A Case Report and Review of the Literature: Infectious Aneurysm Formation in the Pulmonary Arteries—A Rare but Perilous Sequela of Persisting Infection With Klebsiella pneumoniae

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Septic aneurysms of the pulmonary artery are rare conditions, with few cases having been reported worldwide. They are assumed to result from septic emboli that cause a local inflammatory reaction of the arterial wall, ultimately leading to degenerative changes. We report the case of a 63-year-old female patient presenting with Klebsiella pneumoniae urosepsis and first diagnosis of diabetes mellitus, who developed a life-threatening infectious pulmonary artery aneurysm secondary to bacteremia with Klebsiella pneumoniae. The patient required a lobectomy due to pulmonary hemorrhage. We review the clinical hallmarks of Klebsiella pneumoniae related septic pulmonary embolic disease and summarize currently known risk factors for the development of infectious aneurysmatic disease including diabetes mellitus and other states of immunosuppression. The featured case aims to increase the awareness for this seldom but life-threatening complication of infectious diseases such as Klebsiella pneumoniae urosepsis.

Keywords: infectious aneurysm, haemoptoe, urosepsis, Klebsiella pneumoniae, pulmonary artery

INTRODUCTION

The term pulmonary infectious (former mycotic) aneurysm describes an infection-related ectasia of the pulmonary artery due to prolonged bacterial destruction of the arterial wall (Majeed and Ahmad, 2021). The term “mycotic” is actually a historical misnomer, first introduced by Sir William Osler in 1885 (Osler, 1885), as mostly bacterial and not mycotic pathogens trigger infectious aneurysms. Aneurysm formation of the pulmonary arteries is a rare event with a high fatality rate (Barter et al., 1988). It may result from bacteremia or septic emboli that cause degenerative changes in the arterial wall (Brown et al., 1984; Macbeth et al., 1984; Qureshi et al., 1999; Oderich et al., 2001). In this regard, risk factors and biomarkers for aneurysm formation under persistent infection despite systemic antibiotic treatment are missing, leaving the patient...
at risk for undetected disease progression and critical pulmonary complications. The following case highlights these mentioned difficulties and gives brief overview of the literature on *Klebsiella pneumoniae* (KP) related septic pulmonary disease and its role in aneurysm formation.

**CASE-REPORT**

A 63-year-old Caucasian female was admitted to the intermediate care unit with a first diagnosis of diabetes mellitus and an infection-associated metabolic (keto)-acidosis (pH 7.16). Initially, the patient was hypovolemic, disoriented, tachypneic, hypoxemic, complained about epigastric pains, and loss of vision on the left eye. The serum glucose and the inflammation markers were markedly elevated (Table 1). The initial management included fluid resuscitation, insulin therapy and an empiric antibiotic treatment with piperacillin/tazobactam after blood- and urine-culture sampling. Due to persisting hypoxemia high-flow oxygen supplementation therapy was necessary and a CT was performed. This revealed signs of pyelonephritis, a renal vein thrombosis and bilateral subsegmental lung emboli co-localizing with multiple round and wedge-shaped infiltrates and a large consolidation in the right lower lobe (Figures 1, 2).

As the majority of the multifocal nodules either depicted an “afferent-vessel sign”, characterized by close topographic proximity to the arterial flow area, septic pulmonary embolism (SPE) was suspected. A duplex sonography revealed a marginal circumferential thrombosis of the right common femoral vein, and a therapeutic anticoagulation regimen with tinzaparin (16,000IE) was initiated.

During ophthalmologic examination, the left amaurotic eye presented markedly swollen, while cranial magnet-resonance imaging with orbital cross-sectional imaging detected a choroidal effusion in the left bulbus, leading to the diagnosis of endogenous endophthalmitis.

Blood and urine culture tested positive for pan sensible KP (Table 1), and a diagnosis of pyelonephritis with consecutive urosepsis was made. The anti-infective regimen was subsequently de-escalated to ceftazidime in accordance with (KP) endogenous endophthalmitis.

**TABLE 1 | Blood parameters at admission.**

| Parameter                  | Value     |
|----------------------------|-----------|
| pH (ven.)                  | 7.16      |
| pCO2 (ven.)                | 23 mmHg   |
| HCO3- (ven.)               | 8 mmol/L  |
| Base Excess                | – 19.8 mmol/L |
| Serum-Glucose              | 33 mmol/L |
| Serum-Hba1c                | 14.2%     |
| Plasma-ketone bodies       | 557.8 µmol/L |
| C-Peptide                  | 3.28 ng/ml |
| Insulin                    | 6 mL/ml   |
| Anti-GADA                  | Negative  |
| Anti-IA2                   | Negative  |
| Anti-Inulin                | Negative  |
| Serum-CRP                  | 387.7 mg/L |
| Serum-PCT                  | 3.9 µg/L  |
| BSS 1 h                    | 77 mm     |
| BSS 2 h                    | 81 mm     |
| WBC                        | 27,500/µL |
| Neutrophils                | 87.5%     |
| Eosinophils                | 0%        |
| Basophils                  | 0.3%      |
| Monocytes                  | 4.9%      |
| Lymphocytes                | 4.9%      |
| Urin-pH                     | 5.0       |
| Urin-Glucose               | ++ +      |
| Urin-Ketonbodies           | ++ +      |
| Anti-neutrophilicantibodies| 1:80 speckled |

