Comparative genomics among members of the Streptococcus bovis | Streptococcus equinus complex

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Abstract

Background: Today, only one streptococcal species, i.e. Streptococcus thermophilus is recognized as food-grade. Interestingly, other streptococci like Streptococcus macedonicus and Streptococcus infantarius belonging to the Streptococcus bovis/Streptococcus equinus complex (SBSEC) are also found in food matrices. However, these species are phylogenetically related to Streptococcus gallolyticus and Streptococcus pasteurianus that have been linked to endocarditis, bacteremia and colon cancer.

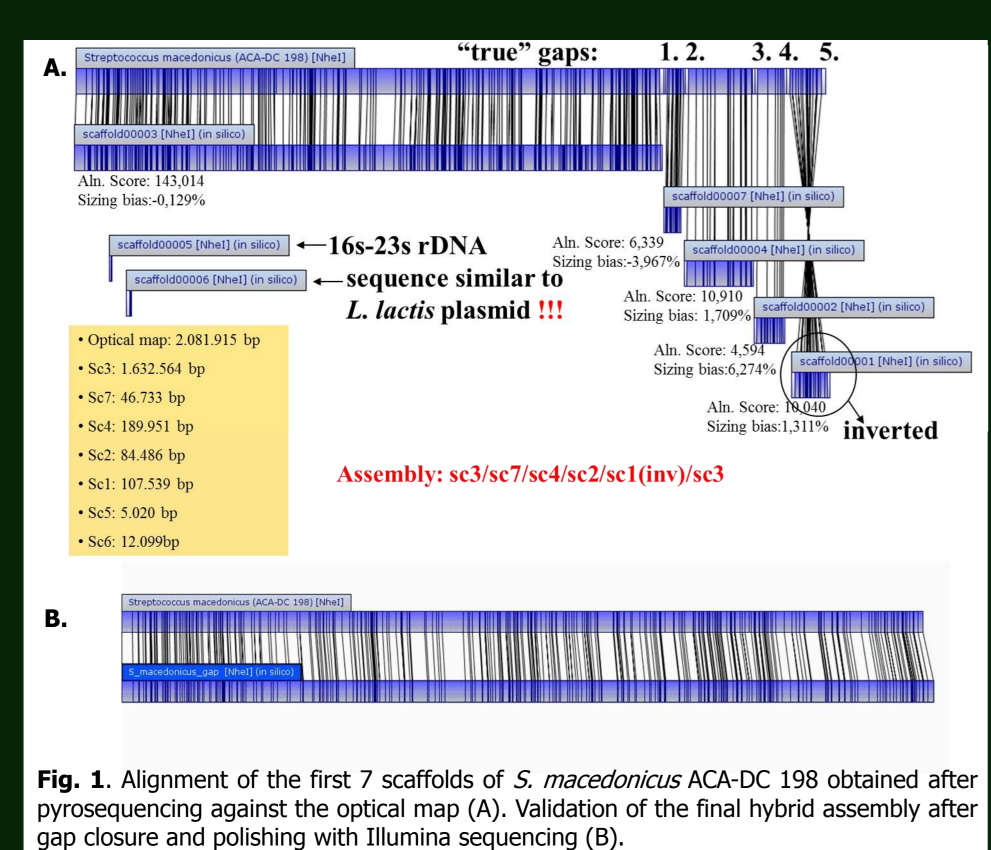
Objectives: To compare the available genomes of the members of the SBSEC in order to shed light onto their evolution and phylogenetic relation and to assess in silico their pathogenic potential.

Methods: Comparative genomics analysis including full chromosome and CDS alignments, whole genome phylogeny and evaluation of gene content (e.g. core genome, singletons, etc.) was performed with appropriate bioinformatics tools.

Conclusions: Despite the fact that the four species of the SBSEC were found tightly related based on whole genome phylogeny, there were two different patterns of evolution among them. Streptococcus pasteurianus, S. macedonicus and S. infantarius seem to have undergone a reductive evolution process that resulted in significantly diminished genome sizes and increased percentages of potential pseudogenes when compared to S. gallolyticus. In addition, S. pasteurianus, S. macedonicus and S. infantarius seem to have lost several genes previously linked to the ability of *S. gallolyticus* to survive in the gastrointestinal tract of herbivores and to its pathogenicity. Our findings indicate differences in the ecological niche and the pathogenic potential among the four species.

