A Review of Pseudorabies Virus Variants: Genomics, Vaccination, Transmission, and Zoonotic Potential

Zongyi Bo\textsuperscript{1,2*} and Xiangdong Li \textsuperscript{2, *}

\textsuperscript{1} Joint International Research Laboratory of Agriculture and Agri-Product Safety, The Ministry of Education of China, Yangzhou University, Yangzhou 225009, China; zybo@yzu.edu.cn

\textsuperscript{2} Jiangsu Co-Innovation Center for the Prevention and Control of Animal Infectious Disease and Zoonoses, College of Veterinary Medicine, Yangzhou University, Yangzhou 225009, China

* Correspondence: 007352@yzu.edu.cn

Abstract: Pseudorabies virus (PRV), the causative agent of Aujeszky’s disease, has a broad host range including most mammals and avian species. In 2011, a PRV variant emerged in many Bartha K61-vaccinated pig herds in China and has attracted more and more attention due to its serious threat to domestic and wild animals, and even human beings. The PRV variant has been spreading in China for more than 10 years, and considerable research progress about its molecular biology, pathogenesis, transmission, and host–virus interactions has been made. This review is mainly organized into four sections including outbreak and genomic evolution characteristics of PRV variants, progresses of PRV variant vaccine development, the pathogenicity and transmission of PRV variants among different species of animals, and the zoonotic potential of PRV variants. Considering PRV has caused a huge economic loss of animals and is a potential threat to public health, it is necessary to extensively explore the mechanisms involved in its replication, pathogenesis, and transmission in order to ultimately eradicate it in China.

Keywords: pseudorabies virus; variant strain; genomics; vaccination; transmission; zoonosis

1. Introduction

Pseudorabies Virus, also called Aujeszky’s disease virus or Suid alphaherpesvirus 1, belongs to genus Varicellovirus, Alphaherpesvirinae subfamily within the family Herpesviridae, and is a double-strand linear DNA with 143 kb and can encode more than 70 proteins \cite{1}. PRV was first documented as the causative pathogen of Aujeszky’s disease in 1902 by a Hungarian veterinarian \cite{2}. Different clinical symptoms of pigs can be induced dependent on their housing stages: infected piglets always show fatal and central nervous system disorders, fattening pigs have respiratory symptoms with low mortality, and pregnant sows have abortions with the death of the fetuses \cite{3}.

The first case report of PRV in China could be traced back to 1947 in a domestic cat \cite{4}. After that, PRV circulated in many swine herds of China due to the lack of an available PRV vaccine. The Bartha vaccine strain, attenuated through many passages of a virulent strain in culture cells and embryos, was introduced into China in the 1970s, and it provided an ideal protection effect to the early prevalent strains (classical strains) in China \cite{5–7}. In late 2011, PRV occurred in many Bartha K61-vaccinated swine herds \cite{8}. After viral isolation and genome sequencing, the results showed that the newly emerging PRV clustered in an independent branch from the previously isolated strains in China \cite{9,10}. To differentiate it from the classical strains isolated before 2011 in China, the newly isolated PRV was designated as a PRV variant.

Though pigs are the only reservoir of PRV, it can infect many species of animals, including sheep, dogs, foxes, tigers, bears, etc. \cite{11,12}. Previously, experimental studies in nonhuman primates showed that rhesus monkeys and marmosets are susceptible to PRV infection, while other higher-order primates, such as chimpanzees and humans are not
susceptible to PRV infection [13]. Most recently, more than 20 human severe pseudorabies encephalitis cases were reported [14]. All these patients infected with PRV variants had close contact with pigs which suggested pigs might be the etiological source of PRV for human infection. So, it is time to pay attention to the public threat induced by PRV variants and the transmission of it between different species of animals and human beings.

In this review, we will briefly summarize the studies of PRV variants from the aspects of its outbreak, genomic characteristics, vaccine development, transmission in different animals, and its significance for public health.

2. The Outbreaks and the Genomic Evolution of PRV Variants

In late 2011, a large-scale occurrence of severe disease with anorexia, neurologic symptoms, high fever, and respiratory distress in piglets, and a high percent of abortion in sows happened in many swine farms of China. Pathological examination showed gross lesions in the lungs, and yellow-white necrosis in the kidneys. Through multiple kinds of diagnostic methods, including ELISA, PCR, viral isolation, immunohistochemical staining, and gene sequencing, the PRV variant was finally recognized as the causative pathogen for these severe clinical diseases [8,15].

