Introduction

Obesity in horses produce serious health concerns that occurs due to excessive accumulation of body fats (Huff, 2007; Henry and Clarke, 2008; Huff et al., 2010). Adipose tissue secretes adipokines and inflammatory proteins, which impact health. Leptin is a 16-kDa polypeptide hormone secreted from the white adipose tissue in response to the change in body fat or energy status (Chilliard et al., 2005). It regulates several immune functions and fertility (Akumbo and Zanwa, 2017). Leptin increased in the plasma of obese horses (Gentry et al., 2002), and was used as an important biomarker to estimate body fat mass (Van Weyenberg et al., 2008). In horses, leptin concentration was related to body condition, age, gender, and season (Fitzgerald and McManus, 2000; Gentry et al., 2002; Cartmilk, 2004). The concentration of leptin increased with the increase of body fat percentage (Chilliard et al., 2000), and body condition (Gentry et al., 2002; Cartmilk et al., 2003). High body condition score (BCS) was associated with perturbations in leptin secretion and altered both ovarian activity and the estrous cycle (Waller et al., 2006). The restriction and excess of calorie in the diet motivated leptin to repress gonadotroph signaling (Gao and Horvath, 2008). The regional adiposity, hyperinsulinemia, hypertriglyceridemia, insulin resistance, and recurrent laminitis were the main phenotypic markers of equine metabolic syndrome (Geor and Frank, 2009; Frank et al., 2010).

Leptin gene is located on chromosome 4 in the horse and consists of 3 exons and 2 introns, with only two exons are (expressed) translated into protein (Huff et al., 2009). The relation between leptin gene polymorphisms and with circulating leptin level and obesity were identified (Amiratul et al., 2018). In equine, polymorphism of the leptin gene was related to the reproductive performance (Amiratul et al., 2018), and showed no significant difference between cortisol concentration and age in stressed horses during endurance competition (Rezadet et al., 2016). The aim of this study was to investigate the relation between adiposity characterized by increased rump fat (RF) and BCS in mares with circulating leptin, insulin, ovarian hormones, and leptin gene polymorphism.

Materials and methods

Animals

Thirty six mares from different breeds (Arabian, European, etc.) were included in the study. The mares were categorized based on RF thickness estimated by the ultrasonography and body condition score (BCS) into obese (RF>5mm; BCS >5), moderate (RF>3≤5 mm; BCS >3≤5) and lean animals (RF<3mm; BCS<3). The circulating estradiol, progesterone, insulin, and leptin were analyzed. The polymorphism of the leptin gene was performed and compared for obese, moderate and lean mares. Depending on increased RF, the obesity was associated with increased BCS (P<0.0001) and leptin (P<0.0001), insulin (P<0.01), and P4 (P<0.0001). Obese mares of leptin genotype BB had the highest leptin (1146±420 pg/ml), insulin (7.42±0.89 μU/l) and progesterone (22.69±9.34 ng/ml), but minimum estradiol (22.69±9.34 ng/ml) concentrations. Mares of leptin genotype AA had the highest RF (P<0.0001). The interaction of leptin genotype and obesity had affected RF and all hormone concentrations with no significant effect on the BCS. In conclusion, obesity in cyclic mares altered ovarian hormones, insulin and leptin concentrations. The hyperleptinemia, and hyperinsulinemia were associated with the leptin genotype BB but neither to the adiposity (RF) nor BCS.

ABSTRACT

Equine metabolic syndrome is a growing concern usually associated with increased insulin and leptin concentrations. To investigate the relations of obesity determined by measuring the rump fat (RF) with ovarian hormones, leptin, and leptin gene polymorphism, cyclic mares (n=36) were categorized based on RF thickness estimated by the ultrasonography and body condition score (BCS) into obese (RF>5mm; BCS >5), moderate (RF>3≤5 mm; BCS >3≤5) and lean animals (RF<3mm; BCS<3). The circulating estradiol, progesterone, insulin, and leptin were analyzed. The polymorphism of the leptin gene was performed and compared for obese, moderate and lean mares. Depending on increased RF, the obesity was associated with increased BCS (P<0.0001) and leptin (P<0.0001), insulin (P<0.01), and P4 (P<0.0001). Obese mares of leptin genotype BB had the highest leptin (1146±420 pg/ml), insulin (7.42±0.89 μU/l) and progesterone (22.69±9.34 ng/ml), but minimum estradiol (22.69±9.34 ng/ml) concentrations. Mares of leptin genotype AA had the highest RF (P<0.0001). The interaction of leptin genotype and obesity had affected RF and all hormone concentrations with no significant effect on the BCS. In conclusion, obesity in cyclic mares altered ovarian hormones, insulin and leptin concentrations. The hyperleptinemia, and hyperinsulinemia were associated with the leptin genotype BB but neither to the adiposity (RF) nor BCS.
Animals were in good health and with clinically normal reproductive tracts and their ages ranged from 4 to 18 years old. Mares were raised on the Equestrian Club (El Basateen), belongs to the Police Department, Ministry of Interior, Egypt. Animals were fed their requirements of pelleted ration with free access to mineral salt blocks, water, regularly exercised and dewormed. Animals were examined and sampled in accordance to the Institutional Animal Care and Use Committee of the National Research Centre, Egypt.

