Detection of Evolutionarily Distinct Avian Influenza A Viruses in Antarctica

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ABSTRACT Distinct lineages of avian influenza viruses (AIVs) are harbored by spatially segregated birds, yet significant surveillance gaps exist around the globe. Virtually nothing is known from the Antarctic. Using virus culture, molecular analysis, full genome sequencing, and serology of samples from Adélie penguins in Antarctica, we confirmed infection by H11N2 subtype AIVs. Their genetic segments were distinct from all known contemporary influenza viruses, including South American AIVs, suggesting spatial separation from other lineages. Only in the matrix and polymerase acidic gene phylogenies did the Antarctic viruses form a sister relationship to South American AIVs, whereas distant phylogenetic relationships were evident in all other gene segments. Interestingly, their neuraminidase genes formed a distant relationship to all avian and human influenza lineages, and the polymerase basic 1 and polymerase acidic formed a sister relationship to the equine H3N8 influenza virus lineage that emerged during 1963 and whose avian origins were previously unknown. We also estimated that each gene segment had diverged for 49 to 80 years from its most closely related sequences, highlighting a significant gap in our AIV knowledge in the region. We also show that the receptor binding properties of the H11N2 viruses are predominantly avian and that they were unable to replicate efficiently in experimentally inoculated ferrets, suggesting their continuous evolution in avian hosts. These findings add substantially to our understanding of both the ecology and the intra- and intercontinental movement of Antarctic AIVs and highlight the potential risk of an incursion of highly pathogenic AIVs into this fragile environment.

IMPORTANCE Avian influenza viruses (AIVs) are typically maintained and spread by migratory birds, resulting in the existence of distinctly different viruses around the world. However, AIVs have not previously been detected in Antarctica. In this study, we characterized H11N2 viruses sampled from Adélie penguins from two geographically different sites in Antarctica and show that the segmented AIV genome diverged between 49 and 80 years ago from other AIVs, with several genes showing similarity and shared ancestry with H3N8 equine influenza viruses. This study provides the first insight into the ecology of AIVs in Antarctica and highlights the potential risk of an introduction of highly pathogenic AIVs into the continent.

Wild aquatic birds such as dabbling ducks, gulls, and other shorebirds are considered the natural reservoir for avian influenza viruses (AIVs). The ecology and migratory patterns of these birds therefore have a direct effect on the global distribution and diversity of AIVs (1). The ecology of AIVs in wild birds has been well studied in many regions of the Northern Hemisphere, but considerably less is known in the Southern Hemisphere. Of the publicly available AIV sequences (n = 19,784), only 5.7%, 1%, and 0.1% are from Africa, Oceania, and South America, respectively, and none have previously been described from Antarctica (GISAID, http://www.gisaid.org). Wild migratory birds play a key role in the spread of AIVs on a local, regional, and intercontinental scale via broadly established flyways (1) (see Fig. S1 in the supplemental material). As a result, spatially segregated birds of the American and Eurasian landmasses harbor distinct lineages of AIVs (1). Traditionally, AIVs have been broadly separated into North American or Eurasian lineages, although this appears to be an oversimplification and less applicable to AIVs from the Southern Hemisphere (2). Recent genetic analysis of AIVs in Australia, located at the southern end of the East Asian-Australian flyway, showed some divergence from Eurasian strains, demonstrating that, although new strains have been introduced frequently from Europe and Asia, there is a high degree of endemic evolution that has produced a group of viruses that are genetically distinct from those circulating in Asia (2–4). Similar evolutionary divergence has also been observed in some South American AIVs, where internal gene segments form unique “South American-like” clades, although the hemagglutinin (HA) and/or neuraminidase (NA)
gene segments are often “North American-like” (5–8). It is therefore possible that evolutionary divergence of AIVs may increase with further advancement south and that spatially segregated regions and continents such as Southern Patagonia and Antarctica may act as evolutionary “sinks” harboring highly diverged AIVs. The major migratory flyways are not thought to extend to Antarctica, although each spring, over 100 million birds breed around the rocky Antarctic coastline and offshore islands (9). These include birds such as the Arctic tern (Sterna paradisaea) (10) and south polar skua (Stercorarius maccormicki) (11) that conduct transhemispheric migrations to Antarctica and thereby have the potential to facilitate movement of avian influenza viruses into and out of Antarctica. A small number of earlier AIV studies in Antarctica have detected influenza A antibodies in penguins and other birds (12–16), but none have detected or isolated any influenza viruses. The detection and subsequent analysis of AIVs from Antarctica would provide insight into the frequency and likely routes of AIV movement into and within the continent and help assess the potential risk of an incursion of highly pathogenic AIVs from neighboring regions.

