# Identification of Streptococci to Species Level by Sequencing the Gene Encoding the Manganese-Dependent Superoxide Dismutase

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We have used a PCR assay based on the use of degenerate primers in order to characterize an internal fragment ( $sodA_{int}$ ) representing approximately 85% of the genes encoding the manganese-dependent superoxide dismutase in various streptococcal type strains (S. acidominimus, S. agalactiae, S. alactolyticus, S. anginosus, S. bovis, S. constellatus, S. canis, S. cricetus, S. downei, S. dysgalactiae, S. equi subsp. equi, S. equi subsp. zooepidemicus, S. equinus, S. gordonii, S. iniae, S. intermedius, S. mittis, S. mutans, S. oralis, S. parasanguis, S. pneumoniae, S. porcinus, S. pyogenes, S. salivarius, S. sanguis, S. sobrinus, S. suis, S. thermophilus, and S. vestibularis). Phylogenetic analysis of these  $sodA_{int}$  fragments yields an evolutionary tree having a topology similar to that of the tree constructed with the 16S rRNA sequences. We have shown that clinical isolates could be identified by determining the positions of their  $sodA_{int}$  fragments on the phylogenetic tree of the  $sodA_{int}$  fragments of the type species. We propose this method for the characterization of strains that cannot be assigned to a species on the basis of their conventional phenotypic reactions.

The genus Streptococcus could be taxonomically divided into six major clusters which included at least 31 species (4, 8, 17, 18, 32, 34–36). These are (i) the pyogenic group, which includes S. agalactiae, S. canis, S. dysgalactiae, S. equi, S. iniae, S. porcinus, and S. pyogenes; (ii) the bovis group, which includes S. bovis, S. equinus, and S. alactolyticus; (iii) the salivarius group, which includes S. salivarius, S. thermophilus, and S. vestibularis; (iv) the mutans group, which includes S. cricetus, S. downei, S. mutans, and S. sobrinus; (v) the anginosus group (also referred to as the milleri group), which includes S. anginosus, S. constellatus, and S. intermedius; and (vi) the mitis group, which includes S. mitis, S. oralis, S. pneumoniae, S. sanguis, S. parasanguis, and S. gordonii. No single system of classification suffices for the differentiation of this heterogeneous group of organisms. Instead, classification depends on a combination of features including patterns of hemolysis observed on blood agar plates, antigenic composition, growth characteristics, biochemical reactions, and more recently, genetic analysis (3, 14, 18, 28).

In clinical laboratories, the current means of identification of streptococci rely on phenotypic tests such as those developed for the API ID 32 Strep system. However, the potential problems inherent to the use of phenotypic tests are that not all strains within a given species may be positive for a common trait (3, 18) and that the same strain may exhibit biochemical variability (15, 30). Moreover, small alterations in the realization of a test may give false results. Consequently, the routine technique based on phenotypic tests do not allow for an unequivocal identification of certain streptococcal species, in particular, those belonging to the milleri, the mutans, and the mitis groups (2, 3, 10, 18,

19). Nucleic acid-based technologies such as DNA hybridization (1, 16, 29) or amplification of selected targets (25, 27, 33) have been developed in recent years to complement and improve the identification of streptococci. We previously described a PCR assay based on the use of degenerate primers which enabled amplification of an internal fragment representing approximately 85% of the sodA gene encoding a manganese-dependent enzyme (manganese-dependent superoxide dismutase [Mn-SOD]) in various gram-positive bacteria including streptococci and enterococci (24). This gene has been identified as a target for the identification of mycobacteria at the species level by PCR (37), and we investigated in this study the sequencing of the sodA PCR product as an approach to the genotypic identification of 29 different streptococcal species including those constituting the milleri, mitis, and mutans groups.

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## MATERIALS AND METHODS

Bacterial strains and culture conditions. The main characteristics of the streptococcal strains used in this study, including the type strains, are listed in Tables 1 and 2. All strains were grown at 37°C on Columbia horse blood agar (bio-Mérieux, Marcy l'Etoile, France) in an anaerobic atmosphere. Phenotypic identifications were performed with the rapid ID 32 Strep System (API-bio-Mérieux, Marcy l'Etoile, France) according to the manufacturer's instructions. The API profiles were interpreted from the computer database for identification.

DNA manipulations. Rapid extraction of bacterial genomic DNA was performed as described previously (6), and primers d1 (5'-CCITAYICITAYGAYG CIYTIGARCC-3') and d2 (5'-ARRTARTAIGCRTGYTCCCAIACRTC-3') were used to amplify an internal fragment representing approximately 85% of the sod4 genes of the bacterial strains. PCRs were performed with a Gene Amp System 9600 instrument (Perkin-Elmer Cetus, Roissy, France) in a final volume of 50 μl containing 250 ng of DNA as template, 0.25 μM (each) primer, 200 μM (each) deoxynucleoside triphosphate, and 1 U of Taq DNA polymerase in a 1× amplification buffer (10 mM Tris-HCl [pH 8.3], 50 mM KCl, 1.5 mM MgCl<sub>2</sub>). The PCR mixtures were denatured (3 min at 95°C) and were then subjected to 35 cycles of amplification (90 s of annealing at 37°C, 90 s of elongation at 72°C, and 30 s of denaturation at 95°C) and to a final elongation cycle of 72°C for 10

