1	Enhanced pathogenicity and neurotropism of mouse-adapted H10N7 influenza
2	virus are mediated by novel PB2 and NA mutations
3	Xuxiao Zhang, <sup>a</sup> Guanlong Xu, <sup>b</sup> Chenxi Wang, <sup>a</sup> Ming Jiang, <sup>a</sup> Weihua Gao, <sup>a</sup> Mingyang
4	Wang, <sup>a</sup> Honglei Sun, <sup>a</sup> Yipeng Sun, <sup>a</sup> Kin-Chow Chang, <sup>c</sup> Jinhua Liu, <sup>a</sup> and Juan Pu <sup>a*</sup>
5	
6	Key Laboratory of Animal Epidemiology and Zoonosis, Ministry of Agriculture,
7	College of Veterinary Medicine, and State Key Laboratory of Agrobiotechnology,
8	China Agricultural University, Beijing, China <sup>a</sup> . China Institute of Veterinary Drug
9	Control, Beijing, China <sup>b</sup> . School of Veterinary Medicine and Science, University of
10	Nottingham, Sutton Bonington Campus, Loughborough, United Kingdom <sup>e</sup> .
11	
12	Running title: Pathogenicity and neurotropism of H10N7 virus in mice
13	
14	* Address correspondence to Juan Pu, pujuan@cau.edu.cn.
15	
16	
17	Word counts: Abstract: 238; Main text: 4151.
18	
19	
20	
21	
22	

#### 23 Abstract

Recent human fatalities from avian-origin H10N8 influenza virus infection raise 24 25 concerns about the threat of this virus subtype to public health. To investigate genetic adaptation of H10 avian influenza viruses in mammals, we generated 26 27 a mouse-adapted avian H10N7 variant (A/mallard/Beijing/27/2011-MA, [BJ27-MA]) through nine serial passages in mice. Mice infected with BJ27-MA virus died by 6 28 days post-infection and showed neuronal infection in contrast to parental virus which 29 elicited no overt symptoms. Sequence analysis showed the absence of the widely 30 recognized mammalian adaptation markers of E627K and D701N in PB2 in the 31 mouse-adapted strain; instead five amino acid mutations were identified: E158G and 32 M631L in PB2, G218E in HA (H3 numbering), and K110E and S453I in NA. 33 34 Neurovirulence of BJ27-MA virus necessitated the combined presence of the PB2 and NA mutations. Mutations M631L and E158G of PB2 and K110E of NA were required 35 to mediate increased virus replication and severity of infection in mice and 36 mammalian cells. PB2-M631L was functionally the most dominant mutation in that it 37 strongly up-regulated viral polymerase activity and played a critical role in the 38 enhancement of virus replication and disease severity in mice. K110E mutation in NA, 39 on the other hand, significantly promoted NA enzymatic activity. These results 40 indicate that the novel mutations in PB2 and NA genes are critical for the adaptation 41 of avian H10N7 influenza virus in mice, which could serve as molecular signatures of 42 43 virus transmission to mammalian hosts including humans.

44

#### 45 **Importance**

The increasingly prevalent H10 subtype of avian influenza virus in China has 46 recently been a source of human fatalities. We demonstrated that an avian H10N7 47 virus can readily be adapted to become highly pathogenic and neurotropic in mice. 48 Mutations in PB2 and NA from the mouse-adapted virus (BJ27-MA) were the major 49 determinants of enhanced pathogenicity of which mutation PB2-M631L was 50 functionally most dominant. Although BJ27-MA virus lacked the well-known 51 mammalian adapted mutations (namely PB2-E627K and PB2-D701N), PB2-M631L 52 mutation enhanced viral polymerase activity, replication and pathogenicity of 53 54 BJ27-MA virus in mice, indicating a novel adaptation strategy. These observations affirm the public health threat of avian H10 subtype influenza viruses and have 55 implications in the assessment of potential mutant viruses that may cause severe 56 infections in humans. 57

58

#### 59 Introduction

Presently, avian influenza viruses (AIVs) cause great economic losses to the global 60 61 poultry industry, which historically were major contributors to the 1918 H1N1, 1957 H2N2 and 1968 H3N2 virus pandemics (1). H5N1 and H9N2 influenza viruses, as the 62 two principal subtypes circulating in poultry, are high on the list of candidates that 63 could potentially cause another major human influenza outbreak (2, 3). However, 64 recent human cases of emergent avian H7N9 virus infection challenge our 65 understanding of the main subtypes of possible future pandemic human virus (4). 66 67 Thus, contingency planning in the prevention and management of avian influenza virus infections in human should be based on a broad range of possible subtypes. 68

Between November 2013 and February 2014, two fatal and one severe cases of 69 70 human infections with a novel reassortant H10N8 virus in Jiangxi, China, were reported for the first time (5-7). Avian H10 virus subtype was firstly isolated from 71 chickens in Germany in 1949 (8, 9); subsequently viruses bearing H10 hemagglutinin 72 (HA) and different neuraminidase (NA) subtypes have become widely prevalent in 73 wild birds and domestic poultry around the world (10-12). Since 1984, repeated 74 infections or deaths of mammals with this subtype have been reported, such as the 75 outbreaks of H10N4 virus in minks (13), H10N5 virus in domestic pigs (14), H10N8 76 virus in feral dogs (15) and H10N7 virus in harbor seals (16-18). Human cases of H10 77 virus infections had occurred sporadically in several other countries. For example, 78 H10N7 viruses had caused a number of human infections in Egypt in 2004 (19). In 79 March 2010, H10N7 virus infection was identified in two abattoir workers in a 80

commercial poultry farm in Australia who showed conjunctivitis and minor upper respiratory tract symptoms (12). In the USA, serological evidence of exposure to H10 virus subtype was confirmed in turkey workers (20). The repeated human cases of H10 virus infections coupled with the prevalence of H10 viruses in birds raise concerns that this particular subtype could pose increasing threat to human and animal health. However, the molecular adaptations of H10 influenza viruses in mammals are largely unknown.

Adaptation is considered to be a primary driver in evolution, and the process of 88 89 natural selection of influenza A viruses in experimental mice appears also to hold true 90 for humans (21). Several adaptation studies of human H3N2, pandemic 2009 H1N1, avian H9N2 and H6N6 influenza viruses in mice have provided better understanding 91 92 of molecular determinants of virus pathogenicity in mammals including humans (22-26). Here, to explore the genetic adaptations of H10 AIV subtype in mammals, we 93 serially passaged a low-pathogenicity avian-derived H10N7 virus in mice. We found 94 that mouse-adapted H10N7 virus acquired high pathogenicity status causing fatal 95 infection and neurovirulence. The well-known mammalian adaptation markers 96 PB2-E627K and PB2-D701N (27, 28) were not found in the mouse-adapted strain, but 97 amino acid substitution PB2-M631L was a dominant contributor to virus virulence. 98

99

#### 100 Materials and methods

#### 101 Ethics statement

102 All animal work was approved by the Beijing Association for Science and

Technology (approval ID SYXK [Beijing] 2007-0023) and conducted in accordance
with the Beijing Laboratory Animal Welfare and Ethics guidelines, as issued by the
Beijing Administration Committee of Laboratory Animals, and in accordance with the
China Agricultural University Institutional Animal Care and Use Committee
guidelines (ID: SKLAB-B-2010-003).

