INTRODUCTION

Streptococcus suis, an emerging zoonotic pathogen, causes invasive diseases in pigs, including sepsis, meningitis, endocarditis, pneumonia, and arthritis. Importantly, similar pathologies are reported in human S. suis infections. In previous work, the locus SSU0375 of S. suis strain P1.7 had been identified as a conditionally essential gene by intrathecal experimental infection of pigs with a transposon library of S. suis. This study aimed to identify the function of the corresponding gene product. Bioinformatics analysis and homology modeling revealed sequence and structural homologies with the Streptococcus pneumoniae mid-cell-anchored protein Z (MapZ) that is involved in cell division in different bacterial species. Indeed, depletion of this locus in S. suis strain 10 revealed a growth defect as compared to the wild type. Electron microscopy analysis of the corresponding mutant demonstrated morphological growth defects as compared to the wild-type strain, including an irregular cell shape and size as well as mispositioned division septa. Light microscopy and subsequent quantitative image analysis confirmed these morphological alterations. In the genetic rescue strain, the wild-type phenotype was completely restored. In summary, we proposed that SSU0375 or the corresponding locus in strain 10 encode for a S. suis MapZ homolog that guides septum positioning as evidenced for other members of the Streptococci family.

KEYWORDS
cell division, electron microscopy, MapZ, quantitative image analysis, Streptococcus suis
in swine and humans are caused by serotype 2 strains (Goyette-Desjardins et al., 2014). However, infections with serotype 9 strains are becoming increasingly important in several countries, particularly in Western Europe (Goyette-Desjardins et al., 2014; Schultz et al., 2012). Besides, the number of antibiotic-resistant strains in S. suis isolates has increased over the past years (Palieri et al., 2011; Tan et al., 2021). Therefore, new therapeutics against streptococcal infections are needed and cell division proteins might be interesting targets for drug development (Shi et al., 2014).

Bacterial binary fission is achieved by accurate placement of the division machinery (Adams & Errington, 2009). For that, bacteria have evolved different mechanisms to identify the cell middle and to position the division machinery, of which the first component is the tubulin-like protein FtsZ (filamentous temperature-sensitive protein Z) (reviewed in (Mahone & Goley, 2020)). After binding to GTP FtsZ assembles into protofilaments to form a contractile ring, the Z-ring. This ring eventually constricts and gives rise to the newborn cells. Two systems, the nucleoid occlusion, and the Min system, earlier emerged to explain the positioning of the Z-ring or mechanisms to control its formation (Bramkamp & van Baarle, 2009; Rothfield et al., 2005). However, several bacteria lack these systems. In recent years, alternative positive regulatory systems were discovered, including the protein MapZ and the serine/threonine kinase StkP. MapZ was first identified in Streptococcus pneumoniae (Fleurie et al., 2014; Holeckova et al., 2014), but it is conserved in Enterococcaceae and Streptococcaceae, the two main families of Lactobacillales (Garcia et al., 2016). MapZ was identified to be a substrate of the Ser/Thr kinase StkP (Manuse, Fleurie, et al., 2016). In newborn cells, MapZ colocalizes with FtsZ at mid-cell before it splits into two rings that are pushed apart by nascent peptidoglycan to the cell equators of the future daughter cells. There MapZ positions the FtsZ ring through direct protein-protein interactions. At the same time, a third MapZ ring positions itself at the constriction site and remains until cell division is completed. Thereby, MapZ serves as a permanent beacon of the cell division site (Fleurie et al., 2014; Holeckova et al., 2014). Recently, Zamakhaeva et al. (2021) showed that in Streptococcus mutans MapZ recognizes peptidoglycan decorated with immature serotype c carbohydrates (SCCs), which is primarily located in the equatorial rings and cell poles. Thus, it is being proposed that immature SCCs could serve as a point of reference for the correct positioning of MapZ during cell division.

The MapZ protein consists of a C-terminal extracellular domain and an N-terminal cytoplasmic domain connected by a transmembrane domain (Manuse, Jean, et al., 2016). The intracellular domain belongs to the intrinsically disordered proteins and directly interacts with FtsZ and liposomes in vitro (Hosek et al., 2020). The extracellular domain is divided into two subdomains separated by a serine-rich flexible linker (SRL). The N-terminal subdomain of MapZ<sub>extra</sub>, and the SRL behave as a pedestal for the function of the C-terminal subdomain, which interacts with peptidoglycan. This C-terminal subdomain includes seven conserved amino acids that are essential for MapZ positioning at the cell equator (Manuse, Jean, et al., 2016).

