# original article

# How Does a Photocatalytic Antimicrobial Coating Affect Environmental Bioburden in Hospitals?

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background. The healthcare environment is recognized as a source for healthcare-acquired infection. Because cleaning practices are often
 erratic and always intermittent, we hypothesize that continuously antimicrobial surfaces offer superior control of surface bioburden.

8 objective. To evaluate the impact of a photocatalytic antimicrobial coating at near-patient, high-touch sites in a hospital ward.

9 setting. The study took place in 2 acute-care wards in a large acute-carehospital.

10 methods. A titanium dioxide-based photocatalytic coating was sprayed onto 6 surfaces in a 4-bed bay in a ward and compared under normal

11 illumination against the same surfaces in an untreated ward: right and left bed rails, bed control, bedside locker, overbed table, and bed 12 footboard. Using standardized methods, the overall microbial burden and presence of an indicator pathogen (*Staphylococcus aureus*) were 13 assessed biweekly for 12 weeks.

results. Treated surfaces demonstrated significantly lower microbial burden than control sites, and the difference increased between treated and untreated surfaces during the study. Hygiene failures (>2.5 colony-forming units [CFU]/cm<sup>2</sup>) increased 2.6% per day for control surfaces (odds ratio [OR], 1.026; 95% confidence interval [CI], 1.009–1.043; P = .003) but declined 2.5% per day for treated surfaces (OR, 0.95; 95% CI, 0.925–0.977; P < .001). We detected no significant difference between coated and control surfaces regarding *S. aureus* contamination.

conclusion. Photocatalytic coatings reduced the bioburden of high-risk surfaces in the healthcare environment. Treated surfaces became
 steadily cleaner, while untreated surfaces accumulated bioburden. This evaluation encourages a larger-scale investigation to ascertain whether
 the observed environmental amelioration has an effect on healthcare-acquired infection.

Infect Control Hosp Epidemiol 2018;00(0):1-7

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Increasing microbial antibiotic resistance has given new impetus 24 to keeping hospitals clean.<sup>1</sup> Hospital-acquired infection (HAI) is 25 rightly seen as an unacceptable burden on the patient, as well as 26 inflating hospital costs.<sup>1</sup> While there is general agreement on the 27 need to control HAI, there is diversity of opinion regarding the 28 best solution. A major problem is the difficulty of conclusively 29 establishing a causal link between surface contamination and 30 HAI,<sup>2</sup> compounded by the lack of universally accepted stand-31 ards for measuring cleanliness.<sup>3</sup> Nevertheless, it is plausible to 32 assert that there is such a link,<sup>4</sup> allowing us to debate the most 33 cost-effective method for reducing contamination in 34 the healthcare environment. 35

Current decontamination strategies include daily detergentbased and disinfectant-based cleaning. Enhanced disinfection methods are available for rooms housing HAI patients and when an outbreak occurs.<sup>5</sup> Powerful disinfectants require caution because few have been properly evaluated under actual conditions of use, and they may ultimately be no better than traditional 41 detergent-based cleaning.<sup>6,7</sup> Manual cleaning has deficits, usually 42 attributed to personnel rather than product, and recontamination 43 inevitably begins immediately after the cleaning.<sup>8,9</sup> 44

Among recent technologies are photocatalytic antimicrobial 45 coatings.<sup>10</sup> They kill microbes by generating powerful oxidiz-46 ing radicals on a semiconductor surface following light 47 absorption in the presence of  $O_2$  and  $H_2O$ . The most impor-48 tant photocatalytic material is titanium dioxide (titania) 49 because the bandgap of the semiconductor overlaps suffi-50 ciently with the spectrum of natural and common artificial 51 light sources. The band edges are positioned appropriately for 52 generating the radicals, and the material is stable with respect 53 to self-destruction.<sup>10,11</sup> The illuminated semiconductor acts as 54 a source of reactive oxygen species (ROS), which are known 55 to be highly effective microbicides,<sup>12</sup> and the mechanism 56 of antimicrobial destruction is believed to involve bacterial 57

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Received July 7, 2017; accepted November 27, 2017

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58 cell-wall damage.<sup>13</sup> Those ROS generated by illuminated tita-

