

Dairy streptococcal cell wall and exopolysaccharide genome diversity

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Abstract

The large-scale and high-intensity application of *Streptococcus thermophilus* species in milk fermentation processes is associated with a persistent threat of (bacterio)phage infection. Phage infection of starter cultures may cause inconsistent, slow or even failed fermentations with consequent diminished product quality and/or output. The phage life cycle commences with the recognition of, and binding to, a specific host-encoded and surface-exposed receptor, which in the case of *S. thermophilus* can be the rhamnose-glucose polysaccharide (RGP; specified by the *rgp* gene cluster) or exopolysaccharide (EPS; specified by the *eps* gene cluster). The genomic diversity of 23 *S. thermophilus* strains isolated from unpasteurized dairy products was evaluated, including a detailed analysis of the *rgp* and *eps* loci. In the present study, five novel *eps* genotypes were identified while variations of currently recognized *rgp* gene cluster types were also observed. Furthermore, the diversity of *rgp* genotypes amongst retrieved isolates positively correlated with phage diversity based on phageome analysis of eight representative dairy products. Our findings therefore substantially expand our knowledge on *S. thermophilus*' strain and phage diversity in (artisanal) dairy products and highlight the merit of phageome analysis of artisanal and traditional fermented foods as a sensitive marker of dominant microbiota involved in the fermentation.

DATA SUMMARY

Genome sequences used in this study have been deposited in National Center for Biotechnology Information (NCBI) and the accession numbers are listed in Table S1 (available in the online version of this article). The authors confirm all supporting data, code and protocols have been provided within the article or through supplementary data files. Supplementary Material can be found in Figshare: <https://doi.org/10.6084/m9.figshare.17080283> [1].

INTRODUCTION

Streptococcus thermophilus is one of the most widely exploited species of lactic acid bacteria (LAB) in dairy fermentations [2, 3]. *S. thermophilus* has been granted the generally regarded as safe (GRAS) status by the U.S. Food and Drug Administration [4, 5] and the 'Qualified Presumption of Safety' (QPS) status in the European Union [6]. Bacteriophage (or phage) infection poses a constant

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Abbreviations: CRISPR-Cas, clustered regularly interspaced short palindromic repeats and CRISPR-associated genes; EPS, exopolysaccharide; GRAS, generally regarded as safe; LAB, lactic acid bacteria; PHASTER, PHAge search tool enhanced release; QPS, qualified presumption of safety; RGP, rhamnose-glucose polysaccharide.

Accession number (strain/sample name): JAHBRN000000000 (Moz111); JAHBR000000000 (Brie16); JAHBRP000000000 (Moz86); JAHBRQ000000000 (Moz83); JAHBRI000000000 (Rico66); JAHBU000000000 (Brie28); JAHBRD000000000 (Moz76); JAHBEO000000000 (FDL19); CP075363 (Moz109); JAHBRJ000000000 (Vach57); JAHBRK000000000 (Vach60); JAHBRL000000000 (Rico65); JAHBRM000000000 (Strac48); JAHBUP000000000 (Nect1); JAHDUQ000000000 (Scam27); JAHBUR000000000 (Strac42); JAHBUS000000000 (Racle124); JAHBUT000000000 (Nect13); JAHBRF000000000 (FDL17); JAHDUU000000000 (Moz77); JAHDUV000000000 (Moz74); JAHBRG000000000 (Roque89); JAHRBH000000000 (Douc24); PRJNA731044 (Brie); PRJNA729649 (Mozzarella B); PRJNA731045 (Ricotta); PRJNA731053 (Straciatella); PRJNA731055 (Vacherin); PRJNA731056 (Semi-soft cheese D); PRJNA731057 (Semi-soft cheese A); PRJNA731046 (Blue cheese).

Data statement: All supporting data, code and protocols have been provided within the article or through supplementary data files. Three supplementary figures and five supplementary tables are available with the online version of this article.

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Impact Statement

Streptococcus thermophilus is one of the most important LAB species in dairy fermentations. This species, however, is subject to a persistent threat of phage infection, which may negatively affect the output and quality of the final fermentation product. It is known that *S. thermophilus* strains possess distinct saccharidic phage receptors that are linked to their phage sensitivity profiles. In the current study *S. thermophilus* strains were isolated from various unpasteurized dairy products, and the diversity of gene clusters involved in rhamnose glucose polysaccharide and exopolysaccharide biosynthesis was assessed. The distribution of dairy streptococcal phages in fermented dairy products was also assessed. The outcome of the present body of work provides insights for the dairy industry to assist in developing optimal and defined starter culture strain mixtures possessing distinct phage receptors/sensitivities, which in turn will minimize the risk of fermentation failure so as to ensure yield, consistency and quality of dairy products.

threat to industrial fermentations and can result in inconsistent, delayed or even failed fermentations and reduced product quality and output [7] with a considerable associated economic impact on the agri-food sector. Phages infecting *S. thermophilus* are currently differentiated into five groups termed the *cos* (or *Moineauvirus*) and *pac* (or *Brussowvirus*) [8], 5093 [9], 987 [10] and P738 [11] groups. Limited studies on the biodiversity of *S. thermophilus* phages have been undertaken [12–14], though phageome studies of cheese and cheese-associated (by)products have recently garnered increasing attention [15, 16]. To elucidate the phage–host interactome, it is essential to understand the biodiversity of both phages and their hosts that are applied in an industrial context, and of the phage- and host-encoded components that contribute to these interactions. Recently, protocols have been optimized for the extraction and analysis of phageomes relating to cheese and cheese wheys [15, 16]. Phageome analysis of fermented foods and their by-products provides useful insights into the prevalence, diversity and abundance of phages present in these foods, particularly where undefined starter culture mixtures are applied. While limited studies of the phageome of fermented dairy products have been reported to date, they are expected to provide a highly useful and complementary accompaniment to traditional phage-screening studies.

Phage infection commences with the recognition of, and binding to, a suitable host-encoded and surface-exposed receptor moiety. In many cases, the receptor moieties in Gram-positive ovococoid bacteria are saccharidic and include the rhamnose-glucose polysaccharide (RGP, whose biosynthetic machinery is encoded by the *rgp* cluster) [17] and exopolysaccharides (EPS, specified by the *eps* gene cluster) in *S. thermophilus* [18–20]. Currently, five *rgp* (types A to E) and six *eps* genotypes (types A to F) are known to exist based on comparative sequence analysis [20]. The chemical structure and/or composition of individual EPS and RGP molecules isolated from a small number of strains have been defined [18, 21, 22], findings that have highlighted the chemical diversity and complexity of these structures. Furthermore, a multiplex PCR system has been devised to reliably classify *S. thermophilus* strains based on their *rgp* genotypes which can, in principle, be applied to establish robust strain blends and/or rotations by incorporating strains with distinct RGPs [19, 20, 22]. Therefore, while it is widely accepted that *S. thermophilus* as a species lacks genetic diversity [23], the *rgp* and *eps* clusters appear to represent localized regions of genomic diversity, which affect its sensitivity to phage predation. In this context it should be noted that the host range of most dairy streptococcal phages is very narrow, an observation that may, at least partially, be linked to the diversity of phage receptors and the corresponding genetic diversity of their biosynthetic machinery.