Furthermore, MapZ acts as a guiding track for treadmilling FtsZ filaments and controls the correct position and angle of the FtsZ ring (Li et al., 2018). Whether MapZ-FtsZ-interaction is sensitive to FtsZ polymerization needs more evaluation since there are contrary findings in the literature (Feng et al., 2019; Hosek et al., 2020). In contrast to former studies, van Raaphorst et al. (2017) provided evidence that MapZ indeed sets the correct division plane in pneumococci, but it is involved neither in the selection of the division site nor in the accurate timing of Z-ring assembly. Mutants devoid of MapZ show an aberrant cell shape and size with mispositioned division septa, formation of cell clusters, and an increase in generation time (Fleurie et al., 2014; Holeckova et al., 2014).

In a recent study using intrathecal infection of pigs with a transposon library of S. suis, the locus SSU0375 was identified as conditionally essential for survival in the blood (Arenas et al., 2020). Its predicted protein sequence shows homologies with the mid-cell-anchored protein Z of S. pneumoniae. Therefore, the objective of this work was to analyze whether locus SSU0375 (or its corresponding locus in strain 10) codes for S. suis MapZ.

2  |  MATERIAL AND METHODS

2.1  |  MapZ in silico analysis

MapZ sequences were extracted from the GenomeNet database KEGG. The sequences were aligned with T-Coffee (Notredame et al., 2000) and the alignment was subsequently visualized using TEXshade (Beitz, 2000b). The schematic topology model of the MapZ protein of S. suis (Figure A2a) was predicted with the TMHMM Server v. 2.0 (http://www.cbs.dtu.dk/services/TMHMM, 18.05.21) and visualized with TEXtopo (Beitz, 2000a). Disordered regions in the protein were predicted using PrDOS (Protein DisOrder prediction System) (Ishida & Kinoshita, 2007) (http://prdos.hgc.jp/cgi-bin/top.cgi, 19.03.21). 3D-homology modeling of the extracellular domain was done using the molecular graphics package UCSF Chimera 1.15 (https://www.rbvi.ucsf.edu/chimera) (Pettersen et al., 2004) and is based on the two known subdomain structures of the extracellular domain of the pneumococcal MapZ protein (2nd9.pdb and 2nda.pdb). Alignment and visualization of the structures were performed with UCSF Chimera.

2.2  |  Bacterial strains

The virulent S. suis serotype 2 strain 10 was kindly provided by Smith et al. (1999) (Lelystad). Its isogenic mapZ-deficient mutant, hereafter called 10∆mapZ, was constructed by the insertion of a spectinomycin cassette in the corresponding putative mapZ gene (locus tag GPW51_RS02050). The generation of mutants in strain 10 has been previously described (Arenas et al., 2020). In brief, a spectinomycin-resistance cassette and the flanking regions of the mapZ gene were amplified from the genomic DNA of strain 10. A new PCR reaction was performed with the spectinomycin-resistance cassette. The cassette was fused to DNA segments of the flanking genes, which were amplified using primer pairs described in Table A1. The final PCR products were...
used to transform strain 10 using a competence peptide protocol (Zaccaria et al., 2014). The resulting transformants were selected on Todd-Hewitt Broth (THB; Bacto™, Becton Dickinson) agar plates supplemented with 100 µg/ml spectinomycin (Sigma-Aldrich), and the correct mutants were identified by PCR. To produce the MapZ protein in the 10ΔmapZ mutant, the mapZ gene from strain 10, including a chloramphenicol-resistance cassette, was reintroduced on the chromosome (Figure A1a). The resulting strain was called cMapZ. For this, mapZ and the flanking genes were amplified from the genomic DNA of strain 10. The PCR product was ligated into the pJET1.2/blunt vector using the CloneJET PCR Cloning Kit (Thermo Fisher Scientific) and introduced into Escherichia coli DH5α (Thermo Fisher Scientific) by conventional transformation methods. A chloramphenicol-resistance cassette was included upstream of the mapZ gene via in vivo assembly cloning (García-Nafria et al., 2016). Plasmids and PCR products were purified by commercial kits (NEB, New England Biolabs) and confirmed by sequencing. Then the genes of interest and the chloramphenicol-resistance gene were amplified again. The resulting amplicon was purified and used to transform 10ΔmapZ as previously described (Zaccaria et al., 2014). The transformants were selected on THB agar (Becton Dickinson) plates supplemented with 4 µg/ml chloramphenicol (Carl Roth). All PCR reactions were conducted with Phusion polymerase (New England Biolabs). Positive clones were identified by colony PCR. Bacterial colonies were resuspended in water and then incubated in a microwave for 10 min at 100 W. Regular PCR reactions were performed with HotStartTaq Plus DNA polymerase (Qiagen). All primers used in this study are described in Table A1.

