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# 1 Impaired alanine transport or exposure to D-cycloserine increases the susceptibility of

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- 3
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- 22
- **Running title:** Re-sensitization of MRSA to  $\beta$ -lactams

**Abstract.** Prolonging the clinical effectiveness of  $\beta$ -lactams, which remain first-line antibiotics 24 for many infections, is an important part of efforts to address antimicrobial resistance. We 25 report here that inactivation of the predicted D-cycloserine (DCS) transporter gene cycA re-26 27 sensitized MRSA to  $\beta$ -lactam antibiotics. The cycA mutation also resulted in hypersusceptibility to DCS, an alanine analogue antibiotic that inhibits alanine racemase and D-28 29 alanine ligase required for D-alanine incorporation into cell wall peptidoglycan (PG). Alanine transport was impaired in the cycA mutant and this correlated with increased susceptibility 30 to oxacillin and DCS. The cycA mutation or exposure to DCS were both associated with the 31 accumulation of muropeptides with tripeptide stems lacking the terminal D-ala-D-ala and 32 33 reduced PG crosslinking, prompting us to investigate synergism between  $\beta$ -lactams and DCS. 34 DCS re-sensitised MRSA to  $\beta$ -lactams *in vitro* and significantly enhanced MRSA eradication by oxacillin in a mouse bacteraemia model. These findings reveal alanine transport as a new 35 therapeutic target to enhance the susceptibility of MRSA to  $\beta$ -lactam antibiotics. 36

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#### 41 Introduction

42 Whilst many bacteria can exhibit resistance to select antimicrobials, isolates of the human 43 pathogen Staphylococcus aureus can express resistance to all licensed anti-staphylococcal drugs. This results in significant morbidity and mortality, with up to 20% of patients with 44 systemic methicillin resistant S. aureus (MRSA) infections dying, despite receiving treatment 45 46 with anti-staphylococcal drugs [1]. As part of our efforts to identify improved therapeutic approaches for MRSA infections, we recently described the novel use of  $\beta$ -lactam antibiotics 47 to attenuate the virulence of MRSA-induced invasive pneumonia and sepsis [2]. We 48 demonstrated that oxacillin-induced repression of the Agr quorum-sensing system and 49 50 altered cell wall architecture resulted in downregulated toxin production and increased MRSA killing by phagocytic cells, respectively [2]. Supporting this in vitro data, a randomised 51 controlled trial involving 60 patients showed that the  $\beta$ -lactam antibiotic flucloxacillin in 52 combination with vancomycin shortened the duration of MRSA bacteraemia from 3 days to 53 1.9 days [3, 4]. 54

Because expression of methicillin resistance in *S. aureus* impacts fitness and virulence and is 55 a regulated phenotype, further therapeutic interventions may also be possible. The 56 complexity of the methicillin resistance phenotype is evident among clinical isolates of MRSA, 57 which express either low-level, heterogeneous (HeR) or homogeneous, high-level methicillin 58 resistance (HoR) [5-7]. Exposure of HeR isolates to  $\beta$ -lactam antibiotics induces expression of 59 60 mecA, which encodes the alternative penicillin binding protein 2a (PBP2a) and can select for mutations in accessory genes resulting in a HoR phenotype, including mutations that affect 61 the stringent response and c-di-AMP signalling [8-12]. Because accessory genes can influence 62 the expression of methicillin resistance in MRSA, targeting the pathways associated with such 63 genes may identify new ways to increase the susceptibility of MRSA to  $\beta$ -lactams. To pursue 64 this, we performed a forward genetic screen to identify loci that impact the expression of 65 resistance to  $\beta$ -lactam antibiotics in MRSA. Using the Nebraska Transposon Mutant Library, 66 which comprises 1,952 sequence-defined transposon insertion mutants [13], inactivation of 67 a putative amino acid permease gene, cycA, was found to reduce resistance to cefoxitin, the 68 β-lactam drug recommended by the Clinical and Laboratory Standards Institute for measuring 69 mecA-mediated methicillin resistance in MRSA isolates. Amino acid transport and 70

susceptibility to oxacillin and D-cycloserine (DCS) were compared in the wild-type and cycA 71 mutant grown in chemically defined media (CDM), CDM supplemented with glucose (CDMG) 72 and other complex media. The activity of DCS and  $\beta$ -lactams, alone and in combination, 73 74 against MRSA was measured in vitro and in a mouse model of bacteraemia. Peptidoglycan 75 analysis was performed to compare the impact of the cycA mutation or exposure to DCS on cell wall structure and crosslinking. Our experiments suggest that therapeutic strategies 76 targeting alanine transport, which was required for resistance to  $\beta$ -lactams and DCS, and a re-77 evaluation of DCS may be important as part of efforts to restore the efficacy of  $\beta$ -lactam 78 antibiotics against MRSA. 79

#### 80 Results

Mutation of cycA increases the susceptibility of MRSA to β-lactam antibiotics and D-81 cycloserine. To identify new ways of controlling expression of methicillin resistance, we 82 83 sought to identify novel mutations involved in this phenotype. An unbiased screen of the NTML to identify mutants with increased susceptibility to cefoxitin identified NE810 84 (SAUSA300 1642) (Fig. S1A), which also exhibited a >128-fold increase in susceptibility to 85 oxacillin (Fig. S1B). NE810 was previously identified among several NTML mutants reported 86 to be more susceptible to amoxicillin [14], but was not investigated further. Expression of 87 88 mecA was not affected in NE810 (Fig. S1C) and genome sequence analysis revealed an intact 89 SCCmec element and the absence of any other mutations. NE810 was successfully 90 complemented (Fig. S1B), and transduction of the SAUSA300 1642 allele into several MRSA strains from a number of clonal complexes and with different SCCmec types was also 91 92 accompanied by increased cefoxitin and oxacillin susceptibility (Table 1).

