

Simultaneous Detection of Streptococcus pneumoniae, S. mitis, and S. oralis by a Novel Multiplex PCR Assay Targeting the gyrB Gene

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A multiplex PCR (mPCR) protocol was developed for simultaneous detection of the gyrB gene in Streptococcus pneumoniae, Streptococcus mitis, and Streptococcus oralis, and the specificity was evaluated using 141 coccus strains. Genomic DNAs purified from S. pneumoniae, S. mitis, and S. oralis strains were efficiently detected with size differences, whereas no PCR products were amplified from any of the reference strains tested. A pilot study of 47 human oral swab specimens was conducted in parallel, and the mPCR assay identified S. pneumoniae in 1 sample, S. mitis in 8 samples, and S. oralis in 2 samples, providing a powerful means for characterization at the level of species compared with traditional culture analysis. Our results suggest that the mPCR protocol presented here is a sensitive and promising tool for the rapid detection and discrimination of S. pneumoniae, S. mitis, and S. oralis from clinical specimens.

treptococcus pneumoniae is an important human pathogen associated with pneumonia, bacterial meningitis, otitis media, and nongonoccal urethritis (1-3). In contrast, two viridans group streptococci (VGS), Streptococcus mitis and Streptococcus oralis, are recognized as important etiological agents of dental caries and subacute bacterial endocarditis and septicemia (4-6), and recently, pancreatic diseases, including pancreatic cancer, have been associated with S. mitis. Thus, the pathogenesis and clinical importance of these cocci are increasing (7).

Precise discrimination among closely related bacterial species is crucial for accurate diagnosis and treatment. Because S. pneumoniae is often isolated with S. mitis, S. oralis, and other commensals in the oral cavity, accurate discrimination among these strains is important for diagnosis and treatment (8). However, identification of these species has traditionally been difficult using current clinical laboratory techniques, because they have a close, common genetic ancestry (5, 9, 10). Moreover, identification of these organisms at the species level by genotypic and phenotypic methods is challenging, since biological and biochemical profiles may be ambiguous due to natural competence and other genetic transfer events (8, 11). Thus, clinical laboratories can encounter "atypical pneumococci" that are optochin resistant, bile insoluble, or unencapsulated (9).

Molecular genetic analyses based on the 16S rRNA gene have provided new insights into the phylogenetic interrelationships of many bacteria (12) and a powerful means for species characterization (13, 14). Unfortunately, the 16S rRNA gene nucleotide sequences from *S. mitis* and *S. oralis* are almost identical to that of *S.* pneumoniae; as such, this gene sequence cannot be used to distinguish between strains at the species level (15).

More-sensitive routine diagnostic methods, such as PCR, could be useful for these bacteria. Several PCR-based methods have been developed for the detection of S. pneumoniae, targeting pneumococcal virulence factors. These factors include autolysin (lytA) (16), pneumolysin (ply) (17), pneumococcal surface antigen A (psaA) (18), manganese-dependent superoxide dismutase (sodA) (19), penicillin-binding protein (20), and a unknown putative gene (21). However, nonpneumococcal isolates containing these genes have also been described, making it difficult to confidently distinguish between S. mitis and S. pneumoniae isolates using these markers (22–24). Furthermore, S. mitis and S. oralis might become evolved from the pathogenic S. pneumoniae by genomic reduction, implying that both inter- and intraspecies recombination events have occurred among these species (25, 26).

Previously, we reported a specific single-step PCR assay to identify S. pneumoniae (27), S. oralis (28), and S. mitis (28). However, simultaneous detection methods specific to S. pneumoniae and the closely related S. mitis and S. oralis strains have not yet been reported. Housekeeping protein-coding genes that are thought to evolve faster than rRNA genes have been proposed as suitable specific markers for the identification and classification of bacteria (29, 30). Among these, DNA gyrase is an essential bacterial enzyme that catalyzes the ATP-dependent negative supercoiling of double-stranded closed-circular DNA. It is composed of two A and two B subunits encoded by the gyrA and gyrB genes, respectively (31). Using the *gyrB* genes for analysis could be more useful than 16S rRNA gene-based methods because there is only a single copy locus in most bacteria, lateral gene transfer of this gene is rare, and the evolution rate is higher than that of the 16S rRNA gene, making it more discriminatory at the species level. In this study, we developed a multiplex PCR (mPCR) protocol targeting the gyrB gene for the rapid, accurate, simple, and species-specific detection of S. pneumoniae, S. mitis, and S. oralis in oral environments and clinical specimens.

