Rhomboid intramembrane protease YqgP licenses bacterial membrane protein quality control as adaptor of FtsH AAA protease

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Abstract

Magnesium homeostasis is essential for life and depends on magnesium transporters, whose activity and ion selectivity need to be tightly controlled. Rhomboid intramembrane proteases pervade the prokaryotic kingdom, but their functions are largely elusive. Using proteomics, we find that Bacillus subtilis rhomboid protease YqgP interacts with the membrane-bound ATP-dependent proces- sive metalloprotease FtsH and cleaves MgtE, the major high-affinity magnesium transporter in B. subtilis. MgtE cleavage by YqgP is potentiated in conditions of low magnesium and high affinity magnesium transporter in sive metalloprotease FtsH and cleaves MgtE, the major high-

Introduction

Rhomboids are intramembrane serine proteases widespread across the tree of life. In eukaryotes, rhomboid proteases regulate signal-

ing via the epidermal growth factor receptor, mitochondrial quality control or invasion of malaria parasites into the host cells (reviewed in Urban, 2016). Rhomboid proteases also pervade the prokaryotic kingdom (Koonin et al., 2003; Kinch & Grishin, 2013), suggesting sufficient biological significance for evolutionary conservation and expansion. However, their functions in bacteria are much less well-understood than in eukaryotes. Notably, in eukaryotes, the rhomboid superfamily includes a number of pseudoproteases of important biological roles, such as iRhoms and Derlins, but the mechanistic and evolutionary aspects of rhomboid pseudoprotease functions have not been clarified (Ticha et al., 2018).

In the Gram-negative eubacterium Providencia stuartii, the rhom-

boid protease AarA is required to process a pro-form of TatA, a component of the twin-arginine translocase secretion apparatus (Stevenson et al., 2007), but this function does not seem to be widely conserved because most bacteria encode a mature form of TatA. In the archaeabacterium Haloferax volcanii, a rhomboid protease is involved in protein glycosylation of the S-layer (Parente et al., 2014; Costa et al., 2018), but the molecular mechanism has not been uncovered yet. Similarly, the main model rhomboid protease GlpG of Escherichia coli, widely distributed in Gram-negative bacteria, is extremely well-characterised structurally and

mechanistically (Strisovsky et al., 2009; Baker & Urban, 2012; Zoll...
et al., 2014), yet its biological function is unknown. A recent report proposed a role of GlpG in the survival of a pathogenic strain of E. coli in the mouse gut (Russell et al., 2017), but the mechanism has not been demonstrated and it is not clear how specific the phenotype is, because rescue experiment has not been reported. Current knowledge of functions of rhomboid proteases in Gram-positives is similarly sketchy. An isolated report indicated that YqgP may play a role in cell division and may be required for glucose export (hence the therein coined alternative name GluP) (Mesak et al., 2004), but rescue experiment was not reported and the involved substrates were not investigated, leaving the molecular mechanisms of these functions unknown.

Therefore, to shed light on one of the significantly populated subclasses of bacterial rhomboid proteases in a model Gram-positive organism, we focused on Bacillus subtilis YqgP and used quantitative proteomics to identify its substrates and interactors. We find that YqgP cleaves the high-affinity magnesium transporter MgtE and that YqgP interacts with the membrane-anchored metalloprotease FtsH. At low extracellular concentration of magnesium cations and high concentration of manganese or zinc cations, cleavage of MgtE by YqgP is potentiated, and the globular N-terminal cytosolic domain of YqgP represents the manganese/zinc-sensing unit. The second molecular role of YqgP is presenting MgtE or its cleavage products as substrates to FtsH, for which the proteolytic activity of YqgP is dispensable but its unoccupied active site is essential. YqgP thus fulfills both protease and pseudoprotease functions in tandem with FtsH, representing an ancestral proteolytic platform dedicated to the regulated degradation of polytopic membrane proteins, functionally equivalent to regulatory ERAD in eukaryotes. Our results shed light on the evolution of membrane proteostasis control in response to environmental stimuli, and on the arising of pseudoproteases, which are surprisingly common in the rhomboid superfamily (Adrain & Freeman, 2012; Freeman, 2014).

Results

Quantitative proteomics reveals candidate interactors and substrates of Bacillus subtilis rhomboid protease YqgP

Bacillus subtilis genome encodes two rhomboid protease genes, ydcA and yqgP [also known as gluP (Mesak et al., 2004)]. No proteolytic activity has been detected for YdCA so far (Urban et al., 2002b; Lemberg et al., 2005), while YqgP is a commonly used model rhomboid protease cleaving a number of synthetic substrates (Lemberg et al., 2005; Urban & Wolfe, 2005; Ticha et al., 2017). YqgP has homologs in a number of Gram-positive bacteria, including Bacilli, but also Staphylococci, Listeriae and others (http://www.ebi.ac.uk/interpro/protein/P54493/similar-proteins), and thus represents an attractive system to explore the cell biology and functions of bacterial rhomboid proteases. To reveal the repertoire of YqgP substrates, we generated a strain deficient in yqgP and re-expressed yqgP or its catalytically dead mutant S288A (Lemberg et al., 2005) in this background from an ectopic chromosomal locus (strains BSS0 and BSS1, respectively, see Table EV2). We then conducted quantitative proteomic analysis of membrane fractions of both of these strains employing SILAC labelling and SDS–PAGE pre-fractionation (GeLC experiment), where we were seeking peptides belonging to proteins migrating at a lower-than-expected apparent molecular weight selectively in the YqgP-expressing strain relative to the S288A mutant expressing one. This analysis yielded the high-affinity magnesium transporter MgtE as the top candidate substrate of YqgP (Fig 1A–C, Dataset EV1, PRIDE dataset PXD014578).

In a complementary approach, we used affinity co-immunopurification and label-free quantitative proteomics to identify proteins associating with YqgP (Fig 1D and E). We ectopically expressed either a functional YqgP-sGFP fusion or catalytically dead YqgP S288A-sGFP fusion as the sole copy of YqgP in the cell. We solubilised the isolated membranes using the NP-40 detergent, isolated YqgP-sGFP by anti-GFP affinity pull-down and analysed the co-isolated proteins by MS-based proteomics. Using this approach, we identified several high-confidence interactor candidates including the membrane-anchored protease FtsH, and ATPase subunits A, D, F and G (Fig 1E, Dataset EV2, PRIDE dataset PXD014566). The only high-confidence overlap between the two proteomic datasets was the high-affinity magnesium transporter MgtE (Fig 1C and E), promoting it to the highest likelihood candidate substrate. MgtE is the main magnesium transporter in B. subtilis, it is essential for magnesium homeostasis (Wakeham et al., 2014), and we thus examined the possible functional relationship between YqgP and MgtE.

YqgP cleaves the high-affinity magnesium transporter MgtE between its first and second transmembrane helices

To validate the results of quantitative proteomics and gain more insight into the role of YqgP in the physiology of B. subtilis, we first examined the proteolytic status of endogenous MgtE by immunoblotting using an in-house generated antibody recognising its N-terminal cytosolic domain (MgtE 1–275). Both endogenous and ectopically expressed YqgP cleaved endogenous MgtE, yielding distinct cleavage products (Fig 2A). The cleavage was abrogated by a rhomboid-specific peptidyl ketoamide inhibitor (Ticha et al., 2017b) at low nanomolar levels, confirming that it was a rhomboid-specific event (Fig 2B). Judging by the apparent molecular size of the N-terminal cleavage fragment compared with the in vitro-translated reference fragments (Lemberg & Martoglio, 2003) of MgtE, we concluded that the cleavage by YqgP occurred within the extracytoplasmic loop between TMH1 and TMH2 of MgtE. This region is close to or within the periplasmic end of TMH2 (Fig 2C and D), which is consistent with the topology and mechanism of a rhomboid protease (Stri-ovsky, 2013, 2016). MgtE transporters function as homodimers with a cytosolic amino-terminal cystathionine-beta-synthase (CBS) domain that senses intracellular Mg2+ and a carboxy-terminal five-transmembrane helical pore for Mg2+ import (Hattori et al., 2007; Takeda et al., 2014). Cleavage of MgtE between TMH1 and TMH2 by YqgP would thus likely inactivate the transporter.

