Protective efficacy of a broadly cross-reactive swine influenza DNA vaccine encoding M2e, cytotoxic T lymphocyte epitope and consensus H3 hemagglutinin

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Abstract

Background: Pigs have been implicated as mixing reservoir for the generation of new pandemic influenza strains, control of swine influenza has both veterinary and public health significance. Unlike human influenza vaccines, strains used for commercially available swine influenza vaccines are not regularly replaced, making the vaccines provide limited protection against antigenically diverse viruses. It is therefore necessary to develop broadly protective swine influenza vaccines that are efficacious to both homologous and heterologous virus infections. In this study, two forms of DNA vaccines were constructed, one was made by fusing M2e to consensus H3HA (MHa), which represents the majority of the HA sequences of H3N2 swine influenza viruses. Another was made by fusing M2e and a conserved CTL epitope (NP147-155) to consensus H3HA (MNHa). Their protective efficacies against homologous and heterologous challenges were tested.

Results: BALB/c mice were immunized twice by particle-mediated epidermal delivery (gene gun) with the two DNA vaccines. It was shown that the two vaccines elicited substantial antibody responses, and MNHa induced more significant T cell-mediated immune response than MHa did. Then two H3N2 strains representative of different evolutional and antigenic clusters were used to challenge the vaccine-immunized mice (homosubtypic challenge). Results indicated that both of the DNA vaccines prevented homosubtypic virus infections completely. The vaccines’ heterologous protective efficacies were further tested by challenging with a H1N1 swine influenza virus and a reassortant 2009 pandemic strain. It was found that MNHa reduced the lung viral titers significantly in both challenge groups, histopathological observation showed obvious reduction of lung pathogenesis as compared to MHa and control groups.

Conclusions: The combined utility of the consensus HA and the conserved M2e and CTL epitope can confer complete and partial protection against homologous and heterologous challenges, respectively, in mouse model. This may provide a basis for the development of universal swine influenza vaccines.

Keywords: Swine influenza, DNA vaccine, Consensus HA, M2e, CTL epitope

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Background
Swine influenza virus (SIV), member of genus Influenza A virus of the family Orthomyxoviridae, is a common and important causative pathogen involved in the porcine respiratory disease. Mortality of SIV-infected pigs is low, but morbidity may approach 100%. Clinical signs of swine influenza include high fever, lethargy, anorexia, coughing, labored breathing and nasal discharge. Synergistic or secondary infections with opportunistic organisms may increase the severity of clinical disease [1]. Beyond the veterinary implications, influenza virus infections in pigs also present an important public health risk. Since pigs express sialic acid receptors for both mammalian and avian strains of influenza viruses on their tracheal epithelial cells [2], they could potentially serve as “mixing vessels” for the generation of new reassortant strains of influenza viruses that have pandemic capacity. The recently emerged 2009 pandemic H1N1 which resulted in over 18,449 deaths is an example (http://www.who.int/csr/don/2010_08_06/en/index.html). Genetic analyses revealed that this pandemic H1N1 influenza virus is a triple reassortant of multiple strains of viruses circulating in the North America and Eurasia swine population [3,4]. Therefore, SIV-infection control would be of benefit to both reduce the economic losses of swine industry and human health.

Vaccination is considered to be the most effective method to control SIVs. Currently, commercially available swine influenza vaccines are adjuvanted, whole-virus killed vaccines. Although the vaccines reduce the severity of disease and the extent of virus shedding in pigs after challenge, they do not provide consistent protection from infection [5]. The vaccines function by targeting the surface glycoprotein hemagglutinin (HA), the most variable influenza virus antigen. Protective efficacy depends on the antigenic match degree between vaccine and circulating strains. However, influenza viruses continuously evolve by increasing the mutations in epitopes (antigenic drift) or by reconstituting the genome with other strains (antigenic shift). The increased incidence of avian-like or human-like SIV reassortants, which exhibit with other strains (antigenic shift). The increased incidence of avian-like or human-like SIV reassortants, which exhibit with other strains (antigenic shift). The increased incidence of avian-like or human-like SIV reassortants, which exhibit with other strains (antigenic shift). The increased incidence of avian-like or human-like SIV reassortants, which exhibit with other strains (antigenic shift).
HI assays were then performed on both homosubtypic and heterosubtypic virus to assess the ability of inducing relevant and cross-reactive functional antibodies. As detailed in Table 1, both of the vaccine immunization groups developed HI antibodies against homologous viruses, but induced very low levels of HI antibodies against heterologous viruses.

