

# **Analysis of linezolid and tigecycline as candidates for local prophylaxis via antibiotic-loaded bone cement**

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25 strength of antibiotic-loaded samples was determined using a Charpy-type impact testing 26 apparatus. Cytotoxicity of eluted antibiotics against MG-63 cells was evaluated using an 27 MTT assay.

28

29 **Results**: Linezolid and tigecycline eluted from bone cement to clinically relevant levels 30 within 1 hour and retained activity over 1 week. Mechanical wear significantly reduced 31 elution of tigecycline but had little effect on elution of linezolid. Linezolid showed low 32 cytotoxicity towards MG-63 cells with  $\leq$  300 mg/mL resulting in  $>$ 50 % cell activity. 33 Cytotoxicity of tigecycline was higher, with an  $IC_{50}$  of 5-10 mg/L.

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35 **Conclusions**: Linezolid and tigecycline retain activity after elution from bone cement. The 36 concentration of tigecycline may need to be carefully controlled due to cytotoxicity. The 37 effect of wear on bone cement may need to be considered if tigecycline is to be used for local 38 delivery. Up to 10% linezolid can be added without affecting the impact strength of the bone 39 cement. These results are promising indications for future investigation of these antibiotics 40 toward use in local antibiotic delivery strategies.

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#### 42 **Introduction**

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44 Prosthetic joint infections present a rare but major complication in arthroplastic surgery. The 45 incidence of infection across all arthroplastic procedures has been reported as ranging from 1  $-46 - 3\%$ .<sup>1-3</sup> Revision surgery to remedy an infected joint prosthesis is associated with increased 47 costs, longer stay in hospital and potential morbidity, compared to revision surgery after 48 aseptic failure.<sup>4-6</sup> The number of arthroplastic procedures and the incidence of infection have 49 increased over the last 10 years, as have the total costs associated with revision surgery.<sup>4,5,7</sup>

50 As the demand for arthroplastic surgery progressively rises, the costs associated with 51 prosthetic joint infection are set to increase greatly. This has led to perioperative antibiotic 52 prophylaxis strategies including the use of antibiotic-loaded bone cement becoming 53 routine. $8,9$ 

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55 The management of a prosthetic joint infection involves removal of the infected prosthesis 56 and radical debridement of the surrounding infected tissue. This is followed by either a one-57 stage revision where a new prosthesis is implanted in a single procedure or a two-stage 58 revision where a temporary spacer is used for several weeks before the new prosthesis is 59 implanted. In both procedures antibiotic therapy is standard practice, commonly combining 60 systemic antibiotic treatment with local delivery using antibiotic-loaded bone cement. 61 Antibiotic-loaded cement is used to cement the prosthesis into place and, in the two-stage 62 revision, is used to form the temporary spacer.<sup>10</sup>

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64 Antibiotic-resistant organisms such as methicillin-, vancomycin- and multidrug resistant 65 strains are increasingly becoming associated with failure of revision surgery. More than 50% 66 of all prosthetic joint infections are caused by staphylococci such as *Staphylococcus aureus* 67 and *Staphylococcus epidermidis* and it has been estimated that around half of all *S. aureus*-68 related periprosthetic joint infections are now methicillin resistant.<sup>1,11-13</sup> The ability of these 69 organisms to acquire antibiotic resistance requires the use of new antibiotics to be explored 70 for use in bone cement.

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72 Here we evaluate linezolid and tigecycline for use in antibiotic-loaded bone cement systems 73 and assess their suitability for this application. There are few studies investigating the inclusion of linezolid in bone cement<sup>14,15</sup> and, to our knowledge, there are no published data



- 100 **MG63 cell culture**
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102 Cells were cultured on Eagles minimal essential medium (EMEM) containing 10 % fetal 103 bovine serum  $(v/v)$ , 2 mM glutamine and 1 % non-essential amino acids  $(v/v)$ . Cells were

- 104 incubated at 37 $^{\circ}$ C (5 % CO<sub>2</sub>) and passaged three times a week.
- 105
- 106 **MTT assay**

