Immunogenicity and Protective Efficacy of Replication-Incompetent Influenza Virus-Like Particles

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Despite the success of influenza virus vaccines in reducing severe illness, their efficacy is suboptimal. We describe here the immunogenicity and protective capacity of replication-incompetent influenza virus-like particles (VLPs) which were generated entirely from cDNAs and lacked either the entire NS gene (encoding both the NS1 and NS2 proteins) or only the NS2 gene. In mammalian cells infected with NS gene-deficient VLPs, the nucleoprotein, but not other viral proteins including hemagglutinin (HA) and neuraminidase (NA), was detected. In contrast, cells infected with VLPs expressing NS1 but not NS2 (NS2 knockout) expressed multiple viral proteins, including HA and NA. When challenged with lethal doses of an antigenically homologous mouse-adapted influenza virus, 94% of mice vaccinated with the NS2-knockout VLPs survived, compared with less than 10% of those given the NS-deficient VLPs. These results demonstrate the potential of replication-incompetent NS2-knockout VLPs as novel influenza vaccines and perhaps also as vectors to express genes from entirely unrelated pathogens.

Influenza A virus causes appreciable morbidity and mortality in humans and domestic animals, resulting in large economic losses worldwide (14). The current method of immunization against influenza is parenteral administration of inactivated influenza virus vaccines. Although associated with a very low incidence of adverse reactions in healthy recipients, such vaccines do not efficiently elicit mucosal immunity or a cytotoxic T-cell response. They exhibit a 70 to 90% efficacy in reducing the incidence of clinical illness but fail to prevent influenza virus infection, warranting efforts to develop alternative vaccines.

Cold-adapted live attenuated vaccines have shown considerable promise in ongoing clinical trials, especially in young children, who are poor responders to inactivated vaccines due to the lack of immune memory of influenza virus (1). However, live vaccines have not consistently proved more efficacious than inactivated vaccines in adults (5), and the limited number of amino acid changes in vaccine strains has led to concern over the emergence of virulent revertants (10), although the phenotype of the cold-adapted vaccine is highly stable in clinical trials (2).

Recently, we (17) and others (8) established a system for generating infectious influenza virus entirely from cDNAs. Transfection of cells with plasmids containing cDNAs encoding all eight viral RNAs (vRNAs) of A/WSN/33 (H1N1) virus, controlled by RNA polymerase I promoter and terminator sequences, results in vRNA synthesis by cellular RNA polymerase I. Cotransfection of cells with plasmids for the synthetic gene-deficient VLPs, all Pol constructs for eight RNA segments, excluding pPolINS, and nine protein expression constructs for structural proteins were transfected (17). Plasmids derived from pH21 for the expression of vRNA are referred to as Pol constructs in this report. The plasmids for the expression of the hemagglutinin (HA) (pEWSN-HA), nucleoprotein (NP) (pCAGGS-WSN-NP0/14), neuraminidase (NA) (pCAGGS-WNA15), and M1 (pCAGGS-WSN-M1-2/1) proteins of WSN virus and the M2 (pEP24c), NS2 (pCANS2), PB1 (pDNA774), PB2 (pDNA762), and PA (pDNA787) proteins of A/Puerto Rico/8/34 (H1N1) virus were described in a previous report (17).

Materials and Methods

Cells and viruses. 293T human embryonic kidney cells (a derivative of the 293 line into which the gene for simian virus 40 T antigen was inserted [4]) and Madin-Darby canine kidney (MDCK) cells were maintained in Dulbecco’s modified Eagle medium supplemented with 10% fetal calf serum and in minimum essential medium (MEM) containing 5% newborn calf serum, respectively. All cells were maintained at 37°C in 5% CO2. Influenza A/WSN/33 (H1N1) (WSN) virus was propagated in 10-day-old embryonated chicken eggs.

Plasmids. All genes of the A/WSN/33 virus containing BomBI sites were cloned into the phH21 vector, which contains the human RNA polymerase I promoter and the mouse RNA polymerase I terminator, separated by BomBI sites, resulting in pPolPA, pPolPB, pPolPB2, pPolNP, pPolHA, pPolNA, pPolIM, and pPolNS (17). The NS mutant, pPolDNSplice, was constructed as described earlier (16). Plasmids derived from pH21 for the expression of vRNA are referred to as Pol constructs in this report. The plasmids for the expression of the hemagglutinin (HA) (pEWSN-HA), nucleoprotein (NP) (pCAGGS-WSN-NP0/14), neuraminidase (NA) (pCAGGS-WNA15), and M1 (pCAGGS-WSN-M1-2/1) proteins of WSN virus and the M2 (pEP24c), NS2 (pCANS2), PB1 (pDNA774), PB2 (pDNA762), and PA (pDNA787) proteins of A/Puerto Rico/8/34 (H1N1) virus were described in a previous report (17).

