

1 Genomic identification of cryptic susceptibility to penicillins and β-lactamase

- 2 inhibitors in methicillin-resistant *Staphylococcus aureus*
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35

# 36 Abstract

37 Antibiotic resistance in bacterial pathogens threatens the future of modern medicine. 38 One such resistant pathogen is methicillin-resistant Staphylococcus aureus (MRSA), 39 which is resistant to nearly all  $\beta$ -lactam antibiotics, limiting treatment options. Here, 40 we show that a significant proportion of MRSA isolates from different lineages, 41 including the epidemic USA300 lineage, are susceptible to penicillins when used in 42 combination with  $\beta$ -lactamase inhibitors such as clavulanic acid. Susceptibility is 43 mediated by a combination of two different mutations in the mecA promoter region 44 that lowers mecA-encoded penicillin binding protein 2a (PBP2a) expression, and in 45 the majority of isolates by either one of two substitutions in PBP2a (E246G or M122I) 46 that increase the affinity of PBP2a for penicillin in the presence of clavulanic acid. 47 Treatment of *S. aureus* infections in wax moth and mouse models demonstrate that 48 penicillin/ $\beta$ -lactamase inhibitor susceptibility can be exploited as an effective 49 therapeutic choice for 'susceptible' MRSA infection. Finally, we show that isolates 50 with the PBP2a E246G substitution have a growth advantage in the presence of 51 penicillin, but the absence of clavulanic acid, which suggests that penicillin/β-52 lactamase susceptibility is an example of collateral sensitivity (resistance to one 53 antibiotic increases sensitivity to another). Our findings suggest that widely available 54 and currently disregarded antibiotics could be effective in a significant proportion of 55 MRSA infections. 56 57 58

59

# 61 Introduction

62	The $\beta$ -lactam family of antibiotics, is one of the most widely used and clinically
63	important groups of antibiotics <sup>1</sup> . Resistance to $\beta$ -lactam antibiotics in
64	Staphylococcus aureus is either mediated by the acquisition of the blaZ gene
65	encoding a $\beta$ -lactamase, or in methicillin-resistant S. aureus (MRSA) from the
66	acquisition of an alternative penicillin binding protein 2a (PBP2a) with a low affinity
67	for $\beta$ -lactam antibiotics, resulting in resistance to most $\beta$ -lactams <sup>2</sup> . PBP2a is
68	encoded by mecA or mecC (PBP2a'/c) carried on a mobile genetic element known as
69	a staphylococcal cassette chromosome mec (SCCmec) <sup>3,4</sup> , and is regulated by two
70	independent regulatory systems (mecl-mecR-mecR2 and blal-blaR) and multiple
71	chromosomal genes <sup>5</sup> .
72	
73	Countering resistance to $\beta$ -lactams was first achieved by the development of $\beta$ -
74	lactamase resistant $\beta$ -lactams such as methicillin $^6$ , and subsequently by the
75	development of $\beta$ -lactamase inhibitors <sup>7</sup> . Unfortunately, MRSA is resistant to both
76	methicillin antibiotics and insensitive to $\beta$ -lactamase inhibitors. However, in the early
77	1990s, $\beta$ -lactams and $\beta$ -lactamase inhibitors were tested against MRSA $^{\text{8-11}},$ with
78	success both <i>in vivo</i> and clinically <sup>12</sup> .
79	
80	Recently, different approaches to restore susceptibility to $\beta$ -lactams have been
81	investigated, including using multiple antibiotics to exploit collateral sensitivity
82	(resistance to one antibiotic increases sensitivity to another) <sup>13-15</sup> . We recently
83	demonstrated that the small subset of MRSA isolates with the mecC gene are
84	susceptible to penicillins and $\beta$ -lactamase inhibitors, as the $\textit{mecC}$ encoded PBP2c
85	does not mediate resistance to penicillins <sup>16</sup> . Here, we demonstrate that
86	unexpectedly, a significant proportion of all clinical mecA-positive MRSA isolates
87	from diverse lineages are susceptible to penicillins and $\beta$ -lactamase inhibitors. These

88 findings could provide the basis of new treatment options for susceptible MRSA

89 infections using already licenced antibiotics.

90

### 91 **Results**

#### 92 Penicillin/β-lactamase inhibitor-susceptible MRSA

As part of an earlier study <sup>16</sup>, we identified an *mecA*-positive MRSA isolate that 93 94 exhibited increased susceptibility to penicillin in the presence of clavulanic acid (a β-95 lactamase inhibitor) (Fig 1a). We screened a selection of whole genome sequenced 96 (WGS) MRSA isolates (n = 110) from different clinically relevant clonal lineages for 97 the same increase in susceptibility (≥10 mm increase in the zone of inhibition in the 98 presence of 15  $\mu$ g/ml clavulanic acid, compared to penicillin alone). Fifty-five (50.0%) 99 isolates from different lineages were susceptible to penicillin-clavulanic acid 100 (Supplementary Table 1). We determined the minimum inhibitory concentration (MIC) 101 for penicillin for a subset of isolates, which showed that the MIC was reduced below 102 the European Committee on Antimicrobial Susceptibility Testing (EUCAST) / Clinical and Laboratory Standards Institute (CLSI) breakpoint ( $\leq 0.125 \,\mu$ g/ml)<sup>17,18</sup> in ten of 103 104 the fourteen susceptible isolates, and none of the ten resistant isolates (Fig. 1b). Two 105 other  $\beta$ -lactam inhibitors, subactam and tazobactam also increased susceptibility to 106 penicillin (Supplementary Fig. 1), suggesting that the effect of clavulanic acid was 107 due to inhibition of the staphylococcal  $\beta$ -lactamase. Next, we evaluated if the 108 increase in susceptibility was penicillin-specific by screening isolates against different 109  $\beta$ -lactam antibiotics in the presence of clavulanic acid (Fig. 1c and d). This revealed 110 that the penicillin-clavulanic acid susceptible isolates showed the greatest increase in 111 susceptibility to penicillins (benzyl- and aminopenicillins), (Fig. 1d) and were broadly 112 more susceptible to cephalosporins (except ceftaroline) and carbapenems than the 113 resistant isolates (Fig 1c). In contrast, the presence of clavulanic acid produced only 114 minor increases in susceptibility in the resistant isolates (Fig 1c).

### 115 Substitutions in PBP2a mediate increased penicillin susceptibility

116 Benzyl- and aminopenicillins bind reasonably well to PBP2a in comparison to the 117 isoxazolyl penicillins (oxacillin) and cephalosporins <sup>19</sup>. As amino acid substitutions in 118 PBP2a mediate resistance to fifth-generation cephalosporins  $^{20}$ , in the absence  $\beta$ -119 lactamase activity, we hypothesised that inversely, increased susceptibility to 120 penicillins might be mediated by PBP2a substitutions. We examined the PBP2a 121 amino acid sequences of the 110 isolates (Supplementary Table 1). This revealed 122 that 80.0% (44/55) of the penicillin-clavulanic acid susceptible isolates had either a E246G (PBP2a<sup>246G</sup>) (n=37) or a M122I (PBP2a<sup>122I</sup>) (n=7) substitution in the allosteric 123 domain of PBP2a<sup>21</sup>. By contrast, only 29.1% (16/55) of the resistant isolates had the 124 PBP2a<sup>246G</sup> substitution. Phylogenetic analysis of PBP2a indicated that 246E is 125 126 present in the S. aureus COL genome (M122I was absent), an early MRSA strain isolated in the 1960s, with origins in the 1940s<sup>22,23</sup>, suggesting this might be the 127 128 ancestral or 'wildtype' form (Supplementary Fig. 2).