After viral isolation and sequencing, there is a large sequence divergence in newly isolated PRV strains from previously classical PRV strains. Here, we constructed a maximum likelihood (ML) phylogenetic tree of 39 strains (Supplemental Table S1) of PRV full-length sequences which have been extracted from NCBI. The results showed that all PRV strains can be phylogenetically clustered into two groups. All foreign strains out of China were clustered into the same group, and nearly all strains isolated from China are located in an independent group with them (Figure 1A). Interestingly, the PRV GD1802 strain, isolated in China in 2018, is clustered into the same group with the strains from foreign countries. We infer that the GD1802 strain might be delivered into China from foreign countries through the introduction of pigs. In addition, when compared with classical strains, such as Ea and Fa, PRV strains isolated after 2011 are located in a relatively dependent branch with them. The gC gene is a major gene which has been commonly used for the phylogenetic analysis of PRV due to its high variability [16,17]. We also analyzed gC gene sequences of these reported PRV strains. As shown in Figure 1B, the gC-based phylogenetic tree reveals that PRV can be phylogenetically divided into two groups, designated as genotype I and genotype II, and genotype II can be further divided into two clades, clade 2.1 and clade 2.2. Clade 2.1 mainly comprises the strains isolated before 2011, and nearly all PRV variants are located at clade 2.2.

Recombination contributes a lot to the genomic divergence of many viruses, such as African swine fever virus [18], porcine reproductive and respiratory syndrome virus [19], classical swine fever virus [20], porcine circovirus [21], and porcine epidemic diarrhea virus [22]. As for PRV, the recombination between different strains has been reported in vivo and vitro [23,24]. So, it is interesting to explore whether the evolution of PRV in China also has a relationship with recombination. As shown in Figure 1B, we found that there are four Chinese-origin strains located at the genotype I cluster, including SC, HLJ-2013, JSY13, and GD1802 strains. Except for GD1802, which might be introduced abroad, the other three strains all have a relationship with recombination. Ye et al. found that SC was a recombinant of an endemic Chinese strain and a Bartha-vaccine-like strain [25]; Bo et al. found that the JSY13 strain was a recombination of the PRV variant JSY7 strain and the Bartha K61 vaccine strain [26]; Liu et al. found that the HLJ-2013 strain is probably a recombination of three origins: a yet unknown parent strain, a European-origin strain, and a Chinese-origin strain [27]. Besides these, Huang et al. found that the Fj62 variant strain was the recombination between the PRV genotype I strain from wild boar and genotype II strain from domestic pig [28]. These reports demonstrated that recombination plays an important role in the evolution of PRV in China.
It was reported that the virulence of PRV variants were higher than the classical strains which were isolated before 2011. So, whether there is a difference in the major virulence-determining genes between PRV variants and classical strains warrants investigation. Therefore, we analyzed gI and gE genes, which are major virulence-determining genes of PRV [29]. The phylogenetic analysis showed that all PRV strains can be divided into two classes, as shown in Figure 2A (gI) and Figure 2B (gE). Compared with PRV from foreign countries or the classical strains isolated in China, there are some typical mutations, insertions, and deletions in the several virulence-determining genes, and non-coding sequences of PRV variants, such as two Aspartate (Asp, D) insertions in gE protein [26,30,31].

Figure 1. The phylogenetic analysis of PRV full-length genome sequences and gC sequences. (A) Phylogenetic analysis of PRV complete genome sequences. (B) Phylogenetic analysis of PRV gC gene sequences. Both maximum likelihood (ML) trees were constructed by using MEGA X software.
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Figure 2. The phylogenetic analysis of PRV gI and gE genes. (A) Phylogenetic analysis of PRV gI gene sequences. (B) Phylogenetic analysis of PRV gE gene sequences. Both maximum likelihood (ML) trees were constructed by using MEGA X software.

