

In Vitro Analysis of Virus Particle Subpopulations in Candidate Live-Attenuated Influenza Vaccines Distinguishes Effective from Ineffective Vaccines[∇]

Philip I. Marcus,^{1*} John M. Ngunjiri,¹ Margaret J. Sekellick,¹ Leyi Wang,^{2,3,†} and Chang-Won Lee^{2,3}

Department of Molecular and Cell Biology and Center of Excellence for Vaccine Research, University of Connecticut, U-3125, Storrs, Connecticut 06269¹; Food Animal Health Research Program, Ohio Agricultural Research and Development Center, The Ohio State University, Wooster, Ohio 44691²; and Department of Veterinary Preventive Medicine, The Ohio State University, Columbus, Ohio 43210³

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Two effective (*vac*⁺) and two ineffective (*vac*⁻) candidate live-attenuated influenza vaccines (LAIVs) derived from naturally selected genetically stable variants of A/TK/OR/71-delNS1[1-124] (H7N3) that differed only in the length and kind of amino acid residues at the C terminus of the nonstructural NS1 protein were analyzed for their content of particle subpopulations. These subpopulations included total physical particles (measured as hemagglutinating particles [HAPs]) with their subsumed biologically active particles of infectious virus (plaque-forming particles [PFPs]) and different classes of noninfectious virus, namely, interferon-inducing particles (IFPs), noninfectious cell-killing particles (niCKPs), and defective interfering particles (DIPs). The *vac*⁺ variants were distinguished from the *vac*⁻ variants on the basis of their content of viral subpopulations by (i) the capacity to induce higher quantum yields of interferon (IFN), (ii) the generation of an unusual type of IFN-induction dose-response curve, (iii) the presence of IFPs that induce IFN more efficiently, (iv) reduced sensitivity to IFN action, and (v) elevated rates of PFP replication that resulted in larger plaques and higher PFP and HAP titers. These *in vitro* analyses provide a benchmark for the screening of candidate LAIVs and their potential as effective vaccines. Vaccine design may be improved by enhancement of attributes that are dominant in the effective (*vac*⁺) vaccines.

Live-attenuated vaccines are considered more effective than their inactive or single-component counterparts because they activate both the innate and adaptive immune systems and elicit responses to a broader range of antigens for longer periods of time (2, 10, 25, 28). Influenza virus variants with alterations in the reading frame of the nonstructural NS1 protein gene (delNS1), which express truncated NS1 proteins, characteristically induce enhanced yields of type I interferon (IFN) relative to the yields of their isogenic parental virus encoding full-length NS1 proteins (11, 13, 21, 33, 39). Many of these delNS1 variants have proved to be effective as live-attenuated influenza vaccines (LAIVs), providing protection against challenge virus in a broad range of species (33, 46), including chickens (39, 44). The IFN-inducing capacity of the virus is considered an important element in the effectiveness of LAIVs (33). In that context, influenza viruses are intrinsically sensitive to the antiviral action of IFN (31, 32, 36), although they may display a nongenetic-based transient resistance (36). In addition, IFN sensitizes cells to the initiation of apoptosis by viruses (42) and by double-stranded RNA (40), which may be spontaneously released in the course of influenza virus replication (14). Furthermore, IFN functions as an adjuvant to boost the adaptive immune response in mammals (3, 4, 11, 26,

41, 43, 46) and in chickens when administered perorally in the drinking water of influenza virus-infected birds (19). This raises the question: does the enhanced induction of IFN by delNS1 variants suffice to render an infectious influenza virus preparation sufficiently attenuated to function as an effective live vaccine? To address that question, we turned to a recent report that described the selection of several variants of influenza virus with a common backbone of A/TK/OR/71-SEPRL (Southeast Poultry Research Laboratory) that contained NS1 protein genes which were unusual in the length and nature of the amino acid residues at the C termini of the truncated NS1 proteins that they expressed because of the natural introduction of a frameshift and stop codon by the deletion in the NS1 protein gene (44). delNS1 variants were isolated from serial low-inoculum passages of TK/OR/71-delNS1[1-124] (H7N3) in eggs (44). Four of these genetically stable plaque-purified variants, each encoding a truncated NS1 protein of a particular length, were tested as a candidate LAIV in 2-week-old chickens. Two of the delNS1 variants were effective as live vaccines (double deletions [D-del] pc3 and pc4) (phenotypically *vac*⁺), and two were not (D-del pc1 and pc2) (phenotypically *vac*⁻) (44), despite only subtle differences in their encoded delNS1 proteins. Why were they phenotypically different?

The present study addresses this question by analyzing and comparing the different virus particles that constitute the subpopulations of these two effective (*vac*⁺) and two ineffective (*vac*⁻) live vaccine candidates. These analyses are based on recent reports in which noninfectious but biologically active particles (niBAPs) in subpopulations of influenza virus particles were defined and quantified (20, 21, 29). The study de-

* Corresponding author. Mailing address: Department of Molecular and Cell Biology, University of Connecticut, U-3125, 91 North Eagleville Rd., Storrs, CT 06269. Phone: (860) 486-4254. Fax: (860) 486-5193. E-mail: philip.marcus@uconn.edu.

† Present address: Department of Infectious Diseases, College of Veterinary Medicine, University of Georgia, Athens, GA.

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TABLE 1. Comparison of biologically active and hemagglutinating particles in allantoic fluid stocks of wild-type TK/OR/71(H7N3) and candidate delNS1 LAIV variants

rgTK/OR/71 designation ^a	No. of aa residues of NS1 protein (1-X...Y) ^b	Particle titer per ml (% of total BAPs ^c)					
		PFPs ^c	IFPs ^c	DIPs ^c	niCKPs ^c	BAPs ^{c,d}	HAPs ^{c,e}
Wild type	1–230	0.83 × 10 ⁸ (4.1)	4.2 × 10 ^{8f} (20.9)	12 × 10 ⁸ (59.7)	3.07 × 10 ⁸ (15.3)	20.1 × 10 ⁸	1,024 × 10 ⁸
D-del pc2 (<i>vac</i> ⁻) ^g	1–115...125	0.022 × 10 ⁸ (0.02)	7.2 × 10 ^{8f} (7.4)	89 × 10 ⁸ (91.7)	0.88 × 10 ⁸ (0.9)	97.1 × 10 ⁸	70 × 10 ⁸
D-del pc1 (<i>vac</i> ⁻) ^g	1–80...90	0.0059 × 10 ⁸ (0.006)	14 × 10 ^{8f} (13.4)	89 × 10 ⁸ (85.5)	1.14 × 10 ⁸ (1.1)	104.1 × 10 ⁸	384 × 10 ⁸
D-del pc4 (<i>vac</i> ⁺) ^g	1–91...93	0.90 × 10 ⁸ (0.4)	3.3 × 10 ^{8h} (1.5)	220 × 10 ⁸ (97.1)	2.30 × 10 ⁸ (1)	226.1 × 10 ⁸	1,024 × 10 ⁸
D-del pc3 (<i>vac</i> ⁺) ^g	1–69...86	0.36 × 10 ⁸ (0.2)	1.3 × 10 ^{8h} (0.74)	170 × 10 ⁸ (97.9)	1.91 × 10 ⁸ (1.1)	173.6 × 10 ⁸	448 × 10 ⁸

^a All viruses were generated using a plasmid-based reverse genetics system (44).

