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Molecular characterization, genetic variability and detection of a functional polymorphism influencing the promoter activity of OXT gene in goat and sheep

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Summary

The purpose of the study reported in this Research Communication was to report the full characterization of the goat and sheep oxytocin-neurophysin I gene (OXT), their promoters and amino acid sequences. Using the genomic DNA as template, we sequenced and compared the whole OXT gene (3 exons), plus 958/960 nucleotides at the 5’ flanking region and 478/477 nucleotides at the 3’ flanking region, in 46 sheep and 24 goats belonging to different breeds/genetic types reared in Italy, Greece and Germany. The comparison of the obtained sequences showed a high degree of genetic variability at these loci. In particular, we focused on the SNP g.438T>C as possible example of trans-specific polymorphism. This SNP alters a putative binding site of the transcription factor
Oct-1 and the set-up of a luciferase assay confirmed that the C variant of this SNP negatively affects the promoter activity of the sheep $OXT$ gene.

The results of this study suggest that the SNP g.438T>C might be useful to promote association studies with traits/physiological processes controlled by this hormone.

**Keywords:** *Capra hircus, Ovis aries, OXT locus, gene polymorphism, luciferase assay*

**Introduction**

The oxytocin is a peptide hormone released into general circulation from the posterior pituitary gland. It is involved in the movement of smooth muscles, inducing uterine contractions during the calving and milk ejection during the lactation (Gimpl & Fahrenholz, 2001). In particular, the oxytocin injected before milking provides a significant increase in milk production, due to an increase in milk flow rate and a decrease in the machine milking time (Lollivier et al. 2002). However, other roles characterize the oxytocin: it controls the oestrous cycle length, the follicle luteinization and the ovarian steroidogenesis. The oxytocin also acts as a neurotransmitter in the central nervous system and plays a role in various processes like cognition, tolerance, adaptation, and complex sexual and maternal behavior (Gimpl & Fahrenholz, 2001).

To date, the oxytocin encoding sequences ($OXT$) have been fully reported for many vertebrates, however incomplete information exists for some ruminants. In particular, no nucleotide or amino acid information is available in goats.

Despite considerable research efforts and interest among the scientific community on the oxytocin gene, so far, a few studies on polymorphism detection, gene expression, and association with specific traits have been carried out. For instance, in ruminants, Cosenza et al. (2007) identified polymorphisms in the river buffalo $OXT$ gene and Pauciullo et al. (2012a) found a significant association of these SNPs with buffalo milk yield and flow.
In the present study, we report the complete sequences of goat and sheep OXT gene, the identification and characterization of a high degree of genetic variability at these loci and we assess their possible effects on the OXT promoter activity.

**Materials and Methods**

**DNA samples**

According to the method of Goossens and Kan (1981), genomic DNA was extracted from individual blood leukocytes of 24 Italian sheep (nine undefined genetic type, 15 Laticauda), 22 German sheep (10 Merino Landschaf, three Rhönshaf, three Texel, three Ostfriesisches Milchschaf and three Grau Gehörnte Heidschnucke), 14 German goats (four Thuringer Waldziege, four Weiße Deutsche, four Bunte Deutsche Edelziege and two Boer goat cross), two Greek and eight Italian goats of undefined genetic type, randomly chosen.

**Primer design, PCR conditions and sequencing**

All primers for amplification and sequencing of sheep and goat OXT were designed by DNAsis-Pro software (Hitachi), using the bovine (EMBL acc. no. AB481096) and ovine (EMBL acc. no. X55131; CP011898) OXT gene sequences as templates (online Supplementary Table S1).

PCR reaction mixture and thermal conditions were accomplished according to Cosenza et al. (2007). PCR products were sequenced on both strands at CEINGE - Biotecnologie Avanzate (Naples, Italy).

**Bioinformatic and statistical analysis**

The BLAST analysis (www.ncbi.nlm.nih.gov/BLAST) was used to confirm the sequencing results in regards to OXT gene. Allelic frequencies and Hardy-Weinberg equilibrium (χ² test) were calculated. Homology searches, comparison among nucleotide and deduced amino acid sequences
and multiple alignments were accomplished using DNAsis-Pro (Hitachi Software Engineering Co.,
Japan).

Construction of reporter plasmids, transient transfections and luciferase assays

To evaluate the role of the SNP g.438T>C on the sheep OXT expression, a 941 bp DNA
fragment including the OXT promoter region was amplified from homozygous T/T and C/C
samples. Furthermore, a nested PCR with two modified internal primers was carried out to provide
restriction sites for the KpnI (GGTAC!C) and SmaI (CCC!GGG) endonucleases. Primers sequences
are reported in online Supplementary Table S1.

