Overview of Complications and Management Approaches of Unilateral Ureteral Obstruction
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ABSTRACT
Unilateral ureteral obstruction (UOO) is a popular speculative version of renal injury, associated with kidney drop. Throughout the time of UOO, nephron damage is started by a boosted intratubular hydrostatic pressure and also secondary ischemia. In the course of the disease, even more, general arbitrators of kidney injury are invited.

Objective: Primary goal of this review was to highlight the complications following the Unilateral ureteral obstruction (UOO) and obstructive uropathy in general, as well as to discuss the therapeutic strategies for this urological condition.

Material and method: We review the Published literature concerning unilateral ureteral obstruction (UOO) up to August, 2017. Search was conducted using electronic databases; Medline, and Embase. Search strategy through mentioned databases was performed using medical subject headings (MeSH) as following, “obstructive uropathy”, “obstructive nephropathy”, “Unilateral ureteral obstruction”, “UOO”. Furthermore, bibliographic of the articles were manually searched for having more relevant studies. Restriction to our search was applied to only English language published studies.

Conclusion: Speculative UOO is an animal version of the human condition, which duplicates not only human-ureteral-tract-obstruction-induced kidney injury, it is largely held that CKD happens consequently of the procedure of damaging fibrosis. UOO is a well-described model of renal fibrosis and the mood of CKD occurrence while UOO persisting.

Keywords: obstructive uropathy, obstructive nephropathy, Unilateral ureteral obstruction, renal injury, chronic kidney disease.

INTRODUCTION
Unilateral ureteral obstruction (UOO) is a popular speculative version of renal injury, associated with kidney failure, discomfort or fever is a urological emergency need of punctual assessment and therapy. Present treatment methods including pharmacological treatment, and decompression by a cystoscopically insertion of ureteral stent or percutaneous nephrostomy (PCN) tube. UOO is well-described version of renal fibrosis and thus is considered a version of chronic kidney disease (CKD). Popular changes in UOO consist of declines in renal feature and also increased fibrosis, tubular apoptosis, as well as cellular spreading leading to renal failure later on.

After UOO, both kidney blood flow (RBF) and glomerular filtering price (GFR) decrease, as well as the reductions are irreversible without intervention. If UOO is managed within a specific duration, both RBF and GFR rise. Prognosis, quality of life and complications are important variables to be considered with UOO. Ureteral stents as well as percutaneous nephrostomy tubes have similar complication prices in the administration of malignant extrinsic ureteral compression.

Renal fibrosis is considered the last usual pathway for the majority of kinds of dynamic kidney diseases as well as entails glomerular sclerosis and/or interstitial fibrosis triggered by...
UO \textsuperscript{1}. In the 1970s, a rabbit UO design demonstrated fibroblast proliferation and changes in the renal parenchyma \textsuperscript{5}. Since a given that searching results, UO animal versions have been considered the useful versions of how obstructive nephropathy especially UO can be easily adjusted by changing the moment, intensity and duration of the speculative setup \textsuperscript{6}. This animal model is used to elucidate the pathogenesis of obstructive nephropathy and the pathological devices moderating kidney fibrosis \textsuperscript{6}.

Our primary goal of this review was to highlight the complications following the Unilateral ureteral obstruction (UO) and obstructive uropathy in general, as well as to discuss the therapeutic strategies for this urological condition.

**METHODOLOGY**

We reviewed the Published literatures concerning unilateral ureteral obstruction (UO) up to August, 2017. Search was conducted using electronic databases; Medline, and Embase. Search strategy through mentioned databases was performed using medical subject headings (MeSH) as following, “obstructive uropathy”, “obstructive nephropathy”, “Unilateral ureteral obstruction”, “UO”. Furthermore, bibliographic of found articles were manually search for more relevant studies. Restriction to our search was applied to only English language published studies.

The study was done after approval of ethical board of King Abdulaziz university.

- Complications of UO:
  - Irreversible renal function damage and Acute kidney injury:

Interstitial extracellular matrix development is the classical histological characteristic of chronic kidney injury as well as is usually the most effective structural correlate of the level of renal function loss and also a high chance of development risk \textsuperscript{7}. Measurable measures of a tubular function or possibly surrogate measures such as proteinuria. Each of these poses technological difficulties in the mouse. In addition, although the widely-used UO model causes considerable parenchymal damages, the intact contralateral kidney compensate functionally.

Numerous methods have actually been used to determine tubular area as well as stability as a different method to evaluate the intensity of renal parenchymal loss related to fibrosis \textsuperscript{8}.