Bacteria have evolved a variety of mechanisms to defend themselves against phage infection, thus bacteria and their associated phages are constantly engaged in an antagonistic co-evolutionary process [24]. *S. thermophilus* incorporates defence mechanisms to limit phage proliferation, through chromosomally- or plasmid-encoded R-M (restriction and modification) systems [25] and also CRISPR-Cas (clustered regularly interspaced short palindromic repeats and CRISPR-associated genes) systems [26]. Strains of this bacterial species are known to harbour up to four CRISPR-Cas systems (CR1 to CR4), with CR1 and CR3 being the most active in spacer acquisition [27, 28]. These systems have the capacity to evolve, adapt and acquire spacers in response to phage (or plasmid) exposure, which provides increased resistance to specific phages [29, 30]. The CRISPR spacer array profiles can be used as unique identifiers of distinct strains [31, 32]. A number of studies have highlighted the presence of dairy streptococci in artisanal fermented foods from a range of geographical locations through metagenomic and/or microbiological approaches [33–37]. Isolates have been characterized largely based on technologically relevant properties including EPS production and structural characteristics [37], antibiotic sensitivity, milk acidification ability and/or gelling capabilities [34]. Within an individual product, it can be challenging to assess if more than one strain of *S. thermophilus* is present due to high levels of sequence conservation across the genomes of strains of this species. CRISPR spacer array profiling is a useful tool to do so but this approach is time consuming and sequence analysis of the CRISPR arrays can be difficult due to the presence of repeats. The establishment of a multiplex PCR system for the rapid differentiation and classification of dairy streptococcal strains based on variable sequences within the *rgp* gene cluster of strains of this species facilitates the identification of strains with distinct phage-sensitivity characteristics. Since the role of RGPs in the initial binding stage of certain *S. thermophilus* phages is a relatively recent observation, it is perhaps unsurprising

that only a limited number of studies have focused on *rgp* gene cluster diversity of *S. thermophilus* strains, particularly when isolated from artisanal dairy products [20, 22].

In the present study, the diversity of *S. thermophilus* strains isolated from 27 dairy products derived from unpasteurized milk was ascertained through CRISPR spacer array analysis and *rgp* genotyping. Twenty three individual strains representing isolates from different food sources and possessing distinct *rgp* genotypes were selected for whole-genome sequencing facilitating a comparative analysis of the *rgp* and the *eps* clusters of the strains. *S. thermophilus* strains were classified based on defining criteria including the size of the genome, presence or absence of the cell-wall associated proteinase-encoding gene (*prtS*), CR2/CR3/CR4, also prophage(s), as proposed by Alexandraki et al. [38]. Furthermore, the phageomes of eight representative food samples were assessed to identify phage prevalence, abundance and diversity, in relation to the diversity and prevalence of *S. thermophilus* isolates obtained from each respective dairy products. The results derived from this investigation provide insights into the plausible correlation between phage and strain diversity/distribution in dairy fermented products derived from unpasteurized milk.

METHODS

Isolation of presumptive *S. thermophilus*

Twenty seven fermented dairy products including raw cow's milk and soft, semi-soft, hard cheeses made with unpasteurized cow's, buffalo's, ewe's and goat's milk (Table 1) were screened for the presence of *S. thermophilus* strains. In total, 5 g of each product was transferred into 45 ml of PBS (Sigma Aldrich, MO, USA) and pummelling for 2 mins at 300 r.p.m. in a stomacher (Stomacher Circular 400; Seward, UK). Serial dilutions of each sample were prepared and plated on LM17 agar [M17 agar (Oxoid, Hampshire, UK) supplemented with 0.5% lactose (Sigma Aldrich)], incubated overnight at 42 °C anaerobically (Anaerocult A – Merck, NJ, USA). A total of 1253 individual colonies exhibiting a creamy-white colour were isolated. The isolates were maintained and stored at –80 °C in LM17 broth supplemented with 30% (v/v) glycerol (Thermo Fisher, MA, USA).

RGP genotyping by multiplex PCR

Presumptive *S. thermophilus* isolates were typed using an established multiplex PCR system based on the *rgp* gene cluster and using five primer pairs (Table S2). PCRs were performed using Phusion Green Hot Start II High-Fidelity PCR Master Mix (Thermo Fisher, Gloucester, UK) employing the following conditions: 98 °C for 10 min followed by 30 cycles of 98 °C for 15 s, 55 °C for 30 s, and 72 °C for 1 min, followed by a final extension step at 72 °C for 10 min. All PCRs were performed using an Applied Biosystems 2720 Thermal Cycler (Thermo Fisher) instrument. Amplicons were visualized on a 1% agarose gel followed by UV transillumination.

Species confirmation using 16S rRNA amplicon sequence analysis

To confirm the identity of presumptive streptococcal isolates identified by the *rgp* genotyping approach, the 16S rRNA gene of presumptive *S. thermophilus* isolates was amplified using the *LucFw* and *LucRv* primers (Table S2) using *Taq* DNA polymerase mastermix (Qiagen, Manchester, UK) under the following conditions: initial denaturation at 94 °C for 10 min, 30 cycles of 94 °C for 30 s, 40 °C for 30 s, 72 °C for 1 min and 30 s, followed by a final extension at 72 °C for 10 min. The amplicons were purified using the GenElute PCR Clean-Up Kit (Sigma Aldrich) according to the manufacturer's instruction and subjected to Sanger sequencing (executed by Eurofins MWG, Waterford, Ireland). The generated sequences were analysed using BLASTN analysis against available sequence data on the National Center for Biotechnology Information (NCBI) database located at the following URL: <https://blast.ncbi.nlm.nih.gov/Blast.cgi>.

DNA extraction, genome sequencing, assembly and annotations

Bacterial DNA was extracted from a fresh 10 ml overnight culture using the Invitrogen PureLink Genomic DNA Mini Kit (Thermo Fisher) according to the manufacturer's instruction with some modifications. The cell pellet was resuspended and incubated in TE buffer containing 25% sucrose (Thermo Fisher) and 30 mg ml⁻¹ lysozyme (Sigma Aldrich). Chromosomal DNA was extracted from each strain and sequenced by Probiogenomics (Parma, Italy) using an Illumina MiSeq platform. Genome libraries were constructed using the TruSeq DNA PCR-Free LT Kit (Illumina) and 2.5 µg of genomic DNA, which was fragmented with a Bioruptor NGS ultrasonicator (Diagenode, USA) followed by size evaluation using Tape Station 2200 (Agilent Technologies, Santa Clara, CA, USA). Library samples were loaded into a Flow Cell V3 600 cycles (Illumina) and sequencing was performed on a MiSeq genomic platform (Illumina, Cambridge, UK) at GenProbio srl (Parma, Italy). Fastq files of the paired-end reads obtained from genome sequencing were used as input for genome assemblies through the MEGAnnotator pipeline in default mode [39]. Open reading frames prediction was performed with Prodigal v2.6 [40]. The MIRA programme v4.0.2 was used for *de novo* assembly of genome sequence data [41]. Following final genome assembly, putative protein-encoding genes were identified using the prediction software Prodigal v2.0 [40]. Protein-encoding genes were automatically annotated using a BLASTP v2.2.26 (cut-off E-value of 0.0001) sequence alignments against the non-redundant protein (nr) database curated by NCBI (<ftp://ftp.ncbi>).