NP147-155 enhanced CTL responses significantly
The ability of the CLT epitope-containing vaccine, MNHa, to induce cellular immune response was determined by IFN-γ ELISPOT assay. A synthesized peptide NP147-155 was used for restimulating splenocytes isolated from 3 mice of each group. Results showed that, after restimulation, the number of activated IFN-γ secreting cells from mice immunized with MNHa was significantly higher than that from mice immunized with MHa and empty vector \((p < 0.001, \text{Figure 3})\).

Both MHa and MNHa vaccination conferred complete protection against homologous challenge, but MNHa showed more significant cross-protection against heterologous challenge than MHa did
Our previous study indicated that the wild type of SwGD164 strain (H1N1) replicates poorly in mice (unpublished data). To make the virus be suitable for vaccine efficacy assessment \textit{in vivo}, we constructed a recombinant virus (r164) by using reverse genetics. r164 contains HA and NA from SwGD164, and the internal proteins from PR8. BALB/c mice were infected intranasally with two homologous strains belonging to different clusters within H3 subtype 21 days after the last immunization [9]. Mice lungs were taken 3 days post infection for virus titration and histopathologic changes observation. Results demonstrated that there was no detectable virus load in all the vaccine-immunized mice, while empty vector control group showed high lung viral titers (Figure 4). In addition, histopathologic observation showed no obvious histopathologic changes in vaccinated mice. In contrast, the empty vector control group exhibited histopathological damages including the dropout of mucous epithelium cells, interstitial edema, hyperemia, hemorrhage, and inflammatory cell infiltration (Figure 5).

Since M2e and NP147-155 are conserved in different subtypes of influenza viruses, we hypothesized that the DNA vaccine can confer partial protection against heterosubtypic challenge. To test this, the vaccines’ protective ability was studied in mice challenged with H1N1 SIVs, which are also circulating in swine population worldwide. The immunized mice were challenged with heterologous G11 and rPan09, respectively. Lung viral titers determination and pathological analysis were performed as described above. Results indicated that all the mice that had been vaccinated with MHa had a detectable virus level, although showed a reduction in mean viral titers in both challenge groups compared with vector control, the reduction did not reach significance \((p = 0.06 \text{ for rPan09 group and } p = 0.67 \text{ for G11 group, Figure 4})\). Histopathological analysis exhibited corresponding results, i.e., mice of MHa group had lung pathology as almost severe as vector inoculated group. In MNHa group, although the mice also exhibited detectable
virus shedding, the mean viral titers had been significantly reduced in both rPan09- and G11-challenged groups compared to vector-immunized control ($p < 0.001$), showing average viral titer reduction of 3.4 and 2.1 logs, respectively (Figure 4). As expected, pathology damage of mice lung of MNHa group was less pronounced than that of MHa and vector control group, moderate histopathological lesions, mainly vascular congestion and various degrees of hemorrhage were observed (Figure 5). Collectively, these findings indicate consensus HA in combination with M2e elicits cross-cluster immunity, and CTL response contributes to the control and clearance of infection and reduces pathogenesis.

**Discussion**

Although the viruses are highly and continuously variable because of antigenic drift and shift, influenza is a vaccine-preventable disease. The conventional inactivated human influenza vaccines are updated annually with the World Health Organization-recommended H1N1, H3N2, and influenza B strains in order to antigenically match the viruses predicted to be the most likely to cause the next epidemic. Unlike human influenza vaccines, strains used for swine influenza vaccines preparation are not regularly replaced, making the vaccines provide limited protection against antigenically diverse SIVs. A universal vaccine designed based on the conserved viral proteins that confers broad protection is an attractive solution to counter the features of highly variable of not only human but also swine influenza viruses. Many studies developed universal vaccines against human influenza, whereas broadly responsive swine influenza vaccines, to our knowledge, have not been reported.

