Role of the Sortase A in the Release of Cell-Wall Proteinase PrtS in the Growth Medium of Streptococcus thermophilus 4F44

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1. Introduction

The lactic acid bacterium Streptococcus thermophilus is widely used in the manufacturing of domestic and industrial fermented dairy products [1], and has obtained the “Qualified Presumption of Safety” and “Generally Recognized as Safe” (GRAS) designations. It belongs to the Streptococcus genus which is mainly pathogenic or commensal [2,3]. S. thermophilus is auxotrophic for certain amino acids and needs to get them from its environment [4,5], whereas milk, the only known habitat of this bacterium, mainly contains proteins and very few immediately assimilable peptides and amino acids. Hence, S. thermophilus needs them for its growth to reach a high cell density of a functional surface [6].

Abstract: Growth of the lactic acid bacterium Streptococcus thermophilus in milk depends on its capacity to hydrolyze proteins of this medium through its surface proteolytic activity. Thus, strains exhibiting the cell envelope proteinase (CEP) PrtS are able to grow in milk at high cellular density. Due to its LPNTG motif, which is possibly the substrate of the sortase A (SrtA), PrtS is anchored to the cell wall in most S. thermophilus strains. Conversely, a soluble extracellular PrtS activity has been reported in the strain 4F44. It corresponds, in fact, to a certain proportion of PrtS that is not anchored to the cell wall but rather is released in the growth medium. The main difference between PrtS of strain 4F44 (PrtS4F44) and other PrtS concerns the absence of a 32-residue imperfect duplication in the prodomain of the CEP, postulated as being required for the maturation and correct subsequent anchoring of PrtS. In fact, both mature (without the prodomain at the N-terminal extremity) and immature (with the prodomain) forms are found in the soluble PrtS4F44 form along with an intact LPNTG at their C-terminal extremity. Investigations we present in this work show that (i) the imperfect duplication is not implied in PrtS maturation; (ii) the maturase PrtM is irrelevant in PrtS maturation which is probably automaturated; and (iii) SrtA allows for the PrtS anchoring in S. thermophilus but the SrtA of strain 4F44 (SrtA4F44) displays an altered activity.

Keywords: Streptococcus thermophilus; sortase A; cell envelope proteinase; LPNTG motif; cell wall-anchoring
cytoplasmic peptidases [8,9]. The **prtS** gene has been probably acquired from a bacterium close to *Streptococcus suis*, displaying the CEP SpA, which is very similar to PrtS (95% identities) [10,11].

PrtS is a subtilisin-like serine protease [12], synthetized and secreted as a pre-proproteinase. Its maturation remains unknown but could be achieved by peptidyl-prolyl isomerases [13]; as for the CEP PrtP and SpeB of *Lactococcus lactis* and *S. pyogenes*, they are maturated by PrtM and PrsA (homologous of PrtM), respectively [14,15]. PrtS is also usually anchored to the cell wall through its LPNTG motif located at its C-terminal extremity. This anchoring is thought to be achieved by the sortase A (SrtA), as the other surface proteins possess the LPXTG motif in Gram-positive bacteria. Sortases are ubiquitous in Gram-positive bacteria. Six classes (A to F) are distinguished, the most well-known being class A with SrtA of *Staphylococcus aureus* (for recent review, see [16]. In pathogen bacteria, sortases are then responsible for the anchoring of surface proteins involved in biofilm formation and/or their pathogenicity, and strategies of the inhibition of their activity are currently being developed [17,18].

After their synthesis as pre-proteins, the Sec-secretion machinery supports the LPXTG surface proteins of Gram-positive bacteria to export them through the cytoplasmic membrane. A signal peptidase cleaves the N-terminal signal peptide and proteins interact with the membrane by their C-terminal extremity. Then, the transpeptidase SrtA, thought to be located at the pole/septum with the Sec secretion system, anchors these proteins to the wall through LPXTG recognition [19,20]. Finally, it must be noted that (i) SrtA contains a non-cleaved signal peptide at its N-terminal extremity for both its exporting and holding onto the plasmic membrane, and (ii) the catalytic domain (triad Cys208, His142, and Arg216) is located in its C-terminal extremity.

Peptidyl-prolyl isomerases (PPIases, foldases, or maturases) maturate certain LPXTG-proteins, such as CEP [13], by catalyzing the cis-trans isomerization of a peptide bond located upstream a prolyl residue of the prodomain. In the case of the CEP SpeB of *S. pyogenes*, such a reaction has been postulated as essential for the stabilization of the pre-protein in a sec-pathway-dependent secretion-competent conformation; for its correct exportation; and for subsequent anchoring [21,22].

Chang et al. [23] detected two forms of PrtS in the strain 4F44: one anchored to the wall and an extracellular soluble one. Both forms are devoid of the 32 amino acids’ imperfect duplication in the prodomain PP, which is thought to be required for PrtS maturation [11], and the soluble form displays an intact LPNTG motif at its C-terminal extremity [23]. Two hypotheses were then proposed. The first implies that because of the absence of the 32 amino acids imperfect duplication in the PP domain, maturation of PrtS is incorrect, leading to an incorrect folding of the CEP and then to its imperfect anchoring, similarly to the SpeB of *S. pyogenes*. The second hypothesis relies on a partial deficiency of the anchoring activity of SrtA of strain 4F44 (SrtA4F44) in spite of only six substitutions between SrtA4F44 and SrtA_{LMD-9}, (sortase A of the strain LMD-9). Indeed, one of the six, the Ile218 substitution, could be important for the recognition of the LPNTG motif [24]. Therefore, this study aims to address the cause of PrtS_{4F44} release in the extracellular medium.

