Management of lower urinary tract fibroepithelial polyps in children

Sybille Rousseau¹, Matthieu Peycelon^{2,3}, Céline Grosos¹, Valeska Bidault², Anna Poupalou⁴, Garance Martin⁵, Éric Dobremez⁶, Luke Harper⁶, Claire Raquillet⁷, Alexis Arnaud⁸, Emmanuel Sapin⁹, Aurélien Scalabre¹⁰, Philippe Buisson¹¹, Guillaume Levard¹², Isabelle Pommepuy¹³, Maguelonne Pons¹⁴, Laurent Fourcade¹, and Quentin Ballouhey¹

Corresponding author:

Doctor Quentin Ballouhey

Q.ballouhey@gmail.com

Service de chirurgie pédiatrique. Hôpital Mère-Enfant. Centre Hospitalier Universitaire de Limoges. 8 Avenue Dominique Larrey 87042, Limoges, France

Doctor Céline Grosos, Dr Sybille Rousseau, and Professor Laurent Fourcade

(1) Service de chirurgie pédiatrique. Hôpital Mère-Enfant. Centre Hospitalier Universitaire de Limoges. 8 Avenue Dominique Larrey 87042, Limoges, France

Doctor Matthieu Peycelon and Dr Valeska Bidault

- (2) Service de chirurgie et urologie pédiatrique. Hôpital Universitaire Robert-Debré, APHP Centre de Référence Maladies Rares « MARVU » ; Université de Paris, Paris, France
- (3) Pediatric Urology, Riley Children's Hospital, Indiana University School of Medicine, Indianapolis, IN, United States

Doctor Anna Poupalou

(4) Service de Chirurgie Pédiatrique, Hôpital HUDERF-ST Pierre (Université Libre de Bruxelles-ULB), Brussels, Belgium

Doctor Garance Martin

(5) Service de chirurgie pédiatrique, Hôpital Trousseau, Paris, France

This is the author's manuscript of the article published in final edited form as:

Rousseau, S., Peycelon, M., Grosos, C., Bidault, V., Poupalou, A., Martin, G., Dobremez, É., Harper, L., Raquillet, C., Arnaud, A., Sapin, E., Scalabre, A., Buisson, P., Levard, G., Pommepuy, I., Pons, M., Fourcade, L., & Ballouhey, Q. (2020). Management of lower urinary tract fibroepithelial polyps in children. Journal of Pediatric Surgery. https://doi.org/10.1016/j.jpedsurg.2020.05.030

Professor Eric Dobremez and Doctor Luke Harper (6) Service de chirurgie pédiatrique, Centre Hospitalier Universitaire de Bordeaux, France **Doctor Claire Raquillet** (7) Service de chirurgie pédiatrique. Centre Hospitalier Ballanger, Aulnay-sous-Bois, France **Doctor Alexis Arnaud** (8) Service de chirurgie pédiatrique, Centre Hospitalier Universitaire de Rennes, France **Professor Emmanuel Sapin** (9) Service de chirurgie pédiatrique, Centre Hospitalier Universitaire de Dijon, France Doctor Aurélien Scalabre (10) Service de chirurgie pédiatrique, Centre Hospitalier Universitaire de Saint Etienne, France **Doctor Philippe Buisson** (11) Service de chirurgie pédiatrique, Centre Hospitalier Universitaire d'Amiens, France Professor Guillaume Levard (12) Service de chirurgie pédiatrique, Centre Hospitalier Universitaire de Poitiers, France **Doctor Isabelle Pommepuy**

(13) Service d'anatomo-pathologie, Centre Hospitalier Universitaire de Limoges, France

(14) Service de chirurgie pédiatrique, Centre Hospitalier Universitaire de Clermont-Ferrand, France

Doctor Maguelonne Pons

Abstract

Introduction

Fibroepithelial polyps (FEP) of the lower urinary tract are relatively common in adults but rare in children, with fewer than 250 cases reported in the literature to date.

