An MDCK Cell Culture-Derived Formalin-Inactivated Influenza Virus Whole-Virion Vaccine from an Influenza Virus Library Confers Cross-Protective Immunity by Intranasal Administration in Mice

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It is currently impossible to predict the next pandemic influenza virus strain. We have thus established a library of influenza viruses of all hemagglutinin and neuraminidase subtypes and their genes. In this article, we examine the applicability of a rapid production model for the preparation of vaccines against emerging pandemic influenza viruses. This procedure utilizes the influenza virus library, cell culture-based vaccine production, and intranasal administration to induce a cross-protective immune response. First, an influenza virus reassortant from the library, A/duck/Hokkaido/Vac-3/2007 (H5N1), was passaged 22 times (P22) in Madin-Darby canine kidney (MDCK) cells. The P22 virus had a titer of >2 × 10^6 PFU/ml, which was 40 times that of the original strain, with 4 point mutations, which altered amino acids in the deduced protein sequences encoded by the PB2 and PA genes. We then produced a formalin-inactivated whole-virion vaccine from the MDCK cell-cultured A/duck/Hokkaido/Vac-3/2007 (H5N1) P22 virus. Intranasal immunization of mice with this vaccine protected them against challenges with lethal influenza viruses of homologous and heterologous subtypes. We further demonstrated that intranasal immunization with the vaccine induced cross-reactive neutralizing antibody responses against the homotypic H5N1 influenza virus and its antigenic variants and cross-reactive cell-mediated immune responses to the homologous virus, its variants within a subtype, and even an influenza virus of a different subtype. These results indicate that a rapid model for emergency vaccine production may be effective for producing the next generation of pandemic influenza virus vaccines.

Before the 2009 H1N1 influenza pandemic occurred, it was widely thought that the next pandemic virus would be the avian H5N1 virus, because hundreds of people worldwide had been infected with the highly pathogenic H5N1 influenza virus since 1997, and 60% of them had died (1–3). However, no pandemic involving the highly pathogenic avian H5N1 influenza virus has occurred. This illustrates how difficult it is to predict which influenza virus strain will give rise to the next pandemic. The seasonal influenza vaccine is produced each year by using a chicken egg-based manufacturing process, which is tedious and time-consuming. This process, which starts with egg preparation and the choice of the seed strain, takes several months to produce an adequate supply of influenza vaccine (4). Therefore, the current vaccine production system would be inadequate to respond to an influenza pandemic, for which a novel rapid emergency vaccine-manufacturing process is required.

All of the genes of mammalian influenza viruses are derived from the avian influenza virus pool (3). In addition, avian influenza viruses are subjected to little immunological pressure in their natural hosts, water fowl. For these reasons, the antigenic sites of the virus are thought to be fairly well conserved in avian and pandemic viruses. Therefore, we hypothesized that all pandemic influenza viruses originate from an avian influenza gene pool and that these viruses express antigens that elicit protective immune reactions from host animals, including humans. The World Organization for Animal Health (OIE) Reference Laboratory for avian influenza at Hokkaido University has established an influenza virus library of all hemagglutinin (HA) and neuraminidase (NA) subtypes and their genes (5–10). The library includes influenza virus strains isolated from natural hosts; these strains have been used to prepare vaccines and have proved to be useful for raising the level of preparedness for future pandemics.

Most conventional influenza vaccines have been manufactured by the embryonated chicken egg-based process. However, the extended time required to produce egg-dependent vaccines might result in too few doses being available to counter a pandemic situation, such as occurred in 2009, or to stop a pandemic originating from a highly pathogenic avian virus, such as an H5N1 virus (11). An alternative to the egg-based process is viral propagation in mammalian cell lines, which also has been used to produce influenza vaccines (12–15). Several cell lines are currently approved for cell culture-based influenza production, and the use of Madin–Darby canine kidney (MDCK) cells and African green monkey Vero cells has been well documented (16–18).

Since a pandemic virus will not always be available from among existing library virus stocks, a cross-protective vaccine de-
sign that can use a stock virus with antigenic properties similar to those of the pandemic virus to generate a vaccine should be established. In animal models, intranasal immunization with an influenza virus split-virion vaccine with mucosal adjuvant [e.g., cholera toxin B, poly(I-C), or poly(γ-glutamic acid) nanoparticles] induces cross-protection and in vivo virus clearance against a drift variant within a subtype and against different subtypes of virus (19–23). Furthermore, intranasal immunization of mice with formalin-inactivated intact virus alone (without adjuvant), but not with an ether-split vaccine alone, induces cross-protection against the homologous virus, intrasubtype variants, and influenza A viruses of different subtypes (24, 25).