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TABLE 1. Streptococcal type strains used in this study

Strain <sup>a</sup>	Other designation <sup>b</sup>	Relevant characteristics	sodA <sub>int</sub> accession no.
S. acidominimus CIP 82.4 <sup>T</sup>	NCDO 2025	Type strain	Z95892
S. agalactiae CIP 103227 <sup>T</sup>	ATCC 13813	Type strain	Z95893
S. alactolyticus CIP 103244 <sup>T</sup>	ATCC 43077	Type strain	Z95894
S. anginosus CIP 102921 <sup>T</sup>	ATCC 33397	Type strain	Z95895
S. bovis CIP 102302 <sup>T</sup>	ATCC 33317	Type strain	Z95896
S. canis CIP 103223 <sup>T</sup>	ATCC43496	Type strain	Z99175
S. constellatus CIP 103247 <sup>T</sup>	ATCC 27823	Type strain	Z95897
S. cricetus CIP 102510 <sup>T</sup>	ATCC 19642	Type strain	Z95898
S. downei CIP 103222 <sup>T</sup>	ATCC 33748	Type strain	Z95899
S. dysgalactiae CIP 102914 <sup>T</sup>	ATCC 43078	Type strain	Z95900
S. equinus CIP 102504 <sup>T</sup>	ATCC 9812	Type strain	Z95903
S. equi subsp. equi CIP 102910 <sup>T</sup>	ATCC 33398	Type strain	Z95901
S. equi subsp. zooepidemicus CIP 103228 <sup>T</sup>	ATCC 43079	Type strain	Z95902
S. gordonii CIP 105258 <sup>T</sup>	ATCC 10558	Type strain	Z95905
S. iniae CIP 102508 <sup>T</sup>	ATCC 29178	Type strain	Z99176
S. intermedius CIP 103248 <sup>T</sup>	ATCC 27335	Type strain	Z95908
S. mitis CIP 103335 <sup>T</sup>	NCTC 12261	Type strain	Z95909
S. mutans CIP 103694	ATCC 35668	Quality control strain for	Z95910
		API product	
S. oralis CIP 10922 <sup>T</sup>	ATCC 35037	Type strain	Z95911
S. parasanguis CIP 104372 <sup>T</sup>	ATCC 15910	Type strain	Z95913
S. pneumoniae CIP 102911 <sup>T</sup>	ATCC 33400	Type strain	Z95914
S. porcinus CIP 103218 <sup>T</sup>	ATCC 43138	Type strain	<b>Z</b> 99177
S. pyogenes CIP 56.41 <sup>T</sup>	ATCC 12344	Type strain	Z95915
S. salivarius CIP 102509 <sup>T</sup>	ATCC 19645	Type strain	Z95916
S. sanguis CIP 55.1328 <sup>T</sup>	ATCC 10556	Type strain	Z95918
S. sobrinus CIP 103230 <sup>T</sup>	ATCC 33478	Type strain	Z95919
S. suis CIP 103217 <sup>T</sup>	ATCC 43765	Type strain	Z95920
S. thermophilus CIP 102303 <sup>T</sup>	ATCC 19258	Type strain	Z95921
S. vestibularis CIP 103363 <sup>T</sup>	ATCC 49124	Type strain	Z95922

<sup>&</sup>lt;sup>a</sup> CIP, Collection de l'Institut Pasteur.

min. The PCR products were resolved by electrophoresis on a 1% agarose gel stained with ethidium bromide.

Cloning and sequencing. Amplification products were purified on a Sephadex S-200 column (Pharmacia, Uppsala, Sweden) and were ligated into the pUC18-SmaI dephosphorylated vector by using the Sure-clone ligation kit (Pharmacia, Uppsala, Sweden). Recombinant plasmids were analyzed by colony-PCR as follows. Twelve randomly chosen clones were amplified by using the universal -21 (5'-GTAAAACGACGGCCAGT-3') and reverse (5'-AACAGCTATGAC CATG-3') primers in a final volume of 50 µl containing 10<sup>3</sup> bacteria, 0.1 µM (each) primer, 200 µM (each) deoxynucleoside triphosphate, and 1 U of Taq DNA polymerase in a 1× amplification buffer (10 mM Tris-HCl [pH 8.3], 50 mM KCl, 1.5 mM MgCl<sub>2</sub>). The PCR mixtures were denatured (10 min at 95°C) and were then subjected to 30 cycles of amplification (90 s of annealing at 45°C, 1 min of elongation at 72°C, and 1 min of denaturation at 95°C). Colony-PCR products were directly sequenced after purification on a Sephadex S-400 column (Pharmacia). The entire nucleotide sequences of both strands of two cloned amplicons obtained from independent PCRs were determined by using the dideoxy chain termination method of Sanger with the dye primer cycle sequencing ready reaction kit on a Genetic ABI PRISM 310 Sequencer Analyzer (Perkin-Elmer, Applied Biosystem Division, Roissy, France).