Objective

The aim of this study was to address the experience of FEP management in children.

Study Design

A retrospective multicenter review was undertaken of children with defined FEP of the lower urinary tract managed between 2008 and 2018. The data at eighteen pediatric surgery centers were collected. Their demographic, radiological, surgical, and pathological information were reviewed.

Results

A total of 33 children (26 boys; 7 girls) were treated for FEP of the lower urinary tract at thirteen centers. The most common presentation was urinary outflow as hematuria (41%), acute urinary retention (25%), dysuria (19%), or urinary infections (28%). A prenatal diagnosis was made for three patients with hydronephrosis. Almost all of the children (94%) underwent ultrasound imaging of the urinary tract as the first diagnostic examination, 23 (70%) of them also either had an MRI (15%), cystourethrography (25%), computerized tomography (6%), or cystoscopy (45%). Two of these children (6%) had a biopsy prior to the surgery. The median preoperative delay was 7.52 (range: 1-48) months. Most of the patients were treated endoscopically, although four (12.1%) had open surgery and two (6.1%) had an additional incision for specimen extraction. The median hospital stay was 1.5 (range: 1-10) days. There were no recurrences and no complications after a median followup of 13 (range: 1-34) months.

Discussion

The main limitation of our study is the retrospective design, although it is the largest one for this pathology.

Conclusion

This series supports sonography as the most suitable diagnosis tool before endoscopy to confirm the diagnosis and to perform the resection for most FEP in children. This report confirms the recognized benign nature in the absence of recurrences.

Level of Evidence

Level V

Keywords: Polyps; Children; Lower urinary tract; Bladder tumor; Ultrasonography; Endoscopic mucosal resection

INTRODUCTION

Urinary tract polyps occur rarely in children [1]. Since the 19th century, approximately 250 cases have been reported, mostly as case reports (Table 1)[1-11]. They are usually discovered in childhood or adolescence, although some authors have also described them in adults [12]. These fibroepithelial polyps (FEP) are congenital tumors of mesodermal origin and they most often occur in males [13]. They only rarely occur in females. They can affect the entire urinary tract, from the renal pelvis to the urethra[14-16]. They are mostly located in the bladder [5] or the urethra. The posterior portion of the urethra is the predominant location [3, 6, 17-20], whereas anterior urethral polyps are only reported rarely [21-24]. They are usually described as a benign pedunculated polyp or bladder mass [4] at sonography (*Figure 1A*). The main differential diagnosis is rhabdomyosarcoma, which is a heterogeneous mass with malignant characteristics. The pathology report typically confirms the presence of a fibroepithelial entity (*Figure 2*) composed of vascular connective tissue [25].

The main features depend on the location of the FEP. As they have a stalk, these polyps are mobile in the bladder or the urethra. At the pathognomonic clinical level, they hence present as an intermittent or acute obstruction of the bladder. They can also cause bladder irritation that manifests as hematuria, dysuria, or urinary tract infections (UTIs). In case of an unusual presentation, the diagnosis can require supplementary preoperative imaging such as MRI (Figure 1B) or endoscopic examination [2, 8, 26]. Surgical management is most commonly achieved endoscopically by transurethral resection. FEP of the lower urinary tract are benign lesions and no recurrences or malignant behavior have been reported to date. Although they are benign tumors, delayed diagnosis can

lead to renal failure as a result of bladder obstruction [27]. Due to the rarity of this condition, no standard management and treatment have been published for this entity. The aim of our study was to report the current management of FEP in children.

METHODS

A multicenter review was carried out to compile cases of FEP of the lower urinary tract in the past ten years. Children operated on for FEP between 2008 and 2018 were considered for this study. This study was approved by the relevant ethics committee, with reference number 301-2019-67.

