Case Description

A 61 year-old male with a history of metabolic syndrome (hypertension, obesity, dyslipidemia), left kidney agenesis, compensatory hypertrophy and previous stone disease of the contralateral kidney, was sent by his general practitioner to our department because the incidental diagnosis of left seminal vesicle cyst lesion, not documented in the annual ultrasound control. He was father of 2 children and did not complain any lower urinary tract symptoms (LUTS) with a satisfied sexual life. He had a previous history negative for urinary tract infections (UTIs), dysuria, hematuria, ejaculatory pain, prostatitis, epididymitis. The examination revealed a patient in good general condition with body mass index of 29.07. The abdomen was soft with no palpable mass. Digital rectal examination was normal. Laboratory tests including urine culture and PSA showed no abnormality. Transabdominal ultrasound scan (US) confirmed left kidney agenesia with compensatory hypertrophy and revealed for the first time (not documented in the transabdominal US performed one year before) multiple hypoechoic formations in the site of the left seminal vesicle, the largest with 3.5 cm in diameter, accompanied by a posterior enhancement evoking impure fluid collection, without any intravesical protrusion. In the absence of symptoms, a conservative treatment was decided, with a strict clinical and ultrasonographic follow-up. The diagnosis of Zinner’s syndrome was made later, by ultrasound and magnetic resonance imaging, with the development of multilocular seminal vesicle cystic lesion, with a rapid onset (within one year). This malformation should be considered in the differential diagnosis of a pelvic cyst in male patients with renal agenesis.
Asymptomatic Zinner’s Syndrome

the first symptoms appear.

Figure 1: A-B: The left seminal vesicle cystic lesion at pelvic ultrasound.

Figure 2: The left seminal vesicle cystic lesion in axial (A) and sagittal (B) MRI image.

Discussion

Zinner’s syndrome is an unusual congenital malformation, first described in 1914 by Zinner, characterised by renal agenesis associated with ipsilateral seminal vesicle cysts and obstruction of the ejaculatory duct[1]. Less than 250 cases are reported in the literature[1]. This malformative association is explained by the common embryogenetic origin of the involved organs (mesonephric Wolffian duct)[2].

The pathophysiology of the malformation is explained by an insult occurred during the 4th and 13th week of gestation. From the distal mesonephric duct arise the ureteral bud which induces the differentiation and development of the kidney. The mesonephric duct will differentiate also to epididymis, ejaculatory duct, vas deferens, seminal vesicle, and hemitrigone. Complete failure of the mesonephric duct results to absence of ipsilateral kidney, ureter, hemitrigone, and seminal vesicle. Anomalous development of the distal mesonephric duct results to atresia of the ejaculatory ducts and abnormal ureteral budding; the latter leads to renal agenesis or dysplasia, the former leads to obstruction and cystic dilatation of the seminal vesicles with development of seminal cysts. The development of seminal vesicle cysts is variable along the time life and this variability is not still well explained[3].

Actually seminal vesicle cysts could be congenital or acquired[4]. Acquired cyst usually are found in adults during 2nd to the 5th decade of life, mainly at the time of greatest sexual and reproductive activity[5-7]. It was not the case of our patient that was older with a sudden and late onset of the seminal vesicle occurrence. A cyst is formed when secretions in the gland owing to insufficient drainage, caused to atresia, causes distention of the seminal vesicles. The cysts that were acquired are often bilateral and are discovered after a history of chronic prostatitis or even a prostate surgery[8].

The natural history of congenital seminal vesicle cysts is not well known. Kanavaki et al documented an 11-year history of this malformation from childhood to adolescence with a slow and gradual increase in size of the cyst, from 0.8 cm up to 1.9 cm[9].

Although acquired seminal vesicle cysts usually develop in elderly, they are often bilateral and occur in patients with history of prostatitis or prostate surgery. In the present case, it
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is unlikely that the cystic lesion is acquired because, despite its rapid onset, there are no obvious causes that might justify its secondary nature.

Usually cyst less than 5 cm in diameter remains asymptomatic and are incidentally discovered by pelvic imaging made for other reasons[10]. The most common symptoms are abdominal, perineal or pelvic pain with or without LUTS, UTIs, and sexual problems. Cysts exceed 12 cm, classified as “giant” cyst, could led LUTS and pelvic pain due to their “mass effect”[10]. Our patient, having a seminal vesicle cystic lesion more than 5 cm but less than 12 cm, never experimented any of these symptoms.

The diagnostic workups include physical examination, transabdominal or transrectal US, computed tomography, and MRI. Both CT and MRI allow to properly investigating renal and seminal vesicles anomalies, which are often combined. MRI exceeds CT in studying abdomen and pelvis not only for the absence of ionizing radiation but also for its imaging capability and soft tissue contrast which are crucial features in the assessment of relations between pelvic organs and structures[11]. In our patient, we performed both transabdominal US and pelvic MRI. Patients without clinical symptoms should not be treated and they can be monitored by transrectal US. Symptomatic seminal vesicle cysts often need mini-invasive surgical treatments such as percutaneous drainage with injection of sclerosing agents[12], laparoscopic[13] or robotic excision[14]. Our case is original as it shows a late and sudden onset of this malformation in the 6th decade of life, usually uncommon, without any symptom.

Conclusion

Zinner’s syndrome is unusual but this syndromic association is known and should be considered in the differential diagnosis of cystic pelvic masses also in males aged over 60 with renal agenesis or dysplasia. Ultrasound is useful for preliminary diagnosis but MRI provides a more detailed analysis and accurate diagnosis. Treatment is considered only for symptomatic patients. A strict follow-up should be proposed for asymptomatic cysts with a sudden onset. We recommend to follow patients with renal agenesis without a previous history of seminal vesicles cysts with a scheduled ultrasound according to patients symptoms. In the case of cyst occurrence we advised to shorten the follow-up (advisable every 6 months).

Conflict of interest: The authors have no conflicts of interest to declare in relation to this study.

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