Premenstrual syndrome and cortisol

Florica SANDRU1,2, Mihai Cristian DUMITRASCU1,3, Eugenia PETROVA1,4, Adina GHEMIGIAN1,4, Nicoleta DUMITRU1,4, Mara CARSOTE1,4, Ana VALEA5,6

1 “Carol Davila” University of Medicine and Pharmacy, Bucharest, Romania
2 Elias Emergency Hospital, Bucharest, Romania
3 University Emergency Hospital, Bucharest, Romania
4 “C.I. Parhon” National Institute of Endocrinology, Bucharest, Romania
5 “Iuliu Hatieganu” University of Medicine and Pharmacy, Cluj-Napoca, Romania
6 Clinical County Hospital, Cluj-Napoca, Romania

Abstract

Premenstrual syndrome (PMS), including the severe subtype premenstrual dysphoric disorder (PMDD), DSM-5 category, represents a challenging combination of hormonal, environmental and neuroendocrine dysfunctions with menstrual cycle-related pattern. Controversies around the role of daily stress and associated anomalies of hypothalamic-pituitary-adrenal axis are related to the fact that stress is all the time, not just a fluctuating element. This is a narrative review on PMS/PMDD and cortisol profile. 46 articles are cited (between 2009 and 2020). PMD/PMDD underlines multiple imbalances and anomalies of the cortisol levels or its secretory pattern may be a few of them, despite the fact that multiple controversies are still present and most of studies are of limited statistical power. Women with PMS may have higher levels of cortisol in relationship to stress independently of the cycle phase, also a delay of CAR (cortisol awakening response) peak and a delayed cortisol slope during day time. It does not seem that CAR pattern is related to the phases of menstrual cycle. CAR anomalies may be associated with pain perception disturbances in PMS females. The most modern area of interest is related to allopregnanolone, a progesterone metabolite with neuroactive profile. The diurnal serum baseline cortisol and the values of cortisol after dexamethasone suppression test may be similar between patients with PMS and without, but the females with PMS that have higher allopregnanolone associate blunted values of cortisol during the night versus control (without PMS) and versus women with low allopregnanolone levels, thus proving a suboptimal response to stress. Allopregnanolone modules GABA receptors on a paradoxical manner inducing anxiety and irritability during luteal phase on women with a specific predisposal configuration of GABA receptor as those confirmed with PMDD. Overall, PMS/PMDD impairs the quality of life, thus the more we understand about its pathogeny, the easier it gets to control it.

Keywords: premenstrual syndrome, cortisol, premenstrual dysphoric disorder, stress, nutrition

Abbreviations

CAR = cortisol awakening response
GABA = acid gama-aminobutiric
GAMSA = GABAA modulating steroid antagonist
S-IgA = salivary immunoglobulin A
PMS = premenstrual syndrome
PMDD = premenstrual dysphoric disorder
SSRI = selective serotonin reuptake inhibitors
TSST = Trier Social Stress Test

Corresponding author:
Mara Carsote
E-mail: carsote_m@hotmail.com

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INTRODUCTION

Premenstrual syndrome (PMS), including the severe subtype premenstrual dysphoric disorder (PMDD), DSM-5 category, represents a challenging combination of hormonal, environmental and neuroendocrine dysfunctions with menstrual cycle-related pattern (1-3). PMDD, a severe, luteal phase-related impairment of emotional and physical status with suggestive symptoms, also displays the complete remission once the menstruation is present, as well as having in common with PMS some uncertainties related to the actual underlying contributor pathways like serotonin, GABA (acid gama-aminobutiric), reproductive steroids, and other brain circuits (1,4). Controversies around the role of daily stress and associated anomalies of hypothalamic-pituitary-adrenal axis are related to the fact that stress is all over described in everyday life, not just a fluctuating element or menses-related and it is difficult to be quantified in specific studies of good statistics quality (5). PMS impairs the quality of life, especially PMDD (6). A combination of physical and psychiatric elements are repeating every menstrual cycle, more severe than most females of reproductive age that experience some symptoms before menstruation (up to 90% of female population having reproductive age) (6). While most of the symptoms may be controlled with lifestyle intervention, PMDD subjects are candidates to SSRI (selective serotonin reuptake inhibitors) and probably longer period of time for second line therapy with oral contraceptives in addition to alternative approaches (7).

OBJECTIVE

We aim to highlight aspects that involve PMS/PMDD and data on cortisol levels upon on different aspects of assays.

METHOD

This is a narrative review. The research words are “premenstrual syndrome”, “premenstrual dysphoric disorder” and “cortisol”. 46 articles are cited (between 2009 and 2020).

