Molecular detection of *Porcine circovirus* type 2 in swine herds of Eastern Cape Province South Africa

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**Abstract**

**Background:** *Porcine circovirus* type 2 (PCV2) remains the main causative viral pathogen of porcine circovirus-associated diseases (PCVAD) of great economic importance in pig industry globally. This present study aims at determining the occurrence of the viral pathogen in swine herds of the Province.

**Results:** The data obtained revealed that 15.93% of the screened samples (54/339) from the swine herds of the studied areas were positive for PCV2; while the severity of occurrence of the viral pathogen as observed at farm level ranges from approximately 5.6 to 60% in the studied farms. The majority (15 out of 17 = 88%) of the analyzed sequences were found clustering with other PCV2b strains in the phylogenetic analysis. More interestingly, two other sequences obtained were also found clustering within PCV2d genogroup, which is presently another fast-spreading genotype with observable higher virulence in global swine herds.

**Conclusion:** This is the first report of PCV2 in swine herds of the Province and the first detection of PCV2b and PCV2d in South African swine herds. It follows the first reported case of PCV2a in an outbreak of porcine multisystemic wasting syndrome (PMWS) in Gauteng Province, South Africa more than one decade ago. This finding confirmed the presence of this all-important viral pathogen in pigs of the region; which could result in a serious outbreak of PCVAD and huge economic loss at the instances of triggering factors if no appropriate measures are taken to effectively curb its spread.

**Keywords:** Porcine circovirus type 2, Eastern Cape Province, South Africa

**Background**

Porcine circoviruses (PCVs) are of the genus *Circovirus* in the family Circoviridae. They are non-enveloped viruses with a single-stranded circular DNA genome [1]. They are the smallest known animal viruses and include *porcine circovirus* type 1 (PCV1) and *porcine circovirus* type 2 (PCV2), with genome sizes of 1759 and 1767/1768 nucleotides, respectively [2]. PCV1 was first detected in 1974 as a contaminant of the porcine kidney cell line PK-15 (ATCC CCL-33) and was determined to be non-pathogenic [3]. However, PCV2 was found in pigs more than two decades later and proved to have the clinical manifestation of post-weaning multi-systemic wasting syndrome (PMWS) [4].

Globally, PCV2 is recognized as an emerging swine pathogen of great economic importance, causing huge losses in the piggery business. The viral pathogen has been the major culprit in cases of PMWS and other clinical disease manifestations in pigs that are generally regarded as *porcine circovirus*-associated diseases (PCVADs) [5]. These include porcine dermatitis and nephropathy syndrome (PDNS), porcine circovirus reproductive disorders, porcine respiratory disease complex, enteritis, acute pulmonary oedema, nervous system lesions, proliferative and necrotizing pneumonia, and a recently resurfacing neonatal congenital tremor [5, 6].

The viral genome of PCV2 is composed of at least five open reading frames (ORFs) that could be transcribed; however, ORF1 (coding for replication proteins) and ORF2 (coding for structural capsid protein) remain the widely sequenced and studied regions [7]. PCV2 generally infects...
7–16-week old weaners and growers, as younger pigs are protected by passive immunity conferred by acquired maternal antibodies [8]. PCV2 infection normally affects the immune system of infected pigs by causing lymphoid tissue depletion, which leads to observable histological lesions. Co-infections with other pathogens normally result in worsened conditions in the infected pigs and is made possible due to the immunosuppression and reduced immunity that result from the PCV2 attacks on the protection system [5].

PCV2 strains have been classified into four main genotypes (PCV2a, PCV2b, PCV2c, and PCV2d) based on phylogenetic analyses performed with their full genomes and ORF2 sequences [9]. Previously, PCV2c was detected only in Denmark from archived materials; however, it was recently found in live feral pigs in Brazil and from field samples taken from sick pigs in China [10–12]. The duo of PCV2a and PCV2b has a worldwide distribution; since 2003 however, there has been an observable worldwide genotypic shift in occurrence from PCV2a to PCV2b in pig herds, which has since made PCV2b the predominant genotype [13]. PCV2d (previously regarded as a mutant PCV2b) is the newly emerging genotype that is currently circulating the globe in swine herds and has an apparent higher virulence [14, 15].

