**Uncommon cause of severe pneumonia: co-infection of influenza B and *Streptococcus***

**Key words**
Influenza B virus; Intensive care; Shock, septic; *Streptococcus*

**Introduction**

Influenza and pneumococcus are two of the most important infectious agents afflicting mankind. Co-infection of these two pathogens poses a particularly serious threat to humans. During the influenza pandemic in 1918, an estimated 40 to 50 million people died as a result and many of the deaths were attributed to co-infection. There are three types of influenza viruses, A, B, and C, of which type C viruses cause only sporadic cases in humans. By contrast, influenza A and B give rise to epidemics; the more devastating epidemics with loss of lives are commonly related to influenza A.1 If influenza A and streptococcal infection co-exist, the mortality is particularly high, whereas with influenza B the infection is usually less severe. To the best of our knowledge, there was only one report of co-infection of influenza and pneumococcus causing severe infection in three young adults known to have had no pre-morbid condition.2

Herein, we report on four young adults infected by a combination of influenza B virus and streptococci, who all experienced fulminant courses with multi-organ failure. It is the first report in Hong Kong of severe bacterial chest infection with septic shock secondary to influenza B infection in previously healthy adults.

**Case reports**

**Case 1**

A 34-year-old previously healthy man was admitted to the hospital because of a severe chest infection in February 2012. He had experienced chills, vomiting, diarrhoea, and severe cough in the 5 days before admission. After admission to the emergency department, his condition deteriorated rapidly with the onset of severe respiratory failure. He was intubated and transferred to the intensive care unit (ICU) for further management.

The initial chest X-ray showed consolidation near the right heart border and left middle zone, which rapidly progressed to bilateral haziness. The complete blood count revealed a normal haemoglobin level (125 g/L), severe leukopenia (0.5 x 10⁹/L), and thrombocytopenia (29 x 10⁹/L). His C-reactive protein level was 219 (reference level, <5) mg/L. He developed acute kidney injury with a serum creatinine level raised to 230 (reference range, 47-82) μmol/L. His liver function test results were normal on admission. Nasopharyngeal aspirate submitted to reverse transcription–polymerase chain reaction was negative for influenza A virus RNA, but positive for influenza B virus RNA. Urinary streptococcal antigen and legionella antigen were negative. Blood culture grew *Streptococcus pyogenes* which was sensitive to penicillin.

The patient developed septic shock with multi-organ failure, and was treated with...
同時感染流感和肺炎球菌會有高發病率和死亡率，起因通常牽涉甲型流感，而乙型流感病毒的感染很少會引致嚴重的情況。文獻記載甲型流感疫情導致多人死亡，特別是在1918年的疫情造成四千至五千萬人死亡。本文報告在香港2011至2012年流感肆虐期間四個乙型流感的病例。四名病人受感染前的健康狀況良好，沒有慢性疾病。他們均因流感症狀被送往醫院，且病情迅速惡化，出現多器官功能衰竭，隨後被確診感染乙型流感及鏈球菌而引致嚴重肺炎。其中三人感染肺炎鏈球菌，另一人則感染化膿性鏈球菌。他們都有白細胞減少症、感染性休克及急性腎損傷。雖然在深切治療部施以積極的抗生素治療和器官功能支持，最終兩人死亡。根據文獻，這是感染乙型流感繼發嚴重侵入性肺炎球菌性肺炎的第二個病例報告。

**Case 2**

A 37-year-old previously well man was admitted to the hospital because of severe pneumonia in March 2012. He had had cough, sputum, and sore throat for 4 days before admission. After admission to the emergency department, his condition rapidly deteriorated and he developed severe respiratory failure, for which he was transferred to the ICU and intubated.

The chest X-ray showed consolidation over right lower lobe, left lingual lobe, and left lower lobe. His complete blood count showed a normal haemoglobin level (153 g/L), normal platelet count (191 x 10^9 /L), and severe leukaopenia (2.4 x 10^9 /L). He also developed acute kidney injury and with a serum creatinine level of 177 μmol/L. His liver function test results were unremarkable. The C-reactive protein level was 143 mg/L. Blood culture grew *Streptococcus pneumoniae* serotype 3. The patient’s condition deteriorated rapidly with the onset of severe shock and respiratory failure. He was intubated and transferred to ICU where he received piperacillin/tazobactam, oseltamivir, and azithromycin.

He developed profound shock with severe metabolic acidosis and disseminated intravascular coagulopathy. Despite treatment with high-dose inotropes and intravenous immunoglobulin, he succumbed on the day of admission to the ICU.

**Case 4**

A 60-year-old man was admitted to the hospital for community-acquired pneumonia, having had breathlessness with cough and sputum for 2 days in January 2012. He had visited Mainland China 20 days earlier. Physical examination yielded crepitations at both lung bases.

A complete blood picture showed a normal haemoglobin level (151 g/L), leukaopenia (0.4 x 10^9 /L), and thrombocytopenia (82 x 10^9 /L). He also had acute kidney injury (creatinine, 110 μmol/L) but liver function test results were unremarkable. The chest X-ray showed consolidation over middle and lower zones of both lungs. The C-reactive protein level was 324 mg/L. Blood and sputum cultures grew *S pneumoniae* serotype 3; the tracheal aspirate for viral culture was positive for influenza B.

