Glycine at Position 622 in PB1 Contributes to the Virulence of H5N1 Avian Influenza Virus in Mice

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ABSTRACT
We isolated two H5N1 viruses, A/duck/Hunan/S4020/2008 (DK/08) and A/chicken/Guangxi/S2039/2009 (CK/09), from live-bird markets during routine surveillance and found that these two viruses are genetically similar but differ in their replication and virulence in mice. The CK/09 virus is lethal for mice with a 50% mouse lethal dose (MLD$_{50}$) of 1.6 log$_{10}$ 50% egg infectious doses (EID$_{50}$), whereas the DK/08 virus is nonpathogenic for mice with an MLD$_{50}$ value of 6.2 log$_{10}$ EID$_{50}$. We explored the genetic basis of the virulence difference of these two viruses by generating a series of reassortant viruses and mutants in the lethal virus CK/09 background and evaluating their virulence in mice. We found that the PB1 gene of the DK/08 virus dramatically attenuated the virulence of the CK/09 virus and that the amino acid at position 622 in PB1 partially impaired the binding of PB1 to viral RNA, thereby dramatically decreasing the polymerase activity and attenuating H5N1 virus virulence in mice. Our results identify a novel virulence-related marker of H5N1 influenza viruses and provide a new target for live attenuated vaccine development.

IMPORTANCE
H5N1 avian influenza viruses have caused the deaths of nearly 60% of the humans they have infected since 1997 and clearly represent a threat to public health. A thorough understanding of the genetic basis of virulence determinants will provide important insights for antiviral drug and live attenuated vaccine development. Several virulence-related markers in the PB2, PA, M1, and NS1 proteins of H5N1 viruses have been identified. In this study, we isolated two H5N1 avian influenza viruses that are genetically similar but differ in their virulence in mice, and we identified a new virulence-related marker in the PB1 gene. We found that the mutation of glycine (G) to aspartic acid (D) at position 622 in PB1 partially impairs the binding of PB1 to viral RNA, thereby attenuating H5N1 virus virulence in mice. This newly identified virulence-related marker could be applied to the development of live attenuated vaccines against H5N1 influenza.

H5N1 avian influenza outbreaks in poultry have become widespread since late 2003, and H5N1 viruses have caused numerous disease outbreaks in domestic poultry and wild birds in many countries throughout Asia, Europe, and Africa (http://www.oie.int). H5N1 virus infection of humans has been reported in 16 countries, with 429 deaths among 784 cases as of 3 March 2015 (http://www.who.int). Several studies have shown that the H5N1 viruses could become transmissible in mammals if they acquired more mutations or reassorted with human influenza viruses (1–5). Thus, the H5N1 viruses circulating in nature pose huge threats to both animals and public health.

The influenza A virus genome comprises eight gene segments, including basic polymerase 2 (Pol II; PB2 gene), basic polymerase 1 (PB1 gene), acidic polymerase (PA gene), hemagglutinin (HA gene), nucleoprotein (NP gene), neuraminidase (NA gene), matrix (M gene), and nonstructural protein (NS gene). These gene segments encode at least 12 proteins, including PB2, PB1, PB1-F2, PA, PA-X, HA, NA, NP, M1, M2, NS1, and NS2. Many studies have investigated the molecular basis of the lethality of H5N1 avian influenza viruses in mammalian hosts, and a series of virulence-related amino acids in different proteins have been identified (6–17). For example, in the PB2 protein, the amino acid substitution at position 627 from glutamic acid to lysine (E627K) and the amino acid substitution at position 701 from aspartic acid to asparagine (D701N) play crucial roles in the ability of H5N1 viruses to replicate and be lethal in mammals (1, 6). Several amino acids in the PA proteins, including those at positions 97, 185, 224, and 383, have been reported to affect the virulence of H5N1 viruses in mice (12, 15, 17). The motif of multiple basic amino acids in the cleavage site of HA is the prerequisite for the virulence of H5N1 virus in both avian and mammalian hosts (6, 18). The amino acids asparagine at position 30 and alanine at position 215 in the M1 protein are necessary for H5N1 virus lethality in mice (13). Moreover, the amino acid serine at position 42 of NS1 is critical for the H5N1 influenza virus to antagonize host cell interferon induction and for the pathogenicity of H5N1 influenza viruses in mammalian hosts (11). During our routine surveillance studies, we isolated two H5N1 viruses, A/duck/Hunan/S4020/2008 (DK/08) and A/chicken/Guangxi/S2039/2009 (CK/09), from birds in live-bird markets...
TABLE 1 Primers used for pBD cDNA construction and for introducing mutations into the PB1 gene of the mutant viruses

