Expression of Putative Stem Cell Markers Related to Developmental Stage of Sheep Mammary Glands

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Introduction

During the entire life cycle of the mammary gland, competent mammary stem cells are presumed to exist to provide for expansion of cell populations and cell turnover or replacement. In fact, mammary glands have high regenerative capacity during pregnancy and lactation (Holland et al., 2003). Because dairy animals undergo reproductive cycles for milk production, they are an excellent model to examine physiologic characteristics of mammary glands.

Based on their ultrastructural analyses, Chepko and Smith (1997, 1999) proposed that small, light staining cells (SLC) within the mammary parenchyma act as multipotent mammary stem cells. Other mammary epithelial cell types were classified as undifferentiated large light cells (ULLC), differentiated large light cells (DLLC), differentiated secretory cells or large dark cells (LDC) and myoepithelial cells. It was suggested that a morphologically indistinguishable subset of SLC act as primary progenitors and give rise to the ULLC. In turn, the ULLC divide multiple times to expand the parenchymal cell population and produce more LDC or myoepithelial cells.

In addition, Ellis and Capuco (2002) strongly supported the hypothesis that the SLC function as the primary proliferative cell populations also in heifers. However, morphologically, it is quite impossible to distinguish the stem cells from the terminally differentiated cells of a particular tissue.

A number of investigators have attempted to isolate molecular markers for mammary stem cells (Li et al., 1998; Welm et al., 2003), but till now, no genetic marker has yet been found to positively identify mammary stem cells in situ. Positive stem cell markers such as c-kit (the receptor for stem cell factor, KIT) and its ligand stem cell factor (KITLG), ATP-binding cassette sub-family G member 2 (ABCG2) and Musashi 1 (MSI1) are good candidates to identify progenitor cells in their niche. Using real-time PCR we showed that KIT, KITLG and MSI1 expressions were up-regulated before lambing and at involution relatively to prepubertal stage. The in situ hybridization analysis for KIT gene confirmed and localized the expression in luminal epithelial cells. The changes in the expression profile of putative stem cell markers in mammary glands of sheep suggest that they modify with the progression of lactation cycle, being up-regulated during differentiation and down regulated during lactation.

Summary

It is thought that the regenerative capacity of the mammary gland following post-lactation involution resides in multipotent stem cells within the luminal tissue. Adult stem cells make up a small percentage of the cells found in mature organ systems, however to define useful markers has long been a challenge. c-Kit (KIT) and its ligand stem cell factor (KITLG), ATP-binding cassette sub-family G member 2 (ABCG2) and Musashi 1 (MSI1) are good candidates to identify progenitor cells in their niche. Using real-time PCR we showed that KIT, KITLG and MSI1 expressions were up-regulated before lambing and at involution relatively to prepubertal stage. The in situ hybridization analysis for KIT gene confirmed and localized the expression in luminal epithelial cells. The changes in the expression profile of putative stem cell markers in mammary glands of sheep suggest that they modify with the progression of lactation cycle, being up-regulated during differentiation and down regulated during lactation.
Localization of ABCG2 (also known as BCRP1 for breast cancer resistance protein), a member of the family of ATP-binding cassette (ABC) transporters, was detected in normal tissues, such as the colon epithelium, placenta syncytiotrophoblast, small intestine epithelium, liver (bile canaliculi), mammary gland (lobules and lactiferous ducts), vein endothelium and capillaries (Maliepaard et al., 2001) and mouse renal tubules (Jonker et al., 2002). ABCG2 expression is highly induced in the apical membrane of alveolar epithelial cells during late pregnancy and especially during lactation (Jonker et al., 2005), and it is involved in the secretion of drugs into milk. ABCG2 is also associated with the side population (SP), a phenotype of stem cells displaying low Hoechst 33342 accumulation (Zhou et al., 2001; Scharenberg et al., 2002), as in human heart, ABCG2 is involved in cellular export of Hoechst 3352. Therefore, ABCG2 may now serve as a marker protein for SP cell selection (Zhou et al., 2002).