#### 108 Viruses and cells

The H10N7 virus A/mallard/Beijing/27/2011 (BJ27) was isolated in Beijing, China 109 and propagated in the allantoic cavities of 10-day-old specific-pathogen-free (SPF) 110 111 embryonated chicken eggs (Merial, Beijing, China) at 37°C for 72 h. Allantoic fluid 112 containing virus was harvested, aliquoted and frozen at -80°C for later use. Viruses were titrated in MDCK cells to determine the 50% tissue culture infectious dose 113 114 (TCID<sub>50</sub>) by the Reed and Muench method (29). Human embryonic kidney (293T), human pulmonary adenocarcinoma (A549), Madin Darby canine kidney (MDCK) and 115 mouse neuroblastoma N2a (N2a) cells were maintained in Dulbecco's modified 116 Eagle's medium (DMEM, Gibco) supplemented with 10% fetal bovine serum (FBS, 117 Gibco), 100 units/ml of penicillin, and 100 µg/ml of streptomycin at 37°C in 5% CO<sub>2</sub> 118 atmosphere. 119

#### 120 Adaptation of the BJ27 virus in mice

Six 6-week-old female BALB/c mice (Beijing Experimental Animal Center) were anesthetized with Zoletil 50 (tiletamine-zolazepam; Virbac S.A., Garros, France) and inoculated intranasally with 50 μl of allantoic fluid containing BJ27 virus. At 3 days post-inoculation (dpi), three mice were euthanized and the lungs were harvested and

homogenized in 1 ml of sterile cold phosphate-buffered saline (PBS). The 125 homogenate was centrifuged at  $6,000 \times g$  for 5 min at 4°C and filtered through a 126 0.22-µm-pore-size cellulose acetate filter (Millipore, USA). Fifty µl of filtered 127 homogenate were used as inoculum per mouse for the next passage (passage 2 [P2]). 128 129 The remaining three mice were monitored daily for clinical symptoms. At 9 passages (P9), the virus in the lung homogenate was subjected to three rounds of plaque 130 purification in MDCK cells, and the cloned virus, designated BJ27-MA, was 131 amplified once in 10-day-old SPF embryonated eggs for 72 h at 37°C, as previously 132 133 described (30).

#### 134 Sequence analysis

Viral RNA was extracted from allantoic fluid containing plaque-purified BJ27-MA.
The eight virus genes were amplified by reverse transcription-PCR (RT-PCR) and
sequenced. Adaptive mutations arising from serial passage were identified by
comparing consensus BJ27-MA and wild type BJ27 sequences.

#### 139 Plasmid construction and virus rescue

The eight gene segments of BJ27 and BJ27-MA were amplified by RT-PCR and cloned into the expression plasmid, PHW2000. Mutations of interest in the PB2, HA and NA gene were introduced by PCR-based site-directed mutagenesis with primer pairs containing point mutations. All constructs were sequenced to confirm mutational changes.

Reassortant viruses between BJ27 and BJ27-MA were generated by reverse
genetics as described previously (31). Briefly, 0.5 μg of each gene segment plasmid

was mixed together and incubated with 8 µl of TransIT-LT1 reagent (Mirus Bio, USA) 147 at 20°C for 30 min. The TransIT-LT1-DNA mixture was transferred to 70% confluent 148 149 293T/MDCK co-cultured monolayers and incubated at 37°C with 5% CO<sub>2</sub>. Six hours post-transfection, the supernatants were replaced with 2 ml of Opti-MEM containing 150 2 µg/ml TPCK-treated trypsin (Sigma-Aldrich). Forty-eight hours post-transfection, 151 the cell supernatants were harvested and inoculated into 10-day-old SPF embryonated 152 eggs and incubated for 72 h at 37°C to prepare a virus stock. Viral RNA was extracted 153 and analyzed by RT-PCR, and each viral segment was sequenced to confirm identity. 154 155 Virus titers were determined by TCID<sub>50</sub> assay on MDCK cells.

#### 156 Mouse experiments

Groups of eleven 6-week-old female BALB/c mice (Beijing Experimental Animal 157 158 Center) were anesthetized with Zoletil 50 (tiletamine-zolazepam; Virbac S.A., Garros, France) and inoculated intranasally with 10<sup>5.5</sup> TCID<sub>50</sub> of viruses in 50 µl PBS. Three 159 mice in each group were euthanized at 3 and 5 dpi; lungs, brains, spleens, kidneys and 160 livers were collected for virus titration in MDCK cells. The remaining five mice in 161 each group were monitored for weight loss and mortality for 14 days. Mice that lost 162 more than 30% of their body weight were humanely euthanized. To determine the 163 fifty percent mouse lethal dose (MLD<sub>50</sub>), groups of three 6-week-old female mice 164 anesthetized with Zoletil 50 and inoculated intranasally with 50 µl of 10-fold serial 165 dilutions of viruses in PBS. The mice were monitored for 14 days. MLD<sub>50</sub> was 166 calculated and expressed in TCID<sub>50</sub>. For histopathology and immunohistological 167 analysis, mouse lungs and brains collected at 5 dpi were fixed in 10% 168

phosphate-buffered formalin, embedded in paraffin, then cut into 5 mm-thick sections
and stained with haematoxylin and eosin (H&E) or immunostained with a mouse
monoclonal antibody specific for influenza A virus NP (Biorbyt, UK).

172 Viral growth kinetics

Selected recombinant viruses were inoculated onto MDCK cell monolayers (at multiplicity of infection [MOI] of 0.01), A549 cell monolayers (at MOI of 0.1) or N2a cell monolayers (at MOI of 0.1) in serum-free DMEM containing 1  $\mu$ g/ml TPCK-treated trypsin and incubated at 37°C with 5% CO<sub>2</sub> atmosphere. Cell supernatants were harvested at 12, 24, 36, 48, 60 and 72 hours post-inoculation (hpi) and titrated on MDCK cells in 96-well plates. Three independent experiments were performed for each virus.

#### 180 **Polymerase activity assay**

The PB2, PB1, PA and NP gene segments of BJ27, BJ27-MA and BJ27-PB2 181 mutants were individually inserted into pCDNA3.1 plasmid. PB2, PB1, PA and NP 182 plasmids (125 ng each) were transfected to sixty percent confluent 293T cells, 183 together with fire-fly luciferase reporter plasmid pYH-Luci (10 ng) and internal 184 control plasmid expressing renilla luciferase (2.5 ng). After 24 hours of transfection, 185 cell lysate was prepared with Dual Luciferase Reporter Assay System (Promega) and 186 luciferase activity was measured using GloMax 96 microplate luminometer 187 (Promega). 188

#### 189 Western blotting

190 PB2 expression levels in different transfection groups were determined by Western

191 blotting. Total cell protein lysates were extracted from transfected 293T cells with CA630 lysis buffer (150 mM NaCl, 1% CA630 detergent, 50 mM Tris base [pH 8.0]). 192 Cellular proteins were separated by 12% sodium dodecyl sulfate-polyacrylamide gel 193 electrophoresis (SDS-PAGE) and transferred to a polyvinylidenedifluoride (PVDF) 194 membrane (Amersham Biosciences, Germany). Each PVDF membrane was blocked 195 with 0.1% Tween 20 and 5% nonfat dry milk in Tris-buffered saline and subsequently 196 incubated with a primary antibody. Primary antibodies used were specific for 197 influenza A virus PB2 (ThermoFisher, USA) and β-actin (Beyotime, China). 198 Secondary antibody was horseradish peroxidase (HRP)-conjugated anti-rabbit or 199 -mouse antibody (Beyotime, China). HRP presence was detected using a Western 200 Lightning chemiluminescence kit (Amersham Pharmacia, Freiburg, Germany), 201 202 following the manufacturer's protocols.

