G1-like M and PB2 genes are preferentially incorporated into H7N9 progeny virions during genetic reassortment

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Abstract

Background: Genotype S H9N2 viruses have become predominant in poultry in China since 2010. These viruses frequently donate their whole internal gene segments to other emerging influenza A subtypes such as the novel H7N9, H5N6, and H10N8 viruses. We recently reported that the PB2 and M genes of the genotype S H9N2 virus, which are derived from the G1-like virus, enhance the fitness of H5Nx and H7N9 avian influenza viruses in chickens and mice. However, whether the G1-like PB2 and M genes are preferentially incorporated into progeny virions during virus reassortment remains unclear; whether the G1-like PB2 and M genes from different subtypes are differentially incorporated into new virion progeny remains unknown.

Results: We conducted a reassortment experiment with the use of a H7N9 virus as the backbone and found that G1-like M/PB2 genes were preferentially incorporated in progeny virions over F/98-like M/PB2 genes. Importantly, the preference varied among G1-like M/PB2 genes of different subtypes. When competing with F/98-like M/PB2 genes during reassortment, both the M and PB2 genes from the H7N9 virus GD15 showed an advantage, whereas only the PB2 gene from the H9N2 virus CZ73 and the M gene from the H9N2 virus AH320 displayed the advantage.

Conclusion: Our findings highlight the preferential and variable advantages of H9N2-derived G1-like M and PB2 genes in incorporating them into H7N9 progeny virions over SH14-derived F/98-like M/PB2 genes.

Keywords: H7N9, H9N2, G1-like M, G1-like PB2, F/98-like M, F/98-like PB2, Advantage, Reassortment

Background

Widespread reassortment of H9N2 viruses in China has created various subtypes that can be phylogenetically grouped into the A-W genotypes [1]. More than one genotype may circulate simultaneously in one region. Some genotypes became dominant over a long period of time [2]. For instance, H9N2 viruses that harbour three polymerase genes and the NP gene from the F/98-like virus plus the remaining four genes from the BJ/94-like virus form a group of viruses designated as F/98-like or H genotype (Fig. 1) [2]. F/98-like (genotype H) viruses were first identified in 1998, became dominant in 2000, and persisted for several years thereafter [2, 3]. Since 2007, the genotype S viruses, which were generated through the reassortment of the F/98-like viruses (genotype H) by substituting their M and PB2 genes with those of the G1-like virus, have emerged (Fig. 1) and gradually dominated in chicken flocks after 2010 [2–5]. Moreover, genotype S H9N2 viruses often donate some or all of the six internal genes to other emerging influenza

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A viruses in China [2, 6] such as the novel H7N9, H10N8, H7N7, and H5N6 viruses [2, 7–9]. That is, the H9N2 and H7N9 viruses currently circulating in China both possess G1-like M and PB2 genes [10].

The S genotype, which differs from F/98-like viruses (genotype H) only in their M and PB2 genes [11, 12], has not been replaced by a new genotype since 2010 [2]. As S genotype viruses carry the genetic backbone of F/98-like viruses (genotype H) plus the M and PB2 genes of the G1-like viruses, it is presumed that the G1-like M and PB2 genes confer better viral fitness over F/98-like counterparts.

Indeed, Pu et al. reported that H9N2 viruses with the G1-like M gene replicate faster in primary chicken embryonic fibroblasts and chickens than do the H genotype viruses. Furthermore, the H9N2 virus with the G1-like M gene exhibit an early surge of viral mRNA and genomic RNA production, suggesting of increased fitness of the virus [13]. Consistently, Hao et al. found that H5Nx and H7N9 viruses harbouring the G1-like PB2 and M genes display better viral fitness than those with F/98-like PB2 or M genes and have high virulence and replication capacity in chickens and mice [11, 12].

