PŮVODNÍ PRÁCE ORIGINAL PAPER

doi: 10.14735/amcsnn2018199

Serum anti-Müllerian hormone levels in multiple sclerosis – a multicenter case-control study

Hladiny anti-Mülleriánského hormonu v séru při roztroušené skleróze – multicentrická případová kontrolní studie

Abstract

Introduction: Multiple sclerosis (MS) is considered as an inflammatory demyelinating disease with a 3-fold risk in females of child-bearing age that imposes deleterious pregnancy-related concerns for patients. We aimed to clarify the correlation of anti-Müllerian hormone (AMH) level with clinical and gynaecological characteristics of relapsing-remitting MS (RRMS) patients. Methods: Fifty consecutive RRMS female patients and 50 age-matched healthy controls from among the hospital staff were enrolled in the study. The serum AMH levels of all subjects were evaluated by ELISA. T-tests and Mann-Whitney U tests were used for the quantitative variables. Pearson's chi-squared test and Fisher's exact test were used for qualifying variables. Regression analysis was performed for AMH values. Results: The mean \pm SD AMH level for the case group was 3.12 \pm 3.6 and for the control group was 3.65 \pm 2.06. Regression analysis detected a significant difference in AMH values between the case and control groups (p = 0.034). There was no significant influence for disease activity and therapy on AMH level in the patients. Age was the only demographic factor that predicted the AMH values in both groups (p = 0.001). Conclusion: The results showed that serum AMH values were significantly lower in MS patients; however, this finding does not necessarily have an effect on fertility.

Souhrn

 $\dot{U}vod$: Roztroušená skleróza (RS) je považována za zánětlivé demyelinizační onemocnění s trojnásobným rizikem výskytu u žen v plodném věku, které u pacientek vzbuzuje nepříznivé obavy související s těhotenstvím. Naším cílem bylo objasnit vzájemný vztah mezi hladinou anti-Mülleriánského hormonu (AMH) a klinickými a gynekologickými charakteristikami pacientek s relaps-remitentní RS (RR RS). Metody: Do studie bylo postupně zařazeno 50 pacientek s RR RS a 50 zdravých kontrolních subjektů odpovídajícího věku z řad nemocničního personálu. U všech subjektů byly vyhodnoceny sérové hladiny AMH metodou ELISA. Pro vyhodnocení kvantitativních proměnných byly použity t-test a Mann-Whitneyův U test. Pro kvalifikaci proměnných byly použity Pearsonův chí-kvadrát test a Fisherův exaktní test. U hodnot AMH byla provedena regresní analýza. Výsledky: Průměrná (\pm SD) hladina AMH u skupiny pacientek činila 3,12 \pm 3,6 a u kontrolní skupiny 3,65 \pm 2,06. Regresní analýza odhalila významný rozdíl v hodnotách AMH mezi případovou a kontrolní skupinou (p = 0,034). U pacientek nebyl zjištěn žádný významný vliv aktivity onemocnění a léčby na hladinu AMH. V obou skupinách byl jediným demografickým faktorem, který predikoval hodnoty AMH, věk (p = 0,001). Závěr: Z výsledků vyplývá, že hodnoty AMH v séru byly výrazně nižší u pacientek s RS, avšak toto zjištění nemá nutně vliv na plodnost.

The authors wish to thank Rasoul-e-Akram Hospital Clinical Research Development Center and Ms Zahra Torabi-Goodarzi (clinical psychologist) for their technically-supported implementation of the project.

The authors declare they have no potential conflicts of interest concerning drugs, products, or services used in the study.

Autoři deklarují, že v souvislosti s předmětem studie nemají žádné komerční zájmy.

The Editorial Board declares that the manuscript met the ICMJE "uniform requirements" for biomedical papers.

Redakční rada potvrzuje, že rukopis práce splnil ICMJE kritéria pro publikace zasílané do biomedicínských časopisů.

