iTRAQ-based proteomic analysis of the reproductive cycle characteristics of Hainan Black Goat, the unique species in China, Hainan Province

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ABSTRACT

Hainan black goats are unique species in China, Hainan province, they are specific mutton goat breed and the tasty meat make the black goat to be one of the four famous dishes in Hainan. However, little knowledge was known about the reproductive cycle of this breed. So, in the study, serum samples of the reproductive period were collected, and high-throughput sequencing were conducted by iTRAQ technology. The results showed the amounts of significantly different proteins between continuous period and the corresponding Go and COG annotation, KEGG enrichment of differential expressed proteins between each two periods were also analyzed. Further, continuously differential expression proteins of the whole cycle and differential expression proteins directly related to reproduction between two adjacent periods were also collected and annotated. This is the first study systematically uncovered reproductive cycle characteristics of Hainan black goats, the study preliminarily uncovered the basical protein information between each reproductive period, and the data could contribute to the built of highly efficient reproduction system of Hainan black goats.

Keywords: Black goat; Reproduction; Proteomics; iTRAQ; Protein

INTRODUCTION

Hainan black goats are unique species in China, Hainan province. With the climatic feature of hot and humid, the distinct dry rainy season, the black goats have been cultivated as resistance of crude feed, strong disease resistance, early sexual maturity and tender meat quality through natural selection (Wang et al., 2017). For the numerous excellent trait and considerable economic value, nowadays, industrialized farming of east goats has become a trend, and the chief problem was the relatively low reproduction rate and low milk production efficiency (Adamala, 2018; Al-zaqri et al., 2017; Kannan et al., 2018; Maddi et al., 2018; Troujeni et al., 2018).

iTRAQ (isobaric tags for relative and absolute quantitation) is the quantitative proteomic technology based on isotope labelling in vitro, which could analyses the differentially expressed protein and the related signal transduction pathway between different period (Howlader et al., 2018; Hasan et al., 2018; Khan, 2018), it is one of the most popular proteomic technology nowadays (Moulder et al., 2017). iTRAQ-based proteomic analysis have been widely applied in the process of livestock production, comparative proteomic analyses of fiber diversity between sheep and goats provided a list of candidate proteins that involved in fiber formation (Li et al., 2018), iTRAQ based technology was also applied to analysis the influence of overgrazing induced alterations in hepatic proteins of sheep (Ren et al., 2016). As in porcine production, iTRAQ-based technology has been used to analysis the in vivo muscle development distinctions during early embryonic development of different pig breeds (Zhang et al., 2016). To investigate the effect of transportation under high temperature conditions, iTRAQ technology was also applied to determine the changes in protein profile of pale, soft and exudative (PSE) – like muscles of broilers (Xing et al., 2017), which provided a further understanding of the mechanism of meat quality changes in response to stress, the technology also surveyed as a feasibility and sensitivity tool to reveal...
the molecular mechanisms underlying brown eggshell pigmentation (Li et al., 2016).

Serum proteomic analysis based on iTRAQ technique has been widely applied, for serum samples are easy to be acquired and biomarker candidates can be screened out more efficiently (Koutroukides et al., 2013, Tonack et al., 2009), i.e. iTRAQ based screen of serum of essential hypertension patients found certain special proteins and the interaction between them, which could better predict the occurrence and illustrate the pathophysiological mechanism of essential hypertension disease (Xu et al., 2017), potential serum biomarkers in locoweed-poisoned rabbits were also efficiently found based on iTRAQ test, which in turn could efficiently diagnosis the disease of locoism (Jiao et al., 2017). The two-dimensional method was also used to evaluate specific characteristics in serum proteins of sheep with mild broncho-pulmonary disease, but compared with the iTRAQ technology (Yun et al., 2017; Chen et al., 2017), the traditional two-dimensional method was obviously less high throughput and with lower accuracy (Chiaradia et al., 2012).

