COMPARATIVE PHYSIOLOGICAL AND MOLECULAR STUDY OF SOME SHEEP BREEDS IN SAUDI ARABIA

1Mohammad S. AL-Harbi, 1Hamed S. Alwagdany and 1,2Sayed A.M. Amer

1Department of Biology, Faculty of Science, Taif University, Taif, 888, Saudi Arabia
2Department of Zoology, Faculty of Science, Cairo University, Giza, Egypt

Received 2013-05-07, Revised 2013-05-17; Accepted 2013-06-06

ABSTRACT

Naemi, Heri, Najdi and Sawakni sheep breeds inhabiting Kingdom of Saudi Arabia were studied on both physiological and molecular bases. Native polyacrylamide gel-electrophoreses for two enzymes Malate Dehydrogenase (Mdh) and Malic Enzyme (Me) recorded 5 isoenzyme loci from which 4 were polymorphic and 1 was monomorphic. Mdh showed three fractions, the first was recorded in Sawakni and Najdi breeds, the second was monomorphic and was recorded in all breeds while the third was recorded in all except Heri. Malic enzyme showed two polymorphic fractions, the first was recorded in Heri and Naemi and the second was recorded in all breeds. Both metabolic enzymes activities were higher in Sawakni than in the native breeds. 607, 227, 498 and 595 bp nucleotides from CO1, ND4, cytb genes and d-loop, respectively, were sequenced with no base substitutions among and/or within breeds except for ND4 gene and d-loop. The molecular tree clustered the three breeds other than Naemi in one group and Naemi was basal. Isoenzymes therefore, discriminate clearly the studied breeds although they are not as accurate as molecular tools indicating their applicability with more efficiency in physiology than in genetics while the d-loop was efficient in discriminating the studied breeds phynotypically.

Keywords: Sheep Breeds, Isoenzymes, Mitochondrial DNA, Control Region

1. INTRODUCTION

Southwest Asia is considered as one of three main areas in which domestication is believed to be occurred very early (Bruford et al., 2003) and domestic sheep (Ovis aries) were among the first domesticated animals. In Saudi Arabia, Naemi, Heri, Najdi and Sawakni breeds are well identified among many local and introduced breeds. The first three are inhabiting the Arabian Peninsula while the fourth one might be introduced from Sudan.

Naemi breed is also called the desert Awassi sheep in Saudi Arabia (Synnot, 1990). Awassi is a local sheep breed from southwest Asia that originated in the Syrian-Arabian desert (Epstein, 1985). Heri breed is related to harat of volcanoes of the north west of Arabian Peninsula. It distributes commonly in different areas of Saudi Arabia mainly Qassim, Hejaz plains, Tehama and Sarawat mountains. Najdi is a well adapted multipurpose breed, used for meat, milk and wool production. Najdi has some unique features such as black hair coat with white head, convex head profile and large, pendulous ears (Pritchard et al., 1977), long legs and fat tailed with coarse fleece (Ali and Al-Noami, 1992). Sawakni is a popular breed imported to the Kingdom from Sudan and thus it is not native to Arabia. This breed is named Sawakni because Sudanese live sheep and sheep meat is recognized in Saudi Arabia markets as ‘Swakni’. It might be the desert Sheep of North Africa and the Horn of Africa (El-Dirani et al., 2009).

Electrophoretic studies were done extensively on the different tissues of various animals from which it reveals that the enzyme exits in multi molecular forms and functions (Markert and Moller, 1959). Malate dehydrogenase is considered as one of the most extensively studied isozyme systems (Lagana et al.,...
2. MATERIALS AND METHODS

2.1. Animals

Twenty three individuals of Naemi, Heri, Najdi and Sawakni sheep breeds from local market of Taif province, Kingdom of Saudi Arabia were used in this study. Blood samples were withdrawn from the jugular vein into heparinized tubes. 300 µL were frozen for the molecular study. The rest of the blood samples were immediately centrifuged at 3000 rpm for 3–5 min and the plasma were decanted for isoenzymatic analyses.