Table 1. Number of isolates of presumptive streptococcal isolates identified by *rgp* typing multiplex PCR ($n=325$). Shaded areas represent a negative result

Products	Product type	Origin	No. of streptococcal isolates*	RGP†		
				A	B	C/E
Raw milk A to D	Raw milk	Cow				
Mozzarella A	Soft cheese	Cow	25	1	2	22
Mozzarella B		Buffalo	45	11	1	33
Soft cheese A	Soft cheese	Goat	1		1	
Soft cheese B		Goat	8			8
Soft cheese C		Goat	4			4
Brie	Semi-soft cheese	Cow	18	3	15	
Camembert		Cow				
Vacherin		Cow	3		3	
Reblochon A		Cow				
Reblochon B		Cow	12		11	1
Stracciatella		Cow	5		4	1
Ricotta		Buffalo	32	1	1	30
Scamorza		Cow	92			92
Cheddar		Cow	25			25
Halloumi		Cow				
Blue cheese	Hard cheese	Sheep	10			10
Semi-soft cheese A		Cow	1		1	
Semi-soft cheese B		Cow	8			8
Semi-soft cheese C		Cow	4			4
Semi-soft cheese D		Cow	32			32
Pecorino		Ewe				
Caciocavallo		Cow				
Hard cheese A	Cow					

*Approximately 48 to 96 creamy, white colonies were isolated from every dairy products, except for raw milk A to D, vacherin, stracciatella and halloumi due to low yield of c.f.u. in the samples.

†No RGP type D isolates were retrieved in this study.

nih.gov/blast/db/). The sequences of the *S. thermophilus* strains sequenced in this study were deposited in the GenBank database and their associated GenBank accession numbers are presented in Table S1.

Genome and comparative analysis

The quality or completeness of genome assemblies was evaluated with the microbial genomes atlas (MiGA) webserver [42]. The genome sequences of each strain was analysed using CRISPRFinder (<https://crispr.i2bc.paris-saclay.fr/Server/>) to retrieve and identify CRISPR repeats and spacer sequences and to establish if each strain was distinct based on unique spacer content within the identified CRISPR arrays. Further analysis was performed to compare the spacer sequences of each strain with phageome sequences of the associated dairy product using BLASTN analysis (<https://blast.ncbi.nlm.nih.gov/Blast.cgi>). The presence of *prtS* was identified in the genomes of the sequenced strains using BLASTN analysis (<https://blast.ncbi.nlm.nih.gov/Blast.cgi>). PHAge search tool enhanced release (PHASTER) [43, 44] was used to screen for prophage-specifying DNA regions within the genome of the *S. thermophilus* strains sequenced in this study. Intact prophages were manually checked to confirm the presence of

genes required to produce a functional phage particle including genes encoding proteins associated with replication functions, packaging, morphogenesis and lysis.

The *rgp* and *eps* gene clusters of *S. thermophilus* strains were identified. The *rgp* cluster was located between two conserved genes, encoding the 30S ribosomal protein and Rex protein, whereas the *eps* cluster was located between two conserved genes, encoding for chloride channel/purine nucleoside phosphorylase and VanZ. To determine and classify the *rgp* and *eps* genotype, the nucleotide sequence of the respective gene clusters of the isolates were compared using BLASTN analysis (<https://blast.ncbi.nlm.nih.gov/Blast.cgi>), with a cut-off of 50% nucleotide identity and query coverage. BLASTP (<https://blast.ncbi.nlm.nih.gov/Blast.cgi>) alignment of the *rgp*- and *eps*-biosynthetic gene clusters were performed to identify homologous gene products/regions between the analysed strains. BLASTP and Pfam v.33.1 [45] were used to assign functions to the individual protein-encoding regions. To identify the closest relatives of the sequenced EPS-specifying genomic regions and to establish their phylogeny based on their overall deduced proteomic content (based on the presence/absence of identified protein families with at least 50% identity over 50% of the protein), the Markov clustering (MCL) algorithm was executed via the mclblastline pipeline v12-0678 as described by Enright *et al.* [46], followed by hierarchical clustering (HCL) analysis performed and viewed in the multi-experiment viewer (MeV) [47]. *rgp* and *eps* genotype groupings were assigned based on similarity to previously defined *rgp* and *eps* type strains defined by Szymczak *et al.* [20] and/or through the identification of distinct groups observed in the heatmap generated through the HCL analysis.

Phageome extraction & data analysis

Phageome extractions were performed on eight representative dairy products screened (brie, mozzarella B, stracciatella, vacherin, semi-soft cheese A, semi-soft cheese D and blue cheese) in this study to determine the presence of phage in the sample. The extraction method followed a combination of elements of previous studies by Dugat-Bony *et al.* [15], Muhammed *et al.* [16] and Milani *et al.* [48] with some modifications. In summary, a given cheese sample (5 g) was homogenized in 30 ml sodium citrate (20 g l⁻¹) solution and pummeling for 2 mins at 300 r.p.m. in a stomacher. Samples were centrifuged at 300 g for 10 mins to remove large aggregates, and subsequently the supernatant was centrifuged for 5000 g for 45 mins at 4 °C. The supernatant was collected and treated with 1 M NaCl (Sigma Aldrich) for 1 h at 4 °C. The supernatant was diluted (1 : 2) in cold SM buffer (10 mM CaCl₂, 100 mM NaCl, 10 mM MgSO₄, 50 mM Tris-HCl at pH 7.5) and the pH was adjusted to ~4.6. Samples were centrifuged at 28000 g for 15 mins, then subjected to double filtration (first with 0.45 µm, followed by 0.2 µm filter). Next, 10% (w/v) PEG 6000 was added to filtered supernatant, then incubated at 4 °C overnight and centrifuged at 15000 g for 15 mins. The pellets were collected and resuspended in 1 ml SM buffer. Samples were treated with DNase I (20 units ml⁻¹, Sigma Aldrich) for 15 mins at room temperature, and then inactivated by incubation at 75 °C for 10 mins. The phage DNA was extracted using Norgen Biotek Phage DNA isolation kit (Norgen Biotek, Thorold, Canada) according to the manufacturer's instructions.

DNA was fragmented to 550–650 bp using a BioRuptor machine (Diagenode, Belgium). Samples were prepared following the TruSeq Nano DNA Sample Preparation Guide (Part no. 15041110Rev.D). Sequencing was performed using an Illumina NextSeq 500 sequencer with NextSeq Mid Output v2 Kit chemicals (Illumina, San Diego, CA 92122, USA). Read mapping of overall raw reads of the eight representative dairy products were performed using the METAnnotatorX bioinformatic platform as described by Milani *et al.* [48] against the NCBI database. Furthermore, read mapping of targeted phage genomes (of 489 streptococcal and lactococcal phages) was performed using MobaXterm server (<https://mobaxterm.mobatek.net/>). Raw reads were quality filtered using IlluQC.pl from the NGS QC Toolkit (v2.3) [49] in order to remove low-quality reads. The overall filtering process yielded 1.05 to 4.75 million reads per sample (Table S3). Quality-filtered reads were mapped for each sample against a panel of in-house database containing published and unpublished streptococcal/lactococcal phage (489 phages in total), using Bowtie2 aligner with default values [50]. Accession numbers for all samples analysed are listed in Table S1.

