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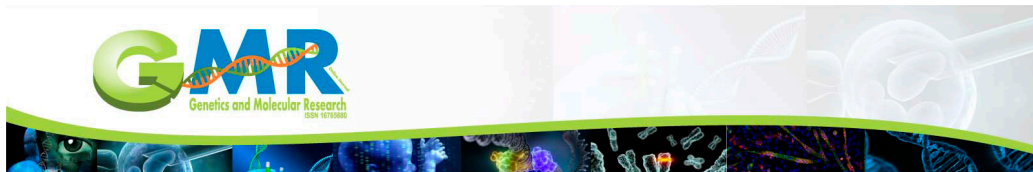


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Molecular phylogeny and a taxonomic proposal for the genus *Streptococcus*

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ABSTRACT. Alternative phylogenies for the genus *Streptococcus* have been proposed due to uncertainty about the among-species group relationships. Here, we performed a phylogenetic analysis of the genus *Streptococcus*, considering all the species groups and also the genomic data accumulated by other studies. Seventy-five species were subjected to a Bayesian phylogenetic analysis using sequences from eight genes (16S rRNA, *rpoB*, *sodA*, *tuf*, *rnpB*, *gyrB*, *dnaJ*, and *recN*). On the basis of our results, we propose a new Phylogeny for the genus, with special emphasis on the inter-species group level. This new phylogeny differs from those suggested previously. From topological and evolutionary distance criteria, we propose that *gordonii*, *pluranimalium*, and *sobrinus* should be considered as new species groups, in addition to the currently recognized groups of *mutans*, *bovis*, *pyogenic*, *suis*, *mitis*, and *salivarius*.

Key words: Bayesian analysis; Phylogeny; *Streptococcus*; Species group

INTRODUCTION

The genus *Streptococcus* is a diverse lineage belonging to the lactic acid group of bacteria. Current taxonomy places this genus, as well as the genus *Lactococcus*, within the family *Streptococcaceae* in the order Lactobacterales (Facklam, 2002). These organisms are Gram-positive, spherical, and catalase-negative, and many are facultative anaerobes (Montes and García-Arenzana, 2007; Lal et al., 2011).

Recent molecular systematic studies based on 16S rRNA sequences have provided evidence to subdivide *Streptococcus* into six “species groups”: pyogenes, anginosus, mitis, salivarius, bovis, and mutans (Kawamura et al., 1995; Facklam, 2002; Rodicio and Mendoza, 2004; Montes and García-Arenzana, 2007; Lal et al., 2011).

In addition to the striking phylogenetic diversification, the ecological features of the members of the *Streptococcus* genus encompass a myriad of environments and life styles. While some species are well known for their clinical importance as pathogens, others have been characterized as members of a range of normal microbiomes in different anatomical structures in animals and humans. This distribution pattern explains their regular occurrence in contaminated biological samples (Montes and García-Arenzana, 2007; Glazunova et al., 2010; Boggs et al., 2012).

The features of pathogenicity associated with the genus *Streptococcus* are also diverse: meningitis, pneumonia, endocarditis, fasciitis, and dental caries are among the better known conditions (Glazunova et al., 2010). Normal human reservoirs of *Streptococcus* include different compartments of the oral cavity and skin, and the respiratory, digestive, gastric, and urinary tracts (Hardie and Whiley, 1997). However, the complete home range for most species of the genus is largely uncertain since this knowledge depends on sampling strategies that are not normally focused on revealing species habitats.

Systematic studies have been performed in the genus. Kawamura et al. (1995) carried out a pioneer study that proposed the current general phylogenetic organization for the genus *Streptococcus*. In that study, 16S rRNA gene sequences from 28 distinct species were analyzed leading to a hypothesis of six species groups that is still generally accepted.

Simmon et al. (2008) examined the viridans group using 16S rRNA, *tuf*, and *rpoB* gene sequences from 22 species and subspecies to evaluate phylogenetic relationships within the genus in a study of samples from patients affected by endocarditis. They demonstrated horizontal gene transfer and different rates of molecular substitutions among DNA fragments in the genus, making phylogenetic inferences a challenging subject for this lineage.

Furthermore, other studies have evaluated the phylogenetic certainty of different gene fragments. For example, Täpp et al. (2003) analyzed the gene *rnpB* in 50 species, in order to obtain phylogenetic information from a source other than 16S rRNA. Similarly, Itoh et al. (2006) proposed the *dnaJ* and *gyrB* genes as good phylogenetic markers, and analyzed 45 species and subspecies. Additionally, the *recN* gene has been shown to render statistically confident clades in an extensive analysis that included 60 species (Glazunova et al., 2010).

To date, the phylogenetic relationships among the known species groups have not been resolved. For example, Itoh et al. (2006) concluded that disagreements between the trees obtained from three different gene fragments could be explained either by horizontal gene transfer or different mutation rates. This also can be seen in the phylogenetic results of Simmon et al. (2008) who described phylogenetic discordances that could be explained

by horizontal gene transfer. In both studies, the analyses were performed using phylogenetic distances; the effects of different phylogenetic methods were not evaluated. Glazunova et al. (2006) and Täpp et al. (2003) used diverse genes and different phylogenetic approaches (distances and Bayesian analysis, respectively) and produced two alternative phylogenetic hypotheses (Figure 1).