MATERIALS AND METHODS

Ethics statement. Oral samples were collected by the Seyee dental clinic (Ulsan, Republic of Korea) and were analyzed in the Department of Microbiology in the College of Medicine at Chung-Ang University. All par-

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TABLE 1 Streptococcus species and other bacteria used in this study

Group	Species	$Strain(s)^a (n = 141)$
	-	
Streptococcus $(n = 115)$	S. pneumoniae S. mitis	KCTC 5080 ^T , CCARM 4009, CCARM 4019, CCARM 4033, CCARM 4109, CCARM 4112, CCARM 4113, CCARM 4114, CCARM 4115, CCARM 4003, CCARM 4005, CCARM 4079, CCARM 4079, CCARM 4084, CCARM 4085, CCARM 4084, CCARM 4106, CCARM 4116, ChDC 4-0134, ChDC 4-0749, ChDC 6-4740, CCARM 4081, CCARM 4091 KCTC 3556 ^T , KCOM 1350, KCOM
		1379, KCOM 1388, ChDC B183, ChDC B186, ChDC B193, ChDC B194, ChDC B227, ChDC B231, ChDC B239, ChDC B242, ChDC B253, ChDC B258, ChDC B260, ChDC B279, ChDC B286, ChDC B303-1, ChDC B315, ChDC B317
	S. oralis	KCTC 13048 ^T , ATCC 9811, DSMZ 20395, DSMZ 20066, ATCC 700233, KCOM 1401, KCOM 1407, KCOM 1408, KCOM 1414, KCOM 1416
	S. pyogenes S. gordonii	KCTC 3984 ^T , KCTC 3096, KCTC 3208 KCTC 3286 ^T , KCOM 1347, KCOM
		1364, KCOM 1369, KCOM 1387, KCOM 1357
	S. infantis	KCOM 1358, KCOM 1375
	S. australis	KCOM 1386, KCOM 1439, KCOM 1441
	S. sinensis	KCOM 1017, KCOM 1427, KCOM1018
	S. sanguinis	KCTC 3284 ^T , KCOM 1567, KCOM 1372, KCOM 1422, KCOM 1428, KCOM 1019, KCOM 1070,
	S. parasanguinis	KCTC 13046 ^T , ChDC B185, ChDC B195, ChDC B215, KCOM 1365, KCOM 1366, KCOM 1370, KCOM 1585
	S. constellatus	ATCC 27823 ^T , ChDC B280, ChDC B284, ChDC B290
	S. intermedius	KCTC 3268 ^T , ChDC KB80, ChDC KB236, ChDC KB717
	S. anginosus	ATCC 33397 ^T , ChDC YA1, ChDC YA3, ChDC YA5, ChDC YA6, ChDC YA7, ChDC YA8, ChDC YA9, ChDC YA10, ChDC YA11, ChDC YA12, ChDC YA13, ChDC B181, ChDC YA201, ChDC B232, ChDC B248, ChDC B252, ChDC B263, ChDC B287, ChDC B311, ChDC B407-1
Other cocci $(n = 26)$	L. lactis subsp. cremoris	DSMZ 40699 ^T
	L. lactis subsp. hordniae	KCTC 3768 ^T
	L. garvieae	LMG 8162 ^T , LMG 8501, LMG 9472, LMG 14494
	L. lactis subsp. lactis E. solitarius	KCTC 3769 ^T , KCTC 3191, KCTC 2013, KCTC 3894, KCTC 3926 KCTC 3553 ^T

TABLE 1 (Continued)

