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Emergence and development of H7N9 influenza viruses in China

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Abstract

The occurrence of human infections with avian H7N9 viruses since 2013 demonstrates the continuing pandemic threat posed by the current influenza ecosystem in China. Influenza surveillance and phylogenetic analyses showed that these viruses were generated by multiple interspecies transmissions and reassortments among the viruses resident in domestic ducks and the H9N2 viruses enzootic in chickens. A large population of domestic ducks hosting diverse influenza viruses provided the precondition for these events to occur, while acquiring internal genes from enzootic H9N2 influenza virus in chickens promoted the spread of these viruses. Human infections effectively act as sentinels, reflecting the intensity of the activity of these viruses in poultry.
Highlights

- H7N9 virus was generated through sequential reassortments in ducks and chickens.
- This virus originated from eastern China and has spread to over 20 provinces.
- Regionally distinct lineages have been established with different genotypes.
- Viruses present at live poultry markets fueled the recurrence of human infections.
- Chickens served as the source of human infections in each outbreak wave.
Main text

Introduction

Transmissions of influenza viruses from wild aquatic birds to poultry have been seen in many different countries almost every year and may lead to sporadic human infections [1]. However, it is rare that an avian influenza virus not yet recognized in poultry is first identified as the causative agent of human infectious diseases. The two examples that stand out are the 1997 Hong Kong H5N1 and the 2013 H7N9 avian influenza outbreaks [2,3].

Human infections with the H7N9 influenza virus total 679 cases from the three outbreak waves (waves I to III) in China (Figs. 1-2), with a fatality rate of about 40% (Food and Agriculture Organization of the United Nations; URL: http://www.fao.org/ag/againfo/programmes/en/empres/H7N9/wave_3/Situation_update_2015_09_29.html). The incidence of human infections was affected by disease control measures, seasonality factors and reflected the activity of the virus in poultry [4-10]. Although the increasing numbers of human infections with the H7N9 influenza viruses have raised concerns of a pandemic threat, the implementation of measures to control the spread of this and related viruses has remained controversial.

In this review we summarize the emergence and epidemiology of human infections with the H7N9 virus, its origin and genesis in birds and its subsequent evolution, dissemination and development. We follow this with a discussion of the emergence of related viruses such as the H10N8 virus and consider the ecological factors in China that favour the emergence of influenza viruses and look at control measures that can be applied in this context.
Emergence and epidemiology of human infections of the H7N9 influenza virus

Even though it was first recognized in Shanghai [3], the 2013 H7N9 avian influenza virus likely emerged from Zhejiang or Jiangsu in the Yangzi River Delta region adjacent to Shanghai. Zhejiang reported the largest number of human infections in wave I (Fig. 1), which might favour it being the source of the H7N9 influenza virus, and reported the index cases of waves II, III and IV [9-12] (World Health Organization; URL: http://www.who.int/entity/influenza/human_animal_interface/avian_influenza/archive/en/index.html). Phylogenetic analyses revealed that the H7N9 viruses from the Yangzi River Delta region had the highest diversity relative to those from other regions and were often closer to the roots of the H7 and N9 phylogenies than viruses from other regions [13]. H7N9 viruses were introduced into Guangdong in April 2013 and then spread rapidly in the Pearl River Delta region in poultry [13-16]. During waves II and III, Guangdong reported the largest number of cases of human infections (Fig. 1).

The direction of dissemination of the wave I virus has been from the eastern provinces (Zhejiang and Jiangsu) to their nearby regions, such as Anhui, Henan, Beijing, Shandong, Fujian, Jiangxi, Hunan and Hubei (Fig. 1). In wave II and III, the viruses disseminated to affect over 20 provinces in eastern, southern and central China and Xinjiang (Fig. 1). Imported cases have also been reported in Taiwan, Macao, Hong Kong, Canada and Malaysia (Fig. 1).

The emergence of human infections with avian influenza viruses is usually associated with the activity of viruses in poultry, including virus prevalence, poultry type and density, mode of exposure and transmission pathway. Although family or hospital clusters of human cases were recognized, it is believed that the H7N9 avian viruses cannot efficiently transmit from human-to-human [4,17-22].
At least two thirds of the human infection cases with the H7N9 viruses had a history of exposure to chickens before disease onset, mostly at live-poultry markets [4-6,9,23]. It was also noticed that after the human infections were confirmed, local authorities could often isolate or detect the H7N9 influenza viruses at live-poultry markets, mainly from chickens or environmental samples [10, 23-25]. Closure of the live-poultry markets in different cities has effectively stopped the disease outbreak afterwards [7,8,24-28]. Unfortunately, due to the asymptomatic infections [29-31] and the lack of timely and effective surveillance in poultry, infections in humans actually act as sentinels of the presence of the virus in birds.