- 108 MG63 cells were seeded at  $2 \times 10^3$  cells per well in 100 µL of EMEM containing the 109 appropriate concentration of antibiotic. Cells were incubated at  $37^{\circ}$ C (5 % CO<sub>2</sub>) for 48 h. 110 After 48 h the medium was removed and fresh medium added. A 12 mM stock solution of 111 MTT was prepared and 10  $\mu$ L added to each well before incubating at 37°C (5 % CO<sub>2</sub>) for 4 112 h. An SDS-HCl (100 mg/mL, 0.01M HCl) stock solution was prepared and 100 µL added to 113 each well before incubating for a further 4 h. Absorbance was measured at 570 nm and 114 compared to positive control cultures containing no antibiotic. 115 116 **Preparation of bone cement**  117 118 Linezolid, tigecycline and gentamicin-containing bone cement samples were prepared by 119 hand-mixing antibiotic powder (3% or 10% wt/wt) with Biomet Bone Cement  $R^{\circledast}$  powder 120 until a homogenous mix was produced. The antibiotic cement powder was then mixed with 121 the appropriate amount of polymethylmethacrylate (PMMA) monomer liquid in a Hi-Vac
- 122 bone cement mixing bowl (Biomet) as per the manufacturer's instructions. Refobacin Bone
- 123 Cement  $R^{\circledast}$  and Bone cement R (Biomet) were also prepared in a Hi-Vac bone cement
- 124 mixing bowl (Biomet) as per the manufacturer's instructions. The bone cement was placed



150 perimeter of the wearing cement sample. A container was placed beneath the assembly and 151 filled with 0.1 M ammonium acetate solution (pH 7.4) until the lower portion of the cement 152 sample was submerged. A magnetic stirrer was used to mix the solution in the container at 153 300 rpm and samples were rotated against the HA counter-face at 60 rpm for 51 h. The HA 154 counter-face was repositioned every 10 - 12 h to ensure a sufficiently abrasive counter-face 155 throughout the experiment. An extension shaft was fitted to the TE-66 to allow simultaneous 156 rotation of an unworn control sample at the same speed. This sample was also partially 157 submerged in a separate container filled with 0.1M ammonium acetate solution (pH 7.4). The 158 experiment was placed in a UV-sealed air-tight container and the temperature and humidity 159 constantly measured during the experiment. At regular intervals, 200 µL aliquots of solution 160 were taken and stored at -20°C before analysis.

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162 **Quantification of antibiotics by LC-MS** 

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164 Detection of linezolid was carried out on a Phenomenex Luna  $C_{18}$  reversed phase column 165 (150 mm x 1 mm) attached to a Finnigan LCQ ESI-MS. The isocratic mobile phase was 0.1% 166 aqueous trifluoroacetic acid (TFA)/acetonitrile (77:23) and the flow rate was 0.05 mL/min. 167 Measurement of linezolid concentration was carried out by monitoring the protonated parent 168 ion at m/z 338.2 and comparing the results to a standard curve. Quantification of tigecycline 169 was carried out as described above except the isocratic mobile phase was 0.1% aqueous 170 TFA/methanol (67:33) and monitoring the protonated parent ion at m/z 586.5. 171

# 173 **Impact strength analysis**



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198 concentration of 0.66 \pm 0.35 mg/L after one hour and then decreased to 0.084 \pm 0.025 mg/L
199 after 24 h and 0.014 \text{ mg/L} \pm 0.013 after 168 h (Fig 2). The initial elution rate of tigecycline
200 from bone cement was calculated as 32.8 \pm 17.2 \mug/hour/g bone cement.
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#### 202 **Effect of wear on elution of bone cement**