The biological functions and meanings of these insertions or deletions in PRV variant strains are yet to be explored. Previously, a study showed that the exchange of gB genes contributes to an immunogenic difference between PRV variant JS-2012 and Bartha-K61, which indicated the gB protein of the PRV variant has the ability to evade the neutralization antibody induced by the Bartha strain [32]. The gC gene, a main host receptor binding protein of PRV, can bind to heparan sulfate proteoglycans in the extracellular matrix. Whether the mutations in the gC gene influence the cell entry stage of PRV and viral transmission among different kinds of tissues, animals, and even human beings is rarely studied. As the PRV variant is more virulent than classical PRV strains, whether the mutations in TK, gI, and gE virulence-determining genes are related to the increased virulence of PRV variants also needs to be addressed in the further studies.

3. PRV Variant-Based Vaccines Are on the Way

As the PRV variant has been circulating in many Bartha K61-immunized pig farms, researchers tried to explore whether the Bartha K61 strain could provide full protection against PRV variants. An et al. used the PRV variant HeN1 strain to challenge the Bartha K61-immunized pigs and sheep, the results showed that HeN1-challenged pigs showed fever and loss of appetite, while no deaths were found; the experiment on sheep showed that the Bartha K61 strain provided full protection against PRV classical SC strain, but
two out of four Bartha K61-immunized sheep challenged with the PRV variant HeN1 strain had clinical signs and died [15]. Luo et al. challenged the Bartha K61-immunized sheep with the PRV variant TJ strain and found that the Bartha K61 vaccine cannot provide complete protection [33]. These experiments demonstrated that the Bartha K61 vaccine can hardly provide full protection against the PRV variant.

Besides the imported Bartha K61 vaccine strain, there are several other licensed PRV live attenuated and inactivated vaccine strains that are based on the PRV strains isolated in China. One of them is the SA125 strain, which is a gE/gI/TK gene-deleted strain based on the cattle-origin Fa strain, and it is the first licensed PRV vaccine in China [34]. Another is the PRV gE/gl/TK gene-deleted HB-2000 strain, which is based on the Ea strain. Besides these two live attenuated vaccines, there is another licensed PRV HB-98 inactivated vaccine which deletes TK/gG genes based on the Ea strain. The existence of the gE gene in the HB-98 vaccine makes it impossible to differentiate the immunized from field strain-infected pigs [35]. Now, there is no direct evidence whether these licensed Chinese-origin PRV strains have better protection than the Bartha K61 vaccine strain to the PRV variant.

The PRV variant has been circulating in China for more than 10 years, and many vaccine candidates that are based on PRV variants have been reported by different research groups. Wang et al. deleted the gE gene of a PRV variant TJ strain and used it to immunize pigs; the pigs had no clinical signs when they were challenged with a PRV variant strain [36]. Wang et al. constructed a TK and gE gene-deleted PRV variant, AH02LA, which could stop the viral shedding, and no clinical signs were observed after being challenged with the PRV variant. By contrast, Bartha K61-immunized pigs showed some mild clinical signs and had viral shedding [37]. Wang et al. used the bacterial artificial chromosome (BCA) manipulation method to construct a PRV variant AH02LA gE gene-deleted strain and found it can provide complete clinical protection against the challenge of the PRV variant [38]. Besides the above-mentioned experimental vaccines, several other PRV variant-based vaccine candidates which have double or triple deletions of TK, gE, or gI genes, also showed complete protection against the challenge of PRV variants [39–41]. Till now, there are two licensed PRV variant-based vaccines. One is the PRV C strain, a gI/gE/Us9/Us2 naturally deleted strain that was isolated in China in 2011 and licensed in 2017 [42]. Another is a gE-deleted inactivated vaccine (HN1201-∆gE), which was certificated in 2019 and could induce a high neutralization antibody titer against both PRV classical and variant strains.

Despite the inactivated and live attenuated vaccines, several other types of vaccines including a subunit vaccine and nucleic vaccine, could also be the alternative for the development of an effective PRV variant vaccine. Among 11 glycoproteins PRV encoded, gB, gC, and gD are the main immunogenetic antigens that can induce neutralization antibodies [7,43]. Previously, Wang et al. used the baculovirus system to express the PRV variant gB protein and challenged the immunized pigs with PRV variant HN1201 strain. The results showed that the gB-based vaccine can provide full protection for the PRV variant [44]. In another report, gB, gC, and gD proteins of the PRV variant were separately expressed using the baculovirus expression system working as subunit vaccine candidates. Pigs were then immunized with each single protein twice and challenged with the PRV variant HNLH strain. The results showed that the survival rates of gB-, gC-, and gD-vaccinated pigs were 100%, 50%, and 87.5%, respectively [45].