**RESULTS**

Identification of influenza virus in penguins. We took combined cloacal/tracheal swabs from 301 Adélie penguins (Pygoscelis adeliae) (adults and chicks), and blood from 270 of those, from two locations on the Antarctic Peninsula during January and February 2013 (Table 1 and Fig. 1). Eight samples (2.7%) (with cycle threshold [CT] values ranging from 25.4 to 36.9) were found to contain influenza virus RNA following a real-time reverse transcription (RT)-PCR assay for the influenza matrix gene. AIV was confirmed by sequence and BLAST analysis of at least one gene sequence (hemagglutinin [HA], nonstructural [NS], or matrix [M]) (Table 1).

<table>
<thead>
<tr>
<th>Location</th>
<th>Age</th>
<th>No. of penguins sampled</th>
<th>No. (%) of samples with AIV RNA detected</th>
<th>No. of samples isolated in eggs</th>
<th>No. of penguins sampled</th>
<th>No. (%) of samples with IAV antibodies detected</th>
<th>No. (%) of samples with A(H11N2) antibodies detected</th>
</tr>
</thead>
<tbody>
<tr>
<td>Admiralty Bay</td>
<td>Adult</td>
<td>110</td>
<td>0</td>
<td>0</td>
<td>96</td>
<td>16 (16.7)</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>Chick</td>
<td>40</td>
<td>1 (2.5)</td>
<td>0</td>
<td>35</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Rada Covadonga</td>
<td>Adult</td>
<td>111</td>
<td>6 (5.4)</td>
<td>3&lt;sup&gt;e&lt;/sup&gt;</td>
<td>99</td>
<td>27 (27.3)</td>
<td>2 (1.8)</td>
</tr>
<tr>
<td></td>
<td>Chick</td>
<td>40</td>
<td>1 (2.5)</td>
<td>1&lt;sup&gt;f&lt;/sup&gt;</td>
<td>40</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Total</td>
<td></td>
<td>301</td>
<td>8 (2.7)</td>
<td>4</td>
<td>270</td>
<td>43 (15.9)</td>
<td>2 (0.7)</td>
</tr>
</tbody>
</table>

<sup>a</sup> No. of samples where the HA, NS, or M gene sequence was obtained and confirmed to be influenza A virus by BLAST analysis.

<sup>b</sup> Egg isolation was attempted only for those samples with a CT value of <38 (n = 14).

<sup>c</sup> Determined using an anti-NP ELISA assay.

<sup>d</sup> Determined using an HI assay with the H11N2 antigen A/peng/Ant/270/13.

<sup>e</sup> Viruses were designated A/Adélie Penguin/Antarctica/226/2013, A/Adélie Penguin/Antarctica/184/2013, A/Adélie Penguin/Antarctica/178/2013.

<sup>f</sup> Virus was designated A/Adélie Penguin/Antarctica/270/2013.

**FIG 1** Map of sampling sites in Antarctica. Adélie penguins were sampled from two locations in Antarctica: Admiralty Bay, King George Island; and Kopaitik Island, Rada Covadonga, Antarctic Peninsula. Inset: An Adélie penguin. The map is courtesy of Marco Villarroel, reproduced with permission.
Six AIV-positive samples were from adult Adélie penguins and two from Adélie chicks, with all samples except for one collected from Rada Covadonga (Table 1). All eight influenza viruses, including the single virus from Admiralty Bay, had a high degree of homology (>99.7% nucleotide sequence identity) between respective genes. Following inoculation of these viruses into embryonated hen's eggs, four viruses were successfully cultured (all from specimens with a Ct value of <33 based on the matrix gene real-time RT-PCR assay). Full genome analysis of the four isolated viruses showed that they were all H11N2 influenza A viruses (GenBank sequence accession numbers KJ729348 to KJ729379). These four viruses were designated A/Adélie penguin/Antarctica/178/2013, A/Adélie penguin/Antarctica/184/2013, A/Adélie penguin/Antarctica/226/2013, and A/Adélie penguin/Antarctica/270/2013 (abbreviated as A/peng/ANT/178/13, A/peng/ANT/184/13, A/peng/ANT/226/13, and A/peng/ANT/270/13, respectively).

