# Novel regulatory logic in the antibiotic resistance response of 1 Enterococcus faecalis against cell envelope targeting antibiotics 2 3 Sali M. Morris<sup>1</sup>, Georg Fritz<sup>2</sup>, Tim Rogers<sup>3</sup>, Susanne Gebhard<sup>1\*</sup> 4 5 6 <sup>1</sup>Life Sciences Department, Milner Centre for Evolution, University of Bath, Claverton Down, Bath, BA2 7AY, United Kingdom; <sup>2</sup>School of Molecular Sciences, The University of 7 Western Australia, WA 6009 Crawley, Australia; <sup>3</sup>Department of Mathematical Sciences, 8 University of Bath, Claverton Down, Bath, BA2 7AY, United Kingdom 9 10 \*Corresponding author. Phone: +44 1225 386421; E-mail: s.gebhard@bath.ac.uk; 11 12 Running title: Antibiotic resistance control in E. faecalis 13 14 Data accessibility statement: The data that support the findings of this study are available in 15 the supplementary material. 16 Funding statement: This work was supported in part by grant MR/N0137941/1 for the GW4 17 BIOMED MRC DTP, awarded to the Universities of Bath, Bristol, Cardiff and Exeter from 18 19 the Medical Research Council (MRC)/UKRI. Conflicts of interest: The authors declare no conflicts of interest. 20 21

### 22 Summary

23 Enterococcal infections frequently show high levels of antibiotic resistance, including to cell envelope-24 acting antibiotics like daptomycin (DAP). While we have a good understanding of the resistance 25 mechanisms, less is known about the control of such resistance genes in enterococci. Previous work 26 unveiled a bacitracin resistance network, comprised of the sensory ABC transporter SapAB, the twocomponent system (TCS) SapRS and the resistance ABC transporter RapAB. Interestingly, components 27 28 of this system have recently been implicated in DAP resistance, a role usually regulated by the TCS 29 LiaFSR. We here explored the interplay between these two regulatory pathways. Our results show the 30 regulation by SapR of an additional resistance operon, *dltXABCD*, and show that LiaFSR regulates the 31 expression of *sapRS*, placing SapRS target genes under dual control: *dltXABCD* expression depends on 32 both antibiotic-induced cellular damage (LiaFSR) and the presence of a substrate drug for the sensory transporter (SapAB). We further show that this network protects E. faecalis from antimicrobials 33 produced by potential competitor bacteria, providing a potential rationale for the evolution of this 34 35 regulatory strategy. The network structure described here can explain why clinical DAP resistance often 36 emerges via mutations in regulatory components, which may ultimately lead to the discovery of new 37 therapeutic targets.

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39 Keywords: antimicrobial peptides, cell envelope stress, antimicrobial resistance, two-

40 component system, signalling

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### 42 Introduction

43 The rise of antibiotic resistant bacteria is one of the greatest current threats to public health, resulting in 670,000 infections a year and 33,000 deaths in Europe alone<sup>1</sup>. Of these infections, the "ESKAPE" 44 organisms<sup>2</sup> (Enterococci spp., Staphylococcus aureus, Klebsiella pneumoniae, Acinetobacter 45 46 baumannii, Pseudomonas aeruginosa, Enterobacter spp.) have driven the rising number of nosocomial and antibiotic-resistant infections in the past decade. Of these bacteria, the enterococci are the second 47 most common causative agent of nosocomial infections in the US; including bacteraemia, endocarditis 48 and urinary tract infections<sup>3-5</sup>. The two species most frequently isolated, *Enterococcus faecalis* and 49 Enterococcus faecium, remain a major infection-control challenge, particularly in healthcare settings. 50

Enterococci became recognised as important nosocomial pathogens due to their high level of intrinsic resistance to several antimicrobials<sup>6</sup> (e.g. penicillin, ampicillin and cephalosporins) and their capacity to acquire further resistance determinants. One such acquired resistance is to the glycopeptide antibiotic vancomycin, which occurs through plasmid acquisition and was first reported in the 1988<sup>7,8</sup>, 30 years after vancomycin was introduced for clinical use<sup>9</sup>. Despite the molecular mechanisms of vancomycin resistance in enterococci being well understood today<sup>10</sup>, infections by vancomycin resistant enterococci (VRE) still result in serious health and economic impacts<sup>11</sup> and are an increasing problem worldwide.

58 One of the last-resort antibiotics used to treat these VRE infections is the lipopeptide antibiotic daptomycin (DAP)<sup>12</sup>. Disappointingly, within 2 years of clinical introduction of the drug in 2003, DAP-59 resistant enterococcal isolates were reported<sup>13</sup>, and in contrast to vancomycin resistance, this occurred 60 through a subtle chromosomal change based on the mutations in genes *liaF*, *cls* and  $gdpD^{14-17}$ . LiaF is 61 62 a transmembrane protein involved in monitoring the integrity of the cell envelope and responding to damage<sup>18</sup>, whereas Cls (cardiolipin synthase) and GdpD (glycerol-phosphodiester phosphodiesterase) 63 are both involved in phospholipid metabolism<sup>14</sup>. Of these, mutation of *liaF* has been proposed to be the 64 first pivotal event towards daptomycin resistance<sup>15,19</sup>. However, the exact role of the *liaF* mutations in 65 66 DAP resistance so far remains unclear.

67 LiaF is part of a three-component regulatory system, LiaFSR, which is important amongst the Firmicutes for coordinating the cell envelope stress response (CESR) against antimicrobial-induced 68 damage<sup>20-23</sup>. The system is comprised of LiaF and a conventional two-component system (TCS): the 69 sensor kinase LiaS and the response regulator LiaR<sup>20,24,25</sup>. LiaF is an inhibitor of LiaS, maintaining the 70 sensor kinase in an inactive conformation<sup>20</sup>. Rather than detecting a specific antimicrobial compound, 71 LiaFSR responds to cell envelope damage, although the exact stimulus is unknown<sup>26</sup>. Upon sensing this 72 73 damage, LiaF releases LiaS, which is then able to phosphorylate LiaR to induce the expression of the 74 system's target operon, *liaXYZ*<sup>19,27</sup>. The *liaXYZ* cluster is involved in sensing and binding antimicrobials at the cell surface to provide resistance $^{27}$ . 75

76 To add to this complexity, the Lia system does not exist in isolation, but is just one of many TCSs involved in monitoring cell envelope integrity, each responding to their own individual stimuli and 77 78 activating a unique set of target genes. The CroRS system, unique to the enterococci, is the main regulator required for cephalosporin resistance<sup>28</sup>, whereas the VicKR (YycFG) system is essential 79 across the low-CG Gram positives and is involved in regulating cell division, lipid biosynthesis, biofilm 80 and virulence<sup>29</sup>. An additional element of the network is the serine/threonine kinase IreK and the 81 82 phosphatase IreP, involved in maintaining cell wall integrity by potentially regulating peptidoglycan biosynthesis and metabolism<sup>30–32</sup>. 83

