Characterisation of a highly pathogenic influenza A virus of subtype H5N2 isolated from ostriches in South Africa in 2004

Celia Abolnik,a Brandon Z. Londt,b Ruth J. Manvell,b Wendy Shell,b Jill Banks,b Gertruida H. Gerdes,a George Akol,c Ian H. Brownb

aARC-Onderstepoort Veterinary Institute, Private Bag x5, Onderstepoort, South Africa. bAvian Virology, Veterinary Laboratories Agency (Weybridge), Addlestone, Surrey, UK. cVeterinary Services, Department of Agriculture, Private Bag x14, Stutterheim, South Africa.

Correspondence: C. Abolnik, ARC-Onderstepoort Veterinary Institute, Private Bag x5, Onderstepoort 0110, South Africa.
E-mail: abolnikc@arc.agric.za

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Abstract

Objectives The HPAI H5N2 strain that caused an outbreak in ostriches of the Eastern Cape Province, South Africa in 2004 was characterized.

Design Haemagglutination inhibition (HI) and agar gel immunodiffusion (AGID) were performed on sera from ostrich farms in the outbreak region, and intravenous pathogenicity (IVPI) tests, reverse-transcriptase-polymerase-chain reaction (RT-PCR), nucleic acid sequencing and phylogenetic comparisons were performed on the HPAI H5N2 virus isolated during the outbreak.

Results The deduced amino acid sequence at the HA0 cleavage site determined by RT-PCR and nucleotide sequencing was PQREKRRKKRGLF and thus the virus fell within the definition of a highly pathogenic virus, but in an IVPI test in chickens on the virus isolated from the index case and a value of 0Æ63 was recorded, which is below the criterion for highly pathogenic viruses in this in vivo test. After a further passage in embryonated eggs a second IVPI was carried out and an elevated value of 1Æ19 was obtained. Cloacal swabs were taken from the initial IVPI birds, inoculated into embryonated chickens eggs and a third IVPI was then performed on the resulting haemagglutinating, infective allantoic fluid. An index of 2Æ73 was recorded.

Conclusions HI tests appeared to be the more sensitive test compared to AGID when testing for antibodies to avian influenza in sera. An ostrich-derived virus with a virulent HA0 cleavage site was not initially virulent in chickens but after passage in the latter the virulence increased. Phylogenetic analyses demonstrated the link between AI viruses carried by wild ducks and those infecting ostriches.

Keywords Highly pathogenic avian influenza, H5N2, intravenous pathogenicity index test, ostrich, phylogeny.

Introduction

The transmissibility and virulence of any avian influenza virus may vary between different species and the clinical signs may be different in different hosts complicating the initial diagnosis.1–3 To date only subtypes H5 and H7 have been reported as causing the severe disease in chickens and other species termed highly pathogenic avian influenza (HPAI) although not all H5 and H7 subtype viruses cause HPAI.

An avian influenza virus infection of ostriches was first diagnosed in South Africa in 1991 4 when high mortality was seen in flocks of young birds, but only an avian influenza virus with low virulence for chickens of subtype of H7N1 was isolated. Subsequent infections of ostriches with low pathogenicity avian influenza (LPAI) viruses of subtypes H5 and H7 were experienced in South Africa, The Netherlands, and Zimbabwe and in birds imported into Denmark and The Netherlands.5–7 Low pathogenicity subtypes H6N8,8 H10N1 and H9N2 have also been isolated in ostriches in South Africa and reports from the USA cite H3N2, H4N2, H4N6, H10N4, H10N7 in rheas and emus as well as subtypes of H5N2, H5N9, H7N1 and H7N3.9 The first report of a natural infection of HPAI virus in rats came from Italy in 2000.10 This occurred during an epidemic of HPAI H7N1 in which hundreds of commercial poultry flocks became infected. The ostriches exhibited nervous signs,
brilliant green urine, loss of appetite and depression before death.

**Objectives**

In July 2004, unexplained deaths in birds showing similar clinical signs to the Italian ostriches occurred in ostrich farms in the Eastern Cape Province of South Africa. This paper reports the characterisation of the causative agent involved in the 2004 outbreak.

