

Original citation: Gupta, A., Low, W.L., Radecka, I., Britland, S.T., Mohd Amin, M.C.I. and Martin, Claire (2016) *Characterisation and In Vitro Antimicrobial Activity of Biosynthetic Silver-loaded Bacterial Cellulose Hydrogels*. Journal of Microencapsulation, 33 (8). pp. 725-734. ISSN Print: 0265-2048 Online: 1464-5246

Permanent WRaP URL: https://eprints.worc.ac.uk/id/eprint/8014

Copyright and reuse:

The Worcester Research and Publications (WRaP) makes this work available open access under the following conditions. Copyright © and all moral rights to the version of the paper presented here belong to the individual author(s) and/or other copyright owners. To the extent reasonable and practicable the material made available in WRaP has been checked for eligibility before being made available.

Copies of full items can be used for personal research or study, educational, or not-for-profit purposes without prior permission or charge, provided that the authors, title and full bibliographic details are credited, a hyperlink and/or URL is given for the original metadata page and the content is not changed in any way.

Publisher's statement:

This is an Accepted Manuscript of an article published by Taylor & Francis in Journal of Microencapsulation on 26 October 2016, available online: <u>https://doi.org/10.1080/02652048.2016.1253796</u>

A note on versions:

The version presented here may differ from the published version or, version of record, if you wish to cite this item you are advised to consult the publisher's version. Please see the 'permanent WRaP URL' above for details on accessing the published version and note that access may require a subscription.

For more information, please contact wrapteam@worc.ac.uk

- Author(s): Abhishek Gupta, Wan Li Low, Iza Radecka, Stephen T. Britland, Mohd Cairul Iqbal Mohd Amin, and Claire Martin
- Article title: Characterisation and *in vitro* antimicrobial activity of biosynthetic silver-loaded bacterial cellulose hydrogels

Article no: IMNC_A_1253796

Enclosures: 1) Query sheet 2) Article proofs

Dear Author,

1. Please check these proofs carefully. It is the responsibility of the corresponding author to check these and approve or amend them. A second proof is not normally provided. Taylor & Francis cannot be held responsible for uncorrected errors, even if introduced during the production process. Once your corrections have been added to the article, it will be considered ready for publication.

Please limit changes at this stage to the correction of errors. You should not make trivial changes, improve prose style, add new material, or delete existing material at this stage. You may be charged if your corrections are excessive (we would not expect corrections to exceed 30 changes).

For detailed guidance on how to check your proofs, please paste this address into a new browser window: http://journalauthors.tandf.co.uk/production/checkingproofs.asp

Your PDF proof file has been enabled so that you can comment on the proof directly using Adobe Acrobat. If you wish to do this, please save the file to your hard disk first. For further information on marking corrections using Acrobat, please paste this address into a new browser window: http://journalauthors.tandf.co.uk/production/acrobat.asp

2. Please review the table of contributors below and confirm that the first and last names are structured correctly and that the authors are listed in the correct order of contribution. This check is to ensure that your name will appear correctly online and when the article is indexed.

Sequence	Prefix	Given name(s)	Surname	
1		Abhishek	Gupta	
2		Wan Li	Low	
3		Iza	Radecka	
4		Stephen T. Britland		
5		Mohd Cairul Iqbal Mohd Amin		
6		Claire	Martin	

Queries are marked in the margins of the proofs, and you can also click the hyperlinks below.

General points:

1. **Permissions:** You have warranted that you have secured the necessary written permission from the appropriate copyright owner for the reproduction of any text, illustration, or other material in your article. Please see http://journalauthors.tandf.co. uk/permissions/usingThirdPartyMaterial.asp.

- 2. Third-party content: If there is third-party content in your article, please check that the rightsholder details for re-use are shown correctly.
- 3. Affiliation: The corresponding author is responsible for ensuring that address and email details are correct for all the coauthors. Affiliations given in the article should be the affiliation at the time the research was conducted. Please see http:// journalauthors.tandf.co.uk/preparation/writing.asp.
- 4. **Funding:** Was your research for this article funded by a funding agency? If so, please insert `This work was supported by <insert the name of the funding agency in full>', followed by the grant number in square brackets `[grant number xxxx]'.
- 5. Supplemental data and underlying research materials: Do you wish to include the location of the underlying research materials (e.g. data, samples or models) for your article? If so, please insert this sentence before the reference section: 'The underlying research materials for this article can be accessed at <full link>/ description of location [author to complete]'. If your article includes supplemental data, the link will also be provided in this paragraph. See <htp://journalauthors.tandf.co. uk/preparation/multimedia.asp> for further explanation of supplemental data and underlying research materials.
- 6. The **PubMed** (http://www.ncbi.nlm.nih.gov/pubmed) and **CrossRef databases** (www.crossref.org/) have been used to validate the references. Changes resulting from mismatches are tracked in red font.

AUTHOR QUERIES

- Q1: Please provide the town and state abbreviation (for US) or town and country of origin (for other countries) identifying the headquarter location for "Acros Organics".
- Q2: A disclosure statement reporting no conflict has been inserted. Please confirm the statement is accurate.
- Q3: Please provide complete details for (Solway and Consalter, 2010) in the reference list or delete the citation from the text.
- Q4: Please check reference Costa and Lobo (2001) in the text citation is changed to Costa and Lobo (2011) as per reference list.
- Q5: Please provide the page range for Ref. (Kim et al. 2009) in the reference list entry.
- Q6: Please provide the page range for Ref. (Murphy & Evans, 2012) in the reference list entry.
- Q7: The ORCID details of the authors have been validated against ORCID registry. please check the ORCID ID details of the authors.
- Q8: Please check and confirm the corresponding author details correct as set in proof.
- Q9: Please resupply Figures 3 and 4 in a format suitable for printing.

How to make corrections to your proofs using Adobe Acrobat/Reader

Taylor & Francis offers you a choice of options to help you make corrections to your proofs. Your PDF proof file has been enabled so that you can mark up the proof directly using Adobe Acrobat/Reader. This is the simplest and best way for you to ensure that your corrections will be incorporated. If you wish to do this, please follow these instructions:

1. Save the file to your hard disk.

2. Check which version of Adobe Acrobat/Reader you have on your computer. You can do this by clicking on the Help" tab, and then About".

If Adobe Reader is not installed, you can get the latest version free from http://get.adobe.com/reader/.

3. If you have Adobe Acrobat/Reader 10 or a later version, click on the Comment" link at the right-hand side to view the Comments pane.

4. You can then select any text and mark it up for deletion or replacement, or insert new text as needed. Please note that these will clearly be displayed in the Comments pane and secondary annotation is not needed to draw attention to your corrections. If you need to include new sections of text, it is also possible to add a comment to the proofs. To do this, use the Sticky Note tool in the task bar. Please also see our FAQs here: http://journalauthors.tandf.co.uk/production/index.asp.

5. Make sure that you save the file when you close the document before uploading it to CATS using the Upload File" button on the online correction form. If you have more than one file, please zip them together and then upload the zip file. If you prefer, you can make your corrections using the CATS online correction form.

Troubleshooting

Acrobat help: http://helpx.adobe.com/acrobat.html Reader help: http://helpx.adobe.com/reader.html

Please note that full user guides for earlier versions of these programs are available from the Adobe Help pages by clicking on the link Previous versions" under the Help and tutorials" heading from the relevant link above. Commenting functionality is available from Adobe Reader 8.0 onwards and from Adobe Acrobat 7.0 onwards.