SAUSA300\_1642 is annotated as a serine/alanine/glycine transporter with homology to CycA 93 in *Mycobacterium tuberculosis* [15, 16], which influences D-cycloserine (DCS) susceptibility in 94 Mycobacteria [15, 16]. In contrast to the observations in Mycobacteria, our data showed that 95 NE810 and several unrelated MRSA strains carrying the cycA mutation were significantly more 96 susceptible to DCS than the wild type JE2 (Fig. S1D, Table 1). The cycA mutation also reduced 97 the DCS MIC of the MSSA strains 8325-4 and ATCC29213 from 32 to 4  $\mu$ g/ml. DCS inhibits 98 alanine racemase (Alr) that converts L-alanine to D-alanine and the Ddl D-alanyl:D-alanine 99 ligase [17]. A mutant in the putative *ddl* SAUSA300 2039 gene is not available in the NTML 100 library, suggesting that it may be essential. However, the *alr* mutant NE1713 was more 101 susceptible to cefoxitin (Fig. S2A; MIC=16µg/ml) and DCS (Fig. S2B; MIC <0.25µg/ml), 102 consistent with an important role for D-alanine in resistance to both antibiotics. 103

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105 **CycA is required for alanine transport and D-ala-D-ala incorporation into the peptidoglycan** 106 **stem peptide.** To investigate the role of CycA in amino acid transport, JE2 and NE810 were 107 grown for 8 h in chemically defined media containing 14mM glucose (CDMG) and amino acid 108 consumption in spent media was measured. Although no growth rate or yield difference was 109 noted between JE2 and NE810 in CDMG (Fig. 1A), alanine uptake by NE810 was significantly 110 impaired compared to JE2 (Fig. 1B). Utilisation of other amino acids by NE810 and JE2, 111 including serine and glycine, were similar (Fig. S3). Impaired alanine transport in the *cycA* 

mutant grown in CDMG correlated with increased susceptibility to oxacillin (1  $\mu$ g/ml) (Fig. 1C) and DCS (1  $\mu$ g/ml) (Fig. 1D). These data demonstrate for the first time that CycA in *S. aureus* is required for alanine transport.

Quantitative peptidoglycan compositional analysis was performed using UPLC analysis of 115 muramidase-digested muropeptide fragments extracted from exponential phase cultures of 116 117 JE2 and NE810 grown for 220 mins in TSB media (Fig. S4). The PG profile of the cycA mutant revealed a significant accumulation of tripeptides compared to wild-type JE2 (Fig. 2A,B), 118 which was associated with a significant reduction in crosslinking (Fig. 2C). In NE810, the dimer, 119 trimer and tetramer fractions were decreased, which was accompanied by a concomitant 120 increase in the monomer fraction (Fig. 2D). Consistent with this data, exposure of JE2 to DCS 121 8µg/ml was also associated with a similar accumulation in muropeptides with tripeptide 122 stems (Fig. 2B), reduced cross-linking (Fig. 2C), increased muropeptide monomers and 123 reduced dimers, trimers and tetramers (Fig. 2D). DCS had a strong dose-dependent effect on 124 the accumulation of muropeptides with tripeptide stems, reduced cross-linking and 125 126 accumulation of monomers (Fig. 2A-D). Sub-inhibitory (0.25 $\times$  MIC) and 4 $\times$  MIC DCS 127 concentrations, were previously shown to be associated with incorporation of an incomplete stem peptide (tripeptide) [17] and reduced D-ala-D-ala levels [18], respectively. These data 128 show that impaired D-ala incorporation in the cycA mutant or following exposure to DCS is 129 accompanied by reduced PG cross-linking and increased  $\beta$ -lactam susceptibility. 130

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Mutation of cycA or exposure to D-cycloserine increases the susceptibility of MRSA to  $\beta$ -132 **lactam antibiotics.** Previously reported synergy between DCS and  $\beta$ -lactam antibiotics [17, 133 19] suggests that impaired alanine uptake in the cycA mutant may have the same impact on 134 cell wall biosynthesis as DCS-mediated inhibition of Alr and Ddl. To further investigate this, 135 we compared the activity of DCS and  $\beta$ -lactam antibiotics, alone and in combination, against 136 JE2 and NE810. Checkerboard microdilution assay fractional inhibitory concentration indices 137 ( $\Sigma$ FICs  $\leq$ 0.5) revealed synergy between DCS and several licensed  $\beta$ -lactam antibiotics with 138 139 different PBP selectivity against JE2 and USA300 FPR3757 (Table 1). Oxacillin and nafcillin 140 were not included in checkerboard assays because measurement of their MICs involves supplementing the media with 2% NaCl, which distorts the MIC of DCS (data not shown). 141 Using the MRSA strains JE2, USA300, DAR173, DAR22, DAR169 and their corresponding cycA 142

143 mutants, the kinetics of killing by DCS, oxacillin and cefoxitin, alone and in combination was

measured over 24h using antibiotic concentrations corresponding to  $0.125 \times$ ,  $0.25 \times$  and  $0.5 \times$ 144 MICs. Recovery of growth in media supplemented with oxacillin or cefoxitin alone was evident 145 after 8 h (Fig. 3), reflecting the selection and expansion of HoR mutants as described 146 147 previously [2, 18, 20]. Recovery of growth in cultures exposed to DCS alone was also evident 148 (Fig. 3), which may correlate with our observation that mutants resistant to DCS (on BHI agar supplemented with 128  $\mu$ g/ml DCS) arise at a rate of approximately 5.5  $\times$  10<sup>-8</sup> per cell per 149 150 generation. Using combinations of DCS and oxacillin or cefoxitin at 0.125× MIC did not achieve a  $\geq$ 2 log<sup>10</sup> reduction in the number of CFU/ml (data not shown). However, at 0.5× MIC for 151 strains JE2, USA300, DAR173 and DAR22, DCS (16 µg/ml)/oxacillin (32 µg/ml) and DCS (16 152  $\mu$ g/ml)/cefoxitin (32  $\mu$ g/ml) combinations achieved a  $\geq$ 5 log<sup>10</sup> reduction in the number of 153 CFU/ml compared to oxacillin, cefoxitin or DCS alone (Fig. 3). For strain DAR169, DCS/β-lactam 154 combinations at 0.25× MIC was sufficient to achieve a  $\geq$ 5 log<sup>10</sup> reduction in CFUs recovered 155 compared to the individual antibiotics (Fig. 3). DCS/ $\beta$ -lactam combinations at 0.5× MIC were 156 also able to achieve  $\geq 5 \log^{10}$  reduction in the number of CFU/ml against the methicillin 157 resistant *S. epidermidis* (MRSE) strain RP62A [21] compared to either antibiotic alone (Fig. S5). 158 Checkerboard experiments with fourteen MRSA strains and MRSE strain RP62A further 159 revealed synergy ( $\Sigma$ FICs  $\leq$ 0.5) between DCS and a range of  $\beta$ -lactam antibiotics with different 160 penicillin binding protein (PBP) specificity, namely cefoxitin (PBP4), cefaclor (PBP3), 161 162 cefotaxime (PBP2), piperacillin-tazobactin (PBP3/ $\beta$ -lactamase inhibitor) and imipenem (PBP1) 163 (Table 1).

