MOLECULAR CHARACTERIZATION OF TUMOR NECROSIS FACTOR-Α (TNFA) GENE IN EGYPTIAN RIVER BUFFALOES

Aboelenin, M.M1; K.F. Mahrous1, M.A. Rashed2 and M.A. Sallam2
1- Cell Biology Dept., National Research Centre, Giza, Egypt
2- Genetics Dept., Faculty of Agriculture, Ain Shams University, Cairo, Egypt

Keywords: Egyptian river buffaloes, Tumor Necrosis Factor-α, PCR-RFLP, Single nucleotide polymorphism

ABSTRACT

Tumor Necrosis Factor-α (TNFα) is a cytokine signaling protein which has an important role in the immune system regulation and affect female reproductive performance. The objective of this study was investigation the TNFα gene polymorphism and its potential effects in female buffaloes fertility. The DNA was extracted from the blood of 81 buffalo females and a 592 bp fragment contains the full coding region of TNFα-exon 4 was amplified by PCR which subsequently treated with Rsal restriction enzyme. The PCR-RFLP pattern showed that all the animals had fixed CC genotype and T allele was not detected. Sequencing of amplified fragment (GenBank accession No. KY885010) flowed by sequence alignment with GenBank database revealed that the river buffalo target sequence was homologues to cattle than goat, sheep, human and mouse on DNA and amino acids levels. Comparison of TNFα amplicon with homologues Bubalus bubalis records in the GenBank detected 3 SNPs in exon 4. Two of these SNPs were synonymous while the third located in the 3’ UTR. Different effects of some of discovered SNPs on RNA cis-regulatory elements and hn mRNA and mature mRNA secondary structures were predicted.

INTRODUCTION

Tumor Necrosis Factor-α (TNFα) is a cytokine signaling protein which is a mediator of inflammation and has several functions in the immune system that include regulation of the immune response against infections, lymphoid tissue development at the lymph nodes and apoptosis induction (Cordero-Coma and Sobrin, 2015). Moreover, many evidence suggested a possible role for TNFα alone induces ovulation in some rat ovary while luteinizing hormone (LH) injection in combination with TNFα increased both of ovulation rate and progesterone secretion compared to LH alone (Brannstrom et al 1995). Furthermore, local injection of TNFα antiserum in the ovarian follicle blocked ovulation in the ewes ovary (Murdoch et al 1997). TNFα gene in river buffalo was assigned to chromosome BBU 2p21-22 (Iannuzzi et al 2015). According to GenBank record with accession number JQ043173.1 the river buffalo TNF-α gene is about 2 kb long and has 4 exons and 3 introns.

Higuchi et al (1999) discovered a C to T SNP within TNFα-exon 4 was recognized by polymerase chain reaction-restriction fragment length polymorphism (PCR-RFLP) method using restriction enzyme Rsal (RFLP- Rsal). The SNP was synonymous and locates in the third nucleotide within a codon codes for Tyrosine (Y). In a study on this SNP in 170 Holstein cattle using the same technique Shirasuna et al (2011) found that CC genotype had earlier ovulation within 3 weeks postpartum and shorter days open than TT genotype. Genotyping of the same SNP in 315 Holstein cows but using the restriction enzyme AfaI (RFLP- AfaI) revealed that the first ovulation after parturition was earlier in the animals with CC (uncut PCR product) and CT genotypes than the animals with TT genotypes (Kawasaki et al 2014).

Although the buffalo (Bubalus bubalis) can adapt to harsh environments and poor quality for-
age, the buffalo has been traditionally regarded as a poor breeder due to its poor fertility in the majority of conditions under which they are raised. The reproductive problems could be manifested mainly as late maturity, long postpartum anoestrous intervals, poor expression of oestrus, poor conception rates and long calving intervals (Perera, 2011). So, in the present study, RFLP assay and DNA sequence analysis were performed to examine genotypes and screen SNPs in exon 4 of TNFα gene in female Egyptian buffaloes.