Firefox users: Firefox's inbuilt PDF Viewer is set to the default; please see the following for instructions on how to use this and download the PDF to your hard drive: http://support.mozilla.org/en-US/kb/view-pdf-files-firefox-without-downloading-them#-w_using-a-pdf-reader-plugin

RESEARCH ARTICLE

2

3

4

6

7

8

9

10

11

12

13

14

15

16

17

18

19

20

21

22

23

24

25

26

27

28

29

30

31

32

33

34

35

36

37

38

39

40

41

42

43

44

45

46

47

48

49

50

51

52

53

54

55

56

57

58

59

60

61

62

63 64

 \odot

 \bigcirc

65 66 67 68 69 70 71 72 73 74 75 76 77 78 79 80 81 82 83 84 85 86 87 88 89 90 91 92 93 94 95 96 97 98 99 100 101 102 103 104 105 106 107 108 109 110 111 112 113 114 115 116 117 118 119 120 121 122 123 124 125 126

127

128

Characterisation and *in vitro* antimicrobial activity of biosynthetic silver-loaded bacterial cellulose hydrogels

Abhishek Gupta^{a,c}, Wan Li Low^{a,c}, Iza Radecka^{b,c}, Stephen T. Britland^{a,c}, Mohd Cairul Iqbal Mohd Amin^d and Claire Martin^{a,c}

^aSchool of Pharmacy, Faculty of Science and Engineering, University of Wolverhampton, Wolverhampton, UK; ^bSchool of Biology, Chemistry and Forensic Science, Faculty of Science and Engineering, University of Wolverhampton, Wolverhampton, UK; ^cResearch Institute in Healthcare Science, Faculty of Science and Engineering, University of Wolverhampton, Wolverhampton, UK; ^dFaculty of Pharmacy, Universiti Kebangsaan Malaysia, Kuala Lumpur, Malaysia

ABSTRACT

Wounds that remain in the inflammatory phase for a prolonged period of time are likely to be colonised and infected by a range of commensal and pathogenic microorganisms. Treatment associated with these types of wounds mainly focuses on controlling infection and providing an optimum environment capable of facilitating re-epithelialisation, thus promoting wound healing. Hydrogels have attracted vast interest as moist wound-responsive dressing materials. In the current study, biosynthetic bacterial cellulose hydrogels synthesised by *Gluconacetobacter xylinus* and subsequently loaded with silver were characterised and investigated for their antimicrobial activity against two representative wound infecting pathogens, namely *S. aureus* and *P. aeruginosa*. Silver nitrate and silver zeolite provided the source of silver and loading parameters were optimised based on experimental findings. The results indicate that both AgNO₃ and AgZ loaded biosynthetic hydrogels possess antimicrobial activity (p < .05) against both *S. aureus* and *P. aeruginosa* and may therefore be suitable for wound management applications.

Introduction

Wound healing is a complex physiological cascade that follows several orderly, but overlapping phases (Guo and DiPietro, 2010; Martin et al., 2014), the rate of which is influenced by the type, size and depth of wound, as well as the presence of infection (Martin et al., 2013). Invasion of endogenous or exogenous pathogenic microorganisms into the wound site (Landis, 2008) can delay normal wound healing and potentially lead to the development of a chronic, non-healing wound. This microbial invasion usually occurs due to the uncontrolled proliferation of opportunistic pathogens amongst the polymicrobial skin microflora: Opportunistic microorganisms (e.g. Streptococcus spp, Corynebacterium spp, Escherichia coli, Bacillus subtilis, Staphylococcus aureus and Pseudomonas aeruginosa) can colonise the wound site and substantially prolong wound healing (Kirketerp-Møller et al., 2008; Landis, 2008). Conventional dressings (e.g. gauze and tulle) form a barrier from the external environment and keep the wound dry, but are unable to directly influence the healing process (Singh et al., 2013; Kamoun et al., 2015). Conversely, moist dressings act as a barrier to infection and also maintain moisture levels around the wound. These dressings are easily removed from the wound site, thus avoiding further trauma during dressing changes, and in the case of hydrogel-based dressings, respond to variations in moisture, thereby facilitating re-epithelialisation (Winter, 1962; Abdelrahman and Newton, 2011; Fu et al., 2013). In addition to absorbing and retaining wound exudate, hydrogel-based dressings also provide a cooling, soothing effect, thereby reducing the sensation of pain (Maneerung et al., 2008; Solway and Consalter, 2010). $\mathbf{Q}^{\mathbf{3}}$

Bacterial cellulose (BC), a biosynthetic homopolymer synthesised by the Gram negative obligate aerobe Gluconacetobacter xylinus (Pinto et al., 2009; Castro et al., 2011) has attracted great interest as a potential wound dressing (Czaja et al., 2006; Solway et al., 2010). Unlike plant cellulose, BC is highly pure and free from contaminants such as lignin, pectin and hemicellulose (Sannino et al., 2009); its highly crystallised microstructure also results in a finer web-like network with higher tensile strength compared to plant cellulose (Pinto et al., 2009). BC is biocompatible, non-pyrogenic, hydrophilic and transparent, all of which make it innately suitable for wound management applications (Czaja et al., 2006; Fu et al., 2013; Abeer et al., 2014). The hydrophilicity and relatively high swelling ratio of BC (Nakayama et al., 2004; Maneerung et al., 2008) allow it to establish a moist microenvironment at the wound interface in addition to maintaining water vapour transmission rate (WVTR) and a constant temperature. Hydrophilicity also enables reversible swelling and de-swelling of the hydrogel, which can be exploited in the fluid management of heavily exuding wounds. In addition, BC hydrogel dressings can protect tissues forming over and around the wound site, as well as promote angiogenesis (Maneerung et al., 2008). These unique properties have led to the successful commercialisation of BC hydrogels (e.g. Dermafill[™], Biofill[®], Bioprocess[®] and Gengiflex[®]) for the treatment of burns, chronic ulcers, skin lesions and periodontal disease, respectively (Czaja et al., 2006; Solway et al., 2010; Fu et al., 2013). Whilst not inherently antimicrobial itself, BC's unique 3-D nanofibrillar network is highly porous and amenable to high loading and controlled release of a range of antimicrobial agents (Fu et al., 2013; Shah et al., 2013; Wu et al., 2014).

ARTICLE HISTORY

Received 18 July 2016 Revised 17 October 2016 Accepted 17 October 2016 Published online