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130 Next, we tested the effect of the PBP2a substitutions experimentally. We deleted mecA (PBP2a) in isolates from two S. aureus lineages: ST398 (EC139<sup>24</sup>) and 131 132 USA300/ST8 (BCVA289<sup>25</sup>) (both PBP2a<sup>246G</sup> and *blaZ* positive) and introduced 133 plasmid-borne copies of three mecA alleles: the putatively resistant 'wildtype' 134  $mecA_{246E}$  and the two alleles associated with susceptibility:  $mecA_{122I}$  and  $mecA_{246G}$ . 135 All the complemented strains were resistant to penicillin alone (MICs >20µg/ml -136 breakpoint >0.125 $\mu$ g/ml) (Fig. 2a). For penicillin-clavulanic acid, the mecA<sub>246E</sub> strains 137 were resistant with MICs of  $1\mu g/ml$  (Fig. 2a). While the strains with mecA<sub>246G</sub> were 138 susceptible, with MICs of 0.125µg/ml. Similarly strains with mecA1221 had MICs of 139 0.125 and 0.25µg/ml for USA300 and ST398 backgrounds, respectively. No 140 difference in susceptibility was seen for cefoxitin (a cephalosporin), confirming that

141 the effect of the substitutions was limited to penicillin susceptibility (Supplementary

142 Table 2).

143

# 144 A combination of *mecA* promoter mutations and PBP2a substitutions mediate

#### 145 susceptibility

146 We reasoned that the PBP2a substitutions might cause an increase in the affinity of PBP2a for penicillin. We tested PBP2a<sup>246E</sup>, PBP2a<sup>246G</sup> and PBP2a<sup>122I</sup> in a bocillin 147 148 competition assay to determine their relative binding affinities for penicillin (Fig 2b). 149 This identified that all three variants had a similar affinity for penicillin, with 50% 150 inhibitory concentrations (IC50) of 10.63  $\mu$ g/ml (standard error of the mean ± 0.53) for PBP2a<sup>246G</sup> and 12.11(± 1.07) for PBP2a<sup>122I</sup> and 9.06 (± 2.70) for PBP2a<sup>246E</sup> (Fig 2b). 151 152 As susceptibility testing had been carried out in the presence of clavulanic acid, we 153 repeated the bocillin binding assays with 15 µg/ml clavulanic acid. In the presence of 154 clavulanic acid, the wildtype PBP2a<sup>246E</sup> had a two-fold higher IC50 of 32.07 (± 8.38) compared to that PBP2a<sup>246G</sup> 16.22  $\pm$  3.19) and PBP2a<sup>122I</sup> (11.73  $\pm$  4.50), which were 155 156 virtually unaffected (Fig 2c).

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158 To confirm the consistency of association between the two PBP2a substitutions and 159 phenotypic susceptibility, we determined the penicillin and penicillin-clavulanic acid 160 MICs for 274 WGS isolates from a range of *S. aureus* lineages (Supplementary 161 Table 3). We then combined the data with the original 24 isolates with MIC data (Fig. 162 1b) and plotted the frequency distributions of the penicillin and penicillin-clavulanic 163 acid MICs (Fig. 2d and e). In the absence of a clinical breakpoint for penicillin-164 clavulanic acid we determined a tentative epidemiological (wildtype) cut-off (ECOFF) using a statistical method based on mixture models <sup>26,27</sup>. This supported setting the 165 166 ECOFF between 2.0 and 3.0 µg/ml (2.449, rounded down to 2.0 µg/ml) for penicillin 167 in the presence of 15 µg/ml clavulanic acid (Fig 2d). The same method supported an 168 ECOFF between 0.19 and 0.25 µg/ml (0.21, rounded down to 0.19 µg/ml) for

169penicillin alone, which is within one doubling dilution of the EUCAST clinical170breakpoint (Fig 2e)<sup>18</sup>. Using the ECOFF of  $\leq 2.0 \,\mu$ g/ml as the cut-off, 213 (71.5%)171isolates were classified as susceptible to penicillin-clavulanic acid, of which only 176172(82.6%) had either a 246G or 122l substitution. This indicated that 37 (17.4%)173isolates were susceptible despite having no PBP2a substitution. Conversely, only 37174(43.5%) of the resistant isolates (MIC >2  $\mu$ g/ml) had a 246G substitution.

175

176 Incomplete congruence between PBP2a substitutions and penicillin-clavulanic acid 177 susceptibility led us to search for further mutations involved in susceptibility. We 178 hypothesised that higher levels of PBP2a expression might overcome the effect of 179 the PBP2a substitutions. We screened the same 298 isolates for mutations in the 180 mecA promoter region, this identified two mutations that correlated with susceptibility. 181 The first mutation was a G to T transversion in the *mecA* ribosomal binding site 182 (RBS), seven nucleotides upstream of mecA start codon (Fig 3a). Isolates with the 183 mecA[-7]:T allele had a median penicillin-clavulanic acid MIC of 0.125 µg/ml (range 184 <0.016 - 6) compared to a median of 8  $\mu$ g/ml (range: 0.023 - 96) for isolates with the 185 'wildtype' mecA[-7]:G. Previous work has demonstrated that despite being in the 186 RBS, the T allele results in lower mecA transcript and PBP2a expression levels <sup>28</sup>. 187 We compared relative levels of mecA expression by reverse transcription quantitative 188 polymerase chain reaction (RT-gPCR) in isolates from phylogenetically separate 189 lineages with the mecA[-7]:G (n=7) allele to isolates with the mecA[-7]:T allele (n=6) 190 (Fig 3b). Isolates with the T allele had a statistically significantly lower relative 191 expression (mean relative expression: 3.24) than isolates with the G allele (mean: 192 7.00) (P=0.0048) (Fig 3b). The second mutation was a C to T transition in the mecA -193 10 box, 33 nucleotides upstream of the mecA start codon (mecA[-33]:T) (Fig 3a). 194 Isolates with this mutation had a median penicillin-clavulanic acid MIC of 0.047µg/ml 195 (range: <0.016 to 0.125). The C to T transition causes the generation of a perfect

palindrome within the Mecl-Blal binding site, which lowers *mecA* transcript and
PBP2a expression levels <sup>28,29</sup>. Three isolates from distinct lineages with the *mecA*[33]:T mutation had a mean relative expression rate of 0.27, suggesting the C to T
transition results in very low levels of *mecA* expression in the tested isolates (Fig 3b).

201 When considering the two promoter mutations (mecA[-7]:G-T and mecA[-33]:C-T) together with the two PBP2a substitutions (PBP2a<sup>246G</sup> and PBP2a<sup>122I</sup>) (Fig 3a) we 202 203 identified six genotypes. These were used to annotate the MIC distributions for 204 penicillin and penicillin-clavulanic acid (Fig 3c and d). For penicillin-clavulanic acid, 205 the genotypes split clearly into the bimodal distribution (Fig 3c). The majority of 206 isolates with mecA[-7]:G (henceforth: Resistant 1) and mecA[-7]:G | E246G 207 (Resistant 2) genotypes were found in the modal peak to the right with MICs above 208 the ECOFF of  $\geq 2 \mu g/ml$  (Fig 3c). The majority of isolates with the other four 209 genotypes: mecA[-7]:G-T (henceforth: Susceptible 1), mecA[-7]:G-T | E246G 210 (Susceptible 2), mecA[-33]:C-T | mecA[-7]:G | E246G (Susceptible 3) and mecA[-211 7]:G | M122I (Susceptible 4) were located in the modal peak to the left with MICs 212 below the ECOFF (≤2.0 µg.ml). Isolates with susceptible genotypes had lower 213 penicillin MICs in the absence of clavulanic acid than those with resistant genotypes 214 (Fig 3c and d). The use of the six genotypes to predict susceptibility using the 215 ECOFF as breakpoint was accurate in 94.6% (282/298) of isolates, with a 0.34% 216 (1/298) very major error (VME) rate (defined as isolates that were phenotypically 217 resistant but genotypically predicted to be susceptible) and a 5.0% (15/298) major 218 error (ME) rate (phenotypically susceptible, genotypically predicted resistant). 219 220 Finally, we investigated if the presence of the six different types of class A 221 staphylococcal ß-lactamases (types A-F), or *blaZ* expression levels might affect penicillin-clavulanic acid susceptibility<sup>16</sup>. There was no association between the ß-222

223 lactamase type and susceptibility that wasn't better explained by the six genotypes

224 (Supplementary Table 3 and Fig. 3a). Nor was there any significant association (P=

0.43) between *blaZ* expression and susceptibility in twenty isolates (11 susceptible, 9

- resistant) as measured by RT-qPCR (Supplementary Fig. 3b).
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#### 228 Clinical prevalence and epidemiology

