Anatomy of the bacitracin resistance network in *Bacillus subtilis*

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Abstract

Protection against antimicrobial peptides (AMPs) often involves the parallel production of multiple, well-characterized resistance determinants. So far, little is known about how these resistance modules interact and how they jointly protect the cell. Here, we studied the interdependence between different layers of the envelope stress response of *Bacillus subtilis* when challenged with the lipid II cycle-inhibiting AMP bacitracin. The underlying regulatory network orchestrates the production of the ABC transporter BceAB, the UPP phosphatase BcrC and the phage-shock proteins LiaIH. Our systems-level analysis reveals a clear hierarchy, allowing us to discriminate between primary (BceAB) and secondary (BcrC and LiaIH) layers of bacitracin resistance. Deleting the primary layer provokes an enhanced induction of the secondary layer to partially compensate for this loss. This study reveals a direct role of LiaIH in bacitracin resistance, provides novel insights into the feedback regulation of the Lia system, and demonstrates a pivotal role of BcrC in maintaining cell wall homeostasis. The compensatory regulation within the bacitracin network can also explain how gene expression noise propagates between resistance layers. We suggest that this active redundancy in the bacitracin resistance network of *B. subtilis* is a general principle to be found in many bacterial antibiotic resistance networks.

Keywords

PspA, phage-shock protein, antibiotic resistance, envelope stress response, signal transduction, gene regulation
Introduction

In their natural environment many microbes are in fierce competition for a limited supply of resources. This frequently involves the production of antimicrobial peptides (AMPs) that suppress the proliferation of competitors (Eijsink et al., 2002). In this biochemical warfare, the cell envelope serves as a prime target, and many AMPs interfere with its biosynthesis and integrity (Breukink and de Kruijff, 2006). To defend against antimicrobial attacks by rival species, it is thus of vital importance for cells to accurately sense these cues and to swiftly mount protective countermeasures, collectively referred to as cell envelope stress response (CESR) (Jordan et al., 2008; Schrecke et al., 2012). In many bacteria the defense against AMPs involves the simultaneous expression of a number of resistance systems that protect cells at various levels. Those include on the one hand specific resistance determinants, such as ABC transporters (Gebhard, 2012) and immunity lipoproteins (Stein et al., 2003; Aso et al., 2005) that transport and/or sequester AMPs from their molecular targets. On the other hand, bacteria induce the production of more nonspecific resistance determinants that alter the charge and composition of the cell envelope to reduce access of AMPs to their sites of action (Revilla-Guarinos et al., 2014) and allow cells to cope with deleterious effects on downstream cell physiology (Joly et al., 2010). While many of the AMP resistance modules have been individually characterized in great detail, our present knowledge about how these modules interact, and how they jointly contribute to the overall AMP resistance of a cell, is still limited. Thus, as for many other bacterial stress responses, the daunting task is to decipher how the cell orchestrates the activity of individual resistance modules into a complex and multi-layered CESR network.

In the present work we approached this question by focusing on the resistance mechanisms of Bacillus subtilis against the peptide antibiotic bacitracin, which is produced by some strains of Bacillus licheniformis and B. subtilis (Azevedo et al., 1993; Ishihara et al., 2002) and is clinically used as broad spectrum antibiotic against Gram-positive bacteria causing skin infections. Bacitracin acts by inhibiting the lipid II cycle of cell wall biosynthesis, which is essential for the translocation of peptidoglycan precursors from the cytosol to the extracytoplasmic space (Fig. 1A). The tight complex formation between bacitracin and the diphosphate lipid carrier undecaprenyl pyrophosphate (UPP) prevents dephosphorylation of UPP to undecaprenyl phosphate (UP) and thereby efficiently blocks recycling of the lipid carrier (Storm and Strominger, 1973; Economou et al., 2013).

To perpetuate progression of the lipid II cycle under bacitracin attack and to protect against cell envelope damage, B. subtilis up-regulates the expression of three major resistance modules (Mascher et al., 2003; Rietkötter et al., 2008): The ABC transporter BceAB (Ohki et al., 2003; Mascher et al., 2003), the UPP phosphatase BcrC (Cao and Helmann, 2002; Ohki et al.,
2003; Bernard et al., 2005) and the phage shock protein (Psp)-like Lial and LiaH proteins (Mascher et al., 2004; Jordan et al., 2006) (Fig. 1A and B). Recent evidence suggests that BceAB confers resistance by clearing UPP from the inhibitory grip of bacitracin (Fritz et al., 2015), but it remains elusive whether bacitracin is transported into the cytoplasm for degradation or whether it is released into the extracytoplasmic space, as suggested previously (Rietkötter et al., 2008; Ohki et al., 2003). Simultaneously, the phosphatase BcrC catalyzes the dephosphorylation of UPP to UP (Fig. 1A) and thereby promotes the progression of the Lipid II cycle. Finally, under cell-envelope perturbing conditions the liaIH operon is induced, and the small membrane anchor protein Lial recruits the cytosolic PspA/IM30 protein family member LiaH into static, membrane-associated patches (Domínguez-Escobar et al., 2014). While the homologous Psp system encoded by the pspABCDE operon of Escherichia coli has been linked to maintenance of the proton motive force under envelope-perturbing conditions (Kleerebezem et al., 1996; Kobayashi et al., 2007), the physiological role of the Lia system in B. subtilis remained elusive: despite its more than ~100-fold induction under bacitracin stress, no increase in bacitracin sensitivity was detected in a liaIH deletion strain (Wolf et al., 2010). While this might suggest that there is no contribution of the Lia system to bacitracin resistance, we reasoned that the presence of the two other bacitracin resistance layers, BceAB and BcrC, could potentially compensate for the lack of LiaIH. However, to date it is not known whether these systems act in fact redundantly, or whether they contribute independently or even cooperatively to bacitracin resistance.

