Multiplex PCR Assay for Detection of *Streptococcus suis* Species and Serotypes 2 and 1/2 in Tonsils of Live and Dead Pigs

C. Marois,1,* S. Bougeard,2 M. Gottschalk,3 and M. Kobisch1

Unité de Mycoplasmologie-Bactériologie1 and Unité d’Épidémiologie Porcine et Assurance Qualité,2 Agence Française de Sécurité Sanitaire des Aliments, 22440 Ploufragan, France, and Groupe de Recherche sur les Maladies Infectieuses du Porc, Faculté de Médecine Vétérinaire, Université de Montréal, St. Hyacinthe, Québec, Canada, J2S7C6

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A PCR assay was developed for the detection of *Streptococcus suis* serotypes 2 and 1/2. This multiplex PCR is based on the amplification of the gene coding for 16S rRNA of *S. suis* and on the amplification of the *cps2J* gene coding for the capsule of *S. suis* serotypes 2 and 1/2. An internal control was constructed and added in this test to monitor the efficiency of amplification in each reaction. To evaluate the specificity of the test, 31 strains of other bacterial species related to *S. suis* or isolated from pigs and 42 strains of *S. suis* serotypes 1 and 3 to 34 were analyzed. The detection threshold of the test was 28 *S. suis* CFU/ml. The specificity and the sensitivity of the multiplex PCR test and the presence of an internal control allowed the analysis of biological samples without a culture step. The PCR assay was then applied to the detection of 14 *S. suis* serotype 1/2 strains, 88 *S. suis* serotype 2 strains isolated from pigs, and 25 *S. suis* serotype 2 strains isolated from humans. This test was also applied to analyze tonsil samples of pigs experimentally infected and carrier pigs without any symptoms.

*Streptococcus suis* is an important pathogen of swine, causing meningitis, arthritis, pericarditis, polyserositis, septicaemia, and sudden death of weaning piglets as well as growing pigs (15). Moreover, *S. suis* may be isolated from healthy pigs, and these animals are a source of *S. suis* transmission in pig herds. This bacterium is also a zoonotic agent responsible for meningitis, septicaemia, arthritis, and endocarditis in humans. Most cases have involved individuals who had occupational exposure to pigs, like butchers, slaughterhouse workers, veterinarians, and pig farmers (12, 18, 28, 29). More recently, several cases of human *S. suis* infection acquired from wild boars have been reported (2, 10, 21, 22).

Thirty-five capsular serotypes of *S. suis* have been described (types 1/2 and 1 through 34) (27). Serotypes 2, 1/2, 9, 7, and 3 are usually isolated in France from diseased or dead pigs, mainly in cases of meningitis, arthritis, and septicaemia (5). Although serotype 2 is considered to be the most-virulent serotype in most countries, strains belonging to other serotypes can also cause disease in pigs (9, 15).

Currently, bacteriological techniques are routinely used to detect *S. suis*. Recently, PCR tests were developed. Monoplex PCR tests, based on sequences of type-specific capsular genes of *S. suis*, were developed to detect specifically serotypes 2 and 1/2, 1 and 14, 7 and 9 (25, 26). Then, these methods were changed into multiplex PCR tests (31). A test based on amplification of the *epf* gene encoding the extracellular factor proteins of virulent serotype 2 was also described previously (30). In 1999, Okwumabua et al. described a PCR assay based on the gene encoding the suilysin of *S. suis* type 2, but this target was not conserved across capsular types or pathogenic strains (19). In 2000, Boye et al. described a method to detect *S. suis* by in situ hybridization with a species-specific probe targeting 16S rRNA (6). This method does not permit the detection of serotypes 32 through 34. In 2001, tRNA intergenic length polymorphism analysis (tDNA-PCR) combined with capillary electrophoresis was described by Baele et al. to identify streptococci, including three *S. suis* strains (1). In 2003, a multiplex PCR test based on *S. suis* *cps* genes specific to serotypes 2 (1/2), 1 (and 14), 7, and 9 and on the *gdh* gene encoding the glutamate dehydrogenase of *S. suis* serotype 2 was developed by Okwumabua et al. (20). This PCR assay allowed the amplification of all serotypes of *S. suis* with the target based on the *gdh* gene. However, this method was only applied to detect or characterize *S. suis* from pure cultures.