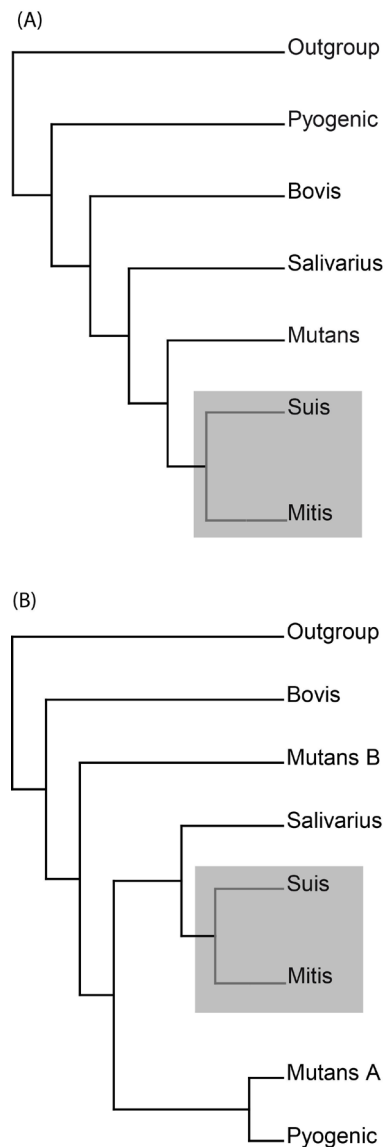


Figure 1. Phylogenetic hypothesis by (A) Glazunova et al. (2006, Figure 1), based on fragment sequences of *recN* gene; and (B) Täpp et al. (2003; Figure 3), using sequences of 16S rRNA and *rnpB*.

As shown in Figure 1, the species groups *suis* and *mitis* are in sister clades in both topologies. This is the only convergence between these phylogenetic hypotheses, as other relationships remained unclear.

Using the available genomic databases it is possible to incorporate all the gene fragment data that has been reported into a single phylogenetic analysis. This approach will enable the development of a new systematic work on this genus, covering all the known species groups. In this article, we attempt to create an extensive phylogeny for the genus *Streptococcus*, and contrast our proposed phylogeny with previously published hypotheses.

MATERIAL AND METHODS

Samples

DNA sequences from the genes *sodA*, *tuf*, *rpoB*, 16s rRNA, *rnpB*, *dnaJ*, *gyrB*, and *recN* were retrieved from GenBank (www.ncbi.nlm.nih.gov/genbank; Table 1). Seventy-five species and subspecies of *Streptococcus* were analyzed. *Lactococcus lactis* subsp. *lactis* and *Lactococcus lactis* subsp. *cremoris* were selected as outgroups, since this genus is one of the four members of the family Streptococcaceae. Although a primary analysis showed that *Pilibacter* is the sister clade of *Streptococcus*, the lack of data for this genus did not allow robust analyses to be performed (Figure 2).

Molecular phylogenetic analyses

Nucleotide sequences were aligned using the software BioEdit V. 7.0.9.0 (Hall, 1999). The alignments were checked and re-edited by eye. In order to avoid the effect of saturation of the phylogenetic signal, especially due to synonymous substitutions, amino acid sequences were also used for the six protein coding genes (*sodA*, *tuf*, *rpoB*, *dnaJ*, *gyrB*, and *recN*). The non-coding genes 16s rRNA and *rnpB* were included in both analyses of nucleotide sequences and amino acid derived sequences. Both nucleotide and amino acid derived sequences were concatenated in the same super-matrix. Absent data were coded as missing data and incorporated in subsequent analyses. This “super-matrix” approach has been shown to perform better than eliminating data, or using non-concatenated matrices, especially when Bayesian methods are used for phylogenetic inference (Wiens and Moen, 2008).

Models of molecular evolution for nucleotide sequences were selected using the Akaike information criterion test (AIC), as implemented in the software Modeltest (Posada and Crandall, 1998) and MrModeltest (Nylander, 2004). Models for amino acid sequences were selected using the PROTTEST v. 3.0 software (Abascal et al., 2005).

Bayesian inference on the complete super matrix was performed using Mr. Bayes v. 3.0b4, specifying four million generations and four independent chains. The resulting tree identified seven subclades that were used as the basis for subsequent analyses (Figure 3 and Table 2).

In order to solve phylogenetic relationships among species groups, a new matrix was built sampling two species belonging to each of the groups. This matrix, called the “mini-matrix”, was used for Bayesian inference using the same conditions and model substitutions as described above for the complete super-matrix. Phylogenetic concordance among the different DNA sequences was assessed by the “homogeneity test” (Farris et al., 1994, 1995) as implemented in Paup, in order to evaluate the phylogenetic compatibility among gene fragments inside the super-matrix.

Table 1. List of species, accession numbers and size (bp) of each gene fragment.