Experimental design

Mares were divided into three groups according to the rump fat measurements (Abo El-Maaty et al., 2017) and body condition score (Henneke et al., 1983); obese (rump fat > 5 mm and BCS > 5), moderate animals (rump fat > 3 ≤ 5 mm and BCS > 3 ≤ 5) and lean animals (rump fat < 3 mm and BCS < 3).

Ultrasound examination

The ultrasound scanner (SonoAce R3, Medison, Samsung, South Korea) was used to measure rump fat depth, count the number and diameter of ovarian follicles, determine the diameter of the corpus luteum and also to exclude pregnancy.

Blood sampling

Blood samples were collected in plain vacuum tubes, then sera were harvested. Blood for leptin Gene polymorphism was collected in vacuum tubes containing EDTA and all were stored at -20°C until molecular genetic analysis and hormone assaying were performed.

Hormone assaying

Horse ELISA leptin and insulin sandwich kits (Bioneovan Co., China), progesterone (P4) and estradiol (E2 DRG, Instruments GmbH, Germany) were assayed with minimum detectable concentrations, which were 35 pg/ml for leptin, 0.3 μU/L for insulin 0.045 ng/ml for P4, and 9.714 pg/ml for E2. The intra-and inter-assay precisions were < 10% and < 12%, for both leptin and insulin; 5.4% and 9.96% for P4, and 6.81 and 7.25% for E2.

Genomic DNA extraction and PCR amplification

Genomic DNA was isolated by the salting out method (Paneto et al., 2008). DNA concentration and purity were evaluated in a spectrophotometer (Nanodrop 2000c, Thermo Scientific, USA) and DNA quality was checked by 1.0% agarose gel electrophoresis before PCR amplification. According to the sequence of the horse leptin gene (GenBank Accession NO NC_009147.2 (Caetano et al., 1999) one pair of polymerase chain reaction (PCR) primers (forward: 5’-GTACACGGATCAATGACAT-3’ and reverse: 5’-AGCCCGAAGTGAAGTGCCA-3’) was designed to amplify a 1900 bp fragment of leptin where exon 2 and 3 separated with intron 2. A PCR cocktail consisted of 1.0 M upper and lower primers, 0.2 mM dNTPs and 1.25U of Taq polymerase. The cocktail was aliquoted into PCR tubes with 50 ng of horse DNA. The PCR thermal profile was as follows: initial denaturation of DNA matrix for 5 min at 94°C, followed by 35 cycles: proper denaturation at 94°C (1 min), primer annealing at 58°C (1 min), PCR product synthesis at 72°C (2 min) and final synthesis at 72°C for 10 min. The PCR products were loaded onto 1.5% agarose to confirm the amplification of the target region using a 100bp marker.

Genotyping

Restriction fragment length polymorphism (RFLP) analysis of the PCR product (Mahrous et al., 2020) was carried out using TaqI enzyme (Fermentas). Ten ul of PCR products were digested with 1ul of FastDigest restriction enzymes for 5 min at 65°C. The digested PCR fragments were subjected to electrophoresis in 2% agarose/ethidium bromide gel (GIBCO, BRL, England) in 1x TBE buffer (0.09 M Tris-boric acid and 0.002 M EDTA). Gels were visualized under UV light and documented in FX Molecular Imager apparatus (BIO-RAD). LEP gene analysis results were elaborated statistically using GENEPOP software package (Raymond and Rousset, 1995). The genetic structure, utility types, the frequency of leptin genotypes (LEP/TaqI) and the frequency of respective alleles were calculated.