| Purpose | Primer(s) (5’–3’) | Reverse |
|---------|------------------|---------|
| PB2 amplification | CCAGCAAAAAAGCAGTTCAATATATTTCA | TTAGTGAACACAAAGGTGTTTTTAAAT (DK/08), TTAGTGAACACAAAGGTGTTTTTAAAC (CK/09) |
| PB1 amplification | CCAGCAAAAAAGCAGTTCAATATATTTCA | TTAGTGAACACAAAGGTGTTTTTAAAT (DK/08), TTAGTGAACACAAAGGTGTTTTTAAAC (CK/09) |
| PA amplification | CCAGCAAAAAAGCAGTTCAATATATTTCA | TTAGTGAACACAAAGGTGTTTTTAAAT (DK/08), TTAGTGAACACAAAGGTGTTTTTAAAC (CK/09) |
| HA amplification | CCAGCAAAAAAGCAGTTCAATATATTTCA | TTAGTGAACACAAAGGTGTTTTTAAAT (DK/08), TTAGTGAACACAAAGGTGTTTTTAAAC (CK/09) |
| NP amplification | CCAGCAAAAAAGCAGTTCAATATATTTCA | TTAGTGAACACAAAGGTGTTTTTAAAT (DK/08), TTAGTGAACACAAAGGTGTTTTTAAAC (CK/09) |
| NA amplification | CCAGCAAAAAAGCAGTTCAATATATTTCA | TTAGTGAACACAAAGGTGTTTTTAAAT (DK/08), TTAGTGAACACAAAGGTGTTTTTAAAC (CK/09) |
| M amplification | CCAGCAAAAAAGCAGTTCAATATATTTCA | TTAGTGAACACAAAGGTGTTTTTAAAT (DK/08), TTAGTGAACACAAAGGTGTTTTTAAAC (CK/09) |
| NS amplification | CCAGCAAAAAAGCAGTTCAATATATTTCA | TTAGTGAACACAAAGGTGTTTTTAAAT (DK/08), TTAGTGAACACAAAGGTGTTTTTAAAC (CK/09) |
| CK/09 PB1D619N mutation | CGAATGGAATTGATGGATGAA | GGAGGATTTGGGCAACACATTCAGAATTTT |
| CK/09 PB1G622D mutation | CGAATGGAATTGATGGATGAA | GGAGGATTTGGGCAACACATTCAGAATTTT |
| CK/09 PB1R635K mutation | CGAATGGAATTGATGGATGAA | GGAGGATTTGGGCAACACATTCAGAATTTT |
| CK/09 PB1D619N/G622D mutation | CGAATGGAATTGATGGATGAA | GGAGGATTTGGGCAACACATTCAGAATTTT |

*The nucleotides that have been changed are underlined and in boldface type.

and found that these two viruses are genetically similar but differ in their replication and virulence in mice. Yet all of the known virulence-related markers are the same in these two viruses, suggesting that other amino acids or motifs may contribute to the difference in lethality between these two viruses in mice. In this study, we used reverse genetics to generate a series of reassortants and mutants of these two viruses in the CK/09 background and tested their virulence in mice. We identified a novel virulence-related marker in the PB1 protein and explored the underlying mechanism for the difference in lethality between these two viruses in mice.

MATERIALS AND METHODS

Ethics statement and facility. The present study was carried out in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the Ministry of Science and Technology of the People’s Republic of China. Studies with highly pathogenic H5N1 avian influenza viruses were conducted in a biosecurity level 3 laboratory approved for such use by the Chinese Ministry of Agriculture. The protocol was approved by the Committee on the Ethics of Animal Experiments of the Harbin Veterinary Research Institute (HVRI) of the Chinese Academy of Agricultural Sciences (CAAS).

