Polymorphism study of MTHFR 677C→T and its correlation with oxidative stress and their influence on female infertility in Erbil – Iraq

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Abstract:
This research includes a study of Methylenetetrahydrofolate reductase gene’s allele 677C→T and its correlation with oxidative stress and their impact on female infertility. Fifty infertile women with the range age (23-42) years and twenty five fertile women with the range age (22-39) years as control group living in Erbil city were selected. The serum level of Malondialdehyde (MDA), superoxide dismutase (SOD), prolactin hormone (PRL), Luteinizing hormone (LH), Thyroid stimulating hormone (TSH), Triiodothyronine hormone (T3), and Thyroxine hormone (T4) were measured, also a body mass index (BMI) was calculated. A restriction enzyme (Hinf1) was used to improve the mutation in DNA bands of infertile women. The results showed significant increases in MDA level, SOD activity, BMI, PRL, LH, TSH, and T4 in patients women compared with the control group. The results show non-significant differences in T3 hormone levels. The results also show a mutation in DNA bands of infertile women compared with fertile control group.

Key words: Female infertility (FI), Methylenetetrahydrofolate reductase (MTHFR) gene, Oxidative stress (OS).

Introduction:
Infertility is the most important clinical problem, affecting people around the world psychosocially and physically. Infertility is a disease of the reproductive system defined by the failure to achieve a clinical pregnancy after 12 months of regular intercourse without contraception [1]. Infertility can be divided into two major types: primary infertility which characterizes women who have never been able to conceive a pregnancy and secondary infertility which describes those who had at least one successful pregnancy but have not been able to obtain another [2]. Causes of female infertility (FI) include Endometriosis, Polycystic ovary syndrome (PCOS), Ovulatory disorders, tubal factors, age factor, body weight and obesity, and lifestyle factors [3].
Oxidative stress (OS) is caused by an imbalance between pro-oxidants and antioxidants, excessive amounts of reactive oxygen species (ROS) generation, nevertheless, may curb the body's characteristic antioxidant protection system, making an environment inappropriate for typical female physiological reactions. This can be promote to a numerous of reproductive ailments comprising endometriosis, PCOS, tubal illnesses, spontaneous abortion and unexplained infertility [4]. The role of ROS and FI has been a subject of large interest and research over the last decade. Antioxidants and ROS seem to have a physiological role in reproductive processes, including, fertilization, oocyte maturation, luteal regression, and endometrial shedding. Macrophages, neutrophils and granulosa cells in Graafian follicles are a source of ROS that are balanced by antioxidants [5]. OS has been recommended to be causal in etiologies such as endometriosis, tubal, peritoneal and unexplained infertility and even PCOS. The irregularities in the tubal peritoneal or endometrial environment which result in infertility are moderated by the generation of excessive pro-oxidants as obvious by elevate levels of ROS from the fluid evaluation, higher concentrations of ROS in these environments may have harmful effects on the spermatozoa, oocytes, sperm-oocyte interaction and embryos, both in the Fallopian tube and the peritoneal cavity, recent studies observing an increase in DNA damage in the leukocytes cells from women with PCOS. There was also an increased capability to OS induced damage in these women with PCOS, OS as is being implicated in the pathogenesis of PCOS may possibly explain the link with long-term complications of PCOS containing cardiovascular disease and malignancy [6]. Methylenetetrahydrofolate Reductase (MTHFR) is a protein-coding gene; it has a role in both folate and homocysteine metabolisms by catalyzing the conversion of 5,10-methylenetetrahydrofolate to 5-methylenetetrahydrofolate, using Flavin adenine dinucleotide (FAD) as a cofactor and regulating the metabolism of folate that affects DNA synthesis and methylation [7]. There are two common alleles of MTHFR gene, 677C→T and 1298 A→C, which led to amino acid substitutions, 677 C→T allele is present in either heterozygous CT in about 40% or homozygous TT in about 5-15% carrier state of individuals. Many studies showed that this allele may cause DNA hypo methylation that has been related to metabolic syndrome (MetS) and its components [8]. Methylation of DNA includes a covalent modulation of DNA and has been found to affect a diversity process which influences DNA function and integrity; furthermore, methylation of DNA has a critical role in the domination of gene activity [9].