As far as we know, there are no system-wide report about the whole reproductive cycle of goats so far (Basak and Gajbhiye, 2018; Basu et al., 2018; Garcia Alegria et al., 2017; Gonzalez-Trinidad et al., 2017; Kylili et al., 2018; Tu et al., 2018). To reveal the characteristics of the changing rule during the reproductive cycle of Hainan black goats, in this study, we used iTRAQ-based technology to reveal the potential serum proteomic information of black goats during continuously reproductive cycle (empty pregnant period, oestrus period, gestation period, lactation period), and certain functional proteins will be described, this is the first report on the changing rule of serum proteins of black goats during the reproductive cycle (Camara et al., 2018; Duan et al., 2018; Fu and Liu, 2017; Zong et al., 2018).

**MATERIALS AND METHODS**

**Serum collection of different reproduction period**

The blood samples (6 mL) were collected from 12 east goats (empty pregnant period, oestrus period, gestation period, lactation period, each period collected 3 individual blood samples as biological repeat), serum was collected from blood samples following the manufacture’s protocol. Briefly, blood sample was incubated at room temperature for 2 h and centrifuged for 10 min at 3000 rpm at 4 °C, and the supernatant was transferred into a clean polypropylene tube and stored at -80 °C for further use.

**Wipe off high-abundant protein**

Add 610 µl rehydration reagent to the bottle of elution reagent to prepare rehydrated elution reagent. Get the serum protein (≥10 mg), centrifuged at 10000 g for 10 min, put off the bottom cap of spin column, transfer spin column to the centrifuge tube without cover, centrifuge 30-60 s at 1000 g, get rid of stock solution and mount the bottom cap of spin column, open the top cap of spin column, add 200 µl wash buffer, incubated at 26 °C for 5 min, then open then bottom cap and transfer spin column to the centrifuge tube without cap, centrifuge 30-60 s at 1000 g, drop the wash buffer and mount the bottom cap. Open the top cap of the spin column and added 100 µl rehydrated elution reagent, mount the top cap, shock slightly for 5 s and incubated at 26 °C for 15 min, open the bottom cap and transfer spin column to the clean centrifuge tube, centrifuge 60 s at 1000 g, label the collect fluid respectively and mount the bottom cap, add DTT to the final concentration of 10 mM and incubated at 56 °C for 1 h, as the temperature returned to RT add the IAM to the final concentration of 55 mM, then added 5 times volume cold acetone, precipitate at -20 °C for 2 h, then centrifuged at 30000 g, 4 °C for 15 min, drop the supernatant and dry the precipitate, added 200 µl lysis buffer 3, ice-bath for 5 min and centrifuged for 15 min at 25000 g, 4 °C, to obtain the protein supernatant.

**Protein extraction and quality control**

The protein concentration was quantified using BCA kit, and the integrity of the samples were evaluated using SDS-PAGE method.

**Protein enzymolysis and peptide fragment labeling**

100 µg protein of each sample were taken out and hydrolyzed at 37 °C for 4 h, with the ratio to enzyme of trypsin as 40:1, then Strata X column was used to desalt the hydrolyzed peptide fragment. Certain iTRAQ labeling reagents were taken out and added 50 µl isopropanol to each of the tube, then 0.5 M TEAB was used to dissolve the peptide fragment, and added into the corresponding labeling reagent, each different peptide fragments were labeled with specific iTRAQ tag, incubated at RT for 2 h.