2.2. Isoenzymes

The isoenzymes were separated in 10 % polyacrylamide gel electrophoresis according to (Stegemann et al., 1985) as follows. A volume of 40 µL plasma was mixed with 20 µl sucrose and 10 µ bromophenol blue, then a volume of 50 µL from this mixture was applied to each well. For Malate Dehydrogenase (MDh) (EC 1.1.1.37), the gel was soaked in 100 ml of 0.05 M Tris HCl pH 8.5 containing 25 mg NBT, 25 mg EDTA, 25 mg NAD, 10 mg malic acid and 3 mg PMS (Wendel and Weeden, 1990). For Malic enzyme (Me) (EC 1.1.1.40), the gel was soaked in 100 ml of 0.05 M Tris HCl pH 8.5 containing 25 mg NBT, 25 mg EDTA, 25 mg NADP, 10 mg malic acid, 100 MgCl2 and PMS (Wendel and Weeden, 1990). The gel was kept in a fixative solution (ethanol and 20 % glacial acetic acid, 9: 11 v/v) for 24 h and then was photographed.

2.3. DNA Extraction and PCR Experiments

Mitochondrial DNA was extracted from 0.5 mL blood samples with QIAGEN spin-column kits according to the manufacturer’s instruction. PCR was conducted in a final volume of 25 µL containing 1 µL DNA template, 0.1 µL of 10 Pmol forward primer, 0.1 µL of 10 Pmol reverse primer of the corresponding genes (Table 1), 12.5 µL PCR master mix (Promega Corporation, Madison, WI) and 11.3 µL autoclaved deionized distilled water. PCR was carried out using a Pcx 0.5 thermal Cycler with the cycle sequence at 94°C for 4 min one cycle, followed by 40 cycles each of which consisted of denaturation at 94°C for one min, annealing at corresponding specific temperature (Table 1) for one min and extension at 72°C for one min with a final strand elongation for one cycle at 72°C was done for an additional 5 min. The PCR products were analyzed in 1% agarose gel electrophoresis in TAE buffer (40 mm Tris, 40mM acetic acid and 1mM EDTA) with ethidium bromide staining. A 100-bp DNA ladder (Biolabs) was used as a molecular marker. Then PCR products were visualized under UV light and photographed. The PCR products were then excised from agarose gels and purified using spin column (BioFlux, Tokyo) according to the manufacturer instructions.

2.4. Sequencing

The purified PCR products were sequenced in an ABI PRISM 3730 µL sequencer (Applied BioSystems) and BigDyeTM Terminator Sequencing Kits with AmpliTaq-DNA polymerase (FS enzyme) (Applied Biosystems) following the protocols supplied by the manufacturer.