### 2. Materials and Methods
#### 2.1. Bacterial Strains and Culture Conditions

The strains and plasmids used in this study are presented in the Table 1. Strains were stored in reconstituted skim milk 10% (m/v) at −80 °C. They were precultured in reconstituted skim milk and then introduced at 1% into M17 medium with lactose 20 g L⁻¹ (LM17) [25]. For the transformation experiments, the strains were first cultivated in LM17 before being inoculated in a chemically defined medium with 20 g L⁻¹ of lactose [26]. The incubation temperature was 42 °C. For the LMD-9_{SrtA}, LMD-9_{PrtS}, and LMD-9_{PrtM} mutants, 5 µg mL⁻¹ of erythromycin was added to the LM17 medium, while for the mutants LMD-9_{srtALMD-9}, LMD-9_{srtA4F44}, LMD-9_{prtS4F44}, and LMD-9_{srtA1le218→val218}, the medium was supplemented with 20 µg mL⁻¹ of streptomycin and 300 µg mL⁻¹ of spectinomycin.
The growth of the strains was assessed for measuring the pH of the medium and the optical density (OD) at 600 nm.

2.2. DNA Extraction, PCR Amplification, Electrophoresis, and Sequencing Conditions

Plasmid DNAs were isolated from *Escherichia coli* using the Miniprep Kit (Fermentas, Villebon-sur-Yvette, France) according to the manufacturer’s instructions. Genomic DNAs were extracted as previously described [27]. Primer3plus software was used to design primers, which were then synthesized by Eurogentec (Seraing, Belgium). Sequences of primers and the sizes of amplicons are reported in Tables S1–S3 (Supplementary Data). Polymerase chain reactions (PCRs) were achieved according to the supplier’s recommendations (Fermentas, Villebon-sur-Yvette, France) in a Mastercycler proS thermocycler (Eppendorf, Hamburg, Germany). Cycle conditions were: 95 °C for 5 min, 35 cycles of 3 steps (95 °C for 30 s; hybridization at appropriate temperatures (Tables S1–S3); 72 °C for 1 min kb⁻¹), and 10 min at 72 °C. For mutant construction, high fidelity Phusion DNA polymerase (Fermentas, Saint Rémy-lès-Chevreuse, France) was used for the amplification of each fragment (final DNA concentration of 5 µg mL⁻¹; extension time 30 s kb⁻¹). For overlapping (OL) PCR, DNA fragments required for the constructions were pooled in equal amount (final concentration of 5 µg mL⁻¹). The mixture also contained 0.5 µmol mL⁻¹ of each primer (forward complementary with 5’ end of the first fragment and reverse complementary with 3’ end of the last fragment); high fidelity Phusion DNA polymerase 4U, each dNTP of 0.2 µmol mL⁻¹, and 5X Phusion HF buffer of 4 µL. The program used was: 2 min at 95 °C, 35 cycles of 3 steps (95 °C 30 s, hybridization for 30 s at the annealing temperatures, and 72 °C 30 s kb⁻¹), and finally 72 °C for 10 min.

The high pure PCR product purification kit (Roche Applied Science, Meylan, France) was used to purify PCR products, taking as an eluent the elution buffer of the kit (for OL PCRs) or ultra-pure water (for sequencing reactions).

PCR products were separated by electrophoresis on 1% (w/v) agarose gel in 0.5× TAE buffer [28] at 100 V. The molecular weight markers used were 1 kb and 100 bp DNA ladders (Fermentas). Sequencing was performed by Beckman Coulter Genomics (Essex, UK) with the Sanger method [29].