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204 The results from three separate experiments to investigate the effect of wear on elution 205 behaviour of cement containing 3 % (wt/wt) tigecycline are shown in Fig. 3. The samples 206 were collected over a 51 h period and the maximum concentration of eluted antibiotic was 207 reached between 5 h and 12 h. The highest concentration overall was seen in the unworn 208 sample 2 after 12 h with a concentration of 2.1 mg/L compared to 0.1 mg/L in the worn 209 counterpart (Fig 3b). Although there is some variability in the maximum concentrations 210 between the three experiments, in all cases a clear trend can be seen with the elution from 211 unworn samples being significantly higher than the worn bone cement samples  $(P < 0.05)$ . After 1 hour the elution of tigecycline from unworn samples was  $9.4 \pm 2.6 \,\mu$ g/hour/cm<sup>3</sup> 212 213 surface and the rate of elution from the worn samples was  $2.3 \pm 2.5 \,\mu$  g/hour/cm<sup>3</sup> surface.

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215 The results from three separate experiments to investigate the effect of wear on elution 216 behaviour of cement containing 3 % (wt/wt) linezolid are shown in Fig. 4. The samples were 217 collected over a 51 h period and the maximum concentration of eluted antibiotic was reached 218 between 24 h and 51 h with concentration continuing to increase in all but one sample. The 219 highest concentration overall was seen in the worn sample 2 after 51 h with a concentration 220 of 53.1 mg/L (Fig 4b). No significant difference can be seen in the elution kinetics between 221 the worn and unworn linezolid samples  $(P = 0.63)$ . After 1 hour the rate of elution from

222 unworn linezolid samples was  $232.5 \pm 22.4 \,\mu$ g/hour/cm<sup>3</sup> surface and the rate of elution from 223 the worn linezolid samples was  $242.4 \pm 24.3 \mu$ g/hour/cm<sup>3</sup> surface. The rates of antibiotic 224 elution from both unworn and worn linezolid samples were > 100-fold higher than that of the 225 worn tigecycline samples and 24.8 and 25.9-fold higher respectively than the unworn 226 tigecycline samples.

#### 227 **Antimicrobial activity of eluted antibiotics**

228 *S. aureus* (SH1000), *S. epidermidis* (DSM 3269) and an *S. epidermidis* strain isolated from an 229 infected prosthesis were used as test organisms to investigate whether the eluted antibiotics 230 retained antimicrobial activity. The MICs of these strains with standard solutions of the 231 antibiotics are shown in Table S1 in the Supplementary material. Concentration of linezolid 232 and tigecycline eluted at various times from antibiotic-loaded cement samples were 233 determined via LC-MS and the MICs of the eluted antibiotics were determined 234 experimentally (Tables 1 and 2). All eluted tigecycline samples showed activity comparable 235 with the standard solution and established breakpoints<sup>21,22</sup> for all organisms tested (Table 1). 236 The linezolid samples eluted up to 72 h all showed activity comparable to determined MICs 237 and breakpoints against the Gram positive organisms.<sup>21</sup> The linezolid samples eluted over 1 238 week (168 h) showed higher MICs compared to the other samples and the Gram negative *E.*  239 *coli* was not inhibited by any of the linezolid samples, as expected (Table 2).

#### 240 **Cytotoxicity of antibiotics towards MG63 cells**

241 The cytotoxic effects of standard solutions of linezolid and tigecycline against MG63 cells 242 were determined using the MTT assay. The addition of increasing concentrations of 243 tigecycline resulted in a marked reduction in cell activity with an  $IC_{50}$  between  $5 - 10$  mg/L.

244 The addition of linezolid showed a small reduction in activity that was not statistically

245 significant (P > 0.05). Up to 300 mg/L of linezolid resulted in < 50% reduction in cell activity 246 and so an IC<sub>50</sub> for linezolid could not be determined (Supplementary material Fig S1). 247 Comparing these results to the concentrations achieved in the elution experiments (Figures 1- 248 4), it is possible that cellular toxicity of tigecycline may be an issue if the *in vivo* eluted 249 concentrations are comparable to those in this laboratory system, whereas linezolid did not 250 show toxicity to mammalian cells, even at substantially higher concentrations than those 251 achieved in the elution experiments.