Species	<i>sodA</i>		<i>tuf</i>		<i>rpoB</i>		<i>recN</i>	
	bp	Accession No.	bp	Accession No.	bp	Accession No.	bp	Accession No.
<i>S. mutans</i>	612	AE014133.2	1197	AE014133.2	3105	AP010655.1	1249	EU917289.1
<i>S. agalactiae</i>	609	AE009948.1	1197	AL766847.1	3105	AE009948.1	1249	EU917242.1
<i>S. acidominimus</i>	435	Z95892.1	761	AY266992.1	691	AF535181.1	1249	EU917241.1
<i>S. anginosus</i>	453	FJ712177.1	826	AF276257.1	3105	AF535183.1	1249	EU917248.1
<i>S. alactolyticus</i>	435	AJ297185.1	-	-	680	DQ232445.1	1249	EU917226.1
<i>S. australis</i>	435	GU907529.1	761	EU156907.1	680	DQ132983.1	-	-
<i>S. caballi</i>	399	EF364100.1	-	-	-	-	-	-
<i>S. canis</i>	435	Z99175.1	761	EU156908.1	680	DQ232488.1	-	-
<i>S. castoreus</i>	435	EU003820.1	-	-	680	EU003817.1	1249	EU917312.1
<i>S. criceti</i>	435	Z95898.1	821	AF276260.1	680	DQ232485.1	1249	EU917257.1
<i>S. cristatus</i>	435	GU907530.1	761	AF276261.2	2061	AF194513.1, AF194514.1, AB199920.1	1249	EU917258.1
<i>S. devriesei</i>	435	DQ232544.1	-	-	680	DQ232446.1	1249	EU917306.1
<i>S. didelphis</i>	435	DQ232545.1	-	-	680	DQ232447.1	1249	EU917259.1
<i>S. downei</i>	435	Z95899.1	792	AF276262.1	680	DQ132984.1	-	-
<i>S. dysgalactiae</i> subsp <i>equisimilis</i>	486	AP010935.1	1215	AP010935.1	3105	AP010935.1	1249	EU917227.1
<i>S. constellatus</i>	453	FJ712176.1	821	AF276259.1	680	FJ712146.1	1093	FJ712113.1
<i>S. intermedius</i>	453	FJ712187.1	799	AF276267.1	-	-	1249	EU917282.1
<i>S. entericus</i>	435	DQ232547.1	-	-	680	DQ232448.1	1249	EU917228.1
<i>S. equi</i> subsp. <i>equi</i>	486	FM204883.1	1100	FM204883.1	3105	FM204883.1	1249	EU917229.1
<i>S. equinus</i>	435	AJ297213.1	792	AF276258.1	3105	AF535187.1	1249	EU917252.1
<i>S. ferus</i>	435	DQ132986.1	792	AF276265.1	686	AY770000.1	1249	EU917265.1
<i>S. gallinaceus</i>	435	DQ232548.1	-	-	680	DQ232503.1	1246	EU917266.1
<i>S. gallolyticus</i>	486	FN597254.1	1100	FN597254.1	3105	FN597254.1	-	-
<i>S. gordonii</i> str. Challis substr. CH1	486	CP000725.1	1100	CP000725.1	3105	CP000725.1	-	-
<i>S. halichoeri</i>	435	DQ232571.1	-	-	680	DQ232471.1	1249	EU917277.1
<i>S. henryi</i>	399	EF364099.1	-	-	-	-	-	-
<i>S. hyointestinalis</i>	435	DQ232550.1	-	-	680	DQ232449.1	1249	EU917231.1
<i>S. hyovaginalis</i>	435	DQ232551.1	-	-	680	DQ232450.1	1249	EU917232.1
<i>S. infantarius</i> subsp <i>infantarius</i>	435	AJ306980.1	761	EU156915.1	672	EU420169.1	-	-
<i>S. infantis</i>	432	GU907532.1	761	EU156917.1	680	DQ232482.1	1249	EU917280.1
<i>S. iniae</i>	486	EU661272.1	761	EU156918.1	680	DQ232493.1	1249	EU917281.1
<i>S. luteciae</i>	435	AJ297212.1	-	-	-	-	-	-
<i>S. lutetiensis</i>	426	GU991740.1	761	EU156916.1	680	DQ232480.1	1249	EU917279.1
<i>S. macacae</i>	435	DQ232553.1	792	AF276268.1	680	DQ232452.1	1249	EU917285.1
<i>S. macedonicus</i>	435	AJ297187.1	761	EU156913.1	693	AY315156.1	1249	HE613569.1
<i>S. marimammalium</i>	435	EU003821.1	-	-	659	EU003818.1	1249	EU917313.1
<i>S. massiliensis</i>	451	AY769999.1	761	EU156919.1	691	AY769998.1	1249	EU917311.1
<i>S. minor</i>	435	DQ232572.1	-	-	680	DQ232472.1	1249	EU917286.1
<i>S. mitis</i>	486	FN568063.1	1100	FN568063.1	3105	FN568063.1	1249	EU917233.1
<i>S. oligofermentans</i>	435	DQ232554.1	761	EU156926.1	680	DQ232508.1	1249	EU917290.1
<i>S. oralis</i>	435	DQ232576.1	764	AF276270.1	2658	AY695496.1, AB199956.1	1249	EU917234.1
<i>S. orisratti</i>	435	DQ232555.1	-	-	680	DQ232453.1	1249	EU917235.1
<i>S. ovis</i>	435	DQ232556.1	-	-	680	DQ232454.1	1249	EU917236.1
<i>S. parasanguinis</i>	435	GU907536.1	789	AF276271.1	-	-	1249	CP002843.1
<i>S. parauberis</i>	435	AJ544723.1	761	EU156937.1	680	DQ232455.1	1212	EU917293.1
<i>S. pasteurianus</i>	435	DQ232583.1	761	EU156914.1	680	DQ232505.1	1249	EU917273.1
<i>S. peroris</i>	435	DQ232541.1	761	EU156938.1	680	DQ232483.1	1249	EU917294.1
<i>S. phocae</i>	435	AJ547799.1	-	-	680	DQ232456.1	1249	EU917237.1
<i>S. pluranimalium</i>	435	DQ232557.1	761	EU156939.1	680	DQ232457.1	1249	EU917295.1
<i>S. plurextorum</i>	356	AM774231.1	-	-	687	AM774232.1	-	-
<i>S. pneumoniae</i>	486	FM211187.1	1100	FM211187.1	3105	FM211187.1	1249	EU917238.1
<i>S. porcinus</i>	435	Z99177.1	-	-	680	DQ232486.1	1249	EU917296.1
<i>S. pseudopneumoniae</i>	390	AB200048.1	761	EU156954.1	680	EU003819.1	1249	EU917314.1
<i>S. pyogenes</i>	486	CP000262.1	1100	AE009949.1	3105	CP000261.1	1249	EU917297.1
<i>S. ratti</i>	435	DQ232559.1	792	AF276272.1	680	DQ232484.1	1249	EU917298.1
<i>S. salivarius</i>	432	GU907538.1	792	AF276273.1	691	AF535169.1	1249	EU917239.1
<i>S. sanguinis</i>	486	CP000387.1	1100	CP000387.1	2922	CP000387.1	1249	EU917299.1