2.3. Mutant Constructions and Natural Transformation

The *srtA* (locus: STER_RS06195; NCBI reference sequence: NC_008532.1) and *prtM* (locus: STER_RS02415; NCBI reference sequence NC_008532.1) genes of the *S. thermophilus* wild-type (WT) strain LMD-9 have been replaced by a cassette carrying the erythromycin resistance gene *ery*, leading to LMD-9∆*srtA* and LMD-9∆*prtM* mutants using the same strategy. Thus, the upstream (*prtM*-UpX or *srtA*-UpX) and downstream (*prtM*-DownX or *srtA*-DownX) fragments of gene *prtM* or *srtA* were amplified by PCR, as well as by the *ery* gene, which was located on plasmid pG+host 9. The three overlapped PCR amplicons (*prtM*-UpX-*ery*-prtM-DownX, or *srtA*-UpX-*ery*-srtA-DownX) obtained were used to create the recombinant fragment by overlapping PCR, which was then introduced into natural competent cells of *S. thermophilus* LMD-9, allowing for obtainment after its integration through a double crossing-over event, specifically either the LMD-9∆*prtM* mutant or LMD-9∆*srtA* mutant. Primers sequences used to amplify the different fragments are listed in Table S1 (Supplementary data). The *ery* gene of mutant LMD-9∆*srtA* had been replaced by the allele *srtA*<sub>4F44</sub> (taxon: 1308; GenBank: GU459010.1) or *srtA*<sub>LMD-9</sub>, whereas the *ery* gene of mutant LMD-9∆*prtM* had been replaced by the allele *prtS*<sub>4F44</sub>, leading to mutants LMD-9∆*srtA*<sub>4F44</sub>, LMD-9∆*srt*A<sub>LMD-9</sub>, and LMD-9∆*prtS*<sub>4F44</sub>. For that, the upstream (UpX-*srtA* or UpX-*prtS*) or downstream (DownX-*srtA* or DownX-*prtS*) fragments of gene *srtA* or *prtS*, as well as the *srtA*<sub>LMD-9</sub> gene, were amplified from the genomic DNA of the LMD-9 strain. The genes *srtA*<sub>4F44</sub> and *prtS*<sub>4F44</sub> were amplified from the genomic DNA of the 4F44 strain and the *spec* gene (conferring resistance to spectinomycin) was amplified from plasmid pSET4S. According to the mutant anticipated, four appropriate overlapped fragments (UpX-*srtA*, *srtA*<sub>4F44</sub>/srt*A<sub>LMD-9</sub>, *spec*, DownX-*srtA* or UpX-*prtS*, *prtS*<sub>4F44</sub>, *spec*, DownX-*prtS*) were used
to create suitable recombinant fragments, which were then introduced into competent cells of mutant LMD-9<sup>ΔsrtA</sup> or LMD-9<sup>ΔprtS</sup> to produce, after a double crossing-over event, the mutants LMD-9<sup>srtA4F44</sup>, LMD-9<sup>srtALMD-9</sup>, and LMD-9<sup>Δsrt4F44</sup>. Primer pairs used for the different amplifications are also indicated in Table S1 (Supplementary data).

### Table 1. Characteristics of Streptococcus thermophilus strains and plasmids used in this study and PrtS activity.

| Strains/Plasmids | Origin | References | srtA Allele | Presence (+)Absentence (-) of the prtS Gene | PrtS Activity in mAU Cell Surface | Growth Supernatant |
|------------------|--------|------------|-------------|---------------------------------------------|---------------------------------|-------------------|
| LMD-9            | Yogurt | [30]       | srtA<sub>AMM-9</sub> | +                                            | 3110                           | 0                 |
| 4F44             | Cheese | [23]       | srtA<sub>4F44</sub> | +                                            | 5620                           | 3240              |
| LMD-9<sup>ΔsrtA</sup> |       |            |             |                                               |                                |                   |
| LMD-9<sup>ΔprtS</sup> |       |            |             |                                               |                                |                   |
| LMD-9<sup>srtA</sup> |       |            |             |                                               | 3100                           | 0                 |
| LMD-9<sup>srtM</sup> |       |            |             |                                               | 2600                           | 1020              |
| LMD-9<sup>srtM4F44</sup> |       |            |             |                                               | 3124                           | 1070              |
| LMD-9<sup>srtM4F44</sup> |       |            |             |                                               | 0                              | 0                 |
| LMD-9<sup>srtM4F44</sup> |       |            |             |                                               | 3093                           | 0                 |
| LMD-9<sup>srtM4F44</sup> |       |            |             |                                               | 3114                           | 0                 |
| LMD-9<sup>srtM4F44</sup> |       |            |             |                                               | Present                        | 0                 |

| Plasmids | Relevant markers and characteristics | srtA<sup>Ile218</sup>, LMD-9, LMD-9<sup>srtA</sup>, and LMD-9<sup>ΔprtS</sup> were prepared as described previously [26,31] to introduce, by natural transformation, the recombinant DNA fragments obtained by OL PCRs. |
|------------------|---------------------------------------------|---------------------------------------------|---------------------------------------------|---------------------------------------------|---------------------------------------------|---------------------------------------------|
| pG+host9         | Erm<sup>+</sup>, pWV01 derivative, with thermoresistant replication function |                                                |                                            |                                            |                                            |                                            |
| pSET4s           | Spec<sup>+</sup>, replication function of Pg + host3 and pUC19          |                                                |                                            |                                            |                                            |                                            |

