Due to the depletion of primordial follicles, ovarian estradiol secretion ceases at menopause and is followed by very low levels of circulating estradiol generating the typical menopausal symptoms. However, the arrest of ovarian estradiol secretion does not induce an overall loss of estrogen production: After menopause, sex steroids continue to be synthesized in peripheral tissues depending on steroid forming enzymes specific for each tissue [1]. In contrast to the ovarian estradiol secre-
tion, peripheral produced estrogens are inactivated intracellularly without significant release to the blood circulation.

One of the common and stressful symptoms of menopause are hot flushes (HF), which occur in > 75% of menopausal women [2]. The episodic sensations of heat, intense sweating and flushing can recur with varying frequency and intensity [3]. Likewise, the age at onset of HF is varying from woman to woman. Even though the pathophysiology of HF is not entirely understood, several authors propose that HF are due to a changed thermoregulation set point of the hypothalamus evoked by the lowered estrogen levels during menopause [3,4]. Thereby, estrogens seem to interact with several neurotransmitters like norepinephrine, serotonin and endogenous opioids [5,6]. Vasomotor symptoms can also occur in women with abrupt drop in sex steroid hormones such as after removal of the ovaries of premenopausal women or in breast cancer patients with chemically induced menopause [3]. However, estradiol levels do neither explain the presence of HF nor correlate with their frequency and intensity [3,7]. Thus, measurement of estradiol levels during menopause rarely reveals clinical benefits. Even after an adequate restoration of estradiol levels by hormone replacement therapies, women can still experience vasomotor symptoms.

Rarely, high levels of circulating estradiol have been reported in postmenopausal women so far and have often been attributed to estradiol-producing neoplasia. In this case report, we describe a postmenopausal woman with persisting postmenopausal symptoms in whom high estradiol levels have been measured.

**Case Report**

We report the case of a 54-year-old woman of normal weight (167 cm, 60 kg), who presented with extremely high serum levels of estradiol (> 4300 pg/ml). Apart from the typical vasomotor symptoms, she did not have any complaints nor an aberrant physical examination. The blood sample was initiated by her local gynecologist just to confirm her menopausal status. Due to the highly elevated estradiol level, she had been sent to a local hospital for a bilateral adnexectomy expecting an estradiol producing ovarian neoplasia. The ovarian histology was completely unsuspicuous. After surgery, the estradiol levels did not change at all and the patient was sent to our institution for further analysis. Clinically, the patient complained of the typical symptoms of hormone deficiency like hot flushes, night sweats, vaginal dryness, sexual dysfunction and poor performance. Her last menstruation had occurred one year ago. Any hormone, drug or alcohol consumption was denied. Physical examination showed a vaginal atrophy and a postmenopausal vaginal smear. Hormonal analysis was repeated at our institution and revealed a high serum estradiol level above range (> 4300 pg/ml). The first test (immunoassay) was done using a biotinylated rabbit-derived polyclonal antibody for estradiol. However, testing other sex steroids and glycoprotein hormones indicating an ovarian or extra-ovarian production, typical blood levels of a postmenopausal woman were found (estrone 13.4 ng/l; progesterone < 0.10 ng/ml; testosterone < 0.03 ng/ml; FSH 70.8 mIE/ml, LH 30.5 mIE/ml; inhibin B < 10 ng/l; AMH < 0.08 ng/ml).

To increase the selectivity of the antibody-ligand reaction, first, an immunoassay with rabbit-derived monoclonal antibodies was performed. However, detecting still high estradiol levels, we changed the detection system to exclude interferences of the immunoassay label. Applying a radioimmunoassay, lower but still elevated estradiol levels (186 pg/ml) could be detected. Another reason for false positive results are cross-reacting molecules which can be induced by structural similarities of the epitopes [8]. Hypothesizing a cross-reactivity with irregular antibodies, we changed the host and applied sheep-derived antibodies which revealed typical postmenopausal estradiol levels (< 5 pg/ml). Our hypothesis of false positive results has been con-
firmed using a fluorescence enzyme immunoassay (FEIA) with specific IgG antibodies against rabbit tissue. Here, a high level of anti-rabbit IgG antibodies could be detected (> 200 mg/l; reference < 30 mg/l) indicating circulating specific anti-rabbit IgG antibodies in the patient.

Asking for her confined domestic circumstances, she reported about breeding rabbits.

Discussion

Clinical diagnostics require an accurate measurement of hormone levels. In postmenopausal women, even high sensitive methods like liquid chromatography-mass spectrometry (LC-MS) are performed to detect estradiol concentrations at low picomolar levels [9]. To date, a variety of immunoassays have been established, mostly monoclonal or polyclonal antibody assays [10]. However, applying automated immunoassays, non-specific results and method-specific bias have to be considered. Interferences can be induced by structural similarities of different epitopes (in the case of polyclonal primary antibodies), by unspecific binding of the detection system or when endogenously produced molecules exist that are structurally similar to the principal analyte (Fig. 1).

In this case, we report a cross-reactivity caused by Human Anti-Animal Antibodies (HAAA). HAAA are immunoglobulins (Ig) which can be induced by longtime contact to animals. These HAAA are also known to induce a cross-reactivity within immunoassays [11, 12]. Detecting false positive results, a cross-reactivity with the host serum can be assumed and should be excluded by changing the host. If the cross-reactivity is not found any more by changing the host, the presence of irregular antibodies can be assumed and can be confirmed by FEIA.

Apart from method-specific bias of the immunoassays elevated estradiol levels in postmenopausal women can be due to hormone and drug consumption. Rare cases of estrogen producing tumors like sex cord stromal tumors including granulosa cell tumors, thecoma and Sertoli stroma cell tumors are described. Thereof, the most common hormonally active are granulosa cell tumors. Granulosa cell tumors are potentially malignant sex cord stromal tumors of the ovary and account for 2% of all ovarian tumors. They can be divided into adult (95%) and juvenile (5%) types based on histological findings [13]. Due to their hormone production, most of them can be detected in an early stage. Germ cell tumors (ovarian carcinoma) as well as Brenner tumors are only rarely associated with endocrine manifestations [14–16]. Feminizing adrenal tumors are rare [17] and comprise adrenocortical adenoma and carcinoma which can lead to an increase of estradiol levels [18, 19]. Several reports demonstrated the presence of high aromatase enzyme activity and an overexpression of CYP19 mRNA [20, 21]. In adrenocortical adenoma, estradiol peaks could even be found at 120 pg/ml [22]. Apart from neoplasia, liver cirrhosis can provoke an increase in estradiol (up to 60 pg/ml) [23, 24]. Finally, contamination of nutrition (e.g. Fusarium toxin-contaminated maize) can induce hyperestrogenism [25].

Generally, measurement of estradiol levels in postmenopausal women is not considered and rarely reveals clinical benefits. If clinical symptoms are not in line with the detected hormone levels, exclusion of method-specific bias or interferences of the test system should be considered. Otherwise misinterpretation of the results can lead to unnecessary interventions – as observed in our case.

Authors’ Roles

IM drafted the manuscript and performed the data collection. MK participated in the experimental concept and carried out the laboratory tests. MG revised the manuscript. CK participated in editorial support; SES participated in the design of the study and revised the article. All authors read and approved the final manuscript.

Conflict of Interest

None.

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