Musashi 1 (MSI1) has been studied as marker involved in mammary stem cell detection since it was found to be expressed by epithelial progenitors in intestine, gastric mucosa, mammary glands, hair follicles and in human endometrium (Potten et al., 2003; Clarke et al., 2005; Nagata et al., 2006; Sugiyama-Nakagiri et al., 2006; Götte et al., 2008). Recently we found that MSI1 is expressed in a different degree, in mammary gland of ewes from prepubertal stage to involution (Colitti and Farinacci, 2009). Wang et al. (2008) indicated that MSI1 promote the expression of a number of genes associated with cell cycle, development and cell adhesion. In fact, up-regulation of MSI1 is associated with increased activity of Nocth and Wnt signalling, the net result being the stimulation of mammary progenitor cell proliferation.

The present study comprehensively describes, by real‐time PCR, the putative stem cell markers expression profile in mammary glands of sheep at different developmental stages, relatively to prepubertal stage. In addition, the KIT mRNA detected by in situ hybridization, was also localized at each stages.

Materials and Methods

Animals

Tissue was collected from mammary glands of thirty Sardinian sheep that were slaughtered at different developmental stages: prepubertal (30 ± 5 days, group P), 10 days before lambing (group LateP), 30, 60, 150 DIM (groups 30L, 60L, 150L, respectively) and 8 days after the end of lactation (group 8IN). For each of these sampling periods, five animals were randomly selected from a flock of grazing sheep and a clinical examination was conducted in vivo to ascertain animal health and to exclude mastitis. Sardinian sheep are a breed primarily used to produce milk; in this study milk yield ranged from 1600 to 900 g/day at 150 days in milk (DIM). Ewes at 30 DIM were allowed to suckle their lamb; the other groups (60L, 150L) were mechanically milked twice daily and manually 10 min before slaughtering, therefore, just before tissue collection (Pulina et al., 2009).

Two samples of tissue were collected, one was collected in TRIzol® (Invitrogen, Milan, Italy), then frozen in liquid nitrogen and the other one fixed in 10% neutral formalin. The experiment was carried out in accordance with state and local laws and ethical regulations (DL No. 116, 128 27/01/1992).

RNA extraction and primer design

Total RNA was extracted from mammary tissue using TRizol® Plus RNA Purification System (Invitrogen), following the manufacturer’s instructions. The concentration of the extracted total RNA was quantified with a spectrophotometer (NanoDrop 1000 ThermoScientific, Wilmington, DE, USA). RNA integrity was evaluated by the observation of 18S and 28S ribosomal bands after electrophoresis on 1% agarose gel in the presence of ethidium bromide. In sample analysis, β-actin (AF035422) expression was used as an internal control, confirming the integrity of the RNA.

Primer3 Input software (Rozen and Skaletsky, 2000) was used to design the primer sequences encoding for the KIT (AM293661), KITLG (Z50743), ABCG2 (DQ886530), MSI1 (AB240581) 18SrRNA (AM711875). Primers and product lengths for each gene are listed in Table 1 according to HUGO Gene Nomenclature Committee.

Reverse transcriptase-PCR and riboprobe synthesis

RT-PCR reactions were performed using a ‘One step’ RT-PCR (Invitrogen) kit. For each reaction, total RNA (1 µg) from ovine mammary tissue was retro-transcribed (50°C for 30 min) and amplified following a PCR protocol by a MJ thermal cycler (PT-100; MJ Research, Inc., Waltham, MA, USA). PCR reactions carried out with the RNA samples, using this set of primers, did not give any amplification product, ruling out the possibility that the observed bands might be due to the presence of contaminant genomic DNA. The RT-PCR product (205 bp) for KIT was purified from agarose with Wizard® SV Gel Clean-Up System (Promega, Milan, Italy), cloned using a dual promoter TOPO TA cloning kit containing pCRII-TOPO cloning vector (Invitrogen, Groningen, The Netherlands). The insert was sequenced (Primm, Milan, Italy) to verify probe specificity and orientation.