In the present study, we report the development of a multiplex PCR test to detect *S. suis* species and serotypes 2 and 1/2 from tonsillar specimens, sampled from live or dead animals, without a culture step. An internal control was constructed and added in the multiplex PCR to monitor the efficiency of amplification in each reaction. Compared to bacteriological methods, the PCR assay was fast and sensitive. This PCR assay was used to study *S. suis* infections using two *S. suis* serotype 2 strains and one *S. suis* serotype 1/2 strain in experimentally infected specific-pathogen-free (SPF) piglets.

**MATERIALS AND METHODS**

**Bacterial strains.** The specificity of the PCR assay was tested with a collection of 203 strains representing 172 *S. suis* strains belonging to one of the 35 capsular types described as well as 25 bacterial species other than *S. suis* (Table 1). For some experiments, the reference strain S735 was also used. In addition, three French porcine field strains of *S. suis* were used for experimental infections: *S. suis* capsular serotype 2 strains 332 and 347, isolated from septicaemia and from palatine tonsils of clinically healthy pigs, respectively, and *S. suis* capsular serotype 1/2 (strain 353) isolated from tonsils of a clinically healthy pig.

*S. suis* sp. and *Actinobacillus lignieresii* were cultivated on Columbia agar base supplemented with 5% sheep blood (AES Laboratories, Combourg, France), *Mycoplasma haemofelis*, *Mycoplasma hyopneumoniae*, and *Mycoplasma hyorhinis* were cultivated on Fries medium (13). *Campylobacter coli* and *Campylobacter jejuni* were cultivated as previously described (17). The other strains were cultivated on pleuropneumoniae-like organism agar (PPLO agar; Difco, Cergy Pontoise, France) supplemented with nicotinamide dinucleotide (10 μg/
M sodium acetate buffer (pH 5.5) and 400 µl of isooamyl alcohol (25:24:1), vortexed, and centrifuged at 10,000 × g for 15 min. The DNA precipitate was redissolved in 10 µl of distilled water.

Samples were prepared for PCR as described by Kellogg and Ksow (16). Briefly, 1 ml of each initial suspension (15S) were centrifuged (12,000 × g, 4°C, 20 min) and the pellets were resuspended in a mixture of 250 µl of 10 mM Tris HCl (pH 8.3), 2.5 mM MgCl 2, and 250 µl of 10 mM Tris HCl (pH 8.3), 2.5 mM MgCl 2. 1% (vol/vol) Tween 20 (Sigma-Aldrich Chimie), 0.01% (vol/vol) Nonidet P-40 (Sigma-Aldrich Chimie), and 1% (vol/vol) Triton X-100 (Sigma-Aldrich Chimie), and proteinase K (120 µg/ml) Sigma-Aldrich Chimie). Samples were incubated for 1 h at 60°C prior to proteinase K heat inactivation at 95°C for 10 min, allowed to cool at room temperature, and kept at −20°C.

When inhibition of the PCR was observed, DNA was reextracted as follows. Four hundred microliters of lysate was placed with 400 µl cool at room temperature, and kept at 60°C for 10 min, allowed to cool at room temperature, and kept at −20°C.

When inhibition of the PCR was observed, DNA was reextracted as follows. Four hundred microliters of lysate was placed with 400 µl of Tris HCl (pH 8.3), 2.5 mM MgCl 2, and 250 µl of 10 mM Tris HCl (pH 8.3), 2.5 mM MgCl 2. 1% (vol/vol) Tween 20 (Sigma-Aldrich Chimie), 0.01% (vol/vol) Nonidet P-40 (Sigma-Aldrich Chimie), and proteinase K (120 µg/ml) Sigma-Aldrich Chimie). Samples were incubated for 1 h at 60°C prior to proteinase K heat inactivation at 95°C for 10 min, allowed to cool at room temperature, and kept at −20°C.