*KlebsiellaPneumoniae*

| Time to positivity | 40 m, 11 h, 15 min |
|--------------------|--------------------|
| Antibiotic         | Ampicillin [R] ≤ 2.0 |
|                    | Ampicillin-Subbactam [S] ≤ 2.0 |
|                    | Piperacillin-Tazobactam [S] ≤ 4.0 |
|                    | Cefuroxin [I] ≤ 1.0 |
|                    | Cefuroxin-Axetil [S] ≤ 1.0 |
|                    | Cefpodoxim [S] ≤ 0.25 |
|                    | Ceftriaxon [S]      |
|                    | Cefotaxim [S] ≤ 1.0 |
|                    | Cefazidim [S] ≤ 1.0 |
|                    | Gentamicin [S] ≤ 1.0 |
|                    | Levofloxacin [S]    |
|                    | Ciprofloxacin [S] ≤ 0.25 |
|                    | Moxifloxacin [S] ≤ 0.25 |
|                    | Meropenem [S] ≤ 0.25 |
|                    | Ertapenem [S] ≤ 0.5 |
|                    | Cotrimoxazol [S] ≤ 20.0 |

With Hba1c, hemoglobin A1c, Anti-GADA, glutamic acid decarboxylase antibodies, Anti-IA2S, tyrosine phosphatase antibodies, CRP, C-reactive protein, PCT, Procalcitonin, BSS, Blood sedimentation speed, WBC, White blood cell count, ENA, Extractable nuclear antigen and ANCA, Anti-neutrophil cytoplasmatic antibody. KP species in urine and blood culture were determined via mass spectrometry under employment of a matrix assisted laser desorption ionization time-of-flight (MALDI-TOF) system (VITEK®, bioMérieux Inc., Nütingen, Germany) and comparison with the companies database (Myla®, bioMérieux Inc., Nütingen, Germany). Minimal inhibitory concentration was assessed with VITEK® 2 (bioMérieux Inc., Nütingen, Germany) following manufacturer’s instructions, to empirically cover pathogenic resistance mechanisms, that may have manifested under the prolonged prior antibiotic regimen, in order to prevent pathogen related aneurysmal rupture. Surgical treatment as minimal invasive video-assisted thoracoscopic lobectomy of the right lower lobe including the...
infectious aneurysm was performed. Histopathological work-up depicted neither signs of malignancy of autoimmune disease nor direct pathogen detection. However, necrotizing and fibrinous purulent, partly gangrenous thromboembolic inflammation was noted (Figure 4). The patient made an uneventful recovery from the surgical procedure. On day 31 after admission the patient was respiratory stable, exhibited no reminiscent clinical or laboratory signs of infection and was discharged to a diabetic rehabilitation center.

**DISCUSSION**

Within the rare occurrences of infectious aneurysms, involvement of the pulmonary arteries is even more rarely seen (Papaioannou et al., 2014; Benhassen et al., 2018), while predictors for manifestation of infectious aneurysms following septic pulmonary embolism are absent. To date, only few cases of this disease have been described, and the reported fatality rate is higher than 50% (Bartter et al., 1988), especially if pulmonary hemorrhage occurs (Papaioannou et al., 2014). Furthermore, Gram-negative infections (like in the case of KP sepsis) were associated with an increased risk for rupture of the aneurysmatic wall (Jarrett et al., 1975).

*Klebsiella pneumoniae* is a member of the *Enterobacteriaceae* family and is a common pathogen in lower urinary tract, soft tissue and pulmonary infections. Thereby, pulmonary pathogenicity of virulent KP species has been recently emphasized by increasing isolation frequencies in respiratory samples over the past decade (Mendez et al., 2020; Braeken et al., 2021). Typically, if rarely, hyper virulent, hypermucoviscous KP species have been interlinked with systemic infection and sequential involvement of multiple organ sites (Fang et al., 2004; Chang et al., 2021), which is likely to have occurred in our patient despite absent genetic in depth characterization of the detected KP isolate. Indeed, pulmonary manifestation of systemic KP infection or more particular septic pulmonary embolism seems uncommon (Chang et al., 2021).
FIGURE 2 | Infectious aneurysm of the pulmonary artery. The initial computed tomography was performed on the day of admission (A) with solely consolidation in the right basal lobe and an accentuated right lower lobe pulmonary artery. Over the course of 20 days an infectious pulmonary arterial aneurysm manifested with a diameter >4 cm (B), coinciding with clinical signs of hemoptysis.