Manganese excess activates cleavage of MgtE by YqgP under conditions of magnesium starvation

It is known that MgtE homologs can also transport Mn2+ (Takeda et al., 2014) under low Mg2+ concentration (Mg starvation) and that high concentrations of Mn2+ are toxic for bacteria, mainly because Mn2+ can mis-metalate Mg2+-binding sites (Hohle & O’Brian, 2014; Chandrangsu et al., 2017). We therefore hypothesised that YqgP
could be involved in the regulation of MgtE under Mn\(^{2+}\) stress, and we tested the influence of manganese on the \textit{in vivo} activity of YqgP. While at high extracellular magnesium concentration (1 mM) the effect of 100 \(\mu\)M MnCl\(_2\) was not detectable, at low extracellular magnesium concentration (0.01 mM), when endogenous MgtE is upregulated, a shift from 1 to 100 \(\mu\)M MnCl\(_2\) activated the YqgP-dependent cleavage of MgtE fourfold (Fig 3A).

We next tested the importance of YqgP for growth in manganese-stress conditions. During growth in minimal medium at low magnesium concentration (10 \(\mu\)M), adding 75 \(\mu\)M manganese at mid-log phase caused growth arrest and lysis of wild-type \textit{B. subtilis}, which was more severe upon deletion of \textit{yqgP}, and fully rescued by ectopic expression of YqgP (Fig 3B). This effect was highly reproducible. Interestingly, all strains resumed growth
at later time points, indicating the possible existence of an adaptive mechanism. This did not appear to involve the second B. subtilis rhomboid YdcA, because deletion of ydcA did not modify the phenotypic behaviour of B. subtilis regardless of the presence or absence of yggP (Fig EV1). Regardless, the yggP-deficient strains showed a marked growth disadvantage even in the latter phase of the growth curve, indicating that the role of YggP for the fitness of B. subtilis in these conditions is significant. The toxic effect of 50 μM manganese sulphate was fully prevented by growth in high (5 mM) magnesium sulphate (Fig 3C), which was consistent with a similar effect on MgtE cleavage by YggP (Fig 3A). In addition, increasing concentrations of YggP inhibitor had a dose-dependent effect on the protective role of YggP both when it was overexpressed or present at endogenous levels.
(Fig 3D), indicating that the pronounced protective effect of ectopically expressed YqgP was a result of its overexpression. Finally, the phenotype manifested also during steady-state growth in rich LB medium supplemented with 75 μM manganese sulphate, where yqgP-deficient B. subtilis ectopically expressing MgtE was delayed in growth compared with the same strain ectopically expressing
YqgP (Fig 3E). Intriguingly, ectopic expression of a proteolytically inactive S288A mutant of YqgP also rescued the phenotype of the parent strain, which was initially surprising and suggested that YqgP may have a role independent of its protease activity. Together, these results reveal that a physiological function of YqgP is the protection from manganese stress by contributing to the degradation of the main magnesium transporter MgtE in *B. subtilis* (Wakeman et al., 2014).

The N-terminal extramembrane intracellular domain of YqgP mediates the manganese-induced activation of MgtE cleavage

To gain insights into the mechanism of MgtE cleavage by YqgP, we turned our attention to the two extramembrane domains of YqgP, the N-terminal cytosolic domain (NTD) and the C-terminal extracellular domain (CTD). While CTD consists of a predicted tandem array of three tetratricopeptide repeat (TPR) motifs (Zeytuni & Zarivach, 2012) and is thus likely to interact with protein or peptide ligands, NTD did not show significant sequence similarity to any protein of known structure, providing no indication of its molecular function. We first investigated the importance of NTD and CTD for YqgP activity. We created *B. subtilis* strains producing YqgP lacking NTD or CTD and measured the *in vivo* “specific activities” of these YqgP variants (i.e. substrate conversions at steady state normalised to enzyme expression level) in cleaving MgtE and the heterologous model substrate TatA (Ticha et al., 2017a). While deletion of CTD (∆A388–507, ∆CTD) seems to have led to about four- to sixfold increase in YqgP activity for both substrates, deletion of NTD (∆A1–178, ∆NTD) reduced YqgP activity for MgtE threefold, but only marginally so for the model TatA substrate (Fig 4A). In addition, deletion of NTD also compromised the activation effect of Mn²⁺ on MgtE cleavage by YqgP, while the deletion of CTD had no effect (Fig 4B–D). Taken together, the existence of such a specific regulatory mechanism involving YqgP NTD further substantiated the idea that the manganese-regulated cleavage of MgtE by YqgP is physiologically significant.

The N-terminal intracellular domain of YqgP binds divalent cations and mediates their sensing by YqgP

Since divalent cations often have related coordination properties, we wondered whether the manganese activation effect on YqgP transmitted via YqgP<sub>NTD</sub> may be a reflection of a more general divalent metal cation effect. We thus inspected a range of divalent metal cations for activation of MgtE cleavage by YqgP, while the deletion of CTD had no effect (Fig 4B–D). Taken together, the existence of such a specific regulatory mechanism involving YqgP NTD further substantiated the idea that the manganese-regulated cleavage of MgtE by YqgP is physiologically significant.

Ni²⁺ cannot be transported but inhibits MgtE transport (Smith et al., 1995). Here, we observed that yqgP deficiency made *B. subtilis* more sensitive to Mn²⁺ and Zn²⁺ but not to Co²⁺ or Ni²⁺ stress at low Mg²⁺, and consistently, re-expression of YqgP rescued this effect to above wild-type levels (Fig 5C). Collectively, this indicates that YqgP<sub>NTD</sub> may act as a sensor for a subset of metal cations in order to prevent cation toxicity due to MgtE permissiveness under low Mg²⁺ conditions.

To understand the basis of the activation of YqgP by divalent cations, we undertook structural analysis of YqgP<sub>NTD</sub>. Domains homologous to YqgP<sub>NTD</sub> occur only in YqgP orthologs among the soil-dwelling bacteria of the genus *Bacillus* (http://www.ebi.ac.uk/interpro/protein/P54493/similar-proteins), bear little sequence similarity to other proteins and show no hits at ProSite. Thus, to understand the structural basis of manganese-induced activation of YqgP, we first determined the 3D structure of NTD by solution NMR (Fig 6A, statistics in Table EV3). The overall structural ensemble of YqgP<sub>NTD</sub> (YqgP 1–177) consists of both α- and β-structural motifs, comprising six β-strand sequences at positions 21–24 (β1), 30–34 (β2), 43–48 (β3), 80–88 (β4), 104–106 (β5) and 109–117 (β6), and five α-helices at positions 3–16 (α1), 54–75 (α2), 96–100 (α3), 121–128 (α4) and 146–168 (α5). The central mixed β-sheet motif is packed in a twisted plane and surrounded by helices α1–4. Notably, negatively charged side chains are clustered into several solvent-exposed regions (Fig 6B). Structural similarity search using Dali server (Holm & Laakso, 2016) identified a number of structurally related proteins, with no conserved functional role but enriched in DNA/RNA polymerases and modifying enzymes (Table EV4).

In particular, the β-stranded central part of YqgP<sub>NTD</sub> adopts a fold characteristic for the type II restriction endonucleases (InterPro and Pfam codes IPR018573 and PF09491, respectively) (Kachalova et al., 2008).

