The extracellular domain of site-2-metalloprotease RseP is important for sensitivity to bacteriocin EntK1

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Enterocin K1 (EntK1), a bacteriocin that is highly potent against vancomycin-resistant enterococci, depends on binding to an intramembrane protease of the site-2 protease family, RseP, for its antimicrobial activity. RseP is highly conserved in both EntK1-sensitive and EntK1-insensitive bacteria, and the molecular mechanisms underlying the interaction between RseP and EntK1 and bacteriocin sensitivity are unknown. Here, we describe a mutational study of RseP from EntK1-sensitive Enterococcus faecium to identify regions of RseP involved in bacteriocin binding and activity. Mutational effects were assessed by studying EntK1 sensitivity and binding with strains of naturally EntK1-insensitive Lactiplantibacillus plantarum—expressing various RseP variants. We determined that site-directed mutations in conserved sequence motifs related to catalysis and substrate binding, and even deletion of two such motifs known to be involved in substrate binding, did not abolish bacteriocin sensitivity, with one exception. A mutation of a highly conserved asparagine, Asn359, in the extended so-called LDG motif abolished both binding of and killing by EntK1. By constructing various hybrids of the RseP proteins from sensitive E. faecium and insensitive L. plantarum, we showed that the extracellular PDZ domain is the key determinant of EntK1 sensitivity. Taken together, these data may provide valuable insight for guided construction of novel bacteriocins and may contribute to establishing RseP as an antibacterial target.

Site-2-metalloproteases (S2Ps) are a family of intramembrane-cleaving proteases involved in regulated intramembrane proteolysis (RIP) (1, 2). In the RIP cascade, an S2P cleaves its substrate, for example, a membrane-bound anti-sigma factor, within the cell membrane, thereby mediating transmembrane signaling to trigger an adaptive response. S2Ps are conserved in all kingdoms of life and are crucial in several biological processes, including stress response, sporulation, cell polarity, virulence, and nutrient uptake (3–9). Due to its vital role in both animal and human pathogens, RseP is regarded as an attractive antimicrobial target. In fact, nature itself targets RseP, which is a known target for antimicrobial peptides belonging to the LsbB family of bacteriocins in selected Gram-positive bacteria (10, 11). Little is known about how these bacteriocins recognize and bind RseP and how this interaction eventually leads to killing of target cells. More insight into these issues is crucial for understanding bacteriocin function and for understanding how RseP may be targeted in antimicrobial therapy.

The hallmarks of the S2P family are the conserved catalytic motifs (HExXH and LDG) located on transmembrane segments (TMSs) of the protease (12). The S2P family of proteases is divided into four subgroups based on membrane topology and domain structure (13). Among the four groups, only a few members have been characterized; these include Escherichia coli RseP (EcRseP) from group I and the group III members MjS2P and SpoVFB from Methanocaldococcus jannaschii and Bacillus subtilis, respectively (12, 14, 15). EcRseP is the most extensively studied S2P and was first identified as a key modulator of stress response (16, 17). When E. coli cells are exposed to stress, a site-1-protease cleaves the membrane-bound anti-σE factor RseA. This primary cleavage triggers a secondary cleavage by the S2P EcRseP, which leads to release of RseA into the cytosol (16–18). RseA is further processed in the cytosol to form the mature σE, which activates genes involved in the stress response (19). It is believed that most S2P signaling pathways follow this same general cascade.

Next to the catalytic motifs, several conserved regions are thought to be involved in substrate interaction and catalysis by EcRseP. These include the membrane-reentrant β-hairpin–like loop (MRE β-loop), the GxG motif, and the PDZ domain (Fig. 1) (20–22). The PDZ domain has been suggested to work as a size-exclusion filter, preventing interaction with the substrate prior to site-1-protease cleavage (21, 23).

In addition, conserved residues near the LDG catalytic motif located in the third transmembrane segment (TMS3) have been implicated in substrate binding and recognition, in particular two asparagines and prolines in the sequence motif NxxxxNxxPxPxLDG (24), here referred to as the extended LDG motif. Despite the identification of these potentially important features, the mechanism of substrate recognition and binding by EcRseP remains somewhat enigmatic.
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Figure 1. Schematic representation of the topology of EfmRseP and alignment of conserved S2P regions. A, schematic representation of the predicted topology of Enterococcus faecium RseP with conserved S2P motifs indicated. TMS1-4 indicates the four predicted transmembrane segments. The GxG motif, MRE β-loop, and the predicted PDZ domain are indicated. The box below shows the predicted active site, consisting of the conserved HEXXH and LDG motifs. B, alignment of the amino acid sequences of active site and the extended LDG motif in RseP from four EntK1-sensitive species (E. faecium, Enterococcus faecalis, Lactococcus lactis, and Staphylococcus haemolyticus) and three EntK1-insensitive species (Lactiplantibacillus plantarum, Staphylococcus arlettae, and Methanocaldococcus jannaschii). Arrow heads indicate residues subjected to alanine substitutions. EntK1; Enterocin K1; MRE β-loop, membrane-reentrant β-hairpin-like loop; S2P, site-2-metalloprotease; TMS, transmembrane segment.

Bacteriocins are antimicrobial peptides produced by bacteria to inhibit other bacteria in competition for nutrition and ecological niches. They are considered promising alternatives and/or complements to antibiotics, mainly due to their potent activity against multidrug resistant pathogens. We have previously demonstrated that enterocin K1 (EntK1), a leaderless bacteriocin belonging to the LsbB family, is especially potent against Enterococcus faecium, including vancomycin-resistant strains (11, 25). Leaderless bacteriocins are synthesized without an N-terminal leader sequence and do not have post-translational modifications, making this group of bacteriocins ideal for synthetic production. Members of the LsbB family are small (30–44 amino acids), cationic, and amphiphilic, with an N-terminal helical structure and a disordered C-terminal end (11, 26). Interestingly, members of the LsbB family of bacteriocins differ in their inhibition spectrum, with LsbB being active only against Lactococcus lactis, while the inhibitory spectrum of EntK1 and enterocin EJ97 (EntEJ97) is broader, including high activity toward E. faecium and Enterococcus faecalis, respectively (11). It has previously been shown that the antimicrobial activity of bacteriocins of the LsbB family depends on RseP being present in target cells (10, 11).

