Avian influenza A (H5N1) infection with respiratory failure and meningoencephalitis in a Canadian traveller

Naheed Rajabali MD1, Thomas Lim MD2, Colleen Sokolowski MD2, Jason D Prevost MD2, Edward Z Lee MD2

In an urban centre in Alberta, an otherwise healthy 28-year-old woman presented to hospital with pleuritic chest and abdominal pain after returning from Beijing, China. After several days, this was followed by headache, confusion and, ultimately, respiratory failure, coma and death. Microbiology yielded influenza A subtype H5N1 from various body sites and neuroimaging was consistent with meningoencephalitis. While H5N1 infections in humans have been reported in Asia since 1997, this is the first documented case of H5N1 influenza in the Western Hemisphere. The present case demonstrated the typical manifestation of H5N1 influenza but, for the first time, also confirmed previous suggestions from human and animal studies that H5N1 is neurotropic and can manifest with neurological symptoms and meningoencephalitis.

Key Words: Encephalitis; H5N1; Influenza A; Meningoencephalitis

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In 1997, a poultry-originating influenza, H5N1 influenza virus, caused an outbreak in Hong Kong, China. This was the first time it had been isolated from a human patient. Since then, >600 people have been infected with the H5N1 virus worldwide in the past 10 years, with a mortality rate of approximately 61% (1). Human H5N1 virus infection generally manifests as severe pneumonia progressing to acute respiratory distress syndrome, but can also present with abdominal pain and diarrhea (1,2). It also demonstrates neurotropism, being isolated in the neurons of animals and asymptomatic individuals (3,4).

CASE PRESENTATION

A healthy 28-year-old woman presented to the Red Deer Regional Hospital (Red Deer, Alberta) emergency department December 28, 2013 with a two-day history of right-sided pleuritic chest pain, mild dyspnea, nausea, right upper quadrant abdominal pain, malaise and chills. Her symptoms began as isolated chest pain during her flight back to Canada after a three-week stay in Beijing, China, and she presented to the hospital the day after her return. Her history was negative for sick contacts, exposure to poultry or birds, and visits to outdoor markets or farms. Her temperature was 39.7°C with a heart rate of 125 beats/min and respiratory rate of 18 breaths/min. The rest of her vital signs were normal. Her physical examination was remarkable only for mild tenderness at the right upper abdominal quadrant. Bloodwork demonstrated an elevated leukocyte count of 12.6×10⁹/L and was negative otherwise, including blood cultures.

The patient was prescribed a five-day outpatient course of levofloxacin with a diagnosis of bacterial pneumonia. She returned to the emergency department in the afternoon of January 1, 2014 with persistent shortness of breath and worsening right-sided chest pain, without cough. She had developed a frontal headache, lightheadness, abdominal pain and several episodes of vomiting. This time, her respiratory rate was 22 breaths/min and she required supplemental oxygen at 2 L/min on nasal prongs to maintain an oxygen saturation of 92%. She had diminished breath sounds in the right lung zones. Her abdomen was diffusely tender, but more so in the right upper quadrant and epigastric region without guarding or distention.

An electrocardiogram showed sinus tachycardia with nonspecific T wave inversions in leads III and aVF. Her initial chest radiograph revealed a subtle rounded consolidation of the right lung apex. A subsequent computed tomography pulmonary angiogram of the chest ruled out pulmonary embolism, but identified a dense rounded consolidation of the right lung apex with surrounding ill-defined ground-glass attenuation. Chest pain was her predominant clinical symptom, and the patient was prescribed a five-day outpatient course of levofloxacin with a diagnosis of bacterial pneumonia.

She returned to the emergency department later that day with fever and headache, with right upper quadrant pain. A chest radiograph at 17:31 showed a small right pleural effusion and worsening of the right apical pneumonia. A chest tube was placed on the right, draining 300 mL of slightly cloudy, yellowish-tan pleural fluid. The patient’s antimicrobial coverage was broadened to piperacillin-tazobactam and azithromycin for nonresolving pneumonia. The abdominal pain was attributed to the parapneumonic effusion because it resolved several hours following...
insertion of a chest tube. The pleural fluid was exudative using Light’s criteria and pathology revealed scant benign mesothelial cells with reactive changes in the background of numerous neutrophils. No bacteria were seen on Gram stain and there was no growth on cultures. An abdominal ultrasound revealed two small liver hemangiomas and was otherwise unremarkable.