2006). This enzyme with lactate dehydrogenase, are very suitable systems for studying several metabolic, genetic, ecological features and they are very useful in systematic studies (Almeida-Val et al., 1992). Malate dehydrogenase and malic enzyme are different enzymatically and structurally (Shows et al., 2007). Malate dehydrogenase catalyzes the transformation of malate to oxaloacetate (Zhao et al., 2010), while malic enzyme reversibly catalyzes the oxidative decarboxylation of malate and is a link between the glycolytic pathway and the citric acid cycle (Shows et al., 1970; Song et al., 2001). Both enzymes are important for the NADPH production in ruminants (Laliotis et al., 2006) and NADH Dehydrogenase subunit 4 (ND4) gene (Guo et al., 2006) and NADH Dehydrogenase subunit 4 (ND4) gene (Bruford and Townsend, 1998). Mitochondrial DNA was extracted from 0.5 mL blood samples with QIAGEN spin-column kits according to the manufacturer’s instruction. PCR was conducted in a final volume of 25 µL containing 1 µL DNA template, 0.1 µL of 10 Pmol forward primer, 0.1 µL of 10 Pmol reverse primer of the corresponding genes (Table 1), 12.5 µL PCR master mix (Promega Corporation, Madison, WI) and 11.3 µL autoclaved deionized distilled water. PCR was carried out using a Pcx 0.5 thermal Cycler with the cycle sequence at 94°C for 4 min one cycle, followed by 40 cycles each of which consisted of denaturation at 94°C for one min, annealing at corresponding specific temperature (Table 1) for one min and extension at 72°C for one min with a final strand elongation for one cycle at 72°C was done for an additional 5 min. The PCR products were analyzed in 1% agarose gel electrophoresis in TAE buffer (40 mm Tris, 40mM acetic acid and 1mM EDTA) with ethidium bromide staining. A 100-bp DNA ladder (Biolabs) was used as a molecular marker. Then PCR products were visualized under UV light and photographed. The PCR products were then excised from agarose gels and purified using spin column (BioFlux, Tokyo) according to the manufacturer instructions.
Table 1. Primers designed and used for PCR amplification and sequencing. Annealing temperature refers to that of the conducted PCR to obtain the amplified fragments.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Primer name</th>
<th>Sequence (5’-3’)</th>
<th>Annealing temperature (°C)</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>CO1</td>
<td>Ovis CO1-F</td>
<td>GCTGGTATCACAATACCTACT</td>
<td>56</td>
<td>This study</td>
</tr>
<tr>
<td></td>
<td>Ovis CO1-R</td>
<td>TAGTCCTAGGAAATGCTGTG</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ND4</td>
<td>Forward-OV11</td>
<td>GAC TCC ACC TCT GAC TTC C</td>
<td>57</td>
<td>Yuncu (2009)</td>
</tr>
<tr>
<td></td>
<td>Reverse-OV11</td>
<td>TGA ATG AGA ATG GCA ACA</td>
<td></td>
<td></td>
</tr>
<tr>
<td>cytb</td>
<td>Ovis cytb-F</td>
<td>AGGCCTATTCCTAGCAATAAC</td>
<td>56</td>
<td>This study</td>
</tr>
<tr>
<td></td>
<td>Ovis cytb-R</td>
<td>TAGTAGCATGGCGCCTAAG</td>
<td></td>
<td></td>
</tr>
<tr>
<td>d-loop</td>
<td>Ovis d-loop-F</td>
<td>CGG ACA TGA GCG TTC ATA AAC</td>
<td>57</td>
<td>This study</td>
</tr>
<tr>
<td></td>
<td>Ovis d-loop-R</td>
<td>GGA TGC TCA AGA TGC AGT TA</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

After reading the targeted genes, the nucleotide sequences have been treated with different software programs (DNASIS, MacClade and PAUP) that enabled to detect genetic relatedness between different samples and breeds. The sequenced genes were tested by BLAST program to check their relatedness to the sequenced genes for sheep in the Genbank database. The same published genes for both sheep and goat were taken from the Genbank with their accession numbers (JN632608, NC-009849, NC-009628, respectively) for the necessary alignments and tree construction.

2.5. Statistical Analyses

The isoenzymatic data were subjected to Student t-test in SPSS package to examine the significance level between each two breeds. The obtained DNA sequences were aligned separately and manually using MacClade v.4. The unalignable and gap-containing sites were deleted and the aligned data were then concatenated so that 1900 bp were left for the analyses. The tree analyses were done by Maximum-Parimony (MP) and Neighbor-Joining (NJ) methods with PAUP* 4.0b10 (Swoford, 2002) by heuristic searches with the TBR branch swapping and 10 random taxon additions, respectively. The bootstrapping replicates were set to be 1000 with simple additions for the two methods.

3. RESULTS

Native polyacrylamide gel-electrophoreses for two enzymes (malate dehydrogenase and malic enzyme) recorded 5 isoenzyme loci from which 4 were polymorphic and 1 was monomorphic. Some of these loci exhibited high activity (shown as thick and dark bands) while others were very thin exhibiting low activity.

Mdh showed three fractions in the electrophoretic pattern (Fig. 1). The first fraction (Mdh-1) was recorded in Sawakni and Najdi breeds only. The second fraction was monomorphic and was recorded in all breeds, while the third was recorded in all breeds except Heri. Table 2 showed the means and standard errors for the percentage amount of the studied isoenzymes in the different sheep breeds. Mdh-2 isoform showed a significant increase (p<0.05) in its activity in Heri than in Najdi. The total enzyme was significantly higher in Sawakni (p<0.01, p<0.05) than in Heri and Naemi breeds, respectively. The mean fractional activity of the total enzyme was 27.32±1.94 in Sawakni, 16.38±0.91 in Heri and 17.72±1.72 in Naemi.

