

This is a repository copy of Two codependent routes lead to high-level MRSA.

White Rose Research Online URL for this paper: <u>https://eprints.whiterose.ac.uk/220292/</u>

Version: Accepted Version

Article:

Adedeji-Olulana, A.F. orcid.org/0000-0001-8009-1755, Wacnik, K. orcid.org/0000-0002-9921-6746, Lafage, L. orcid.org/0000-0002-8105-5083 et al. (17 more authors) (2024) Two codependent routes lead to high-level MRSA. Science, 386 (6721). pp. 573-580. ISSN 0036-8075

https://doi.org/10.1126/science.adn1369

© 2024 The Authors. Except as otherwise noted, this author-accepted version of a journal article published in Science is made available via the University of Sheffield Research Publications and Copyright Policy under the terms of the Creative Commons Attribution 4.0 International License (CC-BY 4.0), which permits unrestricted use, distribution and reproduction in any medium, provided the original work is properly cited. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/

Reuse

This article is distributed under the terms of the Creative Commons Attribution (CC BY) licence. This licence allows you to distribute, remix, tweak, and build upon the work, even commercially, as long as you credit the authors for the original work. More information and the full terms of the licence here: https://creativecommons.org/licenses/

Takedown

If you consider content in White Rose Research Online to be in breach of UK law, please notify us by emailing eprints@whiterose.ac.uk including the URL of the record and the reason for the withdrawal request.



1 Two co-dependent routes lead to high-level MRSA

- 2
- 3 Abimbola Feyisara Adedeji-Olulana^{1†,}, Katarzyna Wacnik^{2,3,†}, Lucia Lafage^{2,3}, Laia
- 4 Pasquina-Lemonche^{1,3}, Mariana Tinajero-Trejo^{2,3}, Joshua A. F. Sutton^{2,3}, Bohdan Bilyk^{2,3},
- 5 Sophie E. Irving^{2,3}, Callum J. Portman Ross^{2,3}, Oliver J. Meacock¹, Sam A. Randerson¹,
- 6 Ewan Beattie¹, David S. Owen^{2‡}, James Florence^{2,3}, William M. Durham¹, David P.
- 7 Hornby^{2,3}, Rebecca M. Corrigan^{2,3,4}, Jeffrey Green^{2,3}, Jamie K. Hobbs^{1,*}, Simon J. Foster^{2,3,*}

8

- ¹ Department of Physics and Astronomy, University of Sheffield, Sheffield, UK.
- ² School of Biosciences, University of Sheffield, Sheffield, UK.
- ³ The Florey Institute, University of Sheffield, Sheffield, UK.
- ⁴School of Medicine, University College Dublin, Dublin, Ireland.
- 13 [†]These authors contributed equally to this work.
- ¹⁴ [‡] Present address. Department of Biosciences and Chemistry, Sheffield Hallam University,
- 15 UK.
- ^{*}Corresponding author. Email : jamie.hobbs@sheffield.ac.uk; s.foster@sheffield.ac.uk.
- 17

18 One Sentence Summary

19 High-level resistance in MRSA requires two pathways that reveal a novel cell division mode.

20 Abstract

21 Methicillin resistant S. aureus (MRSA) is of major clinical concern, in which acquisition of mecA, encoding the cell wall peptidoglycan biosynthesis component Penicillin Binding Protein 22 23 2a (PBP2a), confers resistance to β -lactam antibiotics. In the presence of antibiotics we show that MRSA adopts an alternative cell division mode, with altered peptidoglycan architecture at 24 the division septum. PBP2a can replace the transpeptidase activity of the endogenous and 25 essential PBP2, but not that of PBP1, which is responsible for the distinctive native septal 26 peptidoglycan architecture. Successful division without PBP1 activity, requires the alternative 27 division mode and is enabled by several possible chromosomal, potentiator (pot) mutations. 28 MRSA resensitizing agents differentially interfere with the two co-dependent mechanisms 29 required for high-level antibiotic resistance, providing opportunities for new interventions. 30

31 Introduction

32 Antibiotics are at the heart of modern medicine, but their efficacy is increasingly challenged by the spread of antimicrobial resistance (AMR) (1). MRSA is a so-called AMR "superbug", 33 that causes over 120,000 deaths per annum (2). Methicillin was introduced to circumvent 34 clinical β-lactamase-mediated resistance, but soon became compromised due to the spread of 35 MRSA (3). Resistance in MRSA is primarily based on the acquisition of the mecA gene 36 encoding a novel PBP, named PBP2a, characterised by its low affinity for a broad range of β-37 lactams (3, 4). The mecA gene is carried on a mobile genetic element, the staphylococcal 38 cassette chromosome (SCCmec) (3). SCCmec elements are classified into several types, 39 including I, II, and III, which are primarily hospital-associated clones, and types IV and V often 40 41 identified in community-associated MRSA (3).

PBPs are enzymes that carry out the final stages of assembly of bacterial cell wall 42 peptidoglycan (PG). Cell wall PG is essential for viability of most bacteria and forms a single 43 macromolecule around the cell (the sacculus), made of glycan strands and cross-linked via 44 peptide side-chains (5). High resolution Atomic Force Microscopy (AFM) has recently 45 revealed S. aureus PG to be a porous, heterogeneous hydrogel (6). Its mature surface is an 46 open, disordered mesh with pores that penetrate deep into the wall, whereas the inner surface, 47 where PG is synthesised, is a much denser mesh (6). Another feature of the PG is an outer 48 architecture of concentric rings consisting of long glycan strands that is revealed upon cell 49 scission and is characteristic of the newly exposed septum (6). 50

51 *S. aureus* has four endogenous PBPs of which only PBP1 and 2 are essential for PG synthesis, 52 being able to carry out all the transpeptidase (linking side-chains) functions necessary for cell 53 growth and division (7, 8, 9). PBP1 has multiple roles in cell division, by acting as a 54 coordinator, through interactions with PG and divisome protein partners, and by providing the 55 transpeptidase activity that is thought to be required for the characteristic ring architecture in 56 septal PG (6, 7, 8).

PBP2a is a non-native enzyme in MRSA, acquired from an environmental source, so how it 57 facilitates high-level antibiotic resistance by replacing the transpeptidase activity of 58 endogenous PBPs is intriguing. PBP2a requires the transglycosylase activity of PBP2 to 59 mediate resistance and the two proteins interact, thus demonstrating their functional 60 cooperativity (10). PBP2a can maintain transpeptidase activity with a closed active site 61 62 conformation, thus resisting β -lactam binding while interaction with a second PG substrate molecule at an allosteric site leads to a conformational change that opens the active site for 63 64 catalysis (11).

An interesting feature of many clinical MRSA isolates is that they exhibit heterogeneous 65 resistance, whereby only a very small proportion ($<10^{-4}$) of the population are high-level 66 resistant (>50 μ g ml⁻¹ methicillin) (12). Antibiotics can induce the conversion of the population 67 to homogeneous high-level resistance, that does not revert in the absence of antibiotics. 68 Chromosomal mutations that lead to the conversion to homogeneous resistance, mostly map to 69 genes responsible for the regulation of aspects of cellular physiology and not PBP2a function 70 directly (13). We have named these genes "potentiators" (pot), to differentiate them from 71 72 auxiliary genes (aux), in which mutation leads to decreased resistance (13). We have recently 73 carried out a directed evolution study that provides matched strains enabling the exploration of MRSA resistance mechanisms (14). Development of high-level MRSA is a two-step process 74 75 whereby the presence of mecA is essential but in itself only results in a modest increase in minimum inhibitory concentration (MIC) (low-level MRSA). Acquisition of missense 76 mutations in genes encoding RNA polymerase subunits (rpoB or rpoC), so-called rpo* 77 mutations, potentiate a step-change in resistance levels (high-level MRSA), both in the clinical 78 environment and under laboratory conditions (13, 14). 79

80 Cell wall architecture of MRSA

AFM was used to analyse the nanoscale, PG architecture, where in all cases at least 20 81 82 individual sacculi (i.e. purified cell wall fragments) were examined (see Materials and Methods). AFM analysis (Fig. 1A-B; and fig. S1A-B and S2A-B) showed that low-level 83 resistant MRSA (SH1000 mec A^+ (hereafter designated mec A^+); MIC 2 µg ml⁻¹), in the absence 84 of methicillin, resembled its sensitive parent (SH1000; MIC 0.25 µg ml⁻¹). In both cases, the 85 inner surface of the cell wall in all areas consisted of a dense mesh of PG, the outer surface of 86 the septum, newly exposed after division, exhibited the characteristic septal PG concentric-87 ring architecture, and the PG at the outer surface of the cell, away from the most recent site of 88

division, consisted of an open mesh structure (6) (Fig. 1A-B; and fig. S1A-B, S2A-B). We 89 90 quantified the orientation of individual glycan strands for strains SH1000 and $mecA^+$ in the absence of antibiotic using a custom-made automated image analysis. This revealed that in 91 both cases the outer surface of the septum exhibited a prominent peak in the circumferential 92 direction that is consistent with the concentric-ring architecture (Fig. 1Aiii, Biii). However, no 93 PG concentric rings were apparent at the outer surface of the septum of $mecA^+$ in the presence 94 95 of 1.5 µg ml⁻¹ methicillin (sub-MIC for $mecA^+$). Rather, the outer surface of the septum appeared as a dense mesh structure (Fig. 1Di-iii), while the inner surface displayed a large 96 proportion of long glycan strands that were oriented near the septal centre (fig. S1Dii, see the 97 98 long orange-brown coloured-fibres in fig. S1Diii). Furthermore, the cell wall was thinner after treatment with methicillin (fig. S1F). Under the same conditions (1.5 µg ml⁻¹ methicillin), the 99 parental stain, SH1000, died and cell wall spanning holes were apparent (15) (fig. S2F-H). The 100 cell wall architecture of the high-level MRSA strain (SH1000 mecA+ rpoB* (hereafter 101 designated $mecA^+$ $rpoB^*$); MIC $\geq 256 \ \mu g \ ml^{-1}$), which possessed both mecA and the pot 102 mutation, *rpoB*^{*} coding for a variant of the RNA polymerase β subunit RpoB(H929Q) (14), 103 resembled that of the parental strain $(mecA^+)$ in the absence of antibiotics (Fig. 1C; and fig. 104 S1C and fig. S2C). When treated with 25 µg ml⁻¹ methicillin (sub-MIC for this strain but 105 sufficient to kill both SH1000 and $mecA^+$) the inner surface of the cell wall maintained a dense 106 network of PG mesh, without the appearance of perforating holes (fig. S1Eii). However, in the 107 108 large majority of cases, the septa were thickened with a distinct protuberance, or lump, at the centre (fig. S1Ei). Importantly, although $mecA^+$ $rpoB^*$ was able to grow and divide in the 109 presence of methicillin, there was a total absence of the PG concentric-ring structure on the 110 outer surface of newly divided cells (Fig. 1E). Septal PG concentric rings are a defining feature 111 of PG architecture in several gram-positive bacteria (6, 16). Instead of PG concentric rings, the 112 outer surface of septa obtained from methicillin-treated $mecA^+ rpoB^*$ consisted of a disordered, 113 dense mesh with small pore size (Fig. 1E). As in the absence of antibiotics, the outer surface 114 of the rest of the cell periphery appeared as a more open mesh with larger pore size (fig. S2E 115 and S2I). This open mesh structure is derived from the dense mesh rather than from the 116 concentric ring structure, which remodels as cells divide in different planes during subsequent 117 division rounds (16). An interpretative diagram illustrating these observations is shown in Fig. 118 1F. 119

We then used the clinical, high-level, MRSA strain COL (SCCmec Type I), which possesses 120 both the mecA gene and produces a variant RpoB(A798V, S875L) (14) (MIC >256 µg ml⁻¹) to 121 determine whether the resistance-associated PG architectural changes described above 122 123 (absence of septal PG concentric rings, retention of PG dense mesh without perforating holes) are a common feature of MRSA cells under antibiotic stress. The COL cells were smaller than 124 SH1000 (average cell volume 0.69 ± 0.14 vs $1.22 \pm 0.31 \,\mu\text{m}^3$) as were the cells of mecA⁺ rpoB^{*} 125 (average cell volume $0.60 \pm 0.20 \ \mu\text{m}^3$, fig. S4D). Without antibiotics, COL displayed septal 126 PG concentric rings (fig. S3A), whereas in the presence of 25 µg ml⁻¹ methicillin (sub-MIC), 127 the septal PG of COL exhibited no concentric rings, but rather a disordered, dense mesh, at the 128 septal outer surface (fig. S3G). Treatment of $mecA^+$, $mecA^+$ $rpoB^*$ and COL with sub-MIC 129 concentrations of antibiotics (1.5, 25, and 25 µg ml⁻¹, respectively) led to high levels of PG 130 synthesis at the septum (as observed by ADA-DA incorporation), an increase in cell volume 131 132 and septal abnormalities observed by fluorescence microscopy and transmission electron microscopy (TEM) (fig. S4). 133

134 To demonstrate the wider applicability of our findings we then analyzed representatives of 135 different MRSA lineages and SCC*mec* types (*SCCmec* II (Mu50, MRSA252), III (TW20) and 136 IV (USA300, EMRSA15)) (3, 17-20). All strains had methicillin MICs of >256 μ g ml⁻¹ apart

from EMRSA15 and USA300 (MIC 64 and 1-2 μ g ml⁻¹, respectively) (Table S1). High-level

138 MRSA derivatives (MIC >256 μ g ml⁻¹), of the latter two strains, designated USA300 (HL) and 139 EMRSA15 (HL), were selected by directed evolution on oxacillin gradient plates (see 140 Materials and Methods).

AFM analysis of the clinical strains and high-level resistant derivatives was carried out in the 141 absence and presence of 25 µg ml⁻¹ methicillin (sub-MIC; fig. S3). All untreated strains had 142 septal PG concentric rings at the outer face of the septum (fig. S3). In the presence of 25 µg 143 ml⁻¹ methicillin (sub-MIC), the septal PG of COL, EMRSA15 (HL) and USA300 (HL) had a 144 disordered, dense mesh, at the septal outer surface but Mu50, MRSA252 and TW20 had 145 occasional (10 - 30% of septa) residual PG orientation. Growth of Mu50, MRSA252, and 146 TW20 in 50 µg ml⁻¹ methicillin (sub-MIC) gave rise to disordered mesh at the septal outer 147 surface (fig. S3). Thus, similar adaptations in septal PG architecture in response to antibiotic 148 challenge are conserved across MRSA strains (Fig. 1F). 149

Thus, even though PBP2a, in MRSA backgrounds permits growth and division in the presence of antibiotics, it leads to profound changes to cell wall architecture. This raises the questions as to how PBP2a complements the loss of both essential PBP1 and PBP2 transpeptidase activities, and also how high-level MRSA is able to divide?

154 Mode of cell division underpins high-level MRSA

We have recently suggested that the S. aureus septal PG concentric rings are due to PBP1 155 156 transpeptidase activity (8). Methicillin sensitive S. aureus (MSSA) specifically lacking PBP1 transpeptidase activity is not viable and exhibits aberrant septa (8). However, a high-level 157 MRSA strain with the same site-directed inactivation of PBP1 transpeptidase activity can grow 158 (8), suggesting that PBP2a complements the lack of PBP1 activity, but perhaps without the 159 ability to construct the septal PG concentric-ring structures. We therefore constructed a set of 160 otherwise isogenic strains where, in the absence of the inducer IPTG, only PBP1 without 161 transpeptidase activity (PBP1*) was expressed (Fig. 2A; and fig. S5A and B). Wholly 162 unexpectedly, the presence of PBP2a in this background SH1000 Pspac-pbp1 pbp1* mecA⁺ 163 (hereafter designated $pbp1^* mecA^+$) did not complement the loss of PBP1 transpeptidase 164 activity, demonstrating that PBP2a cannot substitute for the essential transpeptidase function 165 of PBP1 (Fig. 2B). Conversely, a single point mutation in rpoB (resulting in amino acid 166 replacement H929Q; rpoB*), that is required for MRSA with high-level resistance (14), was 167 able to entirely restore the ability of PBP1* to grow in the absence of PBP2a (Fig. 2B; and fig. 168 S5C and D). Growth of P_{spac}-pbp1 pbp1* rpoB* (hereafter designated pbp1* rpoB*) without 169 170 IPTG was associated with septal abnormalities, an increase in cell volume, and alterations to PG synthesis (Fig. 2C and D; and fig. S5E-F and S6A), similar to high-level MRSA grown in 171 the presence of antibiotics (fig. S4B). 172

AFM analysis of the PG architecture of *pbp1* rpoB** with IPTG (PBP1 transpeptidase activity 173 present) revealed open mesh on outer surfaces and septal PG concentric rings as expected for 174 a wild type strain (Fig. 2E (+IPTG); Fig. S6B to D). However, growth without IPTG (no PBP1 175 transpeptidase activity) led to the concentric rings at the septal surface being replaced by a 176 disordered, dense mesh with random glycan strand orientation (Fig. 2E (-IPTG); and fig. S6E 177 to G). Although rpoB* complemented the absence of PBP1 transpeptidase activity, neither 178 rpoB* nor PBP2a, or both combined, could rescue cells lacking the PBP1 protein (fig. S7), 179 180 consistent with the physical presence of PBP1 being necessary for cell division complex assembly. Therefore, the septal PG ring architecture associated with conventional cell division 181 requires the essential transpeptidase activity of PBP1, but S. aureus can adopt an alternative 182 division mode facilitated by rpoB* when PBP1 transpeptidase activity is lost (either by 183 mutation or antibiotic addition; Fig. 1F). This fundamentally different mode of cell division, 184

which lacks the canonical septal PG concentric-ring architecture, is exploited in high-level MRSA, where $rpoB^*$ in combination with *mecA* allows division in the presence of antibiotics.

