





DETECTION OF SNPS IN EXON7 LOCUS OF *CYP19* GENE AND THEIR ASSOCIATION WITH ANESTRUM IN EGYPTIAN BUFFALOES (*BUBALUS BUBALIS*)

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A B S T R A C T

CYP19 is a member of cytochrome P450 aromatase family, which regulates estrogen biosynthesis in animals so it can be used as a candidate gene for marker-assisted selection strategies for fertility in buffaloes. The sequence of this gene in buffalo was recently published but association study has not conducted yet. Therefore, the objective of this study was to detect polymorphisms in a *CYP19* locus containing exon 7, and to determine associations between these polymorphisms and anestrum in Egyptian buffalo. A PCR product of 241bp, containing exon 7 of *CYP19* gene was successfully amplified using PCR and subsequently, subjected to single strand conformation polymorphism (SSCP) followed by sequence analysis to identify its different allelic patterns. There was no any SNP in all examined Egyptian buffaloes as shown by one SSCP pattern and nucleotide sequencing. Consequently, these results indicated that this locus does not influence anestrum in Egyptian buffalo and further studies should be conducted to detect other SNPs associated with fertility in Egyptian buffalo.

Keywords: CYP19, buffalo, PCR, SSCP, sequencing, SNPs, anestrum.

(BVMJ-26(1):151-160, 2014)

1. INTRODUCTION

iver buffalo is a species of great economic potential, especially in developing countries; it is the main source of meat and milk in Egypt (Othman, 2013). According to the last estimate, there are about 195 million buffalo in the world today, 97% in Asia, 2% in Africa mainly in Egypt, and 0.2% in Europe mainly in Italy (FAO, 2013). The limited productivity of Egyptian buffalo is due to many reasons including inbreeding, feeding and health care. However, the major problem seems to be infertility that is much higher than that in cattle. Anestrum (in form of silent heat and long calving interval) have been recognized as major cause of infertility (Ahmed et al., 2010; Derar et al., 2012; Fooda et al., 2011), the animal is non pregnant with smooth

important

examination. One of the physiological causes for true anestrum is a low level of ovarian estrogens (Hafez and Hafez, 2000). This disease is not only lower animal productivity and fertility but also bring economic loss which may be due to extension of the dry period as well as longer calving interval during the life time of the animals (Kumar et al., 2011). Genetic evaluation animal reproductive of depends performance on molecular technology for identifying genes and analysis of the polymorphism of these marker genes whose products are key enzymes in the metabolic pathways of important physiological processes and are

ovaries giving no palpable evidence of

either follicular or luteal activity on rectal

related to phenotypes (Beuzen et al., 2000). The key enzyme in estrogen biosynthesis is cytochrome P450 aromatase, the protein product of CYP19 gene. The role of aromatase is the conversion of androgens to estrogens, and is essential for physiology of reproduction (Jedrzejczak et al., 2011). The bovine CYP19 gene have been mapped to the long arm of chromosome 10, band 2.6 in cattle (Goldammer et al., 1994), chromosome 11 in buffalo (Iannuzzi et al., 2001) and range from 56 kb to 120 kb in different species. It consists of 10 exons. The coding region includes exon II-X with translation start site in exon II (Simpson and Davis, 2001). To detect mutations and single nucleotide polymorphisms, the candidate gene should be amplified in small fragments and these fragment should then be analyzed by rapid technique as single strand conformation polymorphism (SSCP) to locate and identify the nature of mutation polymorphism (Lakhotia or and Somasundaram, 2003); (Orita et al., 1989). It seems that, no study was conducted to identify the CYP19 polymorphisms and association with reproductive their performance in Egyptian buffaloes. Therefore, the aim of this study was to isolate CYP19 gene locus containing exon7 and to detect any polymorphisms associated with anestrum in Egyptian buffalo cows.

2. MATERIAL AND METHODS

2.1. Animals source and grouping

The current study was conducted on 180 buffalo cows and heifers selected from Buffalo Nucleus Herd, Animal Production Research Institute, Ministry of Agriculture. Heifers were naturally served for the first time when they reach 300 to 350 kg of body weight and/or 24 months of age. Buffalo cows should be dried off two months before the expected calving date, and they were served not before two months after calving. Animals were assigned in two main groups: normal fertile and infertile due to anestrum. Buffalo cows with 5 successive calving was grouped as normal fertile, heifers which didn't show oestrus over two years of age and buffaloes which didn't ovulate over sex months after calving was grouped as anoestrus.

