Letter to the Editor

Application of protein profiling of virulent Haemophilus parasuis by MALDI-TOF mass spectrometry

Luisa Z Moreno¹, Givago FR Silva¹, Vasco TM Gomes¹, Carlos EC Matajira¹, Ana Paula S Silva¹, Renan E Mesquita¹, Nicholas P Lotto¹, Thais SP Ferreira¹, Ana Paula G Christ², Maria Inês Z Sato², Yuri Gherpelli³, Michele Dottori³, Paolo Bonilauri³, Andrea Luppi³, Andrea M Moreno¹

¹ Faculdade de Medicina Veterinária e Zootecnia, Universidade de São Paulo, São Paulo, SP, Brasil
² Companhia de Tecnologia de Saneamento Ambiental (CETESB). São Paulo, SP, Brasil
³ Istituto Zooprofilattico Sperimentale della Lombardia e dell’Emilia Romagna (IZSLER). Brescia, Italia

Key words: Haemophilus parasuis, MALDI-TOF MS, serovar, swine.


(Received 09 October 2015 – Accepted 26 November 2015)

Dear Editor,

Haemophilus parasuis is the causative agent of Glasser’s disease and is one of the major opportunistic pathogens in swine intensive production systems, leading to substantial financial losses worldwide [1]. To date, 15 H. parasuis serovars have been described varying in virulence and geographical prevalence. However, the existence of a large number of non-typable isolates represents an important challenge for Glasser’s disease control, as vaccine immunity presents limited cross-serovar protection [2].

H. parasuis is classified as a fastidious organism that is nicotinamide adenine dinucleotide (NAD) dependent, and the standard methods of isolation and biochemical and immunological characterization for this bacterium are considered troublesome and time consuming [3]. Although molecular techniques have succeeded in diagnosing H. parasuis [3], the challenges of achieving proper molecular identification and serotyping at the lowest possible cost still remain.

In recent years, matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI-TOF MS) has become an important bioanalytical diagnostic tool for the detection of protein profiles from whole bacterial cells [4]. This rapid and accurate method can be easily applied to identify bacteria at the genus, species and, in some cases, the subspecies levels [5]. Therefore, MALDI-TOF MS represents a promising alternative to the standard phenotypic and molecular techniques carried out in diagnostic laboratories. In the present study, we assessed the use of MALDI-TOF MS for the characterization of Haemophilus parasuis isolated from different Brazilian states between 2009 and 2014.

A total of 98 H. parasuis isolates were studied. The samples were isolated from 66 animals in 40 herds from nine different Brazilian states (Mato Grosso, Mato Grosso do Sul, Minas Gerais, Paraná, Rio Grande do Sul, Santa Catarina, Distrito Federal and São Paulo) in Brazil from 2009 to 2014. Serotyping was previously performed by the immunodiffusion test using autoclaved antigens [6] at the Istituto Zooprofilattico Sperimentale della Lombardia e dell’Emilia Romagna (IZSLER, Italy). H. parasuis identification was also confirmed by PCR based on species-specific amplification of the 16S rRNA gene, as described by Oliveira et al. [7]. All isolates were also positive for the amplification of the group 1 vtaA gene [8] that has been associated with virulent H. parasuis strains.

For MALDI-TOF MS sample preparation, bacterial proteins were extracted using an ethanol/formic acid protocol [9]. The protein suspension (1 µl) was transferred to a polished steel MALDI target plate (Bruker Daltonik, Bremen, Germany) and allowed to dry at room temperature. The sample was overlaid with 1 µl of matrix (10 mg α-cyano-4-hydroxy-cinnamic acid ml⁻¹ in 50% acetonitrile/ 2.5% trifluoroacetic acid), and mass spectra in the 2–20 kDa range were acquired using a Microflex mass spectrometer (Bruker Daltonik).
Figure 1: Dendrogram showing the relationship among the *H. parasuis* isolates spectral profiles.
Each sample was distributed over six spots and measured four times, resulting in a total of 24 spectra per isolate.

For the MALDI-TOF identification, the spectra were loaded into MALDI BioTyper 3.0 and compared with the manufacturer’s library, which resulted in the log (score) value. Standard Bruker interpretative criteria were applied; scores ≥ 2.0 were accepted for species assignment and scores ≥ 1.7 but ≤ 2.0 for genus identification. For further analysis, at least 20 technical spectral replicates were used to generate a main spectrum for each isolate in Bionumerics 7.5 (Applied Maths NV, Saint-Martens-Latem, Belgium). Cluster analysis was performed using the number of different peaks detected and UPGMA method; principal component analysis (PCA) was applied to analyze the homogeneity of the H. parasuis spectra.

All isolates were identified using MALDI BioTyper with log (score) values > 2.0 for H. parasuis; a mean log (score) value of 2.226 was calculated. Furthermore, a slight variation among replicates for each isolate was observed, which corroborates the importance of using more than one spot per sample and even more than one spot measure to ensure bacterial identification and its reproducibility. Although MALDI-TOF MS has been validated for the identification of several microorganisms [5,10], the technique still presents some limitations with regard to spectral variation and reproducibility. The main causes for this are bacterial growth conditions and protein extraction methods [10].

To avoid interference of the medium used for bacterial isolation and maintenance, the culture broth was initially centrifuged; the pellet was washed with TE (pH 8.0) and then dissolved in deionized water for further protein extraction. In the initial MALDI-TOF MS identification studies, the protein purification or extraction step was only applied when no identification was obtained using the direct colony transfer method. As it has been proven that protein extraction improves identification yield, this step has been integrated into MALDI protocols, especially for intra-species and strain analyses [11,12].

The feasibility of using MALDI-TOF MS for H. parasuis identification was expected, as Kuhnert et al. [9] previously described the MALDI-TOF MS application for Pasteurellaceae identification from animals. However, the comparison of the protein profiles by clustering and PCA has not been reported yet. The protein spectral cluster analysis separated the H. parasuis isolates into two main groups, A and B (Figure 1), and the group A further divides into two subgroups (A1 and A2). The PCA also confirmed this clustering tendency (Figure 2). Although it was possible to observe intra-group similarity among the H. parasuis isolates, no relationship between the spectral clusters and the isolate’s origin and serovar was observed.

As only abundant housekeeping proteins, particularly ribosomal proteins, are detected in MALDI-TOF MS analysis [13] and the ethanol/formic acid protocol results in total bacterial protein extraction, this could explain the lack of association among the spectral clusters and the isolate’s origin and intrinsic characteristics. Moreover, the intra-species variation demonstrated by the spectral cluster analysis has also been found in H. parasuis molecular typing [14, 15]. As the MALDI-TOF MS technique is improving for specific protein detection, it could be used for further H. parasuis applied research particularly for characterizing membrane proteins patterns among serovars due to its association with virulence.

Acknowledgements
This study was supported by FAPESP (grant 2012/19154-5), CAPES and CNPq.

References
population structure and serovar in *Haemophilus parasuis*.


**Corresponding author**
Andrea M. Moreno. FMVZ/USP
Av Prof Dr Orlando Marques de Paiva, 87, Cidade Universitária,
CEP 05508 270. São Paulo/ SP – Brazil.
Phone: +55 021 11 3091-1377
Fax: +55 021 11 3091-7928
Email: morenoam@usp.br

**Conflict of interests:** No conflict of interests is declared.