187 Dual mechanisms for high-level MRSA

High-level MRSA requires two factors; the presence of PBP2a and a potentiator (pot) mutation 188 (as provided by *rpoB**) (13, 14). For high-level MRSA to grow and divide in the presence of 189 β-lactam antibiotics, the essentiality of PBP1 and PBP2 transpeptidase activities must be 190 191 circumvented or enzymatically complemented. Previous studies report that in strain COL the transpeptidase activity of PBP2 can be complemented by the presence of PBP2a (21, 22). 192 However, growth of a COL derivative lacking PBP2 protein is impaired and does not exhibit 193 antibiotic resistance (22). This is because PBP2 transglycosylase activity is required to act 194 cooperatively with PBP2a (22). COL also harbours potentiator rpoB* mutations (A798V, 195 S875L) required for high-level resistance (14). To determine whether there are two co-196 dependent mechanisms that in combination lead to high-level MRSA we investigated the effect 197 of *pbp2* mutations. As expected from previous reports (21, 22) PBP2 is essential and PBP2a 198 and/or rpoB* (H929O) could not compensate for the loss of PBP2 protein in terms of plating 199 200 efficiency and growth (fig. S8A-C). When PBP2 was depleted, with or without the presence of PBP2a, S. aureus stopped dividing, exhibiting decreased septal PG incorporation and altered 201 septal morphology (fig. S9). Loss of PBP2 also led to a decrease in cell size (fig. S9H). 202 Depletion of PBP2 in $rpoB^*$ or $mecA^+$ $rpoB^*$ led to lower growth, decreased septal PG 203 incorporation, altered septal morphology, and death (fig. S9C, D, and G). We could not create 204 PBP2* (transpeptidase mutant) strains in either the parental SH1000 or rpoB* backgrounds, 205 indicating its essentiality. However, strains where only PBP2* is present were viable in both 206 $mecA^+$ and $mecA^+$ $rpoB^*$ (Fig. 3A to D). Both the PBP2 and PBP2* constructs were verified 207 by Western blot and Bocillin labelling (fig. S8D and E). Both strains with PBP2* were able to 208 grow with near parental ($mecA^+$ and $mecA^+$ $rpoB^*$, respectively) cell morphology (fig. S9E 209 and F). All PBP2 and PBP2* constructs demonstrate a diminished cell size compared to 210 SH1000 (fig. S9H). Expression of PBP2* (lacking PBP2 transpeptidase activity) in the mecA⁺ 211 or $mecA^+$ rpoB* backgrounds resulted in septa that exhibited the typical PG concentric-ring 212 architecture, with strands preferentially oriented in the circumferential direction (Fig. 3E and 213 F and fig. S10). We conclude that neither PBP2 nor PBP2a are responsible, even in part, for 214 the PG septal concentric rings associated with conventional cell division. Therefore, there are 215 two factors required for high-level MRSA: (i) PBP2a replaces the essential transpeptidase 216 activity of PBP2, and (ii) a pot mutation (e.g. rpoB*) permits cell division without PBP1 217 transpeptidase activity. 218

219 Potentiator mutations converge on nucleotide signalling

Mutations in *rpoB* and *rpoC* have been associated with clinically important high-level MRSA strains and the conversion from hetero- to homogeneous resistance (13, 14, 23, 24). Other *pot* mutations, such as *rel*, *clpXP*, *gdpP*, *pde2* and *lytH* have been uncovered in laboratory studies and in some cases clinically (13). Whilst other mutations enhanced the MIC of *mecA*⁺, tested in our defined SH1000 background with a single copy of *mecA* in the chromosome, only *rpoB* and *rel* led to high-level resistance (table S1; MIC \geq 256 µg ml⁻¹).

The *rel* gene encodes a key component of the stringent response (25) and whilst the gene is conditionally essential, the *pot* mutant strain (*rel**) has a C-terminal truncation in the regulatory domain of the Rel protein, and likely increases (p)ppGpp levels (26). The stringent response has been previously implicated as having a major role in potentiating high-level MRSA (27) and here we found the presence of $mecA^+$ $rpoB^*$ led to a significant increase in the levels of the stringent response signalling molecules ppGpp and pppGpp (Fig. 4A). To determine the relationship between the stringent response and the dual pathways to high-level MRSA we
investigated its ability to compensate for the loss of PBP1 transpeptidase activity (Fig. 4B).
The *rel** mutation was as effective as *rpoB** in compensating for the absence of PBP1
transpeptidase activity as judged by measurement of plating efficiency (Fig. 4B), implicating
the stringent response in the ability to grow and divide without septal PG concentric rings.

237 Therapeutic development for MRSA

238 To counter the emergence of MRSA, compounds have been identified that resensitize these strains to β -lactams (28). These include clomiphene (29) and norgestimate (30), as well as 239 natural products including epicatechin gallate (ECg) (31) and spermine (32). Their mode of 240 action is mostly unknown and so we tested their effect, at concentrations that resensitize mecA⁺ 241 rpoB* and the other clinical MRSA strains to oxacillin but do not inhibit growth without 242 antibiotic (see Materials and Methods) (Fig. 4C to E). Clomiphene and spermine did not inhibit 243 the plating efficiency of pbp1*rpoB* but did for both $mecA^+ pbp2*$ and $mecA^+ pbp2*rpoB*$, 244 suggesting a link to the activity of PBP2a. Norgestimate impaired the plating efficiency of 245 pbp1*rpoB* and $mecA^+$ pbp2* but not $mecA^+$ pbp2*rpoB*, demonstrating a potential cross-246 247 talk between the co-dependent pathways (i.e., acquisition of mecA and a pot mutation) that lead to resistance. ECg inhibited the plating efficiency of all three strains indicating that it may 248 affect an Aux factor required under all conditions. These observations further differentiate the 249 250 two resistance pathways and provide specific interventions able to dissect the new mode of cell division uncovered here. 251

252 **Discussion**

We have revealed that the high-level resistance to β -lactam antibiotics exhibited by some 253 MRSA strains is linked to an alternative mode of cell division set within the context of wider 254 physiological adaptations (i.e., increased ppGpp and pppGpp) (Fig. 4F). The development of 255 high-level MRSA is a two-step process in which PBP2a compensates for the lack of native 256 257 PBP2 transpeptidase activity in the presence of β -lactam antibiotics (22). PBP2 is an essential enzyme that is required for the synthesis of the dense mesh PG on the inside of the cell wall at 258 both the septum and the cell periphery. It is therefore the major PBP in terms of bulk PG 259 synthesis. PBP2a cannot compensate for the lack of PBP2 protein (specifically its 260 transglycosylase activity (22)). However, as PBPs can form dimers (33), PBP2/2a heterodimers 261 could allow both the multiple protein interactions of PBP2 (34) and PBP2a transpeptidase 262 activity required for PG synthesis. PBP1 has essential transpeptidase activity and operates with 263 its cognate transglycosylase FtsW (35). Here we show that PBP1 activity is responsible for the 264 formation of the concentric rings that are characteristic of septal PG. PBP2a cannot compensate 265 for the lack of PBP1 activity but pot mutations can. The pot mutations permit successful cell 266 division without septal PG rings in the presence of high levels of antibiotics (Fig. 4F). This 267 compensatory mechanism does not involve a replacement of PBP1 activity but rather 268 physiological adaptations that allow division without it. A question arises as to whether the 269 ability to divide without septal PG concentric rings in high-level MRSA strains evolved 270 specifically, in the context of antibiotic use, or whether it is part of a wider physiological 271 272 capability that is deployed under stressful conditions? Mutations in rpo genes are often found associated with antibiotic and stress resistance in S. aureus and many other organisms (24, 36, 273 37). A survey of 1,429 MRSA (ST22) clinical strains revealed that $\sim 10\%$ had at least one point 274 mutation in genes coding for core RNAP subunits or σ factors (24). The current study now 275 links these mutations to the widely conserved stringent response, which is a key component in 276 bacterial responses to stress and growth perturbations (25). Our rpoB* strains exhibit lower 277

- 278 growth rates compared to parental strains (14), which could, at least in-part, facilitate the 279 alternative mode of division.
- Given the array of MRSA SCC*mec* types and clonal lineages, it is likely that the effects of *pot* factors, such as rpo^* , are influenced by the genetic background (13). This provides both complexity in unravelling the interplay between *pot* and *aux* factors but also an opportunity to
- establish those common, underlying principles that underpin resistance. The resensitizing
- agents also provide avenues to probe underlying molecular mechanisms. Our study has
- revealed insights into antibiotic resistance and facets of cell division in S. aureus. It is by
- studying these processes in tandem that we can understand basic mechanisms of the bacterial
- cell cycle and reveal ways to control antibiotic resistance.

288 References

- B. Ribeiro da Cunha, L. P. Fonseca, C. R. C. Calado, Antibiotic discovery: Where have we come from, where do we go? *Antibiot.* 8, 1-21 (2019).
- Antimicrobial Resistance Collaborators, Global burden of bacterial antimicrobial resistance in
 2019: a systematic analysis. *Lancet* 399, 629–655 (2022).
- S. Lakhundi, K. Zhang, Methicillin-resistant *Staphylococcus aureus*: Molecular characterization, evolution, and epidemiology. *Clin. Microbiol. Rev.* 31:10.1128/cmr.00020-18 (2018).
- 4. B. J. Hartman, A. Tomasz, Low-affinity penicillin-binding protein associated with beta-lactam resistance in *Staphylococcus aureus*. J. Bacteriol. 158, 513–516 (1984).
- W. Vollmer, D. Blanot, M. A. De Pedro, Peptidoglycan structure and architecture. *FEMS Microbiology Reviews* 32, 149–167 (2008).
- L. Pasquina-Lemonche, J. Burns, R. D. Turner, S. Kumar, R. Tank, N. Mullin, J. S. Wilson, B.
 Chakrabarti, P. A. Bullough, S. J. Foster, J. K. Hobbs, The architecture of the Gram-positive bacterial cell wall. *Nature* 582, 294–297 (2020).
- 303 7. S. F. Pereira, A. O. Henriques, M. G. Pinho, H. de Lencastre, A. Tomasz, Role of PBP1 in cell division of *Staphylococcus aureus*. J. Bacteriol. 189, 3525–3531 (2007).
- K. Wacnik, V. A. Rao, X. Chen, L. Lafage, M. Pazos, S. Booth, W. Vollmer, J. K. Hobbs, R.
 J. Lewis, S. J. Foster, Penicillin-Binding Protein 1 (PBP1) of *Staphylococcus aureus* has multiple essential functions in cell division. *mBio.* 13, e00669 (2022).
- M. G. Pinho, S. R. Filipe, H. de Lencastre, A. Tomasz, Complementation of the essential peptidoglycan transpeptidase function of penicillin-binding protein 2 (PBP2) by the drug resistance protein PBP2A in *Staphylococcus aureus*. J. Bacteriol. 183, 6525–6531 (2001).
- B. García-Fernández, G. Koch, R. M. Wagner, A. Fekete, S. T. Stengel, J. Schneider, B. Mielich-Süss, S. Geibel, S. M. Markert, C. Stigloher, D. Lopez, Membrane microdomain disassembly inhibits MRSA antibiotic resistance. *Cell* 171, 1354-1367.e20 (2017).
- L. H. Otero, A. Rojas-Altuve, L. I. Llarrull, C. Carrasco-López, M. Kumarasiri, E. Lastochkin,
 J. Fishovitz, M. Dawley, D. Hesek, M. Lee, J. W. Johnson, J. F. Fisher, M. Chang, S.
 Mobashery, J. A. Hermoso, How allosteric control of *Staphylococcus aureus* penicillin binding
 protein 2a enables methicillin resistance and physiological function. *PNAS* 110, 16808-16813
 (2013).
- B. J. Hartman and, A. Tomasz, Expression of methicillin resistance in heterogeneous strains of
 Staphylococcus aureus. Antimicrob. Agents Chemother. 29, 85–92 (1986).
- B. L. Bilyk, V. V. Panchal, M. Tinajero-Trejo, J. K. Hobbs, S. J. Foster, An Interplay of multiple positive and negative factors governs methicillin resistance in *Staphylococcus aureus*. *Microbiol. Mol. Biol. Rev.* 86, e00159 (2022).
- 14. V. V. Panchal, C. Griffiths, H. Mosaei, B. Bilyk, J. A. F. Sutton, O. T. Carnell, D. P. Hornby,
 J. Green, J. K. Hobbs, W. L. Kelley, N. Zenkin, S. J. Foster, Evolving MRSA: High-level βlactam resistance in *Staphylococcus aureus* is associated with RNA Polymerase alterations and
 fine tuning of gene expression. *PLOS Pathog.* 16, e1008672 (2020).
- B. Salamaga, L. Kong, L. Pasquina-Lemonche, L. Lafage, M. von Und Zur Muhlen, J. F.
 Gibson, D. Grybchuk, A. K. Tooke, V. Panchal, E. J. Culp, E. Tatham, M. E. O'Kane, T. E.
 Catley, S. A. Renshaw, G. D. Wright, P. Plevka, P. A. Bullough, A. Han, J. K. Hobbs, S. J.
 Foster, Demonstration of the role of cell wall homeostasis in *Staphylococcus aureus* growth
 and the action of bactericidal antibiotics. *PNAS* 118, 1-8 (2021).

- R. D. Turner, E. C. Ratcliffe, R. Wheeler, R. Golestanian, J. K. Hobbs, S. J. Foster,
 Peptidoglycan architecture can specify division planes in *Staphylococcus aureus*. *Nat. Commun.* 1, 26 (2010).
- S. R. Harris, E. J. Feil, M. T. G. Holden, M. A. Quail, E. K. Nickerson, N. Chantratita, S. Gardete, A. Tavares, N. Day, J. A. Lindsay, J. D. Edgeworth, H. De Lencastre, J. Parkhill, S. J. Peacock, S. D. Bentley, Evolution of MRSA during hospital transmission and intercontinental spread. *Science* 327, 469–474 (2010).
- 340 18. M. T. G. Holden, E. J. Feil, J. A. Lindsay, S. J. Peacock, N. P. J. Day, M. C. Enright, T. J. Foster, C. E. Moore, L. Hurst, R. Atkin, A. Barron, N. Bason, S. D. Bentley, C. Chillingworth, 341 T. Chillingworth, C. Churcher, L. Clark, C. Corton, A. Cronin, J. Doggett, L. Dowd, T. 342 343 Feltwell, Z. Hance, B. Harris, H. Hauser, S. Holroyd, K. Jagels, K. D. James, N. Lennard, A. Line, R. Mayes, S. Moule, K. Mungall, D. Ormond, M. A. Quail, E. Rabbinowitsch, K. 344 Rutherford, M. Sanders, S. Sharp, M. Simmonds, K. Stevens, S. Whitehead, B. G. Barrell, B. 345 346 G. Spratt, J. Parkhill, Complete genomes of two clinical Staphylococcus aureus strains: Evidence for the evolution of virulence and drug resistance. PNAS 101, 9786–9791 (2004). 347
- T. Takano, W. C. Hung, M. Shibuya, W. Higuchi, Y. Iwao, A. Nishiyama, I. Reva, O. E.
 Khokhlova, S. Yabe, K. Ozaki, M. Takano, T. Yamamoto, A new local variant (ST764) of the
 globally disseminated ST5 lineage of hospital-associated methicillin-resistant *Staphylococcus aureus* (MRSA) carrying the virulence determinants of community-associated MRSA.
 Antimicrob. Agents Chemother. 57, 1589–1595 (2013).
- B. A. Diep, S. R. Gill, R. F. Chang, T. H. Van Phan, J. H. Chen, M. G. Davidson, F. Lin, J. Lin,
 H. A. Carleton, E. F. Mongodin, G. F. Sensabaugh, F. Perdreau-Remington, Complete genome
 sequence of USA300, an epidemic clone of community-acquired meticillin-resistant *Staphylococcus aureus. Lancet* 367, 731–739 (2006).
- M. G. Pinho, S. R. Filipe, H. De Lencastre, A. Tomasz, Complementation of the essential peptidoglycan transpeptidase function of penicillin-binding protein 2 (PBP2) by the drug resistance protein PBP2A in *Staphylococcus aureus*. J. Bacteriol. 183, 6525–6531 (2001).
- 360 22. M. G. Pinho, H. de Lencastre, A. Tomasz, An acquired and a native penicillin-binding protein
 361 cooperate in building the cell wall of drug-resistant staphylococci. *PNAS* 98, 10886-10891
 362 (2001).
- 363 23. Y. Aiba, Y. Katayama, T. Hishinuma, H. Murakami-Kuroda, L. Cui, K. Hiramatsu, Mutation
 364 of RNA polymerase β-subunit gene promotes heterogeneous-to-homogeneous conversion of β365 lactam resistance in methicillin-resistant *Staphylococcus aureus*. *Antimicrob. Agents*366 *Chemother.* 57, 4861–4871 (2013).
- A. Krishna, B. Liu, S. J. Peacock, S. Wigneshweraraj, The prevalence and implications of single
 nucleotide polymorphisms in genes encoding the RNA polymerase of clinical isolates of
 Staphylococcus aureus. MicrobiologyOpen 9, e1058 (2020).
- S. E. Irving, N. R. Choudhury, R. M. Corrigan, The stringent response and physiological roles of (pp)pGpp in bacteria. *Nat. Rev. Microbiol.* 19, 256–271 (2021).
- A. T. Deventer, D. Bryson, M. Shortill, A. B. Boraston, J. K. Hobbs, Molecular characterization
 of clinical rel mutations and consequences for resistance expression and fitness in *Staphylococcus aureus. Antimicrob. Agents Chemother.* 66, e00938 (2022).
- 27. C. Kim, M. Mwangi, M. Chung, C. Milheirico, H. de Lencastre, A. Tomasz, The mechanism
 of heterogeneous beta-lactam resistance in MRSA: key role of the stringent stress response. *PLoS One* 8, e82814 (2013).
- 378 28. T. J. Foster, Can beta-Lactam antibiotics be resurrected to combat MRSA? *Trends Microbiol.*379 27, 26–38 (2019).