**Methods**

**Viruses**

Low pathogenicity avian influenza virus A/Egyptian goose/South Africa/04 (H5N2) (EgZA04) was detected by reverse-transcription polymerase-chain reaction (RT-PCR) during active surveillance conducted prior to the HPAI outbreak in the Oudtshoorn area of the Western Cape Province adjacent to the Eastern Cape Province where the HPAI ostrich outbreaks occurred.11 Egyptian geese (Alopochen aegypticus) are resident, locally nomadic birds in South Africa that frequently come into contact with migratory birds in lagoons, water treatment works, dams and rivers but also have direct contact with farmed ostriches when sharing the same pasture.11

**Virus isolation and identification**

Virological examination of tissue and swab material was carried out by the inoculation of 9- to 10-day-old specific pathogen free (SPF) embryonated fowls’ eggs into the allantoic cavity. The alantoic fluids of all eggs were checked for haemagglutination activity using standard techniques.12,13 The virulence of the isolates obtained was determined by the intravenous pathogenicity index test (IVPI) and the derived amino acid sequence at the cleavage site of the precursor haemagglutinin molecule (HA0). The IVPI test was done following standard procedures13 in ten 6-week-old white leghorn chickens hatched from eggs obtained from a commercial SPF flock and reared at the Veterinary Laboratories Agency (VLA), Weybridge, UK, under SPF conditions. Each bird was inoculated intravenously with 0.1ml of infective allantoic fluid from SPF fowls’ eggs inoculated with the virus. Birds were recorded as normal, sick, very sick or dead each day for 10 days. Serological techniques for the detection of antibodies to avian influenza were carried out using agar gel immunodiffusion for specific influenza A antibodies and haemagglutination inhibition (HI) tests for antibodies specific to the H5 subtype using the H5N2 virus from the primary isolation as an antigen and standard techniques.12,13 Ostrich sera were heat inactivated at 56°C for 30 minutes and adsorbed with packed chicken red blood cells.

**RNA isolation and RT-PCR**

Viral RNA was extracted from harvested allantoic fluids using the QIAquick RNA Extraction Kit (Qiagen, Hilden, Germany) or a MagnaPure system (Roche). The HA1 fragment was amplified using gene-specific primers S1 (5’-AGCAGGGGTATAATCTCTCA-3’) and B2a (5’-TTTT-GTCAATGATGTTGACCCATATTGG-3’) using One-step RT-PCR Kits (Qiagen, Germany). Gene-specific primers and procedures used to amplify and sequence internal genes of Egyptian goose (EgZA04) genes are described elsewhere.14 Polymerase chain reaction amplicons were visualised by ethidium bromide staining after 2% w/v agarose gel (Invitrogen, UK) electrophoresis and purified using QIAquick Gel Extraction Kits (Qiagen, Germany).

**Sequencing and phylogenetic analysis**

Reverse-transcriptase-polymerase-chain reaction products were sequenced by cycle-sequencing reactions using ABI Terminator cycle sequencing V1.1 chemistry on a Perkin-Elmer ABI Prism™ 310 DNA sequencer (Applied Biosystems, Foster City, USA). Sequence data was assembled and edited using the Lasergene version 7 seqman software (DNASTAR Inc.). Sequence alignments were performed using the Lasergene version 7 megalign software (DNASTAR Inc). Maximum Likelihood trees were generated using Phylogeny Inference Package (phylip) software version 3.57c15 with the transition to transversion ratio for the data set calculated using tree-puzzle v5.2.16 Nucleotide distance matrices were generated using the Lasergene version 7 megalign software (DNASTAR Inc). For phylogenetic tree construction and comparison, sequences generated from isolates submitted to VLA Weybridge as part of its EU/OIE/FAO Reference Laboratory status or obtained from the Influenza Sequence Database (ISD), Los Alamos, USA17 were included in the trees. Sequences submitted to GenBank were assigned the accession numbers FJ519978–FJ519989.

| Table 1. Comparative serology on ostrich sera using HI and AGP tests |
|---------------------------------------------------------------------|
| **HI** | **Positive** | **Negative** |
| **AGPT** | | |
| Positive | 6 | 1 |
| Negative | 18 | 46 |

AGPT, agar gel precipitin test; HI, haemagglutination inhibition. 
*A positive serum is one with a titre of 24 or more.13. Diagnostic sensitivity 25%. Diagnostic specificity 97.9%.
Results

