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Mono-dimensional Blue Native-PAGE and bi-dimensional Blue Native/Urea-PAGE or /SDS-PAGE combined with nLC-ESI-LIT-MS/MS unveil membrane protein heteromeric and homomeric complexes in *Streptococcus thermophilus*

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Abstract

Protein interactions are essential elements for the biological machineries underlying biochemical and physiological mechanisms indispensable for microorganism life. By using mono-dimensional blue native polyacrylamide gel electrophoresis (1D-BN-PAGE), two-dimensional blue native/urea-PAGE (2D-BN/urea-PAGE) and two-dimensional blue native/SDS-PAGE (2D-BN/SDS-PAGE), membrane protein complexes of Streptococcus thermophilus were resolved and visualized. Protein complex and oligomer constituents were then identified by nLC-ESI-LIT-MS/MS. In total, 65 heteromeric and 30 homomeric complexes were observed, which were then associated with 110 non-redundant bacterial proteins. Protein machineries involved in polysaccharide biosynthesis, molecular uptake, energy metabolism, cell division, protein secretion, folding and chaperone activities were highly represented in electrophoretic profiles; a number of homomeric moonlighting proteins were also identified. Information on hypothetical proteins was also derived. Parallel genome sequencing unveiled that the genes coding for the enzymes involved in exopolysaccharide biosynthesis derive from two separate clusters, generally showing high variability between bacterial strains, which contribute to a unique, synchronized and active synthetic module. The approach reported here paves the way for a further functional characterization of these protein complexes and will facilitate future studies on their assembly and composition during various growth conditions and in different mutant backgrounds, with important consequences for biotechnological applications of this bacterium in dairy productions.

Abbreviations

1D-BN-PAGE, mono-dimensional blue native polyacrylamide gel electrophoresis; 2D-BN/SDS-PAGE, two-dimensional blue native/SDS-PAGE; 2D-BN/urea-PAGE, two-dimensional blue native/urea-PAGE; nLC-ESI-LIT-MS/MS, nano-liquid chromatography coupled with electrospray ionization-linear ion trap tandem mass spectrometry.

1. Introduction

Streptococcus thermophilus is of major importance for the food industry since it is widely used for the manufacture of dairy products; in this context, it is considered as the second most important industrial dairy starter after *Lactococcus lactis* [1,2]. Together with symbiotic *Lactobacillus delbrueckii* subsp. bulgaricus or *Lb. helveticus*, this Gram-positive (G+), lactic acid bacterium is generally used for the production at relatively high process temperatures of yogurt and so-called hard cooked cheeses (e.g., emmenthal, grana) [1,3]. In combination with other lactobacilli, it is also utilized for the manufacture of mozzarella and cheddar cheeses [1].

S. thermophilus is closely related to L. lactis but it is even more strictly related to streptococcal pathogenic species, including S. pyogenes, S. agalactiae, S. pneumoniae S. mutans S. suis and S. equi, which cause different lethal infection diseases or tooth decay [4,5]. Nevertheless, S. thermophilus is generally recognized as a safe bacterium and a multitude of live microorganism cells are ingested annually by humans. The complete genome sequence of various S. thermophilus strains (LMG18311, CNRZ1066, LMD9, JIM8232, ND03 and MNZLW002) was made publicly available (http://www.ncbi.nlm.nih.gov/genome/genomes/420) [6-11]. Its comparison with the genome of streptococcal pathogens highlighted the similitude of this lactic acid bacterium to pathogenic species [6-8], but also revealed that the most significant determinants for pathogenicity are either lacking or present as pseudogenes, except they code essential cellular functions. Comparative genomics also revealed that evolution has shaped the S. thermophilus genome mainly through loss-of-function events, even if lateral gene transfer played an important role [12], disclosing that this bacterium has followed an evolutionary path divergent to that of streptococcal pathogens as result to its adaptation to a specific and well-defined ecological niche, i.e. milk.

To investigate global gene expression changes in *S. thermophilus* during exponential and stationary phases or following adaptation to various environmental stresses, we previously characterized changes of the corresponding cytosolic and/or membrane proteomic repertoires by gel-based and shotgun approaches [13-15]. Global characterization of multi-protein complexes is also an important step to provide an integrative view of multipart polypeptide machineries that are essential for bacterial biological functions and physiology. Recent advancements in high throughput technologies has allowed a direct description of protein-protein interactions; thus, two-hybrid assay [16–23], protein chip [24-26] or co-purification [27-29] procedures have been widely used to characterize bacterial protein-protein interaction networks. Recent considerations on restricted accuracy of deriving results and its labor-intensive nature have limited the application of the first two approaches to the production of large scale protein-protein interaction datasets [30,31]. Thus,

two functional proteomics technologies based on direct MS identification of resolved protein components have been preferentially used for systematic analysis of co-purified hetero-multimeric and/or homomeric complexes following their affinity capture by tagged-protein baits [28] or their direct resolution by 1D-BN-PAGE and 2D-BN/SDS-PAGE [29]. In both cases, protein complex purification has to be performed under native conditions to prevent molecular dissociation. Protein complex affinity capture by tagged-protein baits allowed the characterization of a number of polypeptide machineries, as in the case of Escherichia coli [32-35], Mycoplasma pneumoniae [36], Staphylococcus aureus [37], Rhodopseudomonas palustris and Shewanella oneidensis [38], generating large bacterial protein interaction networks. On the other hand, 1D-BN-PAGE and 2D-BN/SDS-PAGE have found a widespread application for the analysis of bacterial complexomes [29,39,40]. Also in this case, a non-denaturing environment must be kept throughout the first dimension BN-PAGE analysis. It comprises the use of: i) neutral pH-low salt concentration buffers, no reducing/denaturing agents, manipulation at low-temperatures and mild zwitterionic detergents for sample preparation; ii) anionic Coomassie Brilliant Blue G-250 dye and Bis-Tris/imidazole during electrophoresis to impose a net negative charge on protein surfaces, thus facilitating protein complex migration, hampering solute aggregation and stabilizing native gel pH value, respectively. Each multi-protein complex may be then denatured in a second dimension electrophoresis (SDS-PAGE), and the protein alignment within the gel allows the MS-based identification of interactive proteins. Alternatively, each protein complex band from 1D-BN-PAGE can be directly analyzed for its constituents by nLC-ESI-MS/MS [41]. Both approaches were used for high-throughput characterization of: i) membrane protein complexes from Neisseria meningitides [42], Rhodobacter sphaeroides [43], Francisella tularensis [44], Clostridium thermocellum [45], Mycobacterium bovis [46] and Enterococcus faecalis [47]; ii) cytoplasmic protein complexes from Streptomyces coelicolor [48] and Pseudomonas sp. [49]; iii) membrane and cytoplasmic protein complexes from Helicobacter pylori [50,51],E. coli [52,53], Chlorobium tepidum Methanothermobacter therautotrophicus [56]. Results from tagged-protein- and BN-PAGE-based experiments allowed compiling various microbial protein interaction database, such as eNet, MPIDB, STRING, IntAct, DIP, BIOGRID and others [57-61], which now can be searched simultaneously by using the dedicated service PSICQUIC interface [62].

In this study, we report on the combined use of 1D-BN-PAGE, 2D-BN/urea-PAGE, 2D-BN/SDS-PAGE, and nLC-ESI-LIT-MS/MS for the characterization of membrane complexes from *S. thermophilus*. A number of molecular machineries, as obtained from the extraction of bacterial membranes with 0.5% *n*-dodecyl-β-D-maltoside (βDDM), were characterized, describing the

heteromultimeric or homomeric nature of the corresponding protein complexes and discussing their functional properties with respect to organism physiology.

2. Materials and Methods

2.1 Bacterial growth

S. thermophilus strain DSM20617 was grown in M17 medium supplemented with 2% lactose, at 37°C, without shaking [15]. Cells were monitored by measuring the absorbance at 600 nm, collected in their early exponential phase (pH 5.6) and then washed with sterile 20 mM Tris-HCl, pH 7.4, at 4°C. Bacterial cells were harvested by centrifugation at 14,000 x g, at 4°C, and then washed twice in PBS, pH 7.4.

2.2 Protein sample preparation

A biomass corresponding to 5 g of bacterial cells (wet weight) was suspended in extraction buffer (750 mM ε-amino caproic acid, 1 mM PMSF, 50 mM Tris-HCl, pH 7.0) and sonicated in ice with a Labsonic U sonicator, repeating duty cycles of 0.5 sec for 60 sec, for 5 times, with 15 sec intervals. Unbroken cells and cell debris were removed from resulting suspension by centrifugation at 10,000 x g, for 15 min, at 4°C. DNase I (100 µg/mL final concentration) was added to the supernatant; the sample was kept at 25°C, for 1 h, and then centrifuged at 100,000 x g, for 30 min, at 4°C. Membrane pellet was washed once in extraction buffer, at 4°C, and twice in 0.33 M sorbitol, 1 mM PMSF, 50 mM Bis-Tris-HCl, pH 7.0, at 4°C. Finally, membrane pellet was resuspended in resuspension buffer (20% v/v glycerol, 1 mM PMSF, 25 mM Bis-Tris-HCl, pH 7.0) at 4°C, and quantified using the DC protein assay (Bio-Rad Laboratories, Hercules, CA). Under continuous mixing, equal volumes of sample suspension and resuspension buffer containing 1% BDDM (Sigma-Aldrich, St. Louis, MO) were mixed. Solubilization of membrane protein complexes was allowed to occur on ice, for 3 min. Sample was then centrifuged at 100,000 x g, for 30 min, and the supernatant (containing membrane multiprotein complexes) was resolved by 1D-BN-PAGE. A schematic representation of the most important experimental steps used in this study are reported in Supplementary Figure S1.

2.3 Electrophoresis

1D-BN-PAGE was carried out as described by Schagger and von Jagow [63], with some modifications. Different acrylamide gradients were tested to improve protein complex separation;

thus, 4-14% and 7-14% linear gradients were used for final experiments. Anode buffer contained 50 mM Bis-Tris-HCl, pH 7, while cathode buffer was 15 mM Bis-Tris, 50 mM tricine, supplemented with 0.01% Coomassie Blue Brilliant G (Sigma). Before electrophoresis, samples were mixed with 0.1 vol of 100 mM Bis-Tris-HCl, pH 7.0, 30% (w/v) sucrose, 5% w/v Coomassie Blue Brilliant G and run in a mini-vertical unit (Hoefer, Inc. Holliston, MA, USA) (110 × 100 mm, 0.75 mm thick) at 4°C, by applying a constant voltage of 50 V, overnight, which was then gradually increased up to 200 V until completion. For visualization and further sampling for MS-based protein identification, gel lanes were stained using the blue-silver protocol [64]. Apparent molecular mass of bands was determined by using the Native Mark Unstained kit (Invitrogen Life Technologies, USA)

Non-stained gel lanes from 1D-BN-PAGE were cut out immediately and further subjected to a second dimension run by urea-PAGE or SDS-PAGE separation. For urea-PAGE, gel lanes were equilibrated in 6 M urea, 30% (w/v) glycerol, 4% (w/v) SDS, 2% (w/v) DTT, 150 mM Tris-HCI, pH 6.8, for 15 min, and then reacted with 2.5% (w/v) iodoacetamide solved in the same buffer but depleted of the reducing agent, for additional 15 min. Gel lanes were then rinsed in equilibration buffer for 2 min and finally loaded onto the second dimension 12% T gel (1 mm thick). For SDS-PAGE, gel lanes were equilibrated in 150 mM Tris-HCl pH 6.8, containing 10% (w/v) glycerol, 2% (w/v) SDS, 2% (w/v) DTT for 15 min, followed by a second incubation with the same buffer depleted of the reducing agent but containing 2.5% (w/v) iodoacetamide for 15 min. Gel lanes were then rinsed in equilibration buffer for 2 min and finally loaded onto the second dimension 9-16% T gradient gel (1 mm thick). In both cases, proteins were resolved at a constant current (25 mA) and visualized by using a blue-silver-based staining protocol [64].