#### 252 **Impact testing to assess physical strength of bone cements samples**

253 A Charpy type impact test machine was used to evaluate the impact strength of the antibiotic 254 loaded bone cement. Separate bone cement samples loaded either with tigecycline or 255 linezolid at 3 % and 10 % wt/wt were tested, and the results compared to both bone cement without antibiotic and a commercially prepared gentamicin-loaded bone cement, Refobacin<sup>®</sup> 256 257 Bone Cement R (Table 3). There was no significant difference in the impact strength of the 258 tigecycline-loaded cement samples at either concentration, compared to the control without 259 antibiotic. The 10% (wt/wt) tigecycline-loaded cement was the only cement that had an 260 impact strength that appeared slightly lower than the bone cement without antibiotic, 261 however that difference was not statistically significant. Further, there was no significant difference between the linezolid-loaded samples at either concentration and the Refobacin® 262 263 Bone cement R samples (P > 0.05). The impact strength of both the 3% and 10% (wt/wt) 264 tigecycline cement samples were significantly less (P < 0.05) than, though still comparable 265 to, the commercially available Refobacin<sup>®</sup> Bone Cement R.

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#### 268 **Discussion**

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270 The results presented here indicate that tigecycline and linezolid can be included within bone 271 cement and that the elevated temperatures that occur during the curing stage do not 272 compromise their antimicrobial and biocompatibility properties. Both antibiotics elute to 273 clinically relevant concentrations within the first hour in our laboratory elution system (Fig 1 274 and 2) and retain antimicrobial activity up to one week later. The concentrations of eluted 275 tigecycline peaked around 1 h (Fig 2) and then declined, presumably due to decomposition of 276 the antibiotic. The MICs for eluted tigecycline based upon the concentrations measured by 277 LC-MS showed results comparable with those determined using standard antibiotic solutions 278 (Table 1; Supplementary material Table S1). The MICs of eluted linezolid, the concentration 279 of which increased progressively throughout the experiment (Fig 1), were comparable with 280 those determined using standard antibiotic solutions over the first 72 h. After 1 week, eluted 281 linezolid showed approximately 5-20-fold higher MICs than the standard linezolid (Table 2; 282 Supplementary material Table S1) , which may indicate slow decomposition of the eluted 283 antibiotic that was not revealed by LC-MS. Previously, Anagnostakos *et al.* reported elution 284 of 1% of total linezolid from bone cement, compared to 3% for gentamicin loaded cement 285 over 8 days and Jackson *et al.* reported up to 3% elution over a 4 week period.<sup>14,15</sup> Cement 286 containing linezolid and gentamicin has shown inhibited growth of methicillin-resistant 287 S.*aureus* for up to 8 days.<sup>14</sup> However as this previous study is in conjunction with gentamicin 288 it does not necessarily confirm the activity of the linezolid on its own.

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290 The effect of wear on the tigecycline-loaded bone cement samples significantly reduces the 291 elution of tigecycline. After 1 hour there was > 4-fold reduction in the elution rate from the 292 worn sample, compared to the unworn control (Fig 3). Conversely, wear has very little effect 293 on the elution of linezolid from the bone with similar elution rates and profiles for both worn 294 and unworn samples (Fig 4). This may be relevant in the clinical application of these systems 295 where the cement surface experiences wear. Previously we have reported similarly 296 contrasting results with gentamicin and daptomycin–loaded bone cements where elution of 297 gentamicin was significantly reduced by wear, yet elution of daptomycin was not affected.<sup>16</sup> 298 In this study it was suggested that crystal size and distribution were the two main factors 299 influencing this difference in elution characteristics between the two antibiotics. It was 300 observed that the larger crystals of gentamicin within the orthopaedic cement created voids 301 on the surface upon contact with the aqueous solution, thus allowing greater deformation of 302 the bone cement surface due to wear. It was further proposed that this deformation prevented 303 the solution from penetrating deep into the bone cement, thereby limiting the amount of 304 antibiotic that can be eluted. In the current study we have shown that the crystals of 305 tigecycline are smaller than the linezolid crystals and so crystal size appears not to be the 306 main factor determining the reduced elution from worn bone cement samples here 307 (Supplementary material Fig S1). However there is a much greater tendency for the 308 tigecycline crystals to aggregate within the cement compared to the linezolid. The surface of 309 the tigecycline loaded cement showed areas of aggregated tigecycline crystals, which may 310 also produce voids upon contact with the aqueous solution and so increase the deformation of 311 the bone cement surface (Supplementary material Fig S2, S3).

