Identification of Bulge Stem Cells in Mouse and Human Hair Follicles

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1. Introduction

The epithelium of mammalian skin is known to contain permanently residing stem cells (SC) that are able to maintain multiple differentiated lineages in the interfollicular epidermis (IFE), hair follicle (HF), and sebaceous gland (SG) [1].
In this regard, the HF is one of the most complex miniorgans in the bodies of mammals [2] [3]. The adult HF consists of an upper portion that is permanent and a lower portion that constantly remodels during the hair’s cycle of: growth (anagen), degeneration (catagen), and rest (telogen), and the cyclical nature of hair growth is thought to be dependent on HF stem cells [4] [5]. They are located in specific regions of tissue called “niches” and remain in a state of quiescence, to be activated by disease or alterations of physiological processes caused by tissue injury. Stem cell niches are discrete, dynamic, and functional microenvironments, domains that influence SCs’ ability to govern tissue homeostasis and repair throughout the lifetime of an organism [6]. The niches’ main function is to maintain and repair the tissue or organ in which they are located [7]. The mammalian HF has a niche for SC called the “bulge” region is a segment of the outer root sheath (ORS) adjacent to the site of attachment of the arrector pili muscle [8]. Stem cells located in the HF bulge area give rise to the follicle structures during each anagen phase [9].

Several putative biomarkers have been suggested for identifying SC in murine and human HF; however, these results show discrepancies. The purpose of our study was the identification of bulge region in mouse and human HF, and to determine the patterns of biomarker expression of proteins CK15, CD34 and CD200 by immunohistochemistry and immunofluorescence in order to find the reliable molecular markers. The potential biomarkers could be useful in characterizing and isolating follicular SC in mice and humans.

2. Materials and Methods

2.1. Tissue Samples

Mice, 2 - 3 days old (n = 7), were sacrificed, and follicles were obtained from the skin of the body and from the vibrissae. Healthy human scalp skin was obtained from facelift patients (n = 7) with the approval of the ethical committee of the Faculty of Sciences of the Central University of Venezuela.

2.2. Histopathology

In order to observe the morphology of bulge in vivo in mouse and human, the skins with HF were fixed overnight in 4% neutral-buffered paraformaldehyde and transferred to 70% ethanol. They were then embedded in paraffin, sectioned at 8 μm, and stained with blue toluidine and hematoxylin eosin (H&E).

For the immunohistochemistry and immunofluorescence studies, the skin with HF was immediately frozen in Thermo OCT compound (Sakura Finetek, Tokyo, Japan). Consecutive 8-μm-thick sections were sliced with a cryostat and mounted serially on separate slides for three different immunostains against (cytokeratin 15, clone LHK15; Thermo, Lab Vision, Fremont, CA), cluster of differentiation CD34 (also known as mucosalin) (BD Biosciences, San Jose, CA), and CD200 (known as OX2) (BD Biosciences).
2.3. Immunohistochemistry

Frozen sections were defrosted, air dried for 3 hours, and fixed with acetone for 10 min. at 20°C. They were then incubated with primary antibodies (diluted 1:100) overnight at 4°C. The next day, the sections were rinsed for 3 - 5 minutes to remove unbound primary antibodies. Subsequently, they were incubated for 30 min. at room temperature with biotinylated secondary antibodies (anti-rabbit and anti-mouse immunoglobulins), which form a complex with peroxidase-conjugated streptavidin molecules. Signal was detected using the Dako Cytomation (Carpinteria, CA), with diaminobenzidine as the chromagen, following the manufacturer’s protocol. Tissues were counterstained in Harris hematoxylin (Sigma-Aldrich, St. Louis, MO). The specificity of staining was evaluated by negative controls in the absence of primary antibody. Labeled cells were observed, using an inverted microscopy-enable phase contrast and epifluorescence microscope (Axiolab 40, Carl-Zeiss, Göttingen, Germany).

2.4. Immunofluorescence

Frozen sections in OCT compound (Thermo Fisher Scientific, Germany) were sliced with a cryostat and mounted on slides. Sections were blocked with 10% horse serum, diluted 1:30, at room temperature for 25 min and then incubated with the first primary antibody (anti-mouse CK15, anti-mouse CD34 IgG [in mice], and anti-human CD200 IgG [in humans], diluted 1:100) at room temperature for 90 min. After three consecutive washes, they were incubated with the second antibody (goat anti-mouse IgG conjugated with FITC for CK15 [in mouse and human], anti-rat IgG biotin [in mouse], and anti-mouse IgG biotin [in human]), diluted 1:100 at room temperature for 30 min. Slides were washed three times and then incubated with Streptavidin PE for 30 min for CD34 and CD200. Slides were washed, counterstained, and treated with mounting medium containing 4,6-diamino-2-phenylindole (DAPI, Vector Laboratories, Burlingame, CA). Labeled cells were identified using an Olympus FluoView FV1000 confocal microscope (Tokyo, Japan) equipped with a video camera.