Since Mn²⁺ but not Zn²⁺ provided reliable data in ITC titrations, we focused on Mn²⁺ in further structural analyses aiming to understand the structural aspect of divalent cation-mediated activation via YqgP<sub>NTD</sub>. NMR titrations of YqgP<sub>NTD</sub> by the paramagnetic Mn²⁺ induced peak intensity reductions and chemical shift perturbations in a number of residues along NTD. N-H chemical shifts could be caused by direct interaction with the metal ion, or by secondary structural changes, while peak intensity could be also diminished by the paramagnetic effect of Mn²⁺. The most strongly affected regions were amino acids 27–30, 35–37, 48–63 and 89–98 of the 177 amino acid NTD (Fig 6C and D, full titration data are available in Dataset EV3). To identify potential residues involved in binding Mn²⁺, we next focused on amino acids typically involved in chelating divalent cations (D, E, H) from each of the regions affected in NMR titrations. We analysed the effect of alanine substitutions in D29, D37, H49, D50, D52, D60 and E90 on the ability of these mutants to elicit Mn-induced activation of YqgP in *in vivo* (Fig 6E). This analysis indicated that the hotspot in which mutations interfered with Mn-induced activation of YqgP was centred around D50, and D50A mutation abrogated Mn activation completely. Interestingly, several of the residues affected in NMR titrations clustered in one apex of NTD structure, with D50 being central in this region (Fig 6F), representing the likely Mn-binding region in NTD (Fig 6G). In summary, deletion of YqgP<sub>NTD</sub> or mutation of chelating residues from the identified Mn²⁺-binding surface abrogated the Mn²⁺-induced activation of MgtE.
cleavage by YqgP in vivo showing that NTD is the Mn$^{2+}$-sensing device of YqgP.

YqgP works in tandem with the AAA protease FtsH to degrade MgtE in response to the environmental divalent cation concentrations

to explore more precisely MgtE proteolysis in vivo, we decided to construct a yqgP mutant strain complemented with a ectopic mutated version of yqgP that encodes a catalytically dead YqgP.S288A protein. Surprisingly, we observed that the production of this inactive YqgP.S288A in the yqgP-deficient background resulted in the production of an alternative band with lower mobility on SDS–PAGE, i.e. a higher molecular weight, than that of the cleavage product generated by YqgP (Fig 7A). Since the S288A variant of YqgP lacks the catalytic serine nucleophile and is thus proteolytically completely inactive (Lemberg et al., 2005; Wang et al., 2006; Baker & Urban, 2012), these results suggested that another proteolytic enzyme may participate in MgtE processing under these conditions. Co-immunoprecipitation experiments (Fig 1E) showed that YqgP interacts with the ATP-dependent, membrane-bound, processive zinc metalloprotease FtsH, and we thus hypothesised that FtsH is the responsible enzyme. Indeed, deletion of ftsH led to the disappearance of this alternative cleavage product (Fig 7A).
**Figure 5. The N-terminal intracellular globular domain of YqgP binds divalent cations.**

A Detection (left panel) and quantification (right panel) of steady-state conversion of endogenous MgtE by YqgP (strain BS72, Table EV2) grown in minimal M9 medium containing 10 μM MgSO4 and 1 μM each of MnCl2, ZnCl2, CoCl2, NiCl2 and CaCl2 (reference conditions) with the additions of 10 μM or 100 μM of a given divalent cation salt solution (MnCl2, ZnCl2, CoCl2 or NiCl2), or 50 μM and 500 μM for CaCl2. Western blots were quantified by near-infrared fluorescence detection (left), quantified by densitometry and displayed as relative specific activity (graph on the right), which is substrate conversion normalised to enzyme expression level. Black arrow denotes full-length substrate (MgtE), and red arrow denotes the N-terminal cleavage product(s) generated by YqgP. All bands originate from the same Western blot and identical treatment series. Irrelevant lanes have been cropped out for clarity, and the source blot is available online as Source Data.

B Isothermal titration calorimetry of purified recombinant YqgPNTD and selected divalent cations.

C Cation toxicity assays and their relationship to YqgP activity. Wild-type *Bacillus subtilis* (BTM2, Table EV2), its variant lacking YqgP (ΔyqgP, BTM78, Table EV2) and the rescue strain ectopically expressing YqgP (BTM501, Table EV2) were cultivated in minimal M9 medium containing 10 μM MgSO4 and 1 μM each of MnCl2, ZnCl2, CoCl2, NiCl2 and CaCl2 (i.e. reference conditions), and in mid-exponential phase were stressed by the addition of either 70 μM MnCl2, 500 μM ZnCl2, 25 μM CoSO4 or 400 μM NiCl2 (pale blue area). YqgP activity specifically improved cell fitness during manganese and zinc stress, while it had no effect on growth of cells cultivated in the presence of high cobalt and nickel concentrations. Representative experiments of 2–3 independent replicates are shown.

Source data are available online for this figure.
Figure 6. Solution NMR analysis reveals manganese-binding region in the N-terminal domain of YqgP.

A Solution NMR structure of cytosolic N-terminal domain of YqgP (YqgP\textsubscript{NTD}, amino acids 1–177), showing ensemble of 30 structures with lowest energy, and the structure with minimum energy displayed as cartoon.

B Electrostatic surface of YqgP\textsubscript{NTD} structure reveals highly negatively charged areas. Left view is oriented as the left view in panel (A).

C Chemical shift perturbation values and relative peak intensities for backbone resonances (\textsuperscript{15}N and \textsuperscript{1}H) of YqgP\textsubscript{NTD} calculated for complexes of 400 μM YqgP\textsubscript{NTD} with 10 μM MnCl\textsubscript{2} (light grey, light orange) or 40 μM MnCl\textsubscript{2} (dark grey, dark orange bars). Only residues 20–100 of YqgP are displayed for simplicity. Residues identified as potential manganese-binding region based on their chemical shifts and intensity changes upon NMR titrations are depicted in orange. The X sign marks residues whose chemical shifts were not calculated due to the lack of resonance signal.

D Details of 2D \textsuperscript{15}N/\textsuperscript{1}H HSQC spectra of free (green) and Mn-titrated YqgP\textsubscript{NTD} (10 μM MnCl\textsubscript{2} in red and 40 μM MnCl\textsubscript{2} in blue) for the residues marked in orange in panel (C). As a reference, residues marked in black do not display spectral changes upon titration, e.g. peak intensity does not change for S\textsubscript{88}N-H upon Mn titration (overlay of green, blue and red contours), but peak intensity is severely depleted for D29N-H upon Mn titration (absence of red contours). Full titration data are available in Dataset EV3.

E, F Detection (E) and quantification (F) of steady-state conversions of endogenous MgtE by YqgP\textsubscript{variant} bearing single-point mutations in putative Mn-binding region (BS\textsubscript{184}; 187; 196–203, Table EV2), cultivated in modified M9 minimal medium supplemented by low (1 μM) or high (100 μM) MnCl\textsubscript{2}. Black arrow marks full-length MgtE, and red arrow marks its N-terminal cleavage product by YqgP.

G Overview of residues affected by Mn\textsuperscript{2+} binding to YqgP\textsubscript{NTD} mapped onto its solution NMR structure. The view is oriented as the left view in panels (A and B).
bacterial rhomboid protease in the regulation of key membrane protein proteostasis under environmentally restricted conditions in a process similar to the eukaryotic ERAD.