To gain deeper insight into how these modules interact and form an efficient bacitracin stress response network, we here systematically studied their functional and regulatory interactions in a comprehensive set of mutants deficient in the three resistance determinants. Our analysis reveals a hierarchy among resistance modules, which we find reflected in marked anti-correlations between the expression of primary (drug-sensing) and secondary (mostly damage-sensing) layers of bacitracin resistance. This means that the increased expression of the primary resistance layer reduced the expression of the secondary layer and vice versa. Strikingly, these anti-correlations can also explain how gene expression noise propagates between the different resistance modules at the single cell level, as revealed by flow cytometry analyses. Moreover, our study underpins the importance of the UPP phosphatase BcrC for cell wall homeostasis in the absence of bacitracin stress and provides novel clues about the physiological stimuli triggering the induction of the modules in the bacitracin resistance network.
**Results**

**Contributions of CESR modules to antibiotic resistance**

First, we studied whether the three CESR modules protect the cell in a redundant, independent or in a cooperative manner. To this end we constructed mutants deficient in one, two or in all three resistance determinants and determined their sensitivity towards bacitracin using the E-test® agar gradient diffusion method (Fig. 2A). Compared to the minimal inhibitory concentration (MIC) of bacitracin for wild type cells (256 μg/ml), mutants deficient in only one of the resistance modules displayed a clear hierarchy in their sensitivity towards bacitracin: While the MIC of the ΔliaIH mutant was identical to that of the wild type, the ΔbcrC mutant displayed a 5-fold and the ΔbceAB mutant an 85-fold increase in bacitracin susceptibility, suggesting that BceAB acts as the primary resistance determinant under these growth conditions. Interestingly, in a mutant background devoid of bceAB, the additional deletion of either of the other two resistance modules had a significantly stronger impact on the MIC than observed in the single mutants. Here, the ΔbceAB ΔliaIH double mutant had a 6-fold lower MIC than the ΔbceAB mutant, thereby revealing the first phenotype of LiaIH in the bacitracin stress response. Hence, we suggest that the previously reported lack of a ΔliaIH phenotype upon bacitracin stress (Rietkötter et al., 2008) might be explained by a redundant organization of the bacitracin stress response network, in which resistance conferred by BceAB masks the weaker contribution of the LiaIH module. Moreover, the ΔbceAB ΔbcrC double mutant was 24-fold more sensitive than the ΔbceAB reference strain, suggesting that BceAB also partially masks the contribution of BcrC. Please note that we did not observe a similar “masking effect” between the secondary resistance modules, as the MIC of a ΔbcrC mutant (48 μg/ml) was identical to that of a ΔbcrC ΔliaIH mutant (Fig. 2A). Only when compared to a ΔbceAB ΔbcrC double mutant, we found that a ΔbceAB ΔbcrC ΔliaIH triple mutant showed an ~3-fold increased bacitracin sensitivity (Fig. 2A). In summary, these results show that the secondary resistance modules do in fact protect the cell against bacitracin, but also reveal that the contributions of the secondary resistance modules are masked by the much stronger resistance conferred by the Bce system.

Next, we asked whether the increased bacitracin susceptibility of the mutants above was in fact due to the lack of the respective resistance modules, or whether those mutants exhibited a general growth defect that might result in increased bacitracin susceptibility. For instance, it is known that the BcrC phosphatase is also involved in Lipid II cycle progression under normal growth conditions (Bernard et al., 2005), but the extent to which the cytosolic UPP phosphatase UppP (formerly YubB) could compensate for the deletion of BcrC was controversial (Cao and Helmann, 2002; Bernard et al., 2005). To quantitatively test the fitness of the different mutants, we measured their doubling times in LB medium at 37 °C in a microplate reader (Fig. 2B). In the
absence of bacitracin, the wild type and ΔliaIH mutant grew at similar doubling times of \( t_d = 23.0 \pm 2.2 \) min, and \( t_d = 24.7 \pm 2.1 \) min, respectively, while the ΔbbeAB mutant grew slightly faster (\( t_d = 20.0 \pm 0.3 \) min) and the ΔbcrC mutant significantly slower (\( t_d = 28.7 \pm 1.7 \) min) than wild type (P value of unpaired Student’s t-test = 0.024). Moreover, we observed that under these conditions of rapid growth, the ΔbcrC mutant was about as sensitive as the ΔbbeAB mutant, which displayed killing at 10 µg/ml bacitracin and higher (Fig. 2B). This suggests that at high growth rates the deletion of bcrC can only be partially compensated for by the activity of the second UPP phosphatase UppP,, implying that UPP dephosphorylation might become the bottleneck for cell wall biosynthesis and hence for cell growth. Thus, we conclude that the increased bacitracin sensitivity of the ΔbcrC mutant can – at least partially – be attributed to a general growth defect incurred by reduced rates of UPP dephosphorylation.

**Regulatory interactions between the CESR modules**

The redundant contributions of the CESR modules to bacitracin resistance described above provoked the question of the extent to which deletion of one resistance module would affect the expression of the other resistance modules. To study these regulatory interactions, we fused the target promoter of each module to the luxABCDE cassette derived from *Photorhabdus luminescens* (Schmalisch et al., 2010; Radeck et al., 2013) and integrated the resulting reporter plasmids into the chromosome of wild type and mutants deficient in one of the three resistance modules (Table S1). Subsequently, exponentially growing cultures (OD

**Quantitative behavior of the unperturbed CESR network.** In the wild type strain (Fig. 3, black data), \( P_{bbeA} \) (Fig. 3A) displayed low activity (10^4 RLU/OD) in the absence of bacitracin and responded already at low bacitracin concentrations of \( \geq 0.01 \mu g/m^l \). This response gradually increased with rising bacitracin levels and reached its maximum about 300-fold over background at 30 µg ml^-1 bacitracin. Recently, we showed that this gradual response over a high input-dynamic range is the result of negative feedback regulation in the Bce system, in which a flux-sensing mechanism homeostatically adjusts the rate of de novo transporter synthesis to the level needed for cell protection (Fritz et al., 2015). In contrast to \( P_{bbeA} \), \( P_{bcrC} \) (Fig. 3B) already had a high basal activity (7x10^5 RLU/OD) and only responded at much higher bacitracin concentrations (1 µg ml^-1) with a maximum 3-fold induction over background at 30 µg ml^-1. The strong \( P_{bcrC} \) activity in the absence of antibiotic treatment is consistent with the notion that BcrC is an important player in lipid II cycle progression under exponential growth conditions, as noted
above. Similar to $P_{bceA}$, $P_{liaI}$ (Fig. 3C) displayed a low basal activity and a strong (400-fold) induction at high bacitracin levels, but its input-dynamic range was much narrower (0.1 to 10 µg ml$^{-1}$ bacitracin) than seen for $P_{bceA}$ (0.01 to 30 µg ml$^{-1}$ bacitracin). Hence, production of the primary resistance determinant BceAB is induced already at lower antibiotic concentrations than expression of the secondary resistance modules. This suggests that the primary layer might “buffer” against cell envelope stress at low bacitracin levels, while the demand for further protective measures only occurs at higher antibiotic concentrations.

