Oral Desmopressin in the Management of Adults with Nocturia

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Abstract

We investigated the efficacy of oral desmopressin in the treatment of adult nocturia. In an analytical study between 2007-2009 in Zahedan-Iran, Thirty patients ≥55 years with verified nocturia (≥two voids/night) were enrolled. Patients with a history of an obstructive cause of nocturia, those with diseases getting worse by the anti-diuretic affect of desmopressin and those with well-defined curable causes (e.g. cystitis) were excluded. Patients received 0.2 mg of oral desmopressin at bed time for a period of 3 weeks. p < 0.05 was taken as the significant level. All 30 patients enrolled completed the trial. Fourteen (47 %) patients receiving desmopressin had fewer than half the number of nocturnal voids relative to base line (p < 0.001). The mean number of nocturnal voids decreased from 4.6 to 2.4 (p < 0.001). Fatigue (10%), headache (3%) and dizziness (3%) were reported. All adverse events were of mild intensity and there were no instances of hyponatremia. Oral desmopressin is an effective treatment in patients with nocturia and is well-tolerated.

Keywords: Nocturia, Desmopressin, Arginine Vasopressin

1. Introduction

Nocturia is defined as waking at night to void [1]. Although by definition even a single episode of awakening to urinate is nocturia, most clinicians consider nocturia to have significance when a patient voids two or more times nightly [2]. Nocturia is a very common and bothersome symptom [3]. The prevalence of nocturia increases with age and affects quality of life in both men and women [4]. Rising at night to void increases the risk of personal injury, particularly in the elderly where the condition is associated with a greater risk of falls [5]. Nocturia may be due to either polyuria, nocturnal polyuria or low bladder capacity [6]. As the diurnal variation in AVP release is absent in many older subjects [7], nocturnal polyuria, or the over-production of urine at night, is an important cause of nocturia [8], that may be due to age-related changes in the secretion and action of arginine vasopressin (AVP) [9,10]. Treatments currently licensed to improve bladder function do not control nocturnal polyuria [11]. Desmopressin (1-deamino-8-D-arginine vasopressin, ddAVP), a synthetic analogue of AVP is used for treatment of adult nocturia [12], and when administered at bed-time, decreases night-time urine production [13]. Effectively treating nocturia with desmopressin has been shown by some others [11,14,15]. The primary aim of the present study was to investigate the efficacy of oral desmopressin in a 3-week treatment period, in reduction in the number of voids at night.

2. Materials and Methods

In a analytical study, included 30 patients aged ≥ 55 years between 2007-2009 in Zahedan-Iran, with complaint of two or more episodes of nocturnal voids during night (night is defined as the period between going to bed for sleeping and waking for rising) and nocturnal diuresis > 0.9 ml/min [5]. Screening included a precise medical history (including other lower urinary tract symptoms, past medical diseases, fluid intake and medications), physical examination (including measuring supine and standing blood pressure), serum electrolytes, renal function tests and urine analysis. The upper and lower urinary tract ultrasonography with careful attention to post-void residual urine volume was also done. Patients with a serum sodium level below the normal range,
epilepsy, clinically significant renal, hepatic or cardiovascular disease, uncontrolled hypertension, diabetes insipidus, multiple sclerosis, primary polydipsia, genitourinary tract infection, urge incontinence, and untreated BPH (benign prostatic hyperplasia) were excluded. The number of nocturnal voids and nocturnal urine volume were recorded in the questionnaire before commencing the treatment. The patients received 0.2 mg of oral desmopressin (Minirin®, Ferring pharmaceuticals, Sweden) at bed-time for three weeks.

The patients were instructed to drink only to satisfy their thirst until 8 hours after taking the tablets and to stop the treatment and inform their doctor if side-effects such as headaches not relieved by acetaminophen, vision problems, faster heart rate, rapid weight gain, nausea and vomiting, confusion or seizures occur. All patients were visited at the end of the first week of treatment to check their serum sodium levels and probable adverse affects. The primary efficacy endpoint was the proportion of patients who had a reduction by more than half in the mean number of nocturnal voids after treatment compared with baseline. Changes in nocturnal urine volume and number of nocturnal voids, serum sodium levels, incidence of adverse events and patient’s satisfaction of desmopressin treatment were also assessed. Statistical analysis was performed by SPSS software (version 15) and results were presented using P values based on the paired T-test. \( p < 0.05 \) was taken as the significant level.

