Streptococcus suis serotyping by matrix-assisted laser desorption/ionization time-of-flight mass spectrometry

Chadaporn Chaiden1, Janthima Jaresitthikunchai2, Anusak Kerdsin3, Nattakan Meekhanon4, Sittiruk Roytrakul2, Suphachai Nuanualsuwan1,5

1 Department of Veterinary Public Health, Faculty of Veterinary Sciences, Chulalongkorn University, Bangkok, Thailand, 2 Functional Proteomics Technology Laboratory, Functional Ingredients and Food Innovation Research Group, National Center for Genetic Engineering and Biotechnology, National Science and Technology for Development Agency, Pathum Thani, Thailand, 3 Faculty of Public Health, Kasetsart University Chalermprakiat Sakon Nakhon Province Campus, Sakon Nakhon, Thailand, 4 Department of Veterinary Technology, Faculty of Veterinary Technology, Kasetsart University, Bangkok, Thailand, 5 Food Risk Hub, Research Unit of Chulalongkorn University, Bangkok, Thailand

* suphachai.n@chula.ac.th

Abstract

Streptococcus suis, particularly S. suis serotype 2 (SS2), is an important zoonotic pathogen causing meningitis in humans worldwide. Although the proper classification of the causative and pathogenic serotype is salutary for the clinical diagnosis, cross-reactions leading to the indistinguishability of serotypes by the current serotyping methods are significant limitations. In the present study, matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI-TOF-MS) analysis of extracted peptides was developed to improve the classification of serotype of S. suis. The peptide mass fingerprint (PMFs) database of S. suis was generated from the whole-cell peptides of 32 reference strains of S. suis isolates obtained from pigs. Thirty-two human S. suis isolates from clinical cases in Thailand were used to validate this alternative serotyping method in direct comparison to the multiplex (m) PCR approach. All reference strains, representing 32 serotypes of S. suis, exhibited their individual PMFs patterns, thus allowing differentiation from one another. Highly pathogenic SS2 and SS14 were clearly differentiated from the otherwise serologically closely related SS1/2 and SS1, respectively. The developed MALDI-TOF-MS serotyping method correctly classified the serotype in 68.8% (22/32) of the same serotype isolates generated from the PMFs database; while the validity for the clinical human isolates was 62.5% (20/32). The agreement between the MALDI-TOF-MS and mPCR serotyping was moderate with a Kappa score of 0.522, considering that mPCR could correctly serotype up to 75%. The present study demonstrated that PMFs from the developed MALDI-TOF-MS-based method could successfully discriminate the previously indistinguishable highly pathogenic SS2 and SS14 from SS1/2 and SS1, respectively. Moreover, this serotyping method distinguished pathogenic SS6, and so is an alternative approach of choice to rapidly and reliably serotype clinically pathogenic S. suis isolates.
Introduction

Streptococcus (S.) suis is a zoonotic foodborne pathogen [1] that inhabits the nasal cavity and particularly the tonsils of pigs. People who work closely with live pigs can potentially become infected with this pathogen via open skin lesions leading to the clinical symptoms of septicemia, endocarditis, peritonitis, pneumonia, and meningitis [1, 2]. In the past, this bacterium has been classified by its capsular polysaccharide into up to 35 serotypes [1, 2], of which currently 29 serotypes are recognized [3, 4]. Of these, S. suis serotype (SS2) is the most significant serotype as it is predominantly associated with both diseased pigs and human clinical cases, and it has been reported as the causative agent in more than 70% of human clinical cases. Whereas SS14 and some other pathogenic serotypes are far less significant and account for only up to 3% of human clinical cases worldwide [5].

Although the prompt confirmation of the causative serotype as part of the clinical diagnosis is pivotal to the successful treatment rate of S. suis infection, more than 20% of human cases have gone undiagnosed due to inherent drawbacks in the current serotyping methods [5]. Initially, microbial identification of S. suis is presumed following biochemical tests. Subsequently, polymerase chain reaction (PCR) assay targeting the presence of the recombinant/repair protein gene recN [6] is then used to confirm S. suis. Serological methods have been considered as the standard procedures for serotyping S. suis. However, cross-reaction among pairs or groups of serotypes, such as SS1 and SS14; SS2 and SS1/2; SS2 and SS22; SS6 and SS16; and SS1, SS2, and SS1/2; can lead to the inconclusive or erroneous serotyping of S. suis. These indistinguishable serotypes are problematic since pathogenic serotypes in both pigs and humans, such as SS2, SS6, SS14, and SS16, are found among all the inconclusive pairs (or groups) of cross-reactions [5]. In addition, serotyping with all the typing antisera is laborious, time-consuming, and expensive, and preparing the antisera is difficult due to the high cost and labor associated with its production.

Even though multiplex (m)PCR has recently been developed and gained popularity as an alternative technique to serotype S. suis isolates, mPCR does not differentiate SS2 from SS1/2, or SS14 from SS1 due to the high capsular gene cluster similarity. So, additional or alternative methods are required to solve this issue. Lately, matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI-TOF-MS) has been increasingly endorsed and adopted as an alternative approach to microbial identification [7], including S. suis identification [8, 9]. In this method, cellular proteins (and peptides) of S. suis are extracted and then used for species identification via peptide mass fingerprints (PMFs) formed using MALDI-TOF-MS.