Our present study aims to determine if G1-like M and PB2 genes hold a competitive advantage during genetic reassortment, whether they play a role in maintaining the stability of “gene cassette” in H7N9 viruses. Several representative H7N9 and H9N2 viruses were chosen to test the relative copy number of G1-like M/PB2 genes and F/98-like M/PB2 genes in reassortant viruses on the H7N9 genetic background. The TaqMan-MGB quantitative realtime PCR (qRT-PCR) approaches for accurately quantifying the heteroplasmia level of G1-like M/PB2 and F/98-like M/PB2 were introduced in our study. The MGB probes had higher melting temperature. Therefore, they are much shorter than the traditional TaqMan probes, which makes MGB probes more sensitive to single base mismatches [14, 15]. Multiple studies have demonstrated that TaqMan-MGB qRT-PCR is an accurate technique with high specificity, sensitivity and remarkable reproducibility and is quite attractive for use in SNP (single nucleotide polymorphism) detection and allelic discrimination [16, 17].

Our results suggests that the G1-like M and PB2 genes are more likely to be incorporated into the novel H7N9 viruses than that of SH14 virus derived F/98-like M/PB2 genes; G1-like PB2/M genes derived from different virus strains display variable competitive advantages during virus reassortment.

### Results

#### The sensitivity and specificity of duplex real-time RT-qPCR assay

We first evaluated the sensitivity of duplex RT-qPCR by using ten-fold serially diluted plasmid mixtures as the templates in the amplification reaction. As shown in Supplementary Fig. S1A & B and Fig. S2A & B, each gene could be readily amplified with approximately 10 copies of templates when crossing point was less than 35(cp < 35). The standard curves revealed excellent correlation coefficient and amplification efficacy (Fig. S1 C-F, Fig. S 2 C-F and Fig. S 3 A-D) when cp < 35. Moreover, there was no significant difference in amplification efficiency among the probes. To determine the
specificity of duplex RT-qPCR used in this study, the G1-like M/PB2 plasmids from GD15, CZ73, AH320 viruses and the F/98-like M/PB2 plasmids from genotype H H9N2 SH14 virus were used as templates for the amplification reactions. As shown in Supplementary Fig. S4, the M/PB2 genes from GD15, CZ73 and AH320 could not be detected with SH14-Mprobe974 or SH14-PB2probe713 whereas M/PB2 genes of SH14 virus could not be detected by using GD15-Mprobe974 or PB2probe974RC. G1-like M and PB2 genes from GD15 (H7N9) virus hold a competitive advantage during reassortment

To determine whether the G1-like M segment was preferentially incorporated into reassorted virus progeny when it was in competition with F/98-like M gene, 293 T cells were co-transfected with eight plasmids of the GD15 virus plus the ninth one encoding FSH14-M gene (500 ng/plasmid) (Fig. 2a) [18]. After incubation for 72 h, the conditioned media of transfected cells, which contained the reassorted progeny virions, were used to inoculate into embryonated chicken eggs. Quantitative RT-qPCR (qRT-PCR) analysis of allantoic fluids revealed that the copy number of the g1GD15-M gene was significantly greater than that of the F98SH14-M gene (Fig. 2b). However, when just 250 ng of the g1GD15-M plasmid was repeated for the above mentioned 9-plasmid transfection, approximate level of gene copies was displayed between g1GD15-M and F98SH14-M (Fig. 2c).

We next determined whether the G1-like PB2 gene also exhibited competitive advantage during genetic reassortment. Co-transfection experiments with the nine-plasmid system were similarly carried out as above. As shown in Fig. 2d, the copy number of the g1GD15-PB2 gene was significantly higher than that of the F98SH14-PB2 gene in the viruses rescued from the cells transfected with 500 (Fig. 2e) or 250 ng (Fig. 2f) plasmid each. The competitive advantage of the G1-like PB2
gene from GD15 virus was more prominent than that of the G1-like M gene.

**Variable advantage of G1-like M/PB2 genes from different strains during reassortment**

We next investigated if the competitive advantage of the M and PB2 genes from G1-like viruses in reassortment was strain-specific. The G1-like M and PB2 genes from two additional S genotype H9N2 viruses, A/Chicken/Jiangsu/CZ73/2014 (CZ73) and A/Chicken/Anhui/AH320/2016 (AH320), were used in co-transfection experiments as described above. As shown in Fig. 3a, b, the copy number of the g1CZ73-PB2 gene was significantly higher than that of the f98SH14-PB2 gene in the rescued viruses (Fig. 3c). However, there was no significant difference in the copy number of the M gene in reassortant viruses rescued from co-transfection with the g1CZ73-M and f98SH14-M genes (Fig. 3d). We then investigated the effect of the internal gene cassette of the CZ73 virus on virus reassortment. Co-transfections with six plasmids encoding the internal genes of CZ73 virus plus the plasmid encoding the f98SH14-PB2 or f98SH14-M (500 ng/plasmid) (Fig. 3e, f) revealed that the copy number of g1CZ73-PB2 genes was significantly higher than that of f98SH14-PB2 gene (Fig. 3g), although g1CZ73-M genes did not exhibit any competitive advantages (Fig. 3h).