Patients and Methods:
A total number of 50 samples of infertile women (primary infertility) with the range age (23-42) years, that already diagnosed with female infertility by the doctors in Maternity teaching hospital for fertility and I.V.F center in Erbil governorate-Iraq, and 25 samples of fertile women with the range age (22-39) years, as a control group. The period of the work took about six months (from April to October) 2016, with all required information's from both infertile and fertile women. Up to (4-5 ml) of blood was drawn and collected and divided into pre-labelled two groups of tubes, up to 1ml of blood collected in K3-EDTA tubes for DNA extraction and stored at refrigerator, and up to 3-4 ml collected in clot activator and gel tubes for blood serum, the blood samples were cooled on ice until the end of the experiment. Blood separation process was achieved by taking the
clotting blood samples and then centrifuged at 4000 rpm for 5-6 min. Then the blood serum was collected and stored at deep freezing (-8 - -20 °C) for later tests.

Laboratory Measurements:
Estimating the activity of superoxide dismutase (SOD) enzyme:
The method depends on the ability of SOD to inhibit the autoxidation of epinephrine to adrenochrome, the reaction occurs at 30 °C, pH= 10.2, and SOD measured at wavelength 480 nm, absorbance was measured using Spectrophotometer (G10S UV-VIS, Thermo scientific, USA) [10].

Estimate of serum Lipid peroxide (Malondialdehyde (MDA)) levels:
Thiobarbituric acid (TBA) reacts with MDA, which is the final product of lipid peroxidation process, at 100 °C and acidic medium, to produce a pinkish complex product. The product is measured at 532 nm, absorbance was measured using Spectrophotometer (G10S UV-VIS, Thermo scientific, USA) [11].

Hormones measurements:
The hormones levels of Triiodothyronine hormone (T₃), Thyroxine hormone (T₄), Luteinizing hormone (LH), Thyroid stimulating hormone (TSH), and prolactin were measured by (immulite xpi 2000) analyzer.

Genomic DNA extraction:
DNA was extracted from the blood samples using a method described by Geneaid Biotech Ltd., [12].

Genomic DNA concentration and purity:
All the extracted DNA samples were tested by Thermo scientific Nanodrop 1000 spectrophotometer.

Amplification experiment:
All DNA samples were amplified by PCR technique, using a master premixture kit, manufactured by Bioneer corp. [13]. Specific oligonucleotide primers are used for MTHFR 677 C→T gene, the sequence of the Forward primer is 5- CGA AGC AGG GAG CT TGA GGC TG-3 and Revers primer 5- AGG ACG GTG CGGTGA GAG TG-3. Amplification was run in an automated thermocycler (Cg1-96, Corbett research, Australia). The mixture was initially denatured at 94 °C for 5 minutes, followed by (35) cycles for 30 seconds at 94 °C, 30 seconds at 67 °C, 60 seconds at 72 °C, and 10 minutes for a final extension at 72 °C. PCR products were run on 2% agarose gels (Promega, USA) and stained with 1 µg/ml ethidium bromide. The amplified 233 bp PCR products were digested with (Hinf1) (Roche, Germany) restriction enzyme, according to the manufacturer’s instructions. The digested PCR products were separated on 3% agarose gels and visualized with ethidium bromide staining using the gel documentation system (2500 Proxima, ISO Gen, Netherlands) [14].

Body Mass Index (BMI):
BMI was calculated, by measuring both weight and height of the body. The values were calculated as (Kg/m²).

Results:
Table (1) shows the results of MDA, SOD, and BMI for patients and controls. It shows a significant increase in MDA levels of patients (0.03558 ± 0.02012 µmol/l) compared to controls (0.006 ± 0.00285 µmol/l), at a probability (P ≤ 0.05), respectively. It also showed a significant increase in SOD values of patients (0.601 ± 0.052 µmol/min/ml) compared to controls (0.924 ± 0.066 µmol/min/ml) respectively. It also shows a significant increase in BMI values of patients (27.128 ± 2.99 kg/m²) compared to controls (23.904 ± 1.91 kg/m²) at probability (P ≤ 0.05), respectively.
Table (1): Means, SD± of MDA, SOD, and BMI, for patients and control groups.