- M. A. Farha, T. L. Czarny, C. L. Myers, L. J. Worrall, S. French, D. G. Conrady, Y. Wang, E.
 Oldfield, N. C. Strynadka, E. D. Brown, Antagonism screen for inhibitors of bacterial cell wall
 biogenesis uncovers an inhibitor of undecaprenyl diphosphate synthase. *PNAS* 112, 11048–
 11053 (2015).
- 30. Y. Yoshii, K. I. Okuda, S. Yamada, M. Nagakura, S. Sugimoto, T. Nagano, T. Okabe, H.
 Kojima, T. Iwamoto, K. Kuwano, Y. Mizunoe, Norgestimate inhibits staphylococcal biofilm
 formation and resensitizes methicillin-resistant *Staphylococcus aureus* to β-lactam antibiotics.
 npj Biofilms Microbiomes 3, 18 (2017).
- 388 31. P. D. Stapleton, S. Shah, K. Ehlert, Y. Hara, P. W. Taylor, The β-lactam-resistance modifier
 (-)-epicatechin gallate alters the architecture of the cell wall of *Staphylococcus aureus*.
 390 *Microbiology* 153, 2093–2103 (2007).
- 391 32. D. H. Kwon, C. D. Lu, Polyamine effects on antibiotic susceptibility in bacteria. *Antimicrob.* 392 *Agents Chemother.* 51, 2070–2077 (2007).
- 393 33. J. F. Fisher, S. Mobashery, β-Lactams against the Fortress of the Gram-Positive *Staphylococcus* 394 *aureus* Bacterium. *Chem. Rev.* 121, 3412–3463 (2021).
- 34. V. R. Steele, A. L. Bottomley, J. Garcia-Lara, J. Kasturiarachchi, S. J. Foster, Multiple essential
 roles for EzrA in cell division of *Staphylococcus aureus*. *Mol. Microbiol.* 80, 542–555 (2011).
- 35. N. T. Reichmann, A. C. Tavares, B. M. Saraiva, A. Jousselin, P. Reed, A. R. Pereira, J. M.
 Monteiro, R. G. Sobral, M. S. VanNieuwenhze, F. Fernandes, M. G. Pinho, SEDS-bPBP pairs
 direct lateral and septal peptidoglycan synthesis in *Staphylococcus aureus*. *Nat. Microbiol.* 4, 1368–1377 (2019).
- 401 36. L. Ostrer, Y. Ji, A. Khodursky, Identification and characterization of pleiotropic high402 persistence mutations in the beta subunit of the bacterial RNA polymerase. *Antimicrob. Agents*403 *Chemother.* 65, 00522 (2021).
- J. A. Leeds, M. Sachdeva, S. Mullin, S. W. Barnes, A. Ruzin, In vitro selection, via serial passage, of *Clostridium difficile* mutants with reduced susceptibility to fidaxomicin or vancomycin. *J. Antimicrob. Chemother.* 69, 41–44 (2014).
- 407 38. https://doi.org/10.15131/shef.data.26341735
- 408 39. R. M. Corrigan, L. Bowman, A. R. Willis, V. Kaever, A. Grundling, Cross-talk between two nucleotide-signaling pathways in *Staphylococcus aureus*. J. Biol Chem. 290, 5826–5839
 410 (2015).
- 40. R. P. Novick, S. I. Morse, In vivo transmission of drug resistance factors between strains of *Staphylococcus aureus*. J. Exp. Med. 125, 45–59 (1967).
- 41. S. Schenk, R. A. Laddaga, Improved method for electroporation of *Staphylococcus aureus*.
 414 *FEMS Microbiol. Lett.* 94, 133–138 (1992).
- 415 42. J. Schindelin, I. Arganda-Carreras, E. Frise, V. Kaynig, M. Longair, T. Pietzsch, S. Preibisch,
 416 C. Rueden, S. Saalfeld, B. Schmid, J. Y. Tinevez, D. J. White, V. Hartenstein, K. Eliceiri, P.
 417 Tomancak, A. Cardona, Fiji: an open-source platform for biological-image analysis. *Nat.*418 *Methods* 9, 676–682 (2012).
- 419 43. J. G. Beton, R. Moorehead, L. Helfmann, R. Gray, B. W. Hoogenboom, A. P. Joseph, M. Topf,
 420 A. L. B. Pyne, TopoStats A program for automated tracing of biomolecules from AFM
 421 images. *Methods* 193, 68–79 (2021).
- 42. 44. T. Lindeberg, Edge Detection and Ridge Detection with Automatic Scale Selection. Int. J. Comput. Vis. 30, 117–156 (1998).
- 424 45. F. Meyer, Topographic distance and watershed lines. *Signal Processing* **38**, 113–125 (1994).

- 425 46. https://doi.org/10.5281/zenodo.13082796
- 426 47. https://doi.org/10.5281/zenodo.13133978
- 48. S. Berg, D. Kutra, T. Kroeger, C. N. Straehle, B. X. Kausler, C. Haubold, M. Schiegg, J. Ales,
 T. Beier, M. Rudy, K. Eren, J. I. Cervantes, B. Xu, F. Beuttenmueller, A. Wolny, C. Zhang, U.
 Koethe, F. A. Hamprecht, A. Kreshuk, ilastik: interactive machine learning for (bio)image
 analysis. *Nat. Methods* 16, 1226–1232 (2019).
- 431 49. https://doi.org/10.5281/zenodo.13134076
- M. J. Horsburgh, J. L. Aish, I. J. White, L. Shaw, J. K. Lithgow, S. J. Foster, δb modulates
 virulence determinant expression and stress resistance: Characterization of a functional rsbU
 strain derived from *Staphylococcus aureus* 8325-4. *J. Bacteriol.* 184, 5457–5467 (2002).
- 435 51. W. M. Shafer, J. J. Landolo, Genetics of staphylococcal enterotoxin B in methicillin-resistant isolates of *Staphylococcus aureus*. *Infect. Immun.* 25, 902–911 (1979).
- 437 52. B. N. Kreiswirth, S. Löfdahl, M. J. Betley, M. O'reilly, P. M. Schlievert, M. S. Bergdoll, R. P.
 438 Novick, The toxic shock syndrome exotoxin structural gene is not detectably transmitted by a
 439 prophage. *Nature* 305, 709–712 (1983).
- 53. C. Y. Lee, S. L. Buranen, Y. Zhi-Hai, Construction of single-copy integration vectors for *Staphylococcus aureus. Gene* 103, 101–105 (1991).
- 54. G. McVicker, T. K. Prajsnar, A. Williams, N. L. Wagner, M. Boots, S. A. Renshaw, S. J. Foster,
 Clonal expansion during *Staphylococcus aureus* infection dynamics reveals the effect of
 antibiotic intervention. *PLoS Pathog.* 10, e1003959 (2014).
- P. D. Fey, J. L. Endres, V. K. Yajjala, T. J. Widhelm, R. J. Boissy, J. L. Bose, K. W. Bayles, A
 genetic resource for rapid and comprehensive phenotype screening of nonessential *Staphylococcus aureus* genes. *mBio.* 4, 00537 (2013).
- 448 56. R. M. Corrigan, J. C. Abbott, H. Burhenne, V. Kaever, A. Grundling, c-di-AMP is a new second messenger in *Staphylococcus aureus* with a role in controlling cell size and envelope stress.
 450 *PLoS Pathog.* 7, e1002217 (2011).
- V. A. Lund, K. Wacnik, R. D. Turner, B. E. Cotterell, C. G. Walther, S. J. Fenn, F. Grein, A. J.
 M. Wollman, M. C. Leake, N. Olivier, A. Cadby, S. Mesnage, S. Jones, S. J. Foster, Molecular coordination of *Staphylococcus aureus* cell division. *eLife* 7, e32057 (2018).
- 454 58. M. Arnaud, A. Chastanet, M. Débarbouillé, New vector for efficient allelic replacement in naturally nontransformable, low-GC-content, gram-positive bacteria. *Appl. Environ.*456 *Microbiol.* 70, 6887–6891 (2004).

457 Acknowledgements

We are grateful to Nicholas Mullin, Xinyue Chen, Anaam Alomari, Viralkumar V. Panchal, 458 Bartek Salamaga, and Matthew J. Barker for helpful discussions, also Jodi Lindsay (St. 459 George's UCL) and James O'Gara (University of Galway) for provision of strains. Electron 460 Microscopy was carried out at the School of Biosciences Cryo-Electron Microscopy Facility, 461 University of Sheffield. Fluorescence microscopy was performed at the Wolfson Light 462 Microscopy Facility, University of Sheffield. The research was in part carried out at the 463 National Institute for Health and Care Research (NIHR) Sheffield Biomedical Research Centre 464 (BRC). For the purpose of open access, the authors have applied a CC BY public copyright 465 license to any author accepted manuscript version arising from this submission. Funding: This 466 work was supported by the Engineering and Physical Sciences Research Council (grant 467 EP/T002778/1 to J.K.H. and S.J.F), the Wellcome Trust (grants 212197/Z/19/Z to J.K.H and 468 S.J.F and 104110/Z/14/A to J.K.H., S.J.F., J.G. and R.M.C) and the Biotechnology and 469

Biological Sciences Research Council (grant BB/R018383/1 to WMD). Author 470 contributions: A.F.A.-O, and K.W designed the study, performed experiments, analysed, and 471 interpreted data, and wrote the manuscript. (A.F.A.-O.: Figs. 1, 2, 3, Supplementary Materials 472 Figs. S1-S3, S6 & S10; K.W.: Figs. 2-4, Supplementary Materials Figs. S4-S9). L.L., M.T.-T., 473 B.B., S.E.I., J.A.F.S and C.J.P.-R. performed the experiments, analysed, and interpreted the 474 data (L.L. Fig. 1, Supplementary Materials: Figs. S4 & S6; M.T.-T.: Fig. 4, Supplementary 475 Materials Figs. S4, S8, S10; S.E.I.: Fig. 4; J.A.F.S.: Supplementary Materials: Fig. S3 & Table 476 S1 and C.J.P.-R.: Supplementary Materials Fig. S10). L.P.-L. and D.S.O. developed the semi-477 automated macro used to calculate cell volume measurements and L.P.-L analysed the data 478 479 (Supplementary Materials Figs. S4, S5 & S9). Also, L.P.-L. developed the semi-automated macro used to calculate the cell wall pore area (Supplementary Materials Figs. S2, S6, S10). 480 S.A.R and E.B. performed the experiments and analyzed data (Supplementary Materials Fig. 481 482 S2). O.M. and W.M.D. developed Matlab-based software used for the fibre detection (Figs. 1-3). D.J.H., J.F., R.C. and J.G. analysed and interpreted data, and wrote and reviewed the 483 manuscript. J.K.H. and S.J.F. designed the study, interpreted the data, wrote the manuscript 484 and directed the project. Competing interests: The authors declare no competing interests. 485 486 Data and materials availability: The data that support the findings of this study are available in the Online Research Data (ORDA) figshare from the University of Sheffield with the 487 identifier (38). 488

489 Supplementary Materials

- 490 Materials and Methods
- 491 Figs. S1 to S10
- 492 Tables S1 and S4
- 493 References (39 58)

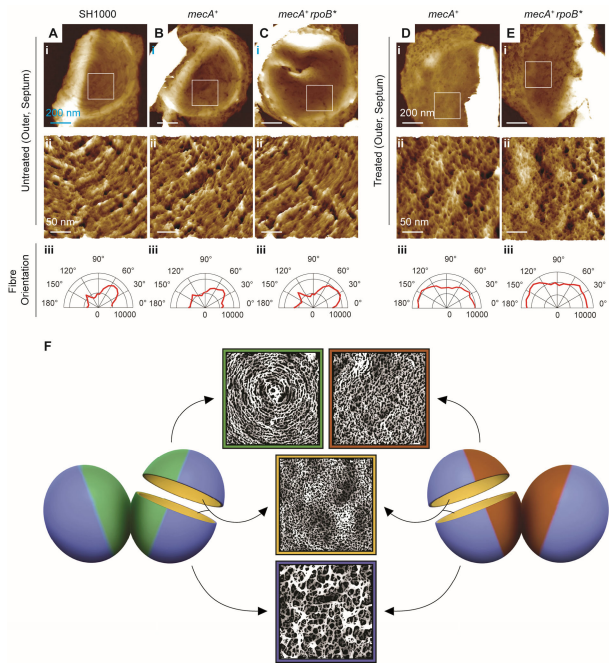
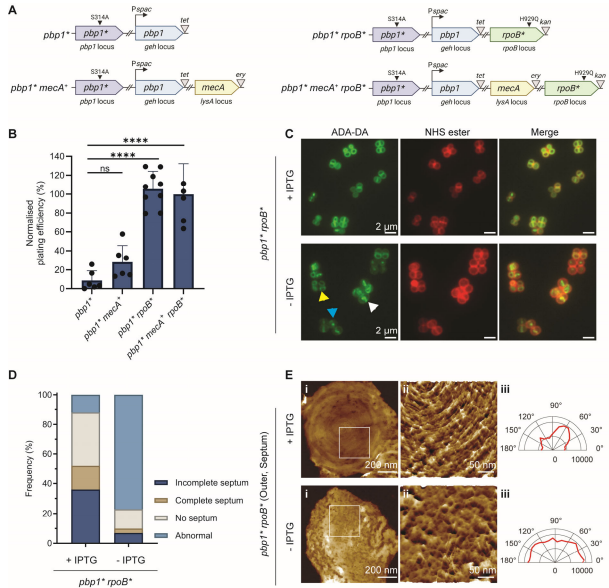


Fig. 1. Methicillin treatment of MRSA alters the architecture of the cell wall. From left to 495 right, (A-C) show the outer surfaces of newly revealed septa, in samples of isolated sacculi of 496 untreated (A) SH1000, (B) $mecA^+$, and (C) $mecA^+$ $rpoB^*$ respectively. (D-E) Show the outer 497 surface of the newly revealed septa of (**D**) $mecA^+$ and (**E**) $mecA^+$ rpoB* treated with methicillin 498 (1.5 and 25 μ g ml⁻¹ respectively). In all columns: (i) shows an individual fragment of sacculus 499 corresponding to the outer surface of the septum. Topographical height (z) range presented in 500 each of these images (from left to right) is 140, 140, 150, 120, and 185 nm. (ii) Shows pseudo-501 three dimensional (3D) high resolution AFM images of the sections indicated by the white 502 503 boxes in (i). Topographical height (z) range presented in each of these images (from left to right) is 7.5, 10, 7.5, 12, and 20 nm. (iii) Represents the combined angular histogram of fibre 504 orientation of AFM high-resolution images similar to those in (ii). The fibre orientation 505 analysis method used for the orientation detection is described in the Materials and Methods 506 section. (F) Shows an interpretative diagram of different architectures (concentric rings, dense 507 mesh, and open mesh) observed by high-resolution AFM on different surfaces (outer surface 508

of newly revealed septa, inner surface of the septa, and outer surface of cell periphery) of untreated (left-hand side) and antibiotic treated (right-hand side) MRSA cell wall. The green colour represents the concentric rings associated with the outer surface of the septum of untreated cells, blue colour shows the open mesh at the cell periphery, yellow colour depicts the dense mesh on the inner wall of the cell and lastly the brown colour represents the dense mesh on the outer surface of the septum after treatment with methicillin. The modified AFM images in Fig. 1F span 400 nm by 400 nm in x and y dimension.



