Prevalence and severity levels of post-radical prostatectomy incontinence: different assessment instruments

Luciana Regina Ferreira da Mata¹
Cissa Azevedo¹
Lívia Cristina de Resende Izidoro²
Darkiane Fernandes Ferreira¹
Fabrícia Eduarda Baia Estevam¹
Fabrícia Moreira Amorim Amaral²
Tânia Couto Machado Chianca¹

ABSTRACT

Objectives: to analyze urinary incontinence prevalence and severity in prostatectomized men assessed by three different instruments. Methods: a cross-sectional study was conducted with 152 men. The pad test, pad used, and International Consultation on Incontinence Questionnaire - Short Form (self-report) were considered. Data were analyzed using Spearman’s correlation, Kappa index, considering a significance level of 0.05. Results: urinary incontinence prevalence was 41.4%, 46.7% and 80.3% according to pad test and self-report. Positive correlations and moderate to poor agreement were found between the instruments. As for severity, most participants had mild incontinence. The largest number of cases of mild and severe incontinence was identified by self-report. Conclusions: the self-report showed higher values for prevalence of mild and severe severity levels. Through the identified differences, we propose that the objective assessment (pad used and pad test) be associated with individuals’ perception (self-report) to better estimate prevalence and severity.

Descriptors: Urinary Incontinence; Prostatectomy; Evaluation Study; Prostatic Neoplasms; Nursing Care.

RESUMO

Objetivos: analisar a prevalência e a gravidade da incontinência urinária em homens prostatectomizados a partir de três instrumentos diferentes. Métodos: estudo transversal, realizado com 152 homens. Foram considerados os instrumentos pad test, pad used e International Consultation on Incontinence Questionnaire - Short Form (autorelato). Os dados foram analisados por correlação de Spearman, Índice Kappa, considerando nível de significância 0,05. Resultados: a prevalência de incontinência urinária foi 41,4%, 46,7% e 80,3% segundo pad used, pad test e autorelato, respectivamente. Constatar-se-ão correlações positivas e concordâncias de moderada a pobre entre os instrumentos. Quanto à gravidade, a maioria dos participantes apresentou incontinência leve. O maior número de casos de incontinência leve e severa foi identificado pelo autorelato. Conclusões: o autorelato apontou valores superiores para prevalência e níveis de gravidade leve e severa. Mediante as diferenças identificadas, propomos que a avaliação objetiva (pad used e pad test) seja associada à percepção do indivíduo (autorelato) para melhor estimativa da prevalência e gravidade.

Descritores: Incontinência Urinária; Prostatectomía; Estudios de Evaluación; Neoplasias de la Próstata; Cuidados de Enfermagem.

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INTRODUCTION

Although the advance in prostate cancer (PC) control is growing, it is still a public health concern worldwide, being responsible for the second leading cause of cancer death among men1-3. Estimates indicate that the highest incidence rates are found in Australia, New Zealand and European countries in northern and eastern regions4-6. In Brazil, 65,840 new cases of PC are expected for each year of the 2020-2022 triennium, which corresponds to a risk of 62.95 new cases for every 100 thousand men7,8.

Among the various treatment modalities for PC, the main strategy considered the gold standard for localized PC is radical prostatectomy (RP). Surgery is based on the surgical removal of the prostate gland, seminal vesicles, part of the vas deferens and, in many cases, the bladder neck9. However, although RP contributes to longer survival, a possible side effect and common after surgery is urinary incontinence (UI), which can significantly compromise quality of life10.

The International Continence Society (ICS) defines UI as a complaint of any UI or involuntary outflow11. All forms of UI are caused by bladder, sphincter dysfunction or a combination of both. In prostatectomized patients, two main types are distinguished: effort (70%) and urgency (30%) UI12. Stress UI is the involuntary UI after performing activities that increase intra-abdominal pressure; urgency UI is the involuntary UI that occurs immediately after a strong urge to urinate13,14.

It is observed that the majority of patients submitted to RP manifest UI in the early postoperative period, immediately after indwelling bladder catheter (IBC) removal, which usually happens between four and 20 days after surgery15. The rehabilitation of urinary continence is gradual and according to the ICS it is estimated that within one month after surgery, 80% of men experience post-radical prostatectomy incontinence (PRPI)16. Another relevant data points out that despite the spontaneous decrease in UI at two years postoperatively, 22% of men can use an absorbent a day and 22% more than an absorbent a day17.

The ICS recommends that for the assessment of an incontinent person it is important to specify UI circumstances, frequency, and severity18. Thus, it is necessary to have specific parameters that assess both UI characteristics and severity19,20. An approach based on the investigation of clinical variables and lifestyle habits related to voiding dysfunction can support the development of strategies aimed at minimizing or resolving them.