M. Mehrpour¹, F. Taherian², P. Arfa-fatollahkhani², A. N. Moghadasi³, S. Mokhtar⁴, H. Keivani⁵

- ¹ Department of Neurology, Firoozgar Hospital, Iran University of Medical Sciences, Tehran, Iran
- ² Department of Neurology, Rasoul-e-Akram Hospital, Iran University of Medical Sciences, Tehran, Iran
- ³ MS Research Center, Neuroscience Institute, Tehran University of Medical Sciences, Tehran, Iran
- ⁴ Department of Infertility, Avicenna Infertility Center, Tehran, Iran
- ⁵ Department of Virology, Iran University of Medical Sciences, Tehran, Iran

\boxtimes

Fatemeh Taherian, MD
Department of Neurology
Rasoul-e-Akram Hospital
Iran University of Medical Sciences
Niyayesh St., Satarkhan St.
Tehran, Iran
e-mail: research.rasoul@gmail.com

Accepted for review: 27. 7. 2017 Accepted for print: 12. 2. 2018

Key words

anti-Müllerian hormone – infertility – multiple sclerosis

Klíčová slova

anti-Mülleriánský hormon – neplodnost – roztroušená skleróza

Introduction

Multiple sclerosis is an inflammatory, neurodegenerative disease affecting the CNS. The first manifestations commonly appear within the second and third decades of life. There is a 3-fold level of risk in females [1–3], making fertility and pregnancy challenging concerns for MS patients.

Limited data are available on fertility in MS patients, but a decrease in the number of children and reduced spontaneous fertility has been reported in several studies. Auto-immunity, sexual disturbances, and ovarian neurologic dysfunction are recognized as suspicious etiologies [3–5]. Moreover, studies evaluating MS patients – treated with immunosuppressive or immunomodulating agents like mitoxantrone and interferon β , resp. – have uncovered diminished follicular reserves and fertility in the treatment groups compared to the agematched healthy controls [6,7].

Recently, endocrine disturbances have been suggested as the pathology of MS infertility. Among the different biomarkers, one of the most reliable predictors of the ovarian reserve is the anti-Müllerian hormone (AMH). This is a dimeric glycoprotein expressed from 36 weeks of gestation that dramatically increases during puberty and persists to the age of 30 years with a gradual decline to menopause. As AMH is a relatively cycle-independent hormone expressed by the antral follicles, there is a strong correlation between serum AMH level and the number of antral follicles [8–11].

Some studies have highlighted the role of AMH in MS infertility [9–11]. One pilot study detected no differences between AMH values in relapsing-remitting MS (RRMS) cases and age-matched healthy controls; however, they found lower AMH levels and ovarian reserves in patients with higher MS activity [10]. Conversely, another recent case control study reported significantly lower AMH values in RRMS patients compared to the control group [11]. The present study aimed to clarify the correlation of AMH levels with clinical and gynaecological characteristics of RRMS patients in greater detail

Materials and methods

This analytic case-control study was carried out from June 2014 through January 2015. Fifty consecutive RRMS female patients – admitted at the Hospitals in Tehran, Iran – were enrolled along with 50 age-matched healthy controls from among the hospital staff.

The exclusion criteria for both groups were a diagnosis of non-RR type MS, being over 35 years old, being pregnant or breastfeeding, undertaking infertility treatments and any autoimmune disease or abnormal thyroid, kidney or liver function.

Study design

All patients were interviewed by an expert to gather complete demographics and the clinical, obstetrical and gynaecological histories of all subjects, including age, body mass index (BMI), cigarette use, age of menarche, menstrual cycle regularity,

dysmenorrhea, amenorrhea, marital status, number of children, desire to bear a child, infertility history and treatment. Also recorded were the method of contraception used, oral contraceptive pill use, abortion history, disease history in terms of duration, type and treatment (disease-modifying therapy; DMT), pulse therapy with immunosuppressive agents, and untreated status (defined as no treatment with a DMT during the 16 weeks prior to AMH evaluation). The Expanded Disability Status Scale (EDSS) score of the patients was obtained by an experienced neurologist.

Sample collection and ELISA

Venous blood samples (3 cc) were collected from the antecubital fossa and maintained in heparinized tubes at room temperature. They were transferred to the ELISA laboratory in less than 1 hour and centrifuged. Laboratory cut points were 0.8–1.5 ng/mL for ages 15–24, 0.8–10 ng/mL for ages 25–29, 0.4–9.3 for ages 30–34, and 0.15–7.3 ng/mL for ages 35–39.