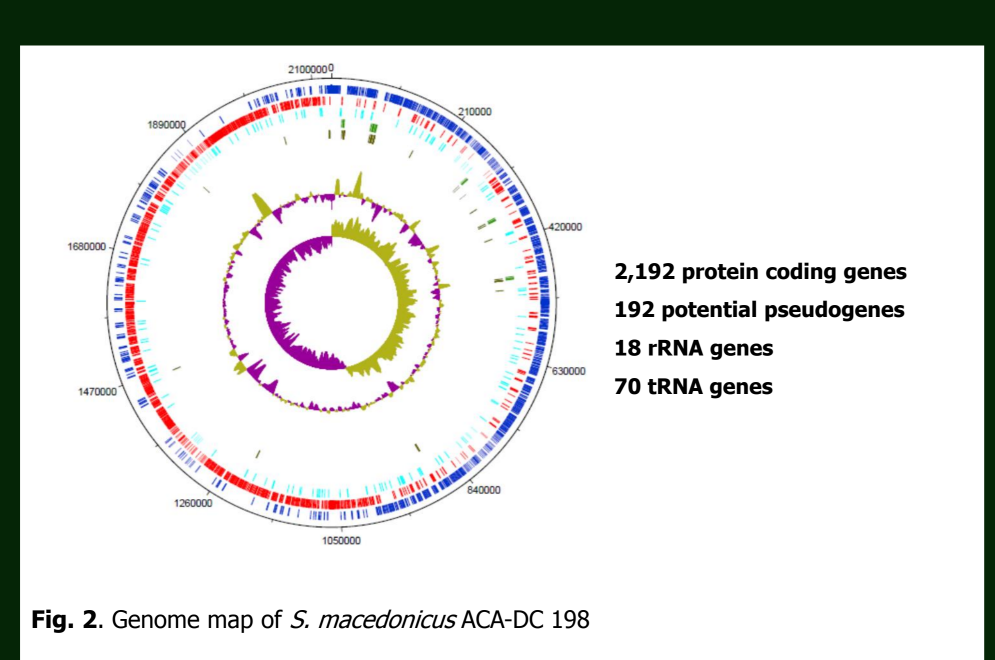
Sequencing the genome of *S. macedonicus* ACA-DC 198

- 1st step: shotgun pyrosequencing with 454 GS-FLX titanium (>100 contigs)
- 2nd step: 3kb paired-end pyrosequencing with 454 GS-FLX titanium (7 scaffolds)
- 3rd step: gap-closure and polishing with Illumina sequencing using the HiSeq 2000 (1 chromosome and 1 plasmid)
- 4th step: validation of the overall assembly (>200X coverage) with an *Nhe*I optical map



Annotating the genome of *S. macedonicus* ACA-DC 198

- 1st step: initial annotation was performed with the BaSys and the RAST pipelines
- 2nd step: annotations were manually compiled in one using Kodon software
- 3rd step: final corrections and quality assessment was performed using GenePRIMP (including predictions for potential



Comparative genomics of *S. macedonicus* **ACA-DC 198**

- The complete genome sequence of *S. macedonicus* offered new opportunities to investigate the properties of the species at the genomic scale
- The inclusion of *S. macedonicus* and *S. pasteurianus* as subspecies of *S. gallolyticus* has been previously suggested (Schlegel et al. Int J Syst Evol Microbiol. 2003), but this taxonomic reappraisal has not been formally accepted due to low DNA-DNA hybridization relatedness values (<70%) (Whiley et al. Int J Syst Evol Microbiol. 2003)