Among the methods of simple, non-invasive and effective measures to assess UI presence and severity, one can mention the pad test and the pad used21. The pad test has been recommended by the ICS since 1988, being a standardized way of measuring UI22. There are different versions of this test according to the duration; however, the one-hour test has been adopted in clinical practice, since the longer the application time, the greater the infeasibility to implement it23. The pad used is widely used for its simplicity, and consists of questioning the number of changes in the pad/liner/diaper in a 24-hour period24.

Since 1997, the ICS recommends that measures related to quality of life be included in all clinical research on UI as an additional complement to traditional clinical parameters25. Thus, in addition to instruments such as those mentioned, there are also validated questionnaires26-28, which consist of precise measures on patients’ perception of UI. In this context, the International Consultation on Incontinence Questionnaire - Short Form (ICIQ-SF) provides a brief and general assessment allowing, in addition to detecting UI, the measurement of its severity and its impact on quality of life29.

It is known that the estimated prevalence of PRPI is between eight and 87% (6,10), which characterizes a high variability. The different strategies for measuring this outcome may result in this variability in prevalence rates30. Therefore, it is important to recognize that subjective and objective methods have different sensitivity for assessing the same clinical condition, in this case UI31. Thus, this study is relevant for comparing UI measurement instruments, in order to answer the following questions: is there a relationship and agreement between the prevalence measures assessed by the pad test, pad used, and ICIQ-SF? How are the distributions of UI severity levels across these three assessment instruments?

OBJECTIVES

To analyze urinary incontinence prevalence and severity in prostatectomized men assessed by three different instruments.

METHODS

Ethical aspects

In compliance with the recommendations of Resolution 466/12 of the Brazilian National Health Council (Conselho Nacional de Saúde) regarding research related to human beings, the study was assessed and approved for implementation by the Research Ethics Committee of Universidade Federal de São João del-Rei. Data collection started by signing the Informed Consent Form (ICF), leaving a copy with participants.

Design, period, place of study

This is a cross-sectional study described from the guidelines for observational studies (Strengthening the Reporting of Observational Studies - STROBE)32. It was carried out from August 2017 to June 2018. The sample consisted of men submitted to RP in a High Complexity Care Unit in Oncology (UNACON - Unidade de Assistência de Alta Complexidade em Oncologia) in Minas Gerais, linked to the Brazilian National Cancer Institute (INCA - Instituto Nacional do Câncer).

Sample; inclusion and exclusion criteria

The sample size calculation was defined using the statistical test of simple random sample for finite population33, estimating a proportion of UI referring to the population of interest equal to 46.7%34. Thus, considering the population of men treated at the institution in a two-year interval equivalent to 242 individuals, with a margin of error of 5% and a confidence level of 95%, the calculation resulted in a minimum size of 149 participants.

Men aged over 18 years who underwent RP in postoperative follow-up for at least two months and maximum two years were selected, with preserved auditory and verbal capacity. Those who were using an IBC and those who reported preoperative UI were excluded.
Study protocol

An invitation to participate in the study was carried out on the medical return day, with presentation of the research objectives and the interview procedures. All men who agreed to participate completed the ICF. Data collection was carried out through an individual interview conducted by one of the researchers before the medical consultation, in a nursing room, in a private environment.

The data collection instruments used were: sociodemographic questionnaire, ICIQ-SF\textsuperscript{(15)} and the clinical tests used pad\textsuperscript{(13)} and one-hour pad test\textsuperscript{(12)}.

For sociodemographic and clinical characterization, an instrument was elaborated that included data such as age, education, individual monthly income, professional and marital status, type of surgery (retropubic or laparoscopic), and post-surgery time.

The ICIQ-SF consists of a simple, short and self-administered instrument that assesses the impact of UI on quality of life and qualifies patients’ UI. It consists of four questions related to UI frequency, severity, impact and self-diagnosis related to the causes or situations of UI experienced, respectively\textsuperscript{(15)}. The total score ranges from zero to 21, and the higher the value, the greater the impact on quality of life. In the present study, items three and four of the ICIQ-SF were considered, for analysis purposes, since they are the items of the instrument related to UI frequency and quantity, i.e., they allow to establish, specifically, UI prevalence and its severity. It is noteworthy that participants were instructed to answer the items based on their current clinical condition with regard to UI.