A survey was sent to 34 centers in order to collect relevant clinical, radiological, and surgical data. These data included the age at presentation, the type of management (endo-surgery versus open surgery), prior medical history, associated anomalies, symptoms, the perioperative course, histopathology findings, and follow-up. Patients were included in case of FEP confirmed by histopathology and operated on between 2008 and 2018. Exclusion criteria were being over 18 years of age, a lack of pathology results, an upper urinary tract FEP, or an absence of follow-up. Descriptive statistics were performed using Fisher's exact test for the categorical variables, the Student's *t*-test for the parametric continuous data (means and the SD are presented), and the Mann-Whitney test for the non-parametric continuous data (medians and the IQR were used). A p-value less than 0.05 was considered significant.

RESULTS

Of the 34 centers, 18 centers replied to the survey. Three centers had unusable data and another one had not encountered cases of FEP. At the 14 remaining pediatric centers, a total of 36 medical files with FEP met the inclusion criteria. All of the patients were managed according to each center's protocols. Three of them were ultimately excluded due to a ureteral position of the FEP. The median age of the patients (26 boys and 7 girls) was 6.2 (range: 1 month-14 years) years of age, and none of them had a relevant prior medical history.

The clinical presentation (Table 2) was non-specific and most of the time comprised symptoms such as hematuria (39%), infections (27%), acute urinary retention (24%), dysuria (18%), hydronephrosis (9%), and pain (3%). For three infants, there was a prenatal diagnosis according to the ultrasonography depiction of hydronephrosis during the third trimester of pregnancy.

Sonographic assessment was used extensively in the diagnostic process, followed by endoscopic evaluation and histological analysis. All but two of the children had ultrasonography as the first diagnostic examination. A total of 13 children (40%) had another complimentary exam, which was either an MRI (15%) (Figure 1B), VCUG (Voiding Cystourethrography) (24%), or a CT scan (6%). Fifteen patients (45%) had a preoperative cystoscopy to sustain the diagnostic modality: five were performed extemporaneously during the same anesthesia to confirm the diagnosis by a brief consultation between two surgeons, and seven were performed systematically before laparotomy (12%) or before direct resection (9%) for FEP protruding through the external urethral meatus. The three remaining patients underwent two distinct cystoscopic procedures: during the first cystoscopy, a biopsy was performed for two of them (6%) because of an unusual endoscopic appearance, and a technical problem occurred in one case (3%). All of the other patients underwent direct surgical excision. The delay between the first symptoms and the surgical

management of the polyp was between one week and 49 months, with a median of 7.52 months.

Endoscopic management with transurethral resection (79%) was performed for 26 patients. For 9 cases (27%), Bugbee electrocautery was used to cut the polyp at its base, and polyp retrieval was performed transurethrally using forceps. A resectoscope was used in 17 cases (51%), and an additional trocar was necessary to stabilize large floating polyp in the bladder for two patients (6%). The specimen (median size 8.5 mm (range: 4-10.2)) was extracted transurethrally in 23 cases (70%) using forceps in twenty cases (60%) or a basket in three cases (18%). One polyp (3%) that was 13 mm in size was extracted by trocar incision, and two specimens (6%) were extracted by cystostomy (polyp sizes of 34 mm and 17 mm, respectively).

In case of large polyps of the bladder neck, an open approach was selected due to exposure difficulties at endoscopy. Seven (21%) patients were treated by open surgery: four (12%) boys by a Pfannenstiel incision after preoperative cystoscopy (the polyp sizes were 22 mm, 14 mm, 20 mm, and 10 mm) and three (9%) girls (*Figure 3*) by direct perineal resection for FEP protruding through the external urethral meatus.

The polyp was located most frequently in the urethra (59%), which in 11 cases included a polyp of the urethral posterior wall (33%), and only one case of location at the urethral anterior wall was reported. The other main location was the bladder (41%). A statistically significant positive association between UTI and urethral localization was found (p < 0.05).

The median size of the polyps was 11.6 mm (range: 4.7-15). All of the specimens were histologically examined, which confirmed the diagnosis of

fibroepithelial polyp (Figure 2). Urethral polyps are cured statistically less frequently by exclusive endoscopic resection (p = 0.02).