^b For example, with D-del pc2, (1–X) represents the N-terminal amino acid (aa) residues 1 to 115 found in the wild-type A/TK/OR/71-SEPRL, and (...Y) represents the C-terminal amino acid residues 116 to 125 which differ from the wild-type sequences due to the introduction of a frameshift and new stop codon in the deleted NS1 protein gene. For details, see Fig. 2 of reference 44.

^c Titers expressed as number of particles per milliliter of allantoic fluid. This includes PFPs (iCKPs) (20), IFPs (21), DIPs (20), niCKPs (20, 29), and HAPs (8, 20).

^d BAP titers ≈ the sum of the titers of PFPs, IFPs, DIPs, and niCKPs.

^e Estimate on the basis of the average number of physical particles in 1 hemagglutinating unit, i.e., 5 × 10⁶ (8, 20).

^f IFPs calculated from type *r* ≥ 1 IFN induction dose-response curves (17, 18; see Materials and Methods and Fig. 1).

^g Arranged in the order of increasing effectiveness as LAIV on the basis of the titer of antibodies against heterologous virus that they induced (44).

^h IFP calculated from type *r* = 2 IFN-induction dose-response curves (17, 18; see Materials and Methods and Fig. 1).

scribed in this report reveals several quantitative and qualitative differences between the particle subpopulations of the four candidate LAIVs, including the different types of IFN-induction dose-response curves, the quantum (maximum) yields (QY) of IFN induced, the efficacy of the interferon-inducing particles (IFPs), the replication efficiency of the virus, and the size of the plaques that they produced. Evidence is presented that the *in vitro* analysis of virus particle subpopulations may be useful to distinguish *vac*⁺ from *vac*⁻ LAIV candidates and provide a basis for identifying and enhancing the performance of particles with desirable phenotypes.

MATERIALS AND METHODS

Cell culture and media. Monolayers of primary chicken embryo cells (CECs) and chicken embryo kidney cells (CEKs) were prepared from cell suspensions of 9-day-old and 18-day-old embryos, respectively, obtained from Charles River SPAFAS, Inc., Storrs, CT. The cells were grown in attachment solution (AS; NCI medium plus 6% calf serum) (38) and incubated at 38.5°C. GMK-Vero and MDCK cells were grown in AS and Eagle’s minimal essential medium (plus 5% calf serum), respectively, and incubated at 37.5°C.

LAIV viruses. In a previous study, four LAIV virus variants (D-del pc1 to pc4) were generated naturally during passage of a stock of A/TK/OR/71-delNS1[1-124] (H7N3) virus in embryonated eggs (44). These viruses were confirmed to be genetically stable through plaque purification, followed by at least 5 passages in 10-day-old embryonated eggs. For the present study, these four variants and an isogenic virus encoding a full-size NS1[1-230] protein, used here and termed wild type, were reconstructed as already described using a reverse genetics system (44). The only differences among these five reconstituted viruses are the length and amino acid composition at the C terminus of the NS1 protein, as described previously (44) and shown in Table 1.

For simplicity, these virus variants are referred to as phenotypically *vac*⁻ (ineffective) or *vac*⁺ (effective) vaccines, on the basis of their capacity to induce an antibody response, prevent virus shedding, and elicit protection against the heterologous virus, A/CK/NJ/150383-7/02 (H7N2) (44). By these criteria, D-del pc1 and pc2 viruses were *vac*⁻ and D-del pc3 and pc4 were *vac*⁺. All stocks were stored at -80°C, thawed rapidly, and held on ice for use.

Interferon induction. Primary CECs were seeded at a high density to produce confluent monolayers after 24 h and aged for 10 days *in vitro* to developmentally mature the IFN system (5, 35, 38). These developmentally aged cells were used to test the IFN-inducing capacities of the viruses. The IFN induction protocol was described recently (20). Briefly, replicate monolayers of developmentally aged CECs (10⁷ cells/50 mm dish) were infected with increasing amounts (multiplicities) of the test virus and incubated at 40.5°C (optimal for IFN induction/production [38]) in medium without serum. The medium was harvested after 20 to 22 h, and the supernatant was assayed for acid-stable type I IFN, as described

earlier (38). A standard preparation of UV-irradiated avian reovirus (45) was used to assess the IFN-inducing capacity of each batch of CECs, and a standard of recombinant chicken alpha IFN (rChIFN-α) (37) was used to calibrate the sensitivity of each CEC preparation to the action of IFN (36, 38).

IFN detection and quantification. The titers of IFN were based on a 50% cytopathic effect (CPE₅₀) assay in 96-well trays of CECs following treatment with IFN for 24 h at 37.5°C, infection with vesicular stomatitis virus (VSV) strain HR, and incubation for 48 h, before determination of the 50% end point photometrically (34, 36, 38).

IFN-induction curves, QYs of IFN, titers of IFPs, and IFP efficiency. Full dose (amount of virus)-response (IFN yield) curves for IFN induction were generated from developmentally aged CECs, as reported previously (18). The type of IFN-induction curves observed and calculation of IFP titers were described in part previously (17, 18, 23). They were based on a Poisson distribution of IFP among the CEC population, $P_{(r)} = (e^{-m} m^r)/r!$ [where $P_{(r)}$ is the fraction of cells that get *r* IFPs when exposed to a multiplicity of *m*, *r* is the actual number of IFPs that enter the cell, *r*! is the factorial of *r*, *m* is the average number of IFPs in the virus suspension per cell in the monolayer, and *e* is the base of the natural logarithm ($e = 2.71828...$)], and the QY of IFN produced by cells in response to infection by IFP. Type *r* ≥ 1 curves best fit a class of cells in which a full yield of IFN is induced in any cell infected with 1 or more IFPs, i.e., $P_{(r \geq 1)} = (1 - e^{-m})$ QY, where e^{-m} is the fraction of cells that get zero IFPs. Type *r* = 2 curves best fit a model in which cells that receive 2 and only 2 IFPs respond by producing a QY of IFN, whereas cells infected with 1 or >2 IFPs induce little or no IFN, i.e., $P_{(r = 2)} = (e^{-m} m^2/2e^m)$ QY.