Direct sequencing of the  $sodA_{int}$  PCR products with either of the degenerate oligonucleotides dI and d2 was performed with the dRhodamine dye terminator sequencing kit (Perkin-Elmer, Applied Biosystem Division), as follows. After purification on a Centricon-100 Concentrator column, 200 ng of the PCR product was mixed with 8  $\mu$ l of terminator reaction mixture and 10 pmol of primer in a final volume of 20  $\mu$ l, and the mixture was subjected to the following thermal cycling: 96°C for 10 s, 50°C for 5 s, and 60°C for 4 min (which was repeated for 25 cycles).

**Sequence analysis.** The nucleotide sequences were analyzed with Perkin-Elmer software (Sequence Analysis, Sequence Navigator, and Autoassembler). Multiple alignment of *sod* genes was carried out by the CLUSTAL X program (31). The construction of the unrooted phylogenetic tree was performed by the neighbor-joining method (26).

**Nucleotide sequence accession numbers.** The sequences were submitted to the EMBL gene bank and were assigned the accession numbers listed in Tables 1 and 2.

### RESULTS AND DISCUSSION

Amplification and sequencing of  $sodA_{int}$  from various streptococcal type strains. By using the primers d1 and d2 in a PCR

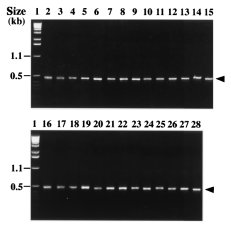


FIG. 1. Amplification of streptococcal type strains with the primers d1 and d2 and separation of the amplicons by 1% agarose gel electrophoresis. Lanes: 1, 1-kb ladder (Gibco, BRL); 2, *S. acidominimus*; 3, *S. agalactiae*; 4, *S. alactolyticus*, 5, *S. anginosus*; 6, *S. bovis*; 7, *S. constellatus*; 8, *S. cricetus*; 9, *S. downei*; 10, *S. dysgalactiae*; 11, *S. equi* subsp. equi; 12, *S. equi* subsp. zooepidemicus; 13, *S. equinus*; 14, *S. gordonii*; 15, *S. intermedius*; 16, *S. mitis*; 17, *S. mutans*; 18, *S. oralis*; 19, *S. parasanguis*; 20, *S. pneumoniae*; 21, *S. porcinus*; 22, *S. pyogenes*; 23, *S. salivarius*; 24, *S. sanguis*; 25, *S. sobrinus*; 26, *S. thermophilus*; 27, *S. suis*; 28, *S. vestibularis*. Arrowheads, 480-bp amplicon.

<sup>&</sup>lt;sup>b</sup> NCDO, National Collection of Dairy Organisms; ATCC, American Type Culture Collection; NCTC, National Collection of Type Cultures.

TABLE 2. Comparative identification of various streptococcal strains

Strain <sup>a</sup>	Relevant characteristics <sup>b</sup>	Species id	lentification	sodA <sub>int</sub> accession no		
Strain	Relevant characteristics	ID 32 Strep <sup>c</sup>	$sodA_{int}^{d}$			
S. agalactiae						
CIP 82.45	Lancefield group B (ATCC 12403)	ND	S. agalactiae (98.6)	Z99178		
NEM318	Lancefield group B	ND	S. agalactiae (97.9)	Z99179		
NEM1317	Lancefield group B	ND	S. agalactiae (98.4)	Z99180		
Milleri group						
NEM1164	Nontypeable, β-hemolytic	S. constellatus (99.6)	S. anginosus (97)	Z99181		
NEM1166	Nontypeable, β-hemolytic	S. anginosus (99.8)	S. anginosus (97.4)	Z99182		
NEM1124	Lancefield group F, β-hemolytic	S. constellatus (99.9)	S. constellatus (98.1)	Z99183		
NEM1162	Lancefield group F, β-hemolytic	S. constellatus (99.6)	S. constellatus (99.1)	Z99184		
NEM1165	Nontypeable	S. constellatus (93.1)	S. constellatus (98.9)	Z99185		
NEM1275	Nontypeable	S. constellatus (63.9)	S. constellatus (99.1)	Z99186		
MG19	Nontypeable	Milleri group	S. constellatus (99.1)	Z99187		
Mitis group						
CIP 103221	ATCC 33399	S. gordonii (?)	S. gordonii (95.9)	Z99189		
BM120	Strain Challis	S. gordonii (99.9)	S. gordonii (98.2)	Z99188		
NEM666	Nontypeable	?	S. gordonii (95.9)	Z99190		
NEM1222	Nontypeable	S. mitis (95.6)	S. mitis (97.8)	Z99192		
NEM1126	Lancefield group C	S. mitis (59.5)	S. mitis (97.8)	Z99192 Z99191		
NEWIII20	Lancencia group C	5. muis (59.5)	5. muis (97.6)	233131		
S. mutans GS-5		ND	C mutana (00.8)	D01027		
	NI 4 1-1 -		S. mutans (99.8)	D01037		
NEM1163	Nontypeable	S. mutans (99.9)	S. mutans (99.5)	Z99193		
S. oralis	1 mag 10 <b>1 = -</b>		g (02 =)	<b>5</b> 00404		
CIP 103216	ATCC 10557		S. oralis (93.7)	Z99194		
NEM1121	Lancefield group C	S. salivarius (99.8)	S. oralis (96.1)	Z99195		
NEM895	Nontypeable	?	S. parasanguis (97.8)	<b>Z</b> 99196		
S. pneumoniae						
NEM667	Serotype 11	ND	S. pneumoniae (100)	Z99246		
NEM1278	Serotype 23F	ND	S. pneumoniae (99.8)	Z99204		
NEM1279	Serotype 23F, Pen <sup>r</sup>	ND	S. pneumoniae (100)	Z99205		
NEM1280	Serotype 23F, Pen <sup>r</sup>	ND	S. pneumoniae (100)	Z99206		
NEM1251	Serotype 16	ND	S. pneumoniae (100)	Z99201		
NEM1252	Serotype 18	ND	S. pneumoniae (100)	Z99202		
NEM1253	Serotype 6	ND	S. pneumoniae (100)	Z99203		
NEM1122	Serotype 23F, Opt <sup>r</sup>	?	S. pneumoniae (99.8)	Z99200		
5. pyogenes						
BM105	Lancefield group A	ND	S. pyogenes (100)	Z49247		
HSC5	Lancefield group A	ND	S. pyogenes (100)	U43776		
5. salivarius						
CIP 102505	ATCC 13419	S. salivarius (?)	S. salivarius (96.3)	Z99197		
NEM1250	Lancefield group D	S. salivarius (?)	S. salivarius (98.6)	Z99197 Z99198		
NEM1257	Nontypeable	S. salivarius (99.5)	S. salivarius (96.3)	Z99198 Z99199		
1412141143/	rvontypeable	5. sauvarius (22.5)	5. suuvarius (30.5)	£33133		

