Introduction

Preeclampsia is associated with high blood pressure and proteinuria during pregnancy. Preeclampsia causes complications in 2-8% of all pregnancies, premature birth in 15% and maternal death in 9-26% of all pregnancies [1]. In preeclampsia patients, decrease in utero-placental blood flow rate as a result of vasoconstriction in the umbilical artery may lead to intrauterine growth restriction, preterm birth, maternal death. The etiology of preeclampsia is not clarified yet. However, it is thought that lipid peroxidation due to the activity of reactive oxygen species (ROS) may be the reason. When increased oxygen demand cannot be met during pregnancy, the increase of oxidative stress factors causes cell damage in fetus due to lipid peroxidation and decrease in perfusion as a result of vasoconstriction in the umbilical artery [2].

Vasoconstriction occurs by increasing the intracellular Ca$^{2+}$ concentration via the Ca$^{2+}$ channel [3]. However, there are some agents that cause vasodilatation in the vascular smooth muscle. One of these agents is leptin, which causes nitric oxide (NO) mediated vasodilatation [4]. Leptin which is an adipokine hormone and leptin receptors are more than adipose tissue in the umbilical cord.
This situation suggests that there should be an important function of leptin and leptin receptors in pregnancy. Several mutations have been identified in the leptin receptor gene. Any mutation in the leptin receptor gene may result in a defective signal transduction. Leptin in the target tissue could not fully perform its task. Most of these mutations result in early onset of obesity, hyperphagia and infertility [5].

The NOS enzyme catalyzes the NO formation reaction as a result of the interaction of L-arginine with the terminal guanidino nitrogen group and molecular oxygen. The eNOS and nNOS isoforms are active when the calcium-calmodulin complex in the cell is bound to the enzyme. Therefore, NOS activity is associated with intracellular calcium level. On the other hand, iNOS always has a firmly bound calmodulin. Therefore, the activity of the iNOS isoform is independent of the calcium level and the enzyme is always in a position to be active [6].

The R223Q polymorphism (rs1137101), one of the LEPR polymorphisms, is formed by the change of arginine to glutamine in codon 223 of exon6. There is a variety of information in the literature about the link between obesity and this polymorphism [7]. Leptin sensitivity in humans is thought to be related to variations in leptin or leptin receptor genes. Polymorphism in leptin receptor genes has not been shown to be associated with hypertension but leptin gene polymorphism has been reported to be associated with hypertension risk [8]. Another LEPR polymorphism is the R109K (rs1137100) polymorphism in codon 109 of exon 4. There are studies showing that obesity and sugar preference are associated with genetic variations. The first report of these was shown by Mizuta et al. [9]. In this study, LEPR Arg223Gln, LEPR Arg109Lys and eNOS -786T>C polymorphisms were investigated in normal and preeclamptic cord tissues.

Materials and Methods

Samples

Normal and preeclamptic pregnant women between the ages of 18-45 gave birth at in Necmettin Erbakan University, Meram Faculty of Medicine, Department of Obstetrics Gynecology (Medical Faculty Dean's Office Ethics Committee approval number is 2018/1273). After delivery, 10-15 cm sections were taken from umbilical cords and put directly into cold Krebs-Henseleit (KHS) solution (NaCl: 119mM, KCl: 4,7mM, MgSO\(_4\): 1,5mM, KH\(_2\)PO\(_4\): 1,2mM, CaCl\(_2\): 6H\(_2\)O: 2,5mM, NaHCO\(_3\): 25mM, Glucose: 11Mm). The umbilical cord samples were stored at +4 °C until use. DNA samples isolated from healthy individuals were used as positive control samples.

DNA isolation

The tissues were homogenized with liquid nitrogen. Homogenized tissues were filled with 2X lysis buffer (10X Lysis buffer: 770mM NH\(_4\)Cl, 10mM EDTA, 46mM KHC\(_6\)O\(_7\)) up to 15ml and were thoroughly mixed for 10 minutes. The tubes were kept on ice for 30 minutes and then centrifuged at 3000 rpm for 10 minutes at +4°C. The supernatant phase was discarded. 0,75ml of SALT/EDTA (75mM NaCl, 25mM EDTA) was added onto pellet and vortexed thoroughly. Then, 0,075ml 10% SDS solution, 37,5µl Proteinase K (10mg/ml) was added and the samples were incubated at 55 °C for 3 hours. 0,75ml phenol (pH 8.0) was added into the tube. The mixture was mixed for 20 seconds and then inverted for 5 minutes. The tubes were centrifuged at 3000rpm for 10min at +4 °C. After centrifugation, the supernatant phase was taken into new sterile tubes and 0,75 ml phenol chloroform: isomyl alcohol (25:24:1) was added. The mixture was mixed for 20 seconds and then inverted for 5 minutes. The tubes were centrifuged at 3000rpm for 10min at +4 °C. At the end of the centrifugation, supernatant phase in the tubes was taken into new tubes and 1/10 of NaAc were added to the supernatant. 95% ethanol was added twice as much as the current volume. The precipitated DNA was taken into the new tube with the glass rod. The pellet was dried to remove the alcohol. The pellet was dissolved with distilled water before use.