Nevertheless, one previous study successfully examined few serotypes of SS using MALDI--TOF-MS still most of the serotypes were not classified especially indistinguishable serotypes by current serotyping methods [8], and so is tackled in this study. The peptide extraction method was modified from Bruker’s recommendation [10], and the extracted peptides of S. suis were then analyzed by MALDI-TOF-MS. The aim of this study was to demonstrate S. suis serotyping by means of the modified peptide extraction method coupled with MALDI--TOF-MS, and in particular to differentiate those inconclusive pathogenic and non-pathogenic serotypes.

Materials and methods

Bacterial strains

In this study, 32 reference strains (SS1-14, SS16-20, SS22-32, SS34, and SS1/2) representing 32 different serotypes of S. suis (Table 1) along with local S. suis SS2 (n = 23) and SS14 (n = 9)
isolates from humans were used, the strains were originally collected from hospital, distributed in 5 regions (north, northeast, central, east, and south) of Thailand. Sequencing of the 16S rRNA gene was used to confirm these 32 reference strains [11]. Bacteria were cultured on Columbia blood agar (Difco Laboratories, Detroit, Mich.) with 5% (v/v) sheep blood at 37˚C in an anaerobic condition for 24 h. Reference strains of \textit{Staphylococcus aureus} ATCC 25923 and \textit{E. coli} DH5 alpha were included to study cross-reaction.

Peptide extraction

Pure colonies of \textit{S. suis} for peptide extraction were prepared on Columbia blood agar as above. Then ethanol was added to 70% (v/v) to precipitate out the bacterial peptides from the whole

| Serotype | Strain   | Source    | MALDI-TOF-MS GenBan acc.no. of 16S rRNA gene |
|----------|----------|-----------|----------------------------------------------|
| 1        | NCTC10237 | Diseased pig | 1 2.187 LSV 594043.1 |
| 2        | NCTC10234 | Diseased pig | 2 2.453 LS483418.1 |
| 3        | 4961      | Diseased pig | 3 2.341 AF009478.1 |
| 4        | 6407      | Diseased pig | - - AF009479.1 |
| 5        | 11538     | Diseased pig | - - AF009480.1 |
| 6        | 2524      | Diseased pig | 6 2.207 AF009481.1 |
| 7        | 8074      | Diseased pig | 7 2.369 AF009482.1 |
| 8        | 14636     | Diseased pig | 8 2.302 AF009483.1 |
| 9        | 22083     | Diseased pig | 9 2.178 AF009484.1 |
| 10       | 4417      | Diseased pig | - - AF009485.1 |
| 11       | 12814     | Diseased pig | - - AF009486.1 |
| 12       | 8830      | Diseased pig | - - AF009487.1 |
| 13       | 10581     | Diseased pig | 13 2.354 AF009488.1 |
| 14       | 13730     | Diseased pig | 14 2.492 AF009489.1 |
| 16       | 2726      | Diseased pig | 16 2.504 AF009491.1 |
| 17       | 93A       | Healthy pig | - - AF009492.1 |
| 18       | NT77      | Healthy pig | 18 2.539 AF009493.1 |
| 19       | 42A       | Healthy pig | 19 2.36 AF009494.1 |
| 20       | 86–5192   | Diseased calf | 20 2.566 AF009495.1 |
| 22       | 88–1861   | Diseased pig | - - AF009497.1 |
| 23       | 89–2479   | Diseased pig | - - AF009498.1 |
| 24       | 88–5299A  | Diseased pig | 24 2.207 AF009499.1 |
| 25       | 89–3576–3 | Diseased pig | 25 2.419 AF009500.1 |
| 26       | 89–4109–1 | Diseased pig | - - AF009501.1 |
| 27       | 89–5259   | Diseased pig | 27 2.304 AF009502.1 |
| 28       | 89–590    | Diseased pig | - - AF009503.1 |
| 29       | 92–1191   | Diseased pig | 29 2.3 AF009504.1 |
| 30       | 92–1400   | Diseased pig | 30 2.256 AF009505.1 |
| 31       | 92–4172   | Diseased calf | 31 2.124 AF009506.1 |
| 32       | EA1172.91 | Diseased pig | 32 2.336 AF009507.1 |
| 33       | 82–2742   | Diseased pig | 34 2.393 AF009509.1 |
| 1/2      | 2651      | Diseased pig | ½ 2.255 AF009476.1 |
| Total    | 32        | 32         | 22 (68.8%) 2.34* |

*Average LSV of correct MALDI-TOF-MS classifications.

https://doi.org/10.1371/journal.pone.0249682.t001

isolates from humans were used, the strains were originally collected from hospital, distributed in 5 regions (north, northeast, central, east, and south) of Thailand. Sequencing of the 16S rRNA gene was used to confirm these 32 reference strains [11]. Bacteria were cultured on Columbia blood agar (Difco Laboratories, Detroit, Mich.) with 5% (v/v) sheep blood at 37˚C in an anaerobic condition for 24 h. Reference strains of \textit{Staphylococcus aureus} ATCC 25923 and \textit{E. coli} DH5 alpha were included to study cross-reaction.