Cells and viruses. Human embryonic kidney cells (293T) and MDCK cells incubated at 37°C in 5% CO₂ were grown in Dulbecco’s modified Eagle’s medium supplemented with 10% and 5% fetal bovine serum, respectively, plus antibiotics. The H5N1 viruses, A/duck/Hunan/S4020/2006 (DK/08) and A/chicken/Guangxi/S2039/2009 (CK/09), were isolated from live-bird markets during routine surveillance. Virus stocks were propagated in 10-day-old specific-pathogen-free (SPF) embryonated chicken eggs and stored at −70°C until they were used for RNA extraction and animal studies.

Sequence analysis. Viral RNA (vRNA) was extracted from allantoic fluid and was subjected to reverse transcription (RT). A set of fragment-specific primers (primer sequences available on request) were used for the PCR amplification and sequence analysis.

Construction of plasmids for virus rescue. Construction of plasmids in viral RNA vRNA-mRNA bidirectional expression plasmid pBD for virus rescue was performed as described previously (8) with the primers shown in Table 1. The constructs were designated pBD/DK/08-PB1, pBD/DK/08-PB2, pBD/DK/08-PA, pBD/DK/08-NS, pBD/DK/08-NA, and pBD/DK/08-PB1, pBD/DK/08-PB2, pBD/DK/08-PA, pBD/DK/08-NS, pBD/DK/08-NA, pBD/DK/08-PB1, pBD/DK/08-PB2, pBD/DK/08-PA, pBD/DK/08-NS, and pBD/DK/08-PB1, pBD/DK/08-PB2, pBD/DK/08-NS, and pBD/DK/08-NA. The nucleotides that have been changed are underlined and in boldface type.

Reassortant or mutant viruses were detected by using a hemagglutination assay, and RNA was extracted and analyzed by reverse transcription-PCR (RT-PCR). Each viral segment was sequenced to confirm the identity of the reassortant viruses.

Animal experiments. For the replication study, groups of three 6-week-old female BALB/c mice (Beijing Experimental Animal Center) were lightly anesthetized with CO₂ and inoculated intranasally with 10⁶ EID₅₀. 50% egg infectious doses of H5N1 influenza virus in a volume of 50 μl were administered orally to all the mice. Lungs, brains, kidneys, and spleens of mice were collected and titrated for virus infectivity in eggs as described previously (22). The 50% mouse lethal dose (MLD₅₀) was determined by inoculating groups of five 6-week-old female BALB/c mice with 10-fold serial dilutions containing 10⁴ to 10⁶ EID₅₀ of the virus in a 50-μl volume.

Polymerase activity. A dual-luciferase reporter assay system (Promega) was used to compare the activities of viral ribonucleoprotein (RNP) complexes. To construct the reporter plasmid, pPol-Luc, the open reading frame of the luciferase gene, flanked by the 5’ and 3’ noncoding regions of the NP gene of the DK/08 virus, was inserted into a pPol 1 plasmid containing the sequence of the human polymerase I promoter. Briefly, 0.5 μg of luciferase reporter plasmid pPol-Luc and the Renilla luciferase expressing plasmid pTK-RL was transfected into 2 × 10⁵ 293T cells together with 0.5 μg of each of the four protein expression plasmids pcDNA3.1-PB2, pcDNA3.1-PB1 (or PB1 mutants), pcDNA3.1-PA, and pcDNA3.1-NP of DK/08 or DK/08. The Renilla luciferase expressed by pTK-RL was used as an internal control to normalize transfection efficiency. Cell extracts were harvested 30 h posttransfection, and luciferase activity was assayed by using the luciferase assay system (Promega). The assay was standardized to the activity of CK/09 (100%). There were two replicates for each complex combination each time, and all experiments were performed three times.

Viral replication in MDCK cells. Virus was inoculated into MDCK monolayers at a multiplicity of infection (MOI) of 0.001. The cells were

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supplemented with Opti-MEM and incubated at 37°C. Virus-containing culture supernatant was collected at various time points (hours postinfection [hpi]) and titrated in MDCK cells. The growth data shown are the virus titration in eggs.

