Diversity and reassortment rate of influenza A viruses in wild ducks and gulls

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Abstract: Influenza A viruses (IAVs) evolve via point mutations and reassortment of viral gene segments. The patterns of reassortment in different host species differ considerably. We investigated the genetic diversity of IAVs in wild ducks and compared it with the viral diversity in gulls. The complete genomes of 38 IAVs of H1N1, H1N2, H3N1, H3N2, H3N6, H3N8, H4N6, H5N3, H6N2, H11N6 and H11N9 subtypes isolated from wild mallard duck and gull habitants of a pond in Moscow city, Russia were sequenced. The sequences were closely related to those of avian IAVs isolated in Sweden and the Netherlands. The analysis of phylogenetic trees showed that stable viral genotypes do not persist from year to year in ducks owing to frequent gene reassortment. For comparison, similar analyses were carried out using sequences of IAVs isolated in the same period from ducks and gulls in the Netherlands. Our results revealed a significant difference in diversity and rates of reassortment of IAVs in ducks and gulls.

Keywords: Avian influenza; Reassortment; Diversity

1. Introduction

Wild aquatic birds of the orders Anseriformes (ducks, geese and swans) and Charadriiformes (gulls, terns, waders and plovers) are natural hosts of IAVs [1]. In these birds, 16 antigenic subtypes of hemagglutinin (HA) and 9 antigenic subtypes of neuraminidase (NA) in various combinations were found [2]. IAVs with the H1- H12, H14 and H15 HA subtypes mainly circulate in ducks. In ducks, these IAVs cause asymptomatic infections, replicate in the intestine and transmit by the fecal-oral route through the water [1]. IAVs with the H13 and H16 HA subtypes are usually found in gulls [3, 4]. The viruses replicate in the intestine and respiratory tract; chicks in densely populated breeding grounds are mainly infected [5]. Thus, the circulation of AIV in gulls differs from circulation in ducks, in particular with respect to this latter point.
IAVs in domestic poultry evolve from IAVs of wild aquatic birds. IAVs of poultry can occasionally transmit and initiate host-specific evolutionary lineages in pigs, humans and other mammals. Long-term adaptation in ducks enables efficient replication and release of the virus into the environment without significant pathogenic effects to the host. In gulls, the virus also replicate asymptotically, while in poultry it may evolves towards increased pathogenicity. In mammals, the influenza virus usually causes non-fatal disease.

The evolution of IAVs is caused by mutations and reassortment of viral genes. Reassortment occurs when a single host cell is infected with different IAVs [6]. Reassortment plays a central role in the microevolution of influenza viruses. In wild ducks, there is a continuous mixing of all eight genome segments due to reassortment. This process may be less intense in secondary hosts.

Reassortment between IAVs of different hosts, including human viruses, contributes to initiation of influenza pandemics [7]. However, once emerged, human IAVs of a novel lineage evolve as a whole, without reassortment. Only intra-subtype reassortants between co-circulating clades are viable and persist in the population on a small spatial-temporal scale [8-12]. Rarely, reassortants between H1N1 and H3N2 were found. For example, prior to the 2009 pandemic, clinical IAV isolates of the H1N2 subtype were described [13-16]. However, all these cases are rather the exception to the rule. Such reassortants are not sufficiently viable [17]. Studies of H1N2 reassortants in the human population showed that they are under negative selection [18, 19].

Reassortment of IAVs in pigs is an example of the formation of new variants. Pigs were called the “mixing vessel” of IAVs because they can be infected with both avian and human viruses and enable their reassortment [20]. The IAVs that had been formed in pigs initiated the 2009 pandemic and, possibly, other pandemics. In recent decades, numerous new lineages of reassortant viruses were discovered in pigs [21-24]. However, IAVs of the same evolutionary lineage can circulate in pigs for many years. The “classical” lineage of swine IAVs, originating from the precursor of the 1918 “Spanish” influenza, remained genetically stable until the 1980s [25, 26]. In horses, two lineages of influenza viruses evolved for decades [27].