A vital research study \textsuperscript{9} used an improved reversible unilateral ureteral obstruction (rUO) model in mice to explore the growth of irreversible loss of kidney function after raising time of obstruction complied with by the release of the obstruction. In normal mice, renal function depended on the moment of obstruction, such that with 1 - 2 days of obstruction, there was the complete function of the obstructed kidney complying with the removal of contralateral kidney function \textsuperscript{9}. With ureteral obstructions of 3 or more days, there was obstruction-time reliant renal failing \textsuperscript{9}.

Immediately adhering to the removal of contralateral (left) kidney function, there was a transient however considerable surge in BUN, which then fell and also supported at a level boosted from standard, in certain, the value of rodent stress is plainly shown by our outcomes. Reversible ureteral obstruction provides numerous essential benefits for researching growth and also a development of CKD.

The transient rise and afterward fall in BUN likely reflects the anticipated shift in blood flow to the formerly obstructed (right) kidney adhered to by hyperfiltration \textsuperscript{6}. In a similar way, in rats, in another study \textsuperscript{10} kidney function in the CLK was raised either 7 days or 3 months in an additional study \textsuperscript{11} after alleviation of UO.

We demonstrated here that the CLK kept a significantly higher percentage of the RBF and GFR at 7 and 14 days’ post relief. In a neonatal rat design, Chevalier et al.\textsuperscript{12} showed the progression of interstitial fibrosis after relief of UO. Twenty-eight days after relief of UO, interstitial collagen, α-smooth muscle actin, and also TGF-β1 were drastically raised in the POK compared with sham-operated rats \textsuperscript{13}. Interstitial fibrosis is a complex procedure that involves synthesis and also the destruction of extracellular matrix proteins, cellular seepage, epithelial-mesenchymal improvement, and also tubular apoptosis and degeneration \textsuperscript{6,11}.

Acute kidney injury (AKI) or Epithelial cell injury, as a consequence of obstruction or otherwise, causes induction of Epithelial to mesenchymal transition (EMT), and regional fibroblast generation \textsuperscript{14}. Many researchers have highlighted the possible role of EMT in the
advancement of fibrosis after kidney injury\textsuperscript{14,15}. EMT is conveniently engaged by a mix of cytokines connected with proteolytic food digestion of basement membrane layers upon which epithelia live. The UUO model rests at the user interface between AKI and also CKD. The severe nature of the insult brings about AKI. Perseverance of the obstruction leads to histological changes and of CKD in one to 2 weeks\textsuperscript{16}.

Total UUO starts a quick series of events in the blocked kidney, leading within 24 h to minimize kidney blood flow and also glomerular filtering rate\textsuperscript{6}. This is adhered to within several days by hydronephrosis, interstitial inflammatory infiltration, and also tubular cell death attributable to apoptosis and necrosis. Tubular epithelial cell death is triggered by a variety of stress factors resulting from UUO, including anemia, hypoxia, oxidant injury, and also axial strain caused by tubular dilatation\textsuperscript{13}.

- Management of UUO

**Radiological diagnosis**

Diagnostic procedures are the first line management in all internal diseases, therefore, we need to discuss the roles of radiology in the diagnosis of UUO. El-Ghar et al\textsuperscript{17} evaluated the role of Magnetic Resonance Urography (MRU) and kidney scintigraphy for the physiological and also useful evaluation of blocked kidneys. An overall of 96 patients with compromised renal function participated in the research study. All included patients underwent improved MRI for kidney scan. Diagnosis of blockage was validated by ureterogram or endoscopy and/or open surgery (gold criterion). Anatomically, MRI found the reason for the blockage in all kidneys with non-calcular obstruction (100% level of sensitivity) and also in 21 kidneys with calcular blockage (70% level of sensitivity).

When integrated with stomach X-ray and ultrasonography, the sensitivity of MRI in discovery of reason for obstruction was 97%. Functionally, as El-Ghar et al.\textsuperscript{17} discovered that the MRU is as precise as renal scan in examining kidney function and also could be used as a single modality for detecting obstruction in cases where patients would not be jeopardized as a result of kidney function contraindications\textsuperscript{17}.

**Pharmacological management**

Rapamycin, also known as sirolimus, is a trienemacrolide antibiotic, which demonstrates anti-fungal, anti-inflammatory, anti-tumor, and also potent immunosuppressive residential or commercial properties.