### 2.3 Growth experiments

*Streptococcus suis* strain 10, its 10ΔmapZ derivative, and cMapZ were cultured overnight at 37°C on Columbia agar supplemented with 7% (v/v) sheep blood (Thermo Fisher Scientific) under aerobic conditions. For liquid cultures, two to three single colonies were inoculated in a starting culture in THB medium supplemented with the corresponding antibiotic and incubated at 37°C overnight. On the next day, bacterial cultures were adjusted to an OD600 of 0.001 in fresh THB medium without antibiotics. The OD600 was measured every 30 min for 8 h. Growth rates were determined by nonlinear regression curve fit.

### 2.4 qRT-PCR

Bacterial strains were prepared as described in 2.3 and grown to an OD_{600} of approximately 0.2. Subsequently, samples were fixed by adding methanol-free formaldehyde (Polysciences) to a final concentration of 5% directly into the culture and stored at 4°C. After washing the samples twice with TE buffer (0.01 M Tris, 1 mM EDTA; pH 6.9), they were adsorbed onto poly-l-lysine coated 12 mm coverslips, left for 10 min, and then fixed with 1% glutaraldehyde for 10 min. After another TE buffer washing step, samples were dehydrated with a graded series of acetone (10%, 30%, 50%, 70%, 90%, and 100%) on ice for 15 min for each dilution. After samples in 100% acetone reached room temperature, they were exposed to critical point drying with liquid CO2 (Leica CPD300). Subsequently, samples were covered with a gold-palladium film (approximately 10 nm) by sputter coating (Bal-Tec SCD500, Balzers Liechtenstein) before examination in a field emission scanning electron microscope (FESEM) Merlin™ (Carl Zeiss). Images were taken with the in-lens SE-detector at an acceleration voltage of 5 kV.

For ultrathin sections, fixed and TE buffer washed bacteria were osmificated with 1% aqueous osmium tetroxide for 1 h at room temperature, washed with TE buffer, and further dehydrated with a graded series of ethanol (10%, 30%, 50%). The 70% step was performed with 4% uranyl acetate in ethanol overnight. Then, dehydration was completed with 90% and 100% ethanol. Infiltration with the embedding resin LRWhite was performed with 1 part resin/1 part 100% ethanol, 2 parts resin/1 part 100% ethanol, followed by pure resin with several changes. Samples were placed into gelatin capsules, filled with pure resin, and polymerized at 50°C for 2 days. Ultrathin sections were cut with a diamond knife. Sections were collected onto butvar-coated copper grids (300 mesh) and counterstained with 4% uranyl acetate for 4 min. Examination of the samples was done using a transmission electron microscope TEM910 (Carl Zeiss) at an acceleration voltage of 80 kV and calibrated magnifications.
2.6 | Light microscopy and image analysis

Bacterial strains were prepared as described in 2.3 and grown to an OD_{600} of approximately 0.3. Then 2 µl of the bacterial culture were spotted on 1% low-melting agarose (GERBU) pads. The samples were analyzed using a Nikon Eclipse Ti-S microscope (Nikon), equipped with the objective Plan Fluor 100×/1.3. Phase-contrast images for quantification were acquired in a blinded manner in 5–15 randomly picked positions per sample.

For segmentation of bacteria, we used STARDIST version 0.6.2 (Schmidt et al., 2018), a deep learning framework based on the popular U-Net architecture to detect and segment individual bacteria. Training data sets were generated by first segmenting a set of training images with the Fiji/ImageJ plugin MicrobeJ (Ducret et al., 2016) and subsequent manual correction of suboptimal masks. Training data were further augmented by 90°-rotation degrees before training. During the training process additional random horizontal and vertical flips, intensity changes, and addition of Gaussian noise were applied to further augment the training data. All images were normalized by simple percentile-based normalization. Eighty percent of the images were used for training and 20% were used for testing. Input image size was 512 x 512 pixels. Details on network architecture and training parameters are summarized in Table A2. The training was performed on an Nvidia Titan RTX GPU. The created masks were used for quantitative image analysis of bacterial cells with the Python library scikit-image (van der Walt et al., 2014). In total, more than 2000 bacteria per strain were analyzed. In some images, obvious abiotic particles were excluded manually from the analysis. Raw data for length and width measurements are provided in the supplementary datasets S1 and S2 (https://doi.org/10.5281/zenodo.5172241). This experiment was performed at least three times with different biological replicates.

2.7 | Van-FL staining

Bacterial strains 10 and 10ΔmapZ were prepared as described in 2.3 and grown until an OD_{600} of approximately 0.2. Then 1 ml of the bacterial culture was transferred into a sterile 1.5 ml Eppendorf tube and 5 µl Vancomycin-FL (stock: 100 µg/ml; Thermo Fisher Scientific) as well as 5 µl vancomycin (stock 100 µg/ml; Carl Roth) were added. The cultures were vortexed and incubated at 37°C for additional 20 min. Afterward, the cultures were centrifuged for 10 min at 1690 x g at 4°C. The pellets were washed with HBSS buffer (Thermo Fisher Scientific), resuspended in PBS buffer (Sigma-Aldrich), and fixed in PBS buffer containing 3% formaldehyde. Bacteria were transferred to poly-l-lysine (Sigma-Aldrich) coated coverslips, mounted with ProLong® Gold Antifade Reagent (Cell Signaling Technology), and stored at 4°C until examination.