When inhibition of the PCR was observed, DNA was reextracted as follows. Four hundred microliters of lysate was placed with 400 µl of phenol–chloroform–isoamyl alcohol (25:24:1), vortexed, and centrifuged at 10,000 × g for 30 s. Then, the supernatant was mixed with 400 µl of chloroform–isoamyl alcohol (24:1), vortexed, and centrifuged, and the supernatant was mixed with 50 µl of 3 M sodium acetate buffer (pH 5.5) and 400 µl of isopropanol for 30 min at 4°C to precipitate the DNA. After centrifugation at 10,000 × g for 15 min, the DNA pellet was washed with 70% ethanol, dried, and resuspended in 50 µl of double-distilled water.

Construction of the PCR IPC. To check for the presence of inhibitors within the PCR mixture, an internal positive control (IPC) was constructed. IPC was synthesized in one PCR. The primers used in this reaction (CI 6-s and CI 7-as) possessed 5’ overhanging ends which were identical to the primers used in the PCR specific for S. suis serotype 2 (cps2J-s and cps2J-as), whereas their 3’ ends were complementary to a predetemined DNA sequence (16S ribosomal DNA [rDNA]) of S. suis of defined length and sequence (Table 2) (AF009477). The IPC sequence was different from the 16S rDNA sequence amplified by the 16S rDNA primers used in the multiplex PCR test.

The IPC was a 16S rDNA fragment of 620 bp from S. suis generated by PCR. The PCR mixture contained PCR buffer II (20 mM Tris HCl [pH 8.4], 50 mM KCl, 1% glycerol, Thermostable AccuPrime protein, 1.5 mM MgCl 2, 200 µM [each] deoxynucleoside triphosphate), 400 nM (each) CI 6-s and CI 7-as primers, 1 U of AccuPrime Taq DNA polymerase (Invitrogen, Cergy Pontoise, France), and 5 µl of a cell lysate of pure culture of S. suis 735 reference strain. Amplification was performed in a Perkin-Elmer Cetus (Courtaboeuf, France) GeneAmp PCR system 9600. The reaction procedure consisted of 40 cycles of denaturation at 94°C for 30 s, primer annealing at 65°C for 25 s, and extension at 72°C for 10 s. The PCR product was purified using a commercially available kit (Life Technologies, Cergy Pontoise, France). This IPC was stored in double-distilled water at −20°C. DNA concentration was determined spectrophotometrically.

Multiplex PCR conditions. The multiplex PCR developed in this study permitted the simultaneous detection of the S. suis species and serotypes 2 and 1/2. The 294-bp PCR product, specific to S. suis, was obtained with the forward primer 16S-195(s) and the reverse primer 16S-489(as) (Table 2) defined on the 16S rDNA sequence (AF009477) (8). The second primer set, detecting serotypes 2 and 1/2, was composed of a forward primer, cps2J-s, and a reverse primer, cps2J-as (Table 2), and enabled the amplification of 459-bp products. These