2.4 Protein digestion and mass spectrometry analysis

Bands from 1D-BN-PAGE or spots from 2D-BN/urea-PAGE and 2D-BN/SDS-PAGE were manually excised from the gels, triturated and washed with water. Proteins were *in-gel* reduced, S-alkylated and digested with trypsin, as previously reported [65]. Protein digests were subjected to a desalting/concentration step on μZipTipC18 pipette tips (Millipore Corp., Bedford, MA, USA). Peptide mixtures were then analyzed by nLC-ESI-LIT-MS/MS using a LTQ XL mass spectrometer (ThermoFinnigan, USA) equipped with a Proxeon nanospray source connected to an Easy-nLC (Proxeon, Denmark) [66]. Peptide mixtures were separated on an Easy C18 column (100 x 0.075 mm, 3 μm) (Proxeon) using a gradient of acetonitrile containing 0.1% formic acid in aqueous 0.1% formic acid, at a flow rate of 300 nL/min; i) for spot identification, acetonitrile was ramped from 5% to 35% over 10 min, from 35% to 95% over 2 min and then remained at 95% over 12 min; ii)

for band identification, acetonitrile was ramped from 5% to 40% over 40 min, from 35% to 80% over 10 min, from 80% to 95% over 2 min and then remained at 95% over 12 min. Spectra were acquired in the range m/z 400-2000. Acquisition was controlled by a data-dependent product ion scanning procedure over the three most abundant ions, enabling dynamic exclusion (repeat count 1 and exclusion duration 1 min). The mass isolation window and collision energy were set to m/z 3 and 35%, respectively.

2.5 Protein identification

nLC-ESI-LIT-MS/MS data were searched by using Mascot (version 2.2.06) (Matrix Science, UK) and Sequest within Proteome Discoverer (version 1.3) software package (Thermo, USA) against an updated *S. thermophilus* database containing available protein sequences (NCBI 24/05/2012, 27333 sequences). As searching parameters, we used a mass tolerance value of 2 Da for precursor ion and 0.8 Da for MS/MS fragments, trypsin as proteolytic enzyme, a missed cleavages maximum value of 2, Cys carbamidomethylation and Met oxidation as fixed and variable modification, respectively. Protein candidates with more than 2 assigned unique peptides with an individual Mascot ion score > 25 and a significant threshold (p < 0.05), and/or a Sequest X_{corr} value > 1.5 (for +1), 2.0 (for +2) and 2.2 (for +3 and higher charges) were further considered for protein identification.

2.6 Bioinformatic analysis

Protein entries from spots in the same vertical line within 2D-BN/urea-PAGE or 2D-BN/SDS-PAGE, and in the corresponding band from 1D-BN-PAGE were analyzed by STRING v. 9.05 (http://string-db.org/), using S. thermophilus LMG18311 as selected organism. GO enrichment for biological processes, molecular functions and cellular components was also performed. The latter option was used to verify the occurrence of identified components as related to a membrane environment. Proteins or protein horthologs within each resulting STRING map were then searched against the eNet database (http://ecoli.med.utoronto.ca/index.php), or against combined MPIDB (http://jcvi.org/mpidb/about.php) [57],IntAct (http://www.ebi.ac.uk/intact) [59], DIP (http://dip.doe-mbi.ucla.edu/dip) [60] and BIOGRID (http://thebiogrid.org) database [61], using the service PSICQUIC interface (http://www.ebi.ac.uk/Tools/webservices/psicquic/view) [62]. This protocol was applied to all components as deriving from 1D-BN-PAGE, 2D-BN/urea-PAGE and 2D-BN/SDS-PAGE experiments, which were then critically evaluated according to available interaction information. A parallel analysis of each protein or protein hortholog for its oligomeric

state as deriving from literature data or crystallographic information at PDB database (http://www.rcsb.org/pdb/home/home.do) was also performed. In this case, hortholog searching was performed by BLASTP analysis (http://blast.ncbi.nlm.nih.gov/Blast.cgi).

2.7 Eps and rgp gene clusters sequencing analysis

The sequence of *eps* and *rgp* gene clusters (accession HG321352 and HG321353) was obtained from a draft genome sequence of strain DSM20617. Partial genome sequencing was obtained from GenProbio s.r.l. (Codorago, Italy) by using the Ion Torrent PGM (Life Technologies, Germany). Functional annotation was performed by the Rapid Annotation using Subsystem Technology server [67] and checked by BLAST analysis [68] to verify and, if necessary, to redefine the start of each predicted coding region or to remove or add coding regions.

2.8 Analysis of exopolysaccharide production

Exopolysaccharide production was evaluated in ruthenium red milk (RRM) plates consisting of 0.5% w/v yeast extract, 10% w/v skim milk powder, 1% w/v sucrose, 1.5% w/v agar, and ruthenium red (0.08 g/l). Ruthenium red stains the bacterial cell wall producing red colonies for nonropy strains and white colonies for ropy strains [69,70].

Bacterial cells were collected by centrifugation, and the resulting pellet was processed for transmission electron microscopy. The extract treated bacterial cells were fixed in 2.5% glutaraldehyde, and later post-fixed with 1% osmium tetroxide (in 0.1 M cacodylate buffer, pH 7.2) for 2 h, at room temperature. After eliminating the remaining osmium tetroxide, the samples were dehydrated in a graduated cold ethanol series (35–100%); each step was performed for about 10–15 min, at room temperature. The fixed cells were embedded in Epon 812. Blocks were cut with an ultramicrotome (Ultracut; Reichert), and collected on nickel grids. Sections were post-stained with 5% uranyl acetate for 5 min at room temperature, and treated with lead citrate for 1 min. Sections were observed and photographed with a Philips CM 12 electron microscope and a Zeiss 900.

3. Results and Discussion

3.1. Isolation and separation of membrane protein complexes

A global prediction of the membrane proteins within the *S. thermophilus* LMG18311 genome already identified 326 sequence entries containing at least one transmembrane helix (TMH) [7]; among that, 220 were predicted to contain more than 2 TMHs and 95 were clearly identified as

transport system (TPS) components, which included 48 ATP-binding cassette (ABC) transporters, 29 secondary transporters, 7 ion channels, 6 F- or P-type ATPases, and 2 sugar phosphotransferase systems (PTS) [7]. Within the ABC transporter group, 30 and 18 were classified as importers and exporters, respectively. Reduced content of sugar importers in *S. thermophilus* genome supported its low capacity for sugar uptake, with respect to other streptococci [6,7,69]. A high percentage of pseudogenes (20%) occurred within the transporter group. Other accessory factors involved in transport or membrane-associated components non-containing TMHs were also identified [7].

In order to optimize the preparation of membrane protein complexes from *S. thermophilus*, different detergents were tested. Those suitable for efficient extraction of membrane components often did not allowed an optimal complex recovery (data not shown). Among non-ionic detergents (βDDM, digitonin and triton-X-100) tested, βDMM generated 1D-BN-PAGE profiles showing the highest abundance of bands putatively ascribed to protein complexes. A further refinement of the concentration range of βDMM to be used for preparative membrane complexes extraction was also obtained (Supplementary Figure S2); thus, a concentration value of 0.5% (w/v) βDMM ensured a sufficient protein extraction power, together with a certain ability to resolve a number of protein complexes in a more or less intact form within a 60-720 kDa mass range. This guaranteed a high protein complex representation within 2D-BN/urea-PAGE and 2D-BN/SDS-PAGE maps (see below and data not shown); however, since each protein complex may have a diverse sensitivity to solubilization, we suggest to test various experimental conditions if it has to be fully characterized for its compositional/stoichiometric properties.

After initial resolution of the protein complexes by 1D-BN-PAGE, two complementary approaches (urea-PAGE and SDS-PAGE) were used for the separation in the second dimension (Figure 1 and 2); in general, a reduced spot diffusion was observed in the first case. This combined procedure ensured confirmative data, but also provided complementary information for specific protein complexes. A similar condition was also verified by cross-relating data from 1D-BN-PAGE, 2D-BN/urea-PAGE and 2D-BN/SDS-PAGE experiments (Figure 1 and 2); in fact, the first analysis highlighted the occurrence of minor protein components that were not sampled in the 2D maps (as result of their migration in faint, diffused spots) or were absent therein as result of their poor solubility within the PAGE matrix. Synergic effect of combining data from 1D and 2D-BN-PAGE has been already underlined in previous studies on complexomes from other prokaryotes and eukaryotes [41,43,49,54,55,71]. Protein information on spots from 2D-BN/urea-PAGE or 2D-BN/SDS-PAGE, and bands from 1D-BN-PAGE always derived from nLC-ESI-LIT-MS/MS analysis of the corresponding *in gel* tryptic digests. In general, 1D-BN-PAGE ensured a higher

number of identified proteins with respect to 2D-BN/urea-PAGE and 2D-BN/SDS-PAGE; this was probably the result of the higher amount of sample loaded for analysis and/or the absence of a second dimension separation. For evaluation/interpretation of the whole experimental results, an integration of 1D-BN-PAGE migration and MS-based identification data with available protein interaction and oligomerization information in other bacteria was achieved. Altogether, these integrated experiments allowed describing 65 heteromeric and 30 homomeric protein complexes where a total of 110 gene expression products were present (Table 1).

A *post-hoc* evaluation of the nature of the proteins identified from 1D-BN-PAGE, 2D-BN/urea-PAGE and 2D-BN/SDS-PAGE experiments generally demonstrated negligible cytoplasmic contaminations, as revealed by the absence of abundant cytosolic proteins already identified in previous dedicated studies on *S. thermophilus*, i.e. transcription elongation factor NusA, Mn-dependent inorganic pyrophosphatase and most ribosomal particle constituents [13,14]. In parallel, various moonlighting proteins were also identified; their occurrence in membranes has been already reported in other bacteria [72,73]. In general, our analysis described a number of protein complexes that are representative of the most important functional modules within the cell membrane. Protein machineries involved in polysaccharide biosynthesis, molecular uptake, energy metabolism, cell division, protein secretion, folding and chaperone activity were highly represented in 1D-BN-PAGE, 2D-BN/urea-PAGE and 2D-BN/SDS-PAGE profiles; a number of homomeric moonlighting proteins were also identified. Their properties will be discussed in detail in the following sections, here organized according to a functional criterion.

3.2 Exopolysaccharide biosynthesis

The ability of *S. thermophilus* to produce exopolysaccharide (EPS) is important for the dairy industry, as it enhances the texture and mouthfeel of certain fermented dairy products. EPS is highly variable among *S. thermophilus* strains and consists of heterosaccharide polymers primarily made of galactose, glucose and rhamnose monomers [74,75]. EPS biosynthesis in *S. thermophilus* involves binding of sugar monomers to a lipid carrier, using amino sugars as precursors. This reaction is performed by a galactose-1-phosphate or glucose-1-phosphate transferase, and subsequent attachment of different monomers is performed by glycosyl transferases. In addition to this, enzymes for polymerization and transmembrane translocation are needed [75,76]. Coding genes for these enzymes are arranged into a main EPS cluster, which generally contains 12-25 gene entries and shows an extremous degree of variability among different bacterial strains [8,69,74,77]; thus, more than 60 different *S. thermophilus* EPS gene clusters have been predicted by restriction

fragment length polymorphism analysis [78]. The modular gene organization is conserved in all EPS clusters and the biosynthesis of EPS is proposed to occur via a common molecular mechanism. Interestingly, the *S. thermophilus* genome also contains a second gene cluster predicted to be involved in rhamnose–glucose polysaccharide (RGP) production. Six conserved genes (rgpA-F) (including two ones coding for molecular ABC exporter components) determine the assembly and secretion of the rhamnose–glucose polysaccharide, while two or more variable genes located upstream (rgpH-I) are required for glucose side-chain coupling, controlling the frequency of branching [7,79]. Many aspects of polysaccharide biosynthesis are still not fully understood, such as the sequence similarity of some enzymes involved in EPS and RGP assemblage, or their eventual, concomitant occurrence in functionally-active machineries.