Malic enzyme showed two polymorphic fractions in the electrophoretic pattern (Fig. 2). The first fraction (Me-1) was recorded in Heri and Naemi breeds only. The second fraction was recorded in all breeds but some samples within breeds did not show this locus. Table 3 showed the means and standard errors for the percentage amount of the studied isoenzymes in different sheep breeds. Me-2 isoform showed a significant fractional activity increase (S<0.05) in Naemi than in Najdi. The mean fractional activity of this isoform was 7.31±0.84 in Naemi and 3.00±0.82 in Najdi. The mean fractional activity of the total enzyme was significantly higher in Heri and Naemi breeds than in the Sawakni (p<0.01, p<0.05) and Najdi (p<0.05, p<0.05) breeds, respectively. The mean values of the total enzyme activity were 16.9±2.54 in Heri, 19.47±2.29 in Naemi, 3.30±.767 in Sawakni and 5.67±1.40 in Najdi.

Unambiguous nucleotides of 607 bp, 227 bp, 498 bp and 595 bp from CO1, ND4, cytb genes and non-coding d-loop, respectively were sequenced for at least three samples from each breed. These data were deposited in NCBI GenBank database with their accession numbers (KC669571- KC669595) for CO1 and d-loop and (KC689756- KC689785) for cytb and ND4 genes. In order to estimate the base composition and frequencies for the obtained sequences, the data were concatenated and the gap-containing sites were deleted except for the d-loop so that 1900 bp were left for analysis. The data showed base frequencies of A = 30.2%, C = 23.8%, G = 15.8% and T = 30.2% of the 1900 nucleotides used for tree analyses, 1675 were constant and 225 were variables.
Table 2. Mean ± SE of the percentage amount for the studied malate dehydrogenase isoenzymes (Mdhs) in plasma of different sheep breeds. The significant level was estimated by Student t-test. *= significant level between Heri and Najdi, + = significant level between Heri and Sawakni, x = significant level between Naomi and Sawakni.

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Isoform (locus)</th>
<th>Heri</th>
<th>Naomi</th>
<th>Sawakni</th>
<th>Najdi</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mdhs</td>
<td>3</td>
<td>--</td>
<td>5.28±1.49</td>
<td>4.68±0.09</td>
<td>6.51±0.880</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>16.29±0.111</td>
<td>13.76±0.55</td>
<td>15.23±0.89</td>
<td>4.68±0.090</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>--</td>
<td>6.88±0.53</td>
<td>5.96±0.620</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Total</td>
<td>16.38±0.111</td>
<td>17.72±1.72</td>
<td>27.32±1.94</td>
<td>21.35±1.29</td>
</tr>
</tbody>
</table>

Table 3. Mean ± SE of the percentage amount for the studied malic enzyme (Me) in plasma of different sheep breeds. The significance level was estimated by Student t-test. *= significant level between Heri and Najdi, + = significant level between Heri and Sawakni, x = significant level between Naomi and Sawakni.

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Isoform (locus)</th>
<th>Heri</th>
<th>Naomi</th>
<th>Sawakni</th>
<th>Najdi</th>
</tr>
</thead>
<tbody>
<tr>
<td>Me</td>
<td>2</td>
<td>4.40±0.111</td>
<td>7.51±0.84</td>
<td>3.30±0.76</td>
<td>3.00±0.82</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>14.25±1.92</td>
<td>12.15±1.82</td>
<td>--</td>
<td>--</td>
</tr>
<tr>
<td></td>
<td>Total</td>
<td>16.92±2.54</td>
<td>19.47±2.29</td>
<td>3.50±2.765</td>
<td>5.67±1.40</td>
</tr>
</tbody>
</table>

Table 4. Pairwise genetic distance among the different sheep breeds. These distances were estimated from the sequenced data in this study. The bolded values refer to the distances within breeds.

<table>
<thead>
<tr>
<th>Breed</th>
<th>Heri</th>
<th>Najdi</th>
<th>Naomi</th>
<th>Sawakni</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heri</td>
<td>0.0016</td>
<td>0.0050</td>
<td>0.0150</td>
<td>0.0045</td>
</tr>
<tr>
<td>Najdi</td>
<td>0.0005</td>
<td>0.0064</td>
<td>0.0169</td>
<td>0.0042</td>
</tr>
<tr>
<td>Naomi</td>
<td>0.0150</td>
<td>0.0169</td>
<td>0.0042</td>
<td>0.0032</td>
</tr>
</tbody>
</table>

About 190 of the variable sites were parsimony-uninformative and 35 were informative under parsimony criterion. The tree that has been constructed showed consistency index (CI = 0.993), homology index (HI = 0.007), retention index (RI = 0.956) and rescaled consistency index (RC = 0.949).