<table>
<thead>
<tr>
<th>Species (Strain(s))</th>
<th>No. of strains tested (n = 203)</th>
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<tbody>
<tr>
<td>Streptococcus suis serotype 1</td>
<td>Reference strain 5428&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Streptococcus suis serotype 1/2</td>
<td>Reference strain 2651&lt;sup&gt;b&lt;/sup&gt; and field strains&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>Streptococcus suis serotype 2</td>
<td>Field strains isolated from pigs&lt;sup&gt;d&lt;/sup&gt;</td>
</tr>
<tr>
<td>Streptococcus suis serotypes 3 to 34</td>
<td>Field strains isolated from humans&lt;sup&gt;e&lt;/sup&gt;</td>
</tr>
<tr>
<td>Streptococcus agalactiae</td>
<td>ATCC 13813</td>
</tr>
<tr>
<td>Streptococcus pyogenes</td>
<td>ATCC 43138 and field strain</td>
</tr>
<tr>
<td>Escherichia coli</td>
<td>Field strains</td>
</tr>
<tr>
<td>Campylobacter jejuni</td>
<td>Field strain</td>
</tr>
<tr>
<td>Campylobacter coli</td>
<td>Field strain</td>
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<tr>
<td>Mycoplasma hyopneumoniae</td>
<td>Field strain</td>
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<tr>
<td>Mycoplasma hyorhinis</td>
<td>Field strain</td>
</tr>
<tr>
<td>Mycoplasma flocculare</td>
<td>Field strain</td>
</tr>
<tr>
<td>Actinobacillus pleuropneumoniae&lt;sup&gt;f&lt;/sup&gt;</td>
<td>ATCC 27088 and field strain</td>
</tr>
<tr>
<td>Actinobacillus lignieresii</td>
<td>ATCC 49236</td>
</tr>
<tr>
<td>Actinobacillus rossii&lt;sup&gt;f&lt;/sup&gt;</td>
<td>ATCC 27072</td>
</tr>
<tr>
<td>Pasteurella multocida</td>
<td>Field strain</td>
</tr>
<tr>
<td>Staphylococcus aureus</td>
<td>Field strain</td>
</tr>
<tr>
<td>Streptococcus agalactiae</td>
<td>ATCC 43765, ATCC 700794, ATCC 700796&lt;sup&gt;f&lt;/sup&gt;</td>
</tr>
<tr>
<td>Field strains isolated from pigs&lt;sup&gt;d&lt;/sup&gt;</td>
<td>3</td>
</tr>
<tr>
<td>Field strains isolated from humans&lt;sup&gt;e&lt;/sup&gt;</td>
<td>25</td>
</tr>
<tr>
<td>Streptococcus porcinus</td>
<td>ATCC 43138 and field strain</td>
</tr>
<tr>
<td>Streptococcus suis</td>
<td>ATCC 12344</td>
</tr>
</tbody>
</table>

<sup>a</sup> Groupe de Recherche sur les Maladies Infectieuses du Porc, Faculté de Médecine Vétérinaire, Université de Montréal, St. Hyacinthe, Québec, Canada (8).
<sup>b</sup> ATCC, American Type Culture Collection, Manassas, Va.
<sup>c</sup> Field strains isolated in France, including strains 332 and 347 used in the experimental infection.
<sup>d</sup> Field strains isolated in France, The Netherlands, Canada, and the United Kingdom.
<sup>e</sup> Field strains isolated in France.
<sup>f</sup> NCDO, National Collection of Dairy Organisms, Shinfield, Reading, United Kingdom.
<sup>g</sup> CCUG, Culture Collection University of Göteborg, Göteborg, Sweden.
primers were defined on the capsule gene cps (AF118389), and they also am-
plified the IPC (620 bp). The multiplex PCR mixture contained PCR buffer (67 mM Tris-HCl, 16 mM (NH₄)₂SO₄, 0.01% Tween 20, 2.5 mM MgCl₂ (pH 8.8)), a 600 μM concentration of each deoxynucleoside triphosphate (Pharmacia Biotech, Orsay, France), 1.1 μM cps2s-1 and cps2s-as primers, 600 nM 16S-19s(s) and 16S-489(as2) primers, 2.5 U of Taq DNA polymerase (Eurobio, Les Ulis, France), 3 fg of IPC, and 5 μl of the DNA template. The DNA template was replaced by double-distilled water for the negative control. Amplification was performed in a Perkin-Elmer Cetus GeneAmp PCR system 9600. The reaction procedure consisted of 40 cycles of denaturation at 94°C for 30 s, primer annealing at 60°C for 30 s, and extension at 72°C for 60 s. The amplified products were separated in a 2% agarose gel in TBE buffer (90 mM Tris, 90 mM borate, 2.5 mM EDTA [pH 8]) for 1 h at a constant voltage of 125 V. Amplified products were stained with ethidium bromide and detected by UV transillumination. The Smart Ladder was used as a molecular size standard (Eurogentec, Angers, France).