Statistical analysis

For comparing allelic and genotype frequencies between the horse breeds and utility types examined, χ2 test was applied to evaluate the significant of differences between the values obtained for particular genotypes and alleles. Simple One Way ANOVA was used and the Duncan’s multiple range tests were performed using SPSS (2016).

Results

The PCR amplification of leptin gene yielded about 1900 bp long fragment (Fig.1). This amplified fragment was digested with the TaqI restriction enzyme and results in homozygous AA genotype with two restriction fragments of ~1040 and ~820 bp, heterozygous AB genotype with four fragments at ~1040, ~820, ~740 and ~300 bp, and homozygous BB genotype with three fragments at ~820,~740 and ~300 bp (Fig. 2). The frequency of allele B (0.734) was higher (P=0.0001)
in the whole population as compared with allele A (0.266). The values of allele A frequency were increasing with the increase of RF and BCS 0.133, 0.294 and 0.357 but those of allele B were decreasing with the increase of RF and BCS (0.869, 0.708, and 0.643) from lean to moderate and obese horses, respectively. The genotypic frequency of the homozygous genotype BB is high (P=0.001) in lean mares (0.82) as compared with moderate (0.576) and obese (0.507) mares. Contrary, the genotypic frequency of the homozygous genotype AA was low (P=0.002) in lean (0.083) in comparison to moderate (0.162) and obese mare (0.221; Table1).

In the obese mares (Table 2), though mares with genotype BB had the lowest (P=0.0001) RF (5.08±0.58), E2 (101±54), but they had the highest P4 (P=0.012; 22.69±9.34), insulin (P=0.0001; 7.42±0.89), and leptin (P=0.0001; 1146±420). The obese mares with genotype AA had the highest RF (P=0.0001; 8.10±1.39), E2 (P=0.0001; 257±22), and BCS (6.65±0.96), the lowest leptin (P=0.0001; 101±55), and nearly similar insulin and P4 to the heterogeneous obese mares of leptin genotype AB (Table 2). Within lean mares, those with leptin genotype AA had the highest (P=0.0001) RF (2.99±0.09), P4 (P=0.027; 17.24±8.18), and leptin (P=0.0001; 234±15), but the lowest E2 (P=0.018; 102±6). When all mares were classified according to their leptin genotype (Table 3), mares with the genotype AA had the highest (P=0.0001) RF (6.06±2.62), BCS (5.92±1.49), and P4 (P=0.048; 19.89±9.92), but low leptin (121±66). Although mares with leptin genotype BB had the minimum RF (P=0.0001; 3.82±1.43), BCS (P=0.0001; 4.97±1.73), P4 (P=0.0458 17.11±8.91), but they had the highest leptin (P=0.0001; 635±593) and insulin (P=0.0001; 5.17±5.72) concentrations.

Regardless leptin genotyping (Table 4), RF thickness descended from obese mares (7.24±8.19) (P=0.0001), to moderate (4.48±1.03), and then to lean mares (2.12±0.53). BCS of obese mares is high (P=0.0001; 6.92±0.83) in contrast to moderate (5.41±1.07), and lean mares (3.43±1.18). Leptin concentrations of obese mares is the highest (P=0.0001; 356±486) when compared to moderate (201±236), and lean mares (196±154). P4 concentrations descended linearly (P=0.0001) from obese (20.97±9.53), to moderate (18.07±9.41), and lean ones (14.28±8.18). Obese mares obtained the lowest E2 concentrations (P=0.0001; 159±143) and the highest insulin concentrations (P=0.01; 4.08±4.28).

Leptin genotype (Table 5) showed significant effects (P<0.0001) on RF (P<0.0001), BCS (P<0.0001), insulin (P=0.0005), P4 (P<0.0001), and E2 (P=0.048) in mares. The obesity depending on increased rump fat showed significant effects (P=0.0001) on RF, BCS, leptin, insulin, and E2. The interactions of leptin genotype and obesity affected RF (P<0.008), leptin (P<0.0001), insulin (P<0.0001), P4 (P<0.0001), and E2 (P<0.0001).

Leptin (r=0.17; P=0.023), correlated with BCS of obese mares and moderate mares (r=0.32; P=0.0001), but has a negative one with RF of obese mares only. (r=0.54; P=0.0001). Insulin correlated negatively with BCS (r=0.33; P=0.0001), RF (r=0.21; P=0.003) of obese mares, and RF (r=0.36; P=0.031) of lean mares but positively with BCS of lean mares (r=0.46; P=0.0001).

