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Identification of enterobacteria in viscera of pigs afflicted with porcine reproductive and respiratory syndrome and other viral co-infections

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ABSTRACT

In order to investigate enterobacteria presence involved in the secondary infections in Porcine Reproductive and Respiratory Syndrome (PRRS) pigs with different viral co-infections, we identified enterobacteria for guiding clinical treatment. Twenty-one diseased pigs were diagnosed with the PRRS virus (PRRSV) and other 7 virus primers by PCR/RT-PCR in the lung and spleen samples. Enterobacteria were isolated using MacConkey agar from 5 visceral samples of PRRS pigs, and identified by 16S rDNA sequencing. PRRSV was positive in 100% of the lung samples and 81.0% of the spleen samples. Seven diseased pigs were diagnosed with only PRRSV infection (33.3%), 7 pigs with PRRSV and 1 or 2 other viruses (33.3%) and 7 pigs with PRRSV and more than 2 types of other viruses (33.3%). PRRSV was more inclined to co-infect pigs with porcine group A rotavirus (PARV) with the co-infection rate of 52.4% (11/21). Approximately 13 types of bacteria were successfully isolated from lung, spleen, liver, kidney and lymph node samples of different PRRS pigs. Enterobacteria were isolated in 100% of lung, liver and lymph samples from pigs infected with PRRSV alone. However, the isolation rates were significantly decreased in the more than 3 viruses co-infection group. *Escherichia coli* was the most prevalent bacterium, followed by *Morganella*, *Proteus*, *Shigella*, *Salmonella*, *Klebsiella* and *Aeromonas*. Most of the isolated enterobacteria were opportunistic pathogens. Therefore, timely combination with antimicrobial agents is necessary for effective treatment of PRRS-infected pigs.

1. Introduction

Porcine reproductive and respiratory syndrome (PRRS) is a swine disease caused by a virus, which poses a significant economic threat to the swine industry worldwide [1]. The PRRS virus (PRRSV) can infect pigs of various ages and cause different clinical symptoms. When the virus infects suckling pigs, it usually causes death, while there may be no significant features except for reproductive failure in some sows when the virus infects adult pigs. When the virus infects nursery pigs, it causes respiratory disorder, but does not directly result in death if no secondary infection occurs, unless pigs infected with some highly pathogenic PRRSV isolates [2]. Therefore, secondary infection is usually one of the main reasons for death in PRRS infected pigs [3,4].

In practical production, farmers focus more attention on ways to prevent and treat infected pigs. Farmers employ vaccine inoculation and isolated rearing to prevent infection [5]. However, there seem to be no effective treatment options. The infection of PRRSV itself could not cause death in nursery pigs. Secondary infection is the direct cause of death, but the pathogens of secondary infection have not been understood to date; thus, there is no effective treatment.

In the present study, to investigate bacteria involved in the secondary infection during the PRRS development, we isolated and identified enterobacteria in different viscera of PRRS pigs, and analysed the enterobacterial proliferation from the gut, a natural strain reservoir, to other viscera of PRRS pigs with different viral co-infections, to provide a reference therapy for secondary infection in pigs with PRRS.
2. Materials and methods

2.1. Samples collection

All the procedures involving animals in this study were carried out in accordance with The Care and Use Guidelines of Experimental Animals established by the Ministry of Agriculture of China, and the animal protocol for this study approved by the Ethics Committee of Jiangxi Agricultural University (protocol number JXAULL-2018004). The clinical manifestations of involved diseased pigs at 40–50 days old included fever (40–41 °C), coughing, purple ears, breathing difficulties, loss of appetite, diarrhoea and other surface symptoms. The autopsy manifestations of the above pigs were lung oedema with fibrinous exudate, groin and mesenteric lymph node enlargement and hemorrhage, mild spleen swelling, renal enlargement and paleness, and petechial hemorrhage. The samples of porcine lung, liver, spleen, kidney and lymph nodes were collected and transported to a lab under refrigerated conditions on the day of collection. Samples were collected from 21 diseased pigs and 2 healthy pigs from 5 different breeding farms in the Jiangxi Province in China.