**Peptide fragment separation and determined by liquid chromatography-mass spectrometry**

Using liquid chromatography (LC-20AB Daqin Japan) and C18 column (5 µm 4.6×250 mm Gemini) to separate the samples, 2 ml mobile phase A (5% ACN pH 9.8) was used to redissolve the dried samples and injected, and the column was gradiently eluted with the speed of 1 ml/min, the fraction was collected every minute under the monitoring wavelength of 214 nm, 20 fractions were obtained in total and freeze dried, then the dried samples were redissolved with mobile phase A (2% ACN 0.1% FA), centrifuged
at 20000 g for 10 min and collected the supernate to inject into the liquid chromatography. The samples first enriched and desalted in the trap column, then connected with self-assembly C18 column (75 μm inner diameter, 3.6 μm column stuff particle size, 15 cm column length), separated with the flow rate of 300 nl/min, and the tail end was directly connected with mass spectrometer triple TOF 5600 (SCIEX, Framingham, MA, USA), the ion source was Nanospray III source and the radiator was the needle made from quartz material. The parameter of mass spectrometer was set as: 2300V Ion source spray voltage, the nitrogen pressure was 30 psi, the atomizer was 15, the temperature of the spray joint was 150 °C, scan as the high sensitive mode, the cumulative time of first level mass spectrum was 250 ms, the mass range of scan was 350-1500 Da, based on the first level scanning information, ranking based on the ion strength of the first level spectrogram from high to low, selecting the ion strength which exceeded 150 cp to be fragmentized and scan the secondary data, the cumulative time of second level mass spectrum was 100 ms. Data collection was based on the type of ITRAQ, and the fracture energy was adjusted based on the ITRAQ reagents, the ion transfer efficiency of second quadrupole Q2 at 100 Da was 100%.

The process of proteomic information analysis
As the original data was collected, the proteomic information analysis was conducted as described in Fig. 1, based on the analysis, the total amount of differential proteins was counted and the annotation of differential proteins between each period were conducted.

RESULTS

Overview of the identified protein amounts
In the study, we have total groups of four, be named as empty pregnant, oestrum, gestation and lactation respectively, to divide the whole reproductive cycle of black goats into these four periods. Each group had three repetitions. Finally, we generated 1011548 secondary spectrum diagram, and under the filter criteria of 1% FDR, we have identified 5745 items of peptide fragments, 5465 items of unique peptide fragments and 1213 items of proteins (Table 1). Fig. 2 showed identified protein results between replicates, it showed that 442 common protein items between each replication. It could be concluded from Fig. 3 that the most abundantly distributed protein mass was above 100 kDa, and the centrally distributed protein mass scope were 10-60 kDa. From Fig. 4, we could find that the most abundantly distribute peptide length scope identified (calculated as the amount of amino acids) were 7-17 amino acids.

Protein quantification based on iTRAQ technology
In this study, we set up six comparison groups: lactation/ gestation, lactation/oestrum, gestation/oestrum, empty pregnant/oestrum, empty pregnant/lactation, empty pregnant/gestation, the significant difference was screened by the criterion of Fold change >1.2 or <0.83 and Q value <0.05, for the data of multiple tests, the final significantly different protein should be defined as significant protein at least one time (>one time). From table 2, the most differential protein amounts in total was the comparison group of empty pregnant/gestation (include both up and down regulated), 201 proteins were significantly regulated between these two periods. The amounts of significantly different proteins of continuous period comparison between empty pregnant/oestrum, gestation/oestrum and lactation/gestation were 141, 170 and 159 respectively, which showed the intrinsic difference between continuous period, corresponding to Fig. 5, from the distribution diagram of CV between repetitions (Fig 6), we could conclude that the comparison values between two periods were credible.

GO and COG enrichment of differential expressed proteins during the reproduction cycle
From Fig 7 it could be concluded that between each two periods, in the category of biological process, the most amounts of proteins enriched were in catalytic activity and binding items, in the category of cellular component, the most amounts of proteins enriched were in cell part and cell items, and in the category of molecular function, the most amounts of proteins enriched were in single-organism process and cellular process items. From the enrichment in COG categories (Fig 8), it could be concluded that the items of translational modification, protein turnover, chaperones and general function prediction were the most concentrated area.