Single neighbor-joining tree (Fig. 3) was obtained from all data sets with reasonable statistical supports for two computational methods (MP and NJ). The tree showed clustering of each breed with each other except for Najdi which showed closed and/or mixed relationship with Heri. Heri, Najdi and Sawakni breeds clustered in one group with strong bootstrapping (100 BP for both MP and NJ methods). Naomi breed came basal to all breeds studied. As shown in Table 4, the genetic distance showed the smallest values within each breed (0.0016, 0.0042, 0.0064 and 0.0032 for Heri, Naomi, Najdi and Sawakni breeds, respectively) except for Najdi where the smallest distance was found between this breed and Heri (D = 0.005). The tree topology also mixed Najdi with Heri. The interpretation of this mixing maybe attributed to the small sampling size or possible hybridization between these two breeds.

For the protein-coding ND4 gene that codes for NADH dehydrogenase subunit 4, 227 bp were sequenced for different samples. The complete length of this mitochondrial gene in O. aries is 1378 bp as deposited in the Genbank database. Based on the alignment, the sequenced fragment is located between base 648 and base 874. The 227 bp showed 4 base substitutions among and within breeds (Fig. 4). All these substitutions were transitions (purines to purines and pyrimidines to pyrimidines). The mutations occurred among all samples either interspecies or intraspecific were in the third position with no amino acids changes. The synonymous changes involved substitutions of adenine with guanine at G678 → A678, adenine with guanine at G750 → A750, thymine with cytosine at T768 → C768 and guanine with adenine at G849 → A849. The numbers below the base letters referred to the exact position of this base inside the complete gene sequence. Approximately 585 bp of the mitochondrial d-loop region were sequenced for three individuals from each of the sheep breeds. The electropherogram of the obtained sequence for the individuals from the same breed were compared and when a bias was found, the most like base was considered (data not shown). The alignment of this fragment with its counterpart of other sheep breeds published in the Genbank indicated that this fragment is located between 15769 and 16287 of the complete d-loop. The complete d-loop of sheep is 1180 bp long, with four or three copies of repeated 75 bp motif in the tRNA-Pro proximal part. The sequenced fragment of the four breeds herein showed approximately these three motifs (Fig. 5). These repeats contain strong stem-and-loop 40 bp (Fig. 6) secondary structure. Sixteen nucleotides of the repeat are two octamer sequences of mirror symmetry (TTAATGTA, TACATTAA) forming the stable stem. In between this stem, there is a loop of 24 bp. All sheep breeds posses two motifs located immediately upstream of this repeat discriminating two haplotypes which are G/ACCCC (haplotype A) and ACCC/TC (haplotype B).
Fig. 1. The electrophoretic profile of *Mdh* isoenzymes in the studied samples. Lanes are as follow: 1-5 (Heri), 6-9 (Naemi), 10-14 (Sawakni) and 15-19 (Najdi).

Fig. 2. The electrophoretic profile of *Me* isoenzymes in the studied samples. Lanes are as follow: 1-5 (Heri), 6-9 (Naemi), 10-14 (Sawakni) and 15-19 (Najdi).

Fig. 3. Neighbor-joining tree constructed from 1900 bp sequenced fragments of CO1, cytb, ND4 genes and d-loop region for the four sheep breeds used in this study. A goat sample represents the outgroup of the tree. Values at nodes refer to the bootstrapping of maximum-parsimony and neighbor-joining analyses, respectively. Values were shown when they were over 50%.
Fig. 4. The aligned nucleotides of the sequenced ND4 gene for different individuals of the studied sheep breeds. Note the high identity among the different samples. The letters inside the boxes are polymorphic among individuals and/or breeds.
Fig. 5. The aligned nucleotides of the sequenced d-loop region for different individuals of the studied sheep breeds. Note the high identity among the different samples. The letters inside the boxes are polymorphic among individuals and/or breeds. The underlined regions refer to the repeats (each symbol of the underlines refer to mirror image repeats of stems and loops).