We then conducted a similar co-transfection experiment by using the M or PB2 gene of another S genotype H9N2 virus, AH320 (Fig. 3i, k). qPCR analysis revealed that the copy number of the g1AH320-M gene in the rescued viruses was significantly higher than that of the f98SH14-M gene (Fig. 3j). The copy number of the g1AH320-PB2 gene in rescued viruses was higher than that of the f98SH14-PB2 gene. However, this was not statistically significant (Fig. 3l).

**Lack of competitive advantage for the g1GD15-M gene incorporation into the H7N9 background over f98SH14-M gene during co-infection**

We next determined if the competitive advantage of G1-like M and PB2 genes during reassortment was in part...
due to faster replication of newly reassorted viruses. We generated four recombinant viruses (WT-GD15, GD15-M\textsubscript{SH14}, GD15-PB2\textsubscript{SH14} and GD15-M\textsubscript{SH14}PB2\textsubscript{SH14}) (Fig. 4a-d). As shown in Fig. 4e, the titers of WT-GD15 virus at 48 h post infection (h.p.i) were significantly higher than three recombinant viruses, which gave similar virus titers in the conditioned media. Quantitative RT-PCR analysing the M gene revealed almost equal g\textsubscript{1}GD15-M and f\textsubscript{98}SH14-M vRNA levels at 48 hpi (Table 1). We then co-infected MDCK cells with GD15-M\textsubscript{SH14} and WT-GD15 viruses, each with 0.01 MOI (Fig. 5a). Again, quantitative RT-PCR revealed that the levels of the g\textsubscript{1}GD15-M and f\textsubscript{98}SH14-M genes in the conditioned media were not significantly different (Fig. 5b). To further investigate whether PB2 genes affected the competitive advantage of M genes at the virus level, we analysed the portion of g\textsubscript{1}GD15-M gene in progeny virions by co-infecting MDCK cells with recombinant GD15-PB2\textsubscript{SH14} and GD15-M\textsubscript{SH14}PB2\textsubscript{SH14} viruses (Fig. 5c). However, the copy number of g\textsubscript{1}GD15-M genes in progeny virions did not demonstrate significantly advantage over the f\textsubscript{98}SH14-M gene (Fig. 5d).

Preference for the g\textsubscript{1}GD15-PB2 gene incorporation into the H7N9 background over f\textsubscript{98}SH14-PB2 gene during co-infection

MDCK cells were co-infected with GD15-PB2\textsubscript{SH14} and WT-GD15 viruses (0.01 MOI each) (Fig. 6a). qRT-PCR analysis revealed that the levels of the G1-like PB2 gene in the progeny viruses was significantly higher than that of the F/98-like PB2 gene (Fig. 6b). Given that these two viruses replicate differentially, 0.005 moi WT-GD15 virus was used in co-infection experiments. We found that at this dosage the two PB2 genes replicated at a comparable rate at 48 hpi (Table 1). Besides, another conserved gene-NP segment were detected to verify whether the changes in the levels of M and PB2 are true. The results showed that at this dosage the number of NP gene copies in progeny viruses were similar among the reassortant viruses, too (Table 1). In the coinfection experiment the number of WT-GD15 virus used was only half of that GD15-PB2\textsubscript{SH14}, the g\textsubscript{1}GD15-PB2 gene still demonstrated significant advantages (Fig. 6 C).

