**Persistent genital arousal disorder: always look beyond the surface - a case report**

Rita Sarabando¹*, Natacha Sousa¹, Ana C Borges¹, Cristina Nogueira-Silva²

¹Department of Obstetrics and Gynaecology, Hospital de Braga, Braga, Portugal
²Department of Obstetrics and Gynaecology, Hospital de Braga, Braga; Life and Health Sciences Research Institute (ICVS), School of Medicine, University of Minho, Braga; ICVS/3B’s - PT Government Associate Laboratory, Braga/Guimarães, Portugal

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*Correspondence:
Dr. Rita Sarabando,
E-mail: sarabandorita@gmail.com

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**ABSTRACT**

Persistent genital arousal disorder is a rare condition characterized by unwanted intrusive symptoms of sexual arousal without specific context. Their possible aetiologies and treatments are multiple and mostly based on case reports. We aim to do a comprehensive review of persistent genital arousal disorder and describe a case of a postmenopausal woman who developed this disease and, during the follow-up, was diagnosed with advanced endometrial cancer, reminding physicians to keep in mind the possibility of multiple diagnosis in the same patient, including malignancy. Although there is no description of this association in the literature, the possible aetiologies of persistent genital arousal disorder are diverse, and we sought this rare case should be disclosed.

**Keywords:** Persistent genital arousal disorder; Endometrial cancer; Malignancy

**INTRODUCTION**

Persistent genital arousal disorder (PGAD) is a rare medical condition characterized by symptoms of sexual arousal in the absence of perceived sexual stimulus. Its prevalence is still not known, but it is estimated to be about 1% of women.¹ The social stigma and the women shame and fear to be misunderstood may contribute to the probable underestimation of this disorder prevalence.¹² The symptoms include genital vasocongestion, increased sensitivity of the genitals and nipples, vaginal lubrication, genital tingling, throbbing, and contractions. Pelvic pain can also be present.²³ In 2001, Leiblum and Nathan originally proposed five criteria for this condition: genital and clitoral arousal that persists for an extended period (hours, days, or months), symptoms do not resolve with ordinary orgasms, and require multiple orgasms over hours or days to remit, genital arousal is unrelated to subjective feelings of sexual desire, persistent genital arousal can be triggered by sexual activity, non-sexual stimuli or by no apparent stimulus at all and a very persistent feelings of genital arousal is intrusive, unwanted and cause at least a moderate degree of distress. Sometimes distress is considered a separate, sixth criteria.⁴ Those criteria have been published by the International society for the study of women’s sexual health (ISSWSH) in the consensus nomenclature for distressing sexual dysfunctions and are included in the recommendations made by the fourth international consensus on sexual medicine.⁵⁶

The aetiology is still not completely known, but it can be related to central or peripheral neurological anomalies (peripheral nerve hypersensitivity or entrapment and neuronal hyperexcitability may be one of the main features of this disorder), Tarlov or sacral spinal cysts, pelvic congestion, pudendal neuralgia, use or withdrawal of medications such as anti-depressants, increased dietary soy intake, or psychological factors such as anxiety.⁴⁷ There seems to be also a relationship between PGAD and restless legs syndrome in a condition classified as restless genital syndrome, in which overactive bladder can also be
As the aetiology is still not completely understood, treatment is also a challenge. Non-pharmacological treatments include avoidance of tight clothing and of prolonged sitting, dietary modifications and counselling, also pelvic floor physiotherapy, nerve modulation or stimulation and surgical options (there are descriptions of removal of masses or spinal meningeal cysts, and embolization of pelvic veins). Pharmacological treatment options that may relieve PGAD symptoms include antiandrogens, anxiolytics, antidepressants, anticonvulsants, dopamine agonists and dopamine antagonists.

This clinical case describes a diagnosis of PGAD and its follow-up, during which the patient was diagnosed with an endometrial cancer in an advanced stage.