516 Fig. 2. Loss of PBP1 transpeptidase activity can be compensated for by *rpoB** but not 517 mecA. (A) Representation of pbp1* genetic constructs. An ectopic pbp1 copy, at the lipase 518 (geh) locus is controlled by the Pspac promoter. The pbp1 gene at its native locus has a point 519 mutation (940T>G) resulting in inactivation of transpeptidase activity (S314A, *pbp1**). The 520 521 $mecA^+$ gene is expressed from its native promoter at the lysA locus. In rpoB*, a point mutation results in an amino acid substitution (H929Q) in RpoB. tet, erv and kan represent tetracycline, 522 erythromycin and kanamycin resistance cassettes, respectively. The graphics were created with 523 BioRender.com. (B) Plating efficiency of *pbp1**, *pbp1*mecA*⁺, *pbp1*rpoB** and *pbp1*mecA*⁺ 524 rpoB* without IPTG. Plating efficiencies were compared to controls grown with IPTG, using 525 a one-way ANOVA with Dunnett's multiple comparison test (ns, not significant; ****, 526 P < 0.0001). Error bars show mean \pm standard deviation (SD). (C) Fluorescence microscopy 527 images of *pbp1* rpoB** grown +/- IPTG for 4 h, labelled with ADA-DA (5 min) and then 528 NHS-ester Alexa Fluor 555 to image nascent PG and cell wall, respectively. Images are z stack 529 average intensity projections. Scale bars = $2 \mu m$. (**D**) Quantification of cellular phenotypes 530 based on ADA-DA incorporation in *pbp1* rpoB** incubated with IPTG (+) or without IPTG 531 (-), n = 511 and 654 (respectively). Examples of cells classified as abnormal with misshapen 532 septal rings (yellow arrowhead), accumulation of ADA-DA at septal centre, 'plug' (blue 533 arrowhead) and mislocalized ADA-DA incorporation (white arrowhead) are shown C. (E) 534

AFM images of newly exposed outer surface of the septum after cell division of pbp1*rpoB*grown +/- IPTG for 4 h, reveal lack of concentric-ring structures in -IPTG. (i) Representative outer septal surfaces with height (z) range of 120 nm and the HS applies to both. (ii) Shows pseudo-3D AFM high resolution images of the region within the white box in (i). Topographical height (z) range (top) = 9.5 nm, and HS (bottom) = 21 nm. (iii) Represents the combined angular histogram of fibre orientation of AFM high resolution images similar to those in (ii).

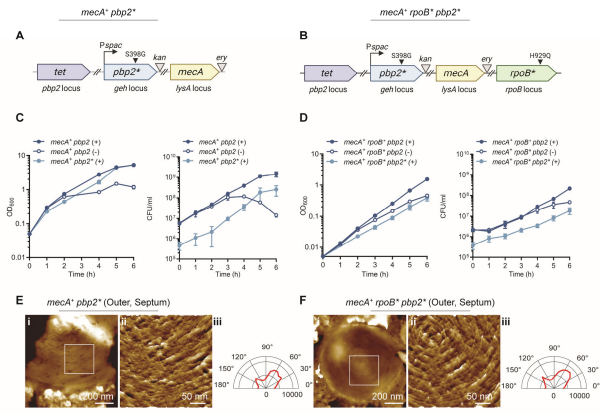


Fig. 3. Loss of PBP2 transpeptidase activity can be compensated for by mecA but not 543 *rpoB**. (A-B) Schematic representation of $mecA^+$ pbp2* (SJF5807) and $mecA^+$ rpoB* pbp2* 544 (SJF5809) genetic constructs. A copy of *pbp2* with an inactive transpeptidase domain (*pbp2**, 545 1191-1192TC>GG, S398G) was placed under the control of the Pspac promoter at the lipase 546 (geh) locus of SH1000, pbp2 at its native locus was then deleted (marked with tet). In both 547 strains, a copy of a *mecA* gene expressed from its native promoter was located at the *lysA* locus. 548 In $mecA^+$ $rpoB^*$ $pbp2^*$ (SJF5809) the rpoB gene has a point mutation which results in H929Q 549 (rpoB*). ery and kan represent erythromycin and kanamycin resistance cassettes, respectively. 550 The graphics in (A-B) were created with BioRender.com. (C) Growth curves of $mecA^+ pbp2$ 551 (SJF5663) grown in the presence (+) or absence (-) of IPTG, and mecA⁺ pbp2* (SJF5807) (+ 552 IPTG). (D) Growth curves of $mecA^+$ $rpoB^*$ pbp2 (SJF5674) grown in the presence (+) or 553 absence (-) of IPTG, and mecA⁺ rpoB* pbp2* (SJF5809) (+ IPTG). (E-F) AFM images of the 554 newly revealed outer surface of septa after cell division of $mecA^+$ pbp2* (SJF5807) and $mecA^+$ 555 $rpoB^* pbp2^*$ (SJF5809), respectively. In both E and F, (i) shows the outer surface of a 556 representative septum. Topographical height (z) range of 130 nm applies to both. (ii) Shows a 557 pseudo-3D high resolution image of the region within the white box in (i). Height range are 12 558 559 nm for E(ii) and 7 nm for F(ii). (iii) Represents the combined angular histogram of fibre orientation of AFM high resolution images similar to images in (ii). 560

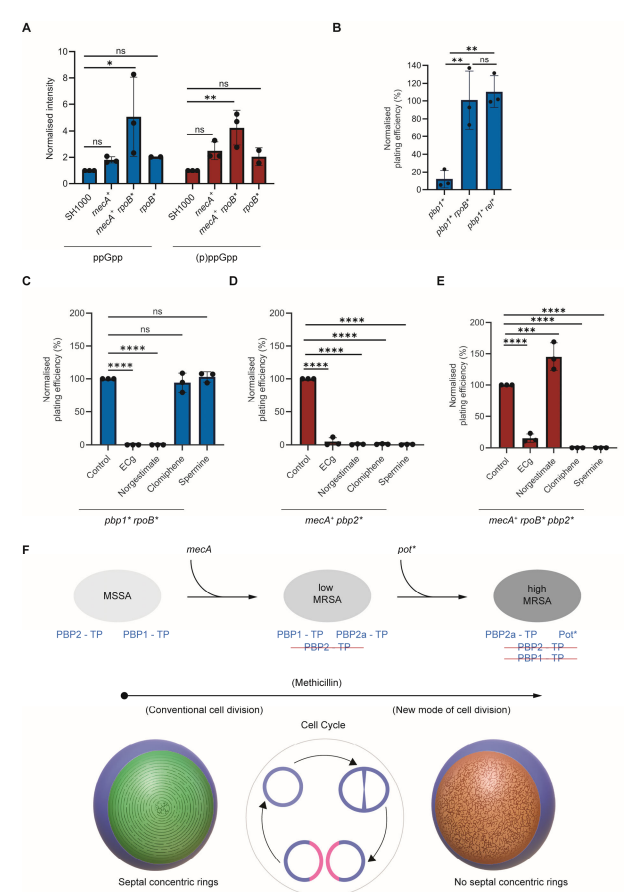


Fig. 4. Dual pathways to high-level MRSA. (A) Measurement of ppGpp and (p)ppGpp levels in SH1000, $mecA^+$, $rpoB^*$ and $mecA^+$ rpoB, normalised to SH1000 and compared using one-

way ANOVA with Dunnet's multiple comparison test (ns, not significant; *, P < 0.05; **, 564 *P* <0.01). *P* values from left to right: 0.8727, 0.0425, 0.8290, 0.1470, 0.0051 and 0.4317. Error 565 bars show the mean \pm SD. (**B**) Plating efficiency of *pbp1**, *pbp1* rpoB** and *pbp1* rel* without 566 IPTG. Plating efficiency values were compared to controls with IPTG, using one-way ANOVA 567 with Tukey's multiple comparison test (**, left to right P = 0.0049 and 0.003). Error bars show 568 mean \pm SD. (C-E) Plating efficiency of (C) pbp1* rpoB*, (D) $mecA^+ pbp2*$ and (E) $mecA^+$ 569 rpoB* pbp2* without IPTG supplemented with ECg, norgestimate, clomiphene or spermine. 570 Data were compared to no inhibitor plates (Control) using a one-way ANOVA with Dunnett's 571 multiple comparison test (ns, not significant; ****, P < 0.0001; ***, P = 0.0004). Error bars 572 show mean \pm SD from three independent biological repeats. (F) Model for high-level MRSA 573 development via acquisition of mecA and pot mutations (including rpo* and rel*), resulting in 574 low-level (low) and subsequently high-level (high) resistance. In MSSA, without methicillin, 575 576 PBP1 and PBP2 transpeptidases are active. In low-level MRSA, at intermediate methicillin levels sufficient to kill MSSA, PBP2 transpeptidase is inhibited but complemented by PBP2a. 577 In high-level MRSA, at methicillin levels sufficient to kill MSSA and low-level MRSA, PBP2 578 and PBP1 transpeptidases are inhibited but complemented by PBP2a transpeptidase and Pot*, 579 580 respectively. PBP1 transpeptidase is responsible for the characteristic septal PG concentric rings, during conventional cell division (green regions on blue cell background). In high-level 581 MRSA, in the presence of methicillin, septal PG concentric rings are replaced by mesh (brown 582 583 regions on blue cell background), revealing a novel mode of cell division requiring both PBP2a and Pot*. 584

1	
	Science
2	AAAS
3	
4	
5	Supplementary Materials for
6	
7	Two co-dependent routes lead to high-level MRSA
8	
9 10 11 12 13	Abimbola Feyisara Adedeji-Olulana ^{1†,} , Katarzyna Wacnik ^{2,3,†} , Lucia Lafage ^{2,3} , Laia Pasquina-Lemonche ^{1,3} , Mariana Tinajero-Trejo ^{2,3} , Joshua A. F. Sutton ^{2,3} , Bohdan Bilyk ^{2,3} , Sophie E. Irving ^{2,3} , Callum J. Portman Ross ^{2,3} , Oliver J. Meacock ¹ , Sam A. Randerson ¹ , Ewan Beattie ¹ , David S. Owen ² , James Florence ^{2,3} , William M. Durham ¹ , David J. Hornby ^{2,3} , Rebecca M. Corrigan ^{2,3} , Jeffrey Green ^{2,3} , Jamie K. Hobbs ^{1,*} , Simon J. Foster ^{2,3,*}
14	
15	Corresponding author. Email: jamie.hobbs@sheffield.ac.uk; s.foster@sheffield.ac.uk.
16	The PDF file includes:
17	Materials and Methods
18	Figs. S1 to S10
19	Tables S1 to S4

20 Materials and Methods

21 <u>Bacterial growth conditions</u>

The strains used in this study are listed in Table S2. S. aureus strains were grown in tryptic soy 22 23 broth (TSB), except for (p)ppGpp experiments which used low phosphate chemically defined 24 medium (39) (CDM), at 37°C with aeration. For solid, Tryptic Soy Agar media (TSA), 1.5% (w/v) agar was added. When necessary, growth media were supplemented with kanamycin 25 $(50 \,\mu g \,m l^{-1})$, tetracycline $(1 \,\mu g \,m l^{-1})$, chloramphenicol $(10 \,\mu g \,m l^{-1})$, erythromycin $(5 \,\mu g \,m l^{-1})$, 26 lincomycin (25 μ g ml⁻¹), methicillin (0.25, 1.5, 2, 25 or 40 μ g ml⁻¹), isopropyl β -D-27 thiogalactopyranoside (IPTG; 50 μ M or 1 mM) or mupirocin (60 μ g ml⁻¹). For mecA⁺ pbp2* 28 (SJF5807) and mecA⁺ rpoB^{*} pbp2^{*} (SJF5809), 1 mM IPTG was added to growth media at all 29 times. 30

31 <u>Construction of plasmids</u>

32 Escherichia coli NEB5a was used for the construction of all plasmids. Correct plasmid

sequences were confirmed by DNA sequencing (Sanger sequencing by Source BioScience).
Plasmids and oligonucleotides used in this study are listed in Table S3 and Table S4,

35 respectively.

36 <u>pKB-*Pspac-pbp2*</u>

A fragment containing the full-length *pbp2* gene and its ribosome-binding site (RBS) was PCR
amplified (from SH1000) using pCQ11-pbp2-F/-R primers and cloned into AscI and NheI cut
pCQ11-FtsZ-SNAP by Gibson assembly, resulting in pCQ11-*pbp2*. Next, the fragment
containing the *Pspac* promoter, RBS and *pbp2* was PCR amplified from *S. aureus* SH1000
genomic DNA using pKB-*Pspac*-pbp1-F and pKB-pbp2-R and cloned into BamHI and EcoRI

42 cut pGM074, giving pKB-*Pspac-pbp2*.

43 <u>pKB-Pspac-pbp2*</u>

A point mutation resulting in PBP2-TP inactivation (S398G) was introduced by site-directed
mutagenesis using pbp2TP-F/-R primers and Q5 Site Directed Mutagenesis kit (New England
Biolabs), creating pKB-*Pspac-pbp2**.

47 <u>pMAD- $\Delta pbp2$ </u>

48 One kb fragments upstream (up) and downstream (down) of *pbp2* were PCR amplified from *S*.

49 *aureus* SH1000 genomic DNA using pMAD-Δpbp2-F and pbp2-up-R, and pbp2-down-F and

50 pMAD- Δ pbp2-R, respectively, while the tetracycline cassette (*tet*) was PCR amplified from

51 pAISH1 using tetR-pbp2-F/-R primers. The up, tet and down fragments were ligated into

52 EcoRI and BamHI cut pMAD by Gibson assembly, resulting in pMAD- $\Delta pbp2$.

- 53 <u>Construction of mutants</u>
- 54 All plasmids were introduced into restrictive-deficient *S. aureus* RN4220 and moved to a final
- 55 S. aureus SH1000 strain by phage Φ 11 transduction (40, 41). Whole genome sequencing was
- 56 provided by MicrobesNG.

57 pbp1*rpoB* and $\Delta pbp1 rpoB*$

58 Strains containing $pbp1^*$ (SJF4656, SH1000 $geh::Pspac-pbp1 pbp1::pbp1^* lacI$) and $\Delta pbp1$ 59 (SJF5106, SH1000 $geh::Pspac-pbp1 \Delta pbp1 lacI$) were transduced with a phage lysate from

60 SJF5046 (SH1000 *lysA::mecA rpoB*^{H929Qkan}), resulting in *pbp1* rpoB** and $\Delta pbp1$ *rpoB**.

61 <u>*pbp2*</u>

The pKB-*Pspac-pbp2* plasmid was used to transform *S. aureus* CYL316. The chromosomal
fragment containing the plasmid integrated in the *geh* locus was moved by phage transduction
into SH1000, resulting in SH1000 *geh::Pspac-pbp2* (SJF4924). SJF4924 was then transformed
with pMAD-Δ*pbp2*. Chromosomal integration of the plasmid at 42°C and excision at 28°C led
to a marked deletion of *pbp2* (*pbp2::tet*). To provide a control of *Pspac-pbp2*, *lacI* was
introduced by transduction using a phage lysate of VF17 (SH1000 *lacI*), resulting in the *pbp2*mutant (SJF5630, SH1000 *geh::Pspac-pbp2 pbp2::tet lacI*).

69 $mecA^+ pbp2$ and $mecA^+ rpoB^* pbp2$

70 Pspac-pbp2 from SJF4924 was transduced into mecA⁺ (SJF4996, SH1000 lysA::mecA) and

71 $mecA^+$ $rpoB^*$ (SJF5003, SH1000 *lysA::mecA* $rpoB^{H929Q}$). Next, *pbp2* was deleted by

transducing *pbp2::tet* from SJF5630. Finally, *lacI* from VF17 was added, creating *mecA⁺ pbp2*(SJF5663, SH1000 *lysA::pmecA geh::Pspac-pbp2 pbp2::tet lacI*) and *mecA⁺ rpoB^{*} pbp2*

(SJF5674, SH1000 lysA::pmecA rpoB^{H929Q} geh::Pspac-pbp2 pbp2::tet lacI).

75 <u>*rpoB* pbp2*</u>

In the *rpoB** mutant (SJF5010, SH1000 *lysA::kan rpoB*^{H929Q}), the kanamycin resistance cassette in the *lysA* locus was swapped for the erythromycin resistance cassette by phage transduction of *lysA::ery* from GMSA015 (SH1000 *lysA::ery*). Next, *Pspac-pbp2* from SJF4924, *pbp2::tet* from SJF5630 and *lacI* from VF17 were added, resulting in *rpoB** *pbp2* (SJF5690, SH1000 *lysA::ery rpoB*^{H929Q} *geh::Pspac-pbp2 pbp2::tet*).

81 $\underline{mecA^+ pbp2^* and mecA^+ rpoB^* pbp2}$

S. aureus CYL316 was transformed with pKB-*Pspac-pbp2**. The chromosomal fragment
carrying the integrated plasmid (*geh::Pspac-pbp2**) was transduced into *mecA*⁺ (SJF4996) and *mecA*⁺ *rpoB** (SJF5003). Next, *pbp2* was deleted by transducing *pbp2::tet* from SJF5630.
Finally, *lacI* from VF17 was added, giving *mecA*⁺ *pbp2** (SJF5807, SH1000 *lysA::pmecA geh::Pspac-pbp2** *pbp2::tet lacI*) and (SJF5809, SH1000 *lysA::pmecA rpoB*^{H929Q} *geh::Pspac-pbp2** *pbp2::tet lacI*).

88 gdpP, lytH, pde2, clpP, clpX and rel

89 SH1000 was transduced with a phage lysate from ANG1959 (SEJ1 gdpP::kan), NE1369 (JE2

90 *lytH::Tn*), NE1208 (JE2 *pde2::Tn*) or NE1714 (JE2 *rel::Tn*), resulting in *gdpP* (SJF5025,
91 SH1000 *gdpP::kan*), *lvtH* (SJF5455, SH1000 *lvtH::Tn*), *pde2* (SJF5454, SH1000 *pde2::Tn*),

91 SH1000 gapP::kan), lyth (SJF5455, SH1000 lyth::In), pae2 (SJF5454, SH1000 pae2::In), 92 clpP (SH1000 clpP::Tn), clpX (SH1000 clpX::Tn) and rel (SJF5457, SH1000 rel::Tn),

 G_{2} cipr (SH1000 cipr.: 1n), cipx (SH1000 cipx.: 1n) and ret (SJF 5457, SH1000 ret.: 1n),

93 respectively.

