Molecular characterisation of the caprine (Capra hircus) lymphocyte function-associated antigen-1 alpha subunit-encoding cDNA

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Abstract

Background: Lymphocyte function-associated antigen-1 (LFA-1, CD11a/CD18, alpha L beta 2) is required for many cellular adhesive interactions during the immune response.

Results: The Capra hircus CD11a-encoding cDNA was sequenced and compared with its human, murine, rat, bovine and ovine counterparts. Despite some focal differences, it shares all the main characteristics of its known mammalian homologues.

Conclusion: Therefore, along with the caprine CD18-encoding cDNA, which has been available for a few months, the sequence data revealed here will allow the Capra hircus LFA-1 expression in vitro as a tool to explore the specificities of inflammation in the caprine species.

Background

Lymphocyte function-associated antigen-1 (LFA-1, αLβ2, CD11a/CD18) is a member of the β2-integrin subfamily of cell surface receptors. Integrins consist of a 120 to 180 kDa α subunit (CD11a in this case) and a 90 to 110 kDa β subunit that are noncovalently associated single-pass transmembrane proteins [1]. The bulk of each integrin subunit is extracellular, where it typically functions as a receptor for extracellular matrix molecules or as a counter-receptor for surface proteins of apposed cells [2]. The heterodimer CD11a/CD18 is expressed on all leukocytes and mediates high affinity adhesion to a variety of cell types that express one or more of the β2-integrins ligands, intercellular adhesion molecules (ICAM-1 to -5) [3-5]. The adhesion process mediated is a critical step of a wide range of immunological activities, including cytosis of target cells, cross-interaction and cross-stimulation between lymphocytes, phagocytosis of complement-coated targets, neutrophils clearance from inflamed tissue, and the regulation of leukocyte traffic between the bloodstream and tissues [6-9]. As the relevance of the goat model for studying leukocyte traffic, diapedesis and pathologic tissue infiltration is well established in such important areas as mastitis [10-13] or lentivirus-associated diseases [14-16], increasing our knowledge about caprine β2 integrins is of great importance to offer new possibilities for research and to provide additional insights into those fields. Along with the caprine CD18-encoding cDNA, which is available for a few months [17], the sequence data provided here will allow the Capra hircus β2-integrin CD11a/CD18 expression in vitro as a tool to examine the specificities of inflammation in the caprine species.
Methods

RNA isolation
Total RNA from phorbol myristate acetate (PMA)-stimulated (25 ng/ml for 15 min) caprine (Boer breed) peripheral blood mononuclear cells (PBMC) was extracted with TRIzol (Invitrogen, USA) as described by the manufacturer. The PBMCs were obtained by density gradient centrifugation with Ficoll-Paque Plus (Amersham, USA) and maintained in RPMI 1640 supplemented with 10% foetal bovine serum (Gibco BRL, USA), penicillin (100 U/ml) and streptomycin (100 μg/ml) at 37°C in a 5% CO₂ atmosphere.

Amplification of cDNA ends
We used SMART RACE technology (Clontech Laboratories Inc., USA) to obtain caprine CD11a (CaCD11a) 5'- and 3'-ends and RT-PCR to amplify full-length CaCD11a CDS. For first-strand cDNA synthesis, and according to the sequence of bovine CD11a available [GenBank: AY267467], gene-specific primers were designed which were expected to give non overlapping ~1 kb RACE products: a sense primer for the 3'-RACE PCR : 5'-TCCGATGTCGCCAGTAAATC-3' (corresponding to nt 2572 to nt 2595) and an antisense primer for the 5'-RACE PCR : 5'-CCGGCCTCCTCTCTGCTCCCCATAG-3' (nt 1470 to nt 1446). Reverse transcription and polymerase chain reactions (PCR) were carried out according to the instruction manual of the SMART RACE cDNA Amplification Kit. The 5'- and 3'-RACE products were gel-purified using the S.N.A.P.™ Gel Purification Kit (Invitrogen, USA), TA-cloned into pCRII-TOPO (Invitrogen, USA) and seeded on kanamycin IPTG plates. Miniprep were obtained from colonies grown in 5 ml LB-kanamycin broth and the clones were sequenced on the ABI-3730 Genetic Analyzer using the Big Dye terminator chemistry (Applied Biosystems, USA).