The strategy of the construction of mutant LMD-9<sup>srtA</sup>Val<sup>218</sup> is presented in Figure A1 (Appendix A). Competent cells of LMD-9, LMD-9<sup>ΔsrtA</sup>, and LMD-9<sup>ΔprtS</sup> were cultured in 200 mL of LM17 until OD<sub>600nm</sub> = 1. Supernatants of centrifuged (3900 g–10 min) cultures were filtered and then stored at 4 °C. Bacterial pellets were washed in Tris-HCl buffer (100 mmol L<sup>−1</sup>, pH 7), then resuspended in 20 mL of the same buffer and kept at 4 °C. The free (supernatant) or bound (pellet) PrtS activity was revealed using the Suc-Ala-Ala-Pro-Phe-pNA substrate (Sigma, St. Louis, MO, USA) [23]. PrtS activity was expressed in arbitrary units, each measure being realized at least in triplicates. Activity values greater than one can be compared since they were established during the same assay and therefore in the same experimental condition for the strains LMD-9<sup>ΔprtS4F44</sup> and LMD-9<sup>ΔsrtF44</sup>, LMD-9<sup>ΔsrtM1</sup>, LMD-9<sup>ΔsrtA</sup>, LMD-9<sup>ΔsrtALMD-9</sup>, and LMD-9<sup>Δsrt4F44</sup> and LMD-9<sup>Δsrt4F44</sup> Activity values greater than one can be compared since they were established during the same assay and therefore in the same experimental condition for the strains LMD-9<sup>ΔprtS4F44</sup> and LMD-9<sup>ΔsrtF44</sup>, LMD-9<sup>ΔsrtM1</sup>, LMD-9<sup>ΔsrtA</sup>, LMD-9<sup>ΔsrtALMD-9</sup>, and LMD-9<sup>Δsrt4F44</sup> and LMD-9<sup>Δsrt4F44</sup> (Table 1) The free activity of PrtS was also demonstrated by zymography with caseins as substrates [7].

2.4. Detection of Extracellular Proteinase Activity

WT strains LMD-9 and 4F44, and mutants LMD-9<sup>ΔsrtA</sup>, LMD-9<sup>Δsrt</sup>ALMD-9, LMD-9<sup>ΔsrtA4F44</sup>, and LMD-9<sup>ΔsrtAIle218 → Val218</sup> were cultured in 200 mL of LM17 until OD<sub>600 nm</sub> = 1. Supernatants of centrifuged (3900 g–10 min) cultures were filtered and then stored at 4 °C. Bacterial pellets were washed in Tris-HCl buffer (100 mmol L<sup>−1</sup>, pH 7), then resuspended in 20 mL of the same buffer and kept at 4 °C. The free (supernatant) or bound (pellet) PrtS activity was revealed using the Suc-Ala-Ala-Pro-Phe-pNA substrate (Sigma, St. Louis, MO, USA) [23]. PrtS activity was expressed in arbitrary units, each measure being realized at least in triplicates. Activity values greater than one can be compared since they were established during the same assay and therefore in the same experimental condition for the strains LMD-9<sup>ΔprtS4F44</sup> and LMD-9<sup>ΔsrtF44</sup>, LMD-9<sup>ΔsrtM1</sup>, LMD-9<sup>ΔsrtA</sup>, LMD-9<sup>ΔsrtALMD-9</sup>, and LMD-9<sup>Δsrt4F44</sup> and LMD-9<sup>Δsrt4F44</sup> (Table 1) The free activity of PrtS was also demonstrated by zymography with caseins as substrates [7].

Molecular modeling simulations were run on a bi-processor AMD Dual Core 280 with 2.4 GHz. Docking and scoring simulations were performed using the LibDock algorithm [34] and the Consensus Score modules of the program-package Discovery Studio version 3.5 (Accelrys, Inc., San Diego, CA, USA), respectively. All molecular mechanics calculations were performed with the CHARMM force field [35]. The Protein Data Bank entry 3FN5 corresponding to the SrTa (Spy1154) of the S. pyogenes serotype M1 strain SF370 was used as the input structure [24]. This was made of two chains per unit cell (length: a—39.8 Å, b—59.46 Å, and c—65.11 Å; angles: α—90°, β—101.96°, and γ—90°). These two chains contained 187 residues corresponding to the catalytic domain. The first 18 residues corresponded to the His-tag region expediting the purification of the protein. The chain A included a 4-(2-Hydroxyethyl)-1-Piperazine Ethanesulfonic Acid (HEPES) entity as an inhibitor. The chain B was picked as the input structure for all simulations.
In our study, the first 18 residues due to the crystallization step were withdrawn from chain B. The resulting structure Sp-SrtA<sub>SF370Δ86</sub> was used as a template to design homology models for St-SrtA<sub>4F44Δ90</sub> and St-SrtA<sub>LMD-9Δ90</sub>. In order to investigate the proteinase PrtS “sorting pattern” binding modes within the SrtA active site, two ligands were designed, namely LPNTG and Ace-QLPNTGEND-NMe. Their possible binding modes within the active site of the SrtA systems (Sp-SrtA<sub>SF370Δ86</sub>, St-SrtA<sub>4F44Δ90</sub>, and St-SrtA<sub>LMD-9Δ90</sub>) were studied through docking simulations using the LibDock algorithm. This method allows the identification of low-energy binding modes of ligands based on polar and apolar interactions sites (hotspots). The SrtA hotspots were defined as the catalytic site which encloses the catalytic residues. To prevent potential interactions, water molecules were removed from the cavity. The structure of the ligand complies following polar and apolar interaction sites of the receptor. Using this docking procedure guarantees that only the highest scoring poses (30 to 100 poses) are kept. The Libdock scoring functions based on simple pair-wise score calculations were used in all simulations. Finally, poses were assessed considering the position and orientation of the sorting pattern within the catalytic cavity and its proximity to the catalytic triad.