In this study, a number of protein complexes made of both eps and rgp gene products were observed; additional complexes made only of eps-coded enzymes were also identified. In particular, epsB-epsC, rgpA-rgpD-rgpE-rgpF-epsI-epsJ, rgpA-rgpD-rgpE-rgpF-epsI, rgpA-rgpD-rgpF-epsIster1438, rgpA-rgpD-rgpF-epsI, rgpA-rgpD-rgpF-ster1438, epsG-epsI-epsJ-epsN-ster1442, epsIepsJ-epsN-ster1442, rgpA-epsG-epsI, epsD-epsN-ster1440 and rgpA-epsI complexes were characterized by combining MS data of samples from 1D-BN-PAGE, 2D-BN/urea-PAGE and 2D-BN/SDS-PAGE (Figure 1 and Table 1). Protein components were identified as deriving from different S. thermophilus strains; their nature strongly reflected the high variability of the corresponding EPS clusters and the absence of genomic information on the DSM20617 strain. Protein redundancies were excluded on the basis of sequence alignment of all identified species. In some cases, complex nature reflected progressive decomposition of higher structures. Generally, their molecular mass value was in good agreement with that expected theoretically. Exceptions regarded poorly-represented epsB-epsC-containing complexes at 602 and 518 kDa, for which the occurrence of additional constituents (in low amounts) escaping a positive MS identification may be hypothesized, as already reported for other poorly-abundant protein complexes from other bacteria [44,45,52,55,57].

To further investigate the arrangement of the genes coding for the proteins reported above in the corresponding clusters and to verify the sequence of those coding for ster-related entries, a successive, dedicated analysis was performed on strain DSM20617. Partial genome sequencing revealed that the EPS locus is composed of 13 genes and one pseudogenes (*epsH**) (Figure 3A), which show a high sequence identity (93-99%) with counterparts from the *S. thermophilus/S. salivarius/S. vestibularis* group. Concerning the genes organization in the EPS locus, only the first part of the cluster (*epsABCDE*) appeared as highly conserved among *S. thermophilus* strains.

Complessively, the entire EPS locus showed a genes order highly similar to that of *S. mitis* NCTC 1, despite a low sequence identity. On the other hand, the RGP locus was composed of 14 genes showing high sequence similarity (99-100%) with orthologs from *S. thermophilus* and *S. parasanguinis* strains (Figure 3B). Both EPS and the RGP locus showed a low GC content (36 and 37% respectively) if compared to that of the whole genome GC (39%), thus suggesting a potential role of horizontal gene transfer events in the acquisition/assembly of these gene clusters.

On the basis on the results reported above, it was possible to ascertain that a number of enzymes coded from genes present in the same cluster establish positive interactions to each others (Table 1 and Figure 3). Their identification in the \(\beta\)DMM-extracted fraction was suggestive for the occurrence of two dedicated biosynthetic machineries as embedded into the lipid bilayer to ensure trafficking of the assembled sugar oligomers from the inner side of the cell membrane toward the bacterial surface, for its incorporation in the bacterial capsular structures or its eventual release in the medium. Our results confirmed previous data on epsB-epsC-epsD binding in S. thermophilus and other pathogenic streptococci, as deriving from co-purification, co-immunoprecipitation or twohybrid assays [76,80] but, at the same time, they also highly expanded the interaction maps of exopolysaccharide biosynthesis enzymes [76,81-83]. On the other hand, the simultaneous occurrence of mixed gene products from EPS and RGP clusters onto independent protein machineries having putative separate oligosaccharide translocation mechanism across membranes was never reported so far; it was highly suggestive of a hierarchical organization of the complexes into a unique, synchronized, functional biosynthetic module. In this context, the occurrence of genes coding for proteins involved in the synthesis of the dTDP-rhamnose precursor have been already demonstrated in the EPS cluster of different pathogenic streptococci and lactobacilli [76,79,84,85], evocating a sort of genetic cross-talk between the corresponding rhamnose- and galactose/glucose-based biosynthetic machineries [75,76]. On the other hand, the functionality of the whole exopolysaccharide biosynthesis module in S. thermophilus DSM20617 was confirmed by a ruthenium red stain assay, which revealed white colonies (unstained) on agar plate (Figure 4A), and previous data [74]. Transmission electron microscopy confirmed the presence of a diffuse polysaccharide matrix on the surface of the bacterial cells (Figure 4B and 4C), highly similar to that reported for the closest neighbor Lactococcus lactis [79]. On the whole, our results can provide original insights for future studies on EPS production in lactic acid bacteria.

3.3 Solute transport systems

A number TMH-containing proteins and TPS components, including ion channels, secondary transporters, sugar PTSs, ABC transporters and ATP synthases, were recognized as constituents of protein complex structures present within distinct bands from 1D-BN-PAGE or as vertical lines of spots in 2D-BN/urea-PAGE and 2D-BN/SDS-PAGE (Figure 1 and Figure 2). Among the porter proteins, the permease LacS (essential for lactose uptake) was found as bound to the HtrA chaperone/protease involved in the folding/degradation of secreted proteins (Table 1). The crucial role of the secretory machinery for the proper localization of folded LacS within the cell membrane has been already reported [86,87]. A similar HtrA-bound condition was also observed for the phosphotransfer-driven group translocator ScrA, involved in sucrose transport. In both cases, our data suggest a specific function of HtrA in assisting the proper folding of these TMH-containing proteins and/or degradation of the corresponding misfolded counterparts. Molecular migration of both complexes in 1D-BN-PAGE was compatible with a dodecameric structure of HtrA containing a single substrate molecule, as already observed in other bacteria [88]. Conversely, native LacS and ScrA were absent within 1D-BN-PAGE, 2D-BN/urea-PAGE and 2D-BN/SDS-PAGE as homodimeric species [89,90] (Figure 1, Figure 2 and Table 1); this was not surprising on the basis of their recalcitrance to be extracted from membranes by soft detergents, as already observed in our laboratory [14].

Three complexes involving members of the phosphoenolpyruvate: glucose/mannose PTS [91], which play a key role in coupling active sugar transport across the cell membrane to a sequential phosphorylation cascade, were also detected by 1D-BN-PAGE. In particular, the mannose PTS system components IIAB (ManL) and IIC (ManM) were identified within a band migrating at about 196 kDa. Together with component IID (ManN), these proteins constitute the glucose/mannose PTS transporter with a 2:1:2 (ManL:ManM:ManN) stoichiometry [47,53,92]. Observed migration was consistent with an intact glucose/mannose PTS transporter. Its functional expression may be related to its regulatory functions more than in glucose transport, as already hypothesized [91]. On the other hand, ManL was also detected as bound to its phosphorylating effector PtsH (Figure 1 and Table 1); measured gel migration was consistent with a dimeric state of both proteins therein, as already revealed by NMR analysis [93]. In parallel, PtsH was also observed to participate in another complex with PtsI (Figure 1 and Table 1). Direct interaction of these expression products from two contiguous genes in the same ORF (stu1264 and stu1265) [91] was already reported in *E. coli* [94]; also in this case, gel migration was in agreement with the presence of protein dimers within the complex [95]. Detection of ManL-PtsH and PtsH-PtsI complexes in 1D-BN-PAGE was very

surprising based on their underlying, relatively weak protein interactions [93,95]. Their observation was putatively ascribed to the relative high concentration of its single constituents, namely ManL and PtsI, as revealed by 2D-BN/urea-PAGE, and the very reduced mass increase of the corresponding complexes due to PtsH contribution ($\Delta M \sim 9 \text{ kDa}$) (Figure 1).

The archetypal member of the aquaporin superfamily, i.e. glycerol facilitator protein, was also detected in 1D-BN-PAGE and 2D-BN/urea-PAGE (Figure 1). Its migration properties (at about 140 kDa) and the absence of known interactors of this porin in the corresponding gel portion, as deduced by eNet and PSICQUIC analysis, strongly supported the occurrence of this protein as tetrameric species, in agreement with previous cryoelectron microscopy and X-ray crystallography studies [96].

Within the ABC transporter group, a number of products from contiguous genes present within the same ORF were identified within distinct bands from 1D-BN-PAGE or as vertical lines of spots in 2D-BN/urea-PAGE and 2D-BN/SDS-PAGE (Figure 1, Figure 2 and Table 1). This was the case ofLmrA1(stu0433)-LmrA2(stu0434), stu0296-stu0297, stu0808-stu0809. Peb1(stu1161)-GlnQ(stu1162), LivG(stu0362)-LivF(stu0363) and PstB2(stu1005)-PhoU(stu1006) complexes; in the latter case, another complex (PstB2-PhoU-FtsZ-Tuf) was also recognized as made of additional cell cytoskeletal proteins. Specific ABC transporter components were also identified in additional macromolecular aggregates migrating in 1D-BN-PAGE at different mass values (Figure 1 and Table 1). Only for Peb1-GlnQ, LivG-LivF, PstB2-PhoU and PstB2-PhoU-FtsZ-Tuf complexes, measured migration properties were in good agreement with what expected on the basis of the corresponding theoretical mass values. For the remaining complexes, additional constituents determining observed migration in 1D-BN-PAGE may have escaped a positive MS identification due to the low amount of protein generally detected, as already reported for other bacteria [44,45,52,55,57]. From their migration in 1D-BN-PAGE, 2D-BN/urea-PAGE and 2D-BN/SDS-PAGE, previous literature data and the organization of the corresponding ABC transporter gene clusters in S. thermophilus, it is tempting to speculate that components at 222 and 135 kDa correspond to intact stu0808-stu0809-stu0810-stu0811 and PstS-PstB1-PstB2-PstC1-PstC2-PhoU complexes, respectively [6,7,44,45,52,55,57]. While information on putative functional efficiency of LmrA1-LmrA2, Peb1-GlnQ, LivG-LivF, PstB2-PhoU and stu0808-stu0809 interactions may be deduced by simple genome analysis, being part of complete multidrug (DrugE2) family ABC exporter and polar amino acid, branched-chain amino acid, phosphate and carbohydrate ABC uptake transporters, respectively [6,7,97,98], it is no clear whenever stu0296-stu0297 are a part of an active protein machinery, being constituents of an incomplete transporter [6,7].

We also observed the larger part of the respiratory chain-related F_1F_0 -ATP synthase complex, which catalyzes ATP synthesis during oxidative phosphorylation and ATP hydrolysis to generate the transmembrane proton electrochemical gradient required for different cell functions [99]. F_1F_0 -ATP synthase contains 8 different subunits in a known stoichiometry ($\alpha_3\beta_3\gamma\delta\epsilon AB_2C_{10-14}$) and exhibits a total molecular mass of about 530 kDa; the complex consists of 2 parts designated as F_0 and F_1 . F_0 is membrane embedded and consists of subunits A, B and C, while F_1 is membrane-extrinsic and consists of 5 subunits, i.e. α , β , γ , δ and ϵ [100]. In this study, we observed only the F_1 part, which was detected within 3 distinct bands in 1D-BN-PAGE or as vertical lines of spots in 2D-BN/urea-PAGE and 2D-BN/SDS-PAGE (Figure 1, Figure 2 and Table 1). The first one (at about 403 kDa) well fitted with the molecular mass of the whole F_1 complex and contained all its subunits. The second (at 371 kDa) and the third one (at 333 kDa) showed progressive disappearance of γ , or γ and ϵ subunits, respectively, which was associated with a partial F_1 complex decomposition. Our results were in good agreement with previous observations on other G+ and G-bacteria [40,43-47,52,53].

3.4 Cell growth and morphology

Bacterial division is generally driven through the formation of a macromolecular machinery (divisome) containing at least a dozen of proteins, which assembles with a defined dependence hierarchy at a specific cell membrane site [101]. The way in which the divisome assembles has been studied extensively in *E. coli* and *B. subtilis*, leading to related assembly pathways that requires the sequential assembly of different subcomplexes [102,103]. In fact, additional proteins associated with cell cytoskeletal structure and peptidoglycan (PG) biosynthesis transiently interact with the divisome machinery depending on division moment. Likewise other bacteria, various genes coding for division proteins in *S. thermophilus* occur within a specific ORF (stu0731-0740), where genes involved in cell wall biosynthesis are also present [6,7]. Additional genes coding for cell morphogenesis proteins and PG synthesis enzymes resides in distinct ORFs.