Confocal microscopy was performed using a TCS SP5 confocal laser scanning microscope equipped with a 63x 1.40–0.60-NA oil HCX Plan Apochromat objective (Leica). Image stacks with a z-distance of 0.13 µm per plane were acquired using a 1-Airy-unit pinhole diameter. Images were deconvolved using Huygens® Essential 20.10 (Scientific Volume Imaging). Maximum intensity projections of 3D-stacks were generated and adjusted identically for brightness and contrast in ImageJ/Fiji (Schindelin et al., 2012).

2.8 | Statistical analysis

Values are expressed as mean ± Standard Deviation (SD) of at least three independent experiments. Growth curves were analyzed via nonlinear regression fit with GraphPad PRISM Software 8.01 for Windows (GraphPad Software). Data in Figure 4 were analyzed using estimation statistics with the DABEST package v0.3.1 (Ho et al., 2019). The generated Cumming plots display the magnitude and robustness of the effect size and its bootstrapped 95% confidence interval (CI).

3 | RESULTS AND DISCUSSION

3.1 | In silico analysis of Streptococcus suis MapZ

The putative mapZ gene of strain 10 had been previously identified as a conditionally essential gene for survival in the blood (Arenas et al., 2020). Alignment of the predicted MapZ protein of strain 10 revealed an overall homology of 40% with S. pneumoniae MapZ (UniProtKB Q8DR55) (Figure 1a). The C-terminal region showed a higher identity with pneumococcal MapZ (47%) than the N-terminal region (30%) and the linker region (lower than 30%). Previous works described the relatively low conservation of the linker region as well as the intracellular domain of MapZ within Enterococccaeae and Streptococcaceae family members (Garcia et al., 2016). Furthermore, the seven amino acids (R429, Y431, N448, Y450, Y470, F471, N474) that were claimed essential for the binding of peptidoglycan (Manuse, Jean, et al., 2016), and consequently the correct positioning of MapZ at mid-cell, were conserved in both proteins (Figure 1a).

In a previous study, S. suis MapZ was identified as a substrate of the serine/threonine kinase STK that phosphorylates threonine 66 (T66) (Zhang et al., 2017). Similar phosphorylation of T67, besides threonine 78, was observed in the MapZ protein of S. pneumoniae and it was suggested that phosphorylation/dephosphorylation of these residues is important for controlling FtsZ function (Fleury et al., 2014; Holecíková et al., 2014). Nonetheless, alignment of the MapZ proteins from S. suis and S. pneumoniae revealed that the second threonine found in the pneumococcal protein is not conserved in S. suis (Figure 1a). Both, the seven amino acids important for MapZ positioning as well as the two phosphorylation sites are conserved in S. mutans (Li et al., 2018). A sequence alignment of MapZ homologs in an earlier study revealed that the two phosphorylation sites are not highly conserved in Streptococci. Some species include both phosphorylation sites (e.g., S. pneumoniae, S. mutans), others include only the first (e.g., S. suis, Streptococcus sanguinis) or the second one (e.g., Streptococcus pyogenes, Streptococcus equis subsp. zooepidemicus) or none of them (e.g., Streptococcus intermedius) (Holecíková et al., 2014; Holecíková et al., 2014). Nonetheless, alignment of the MapZ proteins from S. suis and S. pneumoniae revealed...
et al., 2014). Hosek et al. (2020) described that phosphorylation of MapZ does not affect its structural organization or its interactions with other proteins such as FtsZ. However, the phosphomimetic as well as the phosphoablative form of the pneumococcal MapZ result in cell shape and viability defects. While correct FtsZ positioning is not impaired in these mutants, FtsZ ring structure and numbers are affected (Fleurie et al., 2014). In contrast, Holeckova et al. (2014) did not observe any differences when performing phenotypic analysis of the phosphoablative and the phosphomimetic mutant in comparison to the wild-type strain. Offtarget effects might be the reason for these differences. Here, posttranslational modification of the MapZ protein by phosphorylation might play a role in the division process (Bramkamp, 2015). Altogether, the predicted MapZ protein is divided into a cytoplasmic domain, a transmembrane domain (TM), and an extracellular domain subdivided into two subdomains (Extra 1, 2) which are separated by a flexible linker. Numbers indicate amino acid positions.