59 nia are particularly reactive and it is thought that resistance

<sup>60</sup> against them cannot develop.<sup>12</sup>

Although there have been in vitro investigations of photo-61 catalytic antimicrobial action with titania, very little work in 62 real-life situations has been reported.<sup>10</sup> A commercial titania 63 coating (Altimate EnviroCare Services, Singapore) did not signi-Q4 64 ficantly prevent environmental microbial contamination.<sup>14</sup> This 65 coating was, however, constituted from titania particles dispersed 66 in a binder to ensure their attachment to the coated surfaces; the 67 binder possibly encapsulated the particles and not only scavenged 68 the photogenerated radicals but also formed a physical barrier 69 70 between the particles and the microbes. Titania nanoparticles in suspension have been shown to be effective photocatalytic 71 antimicrobial agents, but they adhere very weakly to most 72 surfaces<sup>10,15</sup> from which they would, therefore, be continuously 73 lost. Petti and Messano<sup>16</sup> dispersed titania nanoparticles in 74 polyvinyl chloride (PVC) and observed antimicrobial action on 75 the surface of blocks made from the polymer, but this approach is 76 obviously unsuitable for retrofitting existing objects. 77

78 We resolved to evaluate a material (MVX, Hitech, Kitakyushu, Japan) that is applied as a dilute aqueous sol of titania 79 nanoparticles and dries to form a tough, adherent monolithic film 80 on the coated surface. Given evidence that photocatalytic anti-81 microbial activity can be synergistically enhanced by the presence 82 of copper or silver,<sup>11</sup> we chose to use a product doped with a 83 small proportion of silver zeolite. While it was tempting to coat all 84 surfaces in a ward due to ease of application (by spraying), we 85 focused on near-patient high-touch surfaces. They were coated 86 immediately after annual deep cleaning of the wards. Following 87 the application, the microbial burden and associated pathogens 88 were monitored over 3 months using standardized methods. 89 90

#### 91 Setting

The coated bay was in an acute-care general medical ward, and 92 93 an untreated control bay was selected in the stroke unit. The decision to spatially separate the treated and control bays, rather 94 than having them in the same ward, was taken to avoid intro-95 ducing a confounding factor in the form of a possible effect of 96 97 the coating on resident staff hands, who potentially have access to all patients on the same ward. Both wards are located in part 98 of the hospital that was constructed in 2004, and architecturally, 99 they are almost identical. The bays have a rectangular shape and 100 a volume of approximately 144 cubic meters. They are naturally 101 102 ventilated with windows along one of the long sides facing north; artificial light is provided during waking hours (dimmed 103 during the hours of sleep) from "daylight" fluorescent lamps. 104 At patient level the illuminance was  $\sim 400 \, \text{lux}$ . 105

## 106 methods

107 Choice of Surface Sites for Coating

108 The following surfaces were coated according to the manu-109 facturer's protocol: (1) left-hand side rails and (2) right-hand

110 side rails of a standard hospital bed; (3) the front face of the bed control panel; (4) the top of the bedside table; (5) the 111 bedside locker (coated in its entirety, but only the top was 112 sampled); and (6) the bed footboard (only the top was 113 sampled). There is consensus about the potential HAI risk 114 from these sites.<sup>17</sup> The furniture (table and locker) was made 115 from laminated wood. Each of these 6 sites was replicated for 116 117 all 4 bed spaces occupying a single bay of the selected ward.

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#### Ward Preparation

Prior to coating, the wards were deep cleaned, which comprises 119 thorough cleaning with a 5,000 ppm solution of Actichlor Plus 120 (a combination of a chlorine-compatible detergent with sodium 121 dichloroisocyanurate, NaDCC, also known as troclosene 122 sodium; Ecolab, Northwich, Cheshire, UK) followed by steam 123 cleaning and, as a final step, enhanced cleaning with hydrogen 124 peroxide vapor (HPV, Deprox, Specialist Hygiene Solutions, 125 Kings Lynn, UK). The stroke ward was deep-cleaned in the week 126 commencing August 1, 2016, and the acute medical ward 127 was deep-cleaned in the week commencing September 10, 2016. 128 No patients were admitted to the ward between deep cleaning 129 and coating. 130