**BceAB is the pacemaker of the CESR network.** If this buffering hypothesis was accurate, the secondary layer should become more sensitive and also more active in the absence of the primary resistance. Indeed, we found that in a $\Delta bceAB$ mutant the $P_{bcrC}$ and $P_{liaIH}$ promoters were activated already at lower bacitracin concentrations and displayed a steeper dose-response behavior than in the wild type (Fig. 3B and C, *blue data*). Note that the activity of $P_{bceA}$ itself remained at a basal level in the $\Delta bceAB$ mutant (Fig. 3A), again highlighting that the transport activity of BceAB is strictly required for activation of the $P_{bceA}$ promoter (Rietkötter et al., 2008; Fritz et al., 2015). To further corroborate the buffering hypothesis, we next tested the effect of different constitutive BceAB levels on the expression of the secondary resistance layer. To this end, we complemented the $\Delta bceAB$ mutant with a xylose-inducible copy of *bceAB*. Strikingly, compared to the highly sensitive $P_{liaI}$ response in the $\Delta bceAB$ mutant (Fig. 4A (ii); *red data*), constitutive expression of *bceAB* at low levels was already sufficient to shift the induction threshold of the $P_{liaI}$ promoter to 3-fold higher bacitracin levels (Fig. 4A (ii); *orange data*). A high constitutive expression level of *bceAB* resulted in a further 10-fold increase of the $P_{liaI}$ induction threshold (Fig. 4A (ii); *light green data*), which could be even further increased by overexpression of *bceAB* in the wild type (Fig. 4A (ii); *dark green data*). Importantly, varying the *bceAB* expression level caused similar shifts in the induction threshold of the $P_{bcrC}$ promoter (Fig. S1). Hence these data show that whenever the production level of BceAB is high, the expression of the two secondary resistance modules is low and *vice versa*. These clear-cut anti-correlations suggest that the ABC transporter actively prevents cell envelope stress and thereby reduces the demand for expression of the secondary layers of the CESR network.

Note that the variation of the *bceAB* expression levels also triggered shifts in the response of the $P_{bceA}$ promoter itself (Fig. 4A (i)). Previously, we showed that this behavior can be rationalized by a flux-sensing mechanism, in which a sensory complex between the ABC transporter BceAB and the histidine kinase BceS detects the rate of bacitracin flux by individual transporters, which in turn activates the $P_{bceA}$ promoter via the response regulator BceR (Fritz et al., 2015). Accordingly, in cells with low BceAB levels the load per transporter saturates already
at low bacitracin levels and triggers full induction of $P_{bceA}$ (Fig. 4A (i); orange curve). Conversely, in cells with higher BceAB levels the load per transporter saturates at significantly higher bacitracin levels, which in turn leads to proportional shifts of the $P_{bceA}$ dose-response characteristic to the right (Fig. 4A (i); green curves) (Fritz et al., 2015).

**BcrC has pleiotropic effects on CESR modules.** The deletion of $bcrC$ triggered a 2- to 3-fold increased activity of its own promoter, $P_{bcrC}$, compared to the wild type (Fig. 3B, green data) – even in the absence of bacitracin stress. Given that the deletion of $bcrC$ slowed down growth by impairing cell wall biosynthesis (Fig. 2B), the elevated $P_{bcrC}$ activity seemed reasonable, because this promoter belongs to the regulon of the alternative sigma factor $\sigma^M$ (Cao and Helmann, 2002). $\sigma^M$ itself responds to a broad spectrum of cell envelope-perturbing agents (Eiamphungporn and Helmann, 2008) and was therefore considered to be a sensor for cell wall integrity (Inoue et al., 2013; Lee and Helmann, 2013). Likewise, the $P_{liaI}$ promoter activity was elevated 3-fold in the $\Delta bcrC$ mutant (Fig. 3C), consistent with the role of the Lia system as a general sensor of cell envelope stress (Wolf et al., 2012). However, it was surprising that the $P_{bceA}$ promoter was also up-regulated 10-fold in the $\Delta bcrC$ mutant (Fig. 3A), since previous reports were consistent with a model in which the Bce system responds to the detoxification flux of the drug and not to downstream damage on cell physiology (cf. Fig. 1C). This curious effect is discussed in more detail below.

To further substantiate that the observed phenotypes specifically arose from the deletion of $bcrC$, we complemented the $\Delta bcrC$ mutant with a xylose-inducible copy of $bcrC$. This complementation indeed returned the elevated activities of $P_{bceA}$ and $P_{liaI}$ back to wild-type levels (Fig. 4B; light green data). Interestingly, the overexpression of $bcrC$ in a wild type background lead to a further decrease of both the $P_{liaI}$ and $P_{bceA}$ activities (Fig. 4B; dark green data), suggesting that an elevated rate of UPP dephosphorylation reduced the cellular susceptibility to bacitracin. Taken together, these data show that the level of BcrC sets the rate of UPP dephosphorylation, which in turn determines how many UPP target molecules the cell displays for binding by bacitracin. Accordingly, low levels of BcrC lead to the accumulation of UPP and make cells vulnerable to bacitracin attack, whereas high BcrC levels keep UPP levels low and make cells more resistant. This pattern is reflected both in the responses of the Lia and BcrC systems, which measure bacitracin-dependent damage of the cell envelope, as well as in the response of the Bce system, which presumably senses the UPP-bound form of bacitracin (see Discussion for more details).
LialH plays a positive autoregulatory role. In contrast to the marked effects the deletions of bceAB and bcrC had on the expression of all resistance modules, the deletion of liaH (Fig. 3, orange data) did not significantly influence the regulation of P$_{bceA}$ and P$_{bcrC}$ (Fig. 3A and B). However, the ΔliaIH mutant displayed up to 7-fold reduced activity of its own promoter (Fig. 3C). This is the first report showing that the expression of the lia operon is not only regulated via the LiaFSR three-component system (Schrecke et al., 2013), but that also the target proteins LiaH play a positive autoregulatory role required for the full Lia response. In fact, when scrutinizing the temporal dynamics of promoter activities, it became evident that the ΔliaIH mutant displayed only a transient P$_{lia}$ induction that reached a peak between 10-20 min after bacitracin addition and declined afterwards, whereas the wild type displayed prolonged P$_{lia}$ activity with a peak at ~40 min after bacitracin addition (Fig. S2).