84 A further TCS involved in monitoring the cell envelope is EF0926/27, which we have now renamed SapRS (Sensor of Antimicrobial Peptides), identified in our previous work<sup>33</sup>. SapRS is part of a 85 86 bacitracin resistance network comprised of the histidine kinase SapS, the response regulator SapR and 87 the Bce-like ABC transporters: SapAB (EF2752/51) and RapAB (Resistance against Antimicrobial Peptides) (EF2050/49)<sup>33</sup>. The use of the sensory transporter SapAB to control the activity of SapRS 88 implements a 'flux-sensing' mechanism to regulate bacitracin resistance, as was shown for the 89 homologous system in Bacillus subtilis<sup>34</sup>. In brief, upon exposure, bacitracin binds to its membrane-90 91 associated target molecule, undecaprenol-pyrophosphate (UPP), blocking the dephosphorylation and recycling of UPP in the lipid II cycle of cell wall biosynthesis<sup>35,36</sup>. The sensory ABC transporter SapAB, 92 based on biochemical evidence from the *B. subtilis* system, forms a sensory complex with  $SapRS^{37}$ . In 93

the absence of bacitracin, the transporter maintains the histidine kinase, SapS, in an 'OFF' state<sup>38</sup>. Upon 94 95 detection of bacitracin-UPP (BAC-UPP) complexes, SapAB switches its role from repressor to activator of SapS, resulting in kinase autophosphorylation<sup>38</sup>. SapS can then phosphorylate SapR, which in turn 96 induces the production of the resistance transporter RapAB, RapAB, again based on evidence from its 97 98 B. subtilis homologue, frees UPP from the inhibitory grip of bacitracin using a target protection mechanism<sup>39</sup>, allowing dephosphorylation of UPP and continuation of cell wall synthesis, rendering the 99 100 cell resistant to bacitracin. The equivalent system in *B. subtilis* is a self-contained module, involving only a single transporter (BceAB) and TCS (BceRS)<sup>40</sup> with no known further genes involved in its 101 signalling or resistance mechanism. However, in *E. faecalis*, we observed that the TCS SapRS is itself 102 transcriptionally induced by bacitracin, however we had not identified the regulatory system that 103 controls this expression<sup>33</sup>. 104

105 Interestingly, recent work by the Arias lab demonstrated that when experimentally evolving E. faecalis 106 for DAP resistance in a  $\Delta liaFSR$  background, mutations were observed in the sensory transporter SapAB<sup>41,42</sup>, suggesting a functional link between the Sap and Lia systems. In *B. subtilis*, the Lia system 107 is known to be one of the main components of bacitracin resistance<sup>43,44</sup>, but there is currently no 108 evidence for a role of LiaFSR in response to bacitracin in E. faecalis. Neither is there any indication of 109 110 a role for the Sap system in responding to DAP exposure. However, this recent evidence suggested an interplay between both the Sap and Lia systems and that, potentially, both systems may be contributing 111 112 to resistance against bacitracin and daptomycin.

113 In accordance with the need to deepen our knowledge of the CESR in *E. faecalis*, in this study we sought to examine the potential interplay between the Sap and Lia systems and investigate the 114 115 involvement of further genes associated with this regulatory network. Utilising mutagenesis and analysis of promoter activity, we provide evidence of the activation of LiaFSR signalling in response 116 to bacitracin and describe a direct functional link between the Lia and Sap systems by demonstrating 117 the regulation of sapR by LiaR in response to antibiotic exposure. We also show the contribution of an 118 additional resistance operon involved in response to bacitracin and unravel a differential response of 119 120 the network between bacitracin and daptomycin treatment. Our data show that interplay between SapRS

and LiaFSR signalling effectively implements a logic 'AND' gate, whereby expression of SapR target
genes is under dual control and requires two separate signalling inputs. These observations also explain
the occurrence of mutations in both regulatory systems during clinical and experimental emergence of
DAP resistance in enterococci found previously, thus providing a compelling model for the complex
CESR network in *E. faecalis*.

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# 127 **Results**

LiaFSR controls expression of SapRS. As mentioned above, from previous work we had identified 128 129 increased expression of *sapRS* under exposure to bacitracin or mersacidin<sup>33</sup>. Interestingly, we also showed that *sapRS* was not autoregulated, in contrast to other two component systems such as CroRS 130 and LiaFSR<sup>20,28</sup>, and regulation did not depend on either SapAB or RapAB<sup>33</sup>. We therefore here first 131 132 aimed to identify the regulator of sapRS. For possible candidates, we considered potential regulators 133 that are common amongst the Firmicutes and are known to respond to bacitracin. A potential candidate was LiaFSR, deemed the 'master regulator' of the CESR in Bacillus subtilis<sup>43</sup> and recently shown to 134 have a possible functional link to the Sap system<sup>41,42</sup>. Because of this evidence, we first aimed to 135 136 examine the relationship between these two systems.

Firstly, we had to test if LiaFSR was able to respond to bacitracin treatment. It is well established that *liaX* is under the control of LiaFSR<sup>27</sup>, and therefore we exposed *E. faecalis* harbouring a  $P_{liaX}$ -*lacZ* transcriptional fusion to increasing levels of bacitracin, as a readout for LiaFSR activity. The results showed a ~10-fold increase in *liaX* expression at 32 µg ml<sup>-1</sup> bacitracin compared with untreated cells

141 (Fig. 1A, black line), indicating that LiaFSR can indeed respond to bacitracin exposure in *E. faecalis*.

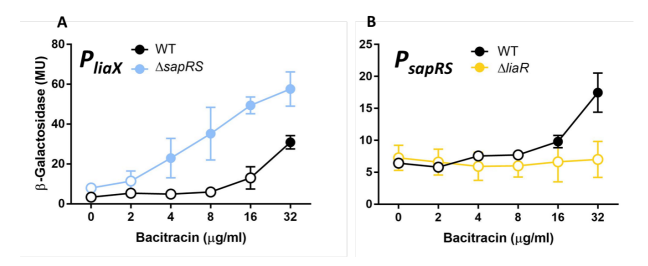


Figure 1. The LiaFSR operon responds to bacitracin exposure and induces the expression of both *liaXYZ* and *sapRS*, with the absence of *sapRS* resulting in the hypersensitivity of the *liaXYZ* promoter. Cells of *E. faecalis* JH2-2 wild-type or isogenic deletion strains harbouring a  $P_{liaX}$ -*lacZ* (panel A) or  $P_{sapRS}$ -*lacZ* (panel B) transcriptional fusion were grown to exponential phase and challenged with increasing concentrations of bacitracin.  $\beta$ -galactosidase activity, expressed as Miller units (MU), was assayed following 1h incubation. Results are means and standard deviations for three biological repeats. The significance of induction relative to untreated cells was calculated for each strain by a one-tailed, pairwise *t*-test analysis. Significance is indicated by a filled symbol (p < 0.05); unfilled symbols represent no significant increases compared to uninduced conditions.

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144 Following this, we tested if LiaR can regulate the expression of *sapRS*. To do this, we monitored the response of a transcriptional PsapRs-lacZ fusion to increasing bacitracin. The results showed that in wild-145 type *E. faecalis* carrying the transcriptional *sapRS-lacZ* fusion, bacitracin concentrations of 16 µg ml<sup>-1</sup> 146 or higher led to significant induction of the reporter, with a three-fold increase compared to uninduced 147 cells at 32  $\mu$ g ml<sup>-1</sup> (Fig. 1B, black line). Deletion of *liaR* in the reporter strain resulted in a complete loss 148 of sapRS induction, with expression remaining at basal levels. This indicated that LiaR indeed regulates 149 the expression of sapRS in response to bacitracin exposure, presenting first evidence of a direct 150 151 functional link between the Lia and Sap regulatory systems.