Generation of the model vRNA. A 290-nucleotide model vRNA was transcribed in vitro by using a cDNA containing the T7 promoter that was used to transfect 293T cells. Protein binding vRNA assay. 293T cells were transfected with a plasmid expressing Flag-tagged, truncated CK/09-PB1 or with plasmid pCAGGS-Flag as a control. The cell lysate proteins captured by protein G, unbound proteins, and vRNA-bound proteins from the vRNA binding protein assay were resolved by means of 15% SDS-PAGE followed by Western blotting.

vRNA binding protein assay. 293T cells were transfected with a plasmid expressing Flag-tagged, truncated CK/09-PB1 or with plasmid pCAGGS-Flag as a control. The cell lysate proteins captured by protein G, unbound proteins, and vRNA-bound proteins from the vRNA binding protein assay were resolved by means of 15% SDS-PAGE followed by Western blotting.

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### RESULTS

**Virus rescue and characterization.** We cloned the cDNAs of each full-length RNA segment of the DK/08 and CK/09 viruses into a

### Table 2 Replication and lethality of H5N1 viruses in mice

| Virus Replication on day 3 p.i. (log<sub>10</sub> EID<sub>50</sub> ± SD) | Lung | Brain | Spleen | Kidney | MLD<sub>50</sub> (log<sub>10</sub> EID<sub>50</sub>)<sup>†</sup> | Attenuation (fold)<sup>‡</sup> |
|------------------|-------|-------|--------|--------|----------------|-------------------|
| DK/08            | 6.0 ± 0.5 | −     | −      | −      | 6.2 | NA |
| CK/09            | 6.7 ± 0.5 | 3.8 ± 0.6 | 4.2 ± 0.4 | 4.0 ± 0.7 | 1.6 | NA |
| R-DK/08          | 6.0 ± 0.7 | −     | −      | −      | 6.2 | NA |
| CK/09-DK/08PB2   | 7.3 ± 0.1 | 4.5 ± 0.0 | 4.5 ± 0.0 | 4.4 ± 0.1 | 1.5 | NA |
| CK/09-DK/08PB1   | 6.5 ± 0.7 | 4.5 ± 0.0 | 4.1 ± 0.7 | 4.4 ± 0.1 | 1.4 | 0 |
| CK/09-DK/08PA    | 5.3 ± 0.4<sup>‡</sup> | −     | 1.5 ± 0.0<sup>‡</sup> | −      | 4.5 | 1,000 |
| CK/09-DK/08HA    | 6.5 ± 0.3 | 4.2 ± 0.4 | 4.2 ± 0.6 | 3.8 ± 0.7 | 1.2 | 0 |
| CK/09-DK/08NP    | 6.8 ± 0.4 | 3.9 ± 0.5 | 4.5 ± 0.0 | 4.2 ± 0.6 | 1.3 | 0 |
| CK/09-DK/08M     | 6.6 ± 0.8 | 4.4 ± 0.1 | 4.4 ± 0.1 | 4.3 ± 0.1 | 1.2 | 0 |
| CK/09-DK/08NS    | 7.1 ± 0.5 | 4.5 ± 0.0 | 4.4 ± 0.1 | 4.5 ± 0.0 | 1.2 | 0 |
| CK/09-DK/08 M    | 6.8 ± 0.6 | 3.4 ± 0.8 | 4.4 ± 0.1 | 4.1 ± 0.5 | 1.4 | 0 |
| Chimera 1        | 6.3 ± 0.4 | 3.5 ± 0.9 | 4.4 ± 0.1 | 4.2 ± 0.6 | 1.8 | 2 |
| Chimera 2        | 4.9 ± 0.3<sup>‡</sup> | −     | 1.4 ± 1.0<sup>‡</sup> | 1.3 ± 0.9<sup>‡</sup> | 4.5 | 1,000 |
| CK/09-PB1D619N   | 6.6 ± 0.1 | 2.9 ± 1.4 | 4.0 ± 0.4 | 3.8 ± 0.4 | 1.4 | 0 |
| CK/09-PB1G622D   | 5.8 ± 0.5<sup>‡</sup> | 1.2 ± 0.8<sup>‡</sup> | 2.9 ± 2.1<sup>‡</sup> | 2.0 ± 1.3<sup>‡</sup> | 4.2 | 501 |
| CK/09-PB1R635K   | 6.5 ± 0.2 | 3.3 ± 0.9 | 3.6 ± 0.6 | 3.6 ± 0.9 | 2.5 | 10 |
| CK/09-PB1D619N+G622D | 5.3 ± 0.4 | −     | 2.3 ± 0.3<sup>‡</sup> | −      | 4.5 | 1,000 |
| CK/09-PB1G622D+R635K | 3.8 ± 0.6<sup>‡</sup> | −     | 3.3 ± 0.5 | 1.3 ± 1.0<sup>‡</sup> | 4.2 | 501 |

<sup>a</sup>Six-week-old female BALB/c mice were used for these studies.