2.2. Total DNA extraction

Blood samples were collected in EDTAcontaining vacutainer tubes (kept in ice box) from jugular veins of 40 animals in each group. The genomic DNA was extracted from the leucocytes using Gene genomic DNA purification JET kit manufacturer following the protocol (Fermentas, #K0721). The concentration of extracted DNA total was spectrophotometrically determined at 230 and 260 nm using Nanodrop then stored at -20°C.

2.3. Polymerase chain reaction (PCR)

The CYP19 locus was amplified by PCR using primers (Table 1) designed by Primer 3.0 software based on the published sequences of Indian buffalo (GenBank accession number, EF126034). The PCR was carried out in a reaction volume of 50 µL, containing 4.0 µL DNA template (approximately 100 ng), 10 µl Dream Taq Green PCR master mix 5x (Fermentas, #K1071, European Union), 2.0 μL (10 umol/L) forward primer, 2.0 µL (10µmol/L) reverse primer, and 32 µL nuclease free water. PCR was performed by employing a follows: PCR program as Initial denaturation step at 94°C for 2 min, then tubes were subjected to 35 cycles of 94°C for 1 min, 60°C for 1 min and 72°C for 1 min, followed by a final extension step at 72°C for 10 min. As a negative control, tubes were prepared with water instead of DNA template. Then PCR products were resolved by electrophoresis on 2% agarose gel in 1X TAE, stained with ethidium bromide and visualized with UV light of Gel Documentation System (Biometra Biomedizinische Analytik, GmbH).

2.4. Single Strand Conformation Polymorphism (SSCP)

The procedure was followed according to kumar et al (2009) with some

modifications. The PCR products (5 µl) were mixed with 5 µl of SSCP dye (95% formamide, 25 mM EDTA, 0.025% xylenecyanole and 0.025% bromophenol blue) in 0.2 ml PCR tubes, then incubated at 95°C for 10 min for denaturation and plunged in ice for 5 min to form single strand conformers then electrophoresed in a 12% polyacrylamide gel (ratio of acrylamide to bis-acrylamide was 39:1 [0.5 gram of bisacrylamide and 19.5 gram of acrylamide was added to 50 ml distilled water and mixed well in water bath at 37°C till complete solubility, 5 ml 10% TBE (contained 108g Tris, 55g boric acid +40 ml 0.5M EDTA [37.22g EDTA in 150 ml of distilled water] and up to 1000 ml deionized water and mixed well), 12 ml of 20 % acrylamide to bis-acrylamide mix were added to 8 ml deionized water, 20 µl of TEMED (tetra methylene diamine) and finally 200 µl of 10% Ammonium persulfate was added and mixed well. The optimal polymerization time was about 4 h. The gel was pre electrophoresed at 160 V for 30 min with 0.5x TBE as electrode buffer. The conformers of PCR products were separated in the gel at 14–16°C using constant voltage of 160 V for 4 h. The DNA fragments in the gel were detected by 500ml of Ethidium bromide 0.5 µg/ml in 1x TBE for 10-30 minutes on a rocking platform for proper fixing of SSCP conformers in the gel. Then the gel was transferred gently in 500ml of sterile distilled water for destaining. The fragment patterns were visualized on the UV Trans-illuminator and photographed by gel documentation system (UVDI Major Science, USA).

2.5. DNA Sequencing

After getting purified PCR products (clones) with expected sizes, the clone was purified using PCR purification kit following the manufacturer protocol (Jena Bioscience # pp-201×s) to remove primer dimmers, primers, nucleotides, proteins, salt, agarose, ethidium bromide and other impurities. The PCR products were sequenced automated in sequencer

(Applied Biosystem, USA) using CYP19 primers. The Sequences were analyzed using the Chromas Lite 2.1 program (http://technelysium.com.au/?page_id=13) and the identity of the sequenced PCR product was examined using Blast search against Genbank database of Indian buffalo and cattle (Bos Taurus) (http://blast.ncbi.nlm.nih.gov/Blast.cgi). The alignments, annotations and assembly of the sequences were performed using 4.8.4 software Geneious http://www.geneious.com/web/geneious/ho me.