This synergy appears to be specific to  $\beta$ -lactams and no synergy ( $\Sigma$ FICs > 0.5) was measured 164 between DCS and several antibiotics that are used topically or systemically for the 165 decolonization or treatment of patients colonized/infected with S. aureus or MRSA 166 (clindamycin, trimethoprim, mupirocin, ciprofloxacin), or several antibiotics to which S. 167 *aureus* isolates commonly exhibit resistance (tobramycin, kanamycin and spectinomycin) 168 (Table S1). Furthermore the cycA mutation had no impact on susceptibility to any of these 169 non- $\beta$ -lactam antibiotics (apart from *ermB*-encoded clindamycin resistance on the 170 171 transposon).

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173 Combination therapy with DCS and oxacillin significantly reduces the bacterial burden in 174 the kidneys and spleen of mice infected with MRSA. The virulence of the NE810 mutant and 175 the therapeutic potential of oxacillin in combination with DCS in the treatment of MRSA

infections were assessed in mice. Treatment with oxacillin or DCS alone significantly reduced 176 the number of CFUs recovered from the kidneys of mice infected with JE2 (Fig. 4). 177 Furthermore the oxacillin/DCS combination was significantly more effective than either 178 179 antibiotic alone and the combination was equally effective in reducing the bacterial burden 180 in the kidneys of animals infected with JE2 or NE810 when compared to no treatment (p≤0.0001) (Fig. 4) demonstrating the capacity of DCS to significantly potentiate the activity 181 of  $\beta$ -lactam antibiotics against MRSA under *in vivo* conditions. Unexpectedly, oxacillin- or DCS-182 mediated eradication of NE810 infections in the kidneys was similar to JE2 (Fig. 4). In the 183 spleen, only oxacillin/DCS combination treatment was associated with a significant reduction 184 in the number of CFUs recovered from mice infected with JE2 or NE810 (Fig. S6). 185

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Alanine transport and resistance to oxacillin and DCS in chemically defined medium are not 187 dependent on cycA. The failure of oxacillin or DCS treatment to enhance the eradication of 188 189 NE810 infections in the mouse bacteraemia model prompted us to further characterise the growth conditions used for the *in vitro* antibiotic susceptibility assays. Specifically we 190 investigated the role of glucose, which we previously reported to increase the growth 191 requirement for amino acids [22], and which we reasoned may be important for CycA-192 dependent alanine transport. Growth of JE2 and NE810 was similar in CDM lacking glucose 193 (Fig. 5A) and uptake of alanine (Fig. 5B) and other amino acids (Fig. S7) was unchanged in 194 NE810 compared to JE2. Furthermore NE810 and JE2 grew equally well in CDM supplemented 195 with oxacillin and DSC (Fig. 5C and D). These data explain in part why the cycA mutant does 196 not exhibit increased  $\beta$ -lactam and DCS susceptibility in the mouse bacteraemia model and 197 further reveal the strong correlation between alanine transport and susceptibility to oxacillin 198 199 and DCS.

#### 200 Discussion

The exploitation of antibiotic re-purposing as part of concerted efforts to address the 201 antimicrobial resistance crisis has been hampered by a lack of mechanistic data to explain 202 203 demonstrated therapeutic potential and the perception that studies attempting to identify 204 new uses for existing drugs are not hypothesis-driven. In this study, we revealed that CycA was required for full expression of resistance to  $\beta$ -lactam antibiotics and DCS. Loss of function 205 of this putative alanine transporter significantly increased the susceptibility of MRSA to  $\beta$ -206 lactam antibiotics, an outcome that could be reproduced through exposure to DCS, which 207 targets the Alr and Ddl enzymes in the early steps of cell wall biosynthesis. 208

The potential of  $\beta$ -lactam/DCS combinations for treatment of MRSA infections follows a 209 recent report that DCS can also potentiate the activity of vancomycin against a laboratory-210 generated vancomycin highly-resistant S. aureus (VRSA) strain in vitro and in a silkworm 211 infection model [23]. The excellent safety profile of  $\beta$ -lactam antibiotics makes these drugs 212 particularly attractive as components of combination antimicrobial therapies. When used in 213 214 the treatment of tuberculosis DCS (trade name Seromycin, The Chao Centre) is typically administered orally in 250 mg tablets twice daily for up to two years. At this dosage, the DCS 215 concentration in blood serum is generally 25-30  $\mu$ g/ml, which is similar to the concentrations 216 used in our *in vitro* and *in vivo* experiments. The known neurological side effects associated 217 with DCS therapy [24, 25] mean that this antibiotic is unlikely to be considered for the 218 treatment of MRSA infections unless alternative therapeutic approaches have been 219 220 exhausted. Oxacillin/DCS combination therapy was significantly more effective than DCS or oxacillin alone over a 5-day therapeutic window suggesting that further studies on using DCS 221 to augment  $\beta$ -lactams as a treatment option for recalcitrant staphylococcal infections are 222 merited. 223