To prepare sense and anti-sense RNA probes, the transcripts were digoxigenin-labelled by in vitro transcription

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In situ hybridization

In situ hybridization was used to localize KIT mRNAs in sheep mammary gland sections using digoxigenin (DIG)-labelled sense and anti-sense KIT probes.

The probes, generated by RT-PCR analysis, were used to hybridize the samples embedded in paraffin and cut at 5 μm. Briefly, the slides were dewaxed with xylene and rehydrated in decreasing concentrations of ethanol till dehydropyrocacbonate (DECP)-treated water. The permeabilization was performed in Tris–EDTA buffer (TE) with 10 μg/ml of RNase free proteinase K at 37°C for 30 min. Post-fixation treatment was performed in a solution of 4% paraformaldehyde at 4°C for 15 min, and after acetylation, the sections were incubated at 37°C for 1 h in prehybridization buffer containing 20% formamide (v/v), 6X SSC, 2X Denhardt’s solution, 400 μg/ml of denatured ssDNA, 5% dextran sulfate (w/v) and 0.5% Tween 20. Hybridization was performed overnight at 55°C in a humidified chamber in the same buffer containing 2 ng/ml digoxigenin-labelled sense or anti-sense probe.

Sections were washed in a shaking water bath in 2X saline-sodium citrate buffer (SSC) at 37°C, 0.5X SSC, 0.25X SSC at room temperature (RT) and then, to digest the unbound RNA probe, incubated in NTE buffer (10 mM Tris–HCl, pH 8.0, 0.5 mM NaCl and 1 mM EDTA) containing 20 μg/ml RNase A. The last washing was performed in 0.1X SSC. After incubation for 30 min with 5% blocking reagent (Roche Diagnostics) in Tris-buffered solution (TBS), the sections were incubated overnight at 4°C in the anti-digoxigenin alkaline phosphatase conjugated antibody (Roche Diagnostics) diluted 1:500 in TBS containing 1% blocking reagent. After washes, alkaline phosphatase was detected by incubation in 5-Bromo-4-Chloro, Nitro Blue Tetrazolium Chloride (NBT/BCIP) containing 0.01 mM levamisole, overnight in the dark. The colour reaction was stopped by washing in water for 5 min. Slides were then counterstained in 0.1% (w/v) of nuclear fast red, mounted using an aqueous solution and photographed with an Olympus DP10 digital camera.

The specificity was verified by substituting the labelled sense riboprobe for the anti-sense probe.

Real-time PCR quantitation

Reverse transcriptions were performed with 1 μg of extracted total RNA by using Improm-II Reverse Transcriptase (Promega) according to the manufacturer’s instructions. The final concentration of cDNA was assumed as 50 ng/μl.

For every gene, an aliquot of each cDNA samples were pooled and standard curves with serial dilution of pool were used to optimize PCR conditions and to calculate the efficiency, fluorescence baseline and threshold.

Real-time RT-PCR reactions were performed in triplicate using Platinum® SYBR® Green qPCR SuperMix-UDG (Invitrogen). PCR conditions were: denaturation at 95°C for 2 min, amplification and quantification programs with 45 cycles of 95°C for 1 s, each of annealing temperature for 30 s, extension at 72°C for 30 s. Melting curve analysis of amplification products was performed at the end of each PCR reaction to confirm that a single PCR product was detected. Reactions were conducted in singleplex in a 96-well spectrofluorometric thermal cycler.

Table 1. Oligonucleotide primer sequences for SybrGreen qRT-PCR (KIT, proto-oncogene tyrosine-protein kinase; KITLG, stem cell factor; ABCG2, ATP-binding cassette sub-family G member 2; MSI1, musashi homolog 1; 18S rRNA, 18S subunit rRNA; ACTB actin beta)