**Evolutionary relationship and divergence time estimates.**

Full genome sequences of the four isolated viruses were compared with all available animal and human influenza virus sequences in public databases to determine the phylogenetic relationships and the evolutionary origin of each gene segment (Fig. 2 and 3). Detailed phylogenetic trees with strain designations are shown in Fig. S2 and S3 in the supplemental material. Unexpectedly, each AIV gene segment from the Adélie penguin viruses was highly different from that of contemporary AIVs circulating either in the Northern or Southern hemispheres (see Fig. S2 and S3 and Table S1 in the supplemental material). Four of the gene segments (HA, PB2, nucleoprotein [NP], and NS) were most closely related to North American avian lineage viruses from the 1960s to 1980s (90 to 96% identity) (Fig. 2 and 3; see also Table S1), while the N2 neuraminidase gene formed a basal position to the North American avian influenza lineage, indicating an early divergence from the known AIV gene pool (Fig. 2; see also Fig. S2). The MP gene of the penguin virus formed a close relationship with North American viruses from 1974 to 2001 but appears to be the closest ancestor to a large number of South American AIVs from Chile, Argentina, and Brazil (Fig. 3; see also Fig. S3E). The PA segments of the penguin AIVs also have a close ancestral relationship with South American viruses, which together appear on their own phylogenetic clade sharing ancestry with equine influenza H3N8 viruses from the 1960s (Fig. 3; see also Fig. S3C). Interestingly, the PB1 segments of the penguin AIVs are also most closely related to equine H3N8 viruses, with significant genetic distance from North American AIVs (Fig. 3; see also Fig. S3B).

Using a molecular clock analysis that incorporates the evolutionary rate of each AIV gene segment, we estimated the time of the most recent common ancestor (TMRCA) of the penguin virus and its most closely related sequence. For the HA, NA, PB2, NP, PA, and NS genes, the TMRCA was between 1957 and 1964, whereas for the MP and PB1 genes, the estimated TMRCA was even earlier at 1933 and 1936, respectively (Table 2). Therefore, there has been an estimated 49 to 80 years of unsampled diversity and evolution leading to these previously unknown influenza viruses (Table 2).

**Molecular markers of host specificity and pathogenicity.**

The HA of the Adélie penguin viruses did not contain the multibasic cleavage site observed in highly pathogenic H5 or H7 AIVs, nor did the genome contain other major mammalian adaptations, such as the PB2 E627K substitution, which is known to alter host range (17) and virulence (18). Infection of embryonated eggs was not pathogenic to the embryo. The receptor binding pocket residues have only one human-like amino acid at position A137 (H3 numbering), with the rest being avian-like with E190, G225, Q226, and G228, suggesting an α2,3-linked sialic acid receptor binding preference (see Fig. S4 in the supplemental material).

Compared with the distantly related H11N9 virus isolated from a wild bird in Australia in 2004, the A/peng/ANT/178/13 HA sequence contained amino acid substitutions at I155T (at the base of the receptor binding site and within contact range to the sialic acid) and A186S and T187S (at the entry of the host receptor pocket), which may alter preferences for different variants of 2,3-linked glycans (Fig. 4). Furthermore, A/peng/ANT/178/13 had a deletion of the A134 residue compared to other wild bird H11 viruses and an insertion of G133 compared to H13 and H16 viruses with known crystal structures (see Fig. S4), which could alter receptor binding slightly due to relative shifts of the adjacent 160 loop. Even though their PB1 and PA genes were phylogenetically related to those of the equine viruses, the Adélie penguin viruses contained typical avian-like amino acid signatures (N375 [PB1], D55 and V100 [PA]), which differ from the equine-like signatures that presumably arose following adaptation to viral replication in horses (19). The viruses contained none of the mutations known to confer resistance to either the adamantane or neuraminidase inhibitor classes of influenza antiviral drugs.