In a type *r* ≥ 1 curve, the dilution of virus that induces 0.63 of the QY of IFN contains, on average, a multiplicity of plaque-forming particles (PFPs) (m_{IFP}) of 1, from which the IFP titer can be calculated as described previously (18). In a type *r* = 2 curve, the dilution of virus that induces a QY of IFN occurs when 0.27 of the cells are infected with an average m_{IFP} of 2. Since virtually all of the virus binds to the cell monolayer during the attachment period and the number of IFN-inducing cells and the dilution of virus required to induce a QY of IFN are known, the IFP titer can be calculated (18).

Assays for PFPs, niCKPs, DIPs, and HAPs. The methods used to quantify subpopulations of PFPs, noninfectious cell-killing particles (niCKPs), defective interfering particles (DIPs), and hemagglutinating particles (HAPs) (in which these particles are subsumed) have been described previously (20, 21, 29). Briefly, PFPs were detected by plaque formation in CEKs (trypsin was not required) or Vero cells (trypsin was required). The number of niCKPs represents the difference between the total number of CKPs, determined by the clonogenic assay (29), and the number of infectious CKPs, where PFPs are infectious CKPs (iCKPs) (thus, total CKPs - iCKPs = niCKPs). DIPs were measured by the reduction in the PFP yield of a DIP-free helper virus (20, 27). The amount of HAPs was estimated on the basis of their capacity to agglutinate chicken red blood cells using a conversion factor (5 × 10⁶) averaged from data from several reports (8, 20) to provide a first approximation of the number of physical particles in 1 hemagglutinating unit. The HAP titers obtained in this way should

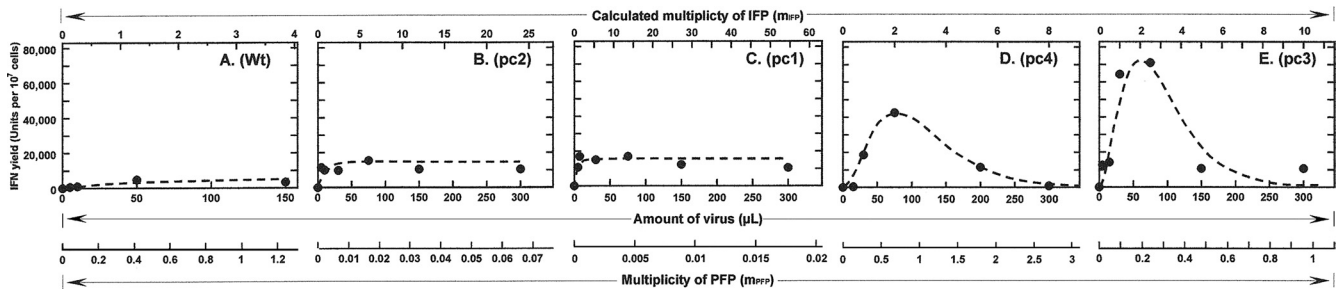


FIG. 1. IFN-induction dose (amount of virus)-response (IFN yield) curves generated by candidate LAIV delNS1 and wild-type (Wt) viruses in developmentally aged CECs at 40.5°C. These have been arranged in the order of increasing quantum (maximal) yields of IFN. Actual data points are represented by closed circles. The dashed lines represent the expected curves following the Poisson distribution of IFPs among cells in the monolayer, i.e., $P_{(r)} = (e^{-m}m^r)/r!$. (A to C) Type $r \geq 1$ curves, where $P_{(r \geq 1)} = (1 - e^{-m})QY$; (D and E) type $r = 2$ curves, where $P_{(r = 2)} = (e^2m^2/4e^m)QY$ (the various terms are defined in the text). Type $r \geq 1$ curves (A to C) show that a cell infected with 1 or more IFPs produces a QY of IFN. Type $r = 2$ curves (D and E) represent the class of cells that, when infected with 2 and only 2 IFPs, produces a QY of IFN; cells infected with 1 or more than 2 IFPs produce little or no IFN.

accurately reflect the relative titers of the different subpopulations, although they do not necessarily represent their sums.

Virus replication. Virus replication was determined over a 48-h period at 40.5°C, which is normothermic for chickens. Confluent monolayers containing 3.0×10^6 CEKs in 50-mm dishes were infected at an m_{PFP} of 0.03. The virus was suspended in 300 μ l of NCI medium and attached to cells for 1 h. Unattached virus was aspirated, and the cells were overlaid with fresh serum-free medium. The medium was harvested at appropriate times, cleared of cell debris by centrifugation, and assayed for PFP and HAP activities.

RESULTS

IFN-induction dose-response curves, IFN QY, IFP titers, and IFP efficiency. IFN-induction dose-response curves (17, 18) were generated by exposing developmentally aged primary CECs (35, 36) to increasing amounts of the four candidate LAIVs derived from TK/OR/71-delNS1 variants that expressed truncated NS1 proteins of different lengths or an isogenic virus that expressed the full-size NS1[1-230] protein (wild type) (Table 1) (44). Figure 1 shows representative responses of the wild type and each of these variants arranged by increasing QY. Two qualitatively different types of IFN-induction dose-response curves were observed. Figure 1A to C shows that wild-type and delNS1 vac^- variants pc1 and pc2 generated IFN-induction dose-response curves that best fit a model based on a Poisson distribution of IFPs in which the fraction of cells infected with ≥ 1 IFP induces a QY of IFN, i.e., the maximum amount of IFN that a cell is capable of producing when ex-

posed to that particular IFP. The type $r \geq 1$ dose-response curve is the most commonly observed with influenza virus and many types of viruses tested (17, 18, 21). In contrast, Fig. 1D and E shows that delNS1 vac^+ variants pc3 and pc4 generated dose-response curves that best fit a model in which cells infected with 2 and only 2 IFPs induce a QY of IFN and cells infected with 1 or >2 IFPs produce little or no IFN. This type $r = 2$ IFN-induction dose-response curve was described in theoretical terms years ago (16) but was observed experimentally for the first time in the present study. Importantly, the type $r = 2$ curves were observed in the same batches of aged CECs in which wild-type, pc1, and pc2 viruses induced the more common type $r \geq 1$ curves. The QY of IFN observed at the plateau of the type $r \geq 1$ curve for wild-type virus averaged 4,000 U/10⁷ CECs, whereas the $r \geq 1$ curves for vac^- pc1 and pc2 were similar, averaging 15,500 U/10⁷ CECs. The QY of IFN for vac^+ variants pc3 and pc4 was observed at the peak of a type $r = 2$ curve (see Materials and Methods) and averaged 57,250 U/10⁷ CECs (Fig. 1; Table 2).

The IFPs averaged for the two vac^- variants, though present in 4.6-fold excess compared to the amount of IFPs in the vac^+ variants (Table 1), were less efficient in inducing IFN on an IFP-per-cell basis. The IFPs in the wild-type subpopulation were even less efficient (Table 2).

