Low levels of 17-β-oestradiol, oestrone and testosterone correlate with severe evaporative dysfunctional tear syndrome in postmenopausal women: a case–control study

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ABSTRACT

Aims To evaluate the role of 17-β-oestradiol, oestrone and total testosterone (TT) deficiency in the pathogenesis of severe evaporative dry eye syndrome (DES), investigating the relationship between tear osmolarity, tear film break-up time (TF-BUT), Schirmer test and serum sex hormones in postmenopausal women.

Methods 44 postmenopausal women were recruited for a case–control study: 22 women with severe evaporative DES (Group A) and 22 without DES (Group B). The tests performed included laboratory blood analysis: fasting plasma profile (17-β-oestradiol, oestrone and TT), glucose level and lipid profile. Detailed eye examinations, including corneal and conjunctival staining, tear osmolarity measurement, tear volume and TF-BUT were performed. The Ocular Surface Disease Index Questionnaire was also administered.

Results Values of Schirmer test and TF-BUT in Group A were significantly lower in comparison with Group B (p<0.001). Serum levels of 17-β-oestradiol, oestrone and TT were significantly lower in Group A compared with Group B (p<0.05). In women with severe evaporative DES, the levels of 17-β-oestradiol, oestrone and TT were inversely correlated with the tear film osmolarity (r=-0.7, -0.88, -0.81, respectively).

Conclusions In postmenopausal women with severe evaporative DES, sex hormone levels are lower than control and that tear osmolarity is negatively correlated with sex hormone levels.

INTRODUCTION

Dry eye syndrome (DES) is the most frequent ocular complaint in elderly. It is characterised by eye irritation symptoms, blurred and fluctuating vision, tear film instability, increased tear osmolarity and ocular surface epithelial disease.1

DES may be caused by any disorder that decreases tear production (aqueous tear deficiency) or increases tear evaporation as in case of tear lipid deficiency.2

The meibomian glands (MGs) secrete the lipid layer in human tears and lack of lipid component increases tear evaporation fourfold.3

Notably, DES affects women much more frequently than men. Women are more likely to experience DES during certain periods of significant hormonal alteration, for example, during pregnancy, lactation, oral contraceptive use and after menopause, suggesting that DES may be related to sex hormones.4-6

Receptors for androgens, oestrogens, progesterone and prolactin have been identified in several ocular tissues, including the lacrimal gland and MGs.7-9 Experimental and human studies have demonstrated that androgen levels are essential for normal lacrimal gland function and structural organisation and that prolactin and oestrogens also play important roles in providing an adequate hormonal milieu for optimal lacrimal production.10-12 However, few data exist on the role of sex hormones on lacrimal composition and MGs. Androgen receptors characterise MGs, a type of sebaceous gland, so it is hypothesised that these glands are under hormonal control.13 This point is particularly noteworthy because there is a significantly increased prevalence of meibomian gland disease (MGD) in postmenopausal women. Androgens act on the acinar epithelial cells, which contain receptor messenger RNA and/or androgen receptor protein. These cells respond to androgens by binding the androgen to a specific lipid-producing area on the cell, which then transcribes specified genes to increase the lipid layer distribution over the ocular surface.12 On the contrary, an androgen deficiency or a topical anti-androgen treatment causes a significant decrease in the ocular lipid profile.13 The administration of sex hormones, such as the androgen precursor dehydroepiandrosterone, to the surface of the eye has been found to stimulate MG lipid production and release and prolong the eye’s tear film break-up time (TF-BUT).10 The oestrogen produced after the menopause is primarily from the peripheral conversion of adrenal androgens to oestrogen and occurs in the liver, kidney, brain, adrenal and peripheral adipose tissue.13 The absolute levels of oestrogen are influenced by weight, sex and age. Obese women have a higher concentration of free oestradiol because of decreased sex hormone-binding globulin and an increased rate of aromatisation.18 In postmenopause, androgens continue to be produced by the ovarian stromal and hilus cells in response to the increased levels of circulating lutein hormone.15 There is a 25% increase in testosterone secretion by the postmenopausal ovary; however, circulating levels of testosterone decrease to slightly less than those of premenopausal women. This
The result is sex hormone deficiency, like that occurring in menopause that may alter not only aqueous lacrimal production but also MG function and lipid production leading to increased evaporation rate and occurrence of DES in its evaporative form.\textsuperscript{14–18}

The aim of this study was to examine the relationship between tear osmolarity, tear volume, TF-BUT and sex hormones serum levels (17-\(\beta\)-oestradiol, oestrone and total testosterone (TT)) in postmenopausal women with severe evaporative DES.

**METHODS**

The study was conducted according to a protocol approved by local ethics committee ‘Azienda Ospedaliera Universitaria Vittorio Emanuele Policlinico Catania’ and conformed to the ethical guidelines of the 1975 Helsinki Declaration and the International Conference of Harmonization (ICH) guideline for Good Clinical Practice. Written informed consent was obtained from each woman before entering the study.