### Coating Procedure

The coating is a dual one, comprising a colorless primer (ie, the 132 primary coating) over which the photocatalytic titania coating 133 MVX is laid. Final coating thickness was approximately 1 µm. 134 The precursors of mix are dilute aqueous solutions of the active 135 ingredients, titania (1.5%) and silver zeolite (0.1%).<sup>18</sup> These 136 solutions, as well as the final coating, are nontoxic to humans.<sup>21</sup> 137 Primary coating (MVX, Hitech) was sprayed onto the selected 138 surfaces and allowed to dry for 20-30 minutes; the ambient 139 temperature in the ward during coating was  $26 \pm 1^{\circ}$ C and the 140 relative humidity was 59  $\pm$  3%. The MVX was then applied 141 likewise by spraying and similarly allowed to dry. After drying, 142 the coating was invisible to the eye, even on mirrors (which are 143 integral on some lockers). All coated objects were discretely 144 fitted with trackers for the TeleTracking Technologies real-time 145 location system (RTLS; Pittsburgh, PA) installed at the hospital 146 as part of the "Safe Hands" program, to ensure that the coated 147 objects could always be unambiguously located, even if clinical 148 exigence (eg, to reduce the risk of falls, or simply to make the 149 patient more visible) led to a patient (with bed and bed-space 150 equipment and furniture) being moved, generally within 151 the ward. 152

#### Sampling Protocol

The approach followed that described by Bogusz et al<sup>19</sup> 154 Starting at 7:00 AM on Tuesdays and Thursdays, for 12 weeks 155 from September 22 to December 21, 2016, after locating the 05 objects with the RTLS, the coated sites and their uncoated 157 equivalents were sampled using double-sided dipslides 158 (Hygiena International, Watford, UK) coated with nutrient and Baird Parker agars, pressing the slides at  $25 \text{ g/cm}^2$ for 5 seconds.<sup>20</sup> Within the sites, the actual locations were determined at random,<sup>21</sup> according to the judgment of the (sole) sampler.

164 Microbiology

Dipslides were incubated for 48-72 hours at  $36 \pm 1^{\circ}$ C according 165 to laboratory protocol, after which the number of aerobic 166 167 colony-forming units (CFU) was determined from the nutrient agar side. Baird Parker agar highlighted potential coagulase-168 positive staphylococci, which were subcultured onto blood agar 169 and identified as methicillin-susceptible or -resistant according 170 to laboratory protocol. The aerobic colony count (ACC) was 171 quantified using a 5-point scale (Table 1).<sup>3,7,19</sup> Staphylococci 172 were classified as either "isolated" or "not isolated." 173

#### 174 Ward Environment

Every day, the ward cleaning team cleaned all items in the patient 175 bed space with Hospec general surface cleaner (containing 176 alcohol ethoxylate as the detergent) (Robert McBride, Middle-177 ton, Manchester, UK), typically during the morning after 178 sampling. No exceptional cleaning (HPV or Actichlor Plus) was 179 requested for the control ward during the study. Actichlor Plus 180 was requested on 3 occasions in the treated ward, but for side 181 rooms away from the treated bay. Unlike the strongly bacteri-182 cidal ionic surfactants, nonionic surfactants are generally consi-183 dered less bactericidal, although they interfere with bacterial 184 membrane fluidity.<sup>22</sup> It is difficult to separate the physical 185 bactericidal effect of the mechanical wiping action from the 186 biochemical bactericidal effect associated with the surfactant,<sup>23</sup> 187 but some attempts at quantification have been made.<sup>7,19</sup> 188

Bed occupancy was high in both treated and control wards, 189 averaging 97.6% for the former and 88.0% for the latter during 190 the study (data for the entire ward). Locally agreed staffing levels 191 are recorded for all wards at the hospital. The stroke ward was 192 generally better staffed than the acute-care ward. Medical staff, 193 allied health professionals (AHP, including physiotherapists, 194 occupational therapists, and speech and language therapists) 195 and domestics were not included, nor were visitor numbers 196 monitored. The degree of dependency (acuity) of the patients 197 occupying the beds was also examined. The median degree was 198 invariably level 1b using the Hurstclassification.<sup>24</sup> 199

The hospital's research and development department determined not to class the study as research but rather as a service evaluation. Therefore, approval from the research ethics committee was not required.

## 204 Statistical Methods

The sampling protocol resulted in a maximum of 102
 bed-space observations for each ward subsequently available for statistical analysis. Each observation produced 6

table 1. Classification of Aerobic Colony Counts (ACCs)

CFU/cm <sup>2</sup>	Name	Numerical Descriptor	Binary score <sup>a</sup>
0	No growth	1	Pass = 1
< 2.5	Very slight growth	2	Pass = 1
2.5-12	Light growth	3	Fail=0
12-40	Moderate growth	4	Fail=0
> 40	Heavy growth	5	Fail=0

NOTE. CFU, colony-forming units.