Group	Species	$Strain(s)^a (n = 141)$
	E. hirae	KCTC 3616 ^T
	E. mundtii	KCTC 3630 ^T
	E. malodoratus	KCTC 3641 ^T
	E. cecorum	KCTC 3642 ^T
	E. saccharolyticus	KCTC 3643 ^T
	E. villorum	KCTC 13904 ^T
	E. moraviensis	KCTC 13911 ^T
	E. phoeniculicola	KCTC 3818 ^T
	E. raffinosus	KCTC 5189 ^T
	E. avium	KCTC 5190 ^T
	E. faecalis	KCTC 3206 ^T
	V. fluvialis	LMG 9464 ^T
	V. salmoninarum	LMG 1149 ^T
	V. lutrae	LMG 19537 ^T

^a Abbreviations: KCTC, Korea Collection for Type Culture (Daejeon, Republic of Korea); CCARM, Culture Collection of Antibiotics Resistant Microbe (Seoul, Republic of Korea); ChDC, Chosun University Dental College (Gwangju, Republic of Korea); KCOM, Korean Collection for Oral Microbiology (Gwangju, Republic of Korea); DSMZ, Deutsche Sammlung von Mikroorganismen und Zellkulturen (Braunschweig, Germany); ATCC, American Type Culture Collection (Manassas, VA); BCCM/LMG, Belgian Coordinated Collections of Micro-organisms (Ghent, Belgium).

ticipants gave written informed consent. For all cases, the Chung-Ang University College of Medicine IRB (protocol 2009-12) approved the collection and analysis of all samples.

Bacterial strains and genomic DNA preparation. A total of 141 bacterial strains that were used in this study are listed in Table 1. The 24 S. pneumoniae strains, 20 S. mitis strains, and 10 S. oralis strains were obtained from the Korean Collection for Type Culture (KCTC; Daejeon, Republic of Korea), the Culture Collection of Antibiotic-Resistant Microbes (CCARM; Seoul, Republic of Korea), the Department of Oral Biochemistry, College of Dentistry, Chosun University (ChDC; Gwangju, Republic of Korea), the Deutsche Sammlung von Mikroorganismen und Zellkulturen (DSMZ, Braunschweig Germany), and the American Type Culture Collection (ATCC, Manassas, VA). These strains were used as positive controls for the mPCR assay. Sixty-one other Streptococcus and 26 other coccus strains were obtained from the Culture Collection of the University of Gothenburg (CCUG; Gothenburg, Sweden), the Belgian Coordinated Collections of Micro-organisms (BCCM/LMG Ghent, Belgium), KCTC, and DSMZ. These strains were used as negative controls. Oral streptococcus strains were grown on 5% sheep blood agar plates (Asan Pharm Co., Seoul, Republic of Korea) at 37°C for 20 h, while the other strains were cultured on brain heart infusion agar plates (Difco Laboratories, Detroit, MI) under the same conditions. Bacterial genomic DNA was prepared by the cetyltrimethylammonium bromide method (32). The purified DNA was quantified using a Nano Quant Infinite M200 spectrophotometer (Tecan, Männedorf, Switzerland) at a wavelength of 260 nm.

Design of multiplex PCR primers. The oligonucleotide primer sets, sL1 (5'-GGCTTAGAGGCTGTTCGT-3'), sR1 (5'-TCACTTCCCACTTT AACCC-3'), sR2 (5'-AGTTTGTTCTAGCCCCTCA-3'), and sR3 (5'-AT CTCACCGTCTGTATAGA-3'), for mPCR amplification of the *gyrB* region were designed according to the *gyrB* nucleotide sequences of *S. pneumoniae* JJA (GenBank accession number NC_012466), *S. pneumoniae* P1031 (NC_012467), *S. pneumoniae* 70585 (NC_012468), *S. pneumoniae* ATCC 700669 (NC_011900), *S. mitis* B6 (NC_013853), *S. oralis* Uo5 (NC_015291), and *S. oralis* Uo5 (NC_015291) published in the NCBI GenBank database, using Oligo V6 software (Molecular Biology Insights, Cascade, CO). The *gyrB* gene of *S. pneumoniae* KCTC 5080^T and the primers are illustrated in Fig. 1. During the mPCR process, amplicon sizes were selected to enable the identification of *S. pneumoniae*, *S. mitis*, and *S. oralis* species, and the amplification conditions were optimized in

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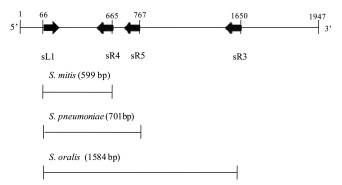


FIG 1 The *gyrB* gene of *S. pneumoniae* KCTC 5080^T and primers sL1, sR1, sR2, and sR3 for multiplex PCR amplification and nucleotide sequence analysis.

terms of primer concentrations, annealing temperatures, and other thermocycling conditions.