Discussion

Metal ions are essential for life because they serve as cofactors in various cellular processes, but their excess poses a lethal threat. Hence, various mechanisms exist to ensure metal ion homeostasis (Chandraganu et al., 2017). Soil bacteria, such as B. subtilis, frequently encounter fluctuations in metal ion availability and need to possess sophisticated regulatory and feedback mechanisms to specifically respond to a limitation or excess of various metal ions (Chandraganu et al., 2017). Magnesium (Mg²⁺) represents the most abundant divalent cation in cells. It is maintained at high free intracellular levels (0.5–1 mM) relative to other cations (Froschauer et al., 2004), and it plays vital roles in membrane integrity, charge neutralisation of nucleic acids and nucleotides, ribosome structure and activities of many metabolic enzymes. To take up environmental Mg²⁺, bacteria usually encode several magnesium transporters of the CorA, MgtA and MgtE families (Groisman et al., 2013). Homologs of MgtE are universally conserved up to humans (SLC41 family), indicating their importance. In B. subtilis, MgtE is the main magnesium transporter (Wakeman et al., 2014) and is tightly regulated at several levels (Groisman et al., 2013). First, low levels of environmental magnesium trigger upregulation of mgtE transcription via a riboswitch mechanism (Dann et al., 2007). Second, the Mg-dependent gating mechanism by the N-terminal cytosolic domain of MgtE ensures metal ion selectivity (Hattori et al., 2009). Third, ATP further regulates MgtE (Tomita et al., 2017), presumably because the cell needs to maintain a balance in magnesium and ATP concentrations as the two are functionally coupled in a number of physiological processes (Pontes et al., 2015, 2016).

Structural principles of ion channel selectivity of MgtE have been studied in detail (Takeda et al., 2014; Maruyama et al., 2018). At low magnesium concentrations (when Mg²⁺ is not a competitive inhibitor), MgtE can also transport Co²⁺, Mn²⁺ and Zn²⁺ to some extent, while Ni²⁺ inhibits MgtE (Smith et al., 1995; Takeda et al., 2014). This implies that at low magnesium concentrations and high concentrations of Co²⁺, Mn²⁺, Zn²⁺ or Ni²⁺, other regulatory mechanisms may be required to maintain the biological function of Mg and/or minimise toxicity of other divalent cations. Indeed, previous observations in other organisms indicate that controlling magnesium transporter selectivity with respect to transition metal ions is a widespread theme. In yeast, excess magnesium prevents manganese toxicity (Blackwell et al., 1997), implying that MgtE may transport manganese when magnesium does not compete with it. In Bradyrhizobium japonicum, a null mutant in the manganese transporter mntH encoding gene does not grow in Mn-limiting medium, but suppressor strains can be selected. One class of suppressors are gain-of-function mutants of mgtE (mutations affecting gating mechanism) (Hohle & O’Brian, 2014). In this system, when the intracellular Mg²⁺ levels are low, the MgtE mutants transport Mn²⁺ with high affinity, which is not the case at high Mg²⁺ and for the wild-type strain. This is particularly relevant in soil, the natural habitat of B. subtilis, where there is a negative correlation between the bioavailability of magnesium and manganese ions. As soil pH
Hence, in the commonly found acidic soils, available manganese becomes more available and magnesium less available (Thomas & Sprenger, 2008; Gransee & Fuhrs, 2013). With stress conditions of low magnesium and high manganese, and the regulation of MgtE by YqgP that we describe is important for fitness in similar conditions (Figs 3 and 5).

Figure 7. The YqgP rhomboid functions as both a protease and a pseudoprotease adaptor of the AAA protease FtsH to maintain proteostasis of MgtE.

A The proteolytically dead mutant YqgP.S288A behaves as a pseudoprotease and acts as a substrate adaptor of FtsH. Pattern of C-terminally truncated forms of endogenous MgtE formed in vivo in the presence of ectopically expressed active YqgP (strain BS72, Table EV2) or its inactive S288A mutant (BS73, Table EV2) in rich LB medium in the presence or absence of endogenous FtsH. Black arrow denotes the full-length MgtE substrate, and red arrows denote its cleavage products generated by YqgP. While the presence of the faster migrating YqgP-dependent products (full red arrows) do not depend on the presence of FtsH, the slower migrating product formed in the presence of YqgP.S288A (open red arrow) does, indicating that FtsH protease is responsible for the alternative cleavage of MgtE. The same arrow symbolics are valid throughout this figure.

B Mapping of the boundary of the FtsH generated proteolytic product of MgtE. Immunoblot comparing the SDS-PAGE mobility of the N-terminal fragment of endogenous MgtE generated by FtsH in the presence of the pseudoprotease version of YqgP (catalytically dead YqgP.S288A, strains BTM78, BTM502, Table EV2) to the mobilities of in vitro-translated reference fragments corresponding to MgtE 1–340, 1–355 and 1–370 shows that YqgP.S288A acts as an FtsH adaptor and that FtsH processing of MgtE in the presence of YqgP.S288A stops at around amino acid 355, which is near or at the cytoplasmic end of TMH 3 of MgtE.

C Schematic depiction of the regions in MgtE defining its remaining cleavage products formed by FtsH and YqgP.

D In vivo cleavage of endogenous MgtE in the presence of ectopically expressed full-length YqgP.S288A (strain BS73), YqgPANTD.S288A (BS185, Table EV2) or YqgPΔCTD.S288A (BS186, Table EV2), in rich LB medium shows that YqgPANTD is necessary for the FtsH-dependent processing of MgtE.

E Analysis of the in vivo kinetics of formation and fate of MgtE cleavage products. Bacillus subtilis strains expressing YqgP or its S288A mutant (strains BTM78, 501 and 502, Table EV2) in the presence or absence of endogenous FtsH protease (strains BTM795, BTM796 and BTM797, Table EV2) were cultured in LB medium and at early exponential phase, after having been expressing YqgP variants for 30 min, were either left grown untreated (as a control with ongoing translation) or treated with 20 μg/ml tetracycline to stop proteosynthesis (translation shut-off). At given time intervals afterwards, all cultures were analysed by α-MgtE Western blotting with near-infrared detection. Equal cell number (judged by OD600) was loaded into each lane. Black arrow marks full-length MgtE, red full arrows mark YqgP-dependent and red open arrows mark YqgP.S288A-dependent cleavage product of MgtE. decreases, manganese becomes more available and magnesium less available (Thomas & Sprenger, 2008; Gransee & Fuhrs, 2013). Hence, in the commonly found acidic soils, B. subtilis has to cope with similar conditions (Figs 3 and 5).
The role of YqgP seems to be important during acute changes in cation concentrations. Several hours after manganese shock, even yqgP-deficient *B. subtilis* starts growing again, albeit at lower rate than wild type or a strain overexpressing YqgP (Fig 3B). This “adaptation” may be based on the described genetic mechanisms related to the regulation of Mn metabolism. It is known that *B. subtilis* responds to Mn\(^{2+}\) excess in two ways. First, the MntR regulator senses excess intracellular Mn\(^{2+}\). It then represses the transcription of two Mn\(^{2+}\)-uptake systems (MntABC and MntH) and activates the expression of efflux systems (MneP and MneS) (Que & Helmann, 2000; Huang et al., 2017). In addition, the two manganese efflux pumps, MneP and MneS, are also activated via Mn\(^{2+}\)-responsive riboswitches (Dambach et al., 2015; Price et al., 2015). These actions take place at the transcriptional level, hence take control to sufficiently allow the cell to recover from an initial Mn\(^{2+}\) shock. As intracellular Mn\(^{2+}\) returns to physiological levels, cells resume growth and YqgP may thus lose its importance for fitness in the later stages of adaptation to these toxic conditions.