<table>
<thead>
<tr>
<th>Gene</th>
<th>Primer sets</th>
<th>Product length (bp)</th>
<th>GeneBank gi-number</th>
</tr>
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<tbody>
<tr>
<td>KIT</td>
<td>Forward: 5’AGGGACTGAAAGGGCACTT 3’</td>
<td>205</td>
<td>110681494</td>
</tr>
<tr>
<td></td>
<td>Reverse: 5’AGAACACCTCTGCTGGTC 3’</td>
<td></td>
<td></td>
</tr>
<tr>
<td>KITLG</td>
<td>Forward: 5’GTGCGCAAAATCTCCAAGA 3’</td>
<td>222</td>
<td>14993549</td>
</tr>
<tr>
<td></td>
<td>Reverse: 5’TGCCCTGACTCCCAAAG 3’</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ABCG2</td>
<td>Forward: 5’GACCTGAGAGAAGCTGA 3’</td>
<td>222</td>
<td>115589695</td>
</tr>
<tr>
<td></td>
<td>Reverse: 5’ATGTTGAGCTCTTCTTGAC 3’</td>
<td></td>
<td></td>
</tr>
<tr>
<td>MSI1</td>
<td>Forward: 5’TTTCTGCTTCCAACCTTTTC 3’</td>
<td>184</td>
<td>124358512</td>
</tr>
<tr>
<td></td>
<td>Reverse: 5’TCCACGGTGGTGTTTACT 3’</td>
<td></td>
<td></td>
</tr>
<tr>
<td>18S rRNA</td>
<td>Forward: 5’AAAACGCTACCACATCCAAG 3’</td>
<td>90</td>
<td>58760943</td>
</tr>
<tr>
<td></td>
<td>Reverse: 5’TCTGTATGTATTTTCGTCAC 3’</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ACTB</td>
<td>Forward: 5’TCCCTGAGAGAGCTCA 3’</td>
<td>102</td>
<td>2182268</td>
</tr>
<tr>
<td></td>
<td>Reverse: 5’AGACCGTGTTGGATAGAG 3’</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
(DNA Engine Opticon 2; MJ Research, Inc., Waltham, MA, USA).

The expression of target genes was normalized using the 18S rRNA gene, which is known to be constitutively expressed (Robinson et al., 2007). The expression level of a given target gene in each experimental group was analysed by the $2^{-\Delta\Delta Ct}$ method (Bustin, 2000; Pfaffl, 2001) where $2^{-\Delta\Delta Ct}$ represents the difference of a given target gene between each group after lambing (groups from LateP to 8IN) versus the group P. More precisely, individual $\Delta Ct$ was calculated for each sample of group (LateP-8IN) as $\Delta Ct = \Delta Ct$ (sample group) – mean $\Delta Ct$ (group P). The $n$-fold expression was calculated as $2^{-\Delta\Delta Ct}$ for up regulated genes, and $- (1/2^{-\Delta\Delta Ct})$ for down regulated genes. The $n$-fold expression of a given target gene was calculated as $\log_2 (2^{-\Delta\Delta Ct})$ (Fig. 2).

**Statistical analysis**

All the recorded variables were submitted to analysis of variance using the ANOVA model to assess significant differences between groups; Duncan’s least significant difference test was used to compare the means (SPSS Inc., 1997).

**Results**

*In situ* hybridization

*In situ* hybridization on histological sections was done using anti-sense and sense cRNA digoxigenin-labelled probes directed against the 205-bp KIT sequence. The hybridization analysis was utilized for studying the expression of the isolated mRNA in the mammary glands of sheep at P, LateP, 60L and 8IN stages. Alkaline-phosphatase-tagged RNA probes were revealed by a reddish-blue colour and the staining indicative of KIT mRNA expression was cytoplasmatic (Fig. 1).

In P mammary tissue the anti-sense probe labelled epithelial cells of differentiating ducts as diffusely punctuate staining (Fig. 1a). Strong reaction was seen in epithelium of alveoli in LateP both in epithelial cells (Fig. 1b, arrows) and in stromal cells lining the alveoli (Fig. 1b, arrow-
heads). Very faint or no positive labelling was observed during lactation (60L), mostly at the apical side of the alveolar cells (Fig. 1c, arrows), whereas in 8IN group weak positive signal was present in the cytoplasm of luminal and stromal cells surrounding shrunken alveoli (Fig. 1d, arrows). The specificity of KIT anti-sense probe was ascertained using the sense probe as negative control (Fig. 1e).