He developed severe respiratory failure on the
day of admission and was intubated, mechanically ventilated, and admitted to the ICU for further management. He was treated with cefotaxime, azithromycin, penicillin, rifampicin, fluconazole, and vancomycin. He also received renal replacement therapy for acute kidney injury, but died 3 days after admission.

**Discussion**

There are three types of influenza viruses, A, B, and C. Influenza A is more common and can cause major epidemics with loss of many lives, whereas influenza B infection is usually milder. According to statistics from the Centre for Health Protection, the incidence of influenza A was about 2 to 3 fold that of influenza B.3

Influenza is often complicated by severe chest infection, if there is co-infection by bacteria that commonly colonise the human upper respiratory tract. The usual causative infecting agent is *S. pneumoniae.*4 It can lead to significant morbidity and mortality; there being 9 to 18 such cases per 1000 adults with invasive pneumonia.5

Influenza B with co-infections is rarely fatal. To the best of our knowledge, only one such case report of severe co-infection with influenza B and bacteria exists in the literature. Three women who were previously healthy had septic shock due to co-infection; one of whom died. All three patients were relatively young (39, 27, and 61 years).2

Pathophysiologically, such severe co-infection is due to several mechanisms. Thus, in animal models the influenza virus can damage respiratory tract epithelium and facilitate bacterial infection.6,7 The virus can also alter and/or impair ciliary function of the epithelium.8 In addition, it can change host immunity and inflammatory responses by impairing bacterial clearance or via the inflammatory cascade.9,10

The virulence of the pneumococcus is mainly due to the polysaccharide capsule, which provides protection from phagocytosis.11 Pneumococci are classified into serotypes according to the structure

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**TABLE.** Clinical details and other characteristics of the four patients*

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Patient 1</th>
<th>Patient 2</th>
<th>Patient 3</th>
<th>Patient 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Onset of symptoms before admission (days)</td>
<td>4</td>
<td>3</td>
<td>3</td>
<td>7</td>
</tr>
<tr>
<td>Days of hospitalisation</td>
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<td>50</td>
<td>2</td>
<td>4</td>
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<td>Admission to ICU on day</td>
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<td>1</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>Days of ICU stay</td>
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<td>40</td>
<td>1</td>
<td>4</td>
</tr>
<tr>
<td>Age (years)</td>
<td>37</td>
<td>34</td>
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<td>60</td>
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<tr>
<td>Septic shock</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
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<tr>
<td>Acute kidney injury</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
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</table>

**Microbiology**

<table>
<thead>
<tr>
<th>Initially isolated bacteria</th>
<th><em>Streptococcus pneumoniae</em> serotype 3</th>
<th><em>Streptococcus pyogenes</em></th>
<th><em>Streptococcus pneumoniae</em> serotype 3</th>
<th><em>Streptococcus pneumoniae</em> serotype 3</th>
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</thead>
<tbody>
<tr>
<td>Influenza A IF†</td>
<td>-ve</td>
<td>-ve</td>
<td>-ve</td>
<td>-ve</td>
</tr>
<tr>
<td>Influenza B IF†</td>
<td>-ve</td>
<td>-ve</td>
<td>+ve</td>
<td>-ve</td>
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<td>Influenza A RT-PCR</td>
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<td>-ve</td>
<td>-ve</td>
<td>NA</td>
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<tr>
<td>Influenza B RT-PCR</td>
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<td>+ve</td>
<td>+ve</td>
<td>NA</td>
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<tr>
<td>4-Fold rise of influenza B titre</td>
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<tr>
<td>Viral culture</td>
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<td>NA</td>
<td>NA</td>
<td>Influenza B</td>
</tr>
</tbody>
</table>

**Laboratory findings**

| Leukocyte count (x 10⁹ /L)                | 2.4                                  | 0.5                    | 0.5                                  | 0.4                                  |
| CRP (mg/L)                               | 274                                  | 219                    | NA                                   | 324                                  |
| Creatinine level (μmol/L)                | 177                                  | 230                    | 335                                  | 110                                  |
| Death on day                             | NA                                   | NA                     | 2                                    | 4                                    |

* ICU denotes intensive care unit, IF immunofluorescence, RT-PCR reverse transcription-polymerase chain reaction, CRP C-reactive protein, NA not applicable, VV-ECMO venovenous extracorporeal membrane oxygenation, +ve positive, and -ve negative
† The IF kit was Light Diagnostics DFA Kit for respiratory syncytial virus, flu A, B, parainfluenza 1, 2, 3, and adenovirus
of the capsule. Hitherto, 90 different serotypes have been described. In our report, three of the patients harboured pneumococci, all of which were of serotype 3, which is the prevalent serotype in Hong Kong. It accounts for 12.3 to 19.8% pneumococcal infections in the age-group of 18 to 64 years, and 14.2 to 20.6% of such infections in the age-group of >65 years. Serotype 3 is covered by the pneumococcal conjugate vaccine 13, which has been available in the market since 2010.13

Our cohort of four patients developed severe streptococcal infection secondary to influenza B infection, whose characteristics and clinical course are described in the Table. Also, their admission chest X-ray films revealed bilateral involvement from the onset (Fig). Worldwide, the more severe co-infection is usually caused by influenza A strain H3N2. Influenza B co-infections cause a milder disease but rarely lead to death.13 Our patients were infected by influenza B, which may represent a new strain that warrants further investigation.

Acknowledgements

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References