<sup>&</sup>lt;sup>a</sup> Data for all strains except S. agalactiae CIP 82.45 (24), S. agalactiae NEM318 (11), MG19 (5), S. gordonii BM120 (20), S. mutans GS-5 (21), S. pneumoniae NEM667 (24), *S. pyogenes* BM105 (24), and *S. pyogenes* HSCS (13) are from this work. <sup>b</sup> Opto<sup>r</sup> and Pen<sup>r</sup>, resistance to optochin and penicillin, respectively.

assay, we amplified an internal fragment representing approximately 85% of the sodA gene encoding a manganese-dependent enzyme (Mn-SOD) in 29 type strains of streptococci (S. acidominimus, S. agalactiae, S. alactolyticus, S. anginosus, S. bovis, S. canis, S. constellatus, S. cricetus, S. downei, S. dysgalactiae, S. equi subsp. equi, S. equi subsp. zooepidemicus, S. equinus, S. gordonii, S. iniae, S. intermedius, S. mitis, S. mutans, S. oralis, S. parasanguis, S. pneumoniae, S. porcinus, S. pyogenes, S. salivarius, S. sanguis, S. sobrinus, S. suis, S. thermophilus, and S. vestibularis). A single amplification product having the expected size of 480 bp was observed with all streptococcal species (Fig. 1 shows the results of part of this analysis). The nucleotide sequences of the  $sodA_{int}$  fragments from these type strains were determined following cloning into pUC18 (Table 1). Analysis of the corresponding deduced amino acid sequences (data not shown) revealed that they all possessed three histidyl residues and one aspartyl residue that supposedly serve as metal ligands at positions characteristic of Mn- or

<sup>&</sup>lt;sup>c</sup> The rapid ID 32 Strep system (API-bio-Mérieux) was used according to the manufacturer's instructions to identify isolates to the species level. The API profiles were interpreted from the computer database for identification. The numbers in parentheses indicate the API identification percentage. ND, not determined; ?, undetermined.

<sup>&</sup>lt;sup>d</sup> The species identification was based on the phylogenic position of the sodA<sub>int</sub> fragment of the strain studied relative to those of the type strains, as indicated in Fig. 1. The numbers in parentheses indicate the percent identity of the sodA<sub>int</sub> fragment with that of the corresponding type strain.

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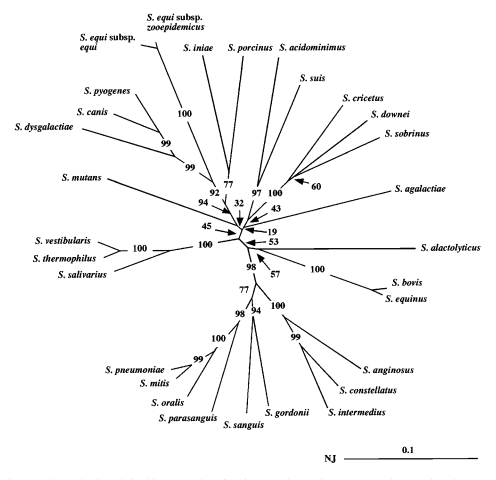


FIG. 2. Phylogenetic unrooted tree showing relationships among the  $sodA_{int}$  fragments from various streptococcal type strains. The tree was established from an analysis of the sequences listed in Table 1 by using the neighbor-joining method (26). The value on each branch is the estimated confidence limit (expressed as a percentage) for the position of the branch as determined by bootstrap analysis. The scale bar (neighbor-joining [NJ] distance) represents 10% differences in nucleotide sequences.