In line with these observations, in a liaIH complementation strain variations of the LiaH production level did not affect the dose-response behavior of the P$_{bceA}$ promoter, but had significant effects on P$_{lia}$ activity itself (Fig. 4C). In the absence of bacitracin, the constitutive expression of liaH triggered a 20-fold increased P$_{lia}$ activity compared to the wild type (Fig. 4C (ii); light green data). At bacitracin concentrations higher than 0.3 µg ml$^{-1}$, however, P$_{lia}$ displayed a weaker activity than in wild type. These data show on the one hand that LiaH has a positive regulatory effect on the P$_{lia}$ promoter even in the absence of externally added antibiotics. On the other hand they show that the inability to up-regulate liaH lead to reduced P$_{lia}$ activity, suggesting that a positive feedback via LiaH might be needed for the full activation of the Lia system in wild type. To rule out that the marker-less deletion of liaIH had polar effects on the expression of the signaling system LiaFSR, we complemented the ΔliaIH mutant with a copy of liaIH under the control of its native promoter (P$_{lia}$), and found that wild type behavior of P$_{lia}$ induction could be restored (data not shown). Moreover, the overproduction of LiaH in a wild type background (Fig. 4C (ii); dark green data) lead to an elevated P$_{lia}$ activity in the absence of bacitracin, while at high bacitracin levels the Lia system was as active as in the wild type. Taken together, these results show that LiaH has no influence on the primary resistance BceAB, but is instead involved in fully activating and perpetuating its own expression by a so far unknown mechanism. In the future, it remains to be clarified whether LiaH is involved in the perception of cell envelope stress, or whether LiaH generates some degree of envelope stress itself.

Single cell induction of CESR modules

The compensatory regulation between the different CESR modules observed at the bulk-level (see above), raises the question of how the bacterial population implements this response at the individual cell level. Do all cells within the population behave uniformly, or is there significant
phenotypic heterogeneity within the population? Given that the excess expression of resistance
determinants is often associated with a fitness cost (Andersson and Hughes, 2010), it is in fact
intriguing to ask whether bacteria evolved to minimize ‘noise’ in resistance gene expression
(adjusting resistance as close as possible to its optimal level), or whether they actively use
heterogeneous gene expression as a means to diversify resistance levels within the population –
a strategy that can be beneficial in fluctuating environments (Fraser and Kaern, 2009).

To scrutinize the expression behavior of the three CESR modules at the single cell level,
we fused their promoters to a plasmid-borne copy of gfp and introduced them into wild type B.
subtilis W168. We then challenged exponentially growing cells with various levels of bacitracin
and quantified GFP fluorescence by flow cytometry one hour after bacitracin addition (Fig. 5). In
the absence of bacitracin the fluorescence distributions of the P_bceA-gfp (Fig. 5A) and the P_liaI-gfp
(Fig. 5C) reporters were identical to the autofluorescence distribution of B. subtilis W168 (data
not shown), while the P_bcrC-gfp reporter activity was ~5-fold higher than background (Fig. 5B),
consistent with the high basal activity of the P_bcrC promoter quantified with the luciferase reporter
above (cf. Fig. 3). This suggests that the gfp reporter is less sensitive than the luciferase reporter,
such that promoter activities below ~10^5 RLU/OD in Fig. 3 are hidden by the
autofluorescence of B. subtilis. However, apart from this difference in reporter sensitivity, the
mean fluorescence values for all promoter-gfp fusions were consistent with the results obtained
for the promoter-lux fusions in Fig. 3.

Next, we compared gene expression noise in the response of the three resistance
modules. As mentioned before, in the absence of bacitracin the fluorescence distributions of
P_bceA-gfp (Fig. 5A) and P_liaI-gfp (Fig. 5C) reporters were identical to the broad autofluorescence
distribution of B. subtilis. In contrast, P_bcrC-gfp reporter displayed a narrow fluorescence
distribution, and also showed low noise levels at all bacitracin levels tested. In the presence of
bacitracin the response of the P_bceA-gfp reporter became almost as homogeneous as the P_bcrC-
gfp reporter. Only the P_liaI-gfp reporter was expressed broadly heterogeneously across the
population when challenged with intermediate concentrations (1-3 µg ml^{-1}) of bacitracin (Fig.
5C), as reported before (Kesel et al., 2013). Indeed, when quantifying gene expression noise by
the coefficient of variation η, we found that at similar mean GFP expression levels the P_liaI
promoter was significantly noisier than the other promoters (Fig. S3A). This broadly
heterogeneous production of LiaIH argues for significant cell-to-cell variability in the downstream
damage perceived by the Lia system in the presence of bacitracin. In contrast, the low noise
levels in the expression of bceAB and bcrC suggest that their expression is subject to a more
stringent control, which might be result of negative feedback regulation within these systems
(see Discussion).
To test whether the noisy Lia response is influenced by the expression of the other two resistance modules, we introduced the $P_{\text{lia}}$-$gfp$ reporter plasmid into $\Delta bceAB$ and $\Delta bcrC$ mutants and determined their single cell response towards bacitracin as above. Strikingly, the Lia response displayed notably less cell-to-cell variability in the $\Delta bceAB$ mutant than in the wild type (Fig. 5D). Also, when comparing their coefficients of variation at similar mean expression levels (Fig. S3B), we found that $P_{\text{lia}}$ is less noisy in the $\Delta bceAB$ mutant than in the wild type, thereby showing that the reduced noise level is not only caused by the stronger and more sensitive $P_{\text{lia}}$ response in this mutant. This suggests that in the unperturbed (wild type) CESR network, the broadly heterogeneous Lia response is directly triggered by heterogeneity in $bceAB$ expression:

At the time of antibiotic treatment there exists a narrow, yet stochastic distribution of BceAB protein levels across the population, such that cells with higher levels of BceAB have sufficient ability to cope with bacitracin, whereas cells with lower levels of BceAB experience more cell envelope damage, which in turn triggers higher LiaIH production levels. Consequently, in the absence of BceAB this model predicts that all cells in the population would experience a similar envelope stress level, consistent with the homogeneous Lia response in the $\Delta bceAB$ mutant.