The nested PCR products were digested with KpnI and SmaI endonucleases and cloned
upstream a reporter in the pGL3 basic vector (Promega) as previously reported (Cosenza et al.
2016). Later, the new products were sequenced using vector primers RVprimer3 and RVprimer4 (as
recommended in the supporting information) to confirm the orientation and integrity of the inserts.
HEK293 cell were then transiently transfected with 200ng of the new plasmids and with 20ng of
pRL-TK vector expressing Renilla luciferase (Promega) using Fugene HD transfection reagent
(Promega). As control, a transfection with pGL3 control vector was also performed. All
transfections were carried out in triplicate and in three independent experiments. After 24 h, the
reporter activity of g.438T and g.438C was measured by luminometry on the Victor luminometer
(PerkinElmer) using the Dual-Glo luciferase assay system (Promega). In all transfection, Firefly
luciferase values were normalized with the Renilla luciferase expressed by the vector pRL-TK co-
transfected as internal control. Promoter activity for each constructs was calculated as ratio of
normalized reporter activity to pGL3 control vector.

Results and Discussion

Structure of goat and sheep oxytocin-neurophysin I gene
We sequenced the genomic DNA of the whole oxytocin-neurophysin I encoding gene (OXT) plus 958/960 nucleotides at the 5’ flanking region and 478/477 nucleotides at the 3’ flanking region respectively in 24 goat (EMBL accession no. LT592266) and 46 sheep (EMBL accession no. LT592265) randomly chosen.

The goat OXT gene was found to extend over 907 bp, including 510 bp of exonic regions and 397 bp of intronic regions. The gene contains only 3 exons, ranging in size from 153 bp (exon 1) to 202 bp (exon 2), and 2 introns of 307 bp (intron 1) and 90 bp (intron 2). The 5’UT (untranslated) region comprises the first 33 nucleotides of the first exon, while the 3’UT region includes the last 102 nucleotides of the 3rd exon.

The ORF (Open Reading Frame) region codes for the peptide leader and for 106 amino acids of the “oxytocin-neurophysin I” complex. The first exon encodes a translocator signal (19 amino acid residues encoded by the nucleotides 34-90), the nonapeptide hormone (from nucleotide 91 to 117), the tripeptide processing signal (GKR) (from nucleotide 118 to 126), and the first 9 out of the 94 residues of the neurophysin I; the second exon encodes the central part of neurophysin I and the third exon (155 bp) encodes the COOH-terminal region of neurophysin I.

The nonapeptide is characterized by a disulphide bridge between the Cys residues 1 and 6. This results in a cyclic part made of six-amino acid and a tail of three-residue α-amidated in the COOH-terminal. The isoleucine in position 3 is essential for stimulating OXT receptors, whereas the Cys-1 and Tyr-2 in the OXT molecule are the principal neurophysin binding residues (Gimpl & Fahrenholz, 2001).

The methionine initiation codon is indicated at nucleotides +34/+36 and the translation stop codon TGA is located at nt 54-56 of the exon 3. All splice junctions follow the 5’ GT/ 3’AG splice rule. Moreover, the polyadenylation signal (AAATAAA) and polyadenylation site were identified in the 3’UT region (3rd exon), located between the nucleotides +76/+78 and +86/92 from the terminator codon, respectively. This structure shows a homology of 98% with the sheep OXT sequence (EMBL LT592265 present work) (online Supplementary Figure S1).
Analysis of the intronic, exonic and 3’ flanking regions

The comparison of the intronic sequences for the 46 sheep under investigation showed only two and three perfect repeats of the monomer 5’-TCCC-3’ downstream the nucleotide 253 of the exon 1. Conversely, the goat OXT showed higher genetic variation being characterised by 6 polymorphic sites located at the intron 1: one transition (g.1151T>C), 4 transversions (g.1194T>A, g.1221C>A, g.1333C>A and g.1381C>G) and one deletion (g.1289_1300delC) (online Supplementary Figure S1). None of the detected mutations affects canonic splicing sites, therefore no influence on the OXT gene expression is expected for these polymorphisms.

The comparison of the exonic sequences of the investigated sheep showed 3 synonymous substitutions. The first was located at the position 186 of the exon 2 (g.1616C>G) and the other two at the position 62 and 147 of the exon 3, respectively (g.1784G>C and g.1879T>A). The last two mutations are located in the 3’ UTR (online Supplementary Figure S1). Since the presence of the cytosine in position 186 of the exon 2 characterizes also other published OXT sequences of sheep (EMBL nos. X55131.1, NM_001009801.2, X16052.1), as well as the sequences of other species, its presence might be indicative of an ancestral condition. For this reason, the sheep OXT allele containing the cytosine was named OXT A, whereas the changed form was named OXT B.