MATERIALS AND METHODS

This investigation was carried out in Cell Biology Department, National Research Centre, Giza, Egypt.

Blood collection and DNA extraction

A total of 81 blood samples were collected from healthy and unrelated Egyptian river buffalo females. 30 samples were collected from the farm of Cattle Information System/Egypt (CISE) and 51 samples were collected from Agricultural Experiments Station (AES), Faculty of Agriculture, Cairo University, Giza, Egypt. 10 ml of blood samples were collected in sterile 15 ml tubes containing 0.5ml of 0.5 M EDTA solution (PH 8.0). Genomic DNA was extracted from the whole blood according to the method described by Miller et al (1988) with minor modifications. The DNA concentration was determined, using Nano Drop1000 thermo scientific spectrophotometer and then diluted to the working concentration of 50 ng/μl.

Animals genotyping and TNFα-exon 4 sequencing

A DNA fragment which includes all the coding region of TNFα-exon 4 and part of its 3’ untranslated region (3’ UTR) were amplified using the primer pair TNF4-F (5’CAGGATGTTGAGAGTAGAAGCTGA3’) and TNF4-R (5’TGAGCCCTAATTCCCTTTCTAA3’) which were designed using Primer3 online software (Yazdani et al 2010, http://primer3.ut.ee/) based on the sequence of GenBank record JQ043173.1. Polymerase Chain Reaction (PCR) were performed in a 25 µl reaction volume, which included 50 ng genomic DNA, 50 ng each primer, 200 µM each dNTP, 2.5 µl 10X PCR buffer and 0.5 µTaq DNA polymerase (Promega, Madison, WI, USA). Amplification was carried out in a thermocycler programmed as follows: An initial start separation cycle at 94°C for 2 min, 35 cycles including a denaturation step at 94°C for 30 sec, an annealing step at 60°C for 30 sec and polymerization step at 72°C for 45 sec. A final extension cycle at 72°C for 10 minute. The PCR products were screened by electrophoresis on 2% agarose gel in 0.5X TBE buffer stained with ethidium bromide and were visualized with UV transilluminator.

PCR product was digested by RsaI restriction enzyme (Thermo Scientific, Dreieich, Germany) at 37°C for 30 min according to the procedure provided by the manufacturer. Digested products were separated by electrophoresis in 2% agarose gel in 0.5X TBE buffer stained with ethidium bromide and were visualized with UV transilluminator.

Purification and sequencing of PCR products

The PCR products of selected samples were purified using GeneJET Gel Elution Kit (Thermo Scientific, Dreieich, Germany). The purified PCR products were sequenced using an automated sequencing service (Macrogen, Korea).

Data analysis

Alignment of the sequences were performed with GenBank database BLAST tool (http://www.ncbi.nlm.nih.gov/BLAST/) to identify the homology between target TNFα region and the other TNFα sequences on the GeneBank database. The effect of discovered SNPs on splicing site strength, RNA cis-regulatory elements and pre and mature mRNA secondary structures were predicted using SplicePort (Dogan et al 2007), RegRNA 2.0 (Chang et al 2013) and RNAasn (Sabarinathan et al 2013) online softwares respectively.

RESULTS AND DISCUSSION

The time needed to resume the ovarian activity after parturition has an important role in fertility postpartum in dairy animals. A cytosine to thymine SNP within TNFα-exon 4 was found to have a statistical correlation with first ovulation postpartum in cattle females (Shirasuna et al 2011 and Kawasaki et al 2014). Depending on high degree of similarity between the buffaloes and cattle genomes (Tantia et al 2011) the association between this SNP and reproductive performance of buffalo females was examined in this study.

PCR-RFLP genotyping

Arab Univ. J. Agric. Sci., 25(2), 2017
The DNA was extracted from 81 buffalo females. The target fragment contains the full coding region of TNFα-exon 4 was successfully amplified by PCR using the designed primers and a PCR product with 592 bp was validated (Fig. 1-A). Subsequently, the PCR product was treated with Rsal and the pattern of PCR-RFLP was analyzed also by electrophoresis in 2% agarose gel (Fig. 1-B).