Mutation of *cycA* increases the susceptibility of MRSA to  $\beta$ -lactam antibiotics and results in hyper-susceptibility to D-cycloserine, whereas a *cycA* point mutation in *M. bovis* contributes, in part, to increased DCS resistance presumably by interfering with transport into the cell [16]. In *E. coli, cycA* mutations can also result in increased resistance or have no effect on DCS susceptibility depending on the growth media [26-30], suggesting that CycA is primarily important for DCS resistance under conditions when its contribution to amino acid transport

is also important. Our data showing that mutation of cycA was not associated with increased 230 DCS resistance strongly suggests that CycA has no role in uptake of this antibiotic in *S. aureus*. 231 Under growth conditions where CycA is required for alanine transport (in nutrient/glucose-232 233 replete media), mutation of cycA or DCS-exposure have similar effects on the structure of S. 234 aureus peptidoglycan (Fig. 8). Consistent with previous studies in S. aureus [17] and in M. tuberculosis [31], our studies showed a dose-dependent accumulation of muropeptides with 235 a tripeptide stem in MRSA exposed to DCS. The cycA mutation was also associated with the 236 increased accumulation of muropeptides with a tripeptide stem. These data indicate that a 237 238 reduced intracellular alanine pool or inhibition of Alr and Ddl is associated with reduced Dala-D-ala incorporation into the PG stem peptide. The increased accumulation of tripeptides 239 240 in turn interferes with normal PBP transpeptidase activity and offers a plausible explanation 241 for increased susceptibility to  $\beta$ -lactam antibiotics. The importance of the terminal stem peptide D-ala-D-ala for  $\beta$ -lactam resistance has previously been reported. Mutation of the 242 *murF*-encoded ligase, which catalyses of the D-ala-D-ala into the stem peptide also increased 243  $\beta$ -lactam (but not DCS) susceptibility [32, 33]. Similarly growth of a HoR MRSA strain in media 244 supplemented with high concentrations of glycine was accompanied by replacement of the 245 D-ala-D-ala with D-ala-gly and decreased methicillin resistance [34]. 246

Impaired uptake of alanine in CDMG correlated with increased susceptibility to oxacillin and 247 DCS, suggesting that alanine utilisation via CycA is important to make D-alanine available for 248 cell wall biosynthesis and consequently resistance to  $\beta$ -lactams. Consistent with this, NE810 249 also exhibited increased oxacillin susceptibility in BHI, TSB and MH media. However no change 250 in alanine transport or susceptibility to oxacillin and DCS was measured in CDM lacking 251 glucose, which may explain the failure of oxacillin and DCS to more efficiently eradicate NE810 252 infections in the mouse bacteraemia model. The availability of nutrients such as glucose and 253 amino acids varies in different niches colonised by S. aureus during infection ranging from 254 glucose-rich in organs such as the liver [35], to glucose-depleted in established abscesses [36]. 255 In turn this impacts the role of amino acids as carbon sources [22, 37], and potentially the 256 activity of CycA in alanine transport and  $\beta$ -lactam susceptibility. Furthermore, normal alanine 257 transport in the cycA mutant grown in CDM indicates that an alternative alanine transport 258 mechanism(s) may be active under these growth conditions (Fig. 6). Identification of this 259 alternative alanine permease may be important in the development of therapeutic strategies 260

- targeting alanine transport to increase  $\beta$ -lactam susceptibility in MRSA, while elucidation of
- the role of glucose in the control of alanine transport should provide new insights into  $\beta$ -
- lactam resistance.

#### 264 Materials and Methods

Bacterial strains, growth conditions and antimicrobial susceptibility testing. Bacterial strains
(Table S2) were grown in Luria Bertoni (LB), brain heart infusion (BHI), Mueller Hinton (MH),
nutrient, sheep blood BHI, chemically defined media (CDM) [38] or CDM 14mM glucose
(CDMG) [38].

Minimum inhibitory concentrations (MICs) were determined in accordance with CLSI guidelines using plate and broth dilution assays in MH, or MH 2% NaCl for oxacillin and nafcillin. Oxacillin MICs were also measured using E-tests (Oxoid) on MH 2% NaCl. Quality control strains ATCC29213 and ATCC25923 were used for oxacillin and cefoxitin MIC assays, respectively.

274 **Identification of cefoxitin susceptible MRSA mutant NE810.** Cefoxitin (30µg) disks (Oxoid) were used to measure susceptibility of NTML mutants. The zone diameter for JE2 was 18mm 275 NE1868 (mecA::Em<sup>r</sup>) was >35mm and NE810 was 22mm. The cycA transposon insertion in 276 NE810 was verified by PCR using the primers NE810 Fwd and NE810 Rev (Table S3). Phage 277  $80\alpha$  was used to transduce the NE810 cycA allele into JE2 and other strains. Genome 278 sequencing was performed by MicrobesNG using the USA300 FPR3757 genome as a 279 reference. To complement NE810, cycA was amplified from JE2 on a 1608 bp fragment using 280 primers NE810F1 Fwd and NE810F1 Rev (Table S3) and cloned into pLI50 using the Clontech 281 In-fusion kit. 282

*mecA* transcription analysis. RT-qPCR was performed on a Roche LightCycler with primers mecA1\_Fwd and mecA1\_Rev for *mecA* and gyrB\_Fwd and gyrB\_Rev for *gyrB* (internal standard) (Table S3), as described previously [2]. Data presented are the average of three experiments with standard errors.

Amino acid transport studies. Amino acid analysis in spent media from cultures grown in CDM or CDMG was performed as described previously [22].

Analysis of peptidoglycan composition in NE810 and JE2 treated with D-cycloserine. Independent quadruplicate 50ml cultures were grown to  $A_{600}$ =0.5, dosed with DCS at 0, 8, 20 or 32 µg/ml for 100 mins, then harvested and resuspended in 5ml PBS (Fig. S4) before peptidoglycan was extracted as described previously [39]. Mass spectrometry was performed on a Waters XevoG2-XS QTof mass spectrometer. Structural characterization of muropeptides

was determined based on their MS data and MS/MS fragmentation pattern, matched with PG
 composition and structure reported previously [34, 40-42].

296 Antibiotic synergy analysis using the microdilution checkerboard assay. Antibiotic synergism 297 was measured using the checkerboard microdilution method in 96-well plates inoculated with 298  $5 \times 10^5$  CFU/ml. Growth or no growth was recorded after 24 h at 37°C. The fractional inhibitory 299 concentration index ( $\Sigma$ FIC) was calculated for each drug combination in triplicate experiments 300 with an FIC index ≤0.5 considered synergistic.