**Origin and genesis of the H7N9 influenza virus**

After the sequences of the H7N9 avian influenza viruses became publicly accessible, initial analyses concluded that this virus was generated by reassortment between viruses from wild bird H7 and N9 viruses (surface genes) and poultry H9N2 viruses (internal genes) [23,32,33].

Systematic analyses of long-term influenza surveillance showed that the H7 subtype of influenza viruses had been transmitted on multiple occasions from migratory birds to domestic ducks during the last twelve years and maintained in domestic ducks [34] (Fig. 3). N9 viruses had similar introductions and evolutionary pathways [34]. Thus, domestic ducks likely played key roles by maintaining wild bird viruses, increasing their prevalence, facilitating the formation of precursor viruses and transmitting viruses to chickens (Fig. 4).

After introduction to chickens, H7 and N9 viruses reassorted with the enzootic H9N2 viruses, adopting their internal gene complex to form the H7N9 virus that emerged in 2013 [32-36] (Fig. 3). Incorporation of the H9N2 internal gene
complex likely increased the ability of viruses with aquatic bird origin surface genes to replicate in chickens, thereby facilitating their spread in a new host.

The detection of H7N7 duck influenza viruses and the genetically related H7N7 chicken viruses, which had internal genes from enzootic H9N2 viruses (Fig. 3), at the same live-poultry market in Zhejiang in April 2013 [34], suggest that this type of interspecies transmission and subsequent reassortment occurs more frequently than previously recognized.

**Evolution, dissemination and development of the H7N9 influenza virus**

Phylogenetic analysis of the HA and NA genes showed that the early H7N9 viruses formed a highly polytomous clade [13,34] (Fig. 2), suggestive of dispersal of near identical viruses from a common source. However, analyses of their internal genes revealed multiple genotypes, with internal genes from different H9N2 sub-lineages [13,33-36] (Fig. 3). This genotypic diversity may be caused by sequential reassortments after the viruses were introduced to different regions or by several distinct reassortment events occurring approximately simultaneously when the viruses were initially generated.

Early H7N9 viruses, with highly similar surface genes, were detected in six provinces or municipalities (i.e., Shanghai, Zhejiang, Jiangsu, Anhui, Shandong, Hunan and Guangdong) within a very short period of time (less than 1.5 months) [13,33-36], and in each wave human and chicken viruses were always clustered together in the phylogenetic trees (Figs. 2-3). This suggested that the H7N9 viruses were transmitted rapidly in the field, likely by transportation of poultry as part of the marketing system. Spatio-temporal aggregation of chicken viruses and human cases was also observed, supporting a market-based dissemination of the H7N9 viruses [37,38].
The outbreaks of waves II and III started in the early autumn of 2013 and 2014 respectively, with the H7N9 influenza viruses developing into different clades and being disseminated over a wider region of China than in wave I [9-13] (Fig. 2). Phylogenetic analyses of H7N9 surface genes showed three distinct clades, A, B and C, that were responsible for all reported human infections in waves II and III where sequences were available [13] (Fig. 2). Viruses in clade A were descendants of those maintained in Eastern China, and were occasionally transmitted to other regions such as Guangdong and Jiangxi [13]. Clade B was formed by viruses exclusively from Guangdong, and was especially localized in the Pearl River Delta region, indicating that these viruses were maintained and diverged in this region, which imports but rarely exports chickens [13]. The root of clade B could be traced to the earliest strains detected in Guangdong in April and May 2013 [13-16]. Clade C was largely isolated from Jiangxi during the Wave II, but also found in human cases imported to Taiwan from Jiangsu [13] and from Xinjiang, the far northeast province (Fig. 2).

Whole genome analysis of waves II and III H7N9 viruses revealed their shift from the original major genotype of wave I to a number of genotypes that predominated in different regions [13]. No viruses of the original major genotype were detected in the subsequent waves [13]. Many of the new genotypic variants acquired internal genes from local H9N2 viruses, resulting in more than 30 genotypes in wave II [13,33,35,36]. Two H7N6 reassortant variants, with N6 genes from H5N6 viruses and all other segments from H7N9 viruses, were detected in Wave II [13]. Wave III viruses emerged from each of the wave II clades (A, B and C), indicating the continuing presence and evolution of the viruses (Fig. 2).