Since genotype S H9N2 strains are generated through the replacement of the M and PB2 genes of F/98-like viruses with those from the G1-like viruses, we wondered if the M or PB2 genes would influence each other’s preference. MDCK cells were coinfected with GD15-M\textsubscript{SH14} and GD15-M\textsubscript{SH14}PB2\textsubscript{SH14} viruses (Fig. 6 D). The copy number of the g\textsubscript{1}GD15-PB2 gene were significantly higher than that of f\textsubscript{98}SH14-PB2 genes in the progeny viruses (Fig. 6e). However, the g\textsubscript{1}GD15-PB2 to f\textsubscript{98}SH14-PB2 ratio in this co-infection experiment was similar to that co-infected with g\textsubscript{1}GD15-PB2\textsubscript{SH14} and WT-GD15 viruses (date not shown), suggesting that the competitive advantage of g\textsubscript{1}GD15-PB2 to f\textsubscript{98}SH14-PB2 during reassortment is not influenced by the M gene.
Discussion

If two homologous gene segments are available in a cell, they will compete with other for incorporation into progeny viruses [19]. We performed co-transfection and co-infection experiments with the M or PB2 gene derived from the G1-like and F/98-like genotypes. During reassortment, the M or PB2 gene from two genotypes would compete with other for their incorporation into progeny virions. Our co-transfection experiments showed that the copy number of \( g_1 \)GD15-M/\( g_1 \)GD15-PB2 genes were higher in progeny virions than that of \( f_98 \)SH14-M/\( f_98 \)SH14-PB2 genes. These observations suggest that there is a biased genetic reassortment between G1-like M/PB2 and F/98-like M/PB2 genes. The advantage of the \( g_1 \)GD15-PB2 but not \( g_1 \)GD15-M gene in incorporating into progeny virions was confirmed in our co-infection experiment. The discrepancy in the results obtained from co-transfection and co-infection experiments are likely due to the differences in the materials and methods used in the study. In addition, the interactions between viruses are more complex than plasmids. Consistent with our observations, Essere and Kawaoka reported that co-transfection and co-infection result in different reassortant genotypes [18, 20, 21]. These authors postulated that some unknown cellular factors may affect genetic reassortment [18].

The advantage of the G1-like M/PB2 genes over that of the F98-like M/PB2 genes were further investigated by using two additional genotype S H9N2 viruses, CZ73 and AH320. We found that the incorporation advantage of the G1-like M/PB2 gene was variable among strains. The findings that only the PB2 gene from the H9N2 virus CZ73 and the M gene from the H9N2 virus AH320 had an advantage suggest that the competitive advantage is not equal for the M and PB2 genes from one virus. We speculate that the internal genes of a novel H7N9 virus are not necessarily from one H9N2 virus but rather from a super recombination of genes from different H9N2 viruses.

The segmented genome of influenza viruses allows for the reassortment of gene segments between viruses when they co-infect same cells [22, 23], resulting in multiple genotypes of influenza viruses [23–25]. Nevertheless, the substitution of the G1-like M and PB2 genes

## Table 1 Number of copies of M/PB2 gene (log_{10})

| Reassortant virus | WT-GD15 (0.01 MOI) | WT-GD15 (0.005 MOI) | GD15-PB2SH14 (0.01 MOI) | GD15-MSH14 (0.01 MOI) | GD15-MSH14PB2SH14 (0.01 MOI) |
|------------------|---------------------|---------------------|-------------------------|------------------------|-------------------------------|
| Copies of PB2 gene (log_{10}) | 11.14 ± 0.15 | 10.33 ± 0.06 | 10.29 ± 0.06 | 10.56 ± 0.05 |
| Copies of M gene (log_{10}) | 11.18 ± 0.12 | 10.74 ± 0.20 | 11.37 ± 0.10 | 11.11 ± 0.07 |
| Copies of NP gene (log_{10}) | 9.92 ± 0.35 | 9.83 ± 0.09 | 10.16 ± 0.10 | 9.68 ± 0.36 |

MDCK cells were infected with each virus alone at 0.01 or 0.005 MOI and incubated for 48 h. The replication level of PB2, M and NP gene were detected by qRT-PCR. Results represent the average copies from three independent infections.
has reduced the genetic diversity of H9N2 viruses [8, 13]; S genotype H9N2 viruses have been predominant in chickens since 2010 [2–4]. We speculate that the uniqueness of the internal gene cassette of the S genotype makes it possible to reach a more ideal equilibrium.