Phage screening and phage–host range determination

Eight of the cheese samples (brie, mozzarella B, ricotta, stracciatella, vacherin, semi-soft cheese A, semi-soft cheese D and blue cheese) were screened for the presence of phages against isolates emanating from the same product. Briefly, individual isolates of *S. thermophilus* obtained from a given sample were grown in 96-well microtitre plates containing 100 µl LM17 broth, incubated at 42 °C anaerobically for 24 h. The resulting culture was used to inoculate [1% (v/v)] two 96-well plates (one plate served as an untreated control) containing 100 µl LM17 broth. Plates were then incubated at 42 °C anaerobically for 1 h. Then, 10 µl of 1M CaCl₂ was added to all isolates in both 96-well plates, after which 50 µl of a filter-sterilized cheese homogenate from the same food sample was added to each well in a 96-well plate. Both microtitre plates were incubated at 42 °C anaerobically overnight.

Individual phages from a streptococcal phage collection (30 phages) (Table S4) were propagated to a high titre (>10⁷ p.f.u. ml⁻¹) on suitable sensitive hosts, and were applied in a host-range survey against the streptococcal strains whose genomes were sequenced as part of this study. Briefly, 10 µl of each phage lysate was spotted against isolates collected in this study via the double layer LM17 agar method, as previously described [51], then the plates were incubated anaerobically for 24 h at 42 °C.

RESULTS AND DISCUSSION

Artisanal dairy products contain multiple *S. thermophilus* strains

From a genomic perspective *S. thermophilus* has long been regarded as a relatively homogenous species with limited genetic variation [38]. However, certain genomic loci allow discernment between distinct strains and permit the identification of specific genotypes within the species. Among these are the loci encoding the biosynthetic apparatus responsible for RGP and EPS production. The present study aimed to screen for and isolate *S. thermophilus*, utilizing the widely accepted approach of using LM17 agar incubated at 42 °C anaerobically. A total of 1253 putative *S. thermophilus* isolates were selected, in order to (i) identify *S. thermophilus* strains associated with 27 unpasteurized dairy products, and (ii) determine the extent of diversity of the *rgp* and *eps* gene clusters (where *eps* clusters can be retrieved from the genome sequence) of the identified *S. thermophilus* strains as an indicator of strain diversity (Fig. S1). No putative *S. thermophilus* isolates were retrieved from 10 of the 27 analysed food samples including four raw milk samples and six cheese samples. From a further eight samples less than ten putative *S. thermophilus* isolates per sample were obtained, while the remaining nine cheese samples possessed more than 10 viable presumptive dairy streptococci (Table 1). Although the viability of *S. thermophilus* was generally observed to be low, the highest counts were observed among the soft and semi-soft cheeses that had not been matured (Table 1).

All isolates were first evaluated by multiplex PCR and for consistency with current literature, the *rgp* genotype classification (based on the PCR system of Kouwen *et al.*, 2019 [22]) was harmonized with that of Szymczak and colleagues [20] throughout this manuscript (where RGP 1=type B; RGP 2=type A; RGP 3=type D; RGP 4=type C or E, according to Kouwen *et al.* [22], and Szymczak *et al.* [20], respectively). Based on this genotyping approach, among the 1253 putative *S. thermophilus* isolates only 325 isolates were confirmed to belong to this species based on the RGP multiplex PCR (Fig. S1). Fifty six representative isolates among the 325 streptococcal isolates were further validated as *S. thermophilus* based on 16S rRNA gene sequence analysis. Of these 325 confirmed streptococcal isolates, the vast majority (83.1% or 270/325) were classified as RGP type C/E, 12% (39/325) as RGP type B, and 4.9% (16/325) as RGP type A (Table 1). This is consistent with a previous study in which type C strains were observed to be the most prevalent (34.8% or 8/23) [20]. The remaining 928 isolates did not produce any dairy streptococcal RGP-related amplicons suggesting that they are not dairy streptococci. In total, 171 isolates were eliminated from further investigation as they were identified as enterococcal species based on their ability to form brown precipitate around the colonies using bile aesculin agar [52]. Forty seven random isolates out of the remaining 757 isolates were evaluated by 16S rRNA gene sequencing and were identified as non-dairy streptococci, lactococci or pediococci (Fig. S1), thus the remaining isolates were discarded as unlikely *S. thermophilus* isolates and were not further characterized. Therefore, while the isolation procedure was selective for lactic acid bacteria it was not specifically selective for *S. thermophilus*.

The *rgp* genotype of strains within each food sample was evaluated to determine if single or multiple strains were likely present in the samples. Through this approach, the scamorza, cheddar, vacherin, blue cheese, soft cheese A to C, and semi-soft cheese A to D samples were observed to harbour isolates of a single dominant *rgp* genotype (Table 1). In contrast, the brie, reblochon B, stracciatella, ricotta and mozzarella A and B samples were observed to harbour strains possessing distinct *rgp* genotypes, highlighting the likely presence of multiple *S. thermophilus* strains in these cheeses. The presence of multiple dairy streptococcal strains in a given dairy product is indicative of thermophilic production regimes where *S. thermophilus* isolates would thrive. Among the analysed samples, there was a prevalence of RGP type C/ E *S. thermophilus* strains (being found in 11 dairy products), while type B strains were prevalent in six cheese products (soft cheese A, brie, vacherin, reblochon B, stracciatella, semi-soft cheese A). The mammalian origin of the milk (i.e. cow, goat, buffalo and ewe) did not appear to correlate with the presence of strains of a specific *rgp* genotype(s) (Table 1).

Comparative genome analysis of *S. thermophilus*

One to two representative isolates with distinct RGP profiles derived from each of 17 products that yielded *S. thermophilus* isolates shown in Table 1 (32 isolates in total) were selected for genome sequencing. Preliminary analysis of the whole genome and CRISPR spacer arrays confirmed that 23 of the 32 strains were distinct (Table 2) and the remaining nine isolates were eliminated from downstream analyses. Since the genome sequence data was typically assembled into multiple contigs, the genomes were checked for completeness based on the presence of 106 essential genes using the webserver MiGA and all strains were predicted to be (100%) complete and of 'excellent' quality (>95%). The genome sizes ranged from 1.78 to 2.05 mega base-pairs (Mbp), with average mol % GC content of 38.63–39.04%, in keeping with the genomes of previously analysed strains of the species.