229 We sought to quantify the frequency of penicillin-clavulanic acid susceptibility in 230 clinical MRSA isolates. We tested a collection of 270 S. aureus isolates (MRSA, n = 231 218; MSSA isolates, n = 52) collected by the Danish Staphylococcus Reference Laboratory as part of bacteraemia surveillance <sup>30</sup> (Supplementary Table 4). Isolates 232 233 were classified as resistant or susceptible based on the ECOFF of  $\leq 2.0 \, \mu g/ml$ 234 penicillin in the presence of 15  $\mu$ g/ml of clavulanic acid. All MSSA isolates, and 235 84.9% (185/218) of the MRSA isolates had an MIC below the ECOFF. The MRSA 236 isolates were from a variety of lineages as inferred from spa-typing including clonal 237 complex (CC)1, CC5, CC8, CC30 and CC80. We then screened 2282 WGS MRSA isolates from Cambridge, UK for the six genotypes (Fig. 4a) <sup>31</sup>. None of the isolates in 238 239 this collection had the Susceptible 4 genotype. Overall, 25.0% of isolates had one of 240 the three remaining putative susceptible genotypes (Fig 4a). The dominant UK 241 MRSA sequence type is ST22 (EMRSA-15), if the CC22 isolates (70.4% of isolates) 242 were excluded then 82.8% (n=610) had one of three susceptible genotypes (Fig. 4a). 243 Lineages with a high abundance of susceptible genotypes included CC1, CC5, CC8, 244 CC30 and CC59 (Fig. 4a). Importantly, 56 of the CC8 isolates (from 24 patients) were USA300 which is the dominant clonal lineage in the United States (USA) <sup>32</sup>. All 245 246 56 USA300 isolates had the Susceptible 2 genotype and a penicillin-clavulanic acid 247 MIC below the ECOFF (Supplementary Table 3). We performed a phylogenetic 248 analysis of 580 CC8 isolates, including 485 USA300 isolates (457 MRSA, 28 MSSA) 249 from across the USA <sup>33,34</sup>. All 457 of the USA300 MRSA isolates had the Susceptible 250 2 (mecA[-7]:G-T | E246G) genotype carried on a SCCmec type IVa (n=455) or IVb

element (n=1) (one isolate was non-SCC*mec* typeable), suggesting that the majority
of the USA300 population is susceptible to penicillin-clavulanic acid (Fig 4b). We then
screened 23 USA300 isolates, distributed across the phylogeny (isolated in New
York <sup>34</sup>) for penicillin-clavulanic acid susceptibility (Fig. 4b). All had an MIC below the
ECOFF (<2 µg/ml) (Supplementary Table 5).</li>

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# 257 Heterogeneity of susceptible populations

258  $\beta$ -lactams resistance in most MRSA is heterogenous, meaning that while most cells in a population have low MICs, a fraction (10<sup>-4</sup>–10<sup>-8</sup>) can survive at much higher MICs 259 260 <sup>35</sup>. We tested representative isolates for all six penicillin-clavulanic acid susceptibility 261 genotypes in a population analysis for their susceptibility profile to penicillin-262 clavulanic acid. Isolates representing the four susceptible genotypes all displayed 263 heterogenous resistance to penicillin-clavulanic acid (Fig 5a). CFUs were drastically 264 reduced at low concentrations of penicillin (0.12 to 1 µg/ml) and susceptible isolates 265 had a median population MIC of 12 µg/ml (range 0.48 to 32) (Fig 5a). Notably, for 266 four isolates from three different genotypes (Susceptible 1, 3, and 4) the entire 267 population was completely inhibited by 4 µg/ml. The remainder of susceptible 268 isolates had 'highly resistant cells' (cells capable of growing in concentrations greater 269 than the ECOFF of  $\leq 2 \mu q/ml$ ) present at a frequency of  $10^{-5} - 10^{-8}$ . In contrast, 270 resistant isolates displayed homogenous resistance to penicillin-clavulanic acid (Fig 271 5a). CFUs of resistant isolates were unaffected at the lower penicillin concentration 272 range (0.12 to 1  $\mu$ g/ml) and had a higher median population MIC of 64  $\mu$ g/ml (range 273 64 to 128), greater than the susceptible isolates (Fig. 5a). Highly resistant cells (MIC 274  $\geq 4 \mu q/ml$ ) were also present at higher frequencies  $(1-10^{-4})$  than susceptible isolates. 275

# 276 Pencillin-clavulanic acid is effective for MRSA treatment *in vivo*

We next sought to demonstrate efficacy in physiologically relevant infection models in
which highly resistant cells would be present. First, we used a wax moth larvae

279 model of infection, larvae were infected with four different MRSA isolates, two with a 280 resistant genotype (both ST22: Resistant 1) and two with a susceptible genotype 281 (ST398 and ST8 (USA300): Susceptible 2). Treatment at approximate human 282 dosages was with penicillin, penicillin-clavulanic acid, amoxicillin, amoxicillin-283 clavulanic acid, vancomycin or PBS (vehicle). Only vancomycin offered any 284 improvement in survival for the two resistant isolates (Fig 5b and c). In contrast, for 285 the two susceptible isolates both penicillin-clavulanic acid and amoxicillin-clavulanic 286 acid increased survival in comparison to penicillin or amoxicillin alone, increasing the 287 median survival times by 40 hours (Fig 5d and e). Given that amoxicillin-clavulanic 288 acid is clinically available, we further tested its efficacy in a more physiologically 289 relevant, higher infective dose  $(1 \times 10^7 \text{ CFU})$  murine thigh infection model, with 290 approximate human dosages using a USA300 strain (strain: MRSA 43484<sup>36</sup> 291 (Susceptible 2, penicillin-clavulanic acid MIC: 0.19 µg/ml, population analysis: 292 Supplementary Fig. 4), USA300 phylogeny: Fig. 4b). Treatment with amoxicillin alone 293 in a dose range of 10 – 100 mg/kg did not reduce the bacterial loads compared to 294 vehicle treatment, whereas 100 mg/kg amoxicillin in combination with 20 mg/kg 295 clavulanic acid significantly reduced the bacterial loads to a similar level as 40 mg/kg 296 vancomycin (Dunnett's multiple comparisons test, p<0.0001) (Fig. 5f). Demonstrating 297 the efficacy of amoxicillin-clavulanic acid as a treatment in a high dose infection 298 model.

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#### 300 **PBP2a** substitutions provide a growth advantage in the presence of penicillin

As the acquisition of *mecA* (PBP2a) in a SCC*mec* element can exhibit significant fitness costs <sup>37</sup>, and affect toxicity <sup>38</sup> and biofilm formation <sup>39</sup>, we hypothesised that the PBP2a substitutions might confer a fitness advantage. We found no significant difference in biofilm formation or toxicity to human monocytic cells between any of the three PBP2a variants in two strain backgrounds (Supplementary Fig. 6). We then investigated the effect of the PBP2a substitutions on general fitness in three lineages

307 (ST22, USA300/ST8) and ST398) assayed by growth in a minimal medium, and in 308 the presence of penicillin. In minimal medium, there were only minor differences (Fig 309 6a, c, e), with the USA300 strain (BCV289) complemented with mecA<sub>246G</sub> growing 310 marginally better in early exponential phase than  $mecA_{246E}$  or  $mecA_{122I}$  (Fig. 6a). In 311 the ST22 (A75) background the  $mecA_{246E}$  strain grew slower during exponential 312 growth than the other backgrounds (Fig 6e). In the presence of penicillin, in all three 313 strain backgrounds the isolates complemented with  $mecA_{246G}$  grew better than the 314 other two variants (Fig 6b, d and f). This was most pronounced in the ST398 315 background (EC139), in which mecA<sub>246G</sub> strain grew considerably better in 316 exponential phase, reached a higher optical density and grew with a reduced 317 doubling time (dt) of 5.33 hours (95% confidence intervals (CI): 5.28 to 5.38) 318 compared to either *mecA*<sub>246E</sub> (dt: 6.65, 95% CI: 6.62 to 6.69) or *mecA*<sub>122I</sub> (dt: 6.92, 319 95% CI: 6.89 to 6.96) (Fig 6d). This demonstrated that complementation with 320 *mecA*<sub>246G</sub> provided a growth advantage in the presence of penicillin, but the 321 magnitude of this effect is influenced by strain background.

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#### 323 Discussion

324 We show that a significant proportion of clinical MRSA isolates are susceptible to a 325 combination of penicillins and a  $\beta$ -lactamase inhibitor. Susceptibility is due to one of 326 two different mutations in the mecA promoter region that both lower mecA (PBP2a) 327 expression, and in the majority of isolates, by an additional substitution in PBP2a 328 (E246G or M122I) that increases the affinity of PBP2a for penicillin in the presence of 329 clavulanic acid. It is not clear how clavulanic acid causes the increased binding 330 affinity of penicillin for PBP2a, as clavulanic acid binds to PBP2a poorly <sup>19</sup>. Modelling 331 of PBP2a shows that position 246 is located near the allosteric site but does not 332 indicate any clear mechanism for the increased affinity for penicillin (Supplementary 333 Fig.7 and Supplementary Discussion). In some isolates, a RBS mutation alone 334 appeared to be sufficient to mediate susceptibility, although given the complexity of

the regulation of  $\beta$ -lactam resistance in MRSA other genes might be involved <sup>5</sup>. Crucially, the PBP2a 246G substitution provides a fitness benefit for growth in presence of penicillin, suggesting that susceptibility to penicillin and  $\beta$ -lactamase inhibitors is a likely a case of collateral sensitivity <sup>40</sup>, which evolved due to selective pressure for maintaining the balance between fitness and resistance.

