Heterogeneity of Moraxella isolates found in the nasal cavities of piglets

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Abstract

Previous studies have shown that the genus Moraxella is commonly present in the nasal microbiota of swine. In this study, 51 isolates of Moraxella were obtained from nasal swabs from 3-4 week old piglets, which represented 26 different fingerprints by enterobacterial repetitive intergenic consensus (ERIC)-PCR. Whole 16S rRNA gene sequencing allowed the identification at species level of the Moraxella spp. isolates. The majority of the field strains were identified as Moraxella pluranimalium, but Moraxella porci was also detected. In addition, a cluster of 7 strains did not group with any described Moraxella species, probably representing a new species. Subsequent phenotypic characterization indicated that strains of Moraxella pluranimalium were mainly sensitive to the serum complement, while the cluster representing the putative new species was highly resistant. Biofilm formation capacity was very variable among the Moraxella spp. isolates, while adherence to epithelial cell lines was similar among selected strains. Additionally, selected strains were tested in phagocytosis assays and again variability was observed in the susceptibility to alveolar macrophages. Antimicrobial tests evidenced the existence of multidrug-resistance in the strains. In summary, phenotypic characterization revealed heterogeneity among Moraxella strains from the nasal cavity of piglets. Strains with pathogenic potential were detected as well as those that may be commensal members of the nasal microbiota. However, the role of Moraxella in porcine diseases and health should be further evaluated.

Background

*Moraxella* is microbiota member of the upper respiratory tract in vertebrates, but
some species may cause opportunistic infections. Best-known Moraxella infections are produced by Moraxella catarrhalis, which causes respiratory infections, including pneumonia and otitis in human [1]. In livestock, Moraxella bovis is well-known as the etiological agent of infectious keratoconjunctivitis in cattle [2]. In swine, the genus Moraxella has representative species, such as Moraxella porci, which was discovered and isolated from meninges [3] and Moraxella pluranimalium, described from an abdominal cavity isolate [4], but the relevance of this genus in swine health is not well established. Nonetheless, Moraxella may play a role in swine as a member of the nasal microbiota, since it has been detected as one of the most abundant genera in the nasal cavity of weaned piglets [5], is abundant in the nasal microbiota of slaughter age pigs [6] and can be found in environmental samples in farrowing buildings [7].

The microbiota plays an important role in host health and disease through different mechanisms, such as maturation of the immune system, improvement of the mucosal barrier and resistance against pathogens [8]. In the case of the nasal microbiota, early colonizers can determine the stability and composition of bacterial community leading to a healthier status, as reported for children early colonized by high abundance of Moraxella, which was associated with less respiratory infections [9]. However, the specific role of Moraxella, among other residents of the respiratory microbiota, in health is still unknown in pigs. In this study, we performed a genotypic and phenotypic characterization of 51 Moraxella fieldisolates from the nasal turbinates of weaned piglets. We assessed their capacity of evasion of the innate immunity by examining their complement and phagocytosis susceptibility, as well as other characteristics closely related to its ecological niche, such as biofilm formation, mucin adhesion and cell adherence. Our results showed genotypic and
phenotypic heterogeneity among the Moraxella isolates.

Results

Bacterial identification and genotyping

Initial identification of bacterial isolates by partial 16S rRNA gene sequencing showed a total collection of 51 Moraxella spp. isolates. Genotyping by enterobacterial repetitive intergenic consensus (ERIC)-PCR determined 26 different ERIC fingerprints, which were not shared among farms (Table 1). Up to 7 different fingerprinting profiles were isolated from the same farm, and up to 3 from a single piglet. One isolate of each fingerprinting (from now on referred as different strains for clarity throughout the text) were selected for further analysis.

Sequencing of approximately 1360 bp of the 16S rRNA gene allowed a more precise identification of the Moraxella isolates. Sequences from the nasal isolates and sequences from Moraxella sp. type strains from the Ribosomal database were used to build an UPGMA (unweighted pair group method with arithmetic mean) tree (Figure 1). Sixteen nasal strains clustered with more than 99.5% identity with Moraxella pluranimalium, while one strain, LL–3, was a bit more divergent, with 99.4% identity. Strain EJ45–1 clustered with Moraxella porci, with 99.2% identity. Strain CR–7A showed similarity to M. porci and Moraxella cuniculi, but did not clearly cluster with any of them. Finally, a group of 7 nasal strains showed homology among them of more than 98%, but did not clustered with any of the already described Moraxella species, indicating that they may represent a new species.