Peptide extraction

Pure colonies of \textit{S. suis} for peptide extraction were prepared on Columbia blood agar as above. Then ethanol was added to 70% (v/v) to precipitate out the bacterial peptides from the whole
bacterial cell. The bacterial suspension was centrifuged at 11,000 g for 5 min, and then the pel-
let was collected. Next, 5% (v/v) trifluoroacetic acid (TFA) in absolute acetonitrile (ACN) was
added to the pellet and the suspension was dissolved by gentle vortex. The bacterial peptides
dissolved in the supernatant were collected after clarification by centrifugation at 11,000 g for
5 min and then kept at -20˚C prior to MALDI-TOF-MS analysis. Lowry assay was used to
quantify the concentrations of extracted peptides [12].

**Peptide analysis by MALDI-TOF-MS**

The extracted peptides from the 32 reference strains of *S. suis* were mixed with the sinapinic
matrix solution [sinapinic acid in 5% (v/v) TFA in absolute ACN] and then spotted onto the
MALDI target plate and allowed to crystalize at room temperature before inserting the MALDI
target plate into the MALDI-TOF-MS instrument. The mass spectrometry (MS) spectra were
collected using an Ultraflex III TOF/TOF (Bruker Daltonik, GmbH) instrument in a linear posi-
tive mode with a mass range between 2–20 kDa. Five hundred shots were reiterated and accu-
mulated with a 50 Hz laser per SS. Likewise, this MALDI-TOF-MS procedure was repeated for
all 32 serotypes of *S. suis*. All MS spectra were analyzed for fingerprint spectra and subjected to
PCA using the FlexAnalysis version 3.4 and ClinProTool version 3.0 software (Bruker Daltonik,
GmbH). The ACTH fragment 18–39 (human), insulin oxidized B chain (bovine), insulin
(bovine), cytochrome C (equine), and apomyoglobin (equine) were used as the external protein
calibrations. Analysis of variance (ANOVA), Student’s t-test and statistics in the software, were
used to assess the statistically significant differences in the PMFs across bacterial strains.

**Generating of *S. suis*’s PMFs database.** The *S. suis* PMFs database was generated by
inserting 20 qualified MS spectra from each of the 32 individual reference serotypes into the
MALDI Biotyper database system according to Bruker’s recommendation. The PMFs of 32
serotypes in the database will be used as the references PMFs for the serotype characteriza-
tion and to evaluate the repeatability, reproducibility and validation.

**Serotype analysis by mPCR**

All reference strains of *S. suis* were serotyped by mPCR [13]. The results from the MALDI-
TOF-MS analysis were compared to those from the mPCR of the same samples in terms of the
repeatability validation and agreement of tests. Four sets of primers targeting the capsule (cps) loci
across 28 serotypes of *S. suis* were used. The first primer sets targeted SS1/2, SS1-3, SS7, SS9, SS11,
SS14, and SS16; the second primer set targeted SS4, SS5, SS8, SS12, SS18, SS19, SS24, and SS25; the
third primer set targeted SS6, SS10, SS13, SS17, SS23, and SS31; and the fourth primer set targeted
SS21 and SS27-30. The mPCR serotyping of *S. suis* were followed as previously described [13].

**Repeatability**

The same isolate of each serotype was repeatedly cultured and extracted in exactly the same
manner. Eight spots from each serotype were applied to determine the precision of this devel-
oped serotyping method Bruker Biotyper. Log score values (LSVs) were used to evaluate the
serotype classifications, where LSVs between 0–1.69 were merely reliable, between 1.70–1.99
were probable, and more than 2.00 were highly probable. Among 8 spots, the matched sero-
types with highest LSVs will be considered as a correct classification.

**Validation**

To assess the validity of this serotyping method against known clinical isolates of *S. suis*, 32 iso-
lates of *S. suis* from human cases in Thailand, comprised of SS2 (*n* = 23) and SS14 (*n* = 9), were
cultured in the conventional anaerobic growth condition, extracted, and characterized by MALDI-TOF-MS as described earlier. Each individual isolate was spotted for eight entries. The serotype classification of the clinical isolates was evaluated by the same Bruker Biotyper LSVs and criteria.

**Reproducibility or culture-condition repeatability**

To evaluate the repeatability of this serotyping method with serotypes cultured under a different condition, then the 32 serotypes of *S. suis* were cultured as before except in an aerobic growth condition, and then peptide extracted and serotype classified by MALDI-TOF-MS using exactly the same MALDI-TOF-MS protocol as described earlier. The serotype classification of was evaluated by the same Bruker Biotyper LSVs and criteria.