2.2. Virus detection

The lung and spleen samples from pigs with PRRS were cut into pieces and homogenised to be used for virus detection, respectively. Viral RNA and DNA were extracted by using TaKaRa MiniBEST Viral RNA/DNA Extraction Kit (TaKaRa Inc, Dalian, China) according to the manufacturer’s instructions. Portion of each extraction was subjected to RNA reverse transcription by using PrimeScript™ – RT reagent Kit with gDNA Eraser (TaKaRa Inc, Dalian, China). The RNA reverse transcription was performed as follows: 1 μL of DNA/cDNA as template, 2 μL of Primer-F and Primer-R (10 μM), 12.5 μL of Premix Taq™ DNA Polymerase and addition of H2O to total 25 μL. A 3-step cycling protocol was used for polymerase chain reaction (PCR) as follows: 94 °C for 5 min, 30 cycles of 94 °C for 30 s, corresponding annealing temperature for 30 s, and 72 °C for 30 s, then 72 °C for 10 min. The PCR productions were examined by agarose gel electrophoresis.

2.3. Enterobacteria isolation and identification

The viscera, including lung, spleen, liver, kidney and lymph nodes from the pigs with PRRS were used for enterobacteria isolation. Firstly, the visceral specimen was removed from the plastic bag to a sterile plate under the biosafety cabinets, and 6 sections were cut by using a sterile blade at different positions of the specimen after the specimen surface was sterilized using 75% alcohol. An inoculating loop was used to scrape the different newly cut surfaces of sections and inoculate the samples onto MacConkey agar (Haibo Inc, Qingdao, China) individually. The enterobacteria from each visceral specimen were cultured on the MacConkey agars at 37 °C for 16 h. The colonies with different morphologies were chosen to inoculate another MacConkey agar for bacterial purification. After the colonies were purified, they were used for DNA extraction. The bacteria were identified using 16S rDNA sequencing (Tsingke Inc, Qingdao, China) with primers 27F and 1492R.

The 16S rDNA primer sequences and PCR fragment sizes are also listed in Table 1 [15–12]. The PCR reaction was performed as follows: 1 μL of DNA/cDNA as template, 2 μL of Primer-F and Primer-R (10 μM), 12.5 μL of Premix Taq™ DNA Polymerase and addition of H2O to total 25 μL. A 3-step cycling protocol was used for polymerase chain reaction (PCR) as follows: 94 °C for 5 min, 30 cycles of 94 °C for 30 s, corresponding annealing temperature for 30 s, and 72 °C for 30 s, then 72 °C for 10 min. The PCR productions were examined by agarose gel electrophoresis.

2.4. Statistical analysis

The statistical software SPSS (version 19.0) (International Business Machines Corporation, New York, USA) was used for data analysis. The statistical software SPSS (version 19.0) was used for data analysis. The viral positive detection rates between lung and spleen samples, and the bacterial positive isolation rates among different viscera were analysed using the chi-square test. The bacterial positive isolation rates among different viral infection groups were analysed using one-way analysis of variance (ANOVA) test. Differences were considered statistically significant at \( p \leq 0.05 \).

3. Results

3.1. Virus detection in lung and spleen samples of suspected PRRS-positive pigs

Forty-six lung and spleen samples from 21 suspected PRRS pigs and 2 healthy pigs were individually examined by PCR or RT-PCR for the presence of PRRSV and other possible viruses. The viral detection results are shown in Table 2. All suspected PRRS pigs showed positive detection for PRRSV in the lung samples, which confirmed that the diseased pigs were indeed infected with PRRSV. No virus was detected in healthy samples. PRRSV was positive in 100% of lung samples, and in 81.0% (17/21) of spleen samples, indicating that PRRSV was significantly easier to be detected in porcine lungs than in spleens (\( p < 0.05 \), Fig. 1).