The virus envelope glycoprotein HA is the most abundant surface protein, antibodies against HA can provide protection by blocking virus attachment and entry. Studies from several groups have demonstrated the antiviral efficacy of consensus HA as an "intra-subtype" universal vaccine. The study of Chen et al. [17] showed that the consensus H5 HA-based DNA vaccine elicited only moderate neutralization activities toward the H5N1 clade 2.1 and clade 2.3 viruses, and provided not complete, albeit significant, protection against clade 2.1 virus, suggesting that consensus HA alone may be not enough to induce complete protection against all strains within one subtype, much less heterosubtypic strains. However, ideally, universal vaccines should be capable of inducing protection against both homosubtypic and heterosubtypic challenges. To this end, we linked M2e and CTL-inducing epitope NP147-155 to consensus H3 HA to construct a universal swine influenza vaccine. The reasons we chose M2e and NP147-155 are: (1) both of them are highly conserved among different SIV strains, NP147-155 are even conserved among human, avian and swine strains.
influenza strains (Table 2), they were added in view of their ability to broaden the protective scope of the vaccine. (2) NP147-155 induces a vigorous CTL response [23]. Studies indicated that specific CTLs provide some level of cross-protection against antigenically distinct viruses of not only the same subtype but also different subtypes [24,25].

Previously we have shown that SwHLJ1 and SwGD164 had evolved into 2 independent lineages based on the phylogenetic analysis of the HA gene. To see whether the two strains were antigenically distinct, cross-HI assay was performed by using r164-specific anti-serum. It was found that the titer of cross-reacting HI antibodies (SwHLJ1-reacting) was significantly lower than that of r164-reacting HI antibodies ($p < 0.001$, data not shown), thus suggesting that they belong to different antigenic clusters. Results of present study demonstrated that both MHa and MNHa vaccination provided complete protection against these two viruses challenges, cross-cluster protections were therefore suggested. While MNHa, but not MHa, induces immune responses partially protect against heterologous influenza infection, suggesting the enhanced T-cell mediated response is not required for homologous protection. Results also further confirm that CTL responses play important roles for heterologous protection, meaning that strengthening CTL responses are promising ways for universal influenza vaccines development.

It was established that the 2009 pandemic H1N1 virus is pathogenic and is readily transmitted in pigs [26-29]. Although the new H1N1 virus is now considered to be post-pandemic, the possibility that it recombines with other influenza viruses in pigs then yielding a novel potential epidemic or pandemic strain still exists[30]. Here in this study, the mice were partially protected from the reassortant 2009 pandemic virus challenge after the immunization of MNHa. The vaccine immunization inhibited the pulmonary viral replication significantly in mice, thus could accelerate the virus clearance, and reduce the potential of transmission and the risk of recombination. Although the heterologous protection elicited

![Figure 5 Histopathology analysis of lungs from the mice inoculated with SwHLJ1 (A, B, C), r164 (D, E, F), rPan09 (G, H, I) and G11 (J, K, L) at day 3 post inoculation, after immunized with MHa (A, D, C, J), MNHa (B, E, H, K) and pCAGGS vector (C, F, I, L).](image)

| Challenge strain | % homology of amino acids |
|------------------|--------------------------|
|                  | HA  | M2e | NP147-155 |
| H1 subtype       |     |     |          |
| rPan09           | 44.3| 79.2| 100       |
| G11              | 44.0| 79.2| 100       |
| H3 subtype       |     |     |          |
| SwHLJ1           | 93.3| 91.7| 100       |
| r164             | 95.1| 79.2| 100       |
by the DNA vaccine was not complete, this methodology has a large potential for improvements. The present study lays the foundation for universal swine influenza vaccine development, and call for further investigations in which the heterologous immune response should be further enhanced, such as the addition of molecular adjuvants [31,32] and/or more copies of conserved viral protein encoding genes [33], and the usage of DNA-prime protein/virus-boost immunization schedule [34,35]. A previous study conducted by Heinen et al. constructed a DNA vaccine containing M2e and full length of NP genes, but HA encoding gene was not included. Their data showed that the M2e-NP DNA vaccination produced enhancement of disease after challenge [36]. Although the mechanism underlying which is unclear, we think the HA may play roles in this discrepancy between the results of Heinen et al.’s and ours. Another possible reason should be taken into consideration is that gene gun immunization induces more strong immune response that the conventional needle injection method, which was used in Heinen et al.’s study [37]. It is necessary to study whether the DNA vaccines we developed here will confer protection in pig models. Also, it would be interesting to study how, if does, the HA protein influence the vaccines’ efficacy.