KEYWORDS

Antimicrobial; bacterial cellulose; hydrogel; silver; silver zeolite

lonic silver (Ag⁺) is a broad spectrum antimicrobial agent with 129 130 activity against yeast, fungi and several antibiotic resistant bacteria including methicillin resistant S. aureus (MRSA) and vancomycin 131 resistant enterococci (VRE) (Murphy and Evans, 2012). Of equal 132 importance, Ag⁺ has relatively low toxicity in human cells at con-133 centrations that are antimicrobially active against pathogenic 134 135 microbes (Copcia et al., 2011; Wilkinson et al., 2011). Since the dawn of the antibiotic era, the use of Ag⁺ has gradually 136 137 decreased, but the upsurge in multidrug resistant microbial strains 138 has led to a resurgence in interest (Chopra, 2007; Radecka et al., 139 2015). Ag⁺ are highly reactive and can bind to multiple intra- and 140 extracellular target sites that alter cellular functionality, structural 141 integrity, permeability and transport systems (Chopra, 2007; 142 Maneerung et al., 2008; Low et al., 2013; Martin et al., 2013; 143 Martin et al., 2015). Once transported into the bacterial cytoplasm, 144 Ag⁺ interact with essential intracellular enzymes and DNA, dis-145 rupting vital cell functions and impairing cell replication, that 146 eventually lead to cell death (Castellano et al., 2007; Low et al., 147 2011; Low et al., 2013; Martin et al., 2015). The multifaceted, broad 148 spectrum mode of action of Ag⁺ can be highly effective at con-149 trolling chronic wound infections even at parts per million con-150 centrations (Brett, 2006). The minimum lethal concentration (MLC) 151 of Aq⁺ against *P. aeruginosa* and *S. aureus* was reported to be 152 $1.59\times 10^{-3\%}$ w/v (equivalent to 15.9 ppm) and $5.08\times 10^{-3\%}$ w/v 153 (equivalent to 50.8 ppm) respectively (Low et al., 2011). Silver has 154 other beneficial wound management functions including anti-155 inflammatory activity resulting from complex formation with met-156 alloproteinases, leading to localised down regulation of inflamma-157 tion, thus creating conditions suitable for re-epithelialisation and 158 angiogenesis (Atiyeh et al., 2007; Leaper, 2012). There are several 159 commercially available topical silver products including dressings 160 (Acticoat[®], Contreet-H[®], Actisorb Silver 220[®], Aquacel-Ag[®], Ag 161 ExtraTM), solutions (Sulfamylon[®]) and creams (Silvadene[®]) 162 FlamazineTM) (Martin et al., 2013). Whilst the most commonly used 163 products contain silver nitrate or silver sulfadiazine as the source 164 of Aq⁺ (lp et al., 2006), an alternative donor of Aq⁺ are silver zeo-165 lites (AgZ): a microporous, non-reactive, crystalline aluminosilicate 166 frameworks containing Ag⁺ (Kwakye-Awuah et al., 2008). The 167 anionic cavities of zeolites are occupied by cations and/or water 168 molecules that have considerable freedom of movement that per-169 mit charge balancing ions to exchange in aqueous media, without 170 affecting the structure of zeolites (Copcia et al., 2011). Cationic 171 Ag⁺ interacts with the anionic zeolite framework to form AgZ; the 172 subsequent release of Ag⁺ from the zeolite structure is achieved 173 by cation exchange (Kwakye-Awuah et al., 2013). The antimicrobial 174 activity of AgZ has been attributed to Ag⁺ release from the zeo-175 lite, and the subsequent formation of reactive oxygen species in 176 the matrix (Matsumura et al., 2003). Kwakye-Awuah et al. (2013) 177 and Matsumura et al. (2003) have demonstrated the antimicrobial 178 activity of AgZ against Gram positive (S. aureus) and Gram nega-179 tive (E. coli and P. aeruginosa) bacteria, all of which are commonly 180 found in infected wounds. 181

The combination of silver and BC is not extensively reported in 182 the literature: Maneerung et al. (2008), Yang et al. (2012) and Wu 183 et al. (2014) reported the antimicrobial activity of silver nanopar-184 ticle-impregnated BC against E. coli (Gram negative) and P. aerugi-185 nosa (Gram positive). Antimicrobial silver-impregnated cellulose 186 (SIC) was microbiocidal against P. putida and M. luteus, and dem-187 onstrated growth inhibition against C. albicans and E. coli (Kim 188 et al., 2009). With the aim of exploring the effect of silver ion 189 donor type on Ag⁺ release and antimicrobial performance, we 190 describe here the characterisation of biosynthetic BC hydrogels 191 loaded with two forms of silver (AgNO₃ or AgZ) and establish 192

their *in vitro* antimicrobial activity against the common wound infecting pathogens, *S. aureus* and *P. aeruginosa*.

Material and methods

Microorganisms

G. xylinus ATCC 23770, P. aeruginosa NCIMB 8295 and S. aureus NCIMB 6571 were obtained from the University Wolverhampton culture collection. All three microorganisms were maintained at -20 °C in a freeze-dried form. Stock cultures of P. aeruginosa and S. aureus were resuscitated on sterile tryptone soy agar (TSA, prepared according to the manufacturer's protocol and sterilised by autoclaving before use) (Sigma-Aldrich, Irvine, UK), and incubated for 48 h at 37 °C. Stock culture of G. xylinus was resuscitated on sterile mannitol agar (composition: yeast extract (5 g/L), peptone (3 g/L), mannitol (25 g/L), agar (15 g/L); all materials were purchased from Lab M, Bury, UK) and incubated at 48 h at 30 °C. Prior to experimental use, overnight broth cultures were aseptically prepared in suitable broth using the stock plates.

Microbiological media

Dextrose, bacteriological peptone, yeast extract for the Hestrin and Schramm (HS) culture media were purchased from Lab M (Bury, UK). HS media was prepared following the standard protocol (Hestrin and Schramm, 1954). TSA, tryptone soya broth (TSB) (prepared according to the manufacturer's protocol and sterilised by autoclaving before use), disodium phosphate and citric acid were purchased from Sigma-Aldrich (Irvine, UK).

Materials

Cellulose (microcrystalline form) was purchased from Sigma-Aldrich (Irvine, UK). Zeolites (13X) were purchased from Laporte Inorganics (Widnes, UK) and AgZ were purchased from Sigma-Aldrich (Irvine, UK). Silver nitrate (AgNO₃) was from Fisher Scientific (Cramlington, UK) and sodium hydroxide from Acros Organics (UK). Ringer solution (1/4 strength) tablets were purchased from Lab M (Bury, UK) and prepared by dissolving one tablet in 500 mL of de-ionised water with constant magnetic stirring prior to sterilisation. Sterile 24-well tissue culture plates were purchased from Sarstedt (Leicester, UK).

Preparation and purification of BC pellicles

BC hydrogels were biosynthesised at 30 °C, under static conditions by *G. xylinus* in freshly prepared sterile HS culture medium. After 14 days, biosynthetic BC pellicles floated on top of the HS growth medium and after harvesting were purified by boiling in 1% w/v sodium hydroxide to obtain pure BC. The pellicles were further boiled in deionised water until the BC became clear and transparent.

Loading of silver in BC pellicles

Silver nitrate (AgNO₃)

The silver content in AgNO₃ used in this investigation was 63.5% w/w. BC pellicles were padded dry and loaded with aqueous AgNO₃ (0.55% w/v) by overnight incubation with constant agitation at 37 °C. The proportion of Ag⁺ in both formulations, i.e. AgNO₃ and AgZ was equivalent.

254

255

256

Silver zeolites

257

267

268

269

270

271

272

273

274

275

276

277

278

279

280

281

282

283

284

285

286

287

288

289

290

291

292

293

294

295

296

297

298

299

300

301

302

303

304

305

306

307

308

309

310

311

312

313

314

315

316

317

318

319

320

The silver content in the commercially available AgZ used in the 258 current study was 35% w/w. After purification, BC pellicles were 259 padded dry on filter paper and loaded with AgZ by overnight 260 immersion in 1% w/v aqueous AgZ suspension at 37 °C under 261 262 static conditions. It was subsequently noticed that some AgZ set-263 tled at the bottom of the vessel during loading, so the above pro-264 cedure was repeated by immersing freshly purified BC in an 265 aqueous AqZ suspension at 37 °C under constant agitation 266 overnight.