Sensitivity of the multiplex PCR. The sensitivity of the PCR test was evaluated using 10-fold dilutions of a culture of S. suis reference strain serotype 2 (7375) at a titer of 2.8 × 10⁶ CFU/ml. Then, each dilution (1 ml) was placed on tonsil biopsy specimens (6 mm²) obtained from S. suis-free animals and reduced into small pieces with a scalpel, and DNA was prepared as mentioned above (16).

Experimental infection. All bacterial strains were prepared under the same conditions: one colony, isolated from an overnight culture on Columbia blood agar base, supplemented with 5% sheep blood, was resuspended in 5 ml of Todd-Hewitt broth (THB) (Difco) and incubated 18 h at 37°C, in 5% CO₂. The bacterial cultures were then diluted in THB with 10% inactivated bovine serum, further incubated 6 h, and adjusted to 10⁸ CFU/ml. For each inoculum, the bacterial cultures were then diluted in THB with 10% inactivated bovine serum, C, in 5% CO₂. The Todd-Hewitt broth (THB) (Difco) and incubated 18 h at 37°C. Agar base, supplemented with 5% sheep blood, was resuspended in 5 ml of sterile water supplemented with NaCl (8.5 g/liter) (SW), and analyzed by classical bacteriological analysis and by PCR.

In each unit, 6 days before infection and on days 8, 16, and 21 p.i., 7 g of dehydrated granulated food, 15 g of feces, and 25 ml of drinking water were collected in four distant places. In each unit, four drag swabs (Sodibox, La Foret, France), previously humidified with 5 ml of SW, were rubbed on the pigs’ clothes, and compulsory showering after visiting the pigs. Food, feces, and dust samples were collected in four distant places. In each unit, four drag swabs (Sodibox, La Foret, France), previously humidified with 5 ml of SW, were rubbed on the pigs’ clothes, and compulsory showering after visiting the pigs.

Serological analysis. Ten microliters of each sample (or IS) was placed onto selective basis Columbia medium supplemented with 5% sheep blood, 15 mg of nalidixic acid/liter, and 10 mg of colistin/liter. Then, the plates were incubated overnight at 37°C in 5% CO₂. S. suis-like colonies were subcultivated on Columbia medium supplemented with 5% sheep blood, identified by PCR, and serotyped by slide agglutination using a type-specific hyperimmune serum (14).

FIG. 1. Experimental design. Negative control animals were in unit A and were divided into two groups: group 1 (six noninfected pigs that received sterile THB) and group 2 (six noninfected pigs that were in direct contact). In unit B, six pigs were infected with S. suis strain 332 (group 3) and six pigs were in direct contact (group 4). In unit C, six pigs were infected with S. suis strain 347 (group 5) and six pigs were in direct contact (group 6). In unit D, six pigs were infected with S. suis strain 353 (group 7) and six pigs were in direct contact (group 8).
control corresponded to a serum from an uninfected SPF pig (tested in quadruplicate), o-Phenylenediamine at 0.4 mg/ml (Sigma-Aldrich Chimie) dissolved in 0.05 M citrate buffer (pH 5.5) with 0.5 M H2O2 was used as a substrate. After 15 min at 37°C in the dark, 25 μl of HCl was added in each well and optical density (OD) was measured at 492 nm on a kinetic microplate reader (Labsystems, Cergy-Pontoise, France). Results were reported as S/P ratios, which were defined as the OD obtained for each serum minus the mean OD of the negative control divided by the mean OD of the positive control.

Statistical analysis of data. The data obtained in experimental study from pig groups were compared simultaneously by Kruskall-Wallis test and in two-by-two tables by Kolmogorov-Smirnov test. Associations between strain used for the multiplex PCR, the different microorganisms listed in Table 1 were used as DNA templates. All S. suis strains, including reference strains as well as S suis serotypes 2 and 1/2 strains, were used for the assay, tonsillar carriership (as determined by external swab, biopsy sample, and whole tonsil), and method applied (PCR and bacteriological analysis) were assessed by Fisher exact test (n ≤ 5) or chi-square test (n > 5) on independence in two-by-two tables. These tests were carried out with Systat 9.0 program for Windows. Differences were estimated to be significant when probabilities (P) were lower than 0.05.