The presence of *prtS* is associated with rapid growth in milk and is typically prevalent in industrial strains, which indicates lateral gene transfer in *S. thermophilus* population [38, 53] and was identified in 12/23 (or 52.2%) of the genomes analysed in this study. Two major clusters of *S. thermophilus* genotypes were proposed by Alexandraki *et al.* [38] in which the genomes of cluster A strains were observed to possess larger genomes (>1.83 Mbp), harbour *prtS*, CRISPR systems CR2, CR3 and CR4 and in which cluster B genomes are typically smaller (<1.83 Mbp), do not harbour *prtS* and often do not harbour CR2, CR3 and CR4. Based on these criteria seven strains (Brie1, Vach57, Vach60, Brie16, Brie28, Scam27, FDL17) analysed in the present study belong to cluster B, while none of the other genomes fulfil all criteria of the proposed cluster A genotype strains. In this study, all strains possessing

Table 2. General genome features of *S. thermophilus* strains sequenced in this study

Strain	Average coverage	No. of bases	Contigs	No. of predicted ORFs	Average GC %	Presence of <i>prfS</i> (Yes/No)	No. of prophage(s)*	Genome completeness (%)†
Moz109	180.8	1 807 378	33	1903	38.87	Yes	four inc.	100
Brie1	242	1 799 678	42	1904	38.93	No	one inc.	100
Vach57	252	1 815 322	41	1931	38.81	No	three inc.	100
Vach60	129.8	1 771 565	57	1877	38.91	No	two inc.	100
Rico65	438.2	1 779 950	39	1867	38.95	Yes	two inc.	100
Strac48	468.1	1 853 447	41	1954	38.81	No	one inc.	100
Moz111	137.8	1 820 649	22	1910	38.95	Yes	two inc.	100
Brie16	154	1 798 726	25	1891	38.87	No	two inc.	100
Moz83	164.5	1 753 946	22	1844	39.04	Yes	three inc.	100
Moz76	126.3	1 807 399	33	1910	38.91	Yes	two inc.	100
Rico66	342.5	1 801 536	34	1871	38.93	Yes	three inc.	100
Brie28	82.6	1 769 315	26	1870	38.90	No	two inc.	100
FDL19	247	1 805 448	29	1873	38.92	Yes	two inc.	100
Nect1	325	1 799 567	36	1896	38.93	Yes	three inc.	100
Scam27	259	1 787 154	27	1903	38.85	No	four inc.	100
FDL17	274	1 790 125	30	1895	38.84	No	four inc.	100
Strac42	251.2	1 796 889	50	1898	38.86	Yes	three inc.	100
Radle124	458.17	1 933 220	67	2048	38.78	No	one intact, two inc., one q.	100
Nect13	222	1 801 872	34	1892	38.92	Yes	two inc.	100
Moz77	191.4	1 786 575	27	1857	38.89	Yes	two inc.	100
Moz74	534.97	1 805 715	26	1898	38.91	Yes	four inc.	100
Roque89	425.4	1 969 840	72	2071	38.63	No	three inc.	100
Douc24	187.1	2 050 171	75	2167	38.81	No	three inc., two q.	100

*No. of prophage(s) predicted, inc., incomplete, and q., questionable.

†Genome completeness as calculated by MiGA webserver considering 106 essential genes.

Table 3. RGP and EPS type classification, number of CRISPR spacers, number of acquired CRISPR spacers compared to the corresponding phageome sequences and source of origin of 23 strains of *S. thermophilus*. Absence of spacers in a given CRISPR locus are shaded grey

Strain	RGP type*	EPS type†	No. of spacers in CRISPR locus				No. of CRISPR spacers that matched phageome sequence‡	Source of origin
			CR1	CR2	CR3	CR4		
Moz109	B	D	17	3	23		7	Mozzarella B_buffalo
Brie1	B	H	49	1	18		0	Brie_cow
Vach57	B	D	16	1	19		0	Vacherin_cow
Vach60	B	H	20	7	19		0	Vacherin_cow
Rico65	B	A	24	1			5	Ricotta_buffalo
Strac48	B	J	15	1	11		15	Straciatella_cow
Moz111	A	A	26	4	18		6	Mozzarella B_buffalo
Brie16	A	C ₂	41	1	34		1	Brie_cow
Moz83	A	C ₂	27	1	28		9	Mozzarella B_buffalo
Moz76	A	I	18	3	9		4	Mozzarella B_buffalo
Rico66	A	C ₁	62	3	20		20	Ricotta_buffalo
Brie28	A	F	30				1	Brie_cow
FDL19	A	C ₁	45	3	20		N/A	Mozzarella A_cow
Nect1	C	J	35	1	26		28	Semi-soft cheese D_cow
Scam27	C	C ₁	19	1	18		N/A	Scamorza_cow
FDL17	C	C ₁	18	1	28		N/A	Mozzarella A_cow
Strac42	C	G	24	11	25		3	Straciatella_cow
Racle124	C	A	18		3		0	Semi-soft cheese A_cow
Nect13	C	D	42	3	19	12	11	Semi-soft cheese D_cow
Moz77	E	C ₁	56	3	29		9	Mozzarella B_buffalo
Moz74	E	C ₂	12	1	6		3	Mozzarella B_buffalo
Roque89	E	C ₁	44	5	36		1	Blue cheese_sheep
Douc24	E	C ₁	63				N/A	Reblochon B_cow

*RGP typing based on multiplex PCR, following classification by Kouwen *et al.* [22].

†Predicted RGP and EPS type followed classification by Szymczak *et al.* [20].

‡Number of spacers with a hit in phageome sequences. N/A means the food sample was not sequenced and analysed for presence of phage genome.

the larger genomes (Strac48, Racle124, Roque89, Douc24) do not possess *prtS* (Table 2). Two strains (Brie28 and Douc24) harbour only CR1 in the genome with absence of *prtS*, nonetheless only Brie28 fits the requirement for cluster B as Douc24 has a relatively large genome (2.05 Mbp) (Tables 2 and 3). Based on our findings, we suggest a genome size cut-off of >1.80 Mbp for cluster A and <1.80 Mbp for cluster B. With this proposed cut-off, the genomes of seven strains could be characterized as cluster A and six as cluster B genotypes. The presence of intact prophage(s) is considerably more rare amongst *S. thermophilus* [38] compared to other species associated with dairy products such as *Lactococcus lactis* [54]. Therefore, the finding that only one (of 23 or 4.3%) genome (Table 2) was predicted to harbour an intact prophage is consistent with the previous literature [55]. This prophage (in the genome of Racle124) has a predicted size of 50.8 kb and harbours genes that encode proteins that are associated with tail and capsid morphogenesis and DNA replication functions. BLASTN analysis of the predicted prophage of Racle124 revealed 93.66% sequence identity over 57% of the genome of the *Brussowvirus* CHPC1248, which was isolated from a French cheese sample [20]. Remarkably, 19 of the 23 analysed genomes harbour an identical prophage remnant of 10.7 kb, while four genomes harbour an identical prophage remnant region of 14.1 kb.

RGP- and EPS-encoding cluster diversity analysis

The *rgp* and *eps* clusters of each *S. thermophilus* strain's genome were identified and compared to establish the diversity of these loci among the isolates. The *rgp* cluster is located between two conserved genes: at the 5' end it is flanked by a gene encoding the 30S ribosomal subunit, while a gene encoding the Rex protein borders the 3' end of the *rgp* gene cluster (Fig. 1). RGP-specifying gene clusters are comprised of two regions: (i) the 3' portion, which is associated with biosynthesis of the rhamnan backbone component and which is believed to be covalently linked to and embedded in the peptidoglycan, and (ii) the 5' portion, which is involved in the biosynthesis of the variable side chain, which is attached to the rhamnan backbone and being exposed at the cell surface (Fig. 1). Comparative analysis of *rgp* gene clusters identified in the sequenced genomes highlighted a high level of intra-group conservation of the clusters (Fig. 1). However, minor modifications were observed among selected isolates. For example, the RGP A strain Brie28 harbours several glycosyltransferase-encoding genes with limited sequence homology to other A type isolates (Fig. 1). Similarly, the RGP type B and E strains Vach57/60 and Moz74, respectively, each harbour a unique glycosyltransferase-encoding gene, suggesting that these strains possess unique glycosidic components in their respective RGP glycan structures relative to members of their genotypic groups. Despite the overall conservation and the lack of apparently novel *rgp* genotypes observed amongst the isolated strains, these clusters exhibit minor modifications, which may have occurred through insertion/deletion events.