RseP of E. faecium (EfmRseP) and EcRseP, both from subgroup 1, shares a 28% sequence identity and has the same predicted membrane topology and conserved domains (Fig. 1). Little is known about the function of RseP in E. faecium; however, recent phenotypic analysis of rseP mutants suggests a role in stress response (25). For E. faecalis, it has been shown that RseP (EfsRseP) is a key regulator of the stress response through RIP-mediated activation of the sigma factor SigV. Deletion of either EfsrseP or sigV increases the susceptibility of E. faecalis to multiple stressors, such as lysozyme, heat, ethanol, and acid (27). In addition, EfsRseP is involved in sex pheromone maturation and is therefore also referred to as Eep (enhanced expression of pheromone) in this organism (28). Lastly, deletion of EfsrseP has been shown to result in severely attenuated virulence in a rabbit endocarditis model and a catheter-associated urinary tract infection model, suggesting an important role for EfsRseP in pathogenesis (7, 29).

Despite the evident role of RseP in enterococcal virulence, critical features of enterococcal RseP, such as the substrate recognition mechanism, remain unknown. The known substrates of RseP-like S2P share no apparent sequence homology; however, amphiphilic helices in the substrates have been indicated as necessary for recognition (20, 30). Considering the helical structure of EntK1, it is conceivable that EntK1 interacts with enterococcal RseP in a similar manner as the native substrates. Therefore, to gain more insight into bacteriocin action and possibly the interaction between RseP and its natural substrates, we have studied the EntK1–RseP interaction, focusing on the role of conserved regions of RseP. The impact of mutations in these regions was assessed by bacteriocin-binding assays and by analyzing bacteriocin sensitivity of strains carrying mutated RseP. The results shed
light on the interaction between EntK1 and RseP, providing insights into bacteriocin specificity and giving valuable information for the design of novel bacteriocins.

Results

Heterologous expression of RseP renders insensitive Lactiplantibacillus plantarum sensitive to EntK1

*L. plantarum* WCFS1 is a Gram-positive bacterium for which pSIP-based vectors have been developed for heterologous protein expression (31, 32). In addition, the bacterium is insensitive to EntK1 despite having an *rseP* ortholog on the chromosome. Together, these properties make *L. plantarum* a suitable host for expressing *E. faecium* RseP for binding and sensitivity studies. As shown in Table 1, expression of RseP from *E. faecium* renders *L. plantarum* sensitive to EntK1, with a minimum inhibitory concentration (MIC<sub>50</sub>) of 0.01 μM, while *L. plantarum* carrying the empty vector (pEV) exhibited a MIC<sub>50</sub> greater than 22 μM, which is considered fully resistant. We also overexpressed *L. plantarum* RseP (LpRseP) in *L. plantarum*, to confirm the inability of LpRseP to be a receptor for EntK1. As expected, the LpRseP-overexpressing strain (LpRseP<sup>His</sup>, see Table 2 for a description of strain names) remained insensitive (i.e., MIC<sub>50</sub> greater than 22 μM) (Table 1). These results suggest that the *L. plantarum* strain is a suitable host for heterologous expression of EfmRseP. Moreover, a pairwise sequence alignment of EfmRseP and LpRseP indicates that subtle sequence differences between EfmRseP and LpRseP define the sensitivity toward EntK1 (Fig. S1).

Of note, while the pSIP vectors used for expression (Table 2) have an inducible promoter, regulated by the inducer peptide SppIP (31), all sensitivity and binding experiments were performed using noninducing conditions. Under inducing conditions (3–30 ng/ml SppIP), the transformants showed aberrant growth on agar plates (data not shown), indicating a cytotoxic effect likely due to the high amounts of the membrane-protein RseP. Noninduced cells appeared to grow normally. The inducible promoter sppA in the pSIP vector has a low basal activity in *L. plantarum*, which permits low expression of rseP genes under noninducing conditions (as demonstrated by the results presented in Table 1).

Antimicrobial activity and binding of EntK1 to sensitive cells depend on RseP

To examine whether the antimicrobial activity observed above is directly linked to the ability of EntK1 to bind target cells, we developed a binding assay for EntK1 to *L. plantarum*. For this assay, EntK1 was chemically synthesized with an N-terminal FITC fluorescent tag. The N-terminal fusion was chosen as the C-terminal half of the LsbB family of bacteriocins and is thought to be necessary for receptor interaction (26). The labeling of EntK1 with FITC reduced the antimicrobial potency, which, however, remained high for *L. plantarum*—expressing *E. faecalis* RseP (Table 1). Fluorescence microscopy of EntK1-sensitive *L. plantarum*—expressing plasmid-encoded *Efm*RseP showed strong fluorescent signals following exposure to FITC-EntK1, consistent with EntK1 binding. In contrast, nonsensitive *L. plantarum* carrying the empty vector (pEV) did not show any visible fluorescent signals under the same conditions, thus confirming lack of EntK1 binding (Fig. 2).