#### 203 Neuraminidase (NA) activity assay with substrate 4-MU-NANA

NA activity assays using the soluble substrate MUNANA (Sigma, Germany) were 204 performed as previously described (32). Briefly, virus was diluted to 10<sup>6</sup> TCID<sub>50</sub>/ml 205 and 50 µl was added to each well of a black 96-well plate (CoStar). Concentrations of 206 MUNANA substrate ranging from 2.0 µM to 200 µM were used. When cleaved by 207 the viral NA, MUNANA produces a fluorescent product. Fluorescence was quantified 208 using a Biotek Synergy H1 plate reader every 3 minutes over the course of 45 minutes. 209 Fluorescence curves were then fitted to the Michaelis-Menton equation to determine 210 values of  $V_{\text{max}}$  and  $K_{\text{m}}$ . Each experiment comprised triplicate samples of each virus. 211

#### 212 Statistical analyses

All statistical analyses were performed using GraphPad Prism Software Version 5.00 (GraphPad Software Inc., San Diego, CA, USA). Statistically significant differences between experimental groups were determined using the analysis of variance (ANOVA). Differences were considered statistically significant at P < 0.05.

#### 217 Nucleotide sequence accession numbers

The nucleotide sequences of the eight gene segments of H10N7 are available from GenBank under accession numbers: KX898962 for PB2, KX898963 for PB1, KX898964 for PA, KX898965 for HA, KX898966 for NP, KX898967 for NA, KX898968 for M, and KX898969 for NS.

222

#### 223 Results

#### 224 Adaptation of avian H10N7 influenza virus in mice

To mimic the adaptation of avian H10 subtype influenza virus in mammalian hosts, 225 A/mallard/Beijing/27/2011 H10N7 virus (BJ27) was serially passaged in murine host 226 by intranasal inoculation of  $10^{5.5}$  TCID<sub>50</sub> of virus per mouse. Mice at passage 1 (P1) 227 infected with wild type BJ27 did not show overt clinical sign. At P5, infected mice 228 showed mild clinical signs, including decreased activity and ruffled coat. At P9 and 229 P10, mice displayed severe clinical symptoms of respiratory distress, inactivity and 230 231 inappetence; all infected mice died by 5 dpi (data not shown), indicating significant 232 increase in pathogenicity. P9 virus from lung homogenate was plaque purified three times in MDCK cells and designated BJ27-MA. 233

### 234 Mouse-adapted H10N7 virus exhibited enhanced pathogenicity and 235 neurovirulence

BALB/c mice were infected, in two groups of eleven mice each, with  $10^{5.5}$  TCID<sub>50</sub> of BJ27 or BJ27-MA virus to compare virus pathogenicity. BJ27-MA virus caused 238 dramatic weight loss in infected mice and all were dead by 6 dpi, while mice infected with BJ27 showed modest weight loss of 8.7% and recovery weight gain from 7 dpi 239 (Fig. 1A and B). To determine whether the differences in pathogenicity between BJ27 240 and BJ27-MA virus were due to altered virus replication, groups of three BALB/c 241 mice were euthanized at 3 and 5 dpi respectively, and virus titers in lung and brain 242 were determined. As shown in Fig 1C and D, mouse-adapted BJ27-MA virus 243 replicated to higher titers in the lungs than wild type BJ27 virus at 3 and 5 dpi. 244 Furthermore, BJ27-MA virus was isolated from brains of infected mice in rising titers 245 246 from mean titer of 2.1 log<sub>10</sub> TCID<sub>50</sub> /ml at 3 dpi to 2.8 log<sub>10</sub> TCID<sub>50</sub> /ml at 5 dpi. No virus was isolated from brains of BJ27 virus infected mice. Therefore, the 247 mouse-adapted BJ27-MA virus has acquired neurotropism which would have 248 249 contributed to the severity of infection in mice.

#### 250 Genetic changes in adapted BJ27-MA virus

To identify potential segments and amino acid substitutions that are responsible for 251 increased pathogenicity and replication of BJ27-MA virus in mice, the consensus 252 sequence of thirty virus clones was determined. Interestingly, the most common 253 mammalian adaptation determinants of PB2-E627K and PB2-D701N (27, 28), did not 254 appear in any of the thirty clones, indicating that other viable adaptations were present 255 256 in the BJ27-MA virus. Here, five conserved amino acid mutations that could be linked 257 to increased pathogenicity were identified in 3 virus segments of the BJ27-MA virus as PB2-E158G, PB2-M631L, HA-G218E (H3 numbering), NA-K110E and 258 NA-S453I. 259

PB2-E158G mutation resides in the amino-terminal NP binding region (1–269aa)
(23), and PB2-M631L lies in the PB2-PB1 and PB2-NP interaction regions (25). HA
G218E is located near the 220-loop of the globular head HA1 domain (33).

NA-K110E and NA-S453I reside in the amino-terminal and carboxyl-terminal region
of NA protein, respectively; both are located in the interface of tetrameric structure of
NA protein (34).

### PB2 and NA segments in BJ27-MA virus conferred increased pathogenicity and replication capacity in mice

To identify virus segments from the BJ27-MA virus that confer increased 268 pathogenicity in mice, a series of recombinant viruses were generated by reverse 269 genetic based on wild type BJ27 (rBJ27) and BJ27-MA (rBJ27-MA) viruses. 270 271 Recombinant viruses rBJ27-PB2, rBJ27-HA and rBJ27-NA were constructed in rBJ27 virus background with the substituted segments of PB2, HA and NA, respectively, 272 from the rBJ27-MA virus. Mice infected with recombinant viruses were monitored 273 274 over 14 days for weight loss and survival rate. As shown in Fig. 2A and B, all of rBJ27 and rBJ27-HA viruses infected mice, similar to wild type BJ27 virus infection, 275 survived with maximum 6.7% and 8.8% weight loss respectively. By contrast, mice 276 infected with rBJ27-MA and rBJ27-PB2 viruses resulted in 25% to 31% weight loss 277 and 100% mortality by 6 dpi. rBJ27-NA virus showed moderate increase in 278 pathogenicity with 40% mortality. The MLD<sub>50</sub> values also showed the same 279 descending order of virus virulence: rBJ27-MA, rBJ27-PB2 (both MLD<sub>50</sub>, 4.75 log<sub>10</sub> 280  $TCID_{50}$  > rBJ27-NA (5.75 log<sub>10</sub> TCID<sub>50</sub>) > rBJ27 and rBJ27-HA (>6.5 log<sub>10</sub> TCID<sub>50</sub>) 281 282 (Table 1). None of these segment recombinants was neurotropic although they were recovered from lungs and extrapulmonary organs (kidney and/or spleen) (Table 1). 283 Thus, the adaptive PB2 and NA segments of BJ27-MA conferred increased virulence 284 285 in wild type BJ27 virus background in mice.

286 Combined PB2 and NA segments of BJ27-MA virus contributed to
 287 neurovirulence

Influenza virus replication in the central nervous system (CNS) often leads to fatal 288 outcome (35-37). Although mouse-adapted BJ27-MA virus was able to efficiently 289 290 replicate in murine brain, none of the above single segment recombinant viruses was found in the brain of infected mice (Table 1). Next, we generated three 291 double-segment recombinant viruses based on the rBJ27 backbone: rBJ27-PB2/HA 292 virus, rBJ27-PB2/NA virus and rBJ27-HA/NA virus. As shown in Table 2, only 293 rBJ27-PB2/NA virus was recovered from infected murine brains at 3 and 5 dpi which 294 produced MLD<sub>50</sub> value and viral loads similar to those of rBJ27-MA virus at each 295 296 time point. Viral NP was readily detected in neurons of mice infected separately with rBJ27-MA and rBJ27-PB2/NA viruses (Fig. 3A). These data demonstrated that the 297 combined PB2 and NA segments of rBJ27-MA contributed to its neurovirulence in 298 299 mice.

The ability of double segment recombinant viruses to replicate in neural tissue was assessed in mouse neuroblastoma N2a cells which has been used to study the replication of neurotropic viruses (38). Only rBJ27-MA and rBJ27-PB2/NA viruses showed up to 15-fold increased virus output relative to rBJ27 virus at 24 and/or 36 hpi (Fig. 3B). The other viruses (rBJ27-PB2/HA and rBJ27-HA/NA) showed no significant difference in virus titers at all time points. Therefore, the combined PB2 and NA segments also enhanced the replication of rBJ27-MA virus in neural cells.

