

RESEARCH ARTICLE

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Genetic and Antigenic Characterization of a Clade 2.3.4.4b Highly Pathogenic H5N1 Avian Influenza Virus Isolated from a Free-Range Layer Duck in India

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Abstract

Highly pathogenic avian influenza (HPAI) H5N1 virus poses a significant threat to the global poultry sector. The report highlights the genetic and antigenic features of a clade 2.3.4.4b HPAI H5N1 virus isolated from a free-range layer duck in the Kuttanad delta of Kerala, India, in 2022. The virus is similar to an H5N1 virus detected in Russia in October 2021 and antigenically distinct from clade 2.3.2, showing a 16-fold reduction in hemagglutination inhibition titer and 23.95% amino acid differences at the HA antigenic sites. The duck virus shares genetic similarities with H5N1 viruses found in Eurasian wild birds, suggesting a role for these birds in its introduction. The study underscores the importance of strengthening avian influenza surveillance to monitor virus evolution and spread.

Keywords: Highly Pathogenic Avian Influenza Virus, H5N1 Subtype, Clade 2.3.4.4b, Free-Range Duck, Phylogenetic Analysis, Antigenic Analysis, India

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INTRODUCTION

HPAI viruses pose a serious threat to India's poultry sector, with more than 500 combined H5N1 and H5N8 outbreaks reported since their first detection in 2006.¹⁻³ Multiple genetic clades of HPAI viruses have emerged over time, including the initial identification of the clade 2.2 H5N1 virus in domestic poultry and sporadic cases in crows.^{4,5} In early 2011, the ancestor of the current clade 2.3.2.1a emerged and spread among poultry, with occasional human cases.^{6,7} The H5N1 clade 2.3.2.1c was detected in poultry during 2014-15, followed by H5N8 clade 2.3.4.4b in wild birds and poultry starting in 2016.^{8,9} In 2020, a new reassortant H5N1 clade 2.3.4.4b caused significant morbidity and mortality in both wild birds and poultry worldwide.¹⁰ Although ducks are known to be silent carriers of HPAI,¹¹ data on HPAI infection in free-range ducks remain limited. Here, we report the genetic and antigenic analysis of a clade 2.3.4.4b H5N1 virus isolated from a free-range layer duck in India.

MATERIALS AND METHODS

Clinical sample processing and animal ethics

We received five free-range layer duck carcasses (12 weeks old) from a single farm in Alappuzha district (Village: Cheruthana, Block: Haripad; Latitude: 9.34139; Longitude: 76.4424) in Kerala State, India, on November 3, 2022. The region lies within the Kuttanad delta region on the west coast of Kerala, where rice is cultivated below sea level.¹² It is home to the Vembanad-Kol wetland (Ramsar site), where many migratory waterfowl from the Central Asian Flyway visit during winter migration (October-March). The birds had a history of corneal opacities, torticollis, and unusual mortality. An autopsy was performed, and organs were collected from all birds. Approximately 100 mg of pooled organ sample was triturated in 900 µL of 1X PBS (pH 7.2) using 2.2 mm bead and tissue homogenizer iRUPT24P (Neuhaus). The homogenizer runs at 3000 rpm for 10 cycles, with a run time and pause time of 15 seconds each and an agitation speed of 5 m/s. The tissue triturate was then centrifuged at 3000 rpm for 5 minutes. Animal experimentation and sample processing were carried out in the animal and laboratory

wings, respectively, of the BLS III containment laboratory in accordance with the Animal Ethics Committee of the ICAR-National Institute of High Security Animal Diseases, Bhopal (M.P.), under approval No. 131/IAEC/NIHSAD/22.