RESULTS

Specificity and sensitivity of the multiplex PCR. In order to assess the specificity of the multiplex PCR, the different microorganisms listed in Table 1 were used as DNA templates. All S. suis strains, including reference strains as well as field strains, showed a fragment of 294 bp corresponding to a part of the 16S rDNA gene. None of the other bacterial species described in Table 1 showed any amplification product. A fragment of 489 bp corresponding to a part of the cps gene coding for the capsule was only obtained with S. suis serotypes 2 and 1/2 strains as expected.

Sensitivity results are shown in Fig. 2. The assay was carried out with 10-fold dilutions of S. suis DNA extracts obtained from a culture of reference strain S735 at a titer of 280 CFU/μl and mixed with a tonsil palatine biopsy specimen from an SPF pig (1 ml of dilution by biopsy). Under the conditions described in the trials (5 μl of DNA extract per assay), the detection limit of the PCR test was 1.4 CFU/assay, corresponding to 280 CFU/ml of tonsil sample for the product specific to serotypes 2 or 1/2 and 0.14 CFU/assay (28 CFU/ml of tonsil sample) for the product shared between all S. suis serotypes. Moreover, the IPC of 620 bp was also noticeable in Fig. 2.

Clinical signs and macroscopic lesions. Data on clinical signs and macroscopic lesions are summarized in Table 3.

(i) Groups 1 and 2. Negative control animals did not exhibit any clinical signs of S. suis infection. Postmortem examinations...
did not reveal any lesions in these pigs. The average daily weight gains (ADGs) were 981 g (group 1) and 993 g (group 2).

(ii) Groups 3 and 4 (S. suis type 2). Rectal temperatures of three piglets infected with strain 332 (group 3) were moderate (40.2 ± 0.3°C) during 24 to 48 h p.i. Under the conditions of this trial (level three biosecurity), the normal temperature of SPF piglets is 39.5°C. All the piglets presented lameness, 24 h p.i. and during 1 to 22 days. Four of them developed arthritis associated with pneumonia in one animal. Contact pigs (group 4) did not develop clinical signs, but four animals showed arthritis at necropsy. The body growth of animals in group 3 was retarded. Differences in ADG between negative control pigs and contact pigs (P < 0.05), although no differences were noted between negative control pigs and contact pigs (P > 0.05).

(iii) Groups 5 and 6 (S. suis type 2). Two piglets infected with strain 347 (group 5) developed hyperthermia (40 ± 0.3°C) during 24 to 48 h, and lameness was observed in four pigs (during 1 to 10 days). Postmortem examinations revealed arthritis in these animals, whereas contact pigs (group 6) had neither symptoms nor lesions. The body growth was affected by the experimental infection (group 5). The ADG was 550 g, significantly different (P < 0.05) from those observed for piglets in negative control groups and group 6 (ADG = 970 g).

(iv) Groups 7 and 8 (S. suis type 1/2). No clinical signs were noticeable in groups 7 and 8, but moderate arthritis was observed at necropsy in three infected and one contact pigs. The ADG were, respectively, 927 and 875 g (groups 7 and 8), and no difference was obtained between these two groups and negative control groups (P > 0.05).

Detection of S. suis. All samples collected from animals in negative control groups and analyzed 16 days post-inoculation (live pigs) and after euthanasia (16 to 30 days p.i.) were negative by PCR and bacteriological analysis. Data are summarized in Tables 3 and 4.

(i) Environmental samples. All environmental samples were negative by PCR and bacteriological analysis.

(ii) Results obtained with live pigs (16 days p.i.) (Table 4). The comparison of PCR as well as bacteriological results, obtained from tonsils in infected or in contact pigs did not show any differences between groups 3, 4, 5, and 6 (P > 0.05). Results from groups 7 and 8 were negative.