Establishment of the multiplex PCR assay. Three reference strains, S. pneumoniae KCTC 5080^T, S. mitis KCTC 3556^T, and S. oralis 13048^T, were used to develop and evaluate the mPCR assay in combination with each other; namely, S. pneumoniae, S. mitis, and S. oralis alone, S. pneumoniae plus S. mitis, S. pneumoniae plus S. oralis, S. mitis plus S. oralis, and S. pneumoniae plus S. mitis plus S. oralis. The mPCR reaction was conducted in a 20-µl-total reaction mixture containing 100 ng genomic DNA template, 1 µl each primer, 2 µl 10× reaction buffer, 0.2 mM deoxynucleoside triphosphates (dNTP)s, 1.5 mM MgCl₂, and 2.5 U Taq polymerase. Primer concentrations were selected as 1, 2, 3, 5, 10, 15, 20, 25, and 30 µM each primer (sL1, sR1, sR2, and sR3). PCR amplification was performed using a TGradient thermal cycler (Biometra, Goettingen, Germany) with the following conditions: 94°C for 5 min, followed by 30 cycles of 94°C for 30 s, annealing temperature for 30 s, and 72°C for 40 s and a final extension at 72°C for 10 min. Ten gradient temperatures, including 49°C, 50°C, 51°C, 52°C, 53°C, 54°C, 55°C, 56°C, 57°C, 58°C, 59°C, and 60°C were used to optimize the annealing temperature. After PCR amplification, 5 μ l of each PCR product was resolved using 1.2% SeaKem LE agarose gel (FMC Bioproducts, Rockland, ME), followed by ethidium bromide staining. A negative-control experiment was performed using sterile water instead of culture or DNA template. Electrophoresis was used to separate the amplicons, and DNA bands were viewed under a GelDoc XR image analysis system (Bio-Rad, Hercules, CA).

DNA sequence analysis. All PCR products were evaluated by sequence analysis. Cycle sequencing was performed using the BigDye terminator version 3.1 cycle sequencing kit, and the sequencing reactions were analyzed using an automated DNA sequencer (model 3730; Applied Biosystems, Foster City, CA). Assembly and editing of nucleotide sequences were performed using BioEdit (33) software and the CLUSTAL_X 1.81 program (34). The nucleotide sequence homologies of each amplified PCR product were evaluated using BLAST searches of the National Center for Biotechnology Information (NCBI) databases.

Specificity and sensitivity of the multiplex PCR assay. The specificity of the mPCR primers was examined using DNA templates from each of the 24 *S. pneumoniae*, 20 *S. mitis*, 10 *S. oralis*, 61 other *Streptococcus*, and 26 other coccus strains. mPCR was performed with 100 ng genomic DNA template in a 20-µl reaction mixture containing 1 µl of each primer (sL1, sR1, sR2, and sR3 at 20 µM, 20 µM, 1 µM, and 2 µM, respectively), 2 µl $10\times$ reaction buffer, 0.2 mM dNTPs, 1.5 mM MgCl₂, and 2.5 U *Taq* polymerase. The sensitivity of the PCR assay was evaluated using a 10-fold series of purified DNA from a mixed culture of *S. pneumoniae* KCTC $5080^{\rm T}$, *S. mitis* KCTC $3556^{\rm T}$, and *S. oralis* $13048^{\rm T}$ (each at $10^{\rm 2}$ CFU/ml). Amplification was conducted using the GeneAmp PCR system 9700 (Applied Biosystems, Foster City, CA) with PCR cycling conditions consisting of an initial denaturation at $94^{\rm e}$ C for 5 min, followed by 30 cycles at $94^{\rm e}$ C for 30 s, $58^{\rm e}$ C for 30 s, and $72^{\rm e}$ C for 40 s, with final extension at $72^{\rm e}$ C for 10