Kill curve assays. Overnight cultures adjusted to 10<sup>7</sup> CFU/ml were exposed to 0.125×, 0.25×, and 0.5× MIC of oxacillin, cefoxitin and DCS alone or in combination, and the number of colony forming units (CFU)/ml enumerated at 0, 2, 4, 8 and 24 h. Data is presented at the antibiotic concentrations where synergy was measured i.e. 0.5× MIC for JE2, USA300, DAR173, DAR22, DAR113, BH1CC, and RP62A, and 0.25× MIC for DAR169. Synergism was defined as a ≥2 log<sup>10</sup> decrease in the number of CFU/ml in cell suspensions exposed to DCS/βlactam combinations compared to the most effective individual drug after 8 h.

308 **Mouse infection experiments.** 6-8 week-old, age matched, outbred CD1 female mice (Charles 309 River, UK) were used in a non-lethal model of bacteremia. JE2 and NE810 cultures were grown to  $A_{600}$ =0.5 in BHI, washed in PBS, adjusted to 1×10<sup>8</sup> CFU/ml. Mice were infected 310 intravenously (via the tail vein) with  $5 \times 10^6$  CFU (n = 10 mice per group). The infections were 311 left untreated (PBS control) or treated with either 75mg oxacillin/Kg/12 hours, 30mg 312 DCS/Kg/12 hours or a combination of both (first antibiotic dose administered 16 hours post 313 infection), before being sacrificed after 5 days. Bacteria present in homogenised spleens and 314 kidneys recovered from the mice were enumerated on blood agar. 315

**Ethics Statement.** Mouse experiments were approved by the UK Home Office (License Number 40/3602) and the University of Liverpool Animal Welfare and Ethics Committee. This study was carried out in strict accordance with the UK Animals (Scientific Procedures) Act 1986. All efforts were made to minimize suffering.

**Statistical analysis**. Two-tailed Student's t-Tests and one-way ANOVA with Kruskal-Wallis test followed by Dunn's multiple comparisons test in the GraphPad Prism application (for the mouse infection experiments) were used to determine statistically significant differences in assays performed during this study. A *p* value <0.05 was deemed significant.

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326

# 327 Footnotes

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# 339 Figure Legends

Figure 1. Mutation of *cycA* impairs alanine uptake. A. Growth of JE2 and NE810 in chemically defined media supplemented with glucose (CDMG). Cell density was measured at  $A_{600}$ . B. Alanine consumption by JE2 and NE810 grown aerobically in CDMG. Residual amino acid was measured in spent media after 2, 4, 6 and 8 h growth. C and D. Growth of JE2 and NE810 cultures for 12 h in CDMG supplemented with 1 µg/ml oxacillin (C) or 1 µg/ml D-cycloserine (D). Cell density was measured at  $A_{600}$ .

- Figure 2. Mutation of cycA or D-cycloserine (DCS) treatment impacts peptidoglycan peptide 346 stem length and reduces cell wall crosslinking. A. Representative UV chromatograms of 347 peptidoglycan from wild-type JE2, NE810 and JE2 treated with increasing concentrations of 348 349 DCS (8, 20 and 32  $\mu$ g/ml). Muropeptides with tripeptide stems are numbered 1-3. The Proposed structures of the three muropeptides with tripeptide stems identified in NE810 and 350 351 DCS-treated JE2 cells. NAG, N-acetylglucosamine; NAM, N-acetylmuramic acid; L-Ala, Lalanine; D-Gln, D-glutamine; D-Glu, D-glutamic; L-Lys, L-lysine. The theoretical and observed 352 353 neutral masses determined by MS are indicated. **B.** Relative abundance of muropeptides with tripeptides in the stem. **C.** Relative crosslinking efficiency. **D.** Relative proportions of cell wall 354 muropeptide fractions based on oligomerization. All errors bars represent 95% confidence 355 interval (n = 4). Significant differences determined using Students t-test are denoted using 356 asterisks (\* p<0.05; \*\* p<0.01; \*\*\* p<0.001). 357
- Figure 3. In vitro kill curves for D-cycloserine (DCS), oxacillin and cefoxitin with JE2, USA300 358 FPR3757, DAR173, DAR22, DAR169 and their isogenic cycA mutants. Antibiotics at the 359 concentrations indicated ( $\mu$ g/ml) were added to suspensions of overnight bacterial cultures 360 adjusted to 10<sup>7</sup> CFU/ml in BHI (A<sub>600</sub>=0.05), incubated at 37°C and the number of CFU/ml 361 enumerated at 0, 2, 4, 8 and 24 h. The data presented are the mean of three independent 362 experiments, and standard error of the mean is shown. Antibiotic synergism was defined as a 363  $\geq 2 \log^{10}$  decrease in the number of CFU/ml in cell suspensions exposed to DCS/ $\beta$ -lactam 364 combinations compared to the most effective individual antibiotic alone. 365

366 Figure 4. Combination therapy with D-cycloserine and oxacillin significantly reduces the bacterial burden in the kidneys of mice infected with MRSA. The number of colony-forming 367 units (CFU) recovered from the kidneys of mice infected by tail vein injection with  $5 \times 10^{6}$  JE2 368 or NE810 (CycA) and left untreated or treated with 75mg of oxacillin (Ox)/kg, 30mg of DCS/kg 369 370 or a combination of both Ox and DCS delivered subcutaneously every 12 hours for 5 days. The first antibiotic dose was given 16 hours after infection. Significant differences determined 371 using one-way ANOVA with Kruskal-Wallis test followed by Dunn's multiple comparisons 372 test are denoted using asterisks (\*p≤0.05, \*\*p≤0.01, \*\*\*\*p≤0.0001). The limit of detection 373 374 (50 colonies) is indicated with a hashed red line.