# 307 PB2-M631L, PB2-E158G and NA-K110E contributed to severe BJ27-MA virus 308 infection

309 To pinpoint the contribution of the single mutations in PB2 and NA to the increased

310 pathogenicity of BJ27-MA, four point mutant viruses were generated with the rBJ27 backbone rBJ27-PB2/E158G, rBJ27-PB2/M631L, rBJ27-NA/K110E and as 311 rBJ27-NA/S453I viruses. Virus rBJ27-PB2/M631L was most virulent in that all 312 infected mice died before 8 dpi (Fig. 2C and D). Virus rBJ27-PB2/E158G and 313 rBJ27-NA/K110E caused moderate weight loss of around 13.3% without fatality. 314 Virus NA-S453I and wild type rBJ27 were least pathogenic and caused little weight 315 loss. MLD<sub>50</sub> was highest with PB2-M631L virus (4.75 log<sub>10</sub> TCID<sub>50</sub>) relative to all the 316 other viruses (>6.5 log<sub>10</sub> TCID<sub>50</sub>) (Table 1). Compared with rBJ27, the viral titers of 317 rBJ27-PB2/M631L, rBJ27-PB2/E158G and rBJ27-NA/K110E in murine lungs were 318 significantly higher at 3 and/or 5 dpi (Table 1). Virus rBJ27-PB2/M631L produced 319 the highest virus titers. However, none of the four point mutation viruses showed 320 321 extrapulmonary infection in liver, spleen, kidney or brain.

Histopathological findings of lung tissues taken at 5 dpi gave a severity picture that 322 was similar to the pathogenicity results (Fig. 2E). Virus rBJ27-PB2/M631L and 323 rBJ27-MA elicited the most severe lung lesions of edema, inflammatory infiltrates, 324 interstitial pneumonia and bronchopneumonia. Lungs from rBJ27-PB2/E158G and 325 rBJ27-NA/K110E virus infection showed less severe bronchopneumonic changes. 326 Almost no lung lesion was detected from infection with rBJ27 and rBJ27-NA/S453I 327 viruses except for some thickening of alveolar wall and mild infiltration of 328 inflammatory cells. 329

We next compared the replication of the four mutant viruses in MDCK and A549 cells, infected at MOI of 0.01 or 0.1 respectively, over 72 h. In MDCK cells (Fig. 4A),

rBJ27-MA virus showed higher output (up to 56-fold higher) than the parental rBJ27 332 from 24 to 72 hpi, and PB2/M631L mutation increased the replication of rBJ27 virus 333 334 at 36 hpi (both P < 0.05). Replication of rBJ27-PB2/E158G, rBJ27-NA/K110E and rBJ27-NA/S453I viruses was similar to that of rBJ27 virus. In A549 cells, the 335 rBJ27-MA virus also showed higher output from 24 to 60 hpi, and rBJ27-PB2/M631L, 336 rBJ27-PB2/E158G and rBJ27-NA/K110E viruses produced more progeny virus at 24 337 or 36 hpi than rBJ27 virus (all P < 0.05) (Fig. 4B). Therefore, in summary, 338 PB2-M631L, PB2-E158G and NA-K110E mutations in rBJ27 virus backbone 339 conferred more severe infection than wild type rBJ27 virus in mice and mammalian 340 cells, with PB2-M631L mutation being the most potent determinant. 341

## 342 PB2-M631L and PB2-E158G mutations enhanced polymerase activity of 343 BJ27-MA virus

PB2 is one of the components of ribonucleoprotein (RNP). RNP polymerase 344 activity has been shown to catalyze viral transcription and genomic replication, which 345 correlate with viral replication and pathogenicity in hosts (39). To evaluate whether 346 the mutations of PB2-E158G and PB2-M631L affect viral polymerase activity, we 347 generated two mutant RNP complexes under the background of the RNP of rBJ27, 348 and measured their polymerase activities in 293T cells by a luciferase minigenome 349 assay (Fig. 5). RNP polymerase activity with single E158G or M631L mutation was 350 28 or 62 times higher, respectively, than that of wild type rBJ27 RNP complex; 351 combined E158G and M631L PB2 mutations induced 75 times higher activity than 352 with the rBJ27 RNP complex (all P < 0.05). Western blotting, based on protein lysates 353

derived from 293T cells transfected with the different PB2 mutant plasmids in RNP 354 polymerase assays, showed comparable PB2 protein expression, which indicated that 355 356 the differences in polymerase activity were not due to levels of protein expression (Fig. 5). Collectively, the raised polymerase activity conferred by PB2-M631L and 357 PB2-E158G mutations correlate with their severity of virus replication in mice and 358 mammalian cells, and suggest that elevated polymerase activity, rather than protein, 359 mediated the increased replication of BJ27-MA virus; the single M631L mutation in 360 PB2 appeared as a major contributor. 361

#### 362 NA-K110E increased NA enzymatic activity

NA enzymatic activity is associated with influenza virus replication and 363 pathogenicity (40). The two amino acid mutations (K110E and S453I) in the 364 365 BJ27-MA NA protein were evaluated for NA enzymatic activity as described previously (32). Based on  $K_m$  values, we found that the NA-K110E mutation caused a 366 significant increase as did the mutant segment (rBJ27-NA with double mutations) in 367 substrate affinity (Table 3). Similarly,  $V_{\text{max}}$ , which was determined by both the specific 368 activity and the amount of enzyme in the reaction mixture, was significantly higher 369 with the K110E mutation than with wild type rBJ27 virus (P < 0.05). The  $V_{\text{max}}$  of 370 rBJ27-NA/S453I virus was increased but not significantly higher than rBJ27 virus. 371 Thus, NA-K110E mutation improved NA enzymatic activity, which would have 372 contributed to the increased replication and pathogenicity of rBJ27-MA virus in mice. 373

374 **Discussion** 

In this study, serial passage of avian H10N7 virus in mice resulted in dramatic

acquisition of pathogenicity in terms of increase virus replication, virus dissemination 376 that extended to the brain and high mortality rate. Five conserved mutations were 377 378 identified in PB2, HA and NA genes of the passaged BJ27-MA virus (PB2-E158G, PB2-M631L, HA-G218E, NA-K110E and NA-S453I). The mutations in PB2 and NA 379 genes significantly up-regulated viral polymerase activity and NA enzymatic activity 380 respectively; their combined presence in BJ27-MA virus was necessary for 381 neurovirulence. In particular, M631L mutation in PB2 was a major molecular 382 determinant for the overall increase in virulence of the mouse-adapted H10N7 virus. 383 PB2 gene plays important roles in the adaptation of influenza viruses from avian to 384 mammals through increasing polymerase activity and viral replication (41). 385 Polymerases of avian-origin generally have impaired function in human and other 386 387 mammalian cells (39). To overcome this natural restriction, avian polymerases need to acquire mutations that lead to improved activity in mammalian hosts. E627K or 388 D701N in PB2 is a common adaptive change of avian influenza viruses that cause 389 mammals and human infections (27, 28, 42). In our mice adaptation study, in place of 390 these reported mutations in the PB2 gene, E158G and M631L were identified to 391 mediate the promotion of polymerase activity, virus pathogenicity and replication in 392 mice. PB2-E158G was reported to be a pathogenic determinant of pandemic H1N1 393 and avian H5 influenza viruses in mice (24). PB2-M631L is a novel and dominant 394 pathogenic mutation not previously described. The structure of PB2 shows that 395 position 631 is close to position 627 and located at the PB2-PB1 and PB2-NP 396

397 interaction regions (22). From the isolates of human infection cases, PB2 sequence

analysis found that nine avian H5N1 and two pandemic H1N1/2009 viruses possessed
PB2-M631L but not E627K or D701N mutation, implying that PB2-M631L could be
functionally important independent of E627K or D701N. Likewise, during the
pH1N1/2009 virus outbreak in humans, the PB2-E627K mutation was absent; instead
PB2-G590S/Q591R mutation was responsible for increased polymerase activity in
human cells (43). Therefore, PB2-M631L could be a novel functional mutation in
H10N7 virus adaptation in mammalian hosts.