3. Results and Discussion

3.1. The Imperfect Duplication of 32 Amino Acid Residues in the Prodomain of PrtS<sub>LMD-9</sub> Is Not Involved in the Anchoring of PrtS to the Cell Wall of S. thermophilus

Delorme et al. [11] postulated that the imperfect duplication of 32 amino acid residues (residues 63 to 90) in the prodomain PP of PrtS of the strain LMD-9 (PrtS<sub>LMD-9</sub>; locus: STER_RS04165; NCBI reference sequence: NC_008532.1) is responsible for the anchoring and/or maturation of this CEP. To evaluate this, the LMD-9<sub>prtS<sub>4F44</sub></sub> mutant strain was constructed by integrating the <sub>prtS</sub> allele of the 4F44 strain (prtS<sub>4F44</sub> (taxon: 1308; GenBank: GU459009.1)), encoding a CEP devoid of such duplication, in the LMD-9<sub>ΔprtS</sub> mutant strain [31]. The proteolytic activity was evaluated at the cells’ surface and in the growth supernatant of this mutant as well as the mutant LMD-9<sub>ΔprtS</sub> served as a negative control. Followed by the growth of both strains in LM17 to an OD<sub>600nm</sub> of 1, cells were harvested by centrifugation and both the filtered supernatants and the cells were subsequently incubated with the synthetic substrate Suc-Ala-Ala-Pro-Phe-pNA. As anticipated, a strong proteolytic activity was noticed at the surface of the cells of LMD-9<sub>prtS<sub>4F44</sub></sub> but not in the filtered supernatant, whereas no such activity was reported for the negative control (Table 1). Hence, it is concluded that any mutation in the sequence of PrtS<sub>4F44</sub>, particularly the absence of the imperfect duplication of 32 amino acid residues in its prodomain PP, is not responsible for the liberation of PrtS<sub>4F44</sub> into the extracellular medium of strain 4F44.

3.2. Role of Maturases in the Maturation of PrtS in S. thermophilus LMD-9

An exploration of data banks allowed for identification of four genes that encode PPIases (prtM, tig or ropA, ppiA, and pplB) in sequenced genomes of S. thermophilus. Among them, two maturases, PplB and PpiA, belong to the cyclophilin family, whereas PrtM and RopA belong to the parvulin and FKBP (FK506 binding protein) families, respectively. An analysis of these four genes in the 4F44 strain revealed that their deduced proteins are identical with the corresponding ones of the LMD-9 strain, except the PrtM protein (taxon: 1308; locus: AMM43147; GenBank: KT809299.1), which displays four differences (D30N, V75A, A78V, and A242T). Hence, to determine its implication in PrtS maturation, the LMD-9<sub>ΔprtM</sub> mutant strain was constructed by deleting the <sub>prtM</sub> gene in the LMD-9 strain. The LMD-9, LMD-9<sub>ΔprtS</sub>, and LMD-9<sub>ΔprtM</sub> strains were grown in milk to evaluate the effect of <sub>prtM</sub> deletion on the growth performance of the mutant. This approach is an indirect measure of the PrtS proteolytic activity considering the relationship between the PrtS activity and S. thermophilus capacity to grow in milk [6,7]. The results revealed that the LMD-9<sub>ΔprtS</sub> mutant showed a delayed growth both in LM17 and milk media contrary to strains LMD-9 and LMD-9<sub>ΔprtM</sub>, which showed a similar growth behavior (Figure 1). In a
similar manner, the proteolytic activity was observed only at the cells’ surface of LMD-9 and LMD-9\textsuperscript{ΔprtM} (Table 1), and not in the culture supernatants.

![Graph showing growth of S. thermophilus LMD-9, LMD-9\textsuperscript{ΔprtM}, and LMD-9\textsuperscript{ΔprtS} in milk (A) or in LM17 (B,C), evaluated by measuring either the extracellular pH (A,B) or OD\textsubscript{600} (C).](image)

Figure 1. Growth of S. thermophilus LMD-9, LMD-9\textsuperscript{ΔprtM}, and LMD-9\textsuperscript{ΔprtS} (a negative control) in milk (A) or in LM17 (B,C), evaluated by measuring either the extracellular pH (A,B) or OD\textsubscript{600} (C).

Consequently, the lack of the putative PrtM maturase in the LMD-9\textsuperscript{ΔprtM} strain did not result in any growth retardation, nor in a decrease in PrtS activity and/or its release into the extracellular medium; it is most probable that the PrtM maturase was not responsible for the PrtS maturation. Several factors support the hypothesis that PrtS could undergo automaturation. First, unlike the \textit{prsa} of \textit{S. pyogenes} and \textit{prtM} of \textit{L. lactis}, which are located upstream of the \textit{speB} and \textit{prtP} genes, respectively and co-transcribed, the \textit{prtM} of \textit{S. thermophilus} is not located near PrtS [9,12,14,36]. Second, when Chang et al. [23] established the N-terminal sequence of the soluble PrtS form of the 4F44 strain, they detected the N-terminal sequence corresponding to the proenzyme (with prodomain PP) as well as the N-terminal sequence of the mature form (without prodomain PP) of PrtS. The same
situation was also detected for anchored PrtS forms present at the cells’ surface of the LMD-9 strain [31]. It was also noticed that the PrtS proenzyme form disappeared progressively in favor of the mature form (unpublished results) preceding a prolonged incubation or high concentration preservation. The fact that PrtS could be automaturated is not inimitable in LAB, e.g., regarding CEP PrtB of *Lb. bulgaricus* [37]. Finally, the fact that the inactivation of *prtM* did not lead to any release of PrtS in the growth medium, associated with the absence of difference between the three other known maturases (PpiA, PplB, and RopA), ruled out the hypothesis of a link between maturation via PrtM especially, correct folding, and anchoring.