Disease investigation and diagnosis

Respiratory signs, swelling and exudates from the eyes, fluorescent green diarrhoea, depression, emaciation, collapse and death were some of the clinical signs observed in ostriches in the flock from which virus was isolated. On post-mortem examination some cases revealed prominent liver damage comprising multifocal to diffuse necrosis and degeneration often together with haemorrhagic lesions of the heart, lungs, kidney, pancreas and intestine. In some cases miliary granulomas in the lungs and fungal growth on the airsac membranes due to infection with Aspergillus fumigatus were observed. Preliminary laboratory results indicated infection with an HPAI virus and sero surveillance was undertaken in the immediate area.

Further virus isolation attempts were carried out resulting in the isolation of an influenza A virus of subtype H5N2.

An IVPI test was done on the virus isolated from the index case (supplied by Stellenbosch Provincial Veterinary Laboratory) at VLA Weybridge and an index of 0.63 was recorded. After a further passage in embryonated eggs a second IVPI was carried out and an elevated value of 1.19 was obtained. Cloacal swabs were taken from the initial IVPI birds, inoculated into embryonated fowls’ eggs and a third IVPI was then performed on the resulting haemagglutinating, infective allantoic fluid. An index of 2.73 was recorded. In each of the first two IVPI tests the chickens that survived had exhibited marked cyanosis of wattles, combs and legs and became depressed, but by the end of the ten day test period the birds had returned to an apparently normal clinical state. The deduced amino acid sequence at the HA0 cleavage site determined by RT-PCR and nucleotide sequencing revealed a virulent cleavage site containing multi basic amino acids PQREKRRKKRGLF for sequences amplified from the A South Africa/Al23/04 (H5N2) (EGZA04) within the Eurasian H5 lineage (Figure 1). osZA04 was incorporated into a cluster containing isolates from Italy in 2004; (A/duck/Italy/1349/2004, dkIT04), Denmark in 2003/2004; (A/duck/Denmark/1356/2003-maDK03), (A/duck/Denmark/150/2004-maDK04) and Germany in 2003; (A/mallard/Germany/Wv1349/03-maDE03). osZA04 was shown to be most closely related to the low pathogenic isolate, dkIT04, with a nucleotide sequence identity of 95.9%.

Phylogenetic analysis

Phylogenetic trees were generated using a 971 nucleotide length of HA1 from H5 sequences (excluding the haemagglutinin cleavage site) which placed the ostrich South African 2004 isolate A/Ostrich/South Africa/N227/04 (H5N2) within the Eurasian H5 lineage (Figure 1). osZA04 was shown to be most closely related to the low pathogenic isolate, dkIT04, with a nucleotide sequence identity of 95.9%.

Genotyping

Genotyping phylogenetic trees were generated from partial internal protein gene sequences produced from swine, avian and human isolates submitted to VLA, Weybridge or obtained from the Influenza Sequence Database (ISD) (data not shown). Phylogenetic analyses of osZA04 internal protein genes placed osZA04 for all gene segments in the Eurasian influenza lineages within clusters containing H7 isolates from Italy in 1999/2001; A/turkey/Italy/4603/1999 (H7N1), A/mallard/Italy/43/01 (H7N3) and A/mallard/Italy/33/01 (H7N3); an H1 virus from Denmark in 2002; A/dk/Denmark/74–68 425 1–3/02 (H1N3).

Ostrich ZA 2004 compared to Egyptian goose ZA 2004

To determine the similarity between the Egyptian goose LPAI virus and the HPAI virus, HA, NA, NS and M sequences amplified from the A/Egyptian goose/South Africa/AI23/04 (H5N2) (EGZA04) were compared to the...
Figure 1. Maximum likelihood phylogenetic tree of partial H5 sequences, generated using Phylogeny Inference Package (phylip) software and including sequences generated from isolates submitted to VLA Weybridge as part of its EU/OIE/FAO Reference Laboratory status or obtained from the Influenza Sequence Database (ISD).
sequences obtained for osZA04. Partial HA1/HA2 amino acid alignments between sequences obtained for osZA04 and EgZA04 identified two amino acid differences (D227N and P333L) and the insertion of basic amino acids to generate a HP cleavage site in the ostrich isolate.