3. Results

Of the patients who reported two or more voids per night, 32 were screened and enrolled. Two patients withdrew consent. The study population comprised 26 men (87%) and 4 women (13%), with a median age of 63.5 (55 - 78) years. A clinical response defined as fewer than half the mean number of nocturnal voids after treatment compared with baseline, was achieved by 14 (47%) patients (\( p < 0.001 \)). If a threshold of \( \geq 40\% \) reduction in nocturnal voids was supposed as a response instead of \( \geq 50\% \), 22 (73%) patients obtained a response to desmopressin. The mean number (SD) of voids per night reduced from 4.6 (0.9) at the baseline to 2.4 (0.7) at the end of treatment period (\( p < 0.001 \)) (Figure 1). And the mean nocturnal urine volume decreased from 640 ± 107.6 ml to 395.3 ± 98.8 ml (\( p < 0.001 \)) (Figure 2). There was a reduction of \( \geq 20\% \) in nocturnal urine volume in 27 (90%) patients. In all, 3 (10%) fatigue, 1 (3%) dizziness and 1 (3%) headache were reported. All of adverse events were of mild intensity. No instances of serious side-effects were detected. The mean serum sodium level decreased from 138 to 136 mmol/L during the first week of treatment (\( p = 0.317 \)) (Figure 3). Serum sodium level <
formed that it is unnecessary to bear this problem any
more. The present study showed desmopressin to have
beneficial effects on nocturia and is well-tolerated; how-
ever, serum sodium testing is suggested in all patients
before and after a few days on treatment.

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130 mmol/L did not occur. Two patients had a decrease
of ≥5 units in serum sodium level during the first week
of treatment, but both remained asymptomatic during the
treatment period and hyponatremia was not reported at
all. Twenty-four (80%) patients were completely satisfied
with the treatment, opposed to 5 (17%) relatively satis-
fied and 1 (3%) not satisfied.

4. Discussion

Adults, particularly the elderly with frequent voiding
episodes at night, often suffer from poor sleep and con-
sequently they are often tired during the day, resulting in
a poor quality of life [16]. In older patients with nocturia,
the day: night urinary output ratio is reduced, and in such
patients there is an increased frequency of nocturnal
micturition. Plasma vasopressin is at undetectable levels
in patients with nocturia, reduces nocturnal diuresis and
the number of nocturnal voids. In summary, oral desmo-
pressin given at bed time is associated with a decrease in
nocturnal urine output. The main aim of the present study
was to investigate the efficacy of oral desmopressin in reduc-
ing episodes of nocturnal voids, which is the main complaint
of subjects with nocturia.

The threshold chosen for clinical response was a ≥
50% reduction in nocturnal void based on previous stud-
ies [11,18]. The mean number of nocturnal voids reduced
by 48% in patients on desmopressin compared with base-
line. In accordance with previous studies, [14,15,19] the
decrease in nocturnal diuresis with desmopressin treat-
ment was associated with reduction in nocturnal voids.

A slight decrease in serum sodium levels during
treatment was seen, which was not statistically signifi-
cant (p = 0.317). The study showed that adverse-effects
associated with desmopressin treatment were of mild
intensity and infrequent, similar to what was previously
reported [11]. Caution is warranted in interpreting the
results as there was no placebo arm. However, the results
accord with those in other trials of desmopressin for this
indication. A potential source of efficacy bias may be the
safety instructions given to patients, to drink only when
thirsty during the night, which could affect fluid intake
and thus urine volume, and the number of nocturnal
voids. In summary, oral desmopressin given at bed time
in patients with nocturia, reduces nocturnal diuresis and
nocturnal voids compared with the baseline. As desmo-
pressin is a potential therapeutic choice for subjects with
nocturia, patients not responding to general advice and
conventional treatments, and those who consider this
suffer as a normal consequence of aging, must be in-
formed that it is unnecessary to bear this problem any