94 $mecA^+ gdpP, mecA^+ lytH, mecA^+ pde2, mecA^+ clpP, mecA^+ clpX and mecA^+ rel$

The mecA⁺ mutant (SJF4996, SH1000 lysA::mecA) was transduced with a phage lysate from
 ANG195 (SEJ1 gdpP::kan), giving mecA⁺ gdpP (SJF5464, SH1000 lysA::mecA gdpP::kan).

SJF5324 (SH1000 geh::mecA lysA::tet) was transduced with a phage lysate from NE1369 (JE2 lytH::Tn), NE1208 (JE2 pde2::Tn) or NE1714 (JE2 rel::Tn), resulting in mecA⁺ lytH (SJF5461, SH1000 geh::mecA lysA::tet lytH::Tn), mecA⁺ pde2 (SJF5460, SH1000 geh::mecA lysA::tet pde2::TN), mecA⁺ clpP (SH1000 geh::mecA, clpP::Tn), mecA⁺ clpX (SH1000 geh::mecA, clpX::Tn) and mecA⁺ rel (SJF5463, SH1000 geh::mecA lysA::tet rel::Tn), respectively.

102 Directed evolution of high-level resistant MRSA strains

103 Using our previous approach (14), EMRSA15 and USA300 (minimal inhibitory concentration, 104 MIC 16-24 and 0.75 μ g ml⁻¹, respectively) were plated on gradients of oxacillin (0-256 105 μ g ml⁻¹) to select for high-level resistant derivatives. High level resistant clones (EMRSA15 106 (HL) and USA300 (HL), MIC >256 μ g ml⁻¹ oxacillin for both) were picked and subjected to 107 whole genome sequencing. This revealed 3 single nucleotide polymorphisms (SNPs) for 108 USA300 (HL) (resulting in protein alterations LysS G429S, LysS R430H and 109 SAUSA300 0212 Q141*) and a total of 13 SNPs in 10 different genes for EMRSA15 (HL).

- 110 <u>*pbp1*rel</u>*</u>
- 111 The *pbp1** mutant (SJF4656, SH1000 *geh::Pspac-pbp1 pbp1::pbp1* lac1*) was transduced
- 112 with a lysate from NE1714 (JE2 rel::Tn), giving pbp1*rel (SJF5513, SH1000 geh::Pspac-
- **113** *pbp1 pbp1::pbp1* lacI rel::Tn*).
- 114 <u>Plating efficiency</u>

Cells were grown from an OD₆₀₀ of 0.1 in TSB supplemented with 10 µg ml⁻¹ chloramphenicol 115 and 50 μ M IPTG to early exponential phase (OD₆₀₀ ~0.3-0.5). Cells were then washed three 116 times in phosphate-buffered saline (PBS) and a dilution series of cell suspensions were plated 117 onto TSA containing 10 µg ml⁻¹ chloramphenicol, with or without 1 mM IPTG. For 118 experiments with sensitizing compounds, pbp1* rpoB* (SJF5306) was grown as described 119 above and plated on TSA containing 10 µg ml⁻¹ chloramphenicol and a sensitizing compound. 120 mecA⁺ pbp2* (SJF5807) and mecA⁺ rpoB* pbp2* (SJF5809) were grown in TSB supplemented 121 with 10 μ g ml⁻¹ chloramphenicol and 1 mM IPTG to exponential phase (OD₆₀₀ ~0.5) and plated 122 on TSA containing 10 μ g ml⁻¹ chloramphenicol, 1 mM IPTG and a sensitizing 123 compound. Relative plating efficiency was expressed as the number of cells on plates without 124 IPTG, compared to the number of cells from plates with IPTG, multiplied by 100. 125

126 <u>Depletion of PBP1 or PBP2</u>

127 Pspac-pbp1 and Pspac-pbp2 containing strains were grown from an OD_{600} of 0.1 to the 128 exponential phase ($OD_{600} \sim 0.3$ -0.5) in TSB supplemented with 10 µg ml⁻¹ chloramphenicol 129 and 50 µM IPTG. Cells were then washed three times by centrifugation and resuspension in 130 TSB and inoculated in fresh TSB containing 10 µg ml⁻¹ chloramphenicol to an OD_{600} of 0.05 131 for phenotypic analysis, or to an OD_{600} of 0.005 for growth studies. Control samples (+IPTG) 132 were grown in TSB supplemented with 10 µg ml⁻¹ chloramphenicol and 1 mM IPTG at all 133 times.

134 Growth in methicillin

High-level resistant, clinical MRSA strains and mecA⁺ rpoB*(SJF5003) were grown overnight 135 in the presence of 25 μ g ml⁻¹ methicillin (or 50 μ g ml⁻¹ where stated), diluted to an initial OD₆₀₀ 136 of 0.05, prior to growth to exponential phase (OD₆₀₀ ~0.5) in the presence of 25 μ g ml⁻¹ 137 methicillin (or 50 μ g ml⁻¹ where stated) at all times. This gives a total of over 8 generations in 138 139 the presence of a high level of methicillin. This concentration of methicillin is sub-MIC for these strains and does not significantly affect growth rate in liquid culture. mecA⁺ (SJF4996) 140 was grown in TSB overnight, diluted to an initial OD₆₀₀ of 0.05, prior to growth in the presence 141 of 1.5 μ g ml⁻¹ methicillin (sub-MIC) to exponential phase (OD₆₀₀ ~0.5). This concentration of 142 methicillin is sub-MIC for this strain and does not significantly affect growth rate in liquid 143 culture. Finally, SH1000 and mecA⁺ (SJF4996) were grown in TSB overnight, diluted to an 144 145 initial OD₆₀₀ of 0.05, prior to growth to exponential phase (OD₆₀₀ \sim 0.5), followed by the addition of 1.5 μ g ml⁻¹ methicillin (supra- and sub-MIC, respectively) and their phenotype 146 followed over the next 3h. 147

148 MIC determination and evaluation of MRSA resensitizing compounds

149 Oxacillin MIC values were determined using E-test MIC Evaluator (Liofilchem or bioMérieux) 150 strips in triplicate. MIC values of resensitizing compounds were determined for SH1000 by growth overnight in liquid TSB, for ECg (200 μ g ml⁻¹), norgestimate (160 μ g ml⁻¹), 151 clomiphene (8 μ g ml⁻¹) and spermine (320 μ g ml⁻¹). Resensitization of mecA⁺ rpoB^{*} to 152 oxacillin (from > 256 µg ml⁻¹ (control) to ≤ 4 µg ml⁻¹ (with resensitizer) was determined using 153 ECg (50 μ g ml⁻¹), norgestimate (10 μ g ml⁻¹), clomiphene (4 μ g ml⁻¹) or spermine (202 μ g ml⁻¹) 154 in TSA, spread with 200 μ l of an overnight culture (diluted normalised to an OD₆₀₀ of ~2 in 155 156 TSB) and overlaid with an E-test strip. The above resensitizer treatments also led to a drop in oxacillin MIC from > 256 μ g ml⁻¹ to $\leq 12 \mu$ g ml⁻¹ for the clinical strains (COL, Mu50, 157 MRSA252, TW20, USA300 (HL) and EMRSA15 (HL), apart for TW20 with norgestimate 158 (170 μ g ml⁻¹) and clomiphene (42 μ g ml⁻¹). The effect of resensitizers on the growth of S. 159 aureus strains was determined using the above sub-MIC concentration of compounds. 160

161 <u>Fluorescence microscopy</u>

162 Fixed and labelled cells were dried onto a poly-L-lysine-coated slide, mounted in SlowFade 163 Gold (Thermo Fisher) and imaged using a Nikon Ti inverted microscope fitted with a 164 Lumencor Spectra X light engine. Images were obtained using a 100× PlanApo (1.4 NA) oil 165 objective using 1.518 RI oil and detected by an Andor Zyla sCMOS camera. The raw data in 166 format .nd2 with three channels (ADA-DA, NHS-ester and brightfield) were used for cell 167 volume and PG synthesis analysis.

168 <u>PFA fixation</u>

169 Cells were treated with 1.6% (wt/vol) paraformaldehyde for 30 min at RT and PFA removed170 by washing cells with water prior to imaging.

171 ADA-DA labelling

- 172 Cells were incubated with 1 mM ADA-DA (azido-D-alanyl-D-alanine, produced as previously
- described (8) at 37°C. Cells were washed, fixed with PFA, and labelled with Alexa Fluor 488
- 174 Alkyne (Merck) (for fluorescent labelling of the ADA-DA azide group) by the click reaction

- 175 (copper(I)-catalysed alkyne-azide cycloaddition) using the Click-iTTM Cell Reaction Buffer
- 176 Kit (Invitrogen) according to the manufacturer's instructions.
- 177 <u>NHS-ester labelling</u>

178 Cells were incubated with 8 μ g ml⁻¹ NHS-ester Alexa Fluor 555 in PBS for 5 min on ice. Cells 179 were then washed in ice cold PBS and fixed with PFA.

180 <u>Fluorescence intensity measurements</u>

Fluorescence intensity of incorporated ADA-DA clicked to Atto488 was measured using Image
 J/Fiji and calculated as counts/pixel.

183 <u>Preparation of whole cell lysates</u>

184 Cells resuspended in PBS were lysed using Lysing Matrix B and FastPrep homogeniser (MP

biomedicals) in 10 cycles of 30 s, at speed of 6.5 m s^{-1} , with a 3-min incubation on ice between cycles. Broken cells were separated from unbroken cells and lysing matrix by centrifugation

 $(5,000 \text{ x g}, 5 \text{ min}, 4^{\circ}\text{C})$. Total protein concentration was established using the BCA protein kit

- 188 (Pierce).
- 189 <u>Western Blot</u>

A total of 50 µg protein was separated on a 10% (w/v) SDS-PAGE gel and transferred onto a 190 nitrocellulose membrane. Membranes were blocked in 5% (w/v) skimmed-milk in TBST 191 (20 mM Tris-HCl, pH 7.6; 17 mM NaCl; 0.1% (v/v) Tween-20) and incubated with polyclonal 192 primary antibodies (1:1,000 dilution for anti-PBP1; 1:2,500 dilution for anti-PBP2) overnight 193 194 at 4°C. For detection, horseradish peroxidase-conjugated goat anti-rabbit IgG (Sigma; 1:10,000 dilution) and Clarity Western ECL Substrate (BioRad) were used according to the 195 manufacturer's instructions. Chemiluminescence was detected using Syngene G:BOX Chemi 196 197 XX9.

- 198
- 199 <u>BocillinFL gel-based analysis</u>

BocillinFL binding was adapted from a published method (35). Whole cell lysates (50 μ g total protein) were incubated with 100 μ M BocillinFL (Invitrogen) for 15 min at 30°C. The reaction was stopped by the addition of 5x SDS-PAGE loading buffer and incubation for 15 min at

was stopped by the addition of 5x SDS-PAGE loading buffer and incubation for 15 min at 30° C. The proteins were separated on a 10% (w/v) SDS-PAGE gel and visualized using a Bio-

- 204 Rad ChemiDoc MP imaging system or a GE Typhoon FLA 9500.
- 205 <u>Transmission electron microscopy (TEM)</u>
- 206 Cell preparation for electron microscopy was performed as described previously (15). Cell

207 pellets (5 ml cultures) were fixed overnight at 4°C in 2.5% (w/v) EM grade glutaraldehyde

208 (Agar Scientific). Samples were washed with PBS and resuspended in 2% (w/v) aqueous 209 osmium tetroxide (Agar Scientific) for secondary fixation (2 h at room temperature). Cells were

washed with PBS and dehydrated by incubating with increasing concentrations of ethanol (60,

washed with FBS and derivated by medbating with increasing concentrations of ethalor (oo, 75, 95 and 100% (v/v) ethanol) 15 min each. Ethanol was removed and samples were incubated

with propylene oxide (Agar Scientific) to complete dehydration. Samples were mixed with a

1:1 mix of propylene oxide and Epon resin (Agar Scientific) and incubated overnight at room 213 214 temperature to allow infiltration. The majority of the resin was removed, and the excess of propylene oxide evaporated at room temperature. Two consecutive incubations of the samples 215 with pure Epon resin (4 h each) were performed and cells were embedded in fresh resin. Resin 216 polymerisation was achieved by incubation at 60°C for 48 h. Thin sections (85 nm) were 217 produced using an Ultracut E Ultramicrotome (Reichert-Jung) and mounted onto 300-square 218 219 mesh nickel TEM grids (Agar Scientific). Sections were stained in 3% (w/v) aqueous uranyl acetate (Polysciences Inc., 21447) for 25 min, washed with dH₂O and stained with Reynold's 220 221 lead citrate for 5 min. The citrate was removed by washing with dH₂O. A FEI Tecnai T12 Spirit Transmission Electron Microscope operating at 80 kV was used for imaging. A Gatan Orius 222 SC1000B bottom-mounted CCD camera recorded the images. EM images were analysed using 223 Fiji (42). Over 300 cells were counted for each repeat, with at least two independent repeats 224 225 per sample/treatment. Cells were categorised according to their phenotypic features as normal 226 and abnormal, dependent on septal misplacement or growth defects.

227 Extraction and purification of PG

PG was extracted as previously described (16). Briefly, cells were grown in the presence or 228 absence of IPTG for 2-4 h for PBP1 and PBP2 depletion experiments or to mid-exponential 229 phase with or without methicillin for methicillin-treatment experiments and boiled at 100°C 230 231 for 15 min to kill the cells. Boiled cells were recovered by centrifugation at 20,000 x g for 3 min and the supernatant was discarded. Next, the pellets were suspended in PBS and transferred 232 233 to the lysing matrix tubes containing 0.1 mm silica beads and broken using FastPrep 24 Homogeniser (10 cycles of 30 s, at speed of 6.5 m s⁻¹, with a 3-min incubation on ice between 234 cycles). Next, the tubes were centrifuged at 170 x g at RT to remove the beads and the 235 236 supernatant, containing the broken cells, was pipetted into new Eppendorf tubes, which were then centrifuged at 20,000 x g at RT for 3 min. PG was resuspended in 5% (w/v) SDS solution 237 and boiled at 100°C for 25 min. Boiling in SDS was repeated one more time for 15 min and 238 239 the PG was serially rinsed with Milli-Q water. Lastly, PG was resuspended in a solution of 50 mM Tris HCl pH 7, containing 2 mg ml⁻¹ of pronase, and incubated at 60°C for 90 min. 240 Afterwards, PG was rinsed thrice with HPLC grade water, and stored at 4°C without any further 241 242 treatment.

243 PG immobilisation and AFM imaging

244 To immobilise purified PG, a mica substrate was coated with Cell-tak (Corning, Netherlands), which is a solution of polyphenolic proteins. Briefly, the mica substrate was incubated with 245 180 µl of Cell-tak solution (171 µl of 100 mM sodium bicarbonate (NaHCO₃) pH 8.0, 3 µl of 246 1 M sodium hydroxide (NaOH), and 6 µl of 1.05 mg ml⁻¹ Cell-tak) for 30 min. Next, the 247 substrate was rinsed three times with HPLC grade water and then dried with nitrogen flow. 248 Fifty microliters of diluted PG solution was then added to the Cell-tak coated substrate, 249 incubated for 1 h and rinsed with HPLC grade water and dried with nitrogen flow. For thickness 250 251 measurements, AFM height topographic images of dehydrated PG were captured in air using AFM tapping mode with Nunano SCOUT 350 - Silicon AFM probe (spring constant: 42 N/m, 252 253 Resonance frequency: 350 kHz) at free amplitude of 10 nm with set point of 70-80% of free 254 amplitude (e.g. 7 nm) on a Dimension FastScan Bio (Bruker, Santa Barbara). For AFM highresolution imaging, all the high-resolution images were acquired in Peak force tapping mode 255 in imaging buffer (10 mM Tris; 200 mM KCl; 10 mM MgCl₂, pH 8.0) with the Bruker 256 257 Fastscan-D cantilevers at the range of 1-3 nN peak force set point on a Dimension FastScan AFM (Bruker, Santa Barbara). The imaging parameters used are as follows; Scan rate: 1 Hz; 258 259 Scan angle: 0°; Peak Force frequency: 2-8 kHz, Peak force amplitude: 80-100 nm, and with

high pixel resolution. Prior to high-resolution imaging, the spring constant and deflectionsensitivity of the cantilevers were calculated using the Sader thermal spectra method.

262 PG thickness measurement data processing

To manually measure the thickness of AFM topographic height images of dehydrated PG were imported into Gwyddion 2.55, masked and levelled using first order polynomial row fit. The thickness of the single leaflet of the PG was measured using the one-dimensional statistical function tool, which computes the average height density across the image frame (or selected area). The height density graph was plotted with their characteristic two peaks. Then, Gaussian functions were fitted on each peak and the background peak was subtracted from the peak of the non-background, to give the average height/thickness of the PG fragment.