Only 15 (45%) of the 33 patients had a postoperative urinary catheter, which was removed at a "median" time of 1.3 (range: 1-7) days postoperatively. In three cases (9%) involving patients who underwent an open approach, the catheter was a suprapubic catheter. No postoperative complications were reported. The mean duration of the hospital stay was 1.3 days (± 2.1 days). Fifteen children (45%) were received treatment as outpatients.

There was no polyp recurrence after an average total follow-up of 13 months (1 - 34), and all of the patients became symptom-free. One child had reflux associated with the polyp, which was still noted during the follow-up. For all of the other patients, there was no reflux, no urinary retention, no hematuria, and no infection following the endoscopic resection.

DISCUSSION

Fibroepithelial polyps are a rare entity that can be encountered during childhood as a pedunculated lesion mostly in the urethral posterior wall (33% in this study). We here report the largest series of lower urinary tract FEP in children. The aim of polyp management is for the children to become symptom-free and to prevent any renal failure. Thus, it is important to identify these lesions and to reduce the preoperative delay. Given the rarity of this lesion, an algorithm for FEP management is proposed (Appendix 1).

The clinical triad of intermittent urinary retention, hematuria, and lower urinary tract symptoms has already been described by Akbarzadeh *et al.* in 2014 [3] as being clearly suggestive of urethral polyps in children. The clinical

presentation of FEP depends on their location. Large posterior urethral polyps protrude and cause outlet obstruction, which can sometimes lead to acute symptoms. Bladder urethral stones have to be kept in mind as a differential diagnosis. In our series, this polyp location was revealed by acute urinary retention in 25% of cases and urethral localization of FEP was associated with larger-sized lesions and a higher incidence of UTI.

Ultrasonography is an excellent and non-invasive method to image and characterize bladder lesions. Urinary ultrasound can be considered to be the first-line and the only morphological examination, revealing a single, spherical, echoic, smooth lesion emanating from the bladder mucosa. A complex image with a grape-like appearance or cystic areas is suggestive of rhabdomyosarcoma. In case of suspected malignancy, contrast MRI provides higher resolution and it can reveal the origin and the local extension of the tumor [28]. In five cases (15%) in our series, an MRI was also performed, thereby confirming the ultrasonography results without providing supplementary information. In eight cases (24%), ultrasonography did not adequately reveal the polyp, and VCUG was hence performed. A diagnosis of a polyp lesion was made in light of a bladder defect. We, therefore, believe that ultrasonography is an adequate assessment tool when a diagnosis of FEP is likely. In case of doubt or no visible mass by ultrasonography, VCUG appears to be the second-line examination. It also has the advantage of excluding posterior urethral valves, which is the differential diagnosis in case of obstructive bladder symptoms in males.

Cystoscopy can be employed both for the diagnosis and for therapeutic purposes. We, therefore, recommend performing cystoscopy to confirm the diagnosis and the treatment at the same time. A typical radiological and

endoscopic presentation allows FEP management with the administration of single anesthesia, as was the case for 30 patients (90%) in our study.

Prenatal diagnosis is extremely rare [7]. In three of our cases (9%), the hypothesis raised in light of hydronephrosis on prenatal ultrasonography. For two (6%) of them, no polyp could be discerned on the postnatal ultrasonography. VCUG was, therefore, performed to rule out vesicourethral reflux and it allowed for a successful diagnosis. Transurethral resection was performed in three of our cases (9%) of neonatal patients without encountering technical difficulties or postoperative complications.

Comparison with the adult population [12, 14, 29, 30] indicates that the clinical presentation of FEP appears to be similar. The management, however, is not entirely the same. Indeed, when there is the possibility of a polyp in adult patients, cystoscopy is performed under local anesthesia to collect biopsies and to probe for the presence of a bladder tumor, without further imaging investigation. In our study, seven cases (21%) were found in girls, which is even rarer than in boys. Most of them exhibited a UTI or hematuria. Three polyps (9% of cases) were located on the urethra and were removed by urethral surgery under direct vision (Figure 3), whereas the four other cases (12%) had a bladder location and were resected endoscopically.