Antimicrobial susceptibility

Moraxella strains from nasal microbiota showed high diversity in antibiotic
susceptibility (Table 2). In general, all the strains were sensitive to amoxicillin+clavulanate and florfenicol, in contrast to trimetoprim + sulfamido and oxytetracycline, where we found a high resistance rate. Other antimicrobials with low resistance rates were ceftiofur and colistin. Six strains showed resistance to amoxicillin, which was abolished with the β-lactamase inhibitor clavulanate. M. pluranimalium UK1–20 was the only nasal strain without antimicrobial resistances, followed by M. pluranimalium LG6–2 with only one resistance. Both strains were isolated from a farm with no antimicrobial use before the sampling time. Multidrug-resistant (MDR), defined as resistance to 3 or more classes of antimicrobials, was found in 20 out of the 26 strains. Although the number of isolates and farms is very low, MDR was associated with the use of antibiotics in the farm of origin (two-tailed Fisher test; \( P = 0.0129 \)).

**Serum susceptibility**

Initially, bacterial suspensions were prepared in PBS obtaining irregular results, due to auto-agglutination by some strains. To overcome this problem, all the strains were resuspended in 20% glycerol in PBS and passed through a needle if necessary; this process did not affect the viability of the bacteria (not shown).

When the strains were assayed for survival after incubation with fresh rabbit serum, 17 of the 26 (65%) strains showed a reduction in viability of more than 3 logarithms after incubation with serum (Figure 2). Within *M. pluranimalium*, 12 out of the 16 strains (75%) showed a reduction of viability of more than 3 logs, similar to the type strain of this species, while the rest 4 (25%) showed a reduction of 2–3 logs. On the other hand, only 2 of the 7 (29%) strains from the cluster representing a putative new *Moraxella* species showed a reduction of more than 3 logs in viability, while the rest of strains in this cluster (5/7; 71%) showed a reduction of viability of less than
2 logarithms (Figure 2), indicating resistance to the serum complement. Moreover, strains showing the highest resistance to serum were observed within the isolates from the cluster of the putative new *Moraxella* species, and belonged to the same farm (VL1-4, VL3-9 and VL1-5; Figure 2). *M. porci* type strain SN9-4M, as well as strain *M. porci* EJ45-1, showed high susceptibility to the treatment with rabbit sera (Figure 2).

**Biofilm assays**

In an attempt to select non-agglutinating variants, we performed several passes of the agglutinating strains UK1-20 and VL6-6 in broth culture. Each passage was performed with bacteria taken from the surface of the culture that did not agglutinate. Curiously, a thick biofilm was observed in the culture tubes of UK1-20 after 27 passes (UK1-20p27); the bacteria in these cultures did not agglutinate but stuck to the glass. In the case of VL6-6, a biofilm in the air-liquid interphase was observed after 27 passes (VL6-6p27) (Figure 3). This observation prompted us to examine the natural level of biofilm formation capacity of the rest of the nasal *Moraxella* sp. strains. We found different rates of biofilm formation that were not dependent on the *Moraxella* species. Observing the results, different levels of biofilm formation could be arbitrarily established: production of biofilm when Abs_{590} ≥ 0.6 and of strong biofilm when Abs_{590} > 1.5. With those definitions, strains showing biofilm production capacity were *M. pluranimalium* GM3-2, GM8-1, EJ44-2A, LG6-2, UK1-20, GM5-1, VL6-4 and VL9-7; *Moraxella* sp. LL-3; *Moraxella* sp. (putative new species cluster) VL6-6; and also the variants UK1-20p27 and VL6-6p27 used as controls (Figure 4). Few isolates (*M pluranimalium* GM3-2 and LG6-2, and *Moraxella* sp. LL-3) were able to form strong biofilms (Figure 4). In general, strains forming biofilm under static conditions formed biofilm also under shaking conditions,
although at different level. Under static conditions, we observed a common trend of decreasing biofilm production when comparing 24 and 48h. In contrast, we observed the opposite effect under shaking conditions, with most of the strains showing a tendency of increasing biofilm production with time. Strains with slow growth (unclassified cluster) showed the same trend in decreasing biofilm production under both conditions. Biofilm formation was not affected by pre-coating the wells with mucin or bovine serum albumin (BSA) (data not shown).

**Cell adhesion**

To explore whether the biofilm formation was associated with cell adhesion capacity, selected strains with different rates of biofilm formation were assayed in PK-15 and A549 cell lines. Although the assayed strains presented adhesion ability to cells, the observed differences in biofilm formation capacity did not correlate with differences in cell adhesion to both PK-15 (not shown) or A549 cells (Figure 5).

**Phagocytosis susceptibility**

Selected strains representing the diversity in phylogenetic, serum resistance and biofilm formation were tested in phagocytosis assays with porcine alveolar macrophages (PAMs). Different levels of association with PAMs were observed among the strains (Figure 6). Within the 4 *Moraxella pluranimalium* strains tested, only LG6–2 presented a clear association with PAMS, similar to the level observed in the susceptible *H. parasuis* strain used as control. *Moraxella porci* EJ45–1, unlike the *M. porci* type strain SN9–4M, showed high levels of association with PAMs. The three isolates from the putative new species cluster presented low association with macrophages, even lower than the *H. parasuis* phagocytosis resistant strain used as control (Figure 6). Levels of phagocytosis did not correlate with biofilm formation capacity or serum resistance (Table 3).
In vitro culture passages affected phagocytosis susceptibility

*In vitro* passages affected the autoagglutination and biofilm formation capacity of *M. plurianimalium* UK1–20 and *Moraxella* sp. VL6–6 as indicated above. Although enhanced adhesion to epithelial cells was not observed in UK1–20p27 (not shown), this strain showed higher association with PAMs than the original strain (Figure 6). In contrast, VL6–6p27 showed the same level of interaction with PAMs as the original strain VL6–6 (Figure 6); this may be due to a different adaptation to laboratory conditions, as observed by the different type of biofilm formed by this strain.