In contrast, our data showed that in a $\Delta bcrC$ mutant noise in the Lia response was markedly increased (Fig. 5D). We suggest that the increased noise in the expression of $bceAB$ in this mutant (Fig. S4) leads to a significant heterogeneity in the downstream damage perceived by the Lia system. However, we cannot exclude that population heterogeneity in other lipid II cycle-associated players factors into the noise properties of $P_{\text{lia}}$ in this highly impaired mutant strain. For instance, stochastic expression of $uppP$, encoding the second, BacA-like UPP phosphatase in $B. subtilis$ (Cao and Helmann, 2002; Bernard et al., 2005; Inaoka and Ochi, 2012), could result in largely variable rates of cell wall biogenesis, which would in turn lead to phenotypic heterogeneity in the susceptibility towards cell wall antibiotics.

**Discussion**

After the discovery of the bacitracin stimulon in $Bacillus subtilis$ (Mascher et al., 2003) and the quantitative characterization of its individual modules (Rietkötter et al., 2008), we here present the first description of the full anatomy of the bacitracin resistance network in $B. subtilis$. Using a systems-level approach we showed that a clear hierarchy exists between resistance modules, allowing us to discriminate between primary (BceAB) and secondary layers (BcrC and LiaIH) of bacitracin resistance. Strikingly, in mutants devoid of the primary resistance layer, the secondary layer was more strongly induced, revealing a high level of redundancy between resistance modules. Accordingly, our data now show for the first time that in the absence of the primary bacitracin resistance module, the deletion of $liaIH$ displays a clear-cut phenotype with a 6-fold
reduction of bacitracin resistance. Hence, we argue that the high level of resistance conferred by
BceAB masks the weaker contribution from the Lia system. This explains previous reports that
noted surprisingly weak phenotypes of a liaIH deletion alone, despite the strong Lia expression
under a variety of cell envelope-perturbing conditions, including lipid II cycle-interfering
antibiotics as well as oxidative stress reagents (Jordan et al., 2006; Rietkötter et al., 2008;
Suntharalingam et al., 2009; Wolf et al., 2010). So far, one of the strongest phenotypes was
found during treatment with the membrane pore-forming lipopeptide daptomycin, where a liaIH
deletion caused a 3-fold reduction of resistance (Hachmann et al., 2009; Wecke et al., 2009).
Notably, B. subtilis features no primary resistance mechanism against daptomycin, again
highlighting that the contribution of the Lia system to antibiotic resistance is strongest if other
resistance layers are lacking. Based on these observations, and in conjunction with the wide
distribution of the PspA/IM30 protein family (of which LiaH is a member) across various bacterial
phyla and even in archaea and eukaryotes (Joly et al., 2010), we speculate that the Lia system
constitutes an ancient resistance module that provides a low level of resistance against a broad
range of cell envelope-perturbing agents. In contrast, the Bce-like resistance modules confer
high levels of protection against a rather narrow range of antimicrobial peptides (Gebhard, 2012)
and are almost exclusively found in Firmicutes bacteria (Joseph et al., 2002; Mascher, 2006;
Dintner et al., 2011), suggesting that these specialized resistance layers were acquired later
during evolution.

The results presented here also shed new light on the role of the UPP phosphatase BcrC
in lipid II cycle homeostasis under antimicrobial peptide attack. First, we showed that the
deletion of bcrC lead to a significant decrease in growth rate - even in the absence of antibiotic
treatment. It appears likely that the second BacA-like UPP phosphatase, UppP, partially
compensates for the loss of BcrC, but that its activity is insufficient to maintain adequate cell wall
synthesis under the rapid growth conditions in LB media. The precise extent to which the uppP
promoter is up-regulated in such a mutant, and how it responds to lipid II cycle-inhibiting
antimicrobial peptides, remains to be elucidated. Second, our data revealed that bcrC deletion
had pleiotropic effects on the expression of all resistance modules and, most notably, triggered
their up-regulation also in the absence of externally added bacitracin. In these highly perturbed
cells, the induction of the Lia system was consistent with its role as a general sensor of cell
envelope stress (Wolf et al., 2010). Likewise, the up-regulation of the σ^M- and σ^X-dependent
P_{bcrC} promoter was not unexpected, because these alternative σ factors were also shown to be
sensors for cell wall integrity (Inoue et al., 2013; Lee and Helmann, 2013). However, it was
surprising to find the P_{bceA} promoter affected in the bcrC mutant, because all previous reports
were consistent with a model in which the Bce system responds to the detoxification flux of the drug and not to downstream damage on cell physiology (Wolf et al., 2012; Fritz et al., 2015).

One possible explanation for the elevated $P_{bceA}$ activity might be that the lack of the phosphatase BcrC causes the accumulation of UPP in the membrane, and thereby provides a surplus of targets for bacitracin. In turn, increased levels of UPP-bacitracin complexes would increase the detoxification flux per BceAB transporter, which then serves as the signal for $P_{bceA}$ activation. While this model can explain the increased $P_{bceA}$ activity in the presence of bacitracin, it is less intuitive why there was also a ~10-fold activation in the absence of bacitracin (cf. Fig. 3A). One possibility is that the accumulation of UPP itself somehow triggers BceAB activity. Interestingly, Kingston and colleagues suggested that BceAB may recognize UPP directly and flip it to the inner face of the membrane, where it may be protected from bacitracin and dephosphorylated by a cytosolically acting UppP (Kingston et al., 2014). Although it is known that BceB directly binds free bacitracin in vitro with high affinity (Dintner et al., 2014), it is conceivable that the physiological substrate of the transporter is the UPP-bacitracin complex in the cell, as suggested previously (Fritz et al., 2015). In this case, the transporter may also be able to interact with both components of the complex separately, i.e., free bacitracin and free UPP, especially when increased amounts of these are present. The increased basal activity of $P_{bceA}$ in the bcrC mutant may then be due to accumulation of UPP. A third alternative explanation might be that one or more of the endogeneously produced antimicrobial peptides activate the Bce system under these conditions. For instance, we recently showed that the endogeneous production of the sporulation delay protein C (SdpC) and the sporulation killing factor A (SkfA) in early stationary phase up-regulate production of BceAB and the paralogous PsdAB transporter in B. subtilis more than 100-fold (Höfler et al., 2016).