3.3. *SrtA* Is Responsible for the Anchoring of PrtS to the Cell Wall of *S. thermophilus* and Is Deficient in Strain 4F44

The hypothesis that, despite few numbers of substitutions found between *SrtA*4F44 and *SrtA*LMD-9, the extracellular liberation of PrtS4F44 could result from a partial deficiency of *SrtA*4F44 implies first providing evidence that *SrtA* is actually responsible for the PrtS anchoring to the cell wall of *S. thermophilus* as in other Gram-positive bacteria. Hence, the LMD-9*ΔsrtA* mutant strain was constructed by replacing the *srtA* gene by an erythromycin resistance gene. Afterwards, a complemented mutant was constructed by reintegrating the *srtA*LMD-9 *ene* to the LMD-9*ΔsrtA* mutant to verify whether the phenotype of wild type LMD-9 strain could be restored (LMD-9*ΔsrtA*LMD-9).

The synthetic substrate Suc-Ala-Ala-Pro-Phe-pNA was used to search proteolytic activity in the filtered growth medium of the LMD-9*ΔsrtA* mutant, complemented mutant, and WT strain (Table 1). It was detected only in the LMD-9*ΔsrtA* mutant (1020 mAU of PrtS activity; Table 1). The same supernatants were then analyzed by a casein-zymogram to detect PrtS (Figure 2A). Three caseinolytic bands (170, 154, and 115 kDa) previously shown to correspond to PrtS [23] were observed (Figure 2A) in the supernatant of mutant LMD-9*ΔsrtA* contrary to WT and LMD-9*ΔsrtA*LMD-9. This corroborates the results obtained using synthetic substrate Suc-Ala-Ala-Pro-Phe-pNA. Unexpectedly, the LMD-9*ΔsrtA* strain displayed a surface PrtS activity (Table 1) despite of *srtA* deletion. To determine whether this resulted from electrostatic or other low-force interactions of PrtS with the cell surface, cells of this mutant and of strains 4F44 and LMD-9*ΔsrtA*LMD-9, used as controls, were suspended in Tris HCl (100 mmol/L, pH 7) buffer and incubated from 0 to 70 h at 4 °C. The cells’ surface-bound PrtS activity was determined at t0 and t70 using a casein-zymogram (Figure 2) and the synthetic substrate Suc-Ala-Ala-Pro-Phe-pNA. After 70 h of incubation, the surface PrtS activity of strains 4F44, LMD-9, and LMD-9*ΔsrtA*LMD-9 was close to that obtained at t0, while that of the LMD-9*ΔsrtA* strain appeared to be undetectable. The PrtS activity was exclusively found in the incubation buffer of this strain (Figure 2C).

To further investigate whether *SrtA*4F44 is partially defective, the LMD-9*ΔsrtA* strain was constructed by introducing the *srtA*4F44 allele into the genome of the LMD-9*ΔsrtA* strain. Hereafter, the soluble PrtS activity was examined in filtered growth supernatants of the strains LMD-9*ΔsrtA*4F44 and 4F44 by using both the Suc-Ala-Ala-Pro-Phe-pNA substrate (Table 1) and a casein-zymogram (Figure 2B). Through analyses being realized in the conditions, levels of PrtS activity could be compared (see Section 2). Twenty-six percentages of total PrtS activity of the mutant LMD-9*ΔsrtA*4F44 were found in its growth supernatant, i.e., a proportion similar to that of strain 4F44 (37%). In addition, the incubation of the cells of this mutant in Tris-HCl (100 mmol L−1, pH 7) buffer during 70 h did not result in a significant increase of extracellular PrtS activity. Therefore, despite the few numbers...
of substitutions found between the SrtA of strains LMD-9 and 4F44, our results provide genetic proof of the dysfunctioning of SrtA<sub>4F44</sub>, which leads to the anchoring to the cell wall of the majority of PrtS molecules and to the release of a fraction of PrtS in the growth medium of the 4F44 strain, as postulated by Chang et al. [23]. Indeed, the fact that after 70 h of incubation of the cells of 4F44 and LMD-9<sub>∆srtA4F44</sub> strains in Tris-HCl buffer the surface PrtS activity remained similar to the initial one strongly suggests that it corresponds to PrtS molecules correctly anchored to the cell wall. Besides, even if the total PrtS activity (bound plus free) appeared in our assays to be higher in strain 4F44 than in strain LMD-9 (8860 mAU against 4194, Table 1), the PrtS release cannot be attributed to a higher expression of its gene in strain 4F44, leading to the saturation of SrtA activity and ultimately to a leakage of non-anchored PrtS molecules in the external environment. Indeed, in the mutant LMD-9<sub>∆srtA4F44</sub>, the gene <i>prtS</i> undergoes the same regulation like in the wild-type LMD-9 strain, as suggested by the PrtS activity levels observed in this mutant (Table 1). Therefore, no saturation of the anchoring activity of sortase SrtA is expected and SrtA<sub>4F44</sub>, which is expressed in this mutant, should anchor all PrtS molecules.