In our study, a number of macromolecular aggregates were identified in 1D-BN-PAGE and 2D-BN/urea-PAGE experiments. In particular, FtsZ-EzrA-DivIVA-MurG, FtsZ-DivIVA-MurG, FtsZ-EzrA-DivIVA, FtsZ-DivIVA, FtsZ-MurG-Pbp2X-SecA-Tuf, FtsZ-DivIVA-SecA-PrtM-Tuf, FtsZ-DivIVA-MurG-Pbp2X-Tuf, FtsZ-SecA-DivIVA-FtsY, MurG-MurM-MurE-Pbp2X, DivIVA-MurG-SecA, MurG-MurM-MurE-Upps, MurG-MurM-MurE and MurG-MurM complexes were characterized, in agreement with previous interaction studies (as also verified by eNet and PSICQUIC analysis) [34,40,47,104]. Considering the possible occurrence of oligomeric proteins

(for FtsZ, DivIVA and SecA) within observed complexes, measured migration properties were in good agreement with what expected on the basis of the corresponding theoretical mass values. Ascertained assemblies often reflected progressive decomposition of higher structures. In general, complexes composition highlighted the simultaneous occurrence of entries uniquely made of elements from the divisome machinery, of enzymes involved in PG biosynthesis, or where mixed elements from both protein classes occurred together. In this context, it has been already suggested that, after initial division stages driven by FtsZ ring formation [105], the divisome locally recruits an assembled multiprotein Mur subcomplex made of enzymes assisting lateral envelope growth [101,102,106]. PG glycosyltransferase MurG was suggested to be a common component of both complexes playing a crucial role for their interaction [107]. Our results were consistent with this scenario. In rod-shaped cells, it has been hypothesized that the cotranslational assembly and localization of the divisome and of the Mur subcomplex, driving the flux of PG precursors toward the septum synthesis machinery, occurs through a genomic channeling mechanism [101].

Ascertained FtsZ-MurG-Pbp2X-SecA-Tuf, FtsZ-DivIVA-SecA-PrtM-Tuf, FtsZ-SecA-DivIVA-FtsY and DivIVA-MurG-SecA complexes also included proteins of the secretory machinery [86,87], namely SecA, FtsY and PrtM. Inclusion of the motor ATPase SecA has been already reported in cell wall biosynthesis and division functional modules of other bacteria [104]; it has been related to the possible role of this protein in directing secretion of the PG synthetic apparatus to regions where PG biosynthesis occur [108]. In fact, many proteins that carry out or mediate PG biosynthesis contain TMHs or membrane anchors linked to large extracellular domains, which are likely exported in a SecA-dependent manner. Bioinformatic analysis of our data confirmed the capability of SecA to interact with FtsZ and MurG [104], thus sanctioning a putative cross-talk of the bacterial secretory machinery with the cell growth- and morphology-affecting complexes mentioned above.

3.5 Protein elongation, secretion and folding

According to *S. thermophilus* genome analysis, components of the secretory machinery include signal recognition particle proteins Ffh and FtsY, trigger factor chaperone RopA, Sec translocase constituents (SecA-SecYEG and YajC), two ortholog proteins of YidC (stu1810 and stu0245) interacting with the translocase, TatA and TatC components of the twin Arg translocation pathway, various signal peptidases (SipA, SipB, LspA, Lgt, Sip and PilD), a PrsA/PrtM peptidylprolyl isomerase (lipoprotein) assisting the folding of the exported proteins and HtrA [7].

In addition to the already-mentioned complexes containing elements from different functional modules, protein assemblies uniquely related to the secretory machinery [86,87] were also recognized on the S. thermophilus membrane. In particular, SecA-FtsY-PrtM, SecA-FtsY-Ffh and SecA-Tuf complexes were identified in 1D-BN-PAGE and 2D-BN/urea-PAGE experiments (Table 1). These results were in good agreement with previous studies on other bacteria where, similarly to what reported here, additional complexes (Ffh-FusA-Tuf, Tuf-GroL-ClpL-PotA, Tuf-RpsA-FusA-DnaK, DnaK-RpsA-ClpL, DnaK-RpsA-Pyk, RpsA-ClpL, GroL-DnaK and Tig-Pyk) related to the elongation cycle of protein biosynthesis and/or involving protein folding-assisting interactions with various chaperones were also observed [32-34,46,50,53,104,109-111]. Ascertained relationships were coherent with that observed following eNet and PSICQUIC analysis. At present, it is not clear if these complexes were related to nascent, unfolded polypeptide chains before their localization close to/within the lipid bilayer or to membrane proteins that are partially unfolded therein. A number of chaperones, elongation factors and ribosomal protein antigens with moonlighting properties were identified as complex constituents, in agreement with previous investigations [72,73,112-115]. Among that, 60 kDa chaperonin GroL that was identified as an abundant homomeric complex migrating at about 518 kDa in 1D-BN-PAGE; its migration properties were not coherent with the characteristic epta- or tetradecameric structure of this chaperone, but strongly resembled those already observed during 2D-BN/SDS-PAGE analysis of membrane proteins from H. pylori [50]. In vitro studies have demonstrated that GroEL can mediate post-translational membrane insertion of lactose permease [116], bacteriorhodopsin [117] and holin [118]. However, in a proteome wide screen no membrane proteins were identified as GroEL substrates [119]. An analogous homomeric condition was observed for the ATPase/protein unfoldase ClpL, which was also observed as a tetrameric species migrating in 1D-BN-PAGE at about 303 kDa, in agreement with previous analytical ultracentrifugation and light scattering experiments [120]. On the basis of what reported above, further studies have to be accomplished to unveil the precise structurefunction relationship of these moonlighting chaperones in a membrane environment and their role in mediating inter-cellular interactions [72].

3.6 Moonlighting proteins with different function

In addition to the already-mentioned chaperones and elongation factors, other moonlighting proteins (GlnA, Eno, GdhA, RpoC, DeoD, Ldh, GapN, Pyk, Pfk, Gapdh, TpiA, Pgi, Pgma and GltX) were also identified as abundant component present on the membrane fraction of *S. thermophilus*; their peculiar localization has been already reported in other bacteria, including

streptococci and lactobacilli [42,72,73,113,114,121]. About an half of them participate in the glycolytic pathway, but has also been reported to mediate adhesion to exogenous plasminogen, fibrinogen and cytoskeletal components as well as to modulate cell signaling processes [72]. Most of these moonlighting proteins occurred as homomeric complexes; this was the case of GlnA, Eno, GdhA, DeoD, GapN, Pyk, Pfk, Gapdh, Ldh and Pgma, which migrated in 1D-BN-PAGE as dodecameric, octameric, hexameric, hexameric, tetrameric, tetrameric, tetrameric, tetrameric, tetrameric, tetrameric and tetrameric species, respectively, in agreement with previous proteomic studies on other bacteria [42,48,53] and protein structures present within the PDB database (http://www.rcsb.org/pdb/home/home.do). These findings suggest the maintenance of the corresponding protein quaternary structures also in a membrane environment.

On the other hand, known moonlighting proteins were also observed to participate in heteromeric complexes, as in the case of the RpoC, which was present in the complexes RpoA-RpoB-RpoC, RpoB-RpoC-stu0256 and RpoC-RplJ-RplS. The occurrence of the RNA polymerase in bacterial membrane is not surprising [53,114]; a band showing a migration compatible with the whole biosynthetic machinery (where only RpoA, RpoB and RpoC were identified) has been already reported in E. coli [53]. In this case and in the current study, remaining RNA polymerase constituents may have escaped MS identification due to the low amount of protein present. RpoB and RpoC were also observed in other complexes; in this context, BLAST analysis identified stu0256 as a putative nucleotide triphosphate pyrophosphatase that hydrolyzes non-standard purines preventing their incorporation into RNA. Analogously, participation into heteromeric complexes was also verified for moonlighting dimeric proteins TpiA and Pgi that, according to their migration in 1D-BN-PAGE, were involved in binding to dimeric Pfl and MetN, respectively. These interaction data were in agreement with previous observations in E. coli [32]. Also in these cases, predicted protein oligomerization in a membrane environment was similar to that observed in aqueous media. Finally, moonlighting protein GltX was observed as bound to FabF and as monomeric species (data not shown), in agreement with previous chromatographic data [122]. This protein was previously identified as a cell wall-associated antigen in S. pneumonia [121].

Other proteins generally reported as cytosolic components were also observed in the membrane fraction of *S. thermophilus*; they included Dpr, Prs1, Prs2, PurB, PlsX and UspA (Figure 1 and Figure 2). A careful evaluation of available literature confirmed their possible occurrence also on bacterial membrane. This was the case of the peroxide-resistance protein Dpr, which was detected on the membrane of *H. pylori* and *E. coli* grown under various environmental conditions [50,123,124]. This protein was suggested to have a scavenging function against reactive oxygen

species and Fe ion misbalance as well as a protective role against DNA damage [125]. Recently, it was proved to influence the attachment of bacteria to abiotic surfaces [126]. Its migration in 1D-BN-PAGE was consistent with a dodecameric structure [125], whose high stability was also appreciated after urea-PAGE (Figure 1) and SDS-PAGE (Figure 1), as already observed in C. thermocellum [45]. On the other hand, two phosphoribosylpyrophosphate synthase isoforms, namely Prs1 and Prs2, were observed to migrate as a vertical line of spots in 2D-BN/SDS-PAGE (Figure 2 and Table 1). Homologue proteins from other bacteria have a functional hexameric structure [127,128]; this information, together with our PSICQUIC analysis that suggested a direct interaction between Prs1 and Prs2, was fully compatible with a heteromeric complex migrating at about 464 kDa, as revealed by 1D-BN-PAGE analysis. Phosphoribosylpyrophosphate synthase was observed as a membrane-bound component in human and rat cells [129,130]. In bacteria, its function has been associated to the biosynthesis of phosphoribosyl-1-pyrophosphate, a central metabolite precursor for cell wall sugar components [131]. Other membrane-associated proteins mentioned above occurred as homomeric complexes. In agreement with data present within the PDB database (http://www.rcsb.org/pdb/home/home.do), PurB, UspA and PlsX migrated in 1D-BN-PAGE as tetrameric, dimeric and dimeric species (Table 1), respectively, thus confirming the maintenance of their protein quaternary structures also in a membrane environment. PlsX was also observed to form a heteromeric complex with LysS and SerS, coherently with data deduced from eNet and PSICQUIC analysis. The occurrence of PurB, PlsX and UspA on the bacterial membrane has been already reported [53,132] and associated with the biosynthesis of fatty acids and membrane phospholipids [133] or with the bacterial response to environmental stresses [134], respectively.

3.7 Proteolytic enzymes and other proteins

Membrane proteases detected in this study included FtsH, PepC and PepB, which migrated in 1D-BN-PAGE as homomeric species present at about 430, 288 and 131 kDa and, respectively. In the first case, observed migration was consistent with the ascertained hexameric crystallographic structure of this membrane-spanning ATP-dependent metalloprotease [135]. It plays a key role in quality and regulatory control within the cell by degrading a unique subset of substrates. In fact, FtsH is able to identify and degrade nonfunctional or damaged membrane proteins by pulling them out of the lipid bilayer, followed by further substrate unfolding and translocation into the proteolytic chamber [111]. On the other hand, PepC is an endopeptidase with moonlighting properties that was observed among the antigenic cell wall-associated proteins of *S. pneumonia*, eliciting protective

immune response in the mouse [121]. Also in this case, 1D-BN-PAGE results were in agreement with the protein hexameric quaternary structure reported in the PDB database. Finally, PepB was never reported as a membrane component so far; its electrophoretic migration was consistent with a dimeric crystallographic structure [136].