Several lineages of highly pathogenic and low pathogenic IAVs circulate in chickens. Asian H5N1 and H7N9 IAVs are reassortants. They all carry a cassette of internal genes derived from chicken H9N2 viruses [28, 29]. However, after initial formation, such variants circulate largely without reassortment. Chicken viruses of the H5 and H9 subtypes circulate in Egypt together, and co-infection with these viruses was often found. Nevertheless, reassortment of these subtypes has not been reported yet [30]. The appearance of highly pathogenic viruses of antigenic subtypes H5N8, H5N3 and H5N6 is associated with the entry of H5N1 viruses into ducks, in which reassortment took place [31-33].

The reassortment of IAVs in their natural animal hosts has a different pattern than in all secondary host species. It is impossible to detect evolutionary lineages of complete genomes for duck IVs, as the viruses circulates in ducks in the form of a large pool of functionally equivalent segments of the genome that form temporary groupings, without strong selective pressure to preserve this composition [34].

Gulls represent a special host of IAVs. IAVs of almost all subtypes were found in gulls. However, the H1-H12 and H14 subtypes were rarely isolated from gulls typically as single specimens and, as a rule, were close to the duck viruses circulating the same location at the same time. All these cases were can be classified as
single cases of spillover to gulls, since these IAVs did not persist in the new host. At the same time, H13 and H16 viruses have almost exclusively been detected in gulls and terns [35-37].

Gulls are usually considered natural hosts of influenza viruses along with ducks. However, there are large differences in the circulation patterns of duck and gull viruses. A seemingly stable core consisting of PB2, PA, M, NS, NP and PB1 gene segments has formed in gulls and may persist for at least four years [38]. [Hall et al., 2013]. All gene segments of gull H13 and H16 IAVs are phylogenetically distant from segments of duck IAVs [35, 36, 39].

Recent studies have shown that reassortment can occur in any living system (in cell culture, chicken embryos, laboratory animals) if the dose of infection is large enough to provide simultaneous access of different viral particles to the host cell [40-42]. However, the emergence of new variants of IAVs via reassortment in some hosts, such as humans, occurs once in about a decade, while in other hosts, such as ducks, it occurs continuously. This notion raises the key question of whether or not the rate of reassortment of IAVs differs significantly between viral host species [6].

The aim of our work was to compare the IAV reassortment rate in wild ducks and gulls. From 2006 to 2019, we isolated IAVs from wild birds resting in a city pond in Moscow, Russia during their autumn migration. Thirty-seven duck IAVs and one duck-origin spillover H6N2 gull isolate were fully sequenced. We did not find stable IAV genotypes perpetuated from year to year. For comparison, a similar study on genome constellations was carried using sequences of duck and gull IAVs isolated in the same years in the Netherlands.

2. Materials and Methods

2.1. Viruses

Fresh feces of gulls and ducks were collected in 2006–2019 on the shore of a city pond in Moscow. Feces were suspended in 1 ml of phosphate-buffered saline (PBS) supplemented with 0.4 mg/ml gentamicin, 0.1 mg/ml kanamycin, 0.01 mg/ml nystatin, and 2% MycoKill AB solution (PAA Laboratories GmbH, Pasching, Austria). The suspension was centrifuged for 10 min at 4000 rpm, and 0.2 ml of the supernatant were inoculated into 10-day-old chicken embryos. Allantoic fluid was collected after 48 h and tested by hemagglutination assay with chicken red blood cells. Positive samples were taken for further passaging. All isolated IAV strains are stored in the virus repository of the Chumakov Federal Scientific Center (Moscow, Russia). Full names, designations of the viruses and GenBank accession numbers are given in Table S1 of supplemental material.

2.2. Sequencing

Viral RNA was isolated from the allantoic fluid using QIAamp Viral RNA mini kit (Qiagen, # 52904) according to the instructions of the manufacturer. Reverse transcription was carried out at 42°C for 1h in a 25 μl reaction mixture containing 8 μl RNA, 1 μl uni12 primer with a concentration of 50 ng/μl (13.5 pmol/μl), 10μl water, 1 μl 10 mM dNTP, 5 μl 5x buffer and 100 units of MMLV (Alpha-Ferment Ltd., Moscow). The resulting cDNA was used in PCR with specific terminal primers to synthesize full-length genome segments [43].
amplified fragments were separated by electrophoresis in 1-1.3% agarose gel in the presence of ethidium bromide and were eluted from the gel with a Diatom DNA Elution kit (Isogen Laboratory Ltd., Russia, # D1031). Sequencing reactions were performed with terminal or internal primers [44]. with the BrightDye™ Terminator Cycle Sequencing Kit v3.1 (Nimagen, the Netherlands), followed by analysis on an ABI PRISM 3100-Avant Genetic analyzer (Applied Biosystems). For assembly and analysis of nucleotide sequences, the Lasergene software package (DNASTAR, Inc.) was used.