![Image of SDS-PAGE zymogram analysis](image)

**Figure 2.** SDS-PAGE zymogram analysis to detect proteolytic activity in the extracellular growth medium of <i>S. thermophilus</i> LMD-9, LMD-9<sub>∆srtA</sub>, LMD-9<sub>∆srtALMD-9</sub> (A), and 4F44, LMD-9<sub>∆srtA4F44</sub> (B), and the proteolytic activity liberated in the incubation buffer (t = 0 and t = 70 h) of the LMD-9<sub>∆srtA</sub> strain (C).

### 3.4. Substitution of the Ile 218 Residue Is Not Responsible for the Deficiency of SrtA<sub>4F44</sub>

As we knew the LPNTG motif was present at the C-terminal extremity of the extracellular soluble form of PrtS<sub>4F44</sub> [23], molecular modeling simulations were performed to determine whether the binding mechanism of the LPNTG motif to the SrtA catalytic site was at least partially altered in SrtA<sub>4F44</sub>. Since <i>S. pyogenes</i> and <i>S. thermophilus</i> belong to the same genus, their respective sortases A were assumed to have similar mechanisms. Thus, the structural models of SrtA<sub>LMD-9</sub> (St-SrtA<sub>LMD-9A90</sub>) and SrtA<sub>4F44</sub> (St-SrtA<sub>4F44A90</sub>) were built from the already defined structure of <i>S. pyogenes</i> SF370 SrtA (Sp-SrtA<sub>SF370A86</sub>) [24]. In order to build the models with the same amino acid residues as in the Sp-SrtA<sub>SF370A86</sub>, it was necessary to delete the first 90 residues of St-SrtA<sub>LMD-9A90</sub> and SrtA<sub>4F44</sub> (St-SrtA<sub>4F44A90</sub>). The percentage of identity/similarity...
between the C-terminal domain of Sp-SrtA\textsubscript{SF370\textgreek{A}86} and St-SrtA\textsubscript{4F44\textgreek{A}90}, and Sp-SrtA\textsubscript{SF370\textgreek{A}86} and St-SrtA\textsubscript{LMD-9\textgreek{A}90} were found to be sufficient (between 70.6% to 97.5% and 87.1% to 99.4%, respectively) to use Sp-SrtA\textsubscript{SF370\textgreek{A}86} as a structural pattern for constructing the St-SrtA\textsubscript{4F44\textgreek{A}90} and St-SrtA\textsubscript{LMD-9\textgreek{A}90} ones. Hence, surimposition of the three structures led to RMSD values below 0.16 Å considering all the 169 C\textalpha{} atoms of the residues 87 to 249. No significant structural difference was observed between the structures, as St-SrtA\textsubscript{4F44\textgreek{A}90} and St-SrtA\textsubscript{LMD-9\textgreek{A}90} models displayed the characteristic structure of sortases, i.e., the eight-stranded β-barrel fold and a long hydrophobic cleft corresponding to the catalytic cavity located at the center of the protein (Figure 3). It was assumed that the residues Cys\textsubscript{208}, His\textsubscript{142}, and Arg\textsubscript{216} compose the catalytic triad (Figure 3). The orientation of these residues is consistent with the model of reverse protonation that has been proposed in biochemical studies of the sortase A of \textit{St. aureus} and \textit{S. pyogenes} [24,40].

These homology models were then used to study possible binding modes of the LPNTG pattern within the sortase catalytic cavity. Enzyme/substrate complexes were generated using docking simulations with two substrates: the LPNTG pattern and a longer pattern, specifically the Ace-QLPNTGEND-Nme pattern. Unfortunately, no significant difference was observed between the complexes Sp-SrtA\textsubscript{SF370\textgreek{A}86}; LPNTG/Ace-QLPNTGEND-Nme; St-SrtA\textsubscript{LMD-9\textgreek{A}90}; LPNTG/Ace-QLPNTGEND-Nme; and St-SrtA\textsubscript{4F44\textgreek{A}90}; LPNTG/Ace-QLPNTGEND-Nme (Supplementary Data and Figures A2 and A3 in Appendix A). Consequently, these results strongly suggested that none of the four substitutions distinguishing St-SrtA\textsubscript{4F44\textgreek{A}90} from St-SrtA\textsubscript{LMD-9\textgreek{A}90} could account for the defective activity of the SrtA\textsubscript{4F44-}

