The Problem of Polypharmacy in Female Patients with Overactive Bladders – Cross-Sectional Study in a Specialist Outpatient Department

Das Problem der Polymedikation bei Patientinnen mit Überaktiver Blase – Querschnittsuntersuchung einer Spezialambulanz

Key words
- overactive bladder
- polypharmacy
- urinary incontinence
- drug-drug interactions

Schlüsselwörter
- Überaktive Blase
- Polymedikation
- Harninkontinenz
- Medikamenteninteraktionen

Abstract

Background: The consumption of certain drugs can cause urinary incontinence. The aim of this study is to determine the frequency of consumption of drugs that can favour incontinence, the incidence of polypharmacy and the incidence of potentially dangerous drug-drug interactions in female patients suffering from overactive bladder (OAB) who presented to a urogynaecological outpatient department.

Methods: We undertook a retrospective case series study. The data from 100 female patients who attended the urogynaecological outpatient department of the Vienna General Hospital [VGH; Allgemeinen Krankenhauses Wien (AKH)] in the period from 20.07.2010 to 30.08.2011 were evaluated. The patients suffered either from an OAB or mixed incontinence with predominantly urge components. Among other factors, we were interested in the drugs taken for longer periods of time as well as the general and the urogynaecological case histories. 15 parameters were recorded: age, BMI, menopausal status, parity, pelvic organ prolapse, DIAPPERS criteria (delirium, infection (urinary), atrophic urethritis and vaginitis, pharmaceuticals, psychological disorders (especially depression), excessive urine output, restricted mobility, stool impaction), drug side effects and drug-drug interactions. A descriptive statistical analysis was performed. The drugs were checked with the help of a drug information system (Intranet-KH [V 6.0]). Of particular interest was the consumption of drugs that could favour urinary incontinence as an adverse side effect. In addition the frequency of polypharmacy and the frequency of potentially health-threatening drug combinations were registered.

Results: 57% of the patients consumed at least one drug that could reinforce urinary incontinence. The frequency of polypharmacy was 38%. In 45% of the patients the possibility for health-threatening drug-drug interactions was the consumption of drugs that could reinforce urinary incontinence as an adverse side effect.

Zusammenfassung

Hintergrund: Die Einnahme bestimmter Medikamente kann die Ursache für Harninkontinenz sein. Das Ziel dieser Studie war die Erhebung der Häufigkeit von Medikamenteneinnahmen, die eine Inkontinenz begünstigen können, die Häufigkeit von Polymedikation sowie die Häufigkeit von potentiell gefährlichen Medikamenteninteraktionen bei Patientinnen einer urogynäkologischen Spezialambulanz, die an Überaktiver Blase (UAB) leiden.


Ergebnisse: 57% der Patientinnen nahmen mindestens ein Medikament ein, das eine Harninkontinenz verstärken kann. Die Häufigkeit von Poly-
Introduction

The overactive bladder (OAB) syndrome is a form of incontinence that has a reversible cause in some cases. One of the most important reversible causes is the consumption of drugs. Certain drugs can trigger incontinence as an adverse side effect and also interactions between drugs can favour the occurrence of OAB [1].

This study therefore examines the relationship between polypharmacy and OAB. Of particular interest were the frequency of consumption of drugs with a potential incontinence-promoting activity, the frequency of polypharmacy and an assessment of the frequency of potentially harmful drug-drug interactions in a population of female patients of the Vienna General Hospital who were suffering from OAB.

The average prevalence of OAB among European women aged 40 years or more amounts to 17% [2]. The prevalence increases with increasing age [2–4]. In the group of over 60-year-olds the prevalence amounts to 45% [3]. The quality of life of the afflicted persons is appreciably impaired and therapy for overactive bladder syndrome leads to a significant improvement in the quality of life [5].

Polypharmacy is one of the reversible causes of overactive bladder [6]. We speak of polypharmacy when five or more drugs are taken regularly [7]. Numerous drugs can favour urinary incontinence and drug-drug interactions can impact on the lower urinary tract [3,8]. Especially elderly persons require polypharmacy [9].

The “overactive bladder” syndrome is defined by the symptom complex polakriopia (≥8 micturitions/24 hours with a normal amount of urine [up to 2.8 l/24 h]), imperative need to urinate (sudden onset of need to urinate accompanied with the danger of involuntary leakage), nocturia (the awakening from sleep at night to pass urine) as well as premicturitional urine leakage (leakage of urine prior to intended urination) [6,10,11].

The clinical picture of overactive bladder can, but need not, be accompanied with incontinence [10].