A 316 nucleotide partial matrix gene alignment of osZA04 with EgZA04 showed a 98.5% identity between the two isolates. A 457 nucleotide partial non-structural (NS) gene alignment of osZA04 with EgZA04 showed a 98.7% identity between the two isolates.

Conclusions

In 2004, South Africa recorded its first isolation of HPAI virus in 43 years when an H5N2 strain caused increased mortalities in ostrich flocks of the Eastern Cape Province. Due to contrasting reports from the field and from past experimental data further assessments for the pathogenicity of avian influenza in ratites have been carried out in an attempt to clarify the role of these viruses in ratite hosts, but the contribution of viral infection to morbidity remains unclear as it is possible that under field conditions secondary bacterial or fungal contamination may have exacerbated the clinical disease picture (A Olivier, personal communication).

The variable values obtained in the IVPI tests were somewhat unusual and emphasise the importance of using both in vivo pathogenicity tests and, for AI viruses of H5 and H7 subtype, molecular estimations of potential virulence as required by the Office International des Epizooties (OIE) definition of HPAI. The initial IVPI value and that obtained after one further passage in eggs were below the HPAI qualifying value of 1.2, although these viruses would still have been considered HPAI for control purposes because of their HA0 cleavage site deduced amino acid sequences. This being confirmed by the IVPI value of 2.73 obtained after this virus had undergone one passage in chickens. It seems likely that passage in ostriches had for some reason resulted in a virus that was attenuated for chickens, but that passages in chickens restored the virulence for chickens produced a ‘chicken-adapted’ strain with higher virulence. Further investigations into this host adaptation effect on virulence would be valuable.

During active surveillance conducted prior to the outbreak, an LPAI H5N2 virus was detected in an Egyptian goose in the adjacent Western Cape Province. Phylogenetic analysis of the H5 genes confirmed that the LPAI H5N2 Egyptian goose virus was the closest relative to the HPAI H5N2 Eastern Cape ostrich outbreak strain. Some of the internal protein genes of the ostrich virus (PB2, PB1 and PA) were closely-related to other wild duck isolates (LP H5N1, H3N8 and H4N8) isolated in South Africa during the same period from another province. This demonstrates the link between AI viruses carried by wild ducks and those infecting ostriches. Southern African ducks and geese are short-to-medium distance migrants and are thus unlikely to be directly involved in the initial introduction of LPAI from Eurasia into southern Africa. However, about two dozen palearctic migrant species of the orders Ciconiiformes and Charadriiformes overwinter annually in Southern Africa. Viruses shed by migrants into the waterbodies could thus be ingested by local ducks and geese and amplified in these reservoir hosts, which move extensively throughout the region. The majority of ostrich farms in South Africa are located in the Karoo area of the Western Cape Province, a semi-arid region with sparse natural grazing and low annual rainfall. Ostrich farms are often concentrated around rivers and riverine areas, and are used by the ostrich farmers in cultivating irrigated lucerne pastures for ostriches and sheep. Wild indigenous and migratory birds are found in abundance in this area and accumulate in vast numbers on ostrich farms, where they graze alongside ostriches and concentrate in great numbers around the watering troughs and feeders, contaminating ostrich water and feed with faecal material. After the 2004 HPAI H5N2 outbreaks, changes in farm management practices were implemented to minimise the contact between free-ranging ostriches and wild waterfowl, for example by fencing off rivers on ostrich farms, preventing contact with wild birds at ostrich feed points and by regular disinfection of feed and water troughs (Adriaan Olivier, personal communication). Hopefully these biosecurity measures will help to minimise the risk of infection, but again the importance of improved surveillance for AIV in wild birds is highlighted.

Following the detection of the HPAI H5N2 virus disease control and eradication programme resulted in the culling of over 26 000 ostriches from 38 farms in the Eastern Cape Province. Monitoring of depopulated ostrich farms that were subsequently restocked did not reveal any seropositive birds indicating that the virus was successfully eliminated by the measures taken. However, caution must be taken in determining the most sensitive test for detecting antibodies in ratite serum given the results obtained when the field sera were tested. Despite the absence of high mortality in ostriches, caution must still be exercised when moving these possible carriers of HPAI in an attempt to prevent the spread of disease from apparently clinically normal birds and flocks.

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