3. RESULTS

The PCR product with the expected size (241bp) was shown by 2% agarose gel electrophoresis (Figure 1). Then, the resulting PCR products were purified using purification technique. gel No polymorphism was detected in CYP19 locus as revealed by presence of only one SSCP banding (monomorphic) pattern (Figure 2). Sequencing was conducted to verify the results of SSCP. The sequences of this CYP19 locus in Egyptian buffaloes (submitted to GenBank with accession number KF957995) showed no polymorphism (Fig. 3). Nucleotide sequences alignment of this CYP19 locus showed 100% identity with Indian water buffaloes (EF178281) (Fig.4) and 98% identity with Bos Taurus (Z69247) (Fig.5).

4. DISCUSSION

Genetic evaluation of animal reproductive performance depends on molecular technology for identifying genes of reproductive efficiency and analysis of the polymorphism of these marker genes (Beuzen et al., 2000). Taking into account various functions and influences of estrogens on the regulation of reproductive processes, it is not surprising that genes, such as CYP19 involved in their synthesis and function are regarded as candidates for the markers of reproductive traits.

Table 1: Forward and reverse primers sequence for *CYP19* locus, annealing temperatures (Ta), size of PCR amplicon (bp) and its localization in the gene.

locus	Primers			Size	Localiza
	Forward (5-3)	Reverse(5-3)	(°C)	(bp)	tion
CYP19	GCAGCAAGGACTTGAAAG	GAACCTGGTGGGCTAC	61	241	I6, E7,
	ATG	AGTC			I7
I = Intron	ntron $E = Exon$				

1	2	3	4	5	6	М
						11
						300bp
						200bp

Fig.1. Ethidium bromide stained 2% agarose gel of unpurified PCR products of *CYP 19* locus (with an average size of 241 bp) before excision from gel (lanes 1-6). M represents 100 bp ladder.

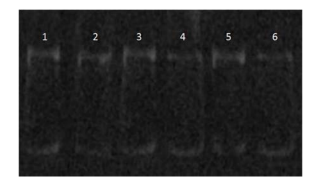


Fig.2. PCR-SSCP patterns of *CYP 19* locus in Egyptian buffalo show one SSCP pattern in this locus.

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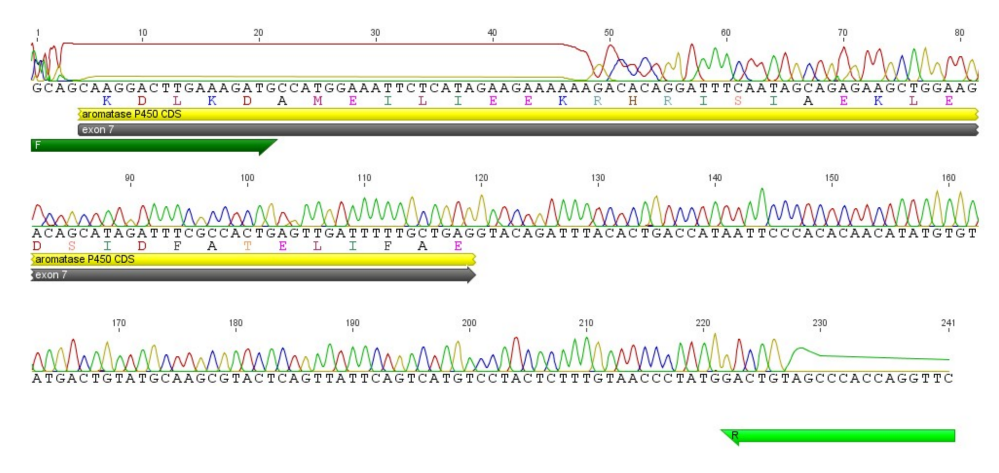


Fig.3. Nucleotide sequences of CYP19 locus and the corresponding amino acid sequences were shown below nucleotide sequences of exon 7.