**Expression analysis**

18S rRNA expression levels were quantified in all samples and resulted in constant expression levels. No significant differences between the groups could be shown in the investigated ovine mammary tissues.

Native agarose gel analysis of total RNA and β-actin amplification were used to verify the integrity of the mRNA extracted. Quantitative measurements of mRNAs encoding KIT, KITLG, ABCG2 and MSI1 in sheep mammary glands were obtained using real-time PCR on total mammary glands RNA samples.

Gene expression results are shown in Table 2. In terms of ΔCt, KIT gene expression significantly differed (P < 0.05), being highest during lactation than at prepubertal stage (P), and during involution (8IN). The KITLG gene abundance was significantly different (P < 0.05) only at LateP. The expression of ABCG2 gene significantly changed (P < 0.05), counter to KIT expression, being at its higher in P group and 8IN group. The expression of MSI1 gene was significantly affected (P < 0.05), being higher in P group and 8IN group. The expression of ABCG2 gene significantly increased (P < 0.05) at LateP in comparison to 30, 60 and 150 days of lactation, and 8IN group. The MSI expression showed the same trend of KIT, but it was significantly transcripted (P < 0.05) in LateP and 8IN groups than in 30L, 60L and 150L groups. The correlation between KIT and MSI1 was equal to 0.95.

**Table 2. Expression level of putative stem cell markers in groups of sheep at different developmental stage. KIT, proto-oncogene tyrosine-protein kinase kit; KITLG, stem cell factor; ABCG2, ATP-binding cassette sub-family G member 2; MSI1, musashi homolog 1**

<table>
<thead>
<tr>
<th>Groups</th>
<th>KIT</th>
<th>KITLG</th>
<th>ABCG2</th>
<th>MSI1</th>
</tr>
</thead>
<tbody>
<tr>
<td>P</td>
<td>6.49±±0.21</td>
<td>6.81±±0.33</td>
<td>3.62±±0.24</td>
<td>12.67±±0.11</td>
</tr>
<tr>
<td>LateP</td>
<td>7.60±±0.53</td>
<td>5.29±±0.29</td>
<td>1.74±±0.59</td>
<td>15.67±±0.68</td>
</tr>
<tr>
<td>30L</td>
<td>9.05±±0.47</td>
<td>7.17±±0.77</td>
<td>1.46±±0.52</td>
<td>18.93±±0.46</td>
</tr>
<tr>
<td>60L</td>
<td>8.67±±0.51</td>
<td>8.46±±0.38</td>
<td>1.41±±0.44</td>
<td>17.97±±1.00</td>
</tr>
<tr>
<td>150L</td>
<td>8.97±±0.92</td>
<td>8.31±±0.79</td>
<td>1.57±±0.63</td>
<td>17.63±±1.14</td>
</tr>
<tr>
<td>8IN</td>
<td>7.09±±0.44</td>
<td>7.61±±0.90</td>
<td>3.19±±0.57</td>
<td>15.10±±0.42</td>
</tr>
</tbody>
</table>

Different superscripts within a column indicate significantly (P < 0.05) different means between groups.

**Discussion**

Development of mammary glands is greatest during puberty with cyclical progression throughout adult life. In ewes, prepubertal mammary glands is composed by developing epithelial ducts, named ‘terminal ductal unit’ (TDU), which are surrounded by densely packed stromal cells embedded in a large adipose tissue (Akers, 2002; Capuco et al., 2002; Colitti and Farinacci, 2009). During pregnancy, branching networks of ducts and lobuloalveolar structures grow and are fully developed at lactation. At involution, extensive restructuring of the tissue includes the elimination of a proportion of the secretory epithelial cells by apoptosis (Colitti et al., 1999). The massive expansion of mammary epithelium during puberty and pregnancy, together with the remarkable regenerative capacity apparent during successive reproductive cycles, implicate stem-like cells (Shackleton et al., 2006). In fact, it was demonstrated that, through serial transplantations, no loss of potency was detected when compared to similar transplantations of the youngest mammary tissue tested (Boulanger and Smith, 2009).