**Degree of test agreement**

The 32 reference strains of *S. suis* were grown anaerobically or aerobically and subjected to both mPCR and developed MALDI-TOF-MS serotyping methods as described. The serotyping results are presented in a $2 \times 2$ contingency table of frequencies with the rows and columns representing the serotype results for both serotyping methods. The degree of agreement between these two methods was assessed using Cohen’s kappa statistic ($\kappa$) [14], as shown in Eq (1);

$$
\kappa = \frac{O_D - E_D}{1 - E_D},
$$

where $n$ is the total observed frequency, $O_D$ is the sum of observed frequencies along the diagonal (Table 2), $E_D$ is the sum of expected frequencies along the diagonal, $P_O$ is the $O_D/n$, and $P_E$ is the $E_D/n$.

**Results**

**Peptide mass spectra of *S. suis***

After extraction, peptide masses between 2–20 kDa were collected and analyzed using the FlexAnalysis version 3.4 and ClinProTool version 3.0 software (Bruker Daltonik, GmbH). All of the extracted peptides delivered an adequate number and intensity (more than $10^4$ a.u.) of peptide mass. Extracted peptides from each individual serotype generated particular PMFs that contained their individual unique mass(es) and so differentiated each serotype from one another (Fig 1A–1D). Overall, 4420, 5337, 5965, 6634, 6748, 6834, and 8260 Da peptide masses were commonly found in most of the SSs. Interestingly, SS1/2, SS11, and SS13, had some other unique masses (2990 and 3005 Da) in common. The former SS32 and SS34 (*S. orisratti*) had a unique PMF pattern with the 4447, 6610, and 6775 Da peptide masses being common to these two serotypes and so Biotyper software classified SS32 and SS34 in another group, as shown in the dendrogram (Fig 2).

| Observed | Multiplex PCR | 
|----------|---------------|
| MALDI-TOF | True serotype | False serotype | Sum |
| True serotype | 59 | 3 | 62 |
| False serotype | 19 | 22 | 41 |
| Sum | 78 | 25 | 103 |

Table 2. Contingency table of identification frequencies to evaluate *S. suis* serotypes using the MALDI-TOF-MS analysis of this study compared to mPCR test.
The ambiguous serotypes by mPCR, which were the highly pathogenic SS2 and SS1 that are serologically associated with SS1/2 and SS14, respectively, were clearly distinguished from one another (Fig 1A and 1B). The 6956 Da peptide mass discriminated between SS2 and SS1/2 ($p < 0.01$), while the 6919 Da peptide mass discriminated between SS1 and SS14 ($p < 0.01$), as shown in Table 3. Likewise, principal component analysis (PCA) revealed that the PMFs of SS2, SS1/2, SS1, and SS14 distinctly clustered around their individual serotypes (Fig 3). Therefore, the PCA results did support that the ambiguity of serotype pairs SS2 and SS1/2; and SS1

Fig 1. Representative MALDI-TOF peptide mass spectra of reference S. suis serotypes (A) SS1-7, (B) SS8-14 and SS16, (C) SS17-20 and SS22-25, and (D) SS26-32 and SS34. The X-axis represents the mass to charge ratio (m/z) and the Y-axis represents the intensity of the spectra.

https://doi.org/10.1371/journal.pone.0249682.g001
and SS14 could be clearly resolved by PMFs mapping using MALDI-TOF-MS. Additionally, cross-reacting serotypes based on serological methods were also classified, where the 6319, 6914, and 5056 Da peptides were index masses that differentiated between SS1 and SS2, between SS6 and SS16; and between SS2 and SS22, respectively, (Table 3).

**Repeatability**

The same isolate of each serotype was repeatedly cultured, extracted, and classified in exactly the same manner to determine the precision of the developed serotyping method. Eight spots from each serotype were applied to classify comparing with the \textit{S. suis} PMFs database.

![Figure 2. Dendrogram based on the PMFs of reference SS1/2, SS1-14, SS16-20, SS22-32, SS34, and SS1/2 grown anaerobically. SS32 and SS34 clustered in a group, while SS1/2, SS11, and SS13 clustered in another group and closer to the main SS PMFs. Staphylococcus aureus ATCC 25923 and E. coli DH5 alpha were included as the out group.](https://doi.org/10.1371/journal.pone.0249682.g002)

| Serotyping method | Ambiguous serotype | Differentiating index mass (Da) by MALDI-TOF-MS |
|-------------------|--------------------|-----------------------------------------------|
| Multiplex PCR     | SS2 vs. SS1/2      | 6956                                          |
|                   | SS1 vs. SS14       | 6919                                          |
| Serological       | SS1 vs. SS2        | 6319                                          |
|                   | SS2 vs. SS1/2      | 6956                                          |
|                   | SS6 vs. SS16       | 6914                                          |
|                   | SS2 vs. SS22       | 5056                                          |
|                   | SS1 vs. SS14       | 6919                                          |

![Table 3. Differentiating between serologically ambiguous serotypes using the index mass obtained from MALDI-TOF-MS analysis of PMFs from anaerobically cultured \textit{S. suis} serotypes.](https://doi.org/10.1371/journal.pone.0249682.t003)
generated as described earlier. The results showed that 22 out of 32 serotypes (68.8%; average LSV of 2.34) were correctly serotype classified. The ambiguous serotypes by mPCR (SS2 with SS1/2, and SS1 with SS14) were also correctly serotyped with LSVs of 2.453, 2.255, 2.187, and 2.492, respectively. The serological cross-reacting serotypes SS1 with SS2, and SS6 with SS16, were correctly classified with LSVs of 2.187, 2.453, 2.207, and 2.504, respectively (Table 1). However, SS22 (cross-reactive with SS2) was unable to be serotyped (distinguished) by this MALDI-TOF-MS approach method.