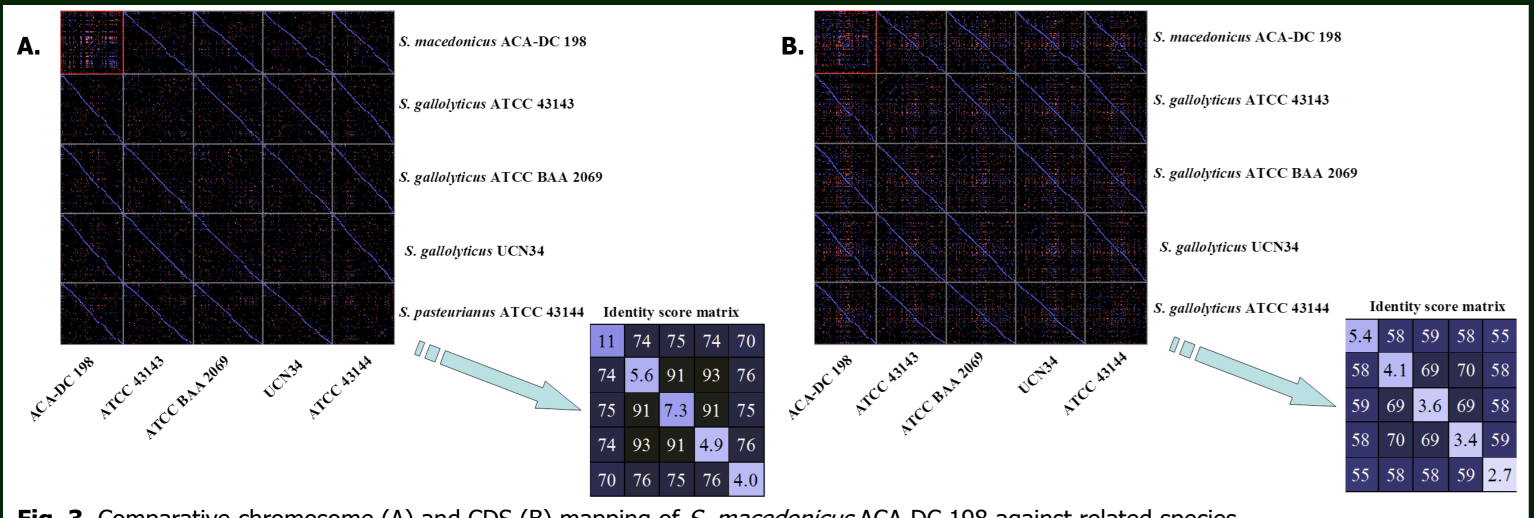


Fig. 3. Comparative chromosome (A) and CDS (B) mapping of *S. macedonicus* ACA-DC 198 against related species

Pairwise alignments of the chromosomes at the nucleotide or the CDS level revealed the degree of synteny between each pair (Fig. 3). The identity score at the nucleotide level of *S. macedonicus* against S. gallolyticus and S. pasteurianus was around 76% and 70%, respectively. Even more, the identity score at the CDS level dropped radically, reaching 58% in the case of *S. macedonicus* against S. gallolyticus and 55% in the case of S. macedonicus against S. pasteurianus. These values can not be used to directly determine the actual taxonomy of the three species. However, it is a fact that they are quiet low and they coincide with the low (≤ 70%) relatedness values of interspecies DNA−DNA hybridization experiments reported previously, reinforcing the notion that *S. macedonicus* and *S. gallolyticus* should remain separate species.