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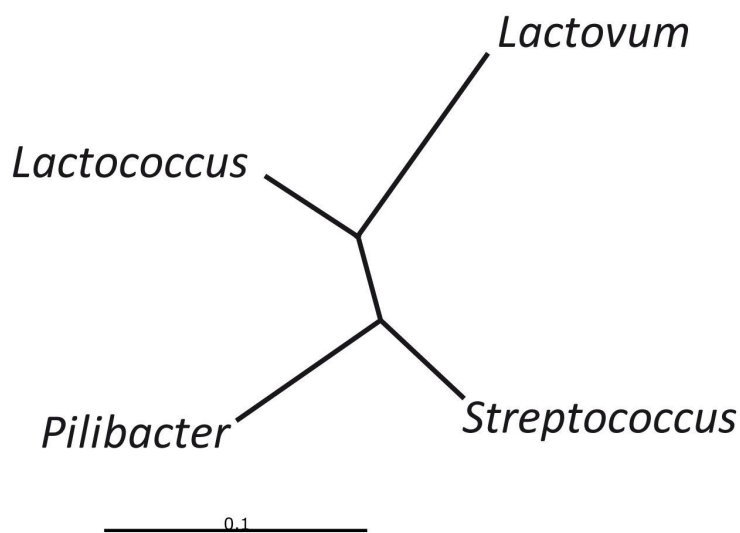
Table 1. Continued.

Species	<i>sodA</i>		<i>tuf</i>		<i>rpoB</i>		<i>recN</i>	
	bp	Accession No.	bp	Accession No.	bp	Accession No.	bp	Accession No.
<i>S. sinensis</i>	435	DQ232560.1	761	EU156962.1	680	DQ232458.1	1249	EU917305.1
<i>S. sobrinus</i>	435	Z95919.1	780	AF276275.1	680	DQ232489.1	-	-
<i>S. suis</i>	486	CP000408.1	1100	CP000408.1	3102	AM946016.1	1249	CP002570.1
<i>S. thermophilus</i>	486	CP000024.1	1100	CP000024.1	3105	CP000024.1	1249	EU917300.1
<i>S. thoralensis</i>	435	DQ232562.1	-	-	680	DQ232461.1	1249	EU917303.1
<i>S. uberis</i>	486	AM946015.1	1100	AM946015.1	3105	AM946015.1	1249	EU917309.1
<i>S. urinalis</i>	435	DQ232561.1	761	EU156963.1	680	DQ232460.1	1249	EU917240.1
<i>S. vestibularis</i>	436	AY687381.1	781	AF276277.1	686	AY687377.1	1249	EU917304.1
<i>L. lactis</i> subsp <i>lactis</i>	481	CP001834.1	1084	CP001834.1	3105	CP001834.1	-	-
<i>L. lactis</i> subsp <i>cremoris</i>	467	CP000425.1	1084	CP000425.1	3105	CP000425.1	-	-
Species	<i>16SrRNA</i>		<i>rmpB</i>		<i>dnaJ</i>		<i>gyrB</i>	
	bp	Accession No.	bp	Accession No.	bp	Accession No.	bp	Accession No.
<i>S. mutans</i>	1542	AB294730.1	380	AJ511678.1	1134	AE014133.2	1953	AP010655.1
<i>S. agalactiae</i>	1537	AF459432.1	374	AJ511673.1	1140	AL766843.1	1953	AE009948.1
<i>S. acidominimus</i>	1334	X58301.1	373	AJ511681.1	967	AB239170.1	900	AB238609.1
<i>S. anginosus</i>	1559	NR_041722.1	370	AJ511731.1	971	AB238697.1	900	AB236189.1
<i>S. alactolyticus</i>	1501	EU728776.1	376	AJ511706.1	-	-	-	-
<i>S. australis</i>	1471	NR_036936.1	-	-	648	GU907541.1	458	EU003771.1
<i>S. caballi</i>	1430	NR_044190.1	-	-	-	-	-	-
<i>S. canis</i>	1482	EU075058.1	374	AJ511684.1	969	AB238698.1	900	AB236191.1
<i>S. castoreus</i>	1508	NR_042215.1	-	-	-	-	458	EU003814.1
<i>S. criceti</i>	1530	EU483241.1	-	-	964	AB238699.1	900	AB238610.1
<i>S. cristatus</i>	1533	NR_042771.1	373	AJ511700.1	896	AB238700.1	900	AB238611.1
<i>S. dentapri</i>	1550	AB469560.1	-	-	-	-	-	-
<i>S. dentirosetti</i>	1543	NR_041460.1	-	-	-	-	-	-
<i>S. devriesei</i>	1471	EU483245.1	-	-	-	-	457	EU003742.1
<i>S. didelphis</i>	1506	AF176107.1	-	-	914	AB238810.1	457	EU003790.1
<i>S. downei</i>	1542	NR_042774.1	376	AJ511699.1	-	-	900	AB238618.1
<i>S. dysgalactiae</i> subsp <i>equisimilis</i>	1492	AP010935.1	374	CP002215.1	1137	AP010935.1	1953	AP010935.1
<i>S. constellatus</i>	1508	AY277942.1	370	AJ511742.1	964	AB238808.1	509	AB441109.1
<i>S. intermedius</i>	1531	GU470908.1	374	AJ511743.1	547	AB441138.1	1947	AB562520.1
<i>S. entericus</i>	1335	NR_025500.1	-	-	-	-	-	-
<i>S. equi</i> subsp. <i>equi</i>	1495	FM204883.1	374	FM204883.1	1137	FM204883.1	1953	FM204883.1
<i>S. equinus</i>	1539	AB362710.1	373	AJ511745.1	916	AB238704.1	-	-
<i>S. ferus</i>	1540	AB259060.1	366	AJ511705.1	902	AB238705.1	903	AB238616.1
<i>S. gallinaceus</i>	1502	NR_025453.1	-	-	-	-	458	EU003785.1
<i>S. galloyticus</i>	1535	AF323911.1	375	AJ511683.1	1143	AP012053.1	1953	AP012053.1
<i>S. gordonii</i> str. Challis substr. CH1	1499	CP000725.1	-	-	1134	CP000725.1	1950	CP000725.1
<i>S. halichoeri</i>	1506	NR_029025.1	-	-	-	-	458	EU003791.1
<i>S. henryi</i>	1430	NR_044189.1	-	-	-	-	-	-
<i>S. hyointestinalis</i>	1504	EU728763.1	372	AJ511696.1	1014	AB238709.1	900	AB238621.1
<i>S. hyovaginalis</i>	928	EF151158.1/ DQ118670.1	376	AJ512493.1	974	AB238710.1	900	AB238622.1
<i>S. ictaluri</i>	1471	DQ462421.1	-	-	-	-	-	-
<i>S. infantarius</i> subsp <i>infantarius</i>	1494	EU163504.1	371	AJ511688.1	1008	AB238812.1	458	EU003810.1
<i>S. infantis</i>	1468	NR_042928.1	387	AJ511687.1	835	AB238711.1	900	AB238623.1
<i>S. iniae</i>	1536	NR_025148.1	378	AJ511708.1	905	AB238712.1	900	GU324259.1
<i>S. luteciae</i>	1461	NR_042051.1	-	-	-	-	-	-
<i>S. lutetiensis</i>	1501	JN713319.1	371	AJ511709.1	-	-	458	EU003729.1
<i>S. macacae</i>	1542	NR_042775.1	373	AJ511702.1	1051	AB238713.1	900	AB238625.1
<i>S. macedonicus</i>	1538	AF459431.1	374	AJ511677.1	-	-	458	HE613569.1
<i>S. marimammalium</i>	1500	NR_025630.1	-	-	-	-	458	EU003815.1
<i>S. massiliensis</i>	1470	NR_043173.1	-	-	-	-	458	EU003813.1
<i>S. merionis</i>	1354	NR_042553.1	-	-	-	-	-	-
<i>S. minor</i>	1497	AY232833.1	-	-	-	-	458	EU003795.1
<i>S. mitis</i>	1540	AY518677.1	387	AJ511694.1	1140	FN568063.1	1947	FN568063.1
<i>S. oligofermentans</i>	1510	NR_029052.1	-	-	-	-	458	EU003768.1
<i>S. oralis</i>	1470	DQ232535.1	387	AJ511698.1	1137	FR720602.1	1947	FR720602.1
<i>S. orisratti</i>	1474	EU075064.1	374	AJ511692.1	1043	AB238813.1	458	EU003798.1