Validation

Thirty-two human isolates of *S. suis* from patients in Thailand, comprised of SS2 (*n* = 23) and SS14 (*n* = 9), were serotyped using the developed MALDI-TOF-MS method. The method correctly serotyped 20 out of 32 serotypes (62.5%) of human isolates, with an average LSV of 2.20 (Table 4). Serotype 2 was correctly classified in 13 out of 23 strains (56.5%, average LSV of 2.19) and SS14 was correctly classified in 7 out of 9 strains (77.8%, average LSV of 2.20).
Reproducibility

Instead of using an anaerobic growth condition, 32 the serotypes of *S. suis* (SS1-14, SS16-20, SS22-32, SS34, and SS1/2) were cultured as before except under an aerobic growth condition. The strains were then peptide extracted and serotype classified by exactly the same MALDI--TOF-MS protocol. The MALDI-TOF-MS approach was found to correctly serotype 18 out of 32 strains of *S. suis* (56.3%) with average LSV of 2.34.

Degree of test agreement

Cohen’s unweighted Kappa statistic was used to elucidate the agreement between the mPCR and this developed MALDI-TOF-MS serotyping method, and gave a Kappa score of approximately 0.522. While the true serotype classifications of the MALDI-TOF-MS and mPCR were not statistically significant (*p* > 0.05) approximately 60% (95% CI: 51%-70%) and 76% (95% CI: 67%-84%), respectively.

Discussion

As a rapid high throughput technique, MALDI-TOF-MS has been increasingly used in microbiological studies. It provides improved accuracy and power of resolution to identify or even classify microbial isolates [7, 15]. In this study, we modified the peptide extraction method coupled with MALDI-TOF-MS to identify *S. suis* and further classify the SSs. Mass spectra yielded from this peptide extraction technique exhibited good qualities, a high intensity, and an adequate number of peptide masses that overall indicated that this method could be applicable for SS classification.

The PMFs presented 4420, 5337, 5965, 6634, 6748, 6834, and 8260 Da peptide masses that were commonly found across all the tested SSs. Two of these masses were close to the 4420 and 8266 Da peptide masses that have recently been reported in previous studies using whole cell extraction of *S. suis*, which mass 4133 and 8367 were reported as the indicative species-specific peaks [8, 9]. This congruent finding may reach a general assumption that the 4420 and 8260 Da peptide masses could act as species-specific markers for *S. suis*. As these two masses were absent in others closely related *Streptococcus* spp. e.g. *S. plurextorum, S. porci* and *S. porcorum* where their species-specific markers corresponded to 6164, 6133 and 4190/8381, respectively [9]. However, some other peptide masses were found to be different from those in previous studies, which can be explained by the different preparation protocols, such as the acid concentration, spotting method, and type of matrices [16–18]. In the present study, the principal difference in the preparation protocol was the modified peptides extraction method, that formic acid was not included, gentle dissolve pellet by vortex and vigorously mix peptides with sinapinic matrix, which likely accounted for some of the dissimilar PMFs. Moreover, dissimilar PMFs could be inherent in the intraspecies discrepancy as well [19].

### Table 4. Validation of MALDI-TOF-MS for serotyping *S. suis* human strains isolated in Thailand compared to the mPCR approach.

| Serotype | Source | MALDI-TOF-MS |
|----------|--------|-------------|
|          |        | Serotype (best match) | LSV |
| 2        | Human  | 2 (13/23)         | 2.19|
| 14       | Human  | 14 (7/9)          | 2.20|
| Total    | 32     | 20 (62.5%)        | 2.20*|

*Average LSV of correct MALDI-TOF-MS classifications.*

https://doi.org/10.1371/journal.pone.0249682.t004
The PMFs of *S. suis*-like serotypes (SS32 and SS34) had a unique pattern with 4447, 6610, and 6775 Da peptide masses commonly found in these two serotypes. The 6775 Da peptide mass was somewhat close to the 6772 Da peptide mass that has previously been reported as a representative mass of *S. porcorum* PMFs [9]. Previous studies have suggested that SS32 and SS34 are likely to be *Streptococcus orisratti* [3]. In accord, the obtained PMFs dendrogram in this study (Fig 2) clustered both SS32 and SS34 together but separate from the other *S. suis* isolates, indicating that these two serotypes possibly possessed a high genetic dissimilarity from the other examined SSs. The phylogenetic tree based on 16s rRNA gene (Fig 4) also supported
this finding as SS32 and SS34 were also clustered separately from the main group of S. suis. Furthermore, SS1/2, SS11, and SS13 revealed some specific peptide masses that exclusively clustered together in the dendrogram (Fig 2). Thus, we presumed that these three reference serotype strains were divergent from the other tested reference serotype strains, in accord with a previous study that reported that SS13 was also divergent from other S. suis strains [20]. However, these divergent characters were not displayed in phylogenetic tree (Fig 4). In terms of the relatedness of PMFs, the cluster of SS1/2, SS11, and SS13 was closer to the main cluster than the cluster of SS32 and SS34 (Fig 2).