In addition to PRRSV, some other viruses that could cause respiratory or digestive symptoms were also detected. These viruses included porcine epidemic diarrhoea virus (PEDV), transmissible gastroenteritis coronavirus (TGEV), porcine group A rotavirus (PARV), classical swine fever virus (CSFV), porcine circovirus type 2 (PCV2), pseudorabies virus (PRV) and porcine parvovirus (PPV). From the results of Table 2, TEGV, CSFV and PRV could not be detected from any of the lung or spleen samples, but PEDV, PARV, PCV2 and PPV had varying extents of positive detection. The positive rates of PEDV and PARV were higher in the spleen (33.3% and 52.4%) than in the lung samples (14.3% and 28.6%), while the positive rates of PCV2 were higher in the lung (47.6%) than in the spleen samples (38.1%). The positive rates of PPV were the same (33.3%) in both types of samples.

We further investigated the co-infection of PRRSV with other related viruses. As long as any of the lung or spleen samples were detected with the virus, the corresponding pig was thought to be infected with PRRSV. Seven diseased pigs were detected with only PRRSV (33.3%), 7 pigs detected with 2 or 3 types of viruses, and 7 pigs detected with 4 or 5 types of viruses. PRRS pigs were more inclined to be co-infected with PARV and PCV2, with co-infection rates of 52.4% (11/21) and 47.6% (10/21), respectively.

3.2. Enterobacteria from the visceral samples of pigs with PRRS

Through the above examination, we confirmed that the diseased pigs...
terium was isolated in healthy samples. Therefore, we mined that some other viruses may infect simultaneously with PRRSV.

PCR. * CSFV, PCV2, PRV and PPV being detected in lung and spleen samples using RT-PCR.

Table 2

| Code | PRRSV(L/S) | PEDV(L/S) | TGEV(L/S) | PARV(L/S) | CSFV(L/S) | PCV2(L/S) | PRV(L/S) | PPV(L/S) |
|------|------------|-----------|-----------|-----------|-----------|-----------|----------|----------|
| 1    | +/+        | -/-       | -/-       | -/—       | -/-       | -/-       | -/-      | -/-      |
| 2    | +/+        | -/-       | -/-       | -/-       | -/-       | -/-       | -/-      | -/-      |
| 3    | +/-        | -/-       | -/-       | -/-       | -/-       | -/-       | -/-      | -/-      |
| 4    | +/+        | -/-       | -/-       | -/-       | -/-       | -/-       | -/-      | -/-      |
| 5    | +/-        | -/-       | -/-       | -/-       | -/-       | -/-       | -/-      | -/-      |
| 6    | +/+        | -/-       | -/-       | -/-       | -/-       | -/-       | -/-      | -/-      |
| 7    | +/-        | -/-       | -/-       | -/-       | -/-       | -/-       | -/-      | -/-      |
| 8    | +/-        | -/-       | -/-       | -/-       | -/-       | -/-       | -/-      | -/-      |
| 9    | +/-        | -/-       | -/-       | -/-       | -/-       | -/-       | -/-      | -/-      |
| 10   | +/-        | -/-       | -/-       | -/-       | -/-       | -/-       | -/-      | -/-      |
| 11   | +/-        | -/-       | -/-       | -/-       | -/-       | -/-       | -/-      | -/-      |
| 12   | +/-        | -/-       | -/-       | -/-       | -/-       | -/-       | -/-      | -/-      |
| 13   | +/-        | -/-       | -/-       | -/-       | -/-       | -/-       | -/-      | -/-      |
| 14   | +/-        | -/-       | -/-       | -/-       | -/-       | -/-       | -/-      | -/-      |
| 15   | +/-        | -/-       | -/-       | -/-       | -/-       | -/-       | -/-      | -/-      |
| 16   | +/-        | -/-       | -/-       | -/-       | -/-       | -/-       | -/-      | -/-      |
| 17   | +/-        | -/-       | -/-       | -/-       | -/-       | -/-       | -/-      | -/-      |
| 18   | +/-        | -/-       | -/-       | -/-       | -/-       | -/-       | -/-      | -/-      |
| 19   | +/-        | -/-       | -/-       | -/-       | -/-       | -/-       | -/-      | -/-      |
| 20   | +/-        | -/-       | -/-       | -/-       | -/-       | -/-       | -/-      | -/-      |
| 21   | +/-        | -/-       | -/-       | -/-       | -/-       | -/-       | -/-      | -/-      |
| 22   | +/-        | -/-       | -/-       | -/-       | -/-       | -/-       | -/-      | -/-      |
| 23   | +/-        | -/-       | -/-       | -/-       | -/-       | -/-       | -/-      | -/-      |

S: Spleen; L: Lung; +: Positive; -: Negative.