KEGG enrichment of differential expressed proteins between each two periods
Two key information should be focus on in Fig 9, protein numbers and rich factor. From Fig. 9A, it could be

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Total spectra</th>
<th>Spectra</th>
<th>Unique spectra</th>
<th>Peptide</th>
<th>Unique peptide</th>
<th>Protein</th>
</tr>
</thead>
<tbody>
<tr>
<td>Black goats (repetition1)</td>
<td>338816</td>
<td>22654</td>
<td>20776</td>
<td>3831</td>
<td>3563</td>
<td>722</td>
</tr>
<tr>
<td>Black goats (repetition2)</td>
<td>335614</td>
<td>23580</td>
<td>21762</td>
<td>3984</td>
<td>3760</td>
<td>746</td>
</tr>
<tr>
<td>Black goats (repetition3)</td>
<td>337118</td>
<td>27958</td>
<td>24944</td>
<td>4316</td>
<td>3983</td>
<td>825</td>
</tr>
<tr>
<td>Total</td>
<td>1011548</td>
<td>74192</td>
<td>67482</td>
<td>5745</td>
<td>5465</td>
<td>1213</td>
</tr>
</tbody>
</table>
Fig 1. Bioinformatics analysis pipeline. The mass spectrometric data was converted into the formatted file type of mgf, and the protein identification software Mascot was used to search and identify the protein species based on the corresponding database. Quality analysis was also conducted to judge whether the data was qualified. As the data was credible, then certain screening threshold was set to obtain the final receivable protein data. Next quantitative analysis based on iTRAQ was conducted and concerned proteins which showed significant difference were screened out. Finally function annotation of GO, Pathway, COG of proteins and enrichment analysis of GO and Pathway of differential proteins were conducted.
concluded that the dot of the highest protein numbers and the most significant rich factor were inconsistent, the pathway related to complement and coagulation cascades, and p53 signaling pathway respectively, as to Fig. 9B, the highest protein numbers and the most significant rich factor were complement and coagulation cascades, prolactin signaling pathway respectively, as to Fig. 9C, the rule was the same with Fig. 9A, Fig. 9D showed the same rule with Fig. 9A in the area of highest protein numbers, but the most significant rich factor located on Fanconi anemia pathway, as to Fig. 9E, the highest protein numbers and the most significant rich factor were metabolic pathways and arachidonic acid metabolism, Fig. 9F showed the complement and coagulation cascades and adipokine signaling pathway as the highest protein numbers and the most significant rich factor respectively.

**Continuously differential expression proteins of the whole cycle**

As seen in Table 3, there are total five proteins which showed significantly differential expression between the adjacent stages: C4b-binding protein alpha chain isoform X4 (C4BP α X4), coagulation factor IX (FIX), leukocyte cell-derived chemotaxin-2 (LECT2), hyaluronan-binding protein 2 isoform X2 (HABP 2 X2) and Apolipoprotein C-II isoform X2 (ApoC-II X2) respectively. The level of C4BP α X4 showed down-regulated rule from empty pregnant to gestation, and then increased in the period of lactation, the level of

<table>
<thead>
<tr>
<th>Compare group</th>
<th>Up regulated</th>
<th>Down regulated</th>
<th>All regulated</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lactation/gestation</td>
<td>58</td>
<td>101</td>
<td>159</td>
</tr>
<tr>
<td>Lactation/oestrus</td>
<td>93</td>
<td>79</td>
<td>172</td>
</tr>
<tr>
<td>Gestation/oestrus</td>
<td>101</td>
<td>69</td>
<td>170</td>
</tr>
<tr>
<td>Empty pregnant/oestrus</td>
<td>66</td>
<td>75</td>
<td>141</td>
</tr>
<tr>
<td>Empty pregnant/lactation</td>
<td>46</td>
<td>60</td>
<td>106</td>
</tr>
<tr>
<td>Empty pregnant/gestation</td>
<td>63</td>
<td>138</td>
<td>201</td>
</tr>
</tbody>
</table>

**Table 2: Protein amounts list with significant difference between each period**

![Fig 2. Venn diagram of protein identification. Venn diagram describing identified protein results between replicates. The blue, red and green color represented replication 1, 2, 3 respectively, and the common part between each replication represented intersection protein items.](image1)