High resolution melting (HRM) curve analysis

The real-time PCR-based HRM technique was used to detect the polymorphisms of the interested genes. Briefly, 50ng DNA samples isolated from each group, 10 µl of 2X HRM mixture, 50 µl forward primer, 50 µl reverse primer and dH\(_2\)O until 20 µl were mixed. The reaction mixtures were placed on the PCR System/BioRad CFX Connect and the HRM protocol consisting of denaturation, amplification and melting steps were applied. The denaturation stage was carried out at 95 °C for 10 minutes and the temperature required for the primers to attach to the target site was chosen as 60 °C. The polymerization of the complementary DNA was performed at 72 °C. The cycle consisting of successive application of these three different temperatures was repeated for 40 times. Following PCR amplification, high-resolution melting was performed: heated to 95 °C for 30 seconds and then cooled to 60 °C for 1 minute and performed from 65 °C to 95 °C, rising at 0.2 °C/s. The graphs were analyzed in the BioRad Precision Melt Analysis program to analyze the possible polymorphisms. In this study, primers belonging to the exon6, exon4 of LEPR gene and the promoter region of eNOS gene are shown in (Tables 1 & 2).

### Table 1: Primers and PCR product lengths of the promoter region of the exon6, exon4 and eNOS gene of the OBRb gene

| Polymorphisms | Forward (5'→ 3') | Reverse (5'→ 3') | Length |
|---------------|------------------|------------------|--------|
| LEPR Gln23Arg | ACCCTTAA AGCTGGTGT TCCAAATAG | AGCTAGCAAAAT ATTTTGTAAGCAATT | 416 bp |
| LEPR Lys109Arg | AAACCTGAC TACCTCTTTTG GCC | AAACCTAGAAGT TACCTGGAA ACAATGGC | 230 bp |
| eNOS -786T>C | ATGCTGCA CCAAGGATCA | GTCCTAGATGC TGACTAGG | 236 bp |

### Table 2: Comparison of the results of HRM and PCR-RFLP analysis

| Genes | HRM Results | PCR-RFLP Results |
|-------|-------------|------------------|
| SNP (+) | SNP (-) | HM-WT | HT | HM-M |
| LEPR109 | 0 | 9 | 9 | 0 | 0 |
| LEPR223 | 1 | 8 | 4 | 3 | 1 |
| eNOS | 1 | 8 | 0 | 0 | 9 |

WT: wild type, MT: mutant type, HT: heterozygous type, HM: homozygous type.
PCR-RFLP analysis

For polymorphism analysis with PCR-RFLP, the target gene sequences were amplified by Polymerase Chain Reaction (PCR) using primers which were also used in HRM analysis to cover each target mutation. All PCR reactions were run with 50ng of DNA and 25μl of PCR mix (1X PCR buffer, 200μM dNTP, 1.5 mM MgCl₂, 0.025 U Taq DNA polymerase, 5 pmol forward and reverse primer). Samples were runned to 35 cycles of PCR protocol for 5 minutes at 95 °C for pre-denaturation, 30sec at 94 °C, 30 seconds at appropriate primer binding temperature and 30 seconds at 72 °C. Then PCR samples were run on agarose 1% gel and genotyped by RFLP technique using the MspI and HaeIII restriction enzymes for each mutation. For this purpose, 10μl PCR sample was cut at appropriate temperature with the appropriate restriction endonuclease enzyme covering each mutation site. Then, the samples were run on 2% agarose gel and genotyping was performed.

Results

HRM analysis results

HRM results showed that there was no change in A>G in exon4 of LEPR gene (Figure 1). In exon6 of LEPR gene, SNP was detected in one patient (Figure 2). The red arrow shows the SNP in that patient. The purple arrow in Figure 3 shows that there was SNP in one patient which is C>T change in the promoter region of eNOS gene.
PCR-RFLP analysis results

Figure 4: Agarose gel image of PCR products for primers used for LEPR exon6.