2.3. Downloading of sequences and evolutionary tree construction

The complete nucleotide sequences of AIV internal genes (PA, PB1, PB2, NP, MP, and NS) and external genes (H1, H3, H4, H5, N1, H6, H11, N1, N2, N3, N6 and N8) were downloaded from the Influenza Research Database (https://www.fludb.org). The selected sequences were aligned by the MUSCLE method using the software package of MEGA X (https://www.megasoftware.net/) [45]. Time-scaled trees were generated for each internal segment with known isolation dates using BEAST under GTR model with 1000 bootstrap replicates [46]. A strict clock model of molecular clock was chosen for all segments.

2.4. Classification of gene variants and detection of gene reassortants

All lineages and subordinate lineages were classified according to the topology of the phylogenetic trees using the approach described in [47], but with more detail. The major clades on each gene tree were defined by strong bootstrap support (>95%) and numbered. The minor clades within the major clade were designated by corresponding number and a letter.

3. Results

During the autumn periods of 2006–2019, feces of wild waterfowl were collected on the bank of a pond in Moscow city, and IAVs were isolated. Over 14 years, about 4,000 samples were collected, and 38 strains of influenza A viruses of subtypes H1N1, H1N2, H3N1, H3N2, H3N6, H3N8, H4N6, H5N3, H6N2, H11N6 and H11N9 were isolated and completely sequenced (Table 1 and Table S1 of supplemental material). All isolates replicated in chick embryos to high titers, were infectious and immunogenic in mice, although these animals were generally not killed. Five viruses tested were non-pathogenic in chickens. [48, 49].

The subtypes of isolates differed in different years. Until 2013, IAVs with hemagglutinins H3 and H4 dominated. These subtypes were no longer isolated after 2014 and were substituted by IAVs of the subtypes H1N2, H1N1 and H11N6. This change in the virus isolation pattern correlated with data from databases on the isolation of IAVs from ducks in Europe.

3.1 Evolutionary relationships of gene segments.

We built phylogenetic trees for internal gene segments of 38 IAVs isolated in Moscow and subclades on each tree were identified and numbered (Figures S1- S6 of supplemental material).
The A/gull/Moscow/3100/2006 (H6N2) was not fundamentally separated from a number of duck viruses; for all gene segments, closely related variants can be found among the duck viruses (Table 1 and Table S2 of supplemental material).

A majority of IAVs isolated from ducks in Moscow had different gene constellations. The gene segments of the same clade/subclade were perpetuated in duck IAVs over several years. For example, PB2 of clade 11 was detected in 2009, 2010, 2013, 2014, 2018 and 2019 (Table 1).

Importantly, although segments of the same clade/subclade were found in duck IAVs isolated in different years, these viruses never preserved their full genome constellations and always contained a unique mixture of segments that belonged to different clades/subclades. Although four pairs of IAVs with identical constellations were detected in our study (Table 1), the viruses of each pair were isolated almost simultaneously and in the same place. These results support previous conclusion of Dugan and colleagues [34] that IAVs circulate in wild birds as a pool of gene segments, which reassort extensively and appear in a new combination each year.

3.2 Diversity of gene segment constellations of IAVs in ducks and gulls

To compare the rate of natural reassortment of influenza viruses in ducks and gulls we compared a set of IAVs isolated from ducks in the Moscow and the Netherlands and viruses isolated from gulls in the Netherlands in the same period of time (2006 - 2019) [39]. The set of gull viruses included 252 viruses of the H13 subtype, 94 viruses of the H16 subtype, and 11 mixed isolates.

The evolutionary trees for the gene segments were generated and the clades and subclades were numbered (Figures S7- S12 of supplemental material.). The data on genome constellations of 228 duck IAVs viruses and 357 gull IAVs are presented in Tables S2-S5 of the supplemental material; selected data are shown in Fig.1 and Tab. 3.

