Complete staghorn calculus in polycystic kidney disease: infection is still the cause

Zhiguo Mao*, Jing Xu, Chaoyang Ye, Dongping Chen and Changlin Mei*

Abstract

Background: Kidney stones in patients with autosomal dominant polycystic kidney disease are common, regarded as the consequence of the combination of anatomic abnormality and metabolic risk factors. However, complete staghorn calculus is rare in polycystic kidney disease and predicts a gloomy prognosis of kidney. For general population, recent data showed metabolic factors were the dominant causes for staghorn calculus, but for polycystic kidney disease patients, the cause for staghorn calculus remained elusive.

Case presentation: We report a case of complete staghorn calculus in a polycystic kidney disease patient induced by repeatedly urinary tract infections. This 37-year-old autosomal dominant polycystic kidney disease female with positive family history was admitted in this hospital for repeatedly upper urinary tract infection for 3 years. CT scan revealed the existence of a complete staghorn calculus in her right kidney, while there was no kidney stone 3 years before, and the urinary stone component analysis showed the composition of calculus was magnesium ammonium phosphate.

Conclusion: UTI is an important complication for polycystic kidney disease and will facilitate the formation of staghorn calculi. As staghorn calculi are associated with kidney fibrosis and high long-term renal deterioration rate, prompt control of urinary tract infection in polycystic kidney disease patient will be beneficial in preventing staghorn calculus formation.

Keywords: Staghorn calculus, Polycystic kidney disease, Urinary tract infection

Background

The incidence of kidney stone in patients with autosomal dominant polycystic kidney disease (ADPKD) is approximately 5 to 10 folds of that in the general population [1-3]. The high incidence of nephrolithiasis is thought to be the consequence of the combination of anatomic abnormality and metabolic risk factors in ADPKD patients [4]. However, complete staghorn calculus in ADPKD is rare and the causes of staghorn calculus in ADPKD remains elusive. As existence of staghorn calculus predicts poor prognosis of kidneys [5,6], the causes for staghorn calculus in ADPKD are of interests of nephrologists.

Here we report a case of complete staghorn calculus in an ADPKD patient induced by repeatedly upper urinary tract infections.

Case presentation

A 37-year-old ADPKD female patient with positive family history was admitted in this hospital for repeatedly urinary tract infection for 3 years. Three years ago, the patient was admitted in community hospital for urinary tract irritation with right flank pain and fever for 2 days, and right upper urinary tract infection was considered. After admission, bilateral enlarged kidneys full of fluid-filled cysts were detected by type-B ultrasound examination, and there were no calculus or crystals in the kidneys detected at that time. ADPKD was diagnosed considering her family history. Since then, she suffered from urinary tract infection every 3–4 months, and urine cultures were negative. Symptoms usually were relieved within a week with antibiotics therapy, but the infections relapsed easily.

One week before admission, this patient’s urinary irritation relapsed with high fever. As oral antibiotics were not responsive, she was admitted in this hospital for further treatment. On admission, her temperature was 39.0°C, blood pressure 110/82 mmHg. Physical examination...
revealed the bilateral enlarged kidneys that were palpable, tenderness existed in right flank area. Full blood cell count showed white blood cell count $11.1 \times 10^9$/l, hemoglobin 12.3 g/dl and platelets $157 \times 10^9$/l. A serum biochemical profile showed her renal function was normal with serum creatinine 0.74 mg/dl. Liver function and electrolytes were unremarkable. Urinalysis revealed +3 leukocytes, +2 blood, and +1 albumin. Chest radiology and electrocardiography were reported to be normal. Abdominal CT scan showed enlarged cystic kidneys and a complete staghorn calculus in the right kidney with no obvious signs of urinary tract obstruction (Figure 1). Wide spectrum antibiotics was administered intravenously and the symptoms relieved in 3 days.

Percutaneous nephrolithotomy (PNL) was suggested, but the patient refused surgery as she felt well after the symptom relieved and worried about the surgical risks. To reduce the urinary tract infection and calculus formation, big volume of water intake was suggested to keep urinary output more than 2 L a day.