NA cleaves sialic acid from glycans on host cell and emerging virions, thus 405 allowing unhindered release of progeny virus from infected cells (40). Several studies 406 found that amino acid mutations or deletions in NA can affect NA enzymatic activity, 407 which correlate with virus replication and pathogenicity in vitro or in vivo (44-46). 408 409 Here, NA-K110E in BJ27-MA virus, acting as a novel mammalian mutation, significantly increased NA activity and viral replication in mice. We found that the 410 two NA mutations (K110E and S453I) are located in the interface of tetrameric 411 structure of NA (34) which may affect the formation of tetramer. 412

Neurovirulence is not commonly observed in the adaptation of avian influenza viruses in mice (21-23, 42). Besides our H10N7 virus, another related virus (H10N8) was reported to acquire neurotropism after two passages in mice (42), suggesting that H10 subtype might be more able to gain the ability to replicate in mammalian brain. Clinically, CNS disease is a common extra-respiratory complication in humans induced by influenza virus. Patients with CNS manifestations are more likely to experience severe illness (35-37). In Australian and Texas, USA, 9.7% and 8.8%

hospitalized children, respectively, infected with pH1N1/2009 virus had neurological 420 complications (47, 48). Ying et al. also found that viruses with high replication ability 421 in murine brain also possess high pathogenicity (49). The collective evidence 422 indicates that high pathogenicity and neurovirulence could cooperate to promote 423 fatalities in human cases of H10N8 virus infection. Molecular mechanism of influenza 424 virus causing infection in the CNS is unclear. HA, NA and PB2 genes have been 425 separately found to be critical to the neurovirulence of H1N1 or H5N1 viruses in 426 mammalian hosts (50-53). However, in our present study, the viruses with single 427 adapted PB2 or NA segment could not cause brain infection although when combined 428 they replicated in the brain to a level comparable with rBJ27-MA virus which 429 indicates that the synergistic effect of PB2 and NA is important for H10N7 430 431 neurovirulence.

In summary, our mouse adaptation study clearly shows that avian H10N7 virus can readily become highly virulent and neurotropic after limited passages in mice. We demonstrated that this enhanced pathogenicity was mediated by specific mutations in PB2 and NA genes; in particular PB2-M631L is a novel and critical determinant of virulence.

437

#### 438 Acknowledgement

This work was supported by the National Key Research and Development Program
(2016YFD0500204) and the National Key Technology R&D Program
(2013BAD12B01).

#### 442 **Reference**

1. Palese P. 2004. Influenza: old and new threats. Nat Med 10:S82-87.

- 444 2. Pu J, Wang S, Yin Y, Zhang G, Carter RA, Wang J, Xu G, Sun H, Wang
- 445 M, Wen C, Wei Y, Wang D, Zhu B, Lemmon G, Jiao Y, Duan S, Wang Q,
- 446 Du Q, Sun M, Bao J, Sun Y, Zhao J, Zhang H, Wu G, Liu J, Webster RG.
- 447 2015. Evolution of the H9N2 influenza genotype that facilitated the genesis of
  448 the novel H7N9 virus. Proc Natl Acad Sci U S A 112:548-553.
- 449 3. Imai M, Watanabe T, Hatta M, Das SC, Ozawa M, Shinya K, Zhong G,
- Hanson A, Katsura H, Watanabe S. 2012. Experimental adaptation of an
  influenza H5 HA confers respiratory droplet transmission to a reassortant H5
  HA/H1N1 virus in ferrets. Nature 486:420-428.
- 453 4. Gao R, Cao B, Hu Y, Feng Z, Wang D, Hu W, Chen J, Jie Z, Qiu H, Xu K,
- 454 Xu X, Lu H, Zhu W, Gao Z, Xiang N, Shen Y, He Z, Gu Y, Zhang Z,
- 455 Yang Y, Zhao X, Zhou L, Li X, Zou S, Zhang Y, Li X, Yang L, Guo J,
- 456 Dong J, Li Q, Dong L, Zhu Y, Bai T, Wang S, Hao P, Yang W, Zhang Y,
- 457 Han J, Yu H, Li D, Gao GF, Wu G, Wang Y, Yuan Z, Shu Y. 2013.
- Human infection with a novel avian-origin influenza A (H7N9) virus. N Engl
  J Med 368:1888-1897.
- 460 5. Chen H, Yuan H, Gao R, Zhang J, Wang D, Xiong Y, Fan G, Yang F, Li
- 461 X, Zhou J, Zou S, Yang L, Chen T, Dong L, Bo H, Zhao X, Zhang Y, Lan
- 462 Y, Bai T, Dong J, Li Q, Wang S, Zhang Y, Li H, Gong T, Shi Y, Ni X, Li J,
- 463 Zhou J, Fan J, Wu J, Zhou X, Hu M, Wan J, Yang W, Li D, Wu G, Feng

- Z, Gao GF, Wang Y, Jin Q, Liu M, Shu Y. 2014. Clinical and
  epidemiological characteristics of a fatal case of avian influenza A H10N8
  virus infection: a descriptive study. Lancet 383:714-721.
- 467 6. Vachieri SG, Xiong X, Collins PJ, Walker PA, Martin SR, Haire LF,
  468 Zhang Y, McCauley JW, Gamblin SJ, Skehel JJ. 2014. Receptor binding
- by H10 influenza viruses. Nature **511:**475-477.
- 470 7. Zhang W, Wan J, Qian K, Liu X, Xiao Z, Sun J, Zeng Z, Wang Q, Zhang
- 471 J, Jiang G, Nie C, Jiang R, Ding C, Li R, Horby P, Gao Z. 2014. Clinical
- 472 characteristics of human infection with a novel avian-origin influenza
  473 A(H10N8) virus. Chin Med J (Engl) 127:3238-3242.
- 474 8. Feldmann H, Kretzschmar E, Klingeborn B, Rott R, Klenk HD, Garten
- W. 1988. The structure of serotype H10 hemagglutinin of influenza A virus:
  comparison of an apathogenic avian and a mammalian strain pathogenic for
  mink. Virology 165:428-437.
- 478 9. Englund L, Hard af Segerstad C. 1998. Two avian H10 influenza A virus
  479 strains with different pathogenicity for mink (Mustela vison). Arch Virol
  480 143:653-666.
- 481 10. Wu H, Lu R, Wu X, Peng X, Xu L, Cheng L, Lu X, Jin C, Xie T, Yao H,
- 482 Wu N. 2015. Novel reassortant H10N7 avian influenza viruses isolated from
  483 chickens in Eastern China. J Clin Virol 65:58-61.
- Vijaykrishna D, Deng YM, Su YC, Fourment M, Iannello P, Arzey GG,
  Hansbro PM, Arzey KE, Kirkland PD, Warner S, O'Riley K, Barr IG,