In order to make the diagnosis of OAB, possibly underlying local, metabolic, neurological or endocrinological pathologies must be excluded [11–13].

Reversible causes

Since the frequency of urinary incontinence increases with age [2,4] and elderly people above all often suffer from several illnesses that can themselves also trigger the symptom of incontinence, the possibility of a temporary and, in such cases, reversible overactive bladder must be taken into account and checked [14].

The possible causes of reversible incontinence were summarised in 1987 by Resnick with the mnemonic DIAPPERS (diaper) [1]. DIAPPERS is an acronym for the following terms: delirium, urinary infection, atrophy of urogenital tract mucous membranes, psychogenic reasons, pharmacotherapy, excessive urinary out-

put, restricted mobility and stool masses in the rectum (constipation). All of these factors can lead to an incontinence that, however, can be markedly alleviated or possibly even completely eliminated by successfully addressing these factors [1].

In particular, the impact of drugs that can trigger incontinence as a side effect or by interactions with other drugs should not be underestimated [8,12].

Antihypertensive agents such as diuretics, alpha-receptor blockers, calcium channel blockers, beta-receptor blockers and ACE inhibitors can favour incontinence for various reasons. The mechanisms have been described in many different studies and vary widely. Diuretics increase urine production [15], alpha-receptor blockers reduce the pressure and thus the resistance of the urethra [16], calcium channel blockers lead to polyuria [17] and beta-receptor blockers result in, via a dominance of the parasympathetic nervous system on the detrusor, to an increased contraction readiness of the bladder wall [18]. Also ACE inhibitors can trigger stress incontinence by way of increased coughing that increases the intra-abdominal pressure [19]. Angiotensin-2 blockers have been described as a possible trigger in one study although, according to the authors, also other antihypertensive agents could have been responsible as possible confounding factors [20].

In addition, a relationship between the consumption of oral oestrogens and an increased risk for the occurrence of incontinence has been found, although the exact mechanism is still unknown [21].

Psychoactive drugs (such as sedatives, hypnotics and antidepressants) can lead to so-called overflow incontinence through the two mechanisms – sedation and urinary retention. Anaesthetics lead to constipation and thus to incontinence [22,23]. It is assumed that atypical neuroleptics, by way of an elevated cholinergic stimulation of the detrusor, can lead to a higher micturition frequency and stronger urge to urinate [24,25].

Anticholinergic drugs, antihistamines and tricyclic antidepressants as well as beta-receptor agonists, anticholinergic drugs for Parkinson’s disease and levodopa can trigger overflow incontinence though urine retention [18,26,27].

Anticonvulsant agents that reduce bladder outlet resistance and thus facilitate micturition can, under certain circumstances, worsen an already existing incontinence [27].

Certain cholinesterase inhibitors that are used in the treatment of dementia can lead to an increased risk for the occurrence of urinary incontinence through an inhibition of the degradation of acetylcholine which, in turn, causes a stronger activation of the detrusor [28]. Prokinetics can also favour incontinence through so-called detrusor instability [29].

Other drugs associated with urinary incontinence are laxatives and muscle relaxants [23].

Table 1 summarises the most important drugs that can favour urinary incontinence.
Also, genital prolapse or, respectively, pelvic organ prolapse (POP) is an independent risk factor for the development of an overactive bladder [30] and was therefore recorded. The aim of the present study was to record and describe the drug consumption habits of female patients with OAB in an urogynaecological specialist outpatient department. We examined
1. the incidence of consumption of drugs that favour incontinence,
2. the incidence of polypharmacy and
3. the incidence of disadvantageous drug-drug interactions.

Materials and Methods

Study design

The present study consists of a retrospective case series design. The study was approved by the ethics committee of Vienna Medical University (No. 19/2013).

The data of 100 consecutive female patients who presented to the urogynaecological outpatient unit of the gynaecology department at Vienna General Hospital between 20.07.2010 and 30.08.2011 were recorded. Inclusion criteria were: age of 18 years or more and the diagnoses “overactive bladder” or “mixed incontinence”. In cases with the diagnosis “mixed incontinence” the predominance of a strain urinary incontinence component represented an exclusion criterion. Further exclusion criteria were the diagnoses of “pure strain urinary incontinence” and “overflow bladder”. The foundations for data collection were the computerised outpatient records in the hospital’s own information systems KIS and AKIM. By means of the stored data the general, gynaecological and urogynaecological case histories including menopausal status, the results of gynaecological and physical examinations as well as DIAPPERS criteria for each and every patient were recorded. All data were extracted from the computerised outpatient files.