Taken together, the work from us and earlier work support the following, multi-layered model of the bacitracin resistance network in B. subtilis: In the presence of bacitracin (Bac) the accumulation of UPP-bacitracin (UPP-Bac) complexes blocks the lipid II cycle of cell wall biosynthesis and, as a consequence, leads to cell envelope damage. UPP-Bac is recognized by the ABC transporter BceAB, which releases UPP from the inhibitory grip of bacitracin by a so far unknown transport mechanism and thereby shifts the binding equilibrium towards the free form of UPP. Expression of bceAB is controlled by a flux-sensing mechanism (Fig. 1C), which homeostatically adjusts the BceAB level such that the transport activity of individual ABC transporters does not exceed a critical threshold (Fritz et al., 2015). At the same time such homeostatic, negative feedback systems are known to reduce gene expression noise (Alon, 2007), fully consistent with the homogeneous response of the Bce system observed at the single cell level. Simultaneously to the action of BceAB, BcrC reduces the concentration of the
bacitracin-target UPP by dephosphorylation to UP, thereby further promoting progression of the lipid II cycle. Under bacitracin stress, transcription of bcrC is controlled by the alternative ECF σ factor σM, which is regulated by the membrane-bound anti-σ factors YhdK/L (Fig. 1C). Previous data showed that either the depletion of UP and/or the depletion of lipid II could be the cues for anti-σ factors YhdK/L (Inoue et al., 2013; Lee and Helmann, 2013; Meeske et al., 2015). This suggests that the end product of the reaction catalyzed by BcrC (UPP → UP) could negatively regulate the expression of bcrC, which would in turn close a negative feedback loop that asserts homeostatic UP level control in the cell. This model is also consistent with all our data, most importantly the elevated PbcrC activity in the bcrC mutant (which we expect to display low UP levels), as well as the low noise level of the PbcrC promoter, which is again characteristic of negative feedback systems. Within our model, the Lia system constitutes the last line of defense that directly responds to and combats cell envelope damage, thereby explaining why the expression of the Lia system did not affect the expression of the other resistance modules in our data.

More generally, we propose that the redundant organization of the bacitracin resistance network of *B. subtilis* described here is a universal principle of many stress response networks within the microbial world, as demonstrated for instance in the oxidative stress responses of *Salmonella enterica* (Hébrard et al., 2009) and *Ralstonia solanacearum* (Flores-Cruz and Allen, 2009) or in the regulation of drug efflux systems in various bacterial species (Grkovic et al., 2002). Here the induction of individual stress response modules typically relieves stress perceived by other modules, which can be interpreted as a coupling between stress response modules via a global negative feedback mechanism. Failure of one of the `nodes` in such a network then triggers compensatory up-regulation of other nodes, which then jointly protect the cell. Interestingly, in the engineering disciplines this concept is known as `active redundancy`, during which the performance of individual devices is automatically monitored and dynamically reconfigured to eliminate performance declines of the system (Pahl and Beitz, 1996). In contrast, `passive redundancy` uses excess capacity to reduce the impact of component failures (Pahl and Beitz, 1996), which would be akin to the constitutive expression of all resistance determinants. In biological stress response networks, we propose that the use of active redundancy serves as an optimal regulation strategy to maximize cellular protection while preventing the direct or indirect costs of excess resistance gene expression.

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**Materials and Methods**

**Bacterial strains and growth conditions**

*Bacillus subtilis* and *Escherichia coli* were routinely grown in Luria-Bertani (LB) medium at 37°C with agitation (200 rpm). Transformations of *B. subtilis* were carried out as described previously (Harwood and Cutting, 1990). All strains used in this study are derivatives of the wild-type strain W168 and are listed in Table S1. Kanamycin (10 mg ml⁻¹), chloramphenicol (5 mg ml⁻¹), spectinomycin (100 mg ml⁻¹), tetracycline (10 mg ml⁻¹) and erythromycin (1 mg ml⁻¹) plus lincomycin (25 mg ml⁻¹) for macrolide-lincosamide-streptogramin B ('MLS') resistance were used for the selection of the *B. subtilis* mutants used in this study. Solid media contained 1.5% (w/v) agar. For complementation studies, full induction of the promoter P<sub>xylA</sub> was achieved by adding xylose to a final concentration of 0.2 % (w/v).

**DNA manipulation**

Plasmids were generated by using standard cloning techniques (Sambrook, Russell 2001) with enzymes and buffers from New England Biolabs (NEB; Ipswich, MA, USA) according to the respective protocols. PCR-DNA amplification for cloning purposes occurred with Phusion<sup>®</sup> or Q5<sup>®</sup> polymerase. Primers used in this study are listed in Table S2 and plasmid descriptions as well as details on their construction are given in Table S3. All plasmids were verified by sequencing of the insert. The integration of plasmids or DNA fragments into the genome, or the presence of a replicative vector, was confirmed by colony PCR. Integration into the thrC-locus was checked by threonine-auxotrophy in minimal medium.

**Determination of minimal inhibitory concentration**

Bacitracin resistance of *B. subtilis* strains was determined using Etest<sup>®</sup> strips on bacterial lawn (bioMérieux, Marcy l’Etoile, France), providing a concentration range from 256 to 0.016 µg ml⁻¹ bacitracin. Briefly, 3 ml of Müller-Hinton (MH) medium (2.1% (w/v) Müller-Hinton broth) were inoculated 1:100 from fresh overnight culture and cells were grown at 37°C with agitation to OD<sub>600</sub> = 0.6-0.8. Subsequently, 30 µl of the cell suspension were added to 3 ml molten MH soft agar (60°C, 0.75% (w/v) agar), mixed and distributed on MH agar plates. After 20 min of solidification, one Etest<sup>®</sup> strip was applied per agar plate. Results were documented after 24 h of incubation at 37°C.