PCV2-infected pigs normally shed the virus through many routes, including urine, faeces, milk, oronasal secretions, colostrum, and semen [16, 17], thereby enhancing the transmission and spread of PCV2 infections among local herds. The global transmission of PCV2 infections is also greatly enhanced through the international trade of live pigs and pig products due to the subtle nature of the disease, which can be in the form of a subclinical nature [13, 18, 19]. Hence, proper diagnostic measures to check for the wild spread of the viral pathogen become imperative. There are two major ways of diagnosing PCV2 infections. Preliminarily, this can be done by observing for clinical signs, but this may not be accurate or reliable in cases of subclinical infections [20]. The confirmatory detection of PCV2 nucleic acids or antigens in samples from infected animals is therefore imperative. This has been achieved over the years through polymerase chain reaction (PCR), immunohistochemistry (IHC), and in situ hybridization techniques [5].

Although PCV2 is rampant in domestic pig populations globally, severe PCVAD may not occur on very many occasions. Not all pigs within the affected herd develop PCVAD because PCV2 is not the only factor required for disease expression [20]. As a multifactorial disease, there have been numerous studies on infectious and non-infectious cofactors in PCV2 infections, which include the co-infection of PCV2 and other pathogens [20]. Non-infectious cofactors of PCV2 infection include the genetic background of the pig, management practices such as high stocking density, and prevailing environmental conditions such as temperature fluctuations within the pen [21, 22].

Being a multifactorial disease, the prevention and control of PCVAD have been achieved over time through all-encompassing measures that consider both infectious and non-infectious contributing agents. Although vaccination has proven effective in preventing PCV2 infection and spread within the herds, the effectiveness is better achieved by combining it with good management practices such as preventing cross-fostering, strict biosecurity practice, careful breed selection, maintaining high standards of hygiene through the effective use of disinfectants, and good housing conditions [23, 24].

Despite the ubiquitous status of the virus, it has recently been revealed that the virus is grossly understudied in sub-Saharan Africa and on the African continent at large [25]. In South Africa, PCV2 was first detected on a commercial breeding farm in 2001 in pigs with clinical manifestations of PDNS and PMWS [26]. To date, however, no large-scale study has been conducted on the prevalence of PCV2 and its associated diseases to ascertain the true infection status of the pigs in that country. The recommendations by Drew et al. [26] and An et al. [27] for further molecular epidemiological studies on PCV2 strains at other premises in South Africa have been grossly neglected.

Therefore, the focus of this study was to validate the current PCV2 status of pigs in South Africa by surveilling for its presence in the swine herds of Eastern Cape Province. This serves as the first surveillance on the pathogen’s occurrence in the swine herds of this region and helps to contribute more South African-generated PCV2 sequences to GenBank, as there are currently very few from the country. In addition, it serves as motivation to conduct further studies in the near future entailing wider geographic regions to accurately document the virus genogroups circulating in the country and the entire sub-Saharan African region.

Results
Characteristics and observable farm management practices on the sampled farms

Three hundred and seventy-five blood, faecal, and nasal-swab samples were collected from seven commercial and communal pig farms in three district municipalities of Eastern Cape Province in the years 2015 and 2016. From the administered questionnaire, it was obvious that virtually all of the managing personnel of the sampled farms were ignorant of PCV2 and its associated diseases. Moreover, the best practices that are required in a pig farming operation to prevent the transmission and outbreak of infectious diseases were essentially absent on most of the communal farms. Only one of the seven sampled farms employed high
biosecurity measures in their piggery operations. It was also highly notable that none of the farms vaccinated their pigs against PCV2, not even the sampled commercial farm. More than half (57%) of the sampled farms were not using disinfectants for routine cleaning of their pens. Furthermore, a majority of the sampled farms had experienced some PCVAD symptoms such as wasting, abortion, and respiratory distress (Table 1).

Discussion
Infectious swine pathogens such as PCV2 have greatly attracted the attention of researchers since the early 1990s due to the economic havoc they produced in many pig-producing countries of the world [18]. Since 1997, PCV2 has been one of the most studied pig pathogens and has been referred to as an everlasting, worldwide, endemic pathogen in swine [28]. Since its first detection, there has been a notable successive change in the predominance of the circulating viral pathogen genogroup at the global level, changing from PCV2a before the year 2003 to PCV2b after that year [13]. Presently, PCV2d is becoming the most rampant and ravaging group with its observably higher virulence [15]. Despite the concern and global efforts to study the epidemiology of the viral pathogen, there has been a seemingly lethargic approach to its study in South Africa and the entire sub-Saharan African countries [25]. However, the scant information available has confirmed the presence of different PCV2 genogroups in the region, as the previously called North American-like strain [PCV2 group 2 (PCV2a)] was initially detected in South Africa in 2001 [26], and the European cluster [PCV2 group 1 (PCV2b)] was found in Ugandan pigs over a decade later [29]. Considering the economic importance of pig production in the developing countries of the world [30], the importance of this study in determining the PCV2 status of pigs on a relatively larger scale cannot be overemphasized.