Despite the large variability of the extracellular region, the predicted MapZ protein of S. suis is composed of four alpha helices while the MapZ protein of S. pneumoniae results from different insertions along the domains (Figure A3a).

Despite the reduced homology between S. suis MapZ and S. pneumoniae MapZ proteins, a comparison of predicted protein topology and structure revealed conservation of the relevant domains with little variation. The linker region in the extracellular domain is predicted to be highly disordered (Figure A2b,c). The domain boundaries in S. suis are indicated in Figure 1b. Streptococcus suis MapZ is 20 amino acids longer than S. pneumoniae resulting from different insertions along the domains (Figure 1a, Figure A3a).

Despite the large variability of the extracellular region, the predicted MapZ protein is composed of four alpha helices while the MapZ protein of S. suis is composed of a combination of two small alpha helices and beta strands interconnected with different loops (Figure A3b, Supplementary Movie 1: https://doi.org/10.5281/zenodo.5172241), as described for pneumococcal MapZ using NMR assays. Disordered regions are unable to form stable 3D structures under physiological conditions (Varadi et al., 2014). However, they carry out important cellular functions (Dunker et al., 2002). As our protein predictions are based on homologies to known structures, the tertiary structure of the linker region was not predicted. Intrinsically disordered regions are composed of mainly polar and charged amino acids conferring high flexibility of the respective
region (Xue et al., 2009). They mainly connect the functional domains and generate flexibility between them. Therefore, they often act as domain linkers (Dunker et al., 2002). Consequently, despite the differences in amino acid composition between both MapZ proteins, a disordered linker domain is conserved in \( S_{suis} \), which probably is important for the function or stability of the protein.

In summary, \textit{in silico} analysis of the protein encoded by the SSU0375 gene locus of \( S_{suis} \) suggests that this is a MapZ homolog to the pneumococcal MapZ protein.

### 3.2 Inactivation of MapZ results in cell division defects and septum misplacement

Our \textit{in silico} studies suggested that SSU0375 codes for \( S_{suis} \) MapZ. To further demonstrate the protein identity, we studied the associated phenotype. Diverse studies on MapZ homologs in other streptococcal species revealed morphological growth defects in mutants lacking MapZ (Fleurie et al., 2014; Li et al., 2018). Therefore, we performed growth experiments and microscopic analysis of strain 10 and its isogenic mutant 10\( \Delta \)mapZ. Growth experiments revealed a longer lag phase and a reduced growth rate of the mutant compared to the wild type. On the chromosome, \textit{mapZ} is placed in an operon. Thus, we wondered whether the observed phenotype could be derived from alterations in flanking genes. To prove that, we tested the gene expression of \textit{mapZ} and its flanking genes using qRT-PCR assays. The results showed alteration of \textit{mapZ} expression but not of flanking genes (Figure A1b). We then wanted to complement the observed phenotype by reintroducing the \textit{mapZ} gene on the chromosome of 10\( \Delta \)mapZ. RT-qPCR assays of strain cMapZ showed restoration of \textit{mapZ} expression (Figure A1b) and observed growth defects (Figure A1c). A similarly reduced growth rate was reported for the pneumococcal \( \Delta \text{mapZ} \) mutant (Fleurie et al., 2014; Holeckova et al., 2014) and \( S_{mutans} \) (Li et al., 2018). Accordingly, we concluded that the observed phenotypes are caused by the defects in MapZ production.

To investigate the morphology of the different strains, bacteria grown to the early logarithmic phase were fixed and analyzed by electron microscopy. Results showed typical elongated ellipsoidal cells for the wild type that divided perpendicular to their long axis into successive parallel planes resulting in the formation of short chains. In
contrast, the MapZ-deficient mutant displayed morphological defects including an irregular cell shape and size of the daughter cells as well as the formation of clusters and small-sized cells (often referred to as minicells; Figure 2). The genetic rescue strain showed a regular cell shape and size comparable to the wild type. Transmission electron microscopy further showed that the division septa were not placed perpendicular to the longitudinal axis in the mutant and that many cells even had several septa located in different axes (Figure 3). These deviations caused a cell division along more than one axis and thereby generated cell clusters. Fleurie et al. reported similar defects for the pneumococcal MapZ mutant (Fleurie et al., 2014). They observed aberrant cell shapes, mispositioned division septa as well as the formation of grape-like clusters (Fleurie et al., 2014). In agreement with our observations, the absence of MapZ in *S. pneumoniae* leads to the formation of minicells that are lacking DNA (Holeckova et al., 2014). The formation of these unproductive cells may explain a reduced growth rate of the mutant with respect to the wild type. Holecková et al. also reported the presence of multiple division septa in some of the daughter cells (Holeckova et al., 2014), which is in concordance with our observations.