3. Results

3.1. Morphology of the Bulge Region in Mouse and Human Skin

Immunohistochemistry and immunofluorescence studies of mouse and human skin samples allowed us to appreciate the organization and distribution of the HF cells and to identify SC of the follicles as well as to estimate the sensitivity of the monoclonal antibodies (CK15 in mouse and human, monoclonal antibody a highly glycosylated, CD34 in mouse, and CD200 in human) which are considered specific markers of SC.

In histological sections of mouse and human skin, the structure and organization of HF were evident, and we were able to distinguish the location of the bulge region (Figure 1). The expression profiles of biomarkers are summarized (Table 1), with differences in longitudinal distribution of expression...
3.2. Differential Expression of Cytokeratin 15 in Mouse and Human Skin

CK15 is an intermediate filament protein expressed in basal keratinocytes of stratified squamous epithelium [10] [11]. It is used as a marker of undifferentiated multipotent cells and is considered a biomarker for mouse and human skin among CK15, CD34 and CD200.

![Figure 1](image_url)

**Figure 1.** Anatomy of mouse and human anagen hair follicles, showing the bulge region. (A). Mouse, bulge region (arrow), hair (HF) (blue toluidine) Scale bars = 100 µm. (B). Human, bulge region (arrow), hair (HF), the SG duct (H & E). Scale bars = 100 µm.

**Table 1.** Summary of immunohistochemical and immunofluorescence staining patterns in the epithelial components of mouse and human scalps and the anagen hair follicles.

<table>
<thead>
<tr>
<th>Epidermis</th>
<th>Hair Follicles</th>
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<tbody>
<tr>
<td></td>
<td>Outer root sheath</td>
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<td></td>
<td>Infundibulum</td>
</tr>
<tr>
<td>Mouse</td>
<td>B</td>
</tr>
<tr>
<td>CK15</td>
<td>+</td>
</tr>
<tr>
<td>CD34</td>
<td>+/-</td>
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<tr>
<td>Human</td>
<td>CK15</td>
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<td>CD200</td>
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The results obtained in this study, using the CK15, CD34, and CD200 antibodies, showed that staining intensities were: (−): no staining; (+/-): partially slightly positive staining; (+): positive staining; and (++): strongly positive staining. B: basal cell, SB: suprabasal cells, ORS: outer root sheath, SG: sebaceous gland.
folicular SC that reside in the bulge [12] [13] [14] [15]. The results of the immunohistochemistry and immunofluorescence analyses (Table 1) for mice and humans showed strong CK15 staining in the basal IFE, moderate staining in the outermost layer of the bulge ORS, the proximal isthmus, infundibulum, and outermost layer of the sub-bulge ORS, suggesting the presence of undifferentiated cells. The controls were negative in the bulge region.

3.3. Expression of CD34 in the ORS Proximal Isthmus, the Infundibulum and the Bulge Region in Mouse Skin

Cluster of differentiation 34 (CD34) is a surface glycoprotein (105 - 120 kDa), expressed on early hematopoietic stem and progenitor cells [16], and functions as an important factor in cell-cell adhesion [17] [18]. Mouse immunohistochemistry and immunofluorescence demonstrated differential expression patterns of CD34 (Table 1), (Figure 2 and Figure 3). We found CD34 staining in some cells in the basal IFE (Figure 2 and Figure 3), and SG duct.

3.4. Expression of CD200 in the Basal IFE and the Bulge Region in Human Skin

The Ox-2 membrane glycoprotein, also called CD200, a member of the immunoglobulin superfamily, is involved in controlling the activation of macrophages and granulocytes [19].