In 2005 Cook and colleagues defined the occurrence of septic pulmonary embolism by the presence of four criteria: (1) uni- or multifocal pulmonary infiltrates with (2) an active extra pulmonary infection site acting as embolic source under (3) exclusion of any malignant or non-malignant differential diagnosis. Finally, resolution of the pulmonary infiltrate under adequate antibiotic treatment is the 4th mandatory feature before the diagnosis can be secured (Cook et al., 2005).

Thus, firstly multiple bilateral nodular opacities or wedge-shaped infiltrates in close proximity to the pulmonary arterial
circulation, depicting the SPE-suggestive “feeding vessel sign”, were present on the patient’s chest CT scan (Lee et al., 2007; Ye et al., 2014; Chang et al., 2016). Although KP related SPE presents with a broad spectrum of radiologic signs including nodules, cavities, ground glass opacities or pleural effusions, the “feeding vessel sign” [11/14 patients, (Chou et al., 2015)] and wedge-shaped peripheral opacities [12/33, (Zhang et al., 2021)] were most frequently noticed in two distinct retrospective case-series.

Secondly, a pyelonephritis that led to urosepsis was identified as the extrapulmonary infection site. KP related lower urinary tract infections are rather frequent events (Paczosa and Mecsas, 2016; Shakya et al., 2017), while permissive hyperglycemia and glucosuria as present in our patients, have been reported as main risk factors (Lee et al., 2016). Additionally, secondary endogenous endophthalmitis is often seen especially in patients of Asian descent with systemic KP infection (Yang et al., 2007; Hagiya et al., 2013; Chang et al., 2016). SPE as pulmonary complication in KP infection is rarely described in Western patients (Cook et al., 2005). However, a systematic review analyzing case reports and case series of patients with SPE, applying the Cook criteria, in patients >14 years of age between 1979 and 2012 in a global perspective, refers to KP as the causative pathogen in 7.3% (11/151 patients) of the reviewed cases (Ye et al., 2014).

Meanwhile, Yang and colleagues found diabetes mellitus to significantly increase the risk for septic metastases in a head-to-head comparison with a non-diabetic control group (Yang et al., 2007). Several case reports have been published wherein KP related SPE manifested in patients with diabetic metabolic decompensation (Zenda et al., 1995; Chang et al., 2015; Ojeda Gómez et al., 2019). A prospective single center cohort study performed by Lee and co-workers describes a significantly increased risk of KP-related SPE in diabetic patients with a Hba1c>9% [OR 5.66, 95%CI (2.01, 15.9)] (Lee et al., 2016). Furthermore, the authors infer that the hyperglycemia induced pathogenicity of KP is related to overexpression of critical genes for the synthesis and assembly of bacterial polysaccharide capsule. Also, neutrophilic phagocytic activity against KP was compromised in vitro following incubation of KP with glucose prior to exposition to patient derived neutrophilic granulocytes (Lee et al., 2016). Referring these findings back to the presented case, our patient showed typical septic manifestation sites of KP related infectious metastatic disease. Moreover, the pronounced
hyperglycemia in our patient is likely to have served as a major risk factor for septic dissemination (Rahimian et al., 2004; Keller et al., 2013).

Thirdly, possible differential diagnoses were ruled out by histopathology having shown no signs of malignancy or granulomatous lung disease.
Fourthly, antibiotic treatment led to a significant improvement of the clinical course, especially in terms of arterial oxygenation and lung function. On the contrary, iterated CT scanning showed no resolution of the initially observed bilateral infiltrates, as would be expected in the case of adequately treated SPE. Instead, our patient developed an aneurysm under anti-infected treatment in only a time period spanning 20 days (Figure 3). Of note, the patient had severe issues in terms of glycemic control in this time period, while diabetes was associated with the adverse outcome and death in patients with infective aortic aneurysms undergoing open surgery (Lau et al., 2015; Sörelius et al., 2019). Thereby, infectious aneurysms are believed to result either from septic thromboembolic occlusion of the pulmonary *vasa vasorum* of the pulmonary arteries from infective endocarditis or distinct septic thrombotic foci with concomitant invasive infection of the arterial wall. Thus, continuous recruitment of neutrophilic granulocytes and release of various proteases leads to progressive thinning of the arterial wall with resultant wall ectasia accompanied with a high risk of hemorrhage (Baddour et al., 2015; Habib et al., 2015; Majeed and Ahmad, 2021). A recent review on published cases with infectious aneurysms of pulmonary arteries between 1949 and 2018 described endocarditis and congenital heart defects as the main predisposing factors in almost half of the cases together (Benhassen et al., 2018). *Staphylococci* or *streptococci* were most frequently isolated, but also Gram-negative bacteria and various fungi have been occasionally reported (Jarrett et al., 1975; Müller et al., 2001; Ebisawa et al., 2018). However, no signs of endocarditis were present in our patient, while pulmonary infectious aneurysm have also been repetitively reported independent of present infectious endocarditis (Dransfield and Johnson, 2003; Benhassen et al., 2018; Alquichire-Luna et al., 2021).