312

313 The impact strength of the linezolid and tigecycline loaded cements produced results 314 comparable to those commercially available bone cements. The lowest impact strength was 315 seen in the 10% tigecycline containing cement suggesting that tigecycline may have some 316 effect on the mechanical strength of the cement. A previous study by Kries et al showed the 317 addition of tigecycline had a detrimental effect on compressive and bending strength of



322 The MTT assay showed that linezolid had low cytotoxicity towards MG63 cells. Up to 300 323 mg/L linezolid concentration resulted in  $\langle 50\%$  loss of cell activity and so an IC<sub>50</sub> was not 324 determined. Tigecycline showed greater cytotoxicity with an  $IC_{50}$  of 5 - 10 mg/L. This result 325 is consistent with the findings of Pina et al.,  $^{24}$  who also found that tigecycline concentrations 326 >10 mg/L severely affected the cell growth of osteoblastic cells.

327

#### 328 **Conclusions**

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330 The antimicrobial activity of linezolid and tigecycline eluted from within bone cement, 331 reaches therapeutically relevant concentrations within the critical perioperative period (based 332 on a typical arthroplasty operation of 1-2 h). Antimicrobial activity is observed up to 1 week 333 later. However, the concentration of tigecycline added to cement may need to be controlled 334 due to the possible cytotoxicity of the eluted antibiotic towards osteoblast cells. The effect of 335 wear in reducing elution of tigecycline in the laboratory reported here is also a factor to be 336 borne in mind if this antibiotic is used in revision surgery. Owing to ongoing antibiotic 337 resistance problems, there is a need to use antibiotics such as linezolid and tigecycline both 338 alone and in conjunction with other antibiotics (such as gentamicin which is included in 339 commercial bone cement preparations currently widely used in arthroplasty surgery). The 340 current study is an *in vitro* assessment of the performance and do not model the conditions *in*  341 *vivo*. Upon implantation the prosthetic comes into contact with extracellular fluid, bone and 342 muscle tissue, all of which will affect elution and the local accumulation of antibiotic. Further



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Fig 1: Concentration of linezolid eluted from bone cement over a 1-week period. Results are shown as the mean of three separate experiments ± standard deviation and have been normalised to 1 g bone cement in 5 mL of buffer.

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Fig 2: Concentration of tigecycline eluted from bone cement over a 1 week period. Results are shown as

the mean of three separate experiments ± standard deviation and have been normalised to 1 g bone cement

in 5 mL of buffer.



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Fig 3) Results from three separate experiments (A, B and C) comparing elution of tigecycline from worn and unworn tigecycline-loaded bone cement. Concentration of antibiotic was quantified by LCMS.

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433 434 435 Fig 4) Results from three separate experiments (A, B and C) comparing elution of linezolid from worn and unworn linezolid-loaded bone cement. Concentration of antibiotic was quantified by LCMS.



Table 1:MICs of tigecycline eluted from bone cement, determined by the broth microdilution method.

Experiments were carried out in triplicate.

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452 Table 2: MICs of linezolid eluted from bone cement, determined by the broth microdilution method.

453 Experiments were carried out in triplicate..



456 Table 3: Impact strength of antibiotic loaded bone cements determined using a Charpy-type testing apparatus.

457 Results are shown as a mean of five separate experiments  $\pm$  standard deviation. Biomet Bone Cement<sup>®</sup> was used

- 458 for all preparations unless stated otherwise.
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