In this study, 54 of the 339 samples screened for the presence of PCV2 were positive for the pathogen, confirming its presence on all of the tested farms. In addition, the farm level severity of occurrence was as high as 60% on one of the surveyed farms. This supports the ubiquitous status of PCV2 in swine herds globally [20]. Some selected high-quality amplicons were sequenced and processed to obtain 17 partial PCV2 genomes that were then analysed. The phylogenetic analyses confirmed that 15 of the PCV2 sequences formed a group together with the PCV2b reference sequences and were closely clustered with the Chinese strains NQY1 (KX247842), FX1102 (JX406426), FJMHO508 (GU247990), 09HuB (HQ395035), RUZHOU (EU418626), and XT1 (KX247844). The remaining two PCV2 sequences were grouped with the PCV2d GenBank sequences, being closest to the Xuancheng (FJ712215) strain from China. The only available reference sequence from South Africa, strain SA1 (AY325495), maintained its position among the other PCV2a reference strains used in this analysis, thus confirming the findings of Drew et al. [26] (Fig. 1).

The topography observed in the phylogenetic clustering was further confirmed through the nucleotide and amino acid sequence alignments (Figs. 2 and 3), in which CHB16 and CHB19 were found to have mutations in the stop codon at the 3’ end of ORF2, leading to an amino acid (Lysine-K) extension. This is a typical feature that distinguishes the mutant PCV2b (mPCV2b, now categorized as PCV2d), which is known to be more virulent than PCV2a and 2b [14]. This outcome agrees with the observed global PCV2 infection pattern, which has had a notable genotypic shift around the world, first from PCV2a to PCV2b, and recently to PCV2d [15, 31, 32]. The same incidence of genotypic shift may have also occurred in South African pigs since Drew et al. [26] first reported the case of a PCV2a outbreak in 2001, which was traced to insemi- nating the affected gilts with imported semen from Iowa, North America. This assertion was made due to the clustering of the first South African PCV2 strain (SA1) with many other sequences from there and Asia.

Alternatively, it is possible that the PCV2b and 2d strains detected in this study have been in circulation within the country all this time, but undetected. This claim might be
| Farm code       | Farm one (2FTP) | Farm two (TSO) | Farm three (MTH) | Farm four (CHA) | Farm five (CHB) | Farm six (ICHQ) | Farm seven (CHD) |
|-----------------|-----------------|----------------|------------------|----------------|----------------|----------------|-----------------|
| Herds population| >5000           | 100            | 80               | 200            | 250            | 50             | 70              |
| Herds composition| Sows, weaners, growers and boars | Sows, weaners, growers and boars | Sows, weaners, growers and boars | Sows, weaners, growers and boars | Sows, weaners, growers and boars | Sows, weaners, growers and boars | Sows, weaners, growers and boars |
| Breeds          | Large white and Landrace | Large white     | Large white      | Large white and Landrace | Large white and Landrace | Large white | Large white |
| Biosecurity measures | High        | Very low       | None             | Very low       | Low            | None           | None            |
| Biocide use     | Virocide       | Nil            | Nil              | Dazzel dip     | Dazzel dip     | Nil            | Nil             |
| Vaccination regime | Nil         | Nil            | Nil              | Nil            | Nil            | Nil            | Nil             |
| Antibiotics use | Applied in cases of infections | Occasionally in cases of infections | Applied in cases of infections | Occasionally in cases of infections | Occasionally in cases of infections | Occasionally in cases of infections | Occasionally in cases of infections |
| Antibiotic /drugs use | Norotrim, Depomycin, Lantrax, Kyroligo, Advocin Ultralet, | Iron, Ivomec / dectomax | Iron, Ivomec / dectomax | Iron, Ivomec / dectomax | Iron, Ivomec / dectomax | Iron, Ivomec / dectomax | Iron, Ivomec / dectomax |
| All-in/All-out practice | Yes         | No             | No               | Yes, but not always | Yes, but not always | No             | No              |
| Disease symptoms and occurrence | Swollen lymph nodes, meningitis, respiratory disorders, red cat, gastric ulcers, scrotal and navel hernia | Meningitis, respiratory disorders, gastric ulcers, hernia | Respiratory disorders, dysentery | Respiratory disorders | Respiratory disorders | Swollen lymph nodes, meningitis, respiratory disorders |
| Prior knowledge of PCV2 infections | Yes | No             | No               | No             | No             | No             | No              |
| Cases of abortion | Not applicable | Occasionally | Occasionally | Occasionally | Occasionally | Occasionally | Occasionally |
substantiated on the basis that the initial detection in 2001 was from a case study of an outbreak in a commercial breeding unit that supplied breeding stock to other smaller farms [26] rather than from a large-scale surveillance study. In that case study, only 4 tissue samples from affected pigs were sent to the United Kingdom for analysis. Hence, an unnoticed transmission of different PCV2 genotypes from one herd to another could have been occurring within the country for many years prior to their being detected. Considering it was present at all of the tested farms, another scenario is that the PCV2b strains detected in this study entered through another route and are now spreading rapidly within the eastern region of the country. The occurrence of PCV2d in this study is highly significant and may represent the generally believed genotype shift from PCV2b to 2d, as has been observed in swine herds elsewhere in the world [15]. However, achieving a better understanding of this requires further prevalence studies and an in-depth molecular characterization of the PCV2 genotypes in circulation within the country.