Figure 5. Alanine transport and resistance to oxacillin and D-cycloserine in chemically
 defined medium are *cycA*-independent. A. Growth of JE2 and NE810 in chemically defined
 medium lacking glucose (CDM). Cell density was measured at A<sub>600</sub>. B. Alanine consumption by

- 378 JE2 and NE810 grown aerobically in CDM. Residual amino acid was measured in spent media
- after 2, 4, 6 and 8 h growth. **C and D.** Growth (cell density at  $A_{600}$ ) of JE2 and NE810 cultures
- for 12 h in CDM supplemented with 1  $\mu$ g/ml oxacillin (C) or 1  $\mu$ g/ml DCS (D).
- **Figure 6.** Proposed model depicting how impaired alanine transport associated with mutation
- of CycA or exposure to DCS can inhibit the D-alanine pathway for peptidoglycan biosynthesis
- leading to increased susceptibility to  $\beta$ -lactam antibiotics.

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**Table 1.** Antibacterial activity (minimum inhibitory concentrations, MIC) and drug synergy (fractional inhibitory concentration indices, ΣFIC) of D-cycloserine (DCS) and several β-lactam antibiotics with different PBP specificity, namely oxacillin (OX; PBP1, 2, 3), nafcillin (NAF; PBP1), cefoxitin (FOX; PBP4), cefaclor (CEC; PBP3), cefotaxime (CTX; PBP2), piperacillin-tazobactin (TZP; PBP3/β-lactamase inhibitor) and imipenem (IMP; PBP1), alone and in β-lactam combinations, against fourteen *S. aureus* strains and *S. epidermidis* RP62A.

Antibiotic $\rightarrow$	0//*		D00*	<b>50</b> //*	DCS/FOX**	050*	DCS/CEC**	<b>CT</b> \/*	DCS/CTX**	<b>T</b> 70*	DCS/TZP**		DCS/IPM**
↓ Strain	UX*	NAF*	DCS*	FUX*	(ΣFIC)***	CEC*	(ΣFIC)***	CIX*	(ΣFIC)***	IZP*	(ΣFIC)***	IPIVI*	(ΣFIC)***
JE2	64	32	32	64	8/8 (0.37)	64	8/2 (0.28)	64	8/4 (0.31)	32	8/0.5 (0.26)	1	ND
NE810 ( <i>cycA</i> )	0.25	0.5	2	8	ND	4	ND	8	ND	2	ND	0.125	ND
USA300	64	32	32	64	8/8 (0.37)	64	8/8 (0.37)	64	8/8 (0.37)	64	8/2 (0.28)	1	ND
USA300 cycA	0.25	1	2	8	ND	4	ND	8	ND	4	ND	0.125	ND
DAR173	128	128	32	256	4/32 (0.25)	128	8/4 (0.28)	512	8/4 (0.25)	128	4/4 (0.15)	64	4/2 (0.15)
DAR173 cycA	0.5	8	4	32	ND	16	ND	16	ND	4	ND	1	ND
DAR22	128	128	32	128	8/8 (0.31)	128	8/8 (0.31)	512	8/4 (0.25)	128	4/4 (0.15)	128	4/1 (0.13)
DAR22 cycA	0.5	8	0.5	16	ND	16	ND	16	ND	4	ND	0.5	ND
DAR169	32	32	32	32	8/2 (0.31)	128	4/32 (0.37)	64	4/8 (0.25)	8	4/1 (0.25)	2	ND
DAR169 cycA	16	4	0.5	16	ND	64	ND	16	ND	2	ND	0.5	ND
COL	512	256	64	512	8/128 (0.37)	128	16/32 (0.5)	>2048	ND	128	32/0.5 (0.5)	256	8/64 (0.37)
DAR113	128	64	32	128	8/8 (0.3)	64	8/4 (0.3)	256	8/8 (0.2)	128	8/4 (0.2)	16	8/0.5 (0.2)
BH1CC	256	512	32	256	8/32 (0.3)	128	8/8 (0.3)	1028	8/16 (0.2)	256	8/4 (0.2)	64	8/1 (0.2)
BH14B(04)	128	64	16	128	4/32 (0.5)	128	4/32 (0.5)	512	4/32 (0.31)	64	4/1 (0.26)	64	2/8 (0.25)
BH8(03)	128	128	32	128	8/32 (0.5)	256	8/64 (0.5)	128	8/16 (0.25)	128	8/8 (0.3)	64	8/0.5 (0.1)
BH6	128	128	32	128	8/16 (0.37)	256	8/64 (0.5)	512	8/32 (0.31)	128	8/4 (0.28)	32	8/0.5 (0.26)
DAR202	64	64	32	128	4/32 (0.37)	128	4/32 (0.37)	64	8/16 (0.5)	64	8/2 (0.28)	4	8/1 (0.5)
DAR45	2	0.5	32	4	ND	32	8/2 (0.31)	4	ND	2	ND	0.5	ND
DAR13	128	32	32	128	4/32 (0.37)	128	8/4 (0.28)	128	4/32 (0.37)	32	8/1 (0.28)	8	4/2 (0.37)
RP62A	128	2	32	64	6/16 (0.5)	64	6/16 (0.5)	32	8/8 (0.5)	8	8/2 (0.5)	32	8/1 (0.2)

\* MIC values for each antibiotic when measured individually;  $\mu\text{g/mI}$ 

\*\* MIC values for each antibiotic when measured in combination, also known as the fractional inhibitory concentration (FIC); µg/ml

\*\*\* FIC indices ( $\Sigma$ FIC) for antibiotic combinations.  $\Sigma$ FIC = FIC A + FIC B, where FIC A is the MIC of DCS in combination with the  $\beta$ -lactam/MIC of DCS alone, and FIC B is the MIC of the  $\beta$ -lactam in combination with DCS/MIC of the  $\beta$ -lactam alone. Combinations are synergistic when the  $\Sigma$ FIC is  $\leq$ 0.5 and indifferent when the  $\Sigma$ FIC is >0.5 to <2. \*ND. Not determined if strain is susceptible (or hyper-resistant) to  $\beta$ -lactam antibiotic or for *cycA* mutants with reduced DCS &  $\beta$ -lactam MICs.



