Research Article

Potential Pathogenicity and Genetic Characteristics of a Live-Attenuated Classical Swine Fever Virus Vaccine Derivative Variant

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Classical swine fever (CSF), caused by CSF virus (CSFV), is a highly contagious disease affecting pigs and causing massive pig production losses with severe global economic recession. The immunization of live-attenuated vaccines is still one of the key measures to CSFV management in endemic countries. However, there are also strong controversies about the usage of live-attenuated vaccines, particularly in pregnant sows and young pigs, such as in Europe, where domestic pigs are routinely not vaccinated until severe outbreaks occur. Here, we report a CSF outbreak in a pig farm in China, which affected more than 90% of the delivery sows and led to ∼45% birth loss. Surprisingly, phylogenetic analysis showed that the CSFV isolate (named CSFV/HeNLY2022, GenBank No. OR195698) was clustered into subgenotype 1.1a, closely together with the live-attenuated vaccine strains. Further genomic analysis also revealed that the isolate CSFV/HeNLY2022 shared the highest nucleotide identity of 99.7% with the C/HVRI vaccine strain (C-strain, GenBank No. AY805221). Moreover, compared to the C/HVRI strain, a total of eight amino acid mutations, distributed in Erns (H436thY and S476thR), E1 (T502thI and P581thT), E2 (M979thK and A1061thS), NS5A (A2980thT), and NS5B (I3818thM), were characterized in the CSFV/HeNLY2022 isolate. Our results suggested that the CSF outbreak was most likely caused by the live-attenuated CSFV vaccine or its derivative. It raises concern that the unscientific application of CSFV vaccines could potentially lead to CSFV spread in pigs. It is needed to perform a more rigorous evaluation of the safety of the C-strain-derived vaccines in combination with other different live-attenuated vaccines.

1. Introduction

Classical swine fever (CSF) remains one of the most economically important viral diseases to pigs worldwide and is notified to the World Organization for Animal Health [1]. The causative agent, CSF virus (CSFV), belongs to the genus Pestivirus (Pestivirus suis species) in the Flaviviridae family and is an enveloped, single-stranded, positive-sense RNA virus [2, 3]. Similar with the other members of the Pestivirus genus, the genome size is about 12.3 kb with 5′ untranslated region (5′ UTR), one open reading frame encoding a single polyprotein, and 3′ UTR. The polyprotein is further cleaved by host or viral proteases into four structural proteins (Core, E1, E2) and eight nonstructural proteins (P7, NS2, NS3, NS4A/B, NS5A/B) [4]. Based on the sequence of 5′ UTR, E2, and NS5B, CSFVs are genetically divided into three genotypes (1, 2, and 3) and at least 11 subgenotypes (1.1–1.4, 2.1–2.3, and 3.1–3.4) [1, 2]. Genotype 2 is predominantly circulating in Asia and Europe [4–6].
FIGURE 1: The clinical information of the pig farm and phylogenetic analysis of the CSFV isolates. (a) Mummies, rotten, and white fetuses were observed when the sows began to deliver. (b) The immune procedure in the pig farm. PPV: WH-1 strain, inactivated vaccine; JEV: SA14-14-2 strain, live-attenuated vaccine; PRV: Batha-K61 strain, live-attenuated vaccine; CSFV: cell culture adapted C-strain, live-attenuated vaccine;
During July 20–August 28, 2022, severe porcine reproductive disorders occurred at a pig farm in Henan province, China (Figure 1(a)). Clinical specimens, including seven tissues (spleen, inguinal lymph nodes, tonsil, and kidney) from stillbirths, newborn piglets, three sera from aborted sows, and two semen samples obtained from boars, were sent to a laboratory for pathogen detection on July 27, 2022. All samples were tested negative for PRRSV, PRV, PPV, and JEV by virus-specific nucleic acids tests. Serum from only one sow tested positive for PCV2 and PCV3. However, 10 of the 12 specimens were CSFV positive, and one semen sample was suspected for CSFV. Then, a survey was conducted by consulting with veterinarians on the pig farm. On March 14, the farm introduced 123 gilts (about 125 kg) from a professional breeding farm, where the pigs are free of several diseases, including CSFV, PRRSV, PRV, JEV, and PPV. The insemination was completed from April 1 to May 7. Meanwhile, live-attenuated vaccines were applied following the immune procedure, including JEV (SA14-14-2 strain) immunized on May 11 and 23, CSFV (cell culture adapted C-strain derivatives) and PRV (Batha-K61 strain) simultaneously immunized during May 28–30 (Figure 1(b)). Interestingly, the sows did not present any abnormal clinical symptoms until delivery. When the sows began to deliver, a large number of mummies, rotten, and white fetuses were observed, accounting for ~33.0% of the newborn piglets, and the mortality rate of living piglets reached about 10% within 7 days. Overall, approximately 90% of the delivery sows showed different degrees of reproductive disorders, and the birth loss was about 45%.

