Relapsing Klebsiella pneumoniae liver abscesses in a non-asian diabetic man

Abcessos hepáticos recidivantes provocados por Klebsiella pneumoniae num homem diabético não asiático

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ABSTRACT

The Klebsiella pneumoniae invasive syndrome, first described in Asia, is being reported in other parts of the world. It causes liver abcsesses, with or without extrahepatic lesions. Diabetes mellitus is the most common hosts' underlying condition. It's frequent among asian people, even outside Asia, appearing to exist genetic factors, not yet established, increasing the risk. We present a case of a 55-year-old portuguese white diabetic man, who had a previous hospital admittion due to Klebsiella penumoniae pneumonia and a two-week latter diagnosis of liver abcsess. 4 years latter he was readmitted with nausea, chills, fever and myalgias. He had elevated inflammatory markers and the CT-scan showed 2 liver abscesses. Klebsiella pneumoniae was isolated in blood and antibiotics were given with complete clinical and imaging resolution. We present this case of relapsing Klebsiella pneumonia liver abscesses in a non-asian man emphasizing the growing incidence of this condition in Europe.

Keywords: Liver abscess; Klebsiella pneumonia; Klebsiella penumoniae invasive syndrome

RESUMO

O síndrome invasivo provocado por *Klebsiella pneumoniae*, descrito pela primeira vez na Ásia, tem vindo a ser reportado em outras partes do mundo. Provoca abcessos hepáticos, com ou sem lesões extra-hepáticas associadas. A diabetes mellitus é a condição predisponente do hospedeiro mais comum. É frequente nos indivíduos asiáticos, mesmo fora da Ásia, parecendo existir factores genéticos, ainda não estabelecidos, que aumentam o risco da infecção. Apresentamos o caso de um homem português de 55 anos, diabético que tinha uma admissão prévia no

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hospital por pneumonia provocada por *Klebsiella pneumoniae*, com diagnóstico de abcesso hepático 2 semanas depois. 4 anos mais tarde, foi readmitido com um quadro clínico de nauseas, calafrios, febre e mialgias. Apresentava parâmetros inflamatórios elevados e a tomografia computorizada (TC) abdominal revelou 2 abcessos. Foi isolada *Klebsiella pneumoniae* em hemoculturas e foram administrados antibióticos com completa resolução clínica e imagiológica. Apresentamos este caso de abcessos hepáticos recidivantes por *Klebsiella pneumoniae* num homem não asiático, enfatizando a incidência crescente desta entidade na Europa.

Descritores: Klebsiella pneumoniae; Abcessos hepáticos; Diabéticos; Relatos de casos

INTRODUCTION

Klebsiella pneumoniae is a Gram negative bacilli that usually causes respiratory and urinary tract infections. Since the 1980's a new invasive syndrome associated with this agent has been described in Asia, causing liver abscess^(1,2) with extrahepatic septic metastases, resulting from haematogenic dissemination^(3,4,5) (endophthalmitis, meningitis, necroting fasciitis) which can be life-threatenig.

The K.pneumoniae invasive syndrome was recently described outside Asia, namely in the United States of America (USA), ^(6,7) and in many countries in Europe ⁽⁸⁻¹⁵⁾. Yet, there is a predominance of asian people involvement, even outside Asia. That remains unclear, but in 2002 a study sugested that some genotypic and phenotypic characteristics identificated in micoorganism from Asian countries are different from the bacteria identified outside Asia⁽¹⁶⁾. More recently, however, there are descriptions of *K.pneumoniae* liver abscess in non-asian patients⁽¹⁷⁾.

That syndrome is usually caused by two specific serotypes of *K. pneumoniae*: K1 and K2⁽¹⁷⁾, which have a hypermucoviscous phenotype confering more virulent properties.

We describe a case of relapsing liver abscess on a non-asian diabetic man.

CASE PRESENTATION

We present a case of a 55-year-old portuguese white man with previous background of type 2 diabetes, with peripheral arterial disease and retinopathy, elevated blood pressure, obesity, dyslipidemia and obstructive sleep apnea syndrome. His

medication was Insuline Lispro, Furosemide, Spirinolactone, Pentoxyphylline, Rosuvastatine, Clopidogrel and Lansoprazole. His alcohol intake was 120mg per day and he is an ex-smoker; no other drug use, travelling history or animal contact was reported.

He was admitted on September 2009 with pneumonia, with *K.pneumoniae* isolation from blood cultures and spuctum. He received antibiotic therapy for 10 days and was discharged. Two weeks latter he was readmitted with an hepatic abscess and he was submitted to surgical drainage. The same micoorganism was isolated on cultures of aspirated pus and he received another course of antibiotics. A 4-month time abdominal CT-Scan was performed with complete resolution of the liver abcsess.