In addition with the fluorescence microscopy, flow cytometry analysis revealed that FITC-EntK1–exposed *L. plantarum*—expressing RseP derived from *E. faecalis* exhibited strong fluorescent signals, while cells containing the empty vector or overexpressing the *Lp*RseP protein showed no signal (Fig. 3). We have previously observed that EntK1 has some antimicrobial activity toward strains of *L. lactis*, *E. faecalis*, and *Staphylococcus haemolyticus* but not strains of *Staphylococcus aureus* and *Staphylococcus arlettae* (33). To confirm that the sensitivity is linked to RseP binding, rseP genes derived from these sensitive and insensitive species were heterologously expressed in *L. plantarum*. Table 1 shows that, indeed, *L. plantarum* strains expressing rseP genes derived from the sensitive strains of *L. lactis* (LIRseP<sup>His</sup>), *E. faecalis* (EfsRseP<sup>His</sup>), and *S. haemolyticus* (ShRseP<sup>His</sup>) were indeed inhibited by EntK1. In addition, Figure 3 shows that these strains had distinctly higher FITC signals than *L. plantarum* strains expressing RseP from the insensitive strains *S. aureus* (SasRseP<sup>His</sup>) and *S. arlettae* (SaRseP<sup>His</sup>). Taken together, these results provide strong evidence that there is a specific interaction between EntK1 and RseP from bacteria that are naturally sensitive to EntK1 but not between EntK1 and RseP from bacteria that are insensitive to EntK1.

Defining the role of conserved S2P motifs in the EntK1:RseP interaction

To define the regions of RseP involved in EntK1 sensitivity, we initially focused on conserved regions that, based on previous studies of other members of the S2P family, seem to be involved in substrate binding and catalysis. In addition to the

### Table 1

<table>
<thead>
<tr>
<th>Strains</th>
<th>Characteristics</th>
<th>MIC&lt;sub&gt;50&lt;/sub&gt; (μM)</th>
<th>FITC-EntK1</th>
</tr>
</thead>
<tbody>
<tr>
<td>EfsRseP&lt;sup&gt;His&lt;/sup&gt;</td>
<td>Expressing RseP from <em>Enterococcus faecalis</em></td>
<td>0.01</td>
<td>0.15</td>
</tr>
<tr>
<td>LpRseP&lt;sup&gt;His&lt;/sup&gt;</td>
<td>Expressing RseP from <em>Lactiplantibacillus plantarum</em></td>
<td>&gt;22</td>
<td>&gt;20</td>
</tr>
<tr>
<td>EfsRseP&lt;sup&gt;His&lt;/sup&gt;</td>
<td>Expressing RseP from <em>Enterococcus faecalis</em></td>
<td>0.04</td>
<td>0.6</td>
</tr>
<tr>
<td>LIRseP&lt;sup&gt;His&lt;/sup&gt;</td>
<td>Expressing RseP from <em>Lactococcus lactis</em></td>
<td>0.09</td>
<td>&gt;20</td>
</tr>
<tr>
<td>ShRseP&lt;sup&gt;His&lt;/sup&gt;</td>
<td>Expressing RseP from <em>Staphylococcus haemolyticus</em></td>
<td>0.17</td>
<td>&gt;20</td>
</tr>
<tr>
<td>SaeRseP&lt;sup&gt;His&lt;/sup&gt;</td>
<td>Expressing RseP from <em>Staphylococcus arlettae</em></td>
<td>&gt;22</td>
<td>&gt;20</td>
</tr>
<tr>
<td>SasRseP&lt;sup&gt;His&lt;/sup&gt;</td>
<td>Expressing RseP from <em>Staphylococcus aureus</em></td>
<td>&gt;22</td>
<td>&gt;20</td>
</tr>
<tr>
<td>pEV</td>
<td>Empty vector</td>
<td>&gt;22</td>
<td>&gt;20</td>
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* Control experiments (Fig. S2) indicated low expression, which may contribute to low sensitivity.
Antimicrobial activity of EntK1 depends on RseP

### Table 2

<table>
<thead>
<tr>
<th>Strain or plasmid</th>
<th>Relevant characteristic(s)</th>
<th>Reference</th>
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<td>Plasmid</td>
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<td></td>
</tr>
<tr>
<td>pLp1261_InvS</td>
<td>Spp-based expression vector, pSIP401 backbone, Ery&lt;sup&gt;e&lt;/sup&gt;</td>
<td>(31, 41)</td>
</tr>
<tr>
<td>Strain</td>
<td></td>
<td></td>
</tr>
<tr>
<td>L. plantarum WCFS1(L)</td>
<td>Template for rseP (LpRseP) and expression host</td>
<td>(53)</td>
</tr>
<tr>
<td>E. faecium P21 (Efm)</td>
<td>Template for rseP (EfmRseP)</td>
<td>(54)</td>
</tr>
<tr>
<td>E. faecalis V583 (Efm)</td>
<td>Template for rseP (EfmRseP)</td>
<td>NCBI:txid226185</td>
</tr>
<tr>
<td>L. lactis IL1403 (L)</td>
<td>Template for rseP (LbRseP)</td>
<td>NCBI:txid272623</td>
</tr>
<tr>
<td>S. aureus ATCC 14458 (Sax)</td>
<td>Template for rseP (SassRseP)</td>
<td>Nofima</td>
</tr>
<tr>
<td>S. arlettae LMGT 4134 (Sae)</td>
<td>Template for rseP (SaeRseP)</td>
<td>LMGT</td>
</tr>
<tr>
<td>S. haemolyticus LMGT 4106 (Shz)</td>
<td>Template for rseP (ShRseP)</td>
<td>LMGT</td>
</tr>
<tr>
<td>E. coli TOP10</td>
<td>Cloning host, Thermo Fisher Scientific</td>
<td></td>
</tr>
</tbody>
</table>

RseP homologs from the respective species are abbreviated with the species initials italicized (e.g., ShRseP is the RseP homolog in *S. haemolyticus*), while the strain names for each *L. plantarum* strain expressing a variant of RseP is not italicized (e.g., *ShRseP* in *L. plantarum* WCFS1 harboring pSIP401 encoding ShRseP). For cases where the species initials are ambiguous, both the first and last letter of the specific name is used (e.g., *E. faecium* and *E. faecalis*). Abbreviations: Em<sup>e</sup>, erythromycin resistance; LMGT, laboratory of microbial gene technology; Nofima, norwegian institute of food, fisheries and aquaculture research.