Fig. 1





Fig. 3



Fig. 4





# SUPPLEMENTARY DATA

Impaired alanine transport or exposure to D-cycloserine increases the susceptibility of MRSA to  $\beta$ -lactam antibiotics

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# **Supplementary Tables**

Table S1. Antibacterial activity (minimum inhibitory concentrations, MIC) and drug synergy (fractional inhibitory concentration indices, ΣFIC) of
D-cycloserine (DCS), and clindamycin (CLI), tobramycin (TOB), trimethoprim (TMP), mupirocin (MUP), ciprofloxacin (CIP), kanamycin (KAN) and
spectinomycin (SPT) alone and in DCS/antibiotic combinations, against fourteen S. aureus strains and S. epidermidis RP62A.

Antibiotic $\rightarrow$		CU*	DCS/CLI**	тор	DCS/TOB	TMD	DCS/TMP	CIP	DCS/CIP	KAN	DCS/KAN	SDT	DCS/SPT	MUD	DCS/MUP
↓ Strain	DCS	CLI	(ΣFIC)***	TOB	(ΣFIC)	TIVIP	(ΣFIC)	CIP	(ΣFIC)	KAN	(ΣFIC)	371	(ΣFIC)	NOP	(ΣFIC)
JE2	32	0.25	ND	1	ND	1	ND	32	16/16 (1)	2	ND	64	16/32 (1)	0.25	ND
USA300	32	2048	32/2048 (2)	0.5	ND	1	ND	32	16/16 (1)	2	ND	64	16/32 (1)	2048	32/2048 (2)
DAR173	32	0.5	ND	512	32/512 (2)	0.5	ND	256	16/128 (1)	512	16/256 (1)	64	16/32 (1)	0.5	ND
DAR22	32	0.5	ND	512	16/256 (1)	0.5	ND	256	16/128 (1)	256	16/128 (1)	64	16/32 (1)	0.5	ND
DAR169	32	0.5	ND	0.5	ND	0.5	4/8 (0.25)	1	ND	2	ND	64	16/32 (1)	0.5	ND
COL	64	0.5	ND	0.5	ND	0.5	ND	1	ND	0.5	ND	64	16/32 (1)	0.5	ND
DAR113	32	0.25	ND	0.5	ND	1	ND	0.5	ND	2	ND	64	16/32 (1)	0.25	ND
BH1CC	32	1024	32/1024 (2)	32	32/32 (2)	256	32/256 (2)	32	16/16 (1)	>1024	ND	>1024	ND	32	16/8 (0.75)
BH14B(04)	16	0.5	ND	0.5	ND	0.5	ND	128	16/128 (1.5)	2	ND	64	8/32 (1)	0.5	ND
BH8(03)	32	1024	32/1024 (2)	0.5	ND	0.5	ND	256	16/128 (1)	8	32/4 (2)	128	16/64 (1)	4	32/4 (2)
BH6	32	0.5	ND	0.5	ND	0.5	ND	>512	16/256 (1)	1	ND	128	16/64 (1)	0.5	ND
DAR202	32	0.5	ND	128	16/64 (1)	0.5	ND	2	ND	>1024	ND	>1024	ND	0.5	ND
DAR45	32	1024	32/1024 (2)	512	16/256 (1)	0.5	ND	128	16/64 (1)	128	16/64 (1)	512	32/1024 (2)	0.5	ND
DAR13	32	0.5	ND	0.5	ND	32	16/16 (1)	32	16/16 (1)	2048	ND	>1024	ND	0.5	ND
RP62A	32	2048	32/2048 (2)	16	16/8 (1)	256	32/256 (2)	0.25	ND	>1024	ND	>1024	ND	0.25	ND

\* MIC values for each antibiotic when measured individually;  $\mu\text{g}/\text{ml}$ 

\*\* MIC values for each antibiotic when measured in combination, also known as the fractional inhibitory concentration (FIC); μg/ml

\*\*\* FIC indices ( $\Sigma$ FIC) for the antibiotic combination.  $\Sigma$ FIC = FIC A + FIC B, where FIC A is the MIC of DCS in combination with the antibiotic/MIC of DCS alone, and FIC B is the MIC of the antibiotic in combination with DCS/MIC of the antibiotic alone. The combination is considered synergistic when the  $\Sigma$ FIC is  $\leq 0.5$ , indifferent when the  $\Sigma$ FIC is > 0.5 to < 2.

<sup>†</sup>ND. Not determined if strain is susceptible (or hyper-resistant) to the antibiotic or for *cycA* mutants with reduced DCS & antibiotic MICs.

Table S2. Bacterial strains and plasmids used in this study

Strains/plasmids	Relevant Details
RN4220	Restriction-deficient laboratory S. aureus
USA300 FPR3757	Community associated MRSA isolate of the USA300 lineage [43]. SCCmec type IV. CC8.
JE2	USA300 cured of p01 & p03. Parent of Nebraska Transposon Mutant Library (NTML).
NE810	JE2 NTML cycA (SAUSA300_1642) mutation. Erm <sup>r</sup> .
USA300 cycA	USA300 FPR3757 cycA. Constructed by transduction of cycA::Tn allele from NE810.
NE1868	JE2 NTML mecA mutation.
NE1713	JE2 NTML <i>alr</i> (SAUSA300_2027) mutation.
BH1CC	MRSA clinical isolate; SCCmec type II; CC8 [44]
COL	MRSA reference strain; SCCmec type I; CC8 [45]
BH14(04)	MRSA clinical isolate; SCCmec type IV; CC22 [44]
BH8(03)	MRSA clinical isolate; SCCmec type IV; CC22 [44]
BH6(03)	MRSA clinical isolate; SCCmec type II; CC8 [44]
DAR113	MRSA reference isolate; SCCmec type IV; CC22 [44, 46]
DAR13	MRSA reference isolate; SCC <i>mec</i> type IV; CC8 [44, 46]
DAR45	MRSA reference isolate; SCCmec type II; CC30 [44, 46]
DAR202	MRSA reference isolate; SCCmec type III; CC239 [44, 46]
DAR173	MRSA reference isolate; SCC <i>mec</i> type IV; CC5 [44, 46]
DAR173 <i>cycA</i>	DAR173 cycA mutant (cycA::Tn allele from NE810)
DAR22	MRSA reference isolate; SCC <i>mec</i> type III; CC5 [44, 46]
DAR22 cycA	DAR22 cycA mutant (cycA::Tn allele from NE810)
DAR169	MRSA Reference strain; SCCmec type I; CC8 [44, 46]
DAR169 cycA	DAR169 cycA mutant (cycA::Tn allele from NE810)
8325-4	NCTC 8325 derivative cured of prophages [47], methicillin susceptible, CC8.
8325-4 <i>cycA</i>	8325-4 cycA mutant (cycA::Tn allele from NE810).
ATCC 29213	Methicillin susceptible S. aureus strain for antibiotic susceptibility testing.
ATCC 29213 cycA	ATCC 29213 cycA mutant (cycA::Tn allele from NE810).
ATCC 25923	Methicillin susceptible S. aureus strain for antibiotic susceptibility testing.
S. epidermidis RP62A	ATCC 35984. Methicillin resistant, biofilm positive. [21]
E. coli	E. coli HST08
Plasmids	
pL150	E. coli-Staphylococcus shuttle vector. Apr (E. coli), Cmr. (Staphylococcus)
pcycA	pLI50 carrying <i>cycA</i> from JE2