Among 228 duck IAVs, we identified 187 distinctive genotypes. Pairs of isolates that matched in all gene segments, as a rule, were isolated on the same day. In only three cases, pairs of IAVs listed below with matching genomes were isolated in different years. They were: d/N/20/2012|H1N1 and d/N/5/2013|H1N1, d/N/48/2011|H7N1 and d/N/31/2012|H7N1 and d/N/71/2008|H10N7 and d/N/1/2009|H10N7.

| Isolation date | Strain name a | Subtype | Clade/subclade b |
|----------------|---------------|---------|-----------------|
| 10/15/2019     | M/5743/2019   | H1N1    | 6 5 2a 10b 4E B11 |
| 10/15/2019     | M/5744/2019   | H1N1    | 6 5 2a 10b 4E B11 |
| 10/01/2013     | M/4970/2013   | H1N1    | 6 5 8a 10c 4D B9d |
| 10/17/2018     | M/5586/2018   | H1N2    | 11 12A 5a 10b 4E 6a |
| 11/01/2018     | M/5662/2018   | H1N2    | 11 12A 5a 10b 4E 6a |
| 10/13/2008     | M/4203/2010   | H3N8    | 4 4C 3a 9B 10 2b |
| 11/26/2010     | M/4238/2010   | H3N6    | 4 4C 3a 9B 10 2b |
| Date         | Isolate Code         | HA Subtype | NA Subtype | MP Subtype | NS1 Subtype | NS2 Subtype |
|--------------|----------------------|------------|------------|------------|-------------|-------------|
| 10/13/2010   | M/4298/2010          | H3N8       | 4          | 9C         | 1           | 10e         | 5e          | 4d          |
| 09/27/2011   | M/4494/2011          | H3N8       | 4          | 4A         | 5b          | 10d         | 7           | 4c          |
| 10/10/2011   | M/4681/2011          | H3N8       | 4          | 4A         | 5b          | 10d         | 7           | 4c          |
| 01/10/2011   | M/4524/2011mix       | H3N2       | 4          | 4C         | 5b          | 10d         | 7           | 4c          |
| 01/10/2011   | M/4524/2011mix       | H3N8       | 4          | 4C         | 5b          | 10d         | 7           | 4c          |
| 01/10/2012   | M/4780/2012          | H3N8       | 4          | 9C         | 5a          | 4           | 3           | B12         |
| 04/07/2012   | M/4788/2012          | H3N8       | 4          | 4C         | 3a          | 10d         | 10          | 4c          |
| 09/04/2009   | M/3806/2009          | H3N8       | 11         | 4C         | 6b          | 10d         | 3           | B11         |
| 01/10/2011   | M/4661/2010          | H3N8       | 12         | 9C         | 6b          | 10d         | 4E          | 3e          |
| 09/19/2009   | M/5370/2009          | H4N6       | 2          | 9C         | 1           | 10e         | 5e          | B11         |
| 09/04/2009   | M/3799/2009          | H4N6       | 2          | 9C         | 1           | 10e         | 1           | B11         |
| 09/19/2009   | M/3735/2009          | H4N6       | 3          | 9C         | 1           | 10d         | 1           | B11         |
| 10/31/2012   | M/4781/2012          | H4N6       | 4          | 4C         | 3a          | 10d         | 7           | 3e          |
| 10/17/2012   | M/4843/2012          | H4N6       | 4          | 4C         | 3a          | 3           | 7           | 4c          |
| 10/10/2012   | M/4771/2012          | H4N6       | 6          | 4A         | 3a          | 3           | 7           | 4c          |
| 10/11/2011   | M/4643/2011          | H4N6       | 10         | 10         | 8b          | 9B          | 4E          | 2c          |
| 10/21/2008   | M/3661/2008          | H4N6       | 12         | 13         | 7           | 12          | 6           | 3d          |
| 10/04/2011   | M/4518/2011          | H4N6       | 12         | 9B         | 5a          | 9B          | 5e          | 2c          |
| 10/04/2011   | M/4528/2011          | H4N6       | 12         | 4C         | 5a          | 10d         | 5e          | 4c          |
| 10/19/2011   | M/4641/2011          | H4N6       | 12         | 4C         | 5a          | 10d         | 5e          | 4c          |
| 11/16/2010   | M/4182/2010          | H5N3       | 11         | 4A         | 2a          | 10d         | 3           | 3d          |
| 11/19/2013   | M/4971/2013          | H5N3       | 4          | 4C         | 6b          | 10d         | 10          | 2b          |
| 11/26/2013   | M/4952/2013          | H5N3       | 11         | 4A         | 5b          | 10b         | 6           | 2b          |
| 01/09/2009   | M/3720/2009          | H6N2       | 6          | 4A         | 2b          | 3           | 12          | B11         |
| 01/09/2010   | M/4031/2010          | H6N2       | 12         | 4A         | 2a          | 13          | 13          | 3c          |
| 10/11/2006   | gull/M/3100/2006     | H6N2       | 1c         | 10         | 7           | 1           | 6           | 6b          |
| 30/06/2008   | M/3641/2008          | H11N9      | 12         | 9A         | 3b          | 5           | 11          | 3d          |
| 10/21/2019   | M/5712/2019          | H11N6      | 11         | 12A        | 5a          | 10b         | 4E          | 3e          |