Sensitivity of LAIV delNS1 variants to action of interferon. The sensitivities of the four candidate LAIV delNS1 variants

TABLE 2. Comparison of quantum (maximal) yields of IFN and relative efficiency of IFN induction

rgTK/OR/71 designation ^a	No. of aa residues of NS1 protein (1-X...Y) ^b	QY of IFN (U/1.0 \times 10 ⁷ cells)	No. of IFPs inducing the QY of IFN	IFN yield (U/10 ⁶ PFP)	Relative IFN induction efficiency ^c
Wild type	1-230	4,000	1.0×10^{7d}	400	1.0
D-del pc2 (vac^-) ^e	1-115...125	14,000	1.0×10^{7d}	1,400	3.5
D-del pc1 (vac^-) ^e	1-80...90	17,000	1.0×10^{7d}	1,700	4.3
D-del pc4 (vac^+) ^e	1-91...93	42,500	5.4×10^{6f}	8,000	20.0
D-del pc3 (vac^+) ^e	1-69...86	72,000	5.4×10^{6f}	13,300	33.3

^a See footnote a in Table 1.

^b See footnote b in Table 1.

^c Calculated relative to the IFN U/10⁶ IFP value for the wild-type virus.

^d The lowest number of IFPs required to infect all (i.e., 1.0×10^7) cells in order to induce a QY of IFN for type $r \geq 1$ dose-response curves.

^e Arranged in the order of increasing effectiveness as LAIV on the basis of the titer of antibodies against heterologous virus that they induced (44).

^f The number of IFPs infecting 27% (i.e., 2.7×10^6) of the cells in order to induce a QY of IFN for type $r = 2$ dose-response curves. Each of these cells is infected with 2 IFPs. The total number of IFPs contributing to the QY equals 5.4×10^6 . See Materials and Methods for calculations.

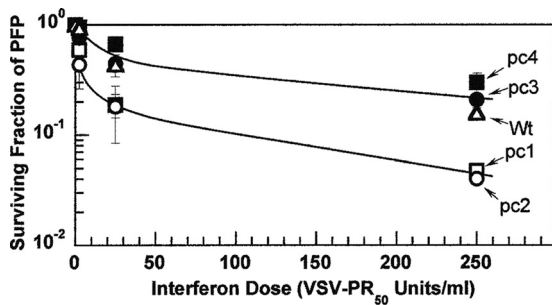


FIG. 2. Sensitivity of candidate LAIV delNS1 and wild-type (Wt) viruses to the action of IFN. Following the exposure of 1-day-old monolayers of primary CEKs to rChIFN- α for 24 h at 38.5°C, the cell monolayers were challenged with PFPs of the respective viruses. An overlay of agarose in medium lacking serum or trypsin was incubated for 3 days before the surviving fraction of PFPs was determined. All viruses revealed biphasic survival curves, with the highest sensitivity occurring at low doses of IFN and a more resistant fraction being found at higher doses. The *vac*⁺ pc3 and pc4 variants, along with the wild-type virus, were experimentally indistinguishable (upper curve) and were less sensitive than the *vac*⁻ pc1 and pc2 variants (lower curve). Error bars represent the standard deviations of two determinations. PR₅₀, the amount of IFN that reduces the PFP titer by 50%.

and the wild type to the action of IFN were determined by exposing monolayers of primary CEKs to increasing doses of rChIFN- α (37) and measuring PFP survival at each dose, as was described earlier for other isolates of influenza virus (6, 36). Figure 2 shows that a single curve accommodates the data points for the two *vac*⁻ variants, pc1 and pc2 (lower curve), and that the data points for the wild-type virus and the two *vac*⁺ variants, pc3 and pc4, can also be accommodated by a single curve (upper curve). From the initial slopes of these curves at low doses of IFN, variants pc1 and pc2 appear to be about 4 times more sensitive to the action of IFN than pc3, pc4, and wild-type virus. At higher doses of IFN, a comparison of the slopes of the PFP survival curves and extrapolation to the ordinate revealed that both sets of viruses contained a large fraction of PFPs that appear to be more refractory to the action of IFN. These biphasic survival curves resemble the triphasic curves observed in two earlier studies in which the resistance to high doses of IFN of a high-pathogenicity virus proved to be transient and was lost upon passage of the virus (36) and in the two original nonisogenic low-pathogenicity

viruses, TK/OR/71-NS1[1-230] (H7N3) and the related virus delNS1[1-124] (6).

DIP titers. Standard PFP yield reduction assays (27), as reported earlier (20), were used to assess the DIP content of the four candidate LAIV variants and the wild type. The initial exponential rate of loss of PFP yield at low multiplicities of DIPs was used to calculate the titers of DIPs, as described previously (20). The four candidate LAIVs contained unusually high titers of DIPs, with the average for *vac*⁺ variants pc3 and pc4 being marginally higher (2.2-fold) than that for the *vac*⁻ variants pc1 and pc2 (Table 1). In contrast, the DIP titer of the wild-type variant was about 12-fold less, similar to the titers observed for several low-pathogenicity type A influenza viruses (LPAIs), including the original TK/OR/71-SEPRV virus that had been subjected to three serial high-multiplicity passages (20).

Titers of infectious and niCKPs. The clonogenic assay was used to determine the total CKP titer, as described previously (29). The titer of niCKPs represents the difference between the total CKP population and the infectious CKP population (iCKPs = PFPs); i.e., the number of niCKPs is equal to the number of CKPs minus the number of PFPs (29). Table 1 shows that the niCKP titers of all four delNS1 variants and the wild type were relatively high, with the average of the titers of *vac*⁻ variants pc1 and pc2 being only 2-fold less than that of *vac*⁺ variants pc3 and pc4 and the wild-type virus. Further analysis revealed that the niCKP/PFP ratios of *vac*⁻ pc1 and pc2 were at least 1 order of magnitude higher than those of *vac*⁺ pc3 and pc4 and the wild-type virus (Table 3). The high niCKP/PFP ratios indicate that virtually all cell killing by the *vac*⁻ variants was due to niCKPs, as was the majority for the *vac*⁺ variants and wild-type virus. Thus, the extensive cytopathic effects noted early upon infection of cells with the delNS1 variants were due almost exclusively to the onset of apoptosis initiated by niCKPs and not PFPs, as was observed earlier with the original TK/OR/71-delNS1[1-124] virus (29).