Molecular cloning of full-length cDNA
Total RNA from PMA-stimulated PBMCs was reverse transcribed using Improm II (Promega, USA). The full-length cDNA was then generated by long distance PCR using Platinum Taq DNA polymerase High Fidelity (Invitrogen, USA) with primers designed from the distal ends of both 5'- and 3'-RACE products : 5'-GTCGCCAGTAAATCCCAAGA-3' (sense, within the 5'-UTR) and 5'-GACGCATCAATCTCCACCAC-3' (antisense, 3'UTR). The procedures recommended by the manufacturer were followed, with these cycling parameters: 5 min at 94°C, then 35 cycles including (i) 30 s at 94°C, (ii) 30 s at 58°C and (iii) 3 min 30 s at 68°C, followed by a final extension at 68°C for 5 min. Resulting PCR products were then processed for sequencing as aforementioned for the RACE products. The CD11a cDNA sequence was deduced from sequences obtained from nine independent clones. Sequence data have been deposited at GenBank under accession No. AV773018 and AV773019.

Bioinformatics
Primers design was performed with Netprimer [18] and Primer 3 [19]. Nucleotidic sequence and similarity analyses were carried out using respectively Chromas v.2.21 [20] and BLAST programs [21]. Alignment of amino acids sequences were drawn by GeneDoc v.2.6.002 [22] following the BLOSUM62 matrix. SignalP v.2.0.b2 [23] and NetNGlyc v.1.0 [24] provided peptide signal and N-glycosylation sites prediction, respectively.

Results & discussion
Characterisation of CaCD11a-encoding cDNA and deduced aa sequence
Two alleles have been identified for the CaCD11a cDNA. The sequence contains ~4200 bp with an ORF of 3498 [Genbank: AV773019] or 3495 bp [Genbank: AV773018] depending on the allele that codes for 1165 or 1164 aa followed by ~600 bp in the 3'-UTR (Fig. 1). The mature CaCD11a contains a 23-aa putative leader peptide, an extracellular domain of 1061 or 1062 residues (24-1084/1085), a single hydrophobic transmembrane region of 24 residues (1085/1086–1108/1109) and a cytoplasmic tail of 57 residues (Fig. 1). Nine N-linked putative glycosylation sites (Asn-X-Thr/Ser) are present in the extracellular domain (Fig. 2). The mature protein contains 19 cysteine residues among which one is located into the cytoplasmic tail (Fig. 2). The extracellular domain also contains an inserted (I) domain of 172 amino acids (residues 153–324) quite similar to those found in all the leukocyte integrin α subunits sequenced to date and located between the β sheets 2 and 3 of a seven bladed β-propeller region [25]. The I-domain is homologous with repeated domains found in von Willebrand factor and cartilage matrix protein [1] and can be expressed as an isolated domain. Its three-dimensional structure consists of a five-stranded parallel β-sheet core surrounded on both faces by α-helices, with a short antiparallel strand occurring on one edge of this sheet [26]. The I-domain contains a metal ion-dependent adhesion site (IDAS) that plays a functional role in ICAM-1 binding domain (Fig. 2). The I-domain also contains an inserted (I) domain of 172 amino acids (residues 153–324) quite similar to those found in all the leukocyte integrin α subunits sequenced to date and located between the β sheets 2 and 3 of a seven bladed β-propeller region [25]. The I-domain is homologous with repeated domains found in von Willebrand factor and cartilage matrix protein [1] and can be expressed as an isolated domain. Its three-dimensional structure consists of a five-stranded parallel β-sheet core surrounded on both faces by α-helices, with a short antiparallel strand occurring on one edge of this sheet [26]. The I-domain contains a metal ion-dependent adhesion site (IDAS) that plays a functional role in ICAM-1 binding (Fig. 2). Three repeats with a divalent cation binding motif are found at amino acid residues 465–473, 527–535 and 587–595 (Fig. 2). All the conserved cysteines and all but one N-glycosylation sites are found outside the I region and divalent cation binding motifs (Fig. 2), consistent with the hypothesis that these regions may undergo conformational changes important in ligand binding [31,32].
suggested by the requirement of Mg²⁺ and Ca²⁺ for binding motifs and the transmembrane region and the respective, with the highest identity for the MIDAS, the cation overall 94%, 98%, 77%, 68% and 55% identity, respectively, with the bovine, ovine, human, murine and rat counterparts shows NW_047562 Capra hircus [34], human [32], murine [35] and rat GenBank: NW_047562 deposited at GenBank under accession No. AY773018 (shown here), respectively without and with Gln-743 (#).

**Figure 1**
The nucleotide and deduced amino acid sequences of *Capra hircus* CD11a cDNA. The putative leader peptide and transmembrane region are underlined. Nine independent clones were sequenced in both directions. Sequence data have been deposited at GenBank under accession No. AY773018 and AY773019 (shown here), respectively without and with Gln-743 (#).