486		Smith GJ, Hurt AC. 2013. The recent establishment of North American H10
487		lineage influenza viruses in Australian wild waterfowl and the evolution of
488		Australian avian influenza viruses. J Virol 87:10182-10189.
489	12.	Arzey GG, Kirkland PD, Arzey KE, Frost M, Maywood P, Conaty S,
490		Hurt AC, Deng YM, Iannello P, Barr I, Dwyer DE, Ratnamohan M,
491		McPhie K, Selleck P. 2012. Influenza virus A (H10N7) in chickens and
492		poultry abattoir workers, Australia, 2012/04/21 ed, vol 18, p 814-816.
493	13.	Berg M, Englund L, Abusugra IA, Klingeborn B, Linne T. 1990. Close
494		relationship between mink influenza (H10N4) and concomitantly circulating
495		avian influenza viruses. Arch Virol 113:61-71.
496	14.	Wang N, Zou W, Yang Y, Guo X, Hua Y, Zhang Q, Zhao Z, Jin M. 2012.
497		Complete genome sequence of an H10N5 avian influenza virus isolated from
498		pigs in central China. J Virol <b>86:</b> 13865-13866.
499	15.	Su S, Qi W, Zhou P, Xiao C, Yan Z, Cui J, Jia K, Zhang G, Gray GC,
500		Liao M, Li S. 2014. First evidence of H10N8 Avian influenza virus infections
501		among feral dogs in live poultry markets in Guangdong province, China. Clin
502		Infect Dis <b>59:</b> 748-750.
503	16.	Zohari S, Neimanis A, Harkonen T, Moraeus C, Valarcher JF. 2014.
504		Avian influenza A(H10N7) virus involvement in mass mortality of harbour
505		seals (Phoca vitulina) in Sweden, March through October 2014. Euro Surveill
506		19.
507	17.	Bodewes R, Bestebroer TM, van der Vries E, Verhagen JH, Herfst S,

508		Koopmans MP, Fouchier RA, Pfankuche VM, Wohlsein P, Siebert U,
509		Baumgartner W, Osterhaus AD. 2015. Avian Influenza A(H10N7)
510		virus-associated mass deaths among harbor seals. Emerg Infect Dis
511		<b>21:</b> 720-722.
512	18.	Krog JS, Hansen MS, Holm E, Hjulsager CK, Chriel M, Pedersen K,
513		Andresen LO, Abildstrom M, Jensen TH, Larsen LE. 2015. Influenza
514		A(H10N7) virus in dead harbor seals, Denmark. Emerg Infect Dis 21:684-687.
515	19.	Pan American Health Organization 2004. Avian influenza virus A (H10N7)
516		circulating among humans in Egypt.
517		http://www.paho.org/hq/dmdocuments/2010/Avian_Influenza_Egypt_070503.
518		<u>pdf</u> .
519	20.	Kayali G, Ortiz EJ, Chorazy ML, Gray GC. 2010. Evidence of previous
520		avian influenza infection among US turkey workers. Zoonoses Public Health
521		<b>57:</b> 265-272.
522	21.	Brown EG, Liu H, Kit LC, Baird S, Nesrallah M. 2001. Pattern of mutation
523		in the genome of influenza A virus on adaptation to increased virulence in the
524		mouse lung: identification of functional themes. Proc Natl Acad Sci U S A
525		<b>98:</b> 6883-6888.
526	22.	Keleta L, Ibricevic A, Bovin NV, Brody SL, Brown EG. 2008.
527		Experimental evolution of human influenza virus H3 hemagglutinin in the
528		mouse lung identifies adaptive regions in HA1 and HA2. J Virol
529		<b>82:</b> 11599-11608.

530	23.	Ping J, Dankar SK, Forbes NE, Keleta L, Zhou Y, Tyler S, Brown EG.
531		2010. PB2 and hemagglutinin mutations are major determinants of host range
532		and virulence in mouse-adapted influenza A virus. J Virol 84:10606-10618.
533	24.	Zhou B, Li Y, Halpin R, Hine E, Spiro DJ, Wentworth DE. 2011. PB2
534		residue 158 is a pathogenic determinant of pandemic H1N1 and H5 influenza
535		a viruses in mice. J Virol 85:357-365.
536	25.	Wang J, Sun Y, Xu Q, Tan Y, Pu J, Yang H, Brown EG, Liu J. 2012.
537		Mouse-adapted H9N2 influenza A virus PB2 protein M147L and E627K
538		mutations are critical for high virulence. PLoS One 7:e40752.
539	26.	Tan L, Su S, Smith DK, He S, Zheng Y, Shao Z, Ma J, Zhu H, Zhang G.
540		2014. A combination of HA and PA mutations enhances virulence in a
541		mouse-adapted H6N6 influenza A virus. J Virol 88:14116-14125.
542	27.	Gao Y, Zhang Y, Shinya K, Deng G, Jiang Y, Li Z, Guan Y, Tian G, Li Y,
543		Shi J, Liu L, Zeng X, Bu Z, Xia X, Kawaoka Y, Chen H. 2009.
544		Identification of amino acids in HA and PB2 critical for the transmission of
545		H5N1 avian influenza viruses in a mammalian host. PLoS Pathog <b>5:</b> e1000709.
546	28.	Yamayoshi S, Fukuyama S, Yamada S, Zhao D, Murakami S, Uraki R,
547		Watanabe T, Tomita Y, Neumann G, Kawaoka Y. 2015. Amino acids
548		substitutions in the PB2 protein of H7N9 influenza A viruses are important for
549		virulence in mammalian hosts. Sci Rep 5:8039.
550	29.	Reed LJ, Muench HA. 1938. A Simple Method of Estimating Fifty Per Cent
551		Endpoints. Am J Epidemiol 27: 493–497.

- 30. Ilyushina NA, Khalenkov AM, Seiler JP, Forrest HL, Bovin NV, Marjuki
  H, Barman S, Webster RG, Webby RJ. 2010. Adaptation of pandemic
  H1N1 influenza viruses in mice. J Virol 84:8607-8616.
- 555 31. Sun Y, Qin K, Wang J, Pu J, Tang Q, Hu Y, Bi Y, Zhao X, Yang H, Shu
- 556 **Y, Liu J.** 2011. High genetic compatibility and increased pathogenicity of 557 reassortants derived from avian H9N2 and pandemic H1N1/2009 influenza 558 viruses. Proc Natl Acad Sci U S A **108**:4164-4169.
- 559 32. Potier M, Mameli L, Belisle M, Dallaire L, Melancon SB. 1979.
  560 Fluorometric assay of neuraminidase with a sodium
  561 (4-methylumbelliferyl-alpha-D-N-acetylneuraminate) substrate. Anal Biochem
  562 94:287-296.
- 33. Wang M, Zhang W, Qi J, Wang F, Zhou J, Bi Y, Wu Y, Sun H, Liu J,
  Huang C, Li X, Yan J, Shu Y, Shi Y, Gao GF. 2015. Structural basis for
  preferential avian receptor binding by the human-infecting H10N8 avian
- 566 influenza virus. Nat Commun **6:**5600.
- 567 34. Sun X, Li Q, Wu Y, Wang M, Liu Y, Qi J, Vavricka CJ, Gao GF. 2014.
- 568 Structure of influenza virus N7: the last piece of the neuraminidase "jigsaw"
  569 puzzle. J Virol 88:9197-9207.
- Mizuguchi M, Yamanouchi H, Ichiyama T, Shiomi M. 2007. Acute
  encephalopathy associated with influenza and other viral infections. Acta
  Neurol Scand Suppl 186:45-56.
- 573 36. McSwiney P, Purnama J, Kornberg A, Danchin M. 2014. A severe

574 neurological complication of influenza in a previously well child. BMJ Case
575 Rep 2014.