<sup>b</sup>Groups of three mice were inoculated intranasally with 10<sup>6.0</sup> EID<sub>50</sub> of the test virus in a 50-μl volume and were killed on day 3 postinoculation (p.i.); organs were then collected for virus titration in eggs. −, no virus was detected in undiluted samples. Virus titers of mice were compared by using the Student-Newman-Keuls test. SD, standard deviation.

<sup>c</sup>The 50% mouse lethal dose (MLD<sub>50</sub>) was determined by intranasally inoculating groups of five mice with 10-fold serial dilutions containing 10<sup>1</sup> to 10<sup>6</sup> EID<sub>50</sub> of virus in a 50-μl volume.

<sup>d</sup>P < 0.05 compared with the titers in the corresponding organs of the R-DK/08-inoculated mice.

<sup>e</sup>Western blotting.

<sup)f</sup>Virulence decrease compared with the R-DK/09 virus. NA, not applicable.

### Nucleotide sequence accession numbers.

The sequence data for the two viruses used in these studies are available in GenBank (accession numbers KT762428 to KT762443).
vRNA-mRNA bidirectional expression plasmid (pBD) as described previously (8) and by using the set of gene segment-specific primers shown in Table 1. All of the constructs were completely sequenced to ensure the absence of unwanted mutations. By using these plasmids, we rescued the DK/08 and CK/09 viruses, designated R-DK/08 and R-CK/09, respectively, grew them in 10-day-old SPF embryonated chicken eggs, and tested their replication and lethality in mice. The rescued R-DK/08 virus, like the wild-type DK/08, replicated only in the lungs and killed only one mouse at the inoculated dose ($10^6$ EID$_{50}$), which was administered intranasally, yielding a 50% MLD$_{50}$ of 6.2 log$_{10}$ EID$_{50}$ (Table 2). However, R-CK/09 replicated systemically and was as highly pathogenic as the original CK/09 virus, with an MLD$_{50}$ of 1.5 log$_{10}$ EID$_{50}$ (Table 2). These results indicate that the rescued viruses maintained the biological properties of the wild-type viruses.

The PB1 gene plays a major role in the difference in pathogenicity in mice between the DK/08 and CK/09 viruses. The two viruses differ by 74 amino acids in their 11 proteins (Fig. 1). We used the "single-gene recombinant" strategy to identify genes that contributed to the virulence of the viruses in mice, as described elsewhere (6, 8–16). However, to avoid any "gain-of-function" concerns, we used only the CK/09 lethal virus as the backbone to generate the reassortants, each containing one gene derived from DK/08, and tested their replication and pathogenicity in mice. Seven viruses that carried the PB2, PA, HA, NP, NA, M, or NS gene of DK/08 virus were as virulent as the CK/09 virus and replicated in all four mouse organs tested (Table 2). However, introduction of the PB1 gene of the DK/08 virus dramatically attenuated the pathogenicity of CK/09 by 1,000-fold (MLD$_{50}$ of 1.5 versus 4.5 log$_{10}$ EID$_{50}$), and the reassortant virus replicated only in the lungs and spleen of infected mice (Table 2).