For SS classification, the PCA results demonstrated that the use of extracted peptides subject to MALDI-TOF-MS analysis was capable of discriminating between the high similarity serotypes pairs of (i) SS2 and SS1/2 and (ii) SS1 and SS14. Notably that, these two pairs of serotypes closely clustered together in phylogenetic tree (Fig 4) but separately in peptide dendrogram (Fig 2). This finding could be used to explained why our modified method could differentiate these two pairs of high similarity serotypes. Importantly, SS2 and SS14 are the two serotypes that accounted for most major S. suis infections [5, 21], where the misidentification of the causative pathogen could delay the appropriate treatment [22]. Moreover, the currently available classification methods are sometimes inconclusive. Hence, this modified peptide extraction method from Bruker’s recommendation coupled with MALDI-TOF-MS analysis could be an alternative approach to discriminate between serologically cross-reactive SSs and so is a potentially significant advantage for S. suis infection diagnosis.

According to the repeatability test, the extracted PMFs coupled with our S. suis database was capable of discriminating 22 out of 32 (68.8%) reference SSs. Given that some serotypes of S. suis share the same structural components [23], then the structural peptides acquired by the present study could be similar and so have led to the misclassification of the other 10 serotypes (31.3%). Moreover, only 8 spots were applied for repeatability test, while 20 qualified spots were used to generate the S. suis PMFs database. Similarly, only 4 spots are sufficient to identify bacteria by means of MALDI-TOF-MS [15]. However, serotype classification is more subtle than bacterial identification thus requiring more replication of MS spectra to differentiate the more closely related serotype within the same species. In order to improve the correct serotype classification rate, we suggested that the repeatability test of the serotype classification, including unknown serotyping, should require at least 8 MS spectra (spots). Likewise, LSVs criteria of correct serotype classification are probable between 1.70–1.99. More PMFs included in the database from each individual serotype would warrant a higher degree of correct classification.

Clinical human isolates of S. suis were included in this study to validate this method. Just over half (56.5%) and three-quarters (77.8%) of human SS2 and SS14 strains, respectively, were correctly classified. This moderate level of validity of the serotyping method implied that the S. suis reference strains and field strains posed a somewhat dissimilar peptide background. Of relevance then is it has previously been reported suggested that different backgrounds of the same serotype of S. suis could result in different phenotype expressions [24]. For further study, more PMFs from field strains of S. suis should be included in the database to improve the reliability of the serotype classification.

The reproducibility or culture-condition repeatability of the serotype classification was merely acceptable (56.3%) when S. suis was cultured in an aerobic condition. The different background derived from different growth conditions of the same isolates could yield the dissimilar peptides, such as in this case, where the anaerobic vs. aerobic growth conditions of S. suis resulted in a mismatched classification. Therefore, the reproducibility of this serotyping method was growth-condition specific, as previously reported for Burkholderia pseudomallei, where altering the incubation condition lowered the identification scores of Burkholderia pseudomallei [18]. Therefore, various growth conditions, including the anaerobic growth of S. suis...
should be recommended and emphasized when using this MALDI-TOF-MS classification coupled with corresponding PMFs database. That only a moderate agreement between the MALDI-TOF-MS and mPCR analyses (unweighted Kappa score of 0.522) pointed out that the results among these methods were controversial. The mPCR is not without limitations, such as ambiguous serotype classifications corresponding to the moderate degree (76%) of the true serotype classification [13], while however, these divergent characters were not displayed in phylogenetic tree SS2 and SS14 from SS1/2 and SS1, respectively. The degree of agreement measured how well the two tests agreed with each other, but not on the reliability to correctly serotype S. suis. Nevertheless, 95% CI of both methods indicated that the true serotype classification rate of both methods was not statistically different. Theoretically, in order to improve the kappa score, more PMFs of S. suis should be included, since the accuracy of the classification depends on the number of reference spectra present in the database [25].

**Conclusions**

In present study, we successfully differentiated the ambiguous serotypes of S. suis. Thus, providing an alternative method for S. suis classification that could be useful for S. suis infection diagnosis and epidemiological study of this zoonotic pathogen.

**Supporting information**

S1 Table. The complete list of m/z of S. suis serotyping by MALDI-TOF-MS identified and used in this study. (XLSX)

**Acknowledgments**

We thank the staffs Center of Excellence for Emerging and Re-emerging Infectious Diseases in Animals, Chulalongkorn University, Bangkok, Thailand for supporting the PCR thermocycler and techniques.