Thus, to protect populations from the next influenza pandemic, a novel rapid emergency vaccine-manufacturing process is urgently needed. For this purpose, we propose a process involving the following steps. (i) Seed viruses for the emergency influenza vaccine that grow well in MDCK cells are made in advance from influenza virus strains in the vaccine library and stored. (ii) The genes of the selected seed viruses are analyzed. (iii) When a pandemic occurs, the seed virus whose HA antigenicity most closely matches that of the pandemic virus is grown in MDCK cells and used to make a formalin-inactivated whole-virion vaccine. (iv) Individuals are immunized intranasally with the vaccine to protect them from the pandemic influenza virus. In the present study, we examined the feasibility of this approach to overcome shortages in the effective influenza vaccine supply in emergency pandemic situations, using a pathogenic H5N1 influenza virus reassortant strain from the influenza virus library.

MATERIALS AND METHODS

Cells. MDCK cells, which are kindly provided by the Kanonji Institute, The Research Foundation for Microbial Diseases of Osaka University (Kanonji, Japan), were maintained in a serum-free medium, Opti-Pro SFM (Invitrogen, Carlsbad, CA) supplemented with 0.3 mg/ml l-glutamine and 8 μg/ml gentamicin. The cells were maintained at 37°C in 5% CO₂ (26).

Virus strain. A low-pathogenicity influenza virus reassortant strain, A/duck/Hokkaido/Vac-3/2007 (Vac-3) (H5N1) (10), was stock in an influenza virus library containing all HA and NA subtypes and their genes. The Vac-3 H5N1 strain was generated originally with A/duck/Hokkaido/101/04 (H5N3) as the HA and NS gene provider and A/duck/Hokkaido/262/04 (H6N1) as the PB2, PB1, PA, NP, NA, and M gene provider. The complete nucleotide sequences of the Vac-3 H5N1 strain are registered in GenBank (accession numbers AB355926 to AB355933). For mouse challenge experiments, mouse-adapted Vac-3 passaged 22 times (P22) (H5N1), mouse-adapted A/Brisbane/59/07 (BR/59/07) (H1N1), A/Hong Kong/483/1997 (HK/483/97) (H5N1), clade 0, which was isolated from a patient with a highly pathogenic avian influenza virus infection in 1997, and A/peregrine falcon/Hong Kong/810/2009 (PF/HK/810/09) (H5N1, clade 2.3.4.), which is a highly pathogenic avian influenza virus, were used. (Alignment of the deduced HA amino acid sequences of the H5N1 strains is shown in Fig. 1). A/Brissbane/59/07 (BR/59/07) (H1N1) was kindly provided by Takato Odagiri (National Institute of Infectious Diseases, Tokyo, Japan). Both Vacc-3 P22 and BR/59/07 were originally nonpathogenic to mice; therefore, mouse-adapted strains for challenge experiments were generated as follows. Mice were inoculated with virus (10⁶ PFU in 40 μl phosphate-buffered saline [PBS]), and their lungs were harvested 3 days later. The lung tissue was homogenized in PBS using Multi-Beads Shocker (Yasui Kikai, Osaka, Japan) (one set of lungs per 1 ml PBS) and centrifuged (10,000 x g for 10 min), and the supernatants were harvested. naive mice were inoculated with the supernatant (40 μl/mouse). These steps were repeated for a total of 10 inoculations. The mouse-adapted viruses were then grown once in MDCK cells, and the mouse 50% lethal dose (LD₅₀) values for the viruses were assessed. Mouse-adapted Vac-3 P22 and mouse-adapted BR/59/07 were grown in MDCK cells, whereas HK/483/97 and PF/HK/810/09 production was in chicken eggs.

Adaptation of the Vac-3 virus to MDCK cells. Vac-3 was used to inoculate MDCK cells cultured at 35°C in minimal essential medium (MEM) supplemented with 0.3 mg/ml l-glutamine, 0.1% bovine serum albumin (BSA), 0.8 μg/ml crystal trypsin (Sigma, St. Louis, MO), and 8 μg/ml gentamicin. Three days after infection, the virus-containing supernatant was collected and was used to measure the virus titer and to inoculate fresh MDCK cells at a 1:1,000 dilution. After 22 passages, the virus-containing supernatant was collected and stored at −80°C. The stock virus, A/duck/Hokkaido/Vac-3/2007 P22 (Vac-3 P22), was used as a seed virus for an inactivated whole-virion influenza vaccine.

