Novel complication of an emerging disease: Invasive *Klebsiella pneumoniae* liver abscess syndrome as a cause of acute respiratory distress syndrome

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Abstract

*Klebsiella pneumoniae* is an increasingly recognized cause of a unique invasive syndrome manifesting as pyogenic liver abscess and hematogenous extrahepatic dissemination to a variety of sites, including the lung. Originally described only in Asia, this entity has now been reported across continents and ethnicities. Intrathoracic complications of invasive *Klebsiella pneumoniae* liver abscess syndrome (IKPLAS) have been characterized sporadically but have not been the subject of a systematic investigation. Review of the English-language literature yields no reports of the acute respiratory distress syndrome as a consequence of IKPLAS. Herein we describe what, to our knowledge, is the first such occurrence.

Case Report

A 38-year-old woman originally from Mexico with a past medical history of type 2 diabetes mellitus (DM) treated with a sulfonylurea presented to a New York City hospital with abdominal pain. She also reported polyuria and polydipsia as well as nausea and vomiting. Her initial vital signs were as follows: blood pressure 93/53 mmHg, pulse 119 beats/min, respiratory rate 28 breaths/min, oxygen saturation 99% while receiving oxygen at 2L/min via nasal cannula, and a temperature of 97.8F. Physical examination revealed Kussmaul breathing, normal cardiopulmonary auscultation, and a soft, non-tender abdomen. She was fully alert and oriented. Initial laboratory evaluation revealed a blood pH of 6.98 with a serum bicarbonate of 5mmol/L (reference range 23-32). The serum anion gap was 22 with a normal serum lactate. The serum glucose level was 506mg/dL, and the urine output was positive for ketones. Urine pregnancy testing was negative. Her initial portable chest radiograph (CXR) showed subtle prominence of the interstitial markings and multiple ill-defined nodular opacities (Figure 1A). In the Emergency Department (ED), the patient received 4 liters of intravenous crystalloid, and an insulin infusion was started. Blood cultures were collected, and she was admitted to the intensive care unit (ICU) for the management of diabetic ketoacidosis.

In the ICU, the patient became progressively more hypoxic and dyspneic despite escalation to a non-rebreather mask. Her temperature rose to as high as 104F, and her level of consciousness declined to obtundation. Diffuse crackles were now heard on lung auscultation. She was endotracheally intubated for hypoxic respiratory failure and placed on volume assist-control ventilation. Repeat CXR showed diffuse bilateral opacities (Figure 1B). While receiving a fraction of inspired oxygen (FiO2) of 70%, the patient’s PaO2 was measured at 59 mmHg for a PaO2/FiO2 (P/F) ratio of 84. Lung-protective ventilation for the diagnosis of ARDS was initiated by reducing the tidal volume to 350ml and increasing the positive end-expiratory pressure (PEEP) to 7.5 cmH2O. Vancomycin and piperacillin/tazobactam were started empirically. Soon thereafter, blood cultures drawn in the ED were reported as growing gram negative bacilli, so piperacillin/tazobactam was switched to imipenem. The patient’s hemodynamics deteriorated rapidly to the point of requiring high-dose norepinephrine infusion along with vasopressin to maintain a mean arterial pressure >65 mmHg. By ICU day #3, her FiO2 requirement had risen to 100%, and the PEEP had been increased to 15 cmH2O. Together with inhaled nitric oxide therapy, these settings were sufficient to maintain an oxygen saturation of >90%. At that time, the gram negative bacilli growing in blood culture were identified as pan-sensitive *Klebsiella pneumoniae*. Antibiotic coverage was narrowed to ceftriaxone. Abdominal ultrasonography performed in search of the source of her bacteremia revealed a complex avascular hepatic mass-like lesion with both cystic and solid components measuring 7.7x5.1x8.0 cm (Figure 1C). Based on the suspicion of liver abscess, interventional radiology was called for bedside placement of a drainage catheter. During the procedure, the patient developed pulsless electrical activity (PEA) with return of spontaneous circulation following 5 minutes of cardiopulmonary resuscitation. The insertion was aborted and therefore abscess specimens could not be sent for culture.

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Thereafter, PEA recurred multiple times, so, after consultation with the patient’s relatives, resuscitative efforts were ultimately aborted, and the patient expired. Her family granted permission for an autopsy.