**Identification of influenza antibodies in penguins.**

Sera from 270 of the 301 penguins, including the eight penguins that were AIV positive, were tested by enzyme-linked immunosorbent assay (ELISA) for cross-reactive NP antibodies. As the NP is highly conserved across all influenza A viruses, the assay is designed to detect any previous influenza A infection, regardless of subtype. In comparison, a hemagglutination inhibition (HI) assay, which detects antibodies to the highly variable HA protein, was used to detect

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**Note:** The image contains a phylogenetic tree of H11 (A) and N2 (B) sequences. Sequences from the viruses isolated from Adélie penguins in Antarctica are highly diverged from other circulating AIVs. The HA phylogenetic tree was constructed using maximum likelihood methodology and shows ancestry with Eurasian avian H11 viruses, while the NA sequence is highly diverged from all other clades but with shared ancestry with North American N2 viruses. Detailed phylogenetic trees with virus designations are shown in Fig. S2 in the supplemental material.
FIG 3 Phylogenetic trees of the internal gene sequences of avian, equine, and mammalian influenza A viruses. Major avian influenza clades are broadly classified into Western, Eastern, or global based on the geographic location of sample collection. Phylogenetic trees were constructed using maximum likelihood methodology. AIV sequences from the penguins are marked as “Antarctica,” with branches colored red. Blue branches indicate South American avian influenza sequences, green branches indicate avian influenza viruses from other regions, and orange branches indicate equine influenza viruses. Detailed phylogenetic trees with virus designations can be found in Fig. S3 in the supplemental material.
TABLE 2 Summary of evolutionary relationships of each gene segment of influenza viruses isolated from Adélie penguins in Antarctica during 2013

<table>
<thead>
<tr>
<th>Gene</th>
<th>% identity&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Lineage&lt;sup&gt;b&lt;/sup&gt;</th>
<th>Mean time of unsampled diversity in yrs (BCI)&lt;sup&gt;c&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>HA gene</td>
<td>90</td>
<td>North American avian</td>
<td>50 (43-58)</td>
</tr>
<tr>
<td>NA gene</td>
<td>90</td>
<td>North American avian/ new lineage</td>
<td>54 (50-60)</td>
</tr>
<tr>
<td>PB2 gene</td>
<td>96</td>
<td>North American avian</td>
<td>50 (41-58)</td>
</tr>
<tr>
<td>PB1 gene</td>
<td>89</td>
<td>Equine H3N8</td>
<td>77 (61-100)</td>
</tr>
<tr>
<td>PA gene</td>
<td>91</td>
<td>Equine H3N8, South American avian</td>
<td>56 (50-69)</td>
</tr>
<tr>
<td>NP gene</td>
<td>96</td>
<td>North American avian</td>
<td>49 (41-58)</td>
</tr>
<tr>
<td>MP gene</td>
<td>97</td>
<td>North American avian, South American avian</td>
<td>80 (56-112)</td>
</tr>
<tr>
<td>NS gene</td>
<td>95</td>
<td>North American avian</td>
<td>56 (45-69)</td>
</tr>
</tbody>
</table>

<sup>a</sup> Percent identity was estimated using BLAST in NCBI influenza virus resources.

<sup>b</sup> Virus lineages were determined using a maximum likelihood phylogeny generated for each gene segment of all influenza virus sequences that were >96% similar. Phylogenies with virus names and accession numbers are presented in Fig. S1A to H in the supplemental material.

<sup>c</sup> Time to the most recent common ancestor (in years) of influenza viruses isolated from Adélie penguins in Antarctica during 2013 and the most closely related virus sequence; represents the mean time of unsampled diversity of this virus (orange bars in Fig. S2 and S3 in the supplemental material). Bayesian confidence intervals (BCI) for the time of unsampled diversity are shown in parentheses.