Analogous considerations on the absence of data concerning protein membrane localization were valid for phosphopantetheine adenyltransferase CoaD, hypothetical protein stu1225 (homologous to short-chain dehydrogenase/reductases), acetolactate synthase Als, CTP synthase PyrG and enoylacyl carrier protein reductase FabK. These proteins were observed to migrate in 1D-BN-PAGE as hexameric, tetrameric, dimeric and dimeric species, respectively, in agreement with available data on their quaternary structure [137-141].

4. Conclusions

Although milk is a rich growth medium for many microorganisms, bacteria that grow and compete well in the milk environment must, at minimum, be able to use lactose as an energy source and milk proteins as a source of amino acids. The adaptation of *S. thermophilus* to the milk environment is reflected by several observations at genomic and transcriptome levels [6-11,142,143], including the detection of specialized systems for metabolizing lactose, the general absence of other carbohydrate metabolic systems, the presence of amino acid and peptide scavenging machinery, and numerous stress response and host defense mechanisms.

In the present study, a combined approach based on 1D-BN-PAGE, 2D-BN/urea-PAGE, 2D-BN/SDS-PAGE and nLC-ESI-LIT-MS/MS was used to investigate membrane protein complexes in *S. thermophilus* cells at their early exponential phase. We were able to reproducibly separate individual proteins and to reveal protein-protein interactions, consistently with the results obtained through independent, traditional biochemical and biophysical procedures. Among the 110 non-redundant components present in the heteromeric/homomeric complexes here reported, 31 corresponded to about 10% of the 326 membrane proteins predicted by *in silico* analysis of the *S. thermophilus* genome [7], while 7 matched about 7% of the 98 secretory proteins analogously envisaged [144]; on the other hand, 44 species were identified as moonlighting proteins [42,72,73] or components whose transient localization on or close to the bacterial membrane has been already reported. Thus, membrane protein machineries involved in essential biochemical processes, such as polysaccharide biosynthesis, molecular uptake, energy metabolism, cell division, protein secretion and folding, were characterized for their constitutive elements. Information on hypothetical proteins were also derived. In general, most (about 84%) of the heteromeric/homomeric complexes reported

PSICQUIC analysis of the corresponding hortolog species or by evaluation of literature data and crystallographic information present within the PDB database. Novel information on protein machineries involved in exopolysaccharide and peptidoglycan biosynthesis, cell division and protein secretion were obtained. The approach reported here paves the way for a further functional characterization of these protein complexes and will facilitate future studies of their assembly and composition during various experimental conditions and in different mutant backgrounds, with important consequences for biotechnological applications of this bacterium in dairy productions.

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predicted by in silico analysis of the S. thermophilus genome [7]; brefers to secretory proteins predicted by in silico analysis of the S. thermophilus genome available in Suppl. Table S1 (for 1D-BN-PAGE), Suppl. Table S2 (for 2D-BN/urea-PAGE) and Suppl. Table S3 (for 2D-BN/SDS-PAGE). Refers to peptides, sequence coverage and identification score are listed. Identification data reported in this table are those with the highest values as obtained from table includes protein abbreviations used within the whole text. [144]; erefers to moonlighting proteins [72,73] or components whose transient localization on or close to the cell membrane has been already reported. This according to eNet and PSICQUIC analysis, literature data and/or crystallographic records present within the PDB database. a Refers to membrane proteins Sequest but not for Mascot searching. Underlined are protein complex components for which interaction/oligomerization information was already available protein identification data where Sequest results are shown; this condition occurred in the cases in which identification parameters were satisfied for PAGE and 2D-BN/SDS-PAGE are indicated with an asterisk and circle, respectively. MS details for the identification data reported in this table are corresponding protein name, gene name in the LMG18311 or LMD-9 genome [6,8], accession number, theoretical Mr value, number of observed unique Band in 1D-BN-PAGE, protein complex experimental migration (as deduced with respect to commercial molecular standards), protein complex description. 1D-BN-PAGE (Figure 1), 2D-BN/urea-PAGE (Figure 1) and 2D-BN/SDS-PAGE (Figure 2). Protein components identified in spots from 2D-BN/urea-Table 1. Protein heteromultimeric and homomeric complexes identified in the membrane fraction of exponentially-growing S. thermophilus cells.

Band	Α						
migration (kDa)	602						
Protein complex	GlnA (dodecamer)	Stu0296-Stu0297		LmrA1-LmrA2		HtrA-LacS	
Protein name	Glutamine synthetase type I – GlnA*° °	Amino acid ABC transporter substrate-binding protein - Stu0296 a, b	Amino acid ABC transporter substrate-binding protein - Stu0297 a.b	ABC-type multidrug (DrugE2) exporter system, ATPase and permease component - LmrA1 ^a	Multidrug ABC exporter ATP binding/membrane-spanning protein - $LmrA2^{a}$	Trypsin-like serine protease – HtrA* ^b	Lactose permease - LacS ^a
Gene name	Stu1776	Stu0296	Stu0297	Stu0433	Stu0434	Stu2024	Stu1398
Accession	54306535	55822277	55822278	116627330	55820521	116628681	38492233
Mass (kDa)	50.1	31.3	32.9	67.1	65.1	42.8	69.1
Unique Peptides	14	2	4	6	3	3	5
coverage	38.26	9.22	9.67	13.41	6.42	14.36	8.68
Mascot score	528	115	58	134	98	175	118

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403			433									446										518				
ATPase A-ATPase C-ATPase H			FtsZ-EzrA-DivIVA	FtsH (hexamer)			RpoA-RpoB-RpoC		Prs1-Prs2			FtsZ-DivIVA-MurG		Stu0296-Stu0297		<u>EpsB-EpsC</u>	GroL (homomer)					FtsZ-EzrA-DivIVA-MurG		<u>EpsB-EpsC</u>		
H^+ -ATPase cytoplasmic F1-part alpha subunit - ATPase $A^{*\circ a}$	Cell division initiation protein – DivIVA*°	Septation ring formation regulator EzrA - EzrA $^{\rm c}$	Cell division protein FtsZ – FtsZ* c	Cell division protein ${\sf FtsH-FtsH}^{ m c}$	DNA-directed RNA polymerase subunit beta' – RpoC*°°	DNA-directed RNA polymerase subunit beta – RpoB*°°	DNA-directed RNA polymerase subunit alpha - RpoA c	Ribose-phosphate pyrophosphokinase - Prs2*°°	Ribose-phosphate pyrophosphokinase - Prs1*°°	pyrophosphoryl-undecaprenol N-acetylglucosamine transferase - MurG ^c	(IDP-N-acetylg/lucosamine-N-acetylmuramy/-(pentaneptide)	Cell division protein FtsZ – FtsZ*°	Amino acid ABC transporter substrate-binding protein - Stu0297 a	Amino acid ABC transporter substrate-binding protein - Stu0296 a	Exopolysaccharide synthesis protein 4C - EpsC ^a	Glycosyl transferase family protein – EpsB*c	60 kDa chaperonin – GroL*°°	pyrophosphoryl-undecaprenol N-acetylglucosamine transferase - $\mathbf{MurG}^{\mathbf{c}}$	UDP-N-acetylglucosamine-N-acetylmuramyl-(pentapeptide)	Cell division initiation protein – DivIVA*°	Septation ring formation regulator EzrA - EzrA °	Cell division protein FtsZ – FtsZ* °	Exopolysaccharide synthesis protein 4C - EpsC ^a	Glycosyl transferase family protein – EpsB*°	Sucrose PTS, EIIBCA - ScrA ^a	
Stu0482	Stu0740	Stu1520	Stu0735	Stu0012	Stu1867	Stu1868	Stu1908	Stu1460	Stu0023	Stu0732	Stu0 /40	Stu0735	Stu0297	Stu0296	Stu1110	Stu1485	Stu0204	Stu0732		Stu0740	Stu1520	Stu0735	Stul 110	Stu1485	Stu1734	
20070091	116627610	55821496	55822702	116626986	55821839	55821840	81558875	116628164	116626993	81559554	11002/010	55822702	55822278	55822277	24637493	55823391	68566260	81559554		116627610	116628215	55822702	24637401	55823391	116628430	
54.5	33.0	65.4	46.5	6.17	135.3	133.3	34.4	35.1	35.1	40.1	0.00	46.5	32.9	31.3	25.5	35.0	6.65	40.1		33.0	61.5	46.5	25.5	0.58	6.66	
4	3	2	4	3	3	3	2	4	4	4	J	. · · · ·	3	2	4	2	16	ω		14	2	3	3	6	3	
25.55	11.34	3.90	10.91	4.73	2.55	3.27	7.37	19.81	17.76	15.17	17.18	10.00	5.67	9.22	19.13	6.82	53.06	11.80		37.11	4.33	7.50	13.91	17.53	6.32	
300	129	51	177	7.268	118	195	4.16^{8}	112	125	82	190	152	50	93	94	128	775	66		515	83	100	6.628	121	78	

50.4	3024365	Stu0229	Aminopeptidase C – PepC*°°	PepC (hexamer)		
	122267209	Stu1497	UvrABC system protein B, excinuclease ABC subunit B - UvrB			
•	55821983	Stu2016	Inosine 5'-monophosphate dehydrogenase – GuaB*°	<u>GuaB-UvrB</u>	288	J
720	81559720	Stu0487	Elongation factor Tu - Tu f $^{ m c}$			
6980	122266980	Stu1730	Protein translocase subunit SecA - SecA a			
8388	116628388	Stu1701	Cell division protein FtsI/penicillin binding protein $2X$ - Pbp $2X^{b}$			
57800	122267800	Stu0732	ODP-N-acetylglucosamme-N-acetylmuramyl-(pentapeptide) pyrophosphoryl-undecaprenol N-acetylglucosamine transferase - MurG°			
702	55822702	Stu0735	Cell division protein FtsZ - FtsZ°	FtsZ-MurG-Pbp2X-SecA-Tuf		
7804	116627804	Stu1013	$4\hbox{-alpha-glucan otransferase}-MalQ*^\circ$	MalQ (homomer)	303	Ι
1590	55821590	Stu1614	ATP-dependent proteinase ATP-binding subunit - ClpL	ClpL (tetramer)		
7327	116627327	Stu0430	Glutamate dehydrogenase – GdhA*°°	GdhA (hexamer)	318	Н
8026	122268026	Stu0481	ATP synthase F1-sector delta subunit - ATPase H*a			
0338	81820338	Stu0484	ATP synthase F1-sector beta subunit - ATPase D*° a			
0091	20070091	Stu0482	H ⁺ -ATPase cytoplasmic F1-part alpha subunit - ATPase A*° a	ATPase A-ATPase D-ATPase H	333	G
0158	62900158	Stu0256	Non-canonical purine NTP pyrophosphatase - Stu0256			
122266859	1222	Stu1867	DNA-directed RNA polymerase subunit beta? - RpoC °			
55821840		Stu1868	DNA-directed RNA polymerase subunit beta - RpoB °	RpoB-RpoC-Stu0256		
7610	116627610	Stu0740	Cell division initiation protein – DivIVA*°			
2702	55822702	Stu0735	Cell division protein FtsZ – FtsZ* °	FtsZ-DivIVA		
722	81559722	Stu0481	ATP synthase F1-sector delta subunit - ATPase H**			
)338	81820338	Stu0484	ATP synthase F1-sector beta subunit - ATPase D*oa			
3568	55820568	Stu0485	ATP synthase F0F1 subunit epsilon - ATPase C*a			
0091	20070091	Stu0482	H^+ -ATPase cytoplasmic F1-part alpha subunit - ATPase $A^{*\circ a}$	ATPase A-ATPase C- ATPase D-ATPase H		
3529	68053529	Stu0635	Enolase – Eno*°°	Eno (octamer)	371	Ŧ
722	81559722	Stu0481	ATP synthase F1-sector delta subunit - ATPase H*oa			
597	81676597	Stu0483	ATP synthase F1-sector gamma subunit - ATPase G*oa			
338	81820338	Stu0484	ATP synthase F1-sector beta subunit - ATPase D*o a			
)094	20070094	Stu0485	H ⁺ -ATPase cytoplasmic F1-part epsilon subunit - ATPase C* a			