A surprising feature of this LiaR regulation was the high concentration required to induce a response, as *liaX* expression only significantly increased at 32  $\mu$ g ml<sup>-1</sup>. One possible explanation for this might be that the response was masked by other components of the bacitracin stress response in *E. faecalis*, for example RapAB, which plays an active role in the removal of bacitracin from UPP<sup>33</sup>, protecting the

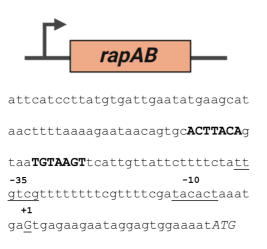
156 cell from damage. In B. subtilis, it was observed that the RapAB equivalent, BceAB, was the primary response to protect the cell from bacitracin exposure and masked the response of the Lia system<sup>45</sup>. To 157 prevent the induction of the main bacitracin resistance genes of *E. faecalis*, controlled by SapRS<sup>33</sup> and 158 thus remove potential interference with Lia signalling, we therefore introduced the  $P_{liax}-lacZ$ 159 160 transcriptional fusion into the  $\Delta sapRS$  background. Compared to the wild type, the expression of the liaX promoter in response to bacitracin challenge of the deletion strain showed increased sensitivity, 161 significantly inducing expression from 4  $\mu$ g ml<sup>-1</sup> bacitracin and reaching overall higher activities (Fig. 162 1A, blue line). This showed that the weak response of Lia signalling to bacitracin in the wild type was 163 indeed due to masking effects of the resistance genes controlled by SapRS, implying the presence of 164 multiple layers of protection, similar to those observed in *B. subtilis*<sup>45</sup>, as well as physiological links 165 between the SapRS- and LiaFSR-dependent components of the CESR in E. faecalis. 166

SapRS controls the expression of the *dltABCD* operon. To expand our understanding of the Lia/Sap 167 168 regulatory network, we next considered if other genes may also be under its control. Currently, the only known gene under SapRS regulation is the resistance ABC-transporter operon rapAB<sup>33</sup>. In B. subtilis, 169 both the Lia and Bce systems act as self-contained modules, controlling the regulation of only a single 170 resistance operon each, *liaIH* and *bceAB* respectively, each encoded adjacently to its regulatory operon 171 on the chromosome <sup>24,40,43</sup>. However, the regulatory setup in *E. faecalis* seems to be much more 172 complex. This is supported by evidence from the literature, with the indication that the Lia system has 173 a larger operon than just itself and *liaXYZ*<sup>27</sup>. There is also a proposal that Lia contributes to DAP 174 resistance through the regulation of *dltXABCD* (*dlt*), although there is no evidence to date that Lia is a 175 176 direct regulator of this operon. Dlt is responsible for the D-alanylation of teichoic acids (TA) on the bacterial cell surface, resulting in a decrease in the negative charge of the cell envelope<sup>46</sup>, a mechanism 177 commonly involved in DAP resistance amongst the low-CG bacteria<sup>47</sup>. 178

Interestingly, recent evidence has suggested a regulatory link between SapRS and *dlt*. When evolving *E. faecium* for DAP resistance in the absence of LiaFSR, mutations occurred in the *sapAB* homologue, *ycvRS*. These mutations correlated with an increase in *dlt* transcription when measured by qPCR and
resulted in an increase in cell surface charge<sup>41,42</sup>. In addition, the *dltABCD* operon is located directly

- downstream from *sapAB* on the chromosome, suggesting a possible functional link between the genes.
- 184 Moreover, the promoter region of *dlt* contains a putative SapR binding site, similar to that of the *rapAB*
- 185 promoter (Fig. 2). This body of evidence therefore led us to investigate the contribution of LiaFSR and
- 186 SapRS to *dlt* regulation.

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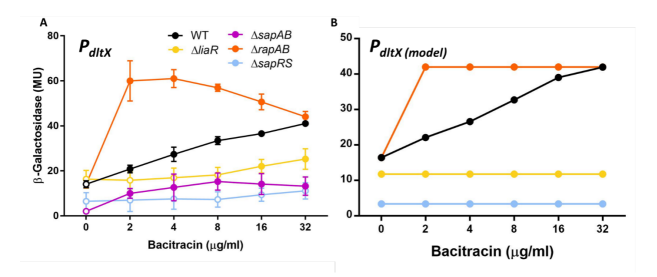
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**Figure 2. Schematic of the promoter regions of both** *rapAB* and *dltABCD*. The sequence 150 base pairs upstream of the ATG start codons are shown. The proposed binding site for SapR on both the *rapAB* and *dltABCD* promoters is in bold and capitalised, and the likely -10 and -35 elements are underlined. The experimentally confirmed transcriptional start sites are capitalised and underlined (+1) and the translational start site is shown in italicised capitals<sup>33,84</sup>.

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       To this end, we first constructed a transcriptional P_{dux}-lacZ fusion to test dlt induction by bacitracin.
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       The results showed that in wild-type JH2-2 carrying the fusion, increasing bacitracin concentrations led
       to a dose-dependent increase in dlt expression, resulting in ~4-fold higher activity at 32 \mug ml<sup>-1</sup>
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       compared to untreated conditions (Fig. 3A, black line), showing that dlt expression is indeed induced
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       in response to bacitracin in E. faecalis. When we tested the response of the reporter in strains carrying
       deletions of either sapRS or sapAB, the results showed a decrease in basal activity in both strains and a
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       complete loss of the promoter's bacitracin response in \Delta sapR (Fig. 3A, blue and magenta lines). Loss
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       of SapAB still resulted in induction compared with uninduced cells, but overall activities were
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       considerably lower than in the wild-type strain. This indicated that SapRS was essential for dlt
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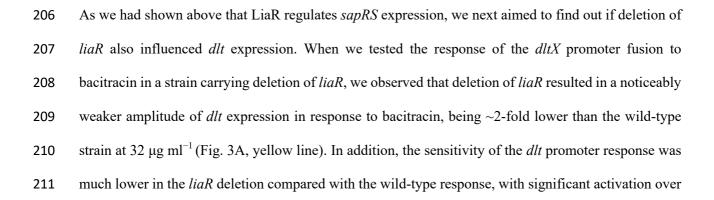
expression in response to bacitracin, but a residual amount of *dlt* induction remained in the *sapAB* deletion. This is consistent with the less direct role of SapAB in signalling, i.e. via controlling SapRS activity and not the target genes directly. The residual bacitracin response observed in the *sapAB* deleted strain can likely be explained by increased production of SapRS due to LiaFSR signalling, which would be expected to lead to increased basal activity of SapRS and thus the *dlt* promoter.



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**Figure 3. Induction of the resistance operon** *dltXABCD* by bacitracin. Panel A indicates the experimental induction of *dltX*. Cells of *E. faecalis* harbouring a *PdltX-lacZ* transcriptional fusion were grown to exponential phase and challenged with increasing concentrations of bacitracin. Beta-galactosidase activity, expressed as Miller units (MU), was assayed following 1h incubation in wild type (WT) and deletion strain backgrounds. Results are means and standard deviations for three biological repeats The significance of induction relative to untreated cells was calculated for each strain by a one-tailed, pairwise *t*-test analysis. Significance is indicated by a filled symbol (p < 0.05), unfilled symbols represent no significant increases compared to uninduced conditions. Panel B indicates the mathematical model of *dltX* induction in the *E. faecalis* strains indicated in panel A.

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baseline occurring at 16  $\mu$ g ml<sup>-1</sup> rather than at 2  $\mu$ g ml<sup>-1</sup> in the wild type. However, there was no effect on basal activity of *dlt*, in contrast to deletion of *sapR* and *sapAB*. These data are consistent with LiaR regulating *sapRS* expression, and SapRS being the actual regulator of *dlt* expression in response to bacitracin.

As we had observed before with the *liaX* and *sapRS* promoters, we also wanted to investigate if the protection provided by the SapRS-target RapAB was dampening the expression of *dlt* in response to bacitracin. To examine this, we tested the response of *dlt* expression in the absence of *rapAB*. In a *ArapAB* background, the *dlt* promoter demonstrated markedly increased sensitivity, and a much stronger response to bacitracin, resulting in a ~3-fold increase in expression at 2 µg ml<sup>-1</sup> when compared to the wild-type (Fig. 3A, orange line). This response demonstrated the presence of a layered protection with RapAB activity moderating *dlt* expression.