The identification of molecular markers that best facilitate the isolation and characterization of stem cell populations has long been a question. Although previous analyses have yielded no definite markers for mammary stem cells, histological analyses have indicated that a lightly staining parenchymal cell population may function as mammary stem cells (Ellis and Capuco, 2002). Other established ability of stem cells is to efflux cellular dyes as Hoechst 33342 under ABCG2 contribution (Zhou et al., 2001; Alvi et al., 2003). Therefore, ABCG2 expression may serve as a new marker for stem cells, not only in hematopoietic cells but also in other types of cells.

In this study, we sought to investigate, by real-time PCR, the expression of some genes identified as stem cells markers in mammary tissue at different developmental stages. In addition, KIT expression was localized by *in situ* hybridization analysis. The quantitative expression analysis was related to prepubertal stage because it seems to be the best candidate stage of mammary gland tissue to found stem cells. In fact, as reported by Capuco (2007), some cells, displaying characteristics of stem-like cells, were identified in prepubertal bovine mammary gland. In this study, KIT, KITLG and MSI1 expressions positively
correlate with prepubertal stage of mammary tissue. Actually, although no evidences are present on the expression of KIT in prepubertal mammary tissue, it is known that KIT is highly expressed in normal mammary human epithelium and that an autocrine stimulation of KIT by its ligand KITLG is important for maintenance of differentiation of mammary epithelium (Ulivi et al., 2004). It has been demonstrated that KIT is essential for full activation of STAT5A (Brizzi et al., 1999) and that it has anti-apoptotic action (Jin et al., 2004) reducing the expression of BAX by inducing FOXO3a phosphorylation (Liu et al., 2009). This is in agreement with our previous results that demonstrated in mammary gland of sheep STAT5A and BAX expressions were significantly down-regulated during lactation (Colitti and Farinacci, 2009). It is likely that the down-regulation of KITLG and KIT expressions during lactation may be related to the physiological state of the gland and contributes to maintain a balanced homeostatic state in the tissue.

The up regulation of ABCG2 is in agreement with Jonker et al. (2005) that found high alveolar expression of the ABCG2 gene in lactating, but not in virgin or nonlactating mammary glands of cow, contributing to the secretion of nutrients into the milk in spite of the coincident risk of contaminating milk with xenotoxins.

The up regulation of MSI1 at prepubertal stage could be due to the activation of Notch signalling – which is known to maintain cells in an undifferentiated state and to inhibit apoptosis – through the translational repression of m-Numb protein (an intracellular Notch antagonist), resulting in a clonal expansion of the immature stem cells. MSI1, a putative stem/progenitor cell marker was observed in the small intestine at the fourth–sixth cell position from the bottom of the crypts and in the cells in the deepest portion of the large intestine (Potten et al., 2003). It is recently shown to be also an important positive regulator of cell proliferation and inhibitor of apoptosis. In fact, gut tumours arise from stem cells expressed at the base of intestinal and colonic crypts where MSI1 is expressed. The involvement of MSI1 in colon cancer cell proliferation, inhibition of apoptosis and mitotic catastrophe, occurs through its role in inhibition of Notch signalling (Sureban et al., 2008). In mammary gland of sheep we found – using immunohistochemistry and in situ hybridisation analyses – that MSI1 was expressed on mammary glands of ewes at different developmental stages and, moreover, that MSI1 fold values resulted strongly correlated (r = 0.96) with the expression of Ki-67 a marker of proliferation assessed for the same groups of ewes (Colitti and Farinacci, 2009). In the present paper, MSI1, in terms of ΔCt and n-fold, positively correlate with lactation stages (R² = 0.93), when it is known that proliferation prevails to apoptosis. The increase of KIT and MSI1 expression in Late P and in 8IN groups could prime mammary stem cells to differentiate, because in those stages new secretory cells are useful to lactation and to remodelling the gland.

In summary, as already underpinned by Welm et al. (2003), it is likely that the final fate of progenitors is in balance between many signalling pathways that act accordingly to regulate the process of differentiation versus self renewal. The present study provides the evidence...
that putative stem cell markers vary their expression during physiological phases of mammary cycle.

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References