RNA extraction and real-time RT-PCR

Viral RNA was extracted from the supernatant of the tissue triturate using the QIAamp Viral RNA Mini Kit (Qiagen, Germany) according to the manufacturer's protocol. Influenza A virus typing and HA and NA subtyping were performed using the GoTaq® Probe 1-Step RT-qPCR System (Promega, USA).^{5,13,14}

Virus isolation

Isolation of the virus was performed in both 9 day-old specific pathogen-free (SPF) chicken eggs and 12 day-old duck eggs as described earlier.¹⁵ The supernatant of the tissue triturate was used as an inoculum for virus propagation in eggs. A total of 800 µL of inoculum was prepared by treating the supernatant with 2X antibiotics (Antimycotic mix and Gentamicin). The eggs were then inoculated with 100 µL of inoculum each via the allantoic cavity route. The eggs were kept in a humid incubator at 37 °C till the death of the embryo. All the eggs were screened every 12 hours, and the dead eggs were chilled at 4 °C. The infected allantoic fluid was harvested aseptically and stored in 1 mL aliquots at -80 °C until further use. Hemagglutination inhibition (HI) assay with H5-subtype-specific sera was used to confirm the viral subtype.¹⁵

Intravenous pathogenicity index

Intravenous pathogenicity index (IVPI) of the virus (harvested from chicken eggs) was estimated using 6 week-old SPF chickens as per the WOA guidelines.¹⁵

RNA extraction, sequencing and analysis

The RNA was extracted from infected allantoic fluid collected from duck eggs using the QIAamp® Viral RNA Mini Kit (Qiagen, Germany) according to the manufacturer's protocol. The cDNA library was prepared separately using Protoscript® First Strand cDNA synthesis Kit (New England Biolabs, USA) with random hexamer and U12 primer,¹⁶ which were mixed after preparation.

Table. Cross-reactivity of hemagglutination inhibition (HI) antibody titers and amino acid substitutions at the antigenic sites of H5N1 viruses

Virus isolates	Clade	HI titer of antiserum against antigens ^a		Amino acid differences at the antigenic site (out of a total 96 amino acids) ^b	
		dk/11CA04/22	ck/03CL488/11	dk/11CA04/22	ck/03CL488/11
dk/11CA04/22	2.3.4.4b	64	16	0 (0%)	23 (23.95%)
dk/04CA05/24	2.3.4.4b	64	8	5 (5.2%)	24 (25%)
ck/03CL488/11	2.3.2	<2	256	23 (23.95%)	0 (0%)

^aTiters are expressed as the reciprocal of the highest dilution of serum that completely inhibits hemagglutination.

Boldface indicates homologous HI titers

^bAmino acid position and antigenic sites²⁷

Abbreviations: dk/11CA04/22, A/duck/India/11CA04/2022 (H5N1); ck/03CL488/11, A/chicken/India/03CL488/2011(H5N1); dk/04CA05/24, A/duck/India/04CA05/2024 (H5N1)

The complete genome sequencing, using Sanger's chemistry, has been determined as described previously.¹⁷ The sequences have been deposited in GenBank under accession numbers PP340455-PP340462. Representative avian influenza virus sequences were taken from public database/GenBank for comparison. The nucleotide sequence alignment was performed using BioEdit (ver 7.7.1),¹⁸ and phylogenetic relationships were estimated using MEGA (ver 7.0.26).¹⁹ Neighbor-joining trees were constructed for all 8 genes [polymerase basic (PB) 2, PB1, polymerase acidic (PA), HA, nucleoprotein (NP), NA, matrix (M), and non-structural (NS)] by using Tamura-Nei model of nucleotide substitution available in MEGA software. Trees were statistically evaluated using the bootstrap method (with 1000 resampling datasets).