Fig. 6. Strong stem-and loop secondary structure of 40 bp inside 75 bp repeat region of sheep breeds d-loop sequenced in this study.

The current breeds are belonging to the haplotype A containing the motif ACCCC or ACCTC. The sequenced fragment of the d-loop region showed 44 substitutions among breeds all of which are transitions except two.
changes in the first repeat which are transversion of adenine to cytosine. Most changes discriminated Naemi breed from others with some individual exceptions occurred in Najdi and Sawakni. The first change among the breeds was found at bases 30-34 (Fig. 6). These four bases can be considered as a key feature discriminating the four breeds from each other. It contains 3 gaps in Naemi, 1 gap in Najdi, GACC in Sawakni and GCCC in Heri. The constructed tree agreed with this finding in discriminating Naemi which came out of the cluster that is containing the other three breeds.

4. DISCUSSION

In the present study, four sheep breeds (Heri, Najdi, Naemi and Sawakni) were investigated biochemically and genetically by studying some metabolic isoenzymes and some mitochondrial DNA traits. The Arabian Najdi and Heri breeds are considered as native breeds. Naemi breed is also called the desert Awassi sheep in Saudi Arabia (Synnot, 1970; Sawakni et al., 1990). Sawakni breed might be the desert Sheep which belongs to seven breeds inhabiting the deserts of Sudan and Somalí (El-Dirani et al., 2009).

Malate dehydrogenase catalyzes the following reversible reaction in the citric acid cycle: L-malate + NAD = oxaloacetate + NADH, while malic enzyme reversibly catalyzes the oxidative decarboxylation of malate and is a link between the glycolytic pathway and the citric acid cycle: L-malate + NADP = pyruvate + CO₂ + NADPH (Showes et al., 1970; Song et al., 2001). It is therefore noteworthy to confirm that Malate Dehydrogenase (Mdh) is enzymatically and structurally distinct from malic enzyme (Showes et al., 1970). Each enzyme can be separated into a cytosolic and a mitochondrial molecular form by cellular location, physicochemical properties and gel electrophoresis (Hsu et al., 1967). Both enzymes are among the principal enzymes that are responsible for the NADPH production in ruminants (Laliotis et al., 2010). The first reaction (catalyzed by Mdh) plays a key part in the malate/aspartate shuttle across the mitochondrial membrane and in the citric acid cycle (Minarik et al., 2002) while the second (catalyzed by Me) plays a role in lipid biosyntheses.

Minarik et al. (2002) stated that in eukaryotic cells, at least two forms of Mdh can be found. One isoform (Mdh-2) is a principal enzyme of the citric acid cycle operating within mitochondria. The other (Mdh-1) is found in the cytosol where it participates in the malate/aspartate shuttle. This shuttle exchanges reducing equivalent across the mitochondrial membranes in the form of malate/oxaloacetate. A third (Mdh-3) isoenzyme was found in the glyoxysomes of yeast, where it converts malate produced from glyoxylate in the glyoxylic cycle (Minard and McAlister-Henn, 1991). All Mdh isoforms are NAD-dependent. The enzymes share a common catalytic mechanism and their kinetic properties are similar, which demonstrates a high degree of structural similarity.

Like goat (AL-Harbi and Amer, 2012), sheep breeds, in the present study, recorded three isoforms of Mdh. The cytosolic Mdh-1 was not shown in Naemi and Heri breeds but it was approximately detected in Najdi and Sawakni breeds and the mitochondrial Mdh-2 was fixed in all breeds. As Mdh enzyme catalyzes the reversible shuttle down of malate-oxaloacetate pathway (Zhao et al., 2010) in the energetic citric acid cycle, it is therefore possible to elucidate that the metabolic rate of the Najdi and Sawakni breeds could be more energetic than the other two breeds.