The *eps* cluster is located between conserved genes encoding chloride channel/purine nucleoside phosphorylase (on the 5' border) and VanZ (on the 3' flank) (Fig. 2). The *eps* cluster is present in each of the 23 assessed genomes and is often interjected by transposons likely contributing to the diversification of these clusters through recombination events (Fig. 2). Eleven distinct *eps* genotypes (Table 3, Fig. 2) were assigned based on HCL analysis, including type A to F based on a previous study by Szymczak *et al.* [20] (Fig. S2). Among the 23 identified *eps* clusters, seven strains (Moz77, Roque89, Douc 24, Scam27, FDL17, FDL19) were classified as type C. Furthermore, based on the observation that certain C-type strains possess a unique gene content while maintaining certain genes that are specific to C-type strains, two C sub-types were arbitrarily defined in this study (C₁ and C₂; Fig. 2). The *eps* loci of Racle124, Moz111 and Rico65 cluster among EPS type A strains; Nect13, Moz109 and Vach57 to D type; and Brie28 to F type (Table 3, Fig. 2). Additional *eps* genotypes were identified in this study in extension to those previously defined by Szymczak *et al.* [20]: Strac42 is proposed as a type G; Brie 1 and Vach60 as type H; Moz76 as type I; and Strac48 and Nect1 as type J (Table 3, Fig. 2). While *rgp* gene clusters show high levels of conservation, the identified *eps* clusters appear highly variable in terms of their genetic composition and locus size (Figs 1 and 2), a finding that is consistent with a previous report [18]. Intact *eps* gene clusters were aligned, revealing that overall the glycosyltransferase-encoding genes are significantly different between groups (<49% identity) and vary moderately between strains of the same groups (for instance, between Brie16 and Moz83, glycosyltransferase-encoding genes share 50–96% sequence identity, with the Moz83 cluster encoding one unique glycosyltransferase).

In agreement with Szymczak *et al.* [20], this study demonstrates that the RGP and EPS-specifying gene clusters harboured by *S. thermophilus* strains are not correlated highlighting the range of combinations of cell-wall polysaccharide biosynthetic gene clusters that may be present among strains of the species (Table 2). However, the one apparent exception to this was the observation that all RGP type E strains, including the reference strain N4L, possess EPS type C (Table 3), which was also observed in a study by Szymczak *et al.* [20] where all four strains (CHPC1042, CHPC1246, CHPC1247, CHPC1248) identified as *rgp* type E possessed *eps* type C. Understanding the diversity (and combinations) of host cell-wall (RGP and EPS) polysaccharide structures and their encoding gene clusters may facilitate predictions of phage sensitivity [56]. Furthermore, it was previously suggested that *S. thermophilus* genomes are homogenous with limited genetic variation [38], but the present study highlights the presence of specific loci that exhibit significant diversity [20]. Significant diversity was observed in the *eps* loci despite a relatively small number ($n=23$) of sequenced/analysed strains (Fig. 2, Table 3). This may represent a biological response to phage (or other external) pressure as a previous study demonstrated that mutations in the *eps* locus resulted in increased resistance to phages [18].

Phageome composition correlation to *S. thermophilus* strains' diversity

To evaluate if phages were present in the food samples that could infect the retrieved isolates, filtered cheese suspensions were tested against the isolates emanating from the respective cheese samples to ascertain if phage–host combinations could be identified. However, using a liquid testing system in 96-well microtitre plates, no such interactions were observed. The screened dairy products did not appear to contain phages that targeted the retrieved dairy streptococcal strains isolated from the corresponding dairy samples. Nonetheless, since it is widely accepted that phages are ubiquitous in food fermentations, it seems unlikely that the foods analysed in the study were devoid of phages. A panel of 30 phages from our in-house collection (listed in Table S4) was used to evaluate the phage robustness of the isolates. Among the 23 strains tested, three of the seven RGP A type strains were sensitive to phages in our collection. Phages SW11 (*cos*), SW6 (*cos*) and SW24 (5093) were able to form plaques on strains Brie28, Moz83 and Rico66, respectively (Table 4). The region encoding the *rgp* gene cluster in the genomes of strains Moz83 and Rico66 displays almost 100% sequence identity across the entire cluster, which suggests that the *cos* phage SW6 and the 5093 phage SW24 require distinct moieties since the RGP composition and structure is likely to be highly similar between these strains. Perhaps Moz83 (EPS C₁ type) and Rico66 (EPS C₂ type) displayed different sensitivity to phages as they possess different EPS genotypes, as it has previously been suggested that EPS represents the receptor for the 5093 phages [10]. The RGP B-type strain Moz109 was shown to be sensitive to infection by the *cos* phages STP1 and SW9, which is consistent with the finding that the deduced Bpp's of these two phages share 96% aa identity across their entire protein