A DNA vaccine can mimic the natural infection in which the immunized antigens could be presented in both major histocompatibility complex classes I and II settings [46]. Previously, E.M.A. van Rooij et al. vaccinated pigs with the plasmids expressing the main immunogenetic antigens gB, gC, and gD, then challenged the immunized pigs with PRV. The results showed that plasmid DNA encoding gB induced the strongest cell-mediated immune responses, whereas plasmid DNA encoding gD induced the strongest neutralizing antibody responses [47]. Furthermore, Hyun A Yoon et al. compared the intramuscular (i.m.) and intranasal (i.n.) immunization routes by immunizing mice with the plasmid that encoded gB. The results showed that immunization can induce the mucosal immunity via
the i.n. route. However, it only induced a low IgG response which cannot protect the mice against the challenge of PRV [48].

A messenger RNA (mRNA)-based vaccine that encodes the immunogenetic antigen of pathogens provides a new vaccine developing platform for multiple viruses [49,50]. As PRV can encode several immunogenetic antigens that can induce the neutralization antibodies, mRNA might be a candidate for PRV vaccine development. Jiang et al. developed a PRV-XJ variant strain gD gene-based mRNA vaccine which was formulated via mRNA encapsulated in the liposomes. Compared with the DNA vaccine that encoded PRV gD protein in a recombinant plasmid pVAX-gD, both mRNA and pVAX-gD plasmids induced a high neutralization antibody titer and antigen-specific B and T cell response in mice. Finally, the PRV variant challenge experiment was performed, and the results showed that one mouse in ten which was immunized with the mRNA vaccine died, whereas there was no death in the pVAX-gD plasmid-vaccinated group [51]. These results demonstrated that multiple types of vaccines besides inactivated and live attenuated vaccines, show protection against the challenge of PRV, and gB and gD proteins play an important role among these major immunogenetic proteins.

Despite many reports showing that the Bartha K61 vaccine cannot provide full protection against PRV variants, there are still several studies showing that Bartha K61 can provide complete protection to pigs against the PRV variant [7]. Previously, Zhou et al. used and then challenged Bartha K61-immunized growing pigs with the PRV variant XJ5 strain. The results showed that all immunized/challenged pigs survived, while the unimmunized pigs all died [52]. Another report also showed that Bartha K61-immunized pigs had no death under the challenge of the PRV variant AH02LA strain [53]. Although there were some mild clinical signs in immunized/challenged growing pigs, no death was found in the above two studies, which demonstrates that the Bartha K61 vaccine strain can provide protection for pigs against the PRV variant challenge. However, as the PRV variant could lead to the deaths of newborn piglets, further experiments still need to be conducted to confirm whether piglets are resistant to the challenge of the PRV variant because they are Bartha K61-immunized or because they got maternal antibodies from the immunized sows.

4. The Pathogenicity of PRV Variant to Different Species of Vertebrates

Although the natural reservoir for PRV is the pig, the first case of PRV was reported in cattle, and the disease was described as “mad itch” [54]. PRV has a broad host range, infecting most mammals and even some avian species. PRV infection can cause different clinical signs dependent on the stages of housing pigs. Besides pigs, PRV infection in other species of animals is always fatal due to the neurologic invasion. Till now, it has been reported that there are about 19 species of animals that can be naturally infected by PRV including domestic/wild pigs [55,56], sheep [57], cats [58], coyotes [59], foxes [60], rats [61], deer [62], bears [63], rabbits [64], dogs [65], horses [66], bats [67], wolves [68], raccoons [69], mice [70], ferrets [71], panthers [72], cattle [73], and chickens [74] (Figure 3). Domestic and wild pigs are the reservoirs for PRV, all other species of animals will die after several days upon the onset of PRV. In the last decade, several animal species including dogs, bovine, mink, foxes, goats, and wolves were found dead due to the infection of PRV variants [75–79]. It is reasonable to infer that some other animal species could be susceptible to PRV variants. The wide host range of PRV can also be demonstrated by its incubation in multiple kinds of cells, such as PK15 cells (Porcine kidney), Vero cells (Green monkey kidney cells), HEp-2 (Human Epithelioma cells), DF-1 cells (Chicken embryo fibroblasts cells), and many other cells. This demonstrates that PRV can bind with its receptor in the surface of these cells, which suggests that PRV might have the possibility to infect multiple kinds of animals.
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Figure 3. The reported animals that can be infected by PRV.