In this study no polymorphism in the target position within TNFα-exon 4 was discovered among all buffalo females. All the animals had a single PCR-RFLP pattern consists of 2 bands had the molecular weights 232 and 360 bp approximately (Fig. 1-B), this pattern represents CC genotype.

This result was confirmed by DNA sequencing of TNFα PCR product of different samples which shows a cytosine nucleotide located in the SNP of interest site and the restriction site of Rsal enzyme (Fig. 1-C).

In contrast to this result which confirms absence of T allele in the tested buffalo females, T allele frequency was 31.36% (Shirasuna et al 2011) and 57.08% (Kawasaki et al 2014) in Holstein cows. As the result to the identified monomorphism in the target SNP site it was not possible find a statistical relationship between this SNP and fertility of buffaloes females.

**Fig. 1.** PCR Product, PCR-RFLP pattern and sequencing result of amplified TNFα-exon 4 fragment. (Fig. 1-A) 2% agarose gel electrophoresis of amplified PCR product of TNFα-exon 4 fragment. M= 100 bp molecular marker. Lanes 1, 4 and 5= Animals from CISE. Lanes 2, 3 and 6= Animals from AES. (Fig. 1-B) PCR-RFLP pattern of TNFα-exon 4 amplicon after digestion by Rsal. M= 100 bp molecular marker. Lanes 1-3= Animals from CISE. Lanes 4-6= Animals from AES (Fig. 1-C) A part of DNA sequencing chromatogram of TNFα-exon 4 amplified fragment. The black box indicates the restriction site of Rsal and the wide arrow pointing to the target C to T SNP position.
Sequencing and analysis of amplified sequence

Sequencing of amplified fragment (GenBank accession No. KY885010) followed by searching for sequence similarity using BLAST software against GenBank database revealed that the amplified 592 bp sequence consists of a part of intron 3 (nucleotides from 1 to 69), full coding region of TNFα-exon 4 (nucleotides from 70 to 488), stop codon (nucleotides from 489 to 491) and 3' untranslated region (3' UTR - nucleotides from 492 to 592). The first 2 nucleotides in the 5' of coding region represent the second and third letters in a codon starts with the last nucleotide in the 3' end of exon 3, while the remaining nucleotides in the coding region of TNFα-exon 4 code for 139 amino acids represent about 59.4% of the total amino acids of buffalo TNFα protein (GenBank accession numbers JQ043173.1 and ABO15707.1).

Nucleotides sequence homology analysis of target sequence in orthologous TNFα genes showed there is a similarity between the buffalo sequenced fragment and cattle (98%), sheep (96%), goat (95%) and human (81%) while mouse TNFα-exon 4 do not show significant homology (Fig. 2-A). The phylogenetic tree of the nucleotides sequences (Fig. 2-B) approved that the TNFα-exon 4 sequence is more homologues between buffalo and cattle while the sequence of sheep is more homologues to goat.

Fig. 2: Multiple sequence alignment and phylogenetic tree of nucleotide sequences of TNFα in some species. Multiple sequence alignment (A) and phylogenetic tree (B) of nucleotide sequences of TNFα-exon 4 in Egyptian buffaloes (Bubalus bubalis), cattle (Bos taurus), sheep (Ovis aries), goat (Capra hircus) and human (Homo sapiens).
On the other hand, the nucleotide sequences of buffalo, cattle, sheep and goat are more similar than the human sequence. These results is in accordance with the fact that both of buffalo cattle belong to subfamily Bovinae while sheep and goat belong to subfamily Caprinae and the 4 species belong to family Bovidae but human belongs to more distance taxa within the taxonomic tree of kingdom Animalia (https://www.itis.gov).

On the level of amino acids sequence the homology between buffalo amino acids sequence and those of cattle, sheep, goat, human and mouse was 100%, 92%, 91%, 82% and 73% respectively (Fig. 3-A). The amino acids sequence of TNFα fragment is completely similar between buffalo and cattle while the sequence of sheep is more similar to goat, and the amino acids sequences of buffalo, cattle, sheep and goat are more homologues than the human and mouse sequences (Fig. 3-B) which are in line with in the taxonomic relationship among these species that mentioned above.

