Case report: Desmopressin and somnambulism

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ABSTRACT

Background: Nocturnal enuresis is caused by a mismatch between nocturnal urine production and bladder capacity. Together with a presumed decreased arousability, this results in an inability to awaken in response to a full bladder. According to recent findings, disrupted sleep might play a role in the pathophysiology of nocturnal enuresis. Case: A 7-year-old boy was diagnosed with primary nocturnal enuresis caused by nocturnal polyuria and a concomitant parasomnia, somnambulism. A polysomnographic study was performed before treating the nocturnal enuresis with the oral lyophilisate formulation of desmopressin (melt). After 1 day of treatment, both nocturnal enuresis and somnambulism disappeared. Treatment was ceased after six months. One week later, the child started to produce more urine and redeveloped nocturnal polyuria. Somnambulism reappeared followed by nocturnal enuresis two weeks later. The same treatment protocol was started up, resulting in disappearance of both nocturnal enuresis and somnambulism. Conclusion: This case report documents the beneficial effect of desmopressin melt on both nocturnal enuresis, caused by nocturnal polyuria, and a concomitant parasomnia in particular somnambulism.

Keywords: Children; Enuresis; Bedwetting; Somnambulism; Sleep; Sleepwalking; Desmopressin; Nocturnal Polyuria; Parasomnia

1. INTRODUCTION

Monosymptomatic nocturnal enuresis (MNE) is caused by a mismatch between the nocturnal diuresis volume and the bladder capacity. It was widely accepted that a decreased arousability, provokes an inability to awaken in response to a full bladder [1]. According to recent findings, disrupted sleep rather than decreased arousability might play an important role in the pathophysiology of nocturnal enuresis (NE) [2]. Sleep of children with NE is characterized by sleep fragmentation. Even more, children with NE clearly have more daytime sleepiness and suffer from sleep deprivation [3]. Mahler et al. demonstrated that sleep deprivation might result in increased nocturnal diuresis volume [4]. During polysomnography, our study group observed increased periodic limb movements (PLMS) and cortical arousals in children with refractory NE [5]. The association among PLMS, arousals and NE still remains unclear.

Nocturnal polyuria (NP) is related to a deficient vasopressin (AVP) secretion overnight. Therefore, the synthetic AVP analogue desmopressin is widely used to treat children with MNE [6]. Its antidiuretic action results from an increased renal reabsorption of water, leading to a reduced volume and more concentrated urine. Desmopressin is recommended by the International Consultation on Incontinence (Grade A, Level 1) for the treatment of MNE.

Somnambulism (sleepwalking) is a well-known sleep disorder, classified as a parasomnia. Electro-encephalographic recording during this event is characterized by a cortical arousal that emerges during the first third of the night when slow wave sleep is present [7,8]. This sudden change in EEG frequency can be caused by PLMS [8] or obstructive sleep apnea [9] or can occur due to brain maturation [10]. Hublin et al. demonstrated retrospectively by questionnaires that 7.7% men and 9.7% women reported sleepwalking “often” or “sometimes”, 17.9% of the men and 17.0% of the women reported “a few times” sleepwalking during childhood (4 - 15 years) [11]. According to Esposito et al. somnambulism is far more presented in children with NE (22.1%) compared to healthy controls (6.7%) [12].

In the following case report we demonstrate a comorbidity of NE and the parasomnia sleepwalking, both resolved by an effective treatment of enuresis.

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2. CASE REPORT

The boy was six years old when his parents consulted a child nephrologist. He experienced diurnal incontinence once a month, daily wet urine spots in underpants, urge incontinence with squatting and giggling incontinence. Moreover, NE was present almost every night. Home recordings revealed that the child had small voided volumes, although drinking schedule was appropriate. NE usually took place during the first hours of the night with documented NP.

After successful treatment of the daytime symptoms (urotherapy, oxybutinin), the picture of NE changed. The boy started “bed-wetting” during sleepwalking episodes. His mother described it as follows: “it looks like he was searching for a toilet and when he thought he found one, he started to pee”. The combination of sleepwalking and “wetting” persisted for a year when the boy and his mother consulted again. Screening showed no longer daytime symptoms. There was a normal bladder volume, but NP still persisted.

The boy got subsequent a video-polysomnographic study. Sleepwalking was not present the night of the polysomnography, but PLMS were clearly observed. After the polysomnography 120 µg of the oral lyophilisate formulation of desmopressin (melt) was given. Not only NP disappeared but both sleepwalking and NE ceased. One night he forgot to take his medication and that night he experienced again NE.

After six months of successful treatment, desmopressin melt was stopped. One week later, the child started to produce more urine and developed NP. His mother reported a more disrupted sleep. Two weeks later the boy started sleepwalking again and after three weeks NE during sleepwalking recurred. He was treated again with desmopressin melt and once more both sleepwalking and NE stopped.

3. DISCUSSION

The recurrence of both enuresis and somnambulism due to the discontinuation of desmopressin melts and the resumption of benefits due to the restart of the drug suggests that the desmopressin therapy not only has an antidiuretic effect [13], but is also responsible for the disappearance of the somnambulism. The positive effect on the somnambulism is likely to be related to the antidiuretic effect. We speculate that the boy suffering from NP without desmopressin, experienced a sense of a full bladder provoking cortical arousals in his sleep. These arousals result in a parasomnium, in this case: somnambulism. This incomplete awakening while searching for a toilet, was previously suggested by Yeung et al. [2]. By using desmopressin, the child no longer had NP and therefore did not experience arousal stimuli from the bladder while sleeping. Probably, as a result, no interrupting arousals caused the parasomnium at that time.

However, a direct effect on the central nervous system (CNS) by desmopressin can not entirely be excluded [14, 15]. The major mechanism of desmopressin in NE results in an increased urinary concentration and subsequent decreased diuresis-rate. Even though, the exact mechanism of our observation is not yet clear.

This case study emphasizes again the important comorbidity of a disrupted sleep (somnambulism, PLMS, cortical arousals) and NE together with NP. Yet, a prospective study is needed to further elucidate the pathophysiologic mechanisms involved.

REFERENCES


ABBREVIATIONS

MNE: monosymptomatic nocturnal enuresis;
NE: nocturnal enuresis;
PLMS: periodic limb movements of sleep;
NP: nocturnal polyuria;
AVP: arginine vasopressin;
EEG: electroencephalography;
CNS: central nervous system.