The autopsy confirmed the presence of a liver abscess with extensive necrosis. Lung sections demonstrated diffuse abscess formation completely obliterating alveolar architecture over a large territory (Figure 2A). Direct examination of the tissue revealed no organisms. In addition, many of the preserved alveoli contained hyaline membranes, the histological hallmark of diffuse alveolar damage (Figure 2B). The post-mortem diagnosis was thus invasive KLA syndrome leading to diffuse lung abscess formation complicated by ARDS.

**Discussion**

Historically, pyogenic liver abscess has represented a localized form of infection caused most commonly by *Escherichia coli* and afflicting patients with underlying hepatobiliary pathology. Its normally localized nature is perhaps the reason why the literature contains no reports of ARDS as a direct sequela of liver abscess. The landscape of this disease has changed over the last several decades, beginning in the 1980s with reports from Taiwan documenting patients with liver abscess caused by *K. pneumoniae* and complicated by bacteremia and dissemination, specifically endophthalmitis. Subsequently, a number of other Asian countries - a region in which fecal *Klebsiella* carriage rates are remarkably high - reported encounters with the so-called IKPLAS and its extrahepatic manifestations, including hematogenous lung seeding. Early on, DM was recognized to be a significant risk factor for this emerging infection. It was also noted that the majority of IKPLAS cases were attributable to the hypervirulent capsular subtypes of *K. pneumoniae* known as K1 and K2, which are especially prevalent in Asian countries. These strains are known for their resistance to phagocytic clearance by macrophages, particularly if they also exhibit the characteristic mucoviscous phenotype conferred by the *rmpA* gene and identified by positivity of the so-called string test (Figure 3). The impairment in phagocytic function typical of the diabetic milieu has been shown to further promote the evasiveness of K1/K2 serotypes and thereby their propensity for entry into the bloodstream and distant spread. In recent decades, experience with IKPLAS has been published by investigators from Europe, the Americas, and Australia describing cases in both Asian immigrants as well as in the native population. Of note, a 2008 series from our institution found a comparable incidence of IKPLAS between Asian-Americans and Hispanic-Americans such as the patient in this report (12 or 60% vs 8 or 40%, respectively). Clinically, fever is a near-universal initial sign, but presentations
are typically non-specific, and development of metastatic infection frequently provides a very delayed clue. Right upper quadrant tenderness may be absent. Klebsiella bacteremia is detected in about 50% of cases and is often what prompts a search for liver abscesses. CT and ultrasound are the preferred imaging modalities for that purpose.

The literature on the scope of intrathoracic manifestations of disseminated KLA is still in evolution; there are no English-language publications comprehensively analyzing chest imaging in this disease. Data on such manifestations are currently limited to case reports and series, studies of KLA in general that report lung findings, as well as descriptions of septic pulmonary emboli (SPE) caused by this infection. Some groups, though not all, have found the lung to be the most common site of extrahepatic seeding, ranging from 16.3% to 43.4% of cases. Among the specific types of intrathoracic complications, septic embolization diagnosed radiologically is the best described. Consolidations, mass-like lesions, interstitial opacities, and pleural effusions, including empyema, have also been reported.

Review of our institutional experience indicates that pleural effusions may actually be the most frequent radiological finding in the chest of invasive KLA patients (40%), followed by consolidations at 20% and masses/nodules at 7.5% (unpublished data). These numbers echo those previously published by a Taiwanese group. No reports known to us have heretofore linked IKPLAS to the development of ARDS.

As is the case with other distant sites of infection that can complicate KLA, we posit that our patient incurred a heavy and progressive burden of hematogenous lung seeding by K. pneumoniae, leading to what is typically described as pulmonary involvement is analogous to septic embolism and is to be distinguished from the typical endobronchial acquisition of conventional pneumonia caused by Klebsiella spp. The septic embolic insult to the lung was so severe that it resulted in the development of catastrophic lung injury in the form of ARDS.

**Conclusions**

In summary, IKPLAS is increasingly being recognized across the globe as a potentially morbid infection that carries a mortality rate of 4-11%, but one that can be effectively treated with prompt drainage and appropriate antibiotics. It is unique among causes of liver abscess in its propensity for hematogenous dissemination to other organs, including the lung. The extent of pulmonary involvement can be sufficient to cause respiratory failure in otherwise normal hosts. Based on our report, clinicians should add IKPLAS to the list of extrapulmonary infections capable of seeding the lungs and resulting in ARDS.

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