![Fig 3. Distribution map of protein molecular weight. The map showed the distribution status of protein molecular weight of the whole identified items. (A) The total protein molecular weight distribution level in replication 1, (B) The total protein molecular weight distribution level in replication 2, (C) The total protein molecular weight distribution level in replication 3. X-axis: The scope of protein molecular weight (kDa), Y-axis: The ratio of corresponding proteins.](image2)
FIX showed down regulated from empty pregnant to oestrus, and then increased in the period of gestation, finally reduced in the lactation period, the level of LECT2, HABP 2 X2 and apoC-II X2 showed the same changing tendency with FIX.

**Differential expression proteins directly related to reproduction between two adjacent period**

As listed in table 4, There are total eight proteins showed differential expression between two adjacent periods which directly related to reproduction, that are serum amyloid A21 precursor (SAA21), anthrax toxin receptor 2 (ANTXR2), von Willebrand factor (VWF), secreted frizzled-related protein 3 (Sfrp 3), Beta-1,4-galactosyltransferase 1 (Beta-1,4-GalT 1), insulin-like growth factor-binding protein 2 precursor (IGFBP-rP2), GTP-binding nuclear protein Ran (GTP-binding protein), retinol-binding protein 4 precursor (RBP4). The ratio of oestrus/empty pregnant showed down-regulated in ANTXR2, Sfrp 3, beta-1,4-GalT 1 and RBP4, and the ratio of gestation/oestrus showed up-regulated in SAA21 and IGFBP-rP2 and showed down-regulated in ANTXR2, VWF, Sfrp 3 and GTP-binding protein, the ratio of lactation/gestation showed up-regulated in VWF and down-regulated in beta-1,4-GalT 1 and IGFBP-rP2, the ratio of empty pregnant/lactation showed up-regulated in GTP-binding protein and down-regulated in IGFBP-rP2.

*Table 3: Continuously differential expression proteins of the whole cycle*

<table>
<thead>
<tr>
<th>Protein items</th>
<th>Oestrum/empty pregnant</th>
<th>Gestation/oestrus</th>
<th>Lactation/gestation</th>
</tr>
</thead>
<tbody>
<tr>
<td>C4BPαX4</td>
<td>↓ 0.74</td>
<td>↓ 0.77</td>
<td>↑ 1.41</td>
</tr>
<tr>
<td>FIX</td>
<td>↓ 0.71</td>
<td>↑ 1.88</td>
<td>↓ 0.87</td>
</tr>
<tr>
<td>LECT2</td>
<td>↓ 0.74</td>
<td>↑ 1.59</td>
<td>↓ 0.74</td>
</tr>
<tr>
<td>HABP2X2</td>
<td>↓ 0.8</td>
<td>↑ 2.75</td>
<td>↓ 0.36</td>
</tr>
<tr>
<td>ApoC-II X2</td>
<td>↓ 0.78</td>
<td>↑ 1.29</td>
<td>↓ 0.6</td>
</tr>
</tbody>
</table>

*Table 4: Differential expression proteins directly related to reproduction between two adjacent period*

<table>
<thead>
<tr>
<th>Protein items</th>
<th>Oestrum/empty pregnant</th>
<th>Gestation/oestrus</th>
<th>Lactation/gestation</th>
</tr>
</thead>
<tbody>
<tr>
<td>SAA21</td>
<td>-</td>
<td>↑ 1.76</td>
<td>-</td>
</tr>
<tr>
<td>ANTXR2</td>
<td>↓ 0.72</td>
<td>↓ 0.65</td>
<td>-</td>
</tr>
<tr>
<td>VWF</td>
<td>-</td>
<td>↓ 0.74</td>
<td>↑ 1.28</td>
</tr>
<tr>
<td>Sfrp 3</td>
<td>↓ 0.72</td>
<td>↓ 0.61</td>
<td>↓ 0.36</td>
</tr>
<tr>
<td>Beta-1,4-GalT 1</td>
<td>↓ 0.56</td>
<td>-</td>
<td>↓ 0.52</td>
</tr>
<tr>
<td>IGFBP-rP2</td>
<td>-</td>
<td>↑ 1.99</td>
<td>↑ 0.65</td>
</tr>
<tr>
<td>GTP-binding protein</td>
<td>-</td>
<td>↑ 0.48</td>
<td>-</td>
</tr>
<tr>
<td>RBP4</td>
<td>↓ 0.70</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