Fe-SODs (22, 23). Moreover, they all contain in the vicinity of the active site four other residues characteristic of Mn-SODs, which suggests that the corresponding enzymes are activated by Mn ions (23). We therefore concluded that the PCR products cloned and sequenced were actual sodA<sub>int</sub> fragments. Multiple alignment of the streptococcal sodA<sub>int</sub> sequences was carried out by the CLUSTAL X program (31), and an unrooted phylogenetic tree was constructed by the neighbor-joining method (26). Domains I and II corresponding to the degenerate oligonucleotides d1 and d2, respectively, and alignment gaps were not taken into consideration for calculations. The topology of the phylogenetic tree obtained (Fig. 2) was evaluated by bootstrap analyses to give the degree of confidence intervals for each node on the phylogenetic tree. The confidence values are given at the branches, which show possibly monophyletic clades of related organisms separated at each node. It is generally accepted that the monophyly of a clade can be accepted if the clade occurs in more than 95% of the bootstrapped trees (9). If this critical value is used, the  $sodA_{int}$ based phylogenetic positions of the streptococcal species studied here were in general agreement with those inferred from an analysis of their 16S rRNA sequences (4, 17), with the following remarkable exceptions: S. agalactiae did not cluster with the species constituting the pyogenic group, and S. mutans was genetically separate from S. cricetus, S. downei, and S.

sobrinus. Pairwise comparison of two given streptococcal species always revealed a lower percentage of sequence identity between their  $sodA_{int}$  fragments than between their 16S RNAs (Table 3 and data not shown). For example, the sequence identities of the 16S RNAs of type strains of S. mitis, S. oralis, and S. pneumoniae are greater than 99% (17), whereas those of their sodA<sub>int</sub> fragments vary from 92% (S. mitis versus S. oralis and S. oralis versus S. pneumoniae) to 96.6% (S. mitis versus S. pneumoniae) (Table 3). These results indicate that the sodA gene might constitute a more discriminative target sequence than the 16S RNA for differentiating closely related bacterial species. On the other hand, it is worth noting that the close structural relationship (98.9% identity) observed between the sodA<sub>int</sub> fragments of S. equi subsp. equi and S. equi subsp. zooepidemicus is consistent with their association in a single species (7).

Species identification of streptococcal clinical isolates by sequencing the  $sodA_{int}$  gene. The design of species-specific oligonucleotides enabling the amplification of a given target DNA constitutes an interesting molecular approach for the identification of bacterial species by PCR (12, 37). The two major problems inherent to these techniques are that (i) species-specific oligonucleotides often cannot be designed for closely related species, and (ii) the number of PCRs necessary for the identification of one isolate is proportional to the num-

TABLE 3. Identity matrix based pairwise comparisons of  $sodA_{int}$  fragments of streptococcal type strains

Strain <sup>a</sup>	% Identity with the following strain <sup>b</sup> :																											
Strain	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24	25	26	27	28	29
1. S. acidominimus	71.3	67.8			69.9											69.7										79.5		
2. S. agalactiae		69.0														69.7												72.6
3. S. alactolyticus			73.1		69.7	,	66.4		69.7		65.7	75.2											,				72.2	71.3
4. S. anginosus				71.7	67.1	86.4		68.5	65.5		66.7	72.2		65.3					75.9					76.6			70.8	70.3
5. S. bovis					72.0											72.9												
6. S. canis						64.1	73.1									67.8											72.0	
7. S. constellatus							65.5		66.0		62.1	,	81.8			,		75.6			69.2					,	71.5	,
8. S. cricetus								84.1	70.3			,				71.0												
9. S. downei									70.1	,	,	,	65.7	,		68.3								68.7				
10. S. dysgalactiae										74.9	,		62.8	, ,		66.9			67.6		72.4				,		73.1	,
11. S. equi subsp. equi											98.9		63.0	,										63.4				
12. S. equi subsp. zooepidemicus												65.7				64.1												66.4
13. S. equinus													71.7	68.5		72.6	,,	72.2	73.3			70.3			69.2		75.9	
14. S. gordonii														64.1	81.1								73.3		68.7	72.9	73.3	,
15. S. iniae															63.9		74.0										70.8	70.1
16. S. intermedius																75.6			77.2					77.0			70.6	,
17. S. mitis																	68.5						76.3		70.8	73.3	75.6	
18. S. mutans																		67.4	67.6	68.5	74.3	75.2	78.4	68.7	72.0	74.5	74.7	75.6
19. S. oralis																			85.3	92.0	66.9	63.0	73.6	79.5	69.4	73.3	74.0	73.6
20. S. parasanguis																				85.7	65.3			79.3			70.6	, 0.0
21. S. pneumoniae																					66.4	66.7	75.4	78.2	71.3	73.6	75.2	74.9
22. S. porcinus																						74.9	71.0	66.4	69.4	71.3	73.1	72.0
23. S. pyogenes																							74.5	63.7	72.6	66.7	71.5	70.3
24. S. salivarius																								72.2	75.6	73.3	89.7	90.1
25. S. sanguis																									67.4	72.0	72.4	71.7
26. S. sobrinus																										75.4	77.0	75.6
27. S. suis																											72.0	72.9
28. S. thermophilus																												96.8
29. S. vestibularis																												

 $<sup>^</sup>a$  The main characteristics of the strains are listed in Table 1.  $^b$  The strain numbers correspond to the strains identified by the numbers on the left.