Polyclonal sera and antigenic analysis

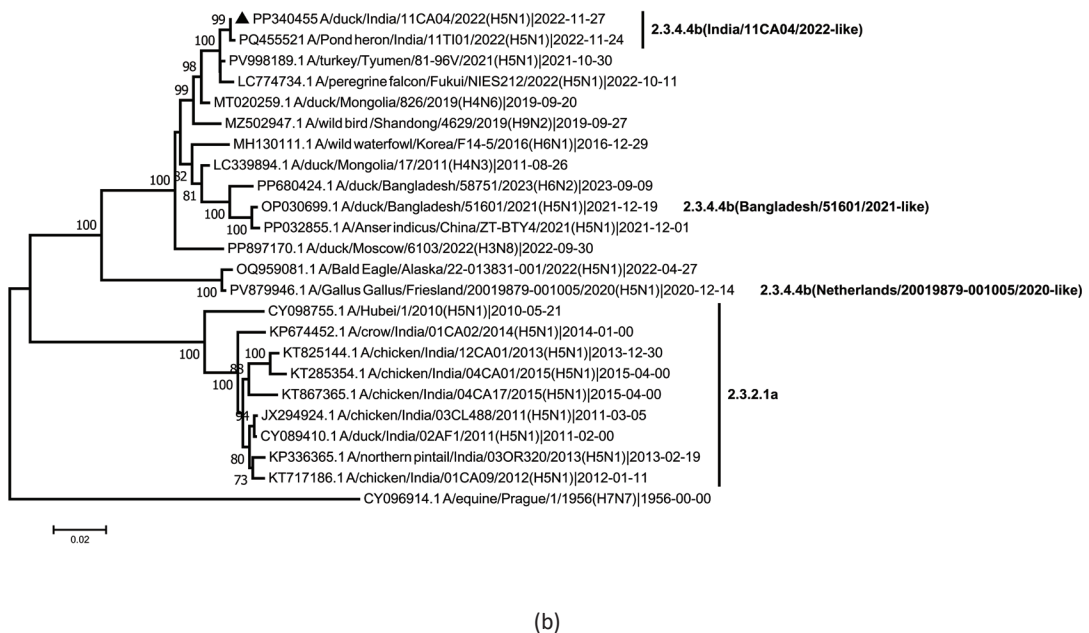
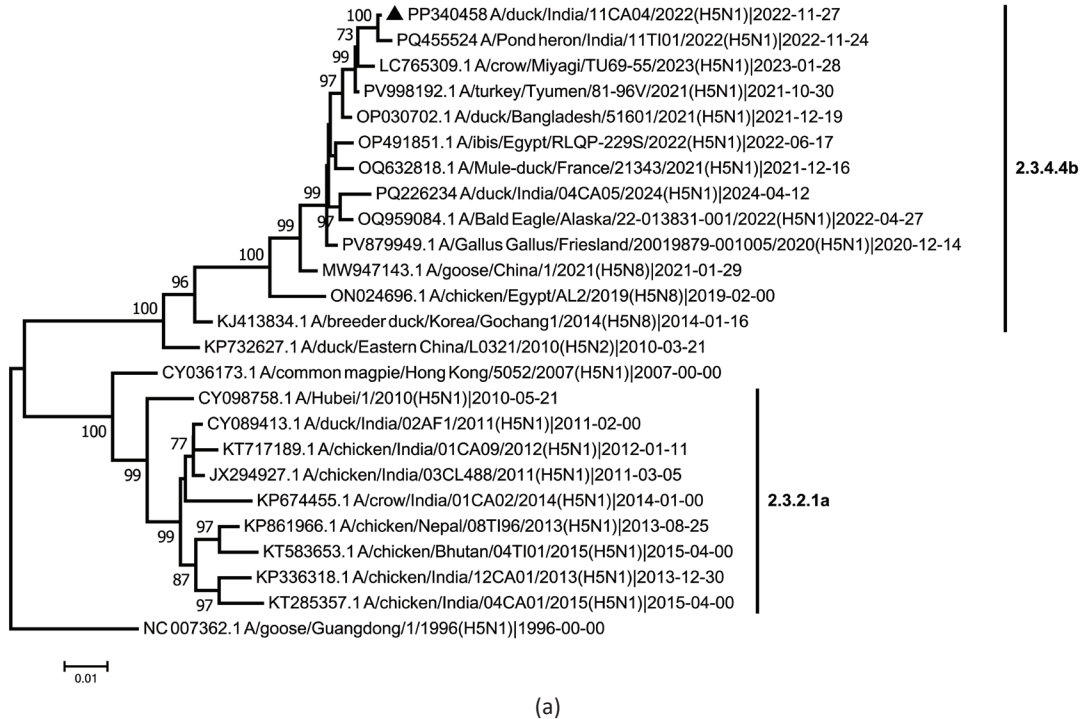
Polyclonal antisera were raised in chickens as described previously.²⁰ The H5N1 virus [A/duck/India/11CA04/2022 (dk/11CA04/22)] harvested from the chicken eggs used in the antigenic analysis was inactivated by β -propiolactone at a final concentration of 0.1%. Three SPF chickens were inoculated with the inactivated virus along with the adjuvant. The first inoculation used Freund's complete adjuvant, followed by a booster dose on the 14th day, with Freund's incomplete adjuvant. The serum was collected on 21st day, aliquoted, and stored at -40 °C until further use. Other serum and viruses, A/Duck/India/04CA05/2024(H5N1) and A/Chicken/India/03CL488/2011(H5N1) used in this study were obtained from the repository