Characterisation of BC and silver loaded BC

Scanning electron microscopy (SEM) and energy dispersive X-ray (EDX) analysis

The samples were freeze dried for 48 h (Christ β 1,8-LSC plus, Martin Christ GmbH, Osterode am Harz, Germany) and coated with an ultrafine gold coating for SEM imaging. The morphology of BC-AgZ loaded under static and agitated conditions as well as BC-AgNO₃ was studied using Zeiss Evo[®]50 EP, SEM (Carl Zeiss AG, Oberkochen, Germany). Samples were also analysed by EDX (Zeiss Evo[®]50 EP, SEM) to confirm the presence of silver. Prior to analysis, each sample was placed on a stub and sprayed with compressed air to remove any traces of dust which can affect analysis.

Fourier transform infrared (FTIR) spectroscopy

Purified BC was analysed using FTIR spectroscopy (Genesis II FTIR[™], Thermo Scientific, Runcorn, UK) by first conducting a background scan followed by a scan of the sample to produce spectra. Pure cellulose, BC-AgNO₃ and BC-AgZ were also investigated in a similar way.

Swelling ratio

Swelling behaviour was investigated with BC pellicles $(9.25 \pm 0.25 \text{ cm} \text{ diameter})$ that were padded dry and the weight (W_x) recorded. Padded dry pellicles were soaked in 200 mL de-ionised water at 37 °C for 24 h under static conditions. After 24 h, the pellicles were withdrawn from water and the swollen BC gently wiped with filter paper prior to reweighing (W_s) . The swelling ratio (conducted in triplicate) was calculated as follows:



Silver release

Similar sized discs of BC-AgZ, BC-AgNO₃ and a negative control of BC loaded with zeolite (BC-Z) (all \approx 7.0 mm diameter) were cut using a corer. BC-Z, BC-AgZ and BC-AgNO₃ discs were transferred into individual wells of a 24-well tissue culture flat bottom plate, and 1 mL of freshly prepared sterile TSB (release media) was added. Discs were then incubated under static conditions at 37 °C for 96 h, and a 1 mL aliquot of release media withdrawn every 24 h; an equal volume (1 mL) of fresh sterile TSB was added at each time point to maintain sink conditions. Silver release was subsequently assessed by Inductively Coupled Plasma (ICP) spectrometry (Agilent Technologies 5100 ICP-OES, Aailent Technologies Inc., Santa Clara, CA). The kinetics of Ag⁺ release from BC-AgNO₃ and BC-AgZ were analysed by fitting the ICP data to zero order, first order, Higuichi and Korsmeyer-Peppas equations (Costa and Lobo, 2011). **9**4

Antimicrobial activity

The antimicrobial activity of BC-AgZ (loaded under both static and constantly agitated conditions) and BC-AgNO₃ was investigated

381

382

383

384

against *P. aeruginosa* and *S. aureus*, using the disc diffusion assay; purified BC and BC-Z (loaded under constant agitation) were used as controls. Discs of BC-AgZ, BC-AgNO₃, BC-Z and BC (\approx 7.0 mm diameter) were aseptically placed on TSA plates seeded with overnight cultures of *P. aeruginosa* or *S. aureus* and following incubation at 37 °C for 24 h, the zone of inhibition (ZOI) was measured. Individual discs were subsequently transferred onto freshly seeded plates (*P. aeruginosa* or *S. aureus*) to ensure consistent, reproducible microbial growth and incubated again under the same conditions for a further 24 h. Results are presented for ZOI at 24, 48, 72 and 96 h.

Statistical analysis

All experiments were performed in at least triplicate and data presented are means ± standard deviation (SD). Data recorded during the course of experiments were analysed statistically by the analysis of variance (ANOVA) using routines of the statistical software "SPSS 20" (SPSS Inc., Chicago, IL) and the differences between means were compared using the Least Significant Differences (LSD) at 5% level of probability (p < .05).

Results

Purification of biosynthetic BC pellicles

BC pellicles harvested after 2 weeks of growth from static HS *G. xylinus* cultures were opaque with extensive brown mottling due to residual media, entrapped bacterial cells and other fermentation debris (Figure 1(a)). After washing once in 1% w/v sodium hydroxide, followed by repeat washing in deionised water, purified pellicles became clear and transparent (Figure 1(b)).

BC-AgZ loading: static versus agitated conditions

Visual inspection of BC-AgZ loaded under static conditions revealed inconsistent loading (Figure 1(c)), where some areas of the BC display a higher AgZ content and others very little. BC-AgZ loaded under constant agitation appears to have a more even distribution of AgZ and hence more consistent loading (Figure 1(d)).

Scanning electron microscopy and EDX

Individual G. xylinus produces ribbons of cellulose that become entangled with each other, thus creating a dense network structure. Untreated BC pellicles reveal the presence of bacteria and other fermentation debris entrapped within the cellulose network (Figures 2(a) and 3(a)), but once purified, no bacteria were visible (Figure 2(b)). The purification process was also confirmed by EDX spectra, which indicate the removal of extraneous material from the BC (Figure 3(a,b)). BC pellicles have interwoven thick mats of cellulose fibres ranging between 0.01 and 0.15 μ m, that following freeze drying, have a dense, fibrous network interspersed with voids (Figure 2(b)). SEM images of freeze dried BC-AgZ (Figure 2(c,d)) and BC-AgNO₃ (Figure 2(e)) revealed AgZ and silver microcrystals entrapped within the dense cellulose network. After loading with AgZ under constant agitation, BC-AgZ displayed more consistent, uniform loading (Figure 2(d)) compared to that prepared under static conditions (Figure 2(c)). The presence of silver in both BC-AgZ and BC-AgNO3 was confirmed by EDX spectra (Figure 3(c,d)). Additionally, the presence of aluminium, silicon as well as elemental traces of calcium, sodium, magnesium and iron was also evident on EDX spectra of BC-AgZ (Figure 3(c)). Clusters of encapsulated AgZ particles ranged from 2.0 to 20.0 µm, with



Figure 2. SEM images of (a) untreated BC, entrapped residual *G. xylinus* highlighted; (b) purified BC, fibre network highlighted; (c) BC loaded with AgZ under static conditions, greater density of AgZ highlighted; (d) BC loaded with AgZ under constant agitation, AgZ highlighted, and (e) BC loaded with AgNO₃ under constant agitation, AgNO₃ impregnated in BC network highlighted.

single AgZ particles (0.5–5.0 μ m) also encapsulated within the BC (Figure 2(c,d)). Conversely, crystals of AgNO₃ entrapped within the BC fibres had an average range of 0.08–3.5 μ m (Figure 2(e)). In summary, the three-dimensional cellulose network consists of a large number of voids that become pores for sequestration of encapsulated silver microcrystals or AgZ when rehydrated.

Fourier transform infrared

The FTIR spectra of a commercially available cellulose sample, purified BC, BC-AgNO₃ and BC-AgZ are shown in Figure 4. Similarity in peak characteristics are shown in the spectra for the commercially available cellulose and purified BC at band regions





of 3330 cm^{-1} , 2894 cm^{-1} , 1641 cm^{-1} , 1370 cm^{-1} , 1159 cm^{-1} and 1056 cm^{-1} (Figure 4(a,b)); these peaks were also observed in both the BC-AgNO₃ and BC-AgZ spectra (Figure 4(c,d)). The characteristic peaks for AgNO₃ (733 cm⁻¹, 803 cm⁻¹ and 1300 cm⁻¹) and AgZ (434 cm⁻¹, 553 cm⁻¹, 665 cm⁻¹ and 980 cm⁻¹) were also observed in the spectra, thereby confirming the encapsulation of AgNO₃ or AgZ within the BC network.