Discussion

*Moraxella* spp from the nasal microbiota of piglets showed heterogeneous characteristics, from antimicrobial resistance profiles to virulent mechanisms and adherence properties. The majority of the isolates belonged to *M. plurianimalium* species, but we also detected *M. porci*, a couple of isolates of uncertain classification and a cluster of strains that may constitute a new *Moraxella* species. This cluster comprised strains with a slower growth rate in broth than the rest of the nasal strains and contained most of the strains with high resistance to the serum complement.

Our results on antimicrobial resistance confirm that the administration of these drugs can promote a rise of resistances, since higher rates of antibiotic resistance were found in strains from farms undergoing antibiotic treatments. MDR strains were broadly detected, and only one strain, UK1–20, from a farm where no antibiotics were used, did not show resistance to any of the antimicrobials tested. It
is important to highlight the presence of four MDR strains of *Moraxella pluranimalium*, with a high number of drug resistances, including colistin. This latter observation is consistent with the recent discovery of colistin resistance genes in *Moraxella* [15–17]. Coding genes of antibiotic resistances can be horizontally transferred between bacteria of the same or even different species, and therefore, the presence of resistance genes in the resident microbiota is of concern since they could be transferred to potential pathogens [18]. Diversity of the nasal *Moraxella* isolates was also evident in the *in vitro* assays, where strains showed a wide heterogeneity.

Biofilms are traditionally considered forms of resistance to environmental conditions and a virulence mechanism in bacteria of clinical importance [19]. As for other characteristics, we observed heterogeneity in biofilm formation capacity of the porcine nasal strains of *Moraxella*. Variability in biofilm formation has been already reported for other species of *Moraxella* from ruminants [20]. Biofilm formation capacity can be associated also to the colonizing ability of the bacteria, as was observed for *H. parasuis* [21]. The interaction of the nasal *Moraxella* strains with the respiratory mucosa needs more study, but attachment to mucin as well as to epithelial cells could play a role, as suggested by our results.

On the other hand, bacteria from the upper respiratory tract can occasionally reach the lower tract, where they will be confronted with other components of the immune system, such as the alveolar macrophages. *In vitro* phagocytosis assays provided insight into the potential of the nasal isolates to survive this first barrier of cell immunity in the lung. Heterogeneity in this feature has been shown for other bacterial species, such as *H. parasuis* [14] or *Klebsiella pneumoniae* [22]. In our case, strains with high association to PAMs, such as LG6–2, were observed, while
other strains did not associate with the phagocytic cells, as the case of the virulent SN9-4M. According to the results, phagocytosis susceptibility is not associated with the phylogeny of the isolates, since heterogeneity in this trait was observed within the distinct species or clusters. One paradigmatic case could be EJ45-1 and SN9-4M from *M. porci*, having high differences between them. The importance of phagocytosis resistance for *in vivo* infection/colonization by *M. pluranimalium* is also supported by the fact that originally the UK1-20 strain showed resistance to PAMs, but lost this trait after *in vitro* culture passages, in agreement with the loss of virulence after laboratory adaptation in other bacteria [23]. Loss of phagocytosis resistance in UK1-20 was concurrent with increase biofilm formation capacity. Thus, biofilm formation may be relevant in environmental persistence, whereas phagocytosis resistance will be important during host infection. However, this phenomenon seems to be strain-dependent, since it was not observed with strain VL6-6, which did not show an adaptation to laboratory conditions that affected its original resistance to PAMs.

Most of the assayed *Moraxella* spp. strains showed sensitivity to the serum complement, with some exceptions, including most of the strains in the cluster representing a putative new species. Thus, most nasal *Moraxella* strains are probably poorly invasive and will be kept on the nasal mucosa, but the new putative species cluster deserves more attention to define their role in disease in pigs.

**Conclusions**

The characterization of *Moraxella* spp. isolates from the nasal cavities of young piglets demonstrated a high heterogeneity within this genus. Most of the isolates were identified as *M. pluranimalium*, and only one as *M. porci*. A group of 7 isolates
clustered together by 16S rRNA gene sequence and did not show homology to any known *Moraxella* species. Most of the *Moraxella* spp. strains were resistant to multiple antimicrobial classes. Heterogeneity of the *Moraxella* strains was also evident in adhesion and virulence-associated assays. Further analyses, including experimental infections, need to be done to explain the function of this genus in swine health and disease.