S. thermophilus: RGP cluster

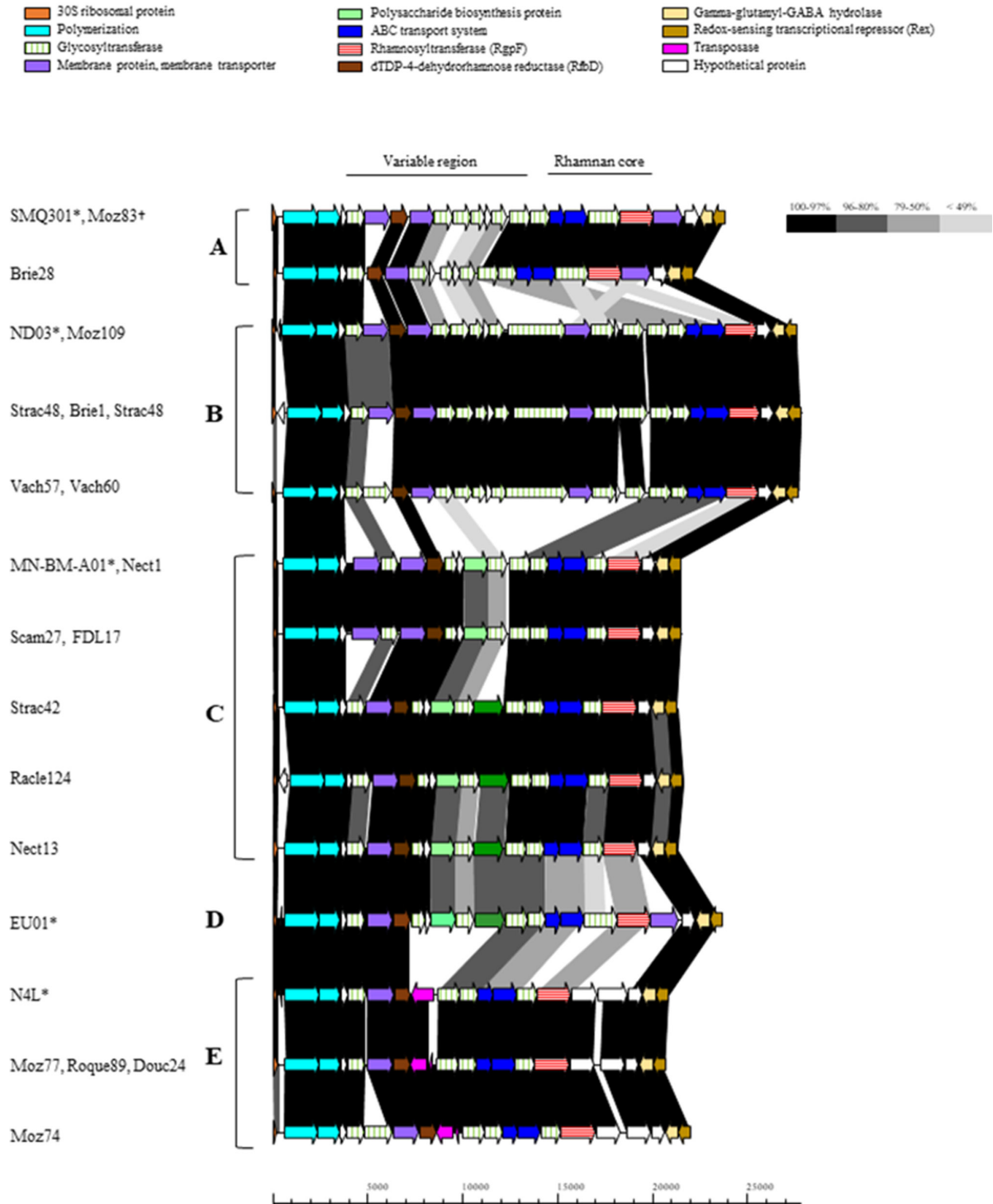


Fig. 1. Schematic overview of the organization and sequence relatedness the RGP-specifying gene clusters of *S. thermophilus* strains (A–E), including five reference strains of *S. thermophilus* (*) for each respective RGP type. Regions of homology (% amino acid identity) are joined by blocks of different shades of grey to black as indicated in the figure. The proposed functions of the individual protein-encoding regions are colour coded and indicated within the figure. Scale bar was measured in base pair (bp). † indicates where representative examples of identical gene clusters are presented; the RGP-encoding operons of Moz111, Moz76, Brie16, FDL19 and Rico66 are identical to SMQ-301 and Moz83.

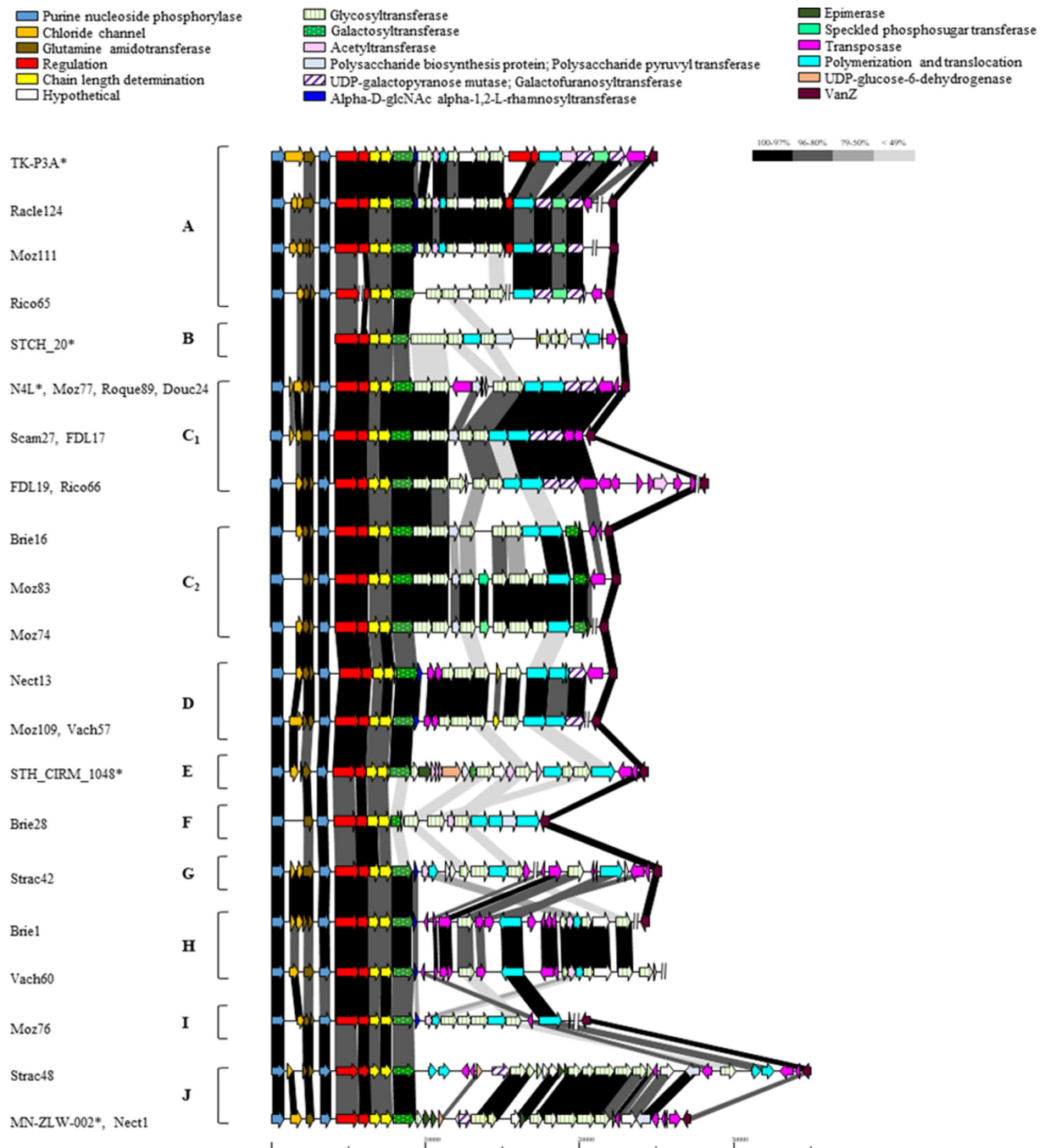


Fig. 2. Schematic overview of the organization and sequence relatedness the EPS-specifying gene clusters of *S. thermophilus* strains (type A to J), including five reference strains of *S. thermophilus* (*). Regions of homology (% amino acid identity) are joined by blocks of different shades of grey to black as indicated in the schematic. The proposed functions of the individual protein-encoding regions are colour coded and indicated above the figure. Scale bar is measured in base pair (bp). In the case where the *eps* cluster was retrieved from different contigs, break symbol (//) was used.

length. The RGP C-type strain Nect13 strain was shown to be sensitive to *cos* phage SW41 (Tables 3 and 4). This analysis confirms the narrow host range that is typical of dairy streptococcal phages and highlights that the majority (18 of 23 strains) of tested strains was insensitive to any of the assessed phages.