Figure 2. Immunohistochemical determination of CD34 in histological sections of mouse anagen hair follicles. Light microscope. (A). CD200 stained section showing brown-colored cells in the basal layer of the IFE (arrow) Scale bar = 20 µm. (B), (C). CD34 positive in the ORS proximal isthmus (arrowhead) and the infundibulum and the bulge region (arrowhead) ((B) Scale bar = 100 µm. (C) Scale bar = 20 µm.). (D). Negative control Scale bar = 100 µm.
Figure 3. Immunofluorescence of longitudinally sectioned mouse vibrissae (A)-(C) and HF (D)-(E) stained for CD34. Confocal microscope. (A) Panoramic view showing the organization of the whiskers in the skin. Image DIC. (B), (C). High CD34 expression in all areas of the bulge region (arrows) and moderate expression in cells of the SG surrounding the whiskers. (C)-(F). High CD34 expression in the bulge region (arrow) and moderate expression in cells of the basal SG. Sum of images (blue and green). Nuclei are counterstained with DAPI. Scale bars = 100 µm.

In the present investigation, immunohistochemistry and immunofluorescence showed slight to moderate CD200 staining in the basal IFE and strong staining in the outermost layer of the bulge ORS, the infundibulum, and the
SG duct (Table 1, Figure 4 and Figure 5). In addition, the controls’ reaction was negative in the bulge region (Figure 5 (A)).

4. Discussion

Our immunohistochemistry and immunofluorescence assessment of follicle sections revealed that SC-associated biomarkers are differentially expressed in the epidermis and anagen hair follicles. Similar results in the immunofluorescence of cytokeratin 15 in mouse and human were also found as a homogeneous staining expressed in epidermal basal keratinocytes in the outermost layer of the bulge ORS, the proximal isthmus, infundibulum, and outermost layer of the sub-bulge ORS, as reported previously [13] CK15 is not a specific of the bulge. However, the positive expression of CK15 is preferably located in the bulge region in the human scalp, and this expression is absent in the basal IFE [12] [20]. A pattern similar to previous reports indicates that CD34 is the best marker to delineate SC in the bulge region in murine models [21] [22] [23]. Also, it has been suggested that the CD34 antibody is an excellent mouse follicular cells and is not present in the cell marker of the human RPFP [24]; coinciding with our results indicating that CD34 give an important role in bulge cells. CD200 was highly expressed in all areas of the bulge region, consistent with previous reports identifying it as a possible SC niche [12] [15] [25] [26], in contrast to mouse bulge cells.

Figure 4. Immunohistochemical determination of CD200 in histological sections of anagen hair follicles of mouse. Light microscope. (A). Negative control. (B). CD200 staining in the bulge (arrowhead) Scale bars = 100 µm.
Figure 5. Immunofluorescence of longitudinally sectioned human hair follicles stained for CD200. Confocal microscope. (A)-(F). Panoramic view showing the organization of the HF in the skin. (A) (D). Image DIC. (B), (C), (E) and (F). High CD200 expression in all areas of the bulge region (arrow). Nuclei are counterstained with DAPI. Scale bars = 100 µm.

This pattern of expression may suggest that follicular cells with a focal immunodetection migrate in the region of prominence from the basement membrane to the bulge region to contribute in tissue regeneration [8] [27] [28], expression being gradually decreased along the follicle. Likewise, the presence of two distinct subsets of multipotent SC, basal and suprabasal layers, has been
demonstrated in mice and humans, within the protrusion. The first, larger than the second [29] [30] [31], showed cells in diverse stages of differentiation, indicating that they may represent transient amplifying cells or progeny of bulge SC.

In general, the techniques described in this study made it possible to compare and determine the pattern of staining more clearly, using confocal fluorescent microscopy that demarcates individual positive cells of a defined tissue. The results of immunohistochemistry and immunofluorescence evaluation carried out in sections of mouse whiskers and HF of mice and humans and were very useful in differentiating cell expression, revealing that the cells associated with CD34 and CD200 antibodies were differentially expressed in the epidermis, the basal SG, the vibrissae, and HF. This perspective suggests that the CD34 and CD200 biomarkers (in mice and humans, respectively) had greater specificity than CK15 in identifying bulge cells. We also suggest that the sensitivity of the CD34 antibody is higher in cells with SC characteristics in mouse HF, while CD200 might have specificity with characteristics for progenitor cells and partially differentiated cells. The location of intrinsic SC within the HF suggested that they are easily accessible. This important attribute makes these cells valuable candidates for use in vitro studies and in regenerative medicine.

5. Conclusion

The Immunohistochemical staining data showed that CD34 in mouse and CD200 in human are reliable markers because they are expressed strongly in the bulge area allowing identification of the niche cells, representing a promising tool for classifying cell populations according to expression levels of stem cell-associated biomarkers.

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Conflicts of Interest

The authors declare no conflicts of interest regarding the publication of this paper.

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