**Author Contributions**

- **Conceptualization:** Sittiruk Roytrakul, Suphachai Nuanualsuwan.
- **Data curation:** Chadaporn Chaiden, Janthima Jaresitthikunchai, Sittiruk Roytrakul.
- **Formal analysis:** Chadaporn Chaiden, Janthima Jaresitthikunchai, Sittiruk Roytrakul.
- **Funding acquisition:** Suphachai Nuanualsuwan.
- **Investigation:** Chadaporn Chaiden, Janthima Jaresitthikunchai, Sittiruk Roytrakul.
- **Methodology:** Janthima Jaresitthikunchai, Sittiruk Roytrakul.
- **Resources:** Anusak Kerdsin, Nattakan Meekhanon.
- **Supervision:** Janthima Jaresitthikunchai, Sittiruk Roytrakul.
- **Validation:** Suphachai Nuanualsuwan.
- **Visualization:** Chadaporn Chaiden.
- **Writing – original draft:** Chadaporn Chaiden.
- **Writing – review & editing:** Chadaporn Chaiden, Janthima Jaresitthikunchai, Anusak Kerdsin, Sittiruk Roytrakul, Suphachai Nuanualsuwan.
References

1. Gottschalk M, Segura M, Xu J. *Streptococcus suis* infections in humans: the Chinese experience and the situation in North America. Anim Health Res Rev. 2007; 8(1):29–45. https://doi.org/10.1017/S146625307001247 PMID: 17692141.

2. Staats JJ, Feder I, Okwumabua O, Chengappa MM. *Streptococcus suis*: past and present. Vet Res Commun. 1997; 21(6):381–407. https://doi.org/10.1023/a:1005870317757 PMID: 9266699.

3. Hill JE, Gottschalk M, Brousseau R, Harel J, Hemmingsen SM, Goh SH. Biochemical analysis, cpn60 and 16S rDNA sequence data indicate that *Streptococcus suis* serotypes 32 and 34, isolated from pigs, are *Streptococcus orisratti*. Vet Microbiol. 2005; 107(1–2):63–9. https://doi.org/10.1016/j.vetmic.2005.01.003 PMID: 15795078.

4. Tien LHT, Nishiibori T, Nishiitani Y, Nomoto R, Osawa R. Reappraisal of the taxonomy of *Streptococcus suis* serotypes 20, 22, 26, and 33 based on DNA-DNA homology and sodA and recN phylogenies. Vet Microbiol. 2013; 162(2–4):842–9. https://doi.org/10.1016/j.vetmic.2012.11.001 PMID: 23245487.

5. Goyette-Desjardins G, Auger JP, Xu J, Segura M, Gottschalk M. *Streptococcus suis*, an important pig pathogen and emerging zoonotic agent—an update on the worldwide distribution based on serotyping and sequence typing. Emerg Microbes Infect. 2014; 3(6):e45. https://doi.org/10.1038/emi.2014.45 PMID: 26038745; PubMed Central PMCID: PMC4078792.

6. Ishida S, Tien le HT, Osawa R, Tohya M, Nomoto R, Kawamura Y, et al. Development of an appropriate PCR system for the reclassification of *Streptococcus suis*. J Microbiol Methods. 2014; 107:66–70. https://doi.org/10.1016/j.mimet.2014.09.003 PMID: 2529648.

7. Stevenson LG, Drake SK, Murray PR. Rapid identification of bacteria in positive blood culture brothsm by matrix-assisted laser desorption ionization-time of flight mass spectrometry. Journal of clinical microbiology. 2010; 48(2):444–7. https://doi.org/10.1128/JCM.01541-09 PMID: 19955282.

8. Pérez-Sancho M, Vela AI, García-Seco T, Gottschalk M, Domínguez L, Fernández-Garayzabal JF. Assessment of MALDI-TOF-MS as alternative method for *Streptococcus suis* identification. Frontiers in public health. 2015; 3. https://doi.org/10.3389/fpubh.2015.00202 PMID: 26347858.

9. Pérez-Sancho M, Vela AI, García-Seco T, Gonzalez S, Domínguez L, Fernández-Garayzabal JF. Usefulness of MALDI-TOF-MS as a Diagnostic Tool for the Identification of *Streptococcus* Species Recovered from Clinical Specimens of Pigs. PLoS One. 2017; 12(1):e0170784. https://doi.org/10.1371/journal.pone.0170784 PMID: 28125697; PubMed Central PMCID: PMC5268416.

10. Alaitoom AA, Cunningham SA, Ihde SM, Mandrekar J, Patel R. Comparison of direct colony method versus extraction method for identification of gram-positive cocci by use of Bruker Biotyper matrix-assisted laser desorption ionization-time of flight mass spectrometry. J Clin Microbiol. 2011; 49(8):2868–73. https://doi.org/10.1128/JCM.01541-09 PMID: 21613431; PubMed Central PMCID: PMC3147777.