Virus growth assay. Confluent MDCK cell monolayers in 6-cm petri dishes were infected with virus at a multiplicity of infection (MOI) of 0.001, in 800 μl of MEM with 0.1% BSA. After being washed with MEM without supplement, the infected cells were incubated at 35°C in 5 ml virus growth medium supplemented with 0.8 μg/ml crystal trypsin. The supernatant was collected at the times indicated in Fig. 2, and the virus titer in the supernatant was determined by plaque assay.

Plaque assay. Viruses diluted 10-fold in 300 μl of MEM with 0.1% BSA were applied to confluent monolayers of MDCK cells in 6-well plates and incubated at 35°C for 1 h. Unbound viruses were removed, and the cells were washed with MEM. The cells were then overlaid with 2 ml virus growth medium containing 0.8% agarose (Sigma) and 0.8 μg/ml crystal trypsin (Sigma). After a 72-h incubation at 35°C, the cells were fixed with 10% formaldehyde and stained with 0.1% crystal violet solution.

Virus gene sequencing. At two time points, i.e., after 2 passages and after 22 passages, the virus-containing supernatant was collected and the virus was purified by plaque cloning. This purification step was repeated 3 times at each time point. The virus-containing supernatants were collected, and the viral RNA was isolated using TRIzol reagent (Life Technologies, Carlsbad, CA) and reverse-transcribed with the Uni-12 primer (27), using a Superscript III reverse transcriptase kit (Life Technologies). The viral genes were then amplified by PCR with KOD-Plus DNA polymerase (Toyobo, Osaka, Japan), according to the manufacturer’s protocol, using the universal primer set (27). The PCR products were purified with a QIAquick gel extraction kit (Qiagen, Venlo, Netherlands). The DNA template was sequenced using a BigDye direct cycle sequencing kit (Life Technologies), according to the manufacturer’s protocol. The sequencing primers were designed according to the NCBI reference sequence, and samples were analyzed using an Applied Biosystems 3500 genetic analyzer (Life Technologies). The DNA sequences were completed and edited using the DNASeqS Pro software package (Hitachi Solutions, Tokyo, Japan).

Vaccine preparation. The seed virus, Vac-3 P22 (H5N1), was grown for vaccine production using a BelloCell cell culture system (CESCO Bioengineering, Taichung, Chinese Taipei). For cell attachment, a bottle was filled with 500 ml medium (10% fetal bovine serum [FBS] in MEM, with the glucose level adjusted to 3 g/liter) and inoculated with MDCK cells at 4 x 10⁹ cells/ml. The bottle was then placed on a BelloStage console, with settings of an up/down speed of 2 mm/s and holding times of 30 s at the top and 0 s at the bottom for the first 3 h. The settings were then changed to an up/down speed of 1 mm/s and holding times of 10 s at the top and 30 s at the bottom, to allow sufficient aeration. The medium was completely replaced after the first 48 h and then either the medium was partially replaced or glucose concentrate was added as needed to maintain a glucose level of 2 g/liter for the MDCK cells. The cell density was measured periodically using a CVD nucleus staining kit (CESCO Bioengineering), by spotting samples taken from the bottle onto disks. Glucose concentrations were measured using a GlucCell meter (CESCO Bioengineering).

When the MDCK cell concentration reached about 2 x 10⁶ to 3 x 10⁷ cells/ml, the cultivation medium was completely aspirated and replaced with 500 ml of PBS. The bottles were placed on the BelloStage and washed for 5 min using the above-described settings for aeration, and then the
PBS was completely aspirated and replaced with infection medium (MEM supplemented with 0.3 mg/ml L-glutamine, 0.1% BSA, 0.8 μg/ml crystal trypsin, and 0.8 μg/ml gentamicin, with the glucose concentration adjusted to 3 g/liter). The cells were then infected at a MOI of 0.001 and incubated using the same settings for the BelloStage as for the cultivation.