- 270 <u>AFM three-dimension (3D) image processing</u>
- 271 The pseudo 3D AFM images in Fig. 1Aii-Eii, Fig. 2Eii (+ IPTG, -IPTG), and Fig. 3Eii-Fii,
- 272 were processed using Nanoscope analysis software. The following image processing
- 273 parameters were used; Pitch = 10, Z-axis aspect ratio = 0.3, image rotation = 0° , plot type =
- height, and projection = parallel.
- 275 Fibre detection and orientation quantification

Peptidoglycan fibres in AFM images were detected and quantified using automated image 276 analysis methods. Our framework was implemented as a Matlab GUI, which accepts outputs 277 from Gwyddion or open-source alternatives such as TopoStats (43). In the first stage, we 278 279 upsampled all images to a uniform resolution (0.1 nm/pixel) to ensure uniformity of analyses across samples. Flattening using a Difference of Gaussians (DoG) filter removed low spatial-280 frequency components, separating the high spatial-frequency PG fibre network from the overall 281 282 geometry of the cell wall. Individual PG fibres were segmented by employing a ridge-detection 283 algorithm (44), the results of which were then fused into a continuous network using a Watershed algorithm (45) to remove small gaps between fibres at crossing points. This rough 284 285 network was then skeletonised and converted to a graph consisting of nodes, representing fibrefibre crossing points, linked by edges, representing fibre bodies. This graph-based network was 286 287 then cleaned based on its topology, removing disconnected fibre sections and fusing adjacent T-shaped configurations of fibres to more accurately capture X-shaped fibre-fibre crossings. 288 Edges and nodes were then assigned to separate fibres based on the original segmentation of 289 the fibre network. Finally, we measured the local orientation of fibres by considering each point 290 291 along the length of the fibre in turn, performing a linear regression on the points assigned to the fibre within a neighbourhood of 10 pixels of the query point and used the resulting 292 293 regression slope as the orientation measurement. From these local orientation measurements, 294 we computed the angular histogram for each image. The fibreFinder codes are publicly 295 available (46).

296 Pore analysis and quantification using Fiji

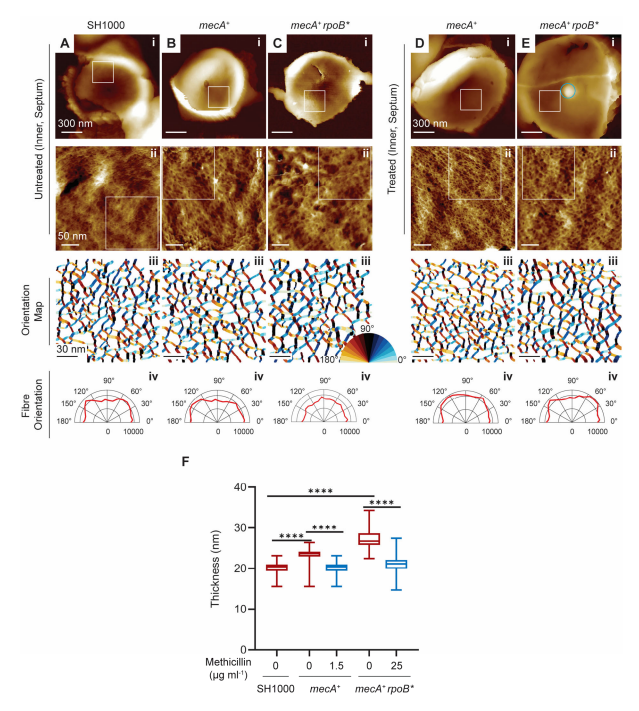
To calculate the area of the pores distributed across both the inner and outer section of the PG, we used a custom-made semi-automated pore analysis macro (AFM_Slicer) in an open-source software ImageJ/FIJI (42). Firstly, the macro pre-filtered the high-resolution image by downscaling the pixel number and removing the noise of the image using the despeckle tool in FIJI. Next, the treated image was simultaneously binarized and sliced into stacks of binary slices where black represented the fibres in the image and white denoted the pores. Lastly, the area of the pores in each slice was calculated using the analyze particle tool, followed by

- calculation of the cumulative fraction of the pore area in each slice. The slice for each imagethat had the maximum number of pores was used for the graphs in figs. S2, S6 and S10 (47).
- 306 Measurement of (p)ppGpp levels

S. aureus strains were grown overnight in low-phosphate CDM (39) at 37°C. Cultures were 307 diluted to an OD₆₀₀ of 0.1 and grown for 2.5 h prior to the addition of 3.7 MBq of [³²P]H₃PO₄ 308 and incubation for a further 3 h at 37°C. Cultures were subsequently normalized for optical 309 denisty, cells recovered by centrifugation $(17,000 \times g \text{ for } 5 \text{ min})$ and suspended in 100 µl of 310 311 600 mM formic acid. Cells were subjected to three freeze/thaw cycles and debris removed by centrifugation $(17,000 \times g \text{ for } 5 \text{ min})$ before the lysate was filtered through a 3 kDa spin 312 313 column. Ten microliters were subsequently spotted on PEI-cellulose F thin-layer 314 chromatography (TLC) plates (Merck Millipore), nucleotides separated, and TLC plates developed using a 1.5 M KH₂PO₄, pH 3.6, buffer. The radioactive spots were visualized using 315 an FLA 7000 Typhoon PhosphorImager, and data were quantified using ImageQuantTL 316 317 software.

318 <u>Cell volume measurements</u>

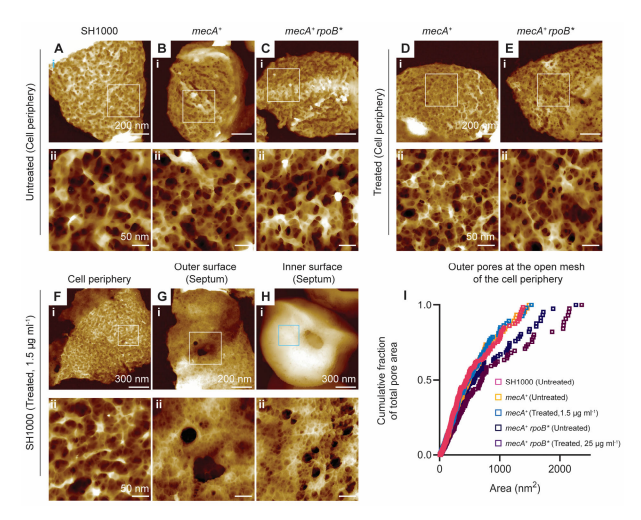
319 The cell volume measurements from the microscopy images were performed using a semi-320 automatic analysis approach (CocciVol) with a combination of FIJI macros (available at https://github.com/Laia-Pasquina/CocciVol) and a user-friendly machine learning interactive 321 open-source software named ilastik (48). First the raw data from the confocal Nikon 322 microscope was obtained in a file format .nd2. The raw data contains a stack of 11 slices across 323 3 µm with the cells focused approximately on the middle with a minimum of three channels: 324 325 ADA-DA, NHS-ester and brightfield. The Macrol V2 Filtering preparing image.ijm file in GitHub opens the raw image and creates a Z stack from the NHS-ester channel of 3 slices 326 around the focus (which needs to be pre-determined by the user by opening the image with FIJI 327 328 and manually finding the focus). Then, the image in .png format is processed in ilastik using 329 two projects. The first ilastik project uses machine learning to run a pixel segmentation routine to learn from the users input in a few cells and then classify the rest of the cells in what pixels 330 331 correspond to cells and what pixels are background. The second ilastik project uses an object classification routine similar to the first one but that classifies the cells into good fit or bad fit 332 for volume analysis as well as applying some watershed filters to distinguish cells that are too 333 334 close together (like diploids). Finally, the ilastik program outputs a table with the analysis of several physical parameters for each cell, which requires further processing to obtain the 335 volume additional processing 336 of the cells. This is carried out with 337 Macro2 Calculate Volume from table.ijm in FIJI (42) to obtain an Excel table with each cell number, volume and the ratio between the short axis and the long axis of the cell. Once one 338 339 image was analysed for one type of sample there is a batch processing option in ilastik which was used to process an average of 5 images per sample. Using this approach, n = 600 images 340 can be analysed in 30 min or less and a similar number of cells were analysed for each sample. 341 342 Several samples were compared to each other in this manuscript. The CocciVol approach was used for the graphs in figs. S4, S5 & S9 (49). 343



345 Fig. S1.

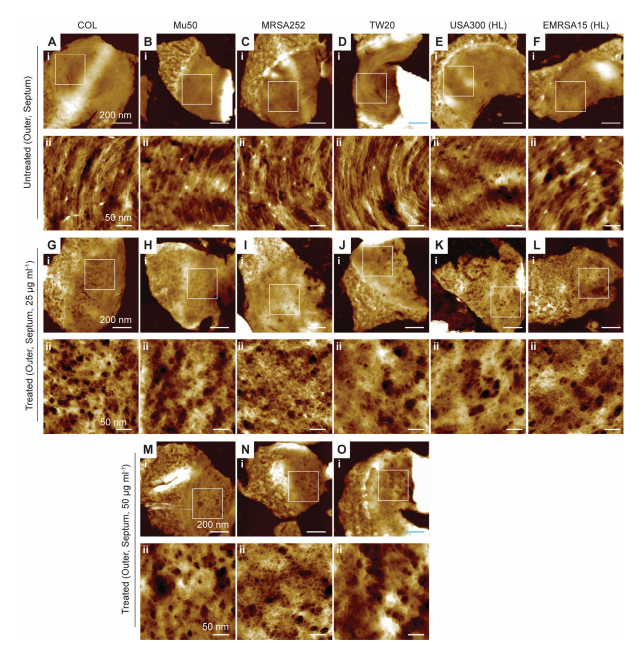
Effect of methicillin on the PG architecture of the inner surface of the septum in MRSA. (A-346 **F**) Parental MSSA strain, SH1000 (A) and MRSA strains, $mecA^+$ (**B**, **D**) and $mecA^+$ $rpoB^*$ (**C**, 347 E) were grown without (Untreated; A-C) or with methicillin (Treated; D, E), at 1.5 and 25 μg 348 ml⁻¹ for **D** and **E** respectively. (i) AFM images of the inner surface of the septum and (ii) 349 350 corresponding higher resolution topographic images of the selected location marked with white square in (i). The topographical height (z) range is as follows; (Ai) 230 nm, (Aii) 13 nm, (Bi) 351 420 nm, (Bii) 11 nm, (Ci) 170 nm, (Cii) 13.5 nm, (Di) 545 nm, (Dii) 11.5 nm, (Ei) 520 nm, 352 (Eii) 8 nm. The circle in (E) indicates the presence of a cell wall lump which is a characteristic 353 feature of $mecA^+ rpoB^*$ when treated with methicillin. The minimum height of the lump is 10 354 nm (n = 10, hydrated sacculi). (iii) Represents the colour-coded orientation map of the fibre 355

chains detected within the white box in (ii), and (iv) depicts the angular histogram of combined orientation of three independent AFM images. The image analysis method used for glycan orientation detection is described in the Materials and Methods section. (F) Plot of the measured thickness of dehydrated sacculi of SH1000, $mecA^+$ and $mecA^+$ $rpoB^*$ with respect to without (red) and with methicillin treatment (light blue). The thickness data were compared using the Mann-Whitney test (****, P < 0.0001, n = 28 for each strain and condition). Data are representative of three independent biological repeats.



364 Fig. S2.

365 Effect of methicillin on the PG architecture of the cell periphery of MRSA strains. (A-C) The parental MSSA strain, SH1000 ((A) and MRSA strains, $mecA^+$ (B) and $mecA^+$ $rpoB^*$ (C) were 366 grown without (Untreated; A-C) or with methicillin (Treated; D, E), at 1.5 and 25 µg ml⁻¹ for 367 **D** and **E**, respectively. (i) AFM images of the outer open mesh of the cell wall periphery and 368 369 (ii) corresponding higher resolution topographic images of the selected location marked with 370 white square in (i). The topographical height (z) range is as follows; (Ai) 80 nm, (Aii) 33 nm, (Bi) 110 nm, (Bii) 45 nm, (Ci) 140 nm, (Cii) 48 nm, (Di) 100, (Dii) 35 nm, (Ei) 95 nm, (Eii) 371 45 nm. (F-H) AFM images of (i), low resolution and (ii) corresponding higher resolution 372 topographic images of the selected location marked with white square in (i) of sacculi of 373 SH1000 treated 1.5 µg ml⁻¹ methicillin. (F) Outer PG mesh at the cell periphery; (G) Outer 374 surface of the septum; (H) Inner surface of the septum. The topographical height (z) range is 375 as follows; (Fi) 70 nm, (Fii) 35 nm, (Gi) 95 nm, (Gii) 43 nm, (Hi) 950 nm, (Hii) 48 nm. (I) 376 Combined plot of cumulative fraction of total pore area, distributed over the outer open mesh 377 378 shown in (A-E) (ii). Data are representative of three independent biological repeats and five 379 AFM-independent images.

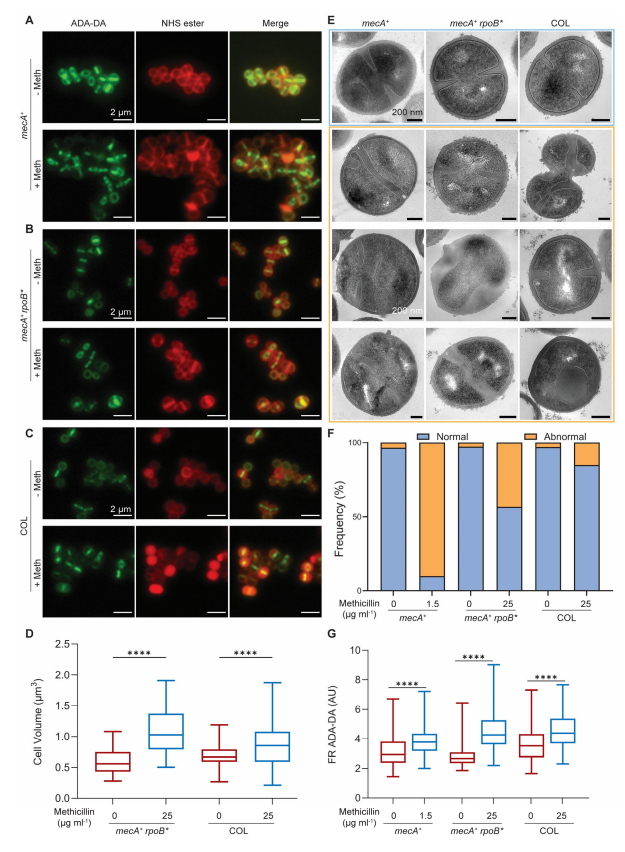


381 Fig. S3.

380

AFM characterization of high level resistant clinical strains. (A-F) Show AFM topographic 382 images of the outer surfaces of newly revealed septa in samples of isolated sacculi of untreated 383 (A) COL, (B) Mu50, (C) MRSA252, (D) TW20, (E) USA300(HL), and (E) EMRSA15(HL). 384 (i) Low-resolution AFM images and (ii) corresponding higher-resolution images of the region 385 386 indicated by the white boxes in (i). The topographical height (z) range is as follows; (Ai) 150 nm, (Aii) 13.5 nm, (Bi) 85 nm, (Bii) 11 nm, (Ci) 75 nm, (Cii) 9 nm, (Di) 65 nm, (Dii) 11.5 nm, 387 (Ei) 80 nm, (Eii) 13.5 nm, (Fi) 105 nm, and (Fii) 11.5 nm. (G-L) Show outer surfaces of the 388 septa of the same strains treated with 25 µg ml⁻¹ methicillin. (i) Low-resolution AFM images 389 and (ii) corresponding higher-resolution images of the region indicated by the white boxes in 390 391 (i). The topographical height (z) range is as follows; (Gi) 95 nm, (Gii) 16 nm, (Hi) 90 nm, (Hii) 18 nm, (Ii) 65 nm, (Iii) 12 nm, (Ji) 88 nm, (Jii) 24 nm, (Ki) 60 nm, (Kii) 25 nm, (Li) 65 nm, 392 and (Lii) 35 nm. (M-O) Show the outer surface of the newly revealed septa of Mu50, 393 MRSA252, and TW20 treated with 50 µg ml⁻¹ methicillin. (i) Low-resolution AFM images and 394 395 (ii) corresponding higher-resolution images of the region indicated by the white boxes in (i).