* With the exception of one gull isolate (gull/M/3100/2006), all IAVs were isolated from ducks. The duck viruses are designated by isolation place (M), strain number and isolation year, for example, M/5743/2019 stands for A/duck/Moscow/5743/2019. The viruses are ordered according to the HA subtype.

* Clade numbers and subclade letters of the indicated segments. IAV isolates containing all gene segments of the same subclade are highlighted in yellow, with the strain names highlighted in orange.
Figure 1. Genome constellations of IAVs isolated from ducks and gulls in the Netherlands in 2007-2009*.

*Colors in each column depict clades/subclades of the indicated gene segment on the phylogenetic tree (see Tables S2-S4 of supplemental material). The names of the strains are not shown in the figure for clarity, they are shown in the tables S2-S5 of supplemental material.
Table 2 shows the diversity of IAVs of different subtypes, calculated as the ratio of the number of mismatched variants of genomes to the number of IAV strains in the group. The diversity is very high among the H1, H2, H3, H4, H5, H6 and H11 subtype IAVs, with 83 to 100% of the virus genomes representing unique genome constellations (Table S2 supplemental material). In contrast, gull IAVs were represented by 84 genome constellations per 252 H13 isolates (33% of genomes unique) and 38 constellations per 94 H16N3 isolates (40% of genome constellations unique) (Table S3, S4 supplemental material).

|            | Duck viruses isolated in Moscow and the Netherlands | Gull viruses |
|------------|---------------------------------------------------|--------------|
|            | H1  | H2  | H3  | H4  | H5  | H6  | H11 | H13 | H16N3 |
| NI         | 24  | 13  | 37  | 31  | 19  | 34  | 9   | 252 | 94    |
| NG         | 20  | 11  | 34  | 29  | 18  | 29  | 9   | 84  | 38    |
| %          | 83  | 85  | 92  | 94  | 95  | 85  | 100 | 33  | 40    |

*NS, number of isolates; NG, number of genotypes; %, percentage of NG relative to NI.

Evolutionary histories of the internal gene segments of H13 and H16 IAVs are shown in Fig. 2 with H16 IAVs colored in red. Non-persistent reassortment events are clearly visible on these trees as single strains of one color surrounded by strains with different color. Transition of gene segment from one subtype to another is also clearly visible as the alternation of groups of different colors on the same evolutionary branch. However, it is difficult to study the constellation of all gene segments on the trees, as this requires simultaneous comparison of patterns on different trees. To facilitate the analysis, we compiled tables in which we designated the clade numbers of the gene segments of all viruses, in the same way as it was done for duck IAV.

Gull virus isolates naturally break down into large groups of closely related viruses, matching by subclades of all genes. Apparently these groups represent variants of the same virus that caused the epidemic, e.g. in the nesting colonies. The genomic constellations of such variants are not very stable, they can vary from group to group and can change from year to year. Nevertheless, in some cases, clusters of the same variants of gene segment persist in the subsequent seasons. For example: HA, NA, PB2, PA and NP of the H13N2/2016 isolates are direct descendants of the H13N2/2014 isolates; PB1, MP and NS of both virus groups are descendants of the common close ancestors, HA, PB2, PB1, PA and NP. H13N2/2009 are descendants of H13N8/2008 (Fig S7-S12 of supplemental material).

The internal gene segments of H16N3 viruses usually do not match the corresponding segments of H13 viruses; however, this rule is not very strict. Some of the exceptions are listed below.