Seventy-two hours after infection, the medium was collected and passed through a 0.22-μm filter (TPP, Trasadingen, Switzerland). The viruses in the medium were then concentrated and purified as follows. First, a tangential flow filtration ultrafiltration set (Millipore, Bedford, MA) was used for the primary concentration (100-fold), using a filter with a 50-kDa cutoff value. The viruses were further concentrated by two centrifugations through a discontinuous sucrose gradient (30% and 60% [wt/vol] sucrose in PBS) at 75,000 × g for 1.5 h at 4°C, in a Himac CP-80WX preparative ultracentrifuge (Hitachi, Japan). The purified viruses were recovered from the interface of the two sucrose layers, transferred to a 1- to 5-ml QuixSep microdialyzer cassette (Orange Scientific, Braine-l’Alleud, Belgium), and dialyzed against 2 liters of PBS at 4°C for 24 h. After dialysis, the protein concentration was measured using a Pierce BCA protein assay kit (Thermo Scientific, Waltham, MA) and adjusted to 1 mg/ml, and the virus was inactivated by incubation with 0.02% formalin for 28 days at 4°C. Virus inactivation was confirmed by several blind passages on MDCK cells, followed by visually checking for plaque formation under a microscope. Finally, the formalin-inactivated vaccine was pressure-dialyzed against PBS using a QuixSep microdialyzer (Orange Scientific). For every batch (4 bottles) with a total volume of 2 liters of

FIG 1 Alignment of deduced HA amino acid sequences of the H5N1 strains used in this study. The alignment was based on published amino acid sequences (GenBank accession numbers BAF76006.1, AAF74330.1, and BAI39636.1). Black blocks, regions with identical amino acids; gray blocks, regions with similar amino acids; white blocks, regions with nonidentical nonsimilar amino acids. Dashes, absent or deleted amino acids.

FIG 2 Enhanced replication of A/duck/Hokkaido/Vac-3/2007 (Vac-3) (H5N1) with repeated passage in MDCK cells. MDCK cells were inoculated at 35°C with Vac-3 in MEM supplemented with 0.3 mg/ml L-glutamine, 0.1% BSA, 0.8 μg/ml crystal trypsin, and 0.8 μg/ml gentamicin. Three days after infection, the virus-containing supernatant was collected and used for the inoculation of fresh MDCK cells at a 1:1,000 dilution, and the viral titer in the supernatant was measured by a virus plaque assay.
infection medium, we obtained 3 to 4 mg of total viral protein after concentration and sucrose gradient centrifugation.

Seed viruses in suspension were inactivated by incubation with 0.2% formalin for 28 days at 4°C. Virus inactivation was confirmed by inoculation into MDCK cells, followed by checking for plaque formation. The formalin-inactivated virus suspension was diluted with PBS to a final concentration of 100 μg total virus protein per ml.

**Vaccination and virus challenge.** BALB/c mice (6 weeks old, female; Japan SLC, Shizuoka, Japan) were anesthetized and inoculated intranasally (inoculation volume, 20 μl) with PBS or PBS containing Vac-3 P22 (H5N1) formalin-inactivated whole-virion vaccine (1, 3, or 10 μg), on days 0 and 21. To assess mouse mortality rates upon virus infection, the mice were inoculated intranasally, 14 days after the final immunization, with 40 μl of PBS containing one of the following: 100 × LD50 of mouse-adapted Vac-3 P22 (106 PFU), 100 × LD50 of HK/483/97 (H5N1, clade 6; 10−1 × 50% egg infectious dose [EID50]), 100 × LD50 of PF/HK/810/09 (H5N1, clade 2.3.4; 10−3.5 × EID50), or 100 × LD50 of mouse-adapted BR/59/07 (H1N1; 102 PFU). The mortality status and weight loss of the mice were assessed daily for up to 14 days thereafter. To measure virus clearance in the lungs of the vaccine-immunized mice, the mice were inoculated intranasally, 14 days after the final immunization, with 20 μl of PBS containing 100 × LD50 of influenza virus. Three days later, the lungs were excised and homogenized using Multi-Beads Shocker (Yasui Kikai, Ibaraki, Japan) were suspended in 1 ml of FBS-free RPMI 1640 medium, the cells were washed 3 times with complete RPMI 1640 medium, the target cells were resuspended at 105 cells/ml in complete RPMI 1640 medium. The target cells were mixed in a 75-cm2 culture flask and cocultured for 6 days at 37°C in 5% CO2, to induce virus-specific cytotoxic T lymphocytes (CTLs) (effector:target cell ratio = 50:1, 17:1, 5:1, and 1.7:1. Following a 4-h incubation at 37°C in 5% CO2, 100 μl of the supernatant was removed to determine the concentration of 51Cr released. All samples were counted in a TopCount NXT microplate scintillation and luminescence counter (PerkinElmer), and percent specific release was calculated as [(experimental release − spontaneous release)/(total release − spontaneous release)] × 100.