*"-" means no significant difference between two adjacent period.*
DISCUSSION

The goat populations in China was the most abundant around the world (Wang et al., 2015a; Wu et al., 2009), Hainan black goat was the specific species in China, Hainan province. The meat-type goat breed has the excellent characteristics of heat resistance (Wang et al., 2014a), in recent years, with the increase in farm size and the number of goats bred, the low farrowing rate and milkability have exerted increasing influence on the scale of Hainan black goat. There were a few reports on the genetic diversity (Zhou et al., 2012), inflammasome components (Zhang et al., 2017) and nutrition regulation methods (Wang et al., 2014b; Wang et al., 2017; Wang et al., 2015b) of Hainan black goat, however, so far, there still no report on the reproductive law of this specific goat species, the study was the first research on the reproduction characteristics of Hainan black goat, we aimed to improve the fertility of this species through uncovering the internal rule of the changing of serum proteins during the whole reproductive cycle.

There were five proteins showed continuously differential expression between two adjacent stages during the whole reproductive cycle, that were C4b-binding protein alpha chain isoform X4, coagulation factor IX, leukocyte cell-derived chemotaxin-2, hyaluronan-binding protein 2 isoform X2 and apolipoprotein C-II isoform X2 respectively. And eight proteins in total directly corrected with reproduction showed significantly differential expression between two adjacent stages, that were serum amyloid A21 precursor, anthrax toxin receptor 2, von Willebrand factor, secreted frizzled-related protein 3, beta-1,4-galactosyltransferase 1, insulin-like growth factor-binding protein 2 precursor, GTP-binding nuclear protein Ran and retinol-binding protein 4 precursor.

We first focused on the five continuously differential expression proteins, based on the go analysis, the role of C4b-binding protein were Innate immune response and complement activation, it has reported that C4BP could inhibit TLR2 and TLR4 induced proinflammatory reaction through interacting with them (Morita et al., 2017), from the study of human serum C4BP, it showed that C4BP could inhibit islet amyloid polypeptide-mediated inflammasome activation and IL-1β production (Kulak et al., 2017), we found that compared with empty pregnant period, the level of serum C4BP in gestation reduced to the ratio of 0.74, and the level was further down-regulated to the ratio of 0.77 compared with oestrum, and then up-regulated to the ratio of 1.41 compared with gestation group, it indicated that during the period of oestrum and gestation, the Hainan black sheep are more sensitive to inflammatory stimulation, and more preventive methods of anti-inflammation should be adopted.

Coagulation factor IX (FIX) is an essential plasma protein for blood coagulation (Ishikawa et al., 2016) and the protein seemed closely related with hemophilia (Tortella et al., 2018, Santagostino, 2016), but recently it also reported that the
effects of FIX on enhancing cell matrix and intercellular adhesion (Mamiya et al., 2016). The ratio of FIX in oestrus down-regulated to 0.71 compared with empty pregnant, and up-regulated to the ratio of 1.88 compared between gestation and oestrus, finally reduced to the ratio of 0.87 compared between lactation and gestation, it indicated that in the period of oestrus and lactation, more hormones and nutrition nutrient needed to pass through the endothelial cells, so lower level of FIX may increase the permeability of endothelial layer and in favour of the transport efficiency of different kinds of small functional molecules, and in the period of gestation, for the protection of the fetus in uterus, the up-regulated level of FIX mean the small molecules become more difficult to penetrate through the endothelial layer, to avoid hazardous substance and pathogenic bacteria as much as possible.