**Methods**

**Isolation and bacteria cultures**

Nasal cavities from 10 piglets of around 3 weeks of age were sampled from 8 commercial farms (Table 1) with standard management practices for husbandry and welfare of the animals. Sampling of piglets was done under institutional authorization and followed good veterinary practices. According to European (Directive 2010/63/EU of the European Parliament and of the Council of 22 September 2010 on the protection of animals used for scientific purposes) and Spanish (Real Decreto 53/2013) normative, this procedure did not require specific approval by an Ethical Committee. Nasal sampling was performed only once to each piglet and is not likely to cause pain, suffering, distress or lasting harm equivalent to, or higher than, that caused by the introduction of a needle in accordance with good veterinary practice (Chapter I, Article 1, 5 (f) of 2010/63/EU). Nasal swabs were transported in Amies medium to the laboratory, where they were plated on chocolate agar (Biomérieux, Marcy l’Étoile, France) to isolate colonies. After 24 and 48 hours of incubation at 37°C and 5% CO₂, different colonies were selected and stored at −80°C in 20% glycerol-Brain Heart Infusion broth (BHI) for further characterization. Bacterial suspensions were also performed in phosphate-buffered
saline (PBS) for DNA extraction.

In addition, reference strains of *Moraxella porci* SN9-4M and *Moraxella pluranimalium* CD12-CA4, both isolated from lesions of diseased pigs, were used in this study. For phagocytosis assay, 2 reference strains of *Haemophilus parasuis* were used as controls.

**Bacterial identification and genotyping**

DNA extraction was performed using Chelex based Instagene™ Matrix (Bio-Rad Laboratories, Hercules, CA, USA) following manufacturer’s instructions. Preliminary identification of isolates was performed by partial sequencing of the 16S rRNA gene using primers 358F (CTACGGGAGGCAGCAGT) and 907R (CCGTCWATTCTTGGAGTTT) [10]. Sequences were analyzed by blasting against the Ribosomal database (http://rdp.cme.msu.edu).

All isolates identified as *Moraxella* were then genotyped by ERIC-PCR [11] with primers ERIC–1F (ATGTAAGCTCCTGGGGATTCAC) and ERIC–2R (AAGTAAGTGACTGGGGGTGAGCG). PCR reaction mixture consisted of 3 mM of MgCl₂, 1.2 µM of each primer, 0.23 mM of dNTPs, 0.75 U of GoTaq® polymerase (Promega, Madison Wisconsin, USA) and 100 ng of DNA sample. Amplification was carried out with an initial denaturation of 94°C for 2 min followed by 30 cycles of 30 sec at 94°C, 1 min at 50°C and 2.5 min at 72°C, and a final extension of 20 min at 72°C. Isolates showing different fingerprinting pattern were considered distinct strains and were chosen for further analysis.

Final identification of the different *Moraxella* strains was performed by 16S rRNA gene amplification with universal primers 8F (AGAGTTTGATCCTGGCTCAG) and 1492R (CGTTACCTTGTACGACTT) [12] and sequencing with 8F, 1492R, 358F and 907R primers. Sequence analysis was performed with Fingerprinting II v3.0 software (Bio-
Antimicrobial susceptibility

Susceptibility to several antimicrobials was tested as previously described [13]. Neo-Sensitabs™ diffusion tablets (Rosco Diagnostica, Taastrup, Denmark) were used for Gentamicin, Ceftiofur, Colistin, Erythromycin, Lincoespectin, Oxytetracycline; Doxycycline, Trimetoprim + Sulfamide (T+S), Enrofloxacin, Amoxicillin and Amoxicillin + Clavulanic acid testing. As no clinical breakpoints are available for these bacteria, inhibition diameters were compared with the manufacturer’s breakpoints for general microorganisms. Marbofloxacin, florfenicol and tulathromycin were tested in chocolate agar plates at different dilutions, from 16 to 0.125 µg/mL in the case of florfenicol and marbofloxacin; and from 64 to 0.125µg/mL in the case of tulathromycin. McFarland suspensions of 0.5 in 0.9% NaCl were prepared for each strain and streaked on the plates with the different antibiotic concentration. Minimal inhibitory concentration (MIC) was determined observing the bacterial growth after 24h of incubation. Isolates with values greater than or less than the diameter or concentration of the breakpoint suggested by the manufacturer were designated ‘sensitive’ and ‘resistant’, respectively.

Serum resistance assay

Serum resistance assay was carried out with rabbit serum (EU Directive 2010/63/EU and Spanish normative Real Decreto 53/2013 were followed). A bacterial suspension of each strain (representative of each ERIC fingerprinting) was prepared in PBS to reach an OD$_{600}$ of 0.3 in a VIS 7200 spectrophotometer (Dinko Instruments, Barcelona, Spain). In duplicate wells, 10 µl of the bacterial suspension (approx. $10^6$ colony forming units [CFU]/mL) were mixed with 90 ul of fresh filtered rabbit serum and mixtures were incubated for 1h at 37°C and 100 rpm. Bacterial survival was
calculated by comparing bacterial counts (obtained by serial dilutions and plating) at time 0 and after 1h incubation. The assay was carried out 3 times for each strain.