Previous infection with either the penguin H11N2 viruses or a distantly related H11N9 virus isolated from a wild bird in Australia in 2004. A total of 15.9% of penguins (43/270) were shown to have influenza A NP antibodies by ELISA. All NP antibody-positive penguins were adults, with a greater proportion from Rada Covadonga than from Admiralty Bay (27.3% compared to 16.7%) (Table 1). Only 2/270 (0.7%) penguins had positive HI titers against the penguin H11N2 virus (Table 1). One of the H11N2 antibody-positive serum samples (with a titer of 40) was from an H11N2 PCR- and isolation-positive penguin that yielded A/peng/Ant/226/13, while the other positive serum sample (with a titer of 80 to 160) was from a penguin that was RT-PCR negative for influenza virus. Neither these sera nor the remaining 268 sera inhibited the unrelated H11N9 wild bird virus from Australia. Overall, 5 of the 8 influenza virus-positive penguins had high NP antibody titers, but only one had specific H11 antibodies. This suggests that the detected NP antibodies may have been induced by prior infection(s), as the remaining influenza virus-positive penguins (which included samples with the highest RNA copy number and two that were culture positive) were NP antibody negative.

**Infectivity in ferrets and receptor binding.** To assess the potential risk of A/peng/Ant/270/13 infection in mammals, we administered the virus to ferrets via intranasal (i.n.) or intratracheal (i.t.) routes and compared infectivity with an Australian wild bird H11N9 virus and a contemporary human H3N2 influenza virus. Intranasal administration of A/peng/Ant/270/13 resulted in little or no viral replication, with detection of virus only on day 2 postinfection (at a low titer of 1 × 10^4 EID<sub>50</sub>/ml) (see Fig. S5 in the supplemental material). The ferrets showed no clinical symptoms or significant weight loss and had no detectable H11 serum antibodies by HI assay on day 14 or at 21 days following an intramuscular boost on day 14. In comparison, i.n. delivery of an Australian wild bird H11 virus and the human H3N2 virus yielded productive infections (shedding virus from days 2 to 5, with peak viral titers of 1 × 10^4.2 and 1 × 10^5.5 EID<sub>50</sub>/ml, respectively) (see Fig. S5) and homologous HI antibody titers of 40 and 2,560, respectively, on day 14 postinfection. Following i.t. delivery of A/peng/Ant/270/13, a low virus titer (1 × 10^1 EID<sub>50</sub>/ml) was detected in one lung lobe but not in the other lobe at day 4, but again no H11 antibodies were detected on day 14. No viral titer was detected in either lung lobe following i.t. administration of ferrets with either the Australian wild bird H11 virus or a human H3N2 influenza virus (see Fig. S5). Using turkey erythrocytes enzymatically modified to contain either terminal SAα2,3Gal (avian-like receptors) or SAα2,6Gal (human-like receptors) linkages on cell surface oligosaccharides, we further dem-
onstrated that the A/peng/Ant/270/13 primarily bound avian-like receptors, with only minor binding to α2,6-linked SA receptors (see Table S2). We also showed that, compared to the human H3N2 virus, the viral infectivity of A/peng/Ant/270/13 in vitro was considerably reduced at 33°C compared to that at 37°C, indicating a lack of adaptation to replication in the cooler mammalian upper airways (see Fig. S6).

**DISCUSSION**

Here, we describe the detection of AIVs in Antarctica, providing the first insights into their ecology in this isolated continent. Previous studies of birds and mammals in Antarctica have failed to detect either live AIV or influenza virus RNA. Although AIV antibodies have been detected in penguins at frequencies ranging from 11.8 to 12.5% in Gentoo penguins (Pygoscelis papua) (12, 15), from 3.7 to 58.4% in Adélie penguins (12, 14, 16), and 10.8% in chinstrap penguins (Pygoscelis antarcticus) (12), several other studies have failed to detect any influenza antibodies in these or other penguin species in Antarctica (14, 16, 20, 21), suggesting that influenza infections may be absent from penguin colonies in Antarctica for long periods of time. The detection of H1N2 virus in 2.7% of Adélie penguins sampled in this study, along with the substantially lower prevalence of penguins with detectable H11 antibody titers (0.7%), suggests either that we sampled during the acute phase of infection before antibodies to H11 had developed or that the H11N2 infections caused only a weak antibody response. The intensity and longevity of antibody responses in penguins infected with AIVs are currently unknown, highlighting the need for longitudinal repeat sampling studies of the same penguin colonies with tagged individuals to enable interpretation of serology data in these species. Although H11 AIVs have never been reported in wild birds from South America, they have been detected in migratory shorebirds in North America (where they were the second most frequently detected HA subtype) (22), as well as in Australia (23) and New Zealand (24). Additionally, H11 AIVs have been commonly detected in ducks in many other regions of the world (25, 26).