We find that in *B. subtilis*, rhomboid protease YqgP is required to cleave and inactivate species of MgtE formed at low environmental Mg\(^{2+}\) and high Mn\(^{2+}\) or Zn\(^{2+}\), which protects *B. subtilis* from toxic Mn\(^{2+}\) or Zn\(^{2+}\) stress. The identity of the MgtE species susceptible to YqgP cleavage in these conditions is unknown. Since their generation requires ongoing translation (Fig 7), we hypothesise that these may be mis-metallated forms at some of the metal-binding sites throughout the MgtE molecule (Foster et al., 2014; Takeda et al., 2014; Maruyama et al., 2018) that can form during translation and folding, and which could interfere with the proper channel function of MgtE. YqgP is thus providing a quality control check for properly functioning MgtE. Cleavage of MgtE by YqgP is more pronounced at low Mg\(^{2+}\) and high Mn\(^{2+}\) or Zn\(^{2+}\) concentrations. This may, in part, result from the higher concentration of susceptible forms of MgtE in these conditions, which is consistent with our translation shut-off chase data (Fig 7). However, metal sensing and cleavage activation are also directly provided by the intracellular N-terminal extramembrane domain of YqgP. This domain binds divalent metal cations at millimolar or submillimolar affinities and approximately 1:1 stoichiometry (Fig 5) along a negatively charged surface of the molecule (Fig 6). How exactly in structural and mechanistic terms Mn\(^{2+}\) or Zn\(^{2+}\) binding to YqgP activates MgtE cleavage is not clear at the moment. It could be through direct interaction with the membrane-embedded MgtE, or by influencing the speed of diffusion of YqgP in the membrane, as recently suggested for several other rhomboid proteases (Kreutzberger et al., 2019).

Interestingly, in addition to the influence on YqgP activity, YqgPNTD is also required for the coupling of YqgP activity to FtsH, the ATP-dependent processive protease that associates with YqgP. More generally, there is emerging evidence from this work and the accompanying back-to-back study by Liu *et al.* that rhomboid proteases can participate in membrane protein quality control in bacteria, where they are very widespread. It remains to be elucidated which structural and dynamic aspects of transmembrane domains rhomboid proteases can recognise in their substrates, to serve as “detectors” of misfolded or orphan protein species and adaptors of downstream degradative proteases. The variability in substrate specificities and various extramembrane domains of rhomboids may be the basis of their diverse functions in bacteria.

The specialised role that YqgP plays in the quality control of an essential transporter protein may seem unexpected, but conceptually, transporters are indeed regulated at a number of levels and very finely using specialised mechanisms. For example, amino acid permeases of the major facilitator superfamily in yeast are downregulated from the cell surface under high nutrient conditions by an increase in ubiquitin E3 ligase activity of Rsp5, which initiates their trafficking from the plasma membrane to multi-vesicular bodies (Lauwers *et al.*, 2010). Other member of the same transporter family in yeast requires a specialised membrane chaperone Sh3, which assists the folding of the transporters and prevents their efficient degradation by ERAD (Kota *et al.*, 2007).

Intriguingly, in addition to the role of YqgP in regulating MgtE, we suspect it can be engaged in other functions. In co-immunoprecipitation experiments with YqgP, we identified a number of ATP synthase subunits with high confidence and abundance by mass spectrometry (Fig 1E). Since we have not identified any of these subunits in our substrate screen (Fig 1C), we did not follow them up. However, this finding raises the possibility that further to its role in MgtE proteostasis, YqgP may be involved in the relationship between magnesium homeostasis and ATP synthesis, which would be consistent with a growing body of evidence supporting a regulatory relationship between magnesium and ATP levels. First, it was reported that a magnesium transporter-defective strain of *B. subtilis* can be rescued by inactivation of ATP synthase-encoding genes (Wakeman *et al.*, 2014), suggesting that lowering ATP levels allows the cell to cope with magnesium deprivation. This is consistent with findings from Groisman and colleagues in *Salmonella enterica,*
where low levels of Mg$^{2+}$ in the cell lead to ATP synthase inhibition and reduced ribosome biogenesis. As a result, the pool of magnesium that is not ATP-bound anymore can be employed to support low levels of ribosome assembly, a priority process that requires large amounts of magnesium (Pontes et al., 2016). Finally, in the same bacterium, ATP synthase activity can be directly inhibited by the MgtC protein, whose expression is upregulated during magnesium deprivation (Lee et al., 2013). Strikingly, MgtC downregulation in the membrane of S. enterica is somewhat reminiscent of how MgtE is controlled by YggP and FtsH in B. subtilis. MgtR, a small hydrophobic peptide, is thought to destabilise MgtC prior to degradation by FtsH (Alix & Blanc-Potard, 2008). Hence, during magnesium deprivation and cation toxicity in B. subtilis, it is tempting to speculate that YggP may also inhibit the activity of ATP synthase to preserve the free Mg$^{2+}$ pool for translation purposes while assisting FtsH for the degradation of MgtE.

A number of native and model rhomboid substrates are single-pass membrane proteins (Urban et al., 2002a; Strisovsky et al., 2009; Riestra et al., 2015; Johnson et al., 2017; Saita et al., 2017). The cleavage of the transmembrane core of MgtE by YggP described here adds to growing evidence that rhomboid proteases do engage in cleaving polytopic membrane proteins under some conditions (Erez & Bibi, 2009), although the features of transmembrane domains that they recognise still need to be properly structurally understood. Our results show that degradation of the susceptible MgtE species requires both YggP and FtsH, with YggP cleavage of the transmembrane core of MgtE being required for full degradation by FtsH. In addition, YggP presents MgtE to FtsH, for which YggP activity is dispensable, but its unoccupied active site is essential, thus behaving partially also as a pseudoprotease. In summary, here we uncover an ancestral membrane-associated degradation mechanism including a substrate receptor/protease and processive degradative dislocase and protease. This is conceptually remarkably similar to the process of ERAD in which rhomboid pseudoproteases Derlins co-operate with ubiquitin E3 ligases such as Hrd1, the AAA ATPase p97/VCP and the proteasome, in dislocating transmembrane proteins from the membranes of endoplasmic reticulum and degrading them (Neal et al., 2018; Ticha et al., 2018). Our work emphasises that even the active rhomboid proteases can have pseudoprotease functions, and suggests that the balance between their protease and pseudoprotease roles may vary. In this context, it is interesting to note that the second rhomboid protease encoded in B. subtilis genome, YdcA, lacks any detectable protease activity despite having all sequence hallmarks of an active rhomboid protease (Urban et al., 2002b; Lemberg et al., 2005). Similarly, mammalian RHBDL1 and RHBDL3 failed to cleave any known model substrates of rhomboid proteases (Johnson et al., 2017), and it ought to be borne in mind that in these or other cases, the pseudoprotease roles may be key to the physiological functions of these rhomboid proteins. More generally, other intramembrane proteases may have physiological roles that do not depend on their catalytic activity, such as the role of SPP in membrane protein dislocation from the ER (Loureiro et al., 2006).

A number of bacteria encode YggP homologs (http://www.ebi.ac.uk/interpro/protein/P54493/similar-proteins), and the function of YggP in magnesium homeostasis control that we describe here might be conserved more widely, because MgtE homologs are also very widespread (https://www.ebi.ac.uk/interpro/protein/O34442/similar-proteins), although it is possible that YggP homologs will have also other functions. This is indeed suggested by two recent reports. First, it was shown that the Staphylococcus aureus YggP homolog Rbd is synthetic lethal with the nucleoid occlusion protein Noc and is involved in the initiation of DNA replication (Pang et al., 2017). Second, in a recent archived report, the same protein (although termed ActH therein) has been identified as a regulatory component of a complex with peptidoglycan amidase LytH and is required for LytH activation (Do et al., 2020). In neither case was it established whether the described function of S. aureus YggP homolog Rbd/ActH depends on its protease activity. It is intriguing to speculate whether these functions will rely on a similar protease or pseudoprotease roles within a degradation machinery such as the one we describe here. It is fully conceivable that YggP homologs can fill a range of functions in diverse bacteria depending on the client protein but using analogous general mechanism as we outline here.