Fig. 1. Detection of different types of viruses in lung and spleen samples of pigs afflicted with PRRS. PRRSV and the other 7 viruses may cause symptoms like breathing difficulties or diarrhoea syndrome, such as PEDV, TGEV, PARV, CSFV, PCV2, PRV and PPV being detected in lung and spleen samples using RT-PCR. *p < 0.05.

were indeed infected with PRRSV, simultaneously. We further determined that some other viruses may infect simultaneously with PRRSV. Nevertheless, the real causes of death among the nursery pigs with PRRS may be the pathogens causing secondary infections [3, 4]. Therefore, we further investigated the enterobacterial proliferation from the gut to the surrounding viscera, such as lung, spleen, liver, kidney and lymph nodes. From the enterobacterial isolation and identification results (Table 3), approximately 13 types of bacteria were successfully isolated from several viscera of different PRRS-positive pigs, and no enterobacterium was isolated in healthy samples.

If only one type of enterobacterium was isolated, the corresponding organ was supposed to be infiltrated by the gut bacteria. Based on this analysis (Fig. 2), we found that the total enterobacterial isolation rate from kidneys (42.9%, 9/21) were significantly lower (p < 0.05), and the total rates from the other 4 viscera were comparative in levels. We also found that when pigs were co-infected with more types of viruses, the lower bacterial isolation rate was obtained (p < 0.05). In the PRRSV alone infection group, enterobacteria were detected with a 100% positive rate in lung, liver and lymph samples, whereas in the co-infection group with more than 3 types of virus species, the isolation rates in lung, liver, kidney and lymph were significantly decreased. Only the rates from spleens were comparative.

Among the isolated enterobacteria, Escherichia coli was the most prevalent bacterium (Fig. 3). The total isolation rate of E. coli from lymph nodes was highest (81.0%, 17/21), while the rate from kidneys was lowest (28.6%, 6/21). Lower E. coli isolation rates were obtained in conjunction with infection with more types of viruses. Besides E. coli, Morganella morganii was found to have a higher positive isolation rate in almost all viscera than other 11 types of bacteria, especially in lymph nodes. Proteus sp., including Proteus mirabilis and Proteus vulgaris, were found to have a sub-higher positive isolation rate in the samples. In addition, some common pathogens, such as Salmonella, Shigella and Klebsiella were detected in the visceral samples (Fig. 4), and the positive detection rate of Shigella in lung samples was up to 23.8%, and the rate of Salmonella in lymph nodes was up to 28.6%. Aeromonas sp. was another bacterium with a positive isolation rate above 10%, although only in lung samples.

4. Discussion

PRRS is a panzootic and economically important disease in pigs. In nursery pigs, PRRS can cause severe respiratory tract symptoms, and may even lead to death. However, pure PRRS infection is not the main reason accounting for porcine death. Multiple organ failure caused by secondary infection may be the etiologic reason [3, 4]. As is well known, the gut is a natural microbiota reservoir, and the opportunistic pathogens in the gut of PRRS pigs may be very easy to proliferate to the surrounding viscera, resulting in the secondary infection. Thus, in the present study, we isolated and identified 13 strains of enterobacteria in different visceral samples from PRRS pigs. Most of them were opportunistic pathogens. We further investigated the association of the bacterial detection with the viral detection results, to try to find a new treatment strategy for PRRS.

In the current study, we first detected 7 other porcine viruses in addition to PRRSV in the lung and spleen samples to investigate the most...
common viruses that tend to co-infect with PRRSV. The result showed that PRRSV could be detected in 100% of lung samples, indicating the easier for PRRSV to invade the respiratory system and then affect other organs, such as the spleen [14]. For the 7 detected viruses, PARV was not detected from any lung or spleen samples. The reason for this may be possibility that more enterobacterial isolation rates from livers and lymph nodes were significantly lower, indicating that the kidney may not be easily colonized for viral co-infections resulted in less enterobacterial isolation, especially decreased. The reason accounted for this may be possibility that more enterobacterial isolation rates from different viral co-infected pigs were paralleled possibly because the spleen, liver, kidney and lymph node samples of PRRS pigs with different viral co-infections.