### The active site

The conserved motifs HExxH and LDG make up the active site of the S2P family (Fig. 1) (14). It has previously been shown that mutations of residues corresponding to EfmRseP His18, Glu19, His22, and Asp372 substantially affect the protease activity of RseP homologs from multiple species (12, 14, 34). To examine whether proteolytic activity of RseP is needed for EntK1 sensitivity, alanine substitutions were introduced in all conserved residues in the active site. Single alanine substitutions in the active site (EfmH18A, EfmE19A, EfmH22A, and EfmD372A) resulted in a slight increase of the MIC<sub>50</sub> from ≤0.002 μM for 6His-tagged WT EfmRseP to 0.01 to 0.7 μM for the 6His-tagged mutants (Table 3). In line with these observations, measurements of the populations with the single alanine substitutions in the binding assay described above showed only a slight reduction in binding with 59 to 80% of the median fluorescence intensity of the *L. plantarum* population expressing the native EfmRseP. The triple alanine substitution (EfmAAxxA) resulted in a considerable increase in the MIC<sub>50</sub> to 2.7 μM (Table 3). However, the triple mutant was still more than 8-fold more sensitive to EntK1 than pEV. The impact of the mutations on the MIC<sub>50</sub> values could be partly due to variation in RseP expression, which was not assessed in detail. For example, it is conceivable that the triple mutant is rather unstable and was produced in lower amounts, leading to a higher MIC<sub>50</sub> value and low EntK1 binding. Nevertheless, the fact that all variants remained sensitive and bound the
bacteriocin clearly shows that the mutant proteins were produced and that the catalytic activity of RseP does not play an essential role in RseP binding and strain sensitivity.

The MRE β-loop and the GxG motif region

Previous studies on EcRseP indicate that the MRE β-loop and the GxG motif region (Fig. 1) interact directly with the substrate (20, 22). To examine the significance of this region for the RseP:EntK1 interaction, residues 39 to 138 encompassing the MRE β-loop and the GxG motif were deleted (Trunc, Fig. S1). The truncation significantly reduced EntK1 sensitivity, as judged by the increase in MIC50 of Trunc to 2.7 μM (Table 3). Using the binding assay, we observed that FITC signals were also significantly reduced to 7.4% compared to the full-length protein (Table 3). Nonetheless, the FITC signal reflecting binding (7.4% versus 0.4%) and the sensitivity towards EntK1 (MIC50 of 2.7 μM versus 22 μM) were higher than that of the empty vector control strain (Table 3). It would thus seem that the MRE β-loop and the GxG motif region are not involved in the RseP:EntK1 interaction.

The extended LDG motif

A conserved motif in TMS3 (NxxPxPxLDG), which includes the LDG catalytic site motif (13), has been suggested as a prime candidate for S2P substrate binding (24). Moreover, previous substrate-binding studies with EcRseP (24) suggest a longer version of the LDG motif, referred as the extended LDG motif (N359xxxxN364xxP367xP369xLD372G in EfmRseP), may be important for substrate binding. The two asparagines and two prolines in the extended motif were individually mutated to alanine. Three of the four mutants remained sensitive to EntK1 and showed strong EntK1 binding (Table 3). However, the alanine substitution of Asn359 in EfmRseP (named EfmN359A) resulted in complete resistance to EntK1, and the binding of the bacteriocin was abolished (Fig. 3 and Table 3).

The absence of EntK1 sensitivity and EntK1 binding could be caused by failure to express the rseP variant. Therefore,
Antimicrobial activity of EntK1 depends on RseP

Table 3

<table>
<thead>
<tr>
<th>Strain*</th>
<th>EntK1 MIC50 (μM)</th>
<th>FITC-EntK1 rMFI % RSD</th>
</tr>
</thead>
<tbody>
<tr>
<td>EfmRseP</td>
<td>≤0.002</td>
<td>100 (8.3)</td>
</tr>
<tr>
<td>pEV</td>
<td>&gt;22</td>
<td>0.4 (4.2)</td>
</tr>
<tr>
<td>LpRseP</td>
<td>&gt;22</td>
<td>0.41 (9.7)</td>
</tr>
<tr>
<td>Active site</td>
<td></td>
<td></td>
</tr>
<tr>
<td>EfmH18A</td>
<td>0.02</td>
<td>61.6 (3.8)</td>
</tr>
<tr>
<td>EfmE19A</td>
<td>0.01</td>
<td>65.4 (3.3)</td>
</tr>
<tr>
<td>EfmH22A</td>
<td>0.01</td>
<td>59 (8.1)</td>
</tr>
<tr>
<td>EfmAAAX</td>
<td>2.7</td>
<td>0.5 (5.2)</td>
</tr>
<tr>
<td>EfmD372A</td>
<td>0.7</td>
<td>80.6 (3.5)</td>
</tr>
</tbody>
</table>

The middle column shows MIC for EntK1 towards L. plantarum strains expressing various RseP variants (see text, Figs. 1 and S1 for details). The strains are named by the protein variant they express. The right column shows the binding of FITC-labeled EntK1 to indicated strains. The FITC signals, indicating binding of the bacteriocin, are presented as the relative median fluorescence intensity (% rMFI) compared to the MFI obtained for EfmRseP6His (100%) with percent relative standard deviations (RSD).

The S2P RseP is highly conserved in multiple species, yet the potency of EntK1 varies considerably between species (11, 33). We investigated the relative median fluorescence intensity (% rMFI) compared to the MFI obtained for EfmRseP6His (100%) with percent relative standard deviations (RSD). All RseP variants contain a C-terminal 6× His-tag.

To further identify regions determining EntK1 sensitivity, we constructed several hybrid proteins in which parts of the RseP from insensitive L. plantarum were replaced with the corresponding parts of RseP from sensitive E. faecium (Hyb1-11, Fig. S3). Previous studies had suggested that residues 328 to 428 in the C-terminal region of RseP from L. lactis (YvjB) determine the sensitivity of L. lactis to LsbB (35). As LsbB and EntK1 target the same receptor, belong to the same bacteriocin family, and have a similar structure (10, 11, 26), we hypothesized that the C-terminal region of RseP from E. faecium would confer EntK1 sensitivity. To test the hypothesis, varying parts of the C-terminal region of LpRseP were replaced with the corresponding region of E. faecium RseP (Fig. S3). Surprisingly, the resulting hybrid proteins, Hyb1 and Hyb2, did not confer sensitivity to EntK1 (MIC50 > 22 μM) nor did they show bacteriocin binding (Table 3). Control experiments with EntEJ97 (Fig. S2) showed that Hyb1 and Hyb2 were produced.