**CASE REPORT**

A 67 year old woman presented in a gynaecological appointment describing vaginal discharges that occurred with a sensation like sexual excitement for the last 3 months, more than 10 times a day and waking her up during the night. The patient denied urinary incontinence or urgency, and the sensation described was present in the absence of sexual desire or sexual activity. She denied ever having had sexual intercourse and referred menopause at the age of 51, with one year of hormone replacement therapy after that. She was asthmatic and was taking daily medication for high blood pressure and dyslipidaemia, with no other relevant illnesses, namely neurological or surgeries. She was not taking any other medications like anti-depressants or over-the-counter pills. On abdominal examination, the patient had no relevant findings, and on gynaecological examination (limited by the fact that the patient did not have coitarche) severe vulvovaginal atrophy was observed, probably related to genitourinary syndrome of menopause (SGUM), with no other relevant findings. The suspected diagnosis was PGAD. She denied other symptoms like paraesthesia and/or dysesthesia, genitopelvic, leg, or lower back pain, bladder symptoms or restless legs. In order to exclude organic pathology, a suprapubic ultrasound was performed and showed a uterus with 85x75x65mm with lobulated contours, heterogeneous texture and multiple myometrial intramural masses compatible with leiomyomas type 4, the biggest of them with 37x29mm (Figure 1). Endometrial cavity measure was 4.86mm (Figure 2). There were no adnexal lesions and no free liquid in the abdominal cavity. She was recommended to do a hysteroscopy in which endometrial biopsies were done and showed endometrial serous carcinoma. In the thoracoabdominopelvic computed tomography done for staging, the uterus was multinodular with lobulated contours and the endometrium was thickened (Figures 3A and 3B). Some lymphadenopathies were seen at peri-aortocaval locations and around iliac vessels, with no other image findings of spread malignancy. She was referred to an oncologic gynaecological service. Hysterectomy with salpingo-oophorectomy, aortic and pelvic lymphadenectomy were performed, and complete resection was achieved (R0). The pathological examination confirmed the endometrial serous carcinoma with invasion of more than half of the myometrium, lymphovascular invasion, and metastases to pelvic and para-aortic lymph nodes (FIGO IIIC2).

Patient is currently under adjuvant chemotherapy, and adjuvant radiotherapy is also planned. Patient is stable and asymptomatic at six months after surgery.

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**Figure 1: Suprapubic ultrasound - uterine leiomyoma.**

**Figure 2: Suprapubic ultrasound - endometrial cavity.**
PGAD is a very perplexing entity since we are still trying to understand all the possible causes for it.\(^3,7\) There are no guidelines or algorithms for its diagnosis and treatment, so the published literature is particularly important in the guidance of the rare cases that physicians are confronted with.\(^2\)

In this case, the first difficulty was to understand the description of the symptoms by the patient, since she was woman with no coitarche or any type of sexual contact, so she first described the symptoms as discharges. After excluding urinary incontinence and asking about more sensations during the episodes we understood that maybe the patient was experiencing unwanted multiple orgasms, triggered by no apparent stimulus, and that were causing an elevated degree of distress, since she had no idea what was happening to her. This postmenopausal woman searched for help after some months having these symptoms and a diagnosis of PGAD was made. At that time, by excluding other possible aetiologies found in the literature, a psychological component was assumed, so an empirical pharmacological treatment with cloxazolam was started.\(^3,7,9\) Also, because of probable SGUM, local treatment with oestrogen and vaginal moisturizer was prescribed. Symptoms started to get better and after some months only very sporadic episodes were reported by the patient.

After three years of illness stabilization, another diagnosis was made - endometrial serous carcinoma at an advanced stage. She got appropriated surgery and is now under chemotherapy. Since the aetiology of PGAD is not completely known, although there was no description of this association in the literature, the possible aetiologies of PGAD were so diverse and still being discovered.\(^3,7\) We sought this rare case should be disclosed. We are still trying to better understand the pathophysiology of PGAD and during the process of diagnosis, every physician goes to the literature and search for what is published. Our aim was to do a comprehensive review of PGAD and describe a rare case of a postmenopausal woman who developed this disease and during the follow-up was diagnosed with another serious disease which was advanced endometrial cancer. This reminds physicians to keep in mind the possibility of multiple diagnosis in the same patient, including the possibility of malignancy and the importance of maintained surveillance and high suspicion in clinical practice. The endometrial cancer was diagnosed in an advanced stage, despite annual gynaecological observation and we could not exclude that the assumed leiomyoma in the first ultrasound could already be the endometrial neoplasm.\(^8\)

CONCLUSION

This case reported a diagnosis of PGAD, which is a very uncommon condition with aetiology and pathophysiology not completely known. We believe to have contributed by sharing this case with the medical community, alerting for the importance of PGAD diagnosis but also to essential ongoing follow-up of every women because other diagnosis may appear, as serious as malignancy.

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