The protein of Leukocyte Cell-derived Chemotaxin 2 (LECT2) seemed to be more correlated with atherosclerosis (Sonmez et al., 2016), it was also reported that LECT2 is associated with the severity of systemic inflammation in patients with sepsis (Ando et al., 2012) and it involved in the regulation of the homeostasis of natural killer T-cell in mammals (Xu et al., 2014), in the
Fig 8. Histogram of COG annotation. X-axis: The protein amounts been classified into corresponding COG annotation, Y-axis: Items of COG.

Fig 9. Statistical graph of significantly enriched pathway. (A) Enriched pathway distribution of lactation/gestation, (B) Enriched pathway distribution of lactation/oestrum, (C) Enriched pathway distribution of gestation/oestrum, (D) Enriched pathway distribution of empty pregnant/oestrum, (E) Enriched pathway distribution of empty pregnant/lactation, (F) Enriched pathway distribution of empty pregnant/gestation. X-axis: Rich factor showed the ratio of items of differential expression proteins in the specific pathway to all the identified proteins annotated in the corresponding pathway, the larger of this value, the greater ratio of differential expression proteins annotated in the corresponding pathway. Y-axis: Certain pathway items. The size of the dot means the amount of differential expression proteins annotated in certain pathway.
period of oestrus and lactation, the level of LECT2 was down-regulated, it indicated that the Hainan black sheep were more sensitive to pathogenic bacteria and virus challenge during this two period, on the contrary, in the period of empty pregnant and gestation, the disease resistant ability was relatively high. Based on GO analysis, the main function of hyaluronan-binding protein is proteolysis.

It is reported that HABP 2 is a Ca2+ dependent serine protease with putative roles in blood coagulation and fibrinolysis (Staveniuter et al., 2017), and hyaluronan degradation is accelerated in inflammatory and neoplastic diseases (Yoshida et al., 2013), HABP also involved in Hyaluronan depolymerization and its metabolism (Shimoda et al., 2017), The ratio of HABP 2 in oestrus reduced to 0.8 compared with empty pregnant, and up-regulated to the ratio of 2.75 compared between gestation and oestrus, compared with gestation, the level of HABP 2 protein up-regulated to the ratio of 0.36 in lactation, it indicated that in the period of gestation and empty pregnant, the inflammatory status are high, and the inflammatory status in the period of oestrus and lactation are low, and anti-inflammatory steps should be taken in the period of gestation and empty pregnant.

Apolipoprotein C –II (apoC-II) is an important co-factor for lipoprotein lipase (Kei et al., 2012), and based on GO analysis, the function of apo C-II were related to positive regulation of triglyceride catabolic process, phospholipid catabolic process and lipoprotein lipase activity. As apoC-II is a small exchangeable apolipoprotein on triglyceride-rich lipoproteins (Wolska et al., 2017), it could hydrolyze triacylglycerol and reduce the corresponding concentration in serum. The ratio of apoC-II in oestrus reduced to 0.78 compared with empty pregnant, and up-regulated to the ratio of 1.29 compared between gestation and oestrus, compared with gestation, the level of apoC-II protein up-regulated to the ratio of 0.6 in lactation, the data showed the concentration of triacylglycerol in serum of empty pregnant and gestation were relatively higher than the period of oestrus and lactation, which indicated that during the period of empty pregnant and gestation, lower energy level diets could be supplied and correspondingly higher energy level diets should be provided in the period of oestrus and lactation.

CONCLUSION

This is the first study systematically uncovered reproductive cycle characteristics of Hainan black goats, the study preliminarily uncovered the basical protein information between each reproductive period, and the data could contribute to the built of highly efficient reproduction system of Hainan black goats.

ACKNOWLEDGMENTS

This study was financially supported by the key technology research and integration application of large-scale healthy breeding of Hainan black goats (Grant No. ZDKJ2016017) and Research and application of meat key gene screening and molecular assistant breeding of the special wild boar in Hainan Province (Grant No. ZDYF2017059).

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