Comparison of PFP and HAP titers, growth rates, and plaque morphology of *vac*⁻ and *vac*⁺ delNS1 variants. Table 1 shows the PFP titers of the four candidate LAIVs used in the vaccine study (44), along with the titer of the isogenic wild type. The average PFP titer in the allantoic fluid of the two *vac*⁻ variants, pc1 and pc2, was low (1.4×10^6 /ml) compared to that of the two *vac*⁺ variants, pc3 and pc4 (63×10^6 /ml)

TABLE 3. Comparison of niBAP subpopulations with infectious particles (PFPs) and total physical particles (HAPs)

rgTK/OR/71 designation ^a	No. of aa residues of NS1 protein (1-X...Y) ^b	Ratio of niBAPs/PFPs			Estimated ratio ^d				
		IFPs/PFPs ^c	DIPs/PFPs ^c	niCKPs/PFPs ^c	HAPs/PFPs	HAPs/IFPs	HAPs/DIPs	HAPs/niCKPs	HAPs/BAPs
Wild type	1-230	5	14	4	1,230	240	85	330	51
D-del pc2 (<i>vac</i> ⁻) ^e	1-115...125	330	4,040	40	3,200	10	0.8	80	0.7
D-del pc1 (<i>vac</i> ⁻) ^e	1-80...90	2,370	15,080	193	65,100	27	4.3	340	3.7
D-del pc4 (<i>vac</i> ⁺) ^e	1-91...93	4	240	3	1,140	310	4.6	440	4.5
D-del pc3 (<i>vac</i> ⁺) ^e	1-69...86	4	470	5	1,240	340	2.6	230	2.6

^a See footnote a in Table 1.

^b See footnote b in Table 1.

^c Based on the titers in Table 1. The ratios are rounded to the nearest whole number.

^d Estimated on the basis of the average number of physical particles in 1 hemagglutinating unit, i.e., 5×10^6 (8, 20); niBAPs \approx sum of IFPs, DIPs, and niCKPs. Total BAPs \approx niBAPs plus PFPs.

^e Arranged in the order of effectiveness as LAIV on the basis of the titer of antibodies they induced against a heterologous virus (44).

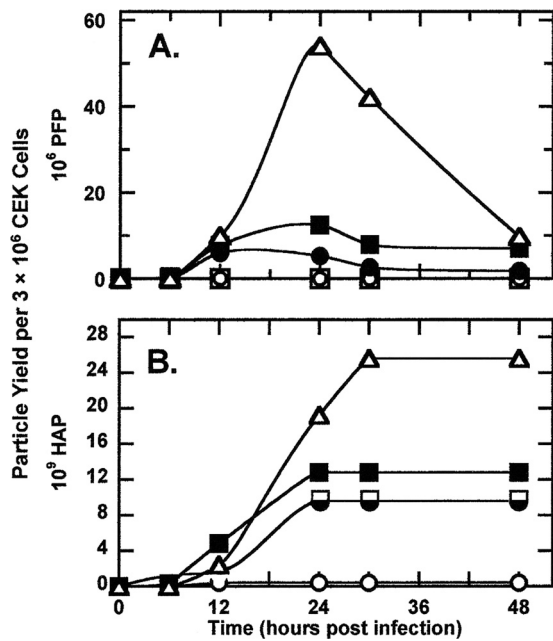


FIG. 3. Growth curves of candidate LAIV delNS1 and wild-type viruses in CEK cells at 40.5°C. Confluent monolayers of primary CEKs were infected with each virus at an m_{PFP} of 0.03, and serum-free NCI medium was added. Replicate dishes were harvested following incubation for the times indicated, and samples were clarified of cell debris and then assayed for PFP (A) and HAP (B) activity as described in Materials and Methods. Symbols: Δ , wild type; \circ , pc2; \square , pc1; \blacksquare , pc4; \bullet , pc3.

(Table 1). The ratio of the PFP titer for vac^+ /PFP titer for vac^- was 45. The PFP titer of the wild type was $83 \times 10^6/\text{ml}$. The low PFP titers of the vac^- variants could not be attributed to the accompanying high DIP content because the vac^+ variants, with their much higher PFP titers, had even more DIPs in the allantoic fluid (Table 1). Consequently, we compared the growth rates of the different delNS1 variants and the wild type (Fig. 3). Confluent monolayers of primary CEKs were infected at 40.5°C, which is normothermic for chickens and the optimal temperature for IFN production in chicken cells (38). CEKs do not require trypsin in the overlay to maintain plaque formation, and as observed from the abundant growth of wild-type virus in the kidneys of 1-day-old chicks (1.33×10^9 50% embryo lethal doses/g of tissue) (6), CEKs are excellent host cells. The highest efficiency of plaque formation was obtained when

the cells were not allowed to age *in vitro* (35, 38). In Fig. 3A, the yield of PFPs per 3×10^6 cells is reported as a function of time. The rapid decline in PFP titer once peak yields are reached is attributed to the cessation of new infectious particle formation and the relatively short half-life (≈ 7 h) of PFPs at 40.5°C (data not shown). Both vac^- variants produced enough infectious particles in CEKs to sustain the formation of small plaques (Fig. 4). In contrast, the vac^+ variants consistently formed larger plaques, though not as large as the plaques formed by the wild type. Figure 4 shows the plaque size and morphology characteristic of each of the variants as they appeared after incubation at 37.5°C for 3 days in CEKs. Although the appearance of plaques means that the yield of PFP per CEK is high enough to sustain plaque formation, the low average yield of PFPs per cell suggests that not all cells in a primary CEK monolayer may serve as hosts. This view is borne out by the intact fibroblast-like cells observed within a plaque and the increase in their number with the age of the monolayer. The small plaques observed for pc2 and pc1 viruses and at a low frequency for pc3 viruses do not appear to reflect the action of IFN, since the number and size of the plaques did not change when 2-aminopurine (30 mM), which is known to block the induction of IFN in CEKs (24), was present in the agarose overlay.

Figure 3B shows that the average peak number of HAPs produced per CEK as a function of time at 40.5°C was in the order wild type > delNS1 pc4 > pc3 = pc1 \gg pc2. This order is essentially the same as that observed for the egg-derived stocks of these candidate LAIVs shown in Table 1. Once they were produced, the HAPs were stable (Fig. 3B). D-del pc2 was the least efficient in producing HAP. Table 3 shows that relative to wild-type virus, a higher fraction of the HAPs in populations of D-del variants is assembled into BAPs.