**General comparison among species**
Overall, the general organization of caprine, bovine [33], ovine [34], human [32], murine [35] and rat GenBank: NW_047562 CD11a proteins is quite similar (Fig. 2). Comparison between mature CaCD11a sequence and its bovine, ovine, human, murine and rat counterparts shows overall 94%, 98%, 77%, 68% and 55% identity, respectively, with the highest identity for the MIDAS, the cation binding motifs and the transmembrane region and the lowest identity for the cytoplasmic tail (Table 1). The high conservation of the MIDAS and the putative cation binding motifs is consistent with an involvement of these regions in the functional activity of LFA-1 α subunit, as suggested by the requirement of Mg²⁺ and Ca²⁺ for CD11a/CD18-dependent cellular interactions [31] or binding to purified ICAM-1 [36,37]. The transmembrane region shows also a high degree of conservation that could be explained by (i) physicochemical, and (ii) functional constraints. Indeed, (i) residues lying in the membrane have to possess rather hydrophobic character to allow liposolubility, which is confirmed by the presence of many leucine residues (figure 2) and (ii) bi-directional integrin signalling (inside-out and outside-in) is accomplished by transmission of information across the plasma membrane [38]. By contrast, the low conservation of the cytoplasmic tail suggests that it is not required to guarantee adequate functioning of LFA-1. This is in agreement with the observation that truncation of the LFA-1 β subunit cytoplasmic domain has no effect on binding to ICAM-1, whereas binding is markedly diminished by β subunit cytoplasmic domain truncation [39]. Residue Glu-332 that is located in the linker following the I-domain and that is known to be critical for communication to the β₁-like domain, rolling, integrin extension and activation by Mn²⁺ of firm adhesion [8] is strictly conserved.

Every cysteine residue in the caprine extracellular portion of mature CD11a is present at the same location in bovine, ovine, human and rat CD11a, which is consistent with a role in maintaining the global structure of the protein whereas two cysteine residues (positions 1009 and 1048) are absent from caprine CD11a and therefore do not seem to be indispensable. The mouse version distinguishes by an additional cysteine residue at
Interestingly, as in sheep [34] and human [GenBank: NM_002209 and AY892236], an allelic variant with a triplet insertion resulting in an additional Glu744 in the extracellular domain was consistently identified, which suggests an allelic polymorphism that might be biologically relevant. Studies of genomic sequences will permit to know if this addition represents two alleles or not.

Finally, one has to note that the lowest between-species percent identities are observed with the rat CD11a sequence which has been derived from an annotated genomic sequence. Cloning and characterisation of rat CD11a from rat PBMCs would probably give a higher identity.

**Conclusion**

This study reports for the first time the isolation and sequencing of the caprine LFA-1 α subunit (CD11a) cDNA, and demonstrates that, despite some focal differences, it shares all the main characteristics of its known mammalian homologues. Along with the caprine CD18-encoding cDNA which is now available [17], the sequence
Table 1: Between-species percent identities of CD11a constitutive blocks. Ca, Bo, Ov, Hu, Mu and Ra: caprine, bovine, ovine, human, murine and rat CD11a, respectively; MIDAS: metal-ion dependent adhesion site.

| Block                        | Ca vs. Bo (%) | Ca vs. Ov (%) | Ca vs. Hu (%) | Ca vs. Mu (%) | Ca vs. Ra (%) |
|------------------------------|---------------|---------------|---------------|---------------|---------------|
| Overall                      | 94            | 98            | 77            | 68            | 55            |
| Putative signal peptide      | 86            | 95            | 60            | 41            | 4             |
| Extracellular domain         | 94            | 98            | 78            | 69            | 60            |
| Transmembrane region         | 95            | 100           | 95            | 79            | 87            |
| Cytoplasmic tail             | 92            | 94            | 58            | 50            | 47            |
| I-domain                     | 95            | 99            | 86            | 72            | 75            |
| MIDAS                        | 100           | 100           | 100           | 85            | 85            |
| Putative cation binding motif 1 | 100           | 100           | 88            | 66            | 55            |
| Putative cation binding motif 2 | 100           | 100           | 77            | 88            | 88            |
| Putative cation binding motif 3 | 100           | 100           | 88            | 77            | 88            |

data provided here will allow the successful expression of caprine LFA-1 in vitro as a tool to examine the specificities of inflammation in the caprine species.

**Competing interests**

The author(s) declare that they have no competing interests.

**Authors’ contributions**

TF carried out cloning and sequencing, participated in the sequence alignment and to the draft of the manuscript. LZ participated in the sequence comparison and to the draft of the manuscript. EB participated in the design of the study. DD conceived the study, and participated in its design and coordination and helped to draft the manuscript. All authors read and approved the final manuscript.

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