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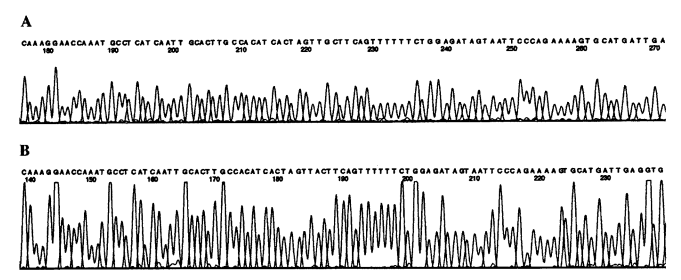


FIG. 3. Electropherograms showing part of the nucleotide sequence of  $sodA_{int}$  from *S. porcinus*. Sequence reactions were carried out with a pUC18 $\Omega sodA_{int}$  recombinant plasmid with the -21 dye primer sequencing kit (A) and  $sodA_{int}$  PCR product with the degenerate oligonucleotide d2 and the dRhodamine dye terminator sequencing kit (B).

ber of bacterial type species that should be considered. Taking into consideration the fact that cloning and sequencing techniques are increasingly used as routine techniques in medical microbiology laboratories, we propose the sequencing of the sodA<sub>int</sub> fragment as a molecular approach to the identification of streptococcal species. In this system, the identification of a clinical isolate is determined by the position of its sodA<sub>int</sub> fragment on the phylogenetic tree of the  $sodA_{int}$  fragments of the type species (Fig. 2). To test the functionality of this approach, various typeable and nontypeable streptococcal isolates were identified by using the ID 32 Strep and/or the sodA<sub>int</sub> systems (Table 2). We also include in this study the sequences of sodA<sub>int</sub> fragments of known streptococcal species present in the databases. The results obtained with the  $sodA_{int}$  system indicated that, as expected, the two group A and the three group B streptococcal strains studied were identified as S. pyogenes and S. agalactiae, respectively. When the streptococcal strains that were unambiguously identified to the species level by the phenotypic method (API identification percentage,  $\geq$ 99.9), the *sodA*<sub>int</sub>-based identification method gave the same results (Table 2). Some discrepancies were observed for the strains the species of which were determined with an API identification percentage of less than 99.9. This was the case for NEM1164 and NEM1121, which were identified with the ID 32 Strep system as S. constellatus and S. salivarius, respectively, but which were identified with the  $sodA_{int}$  system as S. anginosus and S. oralis, respectively (Table 2). The sequence of the sodA<sub>int</sub> fragment of NEM1164 displays 97 and 86% identity with the sequences of the type strains of S. anginosus and S. constellatus, respectively. The sequence of the sodA<sub>int</sub> fragment of NEM1121 displays 96.1 and 73.6% identity with the sequences of the type strains of S. oralis and S. anginosus, respectively. On the basis of these sequence distances, we considered the sodA<sub>int</sub>-based identification of NEM1164 (S. anginosus) and NEM1121 (S. oralis) to be more reliable than the ID 32 Strep system-based identification. Interestingly, certain strains (NEM1275, MG19, NEM666, NEM1126, and NEM895) were identified to the species level with the sodA<sub>int</sub> system but not with ID 32 Strep system (Table 2). Finally, it is important that the intraspecies divergence between the  $sodA_{int}$ 

fragments may vary greatly depending upon the species considered, conceivably because of differences in sequence divergence rates. Consequently, it is not possible to delineate streptococcal species on the basis of the level of DNA homology. In the case of S. oralis, the levels of intraspecies divergence of the  $sodA_{int}$  fragments can exceed those observed between the  $sodA_{int}$  fragments of the type strains of S. oralis, S. mitis, and S. pneumoniae. These results might suggest that the species S. oralis, as defined, is genomically heterogeneous. Surprisingly, the  $sodA_{int}$  fragments from unrelated S. pneumoniae strains were found to display the highest level of intraspecies sequence identity (>99%), which suggests that transformation is not a source of sequence heterogeneity for the sodA gene, at least in pneumococci.