Swelling behaviour

The absorption of surrounding fluids is a vital characteristic of hydrogels for potential application in the management of exudating wounds. After immersion in deionised water, padded dry BC pellicles imbibed an amount of water up to several times its original dehydrated weight, with a swelling ratio of 12.08 ± 0.96 after 24 h.





Figure 5. Silver release (ppm) over 96 h from BC-AqNO3 and BC-AqZ as deter-

Table 1. Mathematical modelling of Ag⁺ release kinetics.

/	Correlation coefficient (R ²)					
$\langle \rangle$	Zero order	First order	Higuichi	Korsmeyer–Peppas		
BC-AgZ	0.93	0.92	0.99	>0.99		
BC-AgNO ₃	0.84	0.69	0.98	0.99		

Lobo, 2011; Dash et al., 2010). Interestingly, Ag⁺ release from BC-AgZ also follows the Higuichi model with a correlation coefficient of >0.97 (R^2 =.99), i.e. diffusional release (Costa and Lobo, 2011). The additional diffusional release of Ag⁺ from AgZ at the surface of the BC may be due to ion exchange with cations present in the TSB media. The correlation coefficients for zero and first order models were both <0.97 (Table 1).

Silver release

mined by ICP (n = 4).

Ag⁺ release from BC hydrogels, as determined by ICP analysis, is presented in Figure 5. Results indicate that Ag⁺ release was greatest after 24 h from BC-AgZ (60.65 ± 4.18 ppm) and BC-AgNO₃ $(36.76 \pm 1.68 \text{ ppm})$. Silver release from BC-AgNO₃ plateaued after 72 h whereas from BC-AgZ release was steady and controlled (Figure 5). Overall, BC-AgZ released more silver $(137.60 \pm 16.17 \text{ ppm})$ over the 96 h compared to BC-AqNO₃ $(64.53 \pm 6.47 \text{ ppm}).$

Mathematical modelling of silver release

When silver release was analysed with zero order, first order, Higuchi and Korsmeyer-Peppas equations, results confirm that BC-AgZ and BC-AgNO₃ adhere to the Korsmeyer–Peppas model (Table 1). With a correlation coefficient of >0.97 ($R^2 \ge .99$ for both BC-AgZ and BC-AgNO₃) and release exponent between 0.45 and 0.89 (BC-AgZ = 0.73; BC-AgNO₃ = 0.56) which indicates that Ag⁺ release is non-Fickian (anomalous transport) (Costa and

Antimicrobial activity

Pure BC and BC-Z exhibited no antimicrobial activity. There was a difference in antimicrobial activity for AgZ loaded BC via static and agitation methods (Figure 6(a,b)). The antimicrobial activity of BC-AgZ loaded under constant agitation was significantly better against P. aeruginosa (BC-AgZ (static): 8.29 mm \leq ZOI \leq 14.15 mm; BC-AgZ (agitated): 16.10 mm \leq ZOI \leq 17.96; p < .05). However, this was not observed in the case of S. aureus when comparing the antimicrobial activity of BC-AgZ loaded under static or agitated condition (BC-AgZ (static): 7.94 mm \leq ZOI \leq 10.31 mm; BC-AgZ (agitated): 7.71 mm \leq ZOI \leq 10.06 mm; p > .05).

The disc diffusion assay for antimicrobial activity indicated that both BC-AgZ and BC-AgNO₃ exhibited higher antimicrobial activity against P. aeruginosa compared to S. aureus (Figure 7(a,b)). Statistical analysis using ANOVA suggests that there is a significant difference between the ZOI of *P. aeruginosa* and *S. aureus* (p < .05) when treated with BC-AgZ. This is also the case for both strains (p < .05) when treated with BC-AgNO₃.

The disc diffusion assay results indicated that BC-AgNO₃ had higher antimicrobial activity against P. aeruginosa up to 24 h when





Figure 6. Antimicrobial activity assessed by ZOI during the disc diffusion assay for BC-AgZ loaded under (a) static and (b) constantly agitated conditions against *P. aeruginosa* and *S. aureus* (n = 6; error bars = SD).

compared to BC-AgZ. After 24 h, the antimicrobial activity of BC-AgNO₃ reduced and BC-AgZ exhibited higher activity against *P. aeruginosa* (Figure 7(a)). Antimicrobial activity against *S. aureus* also had the same trend at 24 h with BC-AgNO₃ exhibiting larger ZOI compared to BC-AgZ. At 48 h, there was no significant difference between BC-AgNO₃ and BC-AgZ against *S. aureus*; following incubation at 72 and 96 h, BC-AgNO₃ had highly reduced antimicrobial activity against *S. aureus* (ZOI =0 mm), whereas BC-AgZ still exhibited some antimicrobial activity at 72 h (ZOI = 7.26±0.53 mm), which was further reduced at 96 h (ZOI = 7.11±0.33 mm) (Figure 7(b)). The difference in activity of BC-AgZ and BC-AgNO₃ against *P. aeruginosa* (Gram negative) and *S. aureus* (Gram positive) bacteria may be due to their structural differences (see section "Discussion" for details).

Discussion

The concept of moist healing has revolutionised the field of wound management and led to an increased interest in the use of hydrogel-based dressings. In the present study, biosynthetic BC hydrogels were loaded with an Ag⁺ donor, AgNO₃ or AgZ, and characterised in vitro for their application as antimicrobial wound dressings. Once purified, these BC hydrogels become clear and transparent which is advantageous in the management of infected, heavily exudating wounds, as it allows the healing process to be monitored without removing the dressing, hence damaging fragile underlying tissue.

FTIR confirmed that the purified biosynthetic BC hydrogel produced by *G. xylinus* closely resembles commercial cellulose. The broad, characteristic peak at 3328 cm^{-1} falls within the region of $3200-3550 \text{ cm}^{-1}$ which is contributed by the stretching of O-H bond (El-Shishtawy et al., 2011). This may also be due to the



Figure 7. Antimicrobial activity assessed by ZOI during the disc diffusion assay for BC-AgZ and BC-AgNO₃ against (a) *P. aeruginosa* and (b) *S. aureus* (n = 9; error bars = SD).

stretching of the intramolecular hydrogen bond of 30…H–O5 within the BC network which has been reported to show a characteristic peak at 3348 cm^{-1} (Oh et al., 2005). The narrow peak at 2900 cm⁻¹ is due to C–H stretching of CH_2 and CH_3 group, whereas the peak at 1640 cm⁻¹ is due to the O-H bending of water molecules (Barud et al., 2008). The peak at 1375 cm⁻¹ (C-H bending) provides an indication for the presence of crystalline regions within the BC structure (Castro et al., 2011). Besides that, the peaks at 1159 cm⁻¹ (asymmetrical C–O–C bridge stretching) and 1055 cm⁻¹ (skeletal vibrations involving C–O stretching) can also be attributed to BC (Barud et al., 2008). The encapsulation of AgNO₃ into the BC is confirmed by the occurrence of the specific peaks for Ag at 733 cm⁻¹ and 803 cm⁻¹ (Valverde-Aguilar et al., 2011). The vibration bands within the region of $1350-400 \,\mathrm{cm}^{-1}$ are attributed to nitrate ions (NO³⁻) (Salim and Malik, 2016); hence the presence of a peak at 1370 cm^{-1} , indicates the availability of the nitrate group within BC-AgNO₃. The peaks of BC-AgZ at 980 cm⁻¹ (asymmetric vibration of Si–O) suggest the presence of a three-dimensional silica phase within the zeolite structure and the peak at 665 cm⁻¹ (symmetric stretch) may be contributed by the internal vibrations of the tetrahedral framework (Shameli et al., 2011). FTIR spectra of zeolites reveal the occurrence of a large, intense band at 986 cm⁻¹ corresponding to the vibration of the Si–O–Si (Hanim et al., 2016), whereas the peak at 676 cm^{-1} may be due to the bending of Al-O bonds within the zeolite structure (Shameli et al., 2011). Additionally, peaks occurring within the region of 420–500 cm⁻¹ may also indicate internal vibrations due to the bending of the T-O tetrahedra, e.g. 477 cm^{-1} corresponding to the internal vibration of (Si, Al)O₄ tetrahedra within the zeolite structure (Karimi-Shamshabadi and Nezamzadeh-Ejhieh, 2016). The presence of silver within the 897

898

899

BC-AgZ network is confirmed by the peak at 553 cm⁻¹, that indicates the Ag–O stretching of AgO and Ag₂O within the 500–600 cm⁻¹ region (Waterhouse et al., 2001; Kim et al., 2013).