Altogether 15 parameters were acquired. These included: age, BMI, menopausal status, parity, pelvic organ prolapse, DIAPPERS criteria (delirium, urinary tract infection, atrophy of the urogenital tract, psychological factors, pharmacotherapy, excessive urine output, restricted mobility and constipation), side effects of the consumed drugs and drug-drug interactions.

Only the DIAPPERS criterion “delirium” could not be extracted from the stored patient records, however, it is assumed that any possible indications for the patient’s disorientation would have been documented by the treating physician. The data were entered into a previously prepared data sheet. If details of medication were lacking, the respective patient was excluded from the study.

The recorded drugs were classified into 35 categories. Each drug was assigned to just one category except when the formulation contained two active substances.

The recorded drugs were evaluated with the help of the drug information system “Medis” (Medis Intranet-KH [V 6.0]). “Medis” is an information system in the Intranet of Vienna General Hospital that provides specialist pharmaceutical information and interaction analyses for drugs. For each drug adverse effects on symptoms that could favour urge incontinence were looked for. These symptoms are pollakiuria, nocturia, incomplete emptying of bladder and urinary incontinence. Each drug exhibiting one or more of these side effects was recorded separately. Furthermore, possible interactions between the drugs taken by an individual were recorded under the function “Rp”. All possible interactions between drugs were documented and summarised in superordinate categories of the affected organ system. The frequency with which each interaction occurred in the described population was recorded. In addition the frequency with which an individual organ system was affected by drug interactions was analysed.

Furthermore, the drugs consumed by the patients were compared with those on a list compiled in cooperation between a...
group of urogynaecologists and a clinical pharmacologist specialising in geriatric medicine. This list contained drugs that favour urinary incontinence and the responsible mechanisms of action (Table 1).

**Results**

We analysed the clinical records of 765 consecutive female patients. Of them 647 were excluded on the basis of their diagnoses, of the 118 patients with the diagnosis OAB 18 were excluded from the analysis because their drug consumption was not or was not completely documented. The data of 100 patients were evaluated.

**Patient characteristics**

Of the 100 patients, 55% were older than 59 years (average ± standard deviation: 59 ± 15, min: 18, max: 89). 73% were already post-menopausal and 25% had a BMI (kg/m²) of 30 or more. Genital prolapse or, respectively, pelvic organ prolapse is defined as lowering of the pelvic organs to just a few centimetres above the hymenal border which was thus designated as a pelvic organ prolapse. 53 patients did not have a genital prolapse. Further patient characteristics are listed in Table 2.

**Drugs**

83% of the subjects took at least one drug on a long-term basis. On average each patient took 4 drugs.

Antihypertensives were the most frequently consumed class of drugs. 41% of the patients took at least one antihypertensive agent per day, the most frequently consumed antihypertensives were beta-receptor blockers (18% of all patients).

The second most frequently consumed class of drugs were hormones. 31% of the patients regularly took hormones with thyroid hormones and local oestrogens being subsumed. Antidepressants occupied the third place among the consumed drugs. 29% of the patients regularly took an antidepressant.

57% of all patients took one or more drugs that could trigger or reinforce urinary incontinence (Fig. 1). Of these drugs antihypertensives and antidepressants were consumed most frequently.

The pathomechanisms that favour urinary incontinence are listed in Table 1. Most frequently taken were drugs that favour urge incontinence through the mechanism of incomplete bladder emptying. 49% of all patients consumed at least one drug that impaired complete bladder voiding thereby possibly leading to so-called “overflow incontinence”. Above all, drugs with anticholinergic activity that are frequently taken by the elderly often lead, on the one hand, through relaxation of the bladder muscles and, on the other hand, to a central nervous action and tranquillising effects to incomplete bladder emptying. Drugs that can trigger incomplete bladder emptying are antidepressants, neuroleptics, anticholinergic agents, antiparkinson drugs, opioids, antihista-mines, prokinetics, beta2-agonists and muscle relaxants [26]. 31% of all patients consumed drugs that have sedative and constipation-inducing effects and so favour urinary incontinence.

The third most frequent mechanism was increased urine output or polyuria that can occur after consumption of, above all, diuretics or calcium blockers [15,17]. 28% of the patients took drugs that increased the production of urine.

Further side effects of the consumed drugs were nocturia (26% of the patients) and polyuria (22% of the patients) that can be triggered, above all, by the consumption of antidepressants [32].

Fig. 2 shows the frequency of consumption of drugs that can favour urinary incontinence as referred to the various age groups. 38% of the patient took five or more drugs daily. Fig. 3 shows the incidence of polypharmacy referred to five different age groups.