\textbf{Figure 3.} Putative structure of St-SrtA4F44\textgreek{A}90. The catalytic triad (in CPK, His\textsubscript{142} in green, Arg\textsubscript{216} in blue, and Cys\textsubscript{208} in red, and its thiolate group in brown) position and the amino acid residues important (according to Race et al. [24]) for the positioning of the LPNTG motif within the cavity (in CPK teal) are indicated. The LPNTG motif is also shown within the cavity of SrtA\textsubscript{SF370\textgreek{A}86}. The structure of Sp-SrtA\textsubscript{SF370\textgreek{A}86} and St-SrtA\textsubscript{4F44\textgreek{A}90} are analogous to that of 4F44. The residue Val\textsubscript{218} corresponds to the residue Ile\textsubscript{218} in St-SrtA\textsubscript{LMD-9\textgreek{A}90}, while the residues Val\textsubscript{194} and Val\textsubscript{218} correspond to residues Ile\textsubscript{214} and Ile\textsubscript{218} in Sp-SrtA\textsubscript{SF370\textgreek{A}86}. For simplicity, the catalytic cavity has been divided into cavity 1 and cavity 2.
To definitively rule out the hypothesis of an eventual role of the substitution of Ile_{218} by Val_{218} in the deficiency of StrA_{4F44} and to validate the modeling prediction, the residue Ile_{218} of strain LMD-9, SrtA_{LMD-9} was replaced by a valine residue (Figure A1). The absence of extracellular PrtS activity (searched using the Suc-Ala-Ala-Pro-Phe-pNA substrate) in the supernatant of this strain confirms the non-involvement of Ile_{218} substitution by Val_{218} in the deficiency of StrA_{4F44}.

To conclude we showed that (i) the 32 amino acid residues’ imperfect duplication located in the prodomain of certain PrtS, such as PrtS_{LMD-9}, was not essential for the correct maturation and subsequent anchoring of PrtS; (ii) the maturase PrtM, homologous to maturases of other CEP, was not responsible for the maturation of PrtS and neither for its correct anchoring to the cell surface; and (iii) SrtA was responsible for the anchoring of PrtS to the cell wall and (iv) SrtA of strain 4F44 was partially defective despite the low number of dissimilar residues (six substitutions), which differentiates it from that of the LMD-9 strain and probably through a subtle mechanism not yet elucidated perhaps because of a lack of a structural model, including the N-terminal part of SrtA.

Supplementary Materials: The following are available online at https://www.mdpi.com/article/10.3390/microorganisms9112380/s1, Table S1: Primers and resulting PCR or Overlapping PCR products used for mutant construction, Table S2: Primers used for the sequencing of PPIases genes, Table S3: Primers used for the verification of mutant sequences.

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Appendix A

**Figure A1.** Strategy to replace the ATT codon specifying the Ile\textsubscript{218} residue of SrtALMD-9 by a GTT codon corresponding to a Valine residue. Two PCRs were carried out using genomic DNA of the LMD-9\textsubscript{srtALMD-9} (LMD-9\textsubscript{srtALMD-9}-g DNA) mutant to obtain (i) an upstream DNA fragment of the \textit{srtA} gene (Up\textit{srtA}) and the first 675 nucleotides of the same gene, and (ii) a DNA fragment containing the second part of the \textit{srtA} gene (87 nucleotides), the \textit{spec} gene (confers resistance to spectinomycin), and a downstream sequence of the \textit{srtA} gene (Down\textit{srtA}) (A). These two overlapped PCR fragments were then used to make the recombinant DNA fragment by overlapping PCR (B). The recombinant DNA amplicon was introduced into competent cells of the LMD-9\textsubscript{\Delta srtA} mutant, thereby permitting obtainment after its integration through a double crossing-over event, specifically the mutant LMD-9\textsubscript{srtA:Ile218\rightarrow Val218} (C). Primer sequences used to amplify different fragments are presented in Table S1.
Figure A2. Interaction between the LPNTG ligand and amino acid residues of St-SrtA<sub>4F44Δ90</sub>. The numbers represent the distance between atoms of the ligand and residues of the cavity of sortase A. Red letters indicate residues of the catalytic triad. The residues Gln<sub>187</sub>, Glu<sub>190</sub>, Val<sub>194</sub>, Ala<sub>211</sub>, and Val<sub>218</sub> correspond to the residues Ala<sub>187</sub>, Arg<sub>190</sub>, Ile<sub>194</sub>, Ile<sub>211</sub>, and Ile<sub>218</sub> of Sp-SrtA<sub>SF370Δ86</sub>, while residue Val<sub>218</sub> corresponds to the residue Ile<sub>218</sub> of St-SrtA<sub>LMD-9</sub>.

Figure A3. Superimposition of the main binding modes of ligand Ace-QLPNTGEND-Nme within the sortase A (here SrtA<sub>4F44Δ90</sub>) catalytic cavity. Ligands denoted here correspond to the best ligand obtained for each complex (Sp-SrtA<sub>SF370Δ86</sub>:Ace-QLPNTGEND-Nme; SrtA<sub>LMD-9Δ90</sub>:Ace-QLPNTGEND-Nme; and SrtA<sub>4F44Δ90</sub>:Ace-QLPNTGEND-Nme) according to (i) the distance between the S atom of the corresponding sortase A Cys<sub>208</sub> residue and the alpha carbon of the threonine residue of the ligand, and (ii) the position of the ligand within the cavity. Thus, the brown ligand corresponds to the ligand of complex Sp-SrtA<sub>SF370Δ86</sub>:Ace-QLPNTGEND-Nme, the purple one to SrtA<sub>LMD-9Δ90</sub>:Ace-QLPNTGEND-Nme, and the blue one to SrtA<sub>4F44Δ90</sub>:Ace-QLPNTGEND-Nme.
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