The findings from this study suggest that the probable cost of neglecting proper PCV2 surveillance in South African pigs, as stated by Mokoele et al. [33], could be grievous and devastating with any instance of a large-scale PCV2 infection outbreak in the country. It is noteworthy that PCV2 was detected in all the sampled farms in the province, indicating a significant level of occurrence of the ‘small but powerful’ viral pathogen in the swine herds. The claim of negligence on the part of stakeholders in the country’s swine industry is further corroborated by the findings of our investigation. The pig farmers in the area had essentially no level of awareness (Table 1), and hence, were currently implementing no preventive measures.

Good management practice has been considered paramount in the prevention and control of PCV2 infections [24]. It is therefore not surprising that PCV2 was detected on all the sampled farms in this study, owing to the fact that the management practices of the majority of the farms was very poor. For example, a large percentage of the farms had little or no biosecurity measures in place that could forestall the farm-to-farm transmission of infectious agents such as PCV2. Furthermore, other important recommendations made in 1997 in Madec’s proposed 20-point plan to lower the impact of the disease [23] were grossly absent on the farms studied, such as the maintenance of good hygiene through the effective use of disinfectants and an all-in/all-out stocking practice. It is noteworthy, however, that despite the observance of good management practices, PCV2 was still present at the commercial farm (farm one) in the study. This observation further stresses the importance of an effective vaccination regime against PCV2 as part of an all-encompassing strategy in preventing infection [23, 24].

As a multifactorial disease having both infectious and non-infectious co-factors, PCVAD prevention and control rely on a prompt and adequate diagnosis, which is a function of efficient surveillance to ascertain the status of circulating PCV2 genotypes in the swine population. This normally informs the adoption of an effective vaccination regime and good management procedures to forestall rapid spread and the incidence of large-scale outbreaks. According to Segalés [34], continuous surveillance for new PCV2 variants is important for a DNA virus that, as Firth et al. report, has evolutionary rates comparable to that of RNA viruses [35]. The observable gross negligence by the stakeholders regarding PCV2 infection in South Africa and sub-Saharan African countries [25] at large could suppress the huge potential suggested for pig production to alleviate poverty and help solve hunger issues in developing countries [30].

**Conclusion**

The detection of PCV2b and its mutant strain (PCV2d) in the swine herds of Eastern Cape Province after first detecting PCV2a in South African pigs more than a decade and half ago is very disturbing considering the economic importance of this viral pathogen. It is therefore extremely important that a

| District Municipalities | Farms (Codes) | Farm types | Total Samples collected | Sampling Year | Samples screened | Positive samples | Positive samples (%) |
|------------------------|---------------|------------|-------------------------|---------------|-----------------|------------------|----------------------|
| AMATHOLE              | 2FTP          | Commercial | 206                     | 2015          | 206             | 19               | 9.22                 |
| O.R. TAMBO            | TSO           | Communal   | 24                      | 2015          | 24              | 2                | 8.33                 |
|                       | MTH           | Communal   | 14                      | 2015          | 14              | 2                | 14.29                |
| CHRIS-HANI            | CHA           | Communal   | 35                      | 2016          | 35              | 21               | 60.00                |
|                       | CHB           | Communal   | 59                      | 2016          | 36              | 2                | 5.56                 |
|                       | CHC           | Communal   | 13                      | 2016          | ND              | ND               | ND                   |
|                       | CHD           | Communal   | 24                      | 2016          | 24              | 8                | 33.33                |