The standard of care for the polyps is transurethral resection. Use of a resectoscope or forceps can achieve satisfactory fulguration of the base of the polyp. This series did not involve use of a laser fiber and there have been no publications of lower urinary tract location in children. Laser therapy is the treatment of choice for ureteral polyps and a number of polypectomies with Holmium have been reported in children [15]. In our opinion, it remains a good treatment option despite the limited resection depth.

In our series, urethral location is associated with less exclusive use of an endoscopic procedure, probably due to exposure difficulties and larger-sized lesions. When the polyp is too large or when it floats into the bladder, a bladder trocar is inserted for stabilization or exposure before endoscopic retrieval. Based on our series, the size of the incision for urethral retrieval appears to be 20 mm; above that size, a dedicated cystostomy appears to be required. Thus, in case of urethral lesion larger than 20 mm, a trocar or a cystostomy can be necessary to support the endoscopic procedure.

In cases of large FEP, fragmentation of the specimen was not considered in this series by the surgeons so as to favor the quality of the definitive pathology examination. This alternative can, however, be an option with an acceptable risk according to the long-term results in case of clear radiological and endoscopic FEP criteria. Such management must be decided at the beginning of the resection, before cutting the base. Indeed, endoscopic fragmentation of a floating lesion can be very difficult.

In our series, 45% of the patients had a postoperative urinary catheter, which was removed after a median of 1.3 days without hematuria. If the surgical procedure is accomplished without any complications, the procedure can be performed as an outpatient (as it was the case for 15 of our patients). No recurrences were reported after a follow-up of more than 12 months, which confirms the data in the literature: recurrence can appear if the stalk of the polyp is not completely excised [31].

The limitations of our study are that it was a retrospective study. Moreover, although if it is the largest study to date for this pathology, only a limited number of patients were included, thereby resulting in a lack of statistical power. However, it allowed the management of this rare disease to be refined.

Conclusion

This series supports the notion that the use of sonography is the most suitable diagnosis tool before endoscopic assessment and resection of FEP in children. In case of an unusual presentation, VCUG is the most informative morphological examination. In case of FEP larger than 20 mm, mini-invasive treatment may also require a bladder trocar for exposure, and sometimes a bladder incision for specimen retrieval. This report also confirms the widely recognized benign nature of FEP with the absence of recurrences.

Competing interests: The authors declare that they have no conflicts of interest.