Besides natural infection, the experimental infections of PRV were also conducted to confirm the susceptibility of different animals, such as domestic pigs [80], wild boars [81], sheep [82], calves [73], dogs [83], cats [84], blue foxes [85], raccoons [86], horses [66], and chickens [74]. The clinical signs after infection mainly include fever, depression, pruritus, self-mutilation, anorexia, neurologic deficits, etc. The viral challenge experiments demonstrated that PRV can be introduced through multiple kinds of routes, including intracranial, intradermal, intramuscular, intranasal, intraocular, intraperitoneal, intratracheal, intravenous, and even foot pads [87]. The animal experiments of PRV can also be performed in some laboratory animals, including mice [88], rats [89], rabbits [90], dogs [90], guinea pigs [91], and rhesus macaques [92]. Due to the wide host range and easy manipulation, PRV can be handled as the model to study many characteristics of herpesvirus, such as molecular biology, pathogenesis, neuroinvasion, and transneuronal spread.

Sometimes the transmission of PRV among different species of animals will lead to the change of PRV pathogenicity. Previously, R E Shope found that PRV was attenuated after passaging through a guinea pig brain [91]. In addition, T F Müller et al. found that PRV strains of wild swine origin might be less virulent than those of domestic pigs [81]. This phenomenon might be explained by the insertions or deletions of multiple segments among different strains, in order to adapt to their host animals.

PRV is an air-borne pathogen. Furthermore, food, water, and excrement can also be intermediate vectors of PRV [93]. The infected pigs or semen are the main origins for PRV infection in swine farms. As for other species of animals, the direct contact of contaminated equipment, or consumption of the infected animals are the main sources for their infection [94]. Thereafter, it is important to keep the animals away from swine farms, especially mice, cats, dogs, etc., in case they are infected with PRV and transmit it to other animals. Furthermore, pig products or byproducts, such as head or offal tissue, which might be polluted by PRV must be kept away from other animals, since they could directly
contact or consume them. Finally, surveillance of PRV not only in pigs, but also in other species of animals should be performed, in order to keep PRV out of the susceptible herds.

5. The Zoonotic Potential of PRV Variants

It has been known that PRV has a wide host range, and the duplication and translocation of sequences from the left end of the genome to the UL-US junction plays an important role for the growth of PRV in different hosts [95]. It was reported that Nectin-1, a host cellular receptor that can bind with PRV glycoprotein gD, plays an important role in PRV entry. Previous studies reported that PRV gD glycoprotein showed a similar binding ability to human nectin-1 protein [96]. Not only human nectin-1, the protein that comes from bats, dogs, cats, cows, sheep, and several other animals also showed conserved functional amino acids binding with PRV gD, which provides evidence of cross-species infection for PRV [14].

The first case report of humans infected with PRV was in 1914, in which two laboratory technicians were infected with PRV after they were exposed to an infected cat. Afterwards, several PRV cases in humans were reported after they came into contact with cast, dogs, cows, or other domestic animals (Summarized in Table 1). The typical symptoms of these cases mainly include pruritis, pain, fever, swelling, sweating, dysphagia, and aphthous stomatitis. Though these PRV infected cases were reported, the etiological confirmation was not convincing due to the lack of a definitely etiological and serological diagnosis.

In 2017, a swine herder suffered from an eye disease in China. Through next-generation sequencing (NGS), real-time PCR, and phylogenetic analysis, the PRV variant was recognized as the causative agent of the disease [97]. This is the first PRV case in humans reported in China. In the following years, several other PRV infection cases in human beings were reported in China (Table 1). Different from the clinical symptoms reported from other countries, the clinical signs of Chinese patients in these cases were mainly encephalitis and endophthalmitis. Of note, almost all PRV infected patients have contact with swine or other PRV-susceptible animals. Among these incidences, a milestone case attracted our attention because one PRV variant, designated hSD-1/2019, was firstly isolated from the cerebrospinal fluid of a PRV-infected patient [98]. This is the first case in the world where PRV was successfully isolated from humans, which proved that humans are susceptible to PRV.