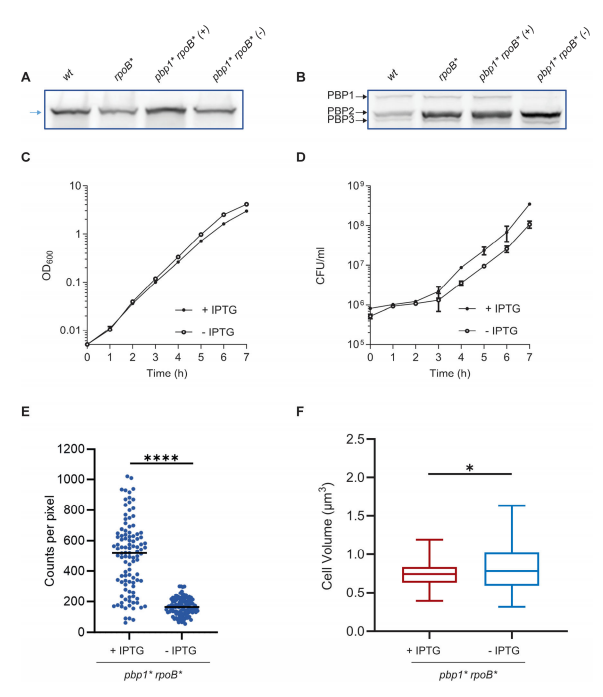
- 396
- The topographical height (z) range is as follows; (Mi) 130 nm, (Mii) 36 nm, (Ni) 105 nm, (Nii) 24 nm, (Oi) 78 nm, and (Oii) 28 nm. Data are representative of two independent biological 397 repeats and three AFM-independent images. 398





401 Effect of methicillin on cell morphology of MRSA strains. (A-C) Fluorescence microscopy 402 images of ADA-DA and NHS-ester labelled MRSA strains (A) $mecA^+$, (B) $mecA^+$ $rpoB^*$, and 403 (C) COL grown without (-Meth, i) or with methicillin (+ Meth, ii), at 1.5, 25 and 25 µg ml⁻¹

404 for A, B and C, respectively. (D) Cell volumes of $mecA^+$ $rpoB^*$ and COL without and with methicillin (1.5 µg ml⁻¹ and 25 µg ml⁻¹), as measured by fluorescence microscopy after NHS-405 ester Alexa Fluor 555 labelling. The P value was determined by Mann-Whitney U tests (****. 406 407 P < 0.0001). The number of cells analysed per sample was $n \ge 300$. (E) Shows TEM images of $mecA^+$, $mecA^+$ $rpoB^*$ and COL grown with methicillin at 1.5, 25 and 25 µg ml⁻¹ respectively. 408 Representatives of cells with normal septa are within a light blue frame while cells with 409 410 abnormal septa are within orange frame. (F) Quantification of cellular phenotypes based on TEM. Cells were categorised as normal and abnormal, depending on the presence of a 411 misplaced, multiple or misshapen septa or other cell cycle defects. Scale bar = 200 nm. Data 412 413 are representative of two independent biological repeats (n > 300). (G) Fluorescence ratio (FR) was calculated by dividing the fluorescence intensity at the septum by the intensity at the cell 414 periphery of $mecA^+$, $mecA^+$ rpoB*, and COL grown without (red, 0 µg ml⁻¹) and with 415 methicillin (light blue) at different concentrations 1.5, 25 and 25 µg ml⁻¹ respectively. 416 Differences are highly significant (****P < 0.0001) with higher FR for cells treated with 417 methicillin (light blue) compared to no treatment (red). Higher values indicate more PG 418 incorporation at the septum whilst lower values mean more peripheral PG. Number of cells 419 420 analysed for each sample was 100.

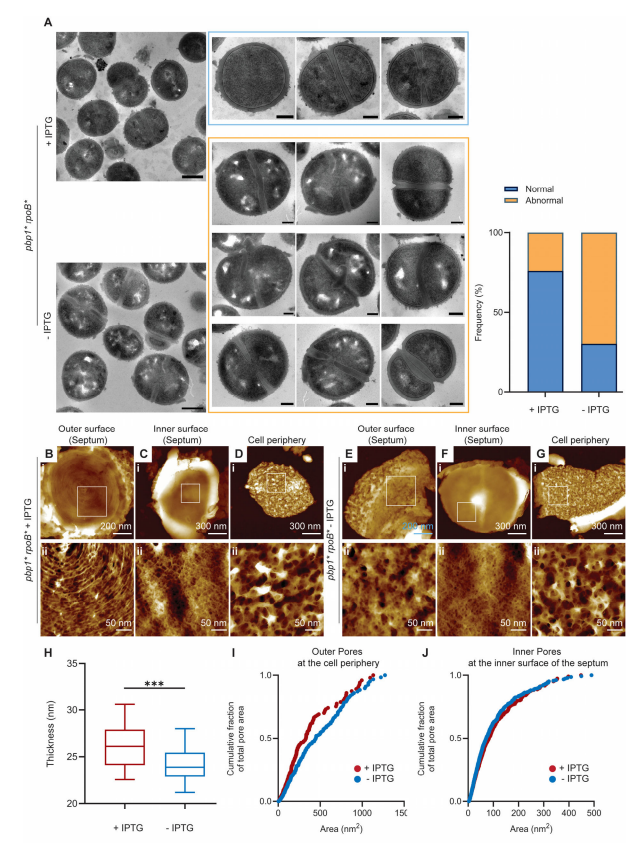






423 Analysis of the role of PBP1 transpeptidase activity. (A) Immunoblot of whole cell lysates of SH1000, *rpoB** and *pbp1** *rpoB** grown with IPTG (+) without IPTG (-) for 4 h analysed using 424 anti-PBP1 antibody. Expected PBP1 and PBP1* size = 83 kDa (blue arrowhead). (B) 425 BocillinFL gel-based analysis of PBPs in SH1000, rpoB* and pbp1* rpoB* grown with IPTG 426 (+) without IPTG (-) for 4 h; the locations of PBPs on the blots are indicated (arrows). (C) 427 Growth curves of *pbp1*rpoB** grown in the presence or absence of IPTG (+ IPTG and - IPTG, 428 429 respectively). Data represent the mean \pm SD. Error bars that are smaller than the data point symbols are not shown. (**D**) CFU counts of *pbp1* rpoB** grown in the presence or absence of 430 431 IPTG (+ IPTG and - IPTG, respectively). Data represent the mean \pm SD. Error bars that are 432 smaller than the data point symbols are not shown. (E) ADA-DA, clicked to Atto488, incorporation over 5 min in pbp1* rpoB* (SJF5306) grown with IPTG (+) and without IPTG 433 (-). Fluorescence intensities were compared using a one-way ANOVA with Tukey's multiple 434

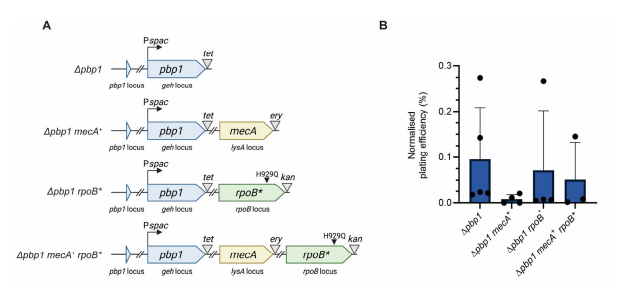
- 435 comparison test (****, P < 0.0001). Number of cells analysed for each sample was n = 110. 436 Each dot represents a single cell. A black line indicates the median of each distribution. Data
- 437 are representative of three independent biological repeats. (F) Cell volumes of pbp1*rpoB*438 after incubation with (+, red, average cell volume is $0.75 \pm 0.16 \,\mu\text{m}^3$) or without (-, light blue,
- 438 after incubation with (+, red, average cell volume is $0.75 \pm 0.16 \,\mu\text{m}^3$) or without (-, light blue, 439 average cell volume is $0.83 \pm 0.30 \,\mu\text{m}^3$) IPTG for 4h, as measured by fluorescence microscopy
- 435 average cent volume is $0.05 \pm 0.50 \,\mu\text{m}$) if 10 for 4n, as measured by hubrescence microscopy 440 after NHS-ester Alexa Fluor 555 labelling. The *P* value was determined by Mann-Whitney U
- 441 tests (*, P = 0.0266). Number of cells analysed for each sample was ≥ 250 .





444 Cell wall morphology of pbpl*rpoB* revealed by TEM and AFM. (A) Left, representative 445 TEM micrographs of pbpl*rpoB* grown for 4 h in the presence and absence of IPTG. Scale 446 bars = 500 nm. Examples of cells classified as normal and abnormal phenotypes are shown

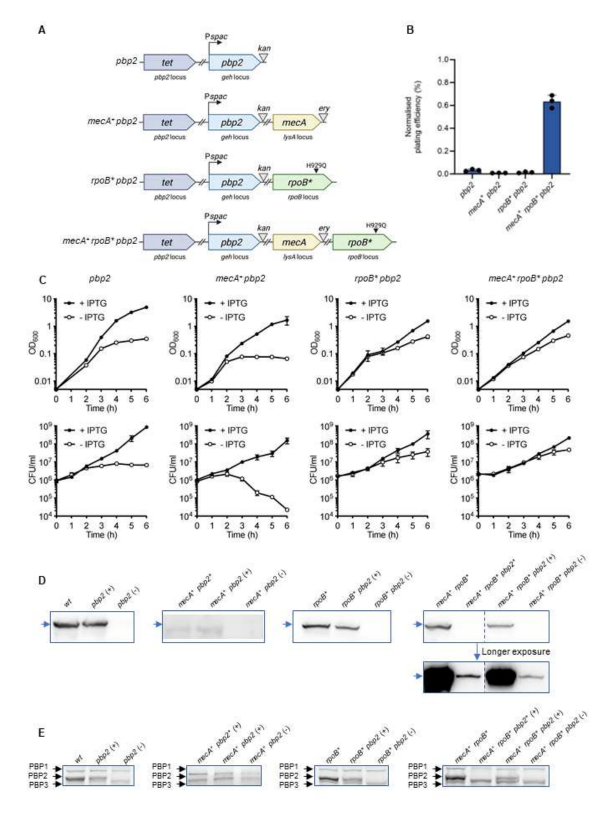
447 within the light blue (+IPTG) and orange frames (-IPTG), respectively. Scale bar = 200 nm. TEM data are representative of two independent biological repeats with $n \ge 300$ per 448 sample. Right, quantification of cellular phenotypes associated with *pbp1* rpoB**, based on 449 450 the TEM data. (B-G) AFM images of the outer surface of the septum (B, E), the inner surface of the septum (C, F), and the open mesh at the cell periphery (D, G) associated with *pbp1** 451 rpoB* grown in the presence of IPTG (+ IPTG; B-D) and absence of IPTG (- IPTG; E-G). For 452 453 (B-G) (i) are individual fragments of sacculus and (ii) are higher resolution topographic AFM images of the selected location marked with the white square in (i). The topographical height 454 (z) range is as follows; + IPTG: 120 nm (Bi), 8.5 nm (Bii), 270 nm (Ci), 13 nm (Cii), 120 nm 455 456 (Di), 55 nm (Dii). -IPTG: 110 nm (Ei), 22 nm (Eii), 290 nm (Fi), 17 nm (Fii), 120 nm (Gi), 48 nm (Gii). Data are representative of three independent biological repeats and more than 10 457 458 AFM independent imaging experiments. (H) Plot of the measured thickness of dehydrated 459 sacculi of *ppbp1* rpoB** after incubation with (+, red) or without (-, light blue) IPTG for 4h. The number of independent fragments measured for each strain was 20. The P value was 460 determined by Mann-Whitney U tests (***, P = 0.0009). (I-J) Cumulative fraction of total pore 461 area as a function of the area of the pores distributed across the open mesh surface of the cell 462 463 wall periphery surface (I) and the inner surface (J) of the septum of *pbp1*rpoB** grown in the presence of IPTG (+ IPTG, red) and absence of IPTG (- IPTG, light blue). 464



466 Fig. S7.

465

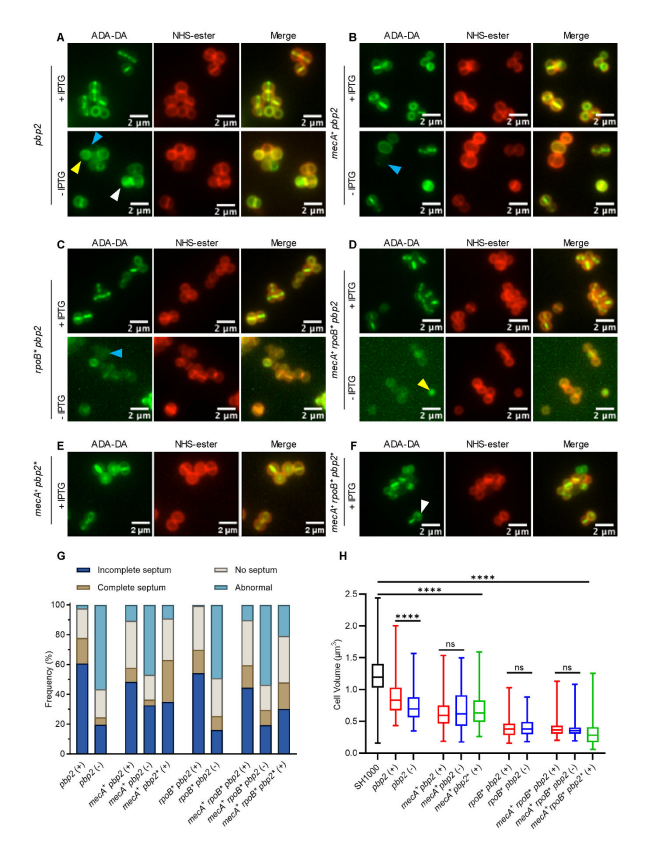
Analysis of the role of PBP1 in growth. (A) Schematic representation of the $\Delta pbp1$ genetic 467 constructs. An ectopic copy of *pbp1* was placed under the control of the Pspac promoter at the 468 469 lipase (geh) locus, while the gene in the native pbp1 locus was deleted ($\Delta pbp1$). In $\Delta pbp1$ 470 $mecA^+$ and $\Delta pbp1 mecA^+ rpoB^*$ a copy of a $mecA^+$ gene under the control of its native promoter is placed at the *lysA* locus. In $\Delta pbp1 rpoB^*$ and $\Delta pbp1 mecA^+ rpoB^*$, the *rpoB* gene has a point 471 472 mutation which results in a single amino acid replacement (H929Q) in RNA polymerase β 473 subunit (rpoB*). tet, ery and kan represent tetracycline, erythromycin and kanamycin 474 resistance cassettes, respectively. The graphics were created with BioRender.com. (B) Plating 475 efficiency of the SH1000 derivatives $\Delta pbp1$, $\Delta pbp1 mecA^+$, $\Delta pbp1 rpoB^*$ and $\Delta pbp1 mecA^+$ rpoB* in the absence of IPTG. Plating efficiency values were compared with the control groups 476 grown in the presence of IPTG. Data were compared using a one-way ANOVA with Dunnett's 477 478 multiple comparison test (ns, not significant, $P \ge 0.05$), $n \ge 3$). Data represent the mean \pm SD.



481 Analysis of the role of PBP2 in growth. (A) Schematic representation of the *pbp2* genetic 482 constructs. An ectopic copy of *pbp2* was placed under the control of the Pspac promoter at the 483 SH1000 lipase (*geh*) locus, while the gene in the native *pbp2* locus was deleted (marked with 484 *tet*). In *mecA*⁺ *pbp2* and *mecA*⁺ *rpoB*^{*} *pbp2* a copy of a *mecA* gene expressed from its native

⁴⁸⁰ Fig. S8.

485 promoter was located at the *lysA* locus. In *rpoB* pbp2* and *mecA⁺ rpoB* pbp2* the *rpoB* gene has a point mutation which results in a single amino acid change (H929Q) in the RNA 486 polymerase β subunit (*rpoB*^{*}). The graphics were created with BioRender.com. (**B**) Plating 487 488 efficiency of the derivatives *pbp2*, *mecA*⁺ *pbp2*, *rpoB*^{*} *pbp2* and *mecA*⁺ *rpoB*^{*} *pbp2* grown in the absence of IPTG. Plating efficiency values were compared with the control groups grown 489 in the presence of IPTG. Data represent the mean \pm SD. (C) Growth curves of SH1000 490 491 derivatives *pbp2*, *mecA*⁺ *pbp2*, *rpoB*^{*} *pbp2*, and *mecA*⁺ *rpoB*^{*} *pbp2* grown in the presence or 492 absence of IPTG (+ IPTG and - IPTG, respectively). Data represent the mean \pm SD. Error bars 493 that are smaller than the data point symbols are not shown. (D) Immunoblots, analysed using 494 anti-PBP2 antibody, of whole cell lysates of SH1000 (wt) and mecA⁺ rpoB^{*}, rpoB^{*}, pbp2, 495 $mecA^+$ pbp2, $rpoB^*$ pbp2 and $mecA^+$ rpoB* pbp2 derivatives grown in the presence (+) or absence (-) of IPTG for 4 h. mecA⁺ pbp2* and mecA⁺ rpoB* pbp2* were grown in the presence 496 of IPTG throughout. Expected PBP2 and PBP2* sizes = 80 kDa are indicated (blue arrowhead). 497 (E) BocillinFL gel-based analysis of PBPs in SH1000 and $mecA^+$ $rpoB^*$, $rpoB^*$, pbp2, $mecA^+$ 498 *pbp2*, *rpoB*^{*} *pbp2*, and *mecA*⁺ *rpoB*^{*} *pbp2* grown in the presence (+) or absence (-) of IPTG 499 for 4 h. $mecA^+$ pbp2* and $mecA^+$ rpoB* pbp2* were grown in the presence of IPTG throughout. 500 501 Data are representative of two (D and E) and three (B and C) independent biological 502 experiments.