The swelling ratio of a hydrogel indicates its ability to absorb 900 fluid and is an important property for dressings, especially those 901 used to manage heavily exudating wounds. Hydrogel wound 902 dressings with high degrees of swelling can be applied to a wide 903 904 variety of wound types, ranging from dry necrotic wounds to full-905 thickness wounds that produce a high volume of exudate (Kokabi et al., 2007). BC is insoluble in water and most organic solvents, 906 but when immersed in aqueous media its fibrous structure 907 imbibes large amounts of fluid and it swells. The swelling behav-908 iour of this biosynthetic BC hydrogel reveal a high swelling cap-909 acity with a swelling ratio of 12.08±0.96 after 24 h, which is in 910 911 agreement with that reported by other groups (e.g. Nakayama et al., 2004; Maneerung et al., 2008; Wei et al., 2011). The high 912 swelling of BC results from the development of hydrogen bonds 913 with water molecules and the network-like structure of the bio-914 polymer itself that allows BC to imbibe and hold onto those mole-915 cules within the crosslinked polymer voids (Maneerung et al., 916 2008). This high water absorptivity enables BC to maintain a moist 917 918 environment at the wound site by donating or receiving fluids, 919 thus maintaining an optimal environment in which healing can progress (Lin et al., 2013). 920

Several methods have been reported for the determination of 921 small molecule antimicrobial release from wound dressings 922 (including metal ions) such as beaker, diffusion cell, paddle over 923 disc and two compartment model (Maneerung et al., 2008; Wei 924 et al., 2011; Jadhav et al., 2012; Peršin et al., 2014; Wu et al., 925 2014). Each method has unique advantages and disadvantages 926 927 and hence no single, standard method has been adopted for the determination of silver release from dressings. In this study, a 928 refined silver release method using 24-well tissue culture flat bot-929 tom plates and TSB as release media was developed. At the 930 wound site, silver release from dressings is dependent on several 931 932 factors including the type of wound, classification and size of 933 dressing, volume of exudate, etc. The authors are mindful that the method designed and employed for the determination of silver 934 release in the current study may not exactly mirror the real 935 wound environment; for that to occur the presence of phago-936 cytes, inflammatory mediators, hydrolytic enzymes, reactive oxy-937 gen species, bacteria and their associated by-products would 938 need to be considered (Ovington, 2007; Martin et al., 2013). 939 Nevertheless, this method has several advantages due to its sim-940 plicity, low cost and biorelevant temperature conditions. Silver 941 release, as guantified by ICP indicated that the Ag⁺ zeolite-loaded 942 hydrogels (BC-AgZ) released more Ag⁺ than their AgNO₃ counter-943 parts (BC-AgNO₃). The microporous zeolite framework allows 944 greater initial Ag⁺ loading and a subsequent ion-exchange medi-945 946 ated release of biologically active Ag⁺ into the surrounding 947 medium (Kwakye-Awuah et al., 2008).

These experiments revealed that BC loaded under conditions of constant agitation demonstrated greater antimicrobial activity, compared to the statically loaded hydrogels. This may be due to the more evenly distributed Ag^+ loading under agitated condition as compared to static incubation, which contributed to a more consistent release of Ag^+ (Figure 2(c,d)).

The prolonged antimicrobial activity exhibited by BC-AgZ against both *P. aeruginosa* and *S. aureus* resulted from the controlled release of Ag⁺ from the zeolite structure. Zeolites are crystalline aluminosilicate cages with cavities occupied by cations or water molecules. The AgZ acts as an inorganic reservoir and silver in the zeolite have considerable freedom of movements (Kwakye-Awuah et al., 2008). In addition to the ionic exchange-based release of Ag⁺ encapsulated within the zeolite, BC matrix was able to provide a second layer of controlled release. In contrast, release of Ag⁺ from BC-AgNO₃ was only controlled by the BC matrix, hence Ag⁺ release was less prolonged when compared to BC-AgZ.

Results revealed that both BC-AgZ and BC-AgNO₃ exhibited antimicrobial activity against the Gram positive and Gram negative strains tested, but that activity was stronger and more prolonged against P. aeruginosa compared to S. aureus. Ag⁺ has strong bactericidal activity (Copcia et al., 2011), and Gram negative bacteria are more susceptible to Ag⁺ than the Gram positive species (Ip et al., 2006; Waghmare et al., 2015) due to differences in their cell wall structure and composition. The Gram positive bacterial cell wall contains a thicker peptidoglycan layer than that of Gram negative strains. Gram positive bacterial cell walls typically lack an outer membrane and are mainly composed of a thick, negatively charged peptidoglycan layer and cytoplasmic phospholipid bilayer. Contrary to this, Gram negative microorganisms have a thin (\approx 2–3 nm) peptidoglycan layer between the outer membrane and the cytoplasmic phospholipid bilayer (Le et al., 2010). The composition of the peptidoglycan layer, which contains teichoic acids, contributes to the overall anionic charge of the Gram positive cell surface (Neuhaus and Baddiley, 2003); this greater net anionic charge may bind more Ag⁺, thus reducing the amount that can reach the plasma membrane and intracellular targets to exert their antimicrobial activity. This finding is in agreement with Le et al. (2010) who reported that E. coli (Gram negative) was more sensitive to the effects of silver nanoparticles (versus S. aureus, a Gram positive) because of differences in the thickness of the peptidoglycan layer (Le et al., 2010). Gram negative microorganisms are generally less sensitive to antibiotics and certain antimicrobial agents due to the selective permeability and protective mechanism of their outer membrane structure (Bomberger et al., 2009; Sperandeo et al., 2009). Nevertheless, Ag⁺ has shown its ability to exert microbiocidal activity against a variety of pathogenic microorganisms, including drug resistant strains. In addition, its multi-target antimicrobial activities are advantageous in limiting the potential development of resistant microbial strains (Brett, 2006; Radecka et al., 2015). Hence the development of a responsive, topical silver formulation would be extremely useful for the treatment of infected chronic wounds.

Conclusions

The present study demonstrates the *in vitro* stability, performance and antimicrobial potential of biosynthetic BC hydrogels loaded with $AgNO_3$ or AgZ against representative wound infecting microorganisms (*P. aeruginosa* and *S. aureus*). The moist and responsive nature of BC hydrogels make them an ideal biomaterial dressing for the management of chronic, infected wounds. Further research aims to optimise the performance of BC hydrogels loaded with microencapsulated silver to provide a responsive, controlled release delivery platform, whilst minimising the toxicity associated with localised high concentrations of topical silver.