**Discussion**

Leptin is one of the mediators that influence the effects of the body weight on fertility. Furthermore, the decreased body fat was associated with decreased leptin levels in horses showing impaired reproductive efficiency (Gentry et al., 2002). Allele frequency in the present study revealed that allele B was higher than allele A in all studied groups. In consistent with our study, the greatest frequency in the BB LEP/TaqI homozygous was also observed with a higher percentage of participation of leptin AB heterozygous animals in comparing with the AA homozygous ones in mares (Kęszka, 2006) and Polish cold-blooded horses (Bohaczyk et al., 2011). Similar to cold blooded Polish horses, the LEP/TaqI frequencies of the gene AA and allele A were the lowest, and the allele B had the highest frequency (Bohaczyk et al., 2011). In the study of Bohaczyk et al. (2011), the frequency of the allele B (0.6108) was higher than the allele A (0.3892) for the whole horse population. In contrast to our results, the frequency of the heterozygous leptin gene AB (0.4897) was the highest, followed by BB (0.3660), whereas the identical allele AA was the lowest (0.1443, Kęszka, 2006).

**Table 1. Leptin genotype and allele frequencies (%) in mare’s populations with different body conditions**

<table>
<thead>
<tr>
<th>Body condition/Rump Fat</th>
<th>Genotypic frequencies</th>
<th>Allelic frequencies</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>AA</td>
<td>AB</td>
</tr>
<tr>
<td>Lean*</td>
<td>0.156</td>
<td>0.221</td>
</tr>
<tr>
<td>Moderate *</td>
<td>0.162</td>
<td>0.263</td>
</tr>
<tr>
<td>Obese*</td>
<td>0.221</td>
<td>0.272</td>
</tr>
</tbody>
</table>

*: P< 0.01; **: P< 0.001
Table 2. Values of rump fat (RF), BCS, leptin and ovarian hormones in obese, moderate and lean mares with different genotypes

<table>
<thead>
<tr>
<th>Genotype</th>
<th>RF (mm)</th>
<th>BCS</th>
<th>Leptin (ng/ml)</th>
<th>Progesterone (ng/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aa</td>
<td>10.1±1.9</td>
<td>6.8±0.9</td>
<td>6.0±0.9</td>
<td>3.7±0.9</td>
</tr>
<tr>
<td>Ab</td>
<td>7.0±0.9</td>
<td>7.5±0.9</td>
<td>5.0±0.9</td>
<td>2.5±0.9</td>
</tr>
<tr>
<td>Bb</td>
<td>5.0±0.9</td>
<td>6.0±0.9</td>
<td>3.0±0.9</td>
<td>1.5±0.9</td>
</tr>
</tbody>
</table>

Data were expressed as Mean±SD
Means with different superscripts (a,b,c) are significantly different at P<0.05; non-significant (NS); rump fat (RF), body condition score (BCS), progesterone.
Table 3. Values of BCS, Rump fat thickness and hormone concentrations in mares with different leptin genotypes

<table>
<thead>
<tr>
<th>Parameters</th>
<th>AA</th>
<th>BB</th>
<th>AB</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rump Fat</td>
<td>6.06±2.62c</td>
<td>3.82±1.43a</td>
<td>4.63±1.21b</td>
<td>0.0001</td>
</tr>
<tr>
<td>BCS</td>
<td>5.92±1.49b</td>
<td>4.97±1.73c</td>
<td>5.39±1.53a</td>
<td>0.0001</td>
</tr>
<tr>
<td>Leptin (pg/ml)</td>
<td>121.0±66.0a</td>
<td>635.0±593.0b</td>
<td>234.0±178.0c</td>
<td>0.0001</td>
</tr>
<tr>
<td>Insulin (µU/L)</td>
<td>3.14±2.97b</td>
<td>5.17±5.72b</td>
<td>2.45±2.12a</td>
<td>0.0001</td>
</tr>
<tr>
<td>P4 (ng/ml)</td>
<td>19.89±9.92b</td>
<td>17.11±8.91c</td>
<td>18.68±8.98ab</td>
<td>0.048</td>
</tr>
<tr>
<td>E2 (pg/ml)</td>
<td>242.0±205.0</td>
<td>191.0±127.0</td>
<td>191.0±170.0</td>
<td>NS</td>
</tr>
</tbody>
</table>