Next, we wanted to quantify these morphological alterations in an unbiased, automatic way. Therefore, we performed a quantitative image analysis. Different cell sizes, morphological changes as well as the formation of aggregates were evident in the MapZ mutant in phase-contrast images while the wild type and the complemented mutant displayed comparable, regular cell shapes (Figure A4). Microscopy pictures were processed via StarDist (Schmidt et al., 2018), a deep learning based application. The mutant showed a slightly reduced cell length with a mean difference of $-0.086 \mu m$ (95% CI: $-0.181, 0.014$; Figure 4a), and significantly higher cell width of $0.075 \mu m$ (95% CI: 0.060, 0.090) than the wild type.
The morphology of the strain cMapZ was very similar to the wild type. The mean length difference between these strains was 0.026 µm (95% CI: −0.075, 0.143; Figure 4a) and the mean width difference was 0.014 µm (95% CI: −0.005, 0.030; Figure 4b). Therefore, quantitative image analysis verified our visual interpretations of the electron microscopy assays. Quantitative image analysis of the pneumococcal MapZ deletion mutants revealed significantly shorter cells compared to the wild type and differences in the cell length distribution (Holeckova et al., 2014). Nevertheless, we could not observe a significant difference between the cell length of the three strains, although indeed we observed minicells in our electron microscopy assays (Figure 2). Probably, the percentage of minicells

**Figure 4** Quantitative image analysis of phase-contrast images of strain 10, 10ΔmapZ, and cMapZ with StarDist. Results are depicted as Cumming plots. The upper part of the plot shows the mean ± SD of cell length (a) or cell width (b) of four independent experiments. Each dot represents the average of one individual experiment and the violin plot indicates the distribution of all data pooled. The lower part shows effect sizes (Δmean) as dots compared to strain 10 with corresponding bootstrap 95% confidence intervals and resampling distributions of the difference in means.

**Figure 5** Van-FL staining visualizes nascent peptidoglycan synthesis. *Streptococcus suis* strain 10 (a) and 10ΔmapZ (b) were grown to the early log-phase and incubated with fluorescent vancomycin for 20 min. Stained cells were analyzed by confocal microscopy. Scale bars represent 2 µm and 1 µm in enlarged insets. Closed arrowheads indicate cell septa located in the middle of the cell; open arrowheads point at fuzzy fluorescent signals.
in the entire bacterial population is reduced in S. suis compared to S. pneumoniae mapZ mutants.

In conclusion, the phenotype observed in the mapZ mutant is in good agreement with that observed for mapZ mutants in other bacteria, confirming that our presumptive mapZ gene encodes for MapZ.

3.3 | Van-FL staining visualizes nascent peptidoglycan

Given the conservation of amino acids important for peptidoglycan binding and to further analyze the role of MapZ during septum formation, we stained the bacteria with fluorescently labeled vancomycin (Van-FL) to visualize sites of nascent peptidoglycan synthesis (Daniel & Errington, 2003).

Van-FL staining of S. suis strain 10 indicated correct peptidoglycan synthesis and positioning. The cell septa were located at the cell center and perpendicular to the longitudinal axis of the cell (Figure 5a). Although peptidoglycan synthesis was also detected in the corresponding mapZ mutant, the staining revealed mispositioned cell septa at an irregular angle and frequently shifted from the cell center. Often only a fuzzy fluorescent signal was detectable (Figure 5b). Similar results were reported for the pneumococcal MapZ-deficient mutant (Holeckova et al., 2014). Thus, the abnormal/misplaced peptidoglycan insertion might be responsible for the abnormal cell division in the mutant strain, which we also observed by electron microscopy. For S. pneumoniae van Raaphorst et al. (2017) revealed that the angle of the Z-ring is significantly more skewed in the mutant strain. The authors suggest that MapZ might have a function in cell wall remodeling and maintaining the perpendicular Z-ring plane due to its large peptidoglycan binding domain. Li et al. observed that changes in the angle of the septum mainly occurred during the early formation of the Z-ring and not during its constriction. Thus, the authors conclude that the MapZ protein of S. mutans acts as a track for FtsZ filaments only during the early stages of cell division (Li et al., 2018). To underline the results of the Van-FL staining and the electron microscopy, direct labeling of the Z-ring in S. suis needs to be addressed in future studies, either by a GFP-fusion or an FtsZ-specific antibody. Thereby not only the positioning but also the angle and structure of the Z-ring can be investigated. This is also interesting in the context of phosphorylation as Fleurie et al. stated effects on FtsZ ring structure and number in phosphomimetic as well as phosphoablative mutants (Fleurie et al., 2014). Altogether, our results confirm previous observations of the pneumococcal MapZ in septum placement and suggest a conserved function within streptococcal species despite the amino acid variability of the protein (Figure 1a). However, the exact mechanism, as well as the role of phosphorylation, need to be elucidated in future studies.