In a generalised summary, this work implicates bacterial rhomboid proteases in the quality control of polytopic membrane proteins in cooperation with other processive proteases, which could be regarded as an evolutionary functional ancestor of the eukaryotic ER-associated degradation (ERAD). In our system, rhomboid plays not only the role of a protease that facilitates dislocation of the membrane protein by the ATP-dependent AAA protease FtsH, but also the role of a substrate adaptor handing over the substrates to FtsH. It is conceivable that with a more efficient and specialised dislocation machinery in higher organisms, the requirement for protease activity of the rhomboid-like protein (Derlin) could become less important, while the adaptor function becoming key. This logic would explain the particularly high occurrence of pseudoproteases in the rhomboid superfamily, which includes Derlins that are well-known essential components of the eukaryotic substrate recognition and dislocation machinery in ERAD (Neal et al., 2018).

Materials and Methods

Materials and chemicals

All common chemicals were from Sigma-Aldrich or VWR unless otherwise indicated. Oligonucleotides were from Sigma-Aldrich and Eurogentec; and restriction and DNA modification enzymes were from New England Biolabs.

Plasmids and DNA cloning

All DNA constructs were created by PCR and isothermal assembly (Gibson, 2011), and verified by Sanger sequencing. All constructs used in this work are summarised in Table EV1.

Bacterial strains, media and growth conditions

All B. subtilis strains used in this study were derived from the 168 trpC2+ strain (Nicolas et al., 2012; Koo et al., 2017), and construction details of the strains are listed in Table EV2. For routine work, B. subtilis strains were streaked onto solid LB agar plate (supplemented with appropriate antibiotic) from −80°C stock solution and

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grown overnight at 37°C. Subsequently, single colony was inoculated into the fresh LB medium and cultivated until the pre-culture reached OD₆₀₀ around 1.0. Culture was then diluted in the fresh growth medium (LB or M9) to starting OD₆₀₀ of 0.01–0.025 and cultivated until the exponential phase was reached at OD₆₀₀ ranging from 0.6 to 1.5. Depending on the experimental setup, protein expression from an ectopic site was induced either by adding 1% (w/v) D-xylene (for constructs under the control of xylene promoter, Pₓyl) or IPTG (Phyperpans) promoter) at the start of cultivation as indicated, at 37°C, until OD₆₀₀ reached 1, unless stated otherwise. For regular growth, both *B. subtilis* and *E. coli* cells were cultivated in LB medium (Merck) at 37°C. For *in vivo* activity assays, *B. subtilis* cells were cultivated in modified M9 minimal medium (low or high Mg²⁺ content) composed of M9 salt solution (6.8 g/l Na₂HPO₄₂H₂O, 3 g/l KH₂PO₄, pH 7.4, 0.5 g/l NaCl, 1 g/l NH₄Cl), 0.5% (w/v) D-glucose, 10 μM (low Mg²⁺) or 1 mM MgSO₄ (high Mg²⁺), 1 mM MnCl₂, 100 μM CaCl₂, 10 μM FeSO₄ and 18 amino acids including all genetically encoded ones except lysine and tyrosine (Harwood & Cutting, 1990). All minimal media in which toxicity assays were performed were prepared in water of ultrapure quality, processed through Milli-Q® Reference Water Purification System (Merck) and having resistivity of at least 18.2 MΩ cm (at 25°C) and TOC values below 5 ppb. For translational shut-off chase experiments, *B. subtilis* cultures were grown to mid-exponential phase, in LB medium at 30°C. At this point, expression of *yqgP* from the *Phyperpans* promoter was induced by adding 0.1 mM IPTG for 30 min. At a given time point, protein synthesis was blocked by the addition of tetracycline to a final concentration of 20 μg/ml. Cation toxicity assays were performed by adding concentrations of MnCl₂, ZnCl₂, CaSO₄ or NiCl₂ into M9 minimal medium at various growth time points, as specified in the result section. *Bacilli* cultures for SILAC-based proteomic experiments were cultured in modified M9 minimal medium (see above) supplemented with either naturally occurring “light” (¹⁵N₁⁴N₂) or “heavy” (¹³C₆¹⁵N₂) isotopic variants of l-lysine (Silantes GmbH, Germany).

Proteins uniformly labelled by ¹⁵N or ¹³C for NMR studies were overexpressed in *E. coli* BL21(DE3) (YggP NTD) or *E. coli* Lemo21(DE3) (YggP CTD). grown in M9 minimal medium containing partial M9 salt solution (6.8 g/l Na₂HPO₄₂H₂O, 3 g/l KH₂PO₄, pH 7.4, 0.5 g/l NaCl), 1 mM MgSO₄, 0.1 mM CaCl₂, 0.5% (w/v) D-glucose (U-¹⁵C, 99%, Cambridge Isotope Laboratories) and 1 g/l NH₄Cl (¹⁵N, 99%, Cambridge Isotope Laboratories).

**Protein expression and purification**

Proteins for structural and calorimetric studies were prepared by overexpression in *E. coli* BL21Star™(DE3) (Invitrogen) and Lemo21(DE3) (Wagner et al., 2008) (New England Biolabs) strains grown in LB medium. This included the C-terminally 6xHistidine-tagged TEV site-containing N-terminal extramembrane domain of YggP [6xHis-TEVsite-YggP(1–177); NTD], and the N-terminally 6xHistidine-tagged TEV site-containing C-terminal extramembrane domain of YggP [YggP(384–507)-TEVsite-6xHis; CTD] expressed from the pET25b or phIS-2 vector, respectively. The N-terminal cytosolic extramembrane domain of MgtE [MgtE(2–275)] for antibody production was overexpressed in *E. coli* Lemo21(DE3) as a GST-6xHistidine-TEVsite-MgtE(2–275) construct from the pGEX6P1 vector.

His₆-YggP(1–170) and His₆-YggP(385–507) for antibody production were expressed in *E. coli* BL21(DE3)pLysS from pRSET-A vector and purified using metal-chelate affinity chromatography.

Typically, protein production was induced with 0.5 mM IPTG at 25°C with aeration and gentle shaking overnight. Bacterial cultures were harvested by centrifugation at 5,000 × g for 10 min at 4°C, and cells were resuspended in isolation buffer composed of 20 mM HEPES, pH 7.4, 300 mM NaCl, 10% (v/v) glycerol, 10 mM imidazole, 1 mM PMSF and 1 × Complete™ EDTA-free Protease Inhibitor Cocktail (Roche). Cells were lysed by three passages through the high-pressure homogeniser Emulsiflex®-C3 (Avestin, Inc.), and lysates were cleared from the cell debris by centrifugation (15,000 × g, 30 min, 4°C). Supernatants were loaded on Ni-NTA Agarose (Qiagen) equilibrated in isolation buffer, and His-tagged protein was eluted specifically with increasing concentration of imidazole in elution buffer (20 mM HEPES, pH 7.4, 300 mM NaCl, 10% (v/v) glycerol). The GST-tagged MgtE(2–275) domain was additionally purified using Glutathione Sepharose® 4 Fast Flow (Merck) and specifically eluted by 10 mM reduced glutathione into 20 mM HEPES, pH 7.4, 300 mM NaCl, 10% (v/v) glycerol. The 6xHistidine tags from YggP domains and GST-6xHistidine tags from MgtE domain were removed by incubation with 6xHistidine-tagged TEV protease at protease-to-substrate ratio of ~1:200 (w/w), in 5 mM Tris–HCl, pH 8, 1 mM EDTA, 1 mM β-mercaptoethanol for 16 h at 25°C. The cleaved protein was then buffer-exchanged into isolation buffer (this paragraph) using PD-10 desalting column (GE Healthcare Life Sciences) and loaded onto Ni-NTA agarose. The flow-through fraction containing tag-free YggP or MgtE domains was collected, concentrated by ultrafiltration (Vivaspin, Sartorius) and used for further applications.