**Phagocytosis assay**

Phagocytosis assay was performed as previously described with PAMs [14]. PAMs were obtained from healthy piglets euthanized by intravenous pentobarbital overdose under institutional authorization. All procedures involving animals followed EU and Spanish normative (Directive 2010/63/EU and Real Decreto 53/2013). Lungs were removed and PAMs were isolated by lung lavage with PBS containing 70 µg/mL of gentamicin. PAMs were collected by centrifugation at 241 x g for 15 min, washed twice with sterile PBS and stored at –150ºC in 10% Dimethylsulfoxide (DMSO) in fetal bovine serum (FBS) until use.

Selected nasal strains of *Moraxella* were tested to determine their susceptibility to phagocytosis by PAMs. *Moraxella porci* reference strain SN9–4M was also included in the assay. In addition of two reference strains of *Haemophilus parasuis*, Nagasaki (virulent) and SW–114 (non-virulent), were included as control for the technique. For the assay, PAMs were plated in Dulbecco’s Modified Eagle Medium (DMEM) supplemented with 10% of FBS and 1% glutamine at a concentration of 5 x 10⁵ cells per well in 6 well-plates. Plates were incubated 1–2 hours to allow the attachment of the macrophages to the bottom of the wells. For each strain duplicate wells were inoculated with 10⁷ CFU of fluorescein isothiocyanate (FITC)-labelled bacteria (bacterial inocula were confirmed by dilutions and plating). After 1h of incubation at 37ºC and 5% CO₂, plates were transferred to an ice bath to stop phagocytosis and were washed twice with PBS to eliminate unbound bacteria. PAMs were then scrapped in PBS and were analyzed by flow cytometry using an EPICS XL-MCLTM
Flow Cytometer (Beckman Coulter, Madrid, Spain) or MACSQuant Analyzer 10 (Myltenyi Biotec, Bergisch Gladbach, Germany). Phagocytosis assay were repeated with PAMs from different animals that were already available at the laboratory. Since each batch of macrophages showed different level of phagocytosis ability, values were harmonized considering the reference *H. parasuis* strain SW114 (phagocytosis sensitive) as 100% phagocytosis in each assay; i.e., using the following calculation:

\[
\text{% PAMs with test bacteria} - \text{% fluorescence background in PAMs without bacteria} / \text{% PAMs with positive control SW114} - \text{% fluorescence background in PAMs without bacteria} \times 100.
\]

**Biofilm assays**

Biofilm assays were performed in 96 well cell culture plates under static and shaking conditions following previous published protocol with some modifications (Bello-Ortí et al. 2014). Bacterial suspensions were made to reach an OD$_{600}$ 0.3 in BHI. Wells in 96 well plates were then inoculated in quadruplicate with a 1:100 dilution of the bacterial suspension in BHI. Plates were incubated at 37ºC, under static conditions with 5% CO$_2$ or under agitation at 100 rpm for 24 and 48h, except for slow growing isolates, which were incubated for 48 and 72 h. After incubation, wells were emptied and rinsed with tap water to remove unattached bacteria. Wells were then stained with 0.1% (w/v) crystal violet (Merck, Darmstadt, Germany) for 2 minutes at room temperature. Wells were washed thrice with tap water to remove the excess of crystal violet and let dried at 37ºC. The dye in the stained biofilms was solubilized with 100 µL of 70% ethanol and quantified at 590 nm in a Powerwave XS Microplate Spectrophotometer (Biotek Instruments Inc., Winooski, VT, USA). Biofilm formation was also assessed on wells pre-coated with 5µg of mucin or
BSA as control.

**Cell adhesion assay**

Porcine epithelial cells PK-15 (ATCC® CCL-33) and human lung A549 (ATCC® CCL-185) cell lines were cultured in DMEM supplemented with 1% glutamine and 5% FBS for PK-15 and 10% for A549 cells.

Concentration of cells per ml was calculated to obtain a monolayer after overnight incubation, approximately 50,000 cells/well in cell culture 96 well plates. After overnight incubation, wells with PK-15 or A549 cells were washed once with sterile PBS. Duplicate wells were inoculated with $10^7$ CFU/mL bacteria from each isolate tested. Bacterial inoculum quantification was confirmed by colony counts. Microplates were then centrifuged 10 min at 100 x g to facilitate the contact between bacteria and cells. After incubation for 1 h at 37ºC and 5% CO₂, wells were washed twice with sterile PBS to remove unattached bacteria, and attached bacteria were released with 0.1% of saponin and pipetting. The resulting suspension was quantified by plating dilutions on agar plates.