<table>
<thead>
<tr>
<th>Variants</th>
<th>Mean ± SD</th>
<th>T-test</th>
<th>P ≤ 0.05</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patients</td>
<td>N=50</td>
<td>Controls</td>
<td>N=25</td>
</tr>
<tr>
<td>MDA µmol/L</td>
<td>0.03558 ± 0.02012</td>
<td>0.006 ± 0.003</td>
<td>7.29</td>
</tr>
<tr>
<td>SOD µmol/min/ml</td>
<td>0.924 ± 0.0066</td>
<td>0.601 ± 0.052</td>
<td>8.84</td>
</tr>
<tr>
<td>BMI kg/m²</td>
<td>27.128 ± 2.99</td>
<td>23.9 ± 1.91</td>
<td>4.8921</td>
</tr>
</tbody>
</table>

Table (2) shows the results of the hormones levels of (PRL, LH, TSH, T₄ and T₃) for patients and control groups. It shows a significant increase in PRL hormone levels of patients (34.2088 ± 14.92 ng/ml) compared to controls (10.3516 ± 5.94 ng/ml) respectively, and likewise presented a significant increase in LH hormone levels for patients (14.33 ± 1.8 ml u/ml) compared to controls (7.912 ± 2.32 ml u/ml), and the results indicate a significant increase in TSH hormone levels of patients (7.69 ± 1.55 µIU/ml) compared to controls (3.5968 ± 1.63 µIU/ml), and the results show a significant increase in T₄ hormone levels of patients (164.74 ± 20.27 nmol/l) compared to controls (108.69 ± 28.83 n mol/l) respectively, for T₃ hormone the results showed non-significant change in the hormone levels of patients (3.32 ± 0.72 nmol/l) compared to controls (3.12 ± 0.58 n mol/l) respectively.

Table (2): Mean, SD± hormone levels, for patients and control groups.

<table>
<thead>
<tr>
<th>Hormones</th>
<th>Mean ± SD</th>
<th>t-test</th>
<th>P ≤ 0.05</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patients</td>
<td>N=50</td>
<td>Controls</td>
<td>N=25</td>
</tr>
<tr>
<td>PRL ng/ml</td>
<td>34.2088 ± 14.92</td>
<td>10.3516 ± 5.94</td>
<td>7.68</td>
</tr>
<tr>
<td>LH ml u/ml</td>
<td>14.33 ± 1.8</td>
<td>7.912 ± 2.32</td>
<td>13.2</td>
</tr>
<tr>
<td>TSH µIU/ml</td>
<td>7.69 ± 1.55</td>
<td>3.5968 ± 1.63</td>
<td>10.58</td>
</tr>
<tr>
<td>T₄ n mol/L</td>
<td>164.74 ± 20.27</td>
<td>108.69 ± 28.83</td>
<td>9.76</td>
</tr>
<tr>
<td>T₃ n mol/L</td>
<td>3.32 ± 0.72</td>
<td>3.21 ± 0.58</td>
<td>1.664</td>
</tr>
</tbody>
</table>

Figure (1) shows three bands at (233bp, 176bp and 57bp) in the DNA of patients women, while control group have only one single band at (233bp) in there DNA. The results show significant differences in genotype distribution in patients women compared to healthy women as controls.

Figure (2) shows one band in the DNA of fertile women with no mutation in the MTHFR gene using gel electrophoresis technique.

Discussion:
In this study a significant increase in MDA levels was observed in infertile women; this elevation in MDA levels is due to the increased generation of ROS due to the excess in oxidative damage generated in these patients. In turn, these oxygen species can oxidize various other important biomolecules including
membrane lipids. This increase in MDA levels is in agreement with a study that reported an increase in MDA levels in patients with FI and especially patients with PCOS [15].

An increase in SOD activity levels may be due to excessive amounts of ROS that were generated through oxidative stress (OS), also the increase in SOD activity levels could be due to the response of the increase in OS in women patients with PCOS, dismutation of superoxide results in the development of (H₂O₂). This lead to altering purines and pyrimidines and cause strand breaks in DNA resulting in DNA damage which may lead to gene mutation. A study indicated that the levels of SOD activity were observed to be higher in blood serum of women patients with PCOS [16].

The increase in BMI values in infertile women may be explained by obesity could intervene with ovarian functions and neuroendocrine that affect oocyte quality and thereby embryo development, implantation, and pregnancy outcome; however, the obesity decreasing both of fertility and ovulatory averages in healthy women [17].