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341 In our susceptibility assays we used Iso-Sensitest media (ISA) rather than Müller-Hinton agar (MHA) as currently recommended by CLSI and EUCAST <sup>18,19</sup>, because 342 343 ISA was the recommended media for penicillin by the British Society for Antimicrobial 344 Chemotherapy (BSAC) at the beginning of the study <sup>41</sup>. Comparison between 345 susceptibility to penicillin-clavulanic acid on ISA and MHA revealed that a number of 346 isolates that were susceptible on ISA remained resistant on MHA (Supplementary 347 table 5), including isolates that responded to treatment in vivo (Fig 5b-e). This 348 suggests that MHA is not the optimum media for the detection of susceptibility of 349 penicillins and  $\beta$ -lactamase inhibitors, which is supported by a recent study which 350 revealed that MHA failed to detect susceptibility to multiple antibiotic classes that 351 were effective *in vivo*<sup>42</sup>. Our data also highlights the risk of using a single antibiotic 352 (e.g. cefoxitin for MRSA) to determine resistance to an entire antibiotic class, 353 potentially missing unexpected susceptibilities. 354 355 Both *in vitro* and *in vivo*, penicillins and clavulanic acid were efficacious at physiologically achievable concentrations <sup>43</sup>. In the absence of a clinical breakpoint, 356

357 pharmacokinetic-pharmacodynamic (PK-PD) breakpoints can be used to infer

358 susceptibility <sup>44</sup>. The tentative ECOFF wildtype cut-off of  $\leq 2.0 \ \mu$ g/ml penicillin in the

359 presence of 15 µg/ml of clavulanic acid, lies in the intermediate susceptibility

360 category (susceptible ≤0.25 µg/ml, resistant >2 µg/ml) of the EUCAST PK-PD

361 breakpoint <sup>18</sup>. Large numbers of isolates had much lower MICs, and had amoxicillin-

362	clavulanic acid zone diameters greater than the breakpoint for other pathogenic
363	species (Supplementary Table 1, 3 and 4) <sup>18</sup> . Previous studies have reported the
364	successful use of penicillins and $\beta$ -lactamase inhibitors for the treatment of MRSA in
365	rabbits and rats, and for human infections <sup>11,45-47</sup> . This work, provides a mechanistic
366	explanation for efficacy in these studies, although there have been previous
367	conflicting reports $^{9}.$ While it is unlikely that penicillins and $\beta$ -lactamase inhibitor
368	combinations would be used as a monotherapy, they would be attractive additional
369	therapeutic option for hard-to-treat infections such as multidrug-resistant MRSA $^{48}$ ,
370	particularly as $\beta$ -lactams synergise with vancomycin and daptomycin $^{49,50}.$ PK/PD
371	modelling studies, including an assessment of highly resistant cells during treatment
372	is now needed to determine the optimum dosing strategy required for sustained
373	efficacy before appropriate clinical trial could be conducted.
374	
375	Our findings demonstrate that cryptic susceptibilities to already licensed and
376	inexpensive antibiotics may emerge within constantly evolving bacterial populations,
377	which then can be exploited for the treatment of antibiotic resistant pathogens.
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#### 389 Material and methods

#### 390 Media and culture conditions

391 Bacterial strains and plasmids used in this study are described in Supplementary 392 Table 5 and 6. For routine culture, Escherichia coli (E. coli) was grown in Lysogeny 393 broth (LB) or on LB-agar (Oxoid, UK) at 37°C. S. aureus was grown on tryptone soy 394 agar (TSA), Columbia blood agar or in tryptone soy broth (TSB) (Oxoid, UK) at 28°C 395 or 37°C accordingly. E. coli and S. aureus media were supplemented with 10 µg/ml 396 chloramphenicol (Cm10) as appropriate. For growth curve studies, S. aureus strains 397 were grown in SSM9PR minimal medium (1 × M9 salts, 2 mM MgSO<sub>4</sub>, 0.1 mM CaCl<sub>2</sub>, 398 1% glucose, 1% casaminoacids, 1 mM Thiamine-HCl and 0.05 mM nicotinamide) at 399 37°C<sup>51</sup>.

400

## 401 Isolate selection

402 For the 110 sequenced isolates in Supplementary Table 1, isolates were selected 403 from sequenced isolates in our laboratory collections to provide a broad as possible 404 selection of isolates as possible from different clinically relevant lineages (Multilocus 405 sequence type (ST)1, 5, 8 22, 36, 45, 80, 88, 239, 398), with an obvious bias towards 406 lineages that dominate in the UK (e.g. 40 isolates from ST22, which is the dominant 407 lineage in the UK). For the additional 274 isolates that were combined with 24 of the 408 original isolates with MIC data (from the previous 110), we again tried to select 409 isolates from our laboratory collections that broadly covered a selection of clinically 410 relevant MRSA lineages including clonal complex (CC) 1, 5, 8 (including 56 UK 411 USA300 isolates), 22, 30, 45, 59, 72, 80, 97, 239, and 398 - amongst others). For 412 lineages with multiple isolates (CC1, n=25, CC5, n=29, CC22, n=91, CC59, n=18, 413 CC45, n=15, CC30, n=20) we used previously generated whole genome phylogenies 414 <sup>31</sup> to select isolates broadly across the phylogeny, as well as selecting isolates from 415 the same clades.

416

## 417 Antimicrobial susceptibility testing

427

418 Inocula were prepared by the growth method. At least four morphologically similar 419 colonies were touched with a sterile loop and transferred into Iso-Sensitest broth 420 (Oxoid, UK). Inoculated broth was incubated at 37°C with 200 rpm shaking until the 421 visible turbidity reached 0.5 McFarland standard. For disk diffusion and Etest method 422 testing, the 0.5 McFarland standard broth was diluted 1:10 in distilled water before 423 spreading onto agar plates. For Microbroth dilution for minimum inhibitory 424 concentrations (MIC), the broth was diluted 1:100 in Iso-Sensitest broth for 425 inoculation. 426

Disk diffusion susceptibility testing was carried out according to BSAC criteria (BSAC

428 Methods for Antimicrobial Susceptibility Testing, version 14, January 2015). 429 Temocillin disks were purchased from MAST group, UK. All other antibiotic disks 430 were purchased from Oxoid, UK. For the clavulanic acid assay, 15 µg/ml clavulanic 431 acid (Sigma-Aldrich, UK) was added to Iso-Sensitest agar (ISA) or Müller-Hinton agar 432 (MHA) (Oxoid, UK), as appropriate. After applying the antibiotic disks, all plates were 433 incubated at 35°C for 20 hours before inhibition zones were measured. Oxacillin disk 434 diffusion was also performed on MHA with 2% NaCl at 30°C for 24 hours. For disk 435 testing with clavulanic acid, susceptibility was defined as  $\geq 10$  mm increase of the 436 zone of inhibition in the presence of 15µg/ml clavulanic acid. For mecA 437 complemented strains. ISA was supplemented with 200 ng/ml anhydrotetracycline 438 (Atc) (Sigma-Aldrich, UK) to induce expression of *mecA* from pXB01, a modified tetracycline-inducible expression vector pRMC2<sup>16</sup>. Microbroth dilution for minimum 439 440 inhibitory concentrations (MIC) was performed according to BSAC guidelines <sup>52</sup>. The 441 antibiotic ranges were prepared in Iso-Sensitest broth a step higher than the final 442 concentrations of 0.015-128 µg/ml for penicillin and 1-32 µg/ml for cefoxitin. A 96 well 443 cell culture plate (Greiner Bio-One, CELLSTAR®) was loaded with appropriate 444 antibiotic dilutions. For each test isolate, 75 µl of each antibiotic dilution were added

to a row of wells and 75 µl diluted test isolate culture was added into the wells. Each
isolate was tested in triplicate. The 96 well plate was then placed in a plastic bag to
minimize evaporation and was incubated at 35°C for 20 hours. Penicillin and
cefoxitin Etest antimicrobial susceptibility testing on selected MRSA isolates was
performed according to manufacturer's guidance. Medium used for Etest was ISA
with or without 15 µg/ml clavulanic acid. Etest strips were purchased from bioMérieux
UK Ltd.