The mutation g.1879T>A is very interesting because it took place inside the polyadenylation site (TTTTTTT). Therefore, it is reasonable to hypothesize that the g.1879T>A transversion might, directly or indirectly, influence the stability of the mRNA and, consequently, the amount of oxytocin produced.

Regarding our investigation in goats, only one polymorphic site was detected at exon level. However, it is a transition g.1814C>T located at the 3’ UTR. The BLAST alignment with the other sequences available in EMBL showed that the cytosine might be the ancestral condition. Therefore, the goat OXT allele containing the cytosine was named OXT A, whereas the changed form was named OXT B.
The analysis of the 3’ flanking region in both the species showed one element 207 bp long (from 2087 to 2293 and from 2086 to 2291 in sheep and goat, respectively, online Supplementary Figure S1) with a similarity of about 80% with the annotated Bov-tA3 sequences (EMBL no. AC150888). The 3’ flanking region also showed genetic variability, in terms of duplication and SNPs (g.1930_1934dupGGCTC and g.2054C>T in goats; g.2052G>C and g.2123G>T in sheep). It is known that the 3’ end is an important region for the control of the gene expression. Therefore, any polymorphism falling in this region may putatively affect binding sites, for example seed sequences for microRNAs (miRNA), which are known to play a fundamental role in the regulation of gene expression in many physiological pathways including the lactation.

**OXT promoter analysis**

Compared to the buffalo (EMBL nos. AM234538; AM234539) and similarly to bovine, the goat and sheep OXT gene promoter sequences (958 and 960 nucleotides, respectively) are characterized by a deletion of 26 bp which occurs between the nucleotides –892/893 (numbering is relative to the first nucleotide of the sheep first exon, +1). Most probably, this event was originated by a 16-nt insertion (TTTCTTTTTTTTTTA) with a 10-nt duplication of an adjacent sequence (AATTATTTTC). Moreover, both the goat and sheep sequences are characterized, unlike that of buffalo and bovine, by the insertion and deletion of two decamers, between nt –506/–495 and –305/–304, respectively (online Supplementary Figure S2).

Apart from the above differences, the organization of the goat and sheep OXT promoter region is very similar to the homologous gene regions in the other two species, including a modified RNA polymerase binding TATA box (TTAAA) located between the nucleotides –24/–29.

The pentamers TGACC and TCACC, known to constitute one half of the estrogen- and the thyroid hormone-responsive element (Beato, 1989), are present in the proximal flanking region of sheep and goat OXT gene (indicated by bold letters in online Supplementary Figure S2).
A highly conserved DNA element exists at about 160 nucleotides upstream of the transcriptional initiation site, a “composite hormone response element” (monomeric orphan receptor binding site, CATAACCTTGACC), conserved across species (Gimpl & Fahrenholz, 2001), including the binding sequence sites of the estrogen response elements (ERE), the COUP-TF sites and the steroidogenic factor-1 (SF-1) (Burbach et al. 2001).

A putative activator protein-1 (AP-1) site (GTGACGAA), with a single difference from the one indicated by Ivell et al. (1990) (GTGACTAA) and from the consensus sequences reported by Spandidos et al. (1988), is indicated by a box at nucleotides –618/–627. Other potential binding sites for AP-1, a CCAAT/enhancer binding protein-alpha (C/EBP alpha) and a Transcription Factor Oct-1 in the OXT promoter sequence (identified by TRANSFAC® 7.0 database) are also boxed in online Supplementary Figure S2.

The comparative analysis of the goat promoter sequences showed 8 polymorphic sites, 7 transitions (g.154G>A, g.156T>C, g.438T>C, g.524A>G, g.579G>A, g.705C>T, g.827G>A) and 1 transversions (g.937G>T) which falls only 2 nucleotides far (-22) from the TATA box.

Similarly, 6 SNPs were identified in the promoter region of sheep OXT gene, 3 transversions (g.171T>A, g.698C>A, g.959G>T) and 3 transitions (g.43A>G, g.438T>C, g.919G>A) in addition to a variability of a mono- (g.97A[6]) and di-nucleotide repeats (g.653AG[2]). Two of these mutations could be responsible for alteration of gene expression. In fact, the first (g.438T>C) would alter the putative binding site of the transcription factor Oct-1, whereas the second mutation (g.959G>T) would involve the CpG- potential first splice donor site.