**In vitro translation**

Reference fragments of *B. subtilis* MgtE encompassing the first 300, 315, 330, 340, 355 and 370 residues of MgtE were generated by PCR amplification of the respective coding regions from *B. subtilis* 168 genomic DNA, and *in vitro*-transcribed and *in vitro*-translated as described previously (Lemberg & Martoglio, 2003; Strisovsky et al., 2009). Briefly, PCR was performed using set of mgtE-specific primers containing SP6 RNA polymerase promoter and ribosome-binding site in the forward primer, and stop codon in the reverse primer. Messenger RNAs were purified by LiCl precipitation and translated using Wheat Germ Extract (Promega). The resulting crude translation mixtures containing the MgtE reference fragments were separated using SDS–PAGE and visualised by polyclonal anti-MgtE(2–275) antibody and immunoblotting.

**Production of primary antibodies**

Purified His₆-YggP(1–170), His₆-YggP(385–507) and MgtE(2–275) were used to raise rabbit polyclonal antibodies (Agro-Bio, France). Forty-two days after immunisation, the sera were collected and purified by affinity chromatography on an Affi-Gel 10 (Bio-Rad) column containing covalently immobilised antigen as described before (Campo & Rudner, 2006). Briefly, approximately 2 mg of purified antigen was dialysed into coupling buffer (20 mM HEPES pH 8, 200 mM NaCl, 10% (v/v) glycerol) and coupled to 1 ml Affi-Gel 10 resin (Bio-Rad) as described by the manufacturer. After coupling,
the antigen resin was washed with 100 mM glycine pH 2.5 and neutralised with 1× PBS to remove all uncoupled protein. Twenty ml of antisera were batch-adsorbed to the antigen resin overnight at 4°C. The resin was loaded into a 5-ml column and washed with 20 ml 1× PBS, 50 ml 1× PBS 500 mM NaCl and 10 ml 0.2× PBS. The affinity-purified antibodies were then eluted with 100 mM glycine pH 2.5. The peak fractions were pooled and dialysed with three changes into 1× PBS with 50% (v/v) glycerol and stored at −20°C.

**Immunoblotting**

*Bacillus subtilis* cultures in exponential phase (OD<sub>600</sub> of ~1) were harvested by centrifugation (5,000 × g, 10 min, 25°C). Cell pellet was resuspended in lysis buffer containing 20 mM Tris–HCl, pH 7.5, 10 mM EDTA, 1 mM PMSF and 1× Complete<sup>®</sup> EDTA-free Protease Inhibitor Cocktail (Roche), 1 mg/ml lysozyme and 2500 U Pierce Universal Nuclease (cat no. 88700, Thermo Fisher) and incubated at 37°C, for 10 min. Reducing SDS–PAGE loading buffer was added, and the mixture was heated to 60°C for 10 min with vigorous shaking. Samples were separated on Tris-glycine SDS–PAGE and electroblotted onto a nitrocellulose membrane (Serva). Equivalent total protein amounts were loaded as estimated by the OD<sub>600</sub> values, and after electrotransfer, total protein in each lane was quantified using the revert total protein stain (LI-COR/C226). The peak fractions were pooled and dialysed with three changes into 1× glycine pH 2.5. The peak fractions were pooled and dialysed with three changes into 1× PBS with 50% (v/v) glycerol and stored at −20°C.

**Rhomboid activity assay in vivo**

*Bacillus subtilis* cells overexpressing particular YqgP variant under the control of P<sub>yif</sub> promoter were cultivated as described above. The amounts of the uncleaved and cleaved forms of MgtE or model substrate TatA were measured by quantitative fluorescence immunoblotting as described in the previous section. Since the integrated intensity of fluorescence signal for a given band is proportional to the molarity of the antigen, substrate conversion at steady state was quantified as the ratio of fluorescence intensities of bands corresponding to the cleavage product (P) divided by the sum of signals for the full-length substrate (S) and the cleavage product (P), i.e. signal(P)/[signal(S) + signal(P)]. Relative “specific activities” at steady state in vivo were then calculated by dividing substrate conversion by the signal from the expressed rhomboid enzyme (YqgP variant) corrected for the total protein signal in that gel lane.

**SILAC-based quantitative proteomics and data evaluation**

For SILAC-based quantitative proteomics, the BS50 and BS51 (Table EV2) strains of *B. subtilis* auxotrophic for lysine were cultivated in the medium described in section Bacterial strains, media and growth conditions. Equal amounts of heavy and light cell cultures (based on optical density at 600 nm, OD<sub>600</sub>) were mixed at harvest and centrifuged. Cell pellets were then resuspended in lysis buffer containing 20 mM HEPES, pH 7.4, 100 mM NaCl, 10% (v/v) glycerol, 1 mM EDTA, 50 μg/ml lysozyme and 0.4× MS-Safe Protease and Phosphatase Inhibitor Cocktail (Merck) and incubated at 37°C, for 15 min and on ice for 15 min, afterwards. Cells were lysed by sonication using SONOPPLUS HD2200 device (BANDELIN) set to 4 × 30 s pulse cycle and 30% power. Cell debris was removed by centrifugation (15,000 × g, 30 min, 4°C). Crude membranes were isolated by ultracentrifugation (100,000 × g, 1.5 h, 4°C) and washed sequentially with 0.1 M Na<sub>2</sub>CO<sub>3</sub> and 1 M NaCl.

*Bacillus subtilis* transmembrane protein-enriched fractions were separated on 4–20% gradient Tris–glycine SDS–PAGE system (Bio-Rad). Two SILAC experiments with swapped labelling were conducted: Experiment 1 [marked “heavy-to-light” or (H/L)]: heavy BS50 (YqgP<sup>g</sup>), light BS51 (YqgP<sup>S288A</sup>); and Experiment 2 [marked “light-to-heavy” or (L/H)]: light BS50, heavy BS51. Both experiments were resolved in separate gel lanes, which were subsequently sliced into five fractions (A–E) each (Fig 1), and individually digested using a standard in-gel digestion protocol. Briefly, every gel slice was first destained with 25 mM ammonium bicarbonate in 50% (v/v) acetonitrile (ACN, Merck), dehydrated and shrunk using acetonitrile (ACN) in a reducing environment of 5 mM 1,4-dithiothreitol (DTT, Merck) and incubated for 30 min at 65°C. Fractions were then alkylated using 12.5 mM iodoacetamide (ACN, Merck), shrunk and digested using 0.1 μg trypsin at 37°C overnight. Tryptic peptides were extracted with 60% (v/v) ACN in 0.5% (v/v) trifluoroacetic acid (TFA, Merck), dried and reconstituted in 20 μl of 2% (v/v) ACN with 0.1% (v/v) formic acid (FA, Merck) and analysed via LC-MS/MS.

The LC-MS/MS analyses were performed using UltiMate 3000 RSLcNANO System (Dionex) coupled to a TripleTOF 5600 mass spectrometer with a NanoSpray III source (AB Sciex). After injection, the peptides were trapped and desalted in 5% (v/v) ACN/0.1% (v/v) FA at a flow rate of 5 μl/min on an Acclaim<sup>®</sup> PepMap 100 column (5 μm, 2 cm × 100 μm ID, Thermo Scientific) for 5 min. The separation of peptides was performed on an Acclaim<sup>®</sup> PepMap 100 analytical column (3 μm, 25 cm × 75 μm ID, Thermo Scientific) using a gradient from 5% (v/v) ACN/0.1% (v/v) FA over 30 min. TOF MS scans were recorded from 350 to 1,250 m/z, and up to 18 candidate ions per cycle were subjected to fragmentation. Dynamic exclusion was set to 10 s after one occurrence. In MS/MS mode, the fragmentation spectra were acquired within the mass range of 100–1,600 m/z.