References

- 1. Demircan M, Ceran C, Karaman A, Uguralp S, Mizrak B: Urethral polyps in children: a review of the literature and report of two cases. *Int J Urol* 2006, 13:841-843.
- 2. Ballard DH, Rove KO, Coplen DE, Chen TY, Hulett Bowling RL: Fibroepithelial polyp causing urethral obstruction: Diagnosis by cystourethrogram. *Clin Imaging* 2018, 51:164-167.
- 3. Akbarzadeh A, Khorramirouz R, Kajbafzadeh AM: Congenital urethral polyps in children: report of 18 patients and review of literature. *J Pediatr Surg* 2014, 49:835-839.
- 4. Kaba M, Kaba S, Kaya TY, Eren H, Pirincci N: A giant pedunculated urothelial polyp mimicking bladder mass in a child: a rare case. *Case Rep Pediatr* 2014, 2014:935850.
- 5. Natsheh A, Prat O, Shenfeld OZ, Reinus C, Chertin B: Fibroepithelial polyp of the bladder neck in children. *Pediatr Surg Int* 2008, 24:613-615.
- 6. Isaac J, Snow B, Lowichik A: Fibroepithelial polyp of the prostatic urethra in an adolescent. *J Pediatr Surg* 2006, 41:e29-31.
- 7. Beluffi G, Berton F, Gola G, Chiari G, Romano P, Cassani F: Urethral polyp in a 1-month-old child. *Pediatr Radiol* 2005, 35:691-693.
- 8. Barzilai M, Shinawi M, Ish-Shalom N, Mecz Y, Peled N, Lurie A: A fibroepithelial urethral polyp protruding into the base of the bladder: sonographic diagnosis. *Urol Int* 1996, 57:129-131.
- 9. Rosenkilde M, Leffers AM: Congenital urethral polyp--a rare cause of infravesical obstruction. A case report. *Acta Radiol* 1995, 36:196-197.
- 10. Gleason PE, Kramer SA: Genitourinary polyps in children. *Urology* 1994, 44:106-109.
- 11. De Castro R, Campobasso P, Belloli G, Pavanello P: Solitary polyp of posterior urethra in children: report on seventeen cases. *Eur J Pediatr Surg* 1993, 3:92-96.
- 12. Walsh IK, Keane PF, Herron B: Benign urethral polyps. *Br J Urol* 1993, 72:937-938.
- 13. H Moch PH, TM Hulbright, VE Reuter: WHO Classification of Tumors of the Urinary System. In: World Health Classification of Tumors. 4th edn. Edited by Cancer IAfRo; 2016.
- 14. Akdere H, Cevik G: Rare Fibroepithelial Polyp Extending Along the Ureter: A Case Report. *Balkan Med J* 2018, 35:275-277.
- 15. Li R, Lightfoot M, Alsyouf M, Nicolay L, Baldwin DD, Chamberlin DA: Diagnosis and management of ureteral fibroepithelial polyps in children: a new treatment algorithm. *J Pediatr Urol* 2015, 11:22.e21-26.
- 16. Ludwig DJ, Buddingh KT, Kums JJ, Kropman RF, Roshani H, Hirdes WH: Treatment and outcome of fibroepithelial ureteral polyps: A systematic literature review. *Can Urol Assoc J* 2015, 9:E631-637.
- 17. Bagley FH, Davidson AI: Congenital urethral polyp in a child. *Br J Urol* 1976, 48:278.
- 18. Downs RA: Congenital polyps of the prostatic urethra. A review of the literature and report of two cases. *Br J Urol* 1970, 42:76-85.
- 19. Foster RS, Garrett RA: Congenital posterior urethral polyps. *J Urol* 1986, 136:670-672.
- 20. Gunther I, Abrams HJ, Sutton AP, Buchbinder MI: Fibroepithelial polyp of the verumontanum: a case report and review of the literature. *J Urol* 1979, 121:525-526.
- 21. Anandan N, Shetty SD, Patil KP, Ibrahim AI: Acute urinary retention caused by anterior urethral polyp. *Br J Urol* 1992, 69:321-322.
- 22. Coleburn NH, Hensle TW: Anterior urethral polyp associated with hematuria in six-year-old child. *Urology* 1991, 38:143-144.
- 23. Moriya K, Kobayakawa H, Yasumoto R, Maekawa M: Anterior urethral polyps. *Br J Urol* 1988, 62:183
- 24. Redman JF, Robinson CM: Anterior urethral polyp in a child. J Pediatr Surg 1977, 12:735-736.

- 25. Levin TL, Han B, Little BP: Congenital anomalies of the male urethra. *Pediatr Radiol* 2007, 37:851-862; quiz 945.
- 26. Stephens FD: Urethral obstruction in childhood; the use of urethrography in diagnosis. *Aust N Z J Surg* 1955, 25:89-109.
- 27. Stueber PJ, Persky L: Solid tumors of the urethra and bladder neck. J Urol 1969, 102:205-209.
- 28. Shelmerdine SC, Lorenzo AJ, Gupta AA, Chavhan GB: Pearls and Pitfalls in Diagnosing Pediatric Urinary Bladder Masses. *Radiographics* 2017, 37:1872-1891.
- 29. Cai Y, Zhang Z, Yue X: Rare giant primary ureteral polyp: A case report and literature review. *Mol Clin Oncol* 2017, 6:327-330.
- 30. Tsuzuki T, Epstein JI: Fibroepithelial polyp of the lower urinary tract in adults. *Am J Surg Pathol* 2005, 29:460-466.
- 31. Juskiewenski S, Miquel H, Fabre J, Vaysse P: [Posterior urethral polyps in boys. Apropos of 3 cases]. *Ann Urol (Paris)* 1972, 6:259-264.