In addition to PRRSV, we isolated and identified the enterobacteria from viscera of the diseased pigs. We found that at least 1 strain of enterobacteria could be isolated from one of the viscera of the PRRS-positive pigs. The isolation rates of enterobacteria in lung and lymph node samples of PRRS pigs were tied highest (81.0%), followed by those in liver and spleen samples (both 76.2%). Only the isolation rate from kidneys was lower, indicating that the kidney may not be easily colonized for enterobacteria from the gut infiltrate. Moreover, we found that more viral co-infections resulted in less enterobacterial colonization, especially when there were 4 or 5 types of co-infecting viruses, and the entero-bacterial isolation rates from livers and lymph nodes were significantly decreased. The reason accounted for this may be possibility that more enterobacterial isolation rates cause more severe symptoms in diseased pigs, so that the enterobacteria may not have enough time to proliferate from the gut to other viscera prior to death. However, the isolation rates in spleens from different viral co-infected pigs were paralleled possibly because the spleen as the largest immune organ in the body was easier for pathogens to infiltrate.

Generally, more than one enterobacterium could be isolated in the
enterobacteria affecting animals and the environment worldwide, diseased pigs or to prevent other pigs in the same pigpen to develop some antibiotic with a quite broader antibacterial spectrum to treat the venting the secondary infection, it is also suggested to try to directly use drug sensitivity analysis can be conducted on these pathogenic bacteria, such as samples were higher, but this is not always the case. For opportunistic pathogens, such as urinary tract infections [17]. Of course, for opportune and effective preventive measures against these bacteria, so as to timely prevent them from spreading to other organs to cause secondary infection when the PRRSV infection is happened [23]. Of course, for opportune and effective preventive measures, it is also suggested to try to directly use some antibiotic with a quite broader antibacterial spectrum to treat the diseased pigs or to prevent other pigs in the same pigpen to develop infection, if there is indeed no enough time to conduct the anti-microbial drug sensitivity analysis.

In the current study, some other opportunistic enterobacteria could also be detected in different viscera. These enterobacteria included Providencia rettgeri, causing diarrhoea [24] or urinary tract infections [25]; Sphingomonas, causing mostly nosocomial, non-life-threatening infections [26]; Acinetobacter, causing various infections in unhealthy individuals [27]; Escherichia fergusonii, causing wound infections and even bacteraemia [28]; Providencia, causing infections associated with gastroenteritis and bacteraemia [29]; Stenotrophomonas, an opportunistic pathogen in highly debilitated patients [30]; and Cronobacter sakazakii, causing bacteraemia, meningitis and necrotising enterocolitis in infants [31]. At the same time, some commensal bacteria were isolated, such as Pasteurella aerogenes, Citrobacter freundii, Enterobacter cloacae and Enterobacter hormaechei. Among the isolated enterobacteria, Sphingomonas and Escherichia fergusonii were firstly reported to be isolated from swine samples. The bacterial identification results demonstrated that timely intervention with antimicrobial agents is necessary for effective treatment of PRRS pigs.

5. Conclusions

Summarily, in pigs with typical PRRS symptoms, PRRSV not only co-infected with other types of viruses, but was also found to result in enterobacterial proliferation from the gut to other viscera, and thus cause the secondary infections that threaten the lives of pigs. In our study, we identified the common enterobacterial pathogens causing secondary infections in PRRS pigs co-infected with other different viruses. Our results may provide some guidance for the clinical treatment of PRRS.

Author statement

Ge Zhao, Liheng Liu, and Junwei Wang were responsible for the conception, and design of the study. Charles Li participated in the experimental design and provided the guidance. Lujie Zhang and Liheng Liu was responsible for samples collection. Na Liu, Jianmei Zhao and Yuehua Li actively identified and analyzed the samples. Ge Zhao, Liheng Liu, and Junwei Wang contributed to drafting this manuscript. All authors have read and approved the final manuscript.

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Declaration of competing interest

The authors declared that they have no conflict of interest to this work.

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