High genetic variability is a common feature of the CSFV genome [1]. To determine the genetic evolution of the CSFV strain in the pig farm, 13 pairs of primers were used to amplify the genomic sequence (Table S1). A near complete genome of the strain (named CSFV/HeNLY2022, GenBank No. OR195698) was acquired except the sequence of 478 bp at the 3’UTR. Based on the encoding sequence of the polyprotein, a phylogenetic tree was constructed by the neighbor-joining method using MEGA11.0. As shown in Figure 1(c), different with traditional CSFV classification, the CSFVs could be divided into two evolutionary groups-GI (includes genotypes 1 and 3) and GII (mainly genotype 2), consistent with the classification of a previous study [4]. Surprisingly, CSFV/HeNLY2022 was clustered into subgenotype 1.1a, closely together with the live-attenuated vaccine strains. Further genomic similarity analysis also revealed that the CSFV/HeNLY2022 strain shared the highest nucleotide identity of 99.7% with the C/HVRI vaccine strain (C-strain/Harbin Veterinary Research Institute, GenBank No. AY805221), but lower identities 94.5%–95.3% with the reference genotype 1.1b and 1.1c strains, and 84.2%–87.9% with the reference genotype 2.1 and 3.2 strains (Figures 2(a) and 2(b)). Moreover, compared to the C/HVRI strain, a total of eight amino acid mutations were found in the CSFV/HeNLY2022 isolate. As shown in Figure 2(c), the substitutions are distributed in E3 (H436S and S476R), E1 (T502I and P581I),
FIGURE 2: Homological analysis of the CSFV isolates. (a) The genomic homology analysis between the isolate CSFV/HeNLY2022 and reference strains. The red rectangular box showed the percent identity of CSFV/HeNLY2022 with live-attenuated strains. (b) Sequence similarity was compared between CSFV/HeNLY2022 and representative CSFV strains. Sequence similarity was performed using the live-attenuated vaccine C/HVRI strain as the query. (c) Amino acid mutation analysis between C/HVRI strain and CSFV/HeNLY2022 isolate.
E2 (M979thK and A1061thS), NS5A (A2980thT), and NS5B (I3818thM). Together, these data suggest that CSFV live vaccines, particularly the cell line Riems, C-strain cell line origin, and Ingelvac® CSF MLV, were further developed to improve the safety and efficacy of protection. These live vaccines played a critical role in the control and eradication of global CSF. Generally, C-strain is genetically stable and safe for pigs of all ages [14], and extensive use data exist mainly for the C-strain vaccines. However, the application data (for example, the genetic variation after cell-adapted culture) of modified live vaccines, particularly the cell line origin, is relatively few.

Here, we observed a severe porcine reproductive disorder in a pig farm in China. Laboratory diagnosis (10 of the 12 clinical samples were CSFV positive) and genetic analysis showed that the pathogenic agent might be CSFV, and the strain sequence has high genomic homology (99.7%) with swine fever attenuated vaccine strain (C/HVRI, GenBank No. AF091507 and AF531433). To the best of our knowledge, this is the first report indicating that CSFV live vaccines (cell culture-adapted C-strain derivatives) may have the potential to induce pathogenicity in immunized pigs under specific conditions. Meanwhile, it is not possible to exclude the conclusion that other unidentified agents or factors could have resulted in the reproductive issues observed (the possibility of this scenario is considered low, since most of the clinical samples were CSFV positive), until the virus has been isolated and clinical disease reproduced by experimental infection.

One possible reason for the reproductive failures could be the intensive nature and timings of the immunization. The pregnant sows had been administered two doses of the JEV live-attenuated vaccine (17 and 5 days ago) before immunized simultaneously with PRV and CSFV live-attenuated vaccines. The attenuated JEV and PRV vaccine strain could change the immune status of pregnant sows, which might make them more susceptible to CSFV. On the other hand, the amino acid substitutions in the polyprotein region also perhaps led to the virulence change. In the future, it is necessary to further explore whether these mutations are associated with the altered pathogenicity of the C-strain vaccine by virus isolation, reverse genetics systems and pig challenge studies. Another disadvantage of the traditional CSFV live vaccines is incompatible with a serological differentiation of infected from vaccinated animals (DIVA), which limits the CSF eradication in many countries [5]. Thus, many efforts have been put into develop novel effective DIVA vaccines. Recently, E2 subunit vaccines have been developed and authorized in Europe and China, which would be very useful to promote the prevention and eradication of CSFV [5, 15, 16].

In conclusion, our study suggests that the live-attenuated CSFV vaccine will most likely cause clinical diseases in sows and raises concern that the unscientific application of CSFV live vaccines in pig immunization procedures could potentially spread CSFV in pig farms. It is needed to perform a more rigorous evaluation of the safety of the C-strain-derived vaccines in combination with different live-attenuated vaccines. Further monitoring of combined vaccination application in pigs is also urgently needed.

Data Availability
All data generated or analyzed during this study are included in this published article and its supplementary information files.

Ethical Approval
All sampling and publication of the data were approved by the farm owners. This study did not involve endangered or protected species. All methods were carried out in accordance with relevant guidelines and regulations.

Disclosure
The funders had no role in study design, data collection, and interpretation or the decision to submit the work for publication.

Conflicts of Interest
All authors declare that they have no potential conflicts of interest.

Authors’ Contributions
Zhenhua Guo has contributed to the conceptualization, data collection and analysis, and writing—original draft preparation. Guangxu Xing has contributed to the data analysis, writing—reviewing and editing. Leyi Wang, Qianyue Jin, and Qingxia Lu have contributed to the writing—reviewing and editing. All authors read and approved the final manuscript. Zhenhua Guo and Guangxu Xing contributed equally to this work.

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Supplementary Materials
Table S1: primers used for genome sequencing. Table S2: the reference sequence information in this study. (Supplementary Materials)

References