Figure 1 Fibroepithelial polyp imaging

A: Ultrasonography typical presentation Image of a 20 x 12-mm-sized fibroepithelial polyp in a 21-month-old boy with intermittent bladder obstruction.

B: MRI features of a fibroepithelial polyp in the bladder

T2 sequence showing a 16 x 12-mm-sized pedunculated lesion in a 21-month-old boy.

Figure 2 Morphometry of a fibroepithelial polyp

A: Photograph of a 12-mm-sized fibroepithelial polyp (preparation with HES). The white arrow indicates the center of the lesion with fibrous connective tissue containing glands, smooth muscles, and nerves (10x magnification). The overlying epithelium is urothelium that contains areas of ulceration (black arrows).

B Simple hierarchical pattern of urothelium with a normal thickness and appearance (40xmagnification).

Figure 3 Perineal aspect of a fibroepithelial polyp

Photograph of a 9-mm-sized fibroepithelial polyp protruding through the urethral meatus of a 14-month-old girl.

Table 1 Cases of fibroepithelial polyps reported in the past 25 years Table 2 Characteristics of the patients

Table 1

Series or case reports of FEP	Numb er of patien ts	Age at diagno sis	Gend er	Symptoms	Diagnost ic evaluati on	Localization	Managem ent	Follo w-up
Current study Ballard [2]	33	7.1	M (26) F (7)	AUR (8), UTI (9), hematuria (15), HN (3) AUR	US, CT Scan, VCU, cystosco py, MRI VCU, CT	bladder (14) urethra (19) verumontan um	endoscopy (26) open surgery (7) endoscopy	3-55 mont hs
Akbarzad eh [3]	18	3.5	M (14) F (4)	AUR (7), UTI (6), dysuria (10), hematuria (14), HN(4), reflux (6)	scan VCU, cystosco py	urethra and bladder	endoscopy (17) endoscopy + cystostom y (1)	3-17 years
Kaba [4]	1	14	M	hematuria	US, CT Scan, VCU, cystosco py		open surgery	
Ala Natsheh [5]	2	3.5 [2- 5]	М	hematuria (1), dysuria (1), AUR (1)	,		endoscopy (2)	1-5 years
Demircan [1]	2	1.75 [1.5-2]	M (1) F (1)	hematuria interlabial mass	VCU (2), US (2), cystosco py (1)	urethra	endoscopy + cystostom y (1) direct surgery (1)	1 mont h
Isaac [6]	1	16	М	AUR	CT Scan, cystosco py	urethra	endoscopy	
Beluffi [7]	1	0.08	М	hydronephr osis	US, VCU	verumontan um	endoscopy + cystostom y	
Barzilai [8]	1	0.8	М	AUR	US, VCU, cystosco py	verumontan um	endoscopy	6 mont hs
Rosenkild e [9]	1	2.5	M	AUR	VCU, cystosco py, US	urethra	cystostom y	1.5 mont hs

Gleason	12	8.9 [1-	М	hematuria	VCU (7),	verumontan	endoscopy	12
[10]		14]	(12)	(5),	US	um (9),	(11),	mont
				obstructive		posterior	endoscopy	hs
				symptoms		urethra (3)	+	
				(4), AUR (2)			cystostom	
							y (1)	
De	17	<2 (6)	М	UTI (4)	US (3)	posterior	endoscopy	12
Castro		2-6 (5)		AUR (4)	VCU (17)	urethra (17)	(17)	mont
[11]		>6 (6)		hematuria	Cystosco			hs
				(7)	py (3)			
				dysuria (9)	' ' '			

