Sildenafil Citrate Can Reduce Postobstructive Diuresis During Electrolyte Excretion After the Management of Bilateral Ureteral Obstruction in a Rat Model

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Abstract

Purpose: The current study aimed at assessing the effect of oral sildenafil citrate on postobstructive diuresis and urinary electrolyte changes after the management of bilateral ureteral obstruction (BUO).

Methods: Twenty-eight adult Munich-Wistar male rats were subjected to the surgery-induced BUO for 24 hours. In intervention group (n = 14) sildenafil citrate 200 mg/kg food was administered from the operation day for 72 hours, while in control group (n = 14) no drug was used. Daily urinary volume, urine sodium (mEq/day), potassium (mEq/day), and osmolality (mOsm/kg H2O) were measured on the day before BUO and for 2 days after BUO release.

Results: In comparison to the control group, postobstructive diuresis was significantly lower in the sildenafil group (P value < 0.001), while urinary sodium and osmolality were both preserved better in this group (P value < 0.001); however, there were no significant difference in potassium level between the groups (P value = 0.285).

Conclusions: Oral sildenafil citrate significantly decreased postobstructive diuresis and preserved urine sodium and osmolality near normal. These 2 effects mean a better preservation of renal function, while preventing massive postobstructive diuresis.

Keywords: Bilateral Ureteral Obstruction, Sildenafil Citrate, Postobstructive Diuresis

1. Introduction

The main renal function is to excrete creatinin and potassium, while preserving sodium and water. This normal renal task can be disturbed when there is an obstruction in the urinary tract, which is called obstructive uropathy (OU). OU is a rare, but curable, cause of acute renal failure; however, after releasing a long lasting obstruction, the kidney response to the volume and the toxin overload maybe destructive per se. This state leads to excretion of a huge amounts of water and sodium and is called pathological postobstructive diuresis, which can cause severe hyponatremia and death. Taking this issue into consideration, measuring urinary output as the main indicator of this phenomenon is outmost important, and several protocols are introduced to overcome this renal malfunction.

During OU, renal blood flow and glomerular filtration rate (GFR) may decrease, causing tubular abnormalities such as urinary concentration defect (1). Moreover, expression of aquaporin is downregulated after long lasting bilateral ureteral obstruction (BUO), suggesting a contribution to long-term polyuria and impairment of urinary concentrating capacity that happen after relief of OU (2-10).

It is reported that sildenafil citrate, a phosphodiesterase type 5 inhibitor (PDE5 inhibitor), restores the downregulated aquaporin channels in kidney caused by OU. This may theoretically improve renal function in such patients. The current study aimed at assessing the effects of sildenafil citrate on renal function after BUO release.

2. Methods

A total of 28 male adult Munich-Wistar rats, weighing 200 - 250 g, were enrolled in the current study. The surgery-induced BUO was performed on all rats and after 24 hours
BUO was released. In the intervention group (n = 14), sildenafil citrate (200 mg/kg food) was given to the rats from the surgery day, while the control group had just standard rat food. Daily urinary volume, urine sodium, potassium, and osmolality were measured every 24 hours from the day before the surgery to the second postsurgical BUO releasing day.

2.1. Induction of BUO

The animals were anesthetized intraperitoneally with 50 mg/kg sodium thiopental. With a midline lower abdominal incision, ureters were removed. Two ligatures, 5 mm apart, were placed in the mid part of the ureter over a section of polyethylene tubing (PE-70) around each ureter. The ureters were occluded by tightening the tubing with a 3-0 silk ligature. After recovering from anesthesia, the animals were returned to their metabolic cages with free access to water and standard rat food based on the specific protocol for each group.

2.2. Collection of the Urine

To collect urine samples and measure daily urine production, metabolic cages were used. The daily urinary output (mL/day) was measured for each rat and urine samples were sent to the laboratory to be evaluated for osmolality and sodium/potassium measures.

2.3. Urine Osmolality

Urine osmolality was determined using a freezing point osmometer (Wescor 5520, USA) and expressed as mOsm/kg H$_2$O.

2.4. Statistical Analysis

SPSS version 22 was used for the statistical analyses. The qualitative data were expressed as mean ± standard deviation (SD) and the quantitative data were shown in relative frequency. Based on the data volume (lower than 30) of each group, the Kolmogorov-Simonov test was used to examine the normal distribution of data. The nonparametric Wilcoxon signed rank test was used, before and after the BUO release, to evaluate the intergroup differences for the parametric variables, while nonparametric Mann-Whitney U test was used to compare intragroup differences for the same variables. P value < 0.05 was considered as the level of significance.

3. Results

BUO is associated with decreased urine osmolality improved with Sildenafil ingestion. Results of urine osmolality and Na/K volumes are presented in Table 1. BUO is a critical condition and may be associated with prolonged subnormal ability to concentrate urine. The current study found an impaired urinary concentrating capacity 48 hours after the release of BUO (group 1), compared with that of the BUO SIL rats (group 2) (P value < 0.001) (Figure 1).

![Figure 1. Mean Urine Osmolality Before and 48 Hours After the Release of BUO](image1)

BUO is associated with increased urine sodium and volume amounts improved with sildenafil ingestion. Rats were studied from 24 hour before BUO to 48 hours after releasing the obstruction; results of the experiments presented a significant increase in urine Na and volume amounts (group 1), compared with those of the BUO SIL rats (group 2) (P value < 0.001) (Figures 2 and 3).