11. Chatellier S, Harel J, Zhang Y, Gottschalk M, Higgins R, Devriese LA, et al. Phylogenetic diversity of *Streptococcus suis* strains of various serotypes as revealed by 16S rRNA gene sequence comparison. Int J Syst Bacteriol. 1998; 48 Pt 2:581–9. https://doi.org/10.1099/00207713-48-2-581 PMID: 9731300.

12. Lowry OH, Rosebrough NJ, Farr AL, Randall RJ. Protein measurement with the Folin phenol reagent. J Biol Chem. 1951; 193(1):265–75. PMID: 14907713.

13. Kerdsin A, Akeda Y, Hatrongjit R, Detchawana U, Sekizaki T, Hamada S, et al. *Streptococcus suis* serotyping by MALDI-TOF-MS using a new multiplex PCR. J Med Microbiol. 2014; 63(Pt 6):824–30. https://doi.org/10.1099/jmm.0.00506-11 PMID: 24696517.

14. Watson PF, Petrie A. Method agreement analysis: a review of correct methodology. Theriogenology. 2010; 73(9):1167–79. https://doi.org/10.1016/j.theriogenology.2010.01.003 PMID: 21038353.

15. Seng P, Drancourt M, Gouriet F, La Scola B, Fournier PE, Rolain JM, et al. Ongoing revolution in bacteriology: routine identification of bacteria by matrix-assisted laser desorption ionization time-of-flight mass spectrometry. Clin Infect Dis. 2009; 49(4):543–51. https://doi.org/10.1086/600885 PMID: 19583519.

16. Schmitt BH, Cunningham SA, Dailey AL, Gustafson DR, Patel R. Identification of anaerobic bacteria by Bruker Biotyper matrix-assisted laser desorption ionization-time of flight mass spectrometry with on-plate formic acid preparation. J Clin Microbiol. 2013; 51(3):782–6. https://doi.org/10.1128/JCM.02420-12 PMID: 23254126; PubMed Central PMCID: PMC3592085.

17. Piamsomboon P, Jaresitthikunchai J, Hung TG, Roytrakul S, Wongvatvatchai J. Identification of bacterial pathogens in cultured fish with a custom peptide database constructed by matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI-TOF-MS). BMC Vet Res. 2020; 16(1):52. https://doi.org/10.1186/s12917-020-2274-1 PMID: 32046727; PubMed Central PMCID: PMC7014616.

18. Nyompanich S, Jaresitthikunchai J, Sirisanga K, Roytrakul S, Tungpradabkul S. Source-identifying biomarker ions between environmental and clinical *Burkholderia pseudomallei* using whole-cell matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI-TOF-MS). PLoS One.
19. Wang J, Zhou N, Xu B, Hao H, Kang L, Zheng Y, et al. Identification and cluster analysis of *Streptococcus pyogenes* by MALDI-TOF mass spectrometry. PLoS One. 2012; 7(11):e47152. https://doi.org/10.1371/journal.pone.0047152 PMID: 23144803; PubMed Central PMCID: PMC3492366.

20. Okura M, Nozawa T, Watanabe T, Murase K, Nakagawa I, Takamatsu D, et al. A locus encoding variable defense systems against invading DNA identified in *Streptococcus suis*. genome Biol Evol. 2017; 9(4):1000–12. https://doi.org/10.1093/gbe/evx062 PMID: 28379509; PubMed Central PMCID: PMC5398294.

21. Kerdsin A, Oishi K, Sripakdee S, Boonkerd N, Polwichai P, Nakamura S, et al. Clonal dissemination of human isolates of *Streptococcus suis* serotype 14 in Thailand. J Med Microbiol. 2009; 58(Pt 11):1508–13. https://doi.org/10.1099/jmm.0.013656-0 PMID: 19661209.

22. Gomez-Torres J, Nimir A, Cluett J, Aggarwal A, Elsayed S, Soares D, et al. Human case of *Streptococcus suis* disease, Ontario, Canada. Emerg Infect Dis. 2017; 23(12):2107–9. https://doi.org/10.3201/eid2312.171005 PMID: 29148386; PubMed Central PMCID: PMC5708224.

23. Perch B, Pedersen KB, Henrichsen J. Serology of capsulated streptococci pathogenic for pigs: six new serotypes of *Streptococcus suis*. J Clin Microbiol. 1983; 17(6):993–6. https://doi.org/10.1128/JCM.17.6.993-996.1983 PMID: 6874916; PubMed Central PMCID: PMC272789.

24. Auger JP, Dolbec D, Roy D, Segura M, Gottschalk M. Role of the *Streptococcus suis* serotype 2 capsular polysaccharide in the interactions with dendritic cells is strain-dependent but remains critical for virulence. PLoS One. 2018; 13(7):e0200453. https://doi.org/10.1371/journal.pone.0200453 PMID: 30001363; PubMed Central PMCID: PMC6042740.

25. Pavlovic M, Huber I, Konrad R, Busch U. Application of MALDI-TOF-MS for the identification of food borne bacteria. Open Microbiol J. 2013; 7:135–41. https://doi.org/10.2174/1874285801307010135 PMID: 24358065; PubMed Central PMCID: PMC3866695.