Conclusions
The combined utility of the consensus HA and the conserved M2e and CTL epitope can confer complete and partial protection against homologous and heterologous challenges, respectively, in mouse models. This may provide a promising strategy for universal swine influenza vaccine development.

Materials and methods
Viruses, cell and mice
Swine influenza viruses of A/Swine/Heilongjing/1/05 (H3N2) [SwHLJ1], A/Swine/Guangdong/164/06 (H3N2) [SwGD164] and A/Swine/Guangdong/1/01(H1N1) [G11] were isolated previously by us [6]. Briefly, clinical samples from diseased pigs were homogenized in phosphate-buffered saline (PBS) containing antibiotics, then inoculated into allantoic cavities of specific pathogen free (SPF) embryos and stored at 80°C until use. Fifty-percent embryo infectious dose (EID50) titers were determined by serial titration in embryos and calculated by the method of Reed and Muench [39]. HEK 293 T cells were maintained in Dulbecco’s modified Eagle’s medium (DMEM, Sigma) supplemented with 10% fetal bovine serum in humidified 5% CO2 atmosphere at 37°C. SPF female BALB/c mice (4–6 weeks of age) were purchased from Shanghai SLAC Laboratory Animal Co., Ltd., and maintained with free access to sterile food and water. All animal studies were conducted in accordance with the ethical guidelines and were approved by the Ethical Committee of Shanghai Veterinary Research Institute, Chinese Academy of Agricultural Sciences.

Plasmid construction
A total of 162 HA and 105 M2 sequences with full-length of H3N2 subtype SIV were, respectively, downloaded from GenBank database and aligned by MegAlign program supplemented in DNASTAR package (DNASTAR Madison, WI). The consensus sequences were created based on the most common amino acid in each position of the alignment. After the consensus M2 sequence was generated, the first 24 amino acids were selected as M2e motif [40]. Then codons of the consensus HA, as well as M2e and an immunodominant cytotoxic T lymphocyte (CTL) epitope of the NP protein, NP147-155 (TYQRTRALV) [41], were optimized for mammalian expression and synthesized by GenScript Corporation (Nanjing, China). The (G4S)3-linked chimeric ORF encoding M2e, CTL epitope and consensus HA was PCR-amplified by introducing 5’ Sma I and 3’ Nhe I restriction sites for ligation into the pcAGGS vector, under the control of the cytomegalovirus (CMV) enhancer and chicken β-actin promoter (designated as MNHAs). Similarly, MHa was designed and constructed, except that the CTL epitope was omitted from the N-terminal end of the consensus HA (Figure 1). After the recombinant plasmids being identified by nucleotide sequencing, they were propagated in E. coli bacteria and purified using Mega purification kit (Qiagen, Valencia, CA) for in vitro transfection as well as in vivo animal immunization. The final DNA preparations were resuspended in nuclease-free water and stored at −20°C until further use.

In vitro expression of recombinant protein
HEK 293 T cells were seeded in 6-well plates and transfected at 80-90% confluence with 4 μg of MHa, MNHAs or empty vector using Lipofectamine 2000 transfection reagent (Invitrogen), as recommended by the manufacturer. After 48 h, the transfected cells were scraped from the culturing plate, washed with PBS, then spotted onto a glass slide, air dried, and fixed with pre-chilled acetone.
Upon removal of the residual solvents from the slides, the cells were incubated with anti-M2e and -HA polyclonal antibodies for 1 h at 37°C. A secondary Alexa Fluor 568-conjugated goat anti-mouse IgG antibody was then used to detect the primary antibody. Fluorescence images were scanned using an inverted microscope after the samples were mounted by glycerol.