It is important to understand the frequency and magnitude of inter- and intraregional movement of Antarctic AIVs. Long-distance migrants such as skuas and giant petrels that traverse large distances both within Antarctica and between continents have been shown to have a high frequency of AIV antibodies (skuas, 1.0 to 11.1% [12–14]; giant petrels, 51.5 to 58.8% [12]). These birds have very close interactions with penguins, often nesting alongside their colonies and attacking penguin chicks, providing a likely route for AIV transmission. The yellow-billed pintail (Anas georgica) is a species of duck that, while most commonly observed in South America, has occasionally been recorded on the Antarctic peninsula, making it the most southerly recorded waterfowl species (27). Given the high rates of AIV carriage in waterfowl compared to those in other orders of birds, it is possible that this species plays a role in AIV ecology in Patagonia, the South Shetland Islands, and the Antarctic Peninsula. The return of migratory birds to Antarctica each year between October and December (start of the austral summer period) coincides with penguins returning to colonies to nest and rear chicks. Because of the high population density of birds, summer is the most likely period for AIV transmission and infection in Antarctica. The large amount of penguin feces in colonies during summer, which in some cases is so significant it can be observed on satellite images (28), presumably facilitates AIV transmission by the fecal-oral route (29). Although the relative susceptibility of different penguin species to AIVs is unknown, we did take swabs from a small number of gentoo (n = 74) and chinstrap (n = 18) penguins in Rada Coudron, but none were AIV positive. The different species typically form exclusive separate colonies, which may explain the lack of cross-infection between the three penguin species at the time of sampling in Rada Coudron.

The detection of this highly divergent H11N2 AIV suggests that its existence in Antarctica is not transient. Although two gene segments share common ancestry with South American AIVs, the other remaining six genes shared ancestry with either North American avian or equine viruses. Antarctica therefore may be the most extreme example of an AIV “evolutionary sink,” where viruses are seeded into the continent on an infrequent basis and then become established, persisting as a local reservoir and evolving independently from AIVs in other areas of the globe due to a lack of interconnectedness. Future studies to understand which species or groups of animals are responsible for maintaining AIVs in Antarctica are needed. Based on all previous studies in Antarctica, the overall AIV antibody prevalence in penguins (3.3%; 82/2,500) (12, 14–16, 20, 21, 30, 31) is considerably lower than it is in skuas and giant petrels (13.3%; 47/353) (12–14), suggesting that the latter species may be more likely to maintain and transmit AIVs in Antarctica. An alternative hypothesis is that AIVs circulate between birds only in summer and then become cryopreserved in ice over winter (32), after which birds are again infected once the ice melts in the following summer. In addition, a number of Antarctic marine mammals, such as seals and whales, may also have the potential to become infected and spread influenza viruses given the previous reports of their susceptibility to influenza infection (33, 34). Although three reported studies of seals (n = 306) in Antarctica did not detect influenza viruses or antibodies (14, 35, 36), these animals appear to be susceptible to both avian and human influenza viruses, as their respiratory tracts contain both α2,3- and α2,6-linked sialic acid receptors (33). Seals have been infected with both AIV (33) and human influenza viruses (37), as well as infecting humans with an H7N7 virus following close experimental contact (38, 39). Whether these animals could act as a “mixing vessel” for avian or human influenza viruses in a natural environment in the same manner as pigs is unknown, although the bidirectional transmission of influenza between seals and humans may be a potential risk for both local wildlife and public health with increasing human activity in Antarctica. The H1N2 Antarctic AIV A/peng/Ant/270/13 appeared to be less infectious in ferrets than has been observed for other AIVs from wild birds. AIVs such as H6N1, H1N9, H9N2, H6N5, and H11N9 (used in this study) have readily infected ferrets and in many cases been able to successfully transmit between cohoused naïve ferrets (40–42). Further work is needed to understand the key residues that may be responsible for these differences in replication.