<sup>a</sup>According to Dancer (2008).<sup>26</sup>

measurements of ACCs, which were allocated a numerical 207 descriptor from 1 to 5 (Table 1). For the statistical analysis, a 208 mean "numerical descriptor" score (ie, arithmetic mean of the 209 6 test sites) was calculated for each bed space. This score was 210 dichotomized into a pass/fail outcome variable (1-2 = "pass")211 and >2-5 = "fail"). Although dichotomizing may lead to a 212 loss of statistical power,<sup>25</sup> it is in concordance with the 213 previously introduced pass-fail dichotomy for bioburden.<sup>3,26</sup> 214 Furthermore, the conventional classification (Table 1) gives a 215 highly nonlinear mapping of ACCs onto a descriptor; by 216 dichotomizing we avoid having to discuss whether to express 217 the results in terms of CFU/cm<sup>2</sup> or in terms of the "degree of 218 219 growth" descriptor.

The difference in pass-fail rates between the 2 wards 220 (experimental and control) was assessed using the  $\chi^2$  independ-221 ence test. Straightforward binary logistic regression analysis was 222 223 used to further explore the probability (odds) of failing the pass-fail test on the 2 wards.<sup>27</sup> Additional factors (introduced as 224 continuous covariates) included the number of days into the 225 226 study (0-90) and the bed occupancy rate (%) for each ward. The 227 multiple regression logit model was fitted using the binary 228 logistic regression analysis option in SPSS software (SPSS, 229 Chicago, IL). The analysis allowed both fixed and categorical factors and continuous covariates to be used as explanatory 230 variables when estimating the probability (or, more correctly, 231 the odds) of failing the test. P < .05 was used as a measure 232 of significance. 233

## results

The overall pass rate for the coated bay was 80.4% (82 passes of 235 102 total samples), while for the control bay it was 52.9% 236 (54 passes of 102 samples). The results of the binary logistic 237 regression analysis, using the control ward as the reference 238 condition, are given in Table 2. The analysis identified no 239 difference in the odds of failing the test between the 2 wards at 240 the beginning of the experiment (odds ratio [OR], 0.993; 95% 241 confidence interval [CI], 0.267-3.69; P = 0.993). However, 242 the odds of failing the test in the control bay increased by 243 2.6% per day (B = 0.026; OR, 1.026; 95% CI, 1.009–1.043; 244 P = .003) but declined by 2.5% per day in the treated bay 245 (B = 0.026 - 0.051; OR, 0.95; 95% CI, 0.925 - 0.977; P < .001).246 These trends are plotted in Figure 1. 247

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For the individual sites, we considered the sampling as a sequence of independent Bernoulli trials with the binary outcome of "pass" or "fail" and an initially unknown probability p of passing, which was found from the maximum of the likelihood of p, given the observed sequence.<sup>28</sup> The results are given in Table 3. Surface treatment with MVX significantly improved microbial cleanliness at every site,

table 2. Factors (Variables) Found to Influence the Probability p of Failing the Test, Estimated Using Binary Logistic Regression, Adopting (Fail vs Pass) as the Dichotomous Response Variable<sup>a</sup>

	B (SE) <sup>b</sup>	P Value <sup>c</sup>	$OR^d$	95% CI
Control ward	0.000		1.00	
Treated ward	-0.007(0.670)	.991	0.993	0.267-3.690
Days into the evaluation (for the control ward)	0.026 (0.009)	.003	1.026	1.009-1.043
Treated ward by days Bed occupancy, % Constant	-0.051 (0.014) 0.076 (0.034) -7.866(3.099)	.000 .026 .011	0.950 1.079 0.000	0.925–0.977 1.009–1.154

<sup>a</sup>Estimated parameters  $B_i$  for the logit model: Log[p/(1-p)] – Constant

+  $B_i$ , where the subscript i = 1 refers to the untreated sites and i = 2 to the treated ones. The control ward was estimated as the baseline constant parameter (at day 0), and the treated ward effect was estimated as a deviation from this constant parameter. The number of days from day 0 and bed occupancy were introduced as continuous covariates.

<sup>b</sup>Slope parameter of the continuous covariate (days), with its standard error in parentheses. <sup>c</sup>Measure of significance.

<sup>d</sup>Odds ratio, equal to  $\exp(B)$ .

