Mosaic Turner syndrome and pituitary microadenoma in patient with polyglandular autoimmune syndrome type II

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Introduction

Polyglandular autoimmune syndrome type II (PGA-II) is the most common immunoenocrinopathy syndrome, characterized by the obligatory occurrence of Addison disease in combination with thyroid autoimmune diseases and/or type 1 diabetes mellitus. This case report presents coexistence of mosaic Turner syndrome and pituitary micoradenoma in patient with PGA-II.

Case report

A 30 year old women underwent in vitro fertilization (IVF) for four times, with no success (always poor ovarian response, double embryo transfer). Hashimoto thyroiditis and subclinical hypothyreoidismus were diagnosed. Levothyroxine substitution was started. Repeatedly elevated prolactin level in the morning, but preserved circadian rhythm and daily values in the referent range. NMR sellar region shown microadenoma in the right half. No adequate cortisol answer in insulin tolerance test, but normal prolactine and growth hormone. In thyrotropin relasing hormone test no paradoxal response. Husband has the oligoasthenospermia, the varicocele surgically removed and normal karyotype (46, XY). Patient's karyotype testing shown a mosaic monosomy X (46, XX/45, X0), with 5% of analyzed cells caracterized by monosomy X, but no syndrome phenotype characteristics, entered puberty at the time, regular menstrual cycles and no echocardiography dysgenetic ovarian caracteristics. During the last IVF no follicle on folitropine stimulation was found, and for the first time higher FSH value and lower AMH value were found, indicating premature ovarian insufficiency; antiovarian antibodies were negative excluding immune-mediated process. A year later, menstrual cycles became irregular.

Case report

Two years later she presented with sings of hypocorticism- artralgia, hyperpigmentation, fatigue, hypotension and low cortisol level, but normal electrolyte level, hydrocortisone substitution was started (Figure 1,2). PTH was in referent range. Positive anti GAD and anti IA2 antibodies were demonstrated, oral glucose tolerance test was normal. Patient is now in oocyte donation process.

Conclusion

Oocyte donation may offer solution to women with multiple autoimmune disorder causing infertility.