						M														T				K			
						222														243				261			
<u>Dpr (dodecamer)</u>						FtsZ-DivIVA-MurG-Pbp2X-Tuf			RpoC-RpIJ-RpIS						RgpA/EpsF-RgpD-RgpE-RgpF- EpsI-EpsJ					FtsZ-DivIVA-SecA-PrtM-Tuf				Tuf-GroL-ClpL-PotA			
Peroxide resistance protein, non-heme iron-containing ferritin – Dpr*°°	Elongation factor Tu - Tuf c	Cell division protein $FtsI/penicillin$ binding protein $2X$ - $Pbp2X^b$	pyrophosphoryl-undecaprenol N-acetylglucosamine transferase - $MurG^{c}$	UDP-N-acetylglucosamine-N-acetylmuramyl-(pentapeptide)	Cell division initiation protein – DivIVA°°	Cell division protein Fts Z - Fts Z^{c}	50S ribosomal protein L19 - RplS	50S ribosomal protein L10 - RpU	DNA-directed RNA polymerase subunit beta' - RpoC°	Polysaccharide biosynthesis protein - EpsJ ^c	Polysaccharide biosynthesis protein – EpsI*°°	Polysaccharide biosynthesis protein – RgpF*°	Glycosyltransferase – RgpE*°	ABC-type polysaccharide/polyol phosphate transport system, ATPase component - RgpD*	Polysaccharide biosynthesis protein EpsF – RgpA/EpsF*°°	Elongation factor Tu – Tuf* ^c	Protease maturation protein precursor - PrtM ^b	Protein translocase subunit SecA – SecA* ^a	Cell division initiation protein – DivIVA*°°	Cell division protein FtsZ - FtsZ°	Spermidine/putrescine import ATP-binding protein PotA - PotA a	ATP-dependent proteinase ATP-binding subunit - ClpL	60 kDa chaperonin - GroL°	Elongation factor Tu — Tuf* °	Protease maturation protein precursor - PrtM ^b	Signal recognition particle receptor (docking protein) - FtsY $^{\rm a}$	
Stu0723	Stu0487	Stu1701	Stu0732		Stu0740	Stu0735	Stu1179	Stu0536	Stu1867	-	-	Stu1467	Stu1468	Stu1469	Stu1472	Stu0487	Stu0456	Stu1730	Stu0740	Stu0735	Stu1538	Stu1614	Stu0204	Stu0487	Stu0456	Stu1432	
116627595	81559720	116628388	122267800		116627610	55822702	62287370	97182027	122266859	24637448	24637447	116628171	116628172	116628173	90655845	81559720	55822430	122266980	116627610	55822702	122267176	116628305	68566260	81559720	55822430	116628140	
19.2	43.8	82.6	40.2		33.0	46.5	13.1	17.5	135.2	38.7	38.2	68.4	66.1	44.6	44.5	43.8	39.9	96.3	33.0	46.5	43.8	77.1	56.9	43.8	39.9	51.0	
7	12	2	2		3	3	2	2	2	2	2	4	14	6	2	7	3	15	17	4	2	4	3	3	2	2	
38.73	39.95	3.44	7.02		12.71	9.32	20.87	13.17	1.73	5.76	7.01	8.09	25.79	13.25	6.67	24.12	9.43	20.85	36.08	12.73	5.99	8.01	5.75	6.78	7.82	5.18	
386	377	$3.8^{\$}$	5.32	5	96	132	4.758	64	3.678	3.98§	5.908	91	291	67	58	302	105	405	508	121	72	141	63	98	79	79	

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														196										216		
				RgpA/EpsF-RgpD-RgpF-EpsI- Ster1438	GapN (tetramer)				FtsZ-DivIVA-SecA-FtsY			ManL-ManM-ManN	Als (tetramer)	PurB (tetramer)				Tuf-RpsA-FusA-DnaK					RgpA/EpsF-RgpD-RgpE-RgpF- EpsI	Dpr (dodecamer)		Stu0808-Stu0809
Cell wall biosynthesis glycosyltransferase - Ster1438*°	Polysaccharide biosynthesis protein - EpsI c	Polysaccharide biosynthesis protein - RgpF*	ABC-type polysaccharide/polyol phosphate transport system, ATPase component - RgpD ^a	Polysaccharide biosynthesis protein EpsF – Rgp A/EpsF* $^{\circ}$	NADP-dependent glyceraldehyde-3-phosphate dehydrogenase - GapN* $^{\circ\circ}$	Signal recognition particle receptor (docking protein) - FtsY a	Protein translocase subunit SecA – SecA* a	Cell division initiation protein – DivIVA*°	Cell division protein FtsZ - FtsZ c	Mannose PTS system component IID - ManN ^a	Mannose PTS system component IIC - ManM ^a	Mannose PTS system component IIAB - ManL ^a	Acetolactate synthase - Als	Adenylosuccinate lyase - PurB °	Chaperon protein DnaK – DnaK* ^c	Elongation factor G - FusA °	30S ribosomal protein S1 – RpsA*°	Elongation factot Tu – Tuf* ^c	Polysaccharide biosynthesis protein - EpsI ^c	Polysaccharide biosynthesis protein – RgpF*	$Polysaccharide\ biosynthesis\ protein/glycosyltrans fer as e-RgpE*$	ABC-type polysaccharide/polyol phosphate transport system, ATPase component - RgpD ^a	$Polysaccharide\ biosynthesis\ protein\ EpsF-Rgp\ A/EpsF^*c$	Peroxide resistance protein, non-heme iron-containing ferritin – Dpr*°°	Carbohydrate ABC uptake transporter ATP-binding protein - Stu0809 a	Hypothetical protein - Stu0808* ^b
Ster1438	-	Stu1467	Stu1469	Stu1472	Stu1263	Stu1432	Stu1730	Stu0740	Stu0735	Stu0331	Stu0332	Stu0333	Stu0923	Stu0045	Stu0120	Stu1789	Stu0592	Stu0487	-	Stu1467	Stu1468	Stu1469	Stu1472	Stu0723	Stu0809	Stu0808
116628177	24637447	116628171	116628173	24637479	116628006	116628140	122266980	116627610	55822702	55820424	55820425	30027111	20976803	55822037	81676627	62286650	161936373	81559720	24637447	116628171	55823378	116628173	24637426	116627595	116627673	55822775
30.3	38.2	68.4	44.6	44.7	50.8	51.0	96.3	33.0	46.5	33.5	27.8	35.8	52.3	49.5	64.8	76.6	43.9	43.8	38.2	68.4	66.0	44.6	44.6	19.2	55.5	35.6
7	5	6	12	4	20	2	4	2	3	3	3	2	2	4	81	5	15	9	4	8	9	9	2	7	2	4
26.36	14.02	13.43	32.00	12.05	55.14	5.18	5.42	7.22	8.86	12.21	12.36	7.58	4.61	11.34	34.27	10.10	40.75	29.65	13.72	18.07	12.30	30.75	6.67	38.73	4.10	12.80
197	79	128	243	115	870	5.188	67	52	129	7.06 [§]	136	72	62	7.688	467	164	430	230	84	156	82	238	72	386	91	88

70	5.25	2	44.6	116628173	Stu1469	ABC-type polysaccharide/polyol phosphate transport system, ATPase			
120	18.85	5	44.0	116628176	Stu1472	Polysaccharide biosynthesis protein EpsF $-$ RgpA/EpsF* $^\circ$ $^\circ$	RgpA/EpsF-RgpD-RgpF- Ster1438		
2165	50.36	34	77.1	55823518	Stu1614	ATP-dependent proteinase ATP-binding subunit - ClpL*°			
164	20.75	7	43.9	161936373	Stu0592	30S ribosomal protein S1 – RpsA*°°			
218	22.41	10	64.8	81676627	Stu0120	Chaperon protein DnaK – DnaK*c	DnaK-RpsA-ClpL		
7.358	9.58	3	38.7	81820355	Stu0301	Methionine import ATP-binding protein - MetN ^a			
497	38	14	49.8	81170506	Stu0194	Glucose-6-phosphate isomerase - Pgi*°°	<u>Pgi-MetN</u>		
514	40	20	69.1	81820361	Stu0200	Proline-tRNA ligase - ProS*°	ProS (dimer)		
448	23.67	15	96.3	122266980	Stu1730	Protein translocase subunit SecA*oa			
177	16.29	6	40.2	122267800	Stu0732	UDP-N-acety/glucosamine-N-acety/muramyl-(pentapeptide) pyrophosphoryl-undecaprenol N-acety/glucosamine transferase – MurG*°°			
808	36.08	17	33.0	116627610	Stu0740	Cell division initiation protein – DivIVA*°	DivIVA-MurG-SecA	167	Q
94	4.42	3	87.0	55823561	Stu1657	Pyruvate formate-lyase, Pfl			
243	25.40	5	26.7	17066728	Stu0488	Triosephosphate isomerase, TpiA ^c	<u>TpiA-Pfl</u>		
118	12.70	3	28.0	116627797	Stu1005	ABC-type phosphate transport system, ATPase component - PstB2*a	PstB2		
69	18.60	5	38.2	24637447	-	Polysaccharide biosynthesis protein - Epsl ^c			
83	8.43	4	68.4	116628171	Stu1467	Polysaccharide biosynthesis protein - RgpF*			
217	26.25	10	44.6	116628173	Stu1469	ABC-type polysaccharide/polyol phosphate transport system, ATPase component - RgpD ^a			
239	22.31	7	44.5	90655845	Stu1472	Polysaccharide biosynthesis protein EpsF – RgpA/EpsF**°	RgpA/EpsF-RgpD-RgpF-EpsI		
215	17.80	7	54.5	161936368	Stu1196	Pyruvate kinase - Pyk °	Pyk (tetramer)		
4.628	3.97	2	82.6	116628388	Stu1701	Cell division protein FtsI/penicillin binding protein 2X - Pbp2 $X^{ m b}$			
820	52.23	20	46.2	13324647	Stul 155	Peptidoglycan branched peptide synthesis protein, alanine adding enzyme - MurM°			
192	30.9	9	40.2	122267800	Stu0732	UDP-N-acetylglucosamine-N-acetylmuramyl-(pentapeptide) pyrophosphoryl-undecaprenol N-acetylglucosamine transferase - MurG*°°			
351	38.32	14	48.7	116627998	Stu1254	Mur ligase - MurE	MurE-MurG-MurM-Pbp2X		
5.638	6.35	2	57.9	55822851	Stu0889	Signal recognition particle protein - Ffh a			
120	10.15	3	51.0	116628140	Stu1432	Signal recognition particle receptor (docking protein) - FtsY a			
379	9.89	6	96.3	122266980	Stu1730	Protein translocase subunit SecA - SecA*o a	SecA-FtsY-Ffh	171	P