programme for development

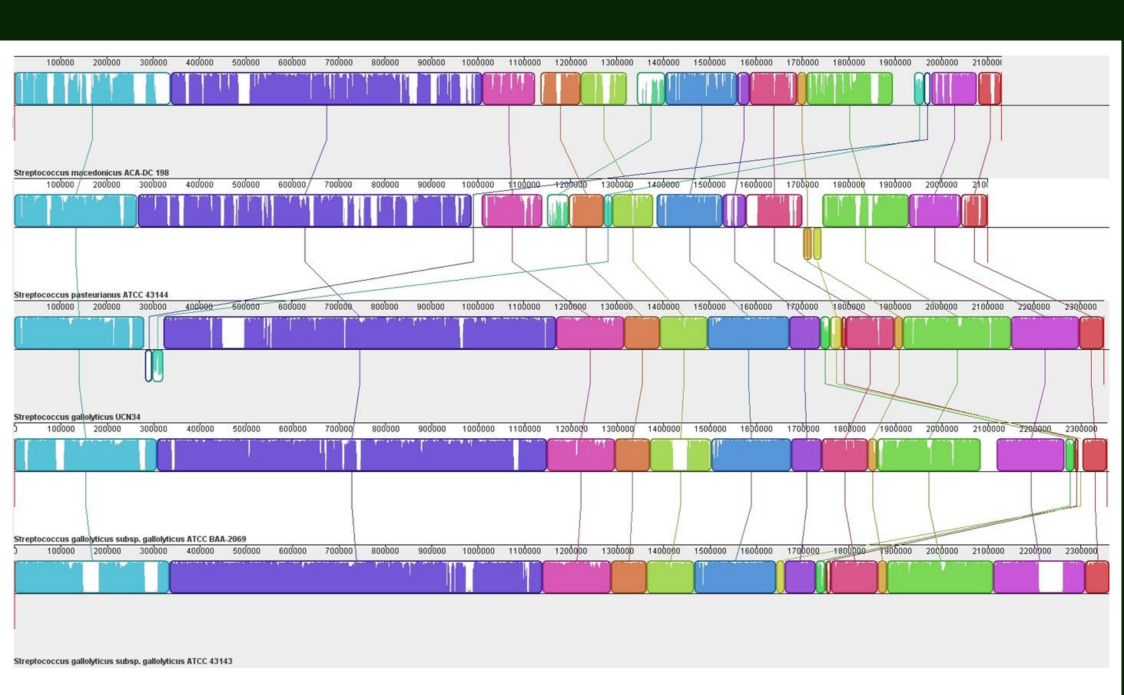
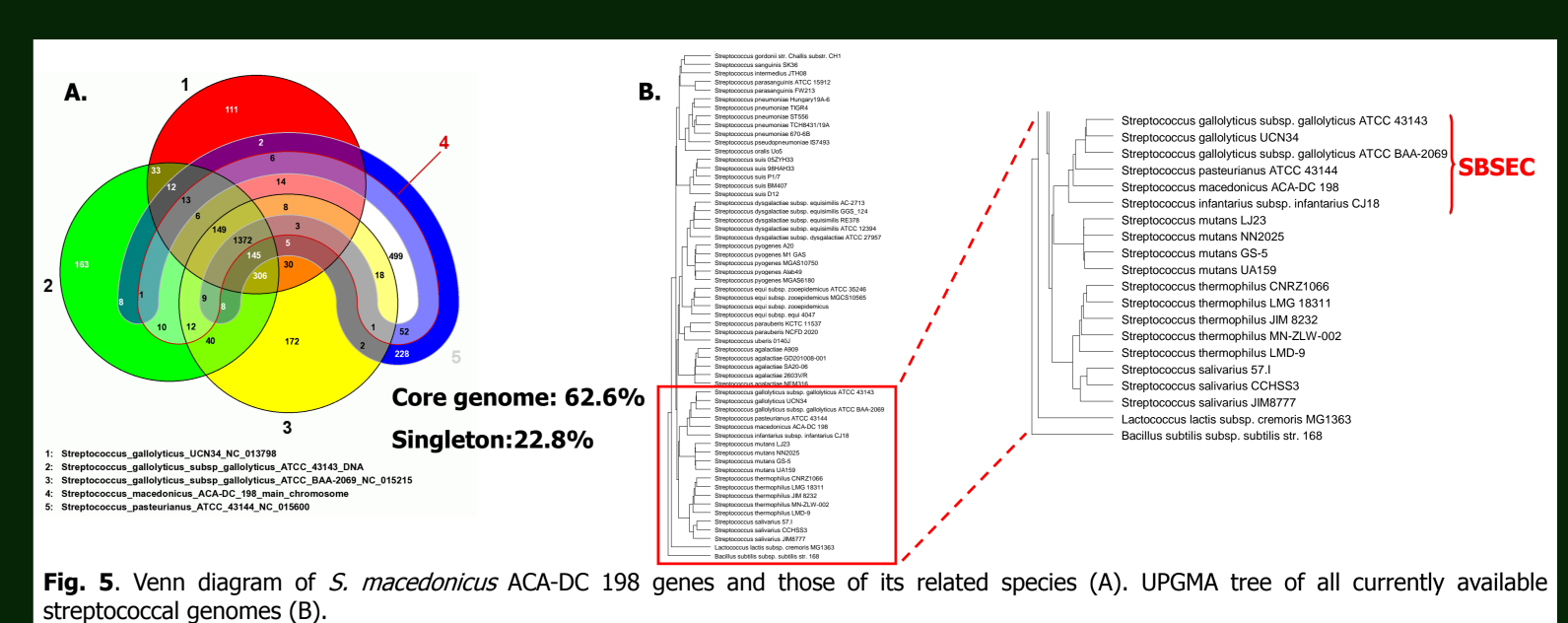


