CASE REPORT

Methicillin Susceptible Panton–Valentine Leukocidin Positive S. Aureus Pneumonia in a Child with Novel Influenza H1N1 Infection

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Summary
The first case in Latvia of Panton – Valentine leukocidin (PVL) positive methicillin susceptible Staphylococcus aureus (MSSA) pneumonia in an adolescent with novel influenza A H1N1 is described. A 15 year old boy was admitted to intensive care suffering from severe respiratory failure with bilateral necrotic pneumonia. The presence of influenza A H1N1 was confirmed by PCR. Invasive S. aureus was spa type t435 and Panton–Valentine leukocidin gene positive. He received therapy with ceftriaxone, oxacillin, clindamycin and oseltamivir phosphate and underwent two chest operations. He was discharged after 58 days in hospital.

Key words: H1N1, S. aureus, PVL; MSSA.

AIM OF THE DEMONSTRATION
Novel influenza type A (H1N1), or pandemic flu was first identified in April 2009 in Mexico and rapidly worldwide. On 11 June, 2009 the World Health Organization formally confirmed H1N1 influenza pandemic. The course of illness was mild or moderate in most cases and hospitalization due to severe influenza was required mainly for persons from the following risk groups: children under 2, pregnant women those with underlying chronic disease and the immuno-compromised (14).

In Latvia the novel pandemic influenza A/ H1N1 virus was first identified in June 2009; and new cases appeared sporadically until November when more than 50 new cases of pandemic flu were identified per week (15). During the period 20 – 26 November, 2009, very high activity was reported in Europe including Latvia, with children up to 15 years of age affected to unusually high degrees (16).

Uncomplicated human influenza virus infection causes transient tracheo–bronchitis due as the virus attaches to tracheal and bronchial epithelial cells. The main complication is extension of viral infection to the alveoli often complicated with bacterial infection resulting in severe pneumonia. Necrotizing S. aureus pneumonia has long been recognized, but the association with PVL has only recently been described. Numerous cases since have been reported worldwide. Panton – Valentine Leukocidin is a bicomponent pore–forming S. aureus exotoxin which mainly acts on neutrophils. PVL producing S. aureus may be either methicillin sensitive or resistant, however mainly associated with community acquired methicillin resistant S. aureus. Here, we report the first case in Latvia of methicillin susceptible Panton – Valentine leukocidin (PVL) positive S. aureus severe pneumonia in an adolescent with influenza A H1N1.

CASE REPORT
A 15 – year old boy was admitted to Daugavpils Regional Hospital on the evening 29 November 2009 with a 4 day history of low–grade to high fever, vomiting and a dry cough with haemoptysis and discomfort behind the sternum on the day of the hospitalization. A chest X–ray, performed on admission, showed total right sided pneumonia, the patients’ CRP (C– reactive protein) was 253,65 mg/l (N 0–7,9 mg/l) with other indicators as follows: HGB 15,8 g/dl, RBC – 5,56 x10⁶ (N 4,5–5,3), WBC – 7,5x10³ (N 4,5–13), PLT 160 x10³ (N 181–521). A nasopharyngeal swab was taken to detect respiratory viruses. Empiric antibacterial therapy with ceftriaxone and metronidazole was commenced. After a few hours the patient was moved to the intensive care unit and subsequently to the main children's hospital due to his progressive respiratory insufficiency. On the morning of November 30 the patient was transferred to the Childrens Clinical University Hospital (CCUH) in Riga. On arrival at CCUH, Riga, the patient had difficulty breathing and had signs of severe respiratory failure; he was sitting in an enforced position, had tachypnea (35 – 40 times per min.) with loud, groaning breathing and intercostal retractions and his blood pressure was raised (200/87 mmHg). Auscultation of the lungs showed unilateral dullness on the right side.
The chest X-ray showed multiple focal shadows on both sides of the lungs and unilateral intensive infiltration in the middle part of the right lung that suggested severe bilateral pneumonia (Pict. 1). Laboratory findings showed significant changes in blood gases – decreased \( pO_2 = 70.6 \) mmHg (N 71–104 mmHg), increased \( pCO_2 = 78.6 \) mmHg (N 32–46 mmHg), base excess was 6 mmol/l (N –5 –5), \( pH = 7.28 \) (N 7.37–7.45) and still elevated CRP – 261.65 mg/l, urea and creatinine levels were normal. His blood count at admission was normal except of “left shift” with 1% of metamyelocytes.

His history was unremarkable except miosis after acute respiratory infection in January, 2008 and recurrent faringitis in the summer of 2008. He was admitted to the intensive care unit and empiric oral antiviral therapy with oseltamivir phosphate (75mg twice daily) and intravenous antibacterial therapy with ceftriaxone and oxacillin in additional to antihypertensives and symptomatic therapy were commenced. One day later clindamycin was added.