Disclosure statement

The authors report no conflicts of interest. The authors alone are responsible for the content and writing of the article.

ORCID

1016

1017

1018

1019

1020

1021

1022

1023

1024

 \mathcal{O}

1025References1026

- 1027 Abdelrahman T, Newton H. Wound dressings: Principles and prac-1028 tices. Surgery, 2011;29(10):491–5.
- 1029Abeer MM, Amin MCIM, Martin C. A review of bacterial cellulose-
based drug delivery systems: Its' biochemistry, current
approaches and future prospects. J Pharm Pharmacol,
2014;66(8):1047–61.
- 1033Atiyeh BS, Costagliola M, Hayek SN, Dibo SA. Effect of silver on1034burn wound infection control and healing: Review of the litera-1035ture. Burns, 2007;33(2):139–48.
- 1036Barud HS, Assunção RMN, Martines MAU, Dexpert-Ghys J, Marques1037RFC, Messaddeq Y, Ribeiro SJL. Bacterial cellulose-silica organi-1038c-inorganic hybrids. J Sol-Gel Sci Technol, 2008;46(3):363-7.
- 1039Bomberger JM, MacEachran DP, Coutermarsh BA, Ye S, O'Toole1040GA, Stanton BA. Long-distance delivery of bacterial virulence1041factors by *Pseudomonas aeruginosa* outer membrane vesicles.1042PLoS Pathog, 2009;5(4):e1000382 [online] Available at: http://1043journals.plos.org/plospathogens/article?id=10.1371/journal.ppat.10441000382.
- 1045Brett DW. A discussion of silver as an antimicrobial agent:1046Alleviating the confusion. Ostomy/Wound Manage, 2006;52(1):104734–41.
- 1048Castellano JJ, Shafii SM, Ko F, Donate G, Wright TE, Mannari RJ,1049Payne WG, Smith DJ, Robson MC. Comparative evaluation of sil-1050ver-containing antimicrobial dressings and drugs. Int Wound J,10512007;4(2):114–22.
- Castro C, Zuluaga R, Putaux JL, Caro G, Mondragon I, Ganán P.
 Structural characterization of bacterial cellulose produced by
 Gluconacetobacter swingsii sp. from Colombian agroindustrial
 wastes. Carbohydr Polym, 2011;84(1):96–102.
- 1056Chopra I. The increasing use of silver-based products as antimicro-
bial agents: A useful development or a cause for concern? J
Antimicrob Chemother, 2007;59(4):587–90.
- 1059 Copcia VE, Luchian C, Dunca S, Bilba N, Hristodor CM.
 1060 Antibacterial activity of silver-modified natural clinoptilolite.
 1061 J Mater Sci, 2011;46(22):7121–8.
- Costa P, Lobo JMS. Modeling and comparison of dissolution pro files. Eur J Pharm Sci, 2011;13(2):123–33.
- 1064Czaja W, Krystynowicz A, Bielecki S, Brown MB. Microbial cellulose1065- the natural power to heal wounds. Biomaterials,10662006;27(2):145–51.
- Dash S, Murthy PN, Nath L, Chowdhury P. Kinetic modeling on drug release from controlled drug delivery systems. Acta Pol Pharm. Drug Res, 2010;67(3):217–23.
- El-Shishtawy RM, Asiri AM, Abdelwahed NA, Al-Otaibi MM. In situ
 production of silver nanoparticle on cotton fabric and its anti microbial evaluation. Cellulose, 2011;18(1):75–82.
- Fu L, Zhang J, Yang G. Present status and applications of bacterial
 cellulose-based materials for skin tissue repair. Carbohydr
 Polym, 2013;92(2):1432–42.
- 1076Guo S, DiPietro LA. Factors affecting wound healing. J Dent Res,10772010;89(3):219–29.
- Hanim SAM, Malek NANN, Ibrahim Z. Amine-functionalized, silverexchanged zeolite NaY: Preparation, characterization and antibacterial activity. Appl Surf Sci, 2016;360(Part A):121–30.
- 1081Hestrin S, Schramm M. Synthesis of cellulose by Acetobacter xyli-1082num. 2. Preparation of freeze-dried cells capable of polymeriz-1083ing glucose to cellulose. Biochem J, 1954;58(2):345–52.
- 1084 Ip M, Lui SL, Poon VK, Lung I, Burd A. Antimicrobial activities of silver dressings: An *in vitro* comparison. J Med Microbiol, 2006;55(Pt 1):59–63.
 1087

1088

- Jadhav H, Joshi A, Misra M, Shahiwala A. Effects of various formulation parameters on the properties of hydrogel wound dressings. Drug Deliv Lett, 2012;2(1):8–13.
- Kamoun EA, Chen X, Eldin MSM, Kenawy E-R. Crosslinked poly(vinyl alcohol) hydrogels for wound dressing applications: A review of remarkably blended polymers. Arab J Chem, 2015;8(1):1–14.
- Karimi-Shamsabadi M, Nezamzadeh-Ejhieh A. Comparative study on the increased photoactivity of coupled and supported manganese-silver oxides onto a natural zeolite nano-particles. J Mol Catal A: Chem, 2016;418–419:103–14.
- Kim J, Kwon S, Ostler E. Antimicrobial effect of silver-impregnated cellulose: Potential for antimicrobial therapy. J Biol Eng, 2009;3(20) [online] doi:10.1186/1754-1611-3-20. Available at: http://www.jbioleng.org/content/3/1/20.
- http://www.jbioleng.org/content/3/1/20. Kim PS, Kim MK, Cho BK, Nam IS, Oh SH. Effect of H₂ on deNOx performance of HC-SCR over Ag/Al₂O₃: Morphological, chemical, and kinetic changes. J Catal, 2013;301:65–76.
- Kirketerp-Møller K, Jensen PO, Fazli M, Madsen KG, Pedersen J, Moser C, Tolker-Nielsen T, Høiby N, Givskov M, Bjarnsholt T. Distribution, organization and ecology of bacteria in chronic wound. J Clin Microbiol, 2008;46(8):2717–22.
- Kokabi M, Sirousazar M, Hassan ZM. PVA-clay nanocomposite hydrogels for wound dressing. Eur Polym J, 2007;43(3):773-81.
- Kwakye-Awuah B, Wemegah DD, Nkrumah I, Williams C, Radecka I. Antimicrobial activity of silver-zeolite LTA on heavily-contaminated underground Ghanaian waters. Int J Sci Res, 2013;2(11):26–31.
- Kwakye-Awuah B, Williams C, Kenward MA, Radecka I. Antimicrobial action and efficiency of silver-loaded zeolite X. J Appl Microbiol, 2008;104(5):1516–24.
- Landis SJ. Chronic wound infection and antimicrobial use. Adv Skin Wound Care, 2008;21(11):531–40.
- Le AT, Tam LT, Tam PD, Huy PT, Huy TQ, Hieu NV, Kudrinskiy AA, Krutyakov YA. Synthesis of oleic acid-stabilized silver nanoparticles and analysis of their antibacterial activity. Mater Sci Eng: C, 2010;30(6):910–16.
- Leaper D. Appropriate use of silver dressings in wounds: International consensus document. Int Wound J, 2012;9(5):461-4.
- Lin WC, Lien CC, Yeh HJ, Yu CM, Hsu SH. Bacterial cellulose and bacterial cellulose-chitosan membranes for wound dressing applications. Carbohydr Polym, 2013;94(1):603–11.
- Low WL, Martin C, Hill DJ, Kenward MA. Antimicrobial efficacy of silver ions in combination with tea tree oil against *Pseudomonas aeruginosa, Staphylococcus aureus* and *Candida albicans*. Int J Antimicrob Agents, 2011;37(2):162–5.
- Low WL, Martin C, Hill DJ, Kenward MA. Antimicrobial efficacy of liposome encapsulated silver ions and tea tree oil against *Pseudomonas aeruginosa, Staphylococcus aureus* and *Candida albicans*. Lett Appl Microbiol, 2013;57(1):33–9.
- Maneerung T, Tokura S, Rujiravanit R. Impregnation of silver nanoparticles into bacterial cellulose for antimicrobial dressing. Carbohydr Polym, 2008;72(1):43–51.
- Martin C, Low WL, Amin MCIM, Radecka I, Raj P, Kenward MA. Current trends in the development of wound dressings, biomaterials and devices. Pharm Patent Anal, 2013;2(3):341–59.
- Martin C, Low WL, Gupta A, Amin MCIM, Radecka I, Britland ST, Raj P, Kenward MA. 2014. Liposomal delivery of antimicrobials. In: Finney L, ed. Advances in liposome research (part of the Biochemistry Research Trends Series). New York, USA: Nova Science Publishers Inc., pp. 27–62.