DISCUSSION

Four candidate LAIVs with different deletions in the NS1 protein gene were naturally selected from serial passages of A/TK/OR/71-delNS1[1-124] in embryonated eggs (44). These deletions in the NS1 protein gene were stable when they were rescued in reassortant viruses (44). Two of the delNS1 variants functioned as effective (vac^+) LAIVs, as reflected by the high titer of antibodies induced against challenge with a heterologous virus and a significant reduction in virus shedding. The other two candidate LAIVs were ineffective as vaccines (vac^-) (44). The *in vitro* analyses of these four delNS1 variants for

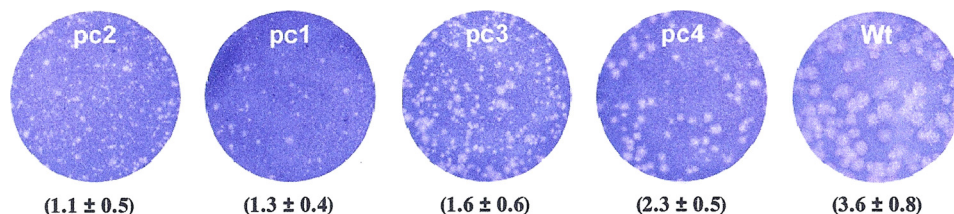


FIG. 4. Characterization of plaque phenotypes of candidate LAIV-delNS1 and wild-type (Wt) viruses. Monolayers of primary CEK cells were infected with each virus and incubated at 37.5°C for 3 days, fixed with formalin, and stained with Giemsa. Plaque sizes or numbers did not change when 2-aminopurine, an inhibitor of IFN induction in chicken embryonic cells (24), was included in the overlay, indicating that any differences in plaque size were not attributable to the IFN system (24). The plaque plates have been arranged in the order of increasing diameter (mean \pm standard deviation mm; $n = 55$).

their content and the activities of their BAPs, namely, PFPs, IFPs, DIPs, and niCKPs, along with the activity of the HAPs (20, 21, 22, 29), show that they can be used to distinguish vac^+ from vac^- candidate LAIVs. Thus, the appropriate *in vitro* tests can be used to screen large numbers of potential LAIVs and provide insight into what viral phenotypes should be enhanced in a search for a more effective vaccine.

The composition of BAPs in subpopulations of influenza virus (20, 21, 29) was measured *in vitro* to determine whether the phenotypes that they express can be used to distinguish vac^+ from vac^- LAIVs. Data presented herein define several parameters measured *in vitro* that differentiate vac^+ from vac^- LAIVs. Prominent are three parameters associated with the IFN system: (i) the capacity to induce IFN, as measured by the quantum yield from full IFN-induction dose-response curves; (ii) the regulation of IFN induction, as revealed by the nature of the type of IFN-induction curve; and (iii) the efficiency of the IFPs. The order of decreasing expression of these parameters is ranked $pc3 > pc4 \gg pc1 \approx pc2 \gg$ wild type, placing the two vac^+ variants first, the two vac^- variants next, and the wild-type virus a distant last (Fig. 1; Table 2).

In a seemingly anomalous situation, the IFP subpopulation in the vac^- variants was, on average, 4.6 times greater than that of the vac^+ variants (Table 1), yet the relative efficiency of IFN induction by the IFPs of the vac^+ variants was, on average, 6.8 times that of the vac^- variants (Table 2). Furthermore, relative to wild-type virus, the efficiency of IFN induction by the D-del variants increases in the same order as the QY of IFN (Fig. 1, Table 2). Notably, the unusual type $r = 2$ nature of the IFN-induction dose-response curves of the two vac^+ variants differed markedly from the usual type $r \geq 1$ nature of the curves that characterized the two vac^- variants and the wild type (Fig. 1), indicating that IFN induction by vac^+ variants was highly downregulated at an m_{IFP} of >2 .

The data also demonstrate that the vac^+ variants are less sensitive to the action of IFN than the vac^- variants (Fig. 2), suggesting that the vac^+ variants, with their increased capacity to induce IFN, may persist longer in the vaccinated host, thereby continuing to induce IFN and prolong the time that it can function as a natural adjuvant to enhance the production of antibody (4, 11, 19) and to help with clearance of the virus. In addition, there is a distinct replication advantage of the vac^+ variants over the vac^- variants (Fig. 3), with the former displaying on average a 45-fold greater PFP titer than the latter in allantoic fluid (Table 1). These differences in replication are reflected in the larger size of the plaques which develop from vac^+ variants in primary CEKs (Fig. 4).

The presence and sizes of the subpopulations of DIPs and niCKPs do not of themselves help to distinguish vac^+ and vac^- variants. However, the contribution of these subpopulations to the vac^+ phenotype has not been ruled out. For example, the unusually high titers of DIPs, which make up over 90% of the BAP population (Table 1), may exert a protective effect through the action of blocking virus replication (9, 20, 27) independent of the virus subtype (9), making them invaluable additions to the LAIVs. The demonstration that DIPs may render virulent avian influenza virus avirulent (1, 7) adds credence to a possible ameliorating role of DIPs in LAIVs. Whether the substantial numbers of niCKPs in a vac^+ LAIV contribute to survival of the host because of the widespread

destruction of cells through apoptosis (29) is not known. However, it is clear that DIPs do not function as niCKPs, nor do they block their action (see Fig. 5 in reference 20).

Quantitative estimates of the HAP population in which all of the above-mentioned BAPs are subsumed were based on published data which compare by various means the numbers of physical particles of influenza virus at the end point of a standardized hemagglutination (HA) test (8, 20). The conversion factor used herein may not be correct in absolute numbers because it has been applied to the native unpurified virus suspended in allantoic fluid used to vaccinate birds (44). However, at the least, the HAP titers shown here should be constant in relative terms for all of the populations of virus described. Although 96% of the HA activity of these allantoic fluid preparations is found in the pellet upon high-speed centrifugation (data not shown), indicative of the absence of soluble HA, it is possible that a small portion of the HA activity resides in the apoptosis-induced cell debris in the size range of influenza virus particles. Nonetheless, it is clear that the sum of the niBAP titers constitutes a significant fraction of the total HAPs in excess of that accounted for by PFPs. Does each BAP represent a particle of a distinct subpopulation, or do some BAP populations display overlapping phenotypes? The answer to this question requires further study. However, data reported thus far, including those obtained by UV target analysis, for the various phenotypes show that BAPs vary in the minimum gene requirements for their expression (20). For some BAPs, like DIPs, it is clear that the vast majority of the particles do not function as niCKPs, nor do they interfere with cell killing (20).

Current studies show that IFN-induction-suppressing particles (ISPs) (16, 21) have the potential to act as IFPs following truncation of the NS1 protein gene (a small gene target) or exposure to low doses of UV radiation (a large gene target) (15, 21). The titers of the DIP subpopulations in the 4 stocks of candidate LAIVs were unusually high, averaging 11.8-fold higher than the titer for the isogenic wild-type virus (Table 1) and 9.9-fold higher than the titer for the original TK/OR/71-delNS1[1-124] virus serially passed 3 times in MDCK cells to maximize the content of DIPs (20). The DIPs of the D-delNS1 variants constituted from 86 to 98% of the BAP population, making them the predominant BAPs (Table 1). On that basis, it is tempting to infer a causal relationship between the truncated NS1 proteins common to the D-del variants and the propensity to generate DIPs, perhaps akin to that reported for the NS2/NEP protein (30) and in keeping with the generally recognized role of the NS1 protein in replication.