452

# 453 **Construction of S.** *aureus* gene deletion and complementation mutants

454 mecA deletion mutants in mecA-MRSA strains (Supplementary Table 7) were 455 generated by allelic exchange with the temperature-sensitive vector pIMAY, as 456 described previously <sup>53</sup>. Primers used for *mecA* deletion are listed in Supplementary 457 Table 8. Upstream sequence (AB) and downstream sequence (CD) of the S. aureus 458 gene to be deleted were amplified with primers A/B or C/D using KOD Hot Start DNA 459 Polymerase (Merck, UK). PCR products AB and CD were used as templates to 460 obtain deletion construct AD with primers A/D in a splicing overlap extension (SOE) 461 PCR. Product AD was digested with restriction enzymes KpnI and SacI and ligated to 462 pIMAY digested with the same enzymes. The resulting plasmids were designated 463 pIMAY $\Delta$ mecA. The plasmids were transformed into E. coli DC10B (a dcm deletion 464 mutant of DH10B), allowing the plasmid to be directly transferred into S. aureus 465 strains <sup>53</sup>. Plasmid DNA extracted from DC10B was then electroporated into recipient 466 strains to create knockout mutants.

467

For complement expression of *mecA*, the genes were cloned into expression plasmid
pXB01, a derivate of tetracycline-inducible expression vector pRMC2 with the *blaZ*gene deleted <sup>54</sup>. The *mecA* gene variants including the ribosome binding site were
amplified from genomic DNA with primers: *mecA*-F-Kpnl / *mecA*-R-Sacl. PCR
products were digested with Kpnl and Sacl and ligated with the pXB01 vector

473 cleaved with the same enzymes, generating plasmids pmecA<sub>246E</sub>, pmecA<sub>246G</sub>, and

474 pmecA<sub>1221</sub>. The plasmids were transformed into *E. coli* DC10B, and plasmid DNA then

- 475 extracted and electroporated into mecA-deletion strains for complementation with
- 476 expression induced with 200 ng/ml Atc.
- 477

# 478 Antimicrobial susceptibility testing of Danish clinical isolates

- 479 Antimicrobial susceptibility testing was performed on a selection of clinical isolates
- 480 obtained from the Danish surveillance of MRSA and *S. aureus* causing bacteraemia.
- 481 The selection was based on a total of 270 isolates including 100 isolates (52 MSSA,
- 482 48 MRSA) from 2011 on which whole genome sequence data were previously
- 483 obtained and 170 consecutively received non-CC398 MRSA isolates in 2016. spa
- 484 types were obtained for all isolates. Antimicrobial susceptibility testing was performed
- 485 as described above except that 0.5 McFarland inoculum was prepared using a
- 486 densitometer and ISA plates with and without 15 µg/ml clavulanic acid was
- 487 purchased as custom made plates (SSI Diagnostica, Hilleroed, Denmark).
- 488

#### 489 Expression and purification of PBP2a variants in *E. coli*

- 490 PBP2A variants (PBP2a<sup>E246</sup>, PBP2a<sup>E246G</sup> and PBP2a<sup>M122I</sup> from MRSA strains RVC5,
- 491 BCVA289 and ARARH150, respectively; residues 26-668, with a G26M mutation)
- 492 were overexpressed using the auto-induction expression method at 25°C <sup>55</sup>. Cells
- 493 were harvested by centrifugation after 20 hours of expression and lysed using
- 494 Bugbuster (Novagen, Merck Millipore) containing 10 U/ml of benzonase nuclease
- 495 (Novagen, Merck Millipore) and Protease Inhibitor Tablets, EDTA-free (Pierce
- 496 Biotechnology, Thermo Fisher Scientific), following the manufacturer's instructions.
- 497 After cell disruption, the lysates were cleared, and the soluble proteins were purified
- 498 using HisPur Ni-NTA Resin columns (Pierce Biotechnology, Thermo Fisher Scientific)
- 499 under native conditions, according to the manufacturer's instructions. The expression
- 500 and purification yields were monitored by SDS-PAGE. The most concentrated elution

501 fractions were buffer exchanged to 20 mM sodium phosphate buffer pH 7.4 using

502 PD-10 Desalting Columns (GE Healthcare Life Sciences), following the

503 manufacturer's instructions. Protein concentrations were assessed using the BCA

504 Protein Assay Kit (Pierce Biotechnology, Thermo Fisher Scientific).

505

# 506 Bocillin FL PBP2a and penicillin G/clavulanic acid binding assays

507 The binding affinity of PBP2a for different antibiotics has been determined using a fluorescent penicillin reporter reagent, Bocillin FL <sup>56-58</sup>. In this study, the affinities of 508 509 PBP2A variants for penicillin and clavulanic acid were determined using the same 510 approach. Briefly, a reaction mixture containing 25 µg/ml of a purified PBP2a variant 511 in 20 mM sodium phosphate buffer (pH 7.4) and various concentrations (0, 0.5, 1, 5, 512 10, 20, 50, 100, 200, 500, 1000, 2000 µg/ml) of penicillin G or clavulanic acid was 513 incubated at 37°C for 20 min. To test if the presence of clavulanic acid affected the 514 affinity for penicillin G, increasing concentrations of penicillin G together with 15 515 µg/ml of clavulanic acid were assayed. A final concentration of 20 µM Bocillin FL 516  $(13.3 \mu g/ml)$  was added to the reaction followed by 10 min incubation at 37°C. The 517 reactions were quenched by adding SDS loading buffer and heating at 95°C for 10 518 min. Samples were visualized using 10% Tris-Glycine-SDS PAGE. Protein gels were 519 washed in distilled water for 10 min and scanned using a 473 nm laser of a Fuji 520 Fluorescent Analyzer TLA-5100. Fluorescent intensity was quantified by ImageJ 521 software and IC<sub>50</sub> was calculated from three independent assays using GraphPad 522 Prism 5 software.

523

## 524 RNA isolation and quantitative real-time PCR (RT-qPCR)

For each selected isolate, 15 ml of early log phase culture with an  $OD_{595nm}$  of 0.3 was treated with 10 µg/ml oxacillin for 1 h at 37 °C to induce *mecA* expression and its respective untreated culture was used as a baseline control. After the induction, the  $OD_{595nm}$  of the control culture was adjusted to be equal to that of the oxacillin induced

529 one if necessary and 10 ml of both cultures were spun for 10 min at 4,500 x g at 20 530 °C. About 0.5 ml supernatant was left behind to re-suspend the pellet and 1 ml of 531 RNAprotect Bacteria Reagent (Qiagen) was added and mixed immediately by 532 vortexing. After incubation for 5 min at room temperature, cultures were spun for 5 533 min at 10,000 x g at 4 °C to pellet the cells. Pellets were snap-frozen and stored at -534 80 °C until RNA isolation. Total RNA was isolated using Qiagen RNeasy Mini Kit 535 (Qiagen) and an additional DNAse treatment was performed with the Ambion 536 TURBO DNA-free kit and cDNA was produced using QuantiTect Reverse 537 Transcription Kit (Qiagen). RT-qPCR was performed using the SensiFast SYBR No-538 ROX Kit (Bioline) on a Rotor-Gene Q machine (Qiagen) using mecA and blaZ 539 primers Gene expression fold-changes in induced cultures were calculated relative to control cultures using the  $\Delta\Delta$ Ct method <sup>59</sup> with *gyrB* as the reference. 540

541

#### 542 Wax moth larva infection and treatment

543 The wax moth larvae assay was based on that previously described by Desbois et al 544 <sup>60</sup>. Galleria mellonella larvae were purchased in bulk from Livefood UK. Larvae were 545 stored at 4°C upon arrival and kept at 37°C during the course of the assay. MRSA 546 strains RVC5, 0081, EC139, BCVA289 were selected for evaluation of antimicrobial 547 activities of penicillin and clavulanic acid in combination. Single bacterial colonies 548 were picked to inoculate 5 ml of TSB, and cultures were grown overnight (~16 hours) 549 at 37°C and 200 rpm shaking. Cultures were then diluted 1:100 into 5 ml of fresh 550 TSB and grown for a further 4 hours at 37°C and 200 rpm shaking. Cultures were 551 then centrifuged at 2,500g for 10 minutes, and pellets resuspended in sterile 552 phosphate buffered saline (PBS) to an OD<sub>595</sub> of 0.2, giving approximately  $1.5 \times 10^8$ 553 CFU/ml. For each strain, six groups of G. mellonella (n=10 in each group) were 554 injected with 10  $\mu$ l aliquots (~1.5 × 10<sup>6</sup> CFU) of resuspended culture behind the rear 555 thoracic segments using a Tridak Stepper Pipette Dispenser (Dymax, UK). Groups of 556 G. mellonella were treated by injection with 50 mg/kg vancomycin, 20 mg/kg penicillin