Fig. 4. Complete alignment of the S. macedonicus ACA-DC 198 genome against those of related species using local collinear blocks (LCBs)

Full chromosome alignments were performed using local collinear blocks (LCBs) among the three species. The analysis revealed a mosaic pattern of homology (Fig. 4). Evidently, a significant portion of the genetic information has been overall conserved, since the majority of the LCBs are shared by all species over most of their genome length. It should be noted that numerous strain-specific lifferences can also be detected. Furthermore, there are LCBs common only among some of the strains, while there are regions divergent enough so as not to be placed within a LCB. These findings indicate that, apart from gene loss through genome decay gene gain events like lateral gene transfer (LGT) must have played a role during the evolution of the three species. In addition chromosomal rearrangements seem to have been rather minimal, as the number of "movable" LCBs was low and their length was short. Inclusion of the S. infantarius genome in the analysis increased significantly the number of LCBs and reduced drastically the level of conservation among the genomes (data not shown), indicating that this particular genome is fairly different from the rest.



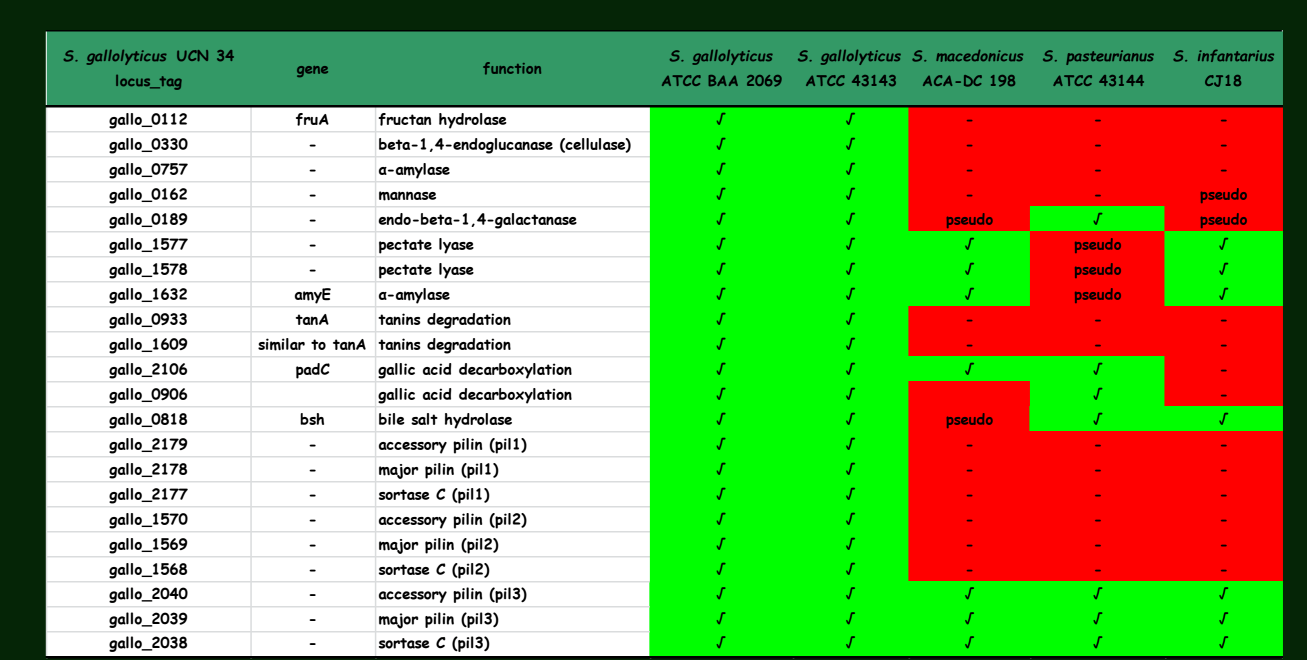
Reciprocal best Blast hits at the gene level also revealed a core genome of only 1,372 genes based on the sequence and the current annotation of the three species. This allows for a significant percentage of variable genes within the species that must have evolved during the adaptation to their specific environment. Still, S. macedonicus, S. gallolyticus, S. pasteurianus and S. infantarius form a single branch in the phylogenetic tree constructed based on the currently available complete streptococcal genome sequences providing extra evidence for the taxonomic integrity of the SBSEC