On July 2013 the patient presented to the emergency department with a two-week history of nausea, chills, fever and myalgias. On physical exam he was vigil, oriented, blood pressure 135/78mmHg, heart rate 112bpm, febrile (38°C), pale, dehydrated, no enlarged lymph nodes, normal heart sounds and chest examinaton; his abdomen was distended, painfull, but no peritoneal reaction; the bowel sounds were normal; no hepatosplenomegaly or palpable masses. The physical exam was otherwise irrelevant.

Laboratory tests showed normocytic normochromic anemia (9,9g/dl), normal leucocyte count (9600/mL) with relative neutrophilia (81%), elevated liver enzymes (aspartate aminotransferase -129UI, alanine aminotransferase - 162UI, gamma glutamyl transferase - 416UI) and elevated inflammatory markers (C-Reactive Protein - 23.5mg/dl; Erytrocyte sedimentation rate 78mm/hr); his urea was slightly elevated, with a normal creatinine.

His urine analysis revealed an elevated leucocyte and erythrocyte count.

A chest radiography was performed and it was normal. The abdominal ultrasonography showed an heterogeneous hepatomegaly, with no focal lesions (Figure 1). The renal ultrasonography revealed enlarged kidneys without obstructive lesions.



Figure 1. Abdominal ultrasonography. The abdominal ultrasonography showed a heterogeneous hepatomegaly, with no focal lesions.

A urinary tract infection was assumed and ceftriaxone 2g ev id was started.

Besides antibiotic therapy, the patient mantained persistent fever and high inflammatory markers, and an abdominal CT-scan was ordered. Imaging findings were consistent with liver abcsesses in IV and VIII segments (7 and 3.5 cm) (Figure 2).

A *K. pneumoniae* was isolated from blood cultures and the urine culture was negative.

Due to the abcsesses localization it was decided to have a conservative approach and parental Piperacillin-Tazobactam and Metronidazol was given. The patient was afebrile 3 days after, with progressive clinical improvement, and was discharged from the hospital at 19th day since admittion. He completed a 4 week course of antibiotherapy. The follow up abdominal CT-scans showed improvement at two-month (Figure 3) time and a normal liver apperance on 6th month (Figure 4).

DISCUSSION

K.pneumoniae invasive syndrome was recently described as an infection of the liver caused by a specific serotype of that microorganism. It can have extrahepatic complications and be very severe.

According to the definition of *K.pneumoniae* invasive syndrome proposed by L Kristopher et al in 2012,⁽¹⁶⁾ the patient that we presented had a clinical probable invasive syndrome at the time of current admition in the hospital

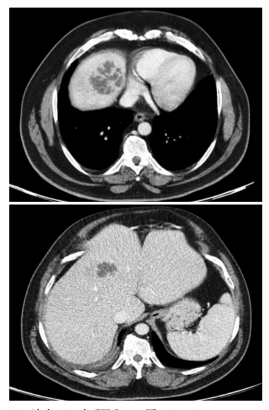


Figure 2. Abdominal CT-Scan. The imaging were consistent with liver abcsesses in IV and VIII segments (7 and 3.5cm).



Figure 3. Abdominal CT-Scan 2 months after treatment. CT-scan showed marked improvement in the abcesses.



Figure 4. CT-scan revealed complete resolution of the abcesses.

as he had a *K.pneumoniae* liver abscess as the sole presenting clinical manifestation. When the liver abscess have extrahepatic complications, especially central nervous system involvement, necrotising fasciitis or endophthalmitis, the syndrome is classified as definite. The same authors also proposed a microbiological definition of the syndrome: it is definite if there is an identification of the K1 or K2 serotypes and it is probable when there is a hypermucoviscous phenotype identificated by the string test.

This syndrome was first described in southest asian countries^(1,2), however there are 38 cases reported in the USA. 50% of the affected patients in USA are asian-descendents. The other half have an other ethnic origin⁽¹⁷⁾.

The most common underlying condition on patients affected by this disorder is diabetes mellitus and it seems to be an association between uncontrolled hyperglicemia and metastatic lesions^(18,19). It hasn't been identified specific human genes predesposing to the infection, but they seem to exist once the disease is more common in Asian people⁽¹⁹⁾. Our patient belongs

to the diabetic risk group and he had a poor glycemic control with macro and mycrovascular complications. Hypothetically, he may have a personal or genetic predesposition once he already had the same clinical presentation four years ago and, at the time, the liver abcsess was resolved.

We present this case report because its rarity outside asian countries as well as the particularity of a relapse of the liver abscess in the same patient four years after the first episode.

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