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										131															146			
EpsI-EpsJ-EpsN-Ster1442				PhoU-PstB2-FtsZ-Tuf			<u>LysS-PlsX-SerS</u>	Pgm (dimer)	DeoD (hexamer)	PepB (dimer)			<u>DnaK-RpsA-Pyk</u>					EpsG-EpsI-EpsJ-EpsN-Ster1442			Ffh-FusA-Tuf				MurE- <u>MurM</u> -Mupps			
Polysaccharide biosynthesis protein - EpsI ^c	Elongation factor Tu – Tuf*° °	Cell division protein FtsZ – FtsZ ^c	ABC-type phosphate transport system, ATPase component - PstB2* a	Phosphate uptake regulatory protein - PhoU * c	Serine-tRNA ligase - SerS	Phosphate acyltransferase - PlsX*°°	Lysine-tRNA ligase - LysS*°	Phosphoglucomutase - Pgm*	Purine nucleoside phosphorylase - DeoD* c	Oligopeptidase - PepB*°	Pyruvate kinase - Pyk*°°	30S ribosomal protein S1- RpsA*°°	Chaperon protein DnaK – DnaK*°	Cell wall biosynthesis glycosyltransferase - Ster1442*c	Polysaccharide biosynthesis protein - EpsN*° c	Polysaccharide biosynthesis protein - EpsJ ^c	Polysaccharide biosynthesis protein - Epsl c	Polysaccharide biosynthesis protein - EpsG °	Elongation factor Tu – Tuf*°°	Elongation factor G - FusA ^c	Signal recognition particle protein - Ffh a	Isoprenyl transferase - Upps	Peptidoglycan branched peptide synthesis protein-alanine adding enzyme-MurM°	UDP-N-acetylglucosamine-N-acetylmuramyl-(pentapeptide) pyrophosphoryl-undecaprenol N-acetylglucosamine transferase – MurG*°°	Mur ligase - MurE	Cell wall biosynthesis glycosyltransferase - Ster1438*c	Polysaccharide biosynthesis protein - RgpF*	component - RgpD ^a
1	Stu0487	Stu0735	Stu1005	Stu1006	Stu0329	Stu0028	Stu0692	Stu0787	Stu1113	Stu0454	Stu1196	Stu0592	Stu0120	Ster1442	-		-	•	Stu0487	Stu1789	Stu0889	Stu0197	Stu1155	Stu0732	Stu1254	Ster1438	Stu1467	
24637447	81559720	55822702	55821038	55737024	122268151	122268399	122267836	116627655	24473734	116627347	116627931	161936373	81676627	116628181	24637452	24637448	24637447	24637427	81559720	62286650	55822851	73920281	13324647	122267800	55823172	116628177	116628171	L
38.2	43.8	46.5	28.0	24.9	48.0	35.5	56.4	63.1	18.2	69.2	54.5	43.9	64.8	37.5	39.0	38.7	38.2	42.6	43.8	76.6	57.9	28.7	46.2	40.2	49.4	30.3	68.4	
9	11	2	3	2	3	5	8	4	3	5	10	9	44	5	2	2	3	2	3	2	2	3	16	5	18	7	3	
27.74	35.68	6.59	15.47	9.63	9.18	14.67	16.13	7.17	24.54	7.99	21.60	29.25	64.91	16.92	7.43	5.76	9.45	7.61	9.30	3.32	6.35	12.05	40.84	18.54	43.85	26.36	6.54	
241	506	67	85	57	9.118	108	229	69	171	107	318	159	1442	81	100	4.458	89	60	64	79	5.638	100	557	190	549	197	68	

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													96												118			
	SecA-Tuf			MurE- <u>MurG-MurM</u>	CoaD (hexamer)			PotA-RpIB-RpID		<u>GroL-DnaK</u>		ManL-PtsH	Stu1225 (tetramer)	Gla (tetramer)		PtsH-PtsI	Ldh (tetramer)			RgpA/EpsF-EpsG-EpsI	PyrG (dimer)	Gapdh (tetramer)	Pfk (tetramer)		<u>RpsA-ClpL</u>			
Elongation factor lu – lut*;	Protein translocase subunit SecA – SecA* a	Peptidoglycan branched peptide synthesis protein, alanine adding enzyme – MurM °	UDP-N-acetylglucosamine-N-acetylmuramyl-(pentapeptide) pyrophosphoryl-undecaprenol N-acetylglucosamine transferase – MurG**c	Mur ligase - MurE	Phosphopantetheine adenylyltransferase - CoaD*°	50S ribosomal protein L4 - RplD*°	50S ribosomal protein L2 - RplB*°	Spermidine/putrescine import ATP-binding protein - PotA a	Chaperon protein DnaK – DnaK*°	60 kDa chaperonin – GroL*°	Phosphocarrier protein HPr - PtsH* ^a	Mannose PTS system component IIAB - ManL*° 1	Oxidoreductase, short chain dehydrogenase/reductase - Stu1225*°	Glycerol uptake facilitator protein - Gla ^a	Enzyme I - PtsI*° a	Phosphocarrier protein HPr - PtsH* ^a	Lactate dehydrogenase - Ldh*° c	Polysaccharide biosynthesis protein – EpsI °	Polysaccharide biosynthesis protein – EpsG °	Polysaccharide biosynthesis protein EpsF – RgpA/EpsF*° °	CTP synthetase - PyrG	Glyceraldehyde-3-phosphate dehydrogenase - Gapdh*°°	6-Phosphofructokinase - Pfk*°°	ATP-dependent proteinase ATP-binding subunit - ClpL*	30S ribosomal protein S1 – RpsA*°	Cell wall biosynthesis glycosyltransferase - Ster1442* c	Polysaccharide biosynthesis protein - EpsN*°°	rorysucciunius onosymusons procem - Epso
Stu0487	Stu1730	Stul 155	Stu0732	Stu1254	Stu1648	Stu1933	Stu1931	Stu1538	Stu0120	Stu0204	Stu1265	Stu0333	Stu1225	Stu1671	Stu1264	Stu1265	Stu1280	-	-	Stu1472	Stu0134	Stu1788	Stu0692	Stu1614	Stu0592	Ster1442	-	
81559720	122266980	13324647	81559554	55821261	116628337	81558868	81820219	122267176	81676627	68566260	55821270	30027111	116627959	55823574	30027107	55821270	122267385	24637447	24637480	90655845	116627092	17066732	13629190	116628305	161936373	116628181	24637452	24057440
43.8	96.3	46.2	40.1	49.4	18.7	22.2	29.9	43.8	64.8	56.9	8.9	35.8	28.5	30.8	63.1	8.9	35.4	38.2	42.8	44.5	59.0	36.0	36.0	77.2	43.9	37.5	39.0	36.7
14	5	9	6	9	5	4	7	9	2	6	3	19	3	2	22	3	23	7	5	8	2	3	13	23	7	3	7	v
41.21	7.54	28.71	16.85	25.95	52.12	21.74	30.69	22.66	5.11	13.17	54.02	68.18	13.73	9.41	48.53	58.62	48.48	24.70	15.45	23.08	3.93	12.20	42.48	40.63	19.50	9.54	21.14	13.03
544	340	312	201	331	286	188	154	289	116	114	101	995	130	78	696	6.188	1209	89	88	128	51	65	423	804	202	54	282	160

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												65														70
	PhoU-PstB2		<u>LivF-LivG</u>	UspA (dimer)	FabK (dimer)		RgpA/EpsF-EpsI	PlsX (dimer)		GlnQ-Peb1		MurG-MurM			<u>GlyS-RpIB-RpIM</u>		<u>Tig-Pyk</u>		GltX-FabF			EpsD-EpsN-Ster1440	Pgma (tetramer)	MetN (dimer)		DapA-DapB
ABC-type phosphate transport system, ATPase component - PstB2* a	Phosphate uptake regulatory protein – PhoU* c	ABC-type branched-chain amino acid transport system, ATPase component - LivG*oa	Branched chain amino acid ABC transporter ATP binding protein - LivF*a	Hypothetical protein - UspA*°°	Trans-2-enoyl-ACP reductase II - FabK*°	Polysaccharide biosynthesis protein – EpsI ^c	Polysaccharide biosynthesis protein EpsF – RgpA/EpsF ^{oc}	Phosphate acyltransferase - PlsX*°°	ABC-type polar amino acid transport system, ATPase component - Peb1*°a	Amino acid ABC transporter periplasmic protein - GlnQ*o a, b	Peptidoglycan branched peptide synthesis protein, alanine adding enzyme – MurM °	UDP-N-acetylglucosamine-N-acetylmuramyl-(pentapeptide) pyrophosphoryl-undecaprenol N-acetylglucosamine transferase – MurG*°°	50S ribosomal protein L13 - RplM	50S ribosomal protein L2 - RplB	Glycine-tRNA ligase beta subunit - GlyS	Pyruvate kinase - Pyk*° c	Trigger factor - Tig	3-Oxoacyl-ACP synthase - FabF	Glutamate-tRNA ligase - GltX°	Cell wall biosynthesis glycosyltransferase - Ster1440*c	Polysaccharide biosynthesis protein - EpsN*°°	Polysaccharide biosynthesis protein - EpsD**	2,3-Bisphosphoglycerate-dependent phosphoglycerate mutase - Pgma*°	Methionine import ATP-binding protein - MetN ^a	4-Hydroxy-tetrahydrodipicolinate reductase - DapB*	4-Hydroxy-tetrahydrodipicolinate synthase - DapA*
Stu1005	Stu1006	Stu0362	Stu0363	Stu1637	Stu0385	-	Stu1472	Stu0028	Stul 161	Stu1162	Stu1155	Stu0732	Stu0093	Stu1931	Stu0507	Stu1196	Stu0132	Stu0388	Stu1814	Ster1440	-	Stul 109	Stu1204	Stu0301	Stu0424	Stu1297
116627797	55737024	116627277	55822340	55823541	116627296	24637447	90655845	122268399	116627887	116627888	13324647	122267800	55822083	81820219	122268006	161936368	122268307	116627299	67461637	116628179	24637452	24473738	81559295	81820355	81559752	122267371
28.0	24.9	27.9	25.5	16.9	33.6	38.2	44.5	35.5	28.1	30.9	46.2	40.2	16.2	29.9	74.4	54.5	46.7	43.5	55.3	27.1	39.0	27.5	26.2	38.7	27.7	33.8
3	2	4	2	8	2	11	14	4	12	4	15	3	3	4	4	7	10	4	2	4	7	4	26	8	10	2
16.67	9.63	14.57	14.83	56.00	11.21	26.52	42.31	12.28	44.71	15.16	40.10	11.52	18.24	19.86	6.34	18.40	26.23	13.17	4.34	21.43	23.71	22.22	74.78	23.10	52.16	7.07
144	88	92	68	217	5.978	386	400	58	491	103	549	91	86	101	101	229	217	189	42	143	204	108	782	235	438	73

Legend to Figures

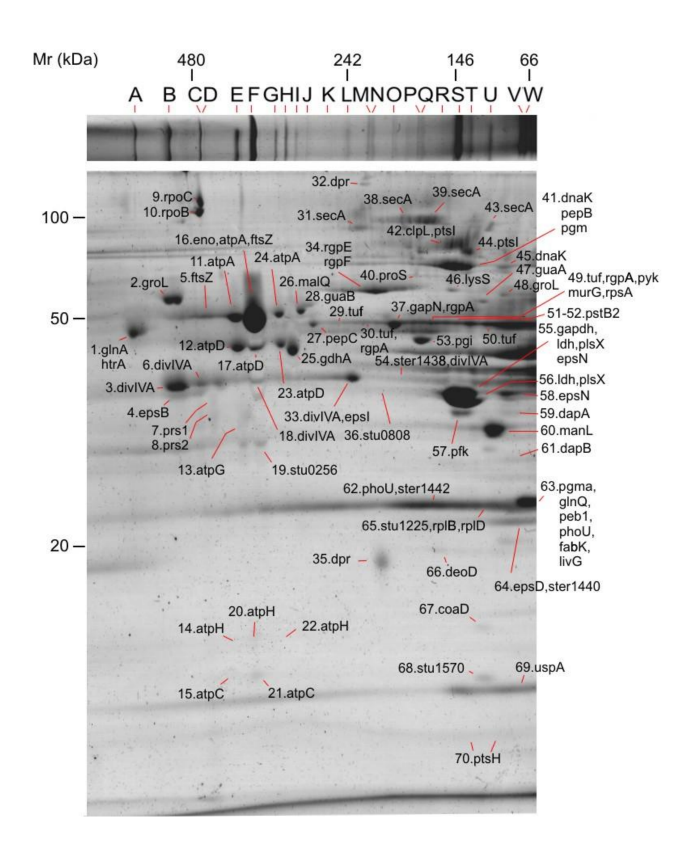
Figure 1. Top. 1D-BN-PAGE separation of membrane protein complexes from *Streptococcus thermophilus* strain DSM20617. About 100 μg of bacterial proteins were analyzed. Electrophoresis was performed on a gel casted with an acrylamide gradient of 4-14% T. Gels were stained by using a Coomassie blue-silver-based procedure. Commercially available molecular mass markers for the 1D-BN-PAGE are indicated at the top. Gel bands subjected to trypsinolysis and nLC-ESI-LIT-MS/MS analysis are indicated. Proteins identified by within each gel band are reported in Table 1 and Supplementary Table S1.