Statistical analysis

Between-group comparisons were applied using the t-test for quantitative variables, which were normally contributed. The Mann-Whitney U test was assigned for abnormally contributed quantitative variables. Pearson's chi-square test and Fisher's exact test were performed to analyse the qualifying variables. Regression analysis was performed on ELISA results. Descriptive analysis of the data was expressed as frequency and mean \pm SD; p < 0.05 was considered statistically significant. Statistical analysis was performed using IBM SPSS (Version 22, SPSS Inc., Chicago, IL, USA) and R software (The R Foundation, Vienna, Austria).

Ethics

This work was approved by the Local Medical Ethics Committee of Iran University of Medical Sciences.

Results

Demographic and gynaecological characteristics

Demographic and gynaecological characteristics of all subjects are listed in Tab. 1. None of the subjects have a history of using illegal drugs or smoking cigarettes. Of the 100 participants, one used an oral contraceptive pill for contraception and one was trying to conceive. Importantly,

Tab. 1. Demographic and gynaecologic characteristics of RRMS patients and healthy controls.

| | group | | - p value |
|---------------------------------|-------------|-------------|-----------|
| | control | case | p value |
| age (years); mean (SD) | 30.12 (3.3) | 28.86 (5.5) | 0.633 |
| menarche age (years); mean (SD) | 13.22 (1.6) | 12.88 (2.5) | 0.672 |
| BMI (kg/m²); mean (SD) | 21.7 (2.63) | 23.9 (3.7) | 0.001 |
| marital status, married, n (%) | 16 (32) | 27 (54) | 0.043 |
| children, n | 9 | 24 | 0.013 |
| abortion, n | 2 | 9 | 0.190 |
| infertility, n (%) | 0 | 3 (6) | 0.242 |
| dysmenorrhea, n (%) | 18 (36) | 19 (38) | 0.836 |
| amenorrhea, n (%) | 5 (10) | 5 (10) | 1.000 |

 $BMI-body\ mass\ index;\ n-number\ of\ patients;\ RRMS-relapsing-remitting\ multiple\ sclerosis;\ SD-standard\ deviation$

Tab. 2. Correlation of demographic and gynaecologic factors and AMH level.

| variable | coefficient | p value |
|---------------|-------------|---------|
| intercept | 8.854 | < 0.001 |
| group (case)* | -1.056 | 0.034 |
| age | -0.172 | 0.001 |
| | | |

AMH – anti-Müllerian hormone * reference category is "control"

none of the healthy controls were infertile, and the infertility rate of the participants was 6%. Nine participants in the case group and two people in the control group had an abortion; however, the Pearson chi-square analysis revealed no significance for both (p = 0.24, p = 0.19, resp.).

The BMI, marital status and number of children were significantly different between the case and control groups. The t-test results revealed that BMI was significantly higher in the case group (p = 0.001). Interestingly, the case group had a higher number of children than their age-matched controls, which was statistically significant in chi-square analysis (p = 0.013). Of the 50 participants 27 (54%) and 16 (32%) were married in the case group and control group, resp., which was significant at p = 0.043 (Tab. 1).

Clinical characteristics of patients

In the case group, three participants were new cases of RRMS in whom the treatment had not commenced. The level of medication was categorized as: 1. low frequency and included Avonex (22 patients); 2. high frequency and included glatiramer acetate, Betaseron and Rebif (22 patients); 3. the last group used fingolimod (3 patients). The EDSS scores of patients had a mean \pm SD of 1.80 \pm 0.95, with a maximum of 4.5 and minimum of 0. The duration of the disease in the case group was 40 days to 14 years (3.7 \pm 3.10 years).