of ICAR-NIHSAD, Bhopal. Antigenic analysis was performed using the hemagglutination inhibition assay.¹⁵

RESULTS AND DISCUSSION

Out of the five samples received, 4 were positive for influenza A (H5N1 subtype) by real-time RT-PCR and virus isolation. One representative isolate, A/duck/India/11CA04/2022 (dk/11CA04/22), was selected for further study. The complete genome of dk/11CA04/22 was amplified, and the virus was characterized as HPAI because it harbors multiple basic amino acids at the HA cleavage site (PLKEKRRKR/G), which was confirmed by an IVPI index of 2.96 (out of 3.0) in chickens.¹⁵ The duck isolate had conserved amino acid residues at HA positions Q222 and G224 (H5 numbering), indicating a preference for avian-like receptors.²¹ The PB2 mammalian virulence (E627K) and host-adaptation (D701N) factors were absent in the isolate. The amino acid markers in M2 (V27 and S31) and M1 (I43V) indicated sensitivity to amantadine and increased virulence of the duck H5N1 virus in mice, chickens, and ducks, respectively.

The dk/11CA04/22 revealed high nucleotide sequence identities (99.5%-100%) with a free-flying bird virus (A/pond heron/India/11TI01/2022 (H5N1)) isolated in India, and a turkey virus (A/turkey/Tyumen/81-96V/2021(H5N1) (99.1%-99.9%) isolated in Russia across all 8 genes, indicating an epidemiological link between the outbreaks. Furthermore, dk/11CA04/22 had over 98.0% homology across