Regardless of the absolute number of physical particles measured as HAPs or the extent to which the activities of BAPs represent discrete or overlapping functions, the biological assays used to define BAP subpopulations of LAIVs *in vitro* can collectively distinguish vac^+ from vac^- populations. On the basis of the findings of the present study, the successful vac^+ LAIV might be expected to display collectively the following characteristics: (i) highly efficient IFPs, (ii) induction of high QYs of IFN, (iii) generation of IFN-induction dose-response curves that reflect tightly regulated IFN induction as a function of m_{IFP} , (iv) enhanced resistance to IFN action, (v) robust levels of virus replication and yields of PFPs, (vi) production of

relatively large plaques, and (vii) high titers of DIPs and niCKPs relative to PFP titers.

Because IFN plays a dominant role in successful LAIVs (11, 12, 13, 33), ameliorates pathogenesis (6), and functions as a natural adjuvant (3, 4, 11, 19, 41, 43), it may be possible to augment the levels of IFN induced endogenously by the vaccine through the exogenous delivery of IFN in drinking water, a procedure used in chickens that allowed the precise control of IFN doses and that is both tactically and economically feasible (19). Peroral use of IFN has also been shown to boost the adaptive immune system in other hosts (26, 41, 43, 46). Low doses of UV radiation enhance even further the IFN-inducing capacity of influenza viruses that express truncated NS1 protein (21) and hence might serve as a second means of augmenting the IFN levels in vaccinated hosts. Assays of BAPs and the attributes of the subpopulations that they generate provide a fingerprint of effective LAIVs, and more will be tested to determine the extent to which their detection and interaction with host cells *in vitro* is predictive of the *vac*⁺/*vac*⁻ phenotypes.

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REFERENCES

1. Bean, W. J., Y. Kawaoka, J. M. Wood, J. E. Pearson, and R. G. Webster. 1985. Characterization of virulent and avirulent A/chicken/Pennsylvania/83 influenza viruses: potential role of defective interfering RNAs in nature. *J. Virol.* **54**:151-160.
2. Belshe, R. B., K. M. Edwards, T. Vesikari, S. V. Black, R. E. Walker, M. Hultquist, G. Kemble, and E. M. Connor. 2007. Live attenuated versus inactivated influenza vaccine in infants and young children. CAIV-T Comparative Efficacy Study Group. *N. Engl. J. Med.* **356**:685-696.
3. Bracci, L., I. Canini, S. Puzelli, P. Sestili, M. Vanditti, M. Spada, I. Donatelli, F. Belardelli, and E. Proietti. 2005. Type I IFN is a powerful mucosal adjuvant for a selective intranasal vaccination against influenza virus in mice and affects antigen capture at mucosal level. *Vaccine* **23**:2994-3004.
4. Bracci, L., I. Canini, M. Vanditti, M. Spada, S. Puzelli, I. Donatelli, F. Belardelli, and E. Proietti. 2006. Type I IFN as a vaccine adjuvant for both systemic and mucosal vaccination against influenza virus. *Vaccine* **24**(Suppl. 2):56-57.
5. Carver, D. H., and P. I. Marcus. 1967. Enhanced interferon production from chick embryo cells aged *in vitro*. *Virology* **32**:247-257.
6. Cauten, A. N., D. E. Swayne, M. J. Sekellick, P. I. Marcus, and D. L. Suarez. 2007. Amelioration of influenza virus pathogenesis in chickens is attributed to the enhanced interferon-inducing capacity of a virus with a truncated NS1 gene. *J. Virol.* **81**:1838-1847.
7. Chambers, T. M., and R. G. Webster. 1987. Defective interfering virus associated with A/chicken/Pennsylvania/83 influenza virus. *J. Virol.* **61**:1517-1523.
8. Desselberger, U. 1975. Relation of virus particle counts to the hemagglutinating activity of influenza virus suspensions measured by the HA pattern test and by the photometric HCU method. *Arch. Virol.* **49**:365-372.
9. Dimmock, N. J., E. W. Rainsford, P. E. Scott, and A. C. Marriott. 2008. Influenza virus protecting RNA: an effective prophylactic and therapeutic antiviral. *J. Virol.* **82**:8570-8578.
10. Gorse, G. J., M. J. Campbell, E. E. Otto, D. C. Powers, G. W. Chambers, and F. K. Newman. 1995. Increased anti-influenza A virus cytotoxic T cell activity following vaccination of the chronically ill elderly with live attenuated or inactivated influenza virus vaccine. *J. Infect. Dis.* **172**:1-10.
11. Hai, R., L. Martínez-Sobrido, K. A. Fraser, J. Ayllon, A. García-Sastre, and P. Palese. 2008. Influenza B virus NS1-truncated mutants: live-attenuated vaccine approach. *J. Virol.* **82**:10580-10590.
12. Kochs, G., I. Koerner, L. Theil, S. Kothlow, B. Kaspers, N. Ruggli, A. Summerfield, J. Pavlovic, J. Stech, and P. Staeheli. 2007. Properties of H7N7 influenza A virus strain SC35M lacking interferon antagonist NS1 in mice and chickens. *J. Gen. Virol.* **88**:1403-1409.
13. Kochs, G., L. Martínez-Sobrido, S. Lienenklaus, S. Weiss, A. García-Sastre, and P. Staeheli. 2009. Strong interferon-inducing capacity of a highly virulent variant of influenza A virus strain PR8 with deletions in the NS1 gene. *J. Gen. Virol.* **90**:2990-2994.
14. Majde, J. A., Z. Guha-Thakurta, S. Chen, S. Bredow, and J. M. Krueger. 1998. Spontaneous release of stable viral double-stranded RNA into the extracellular medium by influenza virus-infected MDCK endothelial cells: implications for the viral acute-phase response. *Arch. Virol.* **143**:2371-2380.
15. Malinoski, C. P., and P. I. Marcus. 2010. Influenza virus: can a single particle express two different phenotypes—IFN induction, or suppression of IFN induction?, p. 280. Abstr. 29th Annu. Meet. Am. Soc. Virol.
16. Marcus, P. I. 1982. Interferon induction by viruses. IX. Antagonistic activities of virus particles modulate interferon production. *J. Interferon Res.* **2**:511-518.
17. Marcus, P. I. 1983. Interferon induction by viruses: one molecule of dsRNA as the threshold for interferon induction, p. 115-180. *In* I. Gresser (ed.), *Interferon 5*. Academic Press, New York, NY.
18. Marcus, P. I. 1986. Interferon induction dose-response curves. *Methods Enzymol.* **119**:106-114.
19. Marcus, P. I., T. Girshick, L. van der Heide, and M. J. Sekellick. 2007. Super-sentinel chickens and detection of low-pathogenicity influenza virus. *Emerg. Infect. Dis.* **13**:1608-1610.
20. Marcus, P. I., J. M. Ngunjiri, and M. J. Sekellick. 2009. Dynamics of biologically active subpopulations in influenza virus: plaque-forming, noninfectious cell-killing, and defective-interfering particles. *J. Virol.* **83**:8122-8130.
21. Marcus, P. I., J. M. Rojek, and M. J. Sekellick. 2005. Interferon induction and/or production and its suppression in influenza viruses. *J. Virol.* **79**:2880-2890.
22. Marcus, P. I., and M. J. Sekellick. 1976. Cell killing by viruses. III. The interferon system and the inhibition of cell killing by vesicular stomatitis virus. *Virology* **69**:378-393.
23. Marcus, P. I., and M. J. Sekellick. 1977. Defective-interfering particles with covalently linked [\pm] RNA induce interferon. *Nature* **266**:815-819.
24. Marcus, P. I., and M. J. Sekellick. 1988. Interferon induction by viruses. XVI. 2-Aminopurine blocks selectively and reversibly an early stage in interferon induction. *J. Gen. Virol.* **69**:1637-1645.
25. Mueller, S. N., W. A. Langley, E. Carnero, A. García-Sastre, and R. Ahmed. 2010. Immunization with live-attenuated influenza viruses expressing altered NS1 proteins results in potent and protective memory CD8⁺ T cells responses. *J. Virol.* **84**:1847-1855.
26. Nagao, Y., K. Yamashiro, N. Hara, Y. Horisawa, K. Kato, and A. Uemura. 1998. Oral-mucosal administration of IFN- α potentiates immune response in mice. *J. Interferon Cytokine Res.* **18**:661-666.
27. Nayak, D. P., T. M. Chambers, and R. K. Akkina. 1985. Defective-interfering (DI) RNAs of influenza viruses: origin, structure, expression, and interference. *Curr. Top. Microbiol. Immunol.* **114**:103-151.
28. Nelson, K. M., B. R. Schram, M. W. McGregor, A. S. Sheoran, C. W. Olsen, and D. P. Lunn. 1998. Local and systemic isotype-specific antibody responses to equine influenza virus infection versus conventional vaccination. *Vaccine* **16**:1306-1313.
29. Ngunjiri, J. M., M. J. Sekellick, and P. I. Marcus. 2008. Clonogenic assay of type A influenza viruses reveals noninfectious cell-killing (apoptosis-inducing) particles. *J. Virol.* **82**:2673-2680.
30. Odagiri, T., K. Tominaga, K. Tobita, and S. Ohta. 1994. An amino acid change in the non-structural NS2 protein of an influenza A virus is responsible for the generation of defective interfering (DI) particles by amplifying DI RNAs and suppressing complementary RNA synthesis. *J. Gen. Virol.* **75**:43-53.
31. Portnoy, J., and T. C. Merigan. 1971. The effect of interferon and interferon inducers on avian influenza virus. *J. Infect. Dis.* **124**:545-552.
32. Richman, D. D., B. R. Murphy, S. Baron, and C. Uhlenhof. 1976. Three strains of influenza A virus (H3N2): interferon sensitivity *in vitro* and interferon production in volunteers. *J. Clin. Microbiol.* **3**:223-226.
33. Richt, J. A., and A. García-Sastre. 2009. Attenuated influenza virus vaccines with modified NS1 proteins. *Curr. Top. Microbiol. Immunol.* **333**:177-195.
34. Rubinstein, S., P. C. Familletti, and S. Pestka. 1981. Convenient assay for interferons. *J. Virol.* **37**:755-758.
35. Sekellick, M. J., W. J. Biggers, and P. I. Marcus. 1990. Development of the interferon system. I. In chicken cells development *in ovo* continues on time *in vitro*. *In Vitro Cell. Dev. Biol.* **26**:997-1003.
36. Sekellick, M. J., S. A. Carra, A. Bowman, D. A. Hopkins, and P. I. Marcus. 2000. Transient resistance of influenza virus to interferon action attributed to random multiple packaging and activity of NS genes. *J. Interferon Cytokine Res.* **20**:963-970.
37. Sekellick, M. J., A. F. Ferrandino, D. A. Hopkins, and P. I. Marcus. 1994. Chicken interferon gene: cloning, expression, and analysis. *J. Interferon Res.* **14**:71-79.
38. Sekellick, M. J., and P. I. Marcus. 1986. Induction of high titer chicken interferon. *Methods Enzymol.* **119**:115-125.
39. Steel, J., A. C. Lowen, L. Pena, M. Angel, A. Solórzano, R. Albrecht, D. R. Perez, A. Garcia-Sastre, and P. Palese. 2009. Live attenuated influenza

