DEPARTMENT OF DEFENSE (AFHSB)
Avian Influenza A (H7N9) Surveillance Summary #77
7 JUN 2017 (next Summary 21 JUN 2017)

CASE REPORT: As of 7 JUN, according to WHO, FAO, China's National Health and Family Planning Commission (National HFPC) and provincial governments within China, the total influenza A (H7N9) human case count since the fifth seasonal epidemic of H7N9 began on 1 OCT 2016 is 889 (+20) cases with at least 239 (+1) deaths. The number of new cases reported each week during the current seasonal epidemic has declined since the peak of activity in JAN 2017, with the Hong Kong Center for Health Protection (CHP) reporting just nine cases per week for epidemiological weeks 21 and 22 (21 MAY - 3 JUN). The number of human cases identified this season accounts for approximately 51% (+1%) of all human cases of H7N9 reported since 2013. The total case count since the disease was first identified in APR 2013 is 1,751 cases with at least 571 deaths, in China, Hong Kong, Macao, Taiwan, Malaysia, and Canada. All cases in Taiwan (6), Hong Kong (21), Macao (2), Malaysia (1), and Canada (2) are believed to have been imported from Mainland China. The overall case-fatality proportion among known cases is 33%; the average age of those affected is 55 years; and at least 452 (+5) (26%) of the cases reported have been female. The most recent known date of onset was 20 MAY. Since 2013, China has reported cases in 24 (+2) provinces and autonomous regions: Anhui, Fujian, Gansu, Guangdong, Guangxi, Guizhou, Hebei, Hubei, Henan, Hunan, Nei Mongol (Inner Mongolia), Jiangsu, Jiangxi, Jilin, Liaoning, Shaanxi, Shandong, Shanxi, Sichuan, Tibet, Yunnan, Zhejiang, and Xingjiang; and four municipalities: Beijing, Chongqing, Shanghai, and Tianjin. Since OCT 2016, cases have been reported from 23 (+2, Nei Mongol (Inner Mongolia) and Shaanxi) different provinces and autonomous regions, and all four municipalities of China. Jiangsu Province has reported the most cases of any single province during the fifth seasonal epidemic, 153 (17%) of 889 cases.

On 31 MAY, the Nei Mongol (Inner Mongolia) Autonomous Region in northern China reported its first human case of H7N9 in a 56-year-old male from Bayannur. Shaanxi Province reported two human cases of H7N9 imported from Yulin, Nei Mongol (approximately 452km west of Bayannur) on 31 MAY and 6 JUN, respectively. The cases were husband and wife, and both reported contact with poultry prior to symptom onset. On 17 MAY, Nei Mongol's HFPC reported the region’s first H7N9 positive poultry and environmental samples from live poultry markets (LPMs) in Xing’an, Tongliao, and Xilingol.

TRANSMISSION: In a CDC study published in APR 2015, H7N9 antibodies were found among 6.7% of case contacts identified between MAR 2013 and MAY 2014 in China, suggesting that human-to-human transmission does occur and could cause mild or asymptomatic infections. Since much of the reporting out of China occurs in monthly batches, with limited information on age, gender, and location, it is possible that only the most severe cases and fatalities are being reported by China. It is unknown how many mild or asymptomatic cases have occurred and how many cases have occurred without laboratory testing. This lack of information coupled with the infrequent reporting makes spatial and temporal cluster analysis difficult. CDC reports that at the conclusion of the fourth wave of H7N9, there had been 26 known disease clusters since the beginning of the outbreak in 2013, and that cluster-associated cases accounted for 7% of total reported cases. As of 7 JUN, AFHSB has identified 11 spatial clusters during this season of H7N9.

In a recent study in Nature Communications, researchers from Hong Kong University identified a mutation in the viral genome of H7N9—a unique nucleotide substitution (NS-G540A) acquired through prior reassortment with low pathogenic H9N2—which is responsible for its unique ability to circulate efficiently in avian hosts as well as replicate in mammalian cells. This particular mutation is also present in human infections with avian influenza subtypes H10N8 and H5N6. A recent Eurosurveillance study reported highly pathogenic H7N9 can mutate to include antiviral resistance during prophylaxis or treatment.

SURVEILLANCE: Reagents for surveillance testing purposes are available via the CDC website. NMRC has produced amplicon H7N9 positive testing control material using the published WHO primers/probes. Kits were sent to AFRIMS, NAMRU-3, NAMRU-6, NAMRU-2, Phnom Penh, NMRC-A, and NHRC for surveillance. Nineteen DoD laboratories were sent diagnostic kits, as have all 50 states, the District of Columbia, Puerto Rico, and more than 60 international labs.
BACKGROUND: On 1 APR 2013, WHO reported three human cases of infection with a novel influenza A (H7N9) virus in China. This was the first time human infection with H7N9 had been detected. CDC believes the H7N9 virus is likely a reassortant of H7N3 viruses from domestic ducks and H9N2 viruses from other domestic poultry. Seasonality has been observed since the beginning of this outbreak with a consistent pattern of declining incidence through the summer months followed by a spike in cases in the winter months. Confirmed avian H7N9 has been rare and subclinical but has been previously identified. H7N9 is usually asymptomatic in birds and many bird owners are likely unaware of infections and the risk of transmission. Detection in birds requires routine active surveillance, which FAO reports has resulted in 4,242 virus-positive samples from the environment and chickens, pigeons, ducks, and wild birds since the beginning of the outbreak in 2013.