Item three of the ICIQ-SF (How often do you lose urine?) aims to assess UI prevalence\textsuperscript{(15)}. Answers range from zero to five, with zero - never; one - once a week or less; two - two or three times a week; three - once a day; four- several times a day; five - all the time\textsuperscript{(15)}. Thus, individuals who reported no involuntary UI (ever) are classified as continents and those who reported any involuntary UI, regardless of frequency, should be classified as incontinent\textsuperscript{(15)}. Item four of the ICIQ-SF (We would like to know how much urine you think you lose) consists of analyzing UI severity\textsuperscript{(15)}. Answers range from zero to six, with zero meaning no loss (continent), two - a small amount (mild UI), three - a moderate amount (mild UI) and six - a large amount (severe UI)\textsuperscript{(15)}.

The pad used aims to quantify the number of pads used by the individual in 24 hours. Thus, UI is classified as mild (when absorbent use is one to two per day), moderate (three to five absorbents per day) and severe (more than six absorbents per day). Men who used no pads were classified as continents\textsuperscript{(11)}.

The one-hour pad test consists of placing a penile pad close to the external urethral meatus to quantify UI by comparing the weight of this pad before and after the hour. During this interval, patients are submitted to a protocol of fluid intake and Activities of Daily Living\textsuperscript{(10,12)}. From the difference in weight of the initial and final absorbent, UIs are classified as: insignificant or continent loss (when the final weight of the absorbent is up to one gram (g), slight loss (1.1 to 9.9 g), moderate loss (10 to 49.9 g) and severe loss (above 50 g)\textsuperscript{(12)}. At the beginning of data collection, patients were instructed on the positioning of the absorbent and to place it at that moment of the interview. Then, 500 mL of water were offered and the activities protocol that should be carried out in an hour was explained.

During the initial 15 to 20-minute interval, according to the pad test protocol, patients should remain at rest; therefore, at this time, the other data collection instruments were applied (sociodemographic questionnaire, ICIQ-SF and pad used). Subsequently, a script of activities was carried out which included: walking in slow steps, sitting and standing, picking up objects on the floor, coughing, washing hands under running water, going up and down stairs. These activities were carried out on the institution’s premises. After the activities were completed, the patient was returned to the nursing room, where a plastic bag was offered to discard the absorbent. Finally, the researcher used a precision scale to measure the weight of the absorbent and complete the test result.

Analysis of results, and statistics

Data were processed and analyzed using the Statistical Package for Social Science (SPSS), version 21.0 for Windows. The Shapiro-Wilk test was applied to test the normality of explanatory variables. The results obtained for explanatory variables (sociodemographic characterization) were analyzed using descriptive statistics with measures of central tendency (mean or median) and variability (standard deviation or interquartile range) for continuous variables, and relative frequency for variables categorical.

PRPI prevalence with the respective 95% confidence interval (confidence interval - 95% CI) was calculated from the findings of the pad test, pad used and item three on the ICIQ-SF scale.

In order to identify possible relationships between the levels of PRPI assessed by the pad test, pad used and items three and four of the ICIQ-SF, Spearman’s correlation test was used. The correlation forces were analyzed considering values between 0.10 and 0.39 as of low magnitude, between 0.40 and 0.69 of moderate magnitude and above 0.70 of strong magnitude\textsuperscript{(20)}.

Then, agreement analysis was performed between PRPI occurrence according to each of the three instruments (pad test, pad used and item three of the ICIQ-SF) using Kappa coefficient with the respective 95% CI. Agreement was analyzed from the parameters: null (k = 0), poor (0.01 - 0.19), weak (0.20 - 0.39), moderate (0.40 - 0.59), substantial (0.60 - 0.79) and almost perfect (0.80 - 1) (21). In all analyzes, a significance level of 0.05 was considered. Histograms were designed to compare the distribution of PRPI severity levels (continent, mild, moderate and severe) according to the three instruments.

RESULTS

The number of participants eligible for the study was 175 men. However, 15 individuals were disregarded due to IBC use, five who reported previous UI and three who did not accept to participate in the study. Therefore, the sample consisted of 152 men. The median age was 67 (62-72.7) years. As for education, participants had a median of four (2-4) years of study. The median individual monthly income was 937 (937-1405) reais (reais is the Brazilian currency). Concerning professional situation, 78.9% were inactive (retired or unemployed) and 80.3% had a partner. Most performed the surgery using the retropubic technique (97.4%) and the post-surgery time varied between 60 and 730 days, with a median of 209 days, i.e., approximately seven months.
PRPI prevalence was 46.7% (CI = 38.7%; 54.7%) according to the pad test, 41.4% (CI = 33.5%; 49.4%) by the pad used and 80.3% (CI = 73.9%; 86.7%) by item three of the ICIQ-SF.