One month after discharge, the patient reported a “tiny stone” passed with urine, and the urinary stone component analysis showed the composition of calculus was magnesium ammonium phosphate. Considering the calculus component and the rapid formation of the staghorn calculus accompanied by the repeated upper urinary tract infections, the main cause of the staghorn calculus was determined as the repeated upper urinary tract infection in ADPKD.

**Conclusions**

The incidence of kidney calculus in ADPKD patients ranges from 8% to 36% [1-3], much higher than in general population, while ADPKD with staghorn calculus is rare. In our center data, there is only one case with complete staghorn kidney calculus among 412 regularly followed ADPKD patients (221 male and 191 female) in the past twelve years.

Calcium oxalate and/or uric acid are the most components of kidney stone in ADPKD, although the percentage of metabolic disturbances was not higher in ADPKD patients with kidney calculus than without [2]. Patients with a larger kidney volume predominant cyst size are at a higher risk of stone formation [2,4]. Numerous expanding kidney cysts distort the intrarenal calyceal system, which leads to urinary stasis and delayed washout of urinary crystals. Hypocitruria, aciduria and low urinary magnesium are common metabolic defects in ADPKD, and these factors facilitate the formation and aggregation urinary crystal.

There are no generally accepted causes for the formation of staghorn calculi in ADPKD. In general population, staghorn calculi were traditionally believed to be synonymous with infection stone, and secondary to of UTI [7]. However, recent data challenged this traditional opinion. In a retrospective analysis with 52 kidneys with complete staghorn calculi, 56% of the kidney stones were metabolic and 44% were infection stones [8]. The rising percentage of metabolic stones were also observed in other urinary stone cases [9,10].

In this case of ADPKD patient, the medical history and kidney stone component analysis supported that this case was a typical UTI induced complete staghorn calculus. This case showed in ADPKD patients, UTI was an important cause of staghorn calculus. As UTI is very common in ADPKD with 30%-50% patients experiencing an episode in their lifetime [11,12], and staghorn calculus was associated with kidney fibrosis [5] and high long term renal deterioration rate [6], prompt control of UTI in ADPKD patient will be beneficial in preventing staghorn calculus formation.

For the treatment of staghorn calculi, PNL was recommended as the first therapy for most patients by AUA guideline in 2005 [7], while shock wave lithotripsy (SWL) monotherapy and open surgery are only indicated on few conditions [13]. In ADPKD patients, the principles of
management of kidney stones are the same as in patients with normal kidneys, however PNL can be challenging because of the distortion of kidney anatomy by cysts compression and the cysts can come in the way of puncture and need to be aspirated before procedure [1,14-16]. As reports on surgical treatment for nephrolithiasis in ADPKD patients are limited, it is reasonable to choose suitable therapeutic modality based on experience and expertise of each center. After removal of kidney stones, risk factors evaluation and control should always be considered to reduce the frequency of recurrent stone disease in such patients.

Consent
Written informed consent was obtained from the patient for publication of this Case report and any accompanying images. A copy of the written consent is available for review by the Editor of this journal.

Abbreviations
ADPKD: Autosomal dominant polycystic kidney disease; UTI: Urinary tract infection.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
ZM, CY and JX treated the patients and collected the data, ZM and DC wrote the manuscript. CM designed the study and helped to draft the manuscript. All authors read and approved the final manuscript.

Acknowledgements
Z. Mao is a Fellow of International Society of Nephrology (ISN). This work was supported by Shanghai Higher Education Outstanding Young Teacher scientific Research Special Fund, Shanghai Municipal Education Commission; Outstanding Young Scholars project, Second Military Medical University and Medical Educational Reforming and Research Grant of Changzheng Hospital. This work was funded by the National Nature Science Fund of China (No. 81000281) and the Key Science and Technology Project of Shanghai Municipal Science and Technology Commission (No. 11431920800).