(iii) Results obtained in euthanized piglets (16 to 30 days p.i.). The results, obtained by PCR and bacteriological analysis from 24 dead animals (Tables 3 and 4), confirmed the presence of S. suis type 2 (strains 332 and 347) infection, respectively, in 14 and 19 samples from joints, muscle, liver, heart, and lungs, but results by the two techniques were not significantly different (P > 0.05). S. suis type 1/2 was isolated in only one pig (group 7) from the joints. Sixteen to thirty days p.i., S. suis type 2 was detected from the tonsils of 24 and 23 pigs by PCR and culture, respectively, whereas nine pigs were identified as positive by PCR and negative by culture in groups 7 and 8 (S. suis type 1/2). In these groups, the difference between the two tests was significant (P < 0.05). The results were not significantly different between the three samples of tonsils (external swab, biopsy sample, and whole tonsil) (Table 4). No significant difference was observed between infected and contact pigs and between groups 3, 4, 5, and 6 (P > 0.05).

Serological results. The ELISA results obtained from negative controls but also from groups 7 and 8 were negative. Pigs infected with S. suis type 2 (strains 332 and 347) developed humoral antibodies detectable by ELISA from 16 days p.i. The level of antibodies increased until the end of the experiment. Contact pigs of both groups were also seropositive with a moderate level of antibodies.

**DISCUSSION**

The multiplex PCR assay described in this study was based on the amplification of a gene fragment coding for 16S rRNA of S. suis and on the amplification a cps2J gene fragment encoding S. suis serotype 2 and 1/2 capsular biosynthesis (24). Furthermore, our test contained an internal PCR control to eliminate false-negative samples due to inhibitors of polymerization. All S. suis strains tested in this study were detected by our multiplex PCR test, while none of the other bacterial species showed a positive reaction.

The 35 capsular types of S. suis were recognized by this PCR test, which was also able to detect S. suis in specimens from live pigs experimentally infected. S. suis isolates belonging to any serotype isolated from tonsil could be potentially virulent. Recently, in our laboratory, S. suis serotype 5 was isolated from a subject with septicemia without any other bacterial isolation. This serotype was also isolated from nursery pigs with serious cases of meningitis on a Canadian farm (9). Some S. suis strains isolated from healthy carrier pigs (particularly in ton-
sils) are able to induce transmission of *S. suis* infection between pigs or between herds. These pigs should be carefully checked, and the detection of *S. suis* species in tonsils is the first step of the diagnosis. Thus, the PCR assay described in this study, based on the 16S rDNA region conserved across capsular types, allowed the detection of *S. suis* species and would be very useful for epidemiological studies. Other PCR tests were previously developed to detect *S. suis* species (6, 19, 31). Okumubaua et al. and Wisselink et al. have also reported two PCR tests to detect two of the *S. suis* virulence factors, the suilysin and the extracellular factor, respectively (19, 30). However, the absence of these proteins in some virulent strains may preclude the routine use of these tests. Moreover, in a previous study we showed that these proteins were present neither in all *S. suis* strains nor in all virulent European strains (5, 19).

In 2003, Okumubaua et al. developed a multiplex PCR based on the *gdh* gene, encoding the glutamate deshydrogenase of *S. suis* serotype 2, allowing the amplification of all *S. suis* serotypes (20). This method was very attractive, but it was only applied to detect or characterize *S. suis* from pure cultures.

Our multiplex PCR assay allowed also the detection of all *S. suis* serotypes and more specifically serotypes 2 and 1/2. A preliminary study was performed to develop a PCR test able to detect serotype 2 alone, because this serotype is the major cause of disease in France and because it is a zoonotic agent that causes septicemia, endocarditis, and meningitis in humans (2, 5, 10, 21, 22, 29). The first step was to show genomic specificities in serotype 2 strains. We sequenced the four potential target genes (*cps2F*, *cps2H*, *cps2I*, and *cps2J*) that code for *S. suis* capsule, in serotypes 2 and 1/2 reference strains. In the original work describing the sequencing of these genes, the serotype 1/2 reference strain was not analyzed (23). The alignment of these nucleotide sequences showed a perfect identity for the *cps2H*, *cps2I*, and *cps2J* genes of the two reference strains. However, a substitution (T to C) was detected in the *cps2F* gene of serotype 2. Since this mutation leads to the disappearance of an SspI restriction site in the serotype 1/2, a PCR-restriction fragment length polymorphism was developed. However, this substitution was not detected in all serotype 2 strains isolated from subjects in the field (data not shown). In this context, we decided to develop a PCR test able to detect simultaneously serotypes 2 and 1/2. These serotypes are the most frequently detected in France (77.3%) (5). Several PCR tests, targeted on the *cps* locus, were previously described to detect these two serotypes (20, 26, 31). We designed original primers from the *cps* locus in order to (i) have a hybridization temperature compatible with the primers based on 16S rDNA, (ii) avoid the appearance of dimers, and (iii) have a PCR product differing in size from the product of 16S-195s and 16S-489 as primer pair.