<sup>256</sup> <sup>e</sup>Confidence intervals for exp(B).

although only borderline significance was achieved for 257 the bed footboard. The left-hand and right-hand bed rails 258 were conceived as internal controls for each other but 259 yielded different probabilities of passing; there may have 260 been physical differences in accessing the bed rails, such 261 as one bed rail being closer to a wall or some other 262 obstruction. 263

Staphylococcus aureuswas isolated from only ~10% of the264dipslides: 97 isolates were recovered from a total of 635 for the265treated surfaces (all sites together), compared with 68 isolates266from a total of 655 for the control surfaces. The low S. aureus267counts rendered the difference insignificant.268

# discussion

The gradual diminution of bioburden on the treated surfaces270occurred even though bed occupancy was higher than in271untreated bay, which would have likely encouraged heavier272microbial contamination on ward surfaces.26This result273abrasion from touching or cleaning, initially considered as a275possibility, did not occur.276

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Among the possible confounding factors considered 277 (ie, Hawthorne effect; bed occupancy; staffing levels; and 278 degree of patient dependency) only bed occupancy differed 279 markedly between the treated and control bays. Although the 280 patients differed between the 2 study bays, we found no 281 evidence for a clinically significant difference with respect to 282 the likelihood of individual patients and attendant staff 283 contributing to the microbial burden in their environment. 284



figure 1. Actual data (open circles) and predicted values (open triangles) for the control sites and treated sites (data: closed blue-grey circles; predicted values: closed triangles) for the duration of the evaluation. The vertical axis is microbial growth according to the 5-point scale (Table 1).

table 3. Success Probabilities p for the (lack of) Aerobic Growth at the Various Sites

Site	р		No. of Observations		s <sup>a</sup>			
	Treated	Control	Treated	Control	Treated	Control	$ p_{\text{treated}} - p_{\text{control}}  / (s_{\text{treated}} + s_{\text{control}})^b$	
Left-side bed rail	.66	.51	98	102	0.05	0.05	1.5	
Right-side bed rail	.82	.44	98	102	0.04	0.05	4.2	
Control panel	.80	.73	99	97	0.04	0.05	0.8	
Bedside table	.86	.75	99	95	0.03	0.04	1.6	
Bedside locker	.95	.79	87	102	0.02	0.04	2.7	
Bed footboard	.51	.48	87	91	0.05	0.05	0.3	
All sites	.77	.61	568	577	0.018	0.020	4.2	

<sup>a</sup>The span s is the square root of the observed formation, which is a measure of the uncertainty of  $p^{28}$ .

<sup>b</sup>The difference between the probabilities divided by the sum of the spans is an index of the significance of the result: the greater the index, the greater the significance.

table 4. Environmental Audits for Housekeeping Compliance With Cleaning<sup>a</sup>

	Monthly "Healt Environmenta Scores,	h Assure" al Audit %	Monthly "Credits for Cleaning" (C4C) Environmental Audit Scores, %		
Month	Treated Ward	Control	Treated Ward	Control	
September	98.2	93.6	99.5 <sup>b</sup>	98.1 <sup>b</sup>	
October	99.1	84.0	98.4°	99.4°	
November	98.2	87.0	99.0 <sup>d</sup>	97.7 <sup>d</sup>	
December	90.0	84.6	98.8	99.6	

<sup>a</sup>The audits do not directly observe the staff actually cleaning but inspect the whole ward environment, including high-touch surfaces. <sup>b</sup>Week commencing September 19.

<sup>c</sup>Week commencing October 24.

<sup>d</sup>Week commencing November 28.

<sup>e</sup>Week commencing January 9.

Environmental audits undertaken to appraise housekeeping compliance with cleaning are reported in Table 4 for the interval of the study. They show little difference between the 288 2 wards.

289 It is interesting to compare the bioburden reduction provided by the photocatalytic coating with conventional 290 detergent or disinfectant application to high-touch 291 surfaces 292 (UK hospitals, generally use detergents, and hospitals in the 293 United States generally use disinfectants). Microbial counts from a wide range of hand-touch sites cleaned with detergent 294 ranged from 2.5 to 40 CFU/cm<sup>2</sup>;<sup>29</sup> detergent cleaning was 295 shown to reduce bioburden from a preclean level of 6.7 to 296 3.5 CFU/cm<sup>2,19</sup> On the other hand, disinfectant reduced 297 median counts for high-touch sites to 0.1–0.6 CFU/cm<sup>2,30</sup> 298 A major difficulty is that sampling methods, surfaces, sites 299 300 (ie, near-patient hand-touch sites host different amounts and 301 types of bioburden than floors or bathroom sites), cleaning 302 agent exposure, and culture techniques are not standardized 303 across studies. Another confounding factor is sampling 304 methodology: greater quantities of bioburden are recovered

from moistened swabs placed in broth then agar methods such as RODAC plates or dipslides.