To the best of our knowledge, only a single case-report on a KP-related infectious aneurysm has been reported by Alquichire-Luna and colleagues in a neutropenic patient with a history of acute myeloid leukemia. In contrast, our patient showed no state of immunosuppression besides decompensated type 2 diabetes. While the KP isolate in the referred case exhibited multiple antibiotic resistances (meropenem, piperacillin-tazobactam) predisposing an adverse clinical course, only pan sensible KP was identified in urine and blood (Table 1) from our patient. However, a recent meta-analysis of infection in patients with manifest type 2 diabetes found an increased risk of antibiotic-resistant infection with predominant involvement of the urinary tract \[OR = 2.42, 95\% CI = (1.83, 3.20)\] and the lung \[OR = 2.35, 95\% CI = (1.49, 3.69)\] (Carrillo-Larco et al., 2022), also predisposing these patients to an adverse clinical course. In our case, secondary formation of a subsegmental pulmonary arterial aneurysm was noted, which occurred most likely due to persistent sterile local inflammatory processes of the arterial wall. Conversely, indicators of systemic infection were tested negative after antibiotic treatment. Unfortunately, no general treatment recommendations on infectious aneurysms are present throughout the literature and in particular clear guidance on selection of the adequate antibiotic agent and treatment duration is missing (Baddour et al., 2015; Habib et al., 2015; Sörelius et al., 2020). Thus, the question after a superior preemptive antibiotic regime than the one administered in the case presented remains unanswered. Data only exists from a nationwide retrospective study from Sweden on treatment of infectious abdominal aortic aneurysms which suggested significant favorability in terms of postoperative mortality at 3 and 5 month of a post-operative antibiotic treatment regimen > 6 month in a multivariate analysis (Sörelius et al., 2016).

However, the applicability of this study for the case presented remains questionable, as patients were treated either with endovascular or open surgical vascular repair including implantation of foreign body material into pre-infected tissue. Likewise, favorability of either microsurgical resection (current case) or interventional management (Alquichire-Luna et al., 2021) of manifested aneurysms in the pulmonary vasculature remains elusive and may rely on the referring center.

The present case-report underscores the dynamic character of local SPE related infection bearing the perilous potential to evolve to an infectious aneurysm. Comorbid diabetes mellitus might serve as an additional risk factor setting the stage for persistent KP infection under antibiotic treatment. Thus, underlying diabetes was present in 62% of patients with KP related liver abscess and subsequent extra hepatic metastatic disease (i.e., pulmonary and ophthalmic involvement) (Han, 1995) and in 68% of patients with KP-related endophthalmitis in a retrospective case series (Yang et al., 2007). Therefore, careful diagnostic work-up is necessary in patients presenting with hemoptysis and persistent signs of systemic infection including CT based angiography. Accordingly, central pulmonary aneurysms can be easily misinterpreted as hilar masses on native imaging techniques, seducing the pulmonologists to an invasive biopsy approach with potential fatal outcome (Dransfield and Johnson, 2003).

**CONCLUSION**

*Klebsiella pneumoniae* may play a significant role in septic pulmonary embolic disease. However, metabolic decompensation of underlying diabetes mellitus, as often seen during infectious diseases, may serve as breeding ground for KP related septic embolisms including pulmonary involvement. CT based angiography should be performed quickly and prior to other invasive diagnostic procedures for avoiding iatrogenic aneurysmal rupture. However, identification of patients at risk for the development of aneurysmal disease resembles a major obstacle, as simple absence of current bacteremia not necessarily rules out persistent infectious or inflammatory reactions in already metastatically involved organs. Meanwhile, the overall manifestation rate following infectious endocarditis or simple bacteremia remains low. Hence, prospective studies to establish standardized guidelines are highly warranted for securing an early diagnosis and adequate treatment of the affected patients. However, such desirable tools are currently missing in the diagnostic quiver, which is why clinicians should be aware of pulmonary sequelae in patients with prolonged infectious disease with
or without hemoptysis. The immunosuppressive capacity of permissive diabetic metabolic decomposition may serve as a perilous pro-infectious catalyst that should not be underestimated.

**DATA AVAILABILITY STATEMENT**

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

**ETHICS STATEMENT**

Ethical review and approval was not required for the study on human participants in accordance with the local legislation and institutional requirements. Written informed consent for participation was not required for this study in accordance with the national legislation and the institutional requirements.

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