Our study has demonstrated an advantage of G1-like M and PB2 genes over F/98-like M and PB2 genes. It is not clear why the internal genes of the S genotype H9N2 virus remain stable as a cassette and has stayed prevalent in H9N2 and H7N9 viruses for many years. The genotype S H9N2 viruses provide their internal genes to various emerging viruses, especially H7N9 viruses [1, 5, 10, 13, 26]. Epidemiological evidence suggests that nearly all human and avian H7N9 isolates possess internal genes that originated from H9N2 viruses [8, 27, 28]. It is likely that the six internal genes of genotype S H9N2 viruses have reached a stable and optimal combination, ensuring the donation of the internal gene cassette to emerging viruses. On the other hand, we also observed gene segments that originated from H9N2 viruses in other viruses, such as H5N2 and H7N7 viruses but not as a whole cassette [4]. The uniqueness of internal genes of H9N2 viruses, especially those of genotype S, warrants further exploration.

**Conclusions**

Our study has demonstrated that G1-like PB2/M genes had competitive advantages over SH14 virus-derived F/98-like PB2/M gene during reassortment. However, their competitive advantage varied among different strains. The competitive advantage of the PB2 gene was more prominent than that of the M gene. Our results suggest that the preferential incorporation of H9N2-derived G1-like M and PB2 genes into progeny virions of H7N9 influenza viruses may help maintain the stability of the internal gene cassette in the novel H7N9 viruses.

**Methods**

**Cells, eggs and viruses**

Human embryonic kidney (293 T) cells and Madin-Darby canine kidney (MDCK) cells were stored by our laboratory and maintained in DMEM supplemented with 10% foetal bovine serum (FBS) and penicillin-streptomycin. The SPF embryonated chicken eggs were
purchased from Beijing Merial Vital Laboratory Animal Technology Company.

The H7N9 viruses A/Chicken/Guangdong/GD15/2016 (GD15, whose internal genes are all genotype S, Fig. 1), the S genotype H9N2 viruses A/Chicken/Jiangsu/CZ73/2014 (CZ73, Fig. 1) and A/Chicken/Anhui/AH320/2016 (AH320, Fig. 1) and the H genotype H9N2 viruses A/Chicken/Shanghai/14/2001 (SH14, Fig. 1) used in this study were isolated and preserved by our laboratory.

Plasmid construct
The 8-plasmid reverse genetic systems of GD15(H7N9) virus, CZ/73(H9N2) virus and AH320(H9N2) virus were constructed in the present study, while the 8 plasmids for SH14(H9N2) virus was constructed as previously reported [29]. All constructs were verified by sequencing and preserved by our laboratory.

Nine plasmid co-transfection
Six plasmids containing PB1, PA, NP, NS, HA, and NA genes of GD15 (H7N9) virus, plus PB2/M plasmid from GD15/CZ73/AH320 virus and PB2/M plasmid from SH14 (F98-like H9N2) virus were cotransfected into 293 T and MDCK cells to examine the competition between G1-like and F98-like PB2. The supernatants were collected after 72 h for inoculation in 9–10 day-old egg. One egg was used for each sample. 36 h post inoculation the allantoic fluid was collected and stored in −70 °C. cotransfection experiment was run in triplicate wells and repeated at least twice for each sample.

Reverse genetics
Reassortant viruses were generated by reverse genetics as previously described [30]. Plasmids were transfected into 293 T and MDCK cells using Polyfect transfection reagent (Qiagen) according to the manufacturer’s instructions. After 72 h, supernatants were harvested and each of the newly generated viruses was plaque purified. The purified viruses were amplified on SPF embryonated chicken eggs to generate the viral stock used in the study.

Viral growth kinetics
The growth properties of reassortant viruses were assessed as follows. Triplicate wells containing MDCK cells were infected with the indicated viruses at a multiplicity of infection (MOI) of 0.01, supplemented with Opti-MEM (catalog no. 31985–070; Gibco) and incubated at 37 °C. Supernatants were collected from each well at 12, 24, 48 and 72 h post infection (h.p.i.) and were stored at −70 °C. Viral titres were subsequently determined as the 50% tissue culture infection dose (TCID50) per 0.1 ml in MDCK cells using the method of Reed and Muench [31].