On the following morning, January 2, she felt worse overall and began to notice blood-tinged sputum with her cough. By the late evening, her oxygen requirements increased to 12 L/min. The chest radiograph demonstrated progressive bilateral lung consolidation. That evening, she had a transient loss of consciousness, with recovery, followed by blurred vision that then improved. She gave a history of intermittent neck pain. A neurological examination was performed, which revealed normal extraocular eye movements, no nystagmus, pupils that were equal and reactive to light symmetrically, normal facial nerves and normal trigeminal nerve sensation. The fundi were not visualized. At this point, the intensive care unit (ICU) was consulted.

In the evening, the patient complained of increasing dyspnea and persisting sharp, right-sided chest pain, again worse with inspiration. She was alert and oriented, denying having a headache, visual changes or neck stiffness. She also denied having a sore throat or abdominal pain.

On examination by the ICU, she showed signs of respiratory distress along with tachypnea at 24 breaths/min saturating at 96% on 8 L/min of inspired oxygen 70%, peak pressure 39 cmH2O, respiratory frequency and respiratory rate of 22 breaths/min, with volumes of 400 mL to 430 mL. Her postintubation arterial blood gases showed a pH 7.37, a partial pressure of carbon dioxide of 36 mmHg, a partial pressure of oxygen of 83 mmHg, a bicarbonate of 21 mmol/L, base deficit 4 and oxygen saturation of 93%.

The chest radiograph revealed right greater than left dense bilateral consolidation, consistent with worsening pneumonia (Figure 1). At approximately 07:00 January 3, the patient became hemodynamically unstable. She was tachycardic with heart rates varying between 120 beats/min and 150 beats/min. Her blood pressure was labile, rising to 220/120 mmHg. Initially, it was believed that she was inadequately sedated and her midazolam and fentanyl infusions were increased. However, her tachycardia and hypertension persisted. When examined, the patient’s pupils were dilated and unresponsive to light. At this point her midazolam and fentanyl infusions were discontinued. Her blood pressure initially normalized, then over the next 1 h to 2 h she became hypotensive requiring inotropic support. She was intravenously given mannitol 20% and furosemide, and hyperventilated to a partial pressure of carbon dioxide of 30 mmHg.

A neurological examination was completed at approximately 4 h and 8 h after sedating infusions were discontinued. She was unresponsive to painful stimuli, had absent cranial nerve responses and failed the apnea test. A nonenhanced computed tomography scan of the head (Figure 2) demonstrated generalized cerebral parenchymal swelling, diffuse sulcal space and cisternal space effacement suggestive of meningoencephalitis.

A magnetic resonance examination of the brain (Figure 3) was obtained, confirming diffuse sulcal space and cisternal space effacement with diffusely increased signal of the extra-axial cerebrospinal fluid (CSF) spaces and ependyma of the lateral ventricles on fluid-attenuated inversion recovery imaging. The increased signal of the CSF spaces on fluid-attenuated inversion recovery imaging indicated proteinaceous debris within the CSF spaces and along the ependymal surface. Diffuse parenchymal swelling and slightly increased T2-weighted signal of the cortex of the temporal lobes, insular cortex and hippocampal regions was noted bilaterally, suggestive of encephalitis. Gadolinium-enhanced magnetic resonance angiography revealed markedly delayed intracranial arterial flow, with partial opacification of the anterior cerebral, middle cerebral and posterior cerebral arteries at 4 min postcontrast injection. Brisk flow to the scalp vessels was noted. Enhancement of the meninges was not observed, likely secondary to diminished intracranial blood flow.

The neurologist on call examined the patient 11 h after sedating medications had been discontinued. The abnormal findings of the earlier
neurological examination were confirmed. The patient met the clinical criteria for brain death and was pronounced dead. A lumbar puncture was performed following brain death and before discontinuation of cardiorespiratory life support.