Overall, these findings were rather surprising, as the signalling pathway appeared remarkably complex 223 224 to result in a relatively simple outcome, i.e. inducing the expression of two resistance genes in response 225 to an antibiotic. To test if our understanding of the regulatory pathway was plausible, we therefore 226 developed a representative mathematical model to see if this would reproduce the behaviour we had observed in the experiments. At the core, this model was based on a simplified form of the flux-sensing 227 mechanism described previously for the *B. subtilis* BceRS-BceAB system<sup>34</sup>, which was then expanded 228 229 upon to reproduce the hypothesised network structure investigated here. In brief, the model considered bacitracin binding to UPP to form UPP-bacitracin complexes with the on-rate dependent on the 230 bacitracin concentration. These complexes then drive transport activity of, in this case, SapAB, 231 according to a soft-switch type functional response (see Methods), which the model translates into 232 233 activation of SapRS and thus *dlt* expression. Importantly, SapRS activity also drives expression of the resistance transporter operon, in this case rapAB, and production of RapAB leads to a reduction in 234 formation of UPP-bacitracin complexes due to the target protection activity of the transporter<sup>34,39</sup>. This 235 236 creates the negative feedback loop that is characteristic of the flux-sensing mechanism and leads to the 237 gradual response behaviour of the output promoters (Fig. 3B, black symbols). To adapt this model to the Lia-Sap regulatory pathway of *E. faecalis*, we additionally considered the activity of the Lia system. 238

This was modelled to also respond to UPP-bacitracin complexes in a switch like manner representing the generation of cellular damage caused by these complexes. The model then linked the Lia and Sap components by modulating the SapRS signalling output (i.e. *dlt* and *rapAB* expression) according to Lia activity, driven by bacitracin. The mathematical details of the model are explained in the Methods section. Fitting to experimental data was performed by matching differential activity of SapRS against both *rapAB* and *dlt* recorded at different bacitracin concentrations.

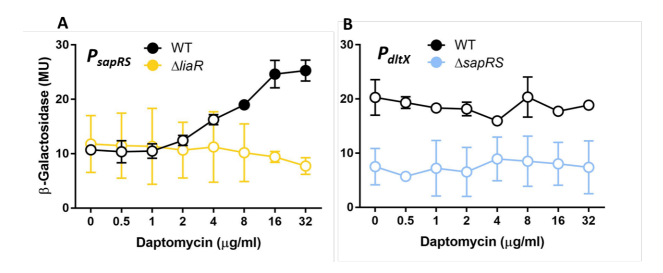
245 This model accurately reflected the behaviour of our experimental strains, depicting the same gradual response to bacitracin of the *dltX* promoter in wild-type *E. faecalis* (Fig. 3B, black symbols), as well as 246 247 the hypersensitive response in the *rapAB* deletion strain, where the negative feedback from RapABdriven removal of bacitracin was missing (orange symbols). Importantly, the model gave the same 248 complete loss of *dltX* activity when *sapRS* was deleted (blue symbols) as observed experimentally, as 249 well as the normal basal level activity but loss of bacitracin-dependent induction when *liaR* was deleted 250 251 (vellow symbols). A sapAB deletion strain was not specifically considered as the model did not differentiate between SapAB and SapRS activities and thus both strains would have been 252 mathematically identical. This close agreement between theoretical and experimental data strongly 253 suggested that our reconstruction of the regulatory pathway and connection between the Lia and Sap 254 255 systems was theoretically plausible and that no further major players were needed to explain the 256 behaviour of the *dltX* target promoter in the various mutant backgrounds analysed here.

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258 Daptomycin induces the expression of sapRS but not dltABCD. We had now established a sequential order of *dlt* regulatory control; in response to bacitracin, LiaFSR induces the expression of *sapRS*, and 259 260 in turn SapRS, activated by its sensory transporter SapAB, induces the expression of *dlt*. However, 261 treatment with bacitracin made it difficult to separate out the individual contributions to regulation by LiaFSR and SapRS, as both signalling systems respond to this antibiotic. Bacitracin forms a complex 262 with UPP, which acts as the input for SapRS signalling via SapAB<sup>39</sup>; at the same time, bacitracin 263 induces cell envelope damage, triggering LiaFSR activation<sup>26</sup>. To allow us to differentiate between the 264 contribution made by each regulator to the network, we instead required an antibiotic that would only 265

trigger signalling by one of the systems. To this end, we turned to daptomycin (DAP), which activates the Lia system by causing damage to the cell envelope<sup>48,49</sup>. However, it does not bind to UPP or the sugar-pyrophosphate moiety of Lipid II, which is the common feature of substrates for BceAB-type transporters<sup>39</sup>. Indeed, the tripartite complex of DAP with Lipid II and phosphatidyl glycerol in the membrane<sup>50</sup> is unlikely to be recognised by SapAB, and thus DAP exposure should not trigger SapRS activation, allowing us to study the specific impact of LiaFSR signalling in the regulatory pathway.

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**Figure 4. Exposure to daptomycin results in the induction of** *sapRS* **but does not induce the expression of** *dltXABCD.* Cells of *E. faecalis* harbouring a  $P_{sapRS}$ -*lacZ* (panel A) or a  $P_{dltX}$ -*lacZ* (panel B) transcriptional fusion were grown to exponential phase and challenged with increasing concentrations of daptomycin. Beta-galactosidase activity, expressed as Miller units (MU), was assayed following 30 minutes incubation in wild type (WT) and deletion strain backgrounds indicated. Results are means and standard deviations for two biological repeats. The significance of induction relative to untreated cells was calculated for each strain by a one-tailed, pairwise *t*-test analysis. Significance is indicated by a filled symbol (P < 0.05), unfilled symbols represent no significant increases compared to uninduced conditions.

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To investigate the effect of DAP exposure on LiaFSR and SapRS signalling and target gene expression, we again utilised transcriptional promoter fusions. The  $P_{sapRS}$ -lacZ transcriptional fusion (LiaFSR target) in the wild-type background showed a ~2-fold induction upon DAP treatment when compared to unexposed levels (Fig. 4A). As we had seen for bacitracin-induction, deletion of *liaR* again

completely abolished *sapRS* activation, showing the induction of *sapRS* by DAP was occurring through
LiaR regulation. This showed that DAP exposure indeed triggered LiaFSR signalling.

281 To test how DAP induction was propagated further down the regulatory pathway, we next exposed the P<sub>dltX</sub>-lacZ transcriptional fusion (SapRS target) to increasing DAP concentrations. Surprisingly, in wild-282 type *E. faecalis*, we found daptomycin treatment did not result in any induction of *dlt*, with expression 283 284 remaining at basal levels across all tested concentrations (Fig. 4B). As seen for the bacitracin challenge, deletion of sapRS resulted in a decrease in basal expression, from 20 MU to 7.5 MU, and again, there 285 286 was no induction of the *dlt* promoter in response to daptomycin exposure. This indicated that although 287 sapRS transcription was induced in response to daptomycin treatment, remarkably, expression of its target gene *dlt* remained at basal levels. This discovery suggested a differential response in the 288 signalling cascade dependent on the inducer, indicating that bacitracin and daptomycin might be 289 differentially sensed by SapRS, as further explained in the discussion. 290