4 | CONCLUSION

In conclusion, our study showed that S. suis locus SSU0375 encodes for a homolog of the MapZ protein. As for other streptococci, MapZ is responsible for proper septum placement and is involved in cell division. The lack of MapZ leads to an increase in generation time as well as morphological growth defects, both causing reduced biological fitness of the mutant in vivo, which may explain the necessity of this protein during infection. Previously, cell division proteins have been proposed as interesting targets for drug development (Shi et al., 2014). The physiological role of MapZ in cell division and its demonstrated relevance during infection make this protein a strong candidate for the development of new therapeutics. Future studies will focus on demonstrating how this gene contributes to the survival of S. suis in its host and the role of phosphorylation in the function of the MapZ protein.

ACKNOWLEDGEMENTS

We would like to thank Hilde E. Smith (Wageningen Bioveterinary Research, Lelystad, The Netherlands) for providing S. suis strain 10 and Ina Schleicher for excellent technical assistance. Part of this work was contributed by M.D. in partial fulfillment of the requirements for a Ph.D. degree from the University of Veterinary Medicine Hannover. This project received funding to P.V.-W. from the European Union’s Horizon 2020 research and innovation program under grant agreement no. 727966.

CONFLICTS OF INTEREST

None declared.

AUTHOR CONTRIBUTIONS

Muriel Dresen: Conceptualization (equal); Formal analysis (equal); Investigation (lead); Writing-original draft (lead); Writing-review & editing (equal). Manfred Rohde: Investigation (supporting); Writing-review & editing (supporting). Jesús Arenas: Resources (supporting); Writing-review & editing (supporting). Astrid de Greeff: Resources (supporting); Writing-review & editing (supporting). Andreas Nerlich: Conceptualization (equal); Formal analysis (equal); Investigation (supporting); Software (lead); Supervision (lead); Writing-original draft (supporting); Writing-review & editing (equal). Peter Valentijn-Weigand: Conceptualization (supporting); Funding acquisition (lead); Supervision (supporting); Writing-review & editing (equal).

ETHICS STATEMENT

None required.

DATA AVAILABILITY STATEMENT

All data generated or analyzed during this study are included in this published article except for the supplementary movie of the homology model, the code for training and analysis of microscopic phase contrast images as well as dataset S1 and S2 that are available in the GitHub repository at https://github.com/AndreasNerlich/Streptococcus_suis_MapZ and in Zenodo: https://doi.org/10.5281/zenodo.5172241.

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How to cite this article: Dresen, M., Rohde, M., Arenas, J., de Greeff, A., Nerlich, A., & Valentijn-Weigand, P. (2021). Identification and characterization of the cell division protein MapZ from Streptococcus suis. MicrobiologyOpen, 10, e1234. https://doi.org/10.1002/mbo3.1234
APPENDIX

FIGURE A1 Characterization of *Streptococcus suis* mutants by colony PCR, growth in THB media, and qRT-PCR. (a) Amplification of mapZ with a fragment size of 1018 bp, genomic DNA served as a positive control, amplification of spectinomycin-resistance cassette with a fragment size of 487 bp, genomic DNA from a *S. suis* strain containing a spectinomycin cassette served as a positive control and amplification of chloramphenicol-resistance cassette with a fragment size of 1018 bp. pSET5s plasmid served as a positive control. Used primers are described in Table A1. (b) The different bacterial strains were grown to the early log phase; RNA was isolated and reverse transcribed followed by qRT-PCR. The expression levels of the tested genes were normalized using the *gyrB* gene of *S. suis* as an internal standard and are presented as x-fold to the control gene. Results show mean ± SD of three independent experiments. nd, not detectable. (c) Growth of S10 wild type (gray), 10ΔmapZ (dark red), and cMapZ (light red) in THB media. Starting OD₆₀₀ was adjusted to 0.001. OD₆₀₀ was determined every 30 min for 8 h. The results show a nonlinear regression curve with mean ± SD of three independent experiments. The generation time of the different bacterial strains grown in THB is displayed in the table on the right. Generation time ± SD is shown.
Figure A2: Prediction of Streptococcus suis MapZ topology. (a) Prediction of topology using TMHMM v2.0. (b) Prediction of disordered regions using PrDOS. Ordered residues are shown in black, disordered residues are in red. (c) Disorder probability of the results in (b). The threshold is indicated by the red line. The analysis was performed with a false positive rate of 5%.
FIGURE A3 Predicted topology and structure of MapZ. (a) Predicted topology of MapZ. MapZ consists of an intracellular domain, a transmembrane domain, and an extracellular domain. Conserved amino acid positions are marked in blue, whereas similar positions are marked in pink. Prediction of membrane protein topology was performed with the TMHMM server and visualization was done with TEXtopo. (b) 3D-homology model of the extracellular domain of MapZ of Streptococcus suis based on structures of the pneumococcal protein. Homology modeling was performed with UCSF CHIMERA. The prediction of the MapZ protein of S. suis is shown in gray and the two known subdomain structures of the extracellular domain of pneumococcal MapZ in green (2nd9.pdb) and red (2nda.pdb). The extracellular domain of MapZ is divided into two subdomains which are separated by a flexible linker.