The quantitative mass spectrometric data files were processed and analysed using MaxQuant (v1.5.2.8) (Cox & Mann, 2008). The search was performed using a UniProt/Swiss-Prot *B. subtilis* database (downloaded 17/05/15) with common contaminants included. Enzyme specificity was set to trypsin, with methionine oxidation as a variable modification. Cysteine carbamidomethylation was considered as a fixed modification. The heavy SILAC label was set to K8, the minimal peptide length to 6, and two missed cleavages were
allowed. Proteins were considered as identified if they had at least one unique peptide, and quantified if they had at least one quantifiable SILAC pair. Transmembrane topology predictions were obtained using Phobius (Käll et al., 2004), and additional sequence visualisations were obtained using QARIP software (Ivankov et al., 2013; Fig 1).

Quantitative proteomics for interactor identification

All procedures were performed essentially as described (Doan et al., 2009). For crude membrane preparation, cells were grown in LB medium, at 37°C. At OD600 of 0.6, 50 ml of culture was harvested for membrane preparation. Membrane proteins were solubilised by the addition of the detergent NP-40 to a final concentration of 1% (v/v). The soluble fraction was mixed with 25 μl anti-GFP antibody resin (Chromotek, Germany) and rotated for 3 h at 4°C. The resin was pelleted at 3,000 × g, and the supernatant was removed. After washes, immunoprecipitated proteins were eluted by the addition of 85.5 μl of SDS–PAGE sample buffer and heated for 15 min at 50°C. The eluted material (the IP) was transferred to a fresh tube, and 2-mercaptoethanol was added to a final concentration of 10% (v/v).

Proteins extracted from the co-immunoprecipitation eluates were resolved using NuPAGE 4–12% system (Thermo Fisher Scientific), stained with Coomassie blue (R250, Bio-Rad) and in-gel digested using modified trypsin (Promega, sequencing grade) as described, previously (Casabona et al., 2013). Of note, samples from replicates 1 and 2 were prepared using oxidation and reduction–alkylation procedures as described (Jaquinod et al., 2012). The peptides were analysed by online NanoLC-MS/MS (UltiMate 3000 and LTQ Orbitrap Velos, Thermo Scientific). In particular, peptides were loaded on a 75 μm × 5 mm PepMap C18 precolumn (Thermo Scientific) and separated on a homemade 75 μm × 150 mm C18 column (Gemini C18, Phenomenex). MS and MS/MS data were acquired using Xcalibur (Thermo Scientific). Mascot Distiller (Matrix Science) was used to produce mgf files before identification of peptides and proteins using Mascot (version 2.6) through concomitant searches against UniProt (B. subtilis strain 168 taxonomy, September 2018 version), classical contaminants database (home-made) and the corresponding reversed databases. The Proline software (http://proline.profiproteomics.fr) was used to filter the results (conservation of rank 1 peptides, peptide identification FDR < 1% as calculated on peptide scores by employing the reverse database strategy, minimum peptide score of 25 and minimum of 1 specific peptide per identified protein group) before performing a compilation, grouping and comparison of the protein groups from the different samples. Proteins were considered as potential partners of the bait if they were identified only in the positive co-IPs with a minimum of three weighted spectral counts or enriched at least five times in positive co-IPs compared with control ones based on weighted spectral counts.

Isothermal titration calorimetry

The binding of divergent cations to the extramembrane domains of YqgP (NTD, CTD) was investigated using titration microcalorimetry. All titration experiments were performed in 50 mM Tris–HCl, pH 7.3, 150 mM NaCl at 25°C using Auto-iTC200 instrumentation (MicroCal, Malvern Pananalytical Ltd). Prior to the ITC measurements, protein samples were dialysed into the same buffer in which divalent cation salt solutions were prepared. Typically, 200 μl of the protein sample was titrated by stepwise 2 μl injections of divalent cation solution. Protein and ligand concentrations were adjusted as follows: 1.5 mM NTD was titrated by 50 mM MnCl2, 50 mM MgSO4, 50 mM ZnCl2 or 50 mM CaCl2, 0.7 mM NTD by 15 mM CoCl2 and 0.25 mM NTD by 7 mM NiCl2-buffered solutions. Control experiments included ligand titration into the buffer only. Concentrations of stock solutions of the analysed proteins were determined by quantitative amino acid analysis, and concentrations of the stock solutions of divalent metal cation salts were determined by elemental analysis. Titration data were processed and analysed using MicroCal Origin 7.0 (MicroCal, Malvern Pananalytical Ltd).

Nuclear magnetic resonance spectroscopy

For structure determination of YqgP N-terminal domain (YqgPNTD), 350 μl samples of 1 mM 15N/13C uniformly labelled protein in 25 mM phosphate buffer pH 6.5 containing 150 mM NaCl and 5% D2O/95% H2O were used. All NMR spectra for backbone assignments and three-dimensional structure determination were collected on 850 MHz Bruker Avance spectrometer equipped with triple resonance (15N/13C/1H) cryoprobe at 37°C. In a series of double and triple resonance, spectra (Renschaw et al., 2004; Veverka et al., 2006) were recorded to determine essentially complete sequence-specific resonance backbone and side-chain assignments. Constraints for 1H–1H distance required to calculate the structure of YqgPNTD were derived from 3D 15N/1H NOESY HSQC and 13C/1H NOESY HMBC, which were acquired using a NOE mixing time of 100 ms. The family of converged structures for YqgPNTD was initially calculated using Dyana 2.1 (Herrmann et al., 2002). The combined automated NOE assignment and structure determination protocol was used to automatically assign the NOE cross-peaks identified in the NOESY spectra and to produce preliminary structures. In addition, backbone torsion angle constraints, generated from assigned chemical shifts using the TALOS+ program (Shen et al., 2009), were included in the calculations. Subsequently, five cycles of simulated annealing combined with redundant dihedral angle constraints were used to produce a set of converged structures with no significant restraint violations (distance and van der Waals violations < 0.2 Å and dihedral angle constraint violation < 5°), which was further refined in explicit solvent using the YASARA software with the YASARA force field (Harjes et al., 2006). The structures with the lowest total energy were selected and analysed using the Protein Structure Validation Software suite (www.proteins.org). Cation-binding experiments were performed in 50 mM Tris–HCl, pH 7.3, 150 mM NaCl, 5% D2O, at 37°C, when the same aliquots of the ligand were added to the protein solution stepwise, after every titration measurement.

Data availability

The mass spectrometry (MS) proteome data have been deposited to the ProteomeXchange Consortium (http://proteomexchange.org) (Deutsch et al., 2017) via the PRIDE (Perez-Riverol et al., 2019) partner repository with the dataset identifiers PXD014578 (http://www.ebi.ac.uk/pride/archive/projects/PXD014578) and PXD014566 (http://www.ebi.ac.uk/pride/archive/projects/PXD014566).
The NMR structural data have been deposited in the Protein Data Bank (PDB) under accession code 6R0J (http://www.rcsb.org/pdb/explore/explore.do?structureId=6R0J) and in the Biological Magnetic Resonance Data Bank (BMRB) under the accession code 34376 (http://www.bmrb.wisc.edu/data_library/summary/index.php?bmrbdId=34376).

Expanded View for this article is available online.

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Author contributions
KS, TD, BC and JBe designed research; JBe, BC, TD, VV, RH, PS, MK, JD, MB, YC and PR performed research and analysed data; and KS and JBe wrote the manuscript with contributions from TD, JBr, BC, VV, AG, RH, PS, YC and MK.

Conflict of interest
The authors declare that they have no conflict of interest.

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