**List of abbreviations**

BHI: Brain Heart Infusion

BSA: Bovine serum albumin

CFU: Colony forming units

DMEM: Dulbecco’s Modified Eagle Medium

DMSO: Dimethylsulfoxide

ERIC: Enterobacterial repetitive intergenic consensus

FBS: Fetal bovine serum

FITC: Fluorescein isothiocyanate
MDR: Multidrug-resistant
MIC: Minimal inhibitory concentration
PAMs: Porcine alveolar macrophages
UPGMA: Unweighted pair group method with arithmetic mean

Declarations

Ethics approval and consent to participate
Consent for sampling from the owners of the animals was obtained. Piglet sampling was part of routine veterinary practice in the farms. Sampling of piglets was done under institutional authorization and followed good veterinary practices. According to European (Directive 2010/63/EU of the European Parliament and of the Council of 22 September 2010 on the protection of animals used for scientific purposes) and Spanish (Real Decreto 53/2013) normative, this procedure did not require specific approval by an Ethical Committee. Nasal sampling was performed only once to each piglet and is not likely to cause pain, suffering, distress or lasting harm equivalent to, or higher than, that caused by the introduction of a needle in accordance with good veterinary practice (Chapter I, Article 1, 5 (f) of 2010/63/EU). Blood sampling of rabbit to obtain serum was performed following good veterinary practices and the above mentioned normative. For PAMs, following Article 18 of 2010/63/EU (Sharing organs and tissues), lungs were obtained from animals killed for other reasons.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests. None of the authors has any financial or personal relationships that could inappropriately influence or bias
the content of the paper.

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**Authors’ contributions**

SLS performed and analyzed the experiments and drafted the manuscript, NGM, MCH and AMPR performed experiments, and VA conceived the study, analyzed results and helped to draft the manuscript. All authors have read and approved the manuscript.

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**Availability of data and material**

The datasets analysed during the current study are available from the corresponding author on reasonable request.

**References**

1. Verduin CM, Hol C, Fleer A, van Dijk H, van Belkum A. Moraxella catarrhalis: from emerging to established pathogen. Clin Microbiol Rev. 2002;15:125–44.

2. Postma GC, Carfagnini JC, Minatel L. Moraxella bovis pathogenicity: an update. Comp Immunol Microbiol Infect Dis. 2008;31:449–58.

3. Vela Al, Sanchez-Porro C, Aragon V, Olvera A, Dominguez L, Ventosa A, Fernandez-Garayzabal JF. Moraxella porci sp. nov., isolated from pigs. Int J Syst
4. Vela Al, Arroyo E, Aragon V, Sanchez-Porro C, Latre MV, Cerda-Cuellar M, Ventosa A, Dominguez L, Fernandez-Garayzabal JF. Moraxella pluranimalium sp. nov., isolated from animal specimens. Int J Syst Evol Microbiol. 2009;59:671-4.

5. Correa-Fiz F, Fraile L, Aragon V. Piglet nasal microbiota at weaning may influence the development of Glasser’s disease during the rearing period. BMC Genomics. 2016;17:404.

6. Weese JS, Slifierz M, Jalali M, Friendship R. Evaluation of the nasal microbiota in slaughter-age pigs and the impact on nasal methicillin-resistant Staphylococcus aureus (MRSA) carriage. BMC Vet Res. 2014;10:69.

7. Cormier Y, Tremblay G, Meriaux A, Brochu G, Lavoie J. Airborne microbial contents in two types of swine confinement buildings in Quebec. Am Ind Hyg Assoc J. 1990;51:304–9.

8. Fung TC, Olson CA, Hsiao EY. Interactions between the microbiota, immune and nervous systems in health and disease. Nat Neurosci. 2017;20:145–55.

9. Biesbroek G, Tsvitsivadze E, Sanders EA, Montijn R, Veenhoven RH, Keijser BJ, Bogaert D. Early respiratory microbiota composition determines bacterial succession patterns and respiratory health in children. Am J Respir Crit Care Med. 2014;190:1283–92.

10. Lane DJ. 16S/23S rRNA sequencing. In: Stackebrandt E, Goodfellow M, eds. Nucleic Acid Techniques in Bacterial Systematics. John Wiley & Sons, England, UK, 1991;pp 115–65.

11. Versalovic J, Koeuth T, Lupski JR. Distribution of repetitive DNA sequences in eubacteria and application to fingerprinting of bacterial genomes. Nucleic
12. Weisburg WG, Barns SM, Pelletier DA, Lane DJ. 16S ribosomal DNA amplification for phylogenetic study. J Bacteriol. 1991;173:697–703.

13. Lorenzo de Arriba M, Lopez-Serrano S, Galofre-Mila N, Aragon V. Characterisation of Bergeyella spp. isolated from the nasal cavities of piglets. Vet J. 2018;234:1–6.

14. Olvera A, Ballester M, Nofrarias M, Sibila M, Aragon V. Differences in phagocytosis susceptibility in Haemophilus parasuis strains. Vet Res. 2009;40:24.

