We performed multilocus sequence typing of 64 North American *Streptococcus suis* serotype 2 porcine isolates. Strains were sequence type (ST) 28 (51%), ST25 (44%), and ST1 (5%). We identified nonrandom associations between STs and expression of the virulence markers slyysin (SLY), muramidase-relased protein (MRP), and extracellular factor (EF). Expression of pili encoded by the *srtF* and *srtG* pilus clusters was also nonrandomly associated with STs. ST1 strains were SLY+ EF+ MRP+ *srtF* pilus+ *srtG* pilus−. ST25 strains were SLY− EF− MRP− *srtF* pilus− *srtG* pilus+, and most ST28 strains were SLY− MRP+ EF− *srtF* pilus+ *srtG* pilus+. ST28 isolates proved essentially nonvirulent in a mouse infection model; ST25 strains showed moderate virulence and ST1 isolates were highly virulent. ST1 is responsible for a high proportion of *S. suis* disease in humans worldwide. Its presence in North America indicates that potential zoonotic *S. suis* outbreaks in this continent cannot be disregarded.

*Streptococcus suis* causes meningitis and septicemia in pigs and is a zoonotic agent (1). In the Western hemisphere, human *S. suis* disease is infrequent and usually affects workers in the swine industry. However, *S. suis* is the most commonly reported cause of streptococcal meningitis in adults in Vietnam and the second in Thailand (2,3). Two outbreaks of human *S. suis* disease have occurred in People’s Republic of China, affecting hundreds of persons and causing 39 deaths (4). Most cases of animal and human *S. suis* infection have been caused by serotype 2 strains (5). The percentage of *S. suis* serotype 2 strains recovered from diseased pigs and the number of cases of human disease is lower in North America than in other parts of the world (6,7).

Multilocus sequence typing (MLST) has shown that *S. suis* serotype 2 strains can be divided into at least 16 sequence types (STs). Closely related STs are grouped in the so-called ST complexes. Although ST complexes 1, 27, and 87 dominate the *S. suis* population, most invasive isolates belong to the ST1 complex (8). For example, most strains isolated from human patients in Japan were ST1 (9), whereas those causing the human outbreaks in People’s Republic of China were ST7, included in the ST1 complex (10,11). However, Takamatsu et al. showed that 80% of the isolates recovered from blood or cerebrospinal fluid of humans in Thailand belonged to STs grouped in the ST27 complex (12).

Most of the *S. suis* serotype 2 strains genotyped so far by MLST originated in Europe and Asia (8–12). Isolates from Canada and the United States have received less attention. In this study, we used MLST to genotype a relatively large collection of US and Canadian *S. suis* serotype 2 strains.

**Materials and Methods**

*S. suis* Field Strains

Sixty-four strains of *S. suis* serotype 2 isolated from pigs with clinical disease in different and nonrelated farms in major swine production areas of Canada and the United States were used. For comparison purposes, 19 porcine and 1 human *S. suis* serotype 2 strains isolated in Japan and
12 human S. suis serotype 2 strains isolated in Thailand were included (12,13). All strains are listed in the online Appendix Table (wwwnc.cdc.gov/EID/article/17/12/11-0609-TA1.htm).

MLST and Phylogenetic Analysis

S. suis genomic DNA was prepared from overnight cultures by using the QIAamp DNA Minikit (QIAGEN, Valencia, CA, USA) following the manufacturer’s instructions. MLST was performed by PCR amplification and DNA sequencing of the 7 loci described above, but the mice received a 10-fold higher dose of ST28 strains 1088563, 1054471, and 1097205. In this second experiment, groups contained 5 mice.

Results

Most of the 64 strains from North America were ST28 (n = 33) or ST25 (n = 28). Together, these 2 STs accounted for 95% of all S. suis serotype 2 strains from North America that were investigated (Table 1). However, a higher ST28 prevalence was true only for the United States; most strains from Canada were ST25. The remaining 3 strains belonged to ST1, which is commonly found in Europe and Southeast Asia.