Data were expressed as Mean±SD
Means with different superscripts (a,b,c) are significantly different at P<0.05; NS: Non significant

Table 4. Values of BCS, rump fat thickness and hormone concentrations in mares with different rump fat thickness

<table>
<thead>
<tr>
<th>Parameters</th>
<th>lean</th>
<th>Moderate</th>
<th>obese&lt;5mm BF</th>
<th>Total</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Back Fat</td>
<td>2.12±0.33a</td>
<td>4.48±1.03b</td>
<td>7.24±2.39c</td>
<td>4.98±2.47</td>
<td>0.0001</td>
</tr>
<tr>
<td>BCS</td>
<td>3.43±1.18a</td>
<td>5.41±1.07b</td>
<td>6.92±0.83c</td>
<td>5.50±1.63</td>
<td>0.0001</td>
</tr>
<tr>
<td>Leptin (pg/ml)</td>
<td>196.±154a</td>
<td>201±236b</td>
<td>356±486b</td>
<td>264±361</td>
<td>0.0001</td>
</tr>
<tr>
<td>Insulin (µU/L)</td>
<td>2.61±1.91a</td>
<td>3.27±4.52b</td>
<td>4.08±4.28b</td>
<td>3.50±4.16</td>
<td>0.01</td>
</tr>
<tr>
<td>P4 (ng/ml)</td>
<td>14.28±8.18a</td>
<td>18.07±9.41b</td>
<td>20.97±9.53c</td>
<td>18.06±9.48</td>
<td>0.0001</td>
</tr>
<tr>
<td>E2 (pg/ml)</td>
<td>229±166a</td>
<td>269±184b</td>
<td>159±143c</td>
<td>219±173</td>
<td>0.0001</td>
</tr>
</tbody>
</table>

Data were expressed as Mean±SD
Body condition score (BCS), rump fat (BF). Means with different superscripts (a,b,c) are significantly different at P<0.05.

Table 5. The effect of obesity, leptin genotype and their interactions on BCS, Rump Fat and circulating hormones in mares

<table>
<thead>
<tr>
<th>Dependent Variables</th>
<th>Leptin Genotype</th>
<th>Obesity</th>
<th>Leptin Genotype * obesity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rump Fat</td>
<td>0.0001</td>
<td>0.0001</td>
<td>0.008</td>
</tr>
<tr>
<td>BCS</td>
<td>0.0001</td>
<td>0.0001</td>
<td>NS</td>
</tr>
<tr>
<td>Leptin pg/ml</td>
<td>0.0001</td>
<td>0.0001</td>
<td>0.0001</td>
</tr>
<tr>
<td>Insulin µU/L</td>
<td>0.005</td>
<td>0.0001</td>
<td>0.0001</td>
</tr>
<tr>
<td>P4 ng/ml</td>
<td>0.0001</td>
<td>NS</td>
<td>0.0001</td>
</tr>
<tr>
<td>E2 pg/ml</td>
<td>0.048</td>
<td>0.0001</td>
<td>0.0001</td>
</tr>
</tbody>
</table>

NS: Non significant

negative effect on gonadal steroidogenesis (Orr, 2016). The hyperleptinemia of obese and moderate mares with leptin genotype BB was associated with minimum E2 level and hypoletinemia in obese and moderate mares with leptin genotype AA associated with high E2 level, which may indicate the presence of a specific and narrow range of leptin concentrations to maintain normal reproductive functions and that concentrations below or above these thresholds interfered in opposing ways with the function of the hypothalamic-pituitary-gonadal axis (Orr, 2016). The differences in ovarian hormones between mares of different rump fat, leptin genotype and concentrations of this study conform with its role in regulating body weight and acting as mediators of many physiological functions (Henson and Castracane, 2003), and agreed with the rise of plasma leptin concentration in horses with high body fat mass (Cartmill et al., 2003).

Conclusion

This study concluded that obese mares have high BCS, back fat, P4 level, and leptin level but low E2 level. The leptin genotype BB was more frequent but not in obese mares with high leptin level, whereas the leptin genotype AA was less frequent but characterized by having high BCS, back fat with low leptin level, suggesting that these polymorphisms has no effect on mares’ body weight.

Conflict of interest

The authors declare that they don’t have any conflict of interest.

References


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