- 576 37. Tanaka H, Park CH, Ninomiya A, Ozaki H, Takada A, Umemura T, Kida
  577 H. 2003. Neurotropism of the 1997 Hong Kong H5N1 influenza virus in mice.
- 578 Vet Microbiol **95:**1-13.
- Sajjanar B, Saxena S, Bisht D, Singh AK, Reddy GM, Singh R, Singh R,
  Kumar S. 2016. Effect of nicotinic acetylcholine receptor alpha 1 (nAChRα1)
  peptides on rabies virus infection in neuronal cells. Neuropeptides 57:59-64.
- 39. Xu G, Zhang X, Gao W, Wang C, Wang J, Sun H, Sun Y, Guo L, Zhang
- **R, Chang K-C.** 2016. Prevailing PA mutation K356R in avian influenza
  H9N2 virus increases mammalian replication and pathogenicity. Journal of
  Virology **90**:8105-8114.
- 586 40. Colman PM. 1994. Influenza virus neuraminidase: structure, antibodies, and
  587 inhibitors. Protein Sci 3:1687-1696.
- 588 41. Fan S, Hatta M, Kim JH, Halfmann P, Imai M, Macken CA, Le MQ,
- Nguyen T, Neumann G, Kawaoka Y. 2014. Novel residues in avian
  influenza virus PB2 protein affect virulence in mammalian hosts. Nat
  Commun 5:5021.
- 592 42. Zhou B, Pearce MB, Li Y, Wang J, Mason RJ, Tumpey TM, Wentworth
- 593 **DE.** 2013. Asparagine substitution at PB2 residue 701 enhances the replication,
- 594 pathogenicity, and transmission of the 2009 pandemic H1N1 influenza A virus.
- 595 PLoS One 8:e67616.

596	43.	Mehle A, Doudna JA. 2009. Adaptive strategies of the influenza viru	IS
597		polymerase for replication in humans. Proc Natl Acad Sci U S A	A
598		<b>106:</b> 21312-21316.	

599 44. Sun Y, Tan Y, Wei K, Sun H, Shi Y, Pu J, Yang H, Gao GF, Yin Y, Feng
600 W, Perez DR, Liu J. 2013. Amino Acid 316 of Hemagglutinin and the
601 Neuraminidase Stalk Length Influence Virulence of H9N2 Influenza Virus in

602 Chickens and Mice. J Virol **87:**2963-2968.

- Hossain MJ, Hickman D, Perez DR. 2008. Evidence of expanded host range
  and mammalian-associated genetic changes in a duck H9N2 influenza virus
  following adaptation in quail and chickens. PLoS One 3:e3170.
- 606 46. Chen H, Bright RA, Subbarao K, Smith C, Cox NJ, Katz JM, Matsuoka
- 607 Y. 2007. Polygenic virulence factors involved in pathogenesis of 1997 Hong
  608 Kong H5N1 influenza viruses in mice. Virus Res 128:159-163.
- 609 47. Wilking AN, Elliott E, Garcia MN, Murray KO, Munoz FM. 2014. Central
- 610 nervous system manifestations in pediatric patients with influenza A H1N1
  611 infection during the 2009 pandemic. Pediatr Neurol 51:370-376.
- 612 48. Khandaker G, Zurynski Y, Buttery J, Marshall H, Richmond PC, Dale
- RC, Royle J, Gold M, Snelling T, Whitehead B, Jones C, Heron L,
  McCaskill M, Macartney K, Elliott EJ, Booy R. 2012. Neurologic
  complications of influenza A(H1N1)pdm09: surveillance in 6 pediatric
  hospitals. Neurology 79:1474-1481.
- 617 49. Zhang Y, Zhang Q, Kong H, Jiang Y, Gao Y, Deng G, Shi J, Tian G, Liu

618		L, Liu J. 2013. H5N1 hybrid viruses bearing 2009/H1N1 virus genes transmit
619		in guinea pigs by respiratory droplet. Science <b>340</b> :1459-1463.
620	50.	Yen HL, Aldridge JR, Boon AC, Ilyushina NA, Salomon R, Hulse-Post DJ,
621		Marjuki H, Franks J, Boltz DA, Bush D, Lipatov AS, Webby RJ, Rehg JE,
622		Webster RG. 2009. Changes in H5N1 influenza virus hemagglutinin receptor
623		binding domain affect systemic spread. Proc Natl Acad Sci U S A
624		<b>106:</b> 286-291.
625	51.	Sun X, Tse LV, Ferguson AD, Whittaker GR. 2010. Modifications to the
626		hemagglutinin cleavage site control the virulence of a neurotropic H1N1
627		influenza virus. J Virol <b>84:</b> 8683-8690.
628	52.	Shinya K, Hamm S, Hatta M, Ito H, Ito T, Kawaoka Y. 2004. PB2 amino
629		acid at position 627 affects replicative efficiency, but not cell tropism, of Hong
630		Kong H5N1 influenza A viruses in mice. Virology <b>320:</b> 258-266.
631	53.	Goto H, Wells K, Takada A, Kawaoka Y. 2001. Plasminogen-binding
632		activity of neuraminidase determines the pathogenicity of influenza A virus. J
633		Virol <b>75:</b> 9297-9301.
634		
635		
636		
637		
638		
639		

640 Figure legends

FIG 1 Pathogenicity and replication of wild type (BJ27) and mouse-adapted 641 (BJ27-MA) H10N7 viruses in mice. Six-week-old female BALB/c mice were 642 inoculated with  $10^{5.5}$  TCID<sub>50</sub> of the indicated viruses or mock infected with PBS. (A) 643 Body weight changes over a 14-day period were plotted as percentage of body weight 644 at 0 dpi (n = 5 per group). Data are presented as means  $\pm$  SD of five individual mice. 645 (B) Survival data expressed as percentage of mice infected with indicated virus. Mice 646 that lost > 30% of their baseline weight were euthanized. BJ27 and BJ27-MA virus 647 titers were determined in lungs (C) and brains (D) of infected mice (n = 3 per group). 648 Data are presented as means  $\pm$  SD of three individual mice.  $\star$ , the value is 649 significantly different from that of BJ27 (P < 0.05, ANOVA). 650

651

FIG 2 Relative virulence of different recombinant and mutant BJ27 (H10N7) viruses 652 in mice. (A and C) Body weight changes in mice (n = 5) infected with  $10^{5.5}$  TCID<sub>50</sub> of 653 indicated viruses over a 14-day period, plotted as percentage of body weight at 0 dpi. 654 Data are presented as means  $\pm$  SD of five individual mice. (B and D) Survival data 655 expressed as percentage of mice infected with indicated viruses. Mice that lost more 656 than 30% of baseline weight were euthanized. (E) H&E examination was performed 657 on the lungs of mice infected with indicated viruses at 5 dpi. Virus rBJ27-PB2/M631L 658 and rBJ27-MA infection caused severe bronchopneumonia; rBJ27-PB2/E158G and 659 rBJ27-NA/K110E infections produced moderate bronchopneumonia; rBJ27 and 660 rBJ27-NA/S453I infection caused almost no lung lesion. Scale bar, 200 µm. 661

FIG 3 Neurovirulence of recombinant BJ27 (H10N7) viruses. (A) Sections of brains taken from mice 5 dpi with  $10^{5.5}$  TCID<sub>50</sub> of indicated viruses were immunostained for viral NP (open arrow). Scale bar, 400 µm. (B) Growth kinetics of recombinant viruses in neuronal N2a cells. Confluent N2a cells were infected with indicated viruses at 0.1 MOI. Data are presented as means ± SD of three independent experiments. \*, the value is significantly different from that of rBJ27 (P < 0.05, ANOVA).

668

FIG 4 Growth kinetics of recombinant H10N7 viruses in MDCK and A549 cells. Confluent MDCK (A) or A549 (B) cells were infected with viruses as indicated at MOI of 0.01 or 0.1 respectively. Data are presented as means  $\pm$  SD of three independent experiments. \*, the value is significantly different from that of rBJ27 (*P* <0.05, ANOVA).

674

FIG 5 Polymerase activity of BJ27 with different PB2 mutations in minigenome assays. Luciferase activities were relative to wild type BJ27 set at 100%. Expression of PB2 and β-actin was detected by Western blotting. Data are presented as means ± SD of three independent experiments. \*, the value is significantly different from that of BJ27 (*P* < 0.05, ANOVA).