It is very interesting to notice that the comparison of sequences for the two examined species showed that both are characterized by the same polymorphism, the transition g.438T>C. This mutation could have been realized following a convergent evolution (allelic lineages evolve similar features independently in separate lineages), or a genetic introgression (allelic lineages are horizontally transferred either from recipient species to donor species or in both directions).
However, another possibility is that this mutation represents a Trans-Specific Polymorphism (TSP), which refers to the occurrence of identical or similar alleles in related species.

TSP is a crucial evolutionary mechanism responsible for sharing adaptive genetic variation across taxa. Although most studies dealing with TSP have concentrated only on the MHC loci, a few examples described in other immune genes suggest that TSP is a common and general evolutionary phenomenon. In fact, it has been documented in ABO blood system, retroviral restriction factor (TRIM5 encoding gene), zinc-finger antiviral protein (ZC3HAV1 gene) and Ladinin-1 encoding gene (Teixeira et al. 2015) in primates, or in complementary sex determiner gene in Hymenoptera (Lechner et al. 2014).

Therefore, the identification of this polymorphism could represent the first report of a DNA-level trans-specific polymorphism in goat and sheep and also the first in a hormone coding gene.

Genotype distribution of goat and sheep for the SNP g.438T>C is reported in Table 1.

Sheep OXT gene expression analysis

To test in vitro the effects of the SNP g.438T>C on the sheep OXT promoter activity, a 941bp region from TT and CC homozygous subjects was amplified and cloned upstream a reporter gene, generating two different constructs. These constructs were used to transiently transfect HEK293 cell and the luciferase activity was measured. As shown in Figure 1, the luciferase activity was lower when the cells were transfected with CC, indicating that the nucleotide C in position 438 negatively affects the promoter activity of the sheep OXT gene.

This polymorphic site falls into a potential binding site for the transcription factor Oct-1. In particular, the presence of cytosine removes the consensus site for this transcription factor (in silico analysis performed by Transfact 7.0 database). The protein Oct-1 is a multifunctional molecule and it belongs to transcription factors of the POU family. Presumably, it participates in the hormonal regulation of various genes, mediating the effect of hormones on gene activation or suppression (Sytina & Pankratova, 2003). In addition, mutations in the consensus sequences of the ubiquitous
Oct-1 transcription factor are reported to reduce hormonal induction in different gene promoters, such as in the mouse β casein encoding gene (CSN2) promoter (Dong & Zhao 2007). Therefore, based on these considerations and keeping in mind the result of the in vitro reporter assay, it is possible to suppose an effective role of this promoter-binding site in the expression of the oxytocin gene.

The knowledge of the transcriptional and translational effects of OXT gene polymorphisms will provide new opportunities to select the best dairy goats and sheep for all the physiological processes controlled by this hormone, such as, for example, uterine smooth muscle contraction during labour, oestrous cycle length, follicle luteinisation and ovarian steroidogenesis, cognition, tolerance, adaptation and complex sexual and maternal behaviour. The detected polymorphisms represent a good opportunity to carry out studies focused on the identification of significant association with milk yield and flow in ovine and caprine species, as already performed in other species for OXT (Pauciullo et al. 2012a), or other genes candidate for quali-quantitative variations of these traits (Pauciullo et al. 2012b).

Acknowledgements

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Conflict of interest

None of the authors has any conflict of interest to declare.

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Table 1. Genotyping data, allele frequency, relative frequencies of the SNP g.438T>C at the promoter region of the OXT gene in sheep and goat

<table>
<thead>
<tr>
<th>Position</th>
<th>Sheep genotype distribution</th>
<th>Allele frequency</th>
<th>Goat genotype distribution</th>
<th>Allele frequency</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>CC</td>
<td>CT</td>
<td>TT</td>
<td>TOT</td>
</tr>
<tr>
<td>g.438T&gt;C</td>
<td>Obs.</td>
<td>1</td>
<td>7</td>
<td>38</td>
</tr>
<tr>
<td></td>
<td>Exp.</td>
<td>0.4402</td>
<td>8.1196</td>
<td>37.4402</td>
</tr>
</tbody>
</table>
Figure 1. The g.438T>C SNP influences the activity of the promoter OXT in sheep. Luciferase activity measured in HEK293 cell transfected with pGL control, g.438T and g.438C. pRL-TK was co-transfected and used for normalizing luciferase activity. Data are shown as mean ± SEM (the mean is presented as the longest horizontal line, whereas the two smaller lines indicate the limit of SEM) and are representative of three independent experiments (circle, square and triangle represent the three replicates of the different conditions). Data elaboration was performed using GRAPHPAD PRISM6 software. *P < 0.05, Student’s t-test.