Samples from the nasopharyngeal swab, endotracheal aspirates and CSF tested positive for influenza A type H5N1. CSF showed 2+ white blood cells, 2+ red blood cells, without bacteria seen. Details of diagnostic specimen analysis and virological investigations were reported by Fonseca (5). Autoimmune serologies obtained later returned positive for perinuclear antineutrophil cytoplasmic antibody. Myeloperoxidase and antiluminal basement membrane antibodies were both negative.

**DISCUSSION**

We report a fatal case of influenza A (H5N1) in a woman, 28 years of age, who presented in January 2014 with pleuritic chest pain, dyspnea, agitation, confusion and coma, after a three-week visit to Beijing, China. She was diagnosed with acute respiratory distress syndrome, secondary to pneumonia and brain death due to meningoencephalitis. The diagnosis of H5N1 was established by isolating the virus from nasopharyngeal swabs, broncho-alveolar lavage and CSF.

Our patient’s demographic is in keeping with what is commonly found in humans infected with H5N1. Most patients with influenza A (H5N1) virus infection have been previously healthy. The median age of patients with influenza A (H5N1) virus infection is approximately 18 years, with 90% of patients ≤40 years of age. Moreover, increases in human cases of influenza A (H5N1) have been observed during cooler months in association with increases in outbreaks among poultry (1,6-8).

Unique to our experience is that this is the first reported case of H5N1 influenza in the Western world because most cases remain endemic in Asia, the Middle East, Europe and Africa. Furthermore, until now, no cases of influenza A (H5N1) illness have been identified among short-term travellers visiting countries affected by outbreaks among poultry or wild birds (9).

Our patient’s initial respiratory manifestation is a common finding in H5N1-infected humans, with the virus primarily affecting the lower respiratory tract causing respiratory failure. Diarrhea is also a common presenting feature of H5N1, occurring in up to 70% of patients. Besides these common presentations, patients have also been noted to complain of vomiting and abdominal pain. These symptoms correlate with findings of viral RNA in fecal samples and viral replication in intestines (1,2,4,10). Our patient complained of vomiting and crescendo-decrescendo abdominal pain, but did not have diarrhea.

Our patient also had mild intermittent headaches, confusion and demonstrated symptoms and a clinical course in keeping with meningoencephalitis. Encephalopathy and encephalitis in humans have been reported in cases associated with seasonal influenza A and B viruses (11,12,13). Only one case of human central nervous system involvement has been suggested by a patient who developed a coma from whom H5N1 was isolated from CSF (10) despite being noted in mammals such as ferrets, mice and felines (8). With respect to the child who developed a coma, imaging of the brain or histological analysis was not performed, and it is uncertain whether the patient experienced encephalopathy or true encephalitis. These reports suggest the H5N1 virus is becoming more neurologically virulent and adapting to mammals. Despite the trend in virulence, the mode of influenza virus transmission remains elusive to date. It is unclear how our patient acquired the H5N1 influenza infection because she did not have any known contact with any animals or poultry.

The nonspecific clinical presentation of H5N1 influenza virus has frequently led to underdiagnoses of subsequently confirmed cases (1). Infection with H5N1 influenza virus should be considered in the differential diagnosis for patients with epidemiological risk factors presenting with systemic manifestations, including the common respiratory and gastrointestinal symptoms, clusters of intermittent abdominal pain and headache, as well as signs of meningoencephalitis.

Having a high index of suspicion for H5N1 infection is essential to the appropriate management of the illness, because treatment with antiviral agents is likely to be beneficial only when it is started early in the course of illness (1,2,14).

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Figure 3) Magnetic resonance examination of the brain revealing diffuse sulcal space and external space effacement with diffusely increased signal of the extra-axial cerebrospinal fluid spaces and ependyma of the lateral ventricles on fluid-attenuated inversion recovery imaging. Diffuse parenchymal swelling and slightly increased T2-weighted signal of the cortex of the temporal lobes, insular cortex and hippocampal regions noted bilaterally suggests encephalitis.