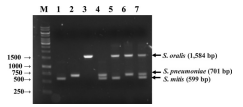


FIG 2 Electrophoretic examination of multiplex PCR products. Lane M, 1 kb plus DNA ladder (Life Technologies, Rockville, ME, U.S.A.); lane 1, *S. mitis* KCTC 3556^T; lane 2, *S. pneumoniae* KCTC 5080^T; lane 3, *S. oralis* KCTC 13048^T; lane 4, *S. mitis* KCTC 3556^T plus *S. pneumoniae* KCTC 5080^T; lane 5, *S. mitis* KCTC 3556^T plus *S. oralis* KCTC 13048^T; lane 6, *S. pneumoniae* KCTC 5080^T plus *S. oralis* KCTC 13048^T; lane 7, *S. mitis* KCTC 3556^T plus *S. pneumoniae* KCTC 5080^T plus *S. oralis* KCTC 13048^T.

min. After PCR amplification, 5 µl of each PCR product was resolved using agarose gel electrophoresis as described above.

Pilot application study using clinical samples. For a pilot application study, 47 samples collected from the oral cavity of outpatients in the Seyee Dental Clinic were used to test the mPCR protocol. The median age of patients was 32 years old (range, 4 to 78 years). All samples were taken using a 3M Quick swab, and genomic DNA was isolated as described previously. One microliter ($10~\mu g/ml$) of each DNA sample was subjected to species-specific detection for *S. pneumoniae, S. mitis*, and *S. oralis*. At the same time, $100~\mu l$ of each dilution was cultured on 5% sheep blood agar plates (Asan Pharmaceutical, Seoul, Republic of Korea) and the 3 oral pathogens were identified according to standard microbiological methods, including the API 20 Strep method (BioMérieux SA, Marcy l'Etoile, France).

RESULTS AND DISCUSSION

Optimization of the multiplex PCR assay. Since mPCR protocols have been developed for the detection of *S. pneumoniae* and other respiratory pathogens, including S. pneumoniae, Haemophilus influenzae, Mycoplasma pneumoniae, and Chlamydophila pneumoniae (35), or for serotyping of S. pneumoniae (36), no protocol has been developed for the simultaneous detection of S. pneumoniae and its closest phylogenetic relatives, S. mitis and S. oralis. In order to determine the optimal conditions for mPCR, genomic DNA extracted from 7 strain combinations, including S. pneumoniae, S. mitis, S. oralis, S. pneumoniae plus S. mitis, S. pneumoniae plus S. oralis, S. mitis plus S. oralis, and S. pneumoniae plus S. mitis plus S. oralis, were used as target templates. The discrimination of S. pneumoniae, S. mitis, and S. oralis is shown in Fig. 2. The amplification generated 3 PCR products as detected on agarose gels, comprising 701-bp, 599-bp, and 1,584-bp amplicons for S. pneumoniae, S. mitis, and S. oralis, respectively. No size variation was detected among the strains investigated. The PCR product showed the highest levels of resolution of specificity when primer concentrations of sL1, sR1, sR2, and sR3 were at 20 µM, 20 μM, 1 μM, and 2 μM and amplification was at an annealing temperature of 56°C compared to the specificity of products obtained under other conditions. The identity of PCR amplicons exactly matched the gyrB gene of each species as determined using nucleotide sequencing and BLAST homology searches.

Specificity of the multiplex PCR assay. The specificity of the mPCR amplification was determined using DNA extracted from 24 *S. pneumoniae*, 20 *S. mitis*, and 10 *S. oralis* strains and 87 other strains. The amplified PCR products from the different bacterial strains resulted in the expected 701-bp PCR product common to the 24 *S. pneumoniae* strains, including type strains and clinical

TABLE 2 Limits of detection for genomic DNA in combination with *S. pneumoniae* KCTC 5080^T, *S. mitis* KCTC 3556^T, and *S. oralis* 13048^T in multiplex PCR

			PCR results ^a		
Lane	DNA concn (ng/µl)	Cell no.	S. pneumoniae	S. mitis	S. oralis
1	6.3×10^{0}	1×10^{7}	+	+	+
2	6.3×10^{-1}	1×10^{6}	+	+	+
3	6.3×10^{-2}	1×10^{5}	+	+	+
4	6.3×10^{-3}	1×10^{4}	+	+	+
5	6.3×10^{-4}	1×10^{3}	+	ND	+
6	6.3×10^{-5}	1×10^{2}	ND	ND	+
7	6.3×10^{-6}	1×10^{1}	ND	ND	ND
8	Negative control	0	ND	ND	ND