- H16N3/2008, H13N2/2009 and H13N2/2010 shared PA
- In 2007, H13N6 and H16N3 had matching PA.
- H13N6/2007 and H16N3/2008 shared NP
- In 2011, H13N6 and H16N3 had matching NP (complete data are given in table S3-S4).
**Figure 2.** Evolutionary history of six internal gene segments of H13 and H16 AIVs.

The trees are based on all published full-length sequences of H13 and H16 gull viruses isolated in the Netherlands. The route is placed on A/Black-headed gull/Netherlands/1/2000 (H13N8). Red color depicts sequences of H16 IAVs. Scale bars, 0.02. The trees were made using FigTree v1.4.4 (http://tree.bio.ed.ac.uk/software/figtree/).

**Table 3.** Genome constellations of IAVs isolated in the Netherlands from black-headed gulls in the years 2007-2016.
In some seasons, large groups of gull IAVs had highly homologous gene segments suggesting that all these IAVs represented descendants of the same precursor circulated during this season. The presence of such
groups in our sets of gull IAVs allowed reliable identification of reassortant viruses, which clearly differed from the other isolates in the group by the origin of either one or a few gene segments (see Table S3 of supplemental material; some of data are given in Tab. 3). As example: among group of H13N6/2014 IAVs, one isolate, bhg/16/2014, contained PB2 from H16N3/2014. Bhg/31/2009/H13N2 strain is a reassortant in which all internal genes coincides with the H13N6 viruses of the same year, and only NA from another clade (Table S3).

Of particular interest are two H13N3 IAVs: bhg/17/2009 and bhg/8/2011, which contained heterologous N3 neuraminidase from H16N3. The former virus shared PB2, PA, NP, and NA with H16N3/2009, the remaining 4 segments is shared with other H13N6/2009 IAVs. Seven gene segments of bhg/8/2011 matched corresponding segments of H16N3/2011, only the HA being shared with H13N8/2011 (Tab. 3 and Table S4).

One can conclude that two H13N3 isolates originated from H16N3 IAVs, in which HA and a number of other genes were replaced by gene segments of H13 IAVs via reassortment. Apparently, such reassortants are not viable enough; they were found in only two isolates despite the fact that there were seven mixed isolates in the same set (Table S4).

Among the H16N3 viruses, there are also reassortants with internal genes from the H13 viruses. In total, among 94 H16N3 viruses, we recognized 17 non-persistent reassortment events for one or several genes, among which 13 reassortants acquired gene segment of H13 subtype (Table S4 supplemental material).

Among 252 H13 viruses, we found 14 non-persistent reassortment events for one or several genes, among which 9 reassortants acquired genes of H16 subtype.

Thus, the percentage of reassortants within one season was close to 10%, which is comparable to about 3% of mixed strains.

Thus, although partial reassortment of gene segments can be observed over years among H13 and H16 IAVs, constellations of several segments remained stable for prolonged periods of time. The observed pattern is very different from what we observed in duck viruses of H1-H6 and H11 subtypes. No stable gene constellations could be detected in the sets of duck IAVs studied.

4. Discussion

Comparison of duck and gull viruses showed a large difference in the detection rate of reassortment in the main duck subtypes (H1, H2, H3, H4, H5, H6, and H11) and the gull subtypes H13 and H16. In gull viruses, reassortment was a fairly frequent, well-recorded phenomenon, while in duck viruses, gene mixing was so intense that it was almost impossible to find viruses with the same genome constellation. There may be several explanations for this difference.

A key difference between gulls and ducks is that gulls breed in dense colonies with much mixing of chicks, whereas ducks are dispersed during breeding. As virus spreads among the gull chicks in nesting colonies, a single newly introduced variant can infect many birds. Large numbers of nearly identical strains can be isolated at this time. In the separated colonies, an outbreak of another variant of the virus may occur. Thus, three clusters of closely related isolates from 2008 (one H16N3 cluster and two H13N8 clusters) represent independent epidemics in three different colonies [5]. Sometimes the virus is carried from one colony to another, leading to emergence of mix-isolates and reassortants. However, these events are rather exceptions than the rule.