Generation of replication-incompetent influenza virus particles. For generation of gene-deficient VLPs, all Pol constructs for eight RNA segments, excluding pPolINS, and nine protein expression constructs for structural proteins were mixed with transfection reagent (2 μl of Trans IT LT-1 [Panvera, Madison, Wis.] per μg of DNA), incubated at room temperature for 15 min, and added to 107
293T cells. Six hours later, the DNA transfection reagent mixture was replaced by Opti-MEM (Life Technologies, Rockville, Md.) containing 0.3% bovine serum albumin and 0.01% fetal calf serum. Forty-eight hours later, VLPs in the supernatant were harvested.

**Immunostaining assay.** Sixteen hours after infection with influenza VLPs, cells were washed twice with phosphate-buffered saline (PBS) and fixed with 3.7% formaldehyde (in PBS) for 20 min at room temperature. Next, they were treated with 0.1% Triton X-100 and processed as described earlier (15). To examine the efficiency of VLP generation, 10^6 cells were infected with 0.1 ml of the culture supernatant of 293T cells and the number of NP-positive cells as detected by the immunostaining assay was determined at 16 h postinfection.

**Immunization and protection tests.** Formalin-inactivated virus was prepared by first propagating WSN virus in MDCK cells. The virus-containing culture supernatant was then treated with 0.1% formalin at 4°C for a week. Inactivation of the virus was confirmed by inoculating the treated material into MDCK cells. BALB/c mice (4-week-old females) were intranasally immunized with 50 or 100μl of 16 hemagglutinating units (HAU) of formalin-inactivated virus or the replication-incompetent VLPs per ml three times at 3-week intervals. On the ninth week, four mice were sacrificed to obtain sera, trachea-lung washes, and nasal washes. One or three months after the last vaccination, immunized mice were challenged intranasally, under anesthesia, with 10 or 100 50% lethal doses (LD50) of the wild-type WSN virus. For determination of virus titers in lungs, lungs were harvested at day 3 and were homogenized and titrated on MDCK cells. The remaining animals were observed for clinical signs and symptoms of infection for 14 days after challenge.

**Detection of virus-specific antibody.** Serum samples were examined for antibody by an enzyme-linked immunosorbent assay (ELISA) (11). In this assay, the wells were coated with purified WSN virus after treatment with 0.05 M Tris-HCl (pH 7.8) containing 0.5% Triton X-100 and 0.6 M KCl at room temperature and diluted in PBS. After incubation of virus-coated plates with test serum samples, bound antibody was detected with rabbit anti-mouse immunoglobulin A (IgA) (Kirkegaard & Perry Laboratories Inc., Gaithersburg, Md.) and goat anti-mouse IgG (Boehringer-Mannheim, Mannheim, Germany) conjugated to horseradish peroxidase.

**RESULTS**

**Generation of replication-incompetent VLPs.** We initially chose to delete the entire NS gene as a means to produce VLPs that would infect cells and express immunogenic proteins without generating progeny particles in infected cells. NS2 protein, which is encoded by a spliced mRNA (12), is essential for influenza virus replication (19), and the absence of this protein was expected to inhibit production of infectious progeny (16). We therefore transfected 293T cells with nine protein expression plasmids as well as with those for the production of viral RNA segments that encoded all genes of the influenza WSN virus, excluding the NS gene (Fig. 1). Forty-eight hours after transfection, supernatants derived from transfected cells were used to infect MDCK cells. Prgeny virus was not produced from the VLP-infected cells.

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ruses by passaging the replication-incompetent VLPs three times in MDCK cells in three separate experiments (data not shown).

The efficiency of VLP generation and the extent of viral protein expression in VLP-infected cells were assessed by fixing cells at 16 h postinfection and examining them for expression of NP, HA, NA, and M1. The titer of the NS gene-deficient VLPs in the culture supernatant of 293T cells was approximately $10^4$ infectious VLPs/ml, measured by counting NP-expressing cells. By immunostaining, the NP protein was readily detected in the infected cells, but HA, NA, and M1 were not (Fig. 2, left panels).

Since NS1 stimulates the translation of viral mRNA (3, 6, 20), we reasoned that its absence might inhibit normal expression of viral proteins. Thus, to produce influenza VLPs expressing immunogenic viral proteins in cells, we next attempted to generate VLPs encoding NS1, but not NS2, which is encoded by spliced mRNA derived from the NS segment (12). To this end, we transfected 293T cells with pPolI-WSN-NS/H9004Splice (Fig. 3), which contained only the NS1 coding region, and the remaining RNA polymerase I constructs and protein expression plasmids. Although no progeny virus was produced from cells infected with the NS2-knockout VLPs (data not shown), they expressed the four viral proteins tested (i.e., NP, HA, NA, and M1) (Fig. 2, right panels). At 48 h posttransfection, the titer of these VLPs, measured by counting NP-positive cells, was approximately $10^5$ infectious VLPs/ml. These results establish the requirement for NS1 protein in translational regulation of viral mRNAs and its importance in viral replication.