Nonrandom Association between STs and Expression of Virulence Markers

SLY (encoded by the sly gene), MRP (mrp gene), and EF (epf gene) are virulence markers that have been used in elaborated genotypic and phenotypic schemes to try to predict the virulence of a given S. suis strain (1,19). For example, Silva et al. designed a multiplex PCR test that can discriminate between at least 6 naturally occurring genetic variants of mrp, named mrp', mrp, mrp*, mrp**, mrp***, and mrp**** (20). We investigated possible associations between STs and these widely used markers in our collection of S. suis serotype 2 strains from North America. To assess whether associations found are independent of the geographic origin of the strains, we included 32 described (12,13) S. suis serotype 2 strains of STs 28, 25, and 1 and isolated in Japan and Thailand (online Appendix Table).

Independently of geographic origin, we found clear, nonrandom associations between STs and expression of virulence markers. All but 2 ST1 strains had the phenotype SLY+MRP+EF+. All ST25 strains were SLY–MRP–EF– and all ST28 strains were SLY–MRP+EF+ (Table 2). Most ST1 strains had an sly+mrp+epf+ genotype.
Table 2. Association of *Streptococcus suis* serotype 2 STs and commonly used virulence markers in isolates from North America*†

<table>
<thead>
<tr>
<th>ST</th>
<th>No. strains</th>
<th>sly</th>
<th>mpr</th>
<th>mpr†</th>
<th>mpr*</th>
<th>mpr**</th>
<th>mpr***</th>
<th>ND</th>
<th>epf</th>
<th>Phenotype</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>11</td>
<td>11</td>
<td>9</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>2</td>
<td>11</td>
<td>11</td>
</tr>
<tr>
<td>25</td>
<td>36</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>2</td>
<td>11</td>
<td>11</td>
</tr>
<tr>
<td>28</td>
<td>49</td>
<td>42</td>
<td>6</td>
<td>1</td>
<td>8</td>
<td>23</td>
<td>3</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

†ST, sequence type; SLY, sulysin; MRP, muraminidase-released protein; ND, no amplification of the *mpr* gene was detected by PCR under the conditions used; EF, extracellular factor.
‡Variants of the *mpr* gene are those described by Silva et al. (20).
§Hemolysis of horse erythrocytes by the strains was considered to be an indication of the expression of SLY.
¶Molecular mass MRP variants identified by Western blotting were in agreement with those expected on the basis of the *mpr* gene variant identified by PCR.

Nonrandom Association between STs and Expression of Pili

Takamatsu et al. reported associations between particular STs and the presence or absence of putative pilus gene clusters, designated *srtBCD*, *srfE*, *srfF*, and *srtG* clusters (13). All ST25 and ST28 strains investigated by these authors were positive by PCR for all genes in the *srfF* and *srfG* pilus clusters (13). Consistently, we found that all ST25 and ST28 strains in our collection of strains expressed the protein (Table 2). In a recent report, all *mpr*+/MRP− strains that were investigated (of various *S. suis* serotypes) had truncations or point mutations in the *mpr* gene that prevented expression of MRP (6). Although we have not sequenced the *mpr* gene in our collection of strains, we hypothesize that similar genetic rearrangements are likely to explain the *mpr*+/MRP− results we observed in ST25 strains in this study. Three *mpr* gene variants were associated with ST28, although variant *mpr* was the most prevalent (85%) among this ST.

Mouse Infection Model

Inasmuch as the MLST data showed that more than half of the strains from North America analyzed were ST28 and the second most represented ST was ST25, we performed a comparison of the virulence of representative ST25 and ST28 strains by using a standardized mouse infection model (18). For comparison, we included the well-characterized and highly virulent ST1 strain P1/7. Most mice in the ST1 group showed severe clinical signs pili (Table 3). However, although all ST25 strains expressed the *srtG* pilus, none produced the *srfF* pilus (Table 3).