ND Not Determined
Fig. 1 Phylogenetic analysis of the partial genomes of the PCV2 from this study with other sequences from GenBank. To further compare the derived sequences in this study with globally reported PCV2 sequences curated in GenBank, a phylogeny was performed using reference sequences from different geographic regions of the world in order to determine the evolutionary origins of the study sequences. The tree was constructed using 64 partial genomes of PCV2 (1041 nt) which comprised of 17 strains from this study and 47 other sequences obtained from GenBank while PCV1 sequence served as an out-group. The construction was done by using a Neighbor Joining algorithm and the percentage of replicate trees in which the associated taxa clustered together in the bootstrap test (1000 replicates) are shown next to the branches. The only PCV2 strain previously submitted to the GenBank from South Africa is indicated with the black square box. Analyses were conducted in MEGA6 [38].
rigorous enlightenment of the farmers about PCV2 and its associated diseases be undertaken in the country. This is highly imperative, as our survey revealed that virtually all of the farmers in the region were ignorant of the pathogen. An effective large-scale vaccination regime should also be initiated to curtail the rapid circulation of the virus and prevent impending future outbreaks. This should be done pari passu with educating the farmers (especially the communal ones) on the biosecurity measures required in piggery operations for optimal performance. Further studies on the prevalence of PCV2 and its co-infecting pathogens in the pigs of Eastern Cape Province and other South African provinces should also be conducted without further delay, with a candid effort geared towards obtaining more of the viral genomes (preferably complete) in circulation within the country. This will provide a thorough and immediate molecular characterization of the circulating strains and better our understanding of the epidemiology of this viral pathogen, both within the country and beyond.

**Methods**

**Sample collection**

A total number of three hundred and seventy five (375) field samples, consisting of blood, faecal and nasal swabs samples were collected from seven (commercial, semi-commercial and communal) farms from three District Municipalities of Eastern Cape Province, South Africa in 2015 and 2016. The fresh samples were randomly collected from both healthy and diseased pigs of the sampled farms; they were processed and kept in −80 °C freezer until when
used. A Questionnaire was also designed (Additional file 1) and administered to each of the sampled farm to obtain information about some farm management practices and level of awareness on PCV2 from the farm managers.

DNA extraction and PCV2 detection

Extraction of total genomic DNA was done from the processed samples by using ReliaPrep™ gDNA Tissue Miniprep System (Promega, Madison, USA) with strict adherence to the manufacturer procedures. Initial screening of the samples was performed through polymerase chain reaction (PCR) by using the primer pair P1 and P2, and subsequently, positive samples were again subjected to a second round of PCR amplification by using a primer pair P3 and P4 (Table 3). The first primer pair amplified a chunk of about 629 base pair (bp) long of ORF1 region (replicase gene) of the viral genome while the second overlapping primer pair amplified the remaining part of the ORF1 region and a portion of ORF2 (capsid gene) totaling about 630 bp in length.

The PCR amplification for PCV2 detection was carried out as earlier described by An et al. [27] with some modifications. The PCR reaction mixtures were made by adding 5 μL of extracted DNA to 45 μL of a reaction mixture containing a final concentration of 1.25 mM MgCl₂, 5X PCR buffer, 0.2 mM dNTPs, 10 pmol of each primer, and 2.5 U of Taq DNA polymerase. PCR amplification was done in a MyCycler™ (Thermocycler 1.065) machine (Bio-Rad, Applied Biosystem, California) with amplification conditions of 95 °C for 4mins (initial denaturation); 35 cycles (final denaturation of 95 °C for 30 s, annealing of 57 °C for 30 s, elongation of 72 °C for 1 min); final elongation of 72 °C for 5mins. PCR products were analyzed by electrophoresis on 1.5% Agarose gels stained with Ethidium bromide (EB) and visualized using an Alliance 4.7 transilluminator (UVitec, Cambridge, UK).