Coinfection of MDCK cells with Reassortant viruses
MDCK cells were coinfected with viruses at 0.01 or 0.005 MOIs. After incubation at 37 °C for 1 h, the virus inoculum was removed, and the cells were washed three times with phosphate-buffered saline (PBS), followed by incubation at 37 °C in Opti-MEM. After 48 h, supernatants were collected from each well for subsequent analysis. Every experiment was run in triplicate wells and repeated at least twice.

Quantitative real-time RT-PCR screening of gene origins
Total RNA was extracted from the allantoic fluid or cell culture supernatant using a Solarbio RNA extraction kit (Solarbio, Shanghao, China). And the extracted RNA for all samples were diluted to 100 ng/ul to guarantee equal amount of total RNA was used for the downstream RT-PCR. DNA was cleared by digesting 2 μl of vRNA with 4 μL of 4× gDNA wiper Mix (Vazyme Biotech, Nanjing, China) and 10 μL of water for 2 min at 42 °C. Next, unique 12 bp primer (′-AGCAAAAGCAGG-3′) was used to perform reverse transcription. To identify gene origins, we designed a pair of primers (M-F948, and M-R1012) in the same sequences of p98SH14-M, p1GD15-M, p1CZ73-M and p1AH320-M, as well as three MGB probes, including SH14-M probe974 (against p98SH14-M), GD15-M probe974 (against p1GD15-M), and M probe975−17 (against p1CZ73-M and p1AH320-M) in the different sequences of p98SH14-M, p1GD15-M, p1CZ73-M and p1AH320-M between the two primers. Three probes were used for quantitative real-time RT-PCR to quantitatively detect gene copies of p98SH14-M, p1GD15-M, p1CZ73-M, and p1AH320-M. Likewise, we designed a pair of primers and three MGB probes, PB2-F650, PB2-R772, SH14-PB2 probe713, GD15-PB2 probe713 and PB2 probe748 (against p1CZ73-M and p1AH320-M), to quantify copies of p98SH14-PB2, p1GD15-PB2, p1CZ73-PB2 and p1AH320-PB2. The specificity of each probe has been verified as early as we received the products. The sequences of the primers and probes are shown in Table 2.

Quantitative real-time PCR was carried out in a 20 μL reaction mixtures containing 1 μL of cDNA, 250 nM each primer, 100 nM FAM probes, 100 nM VIC probe, 10 μL of 2× AceQ U+ Probe Master Mix (Vazyme Biotech, Nanjing, China), and water. The qPCR conditions were as follows: 37 °C for 2 min and 95 °C for 5 min, followed by 45 cycles of 95 °C for 10 s, 60 °C for 30 s and 40 °C for 2 min. The plasmids pHW2000-MSH14, pHW2000-MGD15, pHW2000-MCZ73, pHW2000-MAH320, pHW2000-PB2SH14, pHW2000-PB2GD15, pHW2000-PB2CZ73 and pHW2000-MAH320 were used as standards for the M/PB2 genes of SH14, GD15, CZ73 and AH320 viruses, respectively.
Table 2 Primers and MGB probes used in qPCR in this study and targeting genes they used to detect

| Primer /probe   | Sequence(5’ to 3’)            | Gene detected                             |
|----------------|--------------------------------|------------------------------------------|
| M-F948         | TGGCTGAGTCTATGAGGGAAG           | GD15-M, SH14-M, CZ73-M, AH320-M          |
| M-R128         | ACCATCGTCAACATCCACAG            | GD15-M, SH14-M, CZ73-M, AH320-M          |
| SH14-M probe974| FAM-CGGCAGGAGCAACAGAG-MGB       | SH14-M                                   |
| GD15-M probe974| VIC-CAGCATTCTGCCTGCTTCT-MGB     | GD15-M                                   |
| M probe975-17  | VIC-AGGACTTCTGCTGTCTC-MGB       | CZ73-M, AH320-M                          |
| PB2-F650       | TTGCTCTTTAATGTGGGC              | GD15-PB2, SH14-PB2, CZ73-PB2, AH320-PB2 |
| PB2-R772       | AGGCCCTTGAATCAAATGC             | GD15-PB2, SH14-PB2, CZ73-PB2, AH320-PB2 |
| SH14-PB2 probe713 | FAM-CGGTAGCAGGTGGAGCAA-MGB     | SH14-PB2                                 |
| GD15-PB2 probe713 | VIC-CAGTGCTGAGGGAGGACA-MGB | GD15-PB2                                 |
| PB2 probe974RC | VIC-TTGTCCTCCAGCTACTGG-MGB      | CZ73-PB2, AH320-PB2                      |