It has been shown that one ST25 isolate from Canada, which does not have a discrete *srfF* pilus cluster and is unable to express the *srfF* pilus, is nonetheless PCR positive for each of the individual *srf* genes because PCR amplicons can be generated from homologs of these genes found at various genome locations (13,16). We hypothesized that the ST25 strains analyzed in our study have a genetic organization similar to that ST25 isolate. Consistent with our hypothesis, our attempts to amplify the *srfF* pilus cluster in ST25 strains by using a primer pair annealing to the first and last gene of the *srfF* cluster were unsuccessful (data not shown). All the ST1 strains had the *srfF* cluster genes but, with the exception of 3 strains, not the *srtG* cluster genes. When we assessed the pilus phenotype by Western blotting, all ST1 strains expressed the *srfF* pilus but none expressed the *srtG* pilus (Table 3). The reason(s) the 3 ST1 strains that have the *srtG* cluster genes do not express the corresponding pilus are currently under investigation.

Table 3. Association of *Streptococcus suis* serotype 2 STs and *srfF* and *srtG* pilus clusters in isolates from North America*‡

<table>
<thead>
<tr>
<th>ST</th>
<th>No. strains</th>
<th><em>srfF</em> pilus</th>
<th><em>srfF</em></th>
<th>Pili expression‡, Sfp1</th>
<th><em>srtG</em> pilus</th>
<th><em>srtG</em></th>
<th>Pili expression, Sgp1‡</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>11</td>
<td>11</td>
<td>11</td>
<td>11</td>
<td>3</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>25</td>
<td>36</td>
<td>36</td>
<td>36</td>
<td>36</td>
<td>36</td>
<td>36</td>
<td>36</td>
</tr>
<tr>
<td>28</td>
<td>49</td>
<td>49</td>
<td>49</td>
<td>49</td>
<td>49</td>
<td>49</td>
<td>49</td>
</tr>
</tbody>
</table>

*ST, sequence type.
‡The presence of the genes was detected by PCR by using primers and conditions described by Takamatsu et al. (13).
‡Expression of pilus encoded by the *srfF* and *srtG* pilus clusters was performed by Western blotting by using described antibodies directed against the major subunits of these structures (16,17).
of septicemia, such as depression, swollen eyes, weakness, and prostration during the first 24 hours postinoculation. Several mice died of septicemia during the first 2 days of the trial, and the remaining animals were humanely killed for ethical reasons at day 3 postinoculation (Figure 1). \textit{S. suis} was isolated in pure cultures at high titers (>1 × 10^7 CFU/mL) from blood samples and organs, such as the liver and spleen, of septicemic animals in the ST1 group (>1 × 10^7 CFU/0.5 g of tissue in most animals).

The virulence of ST25 strains was intermediate. They caused moderate clinical signs and relatively low mortality among inoculated mice (Figure 1). Statistical analysis demonstrated that ST25 strains were significantly less virulent than ST1 strains. However, ST25 strains were significantly more virulent than ST28 strains. In fact, no mice in the ST28 group died (Figure 1) or showed clinical signs associated with \textit{S. suis} infection, with the exception of slight depression immediately after inoculation, which subsided after 24 hours postinoculation. Bacteria could not be isolated from the blood of most mice in this group >48 postinoculation, and we could not isolate \textit{S. suis} from different organs at necropsy (results not shown). Given this surprising absence of clinical signs, we repeated the experiment by inoculating 3 additional groups of 5 mice each with the previously used and 2 other ST28 strains by using an infective dose that was 10-fold higher than the one previously used. Despite this increased infective dose, similar low virulence was observed for ST28 strains (Figure 2).

**Discussion**

In this article, we show that most \textit{S. suis} isolates from North America belong to ST28 and ST25 and that strains of these STs are significantly less virulent than ST1 strains. Although ST28 strains were essentially nonvirulent for mice, ST25 strains were of intermediate virulence and able to induce severe disease.

With a population of ≈115 million pigs, Canada and the United States combined are second only to the People’s Republic of China in terms of swine production. Although \textit{S. suis} infections are a main cause of postweanled piglet deaths in North America, the prevalence of \textit{S. suis} serotype 2 strains is much lower on this continent than in other regions of the world (6,7). We show here that in North America the most common STs among \textit{S. suis} serotype 2 strains are ST28 and ST25. By using a mouse infection model, we also show that \textit{S. suis} serotype 2 ST28 and ST25 strains are of lower virulence than ST1 strains. In contrast to Europe and Asia, where >60% of virulent serotype 2 isolates are ST1 (21–23), in North America only a small percentage (5%) of strains belonged to this more virulent ST.