When analyzing possible correlations between the levels of PRPI by the three assessment methods, a positive correlation of moderate magnitude was found between pad test and pad used (r = 0.54/p <0.001), and pad used and item three of ICIQ-SF (r = 0.61/p <0.001). There was also a positive correlation of weak magnitude between the pad test and item three of the ICIQ-SF (r = 0.37/p <0.001).

Concerning the agreement between the three instruments for PRPI prevalence assessment, moderate agreement was identified between pad test and pad used, poor between pad test and item three of the ICIQ-SF, and weak between pad used and item three of the ICIQ-SF (Table 1).

**Table 1 - Agreement between the pad test, pad used and item three instruments of the International Consultation on Incontinence Questionnaire - Short Form regarding urinary incontinence prevalence assessment, Divinópolis, Minas Gerais, Brazil, 2018 (n=152)**

| Assessment methods | Kappa | 95% CI | p      | Agreement |
|---------------------|-------|-------|--------|-----------|
| Pad used and pad test | 0.441 | [0.297; 0.585] | <0.001† | Moderate |
| Item three of the ICIQ-SF and pad test | 0.076 | [0.000; 0.229] | 0.218 | Poor |
| Item three of the ICIQ-SF and pad used | 0.297 | [0.156; 0.437] | <0.001† | Weak |

Note: *International Consultation on Incontinence Questionnaire - Short Form; †p<0.05; 95% Confidence Interval.

The distribution of participants according to PRPI levels of severity (continent, mild, moderate, and severe) by the three assessment methods is shown in Figure 1. The difference in the distribution of participants assessed by ICIQ-SF stands out for the classification continent (n = 30) and “mild” PRPI (n = 99) in relation to the other methods. With this finding, it appears that in the severity assessment by self-report there was a predominance of complaints of loss of a small amount of urine. In the “severe” PRPI classification, self-report (ICIQ-SF) was an instrument that identified a greater number of participants (n = 10) in this category.

**DISCUSSION**

The results of this study allowed us to show the variability in PRPI prevalence and levels of severity before different assessment strategies. For this sample, we estimated a prevalence of PRPI of 46.7% according to the pad test, 41.4% according to the pad used and 80.3% according to item three of the ICIQ-SF. Thus, the prevalence measured by item three of the ICIQ-SF was considerably higher than the prevalence assessed by the pad used and pad test. This finding corroborates the results of research carried out in recent years on methods of assessing UI that highlighted the greater or lesser validity of one method in relation to the other.

A research carried out in Norway identified a 74% prevalence of PRPI after one year of surgery when assessing UI by reporting on the perception of any involuntary UI and 40% according to the pad used (use of at least one absorbent/liner/diaper per day). The authors stressed that this discrepancy can be attributed to the discomfort felt by individuals when they perceive involuntary UI, even if this loss is minimal.

In Spain, scholars have assessed the prevalence of PRPI in 172 men one year after surgery, considering UI as any involuntary UI and identified a prevalence of 23%. They also found that 17.8% used at least one absorbent/day and 11.9% more than one absorbent/day.

In a study carried out in Japan with men after a year of RP, a prevalence of PRPI of 75% was identified by the ICIQ-SF, 33% by the pad used and 36% by the 24-hour pad test. The authors stressed that no method should prevail over the others, and a broad clinical assessment is essential, in addition to considering that the best option will always be the one that can be measured and compared in the pre and postoperative period, whatever this may be. We believe that the differences in prevalence identified in the self-report assessment (ICIQ-SF) in relation to the two objective assessments (pad used and pad test) reinforce the idea that the assessments are complementary for the selection of PRPI control interventions that includes since change in lifestyle and pelvic muscle training for drug or surgical interventions.

Australian scholars have also identified a discrepancy between the subjective impression of prostatectomized patients on continence status and objective criteria. It was found that 34.4% of the 479 men submitted to RP were classified as incontinent through a telephone interview that asked “do you have involuntary UI?”, while only 14.9% of these men were classified as incontinent through the 24-hour pad test. The experience regarding UI may present different interpretations due to the questioning itself, whose focus may be on use or not of an absorbent or the amount of UI itself.

There are also situations in which, although patients experience UI several times a day, they are resistant to using the absorbent because they consider the amount lost to be very small. Thus, we suggest that the amount of UI is what determines use or not of an absorbent and not the frequency of this loss.

The number of absorbents in 24 hours is an important aspect for determining UI; however, this does not accurately define UI, other complementary information such as the size and frequency of changing the absorbent, the absorption capacity of
that absorbent, and the moisture tolerance of each individual being important. Another relevant data refers to the shape of the absorbent, as individuals who use rectangular pads are 40% more likely to change the device more frequently in a 24-hour period when compared to those who use diaper-type pads.