AMH analysis between groups

In the case group, the mean \pm SD for AMH level was 3.12 \pm 3.6 and in the control group was 3.65 \pm 2.06. The serum AMH levels were normal in 38 cases. However, analysis revealed lower AMH levels in nine patients and higher levels in three patients among the case group. Only two patients from the control group had AMH results outside

Tab. 3. Influence of disease activity and therapy on AMH level.

| coefficient | p value |
|-------------|---|
| 2.566 | < 0.001 |
| | |
| -0.776 | 0.758 |
| 1.771 | 0.480 |
| 0.311 | 0.802 |
| 1.393 | 0.629 |
| 0.051 | 0.797 |
| 0.037 | 0.955 |
| | 2.566 -0.776 1.771 0.311 1.393 0.051 |

AMH – anti-Müllerian hormone; EDSS – Expanded Disability Status Scale

the normal range; one was above and one was below the normal range. Regression analysis detected a significant difference in AMH values between the case and control groups (adjusted $R^2 = 0.05403$; p = 0.034). There was no significant difference for age of subgroups between groups (Tab. 2).

Correlation of demographic and gynaecologic characteristics

The number of children, infertility rate, number of abortions and marital status were not statistically significant predictors of AMH values according to regression analysis (p > 0.2). In the backward method, age was the only demographic factor that significantly predicted AMH values in both groups (adjusted $R^2 = 0.05403$; p = 0.001). Analysis showed that advanced age was a significant predictor of AMH level. Details are available in Tab. 2.

Influence of disease activity and therapy

Table 3 shows the results of regression analysis for treatment, duration of disease and EDSS score versus AMH values in the case group. There was no significant influence of the disease activity or therapy on the AMH levels found (adjusted $R^2 = -0.1149$).

Discussion

This cross-sectional case-control study evaluated the relationship of AMH values and demographic, gynaecological and clinical features of RRMS patients in comparison to age-matched healthy controls. The findings showed a significantly lower serum AMH levels in the case group

compared to the controls (p = 0.034). Notably, the results revealed no correlation between the low AMH level and the rate of infertility or abortion in the case group. The results suggest that advanced maternal age and MS were the only significant predictors of decreased AMH level.

A recent cross-sectional pilot study evaluated 76 RRMS patients and 58 controls and reported a lower mean of the AMH levels for the RRMS patients; however, it also detected very low AMH levels in patients not currently under treatment with a DMT [11]. In contrast, the current study revealed no effect for the type of MS therapy on the AMH values. Another recent case-control study found no significant difference in AMH values between the RRMS patients and healthy populations. Moreover, they found that higher disease activity might result in poorer ovarian reserves in RRMS patients [10]. The results of the present study found no potential effect for EDSS score and duration of the disease on AMH levels as a reliable predictor of ovarian reserves [8].

More studies are needed to elucidate the correlation between the clinical features of the disease and fertility in MS patients. In keeping with previous studies, the results illustrated that AMH levels declined significantly with age in both the case and control groups. As shown, AMH levels peaked dramatically during puberty, persisted till the age of 30, and gradually declined to menopause [8].

A higher mean BMI was recorded in the case group as compared with the control group. In line with previous studies, high BMI and obesity are significant and possible risk factors for MS that have deleterious

^{*} reference category is "no drug"

^{**} reference category is "negative"

SERUM ANTI-MÜLLERIAN HORMONE LEVELS IN MULTIPLE SCLEROSIS – A MULTICENTER CASE-CONTROL STUDY

functional effects on patients [12–14]. Surprisingly, and in contrast with previous studies reporting childlessness in MS patients [3,15], the current findings were consistent with a recent pilot study, showing a higher rate of marriage and number of children in the case group [10]. This difference could be attributed to the lifestyle of the healthy controls, who were primarily members of the hospital staff.

This work was limited by a small sample size because of the large number of exclusion criteria that aimed to achieve exact evaluations without the potential effects of co-morbidity on fertility and AMH values of the participants.

Conclusions

This study showed that serum AMH values decreased significantly in the RRMS group; however, this finding might not have a potential impact on fertility. Because this work was one of the first to evaluate AMH as an influential factor on fertility in MS patients, more studies are needed to produce

practical and effective reproductive programmes for MS sufferers.