Table 1. PRV infection cases in humans.

| Year | Numbers | Symptoms                                                                 | Contacted Animals          | Ref.  |
|------|---------|---------------------------------------------------------------------------|----------------------------|-------|
| 1914 | 2       | Swelling, reddening, intense itching                                      | Cat                        | [99]  |
| 1940 | 2       | Pruritis, erythema, and pain around the wound                             | Dog                        | [99]  |
| 1963 | 2       | Throat pain, weakness in the legs,                                        | Dog                        | [99]  |
| 1983 | 1       | Pain in tongue, hypersalivation, dysphagia,                              | Cat                        | [100] |
| 1986 | 2       | Tension in the mouth, nose, and throat; perception of strange smells and taste | Cat, or other domestic animals | [100] |
| 1992 | 6       | Pruritus in the palms, lower and upper arms, shoulders, and back          | Cow                        | [101] |
| 2017 | 1       | Endophthalmitis, fever, headaches                                          | Pig                        | [97]  |
| 2018 | 4       | Encephalitis                                                              | Pig                        | [102] |
| 2019 | 5       | Encephalitis                                                              | Pig                        | [103] |
| 2019 | 1       | Encephalitis                                                              | Pig                        | [104] |
| 2019 | 1       | Encephalitis, fever, headache, seizure                                    | unknown                    | [105] |
| 2019 | 1       | Encephalitis                                                              | Pig                        | [106] |
| 2020 | 1       | Encephalitis                                                              | Pig                        | [107] |
| 2020 | 6       | panencephalitis                                                           | Pig                        | [108] |
| 2021 | 1       | Retinitis                                                                 | Pig                        | [109] |
| 2021 | 1       | Encephalitis, retinal vasculitis, fever                                   | Pig                        | [110] |
| 2021 | 1       | Encephalitis, Endophthalmitis                                             | Pig                        | [111] |
| 2022 | 2       | Encephalitis, seizures, endophthalmitis                                   | Pig                        | [112] |
Of note, the cases of humans infected with PRV have been increasing rapidly in China since 2017. Coincidentally, PRV variants have been the dominant strain in Chinese pig farms since then. Whether PRV variants are more sensitive to humans than the classical PRV warrants further investigation. Among these PRV-infected patients, most of them were reported as having swine-related occupations or had close contact with other infected animals. Furthermore, several patients have injuries to their fingers or other places. Therefore, it is necessary to carry out skin protection for people who have close contact with swine. Till now, all infected humans recovered completely, and the clinical and neurological signs disappeared, though sometimes the clinical signs last for one to several months. No patients died from PRV infection in the above cases. However, similar to other herpesviruses, PRV can induce latent and lytic replication in pigs, and the infected pigs will carry PRV for a long time. Whether the recovered patients still carry PRV is unknown.

6. Discussion and Perspectives

In this review, we summarized the biological characteristics of PRV variants, which have antigenic variation and a higher virulence compared with classical PRV strains in China. Since 2011, PRV variants have become the dominant strains in China and caused huge economic losses for the swine industry. There is still dispute over whether the Bartha K61 vaccine can provide full protection against the PRV variant [52,113]. However, PRV variants happened in many Bartha K61-immunized swine farms, and the phylogenetic analysis showed that PRV variants have a large sequence divergence with the Bartha K61 strain. Therefore, more effective vaccines based on local PRV variants need to be developed. Moreover, multiple kinds of vaccines, including a live attenuated vaccine, inactivated vaccine, subunit vaccine, DNA vaccine, and mRNA vaccine have the potential to provide full protection for pigs against PRV variants. Upon the design of PRV vaccines, it is necessary to follow the strategy of DIVA, which has helped to eradicate PRV in many countries [114].