Because ducks do not breed in colonies, transmission among young ducks is limited during breeding. On the other hand, during the moult and the fall flight ducks from multiple breeding areas may mix, creating
ideal conditions for mixed infections with distinctive influenza viruses. On the pond, where the Moscow duck viruses were isolated, hundreds of mallards accumulate in the fall, which crowd along the edge of the pond, where children throw pieces of bread. Ducks arrive from the north of Europe [50, 51]. Mallards spend about 2 months on this pond. The first birds arrive in mid-September, followed by constantly increasing numbers of birds. As shown by Wille and colleagues, ducks can be infected sequentially by several variants of IAVs, and excrete virus intensively in their feces. Therefore, each introduced virus multiplies and infects other ducks thus promoting multiple infections and reassortments [52].

The second factor affecting the rate of reassortment is the pressure of natural selection, which can sweep unsuccessful combinations. In the secondary hosts, such as chickens and humans, this factor is probably the main reason for the persistence of certain optimal gene constellations over the years. For example, during the epidemics in humans, when two subtypes co-circulate, co-infection and even reassortment is quite possible [13-16], but reassortants are usually less viable than the parental variants and do not become fixed in the population [18,19].

Probably, the same reason explains the stability of the genomes of highly pathogenic influenza viruses. After moving from wild birds to poultry, viruses adopt to a new host and a new route of virus transmission. The adaptation of IAVs to chickens is associated with an increase of the virus pathogenicity [53]. This effect can be explained, at least in part, by the IAV evolution towards efficient transmission in infected poultry owing to cannibalism (that is, pecking and eating of sick individuals). A characteristic feature of chicken influenza is the selection of IAVs with polybasic cleavage site in HA, which enables the virus to infect endothelial cells and to cause generalized infection [54, 55]. Acquisition of new properties requires the coordinated evolution of all genes. In each of the evolutionary branches of IAVs that adapt to chickens, unique concerted changes in the genes could take place, so that exchange of some segments by reassortment can destroy an interplay between the genes and/or their products thus making the virus less viable. Naturally, such reassortants will be swept by natural selection, leading to the persistence of specific constellations of the gene segments.

In duck viruses, stable genome constellations are largely absent [34]. Gull viruses, by contrast, have semi-stable genome constellations, among them, stable combinations of HA and NA. The H16 HA is tightly associated with the N3 NA. The NAs of IAVs with H13 HA (N2, N6 and N8 NAs) separated in the course of evolution from the ancestral viruses of ducks and adapted to the viruses of gulls. The variants with H13 and N3 did not form stable evolutionary lineages, probably, they are not viable enough. The functional balance of HA and NA is an essential element of the viability of IAVs [56]. The receptor specificity of gull IAVs differs from that of duck IAVs. In contrast to duck IAVs, gull IAVs efficiently bind to fucosylated receptors. Unlike all other IAVs of aquatic birds, H16 IAVs recognize both 2-3 and 2-6 sialyl-galactose receptors, being in this respect more similar to swine viruses than to duck viruses [57]. Likely, the neuraminidase of H16N3 viruses is specifically adapted to H16 HA.

The internal gene segment of the H13 and H16N3 viruses, as well as the HA and NA segments, represent separate evolutionary branches adapted to the gull host. The internal genes of the H13 and H16 viruses are still interchangeable, but tend to form relatively stable constellations in each subtype. Authors should discuss the results and how they can be interpreted from the perspective of previous studies and of the working hypotheses. The findings and their implications should be discussed in the broadest context possible. Future research directions may also be highlighted.

5. Conclusions
Duck AIVs represent a unique variant of the symbiosis of the virus with the host, where the virus does not persist as a specific genome, but as a pool of genes, from which new genomic combinations are constantly formed [34].

However, in non-duck species, including gulls, a different evolution scenario is common, when the virus evolves in the host in the form of a whole genome.