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Table 1. Continued.

Species	<i>16SrRNA</i>		<i>rnpB</i>		<i>dnaJ</i>		<i>gyrB</i>	
	bp	Accession No.	bp	Accession No.	bp	Accession No.	bp	Accession No.
<i>S. orisuis</i>	1553	NR_041055.1	-	-	-	-	-	-
<i>S. ovis</i>	1377	NR_026471.1	-	-	-	-	458	EU003799.1
<i>S. parasanguinis</i>	1513	NR_024842.1	373	AJ511704.1	974	AB238798.1	900	CP002843.1
<i>S. parauberis</i>	1541	AF284579.2	374	CP002471.1	890	AB238799.1	900	AB238630.1
<i>S. pasteurianus</i>	1511	EU163495.1	-	-	-	-	458	AP012054.1
<i>S. peroris</i>	1329	EU156772.1	387	AJ511690.1	890	AB238800.1	900	AB238631.1
<i>S. phocae</i>	1510	HM032023.1	371	AJ511670.1	935	AB238801.1	900	AB238632.1
<i>S. pluranimalium</i>	1552	Y18026.1	375	AJ511697.1	-	-	458	EU003801.1
<i>S. plurextorum</i>	1451	NR_042649.1	-	-	-	-	-	-
<i>S. pneumoniae</i>	1511	AM157442.1	386	CP002121.1	1137	AE005672.3	1947	CP000918.1
<i>S. porcinus</i>	1496	NR_024634.1	369	AJ511675.1	-	-	900	AB175052.1
<i>S. pseudopneumoniae</i>	1468	CP002925.1	-	-	-	-	458	EU003816.1
<i>S. pseudoporcinus</i>	1491	DQ303207.1	-	-	-	-	-	-
<i>S. pyogenes</i>	1518	FJ662846.1	374	CP003121.1	1137	AE009949.1	1953	CP000262.1
<i>S. rattii</i>	1444	NR_025516.1	373	AJ511671.1	896	AB238804.1	900	AB238635.1
<i>S. salivarius</i>	1543	AF459433.1	373	FR873482.1	621	GU907550.1	900	AB238640.1
<i>S. sanguinis</i>	1510	AY691542.1	373	AJ511682.1	1134	CP000387.1	1950	CP000387.1
<i>S. seminale</i>	1275	AB370977.1	-	-	-	-	-	-
<i>S. sinensis</i>	1512	AF432855.1	-	-	-	-	458	EU003809.1
<i>S. sobrinus</i>	1551	AB294731.1	373	AJ511707.1	-	-	900	AB238641.1
<i>S. suis</i>	1536	AF009497.1	370	AJ511674.1	1137	CP002633.1	1953	CP002641.1
<i>S. thermophilus</i>	1544	EF990662.1	373	AJ511712.1	1134	FR875178.1	1953	CP002340.1
<i>S. thoralensis</i>	1519	NR_026368.1	-	-	885	AB238806.1	900	AB238637.1
<i>S. uberis</i>	1501	NR_040820.1	372	AJ511693.1	1134	AM946015.1	1953	AM946015.1
<i>S. urinalis</i>	1476	DQ303194.1	373	AJ511680.1	-	-	458	EU003763.1
<i>S. ursoris</i>	1549	AB501126.1	-	-	-	-	-	-
<i>S. vestibularis</i>	1553	NR_042777.1	373	AJ511724.1	-	-	900	AB238643.1
<i>L. lactis</i> subsp <i>Lactis</i>	1539	CP001834.1	-	-	1140	CP001834.1	1956	CP001834.1
<i>L. lactis</i> subsp <i>cremoris</i>	1539	CP000425.1	-	-	1088	CP000425.1	1956	CP000425.1
<i>Lactovum miscens</i>	1456	NR_042140	-	-	-	-	-	-
<i>Pilibacter termitis</i>	1432	NR_042949	-	-	-	-	-	-

**Figure 2.** Unrooted tree obtained from a Bayesian analysis of representative species of the family Streptococcaceae, inferred from their 16sR RNA sequences (see details in Table 1).