While it has previously been postulated that AIV infection caused mortality of Adélie penguin chicks in Antarctica (16), we did not notice any clinical symptoms or below-average weight in the H11N2-infected penguins in this study. Given that highly pathogenic AIVs have been detected in both South America (43) and Australia (44), it is not inconceivable that such viruses could be transferred to the Antarctic continent by migratory birds, potentially resulting in catastrophic mass mortality, such as that seen with other pathogens such as avian cholera (Pasteurella multocida).
Allantoic fluid was harvested after 3 days, and influenza virus was detected in swab specimens by inoculating the swab specimen (diluted 1:1 with phosphate-buffered saline) with universal primers targeting each of the eight influenza segments (53). Influenza viruses were analyzed further by conventional RT-PCR using an Applied Biosystems 7500 Fast real-time PCR system (Life Technologies). Samples with a cycle threshold (CT) value of 42 cycles or cultured in 96-well plates (5% CO2, 37°C). Log10 virus titers were determined using the Reed and Muench method (54).

RT-PCR and sequence analysis. RNA was isolated from cloacal/tracheal swabs using the QIAGEN QIAextractor system. All samples were tested by a one-step RT-PCR assay targeting the influenza matrix genome segment (52), using a SensiFast Probe Lo-ROX one-step kit (Biolonde) on an Applied Biosystems 7500 Fast real-time PCR system (Life Technologies). Samples with a cycle threshold (CT) value of <42 cycles or cultured influenza viruses were analyzed further by conventional RT-PCR using universal primers targeting each of the eight influenza segments (53). Amplified RT-PCR products were sequenced using standard techniques on an Applied Biosystems 3500XL genetic analyzer.

Virus isolation. Virus isolation was attempted from RT-PCR-positive samples by inoculating the swab specimen (diluted 1:1 with phosphate-buffered saline) with 1% neomycin-polymyxin solution [biocidal] into the allantoic cavity of 11-day-old embryonated hen’s eggs. Allantoic fluid was harvested after 3 days, and influenza virus was detected by hemagglutination with turkey erythrocytes. The EID50/mL of each isolate was determined by performing log10 dilutions in PBS containing 1% neomycin-polymyxin solution and infecting the allantoic cavity of 11-day-old embryonated hen’s eggs. Calculation of infectious titers was performed according to the Reed and Muench method (54).

Serological analysis. To test serum samples for broadly reactive influenza NP antibodies, a competitive ELISA with plates coated with the NP protein of A/Hong Kong/4806/97 (H9N2) was used as described by Selleck and Kirkland (55). Serum was considered antibody positive if the sample was inhibited >60% of the monoclonal antibody (MAb) binding. To test serum samples for specific HA antibodies, a hemagglutination inhibition (HI) assay using turkey erythrocytes was used following previously described methods (56).

Dataset preparation and preliminary phylogenetic analysis. To elucidate the evolutionary relationships of viruses isolated in this study, we first carried out a provisional phylogenetic analysis for the hemagglutinin and neuraminidase genes, including all available H11 and N2 subtype influenza gene sequences, and for the internal gene datasets, including viruses of all subtypes, using the maximum likelihood method with a general time reversible (GTR) nucleotide substitution model in FastTree 2 (57). The purpose of these large-scale phylogenetic analyses was to identify broad relationships between the new sequences and established influenza A viruses from multiple hosts. These analyses indicated that the Adélie penguin viruses were distantly related to previously detected avian influenza virus and their derived lineages; hence, for all subsequent analyses, the dataset sizes were reduced by identifying representative sequences at 96% identity using the software CD-HIT (58). Due to the geographical proximity of Antarctica to South America, sequences of all publicly available avian isolates from South America were included in our final datasets.

Maximum likelihood and Bayesian relaxed-molecular clock analyses. For the final datasets, maximum likelihood phylogenies were generated using the GTR model with gamma rate heterogeneity, using MrBayes (59). Statistical supports were estimated using 500 maximum likelihood bootstrap replicates. To estimate divergence times, we applied the lognormal relaxed-clock Bayesian Markov chain Monte Carlo method with an SORD6 codon substitution model (60), as implemented in BEAST version 1.7.5 (61). For each dataset, multiple analyses were conducted for 50 million generations, sampling every 5,000 generations, resulting in 10,000 trees. After convergence was ensured, the final 90% of trees were summarized using TreeAnnotator (61) and produced using Fig tree (http://tree.bio.ed.ac.uk/software/figtree/).