References

- 1. Chen YH, Lin HL, Lin HC. Does multiple sclerosis increase risk of adverse pregnancy outcomes? A population-based study. Mult Scler 2009; 15(5): 606–612. doi: 10.1177/1352458508101937
- **2.** Greer JM, McCombe PA. Role of gender in multiple sclerosis: clinical effects and potential molecular mechanisms. J Neuroimmunol 2011; 234(1–2): 7–18. doi: 10.1016/j.jneuroim.2011.03.003.
- **3.** Cavalla P, Rovei V, Masera S et al. Fertility in patients with multiple sclerosis: current knowledge and future perspectives. Neurol Sci 2006; 27(4): 231–239.
- **4.** Dahl J, Myhr KM, Daltveit AK et al. Planned vaginal births in women with multiple sclerosis: delivery and birth outcome. Acta Neurol Scand Suppl 2006; 183: 51–54.
- **5.** Hellwig K, Brune N, Haghikia A et al. Reproductive counselling, treatment and course of pregnancy in 73 German MS patients. Acta Neurol Scand 2008; 118(1): 24–28. doi: 10.1111/j.1600-0404.2007.00978.x.
- **6.** Cocco E, Sardu C, Gallo P et al. Frequency and risk factors of mitoxantrone-induced amenorrhea in multiple sclerosis: the FEMIMS study. Mult Scler 2008; 14(9): 1225–1233. doi: 10.1177/1352458508094882.
- 7. Cil AP, Leventoğlu A, Sönmezer M et al. Assessment of ovarian reserve and Doppler characteristics in patients with multiple sclerosis using immunomodulating drugs. J Turk Ger Gynecol Assoc 2009; 10(4): 213–219.

- **8.** Bhide P, Homburg R. Anti-Müllerian hormone and polycystic ovary syndrome. Best Pract Res Clin Obstet Gynaecol 2016; 37: 38–45. doi: 10.1016/j.bpobgyn.2016.03.004.
- **9.** Thöne J, Kleiter I, Stahl A et al. Relevance of endoglin, IL-1 α , IL-1 β and anti-ovarian antibodies in females with multiple sclerosis. J Neurol Sci 2016; 362: 240–243. doi: 10.1016/j.jns.2016.01.057.
- **10.** Sepúlveda M, Ros C, Martínez-Lapiscina EH et al. Pituitary-ovary axis and ovarian reserve in fertile women with multiple sclerosis: a pilot study. Mult Scler 2016; 22(4): 564–568. doi: 10.1177/1352458515602339.
- **11.** Thöne J, Kollar S, Nousome D et al. Serum anti-Müllerian hormone levels in reproductive-age women with relapsing-remitting multiple sclerosis. Mult Scler 2015; 21(1): 41–47. doi: 10.1177/1352458514540843.
- **12.** Cambil-Martín J, Galiano-Castillo N, Muñoz-Hellín E et al. Influence of body mass index on psychological and functional outcomes in patients with multiple sclerosis: a cross-sectional study. Nutr Neurosci 2016; 19(2): 79–85. doi: 10.1179/1476830514Y.0000000156.
- **13.** Kvistad SS, Myhr KM, Holmøy T et al. Body mass index influence interferon-beta treatment response in multiple sclerosis. J Neuroimmunol 2015; 288: 92–97. doi: 10.1016/j.jneuroim.2015.09.008.
- **14.** Taylor KL, Hadgkiss EJ, Jelinek GA et al. Lifestyle factors, demographics and medications associated with depression risk in an international sample of people with multiple sclerosis. BMC Psychiatry 2014; 14: 327. doi: 10.1186/s12888-014-0327-3.
- **15.** Nielsen NM, Jørgensen KT, Stenager E et al. Reproductive history and risk of multiple sclerosis. Epidemiology 2011; 22(4): 546–552. doi: 10.1097/EDE.0b013e31821c7adc.

Soutěž o nejlepší práci publikovanou v časopise Česká a slovenská neurologie a neurochirurgie

V roce 2018, stejně jako v předchozích letech, probíhá soutěž o nejlepší článek časopisu ČSNN. Zařazeny budou práce otištěné v číslech 2018/ 1–6. Vítěze vyberou členové redakční rady a bude vyhlášen u příležitosti 32. slovenského a českého neurologického sjezdu v Martine.

Předem děkujeme všem autorům za zaslané příspěvky.