15. AbuOun M, Stubberfield EJ, Duggett NA, Kirchner M, Dormer L, Nunez-Garcia J, Randall LP, Lemma F, Crook DW, Teale C, Smith RP, Anjum MF. mcr-1 and mcr-2 (mcr-6.1) variant genes identified in Moraxella species isolated from pigs in Great Britain from 2014 to 2015. J Antimicrob Chemother. 2017;72:2745–9.

16. Poirel L, Kieffer N, Fernandez-Garayzabal JF, Vela AI, Larpin Y, Nordmann P. MCR-2-mediated plasmid-borne polymyxin resistance most likely originates from Moraxella pluranimalium. J Antimicrob Chemother. 2017;72:2947–9.

17. Kieffer N, Nordmann P, Poirel L. Moraxella Species as Potential Sources of MCR-Like Polymyxin Resistance Determinants. Antimicrob Agents Chemother. 2017;61:pii: e00129–17.

18. Penders J, Stobberingh EE, Savelkoul PH, Wolffs PF. The human microbiome as a reservoir of antimicrobial resistance. Front Microbiol. 2013;4:87.

19. Hall-Stoodley L, Costerton JW, Stoodley P. Bacterial biofilms: from the natural environment to infectious diseases. Nat Rev Microbiol. 2004;2:95–108.

20. Ely VL, Vargas AC, Costa MM, Oliveira HP, Potter L, Reghelin MA, Fernandes AW, Pereira DIB, Sangioni LA, Boton SA. Moraxella bovis, Moraxella ovis and
Moraxella bovoculi: biofilm formation and lysozyme activity. J Appl Microbiol. 2019;126:369–76.

21. Bello-Orti B, Deslandes V, Tremblay YD, Labrie J, Howell KJ, Tucker AW, Maskell DJ, Aragon V, Jacques M. Biofilm formation by virulent and non-virulent strains of Haemophilus parasuis. Vet Res. 2014;45:104.

22. Cavalcanti FCN, Rodrigues JF, Cabral AB, Azarias LCB, Morais Junior MA, Castro C, Lopes ACS. Relationships between phagocytosis, mucoid phenotype, and genetic characteristics of Klebsiella pneumoniae clinical isolates. Rev Soc Bras Med Trop. 2019;52:e20190089B.

23. Gaillard ME, Bottero D, Castuma CE, Basile LA, Hozbor D. Laboratory adaptation of Bordetella pertussis is associated with the loss of type three secretion system functionality. Infect Immun. 2011;79:3677–82.

Tables

Table 1. Sampled farms and nasal Moraxella spp isolated from piglets at weaning (3-4 weeks of age).
| Farm | Selected strains | Isolates with same ERIC PCR profile | Health status | Antibiotic |
|------|------------------|-------------------------------------|--------------|------------|
| GM   | GM3-2            | GM4-1                               | Healthy      | A          |
|      | GM8-1            | -                                   |              |            |
|      | GM5-1            | GM7-2, GM9-1, GM10-1                |              |            |
|      | GM5-5            | GM7-6                               |              |            |
|      | GM5-7            | GM7-8, GM7-9, GM8-4                 |              |            |
| AR   | AR-5A            | AR-5C                               | Healthy      | A          |
|      | EJ43-3A          | -                                   |              |            |
|      | EJ44-2A          | -                                   |              |            |
|      | EJ45-1           | EJ45-5                              |              |            |
| UK/LG| UK1-12           | UK1-32, LG1-4, LG1-6, LG2-2, LG3-7 | Healthy      |            |
|      | UK1-20           | LG5-5, LG8-3                        |              |            |
|      | LG6-2            | -                                   |              |            |
|      | LG6-4            | LG6-6                               |              |            |
|      | LG6-7g           | -                                   |              |            |
| CR   | CR-18            | -                                   | Healthy      | A          |
|      | CR-10            | -                                   |              |            |
|      | CR-7A            | CR-12                               |              |            |
| KD   | KD4-7            | -                                   | Healthy      |            |
| VL   | VL6-4            | -                                   | Healthy      | TUL        |
|      | VL9-7            | VL9-8, VL10-1, VL10-2               |              |            |
|      | VL6-6            | -                                   |              |            |
|      | VL1-4            | VL1-7, VL5-3                        |              |            |
|      | VL3-9            | -                                   |              |            |
|      | VL2-5            | -                                   |              |            |
|      | VL1-5            | VL1-6                               |              |            |
| LL   | LL-3             | -                                   | Nervous signs| Unl        |