Potentially harmful interactions between the consumed drugs were identified in 45% of the patients. 17% of all patients took combinations of drugs with which three or more harmful interactions between the individual drugs could occur.

In 20% of all patients interactions that could have an effect on blood pressure were identified. The most frequent possible interaction in this collective (15% of the entire collective) was a weakening of the blood pressure-lowering action of one drug by another.

Equally frequent (20%) were interactions that could have a reinforcing or a weakening effect on the action of certain other drugs. Most common was the simultaneous consumption of drugs that could reduce the action of thyroid hormone replacement drugs such as, for example, iron and calcium formulations.

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**Table 2** General patient characteristics.

<table>
<thead>
<tr>
<th>General patient characteristics</th>
<th>Number n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Age (n = 100)</strong></td>
<td></td>
</tr>
<tr>
<td>age 0–19</td>
<td>1 (1%)</td>
</tr>
<tr>
<td>age 20–39</td>
<td>9 (9%)</td>
</tr>
<tr>
<td>age 40–59</td>
<td>35 (35%)</td>
</tr>
<tr>
<td>age 60–79</td>
<td>49 (49%)</td>
</tr>
<tr>
<td>age 80–99</td>
<td>6 (6%)</td>
</tr>
<tr>
<td><strong>Obesity</strong></td>
<td></td>
</tr>
<tr>
<td>BMI (kg/m²) ≥ 30</td>
<td>25 (25%)</td>
</tr>
<tr>
<td>BMI (kg/m²) &lt; 30</td>
<td>46 (46%)</td>
</tr>
<tr>
<td>BMI not recorded</td>
<td>29 (29%)</td>
</tr>
<tr>
<td><strong>Menopausal status</strong></td>
<td></td>
</tr>
<tr>
<td>pre-menopausal</td>
<td>22 (22%)</td>
</tr>
<tr>
<td>post-menopausal</td>
<td>73 (73%)</td>
</tr>
<tr>
<td>menopausal status not recorded</td>
<td>5 (5%)</td>
</tr>
<tr>
<td><strong>Parity</strong></td>
<td></td>
</tr>
<tr>
<td>nulliparous</td>
<td>16 (16%)</td>
</tr>
<tr>
<td>1 or 2 births</td>
<td>57 (57%)</td>
</tr>
<tr>
<td>≥ 3 births</td>
<td>22 (22%)</td>
</tr>
<tr>
<td>parity not recorded</td>
<td>5 (5%)</td>
</tr>
<tr>
<td><strong>Pelvic organ prolapse</strong></td>
<td></td>
</tr>
<tr>
<td>prolapse present</td>
<td>46 (46%)</td>
</tr>
<tr>
<td>prolapse beyond vulva level</td>
<td>17 (17%)</td>
</tr>
<tr>
<td>no prolapse present</td>
<td>53 (53%)</td>
</tr>
<tr>
<td>prolapse not recorded</td>
<td>1 (1%)</td>
</tr>
</tbody>
</table>

BMI = body mass index (kg/m²)
12% of the patients took drug combinations that could have a strong effect on the coagulation system. The most probable effect was the possibility of a stronger anticoagulation and thus an increased risk for gastrointestinal bleeding. The combination that was most frequently the reason for a higher risk of bleeding in this collective was the simultaneous consumption of antiplatelet medicines such as Thrombo Ass (aspirin, ASA) or Plavix (clopidogrel) with serotonin re-uptake inhibitors such as Duloxetin, Sertralin or Escitalopram.

Further harmful drug combinations can affect the heart through their intensified cardiodepressive activity with an increased danger for the occurrence of bradycardia or hypotension (11% of the patients). Drug interactions that could alter the patients’ blood glucose or electrolyte households were identified in 6% of the patients. Other, less commonly occurring drug combinations could impact negatively on the neurological and muscular systems. Drug-drug interactions that favour urinary incontinence were not identified.
A further question that remains unanswered by this study is about the actual incidence of the urinary incontinence-promoting side effects of the evaluated drugs.

Practical Conclusions

This study underlines the importance of a critical confrontation with polypharmacy. Polypharmacy is a frequent phenomenon in our society and harbours the risk of illness though over-therapy. In the course of the clarification of urinary incontinence in women, a complete coverage of drug consumption is indispensable. Since many drugs can trigger or promote urinary incontinence, it may be assumed that a discontinuation of the drug in question could lead to an improvement of the incontinence or even to its cure. The omission of a specific drug is, however, often not possible because the drug was prescribed for an important indication. Over-therapy and undesired side effects can only be reduced by a regular inspection of indications and doses in close interdisciplinary cooperation between general practitioners and specialists.

Conflict of Interest

None.

References

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