The increased levels of PRL in infertile women is due to OS and ROS, which led to infertility due to a delay in the development of endometriosis. The increase in PRL levels could lead to dysfunction of the luteal phase, lack of ovulation and menopause, because of the inhibition effect of PRL on the secretion of GnRH hormone. Many factors could lead to an increase in PRL levels such as cardiovascular diseases, pituitary gland tumors and hypothyroidism [18]. The increase in LH hormone levels possibly indicates that the infertile women could have PCOS and this led to irregularities in menstrual which leads to female infertility (FI) or a dysfunction occurs in the pituitary gland [19]. The increase in serum TSH hormone levels in infertile women could be due to thyroid gland dysfunction or to its diseases such as hypothyroidism which is commonly associated with menstrual irregularities, acute hypothyroidism also led to ovulatory dysfunction and this may be due to many interactions of thyroid hormones with the female reproductive system [20]. In this study the increased in T₄ hormone levels in infertile women may be due to thyroid gland diseases such as hyperthyroidism, hypothyroidism and thyroid gland disorders; hence, reproductive and infertility are connected to defects in the endocrine, immune system or both. These systems are also directly related to the thyroid gland since the most thyroid autoimmunity common causes are hypothyroidism in women of reproductive age, the majority of women with thyroid dysfunction incidence menstrual irregularities, increased pregnancy miscarriage and infertility [21].

In this study, the results show that all the patient women had the three bands, one at 233bp and the two other were on 176bp and 57bp after digested by restriction enzyme (Hinf1), which means that all genotypes were heterozygote (CT) and one of the parents had the mutant gene (T), (Fig.1), while in the controls women (Fig. 2), all the samples were located at 233bp which means that all had wild type (CC) which won’t be cut with restriction enzyme. The results in this study are in agreement with a study indicated a gene mutation in alleles C/T polymorphism [22]. MTHFR 677C→T polymorphism was significantly associated with OS which; in turn, leads to FI, OS that can make DNA damage and folate deficiency. This could make a disorder in the structure of DNA. The allele C/T was observed in patients women; furthermore, this study indicates that
MTHFR gene plays a major role in FI [23].

Conclusion:
Oxidative stress is linked to female infertility due to its influence on many causes that lead to female infertility such as PCOS and endometriosis and raised the risk of female infertility. The results show a significant effect of oxidative stress, especially reactive oxygen species on MTHFR gene and exclusively C677T allele, which result in a mutation in the gene in women patients with female infertility.

References
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دراسة متعددة الاشكال للجين MTHFR 677 C→T وارتباطه مع الإجهاد التاكسدي وتأثيرهما على العقم عند النساء في محافظة أربيل – العراق

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خالد فاروق الراوي**

الخلاصة:
تضمن البحث دراسة ل اليل الجين MTHFR 677 C→T وارتباطه مع الإجهاد التاكسدي وتأثيرهما على العقم عند النساء.

تم اختيار 50 امرأة عقية بمتوسط اعمار (22-42) سنة و 25 امرأة سليمة بمتوسط اعمار (39-22) سنة من محافظة أربيل. تم قياس مستوى السيرم لكل من المالون دايالديهايد (MDA)، هرمون البرولاكتين (PRL)، هرمون الليوتيني (LH)، الهرمون المحفز للدرقية (TSH)، هرمون الثايرون ثلاثي اليود (T3)، هرمون الثايروكسين (T4)، هرمون النايتروكسين (NO)، مراقبة سير النتائج باستخدام الانزيم القاطع (Hinf1) للثبت من الطفرة الجينية في الدنا لدى النساء العقيمات.

أظهرت النتائج ارتفاعا معنويا في مستويات كل من الMDA، LH، TSH، PRL، NO، BMI، T4، T3 بين النساء العقيمات ومجموعة السيطرة، حيث لم تظهر النتائج أي تغير في مستويات هرمون الT3، TSH، PRL، NO، BMI، T4، T3.

يعزى ارتباط الإجهاد التاكسدي بالعقم عند النساء لتأثيره على عدة أسباب والتي تؤدي بدورها للإصابة بالعقم مثل الإصابة بمتلازمة التكيس المبيضي والتهاب بطانة الرحم ونسبة الخطرة للإصابة بالعقم عند النساء.

النتائج تأثيرا معنويا للإجهاد التاكسدي وخاصة اصناف الأوكسيجين الفعالة على جين MTHFR و بصورة خاصة على اليل الجين C677T. أن نتيجة هذا التأثير هو حصول طفرة جينية في الجين MTHFR لدى النساء العقيمات.

الكلمات المفتاحية: العقم عند النساء، جين ميثيلين تيتراهايدروفوليت ريدوكتاز، الإجهاد التاكسدي.