Table S3. Oligonucleotide primers used in this study

Target Gene	Primer Name	Primer Sequence (5'-3')
сусА	NE810_Fwd	ACAGAATAGCCACAAATAGCACC
	NE810_Rev	ACAGAATAGCCACAAATAGCACC
сусА	NE810F1_Fwd	GTCTTCAAGAATTCGGCCACAAATAGCACCATTAA
	NE810F1_Rev	CGACTCTAGAGGATCATGTCCCAAGCCCTAAAAC
mecA	mecA1_Fwd	TGCTCAATATAAAATTAAAACAAACTACGGTAAC
	mecA1_Rev	GAATAATGACGCTATGATCCCAA
gyrB	gyrB_Fwd	CCAGGTAAATTAGCCGATTGC
	gyrB_Rev	AAATCGCCTGCGTTCTAGAG

**Supplementary Figures** 



**Figure S1.** Mutation of *cycA* increases the susceptibility of MRSA to β-lactam antibiotics and D-cycloserine. A. Chromosomal location of *cycA* and neighbouring genes *icd* (isocitrate dehydrogenase), *gltA* (citrate synthase), *pyk* (pyruvate kinase) and *pfkA* (6-phosphofructokinase). The locations of transposon insertions in NE810, NE491, NE594 and NE1497 mutants from the Nebraska library are indicated. **B.** E-test measurement of oxacillin minimum inhibitory concentrations (MICs) in JE2 (wild type), NE810 (*cycA*::Tn), NE810 carrying pLI50 (control) and p*cycA*. **C.** Comparison of relative *mecA* gene expression by LightCycler RT-PCR in JE2 and NE810 grown to  $A_{600}$ =3 in BHI media. The data are the average of three independent experiments and standard deviations are shown **D.** Comparison of zones of inhibition around D-cycloserine 30µg disks on lawns of JE2, NE810 (*cycA*::Tn), NE810 pLI50 (control) and NE810 pcycA grown on Mueller Hinton (MH) agar.



**Figure S2. A.** Susceptibility of JE2 (wild type), NE1868 (*mecA*::Tn), NE810 (*cycA*::Tn), and NE1713 (*alr*::Tn) grown on MH agar to cefoxitin (FOX, 30µg disks). **B.** Susceptibility of JE2 (wild type) and NE1713 (*alr*::Tn) grown on MH agar to D-cycloserine (DCS, 30µg disks).



**Figure S3.** Amino acid consumption by JE2 and NE810 grown aerobically in chemically defined media containing 14mM of glucose (CDMG). Residual amino acids were measured in spent media after 2, 4, 6 and 8 h growth.



**Figure S4.** Preparation of cell suspensions for peptidoglycan extraction and structural analysis by UPLC-MS. 50 ml flask cultures were inoculated into fresh BHI media from overnight cultures at a starting cell density of  $A_{600}$ =0.05 and incubated at 37°C. The number of CFU/ml was enumerated every 1-2 h for 6 h and again after 24 hours. For JE2 cultures being dosed with DCS, the antibiotic was added after approximated 2 h ( $A_{600}$ =0.5) and the cells collected after a further 100 mins. Cells from untreated JE2 and NE810 control cultures were collected at the same time point.



**Figure S5.** In vitro kill curves for D-cycloserine (DCS), oxacillin and cefoxitin with methicillin resistant *S. epidermidis strain* RP62A. Antibiotics at the concentrations indicated (equivalent to  $0.5 \times MIC$ ) were added to suspensions of overnight bacterial cultures adjusted to  $10^7$ CFU/ml in BHI, incubated at  $37^\circ$ C and the number of CFU/ml enumerated at 0, 2, 4, 8 and 24 h. The data presented are the mean of three independent experiments. Antibiotic synergism was defined as a  $\ge 2 \log^{10}$  decrease in the number of CFU/ml in cell suspensions exposed to DCS/ $\beta$ -lactam combinations compared to the most effective individual antibiotic alone.



Figure S6. Combination therapy with D-cycloserine and oxacillin significantly reduces the bacterial burden in the spleen of mice infected with MRSA. The number of colony-forming units (CFU) recovered from the spleens of mice infected by tail vein injection with  $5 \times 10^6$  JE2 or NE810 (CycA) and left untreated or treated with 75mg of oxacillin (Ox)/kg, 30mg of DCS/kg or a combination of both Ox and DCS delivered subcutaneously every 12 hours for 5 days. The first antibiotic dose was given 16 hours after infection. Significant differences determined using one-way ANOVA with Kruskal-Wallis test followed by Dunn's multiple comparisons test are denoted using asterisks (\*p $\leq$ 0.05). The limit of detection (50 colonies) is indicated with a hashed red line.



**Figure S7.** Amino acid consumption by JE2 and NE810 grown aerobically in chemically defined media lacking glucose (CDM). Residual amino acids were measured in spent media after 2, 4, 6 and 8 h growth.