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Bubalus bubalis aromatase P450 gene, exon 7 and partial cds Sequence ID: <u>gb|EF178281.1|</u> Length: 241 Number of Matches: 1

Descendent to 244 Company, Complian

Range	Range 1: 1 to 241 GenBank Graphics VNext Match A Previc							
Score		Expect	Identities	Gaps	Strand			
446 bits(241)		1) 5e-122	241/241(100%)	0/241(0%)	Plus/Plus			
Query	1	GCAGCAAGGACTTGAAA	GATGCCATGGAAATTCTCATAG	AAGAAAAAAGACAC	AGGATTT 60			
Sbjct	1	GCAGCAAGGACTTGAAA	GATGCCATGGAAATTCTCATAG	AAGAAAAAAGACAC	AGGATTT 60			
Query	61	CAATAGCAGAGAAGCTG	GAAGACAGCATAGATTTCGCCA		GCTGAGG 120			
Sbjct	61	CAATAGCAGAGAAGCTG	GAAGACAGCATAGATTTCGCCA					
Query	121	TACAGATTTACACTGAC	CATAATTCCCACACAACATATG	TGTATGACTGTATG	CAAGCGT 180			
Sbjct	121	TACAGATTTACACTGAC	CATAATTCCCACACAACATATG	TGTATGACTGTATG				
Query	181	ACTCAGTTATTCAGTCA:	IGTCCTACTCTTTGTAACCCTA	TGGACTGTAGCCCA	CCAGGTT 240			
Sbjct	181	ACTCAGTTATTCAGTCA	IGTCCTACTCTTTGTAACCCTA	TGGACTGTAGCCCA	CAGGTT 240			
Query	241	C 241						
Sbjct	241	Ċ 241						

Fig.4. Nucleotide sequences alignment of CYP19 locus with Indian water buffaloes (EF178281) using BLAST showed 100% identity.

Bos taurus partial cyp19 gene, exon 7

Sequence ID: emb[Z69247.1] Length: 569 Number of Matches: 1

nunge .		ULIU DEN	Bank Graphi		THEAT	: Match 🔺	E-155 Y 11
Score 412 bits(223)		Expect Identities		Gaps	Strand	Strand	
		6e-112	235/241(98%)	0/241(0%)	Plus/F	Plus/Plus	
Query	1	GCAGCAA	GGACTTGAAA	GATGCCATGGAAATTCTC	ATAGAAGAAAAAAGACA	CAGGATTT	60
Sbjct	56	GCAGCAA	GGACTTGAAA	GATGCCATGGAAATTCTC	ATAGAAGAAAAAAGACA	CAGGATTT	115
Query	61	CAATAGO	AGAGAAGCTG	GAAGACAGCATAGATTTC	GCCACTGAGTTGATTTT	GCTGAGG	120
Sbjct	116	CAACAGO	AGAGAAGCTG	GAAGACAGCATAGATTTC	GCCACTGAGTTGATTTT	GCTGAGG	175
Query	121	TACAGAT	TTACACTGAC	CATAATTCCCACACAACA	TATGTGTATGACTGTAT	CAAGCGT	180
Sbjct	176		CICCACIGAC	CATAATTCCCACACAACA	TATGTGTATGACTGTAT	TDDDADDE	235
Query	181	ACTCAGT	TATTCAGTCA	IGTCCTACTCTTTGTAAC	CCTATGGACTGTAGCCC	ACCAGGTT	240
Sbjct	236	ACGCAGT	TGTTCAGTCA	IGTCCTACTCTTTGTAAC	CCTATGGACTGTAGCCC	ACCAGGTT	295
Query	241	C 241					
Sbjct	296	C 296					

Fig.5. Nucleotide sequences alignment of CYP19.7 locus with Bos Taurus (Z69247.1) using BLAST showed 98% identity.

Numerous studies have investigated the between CYP19 association and reproduction in cow; however only scanty studies were performed on such gene in Therefore, this study buffalo. was conducted to isolate a locus of this gene and to study their association with reproductive disorders in Egyptian buffalo.

Detection of single nucleotide polymorphisms (SNPs) can be achieved simply by many techniques including; single strand conformation polymorphism (SSCP) (Orita et al., 1989), RFLP (Cotton et al., 1988; Cotton, 1993; Ganguly and Prockop, 1990; Novack et al., 1986; Youil et al., 1995) and denaturing gradient gel electrophoresis (Myers et al., 1987; Sheffield et al., 1989). Among these techniques, SSCP is very commonly used, cheaper method. For instance a variation C286T in exon7 of the CYP19 gene was observed by SSCP and sequencing in humans (Means et al., 1989; Watanabe et al., 1997). In addition, the search for SSCP polymorphisms in candidate genes associated with genetic variation in traits of economic importance, could lead to the finding of genetic markers useful for improved selection of agricultural populations. Anestrum was the most common single cause of infertility in buffaloes (Ashturkar et al., 1995; Singh and Sahni, 1995), which occurs mainly due to low level of ovarian estrogens (Hafez and Hafez, 2000). The key enzyme in estrogen biosynthesis is cytochrome P450 aromatase, the protein product of CYP19 gene. Therefore, CYP19 gene can play a role in the determination of anestrous animal. Kumar et al. (2009) found a G72C SNP in intron7 of CYP19 in 10% of true anestrous animals (GC) as compared to control animals (GG). This SNP was in the intronic region (noncoding region), and so it is synonymous mutation and does not lead to change in amino acid sequence. In contrast, we did not find such SNP or any other SNP in this locus in all examined Egyptian buffaloes

