ASN-SCSs were found to be strongly effective as an antibacterial agent against MDR pathogens, i.e., against both P. aeruginosa and S. aureus. Following the demonstration of hemocompatibility in vitro, ASN-SCS bandages exhibited superior biocompatibility, which suggests their potential application in chronic wounds such as burns. Due to embedded Ag NPs, ASN-SCSs would possess an increased shelf life and remain sterile for a longer period, supporting less biomedical waste generation. As an outcome, this work presented an easy and sustainable way for developing a novel multifunctional bandage or dressing that has significant therapeutic potential in clinical and biomedical sectors. Therefore, this approach would substantially reduce production costs, simplify the fabrication process, and enable facile commercialization.

Data availability

The data supporting this article is included in the manuscript.

Author contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

Conflicts of interest

The authors declare no competing financial interest.

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