FIGURE A4 Representative phase-contrast images for quantitative image analysis of strain 10 (a), 10ΔmapZ (b), and cMapZ (c). The arrowhead indicates cluster formation of the mutant. Scale bars represent 4 µm.
| Purpose | Primer | Sequence (5′–3′) |
|---------|--------|-----------------|
| 10ΔmapZ mutant | **First PCR** | Fssu0375-L1 CAAGGCCAAATGTGTCAAATC |
| | | Rssu0375-L2 ACGAACGAAAAATCGACCTGCTTCGCACTTCAAAATCTAAAAC |
| | **Second PCR** | Fssu0375-R2 TTAGAAAAACAAATACCTTTGACATTATAACAGCARGGCAAATGGCTA |
| | | Rssu0375-R1 ATGGGCTAATCGACTGCTCTC |
| | **Amplification of Spec cassette** | Fspec GCAGGTCGATTTTGGTCCGT |
| | | Rspec ATGCAAGGTGTTTATTGTTTTCTAA |
| cMapZ mutant | **Cloning of SSU0375 into pJET** | SSU0374_for GTGGGTTGAGGATCAATTGG |
| | | SSU0376_rev CATCACTTGGGGAGGATGTTTC |
| | **Plasmid PCR** | SSU0375_IVA_pJet_f_2 CCGATATGAGGTGCACTTCAATCCGAGGC |
| | | SSU0375_3′_IVA_rev_2 TCGCTGAGATAGTGCCGAAACTTTTCTTTCTTCATCCGAGGC |
| | **Amplification of Cm cassette** | CM_for GGCACCTATCTCAGCGATC |
| | | CM_rev TCCGATAATTGATGTTTCC |
| | **Amplification for transformation** | SSU0374_for GTGGGTTGAGGATCAATTGG |
| | | SSU0376_rev CATCACTTGGGGAGGATGTTTC |
| Colony PCR | **Amplification of SSU0375** | SSU0375_Seq_for ACCGAAGACTTGGGACATAG |
| | | SSU0375_Sall_rev_2 GTGGGTTGAGGATCAATTGG |
| | **Amplification of Spec cassette** | Spec_for AGTCGTCGATATCAGTAAC |
| | | Spec_rev TTCAGCCCTAGTTTGCAGTAAC |
| | **Amplification of Cm cassette** | CM_for GGCACCTATCTCAGCGATC |
| | | CM_rev TCCGATAATTGATGTTTCC |
| qRT-PCR | **Amplification of SSU0374** | SSU0374_for_qRT CCAAGTGATGCAAGGCTATTTC |
| | | SSU0374_rev_qRT TGACTGTCGCCGACATCTTTC |
| | **Amplification of SSU0375** | SSU0375_for_qRT AAGCCATTTCAGCGAATTCAG |
| | | SSU0375_rev_qRT TGGGCTAATCGACTGCTCTC |
| | **Amplification of SSU0376** | SSU0376_for_qRT CTCCCCAGTATGCGCTTGTG |
| | | SSU0376_rev_qRT AGTGGTGACCATGAGCAGA |
| | **Amplification of gyrB** | gyrB_for GAGCCCTGGGTCGCTTACAG |
| | | gyrB_rev AGTGGTGACCATGAGCAGA |
| Parameter                | Value                                           |
|--------------------------|-------------------------------------------------|
| n_rays                   | 32                                              |
| n_channel_in             | 1                                               |
| unet_n_depth             | 4                                               |
| unet_kernel_size         | [3, 3]                                          |
| unet_n_filter_base       | 32                                              |
| net_conv_after_unet      | 128                                             |
| net_input_shape          | [None, None, 1]                                 |
| train_shape_completion   | False                                           |
| train_completion_crop    | 32                                              |
| train_patch_size         | [256, 256]                                     |
| train_dist_loss          | mae                                             |
| train_epochs             | 400                                             |
| train_steps_per_epoch    | 100                                             |
| train_learning_rate      | 0.0003                                          |
| train_batch_size         | 16                                              |
| train_tensorboard        | true                                            |
| train_checkpoint         | weights_best.h5                                 |
| train_reduce_lr          | {'factor': 0.5, 'patience': 40, 'min_delta': 0} |