The serological detection of PRV gE antibodies is always used to evaluate the status of PRV field strain infection, due to the application of gE-deleted PRV vaccines. One study reported that PRV positive rates have decreased to 18.12% in 2020 from 38.20% in 2018 after screening 19,292 pig serum samples by using PRV gE ELISA [115]. Another report analyzed the gE antibodies from 256,326 serum samples, which were collected from 29 provinces in China from 2011 to 2021. The results showed that the average positive rate was 29.87% [31]. Zheng et al. collected 4708 pig serum samples from Henan province during 2018–2019 and found the positive rate of gE antibodies was 30.14% (1419/4708) [116]. Lin et al. collected a total of 18,138 serum samples from 808 PRV-vaccinated pig farms during 2016–2020 in Hunan province and detected the presence of gE-specific antibodies. The results showed that 23.55% (4271/18,138) of the samples were positive for PRV gE-specific antibodies [117]. All these results demonstrate that the infection of the PRV field strain is still very common in Chinese swine farms, though nearly all of them have used PRV vaccines. Therefore, strict biosecurity measures, feeding strategies, daily managements, diagnostic methods, and effective vaccination should be jointly performed in swine farms to decrease the infection of the PRV field strain.

Right now, with the increased number of human cases caused by PRV infection, the exact mechanisms involved in its transmission from animals to human beings are still unknown. One explanation is that the “one health” concept is recognized by more and more researchers, and the interdisciplinary communication is more frequent than before, so the human cases with encephalitis and neurological symptoms, which might be induced by PRV, could be jointly diagnosed by doctors and veterinarians. Another possibility is that the evolution of the PRV variant makes it more susceptible to humans as PRV entry-related proteins gD or gC might have a higher binding ability to its receptor in human cells than before. In contrast, a study showed that 455 persons who participated in heroic self-incubation with PRV via intracutaneous and subcutaneous methods showed no clinical signs, which demonstrated that the infection of PRV in humans was occasionally asymptomatic [99]. We
hypothesize that the infection of PRV in humans might be related to the immune system of people, and immunocompromised patients might be more susceptible to PRV. Till now, the only consolatory thing is that there is no human-to-human PRV transmission case. More studies about the mechanisms evolved in the transmission of PRV from infected animals to humans still need to be explored.

Due to the lack of effective PRV variant vaccines, many researchers have focused on exploring the available compounds that can inhibit the proliferation of PRV in vitro and in vivo. Among them, two kinds of compounds attract people’s attention. One is traditional Chinese herbal medicines, which have a long history of anti-virus effects, but their cellular targets and working mechanisms are unclear. For example, Germacrone, which is extracted from Rhizoma Curcuma, was able to inhibit the proliferation of PRV in the early phase of the PRV replication cycle [118]. Another kind is synthetic chemical compounds, which have a relatively clear cellular target and whose functions are comparatively clear. For example, it was demonstrated that meclizine, a class of H1-antihistamine, could inhibit the replication of PRV at its entry and release stages [119]. The exploration of anti-PRV compounds will contribute to the therapy of PRV infection, especially for the infection of human beings.

Because of the existence of multiple non-essential genes, PRV can be engineered as the vector to express foreign antigens derived from other pathogens. For example, Qiu et al. used PRV as the vector to express the GP5 protein of porcine reproductive and respiratory syndrome virus, and the recombinant virus provided an ideal protection against the challenge of PRRSV [120]. Other proteins, such as the HA gene of the H3N2 subtype of swine influenza virus, and the S protein of porcine epidemic diarrhea virus, were also expressed by PRV, which provided protection to the immunized pigs [121,122]. The advantage of these vectored vaccines are that they provide protection against the infections of other pathogens besides PRV.

In conclusion, as the PRV variant has caused a huge economic loss for the pig industry in China and is a potential threat to public health, it is necessary to pay more attention to the detection and isolation of the PRV variant in different animals, in order to study its epidemiological characteristics. At the same time, it is urgent to develop more safe and efficient DIVA vaccines based on the PRV variants. Furthermore, it is also important to deeply explore the interactions between the PRV infection and host responses, which will not only help to clarify the mechanisms involved in PRV proliferation and pathogenesis, but also contribute to the development of PRV vaccines and antiviral drugs. Most importantly, since PRV has the possibility to infect human beings, researchers need to pay more attention to its cross-species transmission and screen the patients that are suspected of being infected by PRV, especially those who have swine-related occupations.

Supplementary Materials: The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/v14051003/s1, Table S1: The genomic information of 39 strains of PRV.

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