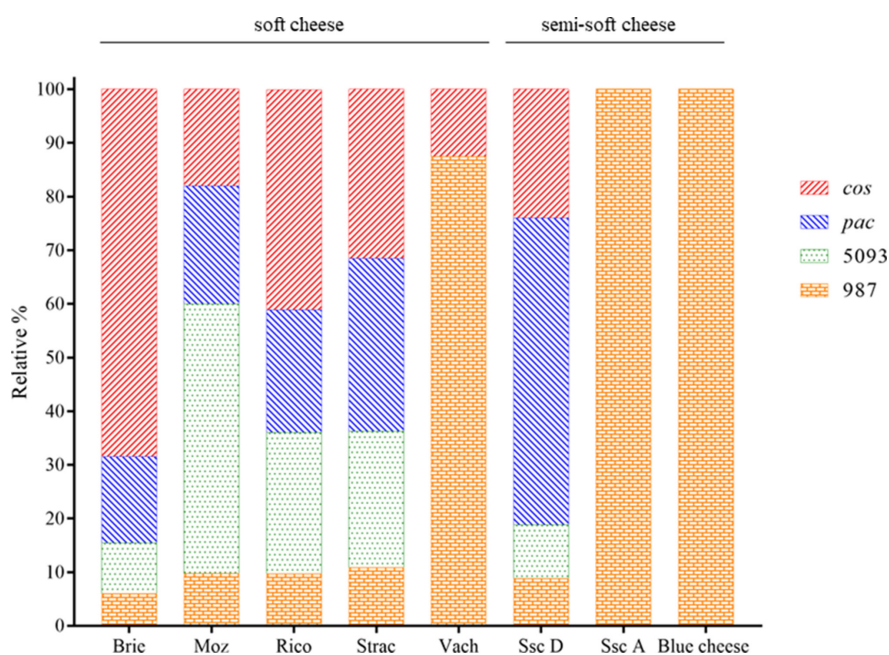
Given the observed diversity among the *rgp* and *eps* loci and their reported role as phage receptors, it was hypothesized that samples containing dairy streptococcal isolates would likely harbour streptococcal phages. To evaluate the presence of phages in the foods and to establish if any correlations could be made between the isolates and the harboured phages, eight representative dairy products (brie,

Table 4. Phage–host relationships identified among the *S. thermophilus* strains. Formation of plaques was shown as '+', whereas absence of plaques was shown as '-'

Strain (RGP type)	SW11 (<i>cos</i>)	SW6 (<i>cos</i>)	SW24 (5093)	SW9 (<i>cos</i>)	STP1 (<i>cos</i>)	SW41 (<i>cos</i>)
Brie28 (RGP A)	+	-	-	-	-	-
Moz83 (RGP A)	-	+	-	-	-	-
Rico66 (RGP A)	-	-	+	-	-	-
Moz109 (RGP B)	-	-	-	+	+	-
Nect13 (RGP C)	-	-	-	-	-	+

mozzarella B, ricotta, stracciatella, vacherin, semi-soft cheese D, semi-soft cheese A and blue cheese) containing strains with distinct *S. thermophilus* RGP profiles were selected for phageome sequencing and analysis. Microbiological analysis of vacherin, semi-soft cheese A and blue cheese yielded strains with limited *rgp* genotype diversity (Table 3). This correlates to a narrow streptococcal phage diversity in the corresponding phageome analysis with only 987 group phages present in semi-soft cheese A and blue cheese, and 987 and *cos* group phages present in vacherin (Fig. 3). Conversely, in brie, mozzarella B, ricotta, stracciatella and semi-soft cheese D, members of four streptococcal phage groups were present, while at least two to four distinct RGP types of *S. thermophilus* are present in these products (Table 3). Therefore, in general, samples possessing streptococci with diverse *rgp* genotypes are correlated with a higher number of phage groups being present in the food. Vacherin, stracciatella, blue and semi-soft cheese A yielded low numbers of viable dairy streptococci ($n \leq 10$ isolates), which may be due to prevalence of other species such as *L. lactis*. Analysis of the overall distribution of phages in the eight selected dairy products revealed a prevalence of lactococcal phages in four of the eight analysed foods (Table S5), i.e. vacherin, stracciatella, blue and semi-soft cheese A (Fig. S3).

Further analysis was conducted to investigate whether the strains had acquired CRISPR spacers that matched the phageome sequence of the respective dairy products they were isolated from. CRISPR has been described as an effective adaptive immunity acquired by bacteria through provision of sequence-specific interference against phages, within the case of *S. thermophilus* novel spacer acquisition occurring mostly in CR1 and CR3, and only rarely in CR4 [57]. Such patterns of CR1 and CR3 being the most active were also observed in this study and most strains, which had acquired spacers that matched respective phageome sequences, had the spacers located mainly in CR1 and CR3 (Table 3). Spacers of short DNA fragments are obtained from infecting phages, thereby providing immunity against subsequent exposure to the same phage [56]. However, Vach57, Vach60 (sourced from vacherin) and Racle124 (sourced from semi-soft cheese A) strains, do not appear to have spacer sequences that correlate to the respective phageome sequences, hence they

**Fig. 3.** Relative % of read mapping to streptococcal phage (*cos*, *pac*, 5093, 987) distribution in eight cheese samples – brie, mozzarella B (moz), ricotta (rico), stracciatella (strac), vacherin (vach), semi-soft cheese D (ssc D), semi-soft cheese A (ssc A) and blue cheese, based on phageome analysis.

do not appear to be recently evolved CRISPR-mediated bacteriophage insensitive mutants (BIMs) (Table 3). Their survival in the dairy products may be due to relative low abundance and less variation of streptococcal phage with only 987 being the dominant phage group in vacherin and semi-soft cheese A (Figs 3 and S3).

CONCLUSIONS

In the present study, we observed a prevalence of *S. thermophilus* strains among soft and semi-soft cheeses. Genome analysis based on RGP and EPS clusters, the presence of prophages and the component CRISPR arrays provided novel insights into strain diversity and the composition of dairy streptococci in dairy foods. Strains isolated in this study exhibited significant diversity within the EPS loci with four novel EPS genotypes identified (G to J) and a subgrouping of the C type EPS cluster genotype (C₁ and C₂) while subtle modifications were observed among the RGP clusters contributing to their diversification (type A to E). Microbiological isolation of surviving strains of *S. thermophilus* has permitted an evaluation of the diversity of strains that may be present in artisanal dairy products. Data generated in this study has provided insights for the dairy industry to follow nature's recipe in developing optimal and mixed defined starter cultures of strains with diverse RGP and EPS types thereby enhancing the production consistency and organoleptic properties of the product.

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Author contributions

Conceptualization, J.M.; data curation, E.P.; formal analysis, E.P., J.M., G.A.L.; funding acquisition, J.M, DvS.; investigation, E.P.; methodology, E.P., B.M.D.; project administration, J.M., DvS.; resources, J.M., DvS., M.V.; software, G.A.L.; supervision J.M., DvS., M.V; validation, E.P., G.A.L.; visualization, E.P.; writing – original draft, E.P.; writing – review and editing, J.M., DvS.

Conflicts of interest

The authors declare that there are no conflicts of interest.

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