It is becoming a foundation for long-lasting immunosuppressive treatment in renal transplantation. With the unique inhibition of a multifunctional serine-threonine kinase, the animal target of rapamycin, rapamycin can control the cell expansion, distinction, survival, movement, and also development\textsuperscript{18}. Lately, it has been reported that rapamycin decreased the cell proliferation in cystic and also non-cystic tubules as well as substantially hindered renal enhancement and cystogenesis, as well as stopped the loss of kidney function in a rat version of polycystic kidney illness\textsuperscript{18}. Moreover, rapamycin has also been shown to prevent the inflammatory and cytotoxic response in murine macrophage cell line and in the rat model of experimental mesangial proliferative glomerulonephritis\textsuperscript{19}.

Recent study\textsuperscript{20} offered proof to sustain that rapamycin has an impressive anti-inflammatory ability. They have actually investigated the impact of rapamycin on UUO-induced renal fibrosis. UUO or sham-operated rats were randomly appointed to rapamycin or lorry as well as were eliminated on days 7 and also 14 after UUO or sham procedure. Rapamycin reduced gross-morphology and cross-sectional modifications in the blocked kidney dramatically. Rapamycin considerably blunted the boost in weight of the blocked kidney, obstructed kidney size, and the obstructed/non-obstructed kidney weight proportion (by 74.6, 42.8, as well as 61.6% on day 14, specifically, all P<0.01).

Ball games for tubular dilatation, interstitial volume, interstitial collagen deposition, as well as \(\alpha\)-smooth muscle mass actin (\(\alpha\)-SMA) after UOO were dramatically lowered by rapamycin. Rapamycin also decreased the number of infiltrative anti-ED1-positive cells and also the gene expression of changing growth variable (TGF)-\(\beta\)1 (84.8 as well as 80.2% on day 7) after UUO (both P<0.01). By dual immunostaining as well as Western evaluation, rapamycin obstructed
the TGF-β1-induced loss of E-cadherin expression and also a fresh increase in the expression of α-SMA in a dose-dependent manner. In conclusion, rapamycin dramatically attenuated tubulointerstitial damage in a UUO-induced rat model of kidney fibrosis, recommending that rapamycin could have the perspective to postpone the progression of tubulointerstitial renal fibrosis 

Surgical management of UUO:

The surgical procedure for creating an animal version of UUO is reasonably uncomplicated if done as a solitary operation in a grown-up rat. Morbidity, as well as death rate, will be decreased by utilizing a temperature-controlled operating table warmed to body temperature level, with the pet anesthetized with isoflurane/oxygen and with making use of a high-quality binocular microscopic lens to envision the operating area. This method is particularly important in mice (especially in the neonatal period), or if the obstruction is to be relieved or turned around by a succeeding operation. Ligation of the ureter is the method made use of most regularly.

Fistulae can create around the band, allowing pee to bypass the blockage (unpublished observations).

Bonds forming around the ligature could increase the difficulty in removing the ligature if healing is to be checked out: this is a greater issue with an adult than neonatal animals. Little vascular clips can be placed in the ureter, but this approach could hurt the ureter, or allow urine to pass otherwise shut appropriately. An item of silastic tubes can be folded up perpendicularly throughout the ureter to produce an obstruction, or the tubing can be slit and also fitted around the ureter longitudinally, forming a sleeve and creating a partial obstruction. Partial UUO could likewise be created by putting the ureter in a surgically developed slit in the underlying psoas muscle. This technique leads to variable (frequently very mild) degrees of partial blockage.

A more reproducible technique of creating variable, reversible partial UUO has been developed in the neonatal mouse. This involves the positioning of a fine band around the ureter and a piece of stainless-steel cord of known diameter, which has been put parallel to the ureter. After ligation, the cable is eloped, leaving a partial blockage with the wanted luminal size. This ligature can be removed at various intervals, allowing the study of healing of the sores.

CONCLUSION

Speculative UUO is an animal version of the human condition, which duplicates not only human-ureteral-tract-obstruction-induced kidney injury, it is largely held that CKD happens consequently of the procedure of damaging fibrosis. UUO is a well-described model of renal fibrosis and also because of this is thought about a design of CKD. Additionally, several crucial procedures in the pathogenesis of AKI as well as CKD, such as tubular cell injury, interstitial inflammation and fibrosis.

Management approaches of UUO can be through several steps first the diagnostic management which will indicate the need of the following two steps of treatment whether pharmacological, as rapamycin substantially undermined kidney interstitial damages in a UUO-induced rat version of renal fibrosis, suggesting that rapamycin could have the possibility to postpone the progression of the transformation of renal mesenchymal and kidney fibrosis of surgical treatment.

REFERENCES