Additional characteristics of the genomes under investigation

Species	Genome size (Mb)	No. of protein coding genes	No. of potential pseudogenes/ (% percentage)
S. gallolyticus ATCC BAA 2069	2.35	2329	nr*/(nr)
S. gallolyticus ATCC 43143	2.36	2287	41(1.8)
S. gallolyticus UCN34	2.35	2251	28/(1.2)
S. macedonicus ACA-DC 198	2.13	2192	192/(8.7)
S. pasteurianus ATCC 43144	2.10	1869	157/(7.7)
S. infantarius CJ18	1.98	1964	nr/(4.6)
* not reported	1.70	1,01	1 / (1.0)

- S. macedonicus, S. pasteurianus and S. infantarius genomes are being shaped by selective pressures that favor extensive gene loss events and genome decay processes when compared to the S. gallolyticus genome
- · This property (i.e. genome decay) has been linked to the adaptation of bacteria to rich in nutrients environments as in the case of S. thermophilus adaptation to the milk environment

Niche-specific and pathogenicity genes presence/absence



Our findings clearly suggest that not only S. macedonicus, but also S. pasteurianus and S. infantarius have deviated from S. gallolyticus in their potential to catabolize complex plant carbohydrates and to cope with the harsh environment of the GI tract of herbivores. Furthermore, in silico analysis of S. gallolyticus has revealed that it contains three pilus gene clusters (pil1, pil2, pil3), which may mediate its binding to the extracellular matrix (ECM), although variations of pilus genes presence/absence within strains have also been reported. Each gene cluster consists of three genes. The first two genes encode two adhesins belonging to the MSCRAMM (microbial surface recognizing adhesive matrix molecules) family, one being the major and one being the minor (or accessory) pilus subunit. Pilus attachment to the peptidoglycan, as well as polymerization of adhesin filaments are catalyzed by a sortase C encoded by the third gene of the cluster. pil1 and pil2 loci are absent in S. macedonicus, S. pasteurianus and S. infantarius indicating a diminished tendency to adhere to ECM that could probably influence their ability to colonize host tissues and to produce infections when compared to *S. gallolyticus*.

Conclusions

- 1. S. macedonicus is most probably a separate species from S. gallolyticus
- 2. In silico analysis of S. macedonicus ACA-DC 198 suggests that:
- The strain is at the process of adapting to a rich in nutrients environment It shows a diminished capacity to live and survive in the GI tract of herbivores
- It has a diminished pathogenic potential compared to *S. gallolyticus*

• Papadimitriou K., S. Ferreira, N. C. Papandreou, E. Mavrogonatou, P. Supply, B. Pot, and E. Tsakalidou (2012) Complete genome sequence of the dairy isolate *Streptococcus macedonicus* ACA-DC 198. J Bacteriol 194:1838-9.

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