Regarding the definition of UI through the number of pads or diapers used, authors have used in their studies the criterion that continent individuals are those who do not use pads or use them only for safety, which is also accepted by the ICS. In contrast, there are scholars who consider it more appropriate and objective to define individuals as continent when they do not use any absorbent/liner/diaper. We understand, therefore, that the existence of different definitions can also impact the fluctuation in the prevalence rates of PRPI.

The results also showed positive correlations between the assessment methods, i.e., an increase in the amount of UI increased the number of absorbents used, as well as individuals’ perceptions of UI frequency. Other studies that assessed the correlation between different methods also identified a positive relationship between them, with a predominance of weak to moderate correlation forces.

As for agreement between instruments, assessed using Cohen’s Kappa coefficient, there was significant agreement between assessment by the pad test and pad used, as well as by the pad used and item three of the ICIQ-SF, with stronger agreement between pad used and pad test. This finding can be justified by the fact that both are objective methods of assessing UI that do not involve man’s perceptions of the frequency of this loss.

In a study that assessed agreement between the one-hour pad test and the ICIQ-SF, a weak agreement (kappa <0.34) was identified between the methods. Although the one-hour pad test has the advantage of a greater chance of adherence by patients, factors such as level of physical effort, use of drugs such as antibiotics and anti-diuretics, and low daily water intake can impact the results. These factors can interfere with the real frequency of UI and, consequently, with the severity and impact of UI on patients’ quality of life.

A research carried out in Germany, which assessed the agreement between pad used and self-report of involuntary UI, showed an agreement of 0.73, 0.70 and 0.64 at three, six and 12 months after surgery, respectively. The greater agreement identified in this study can be justified by the definition adopted for the self-report of involuntary UI. Individuals were considered as continent if they did not use any type of device.

As for comparison of the distribution of UI severity between the three assessment strategies, it is realized that by self-report (ICIQ-SF), most participants were classified as “mild” PRPI. In contrast, severity assessment by the pad test and pad used pointed out that most participants were continent. Moreover, in the “severe” PRPI classification, self-reporting also stood out with the largest number of participants. These findings reinforce the idea that men’s perceptions of UI have a greater impact on the degree of severity compared to severity assessment by measuring the volume of UI and the number of absorbents used. Furthermore, it shows us that subjective assessment is important and complementary and directs nursing actions within the scope of self-perception, coping, and adaptation to health conditions.

**Study limitations**

As a limitation of this study, we evidenced the recruitment of participants in a single institution, which increases the chances of a sample with very similar characteristics in relation to income and education, which can impact hygiene conditions and, consequently, the frequency of changing absorbents. Another limiting factor that we identified is related to the pad test application, since older men had limitations to perform all the daily activities contemplated in the protocol, which made it difficult to standardize the level of physical effort among participants.

**Contributions to nursing**

We know that the differences between prevalence values of PRPI in clinical practice reinforce the importance of the knowledge we need to select assessment instruments, in addition to highlighting the need for new studies that propose specific questionnaires for the male audience.

UI assessment by nurses is an important parameter for planning care. In the case of patients in outpatient care, the severity of this symptom indicates the need for more frequent follow-up, either by telephone contact or home visits. In more complex situations, the possibility of surgical or drug intervention can be discussed with the medical team.

Knowing the specificities of UI measurement instruments favors the professional improvement of nurses in the area of voiding dysfunctions. It is important to understand that nursing is considered the driving force of the health team and is therefore able to change the reality of care related to UI control in the Brazilian health system.

**CONCLUSIONS**

The present study compared PRPI prevalence and severity levels under three different methods, two objectives (pad test and pad used) and one subjective (ICIQ-SF - self-report). We found a higher prevalence of PRPI by the subjective method. As for level of severity, most participants had a mild UI in an assessment using the three instruments. Among the cases of mild and severe UI, self-report assessment showed the largest number of individuals for these classifications.

It is noticed, therefore, that the self-report pointed higher values for prevalence and mild and severe levels. Thus, lower rates indicated by the one-hour pad test and the pad used suggest that such methods are less reliable when assessed in isolation to estimate PRPI prevalence and severity, without considering the perception of individuals who present involuntary UI. Through this difference identified between the instruments, we propose that the objective assessment is associated with the perception of individuals, since the same amount of UI can be perceived differently among men, considering their personal and social perceptions.

Finally, due to the variability in PRPI rates by the methods analyzed and the difficulty in establishing the real magnitude of this involvement, we emphasize the importance of future investigations with proposals for a single instrument that associates objective and subjective data to determine UI prevalence and severity within the male population.
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