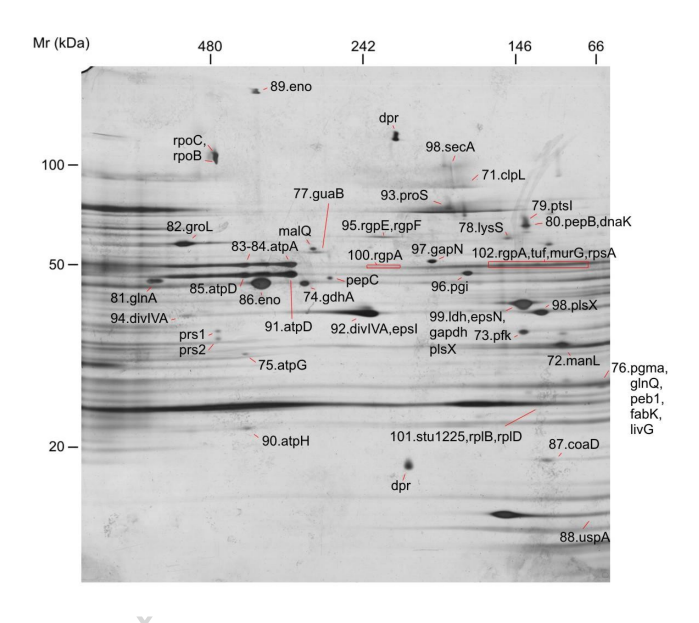
Bottom. 2D-BN/urea-PAGE separation of membrane protein complexes from *Streptococcus thermophilus* strain DSM20617. About 50 µg of bacterial proteins were analyzed. The first dimension (BN-PAGE) was performed on a gel casted with an acrylamide gradient of 4-14% T; the second dimension (urea-PAGE) was performed on a gel casted with 12% T acrylamide. Gels were stained as mentioned above. Molecular mass markers for 1D-BN-PAGE and urea-PAGE are indicated at the top and on the left, respectively. Proteins identified by nLC-ESI-LIT-MS/MS are reported; identification details are specified in Supplementary Table S2. Corresponding heteromeric and homomeric protein complexes identified by combining 1D-BN-PAGE, 2D-BN/urea-PAGE and 2D-BN/SDS-PAGE experiments, followed by nLC-ESI-LIT-MS/MS analysis, are shown in Table 1.

Figure 2. 2D-BN/SDS-PAGE separation of membrane protein complexes from *Streptococcus thermophilus* strain DSM20617. About 50 μg of bacterial proteins were analyzed. The first dimension (BN-PAGE) was performed on a gel casted with an acrylamide gradient of 7-14% T; the second dimension (SDS-PAGE) was performed on a gel casted with an acrylamide gradient of 9-16% T. Gels were stained by using a Coomassie blue-silver-based procedure. Molecular mass markers for 1D-BN-PAGE and urea-PAGE are indicated at the top and on the left, respectively. Proteins identified by nLC-ESI-LIT-MS/MS are indicated; identification details are reported in Supplementary Table S3. Corresponding heteromeric and homomeric protein complexes identified by combining 1D-BN-PAGE, 2D-BN/urea-PAGE and 2D-BN/SDS-PAGE, followed by nLC-ESI-LIT-MS/MS analysis are shown in Table 1.

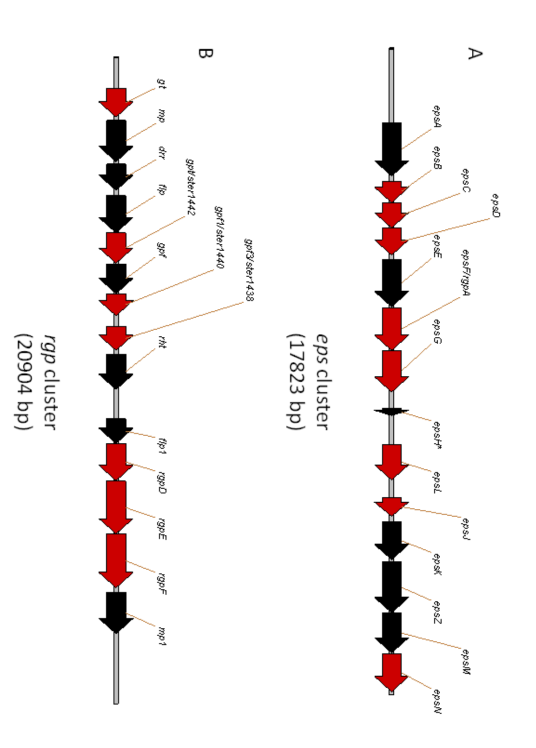
Figure 3. Schematic organization of the *eps* and *rgp* genes cluster in *S. thermophilus* DSM20617. Genes coding for proteins here identified in heteromultimeric and homomeric complexes are reported in red. Shown are genes and related product as deduced by BLAST analysis. Panel A. EPS cluster; epsA, cell envelope-related transcriptional attenuator; epsB, capsular polysaccharide biosynthesis protein; epsC, polysaccharide export protein, MPA1 family, G+ type; epsD, membrane-associated ATPase; epsE, galactosyl transferase; epsF/rgpA, rhamnosyl transferase; epsG, glycosyl transferase; epsH*, exopolysaccharide biosynthesis protein, truncated; *epsL*, β-glycosyltransferase; *epsJ*, glycosyltransferase; epsK, polysaccharide polymerase; epsZ, flippase, assisting in the membrane translocation of lipopolysaccharides; *epsM*, galactopyranose mutase; *epsN*, galactofuranose transferase. Asterisk indicates a truncated gene (*epsH*). Panel B. RGP cluster; *gt*, glucosyltransferase; mp, predicted membrane protein; drr, dTDP-4-dehydrorhamnose reductase; flp, flippase; CDP-glycosylpolyol phosphate:glycosylpolyol gtp/ster1442, glycosylpolyol phosphotransferase; *gpf*, β-1,3-glucosyltransferase; *gpf1/ster1440*, glycosyl transferase; gpf3/ster1438, glycosyl transferase; rht, α-D-GlcNAc-α-1,2-L-rhamnosyltransferase; flp1, rhamnose-containing polysaccharide translocation permease; rgpD, ABC transporter possibly involved in side chain formation of rhamnose-glucose polysaccharide; rgpE, glycosyltransferase possibly involved in side chain formation of rhamnose-glucose polysaccharide; rgpF, α -L-rha- α -1,3-L-rhamnosyltransferase; mp1, predicted membrane protein.

Figure 4. Microscopic analysis of *S. thermophilus* DSM20617. Panel A. White colonies for ropy *S. thermophilus* DSM20617 in ruthenium red milk agar plate. Panel B and C. Transmission electron micrographs showing *S. thermophilus* DSM20617 cells. The black arrows indicate the polysaccharide matrix present on the surface of bacterial cells. The white arrow indicates the bacterial cell wall.

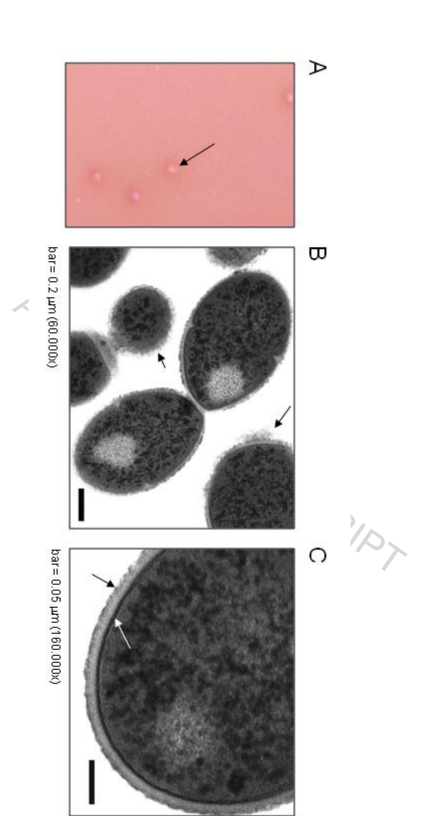




Salzano et al., Figure 2



Salzano *et al.*, Figure 3



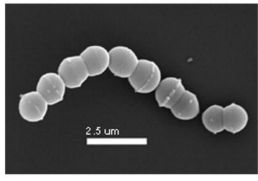
Mono-dimensional Blue Native-PAGE and bi-dimensional Blue Native/Urea- or /SDS-PAGE combined with nLC-ESI-LIT-MS/MS unveil membrane protein heteromeric and homomeric complexes in *Streptococcus thermophilus*

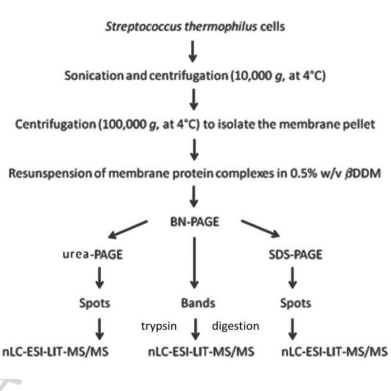
Anna Maria Salzano, Gianfranco Novi, Stefania Arioli, Silvia Corona,
Diego Mora and Andrea Scaloni

SUPPLEMENTARY MATERIAL

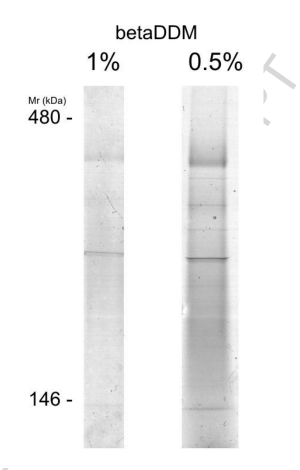
Supplementary Figure 1. A schematic representation of the most important experimental steps used in this study.

Supplementary Figure 2. 1D-BN-PAGE separation of membrane protein complexes from *Streptococcus thermophilus* strain DSM20617 as extracted with different concentrations of βDDM. Electrophoresis was performed on a gel casted with an acrylamide gradient of 5-12% T. Gels were stained by using a Coomassie blue-silver-based procedure. Commercially available molecular mass markers for the 1D-BN-PAGE are indicated on the left.





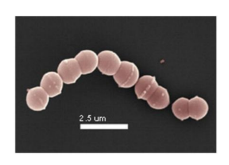
Salzano et al., Supplementary Figure S1

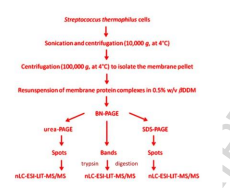


Salzano et al., Supplementary Figure S2

Biological significance

Combined proteomic procedures have been applied to the characterization of heteromultimeric and homomeric protein complexes from the membrane fraction of *Streptococcus thermophilus*. Protein machineries involved in polysaccharide biosynthesis, molecular uptake, energy metabolism, cell division, protein secretion, folding and chaperone activities were identified; information on hypothetical and moonlighting proteins were also derived. This study is original in the lactic bacteria context and may be considered as preliminary to a deeper functional characterization of the corresponding protein complexes. Due to the large use of *Streptococcus thermophilus* as a starter for dairy productions, the data reported here may facilitate future investigations on protein complexes assembly and composition under different experimental conditions or for bacterial strains having specific biotechnological applications.





Graphical abstract



1D-BN-PAGE, 2D-BN/urea-PAGE and 2D-BN/SDS-PAGE resolved membrane complexes of S. thermophilus

Sixty-five heteromeric and 30 homomeric complexes were characterized by nLC-ESI-LIT-MS/MS

Protein machineries involved in various molecular functions were identified

Information on moonlighting and hypothetical proteins was also derived