HA structural homology modeling. Due to the lack of a published H11 crystal structure, numerous models were made based on the H11 viruses based on the crystal structure of the most closely related HAs available in the RCSB protein databank (http://www.rcsb.org). The H11 structures were modeled based on the HA crystal structures of H13 (PDB ID number 4kps), H16 (PDB ID number 4f23), and H2 (PDB ID number 2wr3) in complex with an α2,3Gal-linked sialic acid analogue (avian-like receptor) using MODELLER (62) and YASARA (63).

Ferret infectivity. Ferrets 6 to 18 months old were administered 1 × 10⁸ EID50/mg egg-propagated virus via the i.n. or i.t. routes. Of two ferrets administered virus via the i.n. route, one was nasal washed on days 2, 3, 4, and 5 before being bled and euthanized on day 14, and the other was bled and given a subsequent intramuscular boost with inactivated virus on day 14 and then bled again and euthanized on day 21. Of two ferrets administered virus via the i.t. route, one was nasal washed on days 2, 3, 4, and 5 before being bled and euthanized on day 14, and the other ferret was culled on day 4 to analyze viral titers in lungs. Weights, temperatures, and clinical signs were monitored throughout the experiment.

Receptor binding assay. The receptor specificity of viruses was assessed using methods described by Rogers and Paulson (64). Briefly, viruses were diluted to a hemagglutination titer of 64 hemagglutinating units (HAU)/25 μl in PBS. A 10% suspension of turkey erythrocytes was treated for 1 h at 37°C with 500 μM/ml Vibrio cholerae enzyme (Sigma). Erythrocytes were washed three times with PBS and resuspended to a 20% suspension. The erythrocytes were then mixed with an equal volume of recombinant beta-galactoside-alpha-2,3-sialtransferase (10 μM/μl) or recombinant beta-galactoside-alpha-2,6-sialtransferase (10 μM/μl) (Japan Tobacco, Inc.) and incubated for 3 to 4 h at 37°C. Erythrocytes were washed a further three times with PBS and resuspended to 1%. Hemagglutination assays were conducted using untreated erythrocytes, sialidasetreated erythrocytes (Asialo), and sialidase-treated erythrocytes resialylated to contain either terminal SAα2,3Gal or SAα2,6Gal linkages on cell surface oligosaccharides.

Cell culture infectivity assay at 33°C and 37°C. Madin–Darby canine kidney (MDCK) cells (American Type Culture Collection; CCL-34) were grown to near confluence in 96-well plates (5% CO2, 37°C). Log10 virus dilutions were made in Dulbecco’s modified Eagle’s medium (SAFC Biosciences) containing sodium bicarbonate (3%) with the addition of 2 mM glutamine (SAFC Biosciences), 1% nonessential amino acids (Sigma), 0.05% NaHCO3 (Sigma), 0.02 M HEPES (Sigma), 4% penicillin and streptomycin (Sigma), and 4 μg/ml trypsin (Sigma). Cells were washed twice with PBS before infection with four replicates of 10-fold dilutions of
each virus. After incubation for 5 days at 33°C or 37°C in 5% CO₂, each well was scored for virus growth by cytopathic effect and hemagglutination of turkey erythrocytes, and the dose required to infect 50% of wells (50% tissue culture infectious dose [TCID₅₀]) was determined by the Reed and Muench method (54).

Nucleotide sequence accession numbers. Full genome sequences from the four H11N2 influenza viruses isolated in eggs have been deposited in GenBank under accession numbers KJ729348 to KJ729379.

SUPPLEMENTAL MATERIAL
Supplemental material for this article may be found at http://mbio.asm.org/lookup/suppl/doi:10.1128/mBio.01098-14/-/DCSupplemental.

Figure S1, PDF file, 0.2 MB.
Figure S2, PDF file, 0.3 MB.
Figure S3, PDF file, 2 MB.
Figure S4, PDF file, 1.8 MB.
Figure S5, PDF file, 0.1 MB.
Figure S6, PDF file, 0.1 MB.
Table S1, DOC file, 0.1 MB.
Table S2, DOC file, 0.1 MB.

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