The sensitivity of our multiplex PCR test was evaluated in vitro with SPF pig tonsils experimentally infected with *S. suis* type 2 reference strain and was found to be 28 CFU of *S. suis/ml* of tonsil sample. In the in vitro conditions described in our study, the PCR test was evaluated to be 20 times more sensitive than culture-positive results (28 versus 500 CFU of *S. suis/ml*). Our assay is more sensitive than other PCR tests previously described: the detection threshold of the multiplex test developed by Wisselink et al. (31) was 10 fg of chromosomal DNA in 25 μl of clinical sample (approximately 200 CFU/ml according to our reckoning). This estimate probably lacks accuracy because the test was performed not with tonsils experimentally infected in vitro but with purified DNA. The sensitivity of the PCR tests described by Smith et al. and Okumubaua et al. were not evaluated (20, 25, 26).

The comparison of PCR and bacteriological isolation obtained from tonsils of live pigs (healthy carriers) as well as in other organs in dead pigs, with or without macroscopic lesions, did not show any significant differences, with the exception of groups 7 and 8 infected with serotype 1/2 (*P* < 0.05). It is possible that the number of organisms present in samples from groups 3 to 6 is above the detection limit for both assays. Our multiplex PCR assay, with an internal control and without a culture step, is easy to perform. Ninety-six samples could be analyzed simultaneously in 6 h, whereas bacteriological isolations require at least 4 days (including primary isolation, cloning, biochemical identification, and serotyping). The major difficulty of bacteriological isolation is to locate *S. suis* colonies from multi-infected samples such as tonsils. On the other hand, tonsil specimens appeared to be helpful to detect *S. suis* in live pigs. Our PCR assay used samples directly and was able to detect and identify *S. suis* serotype 2 and 1/2 among suspicious α-hemolytic colonies on blood agar medium. In practice, the main advantages of this test are its abilities (i) to detect *S. suis* from multi-infected samples, (ii) to reduce the time required to identify the bacteria, and (iii) to increase the number of colonies analyzed at the same time.

The comparison of tonsil biopsy specimens, external swabs, and whole tonsils did not show any significant differences. These results differ from those published by Fittipaldi et al. in 2003 (11), with *Actinobacillus pleuropneumoniae* infection. These authors showed that the PCR detection rate was higher with whole tonsils than with tonsil biopsy specimens. However, in that study, conventional naturally infected pigs were tested. External tonsil swabs will be chosen in the future because external tonsil swabs are less traumatizing for pigs.

In this experimental study, clinical signs and macroscopic lesions induced in SPF piglets were moderate except for arthritis (especially for pigs infected with *S. suis* serotype 1/2). These results are different from those of our previous trials carried out under these standardized conditions with different strains of *S. suis* type 2 (3, 4). There was a transmission from infected to contact pigs, since we detected *S. suis* in contact pigs during the trial (on day 16) and at necropsy. At the end of the assay, all contact pigs in units B and C were healthy carriers pigs with no clinical symptoms of the disease (*P* > 0.05).

In conclusion, the assay in the present work may be used routinely to identify pigs carrying *S. suis* serotypes 2 and 1/2 and all other serotypes. It may also be applicable for epidemiological studies and transmission studies of *S. suis* and can contribute to the control of *S. suis* infection.

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