**Statistical evaluations.** Fisher’s exact test, performed using Statcel2 software (OMS, Tokyo, Japan), was used to evaluate the differences between groups in the mortality experiments. To analyze the data in the other experiments, nonparametric Student t tests were used. P values of <0.05 were considered significant.

**RESULTS**

Adaptation of an H5N1 influenza A virus strain from the influenza virus library for MDCK cells and mutations associated with the adaptation to MDCK cells. We first adapted an influenza A virus reassortant strain, A/duck/Hokkaido/Vac-3/07 (Vac-3) (H5N1), from an influenza virus library containing all HA and NA subtypes for growth in MDCK cells by performing serial passages in those cells. The initial viral titers in the first and second passages were 6 × 106 PFU/ml and 3.3 × 106 PFU/ml, respectively (Fig. 2). After 8 passages, the viral titer had increased more than 5 times; after 16 passages, the titer was over 108 PFU/ml. After 22 passages, the viral titer remained the same, at 2.4 × 108 PFU/ml (Fig. 2).

To understand the molecular basis for the 40-fold increase in viral titer caused by adaptation to MDCK cells, twice-passaged virus (P2), in which the amino acids were all the same as in the original strain (data not shown), and 22-times-passaged virus (P22) Vac-3 were biologically cloned by virus plaque purification and sequenced. The sequences of all six P2 clones examined were the same as that of the original strain (data not shown). The sequences of all six P22 clones examined also matched each other, with acquired mutations mapping to the genes for the PB2 and PA proteins but not the HA protein (Fig. 3A). The P22 clone showed much greater growth than the P2 clone in MDCK cells (Fig. 3B); thus, the mutations might be associated with the enhanced viral growth in MDCK cells. Although amino acid changes at PB2 position 627 (from E to K) and PB2 position 701 (from D to N) are known to result in enhanced viral replication (29, 30), these amino acid changes were not found in P2 or P22.

**Cross-protection of mice immunized with inactivated whole-virion vaccine from MDCK cell-adapted A/duck/Hokkaido/Vac-3/07 P22 (Vac-3 P22) against challenges with influenza viruses of the same subtype.** We next produced a formalin-inactivated whole-virion influenza vaccine from the P22 strain of Vac-3 (Vac-3 P22) (H5N1) and determined whether the intranasal immunization of mice with this vaccine induced protection against the homotypic virus. Mice were intranasally inoculated twice with 20 μl of PBS or PBS containing the vaccine. We then examined the survival of the intranasally inoculated mice after infection with 100 × LD50 of Vac-3 P22 or A/Hong Kong/483/1997 (HK/483/97) (H5N1, clade 0). All of the intranasally PBS-treated mice lost more than 20% of their body weight and died. In contrast, all of the mice intranasally immunized with 1, 3, or 10 μg of the vaccine lost less than 10% of their body weight after infection with either virus and survived (Fig. 4A and B).

Next, we examined whether intranasal immunization of mice with the A/duck/Hokkaido/Vac-3/07 P22 formalin-inactivated whole-virion vaccine induced protection against A/peregrine falcon/Hong Kong/810/2009 (PF/HK/810/09) (H5N1, clade 2.3.4), an antigenic variant within the viral subtype. All of the mice intranasally immunized with 3 or 10 μg of the vaccine and 40% of the mice
intranasally immunized with 1 μg of the vaccine lost less than 10% of their body weight after PF/HK/810/09 infection and survived (Fig. 4C).

We next determined the virus clearance against Vac-3 P22 (H5N1) and HK/483/97 (H5N1, clade 0) in mice intranasally immunized with the formalin-inactivated whole-virion vaccine. Intranasally immunized mice were infected with 100 \( \times \) LD\(_{50}\) of the two virus strains 14 days after the final immunization. Three days after infection, the lungs of the infected mice were homogenized, and the viral titers in the lung homogenates were assessed. Intranasal immunization with 1, 3, or 10 μg of the vaccine significantly decreased the titers of both viruses (Fig. 5A and B). In addition, intranasal immunization with 3 or 10 μg of the vaccine significantly decreased the titers of PF/HK/810/09 (H5N1, clade 2.3.4) (Fig. 5C).

Since Vac-3 P22 formalin-inactivated whole-virion vaccine induced cross-protection against PF/HK/810/09, which has the most varied HA in comparison with Vac-3 P22, these results suggest that the vaccine from MDCK cell-adapted Vac-3 P22 would be effective as a pandemic vaccine for H5N1 influenza virus.