Received: 9 June 2013 Accepted: 30 July 2013
Published: 1 August 2013

References
1. Mufti UB, Nalagatla SK: Nephrolithiasis in autosomal dominant polycystic kidney disease. J Endoendr 2010, 24(10):1557–1561.
2. Nishiura JL, Neves RF, Eloi SR, Cintra SM, Ajzen SA, Heilberg IP: Evaluation of nephrolithiasis in autosomal dominant polycystic kidney disease patients. J Urol 2009, 4(4):838–844.
3. Gambaro G, Fabris A, Pulitana D, Lupo A: Lithiasis in cystic kidney disease and malformations of the urinary tract. Urol Res 2006, 34(2):102–107.
4. Torres VE, Wilson DM, Hattery RR, Segura JW: Renal stone disease in autosomal dominant polycystic kidney disease. Am J Kidney Dis 1993, 22(4):513–519.
5. Boonla C, Kriegstein K, Bovonpadungkitt S, Strutz F, Spittau B, Pedanond C, Tosukhowong P: Fibrosis and evidence for epithelial-mesenchymal transition in the kidneys of patients with staghorn calculus. BJU Int 2011, 108(6):1336–1345.
6. Teichman JM, Long RD, Hultberg JC: Long-term renal fate and prognosis after staghorn calculus management. J Urol 1995, 153(5):1403–1407.
7. Preminger GM, Assimos DG, Lingeman JE, Nakada SY, Pearle MS, Wolf JS Jr, Panel AUANG: Chapter 1: AUA guideline on management of staghorn calculus: diagnosis and treatment recommendations. J Urol 2005, 173(6):1991–2000.
8. Vprakasit DP, Sawyer MD, Herrell SD, Miller NL: Changing composition of staghorn calculi. J Urol 2011, 186(6):2285–2290.
9. Mandel N, Mandel I, Fryjoff K, Rejniak T, Mandel G: Conversion of calcium oxalate to calcium phosphate with recurrent stone episodes. J Urol 2003, 169(6):2026–2029.
10. Parks JH, Worcester EM, Coe FL, Evan AP, Lingeman JE: Clinical implications of abundant calcium phosphate in routinely analyzed kidney stones. Kidney Int 2004, 66(2):777–785.
11. Schwab SJ, Bander SJ, Kahr S: Renal infection in autosomal dominant polycystic kidney disease. Am J Med 1987, 82(4):714–718.
12. Hwang JH, Park HC, Jeong JC, Ha Baek S, Han MY, Bang K, Cho JY, Yu SH, Yang J, Oh KH, et al: Chronic asymptomatic pyuria precedes overt urinary tract infection and deterioration of renal function in autosomal dominant polycystic kidney disease. BMC Nephrol 2013, 14:
13. Mishra S, Sabnis RB, Desai MR: Percutaneous nephrolithotomy monotherapy for staghorn: paradigm shift for ‘staghorn morphometry’ based clinical classification. Curr uropol 2012, 22(2):148–153.
14. Umbreit EC, Childs MA, Patterson DE, Torres VE, LeRoy AI, Gettman MT: Percutaneous nephrolithotomy for large or multiple upper tract calculi and autosomal dominant polycystic kidney disease. J Urol 2010, 183(1):183–187.
15. Ng CS, Yost A, Streem SB: Nephrolithiasis associated with autosomal dominant polycystic kidney disease: contemporary urological management. J Urol 2000, 163(3):726–729.
16. Al-Kandari AM, Shoma AM, Eraky L, El-Kenawy MR, Al-Eezri H, El-Kappany HA: Percutaneous nephrolithotomy for management of upper urinary tract calculi in patients with autosomal dominant polycystic kidney disease. Urol 2009, 74(2):273–277.

doi:10.1186/1471-2369-14-168
Cite this article as: Mao et al.: Complete staghorn calculi in polycystic kidney disease: infection is still the cause. BMC Nephrology 2013 14:168.