Zinner syndrome: a rare embryological anomaly of the mesonephric ducts – Report of two cases and literature review

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Abstract
We present the cases of two males in their forties who present with symptoms or previous history of urolithiasis and known unilateral renal agenesis. On computed tomography evaluations of both patients, cystic lesions of the seminal vesicles were seen with further evaluation by transrectal ultrasound, making the diagnosis of Zinner syndrome. This entity has an embryological background that cannot be unseen. The symptoms are very characteristic, but incidental diagnosis based on imaging is very frequent. Good outcomes by laparoscopic treatment have been described.

Key words: Zinner syndrome. Renal agenesis. Seminal vesicle cyst.

Introduction
Zinner syndrome is a rare congenital anomaly usually presented in young men with a variety of symptoms. The image findings are very characteristic and new treatments have been documented in the past few years.

Presentation of clinical cases
Case 1
A 36-year-old male, with a history of known left renal agenesis, underwent a computed tomography (CT) scan on February 2017 for the evaluation of the right renal lithiasis and status of a double-J ureteral catheter seen on plain abdomen X-ray. The CT demonstrated agenesis of the left kidney and increase in volume of seminal vesicles with the left predominance, so it was integrated the diagnosis of Zinner syndrome, transrectal ultrasound was performed where a cystic aspect of the left seminal vesicle is observed (Fig. 1). Two months later, extracorporeal lithotripsy was performed with a torpid response and evolution with self-referral of the patient to private practice hospital on November 2017. No fertility problems were referred by the patient.

Case 2
A 37-year-old male was initially evaluated in December 2016 for presenting with a history of right renal fossa pain of 4 months of duration, suggestive of renal lithiasis; a renal ultrasound was performed where a stone in the distal ureter was observed. Treatment was started by placing a double right J catheter, which is observed in plain abdominal radiography adjacent to the lithium, for posterior realization of the right ureteroscopy; however, the patient suffers again from pain in the right renal fossa, fever, and non-clot forming hematuria, so a new contrast-enhanced CT scan was performed seeing right ureteral lithiasis, left seminal vesicle cyst and homolateral renal agenesis, concluding Zinner syndrome, transrectal ultrasound is performed in which the
The seminal vesicles are best characterized (Fig. 2). The fever yields to antibiotic therapy and is discharged for follow-up by the outpatient clinic, where resection of the left seminal vesicle cyst was done laparoscopically, being discharged after 3 days, on June 2017. The reason of the surgery was no clearly specified. No fertility problems were referred by the patient.

**Discussion**

Seminal cyst was described initially by Smith in 1872 and then the association with ipsilateral renal agenesis described by Zinner in 1914. By 2009, 200 cases have been reported in literature, from that date to 2018, we counted on PubMed 31 cases, without counting these two, giving an approximate of 233 cases in total. In 2009 in Taipei, Seo et al. reported an incidence of 0.0046%. Furthermore, there were reported about 50 cases in pediatric age in literature up to now, only 12/50 were diagnosed in the 1st year, and more than 80% were asymptomatic at long-term follow-up. It is a rare entity usually discovered when imaging evaluation is performed for other purposes, but sometimes is presented with symptomatology, and is man in the third or fourth decade of life the most affected, maybe because they are at the period of greatest reproductive or sexual activity, being found 83% of the cases between 11 and 40 years. We found it very often associated with infertility, painful ejaculation/dysuria or hemospermia, cryptorchidism, obstructed vas deferens mimicking paratesticular mass, and neoplasms in the urinary tract including in this last one malignant degeneration of the cyst.

This syndrome is explained by the common embryologic origin of both organs (kidneys and seminal vesicles) from the mesonephric duct and is due to an insult in embryo genesis between the 4th and 13th weeks of gestation. In the female, it is presented as an obstructed hemivagina and ipsilateral renal anomaly, also called OHVIRA syndrome. The ureteral bud arises from the dorsal part of the distal mesonephric duct and extends dorsocranially to meet and induce differentiation of the metanephric blastema, from which the kidney will develop. The mesonephric duct will differentiate to epididymis, ejaculatory duct, vas deferens, seminal vesicle, and hemitrigone. Complete failure of the mesonephric duct results to the absence of the kidney, ureter hemitrigone, and seminal vesicle. Anomalous development of the distal mesonephric duct results in atresia of the ejaculatory ducts and abnormal ureteral budding; the former leads to obstruction and cystic dilatation of the seminal vesicles with the development of seminal cysts. This is illustrated in Figure 1 and Figure 2.
cyst, the latter leads to renal agenesis or dysplasia. Unilateral renal agenesis is present in 0.1% of newborns and should alert for other genitourinary anomalies present in up to 30-40% of these patients. The ureter can be absent, incomplete, or may have an ectopic course to the seminal vesicle (present in 36% of the cases)\textsuperscript{3,21-23}.

Cyst smaller than 5 cm is usually asymptomatic and is discovered incidentally. Larger cysts can produce a wide variety of urological symptoms; usually, they measure more than 12 cm and can result in bladder or colon obstruction. The most affected side has been reported to be the right with a right: left ratio of 2:1. In both of our patients, the image findings were merely casual due to prostate hyperplasia treatment protocol evaluation through endorectal ultrasound and the other one due to known renal agenesis under CT evaluation. Findings by transrectal ultrasound include anechoic pelvic mass with a thick, irregular wall and occasional wall calcifications, or the mass may contain internal debris reflecting prior hemorrhage or infection. Magnetic resonance imaging (MRI) is paramount in making the definitive diagnosis, appearing as T2-weighted signal hypointensity structures and T1-weighted signal hyperintensity structures\textsuperscript{3,21}.

Differential diagnosis includes Mullerian cyst, ureterocele, dilated ureter, abscess or lymph nodes, and acquired seminal vesicle cysts. The last ones are often bilateral and related to older patients that have undergone prostate surgery or with chronic prostatitis\textsuperscript{3,21}.

Treatment is considered only for symptomatic patients and is surgical. Differential options exist from less invasive transrectal or transperineal aspiration of the cyst and is surgical. Differential options exist from less invasive transrectal or transperineal aspiration of the cyst to the seminal vesicle (present in 36% of the cases)\textsuperscript{3,21-23}. The authors declare that they have no conflicts of interest.

**Ethical disclosures**

Protection of human and animal subjects. The authors declare that no experiments were performed on humans or animals for this study.

Confidentiality of data. The authors declare that they have followed the protocols of their work center on the publication of patient data.

Right to privacy and informed consent. The authors have obtained the written informed consent of the patients or subjects mentioned in the article. The corresponding author is in possession of this document.

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