**Setting and sample**

The case-control study was performed at Neurovisual Science Technology (NEST) Laboratory of Clinical Ophthalmology, Catania, Italy, and at Ophthalmic Clinic, Department of Medical Surgical Specialties, University of Catania, School of Medicine, Catania, Italy. It was conducted during the period from November 2011 to April 2012 on 44 postmenopausal volunteer women aged 52–63 years (mean age 57±3.9), as defined as in amenorrhoea since at least 1 year, with serum follicle stimulating hormone (FSH) levels higher than 12 mIU/mL and oestrogen levels lower than 15 pg/mL. Volunteers were recruited from a hospital clinic who were later subdivided into two groups: 22 women with severe evaporative DES (Group A) and 22 without DES (Group B). The sample size (at least 20 eyes for each group) was determined from the results of our preliminary study, but were instructed not to change the type of drop and not to use it on the visit day.

Twenty-two volunteer women suffering from severe evaporative DES, due to MGD (stage 4 as reported in the classification of Geerling \textit{et al}\textsuperscript{20} as defined by “a chronic, diffuse abnormality of the MGD, commonly characterised by terminal duct obstruction and/or qualitative/quantitative changes in the glandular secretion” and abnormal TF-BUT) constituted Group A.

Twenty-two women with aqueous deficiency and no evidence of MGD were considered as control group (Group B). For each woman, data from right eye were included in the study.

**RESULTS**

All 44 women were of normal and stable weight (mean weight 71.6 kg, range 60.5–80.6, mean body mass index 22.5 kg/m\(^2\), range 20.0–25.0) and there was no significant difference between Group A and Group B in terms of patient age, years since menopause, weight, plasma lipid profile (triglycerides, total cholesterol and HDL-cholesterol), fasting plasma glucose and average daily intake of nutrients (table 1).

Table 2 shows the serum levels of 17-\(\beta\)-oestradiol, oestrone and TT of the two groups. Sex hormones were significantly lower in women affected by DES compared with control women (\(p<0.05\)).

Figure 1 shows the correlation between sex hormone levels and tear osmolarity. The levels of 17-\(\beta\)-oestradiol, oestrone and TT were inversely correlated with tear film osmolarity (\(r\) values=\(-0.7, -0.88, -0.81\), respectively). A positive correlation has been demonstrated between Schirmer values (\(r\) values=0.69, 0.83, 0.74) (figure 2) as well as TF-BUT (\(r=0.76, 0.87, 0.75\)) (figure 3) and sex hormone levels.

Statistical analysis was performed using Systat V13.0 (SSPS, Chicago, Illinois, USA).

At enrolment, physical and clinical examinations were performed and medical, surgical and medication histories were assessed to ensure study eligibility on the basis of inclusion and exclusion criteria. Women with a history of eye disease or previous surgery; use of hormonal, systemic or topical anti-inflammatory and/or antibiotic therapies; use of an experimental drug 30 days or less prior to study entry and use of tobacco were excluded from the study.

The evaluations included the following examinations:

- **Body weight:** patients were weighed after urinating, wearing only underclothing (which was accurate to ±0.1 kg) for all measurements.
- **Laboratory blood analysis:** blood samples were drawn at 8:00 for determination of fasting plasma hormone profile (17-\(\beta\)-oestradiol, oestrone and TT), glucose level and lipid profile (serum triglycerides, total cholesterol and high-density lipoprotein (HDL)-cholesterol).
- **Nutrient analysis:** patients individually completed a questionnaire (Frequency of Food Consumption) composed of a total of 144 foods to establish the type of nutrients usually consumed.

Detailed eye examinations, including corneal staining with fluorescein (evaluated using the National Eye Institute method, a standardised scale (0–3) for each of the five regions of the cornea: central, inferior, nasal, superior and temporal) and conjunctival staining with lissamine green (evaluated using the Oxford Schema (0–4) for three regions of the conjunctiva: central, nasal and temporal). During anterior segment slit-lamp examination, inspection of eyelid margins and conjunctiva was done carefully taking note of hyperaemia of the bulbar and palpebral conjunctiva, lid wiper epitheliopathy, lid parallel conjunctival folds and keratinisation of the MG ductal epithelium (ie, orifice metaplasia) and of the lids.

- **Tear osmolarity:** each patient was seated with head back and eyes upwards towards the ceiling, and 50 mL of tear sample was collected from the lower eyelid and the line of moisture along the inner eyelid margin, near the lateral canthus of both eyes, using Tearlab Osmolarity System Pen (TearLab Inc, San Diego, California, USA), holding a single-use Test Card. Then, the tear film osmolarity of each sample was measured by using the Tearlab Osmolarity System Reader (TearLab Inc). Normal values range between 289 and 308 mOsm/L, in a DES patient range between 312 and 378 mOsm/L.\textsuperscript{19}

- **Schirmer test:** performed for 5 min with anaesthesia with the patient’s eyes closed and calculated in millimetres.