- viruses containing NS1 truncations as vaccine candidates against H5N1 highly pathogenic avian influenza. *J. Virol.* **83**:1742–1753.
40. **Stewart, W. E., II, E. De Clerq, and P. De Somer.** 1973. Specificity of interferon-induced enhancement of toxicity for double-stranded ribonucleic acids. *J. Gen. Virol.* **18**:237–246.
 41. **Takayama, S., K. Iwaki, Y. Nishida, M. Tanaka, M. Fujii, K. Ohashi, M. Ikeda, and M. Kurimoto.** 1999. Effects of oral administration of interferon- α on antibody production in mice with induced tolerance. *J. Interferon Cytokine Res.* **19**:895–900.
 42. **Tanaka, N., M. Sato, M. S. Lamphier, H. Nozawa, E. Oda, S. Nogucho, R. E. Schreiber, Y. Tsujimoto, and T. Taniguchi.** 1998. Type I interferons are essential mediators of apoptotic death in virally infected cells. *Genes Cells* **3**:29–37.
 43. **Tough, D. F., P. Borrow, and J. Sprent.** 1996. Induction of bystander T cell proliferation by viruses and type I interferon *in vivo*. *Science* **272**:1947–1950.
 44. **Wang, L., D. L. Suarez, M. Pantin-Jackwood, M. Mibayashi, A. García-Sastre, Y. M. Saif, and C.-W. Lee.** 2008. Characterization of influenza virus variants with different sizes of the non-structural (NS) genes and their potential as a live influenza vaccine in poultry. *Vaccine* **26**:3580–3586.
 45. **Winship, T. R., and P. I. Marcus.** 1980. Interferon induction by viruses. VI. Reovirus: virion genome dsRNA as the interferon inducer in aged chick embryo cells. *J. Interferon Res.* **1**:155–167.
 46. **Wolf, T., and S. Ludwig.** 2009. Influenza viruses control the vertebrate type I interferon system: factors, mechanisms, and consequences. *J. Interferon Cytokine Res.* **29**:549–557.