Cross-protection of mice immunized with inactivated whole-virion vaccine from MDCK cell-adapted Vac-3 P22 against an influenza virus of a different subtype. We next examined whether intranasal immunization of mice with the Vac-3 P22 formalin-inactivated whole-virion vaccine induced protection against an influenza virus of a different subtype. Mice were intranasally inoculated twice with 20 μl of PBS or PBS containing 3 μg of the vaccine, and their survival after infection with BR/59/07 (H1N1) was examined. All of the intranasally PBS-inoculated mice lost more than 20% of their body weight and died. In contrast, all of the mice intranasally immunized with the vaccine lost less than 10% of their body weight after BR/59/07 (H1N1) infection and survived (Fig. 6A). Furthermore, intranasal immunization with the vaccine significantly decreased the titers of BR/59/07 (H1N1) (Fig. 6B). Thus, intranasal immunization with the inactivated whole-virion vaccine from MDCK cell-adapted Vac-3 P22 induced cross-protection against a different subtype of the influenza virus.

Cross-reactive neutralizing antibodies against several influenza viruses in mice intranasally immunized with the whole-virion vaccine from MDCK cell-adapted Vac-3 P22. Several
studies have demonstrated that intranasal immunization induces both humoral and cell-mediated immunity, suggesting that either or both of them might contribute to cross-protection. Therefore, we next examined how intranasal immunization with Vac-3 P22 formalin-inactivated whole-virion vaccine induced cross-protection. First, we determined the neutralizing antibody titers against several influenza virus strains in mice intranasally immunized with the Vac-3 P22 vaccine. Neutralizing antibodies for Vac-3 P22 (H5N1) and HK/483/97 (H5N1), but not for PF/HK/810/09 (H5N1) or BR/59/07 (H1N1), were detected in both BAL fluids and sera of immunized mice (Fig. 7). These results suggest that the vaccine induced a cross-protective humoral immune response to the homologous virus and to some of the antigenic variants within the influenza virus subtype but not to all variants within the subtype or to viruses of a different subtype.

Cross-reactive cell-mediated immunity against several influenza viruses in mice intranasally immunized with whole-virion vaccine from MDCK cell-adapted Vac-3 P22. We next determined whether a cross-protective cell-mediated immune response contributed to cross-protection against different subtypes of influenza virus, by performing IFN-γ ELISPOT assays. Interestingly, immunization with the Vac-3 P22 formalin-inactivated whole-virion vaccine resulted in equally greater numbers of IFN-γ spots in MNCs restimulated with UV-treated Vac-3 P22 (H5N1), PF/HK/810/09 (H5N1), and BR/59/07 (H1N1) (Fig. 8). Next, we assayed the cytotoxicity of CTLs against influenza virus-infected cells. As shown in Fig. 9, intranasal immunization with the Vac-3 P22 vaccine induced CTL activity not only for Vac-3 P22 (H5N1) but also for BR/59/07 (H1N1). The cross-reactive CTL activity against PF/HK/810/09 (H5N1), PF/HK/810/09 (H5N1), and BR/59/07 (H1N1) (Fig. 8).

Next, we assayed the cytotoxicity of CTLs against influenza virus-infected cells. As shown in Fig. 9, intranasal immunization with the Vac-3 P22 vaccine induced CTL activity not only for Vac-3 P22 (H5N1) but also for BR/59/07 (H1N1). The cross-reactive CTL activity against PF/HK/810/09 (H5N1), PF/HK/810/09 (H5N1), and BR/59/07 (H1N1) (Fig. 8).

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DISCUSSION

In the present study, we have assessed whether our novel emergency vaccine-manufacturing process would be useful for preventing infection in the next influenza pandemic. We have demonstrated that our process, which uses an influenza virus library, cell culture-based vaccine production, and intranasal immunization to induce cross-protection, provides effective protection against homotypic and drift variants within a subtype of influenza virus infections. These results suggest that this vaccine-manufacturing process might be applied effectively for the next generation of pandemic influenza vaccines.

It is currently impossible to predict which influenza virus strains will cause the next pandemic. We previously proposed the use of avian influenza viruses as candidate vaccine strains when antigenically related and low-pathogenicity viruses from humans are not available (6, 7). Therefore, we have been establishing a library of nonpathogenic influenza A virus strains of different subtypes, without using reverse genetics technology (7). Furthermore, Itoh et al. demonstrated that a vaccine prepared from a nonpathogenic influenza virus strain from the influenza library conferred protective immunity against infection by a highly pathogenic avian influenza virus of the same subtype (6). In the present study, we demonstrated that a virus strain from the influenza library was modified to a high-titer vaccine virus by adaptation in MDCK cells and that intranasal immunization with the formalin-inactivated virus vaccine induced cross-protection against homotypic and heterotypic influenza virus infections.