![Figure 2. Mean Urine Volume Before and 48 Hours After the Release of BUO](image2)
### Table 1. Urinary Parameters Before and After the Release of BUO in the Study Groups

| Urinary Parameters | Intervention n = 14 | Control n = 14 | P Value | Intervention n = 14 | Control n = 14 | P Value |
|--------------------|---------------------|----------------|--------|---------------------|----------------|--------|
| **K, mEq/d**       | Before BUO 2.14 ± 0.37 | After the Release of BUO 2.14 ± 0.22 | -0.414 | 0.679 | Before BUO 2.16 ± 0.40 | After the Release of BUO 2.27 ± 0.41 | -2.068 | 0.059 | 0.285 |
| **Na, mEq/d**      | Before BUO 1.01 ± 0.31 | After the Release of BUO 1.07 ± 0.23 | -0.794 | 0.427 | Before BUO 1.01 ± 0.31 | After the Release of BUO 2.74 ± 0.93 | -3.301 | 0.001 | 0.001 |
| **Osmolality**     | Before BUO 941.50 ± 66.71 | After the Release of BUO 942.21 ± 60.94 | -0.974 | 0.330 | Before BUO 943.14 ± 70.89 | After the Release of BUO 606.64 ± 84.10 | -3.296 | 0.001 | 0.001 |
| **Volume, mL/d**   | Before BUO 14.64 ± 1.74 | After the Release of BUO 16.57 ± 2.21 | -2.325 | 0.200 | Before BUO 14.50 ± 0.44 | After the Release of BUO 22.43 ± 0.37 | -3.306 | 0.001 | 0.001 |

Figure 3. Mean Urine Sodium Before and 48 Hours After the Release of BUO

### 4. Discussion

Obstructive uropathy is a condition with many different physiological and pathological consequences that can cause transient or permanent functional disability to kidneys. This impairment in some degrees depends on the duration of obstruction (acute or chronic) (11, 12).

In OU, the initial event is vasodilatation, which lasts 90 minutes; then, more prominent drops occur in renal blood flow, that maybe attributed to release of strong vasoconstrictors, angiotensin II, and thromboxane A2 (7,11, 13). During this period, intraurethral pressure is above normal range and intrarenal blood-flow is redistributed from medullar to cortex (13). These events under BUO conditions can increase renal vascular resistance and then, reduce GFR (14,15).

Following the treatment of OU, some physiologic changes, such as intravascular expansion and subsequently atrial natriuretic peptide (ANP) secretion, occur that finally leading to efferent and afferent arteriole vasodilatation. After these changes, with increasing glomerular capillary pressure and reducing renin production, an absolute increase occurs in GFR, which as a clinical finding, is referred to postobstructive diuresis. As expected, in unilateral ureteral obstruction elimination of harmful vasoactive agents by contralateral kidney through secretion of ANP attenuates or even prevents these harmful changes (7, 11, 15).

In association with postobstructive diuresis, some electrolyte changes are occurred from which the most important ones are the alteration in urinary excretion of sodium and potassium, and changes in urinary osmolality. As expected, in patients with unilateral ureteral obstruction and normal contralateral kidney, compensation mechanism can overcome, though partially, and reabsorb water and electrolytes (12,16).

Under normal conditions, any changes in serum osm-
lality or effective circulating volume may directly affect the secretion of vasopressin from posterior hypophysis. The vasopressin is an antidiuretic agent that acts on the receptors of basolateral membrane of kidney collecting duct cells, which causes the fusion of vesicles containing aquaporins into apical cells of the collecting ducts. As a consequence, an impermeable membrane converts into a water-permeable membrane, through the action of aquaporins AQP2, AQP3, and AQP4 (8-10).

In patients with BUO caused by ischemia, vasopressin hyposensitivity, and downregulation of aquaporin receptors, the low osmolality urine with increased sodium concentration was excreted, which explains long-term polyuria and urinary concentrating defects (2-10, 17).

The current study showed the effects of sildenafil in a BUO model. The results of the current study demonstrated that sildenafil administration prevented postobstructive diuresis 48 hours after the release of a 24-hour BUO. Treatment with sildenafil prevented BUO-related hemodynamic effects, such as decreased urine osmolality and polyuria. In sildenafil-treated rats, urinary osmolality was better than that of the BUO group (group 1). These deleterious effects could be reversed by sildenafil administration and it may imply the clinical application of sildenafil in acute obstructive nephropathy. Further human models are required to better understand the reported mechanism.

This study had several limitations; for instance, the limited number of cases and no enrollment of the sham group. Variables in the study were limited to some urinary parameters, without any serum parameters such as serum sodium and potassium. Maybe the most appropriate way to collect and measure urinary output hourly (rather than daily) in order to confirm pathological post-obstruction diuresis, but due to lower level of urinary output in rats (~0.5-0.75 ml/hour), it was very difficult to collect and measure urinary output hourly; moreover, to the authors’ best knowledge, there is no reliable definition for postobstructive diuresis in rats.

4.1. Conclusions

Sildenafil treatment significantly improved urinary concentration by reversing the BUO-induced downregulation of AQP2. These findings have significant clinical implication in treating obstructive nephropathy.

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