Next, Hyb3 and Hyb4 (inverts of Hyb1 and Hyb2), containing the N-terminal region of EfmlRseP and the C-terminal region of LpRseP were constructed (Fig. S3). Unlike Hyb1 and Hyb2, Hyb3 and Hyb4 conferred sensitivity to EntK1 with MIC50 values of 0.7 μM and ≤0.002 μM, respectively. Hyb3 and Hyb4 also showed binding of the bacteriocin (Table 3). The results obtained with Hyb1-4 show that the N-terminal region of E. faecium RseP (residues 1–324) is involved in EntK1 binding. Although quantitative comparison of MIC50 values is risky due to possible differences in expression, it is worth noting that Hyb4, containing the complete EfmlRseP PDZ domain, was the most sensitive of the four hybrids.

To further narrow down the RseP region needed for EntK1 sensitivity, three additional hybrid proteins containing a decreasing portion of EfmlRseP were constructed (Hyb5-7, Fig. S3). None of these hybrids, all lacking the PDZ domain from E. faecium, could confer sensitivity to or binding of EntK1 (Table 3), indicating that the PDZ domain is required for activity. The control experiments of Fig. S2 showed that Hyb5 and Hyb6 were produced, whereas Hyb7 likely has reduced expression. To confirm the importance of the PDZ region, we constructed four additional hybrid proteins in which different parts of the PDZ domain of LpRseP were replaced with the corresponding sequences of EfmlRseP (Fig. 3, Hyb8-11). Interestingly, only Hyb10 and Hyb11, which contained the entire PDZ domain from EfmlRseP were EntK1-sensitive, with MIC50 values of 0.002 μM and 0.09 μM, respectively (Table 3). Hyb8 and Hyb9, only containing parts of the EfmlRseP PDZ domain, were not sensitive to EntK1 with MIC50 > 22 μM (Table 3). A control experiment showed that both Hyb8 and Hyb9 were highly sensitive to EntEJ97, indicating that these hybrids are produced (Fig. S2).

Importantly, as noted above, all hybrids that did not confer sensitivity or binding to EntK1, except for Hyb7, were sensitive (i.e., inhibition zone > 10 mm for EntEJ97; Fig. S2). This indicates that Hyb1-6 and Hyb8-11 were properly expressed and folded. Moreover, all clones of L. plantarum—expressing recombinant RseP showed growth comparable to EfmlRseP, suggesting that expression of the hybrids had no obvious toxic effect on the host (data not shown).

Discussion

The S2P RseP is highly conserved in multiple species, yet the potency of EntK1 varies considerably between species (11, 33). To further develop EntK1 as a novel treatment option for bacterial infections, a detailed understanding of the determinants of

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*EfmN359A (and all other variants displaying a complete loss of sensitivity, discussed below) were exposed to EntEJ97, another bacteriocin from the LsbB family. EntEJ97 targets RseP but has a different antimicrobial spectrum compared to EntK1 (11), which implies that its interaction with RseP differs from EntK1. Fig. S2 shows that the control pEV clone displayed limited sensitivity towards EntEJ97, while EfmN359A was highly sensitive to the bacteriocin, showing that the alanine substitution did not drastically alter the protein structure nor the expression level and that the removal of the asparagine side chain alone is likely responsible for the alteration in EntK1 sensitivity and binding. Interestingly, Asn359 and the extended LDG motif are highly conserved among both EntK1-sensitive and EntK1-insensitive species (Fig. 1B). Thus, the impact of this residue on EntK1 sensitivity must relate to its interaction with other less conserved regions of the protein.

Mapping the regions involved in EntK1 specificity.

To further identify regions determining EntK1 sensitivity, we constructed several hybrid proteins in which parts of the RseP from insensitive L. plantarum were replaced with the corresponding parts of RseP from sensitive E. faecium (Hyb1-11, Fig. S3). Previous studies had suggested that residues 328 to 428 in the C-terminal region of RseP from L. lactis (YvjB) determine the sensitivity of L. lactis to LsbB (35). As LsbB and EntK1 target the same receptor, belong to the same bacteriocin family, and have a similar structure (10, 11, 26), we hypothesized that the C-terminal region of RseP from E. faecium would confer EntK1 sensitivity. To test the hypothesis, varying parts of the C-terminal region of LpRseP were replaced with the corresponding region of E. faecium RseP (Fig. S3). Surprisingly, the resulting hybrid proteins, Hyb1 and Hyb2, did not confer sensitivity to EntK1 (MIC50 > 22 μM) nor did they show bacteriocin binding (Table 3). Control experiments with EntEJ97 (Fig. S2) showed that Hyb1 and Hyb2 were produced.

Next, Hyb3 and Hyb4 (inverts of Hyb1 and Hyb2), containing the N-terminal region of EfmlRseP and the C-terminal region of LpRseP were constructed (Fig. S3). Unlike Hyb1 and Hyb2, Hyb3 and Hyb4 conferred sensitivity to EntK1 with MIC50 values of 0.7 μM and ≤0.002 μM, respectively. Hyb3 and Hyb4 also showed binding of the bacteriocin (Table 3). The results obtained with Hyb1-4 show that the N-terminal region of E. faecium RseP (residues 1–324) is involved in EntK1 binding. Although quantitative comparison of MIC50 values is risky due to possible differences in expression, it is worth noting that Hyb4, containing the complete EfmlRseP PDZ domain, was the most sensitive of the four hybrids.