Moreover, the sequence identity equal 100% between amino acids sequence of TNFα-exon 4 compared to 98% nucleotides sequences identity which mean that all the 6 SNPs between buffalo and cattle DNA sequences that located in the coding region of TNFα-exon 4 (Fig. 2-A) are synonymous and do not alter the amino acid sequence which is conserved between the 2 species.

**Comparison of the sequence of amplified TNFα fragment with homologues Bubalus bubalis sequences in the GenBank database indicate 3 SNPs among these sequences which listed in Table (1). Interestingly, all the detected SNPs among Bubalus bubalis sequences are similar to 3 out of 11 SNPs were found between river buffalo and cattle sequences (Fig. 2-A). 2 SNPs (A 110G and...
Table 1. The detected SNPs between amplified TNFα sequence and its homologues *Bubalus bubalis* sequences in GenBank database.

<table>
<thead>
<tr>
<th>SNP</th>
<th>Position</th>
<th>Mutated codon</th>
<th>Type of amino acid mutation (position)</th>
<th>Alleles</th>
<th>GenBank accession number</th>
</tr>
</thead>
<tbody>
<tr>
<td>A110G</td>
<td>exon 4</td>
<td>TCA to TCG</td>
<td>Synonymous S to S</td>
<td>A &amp; G</td>
<td>EB-TNFα, NW_005782253.1, XM_006041931.1, XM_006041930.1, AB246789.1, EF562450.1, EF562449.1, EF424255.1, AY221123.1 &amp; KX029323.1</td>
</tr>
<tr>
<td>G383A</td>
<td>exon 4</td>
<td>GGG to GGA</td>
<td>Synonymous G to G</td>
<td>G</td>
<td>EB-TNFα, NW_005782253.1, JQ043173.1, XM_006041931.1, XM_006041930.1, AB246789.1, EF562450.1, EF562449.1, EF424255.1 &amp; AY221123.1</td>
</tr>
<tr>
<td>C510T</td>
<td>3’ UTR</td>
<td>-</td>
<td>-</td>
<td>C &amp; T</td>
<td>EB-TNFα, NW_005782253.1, JQ043173.1, XM_006041931.1 &amp; XM_006041930.1 &amp; AB246789.1</td>
</tr>
</tbody>
</table>

G383A) were located in the coding sequence while the third SNP (C510T) was found in 3’ UTR.

Both of A110G and G383A are synonymous SNPs and did not change the amino acid sequences of river buffalo TNFα protein. Based on the homology between river buffalo and cattle genomes (Tantia et al 2011) we used the cattle codon usage table (http://flowgene.az/) as reference to compare the codon usage in river buffalo. G383A SNP altered the GGG codon to GGA codon which code to amino acid glycine and have similar frequency (25 and 23% respectively). On the other hand, A110G SNP changed TCA codon to TCG codon and both of them code for the amino acid serine but the first codon frequency is 14% compare to second codon which has lowest frequency (7%) among the 6 codons that code for serine. The synonymous codons with different codon usage may lead significant change in protein translation level in the eukaryotic cells (Powell and Dion, 2015).

The strength of intron 3-3’ splicing site was calculated using Splice Port online software (Dogan et al 2007) and the score estimated to be 1.48. Both of G383A and C510T SNPs did not affect the strength of 3’ splicing site while A110G SNP increased the 3’ splicing site score to be 1.79. The DNA sequence was scanned using RegRNA 2.0 online software (Chang et al 2013) to investigate the effect of discovered SNPs in the RNA cis-regulatory elements within the TNFα pre-mRNA. The A110G and G383A SNPs created 2 exonic splicing enhancers are recognized by the splicing regulatory proteins SC35 and ASF/SF2 (Table 2). SplicePort software calculates the score of the splicing site based on ±80bp around the splicing site which explains 3’ splicing site score rising due the exonic splicing enhancer which formed by A110G SNP (about 40 bp upstream 3’ splicing site) compared to exonic splicing enhancer which formed by G383A SNP that did not affect the 3’ splicing site score because it positioned outside this region.