1136

1137

1138 1139

1140

1141

1142

1143

1144

1145

1146

1147

1148

1149

1150

1151

1152

1089

1090

1153

1154

1155

1156

1157

1158

1159

1160

1162

1163

1164

1165

1166

1167

1168

1169

1170

1171

1172

1194

1195

1196

1197

1198

1199

1200

1201 1202

1203

1204

1205

1206

1207

1208

1209

1210

1211

1212

1213

1214

1215

1216

- Martin C, Low WL, Gupta A, Amin MCIM, Radecka I, Britland ST, Raj P, Kenward MA. Strategies for antimicrobial drug delivery to biofilm. Curr Pharm Des, 2015;21(1):43-66.
- Matsumura Y, Yoshikata K, Kunisaki SI, Tsuchido T. Mode of bactericidal action of silver zeolite and its comparison with that of silver nitrate. Appl Environ Microbiol, 2003;69(7):4278-81.
- Murphy PS, Evans GRD. Advances in wound healing: A review of current wound healing products. Plast Surg Int, 2012;2012: Article ID 190436, doi:10.1155/2012/190436.
- 1161 Nakayama A, Kakugo A, Gong JP, Osada Y, Takai M, Erata T, Kawano S. High mechanical strength double-network hydrogel with bacterial cellulose. Adv Funct Mater, 2004;14(11):1124-8.
 - Neuhaus FC, Baddiley J. A continuum of anionic charge: Structures and functions of d-alanyl-teichoic acids in Gram-positive bacteria. Microbiol Mol Biol Rev, 2003;67(4):686-723.
 - Oh SY, Yoo DI, Shin Y, Kim HC, Kim HY, Chung YS, Park WH, Youk JH. Crystalline structure analysis of cellulose treated with sodium hydroxide and carbon dioxide by means of X-ray diffraction and FTIR spectroscopy. Carbohydr Res, 2005;340(15):2376-91.
- Ovington LG. Advances in wound dressings. Clin Dermatol, 1173 2007;25(1):33-8. 1174
- Peršin Z, Maver U, Pivec T, Maver T, Vesel A, Mozetič M, Stana-1175 Kleinschek K. Novel cellulose based materials for safe and effi-1176 cient wound treatment. Carbohydr Polym, 2014;100:55-64. 1177
- Pinto RJ, Margues PA, Neto CP, Trindade T, Daina S, Sadocco P. 1178 Antibacterial activity of nanocomposites of silver and bacterial 1179 or vegetable cellulosic fibers. Acta Biomater, 2009;5(6):2279-89. 1180
- Radecka I, Martin C, Hill DJ. 2015. The problem of microbial drug 1181 1182 resistance. In: Phoenix DA, Harris F, Dennison SR, eds. Novel antimicrobial agents and strategies. Weinheim, Germany: Wiley-1183 VCH, pp. 1–16 1184
- Salim MM, Malek NANN. Characterization and antibacterial activity 1185 of silver exchanged regenerated NaY zeolite from surfactant-1186 modified NaY zeolite. Mater Sci Eng: C, 2016;59:70-7. 1187
- Sannino A, Demitri C, Madaghiele M. Biodegradable cellulose-1188 based hydrogels: Design and applications. Materials, 2009;2(2): 1189 353-73. 1190
- Shah N, Ul-Islam M, Khattak WA, Park JK. Overview of bacterial cel-1191 lulose composites: A multipurpose advanced material. 1192 Carbohydr Polym, 2013;98(2):1585-98. 1193

- Shameli K, Ahmad MB, Zargar M, Yunus WMZW, Ibrahim NA. Fabrication of silver nanoparticles doped in the zeolite framework and antibacterial activity. Int J Nanomed, 2011;6:331-41.
- Singh B, Sharma S, Dhiman A. Design of antibiotic hydrogel wound dressing: Biomedical properties and histological study of wound healing. Int J Pharm, 2013;457(1):82-91.
- Solway DR, Fapwca F, Consalter M, Levinson DJ. Microbial cellulose wound dressing in the treatment of skin tears in the frail elderly. Wounds, 2010;22(1):17-19.
- Sperandeo P, Dehò G, Polissi A. The lipopolysaccharide transport system of Gram-negative bacteria. Biochim Biophys Acta (BBA)-Mol Cell Biol Lipids, 2009:1791(7):594-602.
- Valverde-Aguilar G, García-Macedo JA, Rentería-Tapia VM, Gómez RW, Quintana-García M. Modelling of optical absorption of silver NP's produced by UV radiation embedded in mesostructured silica films. J Nanopart Res, 2011;13(10):4613-22.
- Waghmare SR, Mustopa NM, Suryakant RM, Kailas DS. Ecofriendly production of silver nanoparticles using Candida utilis and its mechanistic action against pathogenic microorganisms. 3 Biotech, 2015;5(1):33-8.
- Waterhouse GI, Bowmaker GA, Metson JB. The thermal decomposition of silver (I, III) oxide: A combined XRD, FT-IR and Raman spectroscopic study. Phys Chem Chem Phys, 2001;3(17):3838-45.
- Wei B, Yang G, Hong F. Preparation and evaluation of a kind of bacterial cellulose dry films with antibacterial properties. Carbohydr Polym, 2011;84(1):533-8.

Wilkinson LJ, White RJ, Chipman JK. Silver and nanoparticles of silver in wound dressings: A review of efficacy and safety. J Wound Care, 2011;20(11):543-9.

Winter GD. Formation of the scab and rate of epithelialization of superficial wounds in the skin of the young domestic pig. Nature, 1962:193:293-4.

- Wu J, Zheng Y, Song W, Luan J, Wen X, Wu Z, Chen X, Wang Q, Guo S. In situ synthesis of silver-nanoparticles/bacterial cellulose composites for slow-released antimicrobial wound dressing. Carbohydr Polym, 2014;102:762-71.
- Yang G, Xie J, Hong F, Cao Z, Yang X. Antimicrobial activity of silver nanoparticle impregnated bacterial cellulose membrane: Effect of fermentation carbon source of bacterial cellulose. Carbohydr Polym, 2012;87(1):839-45.