To further narrow down the RseP region needed for EntK1 sensitivity, three additional hybrid proteins containing a decreasing portion of EfmlRseP were constructed (Hyb5-7, Fig. S3). None of these hybrids, all lacking the PDZ domain from E. faecium, could confer sensitivity to or binding of EntK1 (Table 3), indicating that the PDZ domain is required for activity. The control experiments of Fig. S2 showed that Hyb5 and Hyb6 were produced, whereas Hyb7 likely has reduced expression. To confirm the importance of the PDZ region, we constructed four additional hybrid proteins in which different parts of the PDZ domain of LpRseP were replaced with the corresponding sequences of EfmlRseP (Fig. 3, Hyb8-11). Interestingly, only Hyb10 and Hyb11, which contained the entire PDZ domain from EfmlRseP were EntK1-sensitive, with MIC50 values of 0.002 μM and 0.09 μM, respectively (Table 3). Hyb8 and Hyb9, only containing parts of the EfmlRseP PDZ domain, were not sensitive to EntK1 with MIC50 > 22 μM (Table 3). A control experiment showed that both Hyb8 and Hyb9 were highly sensitive to EntEJ97, indicating that these hybrids are produced (Fig. S2).

Importantly, as noted above, all hybrids that did not confer sensitivity or binding to EntK1, except for Hyb7, were sensitive (i.e., inhibition zone > 10 mm for EntEJ97; Fig. S2). This indicates that Hyb1-6 and Hyb8-11 were properly expressed and folded. Moreover, all clones of L. plantarum—expressing recombinant RseP showed growth comparable to EfmlRseP, suggesting that expression of the hybrids had no obvious toxic effect on the host (data not shown).

Discussion

The S2P RseP is highly conserved in multiple species, yet the potency of EntK1 varies considerably between species (11, 33). To further develop EntK1 as a novel treatment option for bacterial infections, a detailed understanding of the determinants of
bacteriocin sensitivity and binding to RseP is essential. Therefore, in this study, we explored the contribution of conserved S2P motifs to the EntK1:RseP interaction and EntK1 sensitivity. To do so, we first needed to establish a sensitivity and binding assay. Although the antimicrobial activity of EntK1 depends on RseP (11), it remains elusive whether the difference in EntK1 sensitivity between species is solely due to variations in the RseP protein or if other factors, such as cell surface composition and gene expression levels, contribute. To avoid potential problems related to these uncertainties, we expressed rseP from insensitive and sensitive species in the same expression vector (pSIP) and EntK1-insensitive host (L. plantarum). We demonstrated that only rseP from sensitive bacterial species confers EntK1 sensitivity to L. plantarum. Binding of the bacteriocin to the RseP-producing L. plantarum strains was assessed using FITC-labeled EntK1. The levels of FITC-EntK1 signals correlated well with the MIC₅₀ values (Table 3; higher binding correlates with lower MIC₅₀ values). These observations show that subtle differences in the receptor alone likely determine variation in EntK1 sensitivity.

EfmRseP belongs to group I of the S2P family, for which the involvement of several conserved motifs in substrate binding and substrate specificity has been explored to some extent (20, 22, 24, 34). We considered that these motifs could be involved in EntK1 sensitivity and, therefore, we targeted these motifs in the mutagenesis studies to identify their role(s) in RseP as a bacteriocin receptor. We initially focused on the active site of E. faecium RseP, as there were indications in the literature that alterations in the active site of RseP in E. faecalis affects the sensitivity to a member of the LsbB bacteriocin family (11). However, none of the mutations in the catalytic center, including mutations known to abolish protease activity in EcRseP (16, 36), led to EntK1 resistance, demonstrating that the proteolytic activity of RseP is not essential for interaction and the antimicrobial action of EntK1.

Two motifs of EcRseP known to interact with the substrate are the MRE β-loop and a conserved GxG motif located on a membrane-associated region between TMS1 and TMS2 (Fig. 1) (20, 22). If RseP-targeting bacteriocins mimic the interaction of natural substrates with the receptor, these two regions would likely interact with EntK1. Although deletion of the MRE β-loop and the GxG motif led to a significant reduction in EntK1 sensitivity, the removal of these nearly 100 amino acids did not result in total resistance toward EntK1 (Table 3), indicating that neither the MRE β-loop nor the GxG motif is essential for the EntK1:RseP interaction. The reduced sensitivity and binding upon truncation are likely due to global structural changes in the receptor resulting from the large deletion. Of note, the MRE β-loop and GxG motif of EfmRseP are both predicted to be located on the cytoplasmic side of the cell membrane (Fig. 1A); such a location would likely not allow direct interaction with the bacteriocin which attacks target cells from the outside. It should be noted that the predicted topology of EfmRseP and the RseP hybrids was not confirmed experimentally in this study. However, a similar topology for the group 1 S2P EcRseP and SasRseP has been confirmed by the fusion of alkaline phosphatase to specific regions of RseP (6, 15), suggesting that the predicted topology may be conserved among group 1 S2Ps.

Next, we explored the role of the extended LDG motif in EntK1:RseP interaction. Substituting Asn364, Pro367, and Pro369 with alanine in EfmRseP resulted in only minor changes in EntK1 sensitivity and binding (Table 3). This was surprising, as these conserved asparagine and proline residues are known to be important for substrate binding and correct processing in both EcRseP and S2P from B. subtilis, known as SpoIVFB (12, 24, 37, 38). On the other hand, Asn359 was shown to be essential for EntK1 sensitivity and binding (Table 3). Under noninduced conditions, we were not able to detect RseP expression from N359A, or any other clone, using a standard Western blot (Fig. S4). However, when induced, expression levels of N359A and EfmRseP were comparable, suggesting that the insensitivity of the clone was due to the N359A substitution but not due to a failure in expression. Moreover, EfmN359A remained highly sensitive to EntE979, another bacteriocin of the LsbB family targeting RseP, which strongly indicates that the observed changes in sensitivity and binding were not caused by failure to express mutated rseP (Fig. S2). Previous studies have exploited the known substrates of RseP homologs to perform cleavage-based activity assays to confirm proper protein folding and expression following the introduction of mutations (24). However, RseP has no known substrates in E. faecium, which explains why cleavage-based activity assays could not be used. Interestingly, Asn389 in EcRseP, which corresponds to Asn359 in EfmRseP, plays an important role in substrate recognition. When this asparagine was replaced by cysteine, EcRseP showed reduced substrate binding as well as reduced proteolytic activity (24). Despite the evident role of Asn359 in EntK1:RseP binding, it is interesting to note that Asn359 and the surrounding extended LDG domain are highly conserved in the RseP proteins of both EntK1-sensitive and EntK1-insensitive species (Fig. 1B). This suggests that other regions of RseP play a role in bacteriocin binding and sensitivity.