Efficient influenza vaccine production requires a system that can produce a large amount of vaccine virus rapidly and safely. The cell-based method has several advantages over the current process using embryonated chicken eggs, including the following: (i) it enables faster and larger-scale vaccine production; (ii) it avoids the potential selection of variants adapted for chicken eggs, which alters virus antigenicity; (iii) egg-based production requires the selection of high-yield vaccine seed viruses; and (iv) egg-produced viruses may contain allergenic components of eggs (11, 12, 16, 31–34). Due to these advantages, the World Health Organization (WHO) has recommended that mammalian cell culture-based vaccines be established (35). Based on our present results, we propose that the preparation and storage of cell-adapted vaccine viruses from the influenza virus library would be useful for the rapid preparation of an emergency vaccine for an influenza pandemic. However, there are still several problems with the substantiation of faster and larger-scale production of a cell-adapted vaccine that need to be clarified in future studies. In addition, it is true that the virus selected for production in the event of a real
Pandemic might not behave like the current one. This is why many studies on the MDCK cell library strains are ongoing, to establish the cross-protection range for different strains within the same subtype or in different subtypes.

In the present study, MDCK cell-adapted Vac-3 passed 22 times showed >30-times-greater viral growth in MDCK cells than that after two passages (when the amino acid sequence was identical to that of the original strain) (Fig. 3B). Four point mutations were associated with enhanced viral growth, i.e., three in PB2 (V145F, E472K, and M632L) and one in PA (E319Q). A major determinant of viral tropism is the influenza virus polymerase, since the polymerase, which is composed of the viral proteins PB1, PB2, and PA, assembles with viral RNA and nucleoprotein to mediate transcription and replication of the viral genome (36), suggesting that these mutations are associated with enhanced viral growth. Several studies have examined why the repeated passage of influenza viruses through cultured mammalian cells leads to increased viral growth. For example, the amino acid changes at PB2 position 627 (from E to K) (37–42) and PB2 position 701 (from D to N) (29, 30) in several highly pathogenic avian influenza viruses are important for enhancing viral replication in mammals. In addition, Murakami et al. (11) demonstrated that the amino acid change at HA2 position 117 (from N to Y) in human H1N1 influenza virus strain A/Puerto Rico/8/1934 is one of the factors responsible for the enhanced viral growth observed after repeated passage in Vero cells. However, the amino acid at PB2 position 627 was E and that at PB2 position 701 was D in both P2 and P22, indicating that the previously reported amino acid changes did not occur. Thus, the changes we observed in PB2 (V145F, E472K, and M632L) and PA (E319Q) might be novel mutations that promote viral growth in MDCK cells. We are now studying which changes are most responsible for enhanced growth.

Several studies have demonstrated that intranasal immunization with several forms of influenza vaccine induces cross-protection against drift variants within a subtype and against viruses of different subtypes (19–22, 24, 25, 43). Cross-protection with intranasal immunization has been shown to occur mainly through influenza virus-specific humoral immunity (i.e., anti-influenza neutralizing antibody secretion in serum and mucosa) and cell-mediated immunity (i.e., cytotoxic T-cell activity against influenza virus-infected cells). Intranasal immunization with gamma-radiation-inactivated whole-virion influenza vaccine induces influenza virus-specific cross-reactive humoral and cellular immunity (44, 45). On the other hand, formalin-inactivated whole-virion influenza vaccine induces cross-reactive humoral immunity but rarely cellular immunity (24, 45). Furthermore, cross-reactive neutralizing antibodies contribute to the cross-protection caused by intranasal immunization with formalin-inactivated whole-virion vaccine, and cross-reactive neutralization by intranasal immunization with a formalin-inactivated H1N1 influenza virus vaccine includes drift variants within the influenza virus subtype (24). However, Takada et al. demonstrated that formalin-inactivated H5N1 influenza virus whole-virion vaccine in intranasal immunization does not induce cross-protection against different influenza virus subtypes through cross-reactive neutralizing antibodies (25). In the present study, we have demonstrated that intranasal immunization with a formalin-inactivated H5N1 influenza virus vaccine induces cross-reactive humoral and cellular immunity. While cross-reactive neutralizing antibody responses were observed for the homotypic H5N1 influenza virus and its drift variants but not a virus of a different subtype, cross-reactive cell-mediated immune responses were observed for all strains tested, including that of a different subtype. Therefore, formalin-inactivated influenza vaccine might induce cross-reactive cell-mediated immune responses that provide cross-protection against viruses of different subtypes. There are many responses other than the antiviral neutralization response associated with cross-protection, e.g., antibody-dependent cell cytotoxicity. It should be noted that cross-protection against the H1N1 virus is at least partially NA-mediated. Since no challenge with a virus carrying different HA and NA forms has been included, we must clearly limit the description of “cross-reactive” to H5N1 versus H1N1.