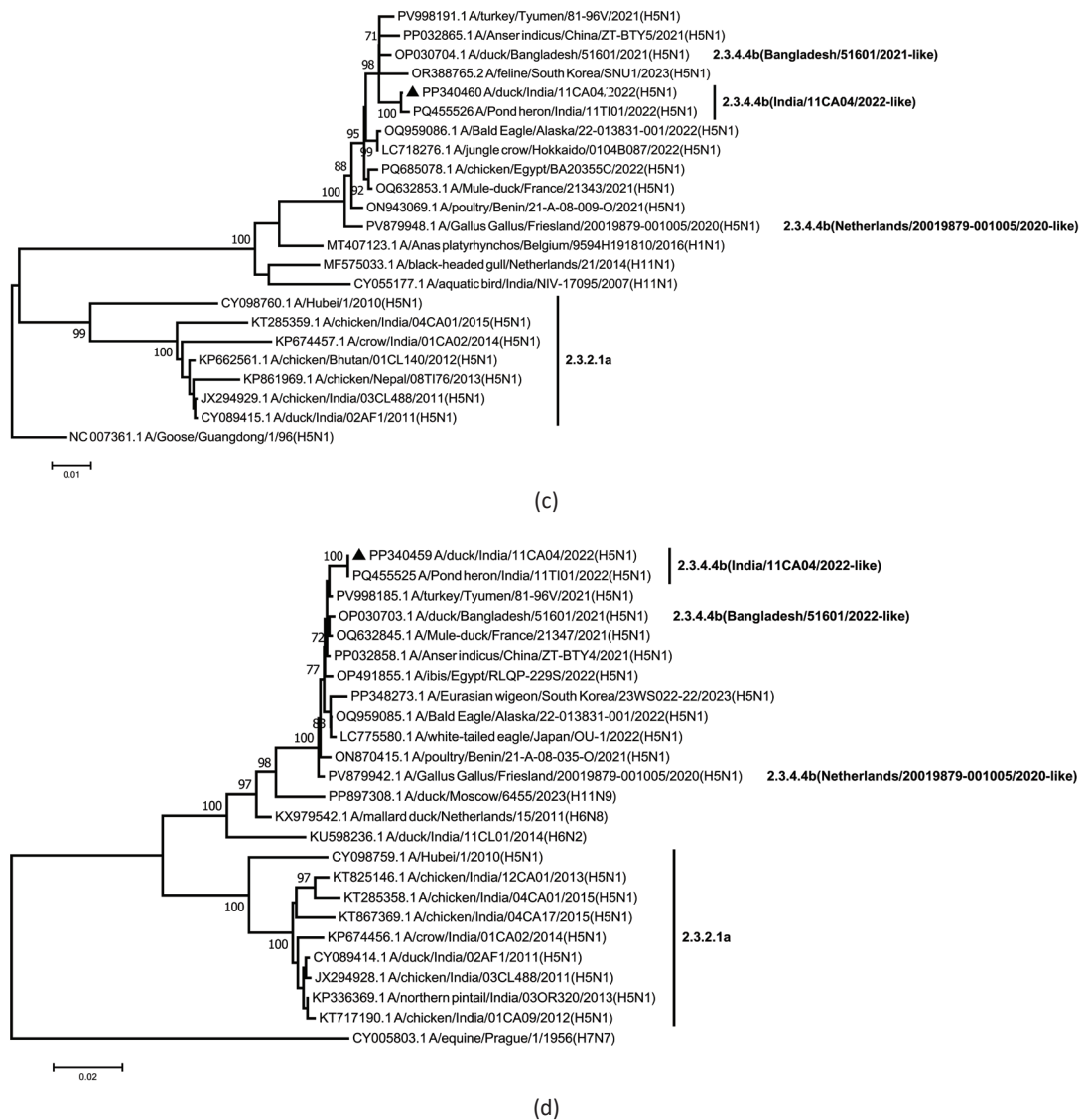


Figure 1. Neighbor-joining phylogenetic trees of (a) Hemagglutinin (nucleotide positions 29-1735), (b) PB2 (nucleotide positions 28-2307), (c) NA (nucleotide positions 21-1433), and (d) NP (nucleotide positions 46-1542) genes of H5N1 and other viruses. Bootstrap values ($\geq 70\%$, out of 1,000 replicates) are indicated near the nodes. The virus sequenced in this study is highlighted by a solid triangle. The scale bar indicates the number of nucleotide substitutions per site. Clade designations are shown to the right. The HA and NA trees were rooted to A/goose/Guangdong/1/1996(H5N1) and the PB2 and NP trees to A/equine/Prague/1/1956(H7N7).