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Our results suggest that the chosen wards were already rather 307 clean, especially with respect to S. aureus; the effect of the 308 photocatalytic coating in lowering bioburden might be more 309 prominent in a less stringently clean hospital. Conversely, a 310 311 recent study of the effect of MVX in the critical-care environment, which is always afforded priority for cleaning (eg, is 312 routinely cleaned with alcohol thrice daily), found no significant 313 microbiological benefit, despite in vitro data from the same 314 coating showing pathogen inactivation.<sup>31</sup> The duration of the 315 study was only 4 weeks, however, which may be inadequate 316 to provide sufficient statistical power to show any significant 317 difference between treatment and control. 318

Although a photocatalytic surface continuously maintains 319 its antimicrobial action, the action is slow. Kinetic laboratory 320 studies, in which surfaces were deliberately contaminated with 321 known amounts of bacteria, suggest that ~1 hour is needed 322 to destroy half the bacteria.<sup>32,33</sup> Hence, if a site had been 323 adventitiously heavily contaminated a few minutes prior to 324 sampling, the result would indicate a high bioburden, whereas 325 sampling 2 hours later might indicate low contamination. 326

327 The ultimate objective for hospitals regarding cleanliness is to reduce the incidence of HAI. At present, the relationship 328 between microbial burden on hospital surfaces and the inci-329 dence of HAI remains unclear. No extant model allows the 330 prediction of the change in HAI incidence as a result of lowering 331 the environmental bioburden by a defined amount, and thus 332 far, no empirical study appears to have tackled this deficit. A few 333 studies have examined the link between standardized measure-334 ments of bioburden and HAI rates but with inconclusive 335 outcomes.<sup>2</sup> Much attention has been given to the proposition 336 that hands are the main vectors for transmission and, therefore, 337 that frequent hand hygiene is the key to reducing HAI, although 338 the limitations of this approach were noted decades ago.<sup>34</sup> 339 Furthermore, although hand hygiene is strongly promoted in 340 the healthcare setting, compliance is still far from ideal but 341 may, nevertheless, have already reached a practical limit.<sup>35</sup> 342 343 In any case, hand contamination is most likely to be transmitted

via the intermediary of high-touch surfaces, such as thoseinvestigated in the present study, rather than directly toanother hand.

"Routine cleaning and disinfection is apparently not suffi-347 cient."<sup>36</sup> Detailed investigation of routine processes may reveal 348 349 weaknesses, in addition to those already discussed, alongside their irreducible intermittency.<sup>9,37</sup> In contrast, a photocatalytic 350 surface is continuously active. Some of the physicochemical 351 changes induced in titania by light persist for many hours or 352 days in the dark, reinforcing this continuity.<sup>38</sup> A photocatalytic 353 coating of the type evaluated here offers a new perspective 354 for overcoming some of the present limitations in cleaning, 355 356 disinfection, and hand hygiene. A further advantage is that the mechanism whereby photocatalytic antimicrobial coatings 357 inactivate microbes is unlikely to lead to the development of 358 resistance,<sup>12</sup> the increase of which is of grave concern to public 359 health authorities. 360

In conclusion, coating high-touch surfaces with a titania-361 based photocatalytic material significantly lowered bioburden 362 compared with a control bay. The trend of continuously 363 364 diminishing bioburden in the treated bay is encouraging, not least in comparison with the untreated control bay, in which the 365 bioburden appeared to continuously increase. A much larger 366 and longer study should now be undertaken with sufficient 367 power to observe whether coating high-touch surfaces with an 368 369 antimicrobial coating reduces the incidence of HAI. Although there is no evidence that nontouch surfaces (walls, ceilings, etc) 370 are reservoirs for microbes, empirically verifying or otherwise 371 the proposition that coating all surfaces with a photocatalytic 372 material reduces the incidence of HAI will be a further useful 373 374 addition to infection prevention efforts.

- 375 acknowledgments
- 376 We thank Lee Turner for performing the dipslide sampling throughout the study.
- 377 We thank Sue Lovegrove of the Microbiology Laboratory, New Cross Hospital for
- 378 processing the dipslides. We thank Dr Khaled Hussein, MVX Hi-tech Company
- 379 (Kitakyushu, Japan), for valuable discussions about the photocatalytic coating
- 380 and its properties, and MVX Hi-tech for providing the photocatalytic materials

and the personnel to spray them onto the chosen surfaces.

- Financial support: The Collegium Basilea (Institute of Advanced Study)
   partly funded this work.
- Potential conflicts of interest: All authors report no conflicts of interest
   relevant to this article.
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