In mammalian tissues three distinct isoforms have been described; a mitochondrial NAD-isofrom and two NADP-dependent isofroms, a first localized in cytosol (Me-1) and a second occurred in mitochondria (Me-2). Cytosolic malic enzyme (Me-1) is considered as a NADP-donor for fatty acid synthesis. It is also involved in the supply of fatty acids with the essential acetyl-coA. Acetyl-coA is produced in mitochondria but it is essential for fatty acids biosynthesis taking place in cytosol (Pearce, 1983). In ruminants, contrary to humans and rodents, the pathway of glucose-pyruvate-acetyl-coA is of little significance, as the principal carbon source for lipogenesis instead of glucose, is the acetic acid produced by the rumen’s microorganisms (Bergen and Mersmann, 2005). In sheep, unlike to other species, two transcripts encoding ovine Me-1 has been reported (Stefos et al., 2009), which may further elucidate possible explanations for the minor role of cytosolic malic enzyme in these species. It is therefore likely to expect the little expression of this enzyme in the studied breeds. The two fractions of Me showed expression in most of individuals of Heri and Naemi while the second fraction was found in all breeds. Since Me has a role in lipid biosynthesis (Zhang et al., 2007), it seems that Heri and Naemi breeds may possess more adaptability of lipid biosynthesis as these two breeds are more desert in their habitat than Sawakni and Najdi.

There was no variation in CO1 gene among all samples and breeds. CO1 data therefore supported the stability of this gene in all breeds studied. Cytochrome C oxidase is the terminal complex of the electron transport chain and is activated to prevent an excessive buildup of reactive oxygen species (Chen et al., 2009). It is also not affected by the variation in the respiratory capacity (Devin and Rigoulet, 2007). These two reasons may
explain the similarity in the sequence of the gene coding for this protein in all sheep breeds. Ahmed et al. (2013) recorded similar finding for the Arabian camel.

Cytb gene also did not show any variation among the studied breeds. The hydrophilic protein of cytochrome b acquires higher mutations in abnormal cases of skeletal muscle weakness and exercise intolerance (Fernandez-Vizarra et al., 2007). It is one of the cytochromes which showed variations when the respiratory capacity changes (Devin and Rigoulet, 2007). It is therefore possible to correlate the identity in the sequence of this gene to the similarity in the respiratory capacity of different sheep breeds.

The only gene which showed slight variations among the breeds was ND4 gene. The inherited variation in mitochondrial genes (ND5 including ND4), in the absence of variation in the nuclear genome and other confounding factors, can influence glucose and lipid metabolism (Houstek et al., 2012). This may be in agreement with the variation in the isoenzyme pattern of both metabolic enzymes of Mdhi and Me and therefore the slight variations within this gene among the breeds may reflect some roles in lipid and glucose metabolism.

The analysis of the sequenced fragment of the d-loop region, in the present study, revealed that most individuals acquired three repeats (Heri and Naemi) and some acquired two repeats (Sawakni and Najdi). Hindleider et al. (1998) found three, four and five repeats in the d-loop of two sheep haplotypes (A and B). The discrepancy in repeat number between this study and that of the authors could be due to that, we sequenced partial part of the d-loop and this repeat is located near to the end of the d-loop. The proposed stem-loop structures within these repeats have been suggested as recognition sites for the arrest of H strand synthesis (Saccone et al., 1991).

Aside from three insertions/deletions and a single transversion, all variable nucleotide positions in sheep breeds control region represent transitions from A to G or C to T and vice versa. The variability among and within breeds was also recorded within these repeats and it was greater among breeds than within them indicating the efficiency of this molecular marker in discriminating among breeds. Such repeat has been described for other vertebrates in this region (Lunt et al., 1998; Brearley and Zhou, 2001) agreed with (Dionne et al., 1991; Rivera et al., 1997) in that there was no significant relationship between d-loop polymorphism and physiological performance. Based on these arguments, we could not able to relate the polymorphism in the d-loop repeat to the difference in physiological performance but to phenotypic differences among the studied breeds.

5. CONCLUSION
From the results, it could be concluded that the genetic variability among sheep breeds is very weak when we use the protein-coding genes where these genes are more impressive for the physiological role. The variability could be clearly addressed when the d-loop region was used where it controls the mtDNA replication. Isoenzymes discriminated clearly among the studied breeds although they are not as accurate as molecular tools indicating their applicability with more efficiency in physiology than in genetics. It is therefore necessary to recommend further molecular study using d-loop region in discriminating among Saudi Arabian sheep breeds and to conduct more isoenzymatic studies to address their physiological adaptations.

6. REFERENCES