all genes, except PB2, with H5N1 viruses detected during 2021-2022 in Asia, Africa, Europe and North America (data not shown). In the PB2 gene, the Indian isolates, along with H5N1 viruses from China, Japan and Russia, showed a close genetic relationship (98.6% homology) with a low pathogenic avian influenza (LPAI) virus from Mongolia (A/duck/Mongolia/826/2019(H4N6),

indicating reassortment. Phylogenetic analysis placed dk/11CA04/22 within clade 2.3.4.4b (Figure 1A). Within this clade, dk/11CA04/22 grouped with recent H5N1 viruses isolated in Asia, Europe, and North America. Since the emergence of the novel H5N1 clade 2.3.4.4b virus in late 2020, it has been predominantly detected in poultry and wild birds in Europe and has spread globally.²² By October

2021, it was detected in the Russia (A/turkey/Tyumen/81-96V/2021(H5N1). It is likely to have been spread to India via winter bird migration along the Central Asian flyway. Barman and co-workers identified the H5N1 clade 2.3.4.4b virus in Bangladesh in 2021.²³ The Indian virus shares seven genes with the Bangladesh virus (A/duck/Bangladesh/51601/2021(H5N1), with only the PB2 gene differing (Figure 1A, B, C, and D [NP gene; representative of other internal genes]). The PB2 gene contributed by LPAI viruses of A/duck/Bangladesh/58751/2023(H6N2)-like and A/duck/Mongolia/826/2019(H4N6)-like, respectively, with Bangladesh and Indian viruses (Figure 1C). A similar reassortant genotype (G10) virus was identified in China, with only the PB2 gene differing from that of the ancestral genotype (G1) virus that first emerged in Europe.²²

Antigenic analysis using chicken antisera in an HI assay was performed to assess cross-reactivity between clade 2.3.4.4b and clade 2.3.2 (the ancestor of clade 2.3.2.1a) H5N1 viruses. Chicken antisera raised against dk/11CA04/22 did not react with the clade 2.3.2 virus A/chicken/India/03CL488/2011(H5N1); however, antisera against the clade 2.3.2 virus reacted with clade 2.3.4.4b viruses, with titers 16- to 32-fold lower, indicating antigenic divergence (Table). Similar inter-clade antigenic diversity among H5 subtypes has been reported in China.²⁴ Although both clade 2.3.4.4b viruses are antigenically similar, they have HA substitutions (23.95-25%) that have caused antigenic drift compared to the clade 2.3.2 virus. In addition to H5N1 HPAI, an H9N2 LPAI virus is circulating among poultry in India, with sporadic transmission to humans.^{25,26}

CONCLUSION

This study details the antigenic and genetic characteristics of an H5N1 clade 2.3.4.4b HPAI virus isolated from a free-range layer duck. The findings suggest that wild migratory birds likely introduced the H5N1 virus to India through interactions with free-range ducks in wetlands. The H5N1 clade 2.3.4.4b virus is antigenically distinct from the clade 2.3.2 virus, a distinction that should inform future control strategies. In addition to H5N1, an H9N2 LPAI virus is circulating among poultry and ducks in India and occasionally infects

humans. The ongoing co-circulation of H5N1 and H9N2 viruses increases the risk of reassortment, potentially producing novel influenza virus(es) with pandemic potential. Therefore, the study highlights the need to strengthen surveillance to track the evolution and spread of these viruses.

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CONFLICT OF INTEREST

The authors declare that there is no conflict of interest.

AUTHORS' CONTRIBUTION

SN, HVM, AS and CT conceived and designed the study. KG, PKN, DKP, SA and MK performed the experiment and sequencing. CT, MK and SN performed data and phylogenetic analysis and wrote the manuscript. All authors reviewed, revised and approved the final manuscript for publication.

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DATA AVAILABILITY

Molecular sequence data generated in this study have been deposited in GenBank under the accession numbers PP340455-PP340462.

ETHICS STATEMENT

This study was approved by the Institutional Animal Ethics Committee (Approval No. 131/IAEC/NIHSAD/22).

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