Of the 11 constructed EfmRseP-LpRseP hybrid proteins, only four (Hyb3, Hyb4, Hyb10, Hyb11) were EntK1 sensitive (Table 3). Importantly, all EntK1-sensitive hybrids contain parts of the EfmPDZ domain, with Hyb4, Hyb10, and Hyb11 containing the entire domain. Of the four sensitive hybrids, hybrids containing the entire EfmPDZ domain exhibited the lowest MIC₅₀ (i.e., most sensitive), underpinning the important contribution of this domain to the EntK1:RseP interaction. Previous studies have shown that the PDZ domain is involved in substrate recognition by RseP-like S2P (21, 23). It has been suggested that the PDZ domain of EcRseP acts as a size-exclusion filter, preventing substrates with large periplasmic domains access to the active site (21). A similar role has been suggested for the PDZ domain of the B. subtilis S2P homolog, RasP (23). Several S2Ps process multiple substrates in vitro and in vivo, yet the substrate specificity of these proteins is poorly understood. We conclude that the PDZ domain of EfmRseP is the defining region for EntK1 binding and thus the major determinant of variation in EntK1 sensitivity.

To better understand the positions of EfmRseP motifs investigated in this study, we predicted the structure of EfmRseP using AlphaFold. AlphaFold is a protein structure prediction technique.
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program based on artificial intelligence that predict protein structures with greater accuracy than any other in silico method (39). As illustrated in Figure 4, AlphaFold predicted that the PDZ domain of EfimRseP forms a pocket which may prevent direct access to core residues. Among the regions investigated in this study, Asn359 in the extended LDG domain is located the closest to the PDZ domain, while the GxG motif and the MRE-β loop appears to be more distant (Fig. 4). Taken together with our experimental data, it is conceivable that the initial docking of EntK1 to the PDZ domain leads to subsequent interactions with core residues such as Asn359.

During the finalization of this article, AlphaFold-Multimer was published (R. Evans et al., Preprint at bioRxiv). AlphaFold-Multimer is an extension of AlphaFold2 using an artificial intelligence model explicitly trained for multimeric input. This allowed us to predict the EntK1-RseP complex, which strikingly predicted the interaction between EntK1 and RseP to primarily involve the PDZ domain and the region near Asn359 (data not shown). However, while most of the residues of RseP in the complex exhibited a high confidence score (pLDDT > 90), most of the residues of EntK1 were ranked poorly (pLDDT < 50). Confidence scores below 50 is a strong predictor of disorder, suggesting that the peptide chain is unstructured at physiological conditions or only structured as part of a complex. Indeed, EntK1 has been shown to be disordered in an aqueous environment by NMR spectroscopy (11). Due to the low confidence scores produced for EntK1 in the complex, these structure predictions are highly speculative and should be used cautiously.

While it remains unknown how the EntK1:RseP complex eventually leads to cell death, the present study reveals molecular details of the interaction of EntK1 with its receptor. Previous studies have shown that bacteriocins of the LsbB family can be engineered to improve both potency and alter the activity spectrum (33). The interpretation of these previous results, as well as future efforts to develop improved RseP-binding bacteriocins, will benefit from the deeper insight into the bacteriocin–receptor interaction that we provide here. Importantly, LsbB family of bacteriocins are attractive not only because they act on vancomycin-resistant strains but also because the bacteriocins are short, synthesized without an N-terminal leader sequence, and contain no posttranslational modification, which enables low-cost synthetic production. The fact that RseP homologs have important roles in virulence in several animal and human pathogens highlights RseP as an attractive antimicrobial target in multiple species (9, 40). The mutational analysis performed in this study combined with the predicted EfimRseP structure may provide a powerful basis for guided construction of novel bacteriocins and may contribute to further development of RseP as a drug target.

Experimental procedures

Bacterial strains and cultivation conditions

Bacterial strains used in this study are listed in Table 2. The following strains were cultivated in Brain heart infusion broth (Thermo Scientific Oxoid): enterococcal strains (37 °C, without agitation), staphylococcal strains (37 °C, 220 rpm), and E. coli (37 °C, 220 rpm). L. plantarum and L. lactis were cultivated without shaking in DeMan, Rogosa and Sharp (MRS) broth (Thermo Scientific Oxoid) at 37 °C and M17 broth (Thermo Scientific Oxoid) supplemented with 0.5% glucose at 30 °C, respectively. Agar plates were prepared by supplementing the appropriate broth with 1.5% (w/v) agar (VWR chemicals). Erythromycin was added to a final concentration of 200 μg/ml for E. coli and 10 μg/ml for L. plantarum when appropriate.

Construction of rseP orthologs, rseP hybrids, and site-directed mutagenesis

Seven orthologs of rseP from EntK1-sensitive and EntK1-insensitive species were expressed in L. plantarum using the pSIP expression system (31, 32) (Table 2). Briefly,
pLP1261_InvS, a pSIP derivative, was digested with NdeI and Acc65I or Xmal (Thermo Fisher Scientific) (41). Genomic DNA from the seven native rseP-containing strains was used as a template for the amplification of rseP. PCR amplification of all rseP variants was performed using Q5 Hot Start High-fidelity DNA polymerase (New England BioLabs) with InFusion primers to yield amplicons with ends complementary to the linearized pSIP vector (Table S1). The amplified PCR fragments were fused with the linearized vector using InFusion HD cloning Kit (Takara Bio) and transformed into competent E. coli TOP10 (ThermoFisher Scientific).