The amino acid glycine at position 622 of PB1 is critical for the virulence of the CK/09 virus in mice. The PB1 proteins of the DK/08 and CK/09 viruses differ by six amino acids (Fig. 2).
pinpoint the amino acid(s) in PB1 that contributes to the virulence of the CK/09 virus, we generated two viruses that expressed chimeric PB1 proteins (Fig. 2) and tested them in mice. When the N-terminal portion of the CK/09 PB1 gene was replaced with the corresponding DK/08 PB1 gene segment (chimera 1), within the context of the remaining genes coming from CK/09, the chimeric virus was as lethal as the wild-type CK/09 virus and replicated systemically in mice (Table 2 and Fig. 2). However, chimera 2 (the CK/09 virus with the C-terminal portion of the DK/08 PB1 gene) was attenuated 1,000-fold, and the virus replicated in multiple organs, but the titers were lower than those of the wild-type CK/09 virus (Table 2). These results indicate that the C terminus of the PB1 gene is important for the virulence of the CK/09 virus in mice.

The two viruses differ by only three amino acids in this region of PB1 at positions 619, 622, and 635 (Fig. 2). To pinpoint which amino acid contributes to the pathogenicity in mice, we generated and tested three mutants, CK/09-PB1D619N, CK/09-PB1G622D, and CK/09-PB1R635K (Fig. 2). All three mutants were detected in all four organs tested in mice, but the titers of CK/09-PB1G622D virus were significantly decreased and the virulence of the mutant was attenuated over 500-fold compared with that of the R-CK/09 virus (Table 2). Addition of the D619N substitution slightly attenuated the CK/09-PB1G622D virus results (MLD50, 4.5 log10 EID50) compared with R-DK/08 (Fig. 3B), which was in agreement with the context of the remaining genes coming from CK/09, the chimeric virus was as lethal as the wild-type CK/09 virus and replicated systemically in mice (Table 2 and Fig. 2). However, chimera 2 (the CK/09 virus with the C-terminal portion of the DK/08 PB1 gene) was attenuated 1,000-fold, and the virus replicated in multiple organs, but the titers were lower than those of the wild-type CK/09 virus (Table 2). These results indicate that the C terminus of the PB1 gene is important for the virulence of the CK/09 virus in mice.

The amino acid at position 622 in the PB1 protein affects viral replication in MDCK cells. We compared the multicycle growth levels of R-DK/08, R-CK/09, and CK/09-PB1G622D in MDCK cells. The R-CK/09 virus grew more rapidly than did the R-DK/08 virus, and the titers of R-CK/09 were significantly higher than those of R-DK/08 (Fig. 3B). The titers of CK/09-PB1G622D were significantly lower than those of R-CK/09 but were higher than those of R-DK/08 (Fig. 3B), which was in agreement with the replication and virulence in mice of these viruses.

The G622D mutation in PB1 partially impairs the ability of PB1 to bind vRNA. PB1 has two regions (the N-terminal 83 amino acids and the C-proximal sequences located downstream of position 493) that bind to vRNA (26–30). The amino acid at position 622 is located in one of the vRNA binding domains of PB1. We therefore tested whether the G622D mutation affects the binding of PB1 to vRNA. 293T cells were transfected with plasmid pCAGGS-3Flags-PB1Δ1-493, which expresses the Flag-tagged, truncated CK/09-PB1Δ1-493 protein, or with plasmid pCAGGS-3Flags-PB1Δ1-493/G622D, which expresses the CK/09-PB1Δ1-493/G622D protein (Fig. 4A), or with plasmid pCAGGS-3Flags as a control. Cell lysates containing 50 μg of total protein were captured by protein G (Life Technology) that had been cross-linked with a mouse anti-Flag monoclonal antibody. The captured PB1 proteins (Fig. 4B) were then used to bind 10 μg of model vRNA that was transcribed in vitro (Fig. 4C). After a 4-h incubation at 4°C, the unbound and protein-bound vRNA was quantified by use of real-time RT-PCR. As shown in Fig. 4D, the amount of vRNA bound by CK/09-PB1Δ1-493/G622D was about 90% of that bound by CK/09-PB1Δ1-493. In the plasmid pCAGGS-3Flags-transfected cell lysate, the truncated PB1 protein was not captured (Fig. 4B) and binding of vRNA was not detected (Fig. 4D). We also performed a vRNA binding protein assay. Ten micro-
grams of biotinylated model vRNA was incubated with cell lysate containing 150 μg of total protein that had previously been transfected with plasmid pCAGGS-3Flags-PB11-493, which expresses the CK/09-PB11-493 protein, or with plasmid pCAGGS-3Flags-PB11-493/G622D, which expresses the CK/09-PB11-493/G622D protein, or with plasmid pCAGGS-3Flags as a control. Cell lysates containing 100 μg of total protein were captured by analysis using protein G (Life Technology). The capture of the PB1 proteins was confirmed by use of Western blotting. We found that the vRNA did not bind any protein in the pCAGGS-3Flags plasmid-transfected cell lysate (Fig. 4E) and that the same amount of vRNA bound to more CK/09-PB11-493 protein than CK/09-PB11-493/G622D protein (Fig. 4E and F). Therefore, the results of these two assays, the protein binding vRNA assay and the vRNA binding protein assay, were in accord and demonstrated that the G622D mutation in PB1 partially impairs the ability of PB1 to bind vRNA.