Statistical analysis
Statistical analyses were conducted by using SAS software, version 9.2 (SAS Institute). Statistically significant differences between the number of the copies of G1-like M/PB2 and F/98-like M/PB2 genes were analysed by using Duncan's multiple range test in ANOVA. Differences were considered significant at P < 0.05.

Supplementary Information
The online version contains supplementary material available at https://doi.org/10.1186/s12917-021-02786-0.

Additional file 1: Figure S1. Sensitivities and standard curves of the duplex TaqMan-MGB qRT-PCR targeting the M genes of SH14 and GD15 viruses. Sensitivities of SH14-M probe975 (A) and GD15-M probe974 (D) were detected by 10-fold serial dilutions of 10^3 copies/ml of SH14-M plasmid and 10^4 GD15-M plasmid, respectively. The detection limit was approximately 10 copies of both gene when cp < 35. The amplification curves and the corresponding standard curve for detection of 10^3 SH14-M (B, C) and 10^4 GD15-M (E, F) showed excellent efficiencies of the duplex TaqMan-MGB qRT-PCR.

Additional file 2: Figure S2. Sensitivities and standard curves of the duplex TaqMan-MGB qRT-PCR targeting the PB2 genes of SH14 and GD15 viruses. Sensitivities of SH14-PB2 probe713 (A) and GD15-PB2 probe713 (D) were detected by 10-fold serial dilutions of 8*10^3 copies/ml of SH14-PB2 plasmid and 10^3 GD15-PB2 plasmid, respectively. The detection limit was approximately 10 copies of both gene when cp < 35. The amplification curves and the corresponding standard curve for detection of 10^3 SH14-PB2 (B, C) and 10^4 GD15-PB2 (E, F) showed excellent efficiencies of the duplex TaqMan-MGB qRT-PCR.

Additional file 3: Figure S3. The amplification curves and the corresponding standard curve for detection of 10^3 CZ73-M and 10^4 AH320-M gene (A, B), and 10^3 CZ73-PB2 and 10^4 AH320-PB2 (CD) gene showed excellent efficiencies of the duplex TaqMan-MGB qRT-PCR.

Additional file 4: Figure S4. The specificity of duplex MGB TaqMan-probe-based real-time RT-qPCR. The M/PB2 genes from GD15, CZ73, and AH320 can’t be detected by SH14-Mprobe974 or SH14-PB2 probe713, and M/PB2 genes from SH14 virus can’t be detected by GD15-Mprobe974, Mprobe975-17, GD15-PB2 probe713 or PB2 probe974RC, either.

Abbreviations
293 T cells: Human embryonic kidney; MDCK cells: Madin-Darby canine kidney cells; DMEM: Dulbecco’s minimum essential medium; FBS: Fetal bovine serum; TCID50: The 50% tissue culture infection dose; MOIs: Multiplicity of infection; PBS: Phosphate-buffered saline

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Not Applicable.

Authors’ contributions
XLL and MG designed this study, XLL, RYG, XXB, XLH, and JM performed the experiments. XLL and XFL drafted the manuscript. MG, JH, SLH, XQW, XLW, SJ, DXP, XAJ, and XFL supervised all the experiments and participated in the data analysis. LXL, MG and XFL discussed and prepared the final report. All of the authors have read and approved the final manuscript.

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Availability of data and materials
The datasets generated and analyses during the current study are available from the corresponding author on request.

Ethics approval and consent to participate
No animal was involved in the experiments. All experiments involving live virulent H7N9 viruses were housed in negative-pressure isolators with HEPA filters in animal biosecurity level-3 facilities in accordance with the institutional bio-safety manual.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

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