Figure 5: Agarose gel image of the fragments by restriction endonuclease enzyme digestion of the PCR products of LEPR exon6.

Figure 6: Agarose gel image of PCR products belonging to primers used for SNP in LEPR exon4.

Figure 7: Agarose gel image of PCR products belonging to primers used for promoter region of enos gene.

The lengths of the regions synthesized by specific primers of the polymorphism of rs1137101 found in LEPR exon6 was determined to be 416bp. All of the samples obtained from 3 positive controls, 3 healthy cord and 3 preeclamptic cord tissues successfully. Then, SNP analysis was performed by the restriction enzyme digestion which we used specifically for this region. As a result, one healthy cord tissue, one preeclamptic cord tissue and one control sample were digested by restriction enzyme. So, these samples were heterozygous for that region. One positive control sample was also digested and it was homozygous for that region as shown in Figures 4-7. The sizes of fragments as heterozygous is 416bp, 292bp and 124bp. The sizes of homozygous fragments are 295bp and 124bp.

The length of the PCR product for the primers used for SNP in exon4 of LEPR gene was found to be 159bp. Then the result of the restriction enzyme digestion, samples were uncut. The length of the PCR product for the primers used for SNP in the promoter region of eNOS gene was 237bp. As a result of the restriction enzyme digestion, no sample was cut, so we suggested that there was no SNP in this gene.

Discussion

In this study, we isolated DNA samples from 3 healthy cord tissues, 3 preeclamptic cord tissues and 3 healthy blood sample as positive control and investigated the polymorphisms in leptin receptor and eNOS genes. In order to evaluate the polymorphisms, we used HRM and PCR-RFLP techniques. HRM analysis was used because it is relatively affordable, rapid and sensitive method. In literature, there are many studies which used HRM technique for SNP analysis [10]. In this study, HRM analysis was performed in order to validate the PCR-RFLP results. In the result of HRM analysis, different genotypes give different denaturation patterns. Because of different homo- and heteroduplex forms complex curves are formed. Homozygous and heterozygous samples could be differentiated by this method [11]. The results of HRM analysis and PCR-RFLP showed different mutation patterns. This difference could be originated from sensitivity degrees of the methods. HRM analysis is more sensitive than PCR-RFLP method and PCR-RFLP method have more steps to be optimized.

HRM results suggest that SNPs in eNOS gene promoter and exon6 of LEPR could be related with preeclampsia. There was no SNP detected in exon4 of leptin receptor gene in preeclampsia patients. This result shows that this SNP is not related with preeclampsia. However, our sample number is not sufficient to suggest this relationship. The increase of the sample number will provide better understanding about the relationship between these SNPs and preeclampsia.

We detected LEPR Gln223Arg polymorphism but no LEPR Lys109Arg polymorphism in our study. Yang et al. showed that there is not a relationship between LEP G2548A and LEPR Gln223Arg polymorphisms in patients with gestational diabetes and normal glucose tolerance [12]. A meta-analysis investigated the relationship between LEPR gene polymorphisms and hypertension risk included a total of 5955 patients with hypertension and 3830 healthy controls. Accordingly, Gln223Arg polymorphism was found to be significantly higher in hypertensive patients than the control group but there was no statistically significant difference in Lys109Arg polymorphism [13]. In another study in North China, Liu et al. investigated whether the Lys109arg, Gln223Arg and Lys656Asn polymorphisms of LEPR gene have a role in the...
susceptibility to essential hypertension (EH). They found that there was no significant relationship between Lys109Arg or Lys656Asn polymorphism and EH risk. However, LEPR Gln223Arg polymorphism has an important role in susceptibility to EH [14]. Pena et al. stated that the association of Gln223 Arg polymorphism with hypertension was not determined [15].

Polymorphism was detected in the promoter of the eNOS gene in our study. In an Iranian study, there is no relationship was found between eNOS and ACE gene polymorphisms and gestational diabetes among pregnant women in the Iranian population when they were investigated whether there is a relationship between this factor [16].

Conclusion

This study is a preliminary step for our next molecular studies although few numbers of the samples were included. Increasing the number of samples would give the statistically significant results. In addition to the present polymorphism analyses, we are planning to investigate the effect of the LEPR (Leptin Receptor) gene, JAK2 and STAT3 genes expression in preeclamptic pregnant women in our next research.

Acknowledgement

None.

Conflict of Interest

No Conflict of Interest.

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