Site-directed mutants, truncations of RseP and RseP hybrids were constructed using splicing by overlap extension PCR. Briefly, two fragments of the rseP sequences were amplified in separate PCR reactions using two primer pairs, each consisting of an inner and outer primer (Table S1). The inner primers generated overlapping complementary ends and acted as mutagenic primers when introducing point-mutations. The overlapping fragments were fused by a second PCR reaction using the outer primers. Fused amplicons containing a mutated rseP gene were purified, fused to the linearized vector, and transformed to E. coli as described above.

All constructed plasmids were verified by DNA sequencing at Eurofins GATC Biotech (Germany) and subsequently transferred into electrocompetent L. plantarum as previously described (42). Fig. S3 shows a schematic representation of all hybrids and the truncated versions of RseP. Protein topology and the PDZ domain were predicted using CCTOP and Pfam, respectively (43, 44).

**Antimicrobial assays**

The bacteriocins used in this study, EntK1, EntEJ97, and FITC-EntK1, were produced by Pepmic Co, Ltd with >95% purity. Bacteriocins were solubilized in 0.1% (vol/vol) TFA (Sigma-Aldrich), except for FITC-EntK1 which was solubilized in dH2O. For semiquantitative assessment of antimicrobial activity, a spot-on-lawn assay was performed. Briefly, an overnight culture was diluted 1:100 in soft-agar and distributed on agar plates containing appropriate antibiotics. Bacteriocins with various concentrations were applied on designated spots on the solidified soft-agar. The agar plates were incubated at appropriate temperatures overnight and inhibition zones were measured the following day. For more accurate quantification, EntK1 sensitivity was determined using a microtiter plate assay to define MIC50 (45). The MIC50 was defined as the lowest bacteriocin concentration needed to inhibit bacterial growth by ≥50%. MIC assays were performed with three biological replicates.

**Binding assays**

Overnight cultures of L. plantarum strains were diluted 50-fold and grown until mid-log phase (4 h), after which cells were harvested by centrifugation at 16,000 g for 3 min and resuspended in sterile 0.9 % (w/v) NaCl to an A600 of 1 (assessed using a SPECTROstar Nano reader, BMG Labtech). Cell suspensions were diluted 20-fold in binding buffer [1 μM FITC-labeled EntK1 in 100 μM triammonium citrate pH 6.5 (Sigma-Aldrich)]. The cells were incubated in the binding buffer on a rotator (Multi Bio RS-24, Biosan, Latvia) at 6 rpm for 20 min at room temperature. After incubation, cells were harvested by centrifugation (16,000g, 3 min) and the binding buffer was discarded. The cell pellets were resuspended in sterile PBS to an appropriate cell density and analyzed using a MACSQuant Analyzer flow cytometer with excitation at 488 nm and emission at 475 to 575 nm (500 V PMT). The instrument was set to trigger on side-scattered light (SSC-A, 370 V PMT) with the threshold set to 8 to reduce false events.

Data and figures were prepared using the CytoExploreR package (v 1.1.0) for the R programming language (v 4.0.5) (https://github.com/DillonHammill/CytoExploreR [accessed June 25, 2022]), https://www.R-project.org/ [accessed June 28, 2022]). All binding assays were performed in triplicate. The median fluorescence intensity (MFI) was calculated as the average of three runs for each strain and expressed as a percent relative to L. plantarum expressing RseP from E. faecium (rMFI). Percent relative standard deviations were calculated as the ratio of the sample SD to the MFI mean multiplied by 100%.

**Phase contrast and fluorescence microscopy**

The cells were stained with the FITC-labeled EntK1 as described for the binding assay. After discarding the remaining binding buffer, cells were resuspended in 25 μl of PBS, spotted on a microscopy slide, and overlayed with 2% low melting agarose in PBS to immobilize the cells. Phase-contrast images and FITC fluorescence images were obtained using a Zeiss Axio Observer with ZEN Blue software and an ORCA-Flash 4.0 V2 Digital CMOS camera (Hamamatsu Photonics) using a 100 × phase-contrast objective. The excitation light source was an HXP 120 Illuminator (Zeiss).

**AlphaFold and structure analysis**

The structure of RseP and complexes between RseP and EntK1 were predicted using the published open source code for AlphaFold according to the instructions by the AlphaFold team (46). All required databases were downloaded on February 10th 2022 and all templates prior to 2022 were included (–max_template_date = 2022-01-01). Interactions present in the predicted complexes were determined by the fully automated protein-ligand interactions profiler (47) and the interactions function implemented in the web-based molecular viewer iCn3D (48, 49). Figures were generated using PyMOL (http://www.pymol.org/pymol). Amino acid sequences used for EfmRseP and EntK1 are presented in Table S2.

**Data availability**

The AlphaFold computations were performed on resources provided by Sigma2 (allocations NN1003K and NS1003K) - the National Infrastructure for High Performance Computing and Data Storage in Norway. For DNA sequence of the
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mutations and flow cytometry, the data will be shared upon request.

Supporting information—This article contains supporting information (50–52).

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Conflict of interest—The authors declare that they have no conflicts of interest with the contents of this article.

Abbreviations—The abbreviations used are: EcRseP, Escherichia coli RseP; EfRseP, Enterococcus faecium RseP; EfRseP, Enterococcus faecalis RseP; EntK1, Enterokinase K1; LpRseP, Lactiplantibacillus plantarum RseP; MFI, median fluorescence intensity; MIC50, minimal inhibitory concentration; MRE β-loop, membrane-reentrant β-hairpin–like loop; RIP, regulated intramembrane proteolysis; S2P, site-2-metalloprotease; TMS, transmembrane segment.

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