#### 291 The network components vary in importance during interactions with antimicrobial producer

292 strains. As stated above, we were surprised by the complexity of the regulatory pathway controlling 293 what in other Firmicutes bacteria is a fairly straightforward response to antibiotic challenge, where each regulatory system controls its own resistance genes. To shed some light on the reasons for the 294 295 complexity of signalling in enterococci, we considered the environments these bacteria can be found 296 in. E. faecalis is a common member of many natural microbial communities, such as in soil and water or the gastrointestinal tract of humans and animals. There, the enterococci reside within the small and 297 large intestine and represent up to 1% of the faecal flora<sup>51–54</sup>. In such environments, *E. faecalis* interacts 298 with other microbes and must defend itself against antimicrobial producing bacteria. Therefore, we next 299 300 wanted to assess the role of the individual components of the resistance network in protecting E. faecalis from antimicrobial activity produced by potential competitor bacteria. To do this, we utilised deferred 301 antagonism assays to simulate relevant environmental pressures from other microbes the enterococci 302 303 may encounter. We used four antimicrobial producing strains: the subtilin producer Bacillus subtilis ATCC6633<sup>55</sup>, bacitracin producer *Bacillus lichenformis* ATCC10716<sup>56</sup>, nisin-A producer *Lactococcus* 304 lactis NZ9000<sup>57</sup> and nisin-P producer Streptococcus gallolyticus AB39<sup>58</sup>. Each producer was spotted 305

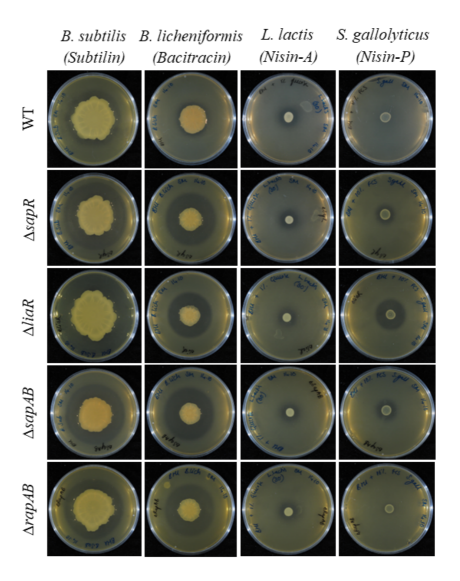
onto a plate, and the antimicrobial was allowed to accumulate over multiple days. This was followed
by the addition of an overlay containing the different *E. faecalis* strains to assess their sensitivity against
the produced compounds based on the size of the zone of inhibition (Fig. 5).

Firstly, we tested the *E. faecalis* strains against the antimicrobial subtilin, produced by *B. subtilis* ATCC6633. Of the components under investigation in this study, the LiaFSR system appeared to contribute most strongly to resistance against subtilin, as the  $\Delta liaR$  mutant displayed the largest increase in zone of inhibition compared to the wild type. The deletions of *sapR*, *sapAB* and *rapAB* also displayed an increased zone of inhibition compared to the wild type, but to a lesser extent than  $\Delta liaR$ . This therefore suggests they play a lesser role in subtilin resistance.

In contrast, when testing the *E. faecalis* strains against the bacitracin producer *B. lichenformis* ATCC10716, deletion of *sapR*, *sapAB* and *rapAB* showed similar increased sensitivity, consistent with their contribution to the bacitracin resistance network that we have examined in this study and previously<sup>33</sup>. Although also presenting increased sensitivity compared with the wild type, deletion of *liaR* resulted in a marginally smaller zone of inhibition compared with the other deletion strains. This is in line with our data presented above, showing the Lia system has a more indirect role in controlling the bacitracin response by regulating *sapRS* expression.

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323



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Figure 5. *E. faecalis* deletion strains show differential zones of inhibition in deferred antagonism tests against antimicrobials produced by other firmicute bacteria. The antimicrobial producer strains indicated at the top were grown overnight and then adjusted to an  $OD_{600}$  of 0.5 with fresh media. Aliquots (5ul) from each producer strain was then spotted and incubated at 25°C for 36-45 hours to allow antimicrobial accumulation. Overnight cultures of the *E. faecalis* strain indicated on the left were then added as a soft agar overlay. Plates were incubated at 25°C for 24 hours and zones of inhibition in the *E. faecalis* lawn were used to assess susceptibility to antimicrobials produced by the central strain.

325

We next tested the *E. faecalis* mutants against the nisin producers, *L. lactis* NZ9000 and *S. gallolyticus* AB39. Although there was some inhibition of wild type *E. faecalis* by the nisin-A producer *L. lactis*, there was no difference in sensitivity between the wild type and deletion strains. This suggests either that the genes of our regulatory pathway do not contribute to resistance against this antibiotic, or that under the chosen conditions the inhibitory activity of *L. lactis* NZ9000 is primarily due to a different
antimicrobial than nisin-A. In contrast, the nisin-P producer *S. gallolyticus AB39*, despite having no
inhibitory effect on wild type *E. faecalis*, was able to strongly inhibit growth of the *liaR* deletion strain.
Deletions of both *sapR* and *sapAB* also resulted in increased sensitivity but to a lesser degree than *liaR*deletion. Deletion of *rapAB* had little to no effect compared to the wild type, suggesting that the other
target genes of the regulatory pathway but not the RapAB transporter are responsible for nisin P
resistance.

337 These findings indicate that in response to different antimicrobials, the various members of this 338 resistance network have differing roles to play and also differ in their relative importance. In response to subtilin and nisin-P, LiaR clearly plays a very important role in resistance, but it is a less important 339 component in response to bacitracin exposure. This reflects LiaR's role as a key regulator in the cell 340 envelope stress response more widely, but with mostly a moderating role in response to bacitracin. The 341 342 sapR, sapAB and rapAB deletions all present with very similar effects in response to the antimicrobials, reflecting the interdependent functional relationship between the three components. The notable 343 344 exception to this is RapAB in the context of nisin-P, where the transporter does not appear to contribute to protection of the cell. 345

346

# 347 **Discussion**

In this study, we aimed to expand our understanding of the cell envelope stress response of *E. faecalis* 348 349 using the previously described bacitracin resistance network as our foundation. Our findings revealed 350 the regulation of the BceRS-type TCS SapRS via the damage sensor, LiaFSR. In addition, we also 351 demonstrated the regulation of the resistance operon *dltXABCD* via SapRS. This work describes a direct 352 functional link between the Lia and Sap signalling systems in *E. faecalis*, which can explain previous 353 observations of suppressor mutations in sapAB when a LiaR-deficient strain was evolved for DAP resistance<sup>41,42</sup>. Moreover, through creation of deletion strains and analysing promoter fusions, we have 354 355 revealed the presence of a differential network response through the damage-sensing LiaFSR and drug-

356 sensing SapAB-SapRS systems, leading to *dlt* expression in response to bacitracin, but not daptomycin. This discovery implies there is distinct control on subsections of the network depending on the inducer. 357 To understand the mechanisms of this differential control, the mode-of-actions of both antibiotics must 358 be considered. Bacitracin is a peptide antibiotic, which forms a complex with undecaprenyl 359 pyrophosphate (UPP), an intermediate of the Lipid II cycle<sup>35,36</sup>. The formation of this complex prevents 360 the dephosphorylation of UPP and slows cell wall formation<sup>59,60</sup>. In contrast, daptomycin is a 361 lipopeptide antibiotic which forms a tripartite complex between lipid II and the membrane phospholipid 362 phosphatidyl glycerol<sup>50</sup>. The formation of this complex leads to a loss of membrane potential and cell 363 364 death. The similarity between both antibiotics is the formation of cell envelope damage, the known stimulus for the activation of the "damage sensor" LiaFSR. However, they differ in their mechanism of 365 action with bacitracin preventing cell wall synthesis and daptomycin directly affecting cell membrane 366 integrity. 367