**Luciferase assays**
Luciferase activities of *B. subtilis* strains harboring pBS3Clux-derivates were assayed using a Synergy™ NEOALPHAB multi-mode microplate reader from BioTek® (Winooski, VT, USA). The reader was controlled using the software Gen5™ (version 2.06). Cells were inoculated 1:1000 from fresh overnight cultures and grown to OD$_{600}$ = 0.1-0.5. Subsequently, cultures were diluted to OD$_{600}$ = 0.01 and split into 100 µl per well in 96-well plates (black walls, clear bottom; Greiner Bio-One, Frickenhausen, Germany). Cultures were incubated at 37°C with linear agitation (intensity, 567 cpm) and the optical density at 600 nm (OD$_{600}$) as well as luminescence was monitored every 5 min. After one hour, freshly diluted Zn$^{2+}$-bacitracin was added to the indicated final concentrations and incubation and monitoring was resumed for 2 hours. Specific luminescence activity is given by the raw luminescence output (relative luminescence units, RLU) normalized by cell density (RLU/OD).

*Flow cytometry assays*

Single-cell fluorescence of *B. subtilis* strains carrying GFP-reporter plasmids was measured using a BD Accuri™ C6 flow cytometer (BD Biosciences, Becton, Dickinson and Company, New Jersey, USA). Cells were inoculated 1:1000 from overnight cultures and grown at 37°C to OD$_{600}$~0.1. Subsequently the culture was split into test tubes, stained with FM® 4-64 (Life Technologies GmbH, USA) to a final concentration of 2 ng ml$^{-1}$ and incubated at 37°C with agitation. After 30 min cells were induced with indicated final concentrations of Zn$^{2+}$-bacitracin and 1 hour after further incubation, culture samples were assayed by flow cytometry. It was controlled by the BD Accuri™ C6 software using the following settings: sample threshold = 11,000 on FSC-H, core size = 5µm, flow rate = 10 µl min$^{-1}$. Noise in the resulting fluorescence distributions (cf. Fig. S3) was quantified by the coefficient of variation η, defined as the ratio of the standard deviation σ to the mean μ. In doing so, we used the geometric mean and variance, because those measures are known to yield more accurate statistics for log-normal distributed values than the arithmetic mean and standard deviation.
References


Kesel, S., Mader, A., Höfler, C., Mascher, T., and Leisner, M. (2013) Immediate and
heterogeneous response of the LiaFSR two-component system of *Bacillus subtilis* to the peptide antibiotic bacitracin. *PLoS ONE* 8: e53457.


Figure Legends

Figure 1. Schematic overview of bacitracin resistance determinants and their regulation in B. subtilis. (A) In the absence of bacitracin, the membrane-associated steps of cell wall biosynthesis in B. subtilis involve the cytosolic attachment of peptidoglycan precursors to the lipid carrier undecaprenyl-phosphate (UP) via MraY and MurG, followed by transport of the resulting lipid II molecule to the extracytoplasmic leaflet of the cytoplasmic membrane via at least two redundant flippases MraY and Amj. After incorporating peptidoglycan precursors into the cell wall by penicillin binding proteins (PBP), the remaining phosphorylated form of the lipid carrier, undecaprenyl-pyrophosphate (UPP), is converted to UP via the phosphatase BcrC, before it can enter the next transport cycle. Bacitracin blocks this essential lipid II cycle by tightly binding to UPP and thereby preventing the recycling of the lipid carrier. Bacitracin resistance is conferred by the increased production of the ABC-transporter BceAB, which removes bacitracin from UPP by a so far unknown transport mechanism, and the increased production of BcrC, which allows the lipid II cycle to progress in the presence of bacitracin. (B) The third player in the bacitracin stress response network is the phage-shock protein-like Lia response. Upon bacitracin challenge, the small membrane anchor LiaH recruits the cytosolic PspA/IM30 protein family member LiaH into membrane-associated patches of unknown physiological function. Potentially, these structures stabilize the membrane underneath damaged areas of the cell wall. (C) Regulation scheme of the bacitracin stress response network in B. subtilis. Expression of bceAB is activated via a flux-sensing mechanism, monitoring the detoxification flux of the ABC transporter BceAB via complex formation between BceAB and the histidine kinase BceS (Dintner et al., 2014; Fritz et al., 2015), which in turn activates transcription via phosphorylation of the response regulator BceR. Expression of bcrC is regulated by the ECF σ-factors σH and σX and their cognate anti σ-factors, which together are considered to be sensors for cell wall integrity (Inoue et al., 2013; Lee and Helmann, 2013). Likewise, expression of liaIH is regulated by the LiaFSR three-component system (Jordan et al., 2006; Mascher, 2006; Schrecke et al., 2013), which has also been shown to be a sensor of cell envelope damage (Wolf et al., 2012).

Figure 2. Contributions of CESR modules to bacitracin resistance. (A) Minimal inhibitory concentration (MIC) of indicated B. subtilis strains as determined by the E-test® agar gradient diffusion method on Müller-Hinton medium. Strains tested were W168, TMB35 (∆bceAB), TMB297 (∆bcrC), TMB1151 (∆liaIH), TMB713 (∆bceAB ∆bcrC), TMB2127 (∆bceAB ∆liaIH), TMB2128 (∆bcrC ∆liaIH) and TMB1829 (∆bceAB ∆bcrC ∆liaIH). Pictures are representative for three biological replicates with a maximal sample deviation of one concentration step; arrows
indicate the fold-change of sensitivity. (B) Doubling times of exponentially growing cells one hour after treatment with indicated bacitracin concentration. Graphs show data for single mutant strains containing the *lux*-reporter, see caption of figure 3. Standard deviation was obtained from at least nine biological replicates.