1 Antibiotics administered before weaning (before sampling)

AMOX, Amoxicillin; ENR, enrofloxacin; CEFT, ceftiofur; TUL, tulathromycin

Table 2. Antimicrobial susceptibility of nasal Moraxella strains isolated from 3-4 week old piglets. As no clinical breakpoints are available for these bacteria, inhibition diameters were compared with the manufacturer’s breakpoints for general microorganisms.
| Strain       | COL | ENR | MARB | OXYTET | DOX | S+T | AMOX | AMC | CEF |
|-------------|-----|-----|------|--------|-----|-----|------|-----|-----|
| M. plurimalium |     |     |      |        |     |     |      |     |     |
| GM3-2       | S   | R   | R    | R      | S   | R   | S    | S   | S   |
| GM8-1       | S   | R   | R    | R      | I   | R   | I    | S   | S   |
| AR-5A       | S   | I   | R    | R      | R   | R   | R    | S   | S   |
| EJ43-3A     | S   | I   | R    | S      | I   | I   | R    | S   | R   |
| EJ44-2A     | R   | R   | R    | R      | R   | R   | R    | S   | R   |
| LG6-2       | S   | S   | S    | I      | S   | R   | I    | S   | S   |
| UK1-12      | S   | S   | S    | S      | S   | S   | S    | S   | S   |
| UK1-20      | S   | S   | S    | S      | S   | S   | S    | S   | S   |
| CR-18       | S   | S   | S    | R      | S   | R   | R    | S   | S   |
| GM5-1       | S   | I   | R    | R      | I   | R   | S    | S   | S   |
| KD4-7       | S   | S   | R    | S      | S   | R   | S    | S   | S   |
| LG6-4       | S   | S   | S    | I      | S   | R   | I    | S   | S   |
| LG6-7g      | S   | S   | R    | I      | S   | R   | I    | S   | S   |
| VL6-4       | R   | R   | R    | R      | R   | R   | R    | S   | S   |
| CR-10       | S   | R   | R    | R      | I   | R   | I    | S   | S   |
| VL9-7       | I   | I   | R    | R      | R   | R   | S    | S   | S   |
| LL-3        | S   | R   | S    | R      | I   | R   | I    | S   | I   |
| M. porci    |     |     |      |        |     |     |      |     |     |
| EJ45-1      | R   | I   | I    | R      | R   | R   | S    | S   | S   |
| Other Moraxella | |     |      |        |     |     |      |     |     |
| GM5-5       | S   | S   | S    | R      | S   | R   | I    | S   | S   |
| VL6-6       | S   | S   | S    | R      | R   | R   | S    | S   | S   |
| VL1-4       | R   | I   | R    | R      | R   | R   | S    | S   | S   |
| VL3-9       | S   | I   | I    | R      | R   | R   | R    | S   | S   |
| VL2-5       | S   | S   | S    | I      | S   | R   | S    | S   | S   |
| GM5-7       | S   | S   | S    | S      | S   | S   | S    | S   | S   |
| VL1-5       | S   | S   | S    | R      | S   | R   | S    | S   | S   |
| CR-7A       | S   | R   | R    | R      | I   | R   | S    | S   | S   |

COL, colistin; GEN, gentamicin; ERY, erythromycin; ENR, enrofloxacin; TET, tetracycline; DOX, doxycycline; S + T, trimethoprim-sulphonamide; AMOX, amoxicillin; CEFT, ceftiofur; AMC, amoxicillin + clavulanate; LI + SP, lincomycin-spectinomycin; MARB, marbofloxacin; TUL, tulathromycin; FLOR, florfenicol.

Table 3. Summary of results from phagocytosis and serum susceptibility and biofilm formation capacity for selected *Moraxella* spp. isolates.
| Strains | Biofilm formation | Serum susceptibility | Phagocytosis |
|---------|------------------|----------------------|--------------|
| M. pluranimalium | | | |
| LG6-2 | + | S | S |
| UK1-20 | + | S | R |
| LG6-7g | - | S | R |
| VL6-4 | + | S | R |
| Moraxella sp. | | | |
| VL6-6 | + | S | R |
| VL1-4 | - | R | R |
| VL1-5 | - | R | R |
| M. porci | | | |
| EJ45-1 | - | S | S |
| SN9-4M | - | S | R |

a Strains in the putative new Moraxella species cluster
b Abs590 > 0.8 in the crystal violet assay
c Viability reduction of more than 3 logarithms

Figures
Figure 1

UPGMA tree built with the 6S rRNA gene sequences of the Moraxella spp. isolates
Serum resistance assay showed diverse CFU reduction among the nasal Moraxella
Effect of culture passages on biofilm formation. Biofilm produced in glass tubes
Biofilm formation by Moraxella spp. isolates. Biofilm formation under static and shaking conditions in microtitre plates was assessed by measuring absorbance at 590 nm (Abs590nm).

**Figure 4**

Bacteria adhesion to A549 cells. Two biofilm forming strains, LG6-2 and UK1-20p27, were compared with non-forming strains UK1-20 and LG6-7g.

**Figure 5**

Bacteria adhesion to A549 cells. Two biofilm forming strains, LG6-2 and UK1-20p27, were compared with non-forming strains UK1-20 and LG6-7g.
Figure 6

Phagocytosis susceptibility of selected Moraxella spp. strains. Selected M. plurani

Supplementary Files

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