368 Since both antibiotics cause cell envelope damage, both lead to induction of LiaR target genes (Figures 369 1&3), including *sapRS*. SapAB, the sensory transporter that controls SapRS activity, however, so far has been shown to be activated by only two antibiotics, bacitracin and mersacidin <sup>33</sup>, both of which bind 370 to lipid II cycle intermediates<sup>35,61,62</sup>. This is consistent with the current working model for BceAB-type 371 transporters, to which SapAB belongs, recognising the complex between a peptide antibiotic and a lipid 372 II cycle intermediate as their substrate<sup>39</sup>. Importantly, the type of antibiotics to which BceAB-type 373 transporters respond all interact with the lipid II cycle intermediates via at least the pyrophosphate group 374 (e.g., bacitracin) or the sugar-pyrophosphate moiety of Lipid II (e.g. the lantibiotics, including 375 mersacidin)<sup>62,63</sup>. In the case of bacitracin, it is therefore the presence of bacitracin-UPP complexes that 376 triggers SapAB activity, leading to SapRS activation via the transport flux-sensing mechanism 377 demonstrated for the *B. subtilis* system<sup>34,38</sup>. Thus, the presence of bacitracin triggers both Lia and Sap 378 signalling simultaneously, inducing the expression of *dlt*. 379

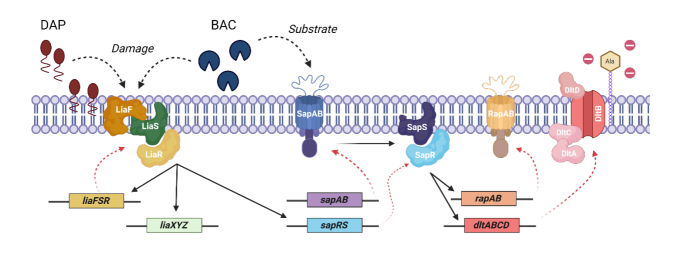
In contrast, DAP interacts with its cellular target in a fundamentally different way, forming a tripartite complex between Lipid II and phosphatidyl glycerol (PG)<sup>50</sup>. It therefore would not be expected to form the type of complexes that could act as substrate for SapAB<sup>39</sup>, and Sap signalling is not activated. BceAB-type transporters have been shown to repress histidine kinase activity in the absence of substrate<sup>38</sup>. Thus, the lack of a SapAB substrate likely leads to the transporter maintaining SapS in an 'OFF' conformation, preventing activation of SapR target genes, i.e. *dlt*. Such a mechanism can explain why in the presence of DAP, when the LiaFSR system activates expression of the *sapRS* operon, no induction of *dlt* expression was observed, because SapAB – unable to detect the tripartite DAP/lipid II/PG complex - would have prevented SapRS signalling.

389 Taken together, these findings indicate the presence of a logic 'AND' gate, whereby two inputs (the 390 damage sensing input provided by LiaFSR and the drug-sensing input provided by SapAB) are needed 391 to obtain the output, which is the induction of *dlt* expression (Figure 6). This mode of regulation offers an explanation for the mutations observed previously in *sapAB* when evolving for DAP resistance in 392 the absence of  $liaR^{41,42}$ . A gain-of-function mutation in sapAB would allow the activation of Sap 393 signalling in the absence of a substrate and induce *dlt* expression, essentially overriding the 'AND' gate 394 395 and providing DAP resistance. This working model also explains the low but statistically significant induction of *dlt* expression by bacitracin we observed in the absence of *sapAB* (Fig. 3). The mutant still 396 397 possesses SapRS and thus bacitracin-challenge leads to activation of sapRS expression via LiaR. This 398 in turn is likely to cause an increase in basal activity of SapRS-signalling and thus moderate dlt 399 activation, even if the full activation via SapAB is missing. A similar effect has been described previously for the second known SapRS target, rapAB, where deletion of sapAB resulted in low 400 promoter activity of *rapAB*, but with a small level of residual bacitracin response remaining<sup>33</sup>. 401

402 The regulatory wiring we have uncovered here between the LiaFSR and the Bce-type SapAB-SapRS systems, resulting in the control of the *dltXABCD* and *rapAB* resistance operons, is remarkably complex 403 404 compared with other Firmicutes bacteria. For example, in B. subtilis LiaFSR controls its own regulon, comprised solely of liaIH(G) and its own encoding genes<sup>24</sup>. In S. aureus, the LiaFSR homologue VraRS 405 has been shown to regulate the expression of multiple genes involved in cell wall biosynthesis<sup>64</sup>, but 406 has not been observed to regulate a BceRS-type TCS. Bce-type systems in most Firmicutes bacteria 407 either solely control their own transporter, as found with BceAB-RS in *B. subtilis*<sup>40</sup>, or can be involved 408 with other Bce-type systems, such as the BraRS-VraDE<sup>65</sup> and GraRS-VraFG<sup>66</sup> systems in S. aureus. In 409

- 410 some cases, Bce-type systems can also control additional genes, for example the GraRS TCS has also
- 411 been shown to regulate the *dlt* operon<sup>67-69</sup>. However, to our knowledge there have been no prior reports
- 412 of direct regulatory relationships that hardwire the Lia and Bce-type pathways together.

413



414

**Figure 6. Proposed model of the antibiotic resistance network.** Schematic illustrating the genes and proteins involved in antibiotic detection and resistance, with transcriptional regulation and translation demonstrated by black and red arrows respectively. Exposure to bacitracin (BAC) causes damage to the cell envelope, which acts as the stimulus for the removal of repression by LiaF, triggering the activation of LiaS. LiaS activity leads to phosphorylation of LiaR, which in turn induces the expression of its target promoters: its own, *liaXYZ* and *sapRS*, indicated by the black arrows. Simultaneously, bacitracin is also detected by SapAB. Communication between SapAB and SapS is indicated by a faded arrow, resulting in the activation of SapS and phosphorylation of SapR. SapR then induces the expression of the resistance genes, *rapAB* and *dlt*, which remove bacitracin from UPP and reduce the positive charge of the cell envelope, respectively. Conversely, exposure to daptomycin (DAP) again results in cell envelope damage, which triggers the phosphorylation of LiaR and induces the expression of *sapRS*. However, daptomycin is not recognised by SapAB, therefore SapS is kept in an 'OFF' conformation and SapR is not phosphorylated. SapR is therefore unable to induce the expression of *rapAB* and *dlt*.

415

416 Our findings reported here show that although the enterococci possess the same regulators as *S. aureus* 417 and *B. subtilis*, evolution has taken these components down different routes. A potential reasoning 418 behind *E. faecalis* maintaining this intricate setup could be that having *dlt* and *rapAB* under Sap control, 419 and *sapRS* itself under Lia regulation, may provide the cell with a layered response. Under this premise,

420 when the cell is experiencing a mild bacitracin attack, signalling in the Lia-Sap pathway is likely to be 421 dominated by the Sap system, which responds to the presence of UPP-bacitracin complexes, even if those do not yet cause cellular damage. Sap signalling then activates both rapAB and dltXABCD. Of 422 these, RapAB provides the main target protection mechanism, especially at lower antibiotic 423 424 concentrations, whereas expression of dlt is likely of minor importance, based on evidence from B. subtilis<sup>39,45</sup>. This is consistent with our observations in *E. faecalis*, where the strength of *rapAB* 425 induction is much higher than that of *dlt*: at  $4\mu g/mL$  bacitracin, *rapAB* expression reaches >100 MU<sup>33</sup>, 426 compared to *dlt*, which reaches a maximum of 25 MU (Fig. 3). However, when the antibiotic threat 427 428 begins to exceed the protection level RapAB can provide, damage to the cell occurs and the Lia system 429 is activated. LiaFSR is then able to boost the signalling cascade by increasing *sapRS* expression. We see this in the absence of *rapAB*, which mimics an extreme failure of target protection, leading to a 430 431 drastic increase in *dlt* expression (Fig. 3). Thus, the regulatory structure between Lia and Sap signalling in *E. faecalis* allows a second line of defence to be mounted if the primary resistance system is no longer 432 433 sufficient to fully protect the cell.