557 sodium salt, 20 mg/kg clavulanic acid, 20 mg/kg penicillin sodium salt combined with 558 20 mg/kg clavulanic acid, 20 mg/kg amoxicillin, 20 mg/kg amoxicillin combined with 559 20 mg/kg clavulanic acid or PBS at 2, 24 and 48 hours after inoculation. The 560 treatments were given blind and the treatment identities not revealed until the 561 experiment was completed. Larvae were considered dead when they did not respond 562 to touch to the head. Survival curves were generated and analysed using GraphPad 563 Prism 6 software. Fig. 5 shows results of a single representative experiment, a 564 replication experiment with broadly similar results is shown in Supplementary Fig. 5. 565

### 566 Murine infection model

567 Fresh overnight colonies from a 5% horse blood agar plate were suspended in saline 568 to an OD<sub>546</sub> of 0.13, giving approximately  $2 \times 10^8$  CFU/ml. Mice (NMRI female mice, 569 26 - 30 gram (Taconic, Denmark), 6-8 weeks old were inoculated intramuscularly 570 with 0.05 ml of the suspension in the left thigh (1 x  $10^7$  CFU). Approximately 0.5 hrs 571 before inoculation, mice were treated orally with 45 µl Nurofen Junior (20 mg 572 ibuprofen/ml - corresponding to 30 mg/kg) for pain relief. Four mice in each group 573 were treated with a single subcutaneous dose of 0.2 ml with 10, 30 or 100 mg/kg 574 amoxicillin (Amoxil, GlaxoSmithKline Middlesex UK) alone or in combination with 2, 6 575 or 20 mg/kg clavulanic acid (Augmentin, Beecham Group Ltd, Middlesex, UK) or 40 576 mg/kg vancomycin (Fresenius Kabi, Halden, Norway) or saline one-hour post 577 infection. Mice were sacrificed at 1 hour for the start of treatment control group and at 578 5 hours post infection for the treatment groups by cervical dislocation and thighs 579 were collected and kept at -80°C. Each sample thigh was homogenized in 5 ml saline 580 using a Dispomix® Drive, and serially diluted in saline and twenty microliter spots of 581 serial dilutions were plated on blood agar plates. All agar plates were incubated for 582 18 - 24 hrs at 35°C. Statistical comparison was carried out using a 1 way ANNOVA 583 and Dunnett's multiple comparison, was performed for treatment groups comparing 584 against the vehicle group in GraphPad Prism software. All animal procedures were

585 carried out at the Statens Serum Institute (SSI) and approved by the Danish Animal 586 Procedure Inspectorate. Ethical approval was granted for the murine thigh infection 587 model (2016-15-0201-01049). The SSI Animal Welfare Committee (SSI-AWC -588 equivalent an Institutional Animal Care and Use Committee (IACUC) requires that 589 each experiment is further approved by the supervising laboratory animal 590 veterinarians who are also part of the IACUC. All animals were randomised on arrival 591 at SSI, and sample sizes were based on a combination of statistical analysis and the 592 principles of 3R, that the minimum number of animals were used that were expected 593 to provide statistically significant difference considering the expected intra-group 594 variability of the infection model used.

595

# 596 **Growth curves**

597 To assess the effect of different mecA variants on the growth of MRSA strains in 598 liquid culture, Bioscreen C optical growth analyzer (Lab system, Finland) was used to 599 monitor the growth rates of deletion mutants A75 $\Delta$ mecA, BCVA289 $\Delta$ mecA and 600 EC139 $\Delta$ mecA complemented with the three mecA variants. Briefly, overnight 601 cultures were diluted 1/1000 into fresh SSM9PR minimal medium with or without 16 602 µg/ml penicillin supplemented with 200 ng/ml Atc to induce the expression of 603 plasmid-borne mecA gene. For each strain, 300 µl of inoculated medium was added 604 into wells of the microplate in triplicate. Fresh medium with Atc was also added to 605 three wells acting as blank controls. Cultures were incubated at 37 °C with 606 continuous shaking for 24 hours and an optical density measurement at OD<sub>600nm</sub> was 607 taken every 30 mins. Growth curves were analysed using the GraphPad Prism 6 608 software and doubling time calculated using non-linear regression using an 609 exponential growth equation with a least square fit, with Y0 constrained at the 610 minimum optical density measured (an OD<sub>600nm</sub> of 0.069).

611

## 612 **Toxicity and biofilm assays**

613	Immortalised human monocyte macrophage THP-1 cell lines were used as described
614	previously <sup>61</sup> . Briefly, the cell line was grown in individual 30 mL suspensions of
615	RPMI-1640, supplemented with 10% heat-inactivated fetal bovine serum (FBS), 1 $\mu$ M
616	L-glutamine, 200 units/mL penicillin, and 0.1 mg/mL streptomycin at 37 $^\circ\text{C}$ in a
617	humidified incubator with 5% $CO_2$ . Cells were routinely viewed microscopically every
618	48–60 h and harvested by centrifugation at 1,000 rpm for 10 min at room
619	temperature and resuspended to a final density of $1-1.2 \times 10^6$ cells/mL in tissue-
620	grade phosphate buffered saline. This procedure typically yielded >95% viability of
621	cells as determined by trypan blue exclusion and easyCyte flow cytometry. To
622	monitor S. aureus toxicity, 20 $\mu L$ of cells were incubated with 20 $\mu L$ of bacterial
623	supernatant and incubated for 12 min at 37°C. For the USA300 strains, supernatants
624	were diluted to 30% of the original volume in TSB as these isolates were
625	considerably more toxic than the single-patient isolates. Cell death was quantified
626	using easyCyte flow cytometry using the Guava viability stain according to
627	manufacturer's instructions. Experiments were done in triplicate, and error bars
628	indicate the average $\pm$ the 95% confidence interval of multiple independent
629	experiments.
630	
631	Biofilm formation was quantified using a 1:40 dilution from overnight cultures into 100

632 µL of fresh TSB supplemented with 0.5% sterile filtered glucose (TSBG) in 96-well 633 polystyrene plate (Costar). Perimeter wells of the 96-well plate were filled with sterile 634 H<sub>2</sub>O and plates were placed in a separate plastic container inside a 37°C incubator 635 and grown for 24 h under static conditions. For the transposon mutants, erythromycin 636 (5 µg/mL) was added to the growth medium. Semi-quantitative measurements of 637 biofilm formation on 96-well polystyrene plates were determined based on the method of Ziebuhr et al <sup>23</sup>. Following 24-h growth, plates were washed vigorously five 638 639 times in PBS, dried and stained with 150 µL of 1% crystal violet for 30 min at room 640 temperature. Following five washes of PBS, wells were re-suspended in 200 µL of

641 7% acetic acid, and optical density at 595 nm was recorded using a Fluorimeter plate
642 reader (BMG Labtech). For this experiment the assays were performed in triplicate
643 on each plate and repeated three times.

644

668

#### 645 **ECOFF determination**

In order to help split isolates into phenotypically "wildtype" and "non-wildtype" based on MIC distributions, a series of mixture models were fitted to the data for each drug independently, ranging from 1 to 5 normal distributions. Each model was fitted to the distribution of log-transformed MIC by maximum likelihood. The likelihood function

650 was based on a multinomial distribution over the set of intervals

 $[0,x_1,...,x_n,\infty]$  where x<sub>i</sub> stands for each of the tested concentrations for the drug

considered. For each drug, the 5 models were compared using Akaike's Information
Criterion to determine the optimal number k of normal distribution components. If the
best model was k=1, the distribution was labelled as unimodal, and no further
analysis was made. For drugs with k≥2, the following analyses were then carried out:
Predicted distribution of MIC per component: for each component, we multiplied
the probability mass at each concentration (= pdf integrated over the immediately

lower interval) by the total number of isolates tested.

659 2. Direct classification of isolates by component: for each MIC, we split the isolates660 into the k components according to their relative probabilities at that point.