Only 3 cases of \textit{S. suis} serotype 2 in locally infected humans have been reported in North America (5). Our results suggest that this low prevalence of human infections might be connected to the lower virulence of the circulating serotype 2 strains among the swine population in North America. In addition to a low prevalence of ST1 strains, we did not find any strains in our collection from North America belonging to STs 101, 102, 103, and 104, which are agents of human disease in Thailand (12). On the basis of its low frequency of isolation, we speculate that the ST1 strains we identified were introduced in North America by importation of animals. Human travel might also contribute to dissemination of ST1 strains, as exemplified by a reported case of human \textit{S. suis} meningitis caused by an ST1 strain involving a patient who contracted \textit{S. suis} in the Philippines but in whom clinical signs appeared only after he returned to the United States (24). The deadly human outbreaks in Asia caused by ST1 complex strains (2–5) and the fact that ST1 strains are replacing at a fast pace STs of lower virulence and causing human disease in countries such as Vietnam and Thailand (21,25,26) highlight that maintaining a low prevalence of ST1 strains among the swine population in North America is crucial for animal and human health. Of note, the only locally acquired human infection in the United States described so far (27) was caused by an ST1 strain (M. Gottschalk, unpub. data).

Another concern for the swine industry and for public health authorities is the presence in North America of \textit{S.}

![Figure 1. Survival of CD1 mice inoculated with \textit{Streptococcus suis} strains of different sequence types (STs). Most animals that received the ST1 strain P1/7 died from septicemia during the first 3 days of the trial. Several animals in this group died from meningitis from day 6 postinfection. Two groups of mice received ST25 strains 89–1591 and 1085543, respectively. Survival of mice in these 2 groups was higher than in the group that received the ST1 strain. However, >40% of the animals in the 89–1591 group and 60% of the animals in the 1085543 group died or were killed for ethical reasons before the end of the trial. In strong contrast, all 15 mice in the ST28 strain group survived the trial. Significant differences in survival were noted between groups (log-rank test, p values indicated in the figure body).](image-url)
In this experiment, the infectious dose was $1 \times 10^8$ CFU/animal, and 2 cases in Canada of S. suis were reported (71, 389). In comparing the virulence of ST25 and ST28 strains, the former are of lower virulence than strains from Eurasia. However, we do not yet understand the reasons for this lower virulence. Our work provides more support to the longstanding hypothesis that S. suis serotype 2 strains in North America are of lower virulence than strains from Eurasia. However, we do not yet understand the reasons for this lower virulence. The genome sequences of several S. suis serotype 2 ST1 and an ST25 strains have been published or made available (25, 29, 30). Genome sequencing of a larger number of additional S. suis strains of these and other STs could help elucidate the genetic basis of virulence differences among strains of this swine pathogen and zoonotic agent.

This study was supported by grants from the Natural Sciences and Engineering Research Council of Canada to M.G. (no. 154280 and Discovery Accelerator Supplement 380299) and Minister of Economic Development, Innovation and Export Trade, China-Quebec Collaboration grants to M.G and J.X. (no. 2008FA31830 and 2008DFA31830, respectively). N.F. is partially supported by the Canadian Institutes of Health Research.

Dr Fittipaldi is a postdoctoral fellow in the Center for Molecular and Translational Human Infectious Diseases Research of the Methodist Hospital Research Institute in Houston, Texas, USA. His primary research interest is the molecular basis of streptococcal pathogen–host interactions.

**References**


Address for correspondence: Marcelo Gottschalk, Groupe de Recherche sur les Maladies Infectieuses du Porc, Faculté de Médecine Vétérinaire, Université de Montréal, 3200 Rue Sicotte, CP5000, St-Hyacinthe, Quebec J2S 7C6, Canada; email: marcelo.gottschalk@umontreal.ca