434 The genes discussed here are all part of the core genome of E. faecalis and thus contribute to intrinsic, rather than acquired drug resistance in this bacterium. To better understand the evolutionary context 435 that appears to have given rise to the unusually complex regulation of the resistance genes, we 436 considered the broader scope of this network - was it able to protect the enterococci from antimicrobial 437 438 challenge from potential competitors in their natural environments? The importance of the role played 439 by LiaFSR in the cell envelope stress response was well demonstrated by the antagonism assays, in which deletion of *liaR* resulted in increased sensitivity against all tested antimicrobial producers, except 440 for L. lactis NZ9000 (nisin-A). In contrast, components of the Sap system only contributed to protection 441 442 from the bacitracin producer B. lichenformis ATCC10716 and the subtilin producer B. subtilis ATCC6633. The only antimicrobial producer against which neither of the genes appeared to give a 443 protective effect was the nisin-A producer L. lactis NZ9000. However, we cannot currently differentiate 444 whether this was because L. lactis did not produce sufficient nisin-A under the chosen growth conditions 445 446 to detect differences, or because E. faecalis uses alternative resistance mechanisms against nisin-A not controlled by either the Lia or Sap systems. Overall, it appears that the regulatory network studied here
plays a key role in protecting *E. faecalis* from antimicrobial peptide producing bacteria that it would
realistically encounter in its natural habitats, offering an explanation for the existence of such a complex
regulatory strategy.

451 One of our most striking results was that daptomycin challenge, although able to activate Lia signalling, 452 was unable to activate the Sap system, potential mechanistic reasons for which are explained above. 453 This was a surprising finding, as *dlt* is known to respond to daptomycin in other Firmicutes, e.g. S. aureus <sup>70–74</sup>. The lack of *dlt* expression in response to DAP may, however, explain the sensitivity to this 454 455 antibiotic observed throughout the *Enterococcus* genus and its effectiveness when treating VRE infections<sup>75</sup>. This is despite the bacteria being in possession of an effective resistance mechanism, except 456 457 this is not induced upon DAP challenge. It is then plausible to see why clinical DAP resistance could result from overriding of the molecular controls investigated here to trigger high *dlt* operon expression, 458 specifically via mutations in genes encoding the components of the LiaFSR system<sup>19</sup>, or experimentally 459 in sapAB as suppressor mutations of a LiaR-deficient strain<sup>41,42</sup>. DAP was introduced for treatment of 460 enterococci infections in 2003<sup>12</sup> and the first resistant isolates were identified already in 2005<sup>13</sup>. It may 461 be that the rapid emergence of DAP resistance is in part derived from a dysregulation of *dlt* expression 462 463 by such mutations of regulatory components.

464 Overall, our findings contribute to an increasing understanding of the regulatory network protecting *E*. 465 *faecalis* from cell envelope attack and provide insights into the regulatory cascade that is involved in 466 controlling resistance gene expression in response to cell envelope targeting antibiotics. Our findings 467 also provide valuable context in which to better understand the emergence of DAP resistance in clinical 468 environments. Ultimately, a system-wide understanding of antimicrobial resistance regulation may lead 469 to identification of an 'Achilles' heel' within the network and identify new therapeutic targets.

470

### 471 **Experimental Procedures**

472 Bacterial strains and growth conditions. All bacterial strains and plasmids used in this study are listed in Table S1 in the supplementary material. E. coli MC1061 was used for cloning with pTCVlac, and 473 strain DH5a was used for all other cloning. E. coli, Bacillus licheniformis and Bacillus subtilis were 474 routinely grown in lysogeny broth (LB) at 37°C with agitation (200 rpm). Lactococcus lactis was grown 475 476 routinely in M17 supplemented with 0.5% lactose at 30°C without agitation. Enterococcus faecalis and Streptococcus gallolvticus were grown routinely in brain heart infusion (BHI) at 37°C without agitation, 477 with media for the latter being supplemented with 5% (v/v) foetal bovine calf serum. Solid media 478 contained 15 g l<sup>-1</sup> agar. Selective media contained chloramphenicol (10  $\mu$ g ml<sup>-1</sup> for *E. coli*; 15  $\mu$ g ml<sup>-1</sup> 479 for *E. faecalis*), kanamycin (50  $\mu$ g ml<sup>-1</sup> for *E. coli*; 1000  $\mu$ g ml<sup>-1</sup> for *E. faecalis*), spectinomycin (100 480  $\mu$ g ml<sup>-1</sup> for *E. coli*; 500  $\mu$ g ml<sup>-1</sup> for *E. faecalis*). For blue-white screening, 5-bromo-4-chloro-3-indolyl-481  $\beta$ -d-galactopyranoside (X-Gal) was used at 120  $\mu$ g/ml. Bacitracin was supplied as the Zn<sup>2+</sup> salt. All 482 483 media for experiments with daptomycin were supplemented with 50 µM CaCl<sub>2</sub>.

484 *E. faecalis* was transformed by electroporation as previously described<sup>76</sup>. *E. coli* was transformed by 485 heat-shock of CaCl<sub>2</sub> competent cells, followed by 1 hour recovery time<sup>77</sup>. Growth was measured as 486 optical density at 600 nm (OD<sub>600</sub>) on a the Biochrom<sup>TM</sup> Novaspec Pro Spectrophotometer using cuvettes 487 with 1 cm light path length or in 96-well plates with 100  $\mu$ L culture volumes on a Spark<sup>®</sup>Microplate 488 reader (Tecan).

489 Construction of plasmids and genetic techniques. All primer sequences used for cloning are listed in 490 Table S2 in the supplementary material. Transcriptional promoter fusions to *lacZ* in *E. faecalis* were constructed in the vector pTCVlac<sup>78</sup>. All fragments were cloned via the EcoRI and BamHI sites of the 491 vector. Constructs for unmarked deletions in *E. faecalis* were cloned into pLT06<sup>79</sup>. For each gene or 492 operon to be deleted, 700- to 1000-bp located immediately before the start codon of the first gene ("up" 493 fragment) and after the stop codon of the last gene ("down" fragment) were amplified. The primers 494 were designed to create a 17- to 20-bp overlap between the PCR products (Table S2), facilitating the 495 fusion of the fragments by PCR overlap extension<sup>80</sup> and were subsequently cloned into the NcoI and 496 BamHI site of the vector pLT06. Gene deletions were performed as previously described<sup>79</sup>. Briefly, 497 following transformation of the parent strain with the temperature-sensitive vector pLT06, overnight 498

499 cultures were grown at 30° containing chloramphenicol and reinoculated 1:1000 into 10 mL BHI the next morning. Cells were then grown at 30° for 2.5 hours, followed by increasing to 42° for a further 500 2.5 hours to force single-site integration. Cells were then serially diluted onto BHI agar containing 501 chloramphenicol and X-Gal and incubated at 42°. Blue colonies growing at 42°C were screened for the 502 503 targeted integration using PCR with primers flanking the site of integration. Positive clones were then serially passaged for two days from overnight culture in BHI medium with no selection at  $30^{\circ}$  to allow 504 a second site recombination event. Cultures were then serially diluted on to MM9-YEG agar<sup>81</sup> 505 containing 10 mM p-chloro-phenylalanine for counter-selection and X-Gal at 37°C. The resulting white 506 colonies were screened for the deletion of the target genes by PCR. All cloned constructs were checked 507 508 for PCR fidelity by Sanger sequencing, and all created strains were verified by PCR using the primers 509 given in Table S2.