Table 2. The effect of detected buffalo TNFα gene SNPs on RNA cis-regulatory elements within TNFα pre-mRNA.

<table>
<thead>
<tr>
<th>SNP</th>
<th>Effect</th>
<th>Motif</th>
<th>Binding factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>A110G</td>
<td>Creat: Exonic splicing enhancer - Homo sapiens</td>
<td>105 gactcGta 112</td>
<td>SC35</td>
</tr>
<tr>
<td>G383A</td>
<td>Creat: Exonic splicing enhancer - Homo sapiens</td>
<td>377 gggaggA 383</td>
<td>ASF/SF2</td>
</tr>
</tbody>
</table>
Molecular characterization of tumor necrosis factor-A (tnfa) gene in Egyptian river buffaloes

The secondary structure of both of mRNA of pre-mRNA and mature mRNA affect several biological processes such as transcription, translation, mRNA splicing and mRNA stability (Wan et al. 2014). Depending on this the local secondary structure of river buffalo TNFα gene transcript was predicted using RNAsnp online server (Sabarinathan et al. 2013).

The program parameters were adjusted to compute the mRNA secondary structure ±100 nts around the SNP, so the effect of A110G SNP on the secondary structure of pre and mature mRNA was calculated separately because this SNP is pointed 40nts upstream exon-4 starting site which mean that the remaining nucleotides (60 nucleotides out of 100) downstream this SNP which used to compute the secondary structure are different between the pre and mature mRNA. The regions ±100 nts around G383A and C510T SNPs were identical in pre and mature mRNA.

The secondary structures of both alleles of A110G (on the pre-mRNA level) and C510T (in pre and mature mRNA levels) SNPs were identical although there were some differences in the calculated minimum free energy (MFE) (Fig. 4-A & D). on the contrary, minor difference within the secondary structure and MFE in the level of one base was detected between the alleles of A110G (on the mature mRNA level) and G383A (in pre and mature mRNA levels) SNPs (Fig. 4-B & C) but those differences were not significant ($P > 0.2$).

Fig. 4. Predicted changes in pre and mature mRNA local secondary structure of river buffalo TNFα gene transcripts due to detected SNPs. A and B: effect of A110G SNP on secondary structure of pre-mRNA and mature mRNA respectively. C and D: effect of G383A and C510T SNPs respectively on mRNA secondary structures which is the same in pre- and mature mRNA. Left panels represent secondary structure of wildtype sequence while right panels represent secondary structure of mutant sequence. MFE= minimum free energy of each structure. The arrows refer to SNP position.
CONCLUSION

A 592 bp fragment was amplified by PCR and PCR-RFLP using Rsal revealed that all the animals had CC genotype. A similarity between the amplified nucleotides sequence in buffalo and cattle (98%), sheep (96%), goat (95%) human (81%) was observed while mouse TNFa-exon 4 do not show significant homology. The homology between the amino acids sequence of buffalo TNFa fragment and those of cattle, sheep, goat, human and mouse was 100%, 92%, 91%, 82% and 73% respectively. Three SNPs were discovered between amplified TNFa fragment and homologues Bubalus bubalis sequences in the GenBank. 2 synonymous SNPs (A110G and G383A) were located in the coding sequence while the third SNP (C510T) was found in 3' UTR. The A110G and G383A SNPs created 2 exonic splicing enhancers. The secondary structures of both alleles of A110G (on the pre-mRNA level) and C510T (in pre and mature mRNA levels) SNPs were identicals, while minor difference within the secondary structure and MFE levels in the level of one base was detected between the alleles of A110G (on the mature mRNA level) and G383A (in pre and mature mRNA levels) SNPs but those differences were not significant (P>0.2).

REFERENCES


http://flowgene.az/
http://primer3.ut.ee/
https://www.itis.gov/