PB1 has several functional domains; its N and C termini interact with PA and PB2, respectively (26, 29–37), and PB1 contains two regions that bind to vRNA (26–30). The amino acid at position 622 is located in one of these vRNA binding domains. The G622D mutation in PB1 resulted in a 10% reduction in the ability of PB1 to bind vRNA in vitro but in a 40-fold decrease in polymerase activity and a 500-fold attenuation of the virulence of the virus in mice. The 10% reduction of PB1/622D binding to vRNA observed in vitro reflects a single round of PB1-vRNA binding; however, the reduction in binding is greater in vivo, as reflected in the polymerase activity analysis and animal studies, because multiple rounds of PB1-vRNA binding would have occurred. The G622D mutation may also alter the interaction of PB1 with PB2 or PA, although this seems unlikely because the PB1 N-terminal PA-interacting helix (residues 1 to 15) and the C-terminal three-helix bundle (residues 685 to 757) that interacts with residues 1 to 35 of PB2 (35, 38) are located far from the amino acid at position 622. Further studies are warranted to determine whether the G622D mutation in PB1 alters its interaction with host proteins.

Many studies have been undertaken to identify the virulence determinants of influenza viruses in different hosts. These studies have identified a number of virulence-related markers by using different virus strains (1, 11, 15, 17, 39, 40). However, the underlying mechanisms of only a few of these markers have been revealed. The D701N mutation in PB2 is reported to have a key role in increasing the replication and virulence of different avian influ-
Table 3 Naturally isolated influenza viruses bearing 622D in their PB1 sequence

| Virus                          | Accession no. of PB1 sequence in GenBank |
|-------------------------------|-----------------------------------------|
| A/mallard/ Sweden/68561/2007 (H4N6) | CY165168                                 |
| A/swine/England/VWL16/1998 (H1N1) | CY037950                                 |
| A/swine/England/266029/1998 (H1N1) | CY116072                                 |
| A/swine/England/WV15/1997 (H1N1) | CY037942                                 |
| A/swine/England/6636804/1996 (H1N1) | CY116250                                 |
| A/swine/England/167655/1997 (H1N1) | CY116002                                 |
| A/duck/Human/S4020/2008 (H5N1) | KT762429                                 |

The present trimeric polymerase assembly (41). Zhu et al. found that deletion of the amino acids at positions 191 to 195 in NS1 attenuates H5N1 influenza virus in chickens by affecting the stability of the NS1 protein and impairing the interaction of NS1 protein with chicken cleavage and polyadenylation specificity factor (40). Jiao et al. further demonstrated that the amino acid serine at position 42 of NS1 is essential for H5N1 influenza virus to antagonize host cell interferon induction by preventing the double-stranded RNA-mediated activation of the NF-kB pathway and the interferon regulatory transcription factor 3 (IRF-3) pathway (11). The present study showed that the G622D mutation in PB1 attenuates the H5N1 influenza virus in mice by partially impairing the binding of PB1 to vRNA. These findings indicate that different genetic changes in influenza virus alter its virulence through different mechanisms.

Although, according to the available sequences in the public database, the glycine at position 622 in PB1 is highly conserved among different subtypes of influenza viruses, a few strains have been detected in nature that have 622D in their PB1 (Table 3). We identified the viruses that were recovered from the mouse organs and found that wild-type DK/08 and the mutants generated in this study stably maintained 622D in their PB1 after they replicated in mice. Thus, the G622D mutation in PB1 could potentially be used in the development of live attenuated vaccines against influenza viruses.

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