Concluding remarks. We have described a method that enables the reliable identification to the species level of groupable and nongroupable streptococci. It consists of (i) a PCR carried out with a single pair of degenerate oligonucleotides for amplification of a streptococcal sodA<sub>int</sub> fragment, (ii) molecular cloning of the resulting amplicon into an Escherichia coli vector, and (iii) sequencing of the insert on both DNA strands. Sequencing of a streptococcal sodA<sub>int</sub> fragment by using this procedure necessitates 72 h following receipt of the bacterial strain; however, we anticipate further simplification and/or automation of various steps of this method. For example, based on the observation that a single abundant PCR product was obtained by using the degenerate sod primers, whatever the species tested (Fig. 1), we successfully tried to sequence directly both strands of the amplified DNA with either of the degenerate PCR primers (Fig. 3 shows part of the results of that analysis). Removal of the cloning step greatly enhances the applicability of the procedure and reduces the delay required for bacterial identification to 24 h. This method might be useful in reference laboratories for the characterization strains that could not be assigned to a species on the basis of their conventional phenotypic reaction. It provides a convenient alternative to the DNA-DNA hybridization method, which constitutes the reference technique for the identification of strains to the species level. We are expanding the applicability of this approach by determining the nucleotide sequence

of  $sodA_{int}$  fragments from other species of streptococci as well as from other related genera such as *Abiotrophia*, *Enterococcus*, *Gemella*, *Leuconostoc*, and *Pediococcus*.

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#### REFERENCES

- Adnan, S., N. Li, H. Miura, Y. Hashimoto, H. Yamamoto, and T. Ezaki. 1993. Covalently immobilized DNA plate for luminometric DNA-DNA hybridization to identify viridans streptococci in under 2 hours. FEMS Microbiol. Lett. 106:139–142.
- Ahmet, Z., M. Warren, and E. T. Houang. 1995. Species identification of members of the *Streptococcus milleri* group isolated from the vagina by ID 32 Strep system and differential phenotypic characteristics. J. Clin. Microbiol. 33:1592–1595.
- Beighton, D., J. M. Hardie, and A. Whiley. 1991. A scheme for the identification of viridans streptococci. J. Med. Microbiol. 35:367–372.
- Bentley, R. W., J. A. Leigh, and M. D. Collins. 1991. Intrageneric structure of *Streptococcus* based on comparative analysis of small-subunit rRNA sequences. Int. J. Syst. Bacteriol. 41:487–494.
- Clermont, D., and T. Horaud. 1990. Identification of chromosomal antibiotic resistance genes in *Streptococcus anginosus* ("S. milleri"). Antimicrob. Agents Chemother. 34:1685–1690.
- de Lamballerie, X., C. Zandotti, C. Vignoli, C. Bollet, and P. de Micco. 1992.
   A one-step microbial DNA extraction method using "Chelex 100" suitable for gene amplification. Res. Microbiol. 143:785–790.
- Farrow, J. A. E., and M. D. Collins. 1984. Taxonomic studies on streptococci
  of serological groups C, G, and L and possibly related taxa. Syst. Appl.
  Microbiol. 5:483–493.
- Farrow, J. A. E., J. Kruze, B. A. Phillips, A. J. Bramley, and M. D. Collins. 1984. Taxonomic studies on *Streptococcus bovis* and *Streptococcus equinus*: description of *Streptococcus alactolyticus* sp. nov. Syst. Appl. Microbiol. 5: 467–482.
- Felsenstein, J. 1985. Confidence limits on phylogeny and approach using the boostrap. Evolution 39:783–791.
- Flynn, C. E., and K. L. Ruoff. 1995. Identification of *Streptococcus milleri* group isolates to the species level with a commercially available rapid test. J. Clin. Microbiol. 33:2704–2706.
- 11. Gaillot, O., C. Poyart, P. Berche, and P. Trieu-Cuot. Molecular characterization and expression analysis of the superoxide dismutase gene from *Streptococcus agalactiae*. Gene, in press.
- Garnier, F., G. Gerbaud, P. Courvalin, and M. Galimand. 1997. Identification of clinically relevant viridans group streptococci to the species level by PCR. J. Clin. Microbiol. 35:2337–2341.
- Gibson, C. M., and M. G. Caparon. 1996. Insertional inactivation of Streptococcus pyogenes sod suggests that prtF is regulated in response to a superoxide signal. J. Bacteriol. 178:4688–4695.
- Hardie, J. M. 1986. Genus Streptococcus, p. 1043–1071. In P. H. A. Sneath, N. S. Mair, M. E. Sharpe, and J. G. Holt (ed.), Bergey's manual of systematic bacteriology. The Williams & Wilkins Co, Baltimore, Md.
- Hillman, J. D., S. W. Andrews, S. Painetr, and P. Stashenko. 1989. Adaptative changes in a strain of *Streptococcus mutans* during colonization of the human oral cavity. Microb. Ecol. Health Dis. 2:231–239.
- Jacobs, J. A., C. S. Schot, A. E. Bunschoten, and L. M. Schouls. 1996. Rapid species identification of "Streptococcus milleri" strains by line blot hybridization: identification of a distinct 16S rRNA population closely related to Streptococcus constellatus. J. Clin. Microbiol. 34:1717–1721.
- Kawamura, Y., X.-G. Hou, F. Sultana, H. Miura, and T. Ezaki. 1995. Determination of 16S rRNA sequences of Streptococcus mitis and Streptococcus