Abbreviations; FEP: fibroepithelial tumor; AUR: acute urinary retention; UTI: urinary tract infection; HN: hydronephrosis; VCU: voiding cystourethrogram; US: ultrasonography

Table 2

Data	Total	Group 1- bladder FEP	Group 2- urethra FEP	Differen ce
Number	33	14	19	
Gender (M/F)	26/7	10/4	16/3	P= 0.42
Age at presentation (years)	7.11 (± 5.34)	7.4 (± 5.7)	6.9 (± 5.2)	p= 0.82
Symptoms (%)				
- AUR	- 8 (24%)	- 2 (14%)	- 6 (32%)	p= 0.42
- UTI	- 9 (27%)	- 1 (7%)	- 8 (42%)	p= 0.04
- Hematuria	- 14 (42%)	- 8 (57%)	- 6 (32%)	p= 0.17
- Pain	- 1 (3%)	- 1 (7%)	- 0	p= 0.40
- Dysuria	- 5 (15%)	- 2 (14%)	- 3 (16%)	p= 1
- Hydronephrosis	- 3 (9%)	- 2 (14%)	- 1 (5%)	p= 0.56
Diagnostic evaluation				
(%)	- 31 (94%)	- 14 (100%)	- 17 (89%)	p= 0.49
- US	- 5 (15%)	- 1 (8%)	- 4 (21%)	p= 0.36
- MRI	- 8 (24%)	- 1 (8%)	- 7 (37%)	p= 0.1
- UC	- 2 (6%)	- 0	- 2 (10%)	p= 0.5
- CT	- 15 (45%)	- 9 (64%)	- 7 (37%)	p= 0.30
- Cystoscopy				
Lesion size (mm)	11.6 (± 8.7)	8.6 (± 8.6)	13.9 (± 8.3)	p= 0.13
Surgery (%)				
- Laparotomy	- 4 (12%)	- 0	- 4 (21%)	p= 0.12
- Perineal	- 3 (9%)	- 0	- 3 (16%)	p= 0.24
approach	- 26 (78%)	- 14 (100%)	- 12 (63%)	p= 0.01
- Endoscopic	• 2	• 0	• 2	• p
Trocar	(8	• 0	(1	=
 Cystosto 	%)		2%	0
my	• 2		• 2	
	(8		(1	5
	%)		2%	• p
)	=
				0
				5
Urinary catheter (%)	15 (47%)	4 (28%)	11 (58%)	p= 0.049
Follow-up (years)	1.1	1.28	0.95	p= 0.46

Abbreviations: US: ultrasonography; UC: urethrocystography; CT: computed tomography; AUR: acute

urinary retention; UTI: urinary tract infection

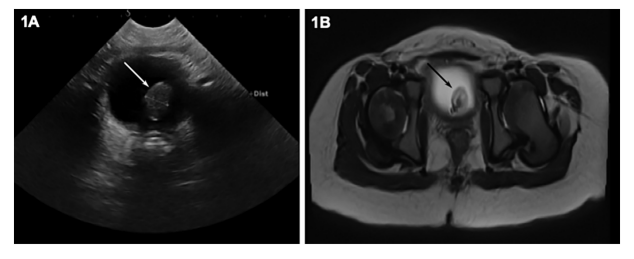


Figure 1

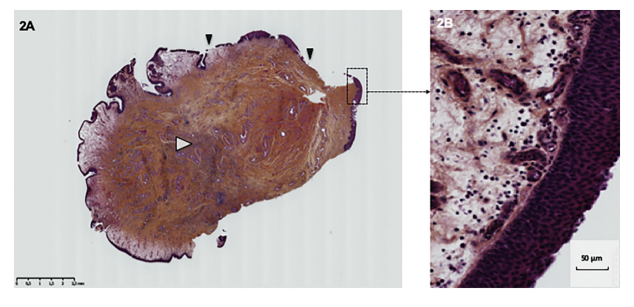


Figure 2



Figure 3