Antimicrobial susceptibility assays. For antibiotic susceptibility assays, minimum inhibitory 510 511 concentrations (MICs) were determined by broth dilution assays in BHI medium in 96-well plates, containing serial two-fold dilutions of antibiotic. Colonies of E. faecalis were suspended in sterile 512 Phosphate Buffered Saline (PBS) to 0.5 McFarland standard turbidity and diluted 1:1,000 in a total 513 volume of 100µL. After 24 h incubation at 37°C, growth was determined by measuring optical density 514 515 (OD<sub>600</sub>) on a Spark<sup>®</sup>Microplate reader (Tecan). The MIC was scored as the lowest antibiotic concentration where no growth was observed following subtraction of the  $OD_{600}$  of a well containing 516 517 sterile medium.

518  $\beta$ -Galactosidase assays. For quantitatively assessing induction of *lacZ* reporter constructs in *E*. *faecalis*, exponentially growing cells ( $OD_{600} = 0.4 - 0.5$ ) inoculated 1:250 from overnight cultures in BHI 519 520 medium were exposed to different concentrations of bacitracin for 1 h or daptomycin for 30 minutes. Cells were harvested via centrifugation and stored at -20°C. β-Galactosidase activities were assayed in 521 permeabilised cells and expressed in Miller units (MU)<sup>82</sup>. For this, cells were resuspended in 1 ml Z-522 buffer (8.04 g Na<sub>2</sub>HPO<sub>4</sub>\*7H<sub>2</sub>O, 2.76 g NaH<sub>2</sub>PO<sub>4</sub>\*H<sub>2</sub>O, 0.123 g MgSO<sub>4</sub>\*7H<sub>2</sub>O and 5 mL 1M KCl in 495 523 mL dH<sub>2</sub>O, pH 7). The samples were adjusted to  $OD_{600} = 0.5$  in a 1 ml volume of Z-buffer and from this, 524 two volumes were taken: 200 µl and 400 µl cells made up to 1 mL each with Z-buffer. This volume 525

corresponds to the 'volume of cells' in the Miller Unit (MU) equation below. Following this, 20 μl 0.1% (w/v) SDS and 40 μl chloroform were added and vortexed for 5 seconds, then rested for 5-10 minutes. Reactions were started by adding 200 μl *o*-nitrophenyl-β-D-galactopyranoside (ONPG) (4 mg mL<sup>-1</sup> in Z-buffer) and incubated at room temperature until yellow colouration was observed. If no colour change was visible, the reaction was incubated for 20 minutes. Reactions were stopped by adding 500  $\mu$ l 1M Na<sub>2</sub>CO<sub>3</sub>, and the time recorded, which corresponds to the 'time' in the Miller Unit (MU) equation below. Absorbance at 420 nm (A<sub>420</sub>) was then read. MUs were calculated using the following equation:

533 
$$Miller Units (MU) = \frac{A420 * 1000}{Time (minutes) * volume of cells (in ml) * 0D600}$$

Computational modelling. Rather than directly modelling the temporal dynamics of the regulatory network as done for the *B. subtilis* Bce system<sup>34,83</sup>, we chose to focus on the (meta) stable state reached in response to challenge with a given bacitracin concentration. The quantities modelled are the concentrations of bacitracin [*bac*], UPP [*UPP*], UPP-bound bacitracin [*UPPbac*], and the effective activities of SapRS [*SapRS*], RapAB [*RapAB*], LiaX [*Lia*], and DltXABCD [*Dlt*]. For the modelling, we made repeated use of a soft-switch sigmoid type function

540 
$$f_{y}(x; y_{\text{base}}, y_{\text{max}}, x_{0}, \mathbf{k}) = f_{\text{max}} \left( 1 + \left( \frac{y_{\text{max}}}{y_{\text{base}}} - 1 \right) e^{-k(x-x_{0})} \right)^{-1},$$

541 where x is the input quantity and y the output which varies between  $y_{base}$  and  $y_{max}$ , with x<sub>0</sub> controlling 542 the switching threshold and k setting the sharpness of the transition. The equations of state for the model 543 are then simply

544 
$$k_{on}([Dlt])([bac] - [UPPbac])[UPP] - k_{off}([RapAB])[UPPbac] = 0$$

 $[Lia] = f_{Lia}([UPPbac])$ 

546 
$$[SapRS] = f_{SapRS}([Lia])$$

547 
$$[Dlt] = f_{dlt}([SapRS])$$

548 
$$[RapAB] = f_{RapAB}([SapRS])$$

549 We modelled the binding and off rates for the UPP / bacitracin interaction as being linearly proportional 550 to [*Dlt*] and [*RapAB*], respectively, with Dlt reducing bacitracin binding and RapAB increasing the off-551 rate. The biological reasoning for this is explained in the Results section.

For [*bac*], actual values from the experimental work were used, and the switching parameters for [*Dlt*] and [*RapAB*] as functions of [*SapRS*] were determined by fitting actual activity levels observed experimentally of these quantities for different bacitracin levels. The remaining parameters were fitted to achieve a description of experimental results for the wild-type strain by the model output. The values for each parameter are given in Table S3. Solving the equations of state for a given input bacitracin level [*bac*] yielded predictions for [*UPP*] and [*UPPbac*], which in turn drive the response curves plotted in Figure 3B.

559 Antagonism assays on solid media. A plate-based assay was utilised to measure growth inhibition between antibiotic producer strains and E. faecalis strains. For each species, cultures of the 560 antimicrobial-producing bacteria were grown overnight in the respective growth medium and 561 562 temperatures stated above and adjusted to  $OD_{600} = 0.5$ . Aliquots (5 µl) from each producer culture were spotted onto the centre of a BHI agar plate and incubated at room temperature (20-25°C) for 3-5 days 563 to allow accumulation of antimicrobial products. Overnight cultures of E. faecalis strains were grown 564 at  $37^{\circ}$ C in BHI medium, inoculated 1:100 into fresh medium and grown to OD = 0.5. Next, 3 mL of 565 566 liquid BHI soft agar (7.5 g/L, 50°C) were inoculated with 30  $\mu$ L culture. The soft agar was then poured onto the plate containing the pre-grown antimicrobial-producing strains and left to dry. The plate was 567 then incubated overnight at room temperature to allow E. faecalis to grow to visualise the zone of 568 inhibition. Results were recorded photographically using a PowerShot G camera attached to a lightbox. 569

570 **Data accessibility.** The numerical data underpinning the results shown in Figures 1, 3 and 4 are 571 available in Table S4. The *Mathematica* file for the model is provided in the supplementary material.

572

#### 573 Acknowledgements

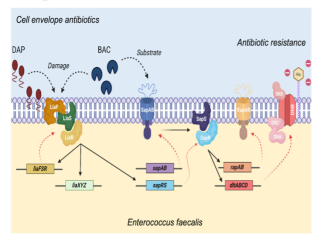
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# 580 Author contributions

- 581 SMM, SG, TR and GF contributed to the conception of the study; SMM and SG designed the
- 582 experimental work; SMM carried out all data acquisition; SMM, TR and SG carried out data analysis
- and interpretation; SMM, SG, TR and GF wrote the manuscript.

584

# 585 **Graphical abstract**



586

# 587 Abbreviated summary

- 588 This study explored a regulatory network controlling resistance against daptomycin and bacitracin in
- 589 *E. faecalis.* We show that the *dltXABCD* operon, which can protect against both antibiotics, is
- 590 controlled by two regulatory systems, LiaFSR and SapRS. Strikingly, this strategy allows *dtlXABCD*
- 591 expression only in response to bacitracin, but not daptomycin, potentially explaining the natural DAP-
- sensitivity of *E. faecalis* and why clinical resistance can emerge via regulatory mutations.

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### 837 Supplementary Material

- 838 Table S1. Vectors, plasmids and strains used in this study.
- 839 Table S2. Primers used in this study.
- 840 Table S3. Parameter values used in the mathematical model.
- **841 Table S4.** Experimental numerical data for Figures 1, 3 and 4.

842 File S1. Mathematical model file for the application *Mathematica*