Our previous study (24) demonstrated that intranasal immunization with formalin-inactivated whole-virion influenza vaccine from human seasonal H1N1 influenza viruses did not induce cross-reactive cell-mediated immunity, and the reason why the formalin-inactivated H5N1 influenza virus whole-virion vaccine,

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**FIG 9** Cross-reactive anti-influenza CTL activity in immunized mice. Mice (5 per group) were inoculated intranasally with PBS (no Vaccine) or with 3 μg of formalin-inactivated whole-virion vaccine from MDCK cell-adapted Vac-3 P22 (Vaccine) or were infected intranasally with 10^3 PFU of Vac-3 P22 (Virus) twice. Fourteen days after the final immunization, spleen MNCs were restimulated with syngeneic mouse spleen cells infected with Vac-3 P22 (H5N1) (A) or BR/59/07 (H1N1) (B), to induce effector cells. 51Cr-labeled target cells infected with Vac-3 P22 or BR/59/07 and the effector cells, restimulated with the same virus as was used to infect the target cells, were added at the indicated effector-to-target cell ratios. Following a 4-h incubation at 37°C in 5% CO2, 100 μl of the supernatant was harvested to determine the concentration of 51Cr released. *P < 0.01 versus the group of mice immunized with PBS.
but not the H1N1 influenza virus whole-virus vaccine, induced cross-reactive cell-mediated immune responses remains to be elucidated. It also is still unclear which formalin-inactivated virus strains can induce cross-reactive cell-mediated immune responses and whether an intranasal formalin-inactivated whole-virus vaccine can induce cross-protection against a virus of a different subtype via a humoral immune response in some cases.

Systemic immunization with formalin-inactivated whole-virus influenza vaccine derived from a pathogenic avian H5N1 influenza virus, but not from human H1N1 influenza viruses, induced CTL activity against the homotypic virus and a highly pathogenic influenza virus (46). Furthermore, Budimir et al. (47) reported that subcutaneous immunization with β-propiolactone and a formalin-inactivated whole-virus vaccine induced cross-protection against an influenza virus of a different subtype, in a CD8+ T-cell-dependent manner. In the present study, we confirmed that intranasal immunization, like subcutaneous immunization, with a formalin-inactivated whole-virus influenza vaccine induced cross-reactive cell-mediated immune responses against homotypic and heterotypic influenza viruses. However, whether the range and magnitude of the cross-reactive cell-mediated immune responses and cross-protection induced by intranasal immunization with a formalin-inactivated whole-virus vaccine are greater than those induced by subcutaneous immunization remains unknown. These procedures will be compared in future studies.

The ELISPOT assay suggests that the responses are T-cell mediated; however, which antigens play a role in the cross-reactive T-cell-mediated immune responses remains elusive. For future studies, we will consider using overlapping peptide libraries covering different viral proteins, which would give a better idea of how much of the T-cell response is directed against HA versus more-conserved internal proteins.

In the present study, we found that intranasal immunization with a formalin-inactivated whole-virus influenza vaccine induced cross-protection against the homotypic virus, drift variants within a virus subtype, and a virus of a different subtype. The cross-protection was due, at least in part, to cross-reactive neutralizing antibodies in the nasal cavity, lungs, and serum and to IFN-γ-secreting cells. Influenza virus vaccines for humans need to induce effective cross-protection with minimal side effects. A cold-adaptive live influenza vaccine for intranasal immunization (FluMist; MedImmune) is available clinically in the United States and Europe but not in Japan. However, it is approved only for individuals 2 to 49 years of age and it can result in viral replication, leading to influenza-like syndromes (24). The formalin-inactivated whole-virus vaccine is licensed clinically only for intramuscular injection; the safety and side effects of intranasal immunization with formalin-inactivated whole-virus vaccine still need to be established. Nevertheless, since viral reaction cannot occur with a formalin-inactivated whole-virus vaccine, this should be a good candidate for intranasal pandemic influenza vaccines for clinical use in the near future.

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