RESULTS AND DISCUSSION

Seven major clades were detected in our analyses; these corresponded to the conventional species groups, except for the group mitis that was split into two non-monophyletic clades (Figure 3). In general, phylogenies inferred either from nucleotide or amino acid derived sequences did not differ either on major-clade relationships or statistical confidence (i.e., posterior probabilities).



Figure 3. Majority consensus tree obtained by a Bayesian analysis of proteic (sodA, tuf, rpoB, dnaJ, gyrB) and nucleotidic (16sr RNA, rnpB) of the entire matrix.

Table 2. Species of *Streptococcus*, and outgroup species, analyzed in this study.

Mutans group	<i>S. ratti</i> , <i>S. ursoris</i> , <i>S. devriesei</i> , <i>S. mutans</i> , <i>S. macacae</i> , <i>S. ferus</i> , <i>S. dentapri</i> , <i>S. downei</i> , <i>S. dentirousetti</i> , <i>S. sobrinus</i> , <i>S. dentirousetti</i> , <i>S. sobrinus</i> , <i>S. criceti</i> , <i>S. orisuis</i> , <i>S. merionis</i> , <i>S. caballi</i> , <i>S. henryi</i> , <i>S. orisratti</i> , <i>S. pluranimalium</i> , <i>S. thoralensis</i> , <i>S. hyovaginalis</i>
Pyogenic group	<i>S. dysgalactiae</i> , <i>S. pyogenes</i> , <i>S. canis</i> , <i>S. castoreus</i> , <i>S. equi</i> subsp <i>equi</i> , <i>S. halichoeri</i> , <i>S. phocae</i> , <i>S. porcinus</i> , <i>S. pseudoporcinus</i> , <i>S. didelphis</i> , <i>S. uberis</i> , <i>S. seminale</i> , <i>S. iniae</i> , <i>S. ictaluri</i> , <i>S. urinalis</i> , <i>S. parauberis</i> , <i>S. marimammalium</i> , <i>S. agalactiae</i>
Bovis group	<i>S. equinus</i> , <i>S. lutetiensis</i> , <i>S. luteciae</i> , <i>S. infantarius</i> , <i>S. gallolyticus</i> , <i>S. pasteurianus</i> , <i>S. macedonicus</i> , <i>S. alactolyticus</i>
Suis group	<i>S. entericus</i> , <i>S. plurextorum</i> , <i>S. suis</i> , <i>S. acidominimus</i> , <i>S. minor</i> , <i>S. ovis</i> , <i>S. gallinaceus</i>
Mitis group A	<i>S. constellatus</i> , <i>S. intermedius</i> , <i>S. anginosus</i> , <i>S. massiliensis</i> , <i>S. cristatus</i> , <i>S. sinensis</i> , <i>S. gordonii</i> , <i>S. sanguinis</i>
Mitis group B	<i>S. pneumoniae</i> , <i>S. pseudopneumoniae</i> , <i>S. mitis</i> , <i>S. oligofermentans</i> , <i>S. infantis</i> , <i>S. peroris</i> , <i>S. oralis</i> , <i>S. australis</i> , <i>S. parasanguinis</i>
Salivarius group	<i>S. vestibularis</i> , <i>S. salivarius</i> , <i>S. thermophilus</i>
Outgroup	<i>Lactococcus lactis</i> , <i>L. cremoris</i>

With regard to the phylogenetic relationships within species groups, it is notable that the mitis group (Kawamura et al., 1995) remained monophyletic but, in agreement with the results of Boggs et al. (2012) (Figure 3), was clearly subdivided into two clades. The suis group was basal to the sister clades.

Here we propose a taxonomic revision, since our results indicate that divergence between the two major clades subdividing the group mitis is in the range of divergence for other species groups (within-mean distances for the “gordonii” and “mitis” clades were 0.039 and 0.044, respectively; between-mean distance was 0.073). Therefore, a new species group was proposed and named gordonii; the group included the species *S. constellatus*, *S. intermedius*, *S. anginosus* [referred to as species group anginosus by Kawamura et al. (1995)], plus *S. massiliensis*, *S. cristatus*, *S. sinensis*, *S. gordonii*, and *S. sanguinis*. Additionally, the group mitis was retained, and included the species *S. australis*, *S. parasanguinis*, *S. oralis*, *S. peroris*, *S. infantis*, *S. oligofermentans*, *S. mitis*, *S. pseudopneumoniae*, and *S. pneumoniae*.

We suggest that gordonii is the basal group of the genus, followed by mitis, suis and finally pyogenic (Figure 3). The most recent divergence was the separation of the sister species groups salivarius and bovis. All the species groups, except mutans, resulted in monophyletic groups (Figure 3). *S. ferus* was basal to the entire genus, and not inside the mutans group as previously reported. Similarly, *S. entericus* and *S. plurextorum* (suis group), *S. marimammalium* and *S. agalactiae* (pyogenic group) were not placed in their previously described groups. The remaining species were clustered inside their formerly ascribed groups.

The diverse analyses performed here clearly show that the majority of the species groups had a stable topology. By changing the parameters of the gene sequences, i.e., using amino acid derived or nucleotide sequences, or by constraining clades, most of the species groups did not change their species composition. In the major part of the non-constrained analysis, the mutans species group resulted in a polyphyletic taxon, compressing at least three different lineages (Figure 3).