A recent CDC MMWR study reported that genetic sequences of 74 virus samples collected from infected persons or live bird markets during the fifth epidemic indicate H7N9 viruses have diverged into two distinct genetic lineages, the Pearl River Delta lineage and the Yangtze River Delta lineage; 69 of these samples were of the Yangtze River Delta lineage. CDC’s preliminary data suggests the Yangtze River Delta viruses are antigenically distinct from earlier H7N9 viruses and from existing candidate vaccine viruses (CVVs). On 2 MAR, experts from the WHO Global Influenza Surveillance and Response System recommended two new CVVs for development: 1) an A/Guangdong/17SF003/2016-like virus, which is a highly pathogenic virus from the Yangtze River Delta lineage; and 2) an A/Hunan/2650/2016-like virus, which is a low pathogenic virus also from the Yangtze River Delta lineage. On 23 MAY, CDC reported that development of the second of these two CVVs, which is matched to the H7N9 virus lineage that has predominated during the fifth seasonal epidemic, has been completed and is being shipped to various manufacturers in China. WHO posted information about the availability of this CVV on 18 MAY.

On 27 FEB, WHO reported a change in the genetic sequence of isolates from two confirmed human H7N9 cases in Guangdong Province indicating the virus is evolving from a low pathogenic to a highly pathogenic avian influenza virus. Similar changes were also found in genetic sequence data from another recent case with travel history to Guangdong Province that was reported by Taiwan to WHO on 4 FEB. Additionally, the Ministry of Agriculture (MOA) of China notified OIE on 21 FEB that genetic sequences of virus samples from LPMs in Guangdong showed changes consistent with highly pathogenic avian influenza virus. As of 7 JUN, FAO and OIE have reported the highly pathogenic form of H7N9 in LPMs from Fujian, Guangdong, Guangxi, and Hunan provinces, and from farms and backyard flocks in Guangxi, Hebei, Henan, Hunan, Shanxi, and Tianjin provinces. WHO has noted that there is no evidence this change from a low pathogenic to a highly pathogenic avian influenza virus has had any impact on the pathogenicity or transmissibility of H7N9 in humans. “Highly pathogenic” and “low pathogenic” refer only to the ability of influenza A viruses to cause disease in avian populations and do not describe disease severity, transmissibility, or pathogenicity of influenza A infections in human populations. However, the recent identification of a highly pathogenic form of H7N9 indicates yet another genetic change in the virus not observed in the previous four seasons and with unknown implications for human populations.

On 1 JUN, Emerging Infectious Diseases released two expedited studies on human infections with the highly pathogenic strain of H7N9. The first study reported that through MAR, there had been a total of eight human cases in three provinces (Guangdong, Guangxi, and Hunan). Following the identification of the first two human infections with highly pathogenic H7N9, provincial China CDC labs reportedly have been conducting genetic analyses of respiratory specimens from all confirmed cases of H7N9 in the country. In the second study, authors noted that highly pathogenic H7N9 viruses may pose a higher risk of zoonotic transmission as these viruses “spread systematically within chicken[s] and are likely to be found at high titer in multiple organs” while low pathogenic avian influenza viruses only infect the respiratory and intestinal tracts of birds.

INTERAGENCY/GLOBAL ACTIONS: On 26 JAN, the U.S. CDC announced a Level 1: Practice Usual Precautions travel advisory for China. U.S. CDC and WHO advise no special screenings at points of entry, and no trade or travel restrictions at this time.

(+xx) represents the change in number from the previous AFHSB Summary 24 MAY 2017.

All information has been verified unless noted otherwise.

For questions or comments, please contact: dhc.ntr.health-surv_list.afhs-bp-alert-response@mail.mil

APPROVED FOR PUBLIC RELEASE
Avian Influenza A (H7N9) Human Cases by Estimated Week of Onset
As of 7 JUN 2017 (N = 1,751)
This map illustrates the geographic distribution of human H7N9 cases and H7N9-positive samples in birds or the environment in China since OCT 2015. Human cases are depicted in the geographic location where they were reported; for some cases, exposure may have occurred in a different geographic location. Precise location of 24 human cases in Anhui (2), Beijing (2), Guangdong (1), Guangxi (1), Hebei (3), Hunan (1), Hubei (2), Jiangsu (1), Jiangxi (6), Sichuan (2) and Zhejiang (3) provinces are currently not known; these cases are therefore not shown on the map.
Cumulative Human Cases of Avian Influenza A (H7N9)
1 APR 2013 - 07 JUN 2017

Cumulative Cases
(N = 1751*)

- No Reported Cases
- 1-15
- 16-25
- 26-115
- >115
- Imported Cases

*47 cases from China were reported without location information in 2015 – 2017

Since the beginning of the avian influenza A (H7N9) outbreak, spikes in cases have been associated with seasonality. These seasonal epidemics typically span 1 OCT to 30 SEP of the following year (see the above maps). These maps only illustrate autochthonous cases in China, not imported cases.