3. Optimal cut-off: we then sought to split each distribution into two modes. First, we estimated candidate cut-off points between each successive component of the mixture model. Each cut-off was computed as the concentration that minimised the risk of misclassification of isolates between the model components (achieved by minimising the difference between the cumulative density functions of the two components considered). For example, with k=3, there are three components centred at MIC values Y1<Y2<Y3; we then calculated two candidate cut-offs: one</p>

separating component 1 from 2+3, and the other separating 1+2 from 3. For each

669 cut-off, we then computed the number of isolates that would be misclassified,

670 using the above direct classification as a reference: e.g. for the first cut-off, the

- number of isolates classified as 2 or 3 with MIC below the cut-off, plus the
- number of isolates classified as 1 with MIC above the cut-off.
- 673

# 674 **Bioinformatics**

675 Whole genome sequences were assembled using the pipeline described previously 676 <sup>62</sup>. For each isolate the sequence reads were used to create multiple assemblies using VelvetOptimiser v2.2.5<sup>63</sup> and Velvet v1.2<sup>64</sup>. The assemblies were improved by 677 scaffolding the best N50 and contigs using SSPACE <sup>65</sup> and sequence gaps filled 678 679 using GapFiller<sup>66</sup>. Presence of PBP2a substitutions were identified by extracting the 680 PBP2a sequence from the assembled genome sequences, aligning the PBP2a sequences using Muscle <sup>67</sup> in Seaview <sup>68</sup> and then identifying the presence of PBP2a 681 682 substitutions using a custom python script. The mecA promoter mutations were 683 identified using a similar approach using *in silico* PCR to identify the *mecA* promoter 684 region and then aligning the mecA promoter sequence and identifying mecA 685 mutations using a custom python script. The presence of the *blaZ* genes was 686 confirmed using BLAST against assemblies, this identified that 273/298 of the 687 previously screened WGS isolates had a single copy of *blaZ* (excluding *blaZ*-688 negative or isolates with two copies or truncated copies of *blaZ*). BlaZ amino acid 689 sequences were then extracted, aligned and amino acids at positions 128 and 216 690 compared to identify the BlaZ type (Type A: 128:T, 216:S, Type B: 128:K, 216:N, 691 Type C: 128:T, 216:N, Type D: 128:A, 216:S, Type E (LGA251): 128:L, 216:S, Type 692 F (a distinct type identified in this work): 128:A, 216:N) as previously described <sup>16,69,70</sup>. 693 Phylogenetic analysis of the CC8 isolates was carried out as previously described <sup>33</sup>, 694 briefly, sequence reads were mapped using SMALT v0.7.4 695 (http://www.sanger.ac.uk/science/tools/smalt-0) to the S. aureus USA300\_FPR3757 696 reference genome (accession: CP000255.1)<sup>71</sup>. A core genome alignment was

697 created after excluding mobile genetic element regions, variable sites associated with recombination (detected with Gubbins<sup>72</sup>) and sites with more than 5% proportion of 698 699 gaps (i.e. sites with an ambiguous base). A maximum likelihood (ML) phylogenetic tree was generated with RAxML v8.2.8<sup>73</sup> based on generalised time reversible 700 701 (GTR) model with GAMMA method of correction for among site rate variation and the 702 phylogenetic tree annotated using Figtree (http://tree.bio.ed.ac.uk/software/figtree/). 703 Phylogenetic analysis of PBP2a sequences was constructed by using PhyML v3.0 in 704 Seaview with a Whelan and Goldman (WAG) substitution model and 100 bootstrap 705 replicates <sup>74</sup>.

706

707

# 708 **Population analysis**

Population analysis was carried out as described by Kim et al <sup>75</sup> with minor 709 710 modifications. Strains were grown overnight in tryptic-soya broth (TSB), and serial 711 10-fold dilutions were plated in triplicate onto Iso-Sensitest Agar (Oxoid) plates 712 containing varying concentrations of penicillin with 15 µg/ml clavulanic acid. Plates 713 were incubated for 24 hours at 35°C. A mean of the three platings were plotted in 714 Figure 5a. Biological replicates for strains BCVA289 and 0081 were generated and 715 plotted in comparison to the original results in Supplementary Figure 4, showing 716 broadly similar results. 717 718 Structural modelling of PBP2a

719 Co-ordinates from representative structures of both the Gly246 (accession code

720 3ZFZ, <sup>21</sup>) and Glu246 (accession code 1VQQ, <sup>76</sup>) forms of PBP2a were used to

compare possible effects induced by sidechain alteration. Figures were prepared

using Chimera <sup>77</sup> and Pymol (The PyMOL Molecular Graphics System, Version

723 1.2r3pre, Schrödinger, LLC).

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## 1016 Author contributions

- 1017 Design of the study: EMH, XB, SJP, MAH. *mecA* deletion and complementation,
- 1018 expression analysis, bocillin assays: XB. Antimicrobial susceptibility testing: XB, BB,
- 1019 NG, KB. Biofilm and toxicity assays: HC and RM. Antimicrobial susceptibility testing
- 1020 of Danish isolates: JL, ARL. Determination of the ECOFF: OR. Structural analysis of
- 1021 PBP2a: AL. Infection and treatment experiments: EMH, XB, CVL. Bocillin binding
- 1022 assays: IRG and RGS. Bioinformatics analysis of whole-genome sequence data:
- 1023 EMH, FC, SR, DJ. Collection of USA300 isolates: ACU, FDL. Wrote bioinformatics
- 1024 scripts: NG. Analysis and interpretation of the data: CUK, GP, MTGH, JP.
- 1025 Coordinated the study and wrote the manuscript: EMH. Responsible for supervision
- 1026 and management of the study: SJP and MAH. All authors read, contributed to and
- 1027 approved the final manuscript.

### 1028 Competing interests

- 1029 CUK is a consultant for the World Health Organization (WHO) Regional Office for
- 1030 Europe, QuantuMDx Group Ldt, and the Foundation for Innovative New Diagnostics,
- 1031 which involves work for Cepheid Inc., Hain Lifescience and WHO. CUK is an advisor
- 1032 to GenoScreen. The European Society of Mycobacteriology awarded CUK the
- 1033 Gertrud Meissner Award, which is sponsored by Hain Lifescience. The Bill & Melinda
- 1034 Gates Foundation, Janssen Pharmaceutica, and PerkinElmer covered CUK's travel
- 1035 and accommodation to present at meetings. The Global Alliance for TB Drug
- 1036 Development Inc. and Otsuka Novel Products GmbH have supplied CUK with
- 1037 antibiotics for *in vitro* research. CUK has collaborated with Illumina Inc. on a number
- 1038 of scientific projects. SJP and JP are consultants to Next Gen Diagnostics Llc. SJP is
- 1039 a consultant to Specific Technologies. All other authors declare no competing
- 1040 financial interest

1041

#### 1042 Data avaliblity

1043 All data generated or analysed during this study are included in this published article1044 (and its supplementry information files).

1045

# 1046 Figure Legends

1047 Figure 1: Penicillin susceptibility in the presence of clavulanic acid. (a) Figure 1048 shows a representative image of two MRSA isolates grown on Iso-sensitest agar or 1049 Iso-sensitest agar with  $15\mu g/ml$  of clavulanic acid with a penicillin E-test strip applied, 1050 the upper susceptible isolate shows increased susceptibility to penicillin in the 1051 presence of clavulanic acid, while the lower resistant isolate shows only a minor 1052 effect. (b) E-test determination of minimum inhibitory concentration of penicillin alone 1053 (black) and penicillin plus 15 µg/ml clavulanic acid (grey) of a selection of penicillin-1054 clavulanic acid susceptible (n=14) and resistant (n=10) MRSA isolates. The red line 1055 indicates the current EUCAST clinical break point of  $\leq 0.125 \mu g/ml$ . The effect of 1056 clavulanic acid on susceptibility as measured by disk diffusion to different  $\beta$ -lactam 1057 antibiotics in a panel of (c) penicillin-clavulanic acid resistant MRSA isolates (n=46), 1058 and (d) penicillin-clavulanic acid susceptible MRSA isolates (n=32). Red bars 1059 indicate isolates grown on Iso-Sensitest Agar (ISA) alone and blue bars indicate 1060 isolates grown on ISA supplemented with 15µg/ml clavulanic acid. Y-axis indicates 1061 the zone of inhibition in millimetres. Error bars indicate the standard deviation of the 1062 mean. Note: OX(MHA) = Oxacillin performed on Müller-Hinton agar (recommended 1063 media by EUCAST and CLSI). 1064