The emergence and development of the H10N8 influenza virus

Coincident with the second outbreak wave of H7N9 in the winter of 2013-2014, an avian H10N8 influenza virus emerged and caused three human infections in
Nanchang, Jiangxi province, a region adjacent to Poyang Lake, a major aggregation site for overwinter migratory birds and the biggest freshwater body in eastern Asia. The first case was reported in November 2013 and the second and third in January and February 2014, respectively [39,40].

Surveillance and phylogenetic analyses revealed a similar genesis pathway to that of the H7N9 influenza virus [40-43]. The H10 subtype of avian influenza viruses was repeatedly introduced from migratory aquatic birds to domestic ducks at sentinel farms on Poyang Lake during the past decade, and was detected in ducks at live-poultry markets in Nanchang [42]. However, neither the H10 nor N8 subtypes of avian influenza viruses had been detected in chickens before August 2013, despite the constant interaction of ducks with chickens [42]. Emergence of the H10N8 virus in chickens was accompanied with the incorporation of the H9N2 internal genes [40-43]. An outbreak of the newly reassorted H10N8 viruses in chickens was observed at the local markets, prior to the appearance of the human infections [42,43]. Subsequently the H10N8 virus reassorted with H5N6 viruses prevailing in multiple types of poultry in southern China to generate H10N6 viruses in chickens [42].

The emergence of the H10N8 and H10N6 viruses in Jiangxi province mirrors the scenario observed in the H7N9 outbreak and the generation of H7N7 and H7N6 viruses in eastern China. However, due to market closures, no H10N8, H10N6, H7N7 and H7N6 viruses have been detected since mid-2014. Only H7N9 viruses have persisted in the field.

**Factors of influenza ecology in China that favour the virus emergence**

China is home to the world’s largest human population. It also hosts the largest numbers of domestic ducks, geese, chickens, minor poultry and pigs. Frequent interactions among all these species and with wild life provide an ideal interface
for transmission of viruses and the introduction of new viral gene components to the hosts.

With rapid economic growth and increasing wealth, farming and trading practices in China have been greatly changed by industrialization and globalization, in the quest for greater production and better quality food sources. Long-distance transportation of poultry and livestock to congregation sites, such as large wholesale markets or distribution centers, allows mixture and exchange of genetic material from viruses of different sources to generate novel variants and permits infectious agents to proliferate in larger, more diverse populations. Redistribution of asymptomatically infected live birds to retail markets, which are found in almost all cities and towns in China, allows a virus to be disseminated on a nationwide scale in weeks or months, leading to extensive human exposure.

As the H9N2, H5 and H6 subtypes of avian influenza viruses have become enzootic in the poultry of China since the mid-1990s [44-47], contribution of well-adapted genes from these viruses may lead to the genesis of novel viruses with increased infectivity and transmissibility in poultry, possibly in humans as well.

A model of the genesis and interspecies transmission of novel influenza viruses in China’s ecosystem (Fig. 4) shows that viruses from the wild aquatic birds, the natural reservoir for influenza A virus, can be occasionally introduced into the poultry production system by their sharing water bodies and habitats with domestic ducks (interface 1). Close interactions between ducks and chickens, either at the farms or live poultry markets (interface 2), allows viruses to enter terrestrial poultry, which can facilitate reassortment with enzootic chicken viruses to generate novel variants that may have the potential to infect humans. Extensive human exposure to such variants at live poultry markets (interface 3) can lead to sporadic human infections or even disease outbreaks as happened with the H7N9
virus. Measures interrupting the live poultry supply chain or the virus transmission chain, especially those that act on the three interfaces, can help to control or prevent the emergence and dissemination of viruses.

**Conclusion**

H7N9 viruses have been persistent in poultry in China which shown a greater transmissibility to humans and other mammalian models than the Asian highly pathogenic H5Nx viruses [48-51], elevating concerns about their pandemic potential. To control H7N9 and other emerging avian influenza viruses, it is necessary to reconsider the management of the agricultural and trading practices currently in place in China. Measures such as, segregation of poultry species during breeding, transportation and trading; implementation of better biosecurity and hygienic controls throughout the entire poultry supply chain; the permanent closure of live poultry retail markets and the implementation of central slaughtering, will help to reduce the risk of novel viruses emerging, disease outbreaks and human exposure to pathogens. Systematic influenza surveillance remains essential for early warning of novel reassortant viruses and interspecies transmission events.