- gordonii and phylogenetic relationships among members of the genus Streptococcus. Int. J. Syst. Bacteriol. 45:406–408.
- Kilian, M., L. Mikkelsen, and J. Henrichsen. 1989. Taxonomic studies of viridans streptococci: description of Streptococcus gordonii sp. nov. and emended descriptions of Streptococcus sanguis (White and Niven 1946), Streptococcus oralis (Bridge and Sneath 1982), and Streptococcus mitis (Andrewes and Horder 1906). Int. J. Syst. Bacteriol. 39:471–484.
- Kilpper-Bälz, R., B. L. Williams, R. Lutticken, and K. H. Schleifer. 1984.
   Relatedness of 'Streptococcus miller' with Streptococcus anginosus an Streptococcus constellatus. Syst. Appl. Microbiol. 5:494–500.
- Le Bouguénec, C., T. Horaud, G. Bieth, R. Colimon, and C. Dauguet. 1984. Translocation of antibiotic resistance markers of a plasmid-free *Streptococcus pyogenes* (group A) strain into different streptococcal hemolysin plasmids. Mol. Gen. Genet. 194:377–387.
- Nakayama, K. 1992. Nucleotide sequence of Streptococcus mutans superoxide dismutase gene and isolation of insertion mutants. J. Bacteriol. 174:4928–4934
- Parker, M. W., and C. C. F. Balke. 1988. Crystal structure of manganese superoxide dismutase from *Bacillus stearothermophilus* at 2.4 Å resolution. J. Mol. Biol. 199:649–661.
- Parker, M. W., and C. C. F. Blake. 1988. Iron- and manganese-containing superoxide dismutases can be distinguished by analysis of their primary structures. FEBS Lett. 229:377–382.
- Poyart, C., P. Berche, and P. Trieu-Cuot. 1995. Characterization of superoxide dismutase genes from gram-positive bacteria by polymerase chain reaction using degenerate primers. FEMS Microbiol. Lett. 131:41–45.
- Rudney, J. D., and C. J. Larson. 1994. Use of restriction fragment polymorphism analysis of rRNA genes to assign species to unknown clinical isolates of oral viridans streptococci. J. Clin. Microbiol. 32:437–443.
- Saitou, N., and M. Nei. 1987. The neighbor-joining method: a new method for reconstructing phylogenetic trees. Mol. Biol. Evol. 4:406–425.
- Saruta, K., T. Matsunaga, S. Hoshina, M. Kono, S. Kitahara, S. Kanemoto, O. Sakai, and K. Machida. 1995. Rapid identification of *Streptococcus pneu-moniae* by PCR amplification of ribosomal DNA spacer region. FEMS Microbiol. Lett. 132:165–170.
- Schleifer, K. H., and R. Kilpper-Bälz. 1987. Molecular and chemotaxonomic approaches to the classification of streptococci, enterococci, and lactococci: a review. Syst. Appl. Microbiol. 10:1–19.
- Schmidhuber, S., W. Ludwig, and K. H. Schleifer. 1988. Construction of a DNA probe for the specific identification of *Streptococcus oralis*. J. Clin. Microbiol. 26:1042–1044.
- Tardif, G., M. C. Sulavik, G. W. Jones, and D. B. Clewell. 1989. Spontaneous switching of the sucrose-promoted colony phenotype in *Streptococcus sanguis*. Infect. Immun. 57:3945–3948.
- Thompson, J. D., D. G. Higgins, and T. J. Gibson. 1994. CLUSTAL W: improving the sensitivity of progressive multiple sequence alignment through sequence weighting, positions-specific gap penalties and weight matrix choice. Nucleic Acids Res. 22:4673–4680.
- Whiley, R. A., and D. Beighton. 1991. Emended descriptions and recognition of Streptococcus constellatus, Streptococcus intermedius, and Streptococcus anginosus as distinct species. Int. J. Syst. Bacteriol. 41:1–5.
- Whiley, R. A., B. Duke, J. M. Hardie, and L. M. C. Hall. 1995. Heterogeneity among 16S-23S rRNA intergenic spacers of species within the 'Streptococcus milleri group.' Microbiology 141:1461–1467.
- 34. Whiley, R. A., H. Y. Fraser, C. W. I. Douglas, J. M. Hardie, A. M. Williams, and M. D. Collins. 1990. Streptococcus parasanguis sp. nov., an atypical viridans Streptococcus from human clinical specimens. FEMS Microbiol. Lett. 68:115–122.
- Whiley, R. A., and J. M. Hardie. 1988. Streptococcus vestibularis sp. nov. from the human oral cavity. Int. J. Syst. Bacteriol. 38:335–339.
- Whiley, R. A., R. R. B. Russell, J. M. Hardie, and D. Beighton. 1988. Streptococcus downeii sp. nov. for strains previously described as Streptococcus mutans serotype H. Int. J. Syst. Bacteriol. 38:25–29.
- Zolg, J. W., and S. Philippi-Schulz. 1994. The superoxide dismutase gene, a target for detection and identification of mycobacteria by PCR. J. Clin. Microbiol. 32:2801–2812.