**Supplementary Materials:** The following are available online at http://www.mdpi.com/1999-4915/10/4/164/s1, Figure S1: Evolutionary tree of the PB2 gene of duck IAV isolated in Moscow and Netherlands in 2006-2019. Branches are numbered as described in the Materials and Methods section., Figure S2: Evolutionary tree of the PB1 gene of duck IAV isolated in Moscow and Netherlands in 2006-2019. Branches are numbered as described in the Materials and Methods section., Figure S3: Evolutionary tree of the PA gene of duck IAV isolated in Moscow and Netherlands in 2006-2019. Branches are numbered as described in the Materials and Methods section. Figure S4: Evolutionary tree of the NP gene of duck IAV isolated in Moscow and Netherlands in 2006-2019. Branches are numbered as described in the Materials and Methods section., Figure S5: Evolutionary tree of the MP gene of duck IAV isolated in Moscow and Netherlands in 2006-2019. Branches are numbered as described in the Materials and Methods section., Figure S6: Evolutionary tree of the NS gene of duck IAV isolated in Moscow and Netherlands in 2006-2019. Branches are numbered as described in the Materials and Methods section., Figure S7: Evolutionary tree of the PB2 gene of gull IAV isolated in Netherlands in 2006-2019. Branches are numbered as described in the Materials and Methods section., Figure S8: Evolutionary tree of the PB1 gene of gull IAV isolated in Netherlands in 2006-2019. Branches are numbered as described in the Materials and Methods section., Figure S9: Evolutionary tree of the PA gene of gull IAV isolated in Netherlands in 2006-2019. Branches are numbered as described in the Materials and Methods section., Figure S10: Evolutionary tree of the NP gene of gull IAV isolated in Netherlands in 2006-2019. Branches are numbered as described in the Materials and Methods section., Figure S11: Evolutionary tree of the MP gene of gull IAV isolated in Netherlands in 2006-2019. Branches are numbered as described in the Materials and Methods section. Figure S12: Evolutionary tree of the NS gene of gull IAV isolated in Netherlands in 2006-2019. Branches are numbered as described in the Materials and Methods section., Table S1: Full names, designations and accession number of genes of IAV isolated in Moscow in 2006-2019., Table S2: Duck IAV isolated in Moscow and Netherlands in 2006-2019. Viruses are grouped by hemagglutinin subtype. The cells contain clade and subclade numbers for the corresponding genes. Colored cells depict strains, all genes of which match by subclade. The names of the strains are marked by orange color, corresponding segments are colored by yellow. If the date of collection coincides, the names of strains is written in black. If the date of collection differed within the same year, the names of strains are written in red. If the strains were collected on different years, the name is written in green. Table S3: Gull IAV of H13 subtype isolated in Netherlands in 2007-2016. The subheadings contain the group names of the viruses. The cells contain clade and subclade numbers for the corresponding genes. The cells with the numbers of the isolates, colored orange, correspond to the strains, all genes of which match by the subclade. The cells colored yellow correspond to the gene variants presented in both H13 and H16 viruses. Light green cells indicate non-persistent reassortment events., Table S4: Gull IV of H16 subtype isolated in Netherlands in 2007-2017. The subheadings contain the group names of the viruses. The cells contain clade and subclade numbers for the corresponding genes. The cells with the numbers of the isolates, colored ocher, correspond to the strains, all genes of which matching by number of subclade. The cells colored yellow correspond to the gene variants presented in both H13 and H16 viruses.
viruses. Light green cells indicate a nonpersistent reassortment events. Table S5: Genome constellations of IAVs isolated from ducks and gulls in the Netherlands in 2007-2017. Colors in each column depict clades of the indicated gene segment on the phylogenetic tree.

**Author Contributions:** Yulia Postnikova, Anastasia Treshchalina, Alexandra Gambaryan, conceived and designed the experiments; Yulia Postnikova, Anastasia Treshchalina, Elizaveta Boravleva, Galina Sadykova, Alexei Prilipov, Alexandra Gambaryan, and Natalia Lomakina performed the experiments; Alexandre Gambaryan, Anastasia Treshchalina, Mikhail Matrosovich and Yulia Postnikova analyzed the data; Yulia Postnikova, Alexandra Gambaryan, Mikhail Matrosovich and Ron A.M. Fouchier wrote the paper.

**Funding:** This research was supported by the research grant 17-04-00148 from the Russian Foundation for Basic Research and by by the Deutsche Forschungsgemeinschaft (DFG, German Research Foundation) - Project number 197785619 – SFB 1021. Publication was supported by Tscha- makov Federal scientific center for the research and development of immune-and-biological products.

**Acknowledgments:** We thank Oanh Vuong, Rachel Scheuer, Josanne Verhagen and Marjolein Poen of Erasmus MC, who generated the Dutch sequence data.

**Conflicts of Interest:** The authors declare no conflict of interest.

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