- **Fluorescein TF-BUT:** measured as the time from a patient’s blink to the appearance of the first dry spot on the cornea and calculated in seconds.

- **Ocular Surface Disease Index (OSDI) Questionnaire:** Patients were authorised to use artificial tears during the study, but were instructed not to change the type of drop and not to use it on the visit day.
Figure 4 shows the comparison between the tear film osmolarities in the two groups. The mean tear osmolarities in Groups A and B were 378.77±8.48 and 309.18±13.47 mOsmol/L, respectively. Tear osmolarity was significantly higher in Group A in comparison with controls (p<0.001; t=20.494). There were statistically significant differences between the two groups in terms of Schirmer test value (p<0.001; t=142.663) (figure 5) and TF-BUT value (p<0.001; t=297.419) (figure 6).

Group A also tended to have worse scores than Group B on self-reported symptoms as measured by OSDI (43.3±16.42 vs 28.5±18.34; p<0.001). Differences in lissamine green, fluorescein staining and hyperaemia on the ocular surface (lid margin, MGs and conjunctiva) between the two groups were statistically significant (p<0.001).

DISCUSSION

Several data in literature suggest a significant influence of gender and sex hormones on the physiology of the lacrimal gland and that they are involved in the pathogenesis of DES. Interestingly, DES occurs mainly in women and its appearance is frequently related to marked variations in serum sex hormone levels like menopause.21

The results of this study showed that the serum levels of 17-β-oestradiol, oestrone and TT were significantly lower in women affected by DES compared with control. Moreover, the levels of all sex hormones investigated were inversely correlated with tear osmolarity (r values = −0.7; −0.88, −0.81, respectively). Tear film hyperosmolality, a recognised component for the definition of dry eye by Dry Eye WorkShop,1 recently was found to be the single best marker of disease severity and marker of disease progression.22

This inverse correlation of sex hormone levels and tear film osmolarity adds further evidence on the role of sex hormone levels in the pathogenesis of DES. In particular, sex hormone deficiency may cause MGD. Meibum lipids promote the tear film stability and prevent tear evaporation, maintaining health and integrity of the ocular surface.23 24 The dysfunction of this gland leads to lipid insufficiency, tear film instability, hyperosmolality and increasing evaporation of the tear film.25

MGD leads to lipid insufficiency, increasing evaporation, instability and hyperosmolality of the tear film. Our findings are...
in accordance with previous evidence indicating that both testosterone and 17-β-Oestradiol influence and regulate MG integrity and functions, enhance the quality and/or quantity of lipids produced by MGs and promote the formation of the tear film’s lipid layer in humans.\(^{26,27}\)

The data from this study suggest that deficiency in sexual hormones may cause not only a reduction in tear production leading to aqueous-deficient dry eye, but also a dysfunction in MG function determining an evaporative dry eye. The mRNA for sex steroid receptors has been demonstrated in MGs in humans suggesting that these glands are target organs for androgens, oestrogens and/or progestins.\(^{7,28}\) Sex steroid deficiency may alter MG function in different ways by altering secretion of the lipid components or also acting throughout an inflammatory

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**Figure 2** Direct correlation between plasma hormone levels (17-β-oestradiol, oestrone and testosterone) and Schirmer test values \((r=0.69; 0.83; 0.74, \text{respectively})\).

**Figure 3** Direct correlation between plasma hormone levels (17-β-oestradiol, oestrone, and testosterone) and tear film break-up time (TF-BUT) values \((r=0.76; 0.88; 0.76, \text{respectively})\).
process. In particular, Suzuki et al demonstrated that testosterone stimulates genes related to lipid metabolic pathways and suppresses genes regulating epithelial keratinisation. In women with complete androgen insensitivity syndrome due to androgen receptor dysfunction, an alteration in neutral and polar lipid patterns of human MG secretion along with a keratinisation of the MG ductal epithelium (ie, orifice metaplasia) and the lid has been observed. Notably, in these women there was a significant increase in signs and symptoms of dry eye. Data in literature show that serum sex hormone levels in postmenopausal women are influenced by age, years since menopause and weight and that oestradiol and androgen levels are negatively correlated with age. Notably, all these parameters were not significantly different in the two groups of women. However, it must be underlined that elevated androgen levels may also be responsible for MGD. In fact, a higher incidence of evaporative dry eye (EDE) has been observed in hyperandrogenic women affected by polycystic ovarian syndrome. This allows us to speculate that lacrimal function as well as other functions in the visual system is profoundly influenced by sex hormones and that a fine equilibrium between androgens and oestrogens is needed for a normal function.

In conclusion, this study demonstrates that postmenopausal women are at risk for DES and that low levels of sex steroids, both androgens and oestrogens, are related to the occurrence of dry eye in its evaporative form. These findings allow for the hypothesis that hormone treatment may be useful to increase tear production and restore the physiologic layer composition of tears.

A randomised clinical trial is needed to demonstrate the effectiveness of this approach.

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