**Gene gun delivery of DNA and virus challenge**

Gene gun immunization was performed as previously described [42,43]. Mice received two nonoverlapping abdominal deliveries of antigen encoding plasmid- or empty vector-coated gold beads (1 μm) at the shaved skin with a 3-wk interval. With each shot, 2 μg of DNA immobilized onto 0.5 mg gold particles was delivered at a helium discharge pressure of 450–500 psi with a Helios gene gun (Bio-Rad). Each test group contained 43 mice with experiments organized as follows: (1) 3 mice of each group were used to complete IFN-γ ELISPOT assays on day 21 post the 2nd immunization; (2) the left 40 mice of each group were divided as 4 subgroups, each comprised of 10 mice, then intranasally challenged with 10^6 EID_{50} of two H3N2 (SwHLJ1 and r164) and two H1N1 (G11 and rPan09) strains, respectively, at 21 days after the final immunization.

**Serologic testing**

Serum samples were collected from orbital bleeds on day 14 post last immunization. M2e and HA antibody titers were measured using indirect ELISA. ELISA plates were coated overnight with synthetic M2e peptides or inactivated SwHLJ1 in 0.1 M carbonate buffer (pH 9.6) at 4°C and blocked with 5% non-fat milk for 2 h at 37°C. A series of two-fold dilution of sera (starting dilution 1:500 for HA and 1:100 for M2e, 100 μl/well) were incubated at 37°C for 1 h, followed by three washes with PBST. Then HRP-conjugated goat anti-mouse IgG (Zhongshan Biotechnology Co., Ltd Beijing, China) and 3,3'-5,5'-tetramethyl benzidine (TMB) was used for M2e- and HA-specific antibody detection and color development, respectively. The resulting optical density (OD) at 450 nm was determined with a plate reader after the reaction was stopped with 2 M H_{2}SO_{4}.

HI assays were performed using 0.5% chicken red blood cells (RBCs) with 4 HA units of homologous (SwHLJ1 and r164) and heterologous (G11 and rPan09) virus and receptor-destroying enzyme (RDE, sigma) treated serum, as previously described [44]. The reciprocal of highest dilution of serum that gave complete inhibition of hemagglutination was considered the HI titer. Each assay was performed in triplicate. A titer of less than 10 (starting serum dilution) was assigned for serum samples that did not inhibit hemagglutination.

**IFN-γ ELISPOT**

The ELISPOT assays were performed using mouse IFN-γ ELISPOT kits following methods recommended by the manufacturer (Dakewe Biotech, PR China). Briefly, single-cell suspensions of freshly isolated spleen lymphocytes were seeded into the plates (10^5/well) pre-coated with anti-IFN-γ monoclonal antibody. Cells were stimulated with synthesized NP147-155 peptide at a final concentration of 5 μg/ml in a 37°C humidified incubator with 5% CO_{2}. Phythoemagglutinin (PHA, 5 μg/ml) and medium alone were used as positive and negative controls, respectively. After a 36 h culture, plates were washed and incubated for 1 h with biotinylated anti-mouse IFN-γ antibody. Streptavidin-horseradish peroxidase was then added, IFN-γ spots were developed with 3-amino-9-ethylcarbazole (AEC) and counted using an automated ELISPOT reader. Results were expressed as spot-forming cells (SFC) per million cells.

**Lung viral titer measurement and histopathological examination**

On day 3 post-infection, mice from each group were sacrificed to collect lungs for virus titration and pathologic examination to determine the protective ability of DNA vaccines. For virus titration, the whole lungs were homogenized in 1 mL of sterile phosphate-buffered saline (PBS) containing 0.1 mg/ml of streptomycin and 100 IU/ml of penicillin. The homogenates were diluted 10-fold serially after being centrifuged, then inoculated into embryonated chicken eggs. Infection within individual eggs was confirmed using a standard hemagglutination assay, the viral titers were determined by the method of Reed and Muench.

Lung tissue histopathologic sections were made as described elsewhere [45]. Briefly, the removed lungs were fixed in 10% neutral buffered formalin, dehydrated and embedded in paraffin. Five micrometer sections were cut and stained with hematoxylin and eosin (H&E), and reviewed for histopathological changes.

**Statistical analysis**

Data of experimental and control groups were presented as the averages ± standard error (SE) and evaluated by ANOVA method, where statistically significant results were defined as having a p value of less than 0.05.

**Competing interests**

The authors declare that they have no competing interests.

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