^a ND, not detected.

isolates. All S. mitis and S. oralis strains yielded specific amplifications for the 599-bp or 1,548-bp PCR products, respectively, showing positive results. This product was specifically detected because no amplifications of PCR products from the 87 strains of other closely related streptococci or Gram-positive cocci were detected (data not shown). In the present study, the following strains were mPCR negative based on gyrB amplification: 61 VGS strains, including 4 Streptococcus constellatus, 4 Streptococcus intermedius, 21 Streptococcus anginosus, 6 Streptococcus gordonii, 2 Streptococcus infantis, 3 Streptococcus australis, 3 Streptococcus sinensis, 7 Streptococcus sanguinis, and 8 Streptococcus parasanguinis; other streptococcus strains; 3 Streptococcus pyogenes strains; 11 lactococcus strains, including 1 Lactococcus lactis subsp. cremoris, 1 Lactococcus lactis subsp. hordniae, 4 Lactococcus garvieae, and 5 Lactococcus lactis subsp. lactis strains; 12 enterococcus strains, including 1 Enterococcus solitaries, 1 Enterococcus hirae, 1 Enterococcus mundtii, 1 Enterococcus malodoratus, 1 Enterococcus cecorum, 1 Enterococcus saccharolyticus, 1 Enterococcus villorum, 1 Enterococcus moraviensis, 1 Enterococcus phoeniculicola, 1 Enterococcus raffinosus, 1 Enterococcus avium, and 1 Enterococcus faecalis strain; and 3 vagococcus strains, including 1 Vagococcus fluvialis, 1 Vagococcus salmoninarum, and 1 Vagococcus lutrae strain.

Because *S. pneumoniae* shares over 99% 16S rRNA gene sequence homology with other VGS, the identification of novel genetic markers specific for *S. pneumoniae* is needed for accurate diagnosis of pneumococcal disease. However, this organism evolved closely with related VGS from a common genetic ancestor (5, 9, 10); therefore, serious specificity problems were noted with these assays, including a false-positive amplification with the *ply* (17) and a putative noncoding region Spn9802 and Spn9828 (21) primer sets from the genomic DNA of *S. mitis*, *S. gordonii*, and *S. sanguinis*, while a few *S. pneumoniae* strains did not amplify with the *lytA* primers (27). Even though the frequency was probably low in the mPCR protocol, 3.3% of *S. mitis* strains shared the *lytA* gene with *S. pneumoniae* (35).

More recently, the *cpsA* gene was identified as a novel genomic marker specific to *S. pneumoniae* and a quantitative PCR (qPCR) assay was developed to detect and enumerate this pathogen (37). In *S. oralis*, the *rgg* gene, known to be involved in the biosynthesis of glucosyltransferase, allowed for the easy and reliable discrimination of *S. oralis* from other streptococci (38). Thus, it should be noted that in our study, we were able to fully discriminate *S. pneumoniae*, *S. mitis*, and *S. oralis* species from one another and from

TABLE 3 Results of the multiplex PCR protocol compared with culture in a pilot application study using human oral samples

	No. of positive samples/no. of samples tested ($n = 47$)		
Species	Multiplex PCR protocol	Culture	
S. pneumoniae	4/47	1/47	
S. mitis	6/47	3/47	
S. oralis	7/47	1/47	
S. pneumoniae + S. mitis	2/47	1/47	
S. pneumoniae + S. oralis	2/47	1/47	
S. mitis + S. oralis	3/47	0/47	
S. pneumoniae $+$ S. mitis $+$ S. oralis	0/47	0/47	
Negative	23/47	40/47	

related streptococci by the current mPCR protocol, promising the rapid diagnosis of streptococcal diseases.