Novel influenza A H1N1 infection was confirmed by PCR and \( S. aureus \) isolated from blood and pleural fluid on the day of admission were methicillin susceptible, Panton – Valentine leukocidin producing and were \( spa \) type \( t435 \). Antibacterial susceptibility was determined according to CLSI standards (M2–A9, M100–S16). The \( lukSF–PV \) genes were detected by PCR (4). Chromatograms of the \( spa \) sequences were analysed by Ridom StaphType software (Ridom GmbH).

Blood analyses that were taken two days later showed elevated inflammatory markers interleukin 6 (IL6) was 172 pg/ml (N <10 pg/ml) and calcitonin prohormone procalcitonin (PCT) level 2– 10 ng/ml (N< 0.5 ng/ml). Eleven hours after admission due to increasing respiratory insufficiency mechanical pulmonal ventilation was started and continued for 15 days. The general condition of the patient remained severe for more than five days.

On the 16\( ^{th} \) day the boy underwent operative therapy with a right side thoracotomy and resection of \( S4, S5 \) of the right lung (Pict.2) because of the severe condition due to pneumothorax and empyema. Further investigations of postoperative material revealed necrosis and inflammation of lung tissues. After the operation his general condition improved and it was decided to continue conservative therapy with antibiotics, but due to a post operative fistula of the right lung, the surgery was repeated after 3 weeks and the fistula was closed. The patient underwent repetitive bronchoscopies and antibacterial therapy with ceftriaxone (14 days), oxacillin (14 days), clindamycin (21 days) and oseltamivir phosphate (5 days). With this treatment blood cultures became negative on the 14\( ^{th} \) day of hospitalization. His general condition improved and after 58 days in hospital the patient was discharged.

**DISCUSSION**

Bacterial infection with *Staphylococcus aureus* is a known cause of severe illness often occurring after, and complicating, viral respiratory infection (9, 17). *In vitro* \( S. aureus \) will adhere mainly to poorly differentiated airway epithelial cells, confirming its tropism for injured and remodelled airway epithelium (7).

Panton – Valentine leukotoxin (PVL) is a pore forming staphylococcal \( y \) toxin encoded by the \( lukSF–PV \) genes (10), and is associated with skin abscesses and necrotizing pneumonia (6). Pneumonia often arises from the blood born spread of organisms from infected tissues and can follow viral respiratory infections, especially influenza (8). From 2002–3 isolates it has been estimated that <2% of \( S. aureus \) in the UK were PVL positive, most were methicillin sensitive, with 65% of them associated with skin and soft tissue infections, 17% with pneumonia (3). Gillet et al (1) compared the clinical features of PVL positive and PVL negative pneumonias and found in contrast to PVL–negative pneumonia patients, those with PVL–\( S. aureus \) were younger and mostly immunocompetent. They presented with influenza like symptoms, high fever, tachycardia, tachypnoe, heamoptysis and bilateral infiltrates, and pleural effusion more often. Other case series confirm the characteristics and severity of PVL – positive infections. (7,4,5,11). The symptoms of the described patient were equal with the described features.

Combined empirical antibacterial therapy of wide spectrum antibiotics is used in life-threatening infections. We used ceftriaxone and oxacillin empirically to treat atypical pneumonia in addition to antiviral therapy with oselatamivir phosphate (12). Oselatamivir phosphate was started due to possibility of influenza infection and severe condition of patient (18). Due to a rapidly worsening general condition and changes in the chest X–ray which was similar to PVL caused pneumonia, clindamycin was added one day after the initial therapy. International guidelines not been published for the therapy of necrotising pneumonia caused by PVL positive \( S. aureus \), however some local guidelines do exist (19), and mostly recommend the use of protein synthesis inhibiting antibiotics including clindamycin – when guided by in vitro susceptibility results. In addition, several publications recommend the addition of clindamycin in the treatment of toxin producing Gr+ bacteria as it may reduce toxin production (13).

Besides conservative therapy, surgery was also used due to the patients severe condition after two weeks, treatment with broad spectrum antibiotics and intensive care therapy. There is controversy over the indications and best timings for surgery in cases of pulmonary necrosis, especially in children. In our case the operation was done successfully despite the postoperative fistula, after the lobectomy, which prolonged the patients stay in hospital.

PVL positive \( S. aureus \) with \( spa \) type \( t435 \) are mostly methicillin susceptible and are spread in Latvia with sporadic cases in Poland, Austria, Romania and Hungary (2). In Latvia PVL positive \( S. aureus \) with \( spa \) type \( t435 \) is spread among children with purulent skin and soft tissues infections (Cupane, in preparation).

Our described case exposes that PVL – positive \( S. aureus \) with \( spa \) type \( t435 \) can complicate influenza in otherwise healthy children, with rapid progression to
severe pneumonia that needs complicated and long management of the illness.

Conflict of interest: None

REFERENCES


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Fig. 1. Chest X-ray on admission with multiple focal shadows on both sides of the lungs and unilateral intensive infiltration in the middle part of the right lung that suggested the severe bilateral pneumonia.

Fig. 2. Chest X-ray when the patient was discharged after resection of the middle part of the right lung with clinical improvement.