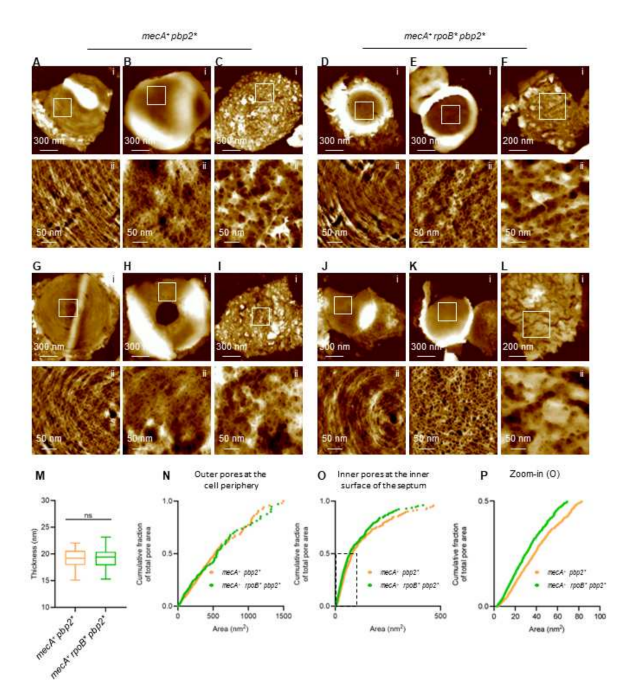






Role of PBP2 in cell morphology. (A-F) Fluorescence microscopy images of the SH1000 derivatives pbp2 (A), $mecA^+ pbp2$ (B), $rpoB^* pbp2$ (C) and $mecA^+ rpoB^* pbp2$ (D) grown in

507 the presence (+ IPTG) or absence (- IPTG) of the inducer for 4 h. mecA⁺ pbp2*(E) and mecA⁺ rpoB* pbp2* (F) were grown in the presence of IPTG at all times. All strains were incubated 508 for 5 min with ADA-DA clicked to Atto488 to show nascent PG, and counter labelled with 509 510 NHS-ester Alexa Fluor 555 to image the cell wall. Images are average intensity projections of z stacks. Cells with dispersed, apparent ADA-DA incorporation (yellow arrowheads), no ADA-511 DA incorporation (blue arrowheads) or mislocalized ADA-DA incorporation (white 512 513 arrowheads) are examples of cells that were classified as abnormal in panel G. (G) Quantification of cellular phenotypes based on ADA-DA incorporation in A-F. From left to 514 right, *n* = 247, 235, 261, 288, 237, 273, 299, 309, 278 and 305. (**H**) Cell volumes of strains in 515 516 A-F measured by fluorescence microscopy after NHS-ester Alexa Fluor 555 labelling. Number 517 of cells analysed for each sample was $n \ge 300$. Data are representative of three independent 518 biological experiments.



520 Fig. S10.

Surface-dependent nanoscale architecture of $mecA^+$ $pbp2^*$ and $mecA^+$ $rpoB^*$ $pbp2^*$ revealed 521 by AFM. AFM topographic images of the outer surface of the septum (A), the inner surface of 522 the septum (**B**), and open mesh structure of the cell periphery (**C**) associated with the cell wall 523 of mecA⁺ pbp2*. (i) Low-resolution AFM images and (ii) corresponding higher-resolution 524 525 images of the region indicated by the white boxes in (i). Topographical height (z) range for (i) are 100 nm, 330 nm, and 85 nm and for (ii) are 11 nm, 16 nm, and 45 nm respectively. (D-F) 526 AFM images of the same cell wall locations for $mecA^+ rpoB^* pbp2^*$; (i) and (ii) as above. The 527 height scales for (i) are 200 nm, 140 nm, and 130 nm. The height scales for (ii) are 16 nm, 9 528 nm, and 55 nm respectively. (G-I) AFM images of the PG structures associated with the 529 following locations on the cell wall of $mecA^+ pbp2^*$; (G) AFM images of the outer surface of 530 531 the septum, (H) inner surface of an incomplete septum, and (I) open mesh structure of the cell

- periphery; (i) and (ii) as above. Topographical heights (z) for (i) are 150 nm, 210 nm, and 85
- 533 nm. Topographical heights (z) for (ii) are 11 nm, 22 nm, and 51 nm respectively. (J-L) AFM 524 increase $\int PC$ structures of the energy location in P^* and 2^* (i) and (ii)
- images of PG structures of the same locations in $mecA^+$ $rpoB^*$ $pbp2^*$; (i) and (ii) as above. Topographical heights (z) for (i) are 140 nm, 200 nm, and 130 nm and for (ii) are 12 nm, 14
- Topographical heights (z) for (i) are 140 nm, 200 nm, and 130 nm and for (ii) are 12 nm, 14 nm, and 70 nm respectively. (**M**), Plot of the measured thickness of dehydrated sacculi of
- min, and 70 min respectively. (W), 1 of the measured interfiess of denythated saccum of $mecA^+$ pbp2* and $mecA^+$ rpoB* pbp2* respectively. The number of independent fragments
- 538 measured for each strain was 20. Data were analysed using the Mann-Whitney non-parametric
- statistical test (ns, not significant = 0.7180). (N-O) Cumulative fraction of total pore area as a
- 540 function of the area of the pores distributed across the open mesh surface of the cell wall
- 541 periphery surface (N) and the inner surface (O) of the septum of $mecA^+$ $pbp2^*$ and $mecA^+$
- 542 $rpoB^* pbp2^*$. (P) Zoomed-in plot of the region highlighted by the dashed box in (O). Data are
- 543 representative of two independent biological repeats and five AFM independent images.

Strain	Oxacillin MIC (µg ml ⁻¹)	Methicillin MIC (µg ml ⁻¹)
SH1000 (SJF682)	≤ 0.25	0.5
<i>mecA</i> ⁺ (SJF4996)	2	4
<i>mecA</i> ⁺ <i>rpoB</i> * (SJF5003)	> 256	> 256
COL (SJF315)	> 256	> 256
Mu50 (SJF5041)	> 256	> 256
MRSA252 (SJF4821)	> 256	> 256
TW20 (SJF6101)	> 256	> 256
USA300 (SJF4703)	0.75	1-2
USA300 (HL) (SJF6109)	> 256	64-128
EMRSA 15 (SJF6025)	16-24	64
EMRSA15 (HL) (SJF6110)	> 256	> 256
<i>geh::mecA</i> ⁺ (SJF5324)	4-8	-
<i>geh::mecA</i> ⁺ <i>rpoB</i> * (SJF5323)	> 256	-
$mecA^+$ lytH (SJF5461)	12	-
$mecA^+ gdpP$ (SJF5464)	≤ 2	-
<i>mecA</i> ⁺ <i>pde2</i> (SJF5460)	3	-
<i>mecA</i> ⁺ <i>rel</i> * (SJF5463)	> 256	-
<i>mecA</i> ⁺ <i>clpP</i> (SJF5459)	12-16	-
$mecA^+ clpX$ (SJF5462)	6-8	-
<i>rpoB* (</i> SJF5010)	≤ 0.25	-
<i>lytH</i> (SJF5455)	≤ 0.5	-
<i>gdpP</i> (SJF5025)	≤ 0.5	-
<i>pde2</i> (SJF5454)	≤ 0.125	-
rel* (SJF5457)	≤ 0.5	-
<i>clpP</i> (SJF5453)	0.25	-
<i>clpX</i> (SJF5456)	0.25-0.38	-

Table S1. 545

- Oxacillin and Methicillin MICs for S. aureus strains. The MICs for oxacillin and methicillin 546 were determined using E-test strips or the microdilution method (respectively) in triplicate as described in Materials and Methods. -, Not determined. 547
- 548

Strain	Genotype and Markers	Source
Escherichia coli		
NEB5a	fhuA2 (argF-lacZ)U169 phoA glnV44 80 (lacZ)M15 gyrA96 recA1 relA1 endA1 thi-1 hsdR17	New England Biolabs
Staphylococcus aureus		
SH1000	Functional <i>rsbU</i> ⁺ derivative of <i>S. aureus</i> 8325-4	(50)
COL	Healthcare acquired MRSA (HA- MRSA)	(51)
Mu50	HA-MRSA	(19)
MRSA252	HA-MRSA	(18)
TW20	HA-MRSA	(17)
USA300	Community-acquired MRSA (CA-MRSA)	(20)
USA300 (HL) (SJF6109)	High-level oxacillin resistant derivative of USA300	This study
EMRSA 15	CA-MRSA	(3)
EMRSA15 (HL) (SJF6110)	High-level oxacillin resistant derivative of EMRSA15	This study
VF17	SH1000 pGL485 (<i>lac1</i>); Cm ^R	(34)
RN4220	Restriction deficient transformation recipient	(52)

CYL316	RN4220 pCL112 Δ 19; Cm ^R	(53)
SJF4924	SH1000 geh::Pspac-pbp2, Kan ^R	This study
SJF5046	SH1000 <i>lysA::mecA rpoB</i> ^{H929Qkan} ; Ery ^R , Kan ^R	(14)
<i>pbp1</i> *(SJF4656)	SH1000 geh::Pspac-pbp1 pbp1::pbp1* lacI; Tet ^R , Cm ^R	(8)
<i>pbp1* mecA</i> ⁺ (SJF5223)	SH1000 geh::Pspac-pbp1 pbp1::pbp1* lacI lysA::mecA; Tet ^R , Cm ^R , Ery ^R	(8)
<i>pbp1*rpoB*</i> (SJF5306)	SH1000 geh::Pspac-pbp1 pbp1::pbp1* lacI rpoB ^{H929Qkan} ; Tet ^R , Cm ^R , Kan ^R	This study
<i>pbp1* mecA⁺ rpoB*</i> (SJF5226)	SH1000 geh::Pspac-pbp1 pbp1::pbp1* lacI lysA::mecA rpoB ^{H929Qkan} ; Tet ^R , Cm ^R , Ery ^R , Kan ^R	(8)
Δ <i>pbp1</i> (SJF5106)	SH1000 <i>geh::Pspac-pbp1</i> Δ <i>pbp1 lacI</i> ; Tet ^R , Cm ^R	(8)
$\Delta pbp1 mecA^+$ (SJF5224)	SH1000 <i>geh::Pspac-pbp1</i> Δ <i>pbp1 lacI</i> <i>lysA::mecA</i> ; Tet ^R , Cm ^R , Ery ^R	(8)
Δ <i>pbp1 rpoB</i> * (SJF5305)	SH1000 geh::Pspac-pbp1 Δpbp1 lacI rpoB ^{H929Qkan} ; Tet ^R , Cm ^R , Kan ^R	This study
$\Delta pbp1 mecA^+ rpoB^*$ (SJF5227)	SH1000 geh::Pspac-pbp1 Δpbp1 lacI lysA::mecA rpoB ^{H929Qkan} ; Tet ^R , Cm ^R , Ery ^R , Kan ^R	(8)
<i>pbp2</i> (SJF5630)	SH1000 <i>geh::Pspac-pbp2 pbp2::tet</i> <i>lacI</i> ; Tet ^R , Kan ^R , Cm ^R	This study

<i>mecA</i> ⁺ (SJF4996)	SH1000 <i>lysA::mecA</i> ; Ery ^R	(14)
$mecA^+ rpoB^*$ (SJF5003)	SH1000 <i>lysA-mecA rpoB</i> ^{H929Q} ; Ery ^R	(14)
<i>rpoB</i> * (SJF5010)	SH1000 <i>lysA::kan rpoB</i> ^{H929Q} ; Kan ^R ,	(14)
<i>mecA</i> ⁺ <i>pbp2</i> (SJF5663)	SH1000 <i>lysA::mecA geh::Pspac-pbp2</i> <i>pbp2::tet lacI</i> ; Ery ^R , Kan ^R , Tet ^R , Cm ^R	This study
<i>mecA</i> ⁺ <i>rpoB</i> * <i>pbp2</i> (SJF5674)	SH1000 <i>lysA::mecA rpoB</i> ^{H929Q} <i>geh::Pspac-pbp2 pbp2::tet lacI</i> ; Ery ^R , Kan ^R , Tet ^R , Cm ^R	This study
GMSA015	SH1000 <i>lysA::ery</i> ; Ery ^R	(54)
<i>rpoB* pbp2</i> (SJF5690)	SH1000 <i>lysA::ery rpoB</i> ^{H929Q} <i>geh::Pspac-pbp2 pbp2::tet lacI</i> ; Ery ^R , Kan ^R , Tet ^R , Cm ^R	This study
<i>mecA</i> ⁺ <i>pbp2</i> * (SJF5807)	SH1000 <i>lysA::mecA geh::Pspac-pbp2* pbp2::tet lacI</i> ; Ery ^R , Kan ^R , Tet ^R , Cm ^R	This study
<i>mecA</i> ⁺ <i>rpoB</i> * <i>pbp2</i> * (SJF5809)	SH1000 <i>lysA::pmecA rpoB^{H929Q}</i> <i>geh::Pspac-pbp2*(TP-) pbp2::tet lacI</i> ; Ery ^R , Kan ^R , Tet ^R , Cm ^R	This study
<i>geh::mecA</i> ⁺ <i>rpoB</i> * (SJF5323)	SH1000 <i>geh::mecA lysA::tet</i> <i>rpoB</i> ^{H929Q} ; Kan ^R , Tet ^R	(14)
<i>geh::mecA</i> ⁺ (SJF5324)	SH1000 geh::mecA lysA::tet; Kan ^R , Tet ^R	(14)
NE1369	JE2 <i>lytH::Tn</i> ; Ery ^R	(55)
<i>lytH</i> (SJF5455)	SH1000 <i>lytH::Tn</i> ; Ery ^R	This study

$mecA^+$ lytH (SJF5461)	SH1000 geh::mecA lysA::tet lytH::Tn; Kan ^R , Tet ^R , Ery ^R	This study
ANG1959	SEJ1 gdpP::kan	(56)
<i>gdpP</i> (SJF5025)	SH1000 gdpP::kan	This study
$mecA^+ gdpP$ (SJF5464)	SH1000 <i>lysA::mecA gdpP::kan</i> ; Ery ^R Kan ^R ,	This study
NE1208	JE2 <i>pde2::Tn</i> ; Ery ^R	(55)
pde2 (SJF5454)	SH1000 <i>pde2::Tn</i> ; Ery ^R	This study
<i>mecA</i> ⁺ <i>pde2</i> (SJF5460)	SH1000 geh::mecA lysA::tet pde2::Tn; Kan ^R , Tet ^R , Ery ^R	This study
NE1714	JE2 <i>rel::Tn</i> ; Ery ^R	(55)
rel (SJF5457)	SH1000 <i>rel::Tn</i> ; Ery ^R	This study
$mecA^+$ rel (SJF5463)	SH1000 geh::mecA lysA::tet rel::Tn; Kan ^R , Tet ^R , Ery ^R	This study
<i>pbp1* rel</i> (SJF5513)	SH1000 geh::Pspac-pbp1 pbp1::pbp1* lacI rel::Tn; Tet ^R , Cm ^R , Ery ^R	This study
<i>clpP</i> SJF5453)	SH1000 <i>clpP::Tn; Ery</i> ^R	This study
<i>mecA</i> ⁺ <i>clpP</i> (SJF5459)	SH1000 <i>geh::mecA, clpP::Tn;</i> Ery ^R Kan ^R	This study
<i>clpX</i> (SJF5456)	SH1000 <i>clpX::Tn; Ery^R</i>	This study
$mecA^+ clpX$ (SJF5462)	SH1000 <i>geh::mecA</i> , <i>clpX::Tn</i> ; Ery ^R Kan ^R	This study

NE912	JE2 <i>clpP::Tn; Ery</i> ^R	(55)
SJF1704	8325-4 <i>clpX::Tn; Ery</i> ^{<i>R</i>}	Lab stock

550 **Table S2.**

551 Strains used in this study.

552	Cm ^R – chloramphenicol,	, Ery ^R – erythromycin, 1	Kan ^R – kanamycin, Tet ^R	- tetracycline
-----	------------------------------------	--------------------------------------	--	----------------

Name	Characteristics	Source
pCQ11- FtsZ-SNAP	pCQ11 derivative containing <i>ftsZ-snap</i> under Pspac; Amp ^R , Ery ^R	(57)
pGM074	pKASBAR-kan derivative with <i>ezrA-psmorange</i> ; Amp ^R , Tet ^R	(54)
pKB-Pspac- pbp2	pGM074 derivative containing P <i>spac</i> , <i>S. aureus pbp2</i> RBS and full length <i>pbp2</i> ; Amp ^R , Kan ^R	This study
pKB-Pspac- pbp2*	pGM074 derivative containing Pspac, S. aureus pbp2 RBS and full length pbp2 with inactivated TP (pbp2*); Amp ^R , Kan ^R	This study
pMAD	<i>E. coli-S. aureus</i> shuttle vector with temperature-sensitive origin of replication in <i>S. aureus</i> and constitutively produced thermostable β -galactosidase encoded by <i>bgaB</i> ; Amp ^R , Ery ^R	(58)
pMAD- Δ <i>pbp2</i>	pMAD containing a deletion cassette for <i>S. aureus pbp2</i> ; Amp ^R , Ery ^R . Tet ^R	This study
pAISH1	TetR derivative of pMUTIN4; Amp ^R ; Tet ^R	(54)

553 **Table S3.**

554 Plasmids used in this study. Amp^{R} – ampicillin, Ery^{R} – erythromycin, Kan^{R} – kanamycin, Tet^{R}

555 – tetracycline

Name	Sequence 5'–3'
pCQ11-pbp2-F	agaaggagatatacatatggagtgaggaccgcgtatgac
pCQ11-pbp2-R	atttattatgcatttagaataggttagttgaatatacctgttaatccac
pKB-pbp2-R	cagctatgaccatgattacgttagttgaatatacctgttaatccac
pKB-Pspac-pbp1-F	cctttttttgccccgggatccgcaaaaagttgttgactttatc
pbp2TP-F	ccctactggtggatctttaaaacc
pbp2TP-R	tgaggatctgttgcttgg
pMAD-Δpbp2-F	ccatggtacccgggagctcgacgatgaaaatacttttaatctaataaaatc
pbp2-up-R	acactatctgcaggtcatacgcggtcctcac
tetR-pbp2-F	gaccgcgtatgacctgcagatagtgtacgtaaaaag
tetR-pbp2-R	tatgttgagtggactctctccccaaagttgatc
pbp2-down-F	ctttgggagagagtccactcaacataaaatcctc
pMAD-Δpbp2-R	ccatggtacccgggagctcgacgatgaaaatacttttaatctaataaaatc

556 Table S4.

557 Oligonucleotides used in this study.