However, a homogeneity analysis revealed incompatibilities among gene sequences within the mutans species group; these incompatibilities might be due to horizontal gene transfer (Table 3) (Abascal et al., 2005; Itoh et al., 2006; Simmon et al., 2008; Boggs et al., 2012). More critical incompatibilities, detected by pairwise analysis, allowed us to identify large genetic discordances between different loci. We took these results into account in subsequent analyses, avoiding incompatibilities and generating different strategies in order to infer phylogenetic relationships within each species group and among groups.

Table 3. Homogeneity test results. P values for all the pairwise combinations of gene and protein fragments are shown for each species group and for the “mini matrix” set of species.

	Mutans group	Pyogenic group	Bovis group	Suis group	Mitis group A	Mitis group B	Mini matrix
All	0.01	0.01	0.27	0.88	0.01	0.01	0.01
sodA x tuf	0.26	0.12	0.56	0.99	0.14	0.01	0.04
sodA x rpoB	0.80	0.42	0.94	0.96	0.71	0.01	0.23
sodA x 16sr	0.02	0.01	0.30	0.53	0.17	0.01	0.03
sodA x mpbB	0.01	0.58	0.05	0.97	0.42	0.01	0.16
sodA x dnaJ	0.01	0.61	0.93	0.99	0.68	0.39	0.01
sodA x gyrB	0.27	0.04	0.88	0.38	0.62	0.03	0.93
sodA x recN	0.01	0.43	1	0.06	0.31	0.01	0.31
tuf x rpoB	0.41	0.37	1	1	0.51	0.01	0.02
tuf x 16sr	0.28	0.45	0.38	0.98	0.07	0.02	0.04
tuf x mpbB	0.01	0.99	0.56	1	0.72	0.01	0.01
tuf x dnaJ	0.01	0.83	1	1	0.77	0.50	0.01
tuf x gyrB	0.87	0.40	0.98	1	0.03	0.05	0.50
tuf x recN	0.01	0.83	0.89	0.97	0.26	0.01	0.23
rpoB x 16sr	0.41	0.08	0.74	0.66	0.51	0.07	0.01
rpoB x mpbB	0.17	0.91	1	1	0.35	0.31	0.01
rpoB x dnaJ	0.03	0.78	1	1	0.21	0.28	0.03
rpoB x gyrB	0.99	0.60	1	0.72	0.49	0.65	0.47
rpoB x recN	0.22	0.90	0.06	0.12	0.19	0.79	0.07
16sr x mpbB	0.05	0.16	0.49	0.98	0.01	0.05	0.09
16sr x dnaJ	0.10	0.29	0.92	1	0.69	0.14	0.01
16sr x gyrB	0.04	0.01	0.76	0.85	0.54	0.24	0.59
16sr x recN	0.01	0.01	0.88	0.41	0.68	0.01	0.33
mpbB x dnaJ	0.29	1	1	1	0.01	0.94	0.01
mpbB x gyrB	0.02	0.42	0.87	1	0.35	0.39	0.07
mpbB x recN	0.01	0.85	0.03	0.95	0.06	1	0.01
dnaJ x gyrB	0.01	0.67	1	1	0.09	0.31	0.02
dnaJ x recN	0.01	0.80	1	0.99	0.01	0.56	0.01
gyrB x recN	0.01	0.77	0.98	1	0.25	0.59	0.87

For example, with regard to the mutans group, only *sodA*, *tuf*, *rpoB*, and *gyrB* could be jointly analyzed, because only these genes shared high homogeneity scores (Table 3). Further analysis of this clade considered the entire matrix with the concatenated sequences of *sodA*, *rpoB*, and *gyrB*. These gene sequences were used in a new sampling strategy, which included all the species from the mutans group, plus two species representing each of the other species groups.

Next, we focused on the unsolved part of the tree: the mutans group and its relationship with salivarius, bovis, and pyogenic groups, using the same gene sequences and parameters as described above. This analysis reinforced the conclusion that the mutans group was not a monophyletic taxon. Rather, it appeared to be constituted by at least three lineages, namely mutans, sobrinus, and pluranimalium. This Bayesian tree resolved most of the relationships between these species (Figure 4).

Subsequently, we focused on unresolved parts of the phylogeny, namely, the relationship of *S. macacae* to the mutans and pyogenic groups, and the relationship between the pluranimalium and sobrinus groups. Homogeneity tests allowed us to select pairs of gene sequences to analyze for this part of the phylogeny (Table 4).

Bayesian analyses of *tuf-recN rpoB-recN* gene sequences placed *S. macacae* inside the mutans group, with *S. ferus*, *S. mutans*, *S. devriesei*, and *S. rattii* (Figure 5a). Bayesian trees of *sodA-rpoB*, 16S rRNA-*mpbB*, and *sodA-recN* gene sequences identified these two groups as sister clades (Figure 5b). Nonetheless, the phylogenetic relationships of *S. caballi* and *S. orisratti* remained unsolved since insufficient nucleotide data were present in the databases. Moreover, there was a lack of phylogenetic signal due to the low rate of nucleotide substitution (i.e., 16S rRNA) and phylogenetic incompatibility between the gene sequences (i.e. between 16S rRNA and *sodA*).

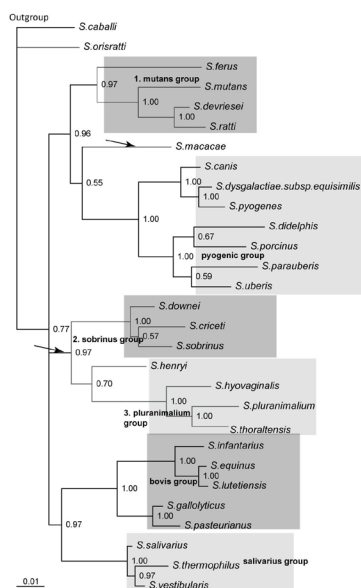


Figure 4. Majority consensus tree obtained from a Bayesian analysis of concatenated sequences of a subtree of *Streptococcus* using 3 proteic fragments (*sodA*, *rpoB*, *gyrB*). Arrows indicate unsolved points of the topology. Species groups *suis*, *gordonii* and *mitis* are excluded from this subtree. The models of protein evolution were WAG (*sodA*) and Rtrv (*rpoB*, *recN*).