**Sequencing of amplified nucleotide sequences, analysis and construction of phylogenetic tree**

Positive PCR products of high quality were selected for sequencing at University of Stellenbosch Central DNA Sequencing Facility using the forward and reverse primers earlier used in PCR amplification. Post PCR clean-ups were performed using Nucleofast 96 well PCR plate (Macherey-Nagel, Düren, Germany) with adherence to the manufacturer’s instructions on a Tecan EVO150 robotic workstation (Tecan Group, Männedorf, Switzerland). The purified products were sequenced with primer pairs used for conventional PCR screening and subsequent nucleotide sequencing of partial PCV2 genomes

| Primer identity | Primer sequence | Amplicon length | Nucleotide Position | Reference |
|-----------------|-----------------|-----------------|---------------------|-----------|
| P1Fw            | 5'-TAATCCCTCGCAAGACGACG-3' | 629             | 116–135             | [27]      |
| P2Rv            | 5'-CGATCACACATCTCTGACG-3' | 629             | 726–745             | [27]      |
| P3Fw            | 5'-CAGAAAGGAGTTGGAAGAGC-3' | 630             | 531–550             | [27]      |
| P4Rv            | 5'-ATGCTGGGACGCTGAGCCCTC-3' | 630             | 1142–1161           | [27]      |
standard Sanger sequencing using the BigDye Terminator V3.1 sequencing kit (Applied Biosystems, Foster City, CA, USA) in line with the manufacturer’s instructions with slight modifications. Sequenced DNA were edited, blasted and assembled using Geneious 10.1.2 [36].

The execution of Basic Local Alignment Search Tool (BLAST) was carried out on the DNA sequences as an initial measure to ascertain that all the sequences were truly PCV2 in comparison with other sequences present in the GenBank. Nucleotide sequence alignment was performed by using Bioedit software [37] while amino acid alignment was done by using ClustalW as implemented in Geneious 10.1.2 software [36]. The Phylogenetic tree was reconstructed using the distance-based neighbor joining algorithm as implemented in Mega 6 [38]. Reliability was evaluated by the bootstrapping method on 1000 replicate of the alignment. All sequences that were obtained from GenBank and used for reconstructing the phylogenetic tree are as follow: PCU49186, AY556474, AY325495, AY322004, AJ223185, AF408635, AF381176, KM924366, KM924364, FJ870968, FJ870967, DQ104423, AF055392, AY691169, AY424405, AY322003, AY321985, KU041850, KU041849, HQ020970, HM038016, FJ870974, FJ870969, AF055394, KX247842, JX406426, EU418626, HQ395035, GU247990, KX247844, AF201311, EU148503, EU148504, EU148505, KU041859, KU041855, KJ680361, KJ680354, KJ511876, KX828241, AY181946, FJ712215, JX535296, KY425815 and KU311021 (see Additional file 2 for details).

### Additional files

**Additional file 1:** Questionnaire on porcine circovirus type 2 (PCV2) study. Designed questionnaire that was used to obtain information about some farm management practices and level of awareness on PCV2 from the managers of the sampled farms. (DOCX 17 kb)

**Additional file 2:** PCV2 sequences in this study and other reference sequences reported previously that were used in the phylogenetic analysis. Details of PCV2 sequences from this study and other reference sequences obtained from GenBank. (PDF 89 kb)

### Abbreviations

**BLAST:** Basic local alignment search tool; DNA: Deoxyribonucleic acid; IHC: Immunohistochemistry; NCBI: National Center for Biotechnology Information; ORF: Open reading frame; PCR: Polymerase chain reaction; PCVAD: Porcine circovirus associated disease; PCVs: Porcine circoviruses; PDNS: Porcine dermatitis and nephropathy syndrome; PMWS: Postweaning multisystemic wasting syndrome

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### Availability of data and materials

The partial genomic sequences of the 17 PCV2 strains reported in this paper are available on the NCBI GenBank database (https://www.ncbi.nlm.nih.gov/nuccore/KY985403) under accession numbers: KY985387 to KY985403.

### Authors’ contributions

BCI, LCO and AIO conceived the idea of the work while KOA also participated in its design; KOA and BCI carried out the sampling. KOA participated in the conceptual aspect of the work, performed the experiments and wrote the manuscript. BCI, LCO and AIO provided consultation and coordination. All authors read and approved the final manuscript.

### Ethics approval and consent to participate

The approval to conduct this research was obtained from the University of Fort Hare’s Research Ethics Committee with Certificate Reference Number: O8012 (Project). Strict compliance was given to the ethics code and the Animals Protection Act, 1962 (Act 71 of 1962) as rightly stipulated. Informed verbal consent had been provided by the animal owners prior to the collection of samples and the questionnaire’s administration.

### Consent for publication

Not applicable.

### Competing interests

The authors declare that there is no competing interest regarding the publication of this paper.

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