RESULTS

PMD/PMDD underlines multiple hormonal imbalances, and anomalies of the cortisol levels or its secretory pattern, may be a few of them, despite the fact that multiple controversies are still present and that the most of studies are of limited statistical significance power (8). A study from 2019 on 61 women with PMDD and a similar number of control subjects evaluated subjective reactivity to stress during luteal phase in addition to assays of salivary cortisol and cortisol awakening response (CAR) showed higher levels of cortisol in relationship to stress independently of the cycle phase, a delay of CAR peak and a cortisol slope during day time which was more flatted than control (2). It has been suggested that women with PMS have a stress-related dysfunction of hypothalamic-hypophysis-adrenal axes (9). For instance, a study on 32 women with PMS (of mean age 22.47 years) versus 36 controls identified their attenuated CAR, during mid-follicular but also during late luteal phase, so it does not seem that CAR pattern is related to the phases of menstrual cycle (9). CAR anomalies may be associated with pain perception which was suggested to be disturbed in PMS females (10,11). One study of 59 females with PMS (aged of 22.2 years) showed that salivary CAR increased during the premenstrual phase and not during the menses (meaning a flat CAR pattern) while salivary cortisol is positively correlated with estradiol, respective progesterone, and pain perception is more severe when ovarian steroids are low (10). A Japanese longitudinal controlled study on 31 women with PMS assessed salivary cortisol and immunoglobulin A (S-Ig A) in association with stress scores during different phased of menstrual cycle, revealing that post-menstrual S-IgA are reduced in PMS women versus control while pre-menstrual levels are higher but cortisol levels did not achieve statistical significant differences, thus S-IgA levels seem better connected with stress scales rather than cortisol in PMS women (12). Another study applied Trier Social Stress Test (TSST) on PMS subjects (13). The data on 36 females with PMS (mean age of 21.69 years) versus 36 age-matched control women revealeded blunt cortisol levels as stress response according to TSST independently to the phases of menstrual cycle (13). Also, based on this study, cortisol rather than heart rate or stress scale, is a better surrogate of PMS while hypo-reactivity of hypothalamic-hypophysis-adrenal axis may be the clue of PMS severity (13). Cortisol anomalies might be the key of both emotional and cognitive processes anomalies in PMS/PMDD (14). One pilot study showed that women with PMS have increased levels of anomalies related to stress perception during luteal phase (14).

Another area of interest is related to allopregnanolone, a progesterone metabolite with neuroactive profile, which is described in another pathway, meaning that diurnal serum baseline cortisol and the values of cortisol after dexamethasone suppression test may be similar between patients with PMS and without, but the females with PMS that have higher allopregnanolone associate blunted values of cortisol during the night versus control (without PMS) and versus women with low allopregnanolone levels (15). Allopregnano-
lone actually seems one of the key players of understanding the mood symptoms that appear to be due to high stress sensitivity during luteal phase (16). Allopregnanolone is a modulator of GABA receptor and in PMS females a poor control of the complex represented by GABA receptor-allopregnanolone is displayed during luteal phase under the control of hypothalamic – pituitary – adrenal axes (16,17). Another collateral theory involving brain status during luteal phase of menstrual cycle may be brain inflammation (18). However, progesterone and allopregnanolone may represent useful therapy for postpartum depression, opposite to their negative role in PMS (19,20). The exposure to progesterone during ovulation may increase the risk of developing PMDD (21,22). Despite the fact that exact etiology of PMDD is still unknown, recently the attention was switched from cortisol to allopregnanolone and its role as modulator of GABA receptor causing sedation if the concentration is increased but paradoxically causing anxiety, irritability, depression and all the other mood elements of PMDD, if the patient with PMS has a particular configuration of GABA receptor which make her more susceptible to have the syndrome (23,24). Therapy with allopregnanolone inhibitor like sepranolone which is a GABAα modulating steroid antagonist (GAMSA) seems promising (25,26,27).

Another pathogenic element, in addition to cortisol anomalies as well as allopregnanolone-GABA system anomalies, is represented by abnormal changes of serotonin pathways in PMS females (28,29). That is why the first line pharmacological therapy is represented by SSRIs (30,31).

**DISCUSSIONS**

Other pathways that are not yet completely approved to be linked with PMS/PMDD involve hypovitaminosis D, otherwise with a large worldwide prevalence (32,33). A meta-analysis of all databased publications until 2018 on this particular topic showed on 16 studies that 25-hydroxyvitamin D was not correlated with PMS criteria as revealed by non-interventional studies, while interventional trials pointed that vitamin D supplementation seem efficient to some extent in controlling the cyclic symptoms, suggesting that nutrition may be a valuable approach of PMS (34). Also, calcium supplementation may be beneficial, despite the fact that dose-intervention trials are still lacking (35,36). Another nutritional issue that was suggested is the level of low serum magnesium as permissive element of PMS (37,38). However, not all authors agree, for instance, a meta-analysis on published papers until 2019 included 13 studies showed that there is no correlation between the level of serum magnesium and PMS symptoms during follicular, respective luteal phase (39). Another aspect is the relationship with high body mass index, obesity been another condition with a massive prevalence anywhere in the world (40,41). Obesity has been associated with multiple conditions of the women within their reproductive years from fertility issues, miscarriage to urinary incontinence etc. (42,43). Leptin levels, also linked with adipose tissue mass and activity, has been found higher in females with PMS based on some studies (44). However, the observation is not universal. For instance, other studies showed that high leptin is correlated with increased caloric intake in women with PMDD and high body mass index (through leptin resistance), while normal weight-ed females with PMDD have a decrease of leptin during late-luteal phase which is translated into overeating behaviors (45). Mood and food intake anomalies in PMDD may also be connected to dysregulation of insulin levels (46).

**CONCLUSION**

Overall, PMS/PMDD impairs the quality of life, especially PMDD, thus the more we understand about its pathogeny, the easier it gets to control it. Recently, cortisol theories shifted to allopregnanolone – GABA system modulation.