Antibody responses of mice immunized with NS gene-deficient or NS2-knockout VLPs. To test the efficacy of replication-incompetent particles as a vaccine, we first performed a pilot experiment by intranasally inoculating mice with 16 HAU of the NS2-knockout VLPs once or twice at 2-week intervals. When immunized mice were challenged with 100 LD$_{50}$ of the wild-type WSN virus 2 weeks after the last vaccination, their survival rates were 25 or 75%, respectively. Based on these results, we intranasally inoculated mice with 16 HAU of the NS gene-deficient or NS2-knockout VLPs three times at 3-week intervals. IgG and IgA were measured in sera, trachea-lung washes, and nasal washes of immunized mice with an ELISA (Fig. 4). Both IgG and IgA production in nasal and trachea-lung washes was significantly higher in mice immunized with NS2-knockout VLPs than with NS gene-deficient VLPs. The IgA response was negligible in serum, regardless of the type of VLPs used for immunization, but IgG production was clearly higher in mice inoculated with the NS2-knockout VLPs. None of the sera from immunized mice had detectable hemagglutination inhibition antibodies. Neither the culture supernatant lacking VLPs (control) nor the inactivated WSN virus elicited a significant antibody response. Thus, antibody responses were induced more efficiently with the NS2-knockout VLPs that produced multiple proteins.

Protective efficacy of replication-incompetent VLPs. Mice immunized with either NS gene-deficient or NS2-knockout VLPs were challenged with 10 or 100 LD$_{50}$ of the wild-type WSN virus 1 month after the last vaccination. In contrast to the fate of control mice and mice receiving inactivated virus or NS gene-deficient VLPs, those immunized with the NS2-knockout
particles were protected against lethal challenge with WSN virus (Table 1). Eight of nine mice in the NS2-knockout group survived even when challenged 3 months after the last vaccination. Moreover, their body weights were not appreciably affected by virus challenge, in contrast to the other vaccination groups, whose weights decreased rapidly postchallenge (Fig. 5). We also determined the virus titers in the lungs of mice. Both control mice and mice immunized with inactivated virus or NS gene-deficient VLPs had more than \(10^7\) PFU in lung tissue after challenge with 10 or 100 LD\(_{50}\) of wild-type WSN virus. In contrast, mice immunized with NS2-knockout VLPs had greater than 100-fold lower titers in lungs after challenge with the same doses (Table 1). We conclude that the NS2-knockout VLPs can effectively protect mice against lethal influenza virus infection.

### DISCUSSION

We have evaluated two types of replication-incompetent influenza VLPs for their immunogenicity in a mouse model. Mice vaccinated intranasally with the NS2-knockout VLPs, but not those given NS gene-deficient VLPs, were protected against lethal infection by wild-type influenza virus. We attribute these results to the greater expression of viral proteins in host cells infected with the NS2-knockout construct, which likely stimulated an immune response sufficient to prevent infection of mucosal surfaces.

The NS2 protein mediates the nuclear transport of viral RNP (16, 24). We have demonstrated that the NS2-knockout VLPs did not produce infectious progeny virus (16), substantiating the importance of this protein in the viral life cycle. In contrast, influenza A virus can replicate in the absence of NS1 in interferon-deficient hosts (9). Although NS1 is shown to translationally regulate viral mRNAs in vitro (3, 6, 20), direct evidence for this effect in the context of influenza virus infection was lacking. Here we show that multiple viral proteins, except for NP, are poorly expressed in cells infected with NS gene-deficient VLPs, in contrast to their abundant expression in cells infected with the NS2-knockout VLPs (Fig. 2). Thus, the NS1 protein clearly enhances the expression of late viral proteins and therefore plays an important role in the life cycle of influenza A viruses. Consistent with our finding, Enami and Enami (7) have reported that translation of HA and M1 proteins was significantly reduced in host cells infected with an NS1 mutant lacking the C-terminal portion of the NS1 protein.

Influenza viruses have been used as carriers to express genes or portions of genes from entirely unrelated infectious agents (13, 21–24). They offer numerous advantages in this regard.
Not only do they stimulate strong cell-mediated and humoral immune responses, but they also afford a wide array of virion surface HA and NA proteins (i.e., 15 HA and 9 NA subtypes plus their epidemic variants), allowing repeated immunization of the same target population. Thus, the replication-incompetent influenza VLPs described here might prove useful as vaccine vectors, permitting expression of foreign proteins or immunogenic epitopes. This potential is especially appealing for vaccination against human immunodeficiency virus, foot-and-mouth disease virus, and other infections, where any reversion of the live virus to a wild type is absolutely unacceptable or where the efficacy of inactivated vaccines may be limited due to limited induction of mucosal immunity and cytotoxic T-lymphocyte responses.

In summary, we show that replication-incompetent NS2-knockout VLPs administered via the respiratory mucosal route can protect mice against otherwise lethal influenza virus challenge. Such particles efficiently elicited mucosal as well as systemic immune responses that were sufficient to reduce initial infection of mucosal surfaces, in contrast to inactivated influenza virus vaccines. The newfound ability to manipulate the viral genome should lead to the development of vaccines that can safely prevent diseases that continue to afflict us with regularity.

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