Figure 2: PBP2a substitutions mediating pencillin susceptibility. (a) Minimium
 inhibitory concentrations (MIC) of penicillin (black) or penicillin in the presence of 15
 µg/ml clavulanic acid (grey) for wildype strains EC139 (ST398) and BCVA289

1068 (USA300), and mecA mutants and complemented mutant strains with either an empty vector (p) or one of the three different alleles of mecA (PBP2a<sup>246E</sup> (p246E), 1069 1070 PBP2a<sup>246G</sup> (p246G), and PBP2a<sup>112I</sup> (p112I). Results presented are the mean of three 1071 independent experiments. (b) Bocillin competition assay to determine the  $IC_{50}$  of penicillin for the three PBP2a variants (PBP2a<sup>246E</sup> – 'WT', PBP2a<sup>246G</sup> and 1072 PBP2a<sup>M122I</sup>). (c) Bocillin competition assay to determine the  $IC_{50}$  of penicillin in the 1073 presence of 15µg/ml clavulanic acid for the three PBP2a variants (PBP2a<sup>246E</sup> – 'WT', 1074 PBP2a<sup>246G</sup> and PBP2a<sup>M122I</sup>). The fluorescence intensity of the bands was plotted as 1075 1076 the percentage of unbound proteins as a function of penicillin G concentration, and 1077 the IC<sub>50</sub> value was calculated from the plot. Data points represent the average of 1078 three replicates and the curve is the predicted nonlinear regression result. (d) Mixture 1079 models of MIC distributions of 298 MRSA isolates for penicillin in the presence of 15 1080 µg/ml clavulanic acid, the best model is a mixture of 4 normal distributions, with a 1081 proposed cut-off at 2.449, and (e) for penicllin alone, the best model is a mixture of 4 1082 normal distributions, with a proposed cut-off at 0.218. The graphs show the proposed 1083 categorisation of the MIC distribution into "wildtype" isolates (low MIC, green hues) 1084 and "non-wildtype" isolates (high MIC, amber hues), and the proposed empirical cut-1085 off as the vertical dashed line. Amber isolates that fall below the cut-off and green 1086 isolates that fall above the cut-off show the expected classification errors by applying 1087 the cut-off.

1088

#### 1089 Figure 3: Genetic basis of MRSA penicillin/clavulanic acid susceptibility. (a)

Summary of PBP2a substitutions and *mecA* promoter mutations. Figure shows a representation of the domain structure of the PBP2a protein and *mecA* promoter with the location of the two PBP2a substitutions and *mecA* promoter associated with penicillin susceptibility indicated and mean penicillin MIC in the presence of 15  $\mu$ g/ml of clavulanic acid for isolates with that genotype. (b) Relative *mecA* expression measured by RT-gPCR. Figure shows the relative *mecA* expression after oxacillin

1096 induction, normalised to qyrB for isolates with mecA[-7]:T (n=3), mecA[-7]:G (n=7) 1097 and mecA[-33]:T (n=6). Error bars indicate the standard deviation of the mean. Data 1098 were analysed with a two-tailed, unpaired t-test. \* P = 0.0048, \*\* P = 0.0016. (c) MIC 1099 distributions for penicillin (upper graph) and penicillin with 15 µg/ml clavulanic acid 1100 (lower graph) with the number of isolates with each genotype combination of PBP2a 1101 substitutions and mecA promoter mutations - indicated by colouring of the histogram 1102 bars. (d) Joint distribution of MIC for 298 MRSA isolates, x axis shows penicillin MIC 1103 and y axis shows pencillin MIC in the presence of 15  $\mu$ g/ml clavulanic acid. Colour 1104 and shape of the plot points indicate the genotype of the isolate.

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# 1106 Figure 4: Prevalence and population genomics of penicillin-clavulanic acid.

(a) Graph shows the percentage abundance in the overall population and by clonal

complex (CC) of penicillin-clavulanic acid susceptible and resistant genotypes in
2282 clinical MRSA isolates from Cambridge, UK (b) Presence of SCC*mec* elements
in clonal complex (CC)8 and USA300 isolates. Figure shows a maximum likelihood
phylogenetic tree constructed from core genome SNPs of 580 CC8 isolates. USA300
isolates are indicated by blue colouring of the branches. Tips of the tree indicate the

1113 SCC*mec* type (red = IVa, blue = IVb = orange = IVc, pink = IVd, purple = IVg, NT

1114 (light blue) = non-typeable, - = negative for SCC*mec*). Isolates that were

1115 phenotypically tested for penicillin-clavulanic acid susceptibility are indicated with an

1116 asterix. Additionally, isolates: BCVA289, which was used for *in vivo* testing (wax

1117 moth) and 43484, which was used for *in vivo* testing (murine thigh) are included for

1118 comparison.

1119

1120 Figure 5: Penicillins and clavulanic acid are efficacious for the treatment of

1121 **susceptible MRSA. (a)** Population analysis of resistance to penicillin and clavulanic

1122 acid. Figure shows the log10 CFU/ml of the different strains at various concentrations

1123 of penicillin G (µg/ml) in the presence of 15 µg/ml clavulanic acid on Isosenstest

1124 plates. Plotted points are the mean of three technical replicates (biological replicates 1125 for BCVA289 and 0081 shown in Supplementary Fig. 4). Survival curves for wax 1126 moth larvae (*Galleria mellonella*) infected with  $\sim 1.5 \times 10^6$  CFU of: (b) 0081 (ST22) 1127 (penicillin-clavulanic acid MIC = 4 µg/ml - resistant). (c) RVC5 (ST22) (MIC = 12 1128  $\mu$ g/ml - resistant), (d) BCVA289 (ST8 - USA300) (MIC = 0.25  $\mu$ g/ml - susceptible), 1129 and (e) (EC139 (ST398) (MIC = 0.25µg/ml - susceptible). Ten larvae in each group 1130 were experimentally infected and then treated at 2, 24, and 48 hours with 1131 vancomycin (50 mg/kg), penicillin (20 mg/kg) clavulanic acid (20 mg/kg), penicillin-1132 clavulanic acid (1:1 ratio - 20 mg/kg), amoxicillin (20 mg/kg), amoxicillin-clavulanic 1133 acid (1:1 ratio - 20 mg/kg) and PBS alone. Presented data are from a single 1134 representative experiment, a replication experiment is presented in Supplementary 1135 Fig. 5. (f) Effect of amoxicillin alone and in combination with clavulanic acid against 1136 MRSA in a murine thigh infection model. Four mice in each treatment group were inoculated with 7 log<sup>10</sup> CFU of MRSA strain 43484 (USA300) and treated 1-hour post 1137 1138 inoculation with a single subcutaneous injection of either vancomycin (40mg/kg), 1139 Amoxicillin (Amox) (10, 30, 100 mg/kg) alone or in combination (5:1 ratio) with clavulanic acid (Clav) (2, 6, 20 mg/kg). The bar indicates the mean. \*\*\*\* indicates a 1140 1141 significant difference (p<0.0001, Dunnett's multiple comparisons test) between 1142 vehicle control. NS indicates there was no significant difference (p=0.0982, Dunnett's 1143 multiple comparisons test) between the combined amoxicillin 100 mg/kg: clavulanic 1144 acid 20mg/kg and vancomycin 40 mg/kg.

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# 1146 Figure 6: PBP2a<sup>246G</sup> substitution provides an increased growth rate in the

**presence of penicillin.** Figure shows growth curves for strain BCV289∆*mecA* grown

- 1148 in (a) SSM9PR minimal medium and (b) SSM9PR with 16  $\mu$ g/ml penicillin, strain
- 1149 EC139∆*mecA* grown in (c) SSM9PR minimal medium and (d) SSM9PR with 16
- 1150 µg/ml penicillin, and strain A75∆mecA grown in (e) SSM9PR minimal medium and (f)
- 1151 SSM9PR with 8 µg/ml penicillin. Lines are coloured depending on the vector the two

- 1152 strains were complemented with: orange = vector only control, green = vector
- 1153 expressing  $mecA^{246G}$ , blue = vector expressing  $mecA^{246E}$ , black = vector expressing
- 1154 *mecA*<sup>122I</sup>. The mean of a minimum of 6 independent replicates are plotted and error
- 1155 bars indicate standard deviation.







Penicillin MIC µg/ml





BCVA289 (ST8-USA300)





EC139 (ST398)