**Figure 3.** Dose-dependent activation of resistance modules in perturbed and unperturbed CESR networks. Target promoter activities of (A) *P*<sub>bceA</sub>-*lux*, (B) *P*<sub>bcrC</sub>-*lux* and (C) *P*<sub>liaI</sub>-*lux* in strains carrying indicated deletions of CESR modules, as given by specific luciferase activity (RLU/OD<sub>600</sub>) one hour after addition of indicated amounts of bacitracin. Measurements were performed during exponential growth phase in LB medium at 37°C in a microtiter plate reader. Data are shown for strains TMB1619, TMB1620, TMB1617 (W168); TMB1623, TMB1624, TMB1621 (Δ*bceAB*); TMB1627, TMB1628, TMB1625 (Δ*bcrC*) and TMB1661, TMB1662, TMB1659 (Δ*liaIH*) containing *P*<sub>bceA</sub>-*lux*, *P*<sub>bcrC</sub>-*lux* or *P*<sub>liaI</sub>-*lux*, respectively, see Table S1. Data points and error bars indicate means and standard deviations derived from at least three biological replicates.

**Figure 4.** Regulatory crosstalk between primary and secondary resistance modules. Target promoter activities of *P*<sub>bceA</sub>-*lux* and *P*<sub>liaI</sub>-*lux* in strains expressing different levels of (A) BceAB, (B) BcrC and (C) LialH, as given by specific luciferase activity (RLU/OD<sub>600</sub>) one hour after addition of indicated amounts of bacitracin. Measurements were performed as described in Figure 3. Colors code for different expression levels of resistance module *X* (*X* = *bceAB*, *bcrC* or *liaIH*), as driven by the xylose-inducible promoter *P*<sub>xylA</sub>: (red) No expression, via deletion of module *X*; (orange) Low constitutive expression, via complementation of the deletion mutant with *P*<sub>xylA</sub>-*X* in the absence of xylose; (light green) High constitutive expression, via complementation of the deletion mutant with *P*<sub>xylA</sub>-*X* in the presence of 0.2% xylose; (dark green) Overexpression in W168 wild type background, via expression of *P*<sub>xylA</sub>-*X* in the presence of 0.2% xylose. The corresponding strains are (A) TMB1619, TMB1623, TMB2590, TMB2594 (*P*<sub>bceA</sub>-*lux*) and TMB1617, TMB1621, TMB2589, TMB2593 (*P*<sub>liaI</sub>-*lux*) (B) TMB1619, TMB1627, TMB2592, TMB2430 (*P*<sub>bceA</sub>-*lux*) and TMB1617, TMB1625, TMB2591, TMB2429 (*P*<sub>liaI</sub>-*lux*) (C) TMB1619, TMB1661, TMB2693, TMB2691 (*P*<sub>bceA</sub>-*lux*) and TMB1617, TMB1659, TMB2692, TMB2690 (*P*<sub>liaI</sub>-*lux*), as listed in Table S1. Error bars indicate the standard deviation between at least three biological replicates.

**Figure 5.** Noise in the response of bacitracin resistance modules. Single cell bacitracin response of wild type strains carrying (A) *P*<sub>bceA</sub>-*gfp*, (B) *P*<sub>bcrC</sub>-*gfp*, and (C) *P*<sub>liaIH</sub>-*gfp* reporter
plasmids (strains TMB2174, TMB2173 and TMB1176, see Table S1), as well as in (D) ΔbceAB and (E) ΔbcrC mutant backgrounds carrying a P_{lux}^{fg}fp reporter plasmid (strains TMB2056 and TMB2057, see Table S1). Fluorescence distributions were quantified using flow cytometry, one hour after treatment of exponentially growing cells (37°C, LB medium) with bacitracin. Fluorescence distributions (colored) were obtained under bacitracin treatment indicated on the right, while transparent overlays (gray) are reference distributions obtained in the absence of bacitracin treatment. In every case one representative dataset of at least two independent biological replicates is shown.
<table>
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<th>genotype</th>
<th>W168</th>
<th>ΔliaIH</th>
<th>ΔbcrC</th>
<th>ΔbceAB</th>
<th>ΔbceAB ΔliaIH</th>
<th>ΔbceAB ΔbcrC</th>
<th>ΔbcrC ΔliaIH</th>
<th>ΔbceAB ΔbcrC ΔliaIH</th>
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<td>0.125</td>
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<tr>
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<td>10^-1</td>
<td>1</td>
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</tr>
<tr>
<td>Bacitracin [μg/ml]</td>
<td>6x</td>
<td>24x</td>
<td>85x</td>
<td>5x</td>
<td>512x</td>
<td>2048x</td>
<td>5450x</td>
<td>2.7x</td>
</tr>
</tbody>
</table>
A

\[ P_{\text{bceA}}^{\text{-lux}} \]

\[
\begin{array}{c}
\text{bacitracin [μg/ml]} \\
0 & 10^{-2} & 10^{-1} & 10^0 & 10^1 \\
\text{luciferase activity [RLU/OD]} \\
10^3 & 10^4 & 10^5 & 10^6 & 10^7 \\
\end{array}
\]

W168, ΔbceAB, ΔbcrC, ΔliaIH

B

\[ P_{\text{bcrC}}^{\text{-lux}} \]

\[
\begin{array}{c}
\text{bacitracin [μg/ml]} \\
0 & 10^{-2} & 10^{-1} & 10^0 & 10^1 \\
\text{luciferase activity [RLU/OD]} \\
10^3 & 10^4 & 10^5 & 10^6 & 10^7 \\
\end{array}
\]

W168, ΔbceAB, ΔbcrC, ΔliaIH

C

\[ P_{\text{liaI}}^{\text{-lux}} \]

\[
\begin{array}{c}
\text{bacitracin [μg/ml]} \\
0 & 10^{-2} & 10^{-1} & 10^0 & 10^1 \\
\text{luciferase activity [RLU/OD]} \\
10^3 & 10^4 & 10^5 & 10^6 & 10^7 \\
\end{array}
\]

W168, ΔbceAB, ΔbcrC, ΔliaIH
Variable BceAB levels

(i) $P_{bceA}$-lux

(ii) $P_{lia}$-lux

bacitracin [µg/ml]

Variable BcrC levels

(i) $P_{bceA}$-lux

(ii) $P_{lia}$-lux

bacitracin [µg/ml]

Variable LiaIH levels

(i) $P_{bceA}$-lux

(ii) $P_{lia}$-lux

bacitracin [µg/ml]
Figure A: 10

Figure B: 10

Figure C: 10

Figure D: 1.2

Gene expression noise $\eta$

mean fluorescence [RFU]

gene expression noise $\eta$

mean fluorescence [RFU]