(normal cyclic or anestrous). This means that this SNP is not conserved in Bubalus bubalis. In cattle CYP19 gene, several SNPs were found, which were located mainly in the promoter regions. The polymorphic site in the promoter (P) 1.1 region of the bovine CYP19 gene was located at 1044 nt (Jedrzejczak et al., 2011; Komisarek and Dorynek, 2002; Vanselow et al., 1999). The P1.1 region also contains 3 SNPs detectable by PCR-RFLP method using restriction enzymes, Cfr13I, namely PvuII, and BseNI (Kowalewska-łuczak, 2010), whilst the P1.2 region contains 2 SNPs detectable by enzymes, namely BseNI and TaiI (Vanselow et al., 1999; Vanselow et al., 2000). Alignment of nucleotide sequences of CYP19 locus in Egyptian buffaloes showed 100% identity with Indian water buffaloes (EF178281) and 98% identity with Bos Taurus (Z69247). This result indicates that the sequence of CYP19 is highly conserved between cattle and buffalo. This is a preliminary study that provides the researchers with raw data, which could be used as a basis for further studies to associate other SNPs in CYP19 with fertility in buffalo.

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التعرف على الطفرات في الإكسون السابع لجين CYP19 وعلاقته بالخصوبة في الجاموس المصري

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الملخص العربى

يعد جين CYP19 واحد من أعضاء عائلة سيتوكروم ب 450 التي تقوم بتنظيم تخليق الأستروجين في الحيوانات لذلك فإنها يمكن أن تستخدم كجينات مساعدة لخطط الانتخاب للخصوبة في الجاموس. لقد تم نشر التتابع النيوكليوتيدي لهذا الجين في الجاموس حديثا ولكن لم يتم عمل در اسات عليه حتى الأن. لذلك كان الهدف من هذه الدر اسة تحدد الطفرات الموجودة في جزء من جين الCYP19 والمشتمل على الإكسون السابع والتعرف علي مدي الارتباط بين هذه الطفرات والخصوبة في الجاموس. تم عزل جزء من جين الCYP19 يصل حجمه إلى 241 زوج من القواعد النيتر وجينية يحتوي على الإكسون السابع باستخدام التفاعل البلمري المتسلسل وطريقة SSCP متبوعا بتحليل التسلسل الجيني للتعرف على الطرز المختلفة. ولكن لم نلاحظ أي طفرات في كل الحيوانات التي تم فحصها ولقد تحقق ذلك بظهور طرز أحادي الشكل وتتابع نيوكليوتيدي واحد في الحيوانات التي تم فحصها ولقد تحقق ذلك بظهور طرز أحادي الشكل وتتابع نيوكليوتيدي واحد في الحيوانات المري لم يتم فيها الشياع. وبالتالي أو ضحت النتائج أن هذا الموقع من الطرز المختلفة. ولكن لم نلاحظ أي طفرات في كل الحيوانات التي تم فحصها ولقد تحقق ذلك بظهور طرز أحادي الشكل وتتابع نيوكليوتيدي واحد في الحيوانات المربيعية والتي لم يتم فيها الشياع. وبالتالي أو ضحت النتائج أن هذا الموقع من الطرز المختلفة. ولكن لم الحصوبة في الجاموس المصري لذلك لا يمكن أن يستخدم في البرنامج الانتخابي لتحسين الجين ليس له تأثير على الخصوبة في الجاموس المصري لذلك لا يمكن أن يستخدم في البرنامج الانتخابي لتحسين الخصوبة في الجاموس المصري.

(مجلة بنها للعلوم الطبية البيطرية: عدد 26(1):151-160, مارس 2014)