		Average virus titer in sample <sup>a</sup>									
		Lung		Brain		Spleen		Kidney		Liver	
Virus	MLD <sub>50</sub>	3 dpi	5 dpi	3 dpi	5 dpi	3 dpi	5 dpi	3 dpi	5 dpi	3 dpi	5 dpi
	(log <sub>10</sub> TCID <sub>50</sub> )										
rBJ27	>6.5	4.5±0.3	4.7±0.1	0/3 <sup>b</sup>	0/3	0/3	0/3	0/3	0/3	0/3	0/3
rBJ27-MA	4.75	6.5±0.3*	6.9±0.3*	$2.1 \pm 0.4^{*}$	$2.8 \pm 0.6^{*}$	2.3,1.8 °*	2.3,1.8*	0/3	$2.9\pm0.3^{*}$	0/3	0/3
rBJ27-PB2	4.75	6.2±0.4*	6.6±0.3*	0/3	0/3	2.3*	0/3	0/3	$2.2 \pm 0.6^{*}$	0/3	0/3
rBJ27-HA	>6.5	5.0±0.4	5.4±0.3	0/3	0/3	0/3	0/3	0/3	0/3	0/3	0/3
rBJ27-NA	5.75	5.5±0.1*	6.1±0.3*	0/3	0/3	1.8*	0/3	0/3	0/3	0/3	0/3
rBJ27-PB2/E158G	>6.5	5.2±0.4*	5.5±0.3*	0/3	0/3	0/3	0/3	0/3	0/3	0/3	0/3
rBJ27-PB2/M631L	4.75	5.7±0.1*	6.3±0.1*	0/3	0/3	0/3	0/3	0/3	0/3	0/3	0/3
rBJ27-NA/K110E	>6.5	5.2±0.3*	5.3±0.4	0/3	0/3	0/3	0/3	0/3	0/3	0/3	0/3
rBJ27-NA/S453I	>6.5	4.9±0.4	5.3±0.1	0/3	0/3	0/3	0/3	0/3	0/3	0/3	0/3

TABLE 1 Pathogenicity and replication of BJ27 (H10N7) recombinant and mutant viruses in mice

<sup>a</sup> Mean virus titer in sample ( $\log_{10} \text{TCID}_{50}/\text{ml}$ ) ± SD. The lower limit of detection was  $10^{0.75} \text{TCID}_{50}/\text{ml}$  for each sample. \*, virus titer of corresponding strains was

significantly higher than that of rBJ27 (P < 0.05, ANOVA).

<sup>b</sup> The number of samples with recovered viruses versus the number of total collected samples.

<sup>c</sup> The number(s) shows the virus titer in an individual infected mouse.

		Average virus titer in brain <sup>a</sup>	
Virus	MLD <sub>50</sub>	3 dpi	5 dpi
rBJ27	>6.5	0/3 <sup>b</sup>	0/3
rBJ27-MA	4.75	2.1±0.4	2.8±0.6
rBJ27-PB2/HA	5.25	0/3	0/3
rBJ27-PB2/NA	4.75	1.8,2.3 °	2.5±0.5
rBJ27-HA/NA	5.5	0/3	0/3

TABLE 2 Pathogenicity and replication of double-segment recombinant H10N7 viruses in murine

TCID<sub>50</sub>/ml in the brain.

brain

<sup>b</sup> The number of samples with recovered viruses versus the number of total collected samples.

<sup>c</sup> The number(s) shows virus titer in an individual mouse.

TABLE 3 NA enzyme kinetics of mutant H10N7 viruses <sup>a</sup>

Virus	$K_m(\mu M)$	V <sub>max</sub>	$V_{\rm max}$ ratio <sup>b</sup>
rBJ27	28.7±4.1	0.53±0.10	1.00
rBJ27-MA	14.8±1.8*	$0.90{\pm}0.09$ *	$1.70^{*}$
rBJ27-NA	15.6±2.3*	$0.86 \pm 0.06$ *	1.62*
rBJ27-NA/K110E	17.8±1.5*	$0.78{\pm}0.02$ *	1.47*
rBJ27-NA/S453I	20.7±2.2	0.66±0.14	1.25

<sup>a</sup> A standardized virus dose of 10<sup>6</sup> TCID<sub>50</sub>/ml was used for the NA kinetics assay. The enzyme kinetics data (standard deviation) were fit to the Michaelis-Menten equation by nonlinear regression to determine the Michaelis constant ( $K_m$ ) and maximum velocity ( $V_{max}$ ) of substrate conversion (in fluorescent units per second). \*, the value of corresponding strains was significantly different from that of rBJ27 (P < 0.05, ANOVA).

<sup>b</sup> The ratio of the recombinant viruses versus rH10N7 virus  $V_{\text{max}}$  values.

**Figure legends** 





**FIG 1** Pathogenicity and replication of wild type (BJ27) and mouse-adapted (BJ27-MA) H10N7 viruses in mice. Six-week-old female BALB/c mice were inoculated with  $10^{5.5}$  TCID<sub>50</sub> of the indicated viruses or mock infected with PBS. (A) Body weight changes over a 14-day period were plotted as percentage of body weight at 0 dpi (n = 5 per group). Data are presented as means ± SD of five individual mice. (B) Survival data expressed as percentage of mice infected with indicated virus. Mice that lost > 30% of their baseline weight were euthanized. BJ27 and BJ27-MA virus titers were determined in lungs (C) and brains (D) of infected mice (n = 3 per group). Data are presented as means ± SD of three individual mice. \*, the value is significantly different from that of BJ27 (*P* < 0.05, ANOVA).



FIG 2 Relative virulence of different recombinant and mutant BJ27 (H10N7) viruses in mice. (A and C) Body weight changes in mice (n = 5) infected with  $10^{5.5}$  TCID<sub>50</sub> of indicated viruses over a 14-day period, plotted as percentage of body weight at 0 dpi. Data are presented as means ± SD of five individual mice. (B and D) Survival data expressed as percentage of mice infected with indicated viruses. Mice that lost more than 30% of baseline weight were euthanized. (E) H&E examination was performed on the lungs of mice infected with indicated viruses at 5 dpi. Virus rBJ27-PB2/M631L and rBJ27-MA infection caused severe bronchopneumonia; rBJ27-PB2/E158G and rBJ27-NA/K110E infections produced moderate bronchopneumonia; rBJ27 and rBJ27-NA/S453I infection caused almost no lung lesion. Scale bar, 200 µm.





**FIG 3** Neurovirulence of recombinant BJ27 (H10N7) viruses. (A) Sections of brains taken from mice at 5 dpi were immunostained for viral NP (open arrow). Scale bar, 400  $\mu$ m. (B) Growth kinetics of recombinant viruses in neuronal N2a cells. Confluent N2a cells were infected with indicated viruses at 0.1 MOI. Data are presented as means  $\pm$  SD of three independent experiments. \*, the value is significantly different from that of rBJ27 (*P* < 0.05, ANOVA).



**FIG 4** Growth kinetics of recombinant H10N7 viruses in MDCK and A549 cells. Confluent MDCK (A) or A549 (B) cells were infected with viruses as indicated at MOI of 0.01 or 0.1 respectively. Data are presented as means  $\pm$  SD of three independent experiments. \*, the value is significantly different from that of rBJ27 (*P* < 0.05, ANOVA).



FIG 5 Polymerase activity of BJ27 with different PB2 mutations in minigenome assays. Luciferase activities were relative to wild type BJ27 set at 100%. Expression of PB2 and  $\beta$ -actin was detected by Western blotting. Data are presented as means  $\pm$  SD of three independent experiments. \*, the value is significantly different from that of BJ27 (P < 0.05, ANOVA).