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Reference:


This is the first report of the emergence of H7N9 influenza viruses in humans.


A study of the epidemiology of human H7N9 infections during the first outbreak wave.


This study provides comparative epidemiological data across the three H7N9 outbreak waves in Zhejiang, one of the most affected regions in China.


Using systematic surveillance and large-scale whole genomic sequencing, the authors provide the most comprehensive dataset and evolutionary analyses on the H7N9 viruses during the first two waves, revealing the establishment of
three geographically distinct virus lineages with at least 48 genotypes. Chickens at live poultry markets were shown to be the major source of human infections.


The authors report the first H7N9 viruses introduced to Guangdong province during the wave 1 outbreak.


A systematic analysis of the human-to-human transmissibility of H7N9 viruses.


   This study compared the infectivity and transmissibility of the H7N9 virus in different types of domestic birds and provides evidence of the asymptomatic infections caused by H7N9 viruses in poultry.


The authors proposed a “sequential reassortment” model for the genesis of the H7N9 virus and discuss the genotypic diversity.


Using a combination of active surveillance, screening of virus archives, and evolutionary analyses, the authors revealed the genesis and source of the H7N9 virus. An H7N7 virus closely related to the H7N9 virus was also detected in Zhejiang province.


The authors analyzed the genetic heterogeneity of the wave 1 H7N9 viruses and proposed a “dynamic reassortment” model for the genesis and evolution of this virus.


This is the first report on the emergence of an H10N8 influenza virus in humans.


This study conducts a longitudinal survey and shows the prevalence, interspecies transmission and evolution of H10 subtype influenza viruses in China. It also reveals the genesis and development of the H10N8 virus that caused human infections in 2013-2014.


Figure Captions

Figure 1. Distribution of officially reported H7N9 infection cases in humans since February 2013. (A) Geographic distribution of the H7N9 influenza viruses in China. Provinces in China with confirmed human H7N9 cases are in blue, those without human cases but positive for H7N9 in poultry or the environment are in light blue, while those without any evidence of H7N9 infections are in grey. (B) Number of humans confirmed to be infected with H7N9 during the first (yellow), second (red) and third (green) waves. Data (as of September 2015) are available at the Food and Agriculture Organization of the United Nations website (URL: http://www.fao.org/ag/againfo/programmes/en/empres/H7N9/wave_3/Situation_update_2015_09_29.html).

Figure 2. Epidemiology and evolution of H7N9 influenza virus. (A) HA phylogenetic tree of the H7N9 lineage. H7N9 viruses from the first, second and third outbreak waves have background colors of yellow, red and green, respectively. Clades A, B and C, identified during the second wave are indicated with red branches on the tree. Human isolates are indicated by blue ovals at node tips. SH1 and SH5 indicate the Shanghai/1/2013 and Shanghai/05/2013 viruses, which are closest to the root of the lineage. The scale bar to the left of the tree represents a branch length of 0.002 substitutions/site. (B) Number of human cases of H7N9 infection (data last updated on September 29, 2015, Food and Agriculture Organization of the United Nations website; URL: http://www.fao.org/ag/againfo/programmes/en/empres/H7N9/wave_3/Situation_update_2015_09_29.html). Major intervals are years; minor intervals are weeks.

Figure 3. Phylogenies illustrating the emergence of H7N9 and H10N8 influenza viruses. Phylogenies of (A) H7 and (B) H10 hemagglutinin genes and (C) PB2 polymerase gene were constructed by the maximum likelihood (ML) method in PhyML v3. The names of human and domestic duck viruses are displayed in red and
purple. Lineages of H7N9, H7N7 and H10N8 viruses in (A) and (B) are indicated by red branches. Transmissions of viruses from domestic ducks to chickens are illustrated by red arrows. The PB2 phylogeny (C) shows the recent H9N2 influenza lineage in China. The tree is compressed for visual clarity, with H7N9, H7N7, H10N8 and H5Nx viruses indicated by colored circles as defined in the legend. Branch length scale bars given under the trees are 0.01 substitutions/site.

Figure 4. The current ecosystem of avian influenza viruses in China. Arrowheads indicate the directions of transmissions.
Figure 1.
Figure 2.
(A) H7 hemagglutinin gene

(B) H10 hemagglutinin gene

(C) PB2 polymerase gene

Figure 3.
Figure 4.