Table 4. Homogeneity test results. P values for all the pairwise combinations of gene and protein fragments are shown for each one of the two pairs of minigroups derived from the mutans group.

	Clade A	Clade B
<i>sodA</i> x <i>tuf</i>	0.56	0.14
<i>sodA</i> x <i>rpoB</i>	0.98	0.87
<i>sodA</i> x 16s	0.72	0.01
<i>sodA</i> x <i>mpB</i>	0.94	0.01
<i>sodA</i> x <i>dnaJ</i>	0.97	0.13
<i>sodA</i> x <i>gyrB</i>	0.54	0.31
<i>sodA</i> x <i>recN</i>	0.98	0.56
<i>tuf</i> x <i>rpoB</i>	0.92	0.68
<i>tuf</i> x 16s	0.75	0.63
<i>tuf</i> x <i>mpB</i>	1	0.14
<i>tuf</i> x <i>dnaJ</i>	1	0.55
<i>tuf</i> x <i>gyrB</i>	0.90	0.33
<i>tuf</i> x <i>recN</i>	1	0.95
<i>rpoB</i> x 16s	0.95	0.02
<i>rpoB</i> x <i>mpB</i>	1	0.04
<i>rpoB</i> x <i>dnaJ</i>	1	0.57
<i>rpoB</i> x <i>gyrB</i>	1	0.56
<i>rpoB</i> x <i>recN</i>	0.85	0.94
16s x <i>mpB</i>	0.98	0.08
16s x <i>dnaJ</i>	0.96	0.06
16s x <i>gyrB</i>	0.54	0.01
16s x <i>recN</i>	0.92	0.01
<i>mpB</i> x <i>dnaJ</i>	1	0.08
<i>mpB</i> x <i>gyrB</i>	0.87	0.01
<i>mpB</i> x <i>recN</i>	1	0.57
<i>dnaJ</i> x <i>gyrB</i>	1	0.06
<i>dnaJ</i> x <i>recN</i>	1	0.78
<i>gyrB</i> x <i>recN</i>	0.94	0.79

Clade A = *Streptococcus pluranimalium*, *S. thoraltensis*, *S. hyovaginalis*, *S. henryi*, and *S. criceti*, *S. sobrinus*, *S. downei*. Clade B = *S. devriesei*, *S. rattii*, *S. mutans*, *S. ferus*, *S. macacae*, *S. canis*, *S. pyogenes*, *S. dysgalactiae*, *S. didelphis*, *S. porcinus*, and *S. uberis*.

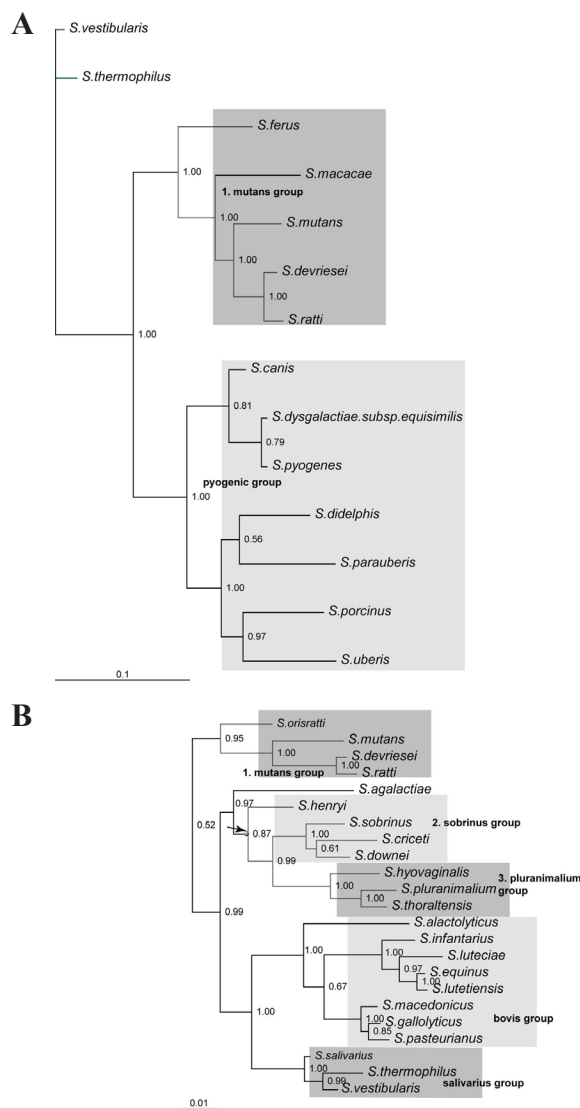


Figure 5. Majority consensus tree obtained of a Bayesian analysis of concatenated sequences. Those matrices were built using (A) 2 aminoacidic fragments (rpoB, recN), in order to solve the phylogenetic position of *Streptococcus macacae*; (B) 2 aminoacidic fragments (sodA, recN) to infer the relationship between sobrinus and pluranimalium groups. The models of protein evolution were WAG (sodA), Rtev (rpoB), and Jones (recN).

We performed a phylogenetic analysis of a mini-matrix comprising two species from each of the nine groups (mutans, sobrinus, pluranimalium, salivarius, bovis, pyogenic, gordonii, mitis, and suis; Figure 6). The results showed that, in contrast to previous studies, the suis, gordonii, and mitis groups were basal clades. In contrast, sobrinus and pluranimalium were sister clades and closely related to the salivarius group. This phylogeny is shown in Figure 7, and represents our hypothesis on the phylogenetic relationships among the species groups of *Streptococcus*.

Conflicts of interest

The authors declare no conflict of interest.

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