Sensitivity of multiplex PCR assay. The sensitivity of mPCR using the *gyrB* gene-based primers was determined by analyzing 10-fold serially diluted genomic DNA (6.3 ng to 6.3 fg) extracted from each *S. pneumoniae* KCTC 5080^T, *S. mitis* KCTC 3556^T, and *S. oralis* 13048^T culture at 10⁷ CFU ml⁻¹. The detection limit of the conventional PCR assay was 630 fg with *S. pneumoniae* DNA, 6.3 pg with *S. mitis* DNA, and 63 fg with *S. oralis* DNA (Table 2).

Genome equivalents were calculated assuming one molecule of S. pneumoniae, S. mitis and S. oralis DNA. Considering a genome size of 2.1 Mb for *S. pneumoniae* and *S. mitis*, a corresponding 2.2 fg of DNA was calculated and 2.1 fg was determined for S. oralis DNA based on a 2.0-Mb genome. These numbers were determined according to the following equation: DNA amount in fg = bp \times 660 Da/bp \times 1.6 \times 10⁻²⁷ kg/Da \times 1 \times 10⁻¹⁸ fg/kg (39). Considering a genome size of 2.1 Mb as determined for S. pneumoniae (GenBank accession number FM_211187), S. mitis (GenBank accession number NC_013853), and S. oralis (GenBank accession number NC_015291) DNAs, the number of genomic copies in the nucleic acid extracts from each strain was determined using the following formula: genome copies = quantity of DNA in extract/2.2 fg or quantity of DNA in extract/2.1 fg. Therefore, we concluded that the minimum limits of detection of 630 fg, 6.3 pg, and 63 fg were equivalent to approximately 286, 2,863, and 30 S. pneumoniae, S. mitis, and S. oralis genomes, respectively.

Application of multiplex PCR in clinical samples. The oral cavity is a reservoir of bacteria comprising more than 700 species or phylotypes, of which approximately 35% have not been cultured (40). The VGS, a major oral streptococcal group, is known for its low pathogenicity and virulence, but several prospective studies have shown positive associations between oral inflammation and increased risk of cardiovascular and cerebrovascular disease (41), preterm birth (42), and certain cancers (43). Recently, additional studies have shown that *S. mitis* is implicated in pancreatic disease, including autoimmune pancreatitis and pancreatic ductal adenocarcinoma (7).

Table 3 shows the results of the current mPCR protocol and traditional culture methods for the 47 oral samples used in our pilot application study. mPCR identified *S. pneumoniae*, *S. mitis*, and *S. oralis* and a combination of these bacteria in all 7 culture-positive and in an additional 40 culture-negative oral samples.

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Among the 47 patients, mPCR and culture techniques identified *S. pneumoniae* in 4 samples (8.5%) and in 1 (2.1%) sample, respectively. *S. mitis* presented in 6 (12.8%) based on mPCR and in 3 (6.4%) based on culture. *S. oralis* presented in 7 (14.9%) based upon mPCR and in 1 (2.1%) based on culture. In the combination with 3 pathogens, *S. pneumoniae* and *S. oralis* were detected in 2 (4.3%) based on mPCR and in 1 (2.1%) based on culture. *S. pneumoniae* and *S. mitis* were revealed in 2 (4.3%) based on mPCR and in 1 (2.1%) based on culture. *S. oralis* and *S. mitis* were revealed in 3 (6.4%) based on mPCR and were not identified in culture. Detection of the three species was not revealed by mPCR or culture in this study.

Because the 16S rRNA gene evolves so slowly, phylogenetic information based on this molecule may not always be sufficient to distinguish closely related species or to resolve their evolutionary relationships. In addition, when several copies of the 16S rRNA gene are present, sequence heterogeneity results in ambiguities in the sequence chromatograms derived from direct sequencing of the PCR products (44). Nucleotide sequences of housekeeping protein-coding genes evolve more rapidly than 16S rRNA and may represent useful alternatives or complements to the 16S rRNA gene (45-47). Otherwise, housekeeping genes have been considered good monitoring tools for bacterial identification and strain typing. The gyrB gene-based mPCR presented in this study may be valuable in clinical microbiology laboratories because it provides a fast and cost-effective means of analyzing large numbers of samples and allows the detection and discrimination of S. pneumoniae, S. oralis, and S. mitis from oral streptococci. Our protocol proved to be more accurate and sensitive than traditional culture methods.

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